A fertile male with cystic fibrosis: molecular genetic analysis

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Abstract
A family study is presented in which the father of a girl with severe cystic fibrosis (CF) was also found to have CF but was mildly affected. He was diagnosed with three positive sweat tests including one after suppression with fludrocortisone. Genetic analysis showed that he is a compound heterozygote with the AF508 CF mutation associated with haplotype B and a second CF mutation associated with haplotype C. In this unusual, fertile CF male, the late age of diagnosis (30 years) and the mild clinical picture suggest that the compound genotype (AF508/other CF mutation) determines a much less severe form of the disease which might have gone unnoticed in the absence of a severely affected child. The implications of these findings for genetic counselling of families with CF are discussed.

It is widely accepted that, with a few exceptions,1-4 cystic fibrosis (CF) males are infertile. This is thought to be largely because of a generalised increase in viscosity of the body fluids (including sperm) resulting from impaired regulation of chloride excretion/resorption.5,6 As the life expectancy of CF patients is increasing with the application of suitable supportive measures, the fertility issue becomes more relevant. In this report the clinical and molecular genetic features of a fertile male with mild CF are described.

Case report
The proband in this study is the father of a severely affected CF girl (figure). He had a history of sinopathy and mild gastrointestinal alterations without any noticeable effect on his nutritional status and quality of life. There was also reference to painful ejaculation and very thick sperm. Repeated sweat testing (including fludrocortisone suppression on one occasion) allowed the diagnosis of CF to be made (figure).

The results of the family study with CF linked probes PT-3, XV2c, KM19, D9, G2, and H809 and oligonucleotide probes for normal and deleted codon 508 of the CFTR gene (ΔF508)10 are shown in the figure. The severely affected girl is homozygous for ΔF508 and haplotype B (−/+ for probes XV2c and KM19, respectively) and her mildly affected father carries the same mutation/haplotype association.

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on one chromosome and another CF mutation associated with haplotype C (+/−) on the other chromosome. The segregation of ApoCII (AC)_n alleles is consistent with II.1 being the father of III.1. The frequency of the haplotype /−−−+CF/(PT-3, XV2c, CS.7, KM19, cystic fibrosis mutation), which is present in both II.1 and III.1, is 0.0056 in the general Portuguese population, giving a 99.44% probability of paternity.

Discussion
The case described above is apparently identical, as far as haplotypes are concerned, to that of an Iranian CF mother described recently. A molecular epidemiological survey of CF in the Portuguese population indicated that a sizeable proportion (22%) of the CF chromosomes belong to haplotype C, while the typically northern European haplotype B accounts for only 52% of CF chromosomes. Similar haplotype distributions have been found in other southern European populations. The fertile male with mild CF presented here is, therefore, a compound heterozygote with one severe and one mild CF mutation. However, another unrelated patient, homozygous for haplotype C and not carrying the ΔF508 mutation, showed a severe clinical picture from birth and eventually died from acute pulmonary heart disease at the age of 7 years (personal observation). These findings suggest a heterogeneous molecular pathology associated with haplotype C chromosomes in the Portuguese population. Only the complete definition of the molecular basis of CF in these two patients (currently under way) will make it possible to establish a genotype/phenotype correlation.

This study raises several points. (1) The diagnosis of almost asymptomatic CF patients in adulthood dramatically changes their risk of bearing affected children. With a carrier partner, 50% of their children will be affected and with a non-carrier partner all offspring will be CF carriers. (2) If a clear genotype/phenotype correlation is found, the prediction of the clinical course of the disease based on the genotype will enable at risk couples to make a more informed decision regarding their progeny. (3) If there is any indication that a parent of a CF child may have some complaint possibly related to CF, a sweat test should be performed on that parent. (4) When general population screening becomes available, should persons identified as being carriers be investigated further in order to determine whether they may actually be compound heterozygotes for CF?

This report raises the possibility that very mild CF cases may remain undiagnosed for life. It remains to be seen to what extent this is so.

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