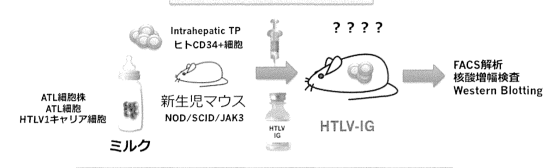
抗HTLV-1ヒト免疫グロブリン によるHTLV-1感染予防法の開発-3 (H26年度)

ヒト化マウスを用いた高力価HTLV-IGの有効性の検討

HTLV-1母乳感染モデルの確立



HTLV-IG製剤開発へ向けた研究

- ・ HTLV-IGの性状・用法・容量の設定
- ・ 抗HTLV-1ヒト免疫グロブリンの認識抗原の同定
- ・ 製造工程中のウイルス混入リスク解析(核酸増幅試験)
- 製造工程中のウイルスクリアランス
- 国家検定に準じた品質管理試験(安全性・物理化学試験)の実施

図8:抗 HTLV-1 ヒト免疫グロブリンによる HTLV-1 感染予防法の開発-3 ヒト化マウスの母乳感染モデルを構築することを試み、感染予防ツールとしての免疫グロブリンの有効性を検討する。また製剤化へ向けた品質管理に関わる予備的検討を行う。

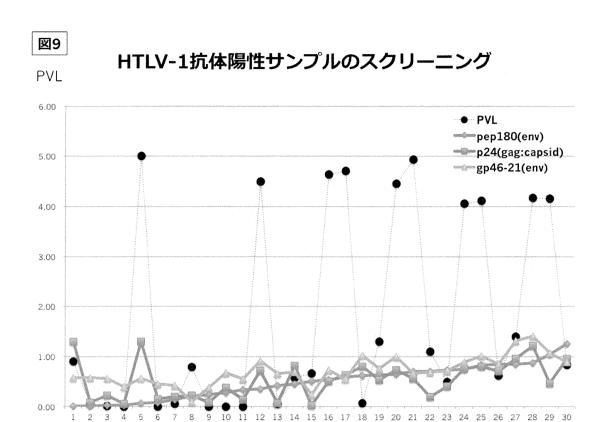


図9:HTLV-1 抗体陽性サンプルのスクリーニング 日本赤十字社に有している抗体陽性血漿をウイルス感染量(PVL)などを指標に分類し、30種類の候補血漿を準備した。

Sample number

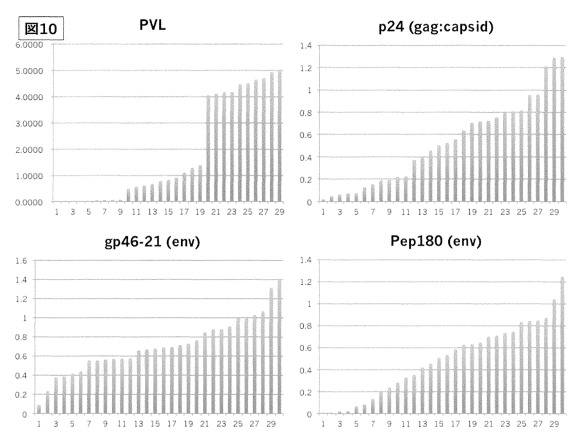


図 10: HTLV-1 抗体陽性サンプルのスクリーニング 日本赤十字社に有している抗体陽性血漿をウイルス感染量(PVL)などを指標に分類し、30 種類の候補血漿を準備した。それらのウイルスに対する結合部位を同定した。

培養細胞を用いたin vitro感染系の構築

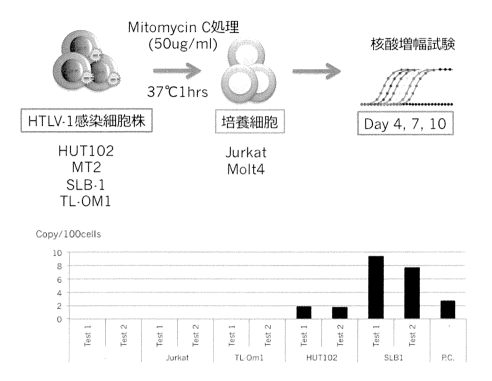


図 11: 培養細胞を用いた in vitro 感染系の構築 HTLV-1 感染細胞である HUT102, MT2, SLB-1, TL-0M1 をマイトマイシン C 処理をし、その後、非感染系細胞である Jurkat 細胞や Molt 4 細胞と共培養し、ウイルス感染を模す in vitro 感染系を構築する。

培養細胞を用いたin vitro感染系の構築

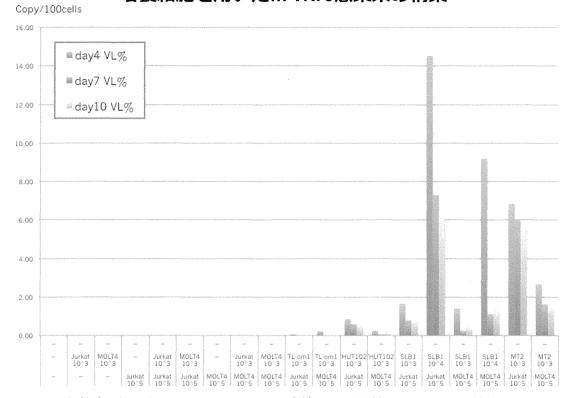
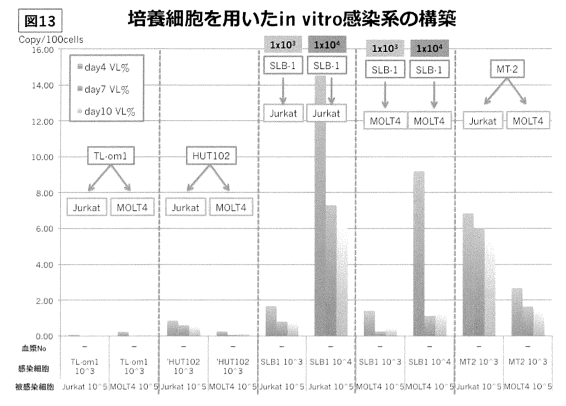


