Epigenotype-phenotype correlations in Beckwith-Wiedemann syndrome


Abstract

Beckwith-Wiedemann syndrome (BWS) is a model imprinting disorder resulting from mutations or epigenetic events involving imprinted genes at chromosome 11p15.5. Thus, germline mutations in \textit{CDKN1C}, uniparental disomy (UPD), and loss of imprinting of \textit{IGF2} and other imprinted genes have been implicated. Many familial BWS cases have germline \textit{CDKN1C} mutations. However, most BWS cases are sporadic and UPD or putative imprinting errors predominate in this group. We have identified previously a subgroup of sporadic cases with loss of imprinting (LOI) of \textit{IGF2} and epigenetic silencing of \textit{H19} proposed to be caused by a defect in a distal 11p15.5 imprinting control element (designated BWSIC1). However, many sporadic BWS patients show biallelic \textit{IGF2} expression in the presence of normal \textit{H19} methylation and expression patterns. This and other evidence suggested the existence of a further imprinting control element (BWSIC2) at 11p15.5. Recently, we showed that a subgroup of BWS patients have loss of methylation (LOM) at a differentially methylated region (KvDMR1) within the \textit{KCQNI} gene centromeric to the \textit{IGF2} and \textit{H19} genes. We have now analysed a large series of sporadic cases to define the frequency and phenotypic correlates of epigenetic abnormalities in BWS. LOM at KvDMR1 was detected by Southern analysis or a novel PCR based method in 35 of 69 (51%) sporadic BWS without UPD. LOM at KvDMR1 was often, but not invariably associated with LOI of \textit{IGF2}. KvDMR1 LOM was not detected in BWS patients with putative BWSIC1 defects and cases with KvDMR1 LOM (that is, putative BWSIC2 defects) invariably had a normal \textit{H19} methylation pattern. The incidence of exomphalos in putative BWSIC2 defect patients was not significantly different from that in patients with germline \textit{CDKN1C} mutations (20/29 and 13/15 respectively), but was significantly greater than that in patients with putative BWSIC1 defects (0/5, p=0.007) and UPD (0/22, p<0.0001). These findings are consistent with the hypothesis that LOM of \textit{CDKN1C} and \textit{H19} results in biallelic \textit{IGF2} expression. BWS patients with embryonal tumours have UPD or a BWSIC1 defect but not LOM of KvDMR1.

This study has further shown how (1) variations in phenotypic expression of BWS may be linked to specific molecular subgroups and (2) molecular analysis of BWS can provide insights into mechanisms of imprinting regulation.

Keywords: Beckwith-Wiedemann syndrome; epigenotype-phenotype correlations; imprinting

Beckwith-Wiedemann syndrome is a prototypic imprinting disorder resulting from mutations or epigenetic alterations (epimutations) in genes from an imprinted cluster at chromosome 11p15.5. \textsuperscript{1,2} Phenotypic expression of BWS is variable, but the major features are pre-/postnatal overgrowth, macroglossia, and abdominal wall defects. Additional features include organomegaly, hypoglycaemia, hypertrophy, genitourinary abnormalities, and a susceptibility to embryonic tumours which occur in 5% of patients. The genetics of BWS are complex, \textsuperscript{3} but three main groups are distinguished: \textsuperscript{4} ~15% have familial BWS, ~2% have cytogenetic abnormalities (duplications or balanced rearrangements) involving chromosome 11p15.5, and the rest are sporadic. Imprinted genes have been implicated in the pathogenesis of both familial and sporadic BWS.

