Mechanism of Action of Secreted Newt Anterior Gradient Protein

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Abstract

Anterior gradient (AG) proteins have a thioredoxin fold and are targeted to the secretory pathway where they may act in the ER, as well as after secretion into the extracellular space. A newt member of the family (nAG) was previously identified as interacting with the GPI-anchored salamander-specific three-finger protein called Prod1. Expression of nAG has been implicated in the nerve dependence of limb regeneration in salamanders, and nAG acted as a growth factor for cultured newt limb blastemal (progenitor) cells, but the mechanism of action was not understood. Here we show that addition of a peptide antibody to Prod1 specifically inhibit the proliferation of blastema cells, suggesting that Prod1 acts as a cell surface receptor for secreted nAG, leading to S phase entry. Mutation of the single cysteine residue in the canonical active site of nAG to alanine or serine leads to protein degradation, but addition of residues at the C terminus stabilizes the secreted protein. The mutation of the cysteine residue led to no detectable activity on S phase entry in cultured newt limb blastemal cells. In addition, our phylogenetic analyses have identified a new Caudata AG protein called AG4. A comparison of the AG proteins in a cell culture assay indicates that nAG secretion is significantly higher than AGR2 or AG4, suggesting that this property may vary in different members of the family.

Introduction

The first member of the anterior gradient protein family, referred to as XAG2, was identified as a marker of anterior non-neural development in Xenopus [1]. The ectopic expression of this protein in embryos induced a supernumerary cement gland, a mucous-secreting element that also expresses XAG2 [2]. It has recently been implicated as a critical secreted signal in Xenopus telencephalon formation [3]. Another amphibian AG protein was encountered in the context of salamander limb regeneration [4]. The newt protein nAG was identified as a binding partner of the salamander-specific protein Prod1, a member of the three-finger protein (TFP)
superfamily that is GPI-anchored at the cell surface [5]. nAG expression was elevated in Schwann cells of the nerve sheath as peripheral axons regenerate into the newt limb blastema, the regenerative zone at the end of the stump [4]. nAG was subsequently expressed in gland cells underlying the specialised wound epidermis of the blastema. Limb regeneration is normally dependent on the presence of regenerating axons, and it is striking that ectopic expression of nAG could rescue regeneration of a denervated blastema. Recombinant nAG acted to promote S phase entry of cultured newt blastemal cells, possibly by binding to Prod1 on the cell surface, although this requirement has not yet been demonstrated [4].

A mammalian homolog of XAG2, generally referred to as AGR2, is presently the focus of considerable interest in different contexts. It is expressed by secretory epithelia and is upregulated in adenocarcinomas, originally shown in breast cancer cells in culture [6], but subsequently a wide variety of tumour cells including those found in the gastrointestinal tract [7,8]. The normal development of both intestine and stomach is perturbed by knockout of the mouse AGR2 gene [9–12], leading to an imbalance of differentiated cell types in both organs, and defects in mucous production. It has been implicated in the metastatic phenotype of breast cancer cells, and interacts with alpha dystroglycan and C4.4A, two markers of metastasis [13]. As is the case for Prod1, C4.4A, more commonly known as Ly6/PLAUR domain-containing protein 3 (Ly6P3), is GPI-anchored and a member of the TFP superfamily. AGR2 is clearly a significant biomarker for human adenocarcinoma and a potential target for drug discovery [8].

AG family members are predicted to consist of a N-terminal signal peptide followed by a single thioredoxin domain, as has now been verified by determination of the 3D structures of human AGR2 [14] and human AGR3 [15]. Most of the AG proteins, with the exception of the majority of AGR3 orthologues, have a single cysteine residue in the canonical thioredoxin active site. It has been suggested that AG proteins may have an intrinsic protein disulphide isomerase activity, and that this could be implicated in functions dependent on the endoplasmic reticulum (ER) localisation [16]. The AG proteins have a non-canonical Lys/Gln/His-Thr/Ser/Ala/Gly-E-L ER-retention sequence. The models for the function of AGR2 propose a role in attenuating ER stress responses [17], or signalling from the ER to regulate gene expression [16], in particular the amphiregulin gene, which may be critical for mediating the action on cultured adenocarcinoma cells [18], and also the EGF receptor gene [19]. On the other hand, the functions of amphibian and fish AGs, in salamander [4], Xenopus [3] and zebrafish [20], involve secretion and extracellular action, and nAG is secreted after transfection of cultured cells [4]. Furthermore AGR2 is found in high concentrations in gastrointestinal mucus [21], and is secreted by human and rat mammary epithelial cells as an O-glycosylated molecule [22].

