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An autopsy case report of adult-onset Krabbe disease: Comparison with an infantile-onset case

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Abbreviations:

·Galactocerebrosidase; GALC

•Hematoxylin and Eosin staining; H&E staining

•Klüver–Barrera staining; KB staining

·Magnetic resonance imaging; MRI

Periodic acid–Schiff; PAS staining

• Triosephosphate isomerase barrel; TIM barrel

Short running title:

An autopsy report of adult-onset Krabbe disease

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1 Abstract

2	Krabbe disease is a lysosomal storage disease caused by a deficiency of the
3	galactocerebrosidase (GALC) enzyme, which leads to demyelination of the central and
4	peripheral nervous systems. Almost all patients with Krabbe disease are infants, and this is the
5	first report of adult-onset cases that describe pathological findings. Here, we present two
6	autopsy cases: a 73-year-old female and a 2-year-old male. The adult-onset case developed
7	symptoms in her late thirties and was diagnosed by the identification of GALC D528N and
8	L634S mutations and by T2-weighted magnetic resonance imaging; she had increased signal in
9	the white matter along the pyramidal tract to the bilateral precentral gyrus, as well as from the
10	triangular part to the posterior horn of the lateral ventricle. Microscopically, Klüver-Barrera
11	staining was pale in the white matter of the precentral gyrus and occipito-thalamic radiation, and
12	a few globoid cells were observed. The GALC mutations that were identified in the present
13	adult-onset case do not completely inactivate GALC enzyme activity, resulting in focal
14	demyelination of the brain.
15	
16	Keywords
17	Krabbe disease, galactocerebrosidase, adult-onset

Introduction 19

20	Krabbe disease (globoid cell leukodystrophy), which was identified by Krabbe in 1916 ¹ ,
21	is an autosomal recessive leukodystrophy with a morbidity of one in two hundred thousand
22	people. It is classified as a lysosomal storage disease and is caused by deficiency of the
23	galactocerebrosidase (GALC) enzyme, which leads to demyelination of the central and
24	peripheral nervous systems ^{2, 3} . Most cases (95%) of Krabbe disease are infantile onset (before
25	the age of 6 months) ⁴ . Clinical manifestations of the disease include hyperirritability,
26	hypersensitivity, stiffness, episodic fever, hypertonicity, decerebrate posturing, blindness, and
27	unresponsiveness. Most patients die by the age of 2 years. Three other onset types of the disease
28	are recognized: late infantile (6 months to 3 years), juvenile (3 to 8 years), and adult (after 20
29	years). The manifestations of adult-onset disease include asymmetric limb weakness, spastic
30	gait, poor coordination for balance, and tremors; the symptoms mimic those of motor neuron
31	disease ^{5, 6} . The disease progresses slowly in some patients, who have a normal life span. This
32	report compares an adult-onset case with an infantile-onset case using both autopsy and
33	pathological findings.
34	

34

35CASE 1

36 CLINICAL SUMMARY

37	This was a female autopsy case that was 73 years old. The patient noticed
38	difficulties with lifting her legs in her late thirties. From 56 to 62 years of age, she used crutches
39	to walk and then progressed to a wheelchair. Neurological manifestations at the age of 67
40	included amyotrophy of the distal muscles in the bilateral limbs (manual muscle test: proximal
41	muscles 2-3, distal muscles 1-2), walking difficulties, jaw jerk reflex, periosteal reflex of the
42	limbs, Babinski reflex were positive, sensation disorder beyond both knees, and axonopathy of
43	the limbs (in a nerve conduction test). On magnetic resonance imaging (MRI; T2-weighted), the
44	cerebral white matter showed increased signals (1) along the pyramidal tract to the bilateral
45	precentral gyrus and (2) from the triangular part to the posterior horn near the lateral ventricle
46	(especially the occipito-thalamic radiation) (Fig. 1A). These findings were unchanged for 13
47	years until her death. The patient was diagnosed with Krabbe disease because of MRI findings
48	and the identification of GALC mutations (D528N and L634S mutations) when she was 71
49	years old. She died of aspiration pneumonia aged 73.
50	
51	PATHOLOGICAL FINDINGS
52	Macroscopically, a white matter lesion around the central sulcus was not obvious
53	(Fig. 1B). Microscopically, the white matter of the precentral gyrus was pale using Klüver-
54	Barrera (KB) staining (Fig. 1C). In this area, mild astrocytosis and a few multinucleated giant

