BMC Genetics



Methodology article

The minisequencing method: a simple strategy for genetic screening of MEN 2 families

Maria João Bugalho*1,2, Rita Domingues¹ and Luís Sobrinho²

Address: ¹Centro de Investigação de Patobiologia Molecular, Instituto Português de Oncologia FG / Lisboa, Portugal and ²Serviço de Endocrinologia, Instituto Português de Oncologia FG / Lisboa, Portugal

 $E-mail: Maria Jo\~{a}o Bugalho*-mjbugalho@ipolisboa.min-saude.pt; Rita Domingues-rdomingues@ipolisboa.min-saude.pt; Lu\'{s} Sobrinho-lsobrinho@ipolisboa.min-saude.pt$

*Corresponding author

Published: 31 May 2002 BMC Genetics 2002, 3:8

Received: 19 November 2001 Accepted: 31 May 2002

This article is available from: http://www.biomedcentral.com/1471-2156/3/8

© 2002 Bugalho et al; licensee BioMed Central Ltd. Verbatim copying and redistribution of this article are permitted in any medium for any purpose, provided this notice is preserved along with the article's original URL.

Abstract

Background: Multiple endocrine neoplasia type 2 is an autosomal dominant disorder. MEN 2A is characterized by medullary thyroid carcinoma, pheochromocytoma and hyperparathyroidism; MEN 2B by medullary thyroid carcinoma, pheochromocytoma and characteristic stigmata. Activating germline mutations of the *RET* proto oncogene are responsible for this hereditary syndrome. Codon 634 mutations are the most common mutations occurring in MEN 2A families whereas a specific mutation at codon 918 is observed in the great majority of MEN 2B families. Analysis of these codons will provide a final diagnosis in the great majority of affected families making unnecessary further studies. To specifically study the codons 634 and 918 we used a minisequencing method as an alternative method to complete sequencing.

Results: Using this mutation detection method we were able to reproduce in all cases, representative of 7 families, the information previously obtained by direct sequencing of PCR products. Depending on the number of primers used in the minisequencing reaction, we were able to interrogate either only one nucleotide of the target codon or the three nucleotides simultaneously.

Conclusions: This technique appears as a simple, rapid and efficient method for genetic screening of MEN 2 families. It can be utilized to seek for unknown mutations at specific codons or to screen for previously identified mutations and is therefore of interest to study index cases or individuals at risk. Results suggest that complete sequencing is unnecessary.

Background

The multiple endocrine neoplasia (MEN) type 2 syndromes comprise three related autosomal dominant cancer syndromes [1]. MEN 2A is characterised by medullary thyroid carcinoma (MTC), pheochromocytoma and parathyroid hyperplasia or adenoma [2]. MEN 2B is similar to MEN 2A except that hyperparathyroidism is rare and

characteristic abnormalities such as marfanoid habitus, mucocutaneous and intestinal tract ganglioneuromatosis are present [3]. In familial medullary thyroid carcinoma (FMTC), medullary thyroid carcinoma is the only manifestation [4].

Germline mutations of the *RET* proto oncogene, which encodes for a receptor tyrosine kinase, are the underlying cause of MEN 2 [5–7].

The majority of mutations in MEN 2A and FMTC families are found in 1 of 5 cysteine codons (609, 611, 618 and 620 in exon 10 and 634 in exon 11) that encode part of the extracellular cysteine rich domain [1,8,9]. Codon 634 mutations are the most common in MEN 2A families occurring in 85% of these families [1]. Mutations in FMTC families are evenly distributed among the 5 cysteine codons [1,8]. Less frequently, mutations are found elsewhere in the RET[10-13]. The great majority of MEN 2B families (95%) present the same mutation (ATG \rightarrow ACG) at codon 918 in exon 16 [1]. In a few cases of MEN 2B families a mutation was found in exon 883 [14]. Analysis of codons 634 and 918 of the RET proto oncogene will therefore provide a final diagnosis in the great majority of 2A and 2B families.

Different methods have been used to identify *RET* proto oncogene mutations from which sequencing is the gold standard. In the present study, we adapted the primer extension technique [15], more frequently known as the minisequencing method [16,17], to mutational analysis of MEN 2 families. The detection of the products obtained was performed on a capillary electrophoresis system.

Results and Discussion

Mutations at codon 634 were identified either by a single reaction or a multiplex reaction. Using only one primer to interrogate the first nucleotide, patients harbouring the TGC→CGC mutation presented two peaks corresponding to T/C reactions as shown in figure 1 in contrast with a single peak observed in normal individuals (fig 2). When the three nucleotides were analysed simultaneously, by a multiplex reaction, patients presented an additional fourth peak corresponding to the mutation (fig 3). Similarly, using only one primer to interrogate the second nucleotide, patients harbouring the TGC

TAC mutation presented two peaks corresponding to G/A reactions (fig 4) in contrast with a single peak observed in normal individuals (fig 5). Simultaneous analysis of the three nucleotides reproduced results documenting a fourth peak corresponding to the mutation (fig 6).

