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## Molecular cloning and characterization of the mouse *Acdp* gene family

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### Abstract

**Background:** We have recently cloned and characterized a novel gene family named ancient conserved domain protein (ACDP) in humans. To facilitate the functional study of this novel gene family, we have cloned and characterized *Acdp*, the mouse homologue of the human ACDP gene family.

**Results:** The four *Acdp* genes (*Acdp1*, *Acdp2*, *Acdp3* and *Acdp4*) contain 3,631 bp, 3,244 bp, 2,684 bp and 2,743 bp of cDNA sequences, and encode deduced proteins of 951, 874, 713 and 771 amino acids, respectively. The mouse *Acdp* genes showed very strong homologies (>90%) in both nucleotide and amino acid sequences to their human counterparts. In addition, both nucleotide and amino acid sequences within the Ancient Conserved Domain (ACD) are highly conserved in many different taxonomic species. Particularly, *Acdp* proteins showed very strong AA homologies to the bacteria CorC protein (35% AA identity with 55% homology), which is involved in magnesium and cobalt efflux. The *Acdp* genes are widely expressed in all tissues tested except for *Acdp1*, which is only highly expressed in the brain with low levels of expression in kidney and testis. Immunostaining of *Acdp1* in hippocampus neurons revealed a predominant localization on the plasma membrane.

**Conclusion:** The *Acdp* genes are evolutionarily conserved in diverse species and ubiquitously expressed throughout development and adult tissues suggesting that *Acdp* may be an essential gene. *Acdp* showed strong homology to bacteria CorC protein and predominantly localized on the plasma membrane. These results suggest that *Acdp* is probably a family of proteins involved in ion transport in mammalian cells

### Background

We have recently cloned and characterized a novel gene family named ancient conserved domain protein (ACDP) which encodes four protein members in humans [1]. We

found that this gene family is evolutionarily conserved in diverse species ranging from bacteria, yeast, *C. elegans*, and *D. melanogaster* to mammals. The sequence conservation and the presence of multiple members within a species

may imply functional importance associated with the genes. To facilitate the functional analysis of the *ACDP* gene family, we cloned and characterized *Acdp*, the mouse homologue of the human *ACDP* gene family.

## Results

### Molecular cloning of the *Acdp* gene family

To clone the mouse *Acdp* genes, the human *ACDP* cDNA and predicted protein sequences were used to search the mouse EST database with the *blastn* and *tblastn* programs. Mouse EST markers corresponding to each *Acdp* member were then identified. For example, EST H3086H12-5 corresponds to *Acdp1*, W98010 for *Acdp2*, 603299135F1 for *Acdp3* and BG083791 for *Acdp4*. A modified oligo-dT with a M13 tail was used for the RT reaction. A forward primer from each EST marker and the M13 primer (oligo-dT tail) were used to amplify the 3' UTR sequence for each corresponding *Acdp* gene from the RT products. To obtain 5'-end coding sequences for the *Acdp* genes, we conducted a series nested PCR with combinations of mouse and human primers. The 5' UTR sequences were identified by directly sequencing BAC DNA containing the corresponding *Acdp* genes. The BAC clones were identified by screening a CITB mouse BAC DNA library (Research Genetics). The 5' UTR sequences obtained from above were further confirmed by RT-PCR. The *Acdp1* gene contains 3,631 bp of nucleotide sequence and encodes a predicted protein with 951 amino acids (AA). The other three *Acdp* genes (*Acdp2*, 3 and 4) contain 3,244 bp, 2,684 bp and 2,743 bp of cDNA sequences, and encode deduced proteins of 874 amino acids, 713 amino acids and 771 amino acids, respectively.

### Tissue distribution

Northern blot analyses of the *Acdp* gene family were carried out using membranes purchased from Origene. A total of 12 mouse tissues were included in the study (Fig. 1). Due to sequence homologies between each *Acdp* member within the conserved domain, probes for Northern blots were PCR fragments from the last exon and the 3' untranslated region sequences. The mouse *Acdp* messages showed almost the same tissue distributions as the human *ACDP* genes. *Acdp1* message is highly expressed in the brain, while kidney and testis also showed low levels of expression. *Acdp2* showed higher expressions in the brain, kidney and liver. However, the *Acdp2* transcript was not present in the skeleton muscle and skin, and it showed very low levels of expression in the rest of tissues. *Acdp3* and *Acdp4* showed different levels of expression in all tissues tested; the highest expressions for *Acdp3* were observed in the brain, kidney, liver and heart, and the highest expressions for *Acdp4* were observed in the kidney, small intestine and testis. The expression levels for *Acdp3* and 4 in skeleton muscle were barely detectable; however,  $\beta$ -actin showed normal expression suggesting that the

results were not a consequence of bad RNA quality (data not shown). The ubiquitous expression pattern may be taken as another indication of the functional importance of *Acdp* proteins in fundamental biological processes in addition to the sequence conservation in evolutionarily divergent species.

