Germline PTEN mutations in Cowden syndrome-like families

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Abstract
Cowden syndrome (CS) or multiple hamartoma syndrome (MIM 158350) is an autosomal dominant disorder with an increased risk for breast and thyroid carcinoma. The diagnosis of CS, as operationally defined by the International Cowden Consortium, is made when a patient, or family, has a combination of pathognomonic major and/or minor criteria. The CS gene has recently been identified as PTEN, which maps at 10q23.3 and encodes a dual specificity phosphatase. PTEN appears to function as a tumour suppressor in CS, with between 13-80% of CS families harbouring germline nonsense, missense, and frameshift mutations predicted to disrupt normal PTEN function. To date, only a small number of tumour suppressor genes, including BRCA1, BRCA2, and p53, have been associated with familial breast or ovarian/cancer families. Given the involvement of PTEN in CS, we postulated that PTEN was a likely candidate to play a role in families with a "CS-like" phenotype, but not classical CS. To answer these questions, we gathered a series of patients from families who had features reminiscent of CS but did not meet the Consortium Criteria. Using a combination of denaturing gradient gel electrophoresis (DGGE), temporal temperature gel electrophoresis (TTGE), and sequence analysis, we screened 64 unrelated CS-like subjects for germline mutations in PTEN. A single male with follicular thyroid carcinoma from one of these 64 (2%) CS-like families harboured a germline point mutation, c.297T→C. This mutation occurred at the last nucleotide of exon 3 and within a region homologous to the cytoskeletal proteins tensin and auxilin. We conclude that germline PTEN mutations play a relatively minor role in CS-like families. In addition, our data would suggest that, for the most part, the strict International Cowden Consortium operational diagnostic criteria for CS are quite robust and should remain in place.

Keywords: PTEN; Cowden syndrome; breast; thyroid carcinoma
Breast and thyroid carcinoma are two frequently occurring neoplasms in the female population. Increased risks for both breast and thyroid cancer are prominent features of Cowden syndrome (CS). The hallmark phenotype of this inherited cancer syndrome is the presence of hamartomas, developmentally incorrect, benign, hyperplastic growths, in multiple organ systems including the skin, gastrointestinal tract, central nervous system, breast, and thyroid. Breast cancer will develop in 25-50% of women with CS and 3-10% of all CS patients will develop thyroid cancer. 4 At present, only four tumour suppressor genes have been associated with familial breast cancer, BRCA1, BRCA2, p53, and PTEN.5 Initially thought to account for over 80% of hereditary breast cancer, 6 germline mutations in BRCA1 and BRCA2 together are now thought to account for 25-50% of all familial breast cancer, 7 thus opening up the possibility of other BRCAX genes. Along these lines, germline mutations in p53 are associated with 70% of cases of Li-Fraumeni syndrome, an autosomal dominant condition comprising breast cancer, brain tumours, sarcomas, and adrenocortical carcinomas. 8-11 Recently, the CS susceptibility gene has been identified as the tumour suppressor gene PTEN, also known as MMAC1 and TEP1. 7-12-14 PTEN maps to 10q23.3 and encodes a 403 amino acid dual specificity phosphatase. 12-15 Germline missense and truncating mutations have been reported in between 13-80% of patients with CS. 16-18 It should be noted that while initial linkage studies of 12 families with CS was highly suggestive of a single locus for CS, 18 a subsequent study proposes that genetic heterogeneity may exist in CS. 18 At the somatic level, PTEN has been shown to be mutated or deleted in a number of human malignancies, including sporadic breast, brain, prostate, and kidney cancer cell lines, as well as in a number of primary tumours including endometrial carcinomas, glioblastomas, malignant melanoma, and thyroid and breast tumours. 20-33 Given the role of PTEN in CS and the relatively large percentage of familial cases of breast cancer that are not caused by germline mutation of BRCA1, BRCA2, or p53, we sought to determine whether PTEN may be mutated in...
Table 1  Phenotypic classification of CS-like families

