



Cover image: 'Catching waves' to diagnose Parkinson's disease? In the presence of pathological blood-derived α -synuclein conformers an amplification assay displays typical waves. From Kluge *et al.* Detection of neuron-derived pathological α -synuclein in blood. Pp. 3058–71. This cover was designed using an image from Freepik.com. https://de.freepik.com/fotos-kostenlos/surfer-faengt-die-grosse-welle_13006291.htm Photo created by wirestock.

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