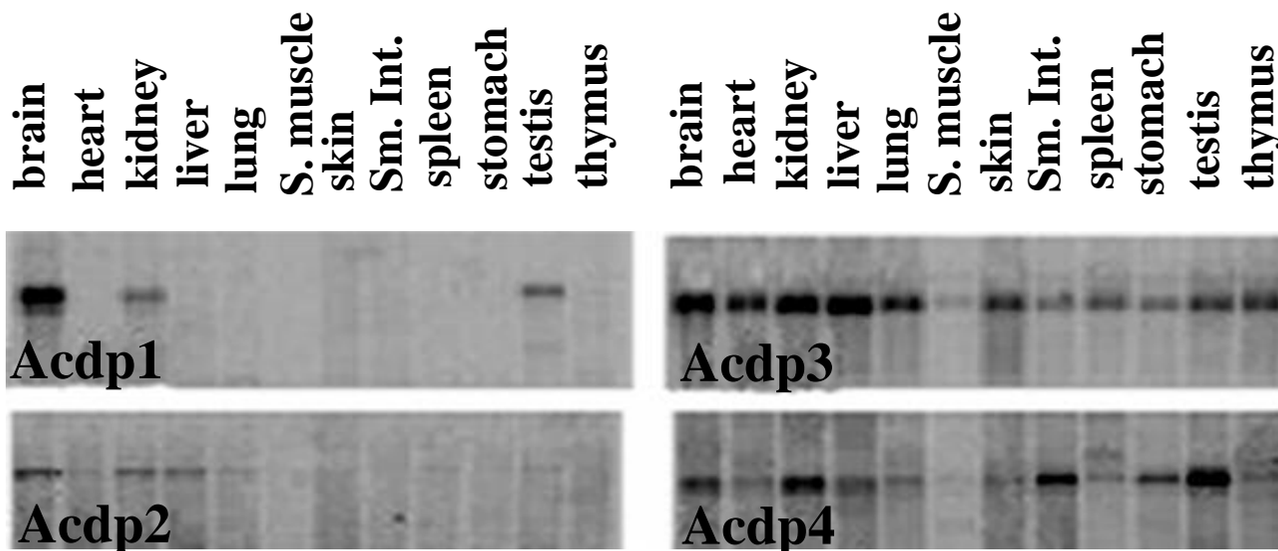
### Chromosomal location

Radiation hybrid mapping indicated that the *Acdp1* gene maps to chromosome 19 between markers *D19Mit119* (34.3 cR proximal) and *D19Mit112* (13.6 cR distal). The *Acdp2* gene maps slightly more distal to the *Acdp1* on chromosome 19 between *D19Mit9* (2.4 cR proximal) and *D19Mit38* (15.1 cR distal). The *Acdp3* and *Acdp4* genes map to chromosome 1 within one BAC clone (RP23-294I17), proximal to marker *D1Mit171* (17.4 cR). These regions are syntenic to the human counterparts.

### Sequence homology and molecular characteristics

The mouse *Acdp* genes showed very strong homologies of both nucleotide and AA sequences to the human *ACDP* genes (Table 1). The highest homologies were observed between the human *ACDP2* and the mouse *Acdp2* gene (91% of nucleotide identity, 97% of AA identity and 99.4% of AA homology). In addition, the 5' UTR nucleotide sequences (20 bp of nucleotides before start codon) also showed high homologies to the human homologs, for example, the *Acdp2* 5' UTR sequence showed 95% identities to its human homolog. However, the homologies in the 3' UTR sequences (20 bp of nucleotides after stop codon) were much lower (40–55%) for all *Acdp* genes except *Acdp4* (90% identity to its human homolog). The ancient conserved domain (ACD) has 55.3% of AA identity and 83.3% of homology between all mouse and human *ACDP* proteins (Fig. 2). The ACD domain is evolutionarily conserved in divergent species ranging from bacteria, yeast, *C. elegans*, *D. melanogaster*, mouse to human (Fig. 3). Particularly, as shown in Fig. 3, *Acdp* proteins showed very strong AA homology to bacteria CorC protein (35% AA identity with 55% homology), which is involved in magnesium and cobalt efflux [7]. High AA homology was also observed between the *Acdp* proteins and the yeast *Amip3* protein (35% AA identity with 56% homology). The *Amip3* is likely to be a homologous to the bacteria CorC protein. The *Amip3* mutants confer resistance to copper toxicity (Personal communication with Dr. V.C. Culotte, John Hopkins Bloomberg, School of Public Health). The evolutionary relationships among those proteins are illustrated by a phylogenetic tree constructed based on the AA homology of proteins (Fig. 4).

