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Nerve fibre degeneration in the brain in amyotrophic lateral sclerosis

Martin R Turner

Editor’s Choice

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Author: Smith MC
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Number of times cited: 95

An advertisement in The New Scientist dated 9 May 1957 read: “Technician or junior technician required for histology in neuropathological work. Some knowledge of photography an advantage. Salary according to experience. Apply in writing stating age and experience to Dr Marion Smith, National Hospital for Nervous Diseases, Queen Square, London WC1.” Three years later the Glasgow trained neuropathologist (1910–1998) and President of the British Neuropathological Society published a paper entitled ‘Nerve fibre degeneration in the brain in amyotrophic lateral sclerosis’, based on detailed postmortem examination of seven cases.

Cerebral involvement in amyotrophic lateral sclerosis (ALS) was already recognised. Seventy years prior, Gowers noted that ‘tractography’, a white matter pathway study that can now be performed in vivo using MRI. Indeed, her published drawings of prominent and consistent corpus callosum as well as corticospinal tract involvement in ALS are strikingly similar to images obtained non-invasively 50 years later using diffusion tensor imaging.2 Her observations on the organisation of internal capsule fibres according to their cortical projections is also reproducible using MRI based tractography.3

Smith acknowledged a key limitation of this type of analysis, persistent to this day, namely that: “…we cannot say categorically that the degeneration starts here and goes there, as we can about the degeneration resulting from circumscribed lesions”. Smith’s study included a long survivor (12 years) in whom there is reported: “…little active degeneration in the cord or lower brain-stem, although there is in the cerebral hemispheres”. It is now recognised that upper motor neuron predominant ALS, as well as the much rarer exclusively upper motor neuron phenotype termed primary lateral sclerosis, may also have significantly slower progression. Her observation of degenerating fibre tracts projecting to the thalamus is in keeping with the in vivo demonstration of particularly prominent microglial activation here.2 She also noted involvement of the basal ganglia, notably including the substantia nigra, in three of the seven brains, in keeping with the recognised clinical overlap of parkinsonism.5

Smith co-authored other important anatomical papers with the Queen Square neurologist Peter Nathan (1914–2002), including a study of the course of the corticospinal tracts and an examination of the anatomical substrate for the Babinski reflex. Prescient of the era of TDP43-defined ALS, she commented at the end of this paper that: “It is possible that essential information relevant to the functional anatomy of the human central nervous system may be derived from the detailed study of routine neuropathological material, apart from the possibility of making observations contributing to an understanding of the aetiology of the disease.”


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