NOVA MUTAÇÃO DO GENE SDHD EM PACIENTES COM PARAGANGLIOMAS DO CORPO CAROTÍDEO

PATIENTS WITH NEW SDHD GENE MUTATION WITH CAROTID BODY PARAGANGLIOMAS

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Received a 10 of May 2017
Accepted a 07 of May 2018

RESUMO

Introdução: Os paragangliomas (PGLs) são neoplasias neuroendócrinas que podem ocorrer em todo o corpo onde exista paraganglia. Representando 0,03% de todos os tumores, os PGLs são extremamente raros. Embora predominantemente benignos e passíveis de cura por ressecção cirúrgica, estima-se que até 6% possam ser malignos. PGLs familiares têm um modo de transmissão autossômico dominante, e uma variabilidade fenotípica significativa. Até ao momento, foram identificados três genes associados aos PGLs familiares. Todas as subunidades de codificação (D, B e C) do complexo enzima succinato desidrogenase (SDH), fazem parte do ciclo de Krebs e da cadeia transportadora de electrões.

Objectivos: Pretende-se descrever uma nova frameshift mutation no gene SDHD identificado numa família com Paragangliomas do Corpo Carotídeo.

Resultados: A análise de mutação do probando revelou a presença de uma nova frameshift mutation, c.549delG (p.L139Ffs), no exão 4 do gene SDHD. Ambos os descendentes herdaram essa mutação patogênica, embora um deles ainda esteja subclínico.

Conclusão: Descrevemos uma nova mutação causal de frameshift na subunidade D do SDH (SDHD) numa família com Paragangliomas do Corpo Carotídeo. Esta descoberta contribui para o conhecimento do espectro mutacional do SDHD, e para ajudar o aconselhamento genético desta família. Destacamos o facto de agora ser possível oferecer a outros familiares, ainda subclínicos, um teste preditivo que permite auxiliar uma vigilância ou intervenção precoce, e consequentemente melhorar o prognóstico.

Palavras-chave
Paraganglioma; Tumor do corpo carotídeo; Paraganglioma familiar; Mutação; SDHD; Succinato desidrogenase; Cirurgia.

ABSTRACT

Paragangliomas (PGLs) are neuroendocrine neoplasms that can occur throughout the body wherever there is paraganglia. Representing 0.03% of all tumours, PGLs are extremely rare. Although predominantly benign and amenable to cure by surgical resection, up to 6% can be malignant. To date, three genes have been identified that are associated with Familial PGLs. All three encode subunits (D, B and C) of the enzyme succinate dehydrogenase complex (SDH), which is part of the Kreb's cycle and the electron transport chain. We report a novel causative frameshift mutation in the subunit D of SDH (SDHD) in a family with Carotid Body Paragangliomas. This finding contributes for extending the known mutational spectrum of SDHD, and to help the genetic counseling of this family. Noteworthy, is now possible to offer to other relatives, still sub-clinical, a predictive test that would eventually aid an early surveillance/intervention for a better prognosis.

Keywords
Carotid body paraganglioma; Familial paraganglioma; Mutation; SDHD; Succinate dehydrogenase; Surgery.

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INTRODUCTION

Paragangiomas (PGLs) are neuroendocrine neoplasms that can occur throughout the body wherever there is paraganglia, non-neuronal cells derived from the neural crest so named for being in close proximity to sympathetic ganglia. There are fundamentally two types of paraganglia: chromaffin or sympathetic made of chromaffin cells, which have primary endocrine functions and non-chromaffin or parasympathetic made of glomus cells, which have primary chemoreceptor functions.\(^1\)

Representing 0.03% of all tumours, PGLs are extremely rare. Although predominantly benign and amenable to cure by surgical resection, up to 6% can be malignant.\(^2\)

About 50% are found in the head and neck region, most commonly as highly vascularised carotid body tumors, which represent approximately 65% of head and neck PGLs.\(^3\)\(^,\)\(^4\) PGLs can also be found as aortic glomus, jugular bulb and vagal and tympanic nerves. Depending on their location, patients with carotid body PGLs may notice a slow growing and painless lateral neck mass, pulse-like sensations or voice changes.

Three different types of Carotid Body PGLs have been described in the literature: Sporadic, Familial and Hyperplastic. The Sporadic form is by far the most common type, representing approximately 85% of these tumors. The prevalence of the Familial type has varied between reported studies from as low as 5% to as high as 50%.\(^5\)

This wide-ranging variability stems from the existence of hidden familial cases. In the latter, diagnosis is usually made earlier (30–35 years) and there’s a higher prevalence of bilateral multifocal PGLs.\(^6\)

The Hyperplastic form is very common in patients with chronic hypoxia, such as those in New Mexico, Peru and Colorado who live at a high altitude (>1524 meters/5000 feet above sea level) and in patients with chronic obstructive pulmonary disease (COPD) or cyanotic heart disease.\(^7\)

Familial PGLs have an autosomal dominant mode of inheritance, therefore all generations of a family can be affected. Nevertheless, due to their exceedingly high phenotypic variability, very different clinical manifestations can occur in patients with the same genotype, even among family members. Genetic heterogeneity, incomplete penetrance and genetic imprinting lie at the root of this phenotypic variability. To date, three genes have been identified that are associated with Familial PGLs. All three encode subunits (D, B and C) of the enzyme succinate dehydrogenase complex (SDH), which is part of the Kreb’s cycle and the electron transport chain. Defective succinate dehydrogenase has been postulated to cause an increase in the intracellular concentration of molecular hypoxia mediators and the vascular endothelial growth factor (VEGF), thus resulting in hyperplasia, angiogenesis, and neoplasia.\(^7\)

Furthermore, mutations in the SDHD gene show a parent-of-origin effect. Consequently, the disease only occurs when the mutation is inherited from the father.\(^8\)

To date, several different mutations in the SDHD gene have been identified, which can be consulted on “The Human Gene Mutation Database” (http://hgmd.cf.ac.uk).