Study of the 2B patient was performed with a single primer designed to screen the most common mutation within exon 16, at codon 918, ATG \rightarrow ACG. The patient was heterozygous for the mutation showing two differently coloured peaks corresponding to T/C reactions (fig 7) in contrast with a single peak corresponding to T reaction observed in a normal individual (fig 8).

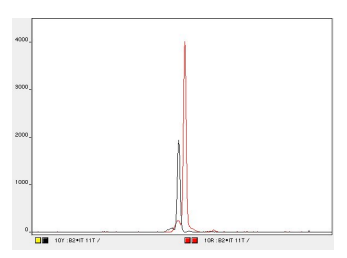


Figure I
Analysis of the first nucleotide of codon 634 by a single reaction – pattern observed in an individual heterozygous for the mutation T/C at the first position. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

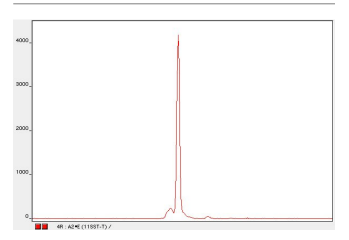


Figure 2
Analysis of the first nucleotide of codon 634 by a single reaction — normal pattern corresponding to the homozygous wild type. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

Using the minisequencing method we were able to recognize all mutations tested (C634R, C634Y, M918T).

Present data support the usefulness of a multiplex reaction to identify the mutation responsible for the disease in a particular family whereas subsequent screening of individuals at risk is possible performing a single reaction.

Conclusions

Germline mutations occurring in MEN 2 families are clustered in a few codons of *RET* proto oncogene. Whereas the great majority of MEN 2A families present different muta-

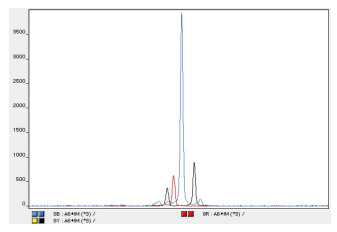


Figure 3Analysis of the three nucleotides of codon 634 by a multiplex reaction – pattern observed in an individual heterozygous for the mutation T/C at the first position. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

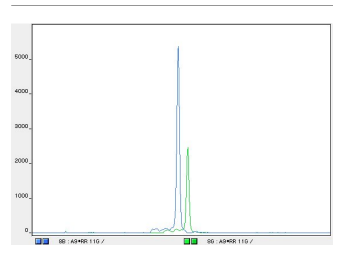


Figure 4
Analysis of the second nucleotide of codon 634 by a single reaction – pattern observed in an individual heterozygous for the mutation G/A at the second position. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

tions at codon 634, the great majority of MEN 2B families present a single mutation at codon 918.

Among different PCR based methods used to identify *RET* proto oncogene mutations in MEN 2 families, complete sequencing is the most reliable procedure.

To specifically study codons 634 and 918 we used a minisequencing method as an alternative method to complete sequencing.

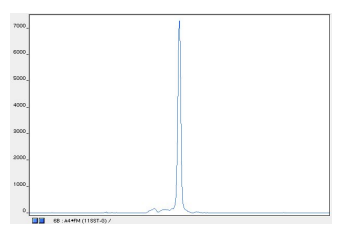


Figure 5
Analysis of the second nucleotide of codon 634 by a single reaction – normal pattern corresponding to the homozygous wild type. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

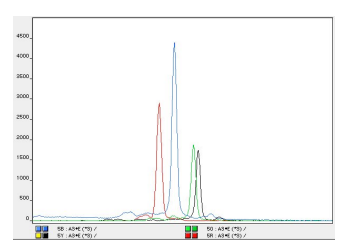


Figure 6
Analysis of the three nucleotides of codon 634 by a multiplex reaction – pattern observed in an individual heterozygous for the mutation G/A at the second position. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

Using this mutation detection method we were able to reproduce in all cases the information previously obtained by direct sequencing of PCR products.

Major advantages of the minisequencing method include its lower cost and speed.

The same procedure may be useful for diagnosis of other genetic disorders associated with point mutations at specific codons.

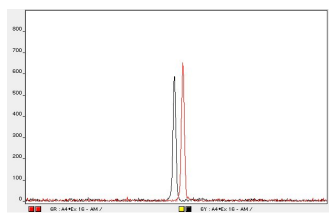


Figure 7 Analysis of the second nucleotide of codon 918 by a single reaction – pattern observed in an individual heterozygous for the mutation T/C at the second position. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

Materials and Methods DNA extraction

High molecular weight DNA from white blood cells was prepared using the guanidine-HCl method adapted from Sambrook [18].

