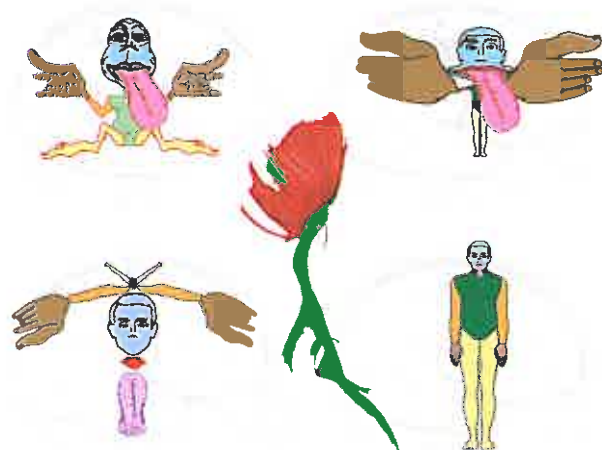


BRAIN

A JOURNAL OF NEUROLOGY



Cover image: The seminal contribution of Penfield and Boldrey published in *Brain* (1937) is revisited in light of current neuroimaging research. From Catani. A little man of some importance. Pp. 3055–3061.

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