図 12: 培養細胞を用いた in vitro 感染系の構築 HTLV-1 感染細胞である HUT102, MT2, SLB-1, TL-0M1 をマイトマイシン C 処理をし、その後、非感染系 細胞である Jurkat 細胞や Molt 4 細胞と共培養し、ウイルス感染を模す in vitro 感染系を構築する。その結果、SLB-1, HUT102, MT-2 で有効なモデルが構築された。



感染効率: Jurkat>Molt4. 感染: MT-2>SLB-1>>>>HUT102>>TL-om1

図 13: 培養細胞を用いた in vitro 感染系の構築 HTLV-1 感染細胞である HUT102, MT2, SLB-1, TL-0M1 をマイトマイシン C 処理をし、その後、非感染系細胞である Jurkat 細胞や Molt 4 細胞と共培養し、ウイルス感染を模す in vitro 感染系を構築する。その結果、SLB-1, HUT102, MT-2 で有効なモデルが構築された。また、感染細胞数依存的に PVL の上昇が認められた。

図14 抗体陽性血漿を用いた感染抑制能の検討

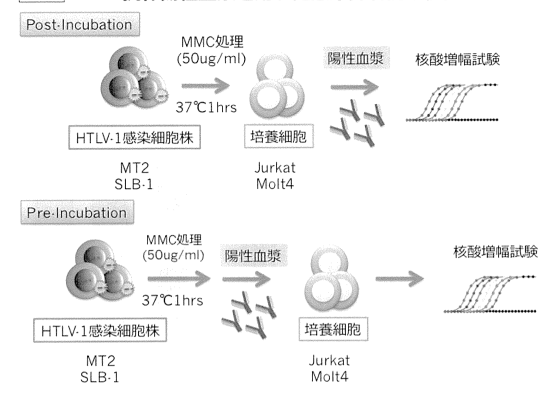


図 14: 抗体陽性血漿を用いた感染抑制能の検討 HTLV-1 感染細胞である HUT102, MT2, SLB-1, TL-0M1 をマイトマイシン C 処理をし、その後、陽性血漿を添加し、非感染系細胞である Jurkat 細胞や Molt 4 細胞における感染の有無を確認する。

抗体陽性血漿を用いた感染抑制能の検討-1

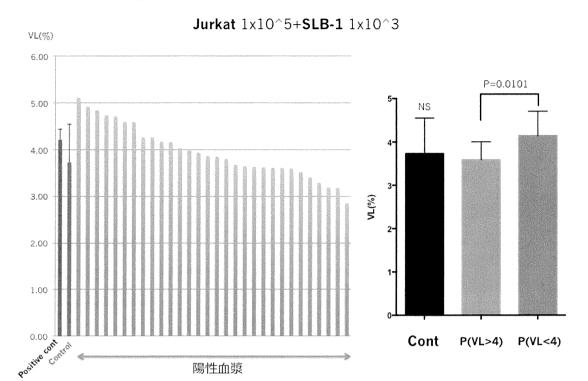


図 15: 抗体陽性血漿を用いた感染抑制能の検討-1 HTLV-1 感染細胞である SLB-1 にマイトマイシン C 処理をし、その後、陽性血漿を添加し、非感染系細胞である Jurkat 細胞と共培養し、4日目における感染の有無を確認した。 SLB-1 の系では 0.1%の投与では有効性は認められなかった。 Positive Control: HTLV-1 標準品, Control (Cont): 未添加サンプル, P: 血漿添加

図16 抗体陽性血漿を用いた感染抑制能の検討-2

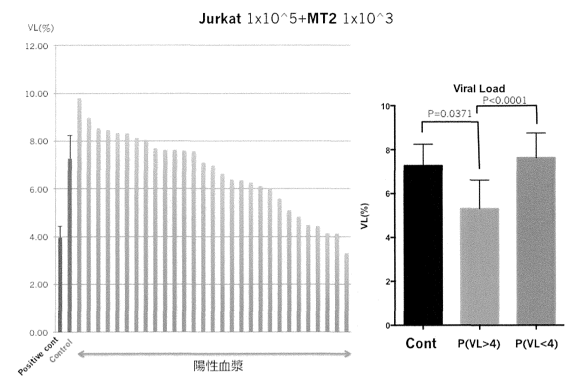


図 16: 抗体陽性血漿を用いた感染抑制能の検討-2 HTLV-1 感染細胞である MT-2 にマイトマイシン C 処理をし、その後、陽性血漿を添加し、非感染系細胞である Jurkat 細胞と共培養し、4 日目における感染の有無を確認した。MT-2 の系では PVL4 以上のサンプルにおいて有為な感染抑制が認められた。 **Positive Control**: HTLV-1 標準品,**Control (Cont)**: 未添加サンプル,P: 血漿添加