We found that all mouse *Acdp* members contain four distinct transmembrane domains (Fig. 5), two CBS domains and a DUF21 domain that are found in bacteria CorC and yeast *Amip3* proteins. CBS domains are small intracellular



**Figure 1**  
Northern blot analyses of the *Acdp* gene family. S. muscle represents skeletal muscle, Sm. Int. represents small intestine. Multiple Choice Northern Blot filters were purchased from Origene.

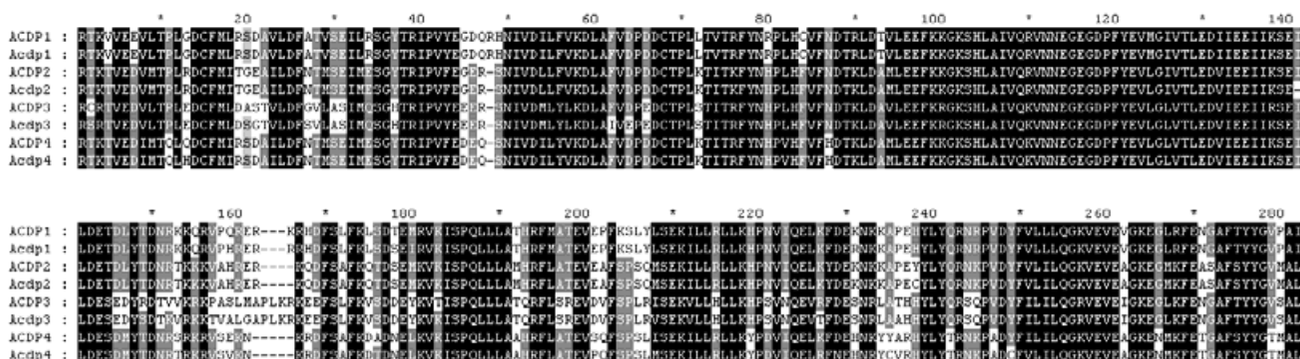
**Table 1: Nucleotide and amino acid homologies (%) between human ACDP and mouse Acdp members.**

Acdp	Acdp1	Acdp2	Acdp3	Acdp4
DNA in coding region	89	91	86	87
AA identity	94	97	86	89
AA homology	98.8	99.4	94	96
5' UTR	85	95	80	80
3' UTR	40	55	50	90

modules that are mostly found in 2 or four copies within a protein. Pairs of CBS domains dimerise to form a stable globular domain [8]. DUF21 (CD: pfam01959.9) is a newly defined domain with unknown function. This domain is a transmembrane region and found to be located in the N-terminus of the proteins adjacent to two intracellular CBS domains <http://www.ncbi.nlm.nih.gov/Structure/cdd/cddsrv.cgi?uid=pfam01959>. A cNMP-binding domain (cyclic nucleotide-monophosphate-binding domain) was found in all *Acdp* members.

In addition, *Acdp1* contains an Alanine-rich region (2–10: AAAAAAAAAA), a Leucine-rich region (204–257: LLRVR-

PRLYGPGDLLPPAWLRLALGALLLALSALF SGLRLSLLSLDPVELRVL), a Proline-rich region (78–130: PGPPVPAAPVPAPSLA PGENGTGDWAPRLVFIEEPPGAG-GAAPSAVPTRPPGP), and two amidation sites (917–920: MGKK; 926–929: SGRK). *Acdp2* has a glycine-rich region (201–222: GAGGSGSASGTVGGKGGAGVAG). *Acdp3* possesses a large alanine-rich region (2–261) and a large leucine-rich region (201–299). *Acdp4* contains a leucine zipper pattern (185–206: LVMVLLVLSGIFSGLNGLMAL) and an amidation site (7–10: GRRR).



**Figure 2**  
 Amino acid sequence homology alignment for all of the ACDP and Acdp genes within the ACD domain. The sequence data for the Acdp genes have been deposited in GenBank under accession number AF202994 (Acdp1), AF216961 (Acdp2), AF216964 (Acdp3) and AF216963 (Acdp4). Identical amino acids or amino acids with very strong homologies among all proteins were shaded black. Identical amino acids or amino acids with very strong homologies in most of the proteins were shaded grey. Dot lines represent gaps for the alignment.

