Novel p.C620L RET Mutation Detected in a Patient with Medullary Thyroid Carcinoma

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ABSTRACT

A patient with an apparent sporadic medullary thyroid carcinoma was tested for RET germline mutations by Sanger sequencing of RET exons 10, 11, and 13 - 16. The patient was heterozygous for two known mutations causative of Multiple Endocrine Neoplasia type 2 disorder, and both mutations were within codon 620 of RET exon 10, c.1859G > T (p.C620F) and c.1860C > G (p.C620W). In order to determine if these adjacent mutations were in cis or in trans, an unlabeled probe method and high-resolution melting analysis were utilized. The mutations were confirmed to occur in cis, representing a novel mutation, c.1859_1860delinsTG (p.C620L). Sanger sequencing of parental samples did not identify any changes at codon 620, so the p.C620L mutation is also de novo. The early age of onset for medullary thyroid carcinoma and the presence of lymph node metastasis in this patient suggests individuals with the p.C620L mutation should be treated and screened (for pheochromocytomas and parathyroid hyperplasia) as Multiple Endocrine Neoplasia type 2 patients with other RET codon 620 mutations (American Thyroid Association risk level B).

Keywords: RET; MEN2; MTC; Medullary Thyroid Carcinoma; High Resolution Melting Analysis; Inherited Thyroid Cancer

1. Introduction

Multiple endocrine neoplasia type 2 (MEN2, OMIM 171400) is an autosomal dominant inherited disorder with the hallmark symptom of medullary thyroid carcinoma (MTC). MEN2 consists of three syndromes: familial medullary thyroid carcinoma (FMTC; have MTC only), MEN2A (MTC, Pheochromocytoma, and parathyroid hyperplasia) and MEN2B (MTC, Pheochromocytoma, and other characteristic findings) [1-3]. MEN2 is caused by pathogenic RET proto-oncogene germline mutations.

The transmembrane RET protein is involved in cell growth and differentiation. In the presence of ligand, the RET monomers dimerize, autophosphorylate, and activate downstream signaling. In RET exon 10 there are four conserved cysteine residues (at codons 609, 611, 618, and 620) that are hotspots for MEN2 causative, gain-of-function mutations [3,4]. These cysteine residues are involved in intramolecular disulfide bonding. It is hypothesized that cysteine residue substitution disrupts the intramolecular disulfide bond, allowing the unpaired partner cysteine to aberrantly create a disulfide bond between the mutant RET monomers, resulting in ligand-independent RET dimerization and activation [5-7]. Mutations at these four codons have been detected in patients with MEN2A or FMTC, and the age of disease onset is variable ranging from childhood to adulthood (see the MEN2 RET database: www.arup.utah.edu/database/MEN2/MEN2_welcome.php [4]).

This report describes the detection of two known pathogenic RET heterozygous changes at exon 10, both within codon 620, by Sanger sequencing in a patient with MTC. High-resolution melting analysis (HRM) and unlabeled probes were used to determine the adjacent mutations occur on the same allele and represent a novel RET mutation, c.1859_1860delinsTG (p.C620L).

2. Materials and Methods

2.1. Samples

This study was conducted under the University of Utah IRB board 00028725, and patient consent was obtained. Patient and parental peripheral blood specimens were
collected and DNA was extracted from blood leukocytes using the MagNa Pure Compact (Roche) procedure following the manufacture recommended protocol.

2.2. Sequencing of the RET Protooncogene

Samples were amplified by PCR for the coding regions and intron/exon boundaries of RET exons 10 - 11, and 13 - 16, then bidirectional Sanger sequencing was performed. Mutation Surveyor® software (SoftGenetics, State College, PA) was used for data analysis.

2.3. Unlabeled Probe and High Resolution Melting Analysis

Sample DNA was amplified by asymmetric PCR, high-resolution melting (HRM) analysis was performed on the HR-1TM (Idaho Technology, Salt Lake City, UT), and data were analyzed with custom software written in LabView (National Instruments, Austin TX) as described previously [8]. Briefly, after PCR, the unlabeled probe hybridizes to the amplicon template which is measured by LCgreen, a DNA intercalating dye. During HRM analysis, the amplicons with mismatches to the probe have a lower melting temperature (Tm) of dissociation than amplicons with complementary sequence to the probe [9].

The PCR primers used for RET exon 10, codon 620 interrogation were (forward 5’GGGCAGCATTTGAGGGAGC3’ and reverse 5’TGGTGGTTGGCGAGCCCGCA3’). The unlabeled probes were 5’GGAGAATGCTCTGCGAGCCCGAAGACATC3’ [wild type sequence at codon 620 (TGC)] and 5’GGAGAATGCTCTTGGAGCCGGCGCGCA3’ [c.1859_1860delinsTG mutation sequence at codon 620 (TTG)]. Unlabeled probes have a blocking 3’ amino modifier (IDT, Coralville, IA) to prevent extension during the PCR step.

3. Results

3.1. Clinical Report

In 2005, the patient presented at 19 years of age with a palpable thyroid nodule. The nodule was found to be 1.3 cm in the left thyroid lobe by ultrasound. The patient did not have dysphagia, voice changes, breathing issues, palpitations, previous radiation exposures, pain, or family history of MTC/MEN2. Fine needle aspiration showed neoplastic proliferation most consistent with medullary thyroid carcinoma (MTC). In concordance with the MTC diagnosis, calcitonin and carcinoembryonic antigen (CEA) levels were elevated: 1181 pg/mL and 53.3 ng/mL, respectively. The patient was screened for additional findings associated with MEN2, but had normal levels of parathyroid hormone, serum calcium, catecholamines, metanephrine and creatinine. Since the patient has no family history of endocrine problems or malignancy and the biochemical examination did not detect other findings of MEN2, sporadic MTC was suspected at the time of the surgery.