抗体陽性血漿を用いた感染抑制能の検討-3

Jurkat 1x10^5+MT2 1x10^3

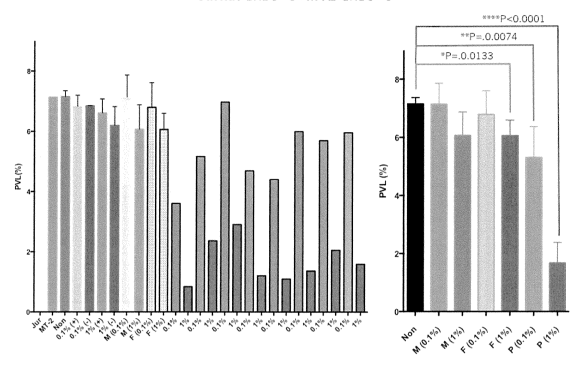


図 17:抗体陽性血漿を用いた感染抑制能の検討-3 HTLV-1 感染細胞である MT-2 にマイトマイシン C 処理をし、その後、陽性血漿を添加し、非感染系細胞である Jurkat 細胞と共培養し、4 日目における感染の有無を確認した。陽性血漿は、非働化、遠心処理をしたもの及び、1%の添加量の3つを追加した。その結果、いずれの血漿においても有為な感染抑制が認められた。 Jur: Jurkat のみ、MT-2: MT-2 のみ、Non: 未添加、M: Male negative plasma, F: Female negative plasma, P: Plasma 添加, +: 遠心処理あり、-: 遠心処理なし

- 1. HTLV-1感染細胞株を用いてHTLV-1感染系を構築した。
- 2. HUT102, TL-om1, SLB-1, MT2を用いて実験を行った結果、SLB-1, MT-2が高効率に感染をすることが明らかとなった。また、被感染細胞としてMolt4, Jurkatなどを検討したが、Jurkatへの感染効率がより良い事が明らかとなり、SLB-1/MT2-Jurkat系の感染システムが感染抑制効果の検証に有効であることが明らかとなった。
- 3. 30種類のHTLV-1陽性血漿を用いて予備的スクリーニングを行った結果、MT-2+Jurkat系ではPVL4以上の陽性血漿で有意な感染阻害が認められた一方、SLB-1-Jurkatでは感染阻害の程度が弱かった。
- 4. 陽性血漿をさらに非動化処理の有無・性別・遠心処理の有無・添加量 などに分類し、条件検討を行った結果、いずれも1%の濃度では有意に 感染抑制効果が認められた。
- 5. 現在、高力価グロブリンの精製を視野に入れてより効率の良い感染阻 害実験系の確立を目指している。また、PBMCでの同様の試験法の標 準化を検討中である。

図 18:考察

図19 本研究の意義と期待される成果・厚生行政への貢献

新規感染予防法 HTLV1感染について、HTLV-IGの感染阻止に関する有効性を示す

ヒト臨床へのTRS ヒト化マウスを用いる事で、ヒト細胞の実際の生体内での感染

予防を確認でき、**ヒトへの外挿**が可能となる

新薬開発/国際貢献 HTLV-1抗体陽性血漿は日本でしか入手できないため、高品質の HTLV-IGは日本発で開発・製造可能。世界的には数千万人の

キャリアがおり、世界的な規模での利用が期待されている。

厚生行政 HTLVワクチン(H23年度 厚生労働科学研究費 長谷川秀樹主任 研究者)が成功すれば、HTLV-IG+Vaccineとの組み合わせで

HBVの母子感染予防のような感染予防策が期待される。

献血の有効利用に繋がる

研究班

代表 水上 拓郎 国立感染症研究所 血液・安全性研究部 第4室室長 分担 浜口 功 国立感染症研究所 血液・安全性研究部 部長

大隈 和 国立感染症研究所 血液・安全性研究部 第3室室長

山口 一成 国立感染症研究所 血液・安全性研究部 客員研究員 佐竹 正博 西東京日本赤十字血液センター

田所 憲治 日本赤十字中央血液研究所

協力 野島 清子 国立感染症研究所 血液・安全性研究部 研究員

松本 千惠子 日本赤十字中央血液研究所

図 19:本研究の意義と期待される成果・厚生行政への貢献

II. 研究成果の刊行に関する一覧表

書籍

著者氏名	論文タイ	トル名	書籍全体の 編集者名	書	籍	名	出版社名	出版地	出版年	ページ
						•				

雑誌

↑ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □					
発表者氏名	論文タイトル	発表誌名	巻号 	ページ	出版年
Takizawa K*, Nakashima T *, Mizukami T *, Kuramitsu M, Endoh D, Kawauchi S, Sasaki K, Momose H, Kiba Y, Mizutani T, Furuta RA, Yamaguchi K, Hama guchi I	Degenerate PC R strategy with DNA microarray for detection of multiple and va rious subtypes of virus in the b lood screening.	Transfusion	In press		2013
Luc S, Luis TC, Boukarabil a H, Macaulay IC, Buza-Vi das N, Bouriez-Jones T, L utteropp M, Woll PS, Loug hran SJ, Mead AJ, Hultqui st A, Brown J, Mizukami T, Matsuoka S, Ferry H, A nderson K, Duarte S, Atkin son D, Soneji S, Domansk i A, Farley A, Sanjuan-Pla A, Carella C, Patient R, de Bruijn M, Enver T, Nerlov C, Blackburn C, Godin I, Jacobsen SE.	The earliest thy mic T cell prog enitors sustain B cell and myel	Nat Immunol.	13	412-419	2012
Kuramitsu M, Sato-Otsubo A, Morio T, Takagi M, To ki T, Terui K, Wang R, Ka nno H, Ohga S, Ohara A, Kojima S, Kitoh T, Goi K, Kudo K, Matsubayashi T, Mizue N, Ozeki M, Masu mi A, Momose H, Takizaw a K, Mizukami T, Yamagu chi K, Ogawa S, Ito E, Ha maguchi I.	Extensive gene deletions in Jap anese patients with Diamond-Bl	Blood	119	2376-238 4	2012

