Cowden syndrome and Lhermitte-Duclos disease in a family: a single genetic syndrome with pleiotropy?

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Abstract
Cowden syndrome is an autosomal dominant condition of multiple hamartomas. Patients with this phakomatosis have an increased risk of breast cancer and thyroid tumours. Lhermitte-Duclos disease is usually a sporadic condition of cerebellar ganglion cell hypertrophy, ataxia, mental retardation, and self-limited seizure disorder. We describe a three generation family with Cowden syndrome and Lhermitte-Duclos disease. Karyotyping performed on the peripheral lymphocytes of the proband and her affected mother showed a 46,XX complement. Single strand conformational polymorphism analysis failed to show any germline p53 mutations as a cause of the syndrome in this family.

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Cowden syndrome, an autosomal dominant genodermatosis of multiple hamartomas involving all three germ layers, is associated with an increased risk of breast cancer and thyroid tumours.1 The aetiology of this syndrome is unclear; neither the susceptibility locus nor the chromosomal location is known. Since the first description of Rachel Cowden, the patient after whom the syndrome is named,2 several publications have added to the clinical features associated with this syndrome.1,3,3 Lhermitte-Duclos disease (LDD) is a rare, usually sporadic, condition in which the normal cells of the cerebellum are replaced by a hamartomatous mass of hypertrophic ganglion cells.4 We report a three generation family with Cowden syndrome and LDD: one family member had Cowden syndrome and histologically documented LDD, two family members had Cowden syndrome and a syndrome clinically consistent with LDD, and another family member may have LDD but his whereabouts are unknown. Cytogenetic and p53 mutation analyses are described.

The proband
A 14 year old girl (III.4, fig 1) had a seven month history of multiple, large masses involving both breasts. Biopsy showed giant fibroadenomas. She has no other symptoms, including haematochezia or melaena. Her IQ has been assessed to be approximately 70 and she attends a special school for the educationally subnormal. Her head circumference is 62 cm (>99th centile). Physical examination showed multiple 2–5 mm perioral and mucocutaneous verrucous papillae and trichoepitheliomatous on the right malar area, right arm (fig 2A), and left anterior chest; a scrotal tongue consisting of multiple papillomatous lesions on the dorsum and frenulum (fig 2B); four café au lait spots (each >5 mm) with irregular margins on her thighs and additional freckling on her groins and upper thighs; and bilateral large, heavy breasts in which multiple solid masses could be palpated, the largest measuring 3 cm in diameter. Thyroid examination was unremarkable. She has mild cerebellar ataxia.

The family
The patient’s maternal grandfather (I.1, fig 1) had well documented Cowden syndrome and biopsy proven Lhermitte-Duclos disease.5 He had prominent mucocutaneous papillomatosis, acral keratoses, a plantar angiolipoma, and nodular reactive lymphoid hyperplasia of the right lung. Neurological features were cerebellar ataxia and increased intracranial pressure. A biopsy showed LDD and carotocavernous fistula. However, he was not mentally retarded and was able to work as a travel agent. He died of a stroke from an intracerebellar haemorrhage in his late 50s.

The proband’s mother (II.4, fig 1) is a 37 year old woman who has a history of resected...
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periareolar fibroadenomas of the left breast and grand mal seizures in her teens. At the age of 30, she experienced intermittent upper abdominal swelling. Evaluation with computerised tomography and angiography showed focal nodular hyperplasia of the liver. Her head circumference was 62 cm (>99th centile). Examination of the skin and mucous membranes showed a few mucocutaneous papillae and acral keratoses, but no scrotal tongue. She has mild to moderate bilateral dysmetria and sways from side to side when walking in tandem, although she does not have truncal ataxia, dysarthria, or Romberg’s sign. She had mild mental retardation with an IQ estimated to be 70. Thyroid examination was unremarkable. Mammography showed bilateral fibroadenomas (fig 3). Magnetic resonance imaging of the brain showed megencephaly but no mass lesions in the cerebellum.

The proband’s uncle, II.2, was described by the family to have macrocephaly and mental retardation. Although he may have LDD as well, we are unable to document this as the family is unaware of his whereabouts. To his family’s knowledge, he has no children. The patient’s sister (III.3), aunt (II.3), an uncle (II.1), and his children (III.1, III.2) are unaffected. II.1 has not consented to a medical examination.

Cytogenetic and molecular genetic studies
Chromosome analyses of cultured lymphocytes from the proband and her mother showed a 46,XX karyotype. In addition, no germline mutation in exons 1-11 of the p53 tumour suppressor gene was detected by single-strand conformational polymorphism (SSCP) analysis, using the most sensitive running conditions for the p53 gene.