Genomic imprinting is an epigenetic mechanism in which gene expression is altered according to the parental origin of the allele. It is estimated that the human genome may contain between 100 and 1000 imprinted genes and most of those identified to date are arranged in clusters. A number of imprinted genes from chromosome 11p15.5 have been implicated in the pathogenesis of BWS, including the maternally expressed \textit{CDKN1C}, \textit{KCQNI}, and \textit{H19} genes and the paternally expressed \textit{IGF2} and \textit{KCQNI} genes. The molecular pathology of familial and sporadic BWS differs. Thus, although germline mutations in the cyclin dependent kinase inhibitor gene \textit{CDKN1C} may be detected in \textgreater 40% of familial cases, \textit{CDKN1C} mutations are infrequent (~5%) in sporadic cases. \textsuperscript{4-9} Among sporadic cases, ~20% have uniparental disomy (mosaic paternal isodisomy) for the 11p15.5 imprinted gene cluster. \textsuperscript{10-12} In addition, many sporadic cases show loss of imprinting (LOI) of \textit{IGF2} resulting in biallelic expression. \textsuperscript{13-15} The epigenetic alterations associated with \textit{IGF2} LOI are heterogeneous. Thus, we described initially a subgroup of sporadic patients with biallelic \textit{IGF2} expression.
silencing of $H19$ expression in which there was $H19$ hypermethylation such that the maternal $H19$ allele showed a paternal epigenotype with $H19$ promoter methylation. Subsequently, we showed that in many sporadic cases $IGF2$ LOI occurs via an $H19$ independent mechanism. These findings suggested that chromosome 11p15.5 contained two imprinting control centres/elements termed BWSIC1 and BWSIC2. Mutations or epimutations at BWSIC1 would result in $IGF2$ LOI and $H19$ silencing, while BWSIC2 defects would be associated with $IGF2$ LOI through an $H19$ independent mechanism.

Investigations of the recipricosity of $IGF2$ and $H19$ imprinting in mice suggested that BWSIC1 was likely to map to the $H19$ upstream region. The finding of $H19$ independent LOI of $IGF2$ in a BWS patient with a chromosomal rearrangement disrupting $KCNQ1$ suggested that BWSIC2 mapped centromeric to $IGF2$ close to (or within) $KCNQ1$. Recently, a differentially methylated region (KvDMR1) within $KCNQ1$ was identified. KvDMR1 shows maternal allelic methylation and the unmethylated paternal allele expresses an antisense RNA ($KCNQ1OT$) (also known as LIT1). We observed that some BWS patients showed loss of methylation at KvDMR1 and (for those that were informative) this was associated with $H19$ independent $IGF2$ LOI (consistent with a BWSIC2 defect).

Previously, we reported significant correlations between the phenotypic variability and molecular pathology in BWS. Thus, exomaphos was not found in BWS patients with UPD or BWSIC1 defects, but was present in the majority of patients with germline $CDKN1C$ mutations. We now report further investigations to define the role of LOM of KvDMR1 (BWSIC2 defects) in sporadic BWS and to elucidate epimutation-phenotype correlations.

**Patients and methods**

**BWS patients**

A diagnosis of BWS was made according to our standard diagnostic criteria; either (1) three major features (anterior abdominal wall defects, macroglossia, and pre- or postnatal growth >90th centile) or (2) two major plus three or more minor features (characteristic ear signs (ear lobe creases or posterior helical ear pits), facial naevus flammeus, hypoglycaemia, nephromegaly, and hemihypertrophy). Patients with normal chromosomes and no evidence of UPD were analysed for KvDMR1 methylation status and germline $CDKN1C$ mutations as described below. Uniparental disomy analysis and $H19$ methylation analysis was performed as reported previously. However, for many BWS patients insufficient DNA was available for Southern analysis and a PCR based approach was developed to identify patients with loss of KvDMR1 methylation. DNA (500 ng) (isolated from whole blood) was digested overnight with the methylation sensitive enzyme NotI (New England Biolabs) according to the manufacturer’s instructions. It was then ethanol precipitated, resuspended in 5 µl of sterile distilled water, and the whole sample subjected to multiplex PCR as set out below. A total of 100 ng of undigested DNA for each sample was PCR amplified in exactly the same way at the same time to act as a normal control for comparison with the digested sample.

Multiplex PCR was carried out using two sets of primers, which were designed using sequence information from PAC DJ74K15. Set 1 crosses the methylation sensitive NotI site, KvLF1-FAM-AGACCTTGGCCCGGT TT, KvLR1- AAAGTGATGTCCGGTG CCT and set 2 acted as a control reaction amplifying an upstream region, KvLF2-HEX-CACCATCTGACCTTATGGAG, KvLR2-CCAGGATGTGGACCTAGAC. The forward primer for each set was tagged with a different fluorescent dye.

