CASE REPORT

A germline \textit{RET} proto-oncogene mutation in multiple members of an Arab family with variable onset of MEN type 2A-associated clinical manifestations

Makia Marafie *, Ibrahim Suliman, Mohammed Dashti, Abdulla Redha, Abdulrahman Alshati

Kuwait Medical Genetics Centre, Maternity Hospital, Sabah Medical Area, P.O. Box 5833, Safat 13059, Kuwait

Received 7 August 2016; accepted 24 August 2016
Available online 17 September 2016

KEYWORDS
Arab; Medullary thyroid carcinoma; MEN2A; Pheochromocytoma; \textit{RET} proto-oncogene

Abstract Background: Multiple endocrine neoplasia type 2A (MEN2A) is a rare cancer associated-syndrome, inherited in an autosomal dominant fashion and caused by germline mutation in \textit{RET} proto-oncogene. Clinical diagnosis depends on the manifestation of two or more certain endocrine tumors in an individual, such as medullary thyroid carcinoma, pheochromocytoma, and parathyroid adenoma or hyperplasia. Prophylactic total thyroidectomy with central neck lymph node dissection is mandatory for mutation carriers, with periodic monitoring of the other concerned organs.

Subjects and methods: We have screened 27 individuals from a large Arab family with multiple affected members. Mutational screening involved the hotspot regions in the most commonly implicated exons 10 and 11 of \textit{RET} proto-oncogene using PCR amplification of the coding and the flanking intronic regions followed by the Sanger sequencing. We aimed for confirmation of the clinical diagnosis and identification of at-risk asymptomatic mutation carriers.

Results: A pathogenic variant c.1901G > T (p.Cys634Phe), in exon 11 of \textit{RET} proto-oncogene was identified in 15 members of different ages.

Conclusion: Genetic counseling plays a key role in the management of such high-risk families and hence helps in avoiding or reducing disease recurrence in their future generations.

1. Introduction

Multiple endocrine neoplasia type 2 (MEN2) is a rare cancer associated-syndrome, inherited in an autosomal dominant fashion and caused by germline mutation in \textit{RET} proto-oncogene. The main manifestation is medullary thyroid
carcinoma (MTC), which is a cancer of the parafollicular C/or calcitonin secreting cells. It is classified into three subtypes: MEN2A, familial medullary thyroid cancer (FMC) and MEN2B. In FMC, MTC is the only presented phenotype and it occurs at a later age of onset with a relatively better prognosis. MEN2B syndrome is characterized by MTC that occurs in childhood, with an increased risk for pheochromocytoma, mucosal neuroma of lips and tongue, ganglioneuroma of the gastrointestinal tract and Marfanoid habitus [1,2]. In MEN2A associated-families, individuals with pathogenic mutations are at increased risk (95%) for development of early adult onset MTC (multifocal or bilateral), which is often associated with C-cell hyperplasia; also risk is increased for pheochromocytoma (50%) and parathyroid adenoma or hyperplasia (20–30%) [2,3]. The prevalence of MEN 2A is estimated to be 1 per 50,000, with usual diagnosis age of 20–30 years. De novo mutations may be responsible for around 5% of MEN2A patients [4].

RET proto-oncogene is located on chromosome 10 (10. q11.2); it comprises 21 exons and encodes a transmembrane tyrosine kinase receptor protein which plays an important role in transferring cell growth and differentiation signals. It is expressed in parafollicular C cells of the thyroid gland, parathyroid glands, adrenal medulla and in the urogenital system [5].

The clinical phenotype depends on type and position of gene mutation [1,6,7]. All reported mutations; their associated phenotypes and the pertinent literature references are shown in the public mutation repository MEN2-RET databases: (http://www.arup.utah.edu/database/MEN2/MEN2_welcome.php); 2016 [accessed 1.8.16]. To date; 166 variants are stored in this database. The most frequently reported mutations are at codons 634 in exon 11, and 620, 618, 611, 609, which are located in exon 10. However, mutations at codon 634 have been reported in 85% of tested individuals, nearly 50% of them are amino acid cysteine to arginine substitution (Cys634Arg). Pathogenic variants at this codon reported to result in a higher incidence of pheochromocytoma, hyperparathyroidism and lichen amyloidosis [3,4,8]. Additionally, the risk of development of the Hirschsprung disease in carriers could reach 7% [9].

Many guidelines have been established for improvement of the diagnosis and for better management of patients with MTC or neuroendocrine tumors [10–13]. Based on the aggressiveness of thyroid tumors and the age of clinical detection, the revised guidelines by American Thyroid Association (ATA) Task Force on Medullary Thyroid Carcinoma has classified RET variants at codon C634 (C634F/G/R/S/W/Y) as high risk mutations (category H) [12]. In general, the recommendations included periodic assessment of the clinical status by thyroid ultrasound and biochemical screening for MTC, CT and MRI for pheochromocytoma, with serum calcium and parathyroid hormone level assessment [10–13].