**Antibody production, Western results and subcellular localization**

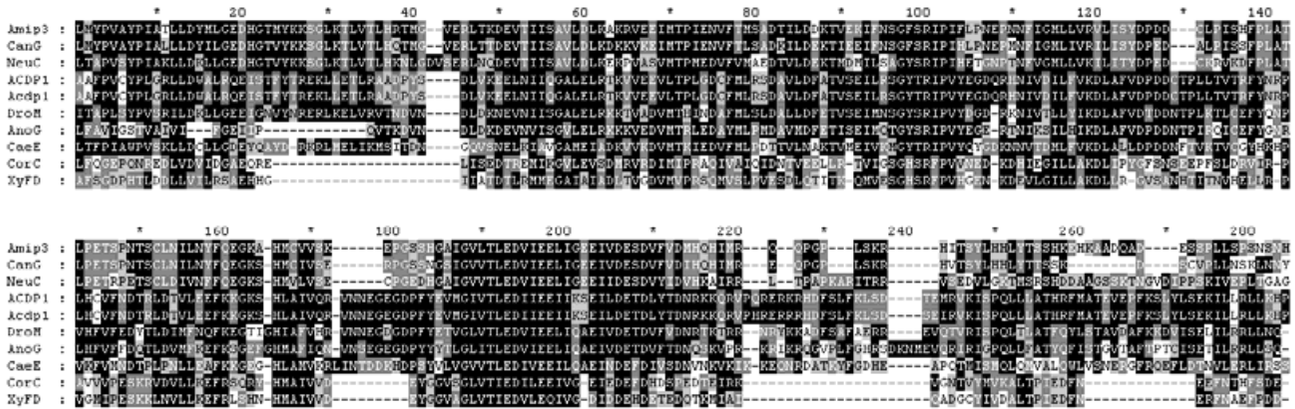
Peptides from Acdp1 N- (TSFLLRVYFQPGPPATAAPVP-SPT) and C- (TQQLTLPAAVPTTR) terminuses, conserved peptide from ACD domain of Acdp1 (HNIVDILFVKD-LAFVDPDDCTPLLTVTRE) were commercially synthesized (Sigma Genosys). These antigenic sites were predicted by software from Sigma Genosys and polyclonal antibodies for each peptide were produced by immunizing rabbits. To test the specificity of the antibodies, we conducted Western-blot analysis of mouse brain tissue extracts. As shown in Fig. 6A, the antibody produced by C-terminal peptide specifically detected Acdp1 (lane 3). The antibody generated by N-terminal peptide recognized Acdp4 in addition to Acdp1, although the reactivity to latter was significantly higher (Fig. 6A, lane 2). As expected, the antibody produced by the conserved sequence peptide detected all Acdp proteins (Fig. 6A, lane 1). To further determine the specificity of the antibody against the Acdp1 C-terminus, we analyzed extracts of HEK293, 3T3 and PC12 cells. The results are shown in Fig. 6B. Apparently, this antibody detected a specific band of Acdp1 in all cell lysates. Of note, shown in Fig. 6B are signals of 10 µg extracts of HEK293 cell lysates, 100 µg extracts of 3T3 and PC12 cell lysates. Thus, the expression levels of Acdp1 in these cell types vary a lot, with the highest expression in HEK293 cells. Nevertheless, these immunoblot results

support our analysis of brain tissue extracts that the antibody against Acdp1 C-terminus specifically recognizes Acdp-1.

The specificity of the Acdp1 C-terminus antibody suggests the possibility of using it to localize Acdp-1 within cells. Since Northern blot revealed almost exclusive expression of Acdp1 in the brain, we examined its subcellular localization in hippocampus neurons isolated from mouse embryos. The neurons were cultured on glass coverslips coated with a confluent monolayer of mouse cortical astrocytes in dishes. Immunostaining was using the Acdp1 C-terminus specific antibody. Confocal imaging revealed that Acdp1 is predominantly localized on the plasma membrane. A series of sections of a cell at the thickness of 0.5 micrometer clearly showed membrane location of Acdp1-immunoreactivity (Fig. 7), which is consistent with the observation of transmembrane domains within the Acdp proteins.

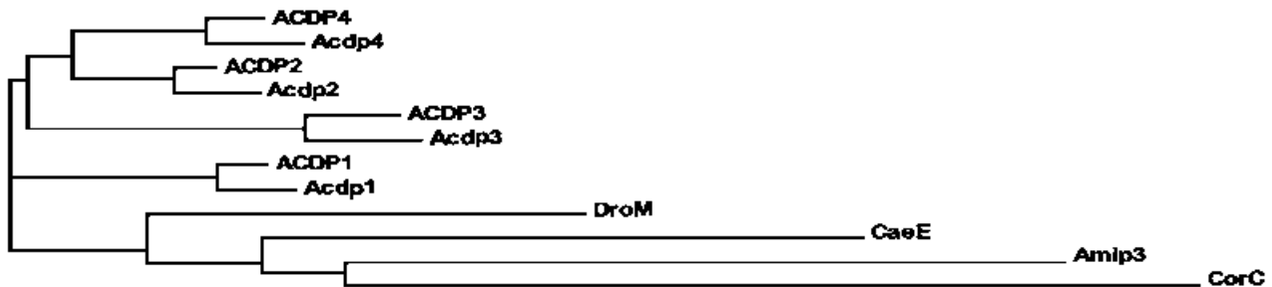
**Discussion**

Although the exact functions of the ACDP gene family are not yet elucidated, several lines of evidence suggest that ACDP genes may play an important role in biological processes. First, these genes are evolutionarily conserved in many phylogenetically divergent species; Second, multiple genes are present in a species; Third, these genes are