55	cells with scant cytoplasm and oval nuclei were identified (Fig. 1D). These multinucleated cells
56	were positive for Periodic acid-Schiff (PAS) stain (Fig. 1E) and CD68 (Supplementary Fig. 1B),
57	and were consistent with globoid cells: the characteristic macrophages of Krabbe disease. There
58	were few T-cells around the vessels. The white matter was strongly positive for neurofilament
59	staining without spheroids; therefore, demyelination is mild (Supplementary Fig. 3A-C). The
60	pyramidal tracts of the midbrain and cervical spinal cord were degenerated and exhibited slight
61	myelin pallor (Supplementary Fig. 1C).
62	
63	CASE 2
64	CLINICAL SUMMARY
65	This was a male autopsy case that was 1 year and 7 months old. The patient had notably
66	delayed milestones of physical development at the ages of 3 and 6 months. At the age of 9
67	months, enlargement of the subarachnoid space was apparent in computed tomography, early
68	closure of the coronal suture was observed by skull X-ray, and delayed myelination was seen on
69	MRI. Furthermore, his cerebrospinal fluid protein was elevated (110 mg/dL). At the age of 11
70	months, he lost eye contact and head control, and brain atrophy and demyelination were noted
71	on MRI (Fig. 2A). At 1 year and 4 months of age, truncal opisthotonus and rigid-spastic

73	1 year and 6 months, he had respiratory failure and apparent weight loss. At 1 year and 7
74	months of age, he was hospitalized with bacterial pneumonia and died of severe respiratory
75	failure. His lysosomal enzyme activity test revealed that he had low GALC activity (Table 1).
76	
77	PATHOLOGICAL FINDINGS
78	Macroscopically, white matter lesions around the internal capsule and cerebral ventricle
79	was noted as light brown and transparent (Fig. 2B). The white matter of the cerebrum and
80	cerebellum was pale in KB staining, but was stained well with Holzer staining, thus suggesting
81	fibrillary gliosis (Fig. 2C). White matter was severely disrupted, and foamy macrophages,
82	globoid cells, and marked astrocytosis were observed. Countless globoid cells had abundant
83	cytoplasm, oval nuclei (Fig. 2D), and PAS-positive inclusions (Fig. 2E). Gliosis was severe
84	with numerous fibrillary astrocytes and gemistocytes (Supplementary Fig. 2B). There was some
85	T-cell infiltration around the vessels. Marked demyelination was observed with no remaining
86	axons in neurofilament staining, but only U-fibers between the gray and the white matter were
87	stained well (Supplementary Fig. 3D-F).
88	
89	Discussion

Compared with the infantile-onset case, the adult-onset case progressed gradually, and the

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91	symptoms were more focused. A table showing the comparisons between the two cases is
92	included (Table 1). The adult-onset case showed a mild disorder of movement, sensation, neural
93	reflex, amyotrophy, and axonopathy in all four limbs (especially in the inferior limbs). In
94	contrast, the infantile-onset case showed rapid development throughout the whole body and
95	involved a developmental delay of psychomotor performance. In both cases, the activity of
96	GALC was unable to be detected by laboratory tests. Microscopically, the convolutional white
97	matter of the precentral gyrus and occipito-thalamic radiation was demyelinated in the
98	adult-onset case, while all white matter-in both the cerebrum and cerebellum-was affected in
99	the infantile-onset case (Fig. 3). Degeneration of the pyramidal tract throughout the midbrain to
100	the spinal cord was observed in both cases. Additionally, only a few globoid cells with scant
101	cytoplasm and oval nuclei were observed in the adult-onset case; in contrast, these cells were
102	present in large quantities with abundant cytoplasm, oval nuclei, and PAS-positive inclusions in
103	the infantile-onset case. Gliosis and demyelination were mild, and axons remained in the
104	adult-onset case, whereas severe gliosis and demyelination lead to marked axonal degeneration
105	in the infantile-onset case.
106	In adult-onset cases, demyelination is generally localized in the precentral gyrus and
107	occipito-thalamic radiation. GALC, which is deficient in Krabbe disease, dissolves important
108	components of myelinization in cells, such as galactosylceramide and psychosine ⁷ . An