We address several of these issues, including the activity of secreted nAG, the dependence on the single Cys residue in the thioredoxin active site, and the variation of ER retention versus secretion in different AG proteins.

Materials and Methods
Site-directed mutagenesis and plasmids
The newt wild type nag sequence was amplified using oligos (Forward 5’ –GCGGCTAG CATCGCTCAACATGGTGAA-3’, Reverse 5’ –CGCGTCAATTCTTCGCTCACA-3’) on newt blastemal cDNA and cloned between the NheI and EcoR1 sites of the vector pCI-neo (Promega) to give pnAGWT. The cysteine at position 72 was mutated to a serine using oligos (Forward, 5’ –CACAGAGATGACTCTCCACACTCTCAGGCTTTGAAGAAAG-3’, Reverse, 5’ –CTTTCTTCAAAGCCTGAGAGTGTGGAGCGTCATCTCTGTG-3’) or an alanine (Forward, 5’ –CACAGAGATGACGCTCCACACTCTCAGGCTTTGAAGAAAG-3’, Reverse, 5’ –CTTTCTTCAAAGCCTGAGAGTGTGGAGGCCTCATCTCTGTG-3’) using the
Quikchange site-directed mutagenesis kit (Stratagene) to create mutant clones nAG-C72S and nAG-C72A respectively.

Myc tags were added at the 3'end of these inserts by PCR using oligos 690 (Forward, 5' -CACAGAGATGACGCTCCACACTCTCAGGCTTTGAAGAAAG-3', Reverse, 5' -CGCTCTAGAAATCAGATCCTCTTCTGAGATGAGTTTTTGTTCCAGTTCAGTTTTCAGATTGTGCT-3') and the resultant fragments were cloned into the EcoRI and XbaI sites of pCI-neo expression vector to give 3 myc+25 constructs. We found that the translation product used a vector stop site, resulting in an expressed protein product with an additional 25 amino acid residues (ISRVDPGGRFPLVRVNASSRHDKIH) at the C terminus which are derived from a sequence in the vector. This appendage proved to be beneficial for the recovery of the protein from the conditioned medium. For derivation of constructs without these additional amino acids, the wild type and mutant constructs were amplified with oligos 690 and primer 5' -CGCTCTAGACTACAGATCCTCTTCTGAGATGAGTTTTTGTTCCAGTTCAGTTTTCAGAAG TTTGAGTGCT-3', and the resultant fragments were cloned into the EcoRI and XbaI sites of pCI-neo vector to give 3 myc-tagged constructs.

The plasmid pEGFP-N2 (Clontec) was used as a negative control in parallel transfections. In some cases a construct pERFP-N2 was used, where the GFP (green fluorescent protein) sequence in pEGFP-N2 is swapped for RFP (red fluorescent protein).

**Animals, cell culture and transfection**

The animal maintenance and procedures of the experiments were approved by the Home Office, UK. Adult newts were obtained from Charles D. Sullivan & Co. TN, USA, and maintained in aquariums at 25°C. The animals were fed with live blood worms throughout the experimental period. Bilateral forelimb amputation was performed under anaesthesia (0.1% tricaine) and the newts were allowed to regenerate. Isolation and culture of limb blastema cells has been described previously [23]. Briefly, the limb blastemas at mid-cone stage of growth (12–14 days after limb amputation) were collected, and the blastema cells were dissociated non-enzymatically from the mesenchyme in amphibian cell culture medium. The cells were plated on to collagen-coated (Type I, Sigma C8919) 96 well microplates and maintained at 25°C with 2% CO2 supply. The cell density varied from 1000–2000 cells per well in various batches. The assay for S phase entry have been described [4]. In brief, the cells were cultured for 72 h, and the medium was changed to include the various additives, either rabbit IgG 683 to Prod1 [5], or affinity purified control rabbit IgG (Sigma 15006), and 10 μM BrdU, followed by a 96 h incubation prior to fixation and antibody staining. The microwells were scanned using a Zeiss Axiovert 200 microscope controlled through Axiovision (Carl Zeiss, Version 4.8.2) software. The images were stitched and exported to Image Pro Plus (Media Cybernetics) for processing and cell counting [4]. Cos7 and HEK 293T cell lines were purchased from ATCC and Invitrogen respectively, and cultured under standard mammalian cell culture protocol in Dulbecco’s Modified Eagle Medium containing 10% fetal bovine serum. The cells were seeded on to 30 or 60 mm culture wells coated with poly(ethyleneimine) solution (1 μg/mL; Sigma, P3143), and transfected at 80% density with nAG-expressing pCIneo plasmids using Lipofectamine 2000 as described [24]. The transfection efficiency in Cos7 cells varied from 40–60%, whereas HEK 293 cells attained 80% transient expression.