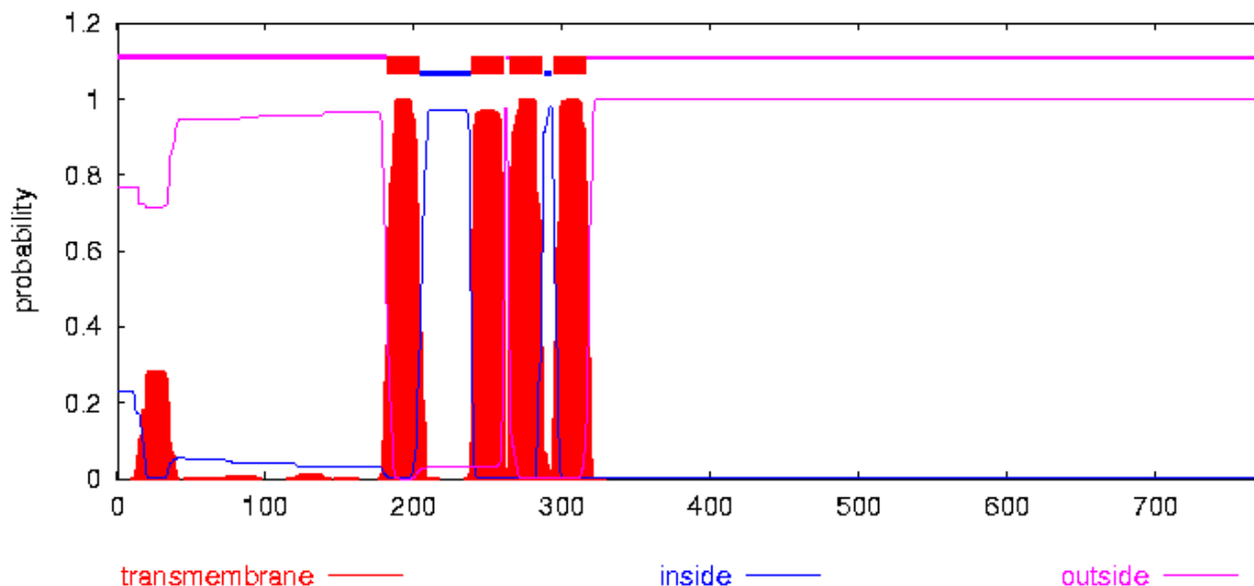
**Figure 3**

Amino acid sequence alignment showing the conservation of ACD domain in various species. Amip3 is a protein from *Saccharomyces cerevisiae* (NP\_014581). CanG is a protein from *Candida glabrata* (AAF33142). NeuC (EAA31204) is a hypothetical protein from *Neurospora crassa*. DroM is a gene product from *D. melanogaster*. The accession number for this gene is CG40084 in BDGP (Berkeley Drosophila Genome Project). AnoG represents a protein from the *anopheles gambiae* str. (EAA01004). CaeE (AAK77203) is a hypothetical protein from the *Caenorhabditis elegans*. CorC represents bacteria magnesium and cobalt efflux protein from the *Shewanella oneidensis*. XyFD is a hypothetical protein from the *Xylella fastidiosa* Dixon (ZP\_00038107).



**Figure 4**

Phylogenetic tree showing relationships among proteins containing the ACD domain from figure 2 and 3. The phylogenetic tree was constructed according to the calculation of the best match for the selected sequences. Abbreviations for each protein are the same as presented in figure 3.



**Figure 5**

Four transmembrane domains within Acdp4 protein. Transmembrane domains were predicted by the TMHMM program <http://www.cbs.dtu.dk/services/TMHMM-2.0/>. The plot shows the posterior probabilities of inside /outside/TM helix. At the top of the plot (between 1 and 1.2) the N-best prediction is shown. The plot is obtained by calculating the total probability that a residue sits in helix, inside, or outside summed over all possible paths through the model.