109	accumulation of these substances causes cell damage and leads to the appearance of globoid
110	cells and demyelination. Thus, frequent myelin turnover can be the main lesion of
111	demyelination ⁸ . The white matter of the precentral gyrus, which was damaged in the present
112	adult-onset case, has neural fibers of the corticospinal tract that travel from Betz cells in the
113	precentral cortex to the lumbo-sacral cord. These are one of the longest and largest neural fibers
114	in the human body ⁹ , and their myelin turnover may be more active than the others. The
115	precentral gyrus might therefore be prone to accumulating substances such as
116	galactosylceramide and psychosine, and thus be more vulnerable to demyelination. White
117	matter lesions of the triangular part of the posterior horn involve a pathway that is related to
118	vision: the occipito-thalamic radiation. Our adult-onset case showed a focal lesion with a few
119	globoid cells in this area, which might have been caused by unknown characteristics of myelin
120	structure or by oligodendrocytes in the occipito-thalamic radiation.
121	Krabbe disease is caused by mutations in the GALC gene that result in degeneration of
122	the GALC protein. GALC consists of 668 amino acids and has three representative domains (the
123	triose-phosphate isomerase [TIM] barrel, β -sandwich domain, and lectin domain) ¹⁰ . These
124	domains combine to form a large substrate-binding pocket. Fig. 4 shows the typical mutation
125	sites in the GALC protein that have been found in adult- or infantile-onset patients with Krabbe
126	disease. The mutation sites vary widely but appear to be concentrated in the TIM barrel and

127	lectin domains. Typical examples of mutations in adult-onset cases include L634S,
128	[I82M+I305V], and G286D ¹¹ , while c.683_694delinsCTC (N228_S232delinsTP), T668P,
129	R220X, 30kDa del, and D528N are more common in infantile-onset cases ¹¹ . In our adult-onset
130	case, L634S and D528N were identified. The L634S substitution occurs in the lectin-binding
131	domain, and has been demonstrated to impair the transport of GALC to lysosomes ¹² .
132	Furthermore, the D528N substitution does not result in a complete loss of enzymatic activity in
133	GALC, but its transport into the lysosome is impaired, resulting in a decrease in enzymatic
134	activity ¹³ . The identified mutations in the present case, L634S and D528N, are characteristic of
135	adult- and infantile-onset cases, respectively; however, the combination of these mutations
136	seems to have resulted in the adult-onset form of the disease. Neither mutation completely
137	inactivates GALC, but they impair its transport to lysosomes, which results in a relatively mild
138	phenotype. Thus, the combination of mutation sites may also be important in determining the
139	type of disease.
140	In the present report, the adult-onset case exhibited a milder clinical course than the
141	infantile-onset case. Additionally, the demyelinated lesions were limited to the precentral gyrus
142	and triangular area of the posterior horn, and were accompanied by just a few globoid cells. The
143	GALC mutations L634S and D528N do not completely inactivate GALC enzyme activity, thus
144	resulting in focal demyelination of the brain.

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Disclosure Statement

None declared.

Ethics Approval

The project was approved by an institutional ethics committee. For human subjects, the investigation was conducted in accordance with the Declaration of Helsinki of 1975.

Author Contributions

All authors contributed to the study conception and design. Material preparation, data collection, and analysis were performed by M.E., M.S., Z.T., and Y.O. The first draft of the manuscript was written by M.E. and M.S. and revised by Z.T. and S.T. All authors commented on previous versions of the manuscript and read and approved the final manuscript.

References

1 Compston A. A new familial infantile form of diffuse brain-sclerosis. *Brain*. 2013; **136**: 2649-51.

2 Suzuki K, Suzuki Y. Globoid cell leucodystrophy (Krabbe's disease): deficiency of galactocerebroside beta-galactosidase. *Proc Natl Acad Sci U S A*. 1970; **66**: 302-9.

3 Love S, Budka H, Ironside JW, Perry A, Greenfield JG. Greenfield's neuropathology. 9th ed edn: CRC Press 2015; 2 v. Chapter 6.

4 Graziano AC, Cardile V. History, genetic, and recent advances on Krabbe disease. *Gene*. 2015; **555**: 2-13.

5 Henderson RD, MacMillan JC, Bradfield JM. Adult onset Krabbe disease may mimic motor neurone disease. *J Clin Neurosci*. 2003; **10**: 638-39.

6 Krieg SI, Krageloh-Mann I, Groeschel S, *et al.* Natural history of Krabbe disease - a nationwide study in Germany using clinical and MRI data. *Orphanet J Rare Dis.* 2020; **15**: 243.

7 Suzuki K. Evolving perspective of the pathogenesis of globoid cell leukodystrophy (Krabbe disease). *Proceedings of the Japan Academy, Series B.* 2003; **79B**: 1-8.

8 McTigue DM, Tripathi RB. The life, death, and replacement of oligodendrocytes in the adult CNS. *J Neurochem*. 2008; **107**: 1-19.

9 Hall JE, Guyton AC. Textbook of medical physiology. 13th ed edn: Elsevier 2016; xix, 1145 p.

10 Deane JE, Graham SC, Kim NN, *et al.* Insights into Krabbe disease from structures of galactocerebrosidase. *Proc Natl Acad Sci U S A.* 2011; **108**: 15169-73.