**Cell lysates, conditioned medium and western blotting**

For the experiments on the nAG mutants, cells were lysed in a buffer containing 1% NP40, 0.15 M NaCl, 1 mM Na3VO4, 0.05 M Tris-HCl pH 7.5 supplemented with protease inhibitors (Complete Cocktail Tablets, Roche). For the experiments on transfection of AG family
members, the cells were lysed in a buffer containing 0.15 M NaCl, 1 mM EDTA, 0.1% SDS, 1% NP40, 1% sodium deoxycholate, 10 mM Tris HCl pH 7.8, supplemented with protease inhibitors. Cell lysates were centrifuged at 5000 g for 10 min, and solubilised in sample buffer prior to analysis by electrophoresis.

After incubating Cos7 or HEK 293T cells for 48 or 72 h, the culture media were collected, adjusted to 10 mM HEPES pH 7.4 and centrifuged at 180 g for 10 min prior to loading onto Vivaspin ultrafiltration units with a 10 kD exclusion limit. The concentrated media samples were solubilised in sample buffer. Cell lysates and concentrated conditioned media samples were analysed by electrophoresis on precast Novex 12% Bis-Tris SDS polyacrylamide gels, and transferred to nitrocellulose by standard methods prior to antibody reaction. For detection of nAG we used the polyclonal rabbit antibody 224 at a dilution of 1:500 and fluorescent secondary goat anti-rabbit conjugated to IR dye emitting at 800 nm (Rockland). For determination of absolute levels of nAG and mutants, we used a protein standard of nAG with C-terminal hexa-histidine tag, produced in insect cells using the baculovirus expression system. For detection of AG4 and AGR2 we used rabbit anti-human AGR2 (Sigma) made against residues 108–157 in the sequence of AGR2, and detected with the same secondary antibody. The blots were analysed in a LICOR Odyssey scanner, and the nAG band intensities were multiplied by the appropriate dilution factors to give estimates for the total secreted and cell-retained protein in the culture dishes. Blots were routinely co-stained with mouse anti-actin (Sigma) and goat anti-mouse emitting at 600 nm. The actin signals were not used in the determination of percentage secretion (since actin is not secreted), but in comparison of lysates to ensure that the nAG, AG4 and AGR2 proteins were expressed at comparable levels and not subject to marked differences in expression after transfection.

Phylogenetic analyses

Tblastn was used to mine nucleotide sequences from databases including NCBI, Ensembl, Sal-site [25], and newt transcriptomes [26, 27] (S1 Table). Multiple sequence alignments were calculated using MUSCLE [28] and MAFFT [29] and inspected and modified using Jalview [30] and ClustalX [31]. GTR+G+I was selected as an appropriate substitution model using ModelGenerator [32]. Trees were computed by maximum likelihood using PhyML3 [33] with 1000 bootstrap replicates and by Bayesian inference with MrBayes 3.2 [34]. For Bayesian calculations convergence was assessed using AWTY [35]. Trees were visualised and figures were prepared using Dendroscope [36].

Results

Activity of the nAG protein on newt blastemal cells

In order to investigate the activity of recombinant nAG, mammalian Cos7 cells were transfected with plasmids expressing either nAG or red fluorescent protein (RFP) as control. The conditioned medium was concentrated and added to microwell cultures of newt limb blastemal cells for 3 days. Affinity purified rabbit IgG against newt Prod1, or control IgG, was added to some wells, and the effect on S phase entry was determined by incorporation of bromodeoxyuridine. The results of assays from three independent preparations of blastemal cells were normalised and are shown in Fig 1. The activity of nAG medium was inhibited to the level of control medium by inclusion of antibody to Prod1 at 20 μg/mL (Fig 1, Lanes 4, 6 and 7), while control IgG had no inhibitory action (Fig 1, Lane 5). The background activity of RFP control medium was not inhibited by addition of either antibody (Fig 1, Lanes 1, 2 and 3). These results are consistent with the hypothesis that the stimulatory activity of nAG on the newt blastemal cells is mediated by interaction with its binding partner Prod1.
Expression and activity of Cys72 mutations