Thirty µl PCR reactions were set up containing 6.6 pmol of each primer, 2 µl dNTPs (2.5 mmol/l), 1.5 mmol/l Mg, 10% DMSO, buffer, and 0.2 µl Taq DNA polymerase. Reactions were overlaid with mineral oil and cycled at the following conditions with the hot lid on: 95°C for 10 minutes once (this step allows for an eight minute hotstart), 94°C, 66°C, 72°C for 30 seconds each ×3, 94°C, 64°C, 72°C for 30 seconds each ×3, 94°C, 62°C, 72°C for 30 seconds each ×3, 94°C, 56°C, 72°C for 30 seconds each ×20.

All samples were separated and analysed using the Genescan software (ABI). Values for the peak heights for each PCR product were taken and used to calculate a single ratio. This ratio measures the reduction in methylation sensitive PCR product after digestion, and is therefore an indication of the levels of methylation in the sample. A methylation ratio of 0.1 or less was chosen as the value indicative of loss of methylation at this site. Patient samples that fell into this category were verified by repetition; if the low reading could not be repeated twice in three tests it was disregarded.

**CDKN1C mutation analysis**

We have previously reported $CDKN1C$ mutation analysis in 54 sporadic BWS patients. To define the molecular pathology of BWS further, we extended this analysis by analysing a further 30 sporadic cases using the methods described previously.

**MOLECULAR GENETIC ANALYSIS**

**KvDMR1 methylation analysis**

Two approaches were used to determine methylation status at KvDMR1. Firstly, Southern analysis was performed on genomic DNA isolated from whole blood as described previously. Southern analysis was performed in nine new cases (eight further cases have been reported previously). However, for many BWS patients insufficient DNA was available for Southern analysis and a PCR based approach was developed to identify patients with loss of KvDMR1 methylation. DNA (500 ng) (isolated from whole blood) was digested overnight with the methylation sensitive enzyme NotI (New England Biolabs) according to the manufacturer’s instructions. It was then ethanol precipitated, resuspended in 5 µl of sterile distilled water, and the whole sample subjected to multiplex PCR as set out below. A total of 100 ng of undigested DNA for each sample was PCR amplified in exactly the same way at the same time to act as a normal control for comparison with the digested sample.

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ported previously, and (3) 15 patients with germline \textit{CDKN1C} mutation (13 of whom were reported previously and two identified in this study). Statistical comparisons were made using Fisher’s exact test and statistical significance was set at 0.05.

\section*{Results}

\subsection*{KvDMR1 Methylation Analysis}

KvDMR methylation status was examined by Southern analysis or a novel PCR based assay or both in 69 patients referred with BWS in whom a cytogenetic rearrangement and uniparental disomy (UPD) had been excluded. Southern analysis was performed in 17 patients (eight reported previously). Loss of KvDMR1 methylation was detected in eight cases (three reported previously). However, in many further cases insufficient DNA was available to perform KvDMR1 methylation analysis using this approach. We therefore developed a novel PCR based technique to detect patients with LOM at KvDMR1 (see Methods).

To validate our PCR based strategy for detecting KvDMR1 LOM, we analysed DNA from 66 normal controls and confirmed that none gave a positive result. We then analysed 67 patients with confirmed or suspected BWS using the PCR based assay. A positive result (indicating loss of KvDMR1 methylation) was obtained in 35 cases (fig 1). KvDMR1 methylation status was analysed by both Southern and PCR based strategies in 15 patients and there was complete concordance between the two approaches.

\subsection*{Loss of Methylation at KvDMR1 and IGF2/H19 Epigenotype}

Thirty three of 69 patients analysed for KvDMR1 methylation by Southern or PCR based techniques had previously been analysed for \textit{H19} methylation status. All of the 23 patients with KvDMR1 loss of methylation in whom \textit{H19} methylation status had been examined showed normal \textit{H19} methylation. Conversely, four patients with \textit{H19} hypermethylation (BWSIC1 defect) displayed normal KvDMR1 methylation. In a previous publication, we reported that the two patients with KvDMR1 demethylation in whom \textit{IGF2} imprinting status had been defined both showed biallelic \textit{IGF2} expression. However, while one patient did show loss of \textit{IGF2} imprinting, the other retained monoallelic \textit{IGF2} expression.

