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副島英伸	第XIV章先天異常・ 奇形 ベックウィ ズ·ヴィーデマン症 候群	編集,福井	ブッ		ンド	中山書店	東京	2011	P679
	第XIV章先天異常・ 奇形 シルバー・ラ ッセル症候群		ブッ		ヽンド	中山書店	東京	2011	P685

#### 雑誌

発表者氏名	論文タイトル名	発表誌名	巻号	ページ	出版年
Higashimoto K, Naka bayashi K, Yatsuki H, Yoshinaga H, Joz aki K, Okada J, Wat anabe Y, Aoki A, Sh iozaki A, Saito S, K oide K, Mukai T, Ha ta K, Soejima H.	Aberrant methylation of H19-DMR acquired after implantation was dissim ilar in soma versus place nta of patients with Bec kwith-Wiedemann syndro me.	Am J Med Genet A			in press
Aoki A, Shiozaki A, Sameshima A, Higa shimoto K, Soejima H, Saito S.	Beckwith-Wiedemann Sy ndrome with Placental Chorangioma due to H1 9-DMR Hypermethylatio n: A Case Report.	J Obstet G ynaecol Res	37(12)	1872-1876	2011
Sato S, Yoshida W, Soejima H, Nakabaya shi K, Hata K.	Methylation dynamics of IG-DMR and Gtl2-DM R during murine embryo nic and placental develop ment.	Genomics	98(2)	120-127	2011
Nakabayashi K, Truji Ilo AM, Tayama C, Camprubi C, Yoshida W, Lapunzina P, Sa nchez A, Soejima H, Aburatani H, Nagae G, Ogata T, Hata K, Monk D.	Methylation screening of reciprocal genome-wide UPDs identifies novel hu man specific imprinted g enes.	Hum Mol Genet	20(16)	3188-3197	2011
Nagae G, Isagawa T, Shiraki N, Fujita T, Yamamoto S, Tsuts umi S, Nonaka A, Y oshiba S, Matsusaka K, Midorikawa Y, Is hikawa S, Soejima H, Fukayama M, Sue mori H, Nakatsuji N, Kume S, Aburatani H.	Tissue-specific demethyla tion in CpG-poor promot ers during cellular differe ntiation.	Hum Mol Genet	20(14)	2710-2721	2011

Kurotaki N, Tasaki S, Mishima H, Ono S, Imamura A, Kiku chi T, Nishida N, To kunaga K, Yoshiura K, Hiroki Ozawa H.	Identification of Novel S chizophrenia Loci by Ho mozygosity Mapping Usi ng DNA Microarray Ana lysis.	PLos One	6(5)	e20589	2011
Oikawa M, Nagayasu T, Yano H, Hayashi T, Abe K, Kinoshit a A, Yoshiura KI.	Intracystic Papillary Carc inoma of Breast Harbors Significant Genomic Alt eration Compared with I ntracystic Papilloma: Genome-wide Copy Number and LOH Analysis Using High-Density Single-Nucleotide Polymorphism Microarrays.	Breast J	17(4)	427-430	2011
Hannibal MC, Bucki ngham KJ, Ng SB, Ming JE, Beck AE, McMillin MJ, Gilders leeve HI, Bigham A W, Tabor HK, Mefford HC, Cook J, Yos hiura K, Matsumoto T, Matsumoto N, Mi yake N, Tonoki H, Naritomi K, Kaname T, Nagai T, Ohashi H, Kurosawa K, Ho u JW, Ohta T, Liang D, Sudo A, Morris CA, Banka S, Black GC, Clayton-Smith J, Nickerson DA, Zack ai EH, Shaikh TH, Donnai D, Niikawa N, Shendure J, Bams had MJ.	Spectrum of MLL2 (AL R) mutations in 110 case s of Kabuki syndrome.	Am J Med Genet A	155A(7)	1511-1516	2011
Kobayashi H, Sakurai T, Takahashi N, Fu kuda A, Obata Y, Sa to S, Nakabayashi K, Hata K, Sotomaru Y, Suzuki Y, Kono T.	Contribution of Intrageni c DNA Methylation in Mouse Gametic DNA M ethylomes to Establish O ocyte-Specific Heritable Marks.	PLoS Genet	8(1)	e1002440	2012
Nakanishi M, Hayaka wa K, Nakabayashi K, Hata K, Shiota K, Tanaka S.	Trophoblast-specific DNA methylation occurs after the segregation of troph ectoderm and inner cell mass in mouse periimpla ntation embryo.	Epigenetics	7(2)	173 - 182	2012
Kobayashi H, Sakurai T, Sato S, Nakabay ashi K, Hata K, Kon o T.	Imprinted DNA methylati on reprogramming during early mouse embryogen sis at the <i>Gpr1-Zdbf2</i> lo cus is linked to long <i>cis</i> -intergenic transcription	FEBS Lette			in press

## IV. 研究成果の刊行物・別冊

### ベックウィズ・ヴィーデマン症候群

Beckwith-Wiedemann syndrome (BMS)

[ICD-10] Q87.3 [ OMIM ] #130650

【特記事項】厚生労働省難治性疾患克服研究事業 研究奨励分野の対象疾患

### **■疫学** 国内推定有病者数/200 人以上

男女比/1:1

■発症に関わる遺伝子 KVDMR1 (11p15.5), CDKN1C (11p15.5), KCNQ1OT1 (11p15.5), H19-DMR (11p15.5), IGF2 (11p15.5), H19 (11p15.5)

■診断 診断基準は主症状3つ以上、または主症状2つと副症状1つ以上とされている が、確定的な基準はない、

主症状:腹壁欠損(臍帯ヘルニア,腹直筋解離,臍ヘルニア),巨舌,過成長(>97パ ーセンタイル),耳垂の線状溝・耳輪後縁の小窩,腹腔内臓器腫大,胎児性腫瘍,

片側肥大, 副腎皮質細胞の腫大(びまん性, 両側性), 腎奇形, 家族歴, 口蓋裂 副症状:妊娠中の羊水過多・胎盤腫大・臍帯肥厚・早産,新生児期低血糖,火焔状母斑, 心肥大・心奇形・心筋症、特徴的顔貌、骨年齢亢進

上記の臨床診断に加え、KVDMR1の低メチル化あるいは H19-DMR の高メチル化や微 小欠失, CDKN1C変異, 11p15.5 父性片親性ダイソミーを検出すると確定診断できる.

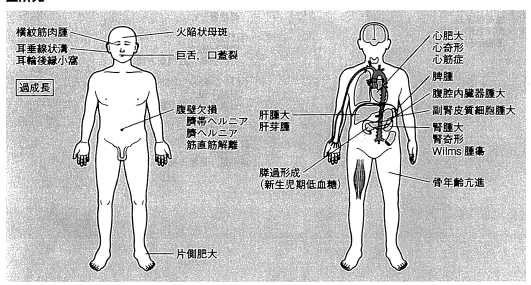
■治療 低血糖については生後の血糖値モニタリング. 臍ヘルニアに対しては外科的根治 術、8歳くらいまでは腹部超音波検査で3か月ごとに腫瘍をスクリーニング、4歳まで は定期的にαフェトプロテイン(AFP)も測定.過度の巨舌に対しては舌縮小術を、片 側肥大による下肢長の左右差(1~2cm 以上)に対しては骨端固定術を考慮する.

■関連語・同義語 ヴィーデマン・ベックウィズ症候群, exomphalos-macroglossiagigantism 症候群 (EMG 症候群)

■ EBM・診療ガイドライン 厚生労働省難治性疾患克服研究事業平成 21 年度研究報告書 ■関連団体・学会 Beckwith-Wiedemann 症候群親の会

■解説 John Bruce Beckwith (1933年生) と Hans-Rudolf Wiedemann (1915-2006) が、臍ヘルニア、巨舌、巨躯を伴う症例を 1969 年に別々に報告した. 11p15.5 のゲノムインプリンティングの異常で発症し、約7~11%に腫瘍を合併する、診断の項 目に記載した遺伝子異常が認められる. (副島英伸)

#### ■所見



【文献】1) Weksberg R, et al: Beckwith-Wiedemann syndrome. Eur J Hum Genet 2010; 18: 8–14.



