Alfons Maria Jakob (1884–1931), Neuropathologist par Excellence

Scientific Endeavors in Europe and the Americas

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Abstract
The study briefly reviews the life and work of Alfons Maria Jakob (1884–1931), a notable representative of pre-war German neuropathology. Today Jakob is mainly remembered by neurologists for the spongiform encephalopathy with progressive dementia and spasticity that he, and Kiel neuropathologist Hans Gerhard Creutzfeldt (1885–1964), described independently. Jakob has left additional contributions to neuroanatomy, neuropathology and neuropsychiatry in the form of original articles and valuable monographs.

First Publications and Academic Career


On December 1, 1909 Jakob became an assistant physician at the psychiatric clinic directed by Emil Kraepelin (1856–1926) in München, where he trained in neuroanatomy with Alois Alzheimer (1864–1915) through May 1, 1911 [1].

Solicited by professor Wilhelm Weygandt (1870–1939), he moved in November 1911 to the psychiatric clinic of the Friedrichsberg State Hospital in Hamburg, where he eventually succeeded Theodor Kaes (1852–1913) as director (Prosektor) of the neuroanatomical lab-
oratory. From the summer of 1915 until the end of World War I, Jakob served as a medical officer at the Belgian front. He was habilitated in neurology and psychiatry at Hamburg University in 1919 and became professor in 1924. He expanded the laboratory with serology, genetics, and experimental psychology sectors, and trained dozens of students from the United States, Russia, Japan, Turkey, Italy, Switzerland, Spain, Portugal and Uruguay [1]. He also kept a reputable medical practice. His brother Franz, also a physician, practiced in Nürnberg [6].

**Creutzfeldt-Jakob Disease**

At the 10th annual meeting of the Society of German Neurologists, held in Leipzig on September 18, 1920, O.B. Meyer presiding, von Weizsäcker read Jakob’s report describing a clinicopathological syndrome in 3 patients (2 female, 51 and 34 years old, and 1 male, 43 years old) with spasticity and progressive dementia associated with cortical, striatal and spinal degeneration [7]. Jakob [8] expanded on those 3 cases in an additional paper. In 1920, Kiel neuropathologist Hans Gerhard Creutzfeldt (1885–1964) had independently published a similar case of a 22-year-old woman [9]. The text that follows gives an overview of what Jakob and additionally Creutzfeldt really described, compared to our current knowledge on spongiform encephalopathies. In-depth accounts have been given by Duckett and Stern [10], Poser and Bruyn [11], and Wolf and Foley [12].

‘Creutzfeldt-Jakob disease’ (CJD) was so named by Spielmeyer [13] and subsequently included amongst the organic psychoses in Bumke’s 1928 *Handbook of Mental Diseases* [12]. The CJD concept is closer to a neuropathological syndrome rather than a single etiologic-nosological entity [10, 14]. Some 84 forms, ‘types’ or variants of CJD have been recorded on the basis of clinical and neuropathological criteria [15].

Today, CJD is classified as a prion disease (‘prionosos’) or transmissible spongiform encephalopathy (TSE), characterized by dementia, pyramidal, and extrapyramidal signs clinically, and by neuronal loss, spongiform changes, and astroglial reaction histologically [11, 12].

Scientific interest in CJD became renewed in 1957 with the description of kuru, a contagious disorder in natives of New Guinea [16] and its similarities to scrapie, a transmissible infectious disease in sheep [17, 18]; the concept of prion diseases was formulated 25 years later [19]. Animal forms of TSE, besides scrapie, include bovine spongiform encephalopathy (‘mad cow disease’) and mink encephalopathy [11].

In all, Creutzfeldt studied 1 case [9, 20], and Jakob 7 cases, including the initial 3 cases [7, 8], case 4 [21], and case 5 described in his book on extrapyramidal disorders [22]. The brains of Jakob’s patients 6 and 7 were further studied and published posthumously by his pupil Kirschbaum [23], who in his monograph reviewed 150 cases of CJD gathered over a 50-year period, including Creutzfeldt’s and Jakob’s original cases. English translations of Creutzfeldt’s first paper [9] and of Jakob’s report on case 4 [21] may be found in Rottenberg and Hochberg [24].

