CLINICAL STUDY

Autosomal dominant neurohypophyseal diabetes insipidus in a Swiss family, caused by a novel mutation (C59Δ/A60W) in the neurophysin moiety of prepro-vasopressin-neurophysin II (AVP-NP II)

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Abstract

Objective: To study clinical, morphological and molecular characteristics in a Swiss family with autosomal dominant familial neurohypophyseal diabetes insipidus (adFNDI).

Participants and methods: A 15-month-old girl presenting with symptoms of polydipsia and polyuria was investigated by water deprivation test. Evaluation of the family revealed three further family members with symptomatic vasopressin-deficient diabetes insipidus. T1-weighted magnetic resonance images of the posterior pituitary were taken in two affected adult family members and molecular genetic analysis was performed in all affected individuals.

Results: The water deprivation test in the 15-month-old child confirmed the diagnosis of vasopressin-deficient diabetes insipidus and the pedigree was consistent with autosomal dominant inheritance. The characteristic bright spot of the normal vasopressin-containing neurophypophysis was absent in both adults with adFNDI. Direct sequence analysis revealed a new deletion (177–179ΔCGC) in exon 2 of the AVP-NP II gene in all affected individuals. At the amino acid level, this deletion eliminates cysteine 59 (C59Δ) and substitutes alanine 60 by tryptophan (A60W) in the AVP-NP II precursor; interestingly, the remainder of the reading frame remains unchanged. According to the three-dimensional structure of neurophysin, C59 is involved in a disulphide bond with C65.

Conclusions: Deletion of C59 and substitution of A60W in the AVP-NP II precursor is predicted to disrupt one of the seven disulphide bridges required for correct folding of the neurophysin moiety and thus disturb the function of neurophysin as the vasopressin transport protein. These data are in line with the clinical and morphological findings in the reported family with adFNDI.

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Introduction

Autosomal dominant familial neurohypophyseal diabetes insipidus (adFNDI) is a form of diabetes insipidus caused by absence of circulating arginine vasopressin (AVP), a nonapeptide derived from a larger single chain vasopressin-neurophysin (AVP-NP II) precursor synthesised by magnocellular neurones of the hypothalamus. Various heterozygous mutations in the coding sequence of the AVP-NP II gene have been identified in patients with adFNDI (1). The AVP-NP II gene is located on chromosome 20p13 and consists of three exons.

Clinically, the deficiency in secretion of the antidiuretic hormone arginine vasopressin (AVP) becomes apparent several months to years after birth and progresses in severity from partial to nearly complete. The postnatal progression has been explained by degeneration of the AVP producing magnocellular neurohypophyseal neurones, as a result of intracellular retention and a cytotoxic effect of the mutated precursor. Several lines of evidence support this concept. Autopsy studies of patients with adFNDI revealed degeneration of magnocellular vasopressinergic neurones in the suprachiasmatic and paraventricular hypothalamic nuclei (2–5). The progressive deficiency in AVP is typically accompanied by the disappearance of the posterior pituitary bright spot signal on T1-weighted magnetic resonance imaging (MRI) (1, 6–9).

We report the results of clinical, morphological and molecular studies of a Swiss family with adFNDI in three generations. Further, co-segregation of clinical
symptoms and the C59Δ/A60W mutation in the AVP-NP II precursor with the absence of the characteristic T1-hyperintense appearance of the neurohypophysis is demonstrated.

Participants and methods

Participants

We studied four members of a pedigree of three generations (Fig. 1). The index patient (III-5), a Swiss female child presented at the age of 15 months with polydipsia (fluid intake of 3–4 litres per day) and polyuria, but on physical examination she was otherwise healthy. She was investigated first by water deprivation test (10).

The family history revealed central diabetes insipidus in three other family members, namely her mother (II-3), one of her uncles (II-2) and her maternal grandfather (I-1) who might be the source of the initial germ-line mutation. Two of these family members (I-1, II-2) had been evaluated some years previously by water deprivation tests (10); member I-1 had developed symptoms at school age and member II-2 at the age of 2–3 years. In member II-3, the diagnosis was established solely on clinical grounds at the age of 18 months because formal testing was refused. All affected family members have now been receiving treatment with 1-desamino-8-arginine vasopressin (DDA VP) for several years, except the index patient (III-5), in whom DDA VP substitution therapy was started only recently, after formal testing. Individuals I-1 and II-2, then 58 and 32 years of age, gave informed consent to undergo T1-weighted MRI of their posterior pituitary glands. However, in the index patient, MRI was unfortunately refused by the parents. Molecular genetic analysis were performed in the index patient (III-5), her parents (II-3 and II-4) and family members I-1 and II-2.

