

Errata

No número anterior da revista (volume 45, no. 6, Dezembro de 2001), por uma falha no processo de composição final do artigo: “**Cytogenetic Analysis and Detection of *KAL-1* Gene Deletion with Fluorescence *in situ* Hybridization (FISH) in Patients with Kallmann Syndrome**”, de autoria de Ericka B. Trar-

bach, Maria T.M. Baptista, Andréa T. Maciel-Guerra & Christine Hackel, a figura 2 (página 555) foi omitida, sendo impressa, em seu lugar, novamente a figura 1.

Abaixo está reproduzida a figura 2 correta. Os Editores lamentam a falha e pedem desculpas aos autores pelo ocorrido.

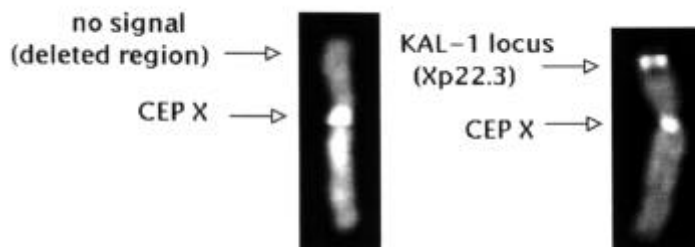


Figura 2. Hybridization of LSI-Kallmann/CEPX probes on human metaphase chromosomes. The LSI Kallmann probe hybridizes to the band Xp22.3 (red signal) and the CEPX to the centromere of the human X chromosome (green signal). In the left image (patient KS-5), only the centromeric region signal is visualized. At right a normal X-chromosome is shown where the marks of the KAL-1 locus and centromeric region can be seen.

Note: Pseudocoloring was performed for image reproduction in black and white.