A historical look using virtual microscopy: the first case report of adrenomyeloneuropathy (AMN)

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Abstract

The history of adrenoleukodystrophy (ALD), adrenomyeloneuropathy (AMN) and other peroxisomal diseases is exemplary for the stunning progress of scientific medicine within the past 50 years. Like many breakthroughs in medicine, the detailed analysis of patients’ pathologically affected tissues was instrumental, resulting in stepwise systematic clarification of what had remained enigmatic until the 1970s. This flashback paper is a recollection of the first neuropathological description of a slowly evolving clinical phenotype, spastic paraparesis with adrenal insufficiency, in a young adult by Budka et al. 1976 [3], using virtual microscopy of the original histologic slides. The clinicopathological presentation derives from the classical cerebral ALD phenotype in boys, where electron microscopy demonstrated the underlying pathological hallmark of characteristic lipid inclusions shared by both phenotypes. Our report allowed the delineation of a new disease type almost simultaneously described in more cases as AMN by Griffin et al. 1977 [4] and Schaumburg et al. 1977 [11]. Moreover, our report indicated clinical heterogeneity in the ALD disease group that, as shown later, extends further to females, to Addison-only, and even to asymptomatic subjects. The gene underlying ALD was discovered in 1993 as a defect in the ABCD1 gene. Yet, it has hitherto remained unclear how the gene defect causes the strikingly broad and unpredictable phenotypic spectrum of ALD/AMN.

Keywords: Adrenoleukodystrophy (ALD), Adrenomyeloneuropathy (AMN), Peroxisomal diseases, Peroxisomes, Neuropathology, History
It’s not what you look at that matters, it’s what you see.
Henry David Thoreau (1817-1862) [16]

The history of what is now established as adrenoleukodystrophy (ALD), adrenomyeloneuropathy (AMN) and other peroxisomal diseases is exemplary for the stunning progress of scientific medicine/medical science within the past 50 years. This journey starts with the mere description of clinico-pathological phenotypes, follows up with the identification of a biochemical abnormality, and culminates by defining the molecular and genetic basis of the disease, by newborn screening, and by promising therapies, as recently reviewed [17]. Instrumental, like in many breakthroughs in medicine, was the detailed analysis of patients’ pathologically affected tissues, resulting in step-wise systematic clarification of what had remained an obscure enigma until the 1970s. Moreover, the history of ALD and AMN is exemplary for the multidisciplinary and international character of contributing studies as a prerequisite for success. This article recollects an early step in our knowledge of the disease group where neuropathology had a pivotal role.

Local background of the first case of AMN

The present (hi)story evolved more than 50 years ago, after I became a trainee at the renowned Neurological Institute, or Obersteiner Institute, in Vienna from late 1971, right after my graduation as an MD. Kurt Jellinger (born 1931) was there in charge of clinical neuropathology, made the preliminary neuropathological autopsy report of an interesting patient who died in late 1972, and suggested me to write up the story. However, I was extremely busy in learning all elements of classical neuropathology and doing clinical training in parallel to become specialist in neurology and psychiatry. Moreover, I was not yet aware of the paramount importance of publishing in science. So I failed to give priority to this task. In addition, my early years were increasingly overshadowed by Kurt’s leave from the Institute in 1975, a severe blow to diagnostic neuropathology and a disaster for me, as I was suddenly expected to bear most of the diagnostic workload myself; I gave a glimpse on those difficult early years elsewhere [2].

So it was May 1976 when the article was finally submitted, and was printed in the September issue of the Journal of Neurology (formerly: Deutsche Zeitschrift für Nervenheilkunde and Zeitschrift für Neurologie) [3] that then had a mixture of German and English-written articles. My co-authors were Elfriede Sluga (1930-2008, Fig. 1) and Wolf-Dieter Heiss (born 1939, Fig. 2), both much more senior than me (Fig. 3).

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**Figure 1.** Elfriede Sluga 1972
(Source H. Budka)

**Figure 2.** Wolf-Dieter Heiss
(Source European Academy of Neurology)

**Figure 3.** Herbert Budka 1972
(Source H. Budka)
Elfriede ("Fritzi") had pioneered the diagnostic use of the electron microscope (EM) at the Neurological Institute, mostly for neuromuscular biopsies, had a dominant clinical interest and later became professor and director of a department of neurology at a municipal hospital in Vienna, in Kurt Jellinger’s footsteps who had done the same earlier. By then, I never had a chance to touch the EM, so Fritzi contributed the essential EM diagnostics of the report. Wolf-Dieter was then a consultant neurologist at the Neurological University Clinic in Vienna, responsible for a ward of some 20 hospital beds where the patient was admitted, and an impressive teacher during my clinical training. He oversaw clinical details of the report and later became professor at the University of Cologne and director of the Max Planck Institute for Brain Research in Köln-Merheim, and is now widely recognized as a top expert on cerebral hemodynamics, including a pioneering role in the establishment of stroke units. Wolf-Dieter, a giant in clinical neurosciences in his own right, originally went to Cologne to succeed another giant, Klaus-Joachim Zülch (1910-1988) who had combined clinical neurology with neuropathology as neuro-oncologist and stroke expert; he is now remembered as main author of the very first edition of the famous WHO Blue Book Series on Histological Typing of Tumours of the CNS [18].

