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fAP1

IDENTIFICATION

Causal gene(s)	APT _X
Repeat size or mutation	pW279X/pW279X; sibling of fAP2
Cell type	Fibroblast

SOURCE & PUBLICATIONS

Links to publications or public resources	The ataxia-oculomotor apraxia 1 gene product has a role distinct from ATM and interacts with the DNA strand break repair proteins XRCC1 and XRCC4 - PubMed https://pubmed.ncbi.nlm.nih.gov/15380105/
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