Autosomal Dominant Familial Neurohypophyseal Diabetes Insipidus Caused by a Novel Mutation in Arginine-Vasopressin Gene in a Brazilian Family

ABSTRACT

Autosomal dominant familial neurohypophyseal diabetes insipidus (adFNDI) is a rare autosomal dominant disorder characterized by polyuria and polydipsia due to deficiency of arginine vasopressin (AVP). More than 50 mutations causing adFNDI have been already reported in the AVP gene. The aim of the present study is to analyze the AVP gene in four generations of one Brazilian kindred with adFNDI. The proband was a 31-year old female with huge hypotonic polyuria (10 L/day) dated from childhood. Molecular analysis included amplification of all exons and exon-intron regions of the AVP gene by PCR and directly sequencing. Sequencing analysis showed a novel point mutation in heterozygous state: G88V (GGC>GTC). All affected patients presented the same mutation also in heterozygous state, while it was absent in four normal members. We expand the repertoire of mutations in AVP describing the novel G88V mutation in one Brazilian kindred with adFNDI. (Arq Bras Endocrinol Metab 2008; 52/8:1271-1275)

Keywords: Antidiuretic hormone; AVP; Familial diabetes insipidus; Arginine-vasopressin gene; Diabetes insipidus, neurogenic; Posterior pituitary gland

INTRODUCTION

 Familial neurohypophyseal diabetes insipidus is a rare disorder characterized by polyuria and polydipsia due to arginine vasopressin (AVP) deficiency (1). This familial disorder generally occurs some months to years after

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RESUMO

Diabetes Insípido Neuro-Hipofisário Familiar Causado por uma Nova Mutação no Gene da Arginina-Vasopressina em uma Família Brasileira.

Diabetes insípido neuro-hipofisário com herança autossômica dominante (adFNDI) é uma doença autossômica dominante rara, caracterizada por poliúria e polidipsia devido à deficiência de arginina-vasopressina (AVP). Mais de 50 mutações causando adFNDI foram descritas no gene AVP. O objetivo deste estudo foi analisar o gene AVP em quatro gerações de uma família brasileira com DINF. O caso-índice é de uma paciente de 31 anos, com volumosa poliúria hipotônica desde a infância (10 L/dia). A análise molecular incluiu amplificação por PCR e sequenciamento automático dos éxons e regiões éxon-intron do gene AVP. A análise do sequenciamento mostrou uma nova mutação de ponto em heterozigose: G88V (GGC>GTC). Todos os pacientes afetados apresentaram a mesma mutação, que não foi encontrada em quatro indivíduos normais da família. Expandimos a lista de mutações no gene AVP, descobrindo a nova mutação G88V em uma família brasileira com adFNDI. (Arq Bras Endocrinol Metab 2008; 52/8:1271-1275)

Descritores: Hormônio antidiurético; Diabetes insípido familiar; Gene AVP; Diabetes insípido neurogênico; AVP; Hipófise posterior
birth, and ongoing progressive AVP deficiency has been described (2,3).

The autosomal recessive inheritance was reported in one family but most patients present an autosomal dominant inheritance and more than 50 mutations, causing autosomal dominant familial neurohypophysial diabetes insipidus, (adFNDI) have been already reported in AVP gene (4,5). The AVP gene is located at chromosome 20 (20p13) and encloses three exons: the first exon encodes a putative signal peptide, the AVP and the aminoterminal portion of neurophysin II (NPII); the second exon encodes the central portion of NPII; and the third exon encodes the carboxiterminal region of NPII and a glycoprotein, copeptin (6).

The NPII is the carrier protein of AVP and has been implicated in sorting the prohormone. The prohormone is synthesized in the magnocellular neurons of the supraoptic and paraventricular nuclei at hypothalamus and is subsequently transported into the nerve terminals in the neurohypophysis, via the regulated secretory pathway. In the secretory granules, processing of the vasopressin prohormone takes place, and upon stimulation of the nerve terminals processed products are released (7-9). Most AVP mutations are within NPII region and consist in substitutions of amino acids implicated in protein structure conformation, like cysteines involved in disulfide-bridge formation or prolines and glycines, which can make turns in polypeptide chains (10-12). Therefore, adFNDI represents the best known inherited endocrine disease caused by prohormone defects (3).