DNA amplification and purification

Seven DNA samples from different individuals representative of 5 MEN 2A families, 1 FMTC family and a *de novo* case of MEN 2B were assayed. All were known to harbour a germline mutation in the *RET* proto oncogene. Four carried a C634R mutation, two a C634Y mutation and one a M918T mutation. Adequate negative controls were included.

Approximately 100 ng of genomic DNA was amplified in a 40 μl reaction containing 20 mM Tris-HCl (pH 8.4), 50 mM KCl, 1.5 mM MgCl₂, 0.2 mM dNTPs, 2 units of Taq polymerase and 1 μM of each primer. To amplify exon 11 (codon 634) the primers used were CRT19A and CRT19B [7]; to amplify exon 16 (codon 918) the primers used were 16F and 16 R [6]. Genomic DNA was denatured for 5 minutes at 94°C prior to 35 cycles of denaturation for 45 seconds at 94°C, annealing for 45 seconds at 62°C and extension for 45 seconds at 72°C followed by a 7 minute 72°C polishing step. The reactions were run on a UNO II thermocycler (Biometra, Göttingen, Germany).

PCR products were purified through low melting point agarose and eluted using the Wizard PCR Preps kit (Promega, Southampton, Hants, UK) according to the manufacturer's protocol.

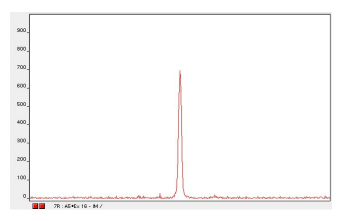


Figure 8
Analysis of the second nucleotide of codon 918 by a single reaction – normal pattern corresponding to the homozygous wild type. Colour is assigned to individual ddNTPs as follows: green/A, black/C, blue/G, red/T.

The minisequencing method

We adapted the ABI PRISM SNaPshot™ ddNTP Primer Extension Kit (Applied Biosystems, USA) to mutational analysis of MEN 2 families. This technique was primarily devised to detect single nucleotide polymorphisms (SNPs). It depends on a primer extension reaction starting from the purified amplified target. A specific minisequencing primer, which is exactly one base short of the polymorphic site, is used for each nucleotide to be studied. Primer extension reactions are carried out using fluolabelled dideoxynucleotides (F ddNTP). Incorporation of the labelled nucleotide is dependent on the genotype. Interruption of the reaction occurs after incorporation of only one terminator. Since purification steps are not necessary, the product obtained is immediately separated electrophoretically; the colour of the peaks obtained makes it possible to identify the SNP or mutation.

In order to search for 634 mutations two different approaches were used. First we interrogated each nucleotide separately, performing 3 reactions and using only one primer in each reaction. Thereafter the three nucleotides were interrogated simultaneously performing a multiplex reaction with three primers. Both single and multiplex reactions were performed according to the manufacturer's instructions. Primers were designed to interrogate the three nucleotides and as so to identify any mutation at this codon: 11SST-T 5' atc cac tgt gcg acg agc tg 3'; 11SST-G 5' ccc cca tcc act gtg cga cga gct gt 3'; 11SST-C 5' aaa aaa aaa cca ctg tgc gac gag ctg tg 3'.

The M918T mutation was easily detected using a specific primer, 16SST-T 5' tcg gat tcc agt taa atg ga 3', one base

short of the second nucleotide of codon 918 in a single reaction.

The separation of the products obtained was performed using the ABI PRISM® 310 Genetic Analyzer (Applied Biosystems, USA) with POP 4 polymer, 47 cm capillary. Run time approximately 12 minutes. The peak signal was analysed with the GeneScan Analysis Software version 3.1.

Authors' Contributions

MJB: Conducted the molecular analysis of MEN 2 families, designed the study and drafted the manuscript. RD: carried out the SnaPshot reactions. LS: participated in coordination of the study and revision of results.

All authors read and approved the final manuscript.

Acknowledgements

This study was partially supported by funds from POCTI/QCAIII, Portugal.