Discussion
There have been over 100 cases of Cowden syndrome reported in English publications to date, describing more than 50 abnormalities observed in patients with this phakomatosis. Since over 70% of these patients have mucocutaneous papillosis, acral keratoses, and breast disease in affected females (both malignant and benign), these features probably characterise Cowden syndrome. Thyroid disease (both benign and malignant), macrocephaly, gastrointestinal hamartomas, and gynaecological problems are believed to be part of the syndrome as well.

Lhermitte-Duclos disease is believed to be a hamartomatous overgrowth of hypertrophic ganglion cells which replace the granular cell layer and Purkinje cells of the cerebellum. Clinically, LDD may manifest with ataxia, signs and symptoms of raised intracranial pressure, seizures, megencephaly, and mental retardation. By 1992, 62 patients with LDD, without Cowden syndrome, had been described; interestingly, some of these patients have been noted to have breast or other cancers. In the family reported in this paper, the proband’s grandfather (I.1) undoubtedly had LDD as the diagnosis was made by cerebellar biopsy. Although no cerebellar mass was noted in the proband’s mother by magnetic resonance imaging, she has the signs of ataxia, megencephaly, seizures, and decreased IQ.
which suggest the clinical diagnosis of LDD. Similarly, the proband has macrocephaly and mild mental retardation. We argue, therefore, that both the proband and her mother may actually have LDD by clinical criteria, although the classic pathognomonic anatomical signs are absent, and that Cowden syndrome and the clinical features of LDD appear to be co-inherited in this family. In patients with LDD, the age that a cerebellar mass becomes evident on imaging studies is unknown. It may be prudent to follow the proband and her mother expectantly to determine if a classic cerebellar hamartoma, with attendant worsening cerebellar signs, will develop with age. In addition, this report calls attention to clinicians to consider the diagnosis of LDD in any patient with Cowden syndrome with meigencphaly or macrocephaly, mental retardation, seizure disorder, or ataxia.

To date, there are four sporadic cases and three families with both Cowden syndrome and LDD (this report). 7,8 11-13 Both these conditions are of unclear aetiology. Given the rarity of both syndromes and the coincident occurrence of both Cowden syndrome and LDD in the four cases and three families, it seems plausible that both are related pathogenetically. 11 Cowden syndrome and LDD appear to be co-inherited in an autosomal dominant manner in some families, as illustrated by the three families (this report). Cowden syndrome is probably the result of a single, variably penetrant, autosomal dominant gene. The germline mutation in Cowden syndrome families with LDD may be different from that in families without LDD. The clinical features of Cowden syndrome also show intrafamilial variation, which may be the consequence of additional modifying genes. Consistent with this, variable expression of the LDD phenotypes in Cowden syndrome was observed in the two families described by Padberg et al. 14

The location of the susceptibility gene(s) for Cowden syndrome and Lhermitte-Duclos disease is unknown. Cytogenetic studies have been unhelpful, and molecular genetic analyses have rarely been performed. Mutation analysis in RAS, HER2/neu, and p52 genes have been performed in one patient with Cowden syndrome. No mutations were found in the DNA from this patient’s breast tumours. The tumour suppressor gene p53 codes for a 53 kDa phosphoprotein involved in cell cycle regulation and is believed to act as a transcription factor. 15 Somatic p53 mutations are the most common abnormalities in human cancer. 15 Over 90% of p53 mutations are found in exons 5–9. Germline p53 mutations have been described in patients with the Li-Fraumeni syndrome, which is a rare autosomal dominant syndrome of breast cancer, sarcoma, and other cancers involving all three germ layers. 16-18 A survey of the constitutional p53 gene was undertaken in our family for two reasons. Firstly, a patient with multiple primary cancers and multiple tricholemmomata of the scalp (as seen in Cowden syndrome) has been previously described to have a C to T mutation in exon 8 of the p53 gene. 19 Secondly, somatic p53 mutations and altered p53 expressions are found in skin tumours. 20 21 However, no somatic p53 mutations were detected by SSCP analysis in the present family with both Cowden syndrome and LDD. Since SSCP analysis using the most sensitive running condition has a sensitivity of 90% to detect p53 mutations, 7 there is a small possibility that an exonic mutation in the p53 gene was not detected. Other explanations, such as intronic mutations, are less likely as the great majority of known p53 mutations are found within exons 5–9. 22 Therefore, to a 90% likelihood, we have excluded mutations in the entire p53 coding sequence as the aetiology of Cowden syndrome in this family. Other genes such as mdm-2 and WT1, whose products are known to interact with p53, 7 23 would be possible candidate genes.

Ideally, mutation analysis of p53 and other candidate genes should be performed in multiple families with Cowden syndrome. However, Cowden syndrome is rare. Thus, it may be prudent for an international collaborative effort both to define the clinical features of Cowden syndrome and to search for the susceptibility gene.

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