III. 研究成果の刊行物・印刷

Extensive gene deletions in Japanese patients with Diamond-Blackfan anemia

Madoka Kuramitsu,¹ Aiko Sato-Otsubo,² Tomohiro Morio,³ Masatoshi Takagi,³ Tsutomu Toki,⁴ Kiminori Terui,⁴ RuNan Wang,⁴ Hitoshi Kanno,⁵ Shouichi Ohga,⁶ Akira Ohara,² Seiji Kojima,⁶ Toshiyuki Kitoh,⁶ Kumiko Goi,¹⁰ Kazuko Kudo,¹¹ Tadashi Matsubayashi,¹² Nobuo Mizue,¹³ Michio Ozeki,¹⁴ Atsuko Masumi,¹ Haruka Momose,¹ Kazuya Takizawa,¹ Takuo Mizukami,¹ Kazunari Yamaguchi,¹ Seishi Ogawa,² Etsuro Ito,⁴ and Isao Hamaguchi¹

Department of Safety Research on Blood and Biological Products, National Institute of Infectious Diseases, Tokyo, Japan; ²Cancer Genomics Project, Graduate School of Medicine, The University of Tokyo, Tokyo, Japan; ³Department of Pediatrics and Developmental Biology, Graduate School of Medicine, Tokyo Medical and Dental University, Bunkyo-ku, Tokyo, Japan; ⁴Department of Pediatrics, Hirosaki University Graduate School of Medicine, Hirosaki, Japan; ⁵Department of Transfusion Medicine and Cell Processing, Tokyo Women's Medical University, Tokyo, Japan; ⁶Department of Pediatrics, Graduate School of Medicine, Styushu University, Fukuoka, Japan; ⁷First Department of Pediatrics, Toho University School of Medicine, Tokyo, Japan; ⁶Department of Pediatrics, Nagoya University Graduate School of Medicine, Nagoya, Japan; ⁹Department of Hematology/Oncology, Shiga Medical Center for Children, Shiga, Japan; ⁹Department of Pediatrics, School of Medicine, University of Yamanashi, Yamanashi, Japan; ⁹Division of Hematology and Oncology, Shizuoka Children's Hospital, Shizuoka, Japan; ⁹Department of Pediatrics, Seirei Hamamatsu General Hospital, Shizuoka, Japan; ⁹Department of Pediatrics, Kushiro City General Hospital, Hokkaido, Japan; and ⁹Department of Pediatrics, Graduate School of Medicine, Gifu University, Gifu, Japan

Fifty percent of Diamond-Blackfan anemia (DBA) patients possess mutations in genes coding for ribosomal proteins (RPs). To identify new mutations, we investigated large deletions in the RP genes RPL5, RPL11, RPL35A, RPS7, RPS10, RPS17, RPS19, RPS24, and RPS26. We developed an easy method based on quantitative-PCR in which the threshold cycle correlates to gene copy number. Using this approach, we were able to

diagnose 7 of 27 Japanese patients (25.9%) possessing mutations that were not detected by sequencing. Among these large deletions, similar results were obtained with 6 of 7 patients screened with a single nucleotide polymorphism array. We found an extensive intragenic deletion in RPS19, including exons 1-3. We also found 1 proband with an RPL5 deletion, 1 patient with an RPL35A deletion, 3 with RPS17 deletions, and 1 with an RPS19

deletion. In particular, the large deletions in the *RPL5* and *RPS17* alleles are novel. All patients with a large deletion had a growth retardation phenotype. Our data suggest that large deletions in RP genes comprise a sizable fraction of DBA patients in Japan. In addition, our novel approach may become a useful tool for screening gene copy numbers of known DBA genes. (*Blood*. 2012;119(10): 2376-2384)

Introduction

Diamond-Blackfan anemia (DBA; MIN# 105650) is a rare congenital anemia that belongs to the inherited BM failure syndromes, generally presenting in the first year of life. Patients typically present with a decreased number of erythroid progenitors in their BM. A main feature of the disease is red cell aplasia, but approximately half of patients show growth retardation and congenital malformations in the craniofacial, upper limb, cardiac, and urinary systems. Predisposition to cancer, in particular acute myeloid leukemia and osteogenic sarcoma, is also characteristic of the disease.²

Mutations in the *RPS19* gene were first reported in 25% of DBA patients by Draptchinskaia et al in 1999.³ Since that initial finding, many genes that encode large (RPL) or small (RPS) ribosomal subunit proteins were found to be mutated in DBA patients, including *RPL5* (approximately 21%), *RPL11* (approximately 9.3%), *RPL35A* (3.5%), *RPS7* (1%), *RPS10* (6.4%), *RPS17* (1%), *RPS24* (2%), and *RPS26* (2.6%).⁴⁻⁷ To date, approximately half of the DBA patients analyzed have had a mutation in one of these genes. Konno et al screened 49 Japanese patients and found that 30% (12 of 49) carried mutations.⁸ In addition, our data showed that 22 of 68 DBA patients (32.4%) harbored a mutation in ribosomal protein (RP) genes (T.T., K.T., R.W., and E.I., unpub-

lished observation, April 16, 2011). These abnormalities of RP genes cause defects in ribosomal RNA processing, formation of either the large or small ribosome subunit, and decreased levels of polysome formation, ^{4-6,9-12} which is thought to be one of the mechanisms for impairment of erythroid lineage differentiation.

Although sequence analyses of genes responsible for DBA are well established and have been used to identify new mutations, it is estimated that approximately half of the mutations remain to be determined. Because of the difficulty of investigating whole allele deletions, there have been few reports regarding allelic loss in DBA, and they have only been reported for *RPS19* and *RPL35A*.^{3,6,13} However, a certain percentage of DBA patients are thought to have a large deletion in RP genes. Therefore, a detailed analysis of allelic loss mutations should be conducted to determine other RP genes that might be responsible for DBA.

