

Open Peer Review on Qeios

Paternal uniparental disomy of chromosome 13

INSERM

Source

INSERM. (1999). Orphanet: an online rare disease and orphan drug data base. <u>Paternal uniparental disomy of chromosome 13</u>. ORPHA:99324

Paternal uniparental disomy of chromosome 13 is an uniparental disomy of paternal origin that most likely does not have any phenotypic expression except from cases of homozygosity for a recessive disease mutation for which only father is a carrier.

Qeios ID: A41RM4 · https://doi.org/10.32388/A41RM4