References

- Eng C, Clayton D, Schuffenecker I, Lenoir G, Cote G, Gagel RF, Ploos van Amstel HK, Lips CJM, Nishisho I, Takai SI, et al: The relationship between specific RET proto-oncogene mutations and disease phenotype in Multiple Endocrine Neoplasia Type 2. JAMA 1996, 276:1575-1579
- 2. Schimke RN: **Genetic aspects of multiple endocrine neoplasia.** *Annu Rev Med* 1984, **35:**25-31
- Schimke RN, Hartman WH, Prout TE, Rimoin DL: Syndrome of bilateral pheochromocytoma, medullary thyroid carcinoma and multiple neuromas: a possible regulatory defect in the differentiation of chromaffin tissue. N Engl J Med 1968, 279:1-7
- Farndon JR, Leight GS, Dilley WG, Baylin SB, Smallridge RC, Harrison TS, Wells SA Jr: Familial medullary thyroid carcinoma without associated endocrinopathies: a distinct clinical entity. Br J Surg 1986, 73:278-281
- Mulligan LM, Kwok JBJ, Healey CS, Elsdon MJ, Eng C, Gardner E, Love DR, Molet SE, Moore JK, Papi L, et al: Germline mutations of the RET 1993, proto-oncogene in multiple endocrine neoplasia type 2A:458-460
- Hofstra RMW, Landsvater RM, Ceccherini I, Stulp RP, Stelwagen T, Luo Y, Pasini B, Höppener JWM, Ploos van Amstel HK, Romeo G, et al: A mutation in the RET proto-oncogene associated with multiple endocrine neoplasia type 2B and sporadic medullary thyroid carcinoma. Nature 1994, 367:375-376
- Eng C, Smith DP, Mulligan LM, Nagai MA, Healey CS, Ponder MA, Gardner E, Scheumann GFW, Jackson CE, Tunnacliffe A, Ponder BAJ: Point mutation within the tyrosine kinase domain of the RET proto-oncogene in multiple endocrine neoplasia type 2B and related sporadic tumours. Hum Mol Genet 1994, 3:237-241
- Mulligan LM, Marsh DJ, Robinson BG, Shuffnecker I, Zedenius J, Lips CJ, Gagel RF, Takai SI, Noll WW, Fink M, et al: Genotype-phenotype correlation in multiple endocrine neoplasia type 2: report of the international RET mutation consortium. J Int Med 1995, 238:343-346
- Komminoth P, Kunz EK, Matias-Guiu X, Hiort O, Christiansen G, Colomer A, Roth J, Heitz PU: Analysis of RET proto-oncogene point mutations distinguishes heritable from nonheritable medullary thyroid carcinomas. Cancer 1995, 76:479-489
- Bolino A, Śchuffenecker I, Luo Y, Seri M, Silengo M, Tocco T, Chabrier G, Houdent C, Murat A, Schlumberger M, et al: RET mutations in exons 13 and 14 of FMTC patients. Oncogene 1995, 10:2415-2419
- Eng C, Smith DP, Mulligan LM, Healey CS, Zvelebil MJ, Stonehouse TJ, Ponder MA, Jackson CE, Waterfield MD, Ponder BAJ: A novel point mutation in the tyrosine kinase domain of the RET proto-oncogene in sporadic medullary thyroid carcinoma and in a family with FMTC. Oncogene 1995, 10:509-513

- Fattoruso O, Quadro L, Libroia A, Verga U, Lupoli G, Cascone E, Colantuoni V: A CTG to ATG novel point mutation at codon 804 in exon 14 of the RET proto-oncogene in two families affected by familial medullary thyroid carcinoma. Hum Mutat 1998, Suppl 1:167-171
- Hofstra RM, Fattoruso O, Quadro L, Wu Y, Libroia A, Verga U, Colantuoni V, Buys CH: A novel point mutation in the intracellular domain of the RET proto-oncogene in a family with medullary thyroid carcinoma. J Clin Endocrinol Metab 1997, 82:4176-4179
- Gimm O, Marsh DJ, Andrew SD, Frilling A, Dahia PL, Mulligan LM, Zajac JD, Robinson BG, Eng C: Germline dinucleotide mutation in codon 883 of the RET proto-oncogene in multiple endocrine neoplasia type 2B without codon 918 mutation. J Clin Endocrinol Metab 1997, 82:3902-3904
- Sokolov BP: Primer extension technique for the detection of single nucleotide in genomic DNA. Nucleic Acids Res 1990, 18:3671
- Pastinen T, Kurg A, Metspalu A, Peltonen L, Syvänen A-C: Minisequencing: a specific tool for DNA analysis and diagnosis on oligonucleotide arrays. Genome Res 1997, 7:606-614
- Syvänen A-C: From gels to chips: "Minisequencing" primer extension for analysis of point mutations and single nucleotide polymorphisms. Hum Mutat 1999, 13:1-10
- Sambrook J, Fritsch EF, Maniatis T: Molecular Cloning. A laboratory manual. Cold Spring Harbor, Cold Spring Harbor Press 1989

Publish with **BioMed** Central and every scientist can read your work free of charge

"BioMedcentral will be the most significant development for disseminating the results of biomedical research in our lifetime."

Paul Nurse, Director-General, Imperial Cancer Research Fund

Publish with BMC and your research papers will be:

- available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- yours you keep the copyright

Submit your manuscript here: http://www.biomedcentral.com/manuscript/