In the present study, we investigated large deletions using our novel approach for gene copy number variation analysis based on quantitative-PCR and a single nucleotide polymorphism (SNP) array. We screened Japanese DBA patients and found 7 patients with a large deletion in an allele in RPL5, RPL35A, RPS17, or RPS19. Interestingly, all of these patients with a large deletion had a phenotype of growth retardation, including short stature and

Submitted July 24, 2011; accepted November 15, 2011. Prepublished online as Blood First Edition paper, January 18, 2012; DOI 10.1182/blood-2011-07-368662.

The online version of this article contains a data supplement.

The publication costs of this article were defrayed in part by page charge payment. Therefore, and solely to indicate this fact, this article is hereby marked "advertisement" in accordance with 18 USC section 1734.

© 2012 by The American Society of Hematology

BLOOD, 8 MARCH 2012 • VOLUME 119, NUMBER 10

Table 1. Primers used for synchronized quantitative-PCR (s-q-PCR) of RPL proteins

Gene	Primer name	Sequence	Primer name	Sequence	Size, bp
RPL5	L5-02F	CTCCCAAAGTGCTTGAGATTACAG	L5-02R	CACCTTTCCTAACAAATTCCCAAT	132
	L5-05F	AGCCCTCCAACCTAGGTGACA	L5-05R	GAATTGGGATGGGCAAGAACT	102
	L5-17F	TGAACCCTTGCCCTAAAACATG	L5-17R	TCTTGGTCAGGCCCTGCTTA	105
	L5-19F	ATTGTGCAAACTCGATCACTAGCT	L5-19R	GTGTCTGAGGCTAACACATTTCCAT	103
	L5-21F	GTGCCACTCTCTTGGACAAACTG	L5-21R	CATAGGGCCAAAAGTCAAATAGAAG	102
	L5-28F	TCCACTTTAGGTAGGCGAAACC	L5-28R	TCAGATTTGGCATGTACCTTTCA	102
RPL11	L11-06F	GCACCCACATGGCTTAAAGG	L11-6R	CAACCAACCCATAGGCCAAA	102
	L11-20F	GAGCCCCTTTCTCAGATGATA	L11-20R	CATGAACTTGGGCTCTGAATCC	109
	L11-22F	TATGTGCAGATAAGAGGGCAGTCT	L11-22R	ATACAGATAAGGAAACTGAGGCAGATT	98
RPL19	L19-02F	TGGCCTCTCATAAAGGAAATCTCT	L19-02R	GGAATGCAGGCAAGTTACTCTGTT	103
	L19-08F	TTTGAAGGCAAGAAATAAGTTCCA	L19-08R	AGCACATCACAGAGTCCAAATAGG	107
	L19-16F	GGTTAGTTGAAGCAGGAGCCTTT	L19-16R	TGCTAGGGAGACAGAAGCACATC	102
	L19-19F	GGACCAGTAGTTGTGACATCAGTTAAG	L19-19R	CCCATTTGTAACCCCCACTTG	106
RPL26	L26-03F	TCCAAAGAGCTGAGACAGAAGTACA	L26-03R	TCCATCAAGACAACGAGAACAAGT	102
	L26-16F	TTTGAGAATGCTTGAGAGAAGGAA	L26-16R	TTCCAGCACATGTAAAATCAAGGA	102
	L26-18F	ATGTTTTAATAAGCCCTCCAGTTGA	L26-18R	GAGAACAGCAAGTTGAAAGGTTCA	102
	L26-20F	GGGCTTTGCTTGATCACTCTAGA	L26-20R	AGGGAGCCCGAAAACATTTAC	104
RPL35A	L35A-01F	TGTGGCTTCTATTTTGCGTCAT	L35A-01R	GGAATTACCTCCTTTATTGCTTACAAG	121
	L35A-07F	TTTCCGTTCTGTCTATTGCTGTGT	L35A-07R	GAACCCTGAGTGGAGGATGTTC	113
	L35A-17F	GCCCACAACCTCCAGAGAATC	L35A-17R	GGATCACTTGAGGCCAGGAAT	104
	L35A-18F	TTAGGTGGGCTTTTCAGTCTCAA	L35A-18R	ATCTCCTGATTCCCCAACTTTGT	102
RPL36	L36-02F	CCGCTCTACAAGTGAAGAAATTCTG	L36-02R	CTCCCTCTGCCTGTGAAATGA	102
	L36-04F	TGCGTCCTGCCAGTGTTG	L36-04R	GGGTAGCTGTGAGAACCAAGGT	105
	L36-17F	CCCCTTGAAAGGACAGCAGTT	L36-17R	TTGGACACCAGGCACAGACTT	114