Plasmids that encode wild-type nAG, or mutants of the protein at Cys72, were transfected in parallel into Cos7 cells or 293T cells in culture. After 2 days the conditioned medium and cell lysates were prepared for analysis by gel electrophoresis and western blotting. The wild-type protein was readily detectable in the conditioned medium of both cell types as an immunoreactive band at 17 kD (Fig 2). The mutant proteins were barely detected in both the conditioned medium (Fig 2) and the lysates. Quantitative analysis of the C72A band (with reference to a wild-type nAG band) indicated that the level of protein in the medium was less than 10% of wild type. The low level of expression of the mutant proteins was observed after transfection of both mammalian cell types. This effect was partially relieved for C72A by culturing the cells in the presence of the proteasome inhibitor MG132 (Fig 2). It was not possible to assay such conditioned medium for activity on cultured blastemal cells because the cells were very sensitive to the inhibitor. Thus the mutation of the Cys72 residue destabilises the nAG molecules and leads to their degradation, probably via the proteasome.

When a myc epitope was introduced at the C terminus of the three nAG protein constructs, they were recovered at significantly higher levels in the conditioned medium of transfected Cos7 cells, even in the case of the wild-type protein (Fig 3A and 3B). We made a chance observation that, when the proteins were further extended for 25 residues after the myc tag, the levels in the medium were even higher, and there was little difference in yield between the wild-type and mutant protein variants (Fig 3A and 3B). The wild-type nAG and its extended variant
(myc+25) had comparable activity in promoting S phase entry of cultured blastemal cells, and thus we were able to compare the activities of the extended versions of the three proteins at the same concentration. The data for three or four assays on independent preparations of cells have been combined and are shown in Fig 4. The magnitude of stimulation by the extended wild-type protein was comparable to that reported for the normal protein [4]. The mutation of Cys72 to either Ala or Ser leads to no detectable activity over the background of control (GFP plasmid-transfected) medium. These data suggest that the Cys72 residue is required for the proliferation-inducing activity of nAG.

Retention and secretion of AG proteins

Through mining of publicly available transcriptomic and genomic databases we have identified a novel group of Caudata (salamander) AG proteins, which we have called AG4. In molecular phylogenetic analyses (this work and [20]), AG proteins are vertebrate-specific and cluster in four groups each of which has a different distribution of vertebrate classes (Fig 5 and S1 Table) and a characteristic thioredoxin active site/C-terminus signature (S1 Fig).

The AGR1 group can be identified by its active site sequence Cys-Pro-His/Tyr-Ser and typically a His/Lys-Thr/Ser-Glu-Leu C-terminal region. This group includes fish, turtle, bird and amphibian sequences. Not all bird genomes contain an AGR1 gene, but its presence in the basal clade Palaeognathae (flightless birds) suggests that the gene was subsequently lost in certain lineages (e.g. Phasianidae). The AGR1 proteins found in Otophysa fish such as zebrafish and the cave fish Astyanax mexicanus are markedly different from the rest and can be recognised by a six-residue C-terminal extension not observed in any other AG protein. The AGR3

Fig 2. Degradation of Cys72 mutants of nAG. Cos7 cells were transfected with plasmids expressing (1) wild-type nAG, (2) C72A nAG, (3) C72S nAG, (4) GFP control, and cultured in the presence or absence of 50 μM MG132 as shown. The conditioned medium was collected, concentrated and analysed by western blotting as described in the Materials and Methods.

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The gene is only present in lobe-fin fishes and tetrapoda (Sarcopterygii). The signature sequences are Cys-Gln-Tyr-Cys/Ser for the active site and Gln-Thr/Ser/Lys-Glu-Leu at the C terminus.

The cluster that includes human AGR2 contains one gene per species from fish to mammals, except for Caudata species which have two genes in this group, nAG and AG4. The

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**Fig 3. Expression of nAG proteins with C terminal extensions.** (A) Wild-type and mutant nAG proteins, with or without (untagged) additional residues at the C terminus, were expressed in Cos7 cells and the conditioned medium was analysed as described in Fig 2. Note that the myc-tagged samples were run on a separate gel to the other samples, and the four lanes were joined to the right hand side. GFP refers to control GFP-transfected cells. (B) The yield of nAG proteins in the medium was quantitated by analysis of western blots for three independent transfections. Note that the largest yield for each of the three proteins is obtained with the myc+25 extension.

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The cluster that includes human AGR2 contains one gene per species from fish to mammals, except for Caudata species which have two genes in this group, nAG and AG4. The
signature sequences for the group are Cys-Pro-His-Ser in the active site, and a Lys-Thr/Ser-Glu-Leu C terminus. At an amino acid level, AG4 sequences are slightly more similar to mammalian AGR2 than nAG (e.g. 75% vs 69% identity between human AGR2 and Notophthalmus AG4 and nAG, respectively). Intriguingly, the AG4 sequences are also the only ones that diverge from the rest of cluster with respect to their thioredoxin active site/C-terminus signature (S1 Fig).