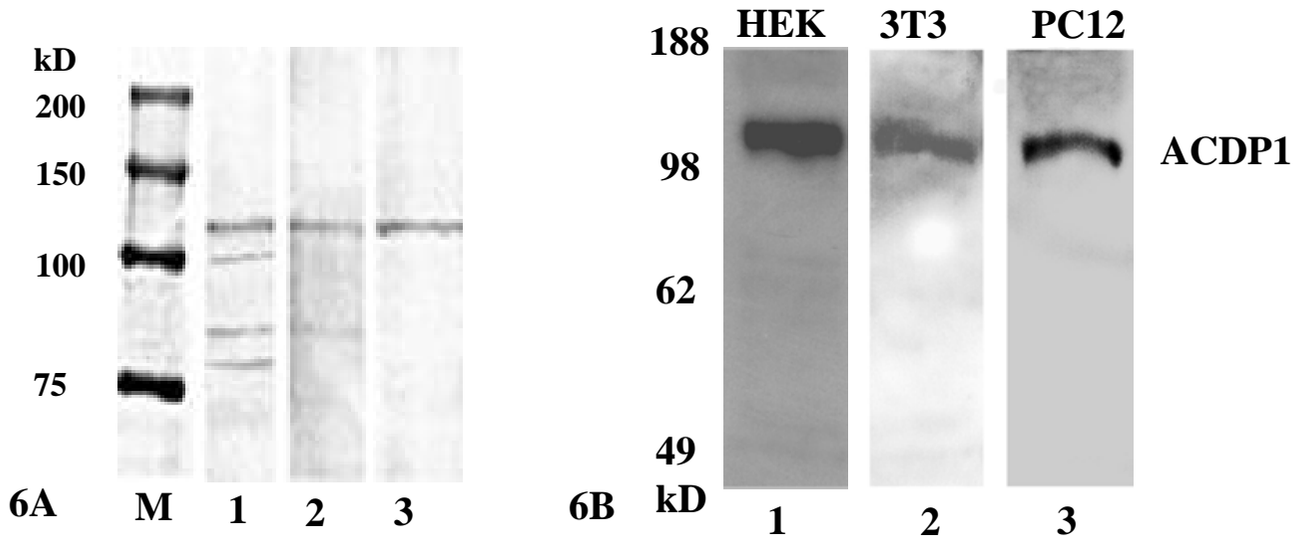
ubiquitously expressed throughout development and adult tissues (unpublished data). The cloning and characterization of the mouse *Acdp* gene family are a very important step towards the elucidation of the functions of this multigene family.

Sequence homology analyses revealed that Acdp proteins shared very strong AA homologies to the bacteria CorC protein and yeast Amip3 protein. CorA, B, C and D belong to a protein family involved in both influx and efflux of magnesium and cobalt. It has been shown that CorA mutants confer resistance to cobalt toxicity [9]. Amip3 appears to be a homologous to CorC protein which is involved in efflux of magnesium and cobalt in bacteria. Amip3 mutants confer resistant to copper toxicity. Acdp proteins also possess the domains that are found in bacteria CorC and yeast Amip3 proteins, such as the CBS domains, DUF21 domain and transmembrane domains. In addition, a cNMP-binding domain was found in all Acdp proteins, which is usually present in ion channels and cNMP-dependent kinases [10-13]. Using antibody

produced by Acdp1 C-terminal peptide, we have shown that Acdp1 is predominantly localized on the plasma membrane in hippocampus neurons. In our previous study, we found human ACDP proteins are predominantly localized in the nucleus in HeLa cells [1]. The discrepancy for Acdp localization in neuronal cells could be caused by non-specificity for previous antibody which was produced by the recombinant ACD domain, or different cells used. In current study, the Acdp1 C-terminus antibody only recognizes Acdp1 in brain tissue extracts as well as in HEK293, 3T3 and PC12 cell lysates, suggesting the specificity of the antibody. Our observations suggested that Acdp might be involved in ion transport in mammalian cells. However, more detailed functional studies are needed to demonstrate the real functions of these proteins.

### Conclusions

Our previous work has described human ACDP genes [1]. The new information of the current work includes: 1). It is the first time to report the existence of multiple Acdp