Creutzfeldt’s case concerned a patient seen in 1913 at Alzheimer’s clinic in Breslau. Because of World War I and Alzheimer’s death, Creutzfeldt finalized his manuscript [9] after his 1919 move to Spielmeyer’s department in Munich. Creutzfeldt published a sequel paper on the case [20].
Both articles were entitled 'On a particular focal illness of the central nervous system', conveying the possibility of having identified a new entity of unknown etiology [10].

The patient first presented with gait ataxia and developmental behavioral abnormalities when she was 16 [12]. The disease progressed irregularly, with remissions. The later clinical picture included pyramidal signs (paresis, spasticity and bilateral Babinski), hyperalgesia, intention tremor, facial hyperkinesia, nystagmus, myoclonus, dementia, mood changes and mutism [11]. Erythema multiforme bullousum, appearing as herpetiform small vesicles, made its appearance in the vicinity of the third ramus of the left trigeminal nerve. Stupor deepened, swallowing became impaired, and death ensued in status epilepticus [9, 10]. Two of the patient’s five siblings had mental deficits and her mother had died of an unknown cause at the age of 56.

The neuropathological finds were focal neuron loss and neuronophagia in the postcentral and precentral gyri (particularly affecting the pyramidal cells of the third and deeper layers), bilateral degeneration of the corticospinal tracts, vascular reaction and astroglial hypertrophy also present in the basal ganglia, thalamus, cerebellum, brainstem and spinal cord [10]. Status spongiosus was not reported [11]. The diagnosis was an acquired polioencephalopathy. Creutzfeldt’s case may actually be the first report of herpes zoster encephalopathy [10].

Jakob’s first 5 cases [7, 8, 21, 22] and Creutzfeldt’s case [9, 20] shared, in Jakob’s opinion, a common neuropathological picture, a polioencephalopathy with lesions in the frontal lobe (deep cortical layers in particular) and changes in the rest of the cortex, striatum, thalamus (medial, ventral and lateral nuclei), substantia nigra, pontine tegmentum, cerebellum, brainstem and spinal cord. Jakob did not regard his first 4 cases as a homogenous group [10] and thought that the bulk of the pathological changes fell on the extrapyramidal system [25]. Diagnosis in cases 1 and 2 was syphilis, in case 3 malaria, and in case 4 chronic alcoholism [7, 8, 21]. In all likelihood, the diagnosis of CJD was correct in Jakob’s case 3, the patient with progressive dementia, leg weakness and pain, ataxia, vertigo and diplopia [7, 8], and in case 5 [22], with diffuse vacuolation of the neuropil throughout cerebral cortical areas and the cerebellar molecular layer, typically seen in TSEs [11]. It is also interesting to note that the sister and maternal grandmother, along with 8 of her siblings, of Jakob’s patient 6, a 44-year-old male, had died of an undeciphered nervous disorder; the brains of patient 6, his sister, and two of his children contained spongiform changes [11].

Jakob gave the disorder a neuropathological rather than a clinical name, ‘spastic pseudosclerosis encephalopathy with disseminated foci of degeneration’ [10]. He designated his subgroup of pseudosclerosis ‘spastic’, owing to the marked corticospinal degeneration [11, 12, 26]. Having access to Creutzfeldt’s slides and the galleys of Creutzfeldt’s chapter [20], Jakob considered Creutzfeldt’s case as another example of spastic pseudosclerosis. However, Creutzfeldt objected to the term ‘spastic pseudosclerosis’ [11], preferring instead the general description ‘progressive focal and diffuse degeneration of the gray matter’ [10] and provisionally leaving the disorder without a specific clinical name [12].