It has been proposed that the dominant negative effect of AVP mutations and the variability in onset of this disease, delayed to several months or years of age, are consequences of abnormal retention of abnormal prohormone in the endoplasmic reticulum, where it fails in folding and/or dimerizing appropriately. This prolonged cytotoxic accumulation eventually leads to degeneration of neurons expressing mutant vasopressin prohormones (13).

In the current study, we present the molecular analyze of the AVP gene in a Brazilian pedigree. Eleven subjects were identified with the adFNDI, which spread for four generations, showing a complete penetrance.

**METHODS**

We studied a Brazilian pedigree with adFNDI followed at our Institution. This study was approved by the Hospital Ethics Committee and written informed patient consent terms were obtained.

**Clinical evaluation**

The proband was a 31 year-old woman who presented clinical features of excessive polyuria and polydipsia. A water deprivation test (WDT) and a magnetic resonance image (MRI) of the hypothalamic-hypophysal region were performed. Nine family members in four generations were supposed to be affected, judging from their symptoms (polyuria and polydipsia) (Figure 1).

**Molecular Analysis**

Genomic DNA was isolated from peripheral blood by standard methods from proband, thirteen relatives alive and a normal control.

**AVP gene PCR**

Three exons and exon-intron boundary regions of AVP were individually amplified by PCR using intronic primers and protocol amplification as previously described (3). Masafumi buffer was essential to improve PCR reaction.

**Sequence analysis**

PCR products were treated with shrimp alkaline phosphatase and exonuclease I (PCR product pre-sequencing kit; Amersham Life Sciences Inc., Cleveland, Ohio, USA) prior to sequencing using the ABI PRISM Big-Dye Terminator kit (Perkin-Elmer Applied Biosystems, Foster City, California, USA). The products were then directly sequenced using an ABI PRISM 3100 Genetic Analyzer automatic DNA sequencer (Perkin-Elmer Applied Biosystems).

**Figure 1.** Pedigree of adFNDI. Black symbols indicate affected subjects; females are indicated by circles and males by squares. Subject indicated by the arrow was the proband. All affected individuals of II, III and IV generations and the normal individuals (II-2, III-5, III-7 and IV-1) were studied. The mutant allele was detected in all affected subjects.
Hpy188 III restriction assays

The mutation G88V generates a restriction site for Hpy188 III enzyme. All relatives were screened for the G88V mutation, using 5 µl of PCR product with 0.5 µl of Hpy188 III enzyme (10 U/µl) at temperature 37ºC, with inactivating at 65ºC. Mutant allele generates two fragments of 154 and 149-bp from PCR product.

RESULTS

Clinical features

The proband presented a characteristic response on WDT (plasma sodium levels enhanced from 146 to 149 mEq/L; specific density changed from 1005 to 1007). MR image displayed an hypoplastic anterior pituitary and posterior pituitary was not visualized (Figure 2). The anterior pituitary function was evaluated by combined test and all pituitary hormonal responses were normal.

Proband relatives alive were three female and five male, with age ranged from 5 to 53 (mean= 32.5). The mean chronological age onset of symptoms in five patients was 4.36 yr (ranging from 4 months to 11 years). In the other patients the precise age onset of symptoms could not be determined, mainly because they did not recognize the huge amount of water intake as anomalous, since many relatives used to drink the same. In most patients the diagnosis was achieved after the family screening evaluation.

Molecular results

Sequencing analysis showed a novel point mutation in heterozygous state in exon 2. The nucleotide 1859 takes part in codon 88 (GGC>GTC) and this substitution results in an amino acid substitution of glycine to valine (G88V) (Figure 2). This codon encodes the glycine in position 57 of neurophysin – NP57.