Table 2. Primers used for s-q-PCR of RPS proteins

Gene	Primer name	Sequence	Primer name	Sequence	Size, bps
RPS7	S7-11F	GCGCTGCCAGATAGGAAATC	S7-11R	TTAGGGAGCTGCCTTACATATGG	102
	S7-12F	ACTGGCAGTTCTGTGATGCTAAGT	S7-12R	ACTOTTGCTCATCTCCAAAACCA	102
	S7-16F	GTGTCTGTGCCAGAAAGCTTGA	S7-16R	GAACCATGCAAAAGTGCCAATAT	112
RPS10	S10-03F	CTACGGTTTTGTGTGGGTCACTT	S10-03R	CATCTGCAAGAAGGAGACGATTG	102
10 DR 915 F	S10-15F	GTTGGCCTGGAGTCGTGATTT	S10-15R	ATTCCAAGTGCACCATTTCCTT	101
	S10-17F	AATGGTGTTTAGGCCAACGTTAC	S10-17R	TTTGAACAGTGGTTTTTGTGCAT	100
RPS14	S14-03F	GAATTCCAAACCCTTCTGCAAA	S14-03R	TTGCTTCATTTACTCCTCAAGACATT	104
	S14-05F	ACAACCAGCCCTCTACCTCTTTT	S14-05R	GGAAGACGCCGGCATTATT	102
	S14-06F	CGCCTCTACCTCGCCAAAC	S14-06R	GGGATCGGTGCTATTGTTATTCC	102
	S14-09F	GCCATCATGCCGAAACATACT	S14-09R	AACGCGCCACAGGAGAGA	102
	S14-13F	ATCAGGTGGAGCACAGGAAAAC	S14-13R	GCGAGGGAGCTGCTTGATT	111
	S14-15F	AGAAGTTTTAGTGAGGCAGAAATGAGA	S14-15R	TCCCCTGGCTATTAAATGAAACC	102
	S14-19F	GATGAATTGTCCTTTCCTCCATTC	S14-19R	TAGGCGGAAACCAAAAATGCT	102
RPS15	S15-11F	CTCAGCTAATAAAGGCGCACATG	S15-11R	CCTCACACCACGAACCTGAAG	108
	S15-15F	GGTTGGAGAACATGGTGAGAACTA	S15-15R	CACATCCCTGGGCCACTCT	108
RPS17	S17-03F	ACTGCTGTCGTGGCTCGATT	S17-03R	GATGACCTGTTCTTCTGGCCTTA	121
	S17-05F	GAAAACAGATACAAATGGCATGGT	S17-05R	TGCCTCCCACTTTTCCAGAGT	114
	S17-12F	CTATGTGTAGGAGGTCCCAGGATAG	S17-12R	CCACCTGGTACTGAGCACATGT	102
	S17-16F	TAGCGGAAGTTGTGTGCATTG	S17-16R	CAAGAACAGAAGCAGCCAAGAG	102
	S17-18F	TGGCTGAATCTGCCTGCTT	S17-18R	GCCTTGTATGTACCTGGAAATGG	103
	S17-20F	GGGCCCTTCACAAATGTTGA	S17-20R	GCAAAACTCTGTCCCTTTGAGAA	101
RPS19	S19-24F	CCATCCCAAGAATGCACACA	S19-24R	CGCCGTAGCTGGTACTCATG	120
	S19-28F	GACACACCTGTTGAGTCCTCAGAGT	S19-28R	GCTTCTATTAACTGGAGCACACATCT	114
41 3 SPEC 25 SPEC (2007 - PA 4)	S19-36F	CTCTTGAGGGTGGTCTGGAAAT	S19-36R	GTCTTTGCGGGTTCTTCCTCTAC	102
	S19-40F	GGAACGGTGTCAGGATTCAAG	S19-40R	AGCGGCTGTACACCAGAAATG	101
	S19-44F	CTGAGGTTGAGTGTCCCATTTCT	S19-44R	GCACCGGGCCTCTGTTATC	104
	S19-57F	CAGGGACACAGTGCTGAGAAACT	S19-57R	TGAGATGTCCCATTTTCACTATTGTT	101
	S19-58F	CATGATGTTAGCTCCGTTGCATA	S19-58R	ATTTTGGGAAGAGTGAAGCTTAGGT	102
	S19-62F	GCAACAGAGCGAGACTCCATTT	S19-62R	AGCACTTTTCGGCACTTACTTCA	102
	S19-65F	ACATTTCCCAGAGCTGACATGA	S19-65R	TCGGGACACCTAGACCTTGCT	102
RPS24	S24-17F	CGACCACGTCTGGCTTAGAGT	S24-17R	CCTTCATGCCCAACCAAGTC	101
	S24-20F	ACAAGTAAGCATCATCACCTCGAA	S24-20R	TTTCCCTCACAGCTATCGTATGG	105
	S24-32F	GGGAAATGCTGTGTCCACATACT	S24-32R	CTGGTTTCATGGCTCCAGAGA	105
RPS26	S26-03F	CGCAGCAGTCAGGGACATTT	S26-03R	AAGTTGGGCGAAGGCTTTAAG	104
	S26-05F	ATGGAGGCCGTCTAGTTTGGT	S26-05R	TGCCTACCCTGAACCTTGCT	102
RPS27A	S27A-09F	GCTGGAGTGCATTCGCTTGT	S27A-09R	CACGCCTGTAATCCCCACTAA	102
	S27A-12F	CAGGCTTGGTGTGCTGTGACT	S27A-12R	ACGTCCATCTTCCAGCTGCTT	103
	S27A-18F	GGGTTTTTCCTGTTTGGTATTTGA	S27A-18R	AAAGGCCAGCTTTGCAAGTG	111
	S27A-22F	TTACCATATTGCCAGTCTTTCCATT	S27A-22R	TTCATATGCATTTGCACAAACTGT	106

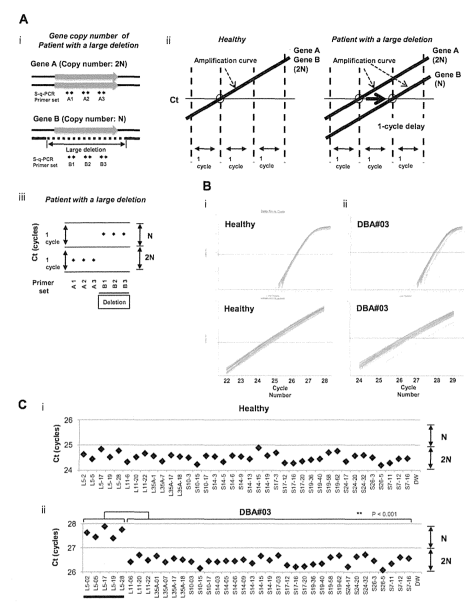


Figure 1. s-q-PCR can determine a large gene deletion in DBA. (A) Concept of the DBA s-q-PCR assay. The difference in gene copy number between a healthy sample and that with a large deletion is 2-fold (i). When all genomic s-q-PCR for genes of interest synchronously amplify DNA fragments, a 2-fold difference in the gene copy number is detected by a 1-cycle difference of the Ct scores of the s-q-PCR amplification curves (ii). Also shown is a dot plot of the Ct scores (iii). (B) Results of the amplification curves of s-q-PCR performed with a healthy person (i) and a DBA patient (patient 3; ii). The top panel shows the results of PCR cycles; the bottom panel is an extended graph of the PCR cycles at logarithmic amplification. (C) Graph showing Ct scores of s-q-PCR. If all specific primer sets for DBA genes show a 1-cycle delay relative to each other, this indicates a large deletion in the gene. Gene primer sets with a large deletion are underlined in the graph. **P< .001.