The term ‘pseudosclerosis’ had been introduced in 1883 by the Berlin neuropsychiatrist Carl Westphal (1833–1890) [27] to describe what would later be identified as a juvenile form (Westphal variant) of Huntington chorea [28]. The eponym Westphal-Strümpell pseudosclerosis was also adopted in the German literature, based on additional cases reported by the Erlangen internist Adolf von Strümpell (1853–1925) in children [29].

A ‘singular’ case of pseudosclerosis with tonic rigidity and high-grade dementia, but without Babinski sign, sensory disturbances or thalamic lesions [30], marked histopathologically by interstitial hepatitis, gliosis in the striatum and globus pallidus, substantia innominata and cerebellar molecular layer, was reported by Economo and Schilder [31] in a 55-year-old man, before Creutzfeldt’s first study [9] in the same journal. Although at some point considered to bear similarities with the classical descriptions of CJD [32], that case was probably Wilson disease [33]. Economo and Schilder [31] had inferred that their case affected the extrapyramidal system, with a distant relationship to paralysis agitans and olivopontocerebellar atrophy. Jakob [7] did observe interesting similarities between his first 3 cases and that of Economo and Schilder [31], but pinpointed that the latter was accompanied by liver damage, and thus found an agreement of his cases in all the substantial points only with the case of Creutzfeldt [9].

The question has been raised whether the disease that Creutzfeldt and Jakob independently described is the same, and the validity of Creutzfeldt’s case report [9] as an instance of ‘classical’ CJD has been disputed [11, 12] on grounds of the patient’s young age, the developmental symptoms, and the absence of extrapyramidal signs. Kirschbaum [23] and Katscher [34] favor ‘Jakob-Creutzfeldt disease’ (or ‘Jakob’s syndrome’ [23]) over CJD, crediting Jakob as the major contributor to CJD, and a majority of authors currently consider Jakob’s cases as the first
true descriptions of CJD [10]. Proof that Jakob’s later patients were true cases of CJD came with the identification of a PRNP gene mutation [35].

Thus, one may conceivably discern a connection between Jakob’s ‘syndrome’ and the seeds for the eventual bestowal of two of the Swedish Academy’s summa cum laude Prizes in Physiology or Medicine: to D. Carleton Gajdusek (and Baruch S. Blumberg) in 1976 ‘for their discoveries concerning new mechanisms for the origin and dissemination of infectious diseases’, and to Stanley B. Prusiner in 1997 ‘for his discovery of prions, a new biological principle of infection’.

**Travels to the Americas**

On 27 March 1924, Jakob embarked on the maiden voyage of the Hamburg–American ocean liner Deutschland from Southampton, arriving in New York on April 6 [6]. He guest lectured at several institutions, including Columbia University [14].

Four years later, he journeyed to Latin America. In May through July 1928, he gave a 20-lecture course in Rio de Janeiro on nervous and mental pathology (fig. 2), with a theoretical and a practical part, using 4,000 microscopic preparations and transparencies [36]. He then travelled to São Paulo and Campinas, Brazil; Buenos Aires, where he met with physiologist Bernardo Alberto Houssay (1887–1971) and visited the neuroanatomical institute of his fellow countryman, Bavarian neuropathologist Christfried Jakob (1866–1956) [37, 38]; over the imposing Cordillera to Santiago de Chile and Valparaiso; and back, over the Andes, to Montevideo, Uruguay. Jakob gave a presentation of his South American impressions to the Hamburg Medical Association on December 11, 1928 [39].