## シルバー・ラッセル症候群

Silver-Russell syndrome (SRS)

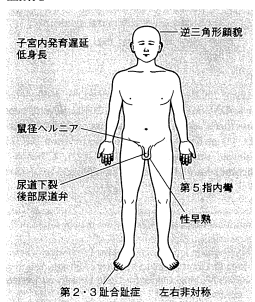
[ICD-10] Q87.1 【 OMIM 】#180860

【特記事項】厚生労働省難治性疾患克服研究事業 研究奨励分野の対象疾患

#### ■疫学 国内推定有病者数/160人以上(実際には非常に多いと推定される) 男女比/1:1

- **■発症に関わる遺伝子** H19-DMR (11p15.5), IGF2 (11p15.5), H19 (11p15.5), CDKN1C(11p15.5), KCNQ1OT1(11p15.5), 7番染色体母性ダイソミー(原因遺伝 子は未同定)
- ■診断 子宮内発育遅延, catch-up growth を伴わない低身長, 逆三角形の顔貌(前額 部突出,小さくとがった下顎,口角下降によるへの字様口唇),身体の左右非対称を主徴 とする. 他に, 第5指内彎, 第2・3趾の合趾症, 性早熟, 尿道下裂, 後部尿道弁, 鼠 径ヘルニアなど多様な症状を呈する. 診断基準が提案されている<sup>1,2)</sup>.
- **■治療** 発達は原則的に正常だが、最終身長は小児期の身長に比例する、哺乳・摂食障害 傾向にあるので、乳児期には適切な食事量を心がける. SGA (small-for-gestational age) 性低身長症の1つとして、基準を満たせば成長ホルモン治療を行う、
- ■関連語・同義語 Russell-Silver 症候群,Silver 症候群
- EBM・診療ガイドライン 厚生労働省難治性疾患克服研究事業平成 21 年度研究報告 書、日本小児内分泌学会・日本未熟児新生児学会「SGA 性低身長症における GH 治療の ガイドライン」「SGA 性低身長症における GH 治療の実施上の注意」
- ■関連団体・学会 SRS 患児・家族の交流を目的としたウェブサイトがある.
- ■解説 Henry K Silver(1918-1991)が先天性片側低形成,低出生体重,低身長,

#### ■所見



尿中ゴナドトロピン増加を伴う症例を 1953年に、Alexander Russell (1914 -2003) が、子宮内発育遅延、上述の特 徴的顔貌, 左右非対称を伴う症例を 1954 年に報告した. 患児の約10%で7番染色 体母性ダイソミーを、約30%で11p15.5 インプリンティング領域の H19-DMR の低 メチル化が認められる.

過成長を示す Beckwith-Wiedemann 症候群では逆に高メチル化がみられる.

また, 11p15.5インプリンティング領 域の母性重複および母性ダイソミーの報告 がある. (副島英伸)

奇形 天異常

【文献】1) Price SM, et al: The spectrum of Silver-Russell syndrome: a clinical and molecular genetic study and new diagnostic criteria. J Med Genet 1999, 36: 837-842.

Rossignol S, et al: Epigenetics in Silver-Russell syndrome. Best Pract Res Clin Endocrinol Metab 2008; 22: 403-414.

# Aberrant Methylation of H19-DMR Acquired After Implantation Was Dissimilar in Soma Versus Placenta of Patients With Beckwith—Wiedemann Syndrome

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Gain of methylation (GOM) at the H19 differentially methylated region (H19-DMR) is one of several causative alterations in Beckwith-Wiedemann syndrome (BWS), an imprinting-related disorder. In most patients with epigenetic changes at H19-DMR, the timing of and mechanism mediating GOM is unknown. To clarify this, we analyzed methylation at the imprinting control regions of somatic tissues and the placenta from two unrelated, naturally conceived patients with sporadic BWS. Maternal H19-DMR was abnormally and variably hypermethylated in both patients, indicating epigenetic mosaicism. Aberrant methylation levels were consistently lower in placenta than in blood and skin. Mosaic and discordant methylation strongly suggested that aberrant hypermethylation occurred after implantation, when genome-wide de novo methylation normally occurs. We expect aberrant de novo hypermethylation of H19-DMR happens to a greater extent in embryos than in placentas, as this is normally the case for de novo methylation. In addition, of 16 primary imprinted DMRs analyzed, only H19-DMR was aberrantly methylated, except for NNATDMR in the placental chorangioma of Patient 2. To our knowledge, these are the first data suggesting when GOM of H19-DMR occurs. © 2012 Wiley Periodicals, Inc.

**Key words:** Beckwith–Wiedemann syndrome; H19-DMR; aberrant DNA methylation; after implantation

#### INTRODUCTION

Beckwith—Wiedemann syndrome (BWS) is an imprinting-related condition characterized by macrosomia, macroglossia, and abdominal wall defects (OMIM #130650). The relevant imprinted chromosomal region in BWS, 11p15.5, consists of two independent imprinted domains, *IGF2/H19* and *CDKN1C/KCNQ1OT1*. Imprinted genes within each domain are regulated by two imprinting control

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Am J Med Genet Part A 9999:1—6.

regions (ICR), the H19-differentially methylated region (DMR) or KvDMR1 [Weksberg et al., 2010]. Several causative alterations have been identified in patients with BWS: loss of methylation (LOM) at KvDMR1, gain of methylation (GOM) at H19-DMR, paternal uniparental disomy (UPD), CDKN1C mutations, and chromosomal abnormality involving 11p15 [Sasaki et al., 2007; Weksberg et al., 2010].

Additional supporting information may be found in the online version of this article.

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Methylation of H19-DMR is erased in primordial germ cells (PGCs) but becomes reestablished during spermatogenesis [Li, 2002; Sasaki and Matsui, 2008]: this methylation regulates the expression of *IGF2* and *H19* by functioning as a chromatin insulator, restricting access to shared enhancers [Bell and Felsenfeld, 2000; Hark et al., 2000]. GOM on the maternal H19-DMR leads to expression of both *IGF2* alleles and silencing of both *H19* alleles. Dominant maternal transmissions of microdeletions and/or base substitutions within H19-DMR have recently been reported in a few patients of BWS with H19-DMR GOM [Demars et al., 2010]. However, when and how the GOM on the maternal H19-DMR occurs is not clear.

Here, we found epigenetic mosaicism in two BWS patients. We also found that GOM at H19-DMR was discordant in blood and skin versus placenta; specifically, methylation levels were lower in placental samples. These findings strongly suggest that aberrant methylation of H19-DMR occurred after implantation. As a result, we expect aberrant de novo methylation happens to a greater extent in embryos than in placentas.

#### MATERIALS AND METHODS

#### **Patients**

Two unrelated patients with sporadic BWS, Patient 1 (BWS047) and Patient 2 (bwsh21-015), were delivered by cesarean in the third trimester of pregnancy. The mothers of both patients conceived naturally. Patient 1 and Patient 2 met clinical criteria for BWS as described by Elliott et al. [1994] and Weksberg et al. [2001], respectively (Table I). The placenta of Patient 1 was large and weighed 1,065 g, but was without any pathological abnormality. The placenta of Patient 2 was also large, weighing 1,620 g, and had an encapsulated placental chorangioma, as reported previously [Aoki et al., 2011]. The standard G-banding chromosome analysis using peripheral blood samples showed no abnormalities in either patient. This study was approved by the Ethics Committee for Human Genome and Gene Analyses of the Faculty of Medicine, Saga University.

#### Southern Blot Analysis

Genomic DNA was extracted from embryo-derived somatic tissues and the placentas of the patients (Fig. 1). Methylation-sensitive

Southern blots with *Bam*HI and *Not*I were employed for KvDMR1, and blots with *Pst*I and *Mlu*I were employed for H19-DMR, as described previously [Soejima et al., 2004]. Band intensity was measured using the FLA-7000 fluoro-image analyzer (Fujifilm<sup>Q2</sup>, Japan). The methylation index (MI, %) was then calculated (Fig. 1). Southern blots with *Apa*I were used to identify the microdeletion of H19-DMR as described previously [Sparago et al., 2004].

## Bisulfite Sequencing and Combined Bisulfite Restriction Analysis (COBRA)

Bisulfite sequencing covering the sixth CTCF binding site (CTS6) was performed. For COBRA, PCR products of each primary imprinted DMR were digested with the appropriate restriction endonucleases and were then separated using the MultiNA Microchip Electrophoresis System (Shimazdu, Japan). The methylation index was also calculated. All PCR primer sets used in this study have been listed in Supplementary Table SI (See Supporting Information online).