Despite the fact that they possess an identical ER-retention signal (Lys-Thr-Glu-Leu), human AGR2 is apparently largely retained in the ER as detailed in the Introduction, while nAG is apparently secreted. We hypothesised that, despite its divergent C-terminal sequence, AG4 might behave in a similar manner to AGR2 and also be an ER-retained protein. In order to provide an initial test of this hypothesis, we transfected 293T cells with constructs expressing wild-type (WT), C72A, or C72S either without a tag (untagged), with a myc-tag plus (myc + 25) or myc-tag minus (myc-tag without 25 aa). The conditioned media were analysed by western blotting and equivalent amounts of protein were added to blastemal cells growing in microwells. The data are given for the number (n) of microwells from 3 (in case of C72S) or 4 (other proteins) independent preparations of blastemal cells, normalised to the value for GFP.

Fig 4. Mutation of Cys72 leads to loss of activity. Cos7 cells were transfected with constructs expressing wild-type (WT), C72A, or C72S either without a tag (untagged), with a myc-tag plus (myc + 25) or myc-tag minus (myc-tag without 25 aa). The conditioned media were analysed by western blotting and equivalent amounts of protein were added to blastemal cells growing in microwells. The data are given for the number (n) of microwells from 3 (in case of C72S) or 4 (other proteins) independent preparations of blastemal cells, normalised to the value for GFP.

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Fig 5. Representative phylogenetic tree highlighting the species distribution of selected anterior gradient genes. For clarity only relevant support values are shown. Note that proteins belonging to the AG4 cluster have, so far, only been found in salamanders while the nAG and XAG2 grouping contains both salamander and anuran sequences. The tree was calculated with MrBayes 3.1 using likelihood model parameters: nst = 6; rates = invgamma. The calculation was run for 10 million generations, sampling every 500 generations; the initial 5,000 trees were discarded. An equivalent clustering of sequences was obtained using a maximum-likelihood approach as implemented in Phyml3 using the LG+G+I model and 1000 bootstrap replicates.

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Table 1. Differential secretion of AG family members by cultured cells.

| Protein  | % secreted protein |
|----------|-------------------|
| nAG (newt) | 10.6 ± 2.7        |
| AGR2 (human) | 1.65 ± 0.5       |
| AG4 (newt)  | 0.5 ± 0.1         |

HEK293T cells were transfected in parallel with plasmids expressing the wild-type (untagged) proteins nAG, human AGR2 or newt AG4. After 72 h the conditioned medium and cell lysate for each dish was processed and analysed by western blotting as described in the Materials and Methods section. The results are expressed as the percentage of the total immunoreactive AG protein in the dish that is secreted, with the results (mean ± SD) averaged over four independent transfections.

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Discussion

Mutation of the single cysteine residue in nAG led to protein degradation after transfection, possibly by the ERAD pathway. This loss was inhibited by extensions of the sequence at the C terminus, probably by interfering with the retrieval to the ER by the KDEL receptors, thus leading to more rapid transit through the secretory pathway. This serendipitous finding allowed us to obtain enough of the mutant nAG proteins to establish that the cysteine residue is essential for the S phase entry activity of the extended wild-type protein. The secreted protein in the conditioned medium predominantly migrated on non-reducing SDS gels as a monomer band, so it is likely that the sulfhydryl group was not involved in disulfide-bonded homodimer formation. This finding was in agreement to what had been reported for recombinant human AGR2 [14], although results using cell lysates suggested otherwise [17]. It is possible that the mechanism of action could involve formation of a heterodimeric disulfide-bonded intermediate with Prod1 on the cell surface, comparable to that observed between AGR2 and mucin [10], although the presence of such a covalent bond has been questioned [21]. Although, as in the case of others [14], we have been unable to detect any PDI activity using conventional biochemical assays, the fact that the proliferation-inducing activity of nAG is abrogated in the absence of Cys72 strongly suggests that this residue is able to form a mixed disulfide bond, perhaps in the presence of other proteins or small molecules yet to be found.

