Subacute sensory neuronopathy and cancer: the identification of paraneoplastic syndromes

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In this landmark paper, Denny-Brown published cases of severe sensory ataxia found at autopsy to have sensory ganglionopathy in association with bronchogenic carcinoma

The concept of paraneoplastic disorders is relatively recent in the clinical neurosciences, and one that remains incompletely understood to this day, particularly in terms of therapy. By means of definition, paraneoplastic disorders have tended to encompass all remote effects of cancer, not attributable to metastases, treatment or the presence of coexistent illness.1 In terms of disease pathophysiology, the cause of paraneoplastic disorders has been linked to a form of immunological cross-reactivity between tumour antigens and neuronal tissue, thereby triggering an immune-based attack on the nervous system.

Early concepts of paraneoplastic phenomenon slowly emerged through the study of neuropathy. Until 60 years ago, the literature on neuropathy itself was relatively sparse. Most neuropathies were described as sensory and motor, starting distally with proximal progression. As Lecturer Dr Derek Ernest Denny-Brown (then later Professor, OBE, MD, FRCP) pointed out in this landmark JNNP paper, sensory neuropathies were generally considered rare and were usually attributed to infection such as tabes or a postinfectious process like Guillain-Barré syndrome.2 Denny-Brown’s paper that launched a concept for paraneoplastic phenomena began by describing two cases (one a boiler cleaner and the other a retired Army Colonel) n which both had initially presented with rapidly progressive, painful, sensory ataxic syndromes. On neurological examination, both patients displayed ataxia, severely reduced sensation and reduced or absent reflexes. Both died within 8 months. Subsequent autopsy identified a severe loss of nerve cells in the dorsal root ganglia.3 Further clinical research has more precisely identified the cause of this specific syndrome to circulating auto-antibodies (anti-Hu) directed against intracellular antigens present in the dorsal root ganglion cells.4 Further clinical research has greatly expanded our understanding of these paraneoplastic syndromes more generally, with the unifying feature linked to an immune-mediated pathogenesis.4 Distinct antigens have led to classifications based around clinical phenotypes and associated autoantibody production and also through the location of the antigen, being either intracellular or extracellular.5 6 In addition, there can be overlap with similar syndromes which may develop in the absence of an identifiable cancer. In those syndromes with autoantibodies that target intracellular antigens, cancer is almost always present.

While there has been improved understanding in disease pathophysiology, treatment approaches still often remain uncertain. Some disorders such as the Lambert-Eaton myasthenic syndrome and myasthenia gravis may respond well to immunosuppression and to treatment of the underlying tumour. In contrast, encephalomyelitis associated with cancer and paraneoplastic cerebellar degeneration usually respond poorly to treatment. Differences in cellular and humoral immunity, as well as the nature of autoantibody target antigens such as receptors and ion channels, likely underlie these differential responses to therapy.7 While the field of paraneoplastic neurology has certainly advanced since the original JNNP landmark study, as Denny-Brown wisely concluded then, more studies on similar cases in the future are indicated to better unlock the precision therapies that will no doubt be required.8

Primary sensory neuropathy with muscular changes associated with carcinoma

Authors: Denny-Brown D
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