Total thyroidectomy with central compartment node dissection was performed. A tumor of 1.2 × 1.1 × 1.0 cm was found in the left thyroid lobe. Multicentric or bilateral involvement was not identified. The tumor did not extend beyond the resection margins and capsular invasion was not identified. One of three lymph nodes tested positive for metastasis. The surgery confirmed the MTC diagnosis with small focus of perithyroidal connective tissue involvement and discovered metastasis to one of three perithyroidal lymph nodes. No unusual findings were found in the parathyroid tissue samples submitted for pathological examination. The left inferior parathyroid was autotransplanted to the left sternocleidomastoid. Calcitonin levels post-surgery remain borderline high at 13 - 15 pg/mL and were rising, suggesting the potential for recurrence.

3.2. RET Proto-Oncogene Mutation Identification

The patient was tested at ARUP laboratories for germline RET mutations in exons 10, 11, and 13 - 16 by clinical Sanger sequencing (Figure 1(a)). Two heterozygous mutations were found, adjacent to each other within codon 620, a known hotspot for MEN2 causative RET mutations. Sanger sequencing of the parental samples, targeting RET exon 10 yielded only wild type sequence at codon 620. So, were these two RET mutations in trans c.[1859G > T;1860C > G] (p.[C620F;C620W]) or in cis c.1859_1860delinsTG (p.C620L)?

Since these two mutations affect adjacent base pairs, phase was determined using HRM and the unlabeled probe method [9] as described previously for RET mutation identification [8]. A probe complementary to the wild type RET sequence for codon 620 (TGC) was used (Figure 1(b)). Two control DNA samples each heterozygous for one codon 620 mutation [c.1859G > T (p.C620F) or c.1860C > G (p.C620W)] demonstrated a decreased Tm (melting temperature) for the mutation alleles compared to their wild type alleles. The patient’s sample demonstrated an allele with an even lower Tm than either of the control alleles harboring single nucleotide mutations. The patient sample also had an allele with the same Tm as the wild type control sample (which had only wild type alleles). In addition, a probe that complemented the c.1859_1860delinsTG mutation sequence at codon 620 (TTG) was used (Figure 1(c)). The patient’s DNA sequence complemented the probe resulting in a variant
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4. Discussion

Since an apparently sporadic MTC (i.e., isolated, non-familial MTC) can be the presenting clinical feature for some MEN2 patients, RET genotyping is often performed for patients with sporadic MTC [10]. The most common areas of MEN2 causative germline RET mutations are exons 10, 11, 13 - 16 [3,11-13]. Molecular sequencing of the RET gene was performed to determine if this patient with apparently isolated MTC had a heritable cause of disease and may require further screening for other MEN2 manifestations. Sequencing detected two adjacent heterozygous RET mutations, each known to be causative of MEN2. It was important to define the chromosomal configuration of these two mutations to aid in risk assessment and genetic counseling for family members. Unlabeled probe high resolution melting analysis methods determined these two changes were on the same allele, and therefore identified the patient’s mutation as c.1859_1860delinsTG (p.C620L). This mutation appears to be novel, since it was not found in NCBI dbSNP135, 1000 Genomes, or other publicly available databases. Also, the p.C620L change was not found in a self-reported MEN2/MTC unaffected cohort (136 people) tested for variations in RET introns and exons 9 through 16 [14]. The patient’s parents had wild type sequence at codon 620, so the mutation is de novo as well.

Any mutation changing the RET 620 cysteine residue to any other amino acid has been reported to result in MTC or MEN2A [4,15], supporting the presumed pathogenicity of the novel p.C620L mutation. As seen with the other RET codon 620 mutations [5-7], it is likely that the p.C620L mutation disrupts intra-cellular disulfide bonding, allowing intermolecular disulfide binding with other mutant RET monomers, leading to constitutive activity and signaling. The youngest age of MTC onset associated with RET codon 620 mutations is 5 years and prophylactic thyroidectomy is general recommended before or at 5 years of age [1,16]. The young age of onset in our patient as well as the detected lymph node metastasis, indicates this mutation should follow the American Thyroid Association’s recommendations for risk level B, which is the classification assigned to other codon 620 mutations [16].

In conclusion, the two adjacent Sanger sequencing detected mutations were determined to be in cis and therefore the patient had the novel and de novo RET mutation c.1859_1860delinsTG (p.C620L). The p.C620L mutation is very likely MEN2 causative due to loss of cysteine at codon 620 and the young age of MTC metastasis in the patient. Since this patient was the only affected family member and had MTC but no other features of MEN2, the specific MEN2 subtype association with this mutation cannot be predicted. Because other RET codon 620 mutations are associated with Pheochromocytomas (young-
est reported age of onset in the literature was of 19 years of age) and the much rarer parathyroid hyperplasia; the patient should be followed for potential development of these additional MEN2 manifestations. The novel p.C620L mutation will be added to our online, publicly available MEN2 RET mutation database [4].

5. Acknowledgements

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REFERENCES


Abbreviations:

HRM: High-Resolution Melting;
MEN2: Multiple Endocrine Neoplasia Type 2;
MTC: Medullary Thyroid Carcinoma;
Tm: Melting Temperature.