

[Cell Lines](#) [MSA/OPCA, PD](#) [Publication](#)

CRICKi-12-A

IDENTIFICATION

Research Resource Identifier (RRID)	CVCL_C5T4
Causal gene(s)	SNCA
Repeat size or mutation	c.G152A mutation in exon 3 of SNCA
Cell type	hiPSC

DONOR INFORMATION

Donor gender	Female
Age at collection (years)	57

SOURCE & PUBLICATIONS

Originating lab / institution	The Francis Crick Institute
Links to publications or public resources	Generation of TWO G51D SNCA missense mutation iPSC lines (CRICKi011-A, CRICKi012-A) from two individuals at risk of Parkinson's disease - PubMed https://pubmed.ncbi.nlm.nih.gov/37336145/ CRICKi012-A u00b7 Cell Line u00b7 hPSCreg https://hpscereg.eu/cell-line/CRICKi012-A