\subsection*{Germline \textit{CDKN1C} Mutation Analysis}

To provide further information on the frequency of germline \textit{CDKN1C} mutations in sporadic cases and to enhance genotype-phenotype comparisons (see later), we analysed a further 30 sporadic cases. Germline \textit{CDKN1C} mutations were identified in two cases (fig 2).

In case 1, a frameshift insertion of T at nucleotide 1618 causes a sequence frameshift. Case 2: mutation 2. T to G transversion at nucleotide 1325 produces a non-conservative leucine to arginine substitution at codon 33.
PAPA repeat and PCNA binding domains. The mutation was not detected in paternal DNA and appeared to have arisen de novo. In case 2, a T to G transversion at nucleotide 1325 caused a non-conservative substitution of leucine to arginine at codon 33 within the cyclin binding domain. This residue is conserved in the mouse cdkn1c gene. The mutation was also present in the mother but not present in paternal DNA.

**Genotype-Phenotype Correlations**

To determine possible genotype-phenotype correlations, we compared the frequency of exomphalos and embryonal tumours in patients with KvDMR1 methylation to that in other well defined molecular subgroups of BWS (see Methods). Of 29 BWS patients with LOM of KvDMR1 for which sufficient clinical information was available, 20 of 29 (69%) cases had exomphalos. Compared to other molecular subgroups, the frequency of exomphalos in patients with LOM of KvDMR1 (BWSIC2 defect) was significantly greater than for patients with BWSIC1 defects (20/29 versus 0/5, p=0.007) or UPD (20/29 vs 0/22, p=0.0001), but was similar to that in patients with germline CDKN1C mutations (20/29 vs 13/15, p=0.28). None of the patients with a mutation in case 2 had UPD (2/22) or a BWSIC1 defect (1/5).

**Discussion**

We have further investigated the molecular pathology of sporadic BWS. Combining our current and previous results for CDKN1C mutation analysis, we have now identified germline mutations in almost 5% (four of 84) of sporadic cases. In these patients, the CDKN1C mutation was maternally inherited in one case, arose de novo in two cases, and was of uncertain origin in one case. The identification of a maternally inherited CDKN1C mutation in an isolated case has important implications for future pregnancies. The putative missense mutation in case 2 caused a non-conservative substitution of an evolutionarily conserved amino acid in the CDK inhibitor domain. To the best of our knowledge, a total of 14 germline CDKN1C mutations have now been described in BWS including three missense mutations (two of which were in the CDK binding domain). 1-4

The finding of KvDMR1 demethylation in a large proportion of sporadic BWS cases suggests that studies of KvDMR1 methylation and UPD status should be the initial investigations in BWS. Our PCR based assay was designed to detect patients with complete KvDMR1 LOM and patients mosaic for demethylation would not be expected to be detected so our results may provide an underestimate. We have confirmed and extended our findings that LOM at KvDMR1 is not associated with changes in H19 methylation, and that BWS patients with H19 hypermethylation and silencing do not show alterations in KvDMR1 methylation. 17 These findings are consistent with the hypothesis that there are two “imprinting control elements” (or imprinting centres) within the 1p15.5 imprinted gene cluster. Thus in humans, BWSIC1 would regulate IGF2 and H19 imprinting while BWSIC2 would regulate IGF2, CDKN1C, KCNQ1, and KCNQ1OT imprinting. 2 The results of transgenic mouse experiments support this model as Cdkn1c and Kvlqt1 (Kcnq1) imprinting appears to be independent of H19. 20 In the mouse, Cdkn1c expression requires a maternal germline imprint, although maternal germline methylation has not yet been found in or near the gene. Transgenic experiments have failed to identify a functional imprinting signal within 35 kb surrounding the Cdkn1c gene, 22 suggesting that Cdkn1c imprinting might be influenced by the BWSIC2 locus. Indeed, in a recent publication, Horikae et al 21 reported that a targeted deletion of KvDMR1 (in a modified mouse monochromosome hybrid system) silenced paternal KCNQ1OT (LIT1) expression and activated expression of KCNQ1 and CDKN1C with no effect on H19 imprinting.