**Figure 6**

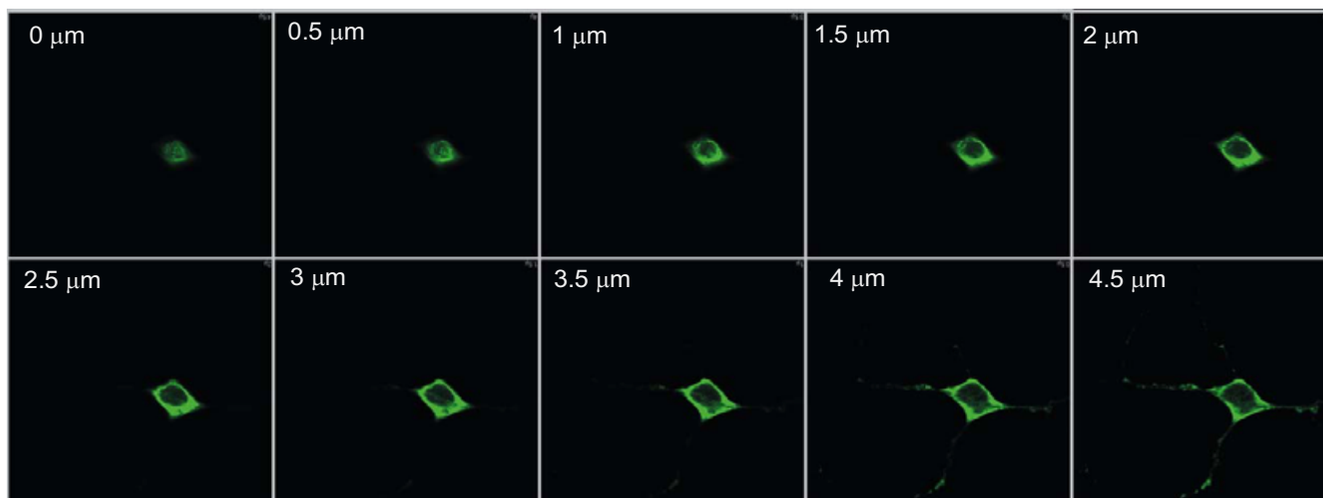
Fig. 6A: Immunoblot analysis of Acdp proteins in brain tissue extracts. Immunoblotting were carried out using a Western blotting detection system (ECL) (PIERCE). Lane 1, probed with antibody generated by conserved peptide. From top to bottom, each band corresponding to Acdp1 (115 kD), Acdp2 (100 kD), Acdp4 (90 kD) and Acdp3 (80 kD). Lane 2 and 3, probed with the Acdp1 antibodies generated by N-terminal and C-terminal peptides, respectively. Fig. 6B: Immunoblot analysis of Acdp1 in HEK293, 3T3 and PC12 cells. The blots were probed with the antibody against the C-terminus of Acdp-1. Lane 1, 10  $\mu$ g of HEK293 cell lysates. Lane 2 and 3, 100  $\mu$ g of 3T3 and PC12 cell lysates.

genes in other mammals in addition to human, while Acdp appears to be a single copy gene in lower organisms such as in *C. elegans*, yeasts and bacteria. We have also suggested the evolutionary relationships of Acdp genes in different species (phylogenetic tree); 2). Molecular cloning and characterization of murine Acdp gene family are essential for study of this novel gene family in animal model, e.g. for generation of knockout or transgenic models; 3). We have demonstrated both DNA and amino acid conservations between mouse and human for each Acdp gene and the whole Acdp gene family, which provide important information for the possibility of functional redundancy or overlap between Acdp members; 4). We have generated antibodies specific for Acdp1 and all Acdp proteins. The Acdp1 C-terminus antibody appears to specifically recognize Acdp1. Using this antibody, we have demonstrated that Acdp1 is predominantly localized on the plasma membrane in hippocampus neurons. These results represent an important step towards the characterization of functions for the Acdp gene family.

## Methods

### cDNA cloning

cDNA cloning of the *Acdp* gene family was performed based on human homologue sequences as previously reported [2]. Briefly, the human *ACDP* cDNAs and predicted AA sequences were used to search the mouse EST database for EST markers corresponding to each Acdp member. A forward primer within the human *ACDP* 5' cDNA coding region (after start codon) and a mouse reverse primer from the mouse EST marker were used to amplify the homologue sequence from mouse cDNA at very low annealing temperature (45–50°C). A nested PCR using an inside reverse primer from the mouse EST sequence and the same human forward primer was then carried out to amplify the specific mouse gene from the first round PCR products at high annealing temperature (62°C). The expected PCR products were directly excised from agarose gel and sequenced by an ABI377 automatic sequencer. The sequence was further confirmed using a forward primer from newly identified sequence and a

**Figure 7**

Subcellular localization of Acdp1 in hippocampus neurons. A series of confocal images from a cultured neuron stained with an anti-Acdp1 antibody. The step of each imaging section is 0.5  $\mu\text{m}$ , from the surface of the neuron (0  $\mu\text{m}$ ) to the middle plan (4.5  $\mu\text{m}$ ).

reverse primer from known sequence. Once most of the coding sequences were identified, partial sequence of exon 1 and the 5' UTR sequences were obtained by BAC DNA sequencing.