#### DNA Polymorphism Analyses

For quantitative polymorphism analyses, tetranucleotide repeat markers (D11S1997 and HUMTH01) and a triplet repeat marker (D11S2362) from 11p15.4–p15.5 were amplified and separated by electrophoresis on an Applied Biosystems 3130 genetic analyzer (Applied<sup>Q3</sup> Biosystems); data were quantitatively analyzed with the GeneMapper software. The peak height ratios of paternal allele to maternal allele were calculated. A single nucleotide polymorphism (SNP) for the RsaI recognition site in H19 exon 5 (rs2839703) was also quantitatively analyzed using hot-stop PCR [Uejima et al., 2000]. Band intensity was measured using the FLA-7000 fluoro-image analyzer (Fujifilm).

#### Mutation Search of H19-DMR

To search for mutations in the binding sites of CTCF, OCT4, and SOX2, we sequenced a genomic region in and around H19-DMR, which included seven CTCF-binding sites, three OCT4 sites, and one SOX2 site.

		Birth weight			Placental weight	Placental—feta
Patient ID	Conception	(gestational age)	Clinical features	Karyotype	and pathology	weight ratio
Patient 1 (BWS047)	Natural	4,506 g (36w2d)	Macrosomia macroglossia abdominal wall defect hypoglycemia	46,XŸ	1,065 g no pathological findings	0.236
Patient 2 (bwsh21-015)	Natural	2,540 g (33w5d)	Macrosomia macroglossia hypoglycemia renal malformation hepatosplenomegaly	46,XX	1,620 g placental chorangioma	0.638

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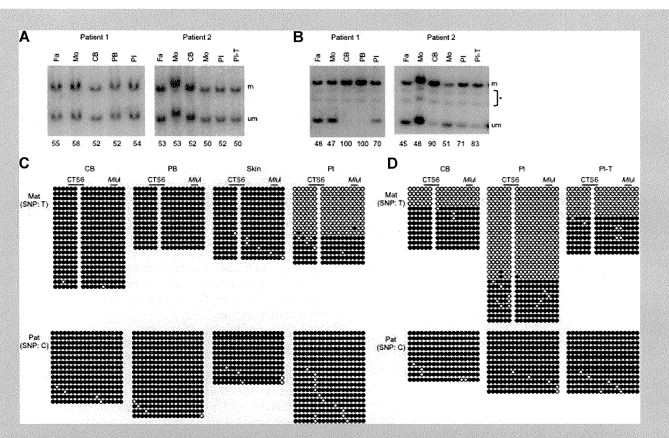


FIG. 1. Methylation analyses of KvDMR1 and H19-DMR. A: Methylation-sensitive Southern blots for KvDMR1. Genomic DNA was extracted from the cord blood, peripheral blood, skin, and placenta of Patient 1 and from the cord blood, placenta, and placental chorangioma of Patient 2. Methylation at KvDMR1 was normal in all samples analyzed. Methylation indices [MI, %] are shown under the figure. B: Methylation-sensitive Southern blots for H19-DMR. The MIs of blood samples were higher than the MIs of placental samples. MI was calculated using the equation  $[M/(M+U)] \times 100$ , where M is the intensity of the methylated band and U is the intensity of the unmethylated band. C: Bisulfite sequencing of H19-DMR in Patient 1. The two parental alleles were distinguishable by differences in SNPs. Both parental alleles were completely methylated in the cord blood, peripheral blood, and skin samples, and the maternal allele, which is normally unmethylated, was partially methylated in the placenta. D: Bisulfite sequencing of H19-DMR in Patient 2. Methylation of the maternal allele was higher in the cord blood than in the placenta or placental chorangioma. These results were consistent with the results of the Southern blot analysis. We confirmed complete methylation of paternal H19-DMR alleles and complete demethylation of maternal H19-DMR alleles in four normal control placentas that were heterozygous for identifiable SNPs (data not shown). Fa, father; Mo, mother; CB, cord blood; PB, peripheral blood; PI, placenta; PI-T, placental chorangioma; m, methylated band; um, unmethylated band; \*, nonspecific bands; Mat, maternal allele; Pat, paternal allele; CTS6, sixth CTCF binding site; M/U, a restriction site approximately 80 bp downstream of CTS6 assayed by methylation-sensitive Southern blot and COBRA.

#### **RESULTS**

We first examined the methylation status of the two ICRs, KvDMR1, and H19-DMR, at 11p15.5 using methylation-sensitive Southern blot analysis. Methylation at KvDMR1 was normal in all samples collected (Fig. 1A); however, methylation at H19-DMR was aberrant (Fig. 1B). In Patient 1, hypermethylation at H19-DMR was complete in cord blood and peripheral blood samples (MI = 100%), and hypermethylation in the placenta was partial (MI = 70%). In Patient 2, H19-DMR was partially hypermethylated in cord blood (MI = 90%) but less so in the placenta and placental chorangioma (MI = 71% and MI = 83%, respectively). For further investigation of differences in methylation between the patients' somatic tissues and placentas, the CTS6 site was subjected

to bisulfite sequencing (Fig. 1C and D). We could distinguish the two parental alleles in each patient sample using informative SNPs (rs10732516 and rs2071094). The maternal allele, which is normally unmethylated, was completely methylated in the cord blood, peripheral blood, and skin from Patient 1. However, in placental samples from Patient 1, the maternal allele was only partially methylated: 36% of all CpGs analyzed were methylated. Similar results were observed in Patient 2: the maternal allele in the cord blood was 68% methylated; however, the maternal allele was only 31% and 55% methylated in the placenta and chorangioma samples, respectively. The paternal alleles, which are normally fully methylated, were fully methylated in all samples. These findings supported the results of the Southern blots. Furthermore, we could not find any microdeletions or mutations in or around H19-DMR,

including seven CTCF-binding sites, three OCT4 sites, and one SOX2 site, indicating that there was no genetic cause of the hypermethylation (Fig. 2A and data not shown).

Next, we analyzed polymorphic markers at 11p15.4–p15.5 to determine whether copy number abnormalities or paternal UPD might be involved in these BWS patients. Although smaller PCR products were more easily amplified, paternal—maternal allele ratios in blood samples were between 0.92 and 1.33, indicating that both parental alleles were equally represented in both patients (Fig. 2B). Therefore, we could rule out copy number abnormality and paternal UPD within the patients' blood. We also investigated

maternal contamination in the placenta. *D11S1997* and *HUMTH01* for Patient 1 and the *RsaI* polymorphism in *H19* (rs2839703) for Patient 2 were used for this investigation because the mothers were expected to be homozygous for such polymorphisms. Thus, we investigated contamination of our samples by assessing the homozygosity of the polymorphisms in the mothers. The paternal—maternal ratios in Patient 1 were 0.94 and 1.03, indicating an equal contribution of both parental alleles and suggesting no contamination (Fig. 2B). In Patient 2, the ratios were 0.77 and 0.78 in the placenta and chorangioma, respectively, suggesting a small amount of contamination (Fig. 2C). However, such contamination was too

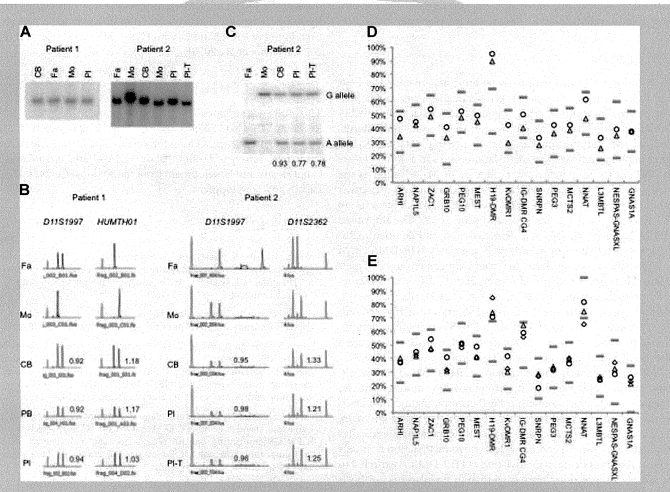


FIG. 2. Microdeletion analysis of H19-DMR, polymorphism analyses, and COBRA of primary imprinted DMRs in embryo-derived and placental samples. A: Southern blots identifying a microdeletion of H19-DMR. A genomic fragment (7.7 kb) generated by Apal digestion, which included the entire H19-DMR, was evident in all samples, indicating that there was no microdeletion in this DMR. B: Microsatellite markers at 11p15.4—p15.5. The peak heights associated with each parental allele in all samples were quantitatively analyzed. The results indicated that both parental alleles were present and equally represented. C: Hot-stop PCR of an Rsal polymorphic site in Patient 2. The ratios of paternal allele to maternal allele are shown under the figure. Although the ratios in the placenta and placental chorangioma are lower than in the cord blood, suggesting a small amount of maternal contamination, this was not enough to affect the results of the methylation analyses. COBRA of cord blood (D) and placentas (E), demonstrating that H19-DMR was hypermethylated. CTS6 is contained within H19-DMR. Methylation at other DMRs was normal in all samples, except for methylation at NNAT, which was aberrant in the placental chorangioma. Cord blood and placentas from 24 normal individuals were used as controls. The upper limit of normal methylation was defined as the higher of these two values: (1) the average of controls + 3 SD, or (2) the average + 15%. Similarly, the lower limit of normal methylation was definite as the lower of these two values: (1) the average of controls - 3 SD, or (2) the average - 15%. The upper and lower limits are indicated by gray bars.  $\bigcirc$ : Patient 1; Patient 2;  $\bigcirc$ : placental chorangioma of Patient 2.