small-for-gestational age (SGA), which suggests that this is a characteristic of DBA patients with a large gene deletion in Japan.

Methods

2378

Patient samples

Genomic DNA was extracted using the GenElute Blood Genomic DNA Kit (Sigma-Aldrich) according to the manufacturer's protocol. Clinical manifes-

tation of patients from a Japanese DBA genomic library are listed elsewhere or are as reported by Konno et al.⁸ The study was approved by the institutional review board at the National Institute of Infectious Diseases and Hirosaki University.

DBA gene copy number assay by s-q-PCR

For s-q-PCR, primers were designed using Primer Express Version 3.0 software (Applied Biosystems). Primers are listed in Tables 1 and 2. Genomic DNA in water was denatured at 95°C for 5 minutes and

immediately cooled on ice. The composition of the s-q-PCR mixture was as follows: 5 ng of denatured genomic DNA, 0.4mM forward and reverse primers, 1× SYBR Premix Ex Taq II (Takara), and 1× ROX reference dye II (Takara) in a total volume of 20 µL (all experiments were performed in duplicate). Thermal cycling was performed using the Applied Biosystems 7500 fast real-time PCR system. Briefly, the PCR mixture was denatured at 95°C for 30 seconds, followed by 35 cycles of 95°C for 5 seconds, 60°C for 34 seconds, and then dissociation curve measurement. Threshold cycle (Ct) scores were determined as the average of duplicate samples. The technical errors of Ct scores in the triplicate analysis were within 0.2 cycles (supplemental Figure 1, available on the Blood Web site; see the Supplemental Materials link at the top of the online article). The sensitivity and specificity of this method was evaluated with 15 healthy samples. Any false positive was not observed in all primer sets in all healthy samples (supplemental Figure 2). We performed direct sequencing of the s-q-PCR products. The results of the sequence analysis were searched for using BLAST to confirm uniqueness. Sequence data were obtained from GenBank (http://www.ncbi.nlm.nih.gov/gene/) and Ensemble Genome Browser (http://uswest.ensembl.org).

Genomic PCR

Genomic PCR was performed using KOD FX (Toyobo) according to the manufacturer's step-down PCR protocol. Briefly, the PCR mixture contained 20 ng of genomic DNA, 0.4mM forward and reverse primers, 1mM dNTP, 1× KOD FX buffer, and 0.5 U KOD FX in a total volume of 25 µL in duplicate. Primers are given in supplemental Figure 3 and Table 2. PCR mixtures were denatured at 94°C for 2 minutes, followed by 4 cycles of 98°C for 10 seconds, 74°C for 12 minutes, followed by 4 cycles of 98°C for 10 seconds, 72°C for 12 minutes followed by 4 cycles of 98°C for 10 seconds, 70°C for 12 minutes, followed by 23 cycles of 98°C for 10 seconds and 68°C for 12 minutes. PCR products were loaded on 0.8% agarose gels and detected by LAS-3000 (Fujifilm).

DNA sequencing analysis

The genomic PCR product was purified by the GenElute PCR clean-up kit (Sigma-Aldrich) according to the manufacturer's instructions. Direct sequencing was performed using the BigDye Version 3 sequencing kit. Sequences were read and analyzed using a 3120x genetic analyzer (Applied Biosystems).

SNP array-based copy number analysis

SNP array experiments were performed according to the standard protocol of GeneChip Human Mapping 250K Nsp arrays (Affymetrix). Microarray data were analyzed for determination of the allelic-specific copy number using the CNAG program, as described previously.¹⁴ All microarray data are available at the EGA database (www.ebi.ac.uk/ega) under accession number EGAS00000000105.

Results

Construction of a convenient method for RP gene copy number analysis based on s-q-PCR

We focused on the heterozygous large deletions in DBAresponsible gene. The difference in copy number of genes between a mutated DBA allele and the intact allele was 2-fold (N and 2N; Figure 1Ai). If each PCR can synchronously amplify DNA fragments when the template genomic DNA used is of normal karyotype, it is possible to conveniently detect a gene deletion with a 1-cycle delay in s-q-PCR analysis (Figure 1Aii-iii).

Table 3. Summary of mutations and the mutation rate observed in Japanese DBA patients

Gene	Sequencing analysis
RPS19	10
RPL5	6
RPL11	The Control of Section 1997
RPS17	1
RPS10	1
RPS26	1
RPL35A	0
RPS24	0
RPS14	0
Mutations, n (%)	22 (32.4%)
Total analyzed, N	68

To apply this strategy for allelic analysis of DBA, we prepared primers for 16 target genes, RPL5, RPL11, RPL35A, RPS10, RPS19, RPS26, RPS7, RPS17, RPS24, RPL9, RPL19, RPL26, RPL36, RPS14, RPS15, and RPS27A, under conditions in which the Ct of s-q-PCR would occur within 1 cycle of that of the other primer sets (Tables 1 and 2). At the same time, we defined the criteria of a large deletion in our assay as follows. If multiple primer sets for one gene showed a 1-cycle delay from the other gene-specific primer set at the Ct score, we assumed that this represented a large deletion. As shown in Figure 1Bii and 1Cii, the specific primer sets for RPL5 (L5-02, L5-05, L5-17, L5-19, and L5-28) detected a 1-cycle delay with respect to the mutated allele of patient 3. This assessment could be verified by simply confirming the difference of the cycles with the s-q-PCR amplification curves.