For the Neurological Institute in Vienna, our report laid the foundation for a new research and diagnostic area that became important not only for the Institute’s neuropathology branch, but still more so for its Division of Neurochemistry, led by Hans ("Hanno") Bernheimer (1930-2016), and mainly involving Brunhilde Molzer (born 1940) [7] with very long chain fatty acids (VLCFA) diagnostic biochemistry for ALD/AMN suspects. In the 1990s Johannes Berger (born 1964) joined the neurochemistry group and expanded its molecular and experimental studies; from 2000, he has done research on peroxisomes as Professor in the newly founded Center of Brain Research in Vienna.

The patient

Our propositus was a relatively healthy – well, not quite – man who succumbed quickly and rather unexpectedly in 1972 at the age of 24, after a slowly evolving disease with spastic paraparesis over 2 years. No definite diagnosis was made during lifetime.
You can make yourself familiar with the neuropathology at autopsy by viewing scanned slides of the right adrenal gland (Fig. 4) and of the medulla oblongata (Figs. 5 & 6). Indeed, I strongly recommend you to stop here and do your virtual microscoping before you continue to read more details. There are of course different types of observers, but my favorite personal style has always been the detective-style approach, starting with a few basic data (sufficient here in the preceding paragraph), and aiming to have a more complete story evolving from the discoveries during microscoping (and earlier autopsy with brain cutting). If you read the more detailed description already at start, you might easily and quickly look at everything, but it may happen that you do not SEE in Thoreau’s sense (see citation at the start of this article). Moreover, the suggested early microscoping reflects the usual neuropathologist’s way of working: ultimately, you learn most from what you initially might NOT see.

Recapitulation of the case report

In our report, we made a detailed description of the clinical course, laboratory findings, and results of a complete autopsy including histology and electron microscopy of various organs. There is nothing new that can be added since then. So an excerpt follows for those who are unable or unwilling to read the original paper.

Our propositus had a family history of an unusual dark skin pigmentation of his mother and was hyperpigmented since childhood. Originally, this was an active young man who used to make bicycle tours. At age 22, progressive weakness was noted in both legs. Subsequently, bladder and anal sphincter dysfunction developed; sexual functions were unimpaired. Seven months before death, he presented at the Neurologic Clinic in Vienna with a deeply bronzed skin, alopecia including axillary and pubic hairs, and a spastic gait with bilaterally exaggerated tendon reflexes of legs, bilateral Babinski’s sign, and loss of abdominal reflexes. There was hypesthesia of both legs for all qualities, with distal accentuation. When Wolf-Dieter Heiss first saw the patient, he immediately suspected Addison’s disease, because of the bronzed skin. However, extensive involvement of endocrinology specialists did not result in such a clear-cut diagnosis during the lifetime of the patient. Thereafter, laboratory results as detailed in our report were interpreted as compatible with primary adrenal insufficiency.

After a visit to a sauna bath 2 weeks before death, the patient deteriorated rapidly, with collapse, vomiting, vertigo, and impairment of consciousness alternating between agitation and sopor. One week before death, hyperthermia and progressive metabolic and circulatory dysregulation developed. Death was caused by cardiovascular failure.

A complete autopsy at the Institute of Pathology in Vienna was performed by the then famous Prof. Lothar (“Rex”) Kucsko (1912-1976) who contributed also otherwise to my career [2]. He described the adrenals as atrophic, firm, chocolate-brown and without discernible structure, but including two small nodules on the right side; his final diagnosis was so-called immune adrenalitis, with an acute Addison crisis as cause of death. He was perfectly right on the latter, less so on the former. Histology of various other organs was characterized by widespread scattered lymphocytic infiltrates.

Under the microscope: what you may see

Fig. 4 Adrenal (HE). The organ structure is disorganized, with intact medulla (bluish stained tissue) but only focal remnants of cortical tissue (reddish stained tissue) with large, sometimes bizarre and ballooned cells with occasional cytoplasmic “stripes”. In addition, there is prominent fibrosis and focal lymphocytic infiltration.

Fig. 5 & 6 Medulla oblongata (LFB, HE). There is symmetric degeneration of pyramids, to a lesser degree of medial and lateral lemnisci, spinocerebellar and Goll’s tracts. In addition to perivascular lymphocytic infiltrates, there are perivascular lymphoid cuffs of histiocytes with epithelialoid appearance and occasional cytoplasmic “striation” (I am not sure how obvious the striation is on 2-D virtual files that don’t allow a 3-D impression, in contrast to old-fashion microscoping with a glass slide when you can play with the focusing screw).
Medico-scientific background

For decades, what is now called ALD has been a rare and poorly understood, but rapidly evolving and fatal demyelinating disease of male kids, a combination of inflammatory diffuse sclerosis of the brain with adrenal insufficiency. Haberfeld and Spieler have been credited with the first clinicopathological description of classical childhood ALD [5], followed by articles by Schilder who gave his name to what he called encephalitis periaxialis diffusa [13][14][15].