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small to affect the results of the methylation analyses. In addition, sequence analysis did not show any mutations in *CDKN1C* (data not shown). These findings indicated that H19-DMR was aberrantly hypermethylated in both BWS patients and their associated placentas, but the aberrant methylation was consistently lower in the placenta, and that the H19-DMR GOM was strictly an isolated epimutation.

Finally, we analyzed the methylation status of 16 primary imprinted DMRs scattered throughout the genome-using COBRA (Fig. 2D and E). Only H19-DMR showed aberrant methylation among all primary DMRs in all samples, except for NNAT DMR, which was abnormal only in the placental chorangioma, indicating that the *IGF2/H19* imprinted domain was targeted for aberrant methylation in both somatic tissues and the placenta.

#### DISCUSSION

Methylation associated with parental imprints are erased in PGC and reestablished during gametogenesis in a sex-specific manner. The paternal pronucleus in the zygote undergoes active demethylation; extensive passive demethylation then ensues on maternal and paternal chromosomes during the pre-implantation period. After implantation, de novo methylation results in a rapid increase in DNA methylation in the inner cell mass (ICM), which gives rise to the entire embryo; in contrast, de novo methylation is either inhibited or not maintained in the trophoblast, which gives rise to the placenta [Li, 2002; Sasaki and Matsui, 2008]. The imprinted DMRs, however, escape these demethylation and de novo methylation events that occur in early embryogenesis. H19-DMR GOM in BWS patients is considered an error in imprint erasure in female PGCs [Horsthemke, 2010]. However, H19-DMR GOM, whether with or without microdeletions within H19-DMR, was partial, indicating a mosaic of normal cells and aberrantly methylated cells [Sparago et al., 2007; Cerrato et al., 2008]. These findings demonstrated that aberrant hypermethylation at H19-DMR was acquired after fertilization, although the precise timing was unknown.

Both participants in this study had isolated GOM at H19-DMR. The partial and variable hypermethylation among samples suggested epigenetic mosaicism. Furthermore, methylation levels in the placentas were lower than those in the blood and skin, suggesting that the aberrant methylation was acquired after implantation—when genome-wide de novo methylation normally occurs. Aberrant de novo methylation at H19-DMR is expected to be more widespread in the embryo than in the placenta, as this is normally the case for de novo methylation [Li, 2002; Sasaki and Matsui, 2008]; this disparity in efficiency could lead to the discordance between hypermethylation in trophoblast-derived placenta and that in embryo-derived blood and skin. This hypothesis is supported by a mouse experiment in which a mutant maternal allele harboring a deletion of four CTCF binding sites was hypomethylated in oocytes and blastocysts, yet was highly methylated after implantation [Engel et al., 2006]. To our knowledge, this is the first evidence demonstrating that aberrant hypermethylation of maternal H19-DMR is acquired after implantation in humans.

We found that of 16 primary imprinted DMRs analyzed, only H19-DMR showed aberrant methylation; even methylation at IG-DMR CG4, another paternally methylated, primary imprinted

DMR, was normal in our patients. Although we only studied two patients, this finding indicated that the *IGF2/H19* imprinted domain in both the embryo and placenta was more susceptible than other imprinted domains to aberrant methylation acquired after implantation.

In conclusion, we found that methylation of H19-DMR was discordant in embryo-derived somatic tissue and placenta, strongly suggesting that the aberrant de novo methylation occurred after implantation. However, the precise mechanism of isolated H19-DMR GOM is still unknown. Since no mutations in *CTCF*, an important trans-acting imprinting factor, were found in these patients with isolated GOM at H19-DMR, the potential for mutations in the OCT and SOX transcription factors should be investigated because mutations of OCT-binding sites have previously been found in a few patients with H19-DMR GOM [Cerrato et al., 2008; Demars et al., 2010].

#### **ACKNOWLEDGMENTS**

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#### REFERENCES

Aoki A, Shiozaki A, Sameshima A, Higashimoto K, Soejima H, Saito S. 2011. Beckwith—Wiedemann syndrome with placental chorangioma due to H19-differentially methylated region hypermethylation: A case report. J Obstet Gynaecol Res 37:1872—1876.

Bell AC, Felsenfeld G. 2000. Methylation of a CTCF-dependent boundary controls imprinted expression of the Igf2 gene. Nature 405:482–485.

Cerrato F, Sparago A, Verde G, De Crescenzo A, Citro V, Cubellis MV, Rinaldi MM, Boccuto L, Neri G, Magnani C, D'Angelo P, Collini P, Perotti D, Sebastio G, Maher ER, Riccio A. 2008. Different mechanisms cause imprinting defects at the IGF2/H19 locus in Beckwith—Wiedemann syndrome and Wilms' tumour. Hum Mol Genet 17:1427—1435.

Demars J, Shmela ME, Rossignol S, Okabe J, Netchine I, Azzi S, Cabrol S, Le Caignec C, David A, Le Bouc Y, El-Osta A, Gicquel C. 2010. Analysis of the IGF2/H19 imprinting control region uncovers new genetic defects, including mutations of OCT-binding sequences, in patients with 11p15 fetal growth disorders. Hum Mol Genet 19:803–814.

Elliott M, Bayly R, Cole T, Temple IK, Maher ER. 1994. Clinical features and natural history of Beckwith–Wiedemann syndrome: Presentation of 74 new cases. Clin Genet 46:168–174.

Engel N, Thorvaldsen JL, Bartolomei MS. 2006. CTCF binding sites promote transcription initiation and prevent DNA methylation on the maternal allele at the imprinted H19/Igf2 locus. Hum Mol Genet 15:2945–2954.

Hark AT, Schoenherr CJ, Katz DJ, Ingram RS, Levorse JM, Tilghman SM. 2000. CTCF mediates methylation-sensitive enhancer-blocking activity at the H19/Igf2 locus. Nature 405:486–489.

Horsthemke B. 2010. Mechanisms of imprint dysregulation. Am J Med Genet C Semin Med Genet 154C:321–328.

Li E. 2002. Chromatin modification and epigenetic reprogramming in mammalian development. Nat Rev Genet 3:662–673.

Sasaki H, Matsui Y. 2008. Epigenetic events in mammalian germ-cell development: Reprogramming and beyond. Nat Rev Genet 9:129–140.

Sasaki K, Soejima H, Higashimoto K, Yatsuki H, Ohashi H, Yakabe S, Joh K, Niikawa N, Mukai T. 2007. Japanese and North American/European patients with Beckwith—Wiedemann syndrome have different frequencies of some epigenetic and genetic alterations. Eur J Hum Genet 15: 1205–1210.

Soejima H, Nakagawachi T, Zhao W, Higashimoto K, Urano T, Matsukura S, Kitajima Y, Takeuchi M, Nakayama M, Oshimura M, Miyazaki K, Joh K, Mukai T. 2004. Silencing of imprinted CDKN1C gene expression is associated with loss of CpG and histone H3 lysine 9 methylation at DMR-LIT1 in esophageal cancer. Oncogene 23:4380–4388.