MTC is the most common cause of death in MEN2-associated families. Therefore prophylactic total thyroidectomy with central neck lymph node dissection is mandatory for RET mutation carriers, with periodic monitoring for residual or recurrent MTC and annual calcium calcitonin stimulation test. Prophylactic surgery is recommended for young individuals of <5 years of age, who are carriers for certain high-risk pathogenic variants in codon 634 [12,13], while biochemical screening for pheochromocytoma and hyperparathyroidism should start at 8 years of age [12].

We have screened the first Kuwaiti family with multiple affected members, for confirmation of the clinical diagnosis and as a measure for identification of at-risk asymptomatic mutation carriers.

2. Subjects and methods

2.1. Family data

The proband (P9) was 35 year old healthy male, approached the cancer genetics clinic seeking genetics counseling and predictive gene testing. Pedigree analysis and medical reports revealed many relatives manifesting thyroid/parathyroid/adrenal associated-diseases consistent with the diagnosis of MEN2A syndrome (Fig. 1, Table 1). Subsequently his high-risk family members (P1–P27) were seen in groups or in separate individualized sessions according to their request. Genetic counseling was provided, the importance of predictive gene test was discussed, and informed consent was obtained for blood collection and testing. The work has been carried out in accordance with The International Code of Medical Ethics of the World Medical Association (Declaration of Helsinki) and Ethical approval of the Kuwait Medical Genetics Center.

2.2. Molecular screening

Blood samples were obtained from 27 individuals in the family. Genomic DNA was extracted from peripheral blood leukocytes using the automatic Maxwell® 16 System DNA purification Kits (Promega, USA) according to manufacturer’s protocol. We initially performed mutational screening for the hotspot regions in the most commonly implicated exons 10 and 11 of RET proto-oncogene using PCR amplification of the coding and the flanking intronic regions, followed by the Sanger sequencing of the PCR product using ABI PRISM® 3100 Genetic Analyzer (Applied Biosytem, USA). We used the previously designed primers and conditions for amplification of exons 10 and 11 [14]. We did not proceed to test other exons; soon the pathogenic variant was identified.

3. Result and discussion

We have identified a heterozygous variant in exon 11 of RET proto-oncogene; c.1901G > T; p.Cys634Phe (or C634F as commonly used in the literature) (Fig. 2); in the proband and 14 members of his family. Mulligan et al. had previously described this variant in 1993 as a disease causing mutation [15]. It was later published as pathogenic mutation by Gene Review, PMID:20301434 (http://www.ncbi.nlm.nih.gov/books/NBK1257), and was reported in ClinVar database, as NM_020975.4 at (http://www.ncbi.nlm.nih.gov/clinvar); 2016 [accessed 1.8.2016]. This is a missense mutation that occurred at the cysteine residue within the extracellular cysteine-rich domain. It substitutes cysteine for phenylalanine. This is a hot-spot gene position for pathogenic mutations, which are commonly found in various populations and ethnic groups including North Africans, but with different frequencies due to different genetic background [3,4,16–20]. Codon 634
mutation was the most commonly detected missense mutation in many studies (55–93%); Cys634Arg being the most prevalent mutation followed by Cys634Tyr and CYs634Gly [3,7,19,20], whereas our family variant Cys634Phe was less commonly reported [14]. However, it has been detected in 2/15 French patients with pheochromocytoma [21], and out of the 31 screened MTC-associated Moroccan patients, only one carried this mutation; he developed MTC and pheochromocytoma at the age of 34 years [22].

In this family seven out of 15 heterozygous carriers had developed clinical manifestations (Fig. 1, Table 1). Among the affected members; we observed intra-familial variability in phenotypes and disease onset, with an average age at onset of 43.6 years; the youngest case was a 16 year old female (P13), while the oldest case was a 70 year old woman (P3). Moreover, two sisters P3 and P5 had developed the full disease spectrum at a later age; having MTC with no local or distal metastasis, parathyroid adenoma and unilateral pheochromocytoma, both had no clinical complaint until the clinical diagnosis ages, which were 70 and 54 years respectively. In P3 patient, pheochromocytoma was the first symptom of MEN2A, causing sudden severe high blood pressure crises that required immediate hospitalization. Their mother died at an old age long time ago due to thyroid cancer as was claimed by her relatives. Another relative (P10), the daughter of (P3), was diagnosed with MTC and parathyroid adenoma at the age of 47 years with no nodal metastasis. Very recently; her 46 year old sister (P7) was diagnosed with MTC and the only member with cutaneous lichen amyloidosis; her 44 year old brother (P8) had sudden severe hypertensive crises, due to pheochromocytoma of left adrenal gland. P11 who is the husband of P10 and her first cousin, was symptomatic and proved to be

![Family pedigree](image_url)

Figure 1 Family pedigree. The proband (P9) is indicated with an arrow. (P1–P27) represent family members that were tested, open symbols for non-symptomatic individuals; diagonal bars for deceased members. Symptomatic individuals are indicated with solid symbols.