The hypothesis that secreted nAG acts on blastemal cells via the GPI-linked Prod1 receptor [4] is supported by our data showing that the S phase entry activity of nAG is inhibited by an antibody to Prod1. It has recently been reported that mouse AGR2 acts via the orphan membrane receptor C4.4A in models of pancreatic adenocarcinoma [37]. The administration of antibodies against C4.4A reduced growth and metastasis in mice [37]. It is possible that the ability of AG proteins to signal through GPI-linked TFP superfamily members is conserved through evolution, but the mechanism of action is a subject for future study.

Previous studies have highlighted that some members of the AG family are not present in amniotes, and that these proteins may contribute to the enhanced regeneration of appendages observed in amphibians [38]. Our results are broadly consistent with this suggestion but, as our phylogenetic analysis encompassed a wider variety of sequences and species, we found two significant differences: first, it is the cluster encompassing nAG and XAG2 that only contains amphibian sequences, and second, the AGR1 cluster does include amniote, but not mammalian, sequences and it may consist of more than one parologue. The Caudata AG4 members are closest in amino acid sequence identity to Amniota AGR2 but their dissimilar active site and ER retention signals suggest that they are experiencing distinct selective pressures to AGR2, perhaps due to the co-existence of multiple AG proteins that might have partially overlapping functions. We speculate that the importance of nAG for regeneration in Caudata, and XAG2 for development in Anura, reflects the evolution of a branch where the AG protein is secreted and acts in the extracellular space (green in Fig 5). The AG4 branch (purple in Fig 5) could maintain the essential ER-related functions proposed for AGR2. It is interesting that the ER-retention signal for AG proteins varies across the different families, suggesting that novel activities might evolve readily by modulating the cellular localisation and/or the degree of retention of these proteins in the ER. Much remains to be understood about the diversification of the AG protein family, and its roles in development, regeneration and cancer.

Supporting Information

S1 Fig. Multiple sequence alignment of representative anterior gradient proteins. The positions are coloured by chemical properties and conservation as defined by the ClustalX scheme. (TIF)
S1 Table. List of anterior gradient nucleotide sequences. The details of anterior gradient nucleotide sequences used in the phylogenetic analyses with their accession codes are listed here.

(XLS)

Author Contributions
Conceived and designed the experiments: KSG AGG PCD JPB. Performed the experiments: KSG AGG JPD JWG AK PBG JPB. Analyzed the data: KSG AGG JPD JPC JPB. Contributed reagents/materials/analysis tools: JPD JWG AK PBG. Wrote the paper: AGG JPB.