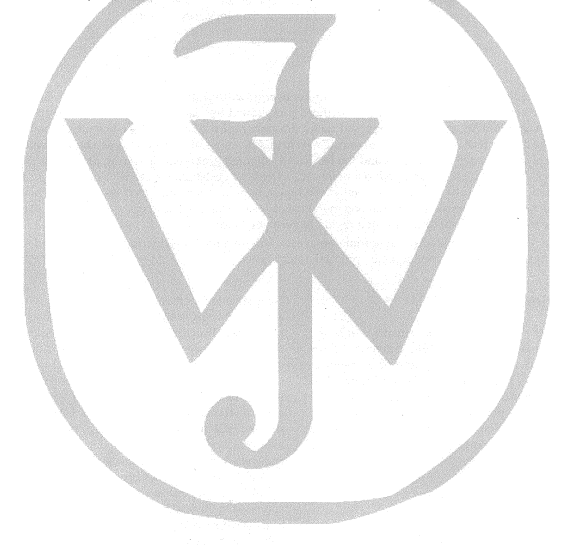
Sparago A, Cerrato F, Vernucci M, Ferrero GB, Silengo MC, Riccio A. 2004. Microdeletions in the human H19 DMR result in loss of IGF2 imprinting and Beckwith–Wiedemann syndrome. Nat Genet 36:958–960.

Sparago A, Russo S, Cerrato F, Ferraiuolo S, Castorina P, Selicorni A, Schwienbacher C, Negrini M, Ferrero GB, Silengo MC, Anichini C, Larizza L, Riccio A. 2007. Mechanisms causing imprinting defects in familial Beckwith–Wiedemann syndrome with Wilms' tumour. Hum Mol Genet 16:254–264.

Uejima H, Lee MP, Cui H, Feinberg AP. 2000. Hot-stop PCR: A simple and general assay for linear quantitation of allele ratios. Nat Genet 25: 375–376.

Weksberg R, Nishikawa J, Caluseriu O, Fei YL, Shuman C, Wei C, Steele L, Cameron J, Smith A, Ambus I, Li M, Ray PN, Sadowski P, Squire J. 2001. Tumor development in the Beckwith–Wiedemann syndrome is associated with a variety of constitutional molecular 11p15 alterations including imprinting defects of KCNQ1OT1. Hum Mol Genet 10:2989–3000.

Weksberg R, Shuman C, Beckwith JB. 2010. Beckwith-Wiedemann syndrome. Eur J Hum Genet 18:8–14.



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# Beckwith–Wiedemann syndrome with placental chorangioma due to H19-differentially methylated region hypermethylation: A case report

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#### **Abstract**

Beckwith–Wiedemann syndrome (BWS) is a common overgrowth syndrome that involves abdominal wall defects, macroglossia, and gigantism. It is sometimes complicated by placental tumor and polyhydramnios. We report a case of BWS, prenatally diagnosed with ultrasonography. A large and well-circumscribed tumor also existed on the fetal surface of the placenta, which was histologically diagnosed as chorangioma after delivery. Polyhydramnios was obvious and the fetal heart enlarged progressively during pregnancy. Because the biophysical profiling score dropped to 4 points at 33 weeks of gestation, we carried out cesarean section. By epigenetic analysis, H19-differentially methylated region hypermethylation was observed in the placental tumor, normal placental tissue, and cord blood mononuclear cells. This is the first report of BWS with placental tumor due to H19-differentially methylated region hypermethylation.

**Key words:** Beckwith–Wiedemann syndrome, epigenetic abnormality, H19-differentially methylated region, hypermethylation, placental chorangioma, prenatal diagnosis.

#### Introduction

The incidence of Beckwith–Wiedemann syndrome (BWS) is estimated to be 1 in 13 700 deliveries. BWS presents characteristic findings, genetic abnormalities, and a higher risk of malignancies. To our knowledge, there are only four reports of BWS with placental chorangioma, but no report was supported by epigenetic analysis. This is the first report of BWS with placental chorangioma, biallelic expression of *IGF2*, and reduced *H19* expression.

#### Case Report

A 27-year-old woman had an uneventful pregnancy (gravida 0, para 0) until 29 weeks of gestation. Assisted reproductive technology had not been performed. She

complained of increased abdominal circumference and a sense of oppression at 29 weeks and 3 days into her pregnancy. Transabdominal sonography showed that the fetus was large for the gestational age. Additionally, a large and well-circumscribed placental tumor measuring about 12 cm in diameter and polyhydramnios were observed. She was diagnosed with preterm labor and was transferred to our hospital. Ultrasonography showed fetal macroglossia (Fig. 1a), enlargement of the liver (Fig. 1b) and the kidneys (Fig. 1c,d), and a large and well-circumscribed placental tumor (Fig. 1e). These clinical physical features suggested a prenatal diagnosis of BWS. Mild mitral and tricuspid regurgitation appeared in the fetal heart at 32 weeks of gestation, and became worse at 33 weeks of gestation. At 33 weeks and 5 days of gestation, biophysical profiling score was dropped to 4 points, and therefore a cesarean section

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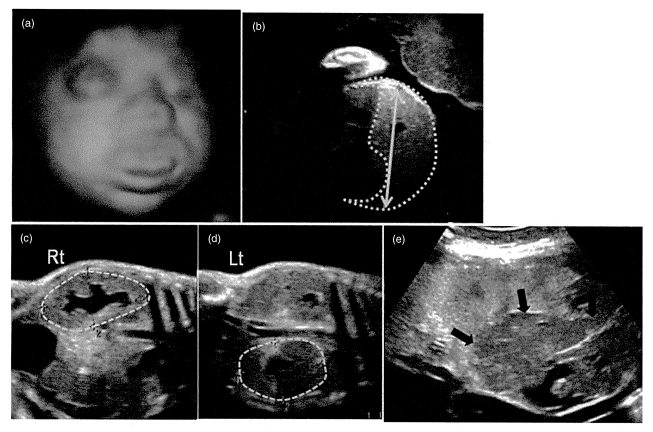
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**Figure 1** (a–e) Fetal and placental findings. Characteristic findings were seen using ultrasonography. (a) Macroglossia. (b) Hepatomegaly, measuring 87 mm (normal range at 29 weeks: 39 mm). (c,d) Enlarged kidneys, measuring 60 × 32 mm in right side, and 50 × 37 mm in the left side (normal range at 29 weeks: 28 × 15 mm). (e) A large, well-circumscribed tumor (12 cm in diameter) was found on fetal side of the placenta (arrows).

was performed for non-reassuring fetal status. The female baby weighed 2540 g and umbilical arterial blood pH was 7.127. Apgar scores were 3 and 7 points at 1 and 5 min, respectively. The baby had solitary, purplered focuses on the body, macroglossia, and distended abdomen. Blood test showed hyperleukocytosis (32 520/mm³), anemia (9.0 g/dL), thrombocytopenia  $(5.8 \times 10^4/\text{mm}^3)$ , and coagulopathy (Table 1). Blood transfusion was done for continuous anemia and low platelet count. For persistent neonatal hypoglycemia (around 20 mg/dL), steroid and glucose were used to keep the blood sugar level normal. Although mitral and tricuspid valve regurgitations existed, the cardiac function was stable. The fetal heart became progressively enlarged on the 8th postnatal day, and the baby suddenly died of cardiogenic shock on the 64th postnatal day (Fig. 2).

The weight of the placenta was 1620 g with a well-capsulated placental tumor measuring 12 cm on the

Table 1 Coagulopathy in the neonatal blood test

Prothrombin time	45.4 s
Prothrombin time %	<20
International normalized ratio	4.52
Activated partial thromboplastin time	>100 s
Fibrinogen	<50 mg/dL
Fibrin degenerative product	109.9 μg/mL
D-Dimer	$35.0  \mu g/mL$

fetal side of the placenta (Fig. 3a). Histological examination showed enlarged villi with an increased number of small blood vessels and the tumor was diagnosed with cellular placental chorangioma (Fig. 3b,c).