Diagnostic tests

A water deprivation test was performed in the index patient according to standard procedures (10) using DDAVP to terminate the 7-h fluid restriction period used for differentiating central from nephrogenic diabetes insipidus (11).

MRI studies

Two affected family members (I-1 and II-2) were studied on a 1.5 T High Resolution MRI-Scanner (Siemens Vision, Erlangen, Germany) using sagittal, coronal and T1-weighted spin–echo sequences (response time 350 ms, echo time 12 ms, slice thickness 3.0 mm, field of view 180×180 mm, Matrix 230×256). Series were acquired without intravenous application of contrast media and without fat suppression.

Molecular genetic analysis

After we had obtained informed consent from the participants, genomic DNA was extracted from peripheral leukocytes (12). All three exons of the AVP-NP II gene were amplified using intronic primers (exon 1 sense: TCGCTCCACGGGAAACACCTGGGAGGATA; exon 1 antisense: AGCAGTCGCTAGTCGGCTGAGATG; exon 2/3 sense: TCGCTGCCCTCCCTCAACCCCTCGACTC; exon 2/3 antisense: CCTCTGCTCCTCTTCCCTCAGAG) with conditions reported previously (13). The PCR products were purified with Centricon 100 columns (Amicon, Beverly, MA, USA), and both strands were sequenced directly using a dRhodamine terminator cycle sequencing kit (Applied Biosystems, Foster City, CA, USA). Sequencing products were analysed on a 377 DNA sequencer (Applied Biosystems). All the data were confirmed by a second sequence analysis from both sides.

Results

Clinical diagnosis

During water deprivation testing the 15-month-old index patient developed moderate dehydration (299 mosmol/kg) without urinary concentration (87 mosmol/kg). AVP was unmeasurably low. Administration of DDAVP stopped urinary flow completely, consistent with the diagnosis of central AVP-deficient diabetes insipidus.

MRI studies

The characteristic high-intensity signal of the posterior part of the pituitary gland in unenhanced T1-weighted spin–echo sequences without fat suppression was completely absent in both patients studied (Fig. 1: I-1, II-2). Two representative sagittal images are shown in Fig. 2 in comparison with the image of a control individual showing the bright spot of a normal neurohypophysis.

Molecular genetic analysis

Analysis of the pedigree is consistent with an autosomal dominant transmission of diabetes insipidus
Sequence analysis of the AVP-NP II coding sequence in all the affected individuals revealed a deletion 177–179 DCGC in exon 2 (Fig. 3; nomenclature according to 14, 15). At the amino acid level, this deletion eliminates cysteine 59 and it substitutes alanine 60 by tryptophan (A60W). Although this deletion affects the second and third base of codon 59, in addition to the first base of codon 60, the remainder of the reading frame remains unaffected (Fig. 4). According to the three-dimensional structure of neurophysin, C59 is involved in a disulphide bond with C65 (1). According to the traditionally used nomenclature with separate numbering for the signal peptide, AVP, neurophysin and copeptin, this alteration corresponds to C28 and A29W (Fig. 4) (1, 15).

**Discussion**

Autosomal dominant familial neurohypophyseal diabetes insipidus is caused by mutations in the AVP-NP II gene (1, 13). The AVP-NP II gene is located on chromosome 20p13 and has three exons, with an open reading frame of 492 bp. Exon 1 encodes a signal peptide, AVP, and the N-terminal portion of NP II; exon 2 encodes the central region of NP II; exon 3 encodes the C-terminal part of NP II and copeptin, a glycoprotein with unknown function (16). About 30 distinct mutations have been reported in the AVP-NP II gene causing adFNDI (13, 15, 17–34). The mutations include small deletions and missense and nonsense mutations that affect the signal peptide, the AVP moiety, or the AVP carrier protein, NP II (1). All these dominant mutations described so far contrast with the recently reported recessive mutation in AVP (35). In this autosomal recessive case, the three affected children, offspring of first-degree relatives, were homozygous for a transition C77T in exon 1, replacing P26L (P7L within AVP) (Fig. 4). The mutated form of AVP was a weak agonist with approximately 30-fold reduced binding to the human AVPR2 receptor. In contrast to patients with adFNDI, who present with a progressive decrease in AVP concentrations, the mutated AVP was increased in all three children and there was a further, 30-fold increase during water deprivation (35).

