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1 Concise Communication

2	Beneficial screening of Fabry disease in patients with hypohidrosis
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26 Abstract

27 Fabry disease (FD), which is a lysosomal storage disease resulting from a deficiency of 28 α-galactosidase A, leads to the accumulation of globotriaosylceramide in various tissues 29 and multiorgan impairment. Early diagnosis is important to improve long-term prognosis. 30 Early clinical manifestations of FD include neuropathic pain, vascular skin lesions, and 31 sweating abnormalities. Hypohidorosis is one of the clinical findings in the early stage of 32 FD. However, there have been no studies on prospective screening of FD in patients with definitive diagnosis of hypohidrosis. We examined a-galactosidase A activity in white 33 34 blood cells in 17 (1 female and 16 male) patients with generalized hypohidorosis. Among 17 patients, 1 male patient (approximately 5.8%) had significantly reduced α-35 36 galactosidase A activity. He presented with a history of hypohidrosis with heat 37 intolerance and neuropathic tingling pain in a warm environment from 6 years ago. He 38 had a few angiokeratoma on the trunk and extremities. Ultrastructural examination of skin biopsy from the angiokeratoma revealed lamellar inclusions in endothelial cells. Kidney 39 40 biopsy revealed swollen podocytes and Gb3 deposition in the glomerulus, and urinalysis 41 revealed mulberry bodies. He was finally diagnosed with FD and started on enzyme replacement therapy with agalsidase alpha in the early stage. In addition, his family 42

- 43 screening led to find the patients of four additional FD. Screening for FD in patients with
- 44 hypohidrosis may lead to efficient early detection of FD.
- 45
- 46 Keywords
- 47 Fabry disease, hypohidrosis, α-galactosidase A activity, screening, early diagnosis

48 INTRODUCTION

49	Fabry disease (FD, OMIM 301500) is an X-linked lysosomal storage disorder caused by
50	mutations in <i>GLA</i> gene that encodes α -galactosidase A, resulting in reduced lysosomal
51	α -galactosidase A activity ¹ . Consecutively, globotriaosylceramide (Gb3) accumulates
52	in various tissues, leading to multiorgan impairment. Because patients with FD often
53	remain undiagnosed until severe complications involving cardiac, kidney, and
54	cerebrovascular lesions develop, early diagnosis is important to improve long-term
55	prognosis through suppressing multiorgan damage progression by several available
56	therapies ¹⁻³ . Early clinical manifestations of FD include neuropathic pain, vascular skin
57	lesions, and sweating abnormalities ⁴ . However, to the best of our knowledge, there
58	have been no studies on prospective screening of FD in patients with a definitive
59	diagnosis of hypohidrosis on the thermoregulatory sweat test. To confirm the benefits of
60	screening for FD in patients with hypohidrosis, we examined α -galactosidase A activity
61	in white blood cells in patients with generalized hypohidorosis.
62	

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63 CASE REPORT

64 One female and sixteen male patients with generalized hypohidrosis confirmed by

65 thermoregulatory sweat test at Kobe University (Kobe, Japan) were included.

66	Generalized hypohidrosis is defined as $\geq 25\%$ of the entire body affected with anhidrosis					
67	or hypohidrosis according to diagnostic criteria for acquired idiopathic generalized					
68	anhidrosis (AIGA) ⁵ . The clinical characteristics of these patients are shown in Table 1.					
69	α -galactosidase A activity was abnormally low in two male patients (Table 1, Pt.					
70	No.12,17). Further examination for definitive diagnosis of FD was performed in these					
71	patients with low α -galactosidase A activity. Notably, one patient (Table 1, Pt. No. 12)					
72	was diagnosed with AIGA without any other symptoms or laboratory abnormalities of					
73	FD, and steroid pulse therapy was effective in improving sweating abnormalities in this					
74	<u>patient</u> . Another patient with markedly reduced α -galactosidase A activity					
75	(0.1 nmol/h/mg) (Table1, Pt. No.17 case report) was definitively diagnosed with					
76	classical FD. Of the other 15 patients, 1 patient had a hypothalamic pituitary tumor that					
77	caused hypohidrosis and 14 patients were diagnosed with AIGA. Detailed information					
78	on Pt. No.17 is described herein.					
79	A 20-year-old man presented with a 6-year history of hypohidrosis with heat intolerance					
80	and neuropathic tingling pain in a warm environment. He had a past medical history of					
81	bronchial asthma and atopic dermatitis. Thermoregulatory sweat test using the iodine-					
82	starch method with sweating provoked by heat stimulation revealed anhidrosis of his					
83	entire body, including palms, soles, face, and axilla. Diseases that include hypohidrosis,					