11 Hossain MA, Otomo T, Saito S, *et al.* Late-onset Krabbe disease is predominant in Japan and its mutant precursor protein undergoes more effective processing than the infantile-onset form. *Gene.* 2014; **534**: 144-54.

12 Shin D, Feltri ML, Wrabetz L. Altered Trafficking and Processing of GALC Mutants Correlates with Globoid Cell Leukodystrophy Severity. *J Neurosci.* 2016; **36**: 1858-70.

13 Lee WC, Kang D, Causevic E, Herdt AR, Eckman EA, Eckman CB. Molecular characterization of mutations that cause globoid cell leukodystrophy and pharmacological rescue using small molecule chemical chaperones. *J Neurosci.* 2010; **30**: 5489-97.

Figure legends

Fig. 1 Radiological and pathological images of the adult-onset case (Case 1).

(A) Magnetic resonance images (T2-weighted images) 13 years before death showing high intensity in the white matter from the triangular part to the posterior horn in the lateral ventricle (arrows) and along the pyramidal tract to the bilateral precentral gyrus (arrowheads); these findings do not change until death. (B) No remarkable findings around the central sulcus (left) and posterior horn (right) in a coronal section macroscopically. (C) The white matter of the precentral gyrus showing myelin pallor in Klüver–Barrera staining. (D) A few globoid cells with scant cytoplasm and oval nuclei, and mild astrocytosis in the demyelinated lesions. (E) Globoid cell positive for Periodic acid–Schiff staining. Scale bars: 50 mm (B), 10 mm (C), 5 μ m (D, E).

Fig. 2 Radiological and pathological images of the infantile-onset case (Case 2).

(A) Magnetic resonance images (T2-weighted images) at the age of 11 months (8 months before death) showing high intensity in the posterior horn of the lateral ventricle (arrows). (B) The white matter around the internal capsule and cerebral ventricle showing brownish and transparent, macroscopically. (C) The white matter around the cerebral ventricle showing myelin pallor in Klüver–Barrera staining and fibrillary gliosis by Holzer staining. (D) Countless globoid cells with abundant cytoplasm and oval nuclei, and marked gliosis. (E) Globoid cell inclusions positive for Periodic acid–Schiff staining.

Scale bars: 50 mm (B, C), 5 µm (D, E).

Fig. 3 Distribution of the lesions.

The demyelinated lesion is colored red, and the secondary degeneration is depicted in blue. The main lesion is localized in the adult-onset case (Case 1) and is widespread in the infantile-onset case (Case 2). Both cases exhibit secondary degeneration of the pyramidal tract.

Fig. 4 Locations of amino acid substitutions in GALC.

The specific mutations for adult-onset disease, such as L634S (our case has this mutation) and G286D,

are colored red. In contrast, mutations in blue are common in infantile-onset disease (e.g.,

c.683_694delinsCTC and T668S).

Tables

	Case 1	Case 2
Age of onset	Mid-30s	3 months
Symptoms	Limb weakness Walking difficulty Spastic gait Amyotrophy of distal muscle, especially bilateral limbs Jaw jerk reflex, limbs periosteal reflex, Babinski's reflex All sensation disorder beyond both knees Axonopathy of limbs	Psychomotor retardation Dilation of circumference and subarachnoid cavity Delay of myelination Cerebrospinal fluid protein abnormality Delay of peripheral nerve conduction velocity Opisthotonos of trunk Rigidity and spasm Massive myoclonus Tonic seizure Respiratory disorder
Disease progression	Slow	Rapid
Galactocerebrosidase Reference value 0.75 ± 0.27 (mol/h/mg)	0.08	Trace
GALC mutation	D528N, L634S	Not examined
Lesions	White matter close to precentral gyrus and posterior horn of lateral ventricle (localized) Secondary degeneration of the pyramidal tract	Cerebrum and cerebellar white matter (extensive) Secondary degeneration of the pyramidal tract
Globoid cell Number Cytoplasm PAS-positive inclusions CD68 staining	A few (one cell in one slide) Scant Oval Few Positive	Countless Abundant Oval Rich Positive
Gliosis	Mild	Severe
Demyelination	Mild	Severe
Axon	Remained	Degenerated
Lymphoid cell infiltration	Scant T-cell infiltration around vessels	T-cell infiltration around vessels

Table 1 Summary of the adult-onset case (Case 1) and the infantile-onset case (Case 2)



Fig. 1 Radiological and pathological images of adult-onset case (Case 1).



Fig. 2 Radiological and pathological images of infantile-onset case (Case 2) .



Fig. 3 Distribution of the legions.



Fig. 4 Locations of amino-acid substitution in GALC.