<table>
<thead>
<tr>
<th>Phenotype of families</th>
<th>No of families</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast and thyroid carcinoma occurring in at least one person</td>
<td>22</td>
</tr>
<tr>
<td>Breast and thyroid carcinoma occurring in different subjects</td>
<td>32</td>
</tr>
<tr>
<td>Breast carcinoma and thyroid disease (eg goitre)</td>
<td>3</td>
</tr>
<tr>
<td>Breast carcinoma/CS-like (eg trichilemmoma), no thyroid involvement</td>
<td>6</td>
</tr>
<tr>
<td>Thyroid carcinoma/CS-like, no breast involvement</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>64</td>
</tr>
</tbody>
</table>

Figure 1  DGGE detection of c.209T→G in the germline of a patient from a CS-like family. Control mutations from CS and BRR families are also included to display the sensitivity of this technique for the detection of PTEN mutations. Lane 1, wild type control (exon 3); lane 2, Y68H (exon 3); lane 3, IVS2-2A→G (exon 3); lane 4, c.209T→G (exon 3); lane 5, wild type control (amplion 5I, representing the 5' half of exon 5); lane 6, Q87X (amplion 5I); lane 7, c.347-351delCAAT (amplion 5I); lane 8, wild type control (amplion 5I, representing the 3' half of exon 5); lane 9, C124R (amplion 5I); lane 10, E157X (amplion 5I); lane 11, wild type control (exon 7); lane 12, R233X (exon 7); lane 13, c.791ATIns (exon 7).

DENATURED GRADIENT GEL ELECTROPHORESIS (DGGE) AND TEMPORAL TEMPERATURE ELECTROPHORESIS (TTGE)

A combination of DGGE and TTGE was performed for all nine exons of PTEN. GC-clamped primer sequences, PCR conditions, and DGGE conditions have been previously described, with the exception of the primers for exons 2 and 4. Exon 2 and 4 primer sequences, with GC clamps added, were as follows: exon 2, 2F, 5'-CGT CCC GGC TTT GAT TGC TGC ATA TTT CAG-3' and 2R, 5'-CCG CCG CCG CCG CCC GGC CCC GTC CCG CCG CCC CCC GTC TAA ATG AAA ACA CAA CAT G-3'; exon 4, 4F, 5'-CGC CGG CCG CCC CCC CCC CCG CCC TGG CCG CCG CCC CCC CCC GAA ATA ATA AAC ATT ATA AAG ATT CAG GCA ATG-3' and 4R, 5'-GAC AGT AAG AAT CAG CAG TCT ATC-3'. Split exon 5 primers with GC clamps and conditions for mutation detection have been previously reported.

TTGE is a mutation detection technique using the basic PCR fragment denaturation principles of DGGE. The major difference between these methods is that a temperature gradient, rather than a chemical gradient of varying urea and glycerol percentages, is used for strand separation of the GC-clamped homoduplex PCR products by generating a linear temperature gradient over the length of the electrophoresis run (Bio-Rad Laboratories, Hercules, CA). One or 0.75 mm thick gels of 10% polyacrylamide:bis (37:5:1) (Bio-Rad Laboratories) and 7 mol/l urea (Bio-Rad Laboratories) were run using the DCode™ Universal Mutation Detection System (Bio-Rad Laboratories). Electrophoresis was performed at 130 V for six hours with a temperature gradient of 46-58°C and a ramp rate of 2°C per hour. TTGE fragments were visualized under ultraviolet transillumination after the gel was stained with ethidium bromide (Bio-Rad Laboratories).

Both DGGE and TTGE have proven high accuracy in detecting mutations in general and specifically in detecting known PTEN mutations from CS patients (fig 1).
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SEQUENCE ANALYSIS
Exons which showed DGGE and TTGE variants underwent direct sequence analysis. The PCR primers and reaction conditions have been described elsewhere. When PCR products were gel isolated and purified using the Wizard PCR Preps DNA Purification System (Promega, Madison, WI). Direct sequencing of these products was performed using the ABI Prism dye terminator cycle sequencing ready reaction kit (Perkin-Elmer Corp., Norwalk, CT). Cycle sequencing products were electrophoresed on 6% Long range gels (FMC Bioproducts, Rockland, ME) and analysed on an Applied Biosystems model 373A automated DNA sequencer (Perkin-Elmer Corp).

