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MUTATION ANALYSIS OF GENES INVOLVED IN SPERM MOTILITY: A STUDY IN PATIENTS WITH TOTAL SPERM IMMOTILITY

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Reduced sperm motility represents one of the major male causes of infertility. The axoneme (Ax) is the flagellar motor of the sperm cell and several mutations in genes involved in the assembly and regulation of the Ax have been proved to be responsible for certain cases of infertility associated with severe sperm immotility. For instance, mutations in the genes CCDC39, CCDC40, AKAP3, and AKAP4 (that are involved in the assembly of outer dynein arms) are associated with primary ciliary dyskinesia (PCD). PCD is an inherited autosomal recessive genetic disorder whose typical diagnostic features include the absence of dynein arms and reduced sperm motility. Fibrous Sheath Dysplasia (FSD) is a flagellar pathology, which causes total sperm immotility, mainly due to hyperplasia and disorganization of the Fibrous Sheath (FS). Previous reports suggested that mutations in AKAP3 and AKAP4 genes (the main components of FS) might contribute to FSD. In a group of five Portuguese patients from Assisted Reproductive Medicine centres that presented totally sperm immotility, transmission electron microscopy revealed several structural defects in sperm flagellum, such as anomalies in dynein arms, microtubules and FS. Given the importance of CCDC39, CCDC40, DNAH5, DHA1, AKAP3 and AKAP4 genes in sperm motility, we decided to screen these genes in our patients. To identify genetic alterations that could explain their phenotype, we initiated the analysis of the exonic regions of these 6 genes by Sanger sequencing. We have already sequenced five genes and DNAH5 analysis is still ongoing (we have already sequenced thirty-five exons that are known to harbour a significant number of mutations, from a total of seventy-nine). Ten variants in CCDC39, twenty-six in CCDC40, two in DHA1, seven in AKAP3, one in AKAP4 and thirty-nine in DNAH5 have been identified. The work’s major contribution was the identification of fourteen new variants in CCDC39, CCDC40, AKAP3 and DNAH5 genes. With this work we expect to be able to offer a differential diagnosis to the patients and find potential genetic markers for individuals with this kind of problem.

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PRENATAL DIAGNOSIS MOSAIC 45, X CASE WITH A MARKER CHROMOSOME

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Introduction: Small supernumerary marker chromosomes (sSMC) are structurally abnormal chromosomes that cannot be identified or characterized unambiguously by conventional banding cytogenetics. Generally the size is about or smaller than a chromosome 20, and molecular cytogenetic techniques are necessary for a comprehensive characterization. Prenatally ascertained sSMCs occur in 0.075%, and 0.044% in subsequently studied postnatal analysis. The overall risk of phenotypic abnormalities in prenatally ascertained sSMCs has been estimated to be about 13%. sSMC in Turner syndrome (sSMC) are very rare in the common population (1:100,000) – however, they can be observed with a 45- and even 60-times higher frequency in infertile or intellectual disability patients, respectively. Even though sSMC derive from one of the sex chromosome in >99% of the cases and the majority form ring chromosome. There are also exceptional reports on sSMC derived from one of the autosomes. Thus, a detailed molecular cytogenetic marker chromosome characterization is needed to provide information on sSMC.

Methods: A 29-year-old primigravida woman underwent amniocentesis at 16 weeks of gestation due to a positive maternal biochemistry screening for trissomy 13 and 18, and the presence of a single umbilical artery. Karyotype (GTL-banding), aCGH (4x180k Agilent Human CGH Microarray) and FISH analysis (CEPX Spectrum Green) were sequentially performed in cultured amniocytes to better characterize this sSMC.

Results: Amniocentesis revealed a karyotype 45,X[22]/46,X,+mar[8]. Among 30 cultured amniocytes colonies, 8 contained the sSMC, whereas the remaining 22 colonies were 45,X. Using DNA extracted from cultured amniocytes, aCGH showed that the sSMC was originated from chromosome X and revealed a 19,81-Mb gene dosage increase at Xp11.21-Xq21.1. FISH analysis showed 41% (41/100) mosaicism for sSMC in cultured amniocytes and confirmed the identification of the sSMC as derivate from chromosome X.

Discussion: Althought sSMC are rare, this is a well-known cytogenetic entity. So far, a detailed molecular cytogenetic characterization of sSMC by aCGH was only performed in a few cases. In this case, we could conclude that sSMC was derived from Xp11.21-Xq21.1 probably as a ring [r(X)p11.21q21.1]] due to absence of telomeric regions.