References
1. Sive HL, Hattori K, Weintraub H (1989). Progressive determination during formation of the anteroposterior axis in Xenopus laevis. Cell. 9: 58:171–180. PMID: 2752418.
2. Aberger F, Weidinger G, Grunz H, Richter K (1998). Anterior specification of embryonic ectoderm: the role of the Xenopus cement gland-specific gene XAG-2. Mech Dev. 72:115–130. PMID: 9539957.
3. Tereshina MB, Ermakova GV, Ivanova AS, Zaraisky AG (2014) Ras-dva1 small GTPase regulates telencephalon development in Xenopus laevis embryos by controlling Fgf8 and Agr signaling at the anterior border of the neural plate. Biol Open. 3:192–203. doi: 10.1242/bio.20147401 PMID: 24570397; PubMed Central PMCID: PMC4001240.
4. Kumar A, Godwin JW, Gates PB, Garza-Garcia AA, Brockes JP (2007) Molecular basis for the nerve dependence of limb regeneration in an adult vertebrate. Science. 318:772–777. doi: 10.1126/science.1147710 PMID: 17975060; PubMed Central PMCID: PMC2696928.
5. Morais da Silva SM, Gates PB, Brockes JP (2002) The newt ortholog of CD59 is implicated in proximo-distal identity during amphibian limb regeneration. Dev cell. 3:547–555. PMID: 12408806.
6. Thompson DA, McPherson LA, Carmeci C, deConinck EC, Weigel RJ (1997) Identification of two estrogen receptor transcripts with novel 5' exons isolated from a MCF7 cDNA library. J Steroid Biochem Mol Biol. 62:143–153. PMID: 939949.
7. Brychtova V, Vojtesek B, Hrstka R (2011) Anterior gradient 2: a novel player in tumor cell biology. Cancer Lett. 304:1–7. doi: 10.1016/j.canlet.2010.12.023 PMID: 21371820.
8. Chevet E, Fessart D, Delom F, Mulot A, Vojtesek B, Hrstka R, et al. (2013) Emerging roles for the pro-oncogenic anterior gradient-2 in cancer development. Oncogene. 32:2499–2509. doi: 10.1038/onc.2012.346 PMID: 22945652.
9. Zhao F, Edwards R, Dizon D, Afrasiabi K, Mastroianni JR, Geyfman M, et al. (2010) Disruption of Paneth and goblet cell homeostasis and increased endoplasmic reticulum stress in Agr2-/- mice. Dev Biol. 338:270–279. doi: 10.1016/j.ydbio.2009.12.008 PMID: 20025862; PubMed Central PMCID: PMC2937056.
10. Park SW, Zhen G, Verhaeghe C, Nakagami Y, Nyguenvu LT, Barczak AJ, et al. (2009) The protein disulfide isomerase AGR2 is essential for production of intestinal mucus. Proc Natl Acad Sci U S A. 106: 6950–6955. doi: 10.1073/pnas.0808722106 PMID: 19359471; PubMed Central PMCID: PMC2678445.
11. Gupta A, Wodziak D, Tun M, Bouley DM, Lowe AW (2013) Loss of anterior gradient 2 (Agr2) expression results in hyperplasia and defective lineage maturation in the murine stomach. J Biol Chem. 288: 4321–4333. doi: 10.1074/jbc.M112.433086 PMID: 23209296; PubMed Central PMCID: PMC3567683.
12. Chen YC, Lu YF, Li IC, Hwang SP (2012) Zebrafish Agr2 is required for terminal differentiation of intestinal goblet cells. PLoS One. 7: e34408. doi: 10.1371/journal.pone.0034408 PMID: 22514630; PubMed Central PMCID: PMC3326001.
13. Fletcher GC, Patel S, Tyson K, Adam PJ, Schenker M, Loader JA, et al. (2003) hAG-2 and hAG-3, human homologues of genes involved in differentiation, are associated with oestrogen receptor-positive breast tumours and interact with metastasis gene C4-4a and dystroglycan. Br J Cancer. 88:579–585. doi: 10.1038/sj.bjc.6600740 PMID: 12592373; PubMed Central PMCID: PMC2377166.
14. Patel P, Clarke C, Barraclough DL, Jowitt TA, Rudland PS, Barraclough R, et al. (2013) Metastasis-promoting anterior gradient 2 protein has a dimeric thioredoxin fold structure and a role in cell adhesion. J Mol Biol. 425: 929–943. doi: 10.1016/j.jmb.2012.12.009 PMID: 23274113.
15. Nguyen VD, Ruddock LW, Salin M, Wierenga RK (2011) Crystal structure of the human anterior gradient protein 3.
16. Gupta A, Dong A, Lowe AW (2012) AGR2 gene function requires a unique endoplasmic reticulum localization motif. J Biol Chem. 287:4773–4782. doi: 10.1074/jbc.M111.301531 PMID: 22184114; PubMed Central PMCID: PMC3281655.
17. Ryu J, Park SG, Lee PY, Cho S, Lee do H, Kim GH, et al. (2013) Dimerization of pro-oncogenic protein Anterior Gradient 2 is required for the interaction with BiP/GRP78. Biochem Biophys Res Commun. 430: 610–615. doi: 10.1016/j.bbrc.2012.11.105 PMID: 23220234.
18. Dong A, Gupta A, Pai RK, Tun M, Lowe AW (2011) The human adenocarcinoma-associated gene, AGR2, induces expression of amphiregulin through Hippo pathway co-activator YAP1 activation. J Biol Chem. 286:18301–18310. doi: 10.1074/jbc.M110.215707 PMID: 21454516; PubMed Central PMCID: PMC3093902.