The epigenetic analysis after delivery of the cord blood, placenta, and a part of the placental tumor showed H19-differentially methylated region (DMR)

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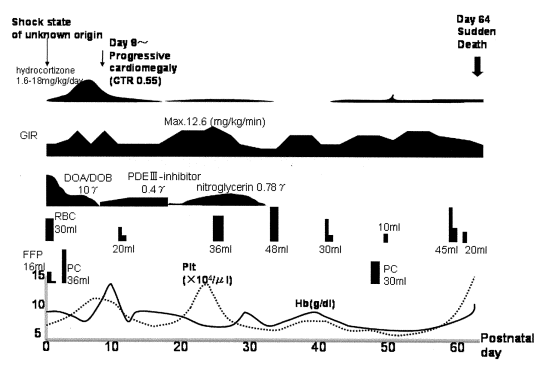
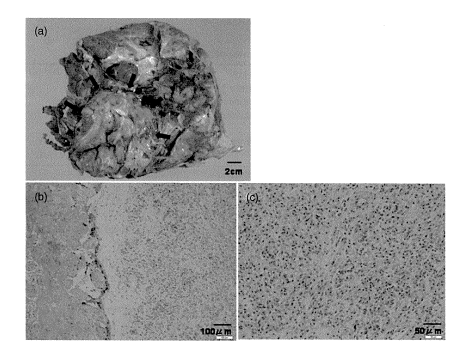


Figure 2 Neonatal clinical course. The baby went into a shock state 2 h after delivery, and was resuscitated with heart massage, cardiotonics and infusion of normal saline and sodium hydrogen carbonate. We performed a blood transfusion for continuous anemia and low platelet count. Because the glucose supply was not enough for persistent hypoglycemia, we administered steroids to keep the blood sugar level normal. The fetal heart was progressively enlarged on the 8th postnatal day, and the baby suddenly died of cardiogenic shock on the 64th postnatal day. CTR, cardiothoracic rate; DOA/DOB: dopamine/dobutamine; FFP, fresh frozen plasma; GIR, glucose infusion rate; Hb, hemoglobin; PC, platelet concentrates; PDE III-inhibitor, phosphodiesterase III inhibitor; Plt, platelet; RBC, red blood cell.

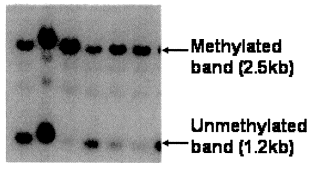


Figures 3 (a–c) Macroscopic and histological appearance of the placental tumor. (a) A well-circumscribed tumor (12.5 × 11.5 cm) was found on the fetal surface of the placenta (arrow). (b) Villi are enlarged and edematous (in the middle). (c) There is diffuse vascular proliferation in the enlarged villi, compatible with chorangioma.

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hypermethylation. The parental blood tests showed no epigenetic abnormalities (Fig. 4). Another candidate methylation alteration site in BWS, KvDMR1, was normally methylated (data not shown). Genetic analyses ruled out paternal uniparental disomy of 11p15.5 and mutation of CDKN1C (data not shown).

## Maternal peripheral blood Paternal peripheral blood Cord blood Maternal peripheral blood Normal placenta



#### 45 46 90 51 71 93 % methylated

Figure 4 Epigenetic analysis. Quantity of H19differentially methylated region (DMR) methylation status was determined by Southern blotting. Three micrograms of genomic DNA was digested with PstI and methylation-sensitive MluI. The blot was probed by the PCR product, which was generated with a primer pair: 5'-CTCCGACTCCGTCTAAGGACA-3' and 5'-GAGTGGAGACTGGCGAGTTTC-3'. KvDMR1 methylation status was also analyzed by Southern blotting with BamHI and methylation-sensitive NotI. The blot was probed as previously described.2 Band intensity obtained from Southern blotting was measured with a FLA-7000. The methylation index was calculated by (intensity of methylated band/[intensity of methylated band + intensity of unmethylated band]). H19-DMR hypermethylations are seen in the cord blood, normal placenta, and a part of the placental tumor (90%, 71%, 83%, respectively). Biallelic expression of Igf2 and suppression of H19 caused placentomegaly, macroglossia, visceromegaly and increased size for gestational age.

#### Discussion

Four BWS cases complicated with placental chorangioma have been reported,<sup>3-6</sup> but none was supported by epigenetic analysis. This is, to our knowledge, the first report proven to have epigenetic abnormality in BWS with placental chorangioma.

Chorangioma is asymptomatic when the size is smaller than 5 cm in diameter, but chorangiomas larger than 5 cm can cause some clinical problems, such as polyhydramnios, intrauterine fetal distress or death, fetal cardiac failure, neonatal anemia and thrombocytopenia, and disseminated intravascular coagulation, triggered by thromboplastic substances released from the small blood vessels in the chorangioma. In our case, the tumor size was 12 cm and the features described above were observed.

Genomic imprinting is an epigenetic modification that inactivates one allele of a gene in a parent-of-origindependent manner.8 Insulin-like growth factor 2 (IGF2)-H19 imprinting control region (ICR), consisting of a methylation-sensitive chromatin insulator on chromosome 11p15.5, is responsible for epigenetic malformations in BWS.9 In the *Igf2-H19* domain, especially in the placenta and tissues of endodermal origin, such as the liver, allele-specific gene expressions of parental origin are regulated by ICR located 2-4 kb upstream of the H19 promoters. This region functions as a methylationsensitive insulator that binds to CCCTC-binding factor (CTCF) on the unmethylated maternal allele. Alternatively, on the paternal allele, DNA methylation of ICR prevents CTCF binding, which permits access of the downstream enhancers to paternal Igf2 promoters.8

In BWS cases, which have H19-DMR methylations in both paternal and maternal alleles (H19-DMR hypermethylation), the enhancers on the maternal allele activates IGF2 promoters, leading to biallelic IGF2 expression and the maternal copy of the H19 gene is silenced because of expanded methylation to its promoter. Expression of imprinting gene, IGF2, depends on whether ICR is methylated or not, and loss of imprinting (LOI) of IGF2 is associated with downregulation of H19 expression, as previously reported.10-12 Using epigenetic analysis, we could prove the existence of H19-DMR hypermethylation in the cord blood, placenta, and the placental tumor. Although we could not show biallelic IGF2 expression because of homozygosity for IGF2 polymorphism in the case (data not shown), H19-DMR hypermethylation would result in biallelic IGF2 expression and associated reduced expression of H19 due to the

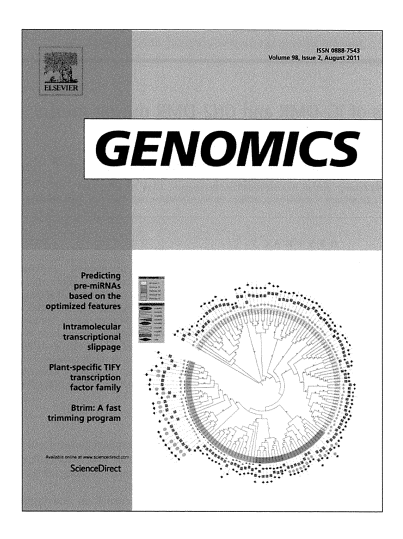
© 2011 The Authors Journal of Obstetrics and Gynaecology Research © 2011 Japan Society of Obstetrics and Gynaecology reason mentioned above. In view of the potential function of *IGF2-H19*, these conditions led to cell proliferation and caused clinical appearances, such as macroglossia, visceromegaly (increased abdominal circumference), and the placental tumor. We explained to the mother that an occasional epigenetic abnormality (H19-DMR hypermethylation) caused BWS in the baby and that it would be extremely rare to recur in the next pregnancy.

#### References

- Thorburn MJ, Wright ES, Miller CG, Smith-Read EH. Exomphalos-macroglossia-gigantism syndrome in Jamaican infants. Am J Dis Child 1970; 119: 316–321.
- Mitsuya K, Meguro M, Lee MP et al. LIT1, an imprinted antisense RNA in the human KvLQT1 locus identified by screening for differentially expressed transcripts using monochromosomal hybrids. Hum Mol Genet 1999; 8: 1209– 1217
- Lage JM. Placentomegaly with massive hydrops of placental stem villi, diploid DNA content, and fetal omphaloceles: possible association with Beckwith-Wiedemann syndrome. *Hum Pathol* 1991; 22: 591–597.
- Drut RM, Drut R. Nonimmune fetal hydrops and placentomegaly: diagnosis of familial Wiedemann-Beckwith syn-

- drome with trisomy 11p15 using FISH. *Am J Med Genet* 1996; **62**: 145–149.
- 5. Takayama M, Soma H, Yaguchi S *et al*. Abnormally large placenta associated with Beckwith–Wiedemann syndrome. *Gynecol Obstet Invest* 1986; **22**: 165–168.
- Drut R, Drut RM, Toulouse JC. Hepatic hemangioendotheliomas, placental chorioangiomas, and dysmorphic kidneys in Beckwith-Wiedemann syndrome. *Pediatr Pathol* 1992; 12: 197–203.
- Fox H. Non-trophoblastic tumors of the placenta. In: Fox H (ed.). Obstetrical and Gynecological Pathology, 3rd edn. Philadelphia: Churchill Livingstone, 1987; 1030–1044.
- Fowden AL, Sibley C, Reik W, Constancia M. Imprinted genes, placental development and fetal growth. *Horm Res* 2006; 65 (Suppl 3): 50–58.
- Choufani S, Shuman C, Weksberg R. Beckwith-Wiedemann syndrome. Am J Med Genet C Semin Med Genet 2010 154C: 343–354.
- Steenman MJ, Rainier S, Dobry CJ, Grundy P, Horon IL, Feinberg AP. Loss of imprinting of *IGF2* is linked to reduced expression and abnormal methylation of *H19* in Wilms' tumour. *Nat Genet* 1994; 7: 433–439.
- 11. Bell AC, Felsenfeld G. Methylation of a CTCF-dependent boundary controls imprinted expression of the *Igf*2 gene. *Nature* 2000; **405**: 482–485.
- Hark AT, Schoenherr CJ, Katz DJ, Ingram RS, Levorse JM, Tilghman SM. CTCF mediates methylation-sensitive enhancer-blocking activity at the H19/Igf2 locus. Nature 2000; 405: 486–489.