The mutation in the pedigree presented here eliminates a cysteine (C59Δ) believed to be involved
in the formation of a disulfide bond with cysteine 65 as predicted by sequential proteolysis and the crystal structure of bovine NP II (18, 36, 37). Interestingly, at the same locus a transition T175C resulting in substitution of C59R has been previously reported in a family with adFNDI (1). However, the mutation in our study family is quite different and unusual because the deletion C59D is followed by a frameshift that affects only the following codon (A60W), but not the remainder of the reading frame, which is entirely preserved (Fig. 3). The structural consequence of this mutation (C59D, A60W) in the neurophysin moiety is expected to alter correct folding of the AVP-NP II precursor by eliminating a disulphide bridge (C59–C65) (1, 18, 36, 37). It is noteworthy that the only known frameshift mutation in the AVP-NP II gene identified to date occurred in the Brattleboro rat and is associated with a recessive mechanism of disease (38).

Incorrect folding by elimination or introduction of cysteine residues is a commonly observed mechanism underlying abnormal folding of AVP-NP II in adFNDI (1, 15). Recently, a report of the expression of mutated AVP-NP II in different transfected cells documented retention of the mutated precursor molecule in the endoplasmic reticulum (17, 39–41). Furthermore, there is evidence for the toxicity of mutant precursors or their degradation products in cultured neuro2A neuroblastoma cells (40). Other mutations in the NP II moiety are believed to affect flexibility or rigidity of the molecule, or they may alter the structure of the binding pocket and thus impair binding of AVP to its carrier protein (1). Mutations in the signal peptide impair or misdirect cleavage of the signal peptide from the N-terminus of AVP (1, 41, 42). In addition to the cytotoxicity caused by the misfolded mutant AVP-NPII precursors, heterodimer formation between the wild-type and mutant precursors may contribute to the pathogenesis of adFNDI through a dominant negative mechanism (43).

Overall, the fact that all the mutations affecting the signal peptide or the neurophysin moiety result in amino acid substitutions or deletions of residues that are important for processing, folding or oligomerization of the precursor suggests a possible unifying molecular mechanism for the development of the disease (1, 18).

In patients with adFNDI who were started on treatment early and have been receiving DDAVP substitution therapy for many years, secretion of endogenously misfolded AVP-NP II might be blocked by the exogenous DDAVP application and, therefore, the suggested cytotoxic effect of this altered peptide could be reversible for a certain period of time, as has been seen in some autosomal dominant forms of GH deficiency (44, 45). The observation that the high-intensity signal of the neurohypophysis was undetectable in both patients I-1 and II-2, who have been receiving treatment with DDAVP for more than 30 years, argues against the possibility of a reversible process in adFNDI.

The observed absence of the so-called bright spot of the posterior pituitary gland in patients with adFNDI is believed to be caused by the absence of AVP-containing neurovesicles in axonal endings, thus reflecting loss of posterior lobe function (6–9, 26).

The findings of our study demonstrate the cosegregation of clinical symptoms and the C59D/A60W mutation in the AVP-NP II precursor, with absence of the characteristic T1-hyperintense appearance of the neurohypophysis in both of the two adult adFNDI patients studied. These findings are in line with the suggested pathomechanism for adFNDI (17, 39–41, 43). The mechanism underlying adFNDI is thus distinct from some forms of autosomal dominant GH deficiency that has preserved inducible GH secretion and absence of cytotoxic degeneration of somatotrophs (44, 45).

In conclusion, the deletion C59D/A60W is predicted to result in an AVP-NP II precursor that is misfolded and thus retained within the magnocellular neurones. In addition, MRI results from the affected patients support the molecular mechanism postulated for the majority of AVP-NP II mutations associated with adFNDI (17, 39–41, 43).
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