PTEN POLYMORPHISM ANALYSIS
A previously identified intronic polymorphic site in PTEN, IVS8+32G/T, was analysed in a single affected member from each CS-like family to investigate hemizygosity at the PTEN locus in mutation negative families. This site is moderately heterozygous, with an earlier report finding 50% of samples to be informative. Potential hemizygosity was assessed by the amplification of exon 8 and flanking intronic sequence and digestion with the restriction endonuclease HincII under conditions suggested by the manufacturer (New England Biolabs, Beverly, MA).

Results

PTEN MUTATION ANALYSIS
A missense point mutation, c.209T→C (L70P), predicted to affect splicing was identified in a single affected patient (1 of 64, 2%) (fig 1). This mutation was not identified in 100 normal alleles. When this occult germline PTEN mutation was identified, the family history was reassessed (fig 2). The subject analysed for this study, III.1, developed follicular thyroid carcinoma at the age of 31. His mother, II.2, had breast adenocarcinoma diagnosed at the age of 49 and again at 53. She also had endometrial carcinoma diagnosed at 63 years. Careful clinical assessment of these two subjects was unable to identify macrocephaly, skin lesions typical of CS, or scrotal tongue. The maternal grandfather, I.1, was diagnosed with leukaemia at the age of 57. Unfortunately, family members other than III.1 were unavailable for analysis. Fresh tumour from III.1, which would have allowed us to study the putative aberrant splicing effect of this mutation, was also unavailable. No mutations were identified in the other 63 unrelated CS-like families.

PTEN POLYMORPHISM ANALYSIS
 Forty-eight percent (30 of 63) of unrelated subjects from PTEN mutation negative CS-like families were found to be heterozygous at the IVS8+32T/G site. This analysis would suggest that, at least in these families, gross germline deletion of PTEN can be excluded.

Discussion
An occult germline PTEN mutation, c.209T→C at the last nucleotide of exon 3 was found in one of 64 (2%) CS-like families. This family's cancers, comprising leukaemia, which may or may not be related, adenocarcinoma of the breast, endometrial carcinoma, and follicular thyroid carcinoma, together do not meet the International Cowden Consortium Criteria used for the diagnosis of CS in this study. However, we cannot exclude the possibility that this family represents a case of low penetrance CS. The family with PTEN mutation in this study contrasts with that in a recent study that reported a PTEN mutation in a family initially classified as having breast and thyroid tumours only but reclassified as CS after mutation analysis led to closer clinical assessment. Closer clinical assessment of the family presented in the current study did not identify additional features of CS.

In the remaining families where no occult germline mutations were identified, it is highly unlikely that these mutations would have gone undetected. Both DGGE and TTGE are highly sensitive mutation detection techniques and both have been shown consistently to detect known PTEN mutations and other sequence polymorphisms (Marsh and Eng, unpublished data, 1998; fig 1). Further, because at least one affected member from nearly half of these mutation negative families was heterozygous at the IVS8+32T/G polymorphism, whole gene deletion is unlikely, at least in these families.

In CS, while missense and truncating mutations are scattered largely along the entirety of PTEN, a mutational “hot spot” exists in exon 5, which contains the PTPase core motif at codons 122-132. Thus, many mutations in CS are predicted to disrupt the phosphatase function of this protein. Interestingly, the mutation identified in exon 3 falls in the N-terminal half of the PTEN protein that has been shown to have some sequence similarities to the cytoskeletal proteins tensin and auxilin.
Specifically, the leucine residue at codon 70 that is altered by this T to C point mutation (L70P) is conserved in both bovine auxilin and chicken tensin. Thus, it is possible that this mutation may be affecting the phosphatase function of this protein, as one may predict if this putative splice site mutation leads to a truncated protein, and may also function to disrupt normal cellular motility and cell-cell interactions.