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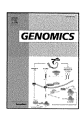
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# Methylation dynamics of IG-DMR and *Gtl2*-DMR during murine embryonic and placental development

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#### ABSTRACT

The *Dlk1-Dio3* imprinted domain on mouse chromosome 12 contains IG-DMR and *Gtl2*-DMR, whose methylation patterns are established in the germline and after fertilization, respectively. In this study, we determine that acquisition of DNA methylation at the paternal allele of the *Gtl2*-DMR is initiated after the blastocyst stage and completed by embryonic day 6.5, and that *Gtl2* (approved symbol: Meg3) is monoallelically expressed from the maternal allele as early as the blastocyst. Therefore, DNA methylation at the *Gtl2*-DMR is not a prerequisite for the imprinted expression of *Gtl2*, which may be involved in the control of proliferation and differentiation of cells during early gestation. We also reveal that a subregion of the IG-DMR exhibits tissue-specific differences in allelic methylation patterns. These results add to the growing body of knowledge elucidating the mechanism whereby parent-of-origin-dependent DNA methylation at the IG-DMR leads to the imprinted expression of the *Dlk1-Dio3* cluster.

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#### 1. Introduction

Genomic imprinting is an epigenetic mechanism that regulates transcription, whereby the expression of a subset of genes is limited to or biased towards one parental allele. To date, over one hundred imprinted genes have been identified in the mouse (http://www.har. mrc.ac.uk/research/genomic\_imprinting). Imprinted genes tend to be clustered on the genome. One of the common features among imprinted loci is that such genomic intervals include one or more differentially methylated regions (DMRs), which exhibit parent-oforigin dependent DNA methylation patterns [1]. DMRs have been classified into two types according to the time at which their DNA methylation patterns are established. Primary (germline) DMRs harbor allelic DNA methylation inherited from the male or the female gamete. Secondary (post-zygotic) DMRs acquire parent-of-origin dependent methylation patterns after fertilization. In mice, germline DMRs are shown to be established during the oocyte growth stage (postnatal days 5 to 20) [2,3] or the prospermatogonia stage (embryonic days 14.5 to newborn) [4-6] by a DNMT3L-dependent mechanism [7-12]. Several germline DMRs have been shown to govern the imprinted expression of genes as well as the methylation of post-zygotic DMRs within chromosomal regions. These germline DMRs, known as imprinting control regions (ICRs), regulate these regions by cis-acting mechanisms. [13-15]. On

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the other hand, little is known about the function of secondary DMRs in the regulation of imprinted gene expression, as well as *cis*-acting mechanisms and *trans*-acting factors that establish DNA methylation at secondary DMRs.

Studies focusing on the regulatory functions of ICRs have revealed a number of different molecular mechanisms that underlie the coordinated and long-range regulation of imprinted genes. In the H19/Igf2 domain, long range chromatin interactions mediated by CTCF between the primary H19-DMR and the secondary Igf2-DMRs play an integral role in the regulation of imprinted gene expression at this locus [13,16]. Another mechanism involves non-coding (nc) RNAs such as Airn in the Igf2r locus and Kcnq1ot1 in the Kcnq1 imprinted gene cluster. These ncRNAs are transcribed from ICRs and are shown to be functionally linked to the silencing of genes in cis through gene- and lineage-specific repressive chromatin modifications [14,17,18]. These two mechanisms are likely to be involved in the regulation of many other imprinted loci as well. However, the sequence of events leading to the establishment and maintenance of imprinted expression for a cluster of genes remains largely elusive for many imprinted loci.

The *Dlk1-Dio3* imprinting cluster on mouse distal chromosome 12 contains the intergenic germline-derived DMR (IG-DMR) and the *Gtl2-DMR*, whose methylation patterns are established in the germline and after fertilization, respectively [19,20]. The cluster consists of at least three paternally expressed protein-coding genes (*Dlk1*, *Rtl1*, and *Dio3*), and four maternally expressed ncRNAs (*Gtl2*, *Anti-Rtl1*, *Rian* and *Mirg*). The IG-DMR is shown to function as the ICR of this imprinted gene cluster [15,21]. A targeted disruption study of the IG-

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DMR [15] has revealed that the maternally inherited IG-DMR, which is unmethylated, is essential in the embryo to maintain the unmethylated status of the *Gtl2*-DMR, the expression of the ncRNAs, and the repression of the protein-coding genes, on the maternal allele. However, the principal mechanism whereby the allele-specific methylation at the IG-DMR leads to the imprinted expression of the cluster of genes on chromosome 12 is unknown. It has been also demonstrated that, in the placenta, the absence of the maternally inherited IG-DMR results in the activation of protein-coding genes but only partial repression of the ncRNAs, and leads to no phenotypic consequence [21]. Therefore, mechanisms underlying the imprinted expression of the maternally-expressed ncRNAs are different between the embryonic and the extra-embryonic tissue lineages.

Among known secondary DMRs, the Gtl2-DMR is unique in that it has been demonstrated to possess an essential long-range imprinting regulatory function. A neonatal patient showing a paternal uniparental disomy 14-like phenotype in the body but not in the placenta was identified to have a maternally-inherited heterozygous microdeletion that encompasses the MEG3-DMR (the human orthologue of the mouse Gtl2-DMR) but not the IG-DMR. In this patient, the maternal allele of DLK1 has been shown to be reactivated [22]. Recent studies have used knockout mouse models with targeted deletions of the Gtl2 locus, spanning the Gtl2-DMR. These studies have also suggested that Gtl2 and/or Gtl2-DMR could regulate the expression of maternally expressed genes, indicating that the methylation of the Gtl2-DMR is a critical element in the Dlk1-Dio3 imprinted domain [23,24]. In light of the critical roles that Gtl2 and Gtl2-DMR may play in the imprinted regulation of this region, understanding the epigenetic mechanisms that govern them during early development is expected to further elucidate the mechanisms regulating the Dlk1-Dio3 imprinted domain.

Recently, Stadtfeld et al. [25] reported that mouse induced pluripotent stem cells (iPSC) with repressed expression of maternally expressed ncRNAs in the *Dlk1-Dio3* domain contributed poorly to chimeras and failed to generate all-iPSC mice. In contrast, iPSCs with normal ncRNA expression patterns contributed to high-grade chimeras and produced all-iPSC mice. Hypermethylation of both the IG-DMR and the *Gtl2-DMR* was found to be associated with the reduced expression of ncRNAs in the iPSCs exhibiting poor contribution to chimeras [25]. This epimutation is considered to be caused by the iPSC reprogramming, rather than existing aberrant methylation patterns in the DMRs of the somatic cell of origin [25]. Therefore, a better understanding of the epigenetic regulation of these DMRs may eventually lead to improved reprogramming strategies of iPSC.

In this study, we determined the allelic DNA methylation patterns at the IG-DMR and the *Gtl2*-DMR, as well as the allelic expression patterns of *Dlk1* and *Gtl2* at early developmental stages (embryonic days 3.5 to 7.5) in embryonic and extra-embryonic tissues.