- An affected male.
- An affected female.

<table>
<thead>
<tr>
<th>P</th>
<th>Age 1 (years)</th>
<th>Age 2 (years)</th>
<th>Sex</th>
<th>Manifestation</th>
<th>Intervention</th>
</tr>
</thead>
<tbody>
<tr>
<td>P3</td>
<td>72</td>
<td>70</td>
<td>F</td>
<td>MTC, Parath, Pheo</td>
<td>Thyroidectomy, parathyroidectomy, Rt adrenalectomy</td>
</tr>
<tr>
<td>P5</td>
<td>70</td>
<td>54</td>
<td>F</td>
<td>MTC, Parath, Pheo</td>
<td>Thyroidectomy, parathyroidectomy, Rt adrenalectomy</td>
</tr>
<tr>
<td>P7</td>
<td>46</td>
<td>46</td>
<td>F</td>
<td>MTC, Lichen cutaneous amyloidosis</td>
<td>Planned for thyroidectomy</td>
</tr>
<tr>
<td>P8</td>
<td>44</td>
<td>44</td>
<td>M</td>
<td>Pheo,</td>
<td>Planned for thyroidectomy, adrenalectomy</td>
</tr>
<tr>
<td>P9*</td>
<td>–</td>
<td>–</td>
<td>M</td>
<td>No</td>
<td>UCI</td>
</tr>
<tr>
<td>P10</td>
<td>47</td>
<td>45</td>
<td>F</td>
<td>MTC, Parath</td>
<td>Thyroidectomy, parathyroidectomy</td>
</tr>
<tr>
<td>P11</td>
<td>45</td>
<td>30</td>
<td>F</td>
<td>C-cell hyperplasia</td>
<td>Prophylactic thyroidectomy</td>
</tr>
<tr>
<td>P12</td>
<td>47</td>
<td>–</td>
<td>M</td>
<td>No</td>
<td>UCI</td>
</tr>
<tr>
<td>P13</td>
<td>30</td>
<td>16</td>
<td>F</td>
<td>C-cell hyperplasia</td>
<td>Prophylactic thyroidectomy</td>
</tr>
</tbody>
</table>

MTC: medullary thyroid carcinoma, Parath: parathyroid hyperplasia/or adenoma, Pheo: pheochromocytoma. M: male, F: female, P = patient's number as in Fig. 1, P9* = proband, Age 1 = age at molecular test, Age 2 = age at first clinical manifestation, UCI = under clinical investigation.
a heterozygous carrier. However, three of their children (P22, P23, and P25; aged 16, 12 and 5 years) proved to be asymptomatic heterozygous carriers. One of the carriers (P25) was a 5 year old girl with Down syndrome, who is currently asymptomatic. Additionally, the proband (P9) and four more relatives were asymptomatic; being an adult (P12) and three other children (P20, P26 and P27) who were below 16 years of age. All were referred to the endocrine clinic for further clinical management.

Distant metastasis was significantly associated with C634R mutations than with C634Y or C634W mutations, while individuals harboring C634R, seem to have more aggressive disease, as demonstrated by more frequent distal metastasis at diagnosis and at an earlier ages according to the Kaplan–Meier estimate of cumulative lymph node and distant metastases rates [23]. In contrast, the C634Y genotype appears to have an indolent behavior [7,23]. In the current family, it seems that variant c.1901G > T (p.Cys634Phe)/or (C634F) is of low aggressive effect or of slow-growing phenotype in comparison to other reported point mutations at this codon. This is because most of the affected members were diagnosed after the age of 40 years (5/7), and 2/7 had underwent prophylactic thyroidectomy at the ages of 16 and 30 due to c-cell hyperplasia. Also all the thyroid tumors in the affected members were

Figure 2  Partial DNA sequence for exon 11 of RET proto-oncogene, showing (A) c.1901G > T heterozygous variant found in the proband (arrow), (B) normal control DNA sequence of RET gene showing the wild type at same position (arrow).
not associated with local or distal metastasis. In conclusion; MEN2A is a serious hereditary disease that involves not only adults, but also young children. Therefore these genetically predisposed families should be educated about the importance of testing their at-risk members; including children as young as few months of life in an attempt to detect the disease at an early age and proceed with preventive surgical intervention at the proper time.

Conflict of interest

There is no conflict of interest to the publication of this article.

Funding

No funding body was involved.

References