Study of large gene deletions in a Japanese DBA genomic **DNA library**

Sixty-eight Japanese DBA patients were registered and blood genomic DNA was collected at Hirosaki University. All samples were first screened for mutations in RPL5, L11, L35A, S10, S14, S17, S19, and S26 by sequencing. Among these patients, 32.4% (22 of 68) had specific DBA mutations (Table 3 and data not shown). We then screened for large gene deletions in 27 patients from the remaining 46 patients who did not possess mutations as determined by sequencing (Table 4).

When we performed the s-q-PCR DBA gene copy number assay, 7 of 27 samples displayed a 1-cycle delay of Ct scores: 1 patient had RPL5 (patient 14), 1 had RPL35A (patient 71), 3 had RPS17 (patients 3, 60, 62), and 2 had RPS19 (patients 24 and 72; Figure 2 and Table 4). Among these patients, the large deletions in the RPL5 and RPS17 genes are the first reported cases of allelic deletions in DBA. From these results, we estimate that a sizable number of Japanese DBA patients have a large deletion.

Based on our findings, the rate of large deletions was approximately 25.9% (7 of 27) in a category of unspecified gene mutations. Such mutations have typically gone undetected by conventional sequence analysis. We could not find any additional gene deletions in the analyzed samples.

Confirmation of the gene copy number for DBA genes by genome-wide SNP array

We performed genome-wide copy number analysis of the 27 DBA patients with a SNP array to confirm our s-q-PCR results. SNP array showed that patient 3 had a large deletion in

Table 4. Characteristics of DBA patients tested

Patient no.	Age at diagnosis	Sex	Hb, g/dL	Large deletion by s-q-PCR	Large deletion by SNP array	Inheritance	Malformations	Response to firs steroid therapy
Patients with	a large deletion in RF	genes		100 mm	Translate all the			1000
3*†	1 y	М		RPL5	RPL5	Sporadic	Short stature, thumb anomalies	Response
14*	5 y	M	5.5	RPS17	RPS17	Sporadic	White spots, short stature	Response
24*†	1 mo	F	5.5	RPS19	ND	Sporadic	Short stature, SGA	Response
60*†	2 mo	F	2.4	RPS17	RPS17	Sporadic	SGA	NT
62*†	1 mo	F	6.2	RPS17	RPS17	Sporadic	Small ASD, short stature, SGA	Response
71			5.3	RPL35A	RPL35A	Sporadic	Thumb anomalies, synostosis of radius and ulna, Cohelia Lange-like face, cleft palate, underdescended testis, short stature, cerebellar hypoplasia, fetal hydrops	NT
72†	Оу	M	2	RPS19	RPS19	Sporadic	Thumb anomalies, flat thenar, testicular hypoplasia, tetal hydrops, short stature, learning disability	No
	nout a large deletion in				. 115 Sign	0 .	ND.	
5*	1 y	F	3.1	ND ND	ND ND	Sporadic	ND	Response
15*	1 mo	F	1.6	ND %	ND ND	Sporadic	ND	Response
21*	1 y	F 2	2.6	ND ND	ND ND	Sporadic	ND	Response
26*	1 y 1 mo	F	8	ND	M [®] ND °	Sporadic	Congenital hip dislocation, spastic quadriplegia, hypertelorism, nystagmus, short stature, learning disability	Response
33*	2 mo	F	1.3	ND	ND	Sporadic	ND	Response
36*	0 у	M	8.2	ND	ND	Familial	ND	Response
37*	4 y	М	6.1	ND	ND	Sporadic	Hypospadias, underdescended testis, SGA	NT
45*	5 d	M	5.1	ND	ND	Sporadic	Short stature, microcephaly, mental	Poor
							retardation, hypogammaglobulinemia	
50*	2 m	F	3.4	ND	ND	Familial	ND	Response
61*	9 m	M	4	ND	ND	Sporadic	ND	Response
63*	0 у	М	6.8	ND	ND	Sporadic	Micrognathia, hypertelorism, short stature	Response
68	1 y 4 mo	M	5.9	ND	ND	Sporadic	ND	NT (CR)
69	1 y	М	9.3	ND	ND	Sporadic	ND	Response
76	Oy	M	4	ND	ND	Sporadic	ND	Response
77	0 y	М	7.8	ND	ND	Familial	Short stature	No
83	9 mo	F	3	ND ND	ND	Sporadic	ND	NT
90	10 mo	М	9	ND	ND	Sporadic	ND	No
91	0 y	F	3.8	ND	ND	Sporadic	ND	Response
92	2mo	M	3.7	ND	ND	Sporadic	ASD, PFO, melanosis, underdescended testis, SGA, short stature	Response
93	11 mo	M	2.2	ND.	ND.	Sporadic	White spots, senile face, corneal opacity, underdescended testis, syndactyly, ectrodactyly, flexion contracture, extension contracture	Response

ND indicates not detected; NT, not tested; CR, complete remission; ASD, atrial septal defect; and PFO, persistent foramen ovale.

^{*}Status data of Japanese probands 3 to 63 is from a report by Konno et al.8

[†]Large deletions of the parents of 5 DBA patients (3, 24, 60, 62, and 72) were analyzed by s-q-PCR, but there were no deletions in DBA genes in any of the 5 pairs of parents.