In the 1970s, most research on ALD was concentrated at the Albert Einstein College of Medicine in Bronx, New York. The Department of Pathology there was a hotbed of neuropathology, with eminent researchers like Robert D. Terry and Henryk M. Wisniewski who contributed landmark studies in Alzheimer disease, and John W. Prineas and Cedric Raine who did the same in multiple sclerosis. A young trainee, James M. Powers, teamed up with Herbert H. Schaumburg, an expert in neurotoxicology and peripheral nerves, Ann Johnson and Kunihiko Suzuki, to study a disease that just recently had been termed ALD [1]; their work over the next few years elucidated most of its (neuro)pathology and biochemistry. Powers and Schaumburg demonstrated characteristic lipid inclusions in ALD adrenals by EM examination [9], followed by their description in brain [12] and other organs [10]. Subsequent biochemical studies showed a striking excess of VLCFA in brain cholesterol esters as substrate of ALD 

ALDP/ABCD1 gene [8] encoding for a peroxisomal membrane protein and member of the ATP-binding cassette (ABC) transporter family. On top of that, Kennedy-Krieger contributed much more that is by now known in peroxisomal diseases. Moreover, they had a major role in development and assessment of therapies such as “Lorenzo’s Oil” of Hollywood fame, bone marrow and hematopoietic stem cell transplantation, and gene therapy [17]. Still more impressively, Hugo and Ann Moser established an absolutely exemplary relation to their ALD/AMN patients and their families who returned their love and affection to these two outstanding advocates for persons suffering from neurological disabilities.

AMN (neuro)pathology

The (neuro)pathology of AMN is typical but not specific without appropriate clinical information including family history, VLCFA biochemistry and/or EM findings. Nowadays genetic testing for the ALDP/ABCD1 gene is the mainstay of definite diagnostics. Neuropathology, as seen in the virtual slides, is characterized by two hallmarks: first, degeneration of long tracts of brainstem and spinal cord in a pseudosystemic distribution, and second, the presence within degenerating long tracts of perivascular sleeves of lipid-filled histiocytes that rarely may, or may not, give a “striated” impression. This striation might be the light microscopical appearance of the circumscribed packaging of lamellar lipid aggregates with clear clefts as seen by EM. In parallel, such degenerative changes may also be present in the peripheral nervous system. The scattered lymphocytic infiltrates are to be interpreted as secondary to the adrenal insufficiency that may, or may not, be manifest in an otherwise typical AMN. The pathology of the adrenal glands reflects the degree of adrenal insufficiency, with storage of lipids in enlarged cortical cells.

Originally, in addition to histology, EM examination was important to document peculiar lipid inclusions with a lamellar “leaflet” structure in perivascular histiocytes of the oblongata pyramids, as described and illustrated on two pages of our report, and found earlier as characteristic of ALD by Powers and Schaumburg [9] and Schaumburg et al. [12]. We
suggested the ultrastructural inclusions in the CNS to derive from degenerated myelin in the affected tracts, linking ALD/AMN to pathological storage of a myelin degradation product [3].

Significance

Our study [3] allowed the delineation of what became a new disease type, a little bit later termed AMN [4][11]. The presentation deviated from the classical cerebral ALD phenotype in boys. However, histology and electron microscopy demonstrated the underlying pathological hallmark of characteristic inclusions to be shared by both phenotypes. Moreover, as the first pathological description of slowly evolving spastic paraparesis with adrenal insufficiency in young adults, it indicated clinical heterogeneity in the ALD disease group that, as shown later, extends further to females [7] with the same heterogeneity as in males, to Addison-only, and even to asymptomatics [17]. Neonatal ALD is now considered another disease with different genetic defects. Despite all progress, however, it has remained unclear how defects of the ABCD1 gene cause the strikingly broad, and unpredictable, phenotypic spectrum of ALD/AMN.

Our publication has met moderate interest, at least in scientometric terms: by Sept. 3, 2023, Google Scholar listed 125 citations. Indeed, in early years after the publication citations were rather scarce, until Hugo Moser started to cite it in his reviews on the history of the field. It was a great pleasure and honor when he was visiting our Institute in Vienna a few times, and I was greatly impressed by his personality, openness and kindness, knowledge and willingness to co-operate on behalf of his patients.

I was also happy to have met Jim Powers a few times, a kind and experienced colleague and now the foremost neuropathologist of ALD/AMN. We discussed our mutual interests, and it is clear that he – with Herb Schaumburg, Cedric Raine, Jack Griffin, Peter Spencer, John Prineas and others – published their AMN cases in the December 1977 issue of Neurology [11] without being aware of our preceding publication. Indeed there was no Pubmed or Internet at that time. Anyway, our stories are another example that scientific progress may occur simultaneously and independently in distant locations and settings.

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