**Authored Works**

Jakob published 80 articles on diverse topics [1–3], including cerebellar tumors (1910), trauma and secondary fiber degeneration (1912), cerebellar ataxia [40], multiple sclerosis (1913), epilepsy (1914), diffuse infiltrating encephalomyelitis (1914), spinal cord concussion (1919), endarteritis syphilitica (1920), paralysis and tabes (1922), megalencephaly (1925), miliary gummata [41], and yellow fever [42], the latter co-authored with Amadeu Fialho and Eudoro Libanio Villela and presented at the 19th annual meeting of the Society of German Neurologists on 20 September 1929 in Würzburg, chaired by Otfrid Förster (1873–1941). He published a review in Spanish on anatomo-psychiatric correlations [43] and one in Portuguese on multiple sclerosis [44]. Jakob was one of the proponents of the value of histopathology for elucidating the research problems of psychiatric diseases [25, 45].

Jakob wrote the cerebellar chapter for Mollendorff’s 1928 Handbook of Microscopic Anatomy, and the neurosyphilis chapter for Bumke’s 1930 Handbook of Mental Diseases [2, 3]. He authored a monograph on extrapyramidal disorders [22], and contributed two scholarly volumes to Aschaffenburg’s Handbook of Psychiatry, on normal and pathological neuroanatomy and neurohistology [46] and on special cerebral histopathology [47].

Jakob reviewed the Cytoarchitectonics of the Adult Human Cerebral Cortex of Economo and Koskinas [48] and the Icones Neurologicae of Strümpell and C. Jakob [49]. Strümpell and C. Jakob had produced the original epitome of Icones Neurologicae in 1897 [50]; the plates were...
re-edited by Müller and Spatz in 1926 with 13 folded plates, 11 of them 106 × 140 cm in size and two 140 × 212 cm [51].

Jakob called the Economo and Koskinas Cytoarchitectonics ‘a masterpiece unique in the international medical literature’, its 112 plates ‘brilliant achievements in scientific microphotography’, and its text a ‘joy to the reader’ [48]. Jakob found the nomenclature of Economo and Koskinas ‘fully meaningful’ in its departure from the methods of Vogt and Brodmann. In his own textbook on Cerebral Anatomy and Histology, Jakob [46] devoted two-thirds of the chapter on the architectonic organization of cortical fields to the findings of Economo and Koskinas and reproduced 23 of their original figures, arguing that ‘at last we have acquired a perfect and complete map of the cerebral cortex by areas’ [48].

Jakob is described as a ‘private and disciplined individual, excellent and knowledgeable teacher and charismatic leader’ [10]. Besides his own publications, Jakob conceived, and in part dictated, an estimated further 80 works by his students, in which he does not appear as a co-author [1].

**Alpers Disease**

At the third session of the 20th annual meeting of the Society of German Neurologists in Dresden on 20 September 1930, H. Curschmann presiding, Jakob presented a yet another new entity [52], studied in conjunction with his pupils Somoza, Freedom [53] and Alpers [54]. ‘Alpers disease’ or progressive infantile poliodystrophy (also called spongy glio-neuronal dystrophy [55]) is a mitochondrial disorder with autosomal recessive inheritance appearing in early infancy [56–58], leading to marked dementia, prominent seizures, spasticity and opisthotonus, and accompanied by liver failure [59, 60]. The histological hallmarks include almost total neuronal loss in the cortical gray matter with spongy changes and astrogliosis, and the molecular genetic defect is associated with mutations in the polymerase-γ (POLG) gene of the mitochondrial DNA (mtDNA) [61–63].

**Postscript**

Jakob gave his ‘swan-song’ presentation, on the nosology and localization of torsion dystonia with cinematographic and anatomical demonstrations, at the First International Neurological Congress in Berne [64], during the clinical-pathogenetic section of the afternoon of 3 September 1931, chaired by Gheorghe Marinescu (1863–1938).

Shortly after the Berne Congress, neurology lost two of its protagonists: Alfons Jakob on October 17, 1931 in Hamburg, at the age of 47, after an operation to contain complications of streptococcal osteomyelitis, from which he had been suffering for 7 years; and, within 4 days, Constantin von Economo in Vienna, at the age of 55. Three years later to the day, on October 17, 1934, the neuroscience world would lose yet another of its greatest, the venerated Santiago Ramón y Cajal.

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