The screening of this novel mutation was performed in all alive relatives using the restriction enzyme Hpy188 III confirming the same mutation in all other nine affected members, while it was absent in 4 normal members and a normal control.

DISCUSSION

In this report, we studied a large family with clinical, laboratorial and molecular diagnosis of adFNDI. The proband patient presented a huge polyuria which motivated him to search medical help. The fluid deprivation test was characteristic of AVP deficiency as well as the absence of the hyper signal in the posterior pituitary. Family adjustment to polyuria and, consequently, polydipsia is an interesting finding in these cases and can be associated to progressive establishment of disease, which is corroborated by previous reports on gradual decline in vasopressin secretion (2). Most of the relatives have not realized that they had polydipsia because this was the familiar pattern of water ingestion, and they were studied only after the proband was investigated in our Service.

Sometimes it is possible that patients can present a suggestive history of DI associated to an inconclusive WDT.
For them, the measurement of AVP and molecular analysis of AVP is crucial for the diagnosis of DI (15).

Abnormalities in the pituitary are described in the MRI of patients with adFNDI, such as absence of posterior pituitary bright spot (15). Our patient presented a hypoplastic anterior pituitary on MRI and this finding did not correlate with clinical and laboratorial evaluation, revealed as normal. The cause of this structural abnormality is still unclear, and emphasizes the importance to monitor the anterior pituitary function in this patient.

Some technical difficulties in PCR amplification and direct sequencing of AVP were encountered because this gene is extremely rich in G+C, especially in the last half of the gene where these nucleotides encompass more than 70% of the sequence, as already reported by other researchers (3).

In the present study, we have identified a novel point mutation in AVP gene (G88V). This is the third mutation described in codon 88 that encodes one glycine (16). Two different mutations in the same codon (G88S and G88R) have been already described in patients with a similar phenotype. Glycine in codon 88 displays full interspecies conservation (6). In addition, glycine and valine are both amino acids non polar and neutral, otherwise, valine is more than ten times more hydrophobic than glycine (17).

Nijenhuis and cols. performed functional analyses of five mutations in the AVP gene, three of them in glycine residues (G45R, G96V and G88S, respectively NP14G, NP65G and NP57G positions on neurophysin). The analyzes of mutation G88S displayed a relatively efficient processing and secretion of protein, an aberrant distribution of the mutant prohormone within the endoplasmic reticulum (ER) and a modification in ER distribution in cells (13). Therefore, they propose that the accretion and the consequent disturbance of the ER are deleterious to the cell and will decrease the functionality and/or the viability of the magnocellular neurons, which express high amounts of vasopressin prohormone in vivo (13). This hypothesis would explain both the dominant inheritance of human adFNDI and the delayed onset of the symptoms, as observed in some patients here described.

Another study showed that mutation of residues NP14G or NP65G, which are both located in one of the β-pleated sheet of the NPII, virtually abolished efflux of phrohormone from the ER (18). Mutation of residue NP57, which is located in the loop connecting the α-helix with the C-terminal β-pleated sheet of NPII, provokes only a decrease in the efflux of prohormone (19). The grade of severity of ER retention for different vasopressin prohormones suggests that mutations in the β-strands of neurophysin are most deleterious for folding of the vasopressin prohormone, followed by mutations in the α-helix and then by mutations in less ordered structures like the loop from residues 50–58 (19). These findings were reinforced by analyzes in vitro performed by Eubansks and cols. (20) demonstrating that the mutation G88S has weaker free energy of internal bonding between hormone and NPII than in the WT precursor.

One important practical issue to detect AVP mutation and confirm the diagnosis of adFNDI is to be able to predict the development of adFNDI in asymptomatic members, since the penetrance of this disease appears to be 100% (15).

In conclusion, we expanded the repertoire of mutations in AVP describing the novel G88V mutation in one Brazilian kindred with adFNDI.

No potential conflict of interest relevant to this article was reported.

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