19. Dong A, Wodziak D, Lowe AW (2015) Epidermal growth factor receptor (EGFR) signaling requires a specific endoplasmic reticulum thioredoxin for the post-translational control of receptor presentation to the cell surface. J Biol Chem. 290: 8016–8027. doi: 10.1074/jbc.M114.623207 PMID: 25666625; PubMed Central PMCID: PMC4375459.
20. Ivanova AS, Shandarin IN, Ermakova GV, Minin AA, Tereshina MB, Zaraisky AG (2015). The secreted factor Agf1 missing in higher vertebrates regulates fins regeneration in Danio rerio. Sci Rep. 5: 8123. doi: 10.1038/srep08123 PMID: 21118959; PubMed Central PMCID: PMC3220234.
21. Bergstrom JH, Berg KA, Rodriguez-Pineiro AM, Stecher B, Johansson ME, Hansson GC (2014) AGR2, an endoplasmic reticulum protein, is secreted into the gastrointestinal mucus. PLoS One. 9: e104186. doi: 10.1371/journal.pone.0140186 PMID: 25111734; PubMed Central PMCID: PMC4128659.
22. Clarke C, Rudland P, Barraclough R (2015) The metastasis-inducing protein AGR2 is O-glycosylated upon secretion from mammary epithelial cells. Mol Cell Biochem. 408: 245–252. doi: 10.1007/s11010-015-2502-3 PMID: 26169982.
23. Kumar A, Godwin JW (2010) Preparation and culture of limb blastema stem cells from regenerating larval and adult salamanders. Cold Spring Harb Protoc. pdb prot5367. doi: 10.1101/pdb.prot5367 PMID: 20150126.
24. Blassberg RA, Garza-Garcia A, Janmohamed A, Gates PB, Brockes JP (2011) Functional convergence of signalling by GPI-anchored and anchorless forms of a salamander protein implicated in limb regeneration. J Cell Sci. 124: 47–56. doi: 10.1242/jcs.076331 PMID: 21118959; PubMed Central PMCID: PMC3001407.
25. Baddar NW, Woodcock MR, Khatri S, Kump DK, Voss SR (2015) Sal-Site: research resources for the Mexican axolotl. Methods Mol Biol. 1290:321–336. doi: 10.1007/978-1-4939-2495-0_25 PMID: 25740497.
26. Looso M, Braun T (2015) Data mining in newt-omics, the repository for omics data from the newt. Meth-ods Mol Biol. 1290:337–351. doi: 10.1007/978-1-4939-2495-0_26 PMID: 25740498.
27. Abdullayev I, Kirkham M, Bjorklund AK, Simon A, Sandberg R (2013) A reference transcriptome and inferred proteome for the salamander Notophthalmus viridescens. Exp Cell Res. 319:1187–1197. doi: 10.1016/j.yexcr.2013.02.013 PMID: 23454802.
28. Edgar RC (2004) MUSCLE: a multiple sequence alignment method with reduced time and space complexity. BMC Bioinformatics. 5: 113. doi: 10.1186/1471-2105-5-113 PMID: 15318951; PubMed Central PMCID: PMC3177056.
29. Katoh K, Standley DM (2014) MAFFFT: iterative refinement and additional methods. Methods Mol Biol. 1079:131–46. doi: 10.1007/978-1-62703-646-7_8 PMID: 24170399.
30. Waterhouse AM, Procter JB, Martin DM, Clamp M, Barton GJ (2009) Jalview Version 2—a multiple sequence alignment editor and analysis workbench. Bioinformatics. 25:1189–1191. doi: 10.1093/bioinformatics/btp333 PMID: 1915095; PubMed Central PMCID: PMC2672624.
31. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, et al. (2007) Clustal W and Clustal X version 2.0. Bioinformatics. 23: 2947–2948. doi: 10.1093/bioinformatics/btm404 PMID: 17846036.
32. Keane TM, Naughton TJ, McNemey JO (2004) ModelGenerator: amino acid and nucleotide substitution model selection. National University of Ireland, Maynooth, Ireland.
33. Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O (2010) New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. Syst Biol. 59: 307–321. doi: 10.1093/sysbio/syq010 PMID: 20525638.
34. Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Hohna S, et al. (2012) MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. Syst Biol. 61:539–542. doi: 10.1093/sysbio/sys029 PMID: 22357727; PubMed Central PMCID: PMC3329765.
35. Nylander JA, Wilgenbusch JC, Warren DL, Swofford DL (2008) AWTY (are we there yet?): a system for graphical exploration of MCMC convergence in Bayesian phylogenetics. Bioinformatics. 24:581–583. doi: 10.1093/bioinformatics/btm388 PMID: 17766271.

36. Huson DH, Scornavacca C (2012) Dendroscope 3: an interactive tool for rooted phylogenetic trees and networks. Syst Biol. 61:1061–1067. doi: 10.1093/sysbio/sys062 PMID: 22780991.

37. Arumugam T, Deng D, Bover L, Wang H, Logsdon CD, Ramachandran V (2015) New Blocking Antibodies against Novel AGR2-C4.4A Pathway Reduce Growth and Metastasis of Pancreatic Tumors and Increase Survival in Mice. Mol Cancer Ther. 14: 941–951. doi: 10.1158/1535-7163.MCT-14-0470 PMID: 25646014.

38. Ivanova AS, Tereshina MB, Ermakova GV, Belousov VV, Zaraisky AG (2013) Agr genes, missing in amniotes, are involved in the body appendages regeneration in frog tadpoles. Sci Rep. 3:1279. doi: 10.1038/srep01279 PMID: 23412115; PubMed Central PMCID: PMC3573343.