84	such as Sjögren's syndrome and hypothyroidism, were excluded by laboratory and
85	physical examinations. As screening for FD, α -galactosidase A activity in white blood
86	cells was examined, and it was significantly reduced (0.1 nmol/h/mg). The clinical
87	examination revealed the presence of a few erythematous-purple papules on the trunk
88	and extremities (Fig. 1a). This symptom was noticed about 4-5 years ago. Abnormal
89	biochemical findings other than a decrease in α -galactosidase A activity were not
90	detected. His enzymatic test showed low levels of plasma alpha-galactosidase A activity
91	(0.1 nmol/h/mg), high levels of lyso-Gb3 (147.2ng/ml). Ultrastructural examination of
92	skin biopsy specimen from the angiokeratoma of the left thigh revealed lamellar
93	inclusions in endothelial cells (Fig. 1a, b). Kidney biopsy revealed swollen podocytes
94	and Gb3 deposition in the glomerulus. Corneal opacity (cornea verticillata) was faintly
95	observed, and urinalysis revealed mulberry bodies (Fig. 1c). Targeted sequencing by
96	next-generation and sanger sequencing revealed a hemizygous mutation of c.928 C>T;
97	p.(Leu310Phe) in exon 6, which has been reported previously as a pathogenic variant of
98	GLA encoding α -galactosidase A ⁶ . The patient was started on enzyme replacement
99	therapy (ERT) with agalsidase alpha 0.2 mg/kg body every 2 weeks. Furthermore, an
100	interview with the patient about his familial history revealed that his mother and three
101	brothers with neuropathic tingling pain and hypohidrosis from childhood. And his

102 mother was suffering from left ventricular hypertrophy of unknown origin. They were
103 finally diagnosed with FD and started on ERT (Fig. 2). <u>The patient and his two younger</u>
104 <u>brothers noticed an improvement in sweating after ERT.</u>

105

106 DISCUSSION

107 Early diagnosis is important to improve long-term prognosis of FD. Newb	oorn screening
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108 and high-risk screening have been attempted for early diagnosis. A newborn screening

study revealed that the frequency of FD was estimated to be 1:7683 in 599,711

110 newborns in Japan and 1:10,585 (2/21,170) in Washington, United States ^{7,8}. High-risk

- screening studies have found 7 patients with FD among 230 males (approximately 3%)
- 112 with left ventricular hypertrophy 9 and between 0.16 and 1.2 % have FD from dialysis
- 113 patients ¹⁰. The main causes of death for FD are renal failure, heart disease or stroke at
- around the age of 50 years for hemizygous men and 70 years for obligate carrier
- 115 women^{11,12}. Therefore, earlier screening before serious organ damage is important to
- 116 improve the long-term prognosis of FD. In contrast, 1 patient with FD was found among
- 117 17 patients with hypohidrosis (approximately 5.8%) in our small study; this high rate
- 118 indicates that screening by hypohidorosis might be more efficient, although further

119 studies need.

120	Dermatologists play an important role in early diagnosis because the cutaneous findings
121	like a neuropathic pain, angiokeratomas, and sweating abnormalities appear in early
122	stage of FD. Orteu et al ¹³ have documented the dermatological features of this disease
123	with reference to data from 714 patients (345 males, 369 females) registered on the
124	Fabry Outcome Survey (FOS), a multicentre European database. They showed that
125	78% of males and 50% of females of FD had one or more dermatological abnormality,
126	the commonest being angiokeratoma (66% males, 36% females), hypohidrosis (53%
127	males, 28% females), and telangiectasia (23% males, 9% females) ¹³ . Although
128	angiokeratoma is the most frequent manifestation, hypohidorosis is also an important
129	sign that appears with a high probability, especially in men. It is widely assumed that
130	the hypohidrosis of FD results principally from an autonomic peripheral neuropathy,
131	although Gb3 deposition in sweat gland cells have also been reported ¹⁴ , and the sweat
132	gland dysfunction may play a role.
133	Collectively, screening for FD in patients with hypohidrosis may lead to efficient early
134	detection of FD. Thus, the important role of dermatologists in the early detection of FD
135	should be reaffirmed.
136	

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- 185 eccrine sweat glands in a Fabry disease patient with hypohidrosis. *J Dermatol*
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187 Figure legends

- 188 Figure 1 (a, b): Angiokeratoma of the left thigh. (a) Clinical image shows multiple,
- raised, red-to-purple, hyperkeratotic papules on the thigh. (b) Electron microscopy
- 190 shows lamellar inclusions in endothelial cells on ultrastructural examination. (c):
- 191 Urinalysis revealed mulberry bodies.
- **192** Figure 2: Family pedigree: the index patient is marked with the arrowhead.
- 193

194 Table 1. α -galactosidase A activity in white blood cells of 17 patients diagnosed with

Patient	Age	Sex	Medical history	α -galactosidase A activity	Diagnosis
No.				in white blood cells	
				(normal range: 49.6–	
				116 nmol/h/mg)	
1	46	М	Chronic kidney disease,	50.2	AIGA
			Hypertension, Atrial fibrillation		
2	41	М	Atopic dermatitis-	103.3	AIGA
3	31	М	Systemic lupus erythematosus,	89.1	AIGA
			Lupus nephritis		
4	23	М	-	80.9	AIGA
5	29	М	-	72.7	Hypothalamic-
					pitutary tumor
6	50	М	-	62.5	AIGA
7	43	М	Central diabetes insipidus	83.2	AIGA
8	50	М	-	80.4	AIGA
9	25	М	-	52.5	AIGA
10	18	М	Asthma	69.1	AIGA
11	30	F	Gender dysphoria	66.3	AIGA
12	26	М	-	<u>30.6</u>	AIGA
13	39	М	-	51.2	AIGA-
14	17	М	-	79.6	AIGA
15	43	М	-	70.2	AIGA
16	38	М	-	72.7	AIGA
17	20	М	Atopic dermatitis, Asthma	<u>0.1</u>	FD

195 generalized hypohidrosis