We report a family affected by hereditary PGL in which a novel mutation in the SDHD gene was identified.

CASE PRESENTATION

The family pedigree is presented in Figure 1.

Figure 1 Pedigree of the PGL family studied. Genetic tests were performed on individuals indicated by an asterisk.

The index case (II.1) was a 58-year-old man with a known history of bilateral PGL of the carotid body. The diagnosis had been established 35 years earlier when he first presented to the Vascular Surgery outpatient clinic for evaluation of a left latero-cervical mass. Angiography suggested a carotid body PGL that was surgically removed through subadventitial dissection. A similar mass was detected in the contralateral side 9 years later and it was also removed through subadventitial dissection. In both cases, the paraganglioma diagnosis was confirmed histologically. The current mutation analysis was conducted in the context of a systematic review of paraganglioma cases operated in the Vascular Surgery Department of Centro Hospitalar e Universitário de Coimbra (CHUC). DNA was isolated from peripheral blood using the DNA isolation kit and all the coding region of the SDHD gene (1-4 exons) was PCR amplified. The presence of the pathogenic mutation was always confirmed on a separate amplification with subse-
In the present study, the mutation analysis for SDHD gene has been performed in a family with PGLs of the carotid body. An heterozygous mutation, c.549delG; p. L139Ffs in the SDHD gene has been identified in the index case and in his two offsprings (one still unaffected). This frameshift mutation was novel, not previously described (http://hgmd.cf.ac.uk) and was located in exon 4 of the SDHD gene (Figure 3).

DISCUSSION

In the present study, the mutation analysis for SDHD gene has been performed in a family with PGLs of the carotid body. An heterozygous mutation, c.549delG; p. L139Ffs in the SDHD gene has been identified in the index case and in his two offsprings (one still unaffected). This frameshift mutation was novel, not previously described (http://hgmd.cf.ac.uk) and was located in exon 4 of the SDHD gene (Figure 3).
The genetic testing for this one base pair deletion was extended to the sister of the index patient and in accordance with her phenotype, no mutation has been found. As illustrated in this family, PGLs should always be considered in patients presenting with swelling at the lateral neck, cranial nerve palsies, voice changes or auditory defects. A detailed family history and thorough clinical examination are of paramount importance. Ultrasound of soft tissue and vessels of the neck is mandatory. MR angiography should also be performed. The successful management of PGLs hinges on early diagnosis. The familial type has been associated with age at diagnosis <40 years, bilateral multifocal tumors, preceding pheochromocytoma and PGL family history. That said, due to the disease’s striking phenotypic variability, individuals with Familial PGLs can erroneously be clinically ascertained as having sporadic appearing PGLs. Furthermore, it has been reported than up to 30% of apparently sporadic PGLs are caused by germline mutations. Familial PGLs are thus underestimated. Diagnosis is often established when a patient seeks clinical advice because of a palpable mass. At that stage, the tumor is often large enough to pose significant intra-operative challenges. Since the only cure for PGLs is complete surgical excision, early diagnosis is critical for the operative outcome. The incidence of additional vascular procedures and nerve lesions has in fact been reported to be significantly lower in mutation carriers. The timely detection of both unilateral and contralateral PGLs in these patients enabled them to be operated upon earlier, which translated into less surgical complications, consequence of the more favorable operative outcome associated with smaller tumors. Mutation carriers should be screened clinically and radiologically. Ultrasound and additional MR angiography scans of the head and neck regions, thorax, abdomen and pelvis should be performed so that PGLs can occur throughout the entire vegetative system. Genetic testing should be extended to children. Accordingly to the European-American Paraganglioma Study Group, screening should start at 10 years of age, with follow-up screens every 2 years. The tumor’s sluggish growth, with doubling times varying from 4 and 14 years, obviates the need for annual examinations. Non-carrier family members don’t need to be screened. Population-specific variations in the prevalence, penetrance, and phenotypic expression of SDH subunit gene mutations have been postulated. The well-documented association of SDH-D mutations with head and neck PGLs may in fact vary across different populations. Further genetic studies are necessary.

We report a family with a novel heterozygous frameshift mutation in the SDHD gene, expanding the clinical and genetic heterogeneity of hereditary PGLs. This study highlights the importance of mutation analysis for individual follow-up strategy, in particular for mutation carriers who should undergo regular clinical examination. This procedure will certainly provide an early detection of PGLs in a context of genetic counseling and thereby improve the operative outcome of these patients.

REFERENCES