#### Northern blot analyses

Multiple Choice Northern Blot filters containing 12 different mouse tissues were purchased from Origene. The filters were probed for each Acdp gene with a PCR fragment (around 350 bp) from last exon and the 3' untranslated region labeled with  $\alpha$ - $^{32}\text{P}$  dCTP using the random primer extension system (Life Technologies). Hybridizations were carried out overnight. The filters were washed twice with washing buffer I (2 $\times$ SSC, 0.1% SDS) at 42°C for 15-min, and then washed twice with washing buffer II (0.25 $\times$ SSC, 0.1% SDS) at 65°C for 15-min. Washed filters were exposed to X-ray films for overnight or longer [3,4].

#### Antibody production and Western blot analyses

Peptides linked to KLH (keyhole limpet hemacyanin) from Acdp1 N- and C-terminals and the conserved domain (ACD) were used for generation of antibodies specifically for Acdp1 and all Acdp members as reported, respectively [5]. Western blots were carried out Using ECL (PIERCE) as described previously [1]. The membranes were washed extensively after incubation with primary and secondary antibodies and were then developed with X-ray films with optimal exposure time.

#### Chromosome localization

The T31 mouse radiation hybrid panel from Research Genetics was used to map the chromosome location of each Acdp member. Primers from 3' UTR of each Acdp member were used to amplify the 100 radiation hybrid clones representing the mouse genome. The data were submitted to the Jackson Laboratory Mouse Radiation Hybrid Database for analysis.

#### Sequence analyses

Sequence assembly was performed with program Sequencher (Gene Codes Corp). Protein and DNA homology searches were carried out with tblastn, tblastx, blastp and blastn programs <http://www.ncbi.nlm.nih.gov/BLAST/>. Multiple sequence alignments were performed with GeneDoc and pairwise sequence alignment <http://pir.georgetown.edu/pirwww/search/pairwise.html>. Multiple programs including BCM Search Launcher <http://searchlauncher.bcm.tmc.edu/>, ProfileScan <http://hits.isb-sib.ch/cgi-bin/PFSCAN>, sequence motif search <http://motif.genome.ad.jp/>, ExPASy <http://www.expasy.ch/> and 3Dpssm <http://www.bmm.icnet.uk/~3dpssm/> were used for searching sequence features of known protein. Phylogenetic tree was constructed by Clustalw program (version 1.81) using UPGMA (Unweighted Pair Group Method using Arithmetic averages) algorithm [14].

### Neuronal cell preparation and immunostaining

Hippocampal neuron cultures were prepared as previously reported [6]. In brief, the hippocampuses were dissected out from mouse embryos at 16 days *in utero*. The tissues were then incubated for 20 min at 37°C in MEM (minimum essential medium) modified for suspension culture (Life Technologies) plus 0.25% trypsin (Life Technologies). The dissociated hippocampal neurons were plated on glass coverslips coated with a confluent monolayer of mouse cortical astrocytes obtained as described below. The neurons were maintained at 37°C in a humidified atmosphere with 5% CO<sub>2</sub>. Cortical astrocytes dissociated from newborn mouse cortices were grown in culture flasks at 37°C in a humidified atmosphere with 5% CO<sub>2</sub> until confluent. The cells were exposed to 10<sup>-5</sup> M cytosine arabinoside (Sigma) and cultured for additional 12–24 hrs at 37°C. After remove of the media with cellular debris, the cells were used for coating coverslips.

For immunostaining, neuronal cells on the coverslips were first fixed in PBS containing 4% paraformaldehyde (PFA) for 12 hrs at 4°C and then incubated in a solution containing 4% PFA and 0.4% Triton X-100 at 4°C for 1 hr. After washing with PBS three times, the cells were incubated with a blocking solution containing 1:30 normal goat serum, and subsequently incubated with a rabbit polyclonal anti-ACDP antibody (1:3000) overnight at 4°C. After extensive washing with 1% goat serum PBS solution, the cells were incubated with an Alex 488 conjugated secondary antibody (1:100 in 1% goat serum PBS solution, Molecular Probes) for 3 hrs at room temperature. Following final washes with 1% goat serum PBS solution, the neuronal cells on the coverslips were covered with a glycerol-based anti-photobleach medium. The cells were viewed under a confocal microscope (Carl Zeiss). Images were captured with a CCD camera and acquired by the Scion Image software (Scion Corporation, Frederick, MD).

### Gene bank accession number

The cDNA sequences for the *Acdp* gene family have already been deposited in gene bank with accession numbers AF202994 (*Acdp1*), AF216961 (*Acdp2*), AF216964 (*Acdp3*) and AF216963 (*Acdp4*).

### Authors' contributions

This study was designed by CYW and JXS. The study was performed as follows: CYW, PY, JDS and HA contributed to the molecular cloning part of the manuscript. CYW and SP were responsible for the *Acdp* antibody production. JGG and JLG did immunostaining for *Acdp1* localization in neuronal cells. CYW, DG and ZD carried out the Western analyses.

### Acknowledgements

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