Whether germline PTEN mutations are associated with CS and related inherited hamartoma syndromes (Bannayan-Ruvalcaba-Riley syndrome, [BRR, MIM 153480] and juvenile polyposis syndrome [JPS, MIM 174900]), as well as syndromes comprising partial CS phenotypes, is largely unknown. Before the identification of PTEN as the CS gene, it was not inconceivable that the three related hamartoma syndromes and CS-like syndromes were all associated with different mutations in a single gene. We have shown that germline PTEN mutations are associated with the great majority, approximately 80%, of classical CS families.\(^\text{18}\) Nelen et al\(^{17}\) identified PTEN mutations in 47% of CS cases studied. One other study of 23 CS families identified only 13% of families with germline PTEN mutation.\(^{18}\) This was perhaps not surprising as limited linkage information in these families suggested the possibility of genetic heterogeneity in CS, even though initial studies of a group of 12 CS families showed no evidence for heterogeneity.\(^\text{19}\)

We have also shown that germline PTEN mutations account for at least a proportion of BRR, which is characterised by macrocephaly, lipomatosis, thyroid dysfunction, hamartomatous polyps of the gastrointestinal tract, and pigmented macules of the glans penis, but without a known predisposition to breast and thyroid cancer.\(^{19,20}\) How mutations in a single gene, at times identical,\(^{19,20}\) can function to predispose to two overlapping but apparently distinct syndromes, one with malignancy and one without, remains to be elucidated.

Disparate reports concerning the third hamartoma syndrome, JPS, and PTEN mutation or deletion have recently been published.\(^{36,39,41}\) A putative JPS locus, J1P, at 10q22-24 was initially thought to encompass PTEN, although fine structure mapping placed this locus slightly centromeric of PTEN.\(^{42}\) Subsequently, the 10q22-24 region was excluded as a putative JPS locus by linkage analysis in eight JPS families.\(^{35}\) Screening of PTEN in 21 classical JPS families and 16 cases of sporadic JPS did not identify any germline mutations.\(^{36,41}\) In contrast, PTEN mutation has been reported in four patients with "juvenile polyposis".\(^{36,41}\) Although the clinical diagnosis of classic juvenile polyposis in these cases is questionable. Given these genetic data and the phenotypic overlap of these syndromes, we can say with some confidence that if a germline PTEN mutation were detected in a person previously thought to have "juvenile polyposis", then the diagnosis needs to be revised, as that person is likely to have either CS or BRR. Along the same lines, we have now investigated a cohort of families, each of which contains some of the component tumours of CS but do not meet the Consortium diagnostic criteria for CS. Only one such family was found to have an occult germline PTEN mutation, arguing that such germline alterations play a minor role in families that do not meet the strict CS diagnostic criteria. Nonetheless, this finding is significant for three reasons. Firstly, it suggests that the operational diagnostic criteria for CS established by the International Cowden Consortium are, for the most part, robust and are useful for identifying PTEN mutation positive CS families. Secondly, we must also conclude from our data that other genes are involved which lend susceptibility to a CS-like disease and to site specific breast and non-medullary thyroid cancer. Thirdly, for non-CS patients identified with occult PTEN mutations, albeit uncommonly, there are important implications for future hamartoma/cancer development that should impact on surveillance.

Unanswered questions remain, however. For example, are CS-like families without germline PTEN mutations at any less risk of cancer than those with mutations? Preliminary genotype-phenotype analyses suggest that classical CS families without germline PTEN mutations are at lower risk of developing malignant breast disease compared to their PTEN mutation positive counterparts.\(^{18}\) By extrapolation, it would seem that PTEN mutation negative CS-like families should be at decreased risk of developing breast cancer. Unfortunately, this study was unable to confirm this clinically relevant extrapolation. We can conclude, however, that in the majority of cases, germline PTEN mutations lead specifically to a CS or BRR phenotype and that the phenotype of CS-like families is, for the most part, caused by unknown mechanisms.

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