#### 2. Results

2.1. Developmental dynamics of allelic DNA methylation patterns at IG- and Gtl2-DMRs in sperm, blastocysts, and post-implantation embryos

We examined allelic DNA methylation patterns at the IG-DMR and the *Gtl2*-DMR in whole embryos at embryonic day 3.5 (E3.5) and E5.5 as well as their methylation status in sperm. We regarded the genomic intervals defined by Kobayashi et al. [26] and Takada et al. [20] as the IG-DMR and the *Gtl2*-DMR, respectively. Three regions within the IG-DMR and the two regions within the *Gtl2*-DMR were chosen as targets for bisulfite sequencing (Fig. 1A). All five regions contain at least one single nucleotide polymorphism (SNP) between C57BL/6 (B6) and JF1/Ms (JF1) strains that can distinguish parental alleles in F1 hybrid materials (see details in Section 4.2 in the Materials and methods).

The IG-DMR was heavily methylated in all three regions (methylation percentage 81.3-95.8%) in sperm (Fig. 1B) as shown previously [20]. In blastocysts (E3.5), all three regions within the IG-DMR were maternally unmethylated (0-1.8%), yet paternally methylated (43.1-71.8%) (Fig. 1B). The observed levels of paternal methylation at E3.5 were significantly lower than those observed in sperm, implying that the paternal IG-DMR partially loses methylation at CpG dinucleotides after fertilization. This loss of methylation may be caused by the active and the passive demethylation of the paternal genome in pronucleus and preimplantation embryos, respectively [27,28]. At E5.5, the maternal allele of IG-DMR was found to be hypomethylated (1.9-18.3%), and the paternal allele to be hypermethylated (80.2-91.4%; Fig. 1B). The methylation level of the paternal allele was consistently higher at E5.5 than at E3.5. Additionally, it was almost fully methylated at E5.5 in all three regions examined, suggesting that de novo methylation events occur on the paternal allele of the IG-DMR during the developmental period between E3.5 and E5.5. These are the first results to illustrate the developmental dynamics of paternal methylation levels at the IG-DMR around the implantation period.

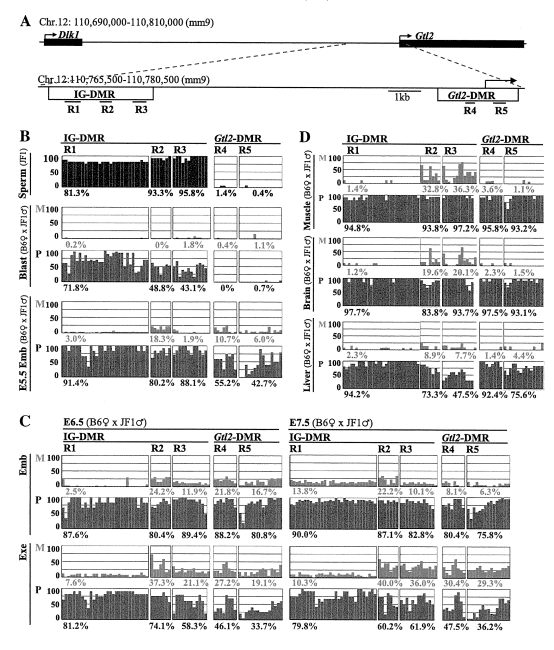
The differential methylation of the Gtl2-DMR on the paternal allele has been shown to be established in E13.5 embryos [20]. However, the post-zygotic stages at which the region's paternal methylation is initiated and completed remain unknown. Additionally, the relationship between the imprinted expression of Gtl2 and DNA methylation at the Gtl2-DMR has not been elucidated. We confirmed that the Gtl2-DMR was unmethylated in sperm, and found that it was unmethylated on both parental alleles in blastocysts (Fig. 1B). In E5.5 embryos, the maternal allele remained hypomethylated (6.0 and 11.7%), while the paternal allele became partially methylated (55.2% in R4 and 42.7% in R5 regions) (Fig. 1B). In E6.5 and E7.5 embryos, the paternal allele of the Gtl2-DMR was found to be heavily methylated (75.8% or higher) (Fig. 1C). These data demonstrate that paternal methylation of the Gtl2-DMR is initiated after the blastocyst stage and is completed by E6.5 stage in the embryonic lineage.

2.2. Allelic DNA methylation patterns at IG- and Gtl2-DMRs in early and late gestational stages

It has been reported that, in both human and mouse placenta, the IG-DMR maintains its allele-specific methylation patterns, whereas the Gtl2-DMR does not show differential methylation between parental alleles [21,29,30]. To determine the developmental stage at which the allelic methylation patterns at the Gtl2-DMR diverge between embryonic and extra-embryonic lineages, we examined the DNA methylation status of the Gtl2-DMR as well as the IG-DMR in E6.5 and E7.5 tissues. In extra-embryonic tissues at both E6.5 and E7.5 stages, the Gtl2-DMR was partially methylated on both parental alleles, whereas its differential methylation was well maintained in embryonic tissues (Fig. 1C). The Gtl2-DMR was previously shown to be partially methylated on both parental alleles in late gestation (E16.5) placentas [21,29]. Our results demonstrate that the allelic methylation pattern observed in E16.5 placenta is already present in the extraembryonic lineage at E6.5 stage.

Unexpectedly, we observed loss of differential methylation at the R2 and R3 regions within the IG-DMR in extra-embryonic tissues, although the R1 region maintained its differential methylation. The loss of differential methylation was more evident in E7.5 stage than in E6.5 stage (Fig. 1C). To assess whether the loss of differential methylation at the R2/R3 regions as well as the R4/R5 regions was specific to the extra-embryonic lineage, we examined the allelic methylation patterns of the R1–R5 regions in fetal tissues from late gestation time points. E16.5 skeletal muscle, E15.5 brain, and E16.5 liver were analyzed since they represent tissues derived from the

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**Fig. 1.** Allelic DNA methylation patterns at the IG-DMR and the *Gtl2*-DMR during embryonic and extra-embryonic development. (A) A schematic diagram of the locus containing the IG-DMR and the *Gtl2*-DMR (shown as open boxes). The bars under the open boxes indicate the regions (R1-R5) analyzed by bisulfite sequencing. The arrow indicates the transcription start site of *Gtl2*. Scale bar = 1 kb. (B-D) Graphical representation of the methylation percentage at each CpG site in sperm, blastocysts at E3.5 (Blast), and embryos at E5.5 (E5.5\_Emb) (B), in embryonic (Emb) and extra-embryonic (Exe) tissues at E6.5 and E7.5 (C), and in fetal tissues (E16.5 skeletal muscle, E15.5 brain, and E16.5 liver) (D). The vertical bars represent the percentage ratio of methylated cytosine at each CpG site, which were determined from the data of clone-based bisulfite sequencing (Supplementary Fig. 1). Overall methylation percentage for each region (the number of methylated CpGs per the number of total CpGs) is shown under each panel. As described in the Materials and Methods (Section 4.2), methylation percentage for each CpG site and each region was calculated using bisulfite sequencing data for a single sample (sperm and E15.5/16.5 fetal tissues) or two independent samples (E3.5 to E7.5 samples). M and P denote maternal and paternal alleles, respectively.

mesoderm, the ectoderm, and the endoderm, respectively. We found that differential methylation at the R1 region of the IG-DMR and the R4/R5 region of the *Gtl2*-DMR was strictly conserved. In contrast, differential methylation at the R2/R3 regions of the IG-DMR was partially lost to varying degrees in these tissues (Fig. 1D). Partial gain of methylation on the maternal allele (most notably observed in skeletal muscle) and partial loss of methylation on the paternal allele (most remarkably observed in the liver) were detected. Taken together, our data demonstrate that only a subregion of the IG-DMR containing the R1 region strongly maintains allele-specific differential methylation during embryonic development, whereas the rest of region containing the R2/R3 regions exhibits various tissue-specific allelic methylation patterns.

## 2.3. Allelic expression patterns of Dlk1 and Gtl2 during embryonic and extra-embryonic development

We subsequently assessed the expression levels and the allelic expression patterns of *Dlk1* and *Gtl2*. Although the expression of *Gtl2* is shown to be detectable as early as the pre-implantation stage [31], previous studies have not assessed the expression levels of these genes in a quantitative manner and have not determined their allelic expression patterns during early gestation (E3.5 to 7.5). Therefore, we performed both quantitative RT-PCR and pyrosequencing to quantify the allelic expression of these transcripts.

To determine the relative expression levels of Gtl2 and Dlk1, quantitative RT-PCR was performed using E3.5 to E7.5 tissues and