The results of the current study are compatible with our previous suggestion that most sporadic BWS patients have epigenetic alterations in imprinted genes. They also confirm that BWSIC2 defects are more frequent than those at BWSIC1. 12 14 17 18 26 Imprinting centre defects are well defined in Prader-Willi and Angelman syndromes (PWS/AS). 25-27 In these disorders the detection of germline deletions has enabled the precise localisation of the sequences involved in imprint control. To date, no equivalent germline deletions have been identified in BWS patients. In patients with putative BWSIC1 defects, no evidence of H19 gene deletions or mutations have been identified. 12 14 26 However, mutations could be present in one or more of the putative BWSIC1 loci. They have yet to be analysed. 26 30 We note that in PWS/AS cases, epigenetic errors (epimutations) rather than mutations predominate in sporadic cases 27 and BWS patients with putative BWSIC1 and BWSIC2 defects may also result from epimutational events. Although limited Southern analysis has not yet shown evidence of germline deletions in BWS patients with LOM at KvDMR1, further investigations are required.

KvDMR1 is close to the transcription initiation site of a paternally expressed antisense transcript (KCNQ1OT or LIT1). Patients with loss of methylation at KvDMR1 show biallelic expression of KCNQ1OT. It has been proposed that KvDMR1 may be a boundary element that separates CDKN1C and KCNQ1 promoters from a downstream enhancer and that the boundary is open when methylated. 2 This model would predict that loss of KvDMR1 methylation would close the boundary leading to reduced expression of KCNQ1 and CDKN1C. 2 Previously we had reported that two patients with KvDMR1 LOM showed biallelic IGF2 expression. These results contrasted with those of Lee et al 28 who reported monoallelic IGF2 expression in four of five patients with LOI of KCNQ1OT. In this study
we identified two further BWS patients with KvDMR1 methylation in whom IGF2 imprinting status was defined. Intriguingly, one showed biallelic IGF2 expression and one monoallelic IGF2 expression. These results reconcile the differences between our previous findings (in which all patients with KvDMR1 LOM had biallelic IGF2 expression) and those reported by Lee et al.18 It appears that LOM of KvDMR1 is variably associated with the loss of imprinting of IGF2. The precise reasons for this variability is of great interest and could reflect the precise nature of the mutation or epimutation causing LOM of KvDMR1. Thus, a patient with a BWSCR1 breakpoint showed H19 independent LOI of IGF2,16 so the presence of IGF2 LOI may indicate a more complete defect in BWSIC2 function. The precise mechanism by which LOM at KvDMR1 would cause LOI of IGF2 is unknown. One possibility is that KvDMR1 LOM is associated with higher order chromatin modification that would alter imprinting over a large region. However, although loss of asymmetry of replication timing was observed in a BWS patients with a chromosome 11p15.5 inversion breakpoint and H19 independent LOI of IGF2,16 Squire et al19 reported normal asynchronous replication patterns in another BWS patient with a chromosome 11 inversion and in a patient with LOI of IGF2 and normal H19 methylation (KvDMR1 methylation was not assessed). A further possibility is that LOM of KvDMR1 results in closure of a boundary element between CDKN1C and a telomeric enhancer, which might then allow the enhancer to interact with the maternal IGF2 allele. We note the recent report of IGF2 control regions such as DMR1 which can influence IGF2 imprinting via a H19 independent pathway.48

BWS shows clinical variability and molecular heterogeneity. We have identified phenotypic differences between subgroups of BWS with defined molecular pathology.4,11,12 We have now extended these correlations by showing that exomphalos is frequent in patients with germline CDKN1C mutations and in patients with putative BWSIC2 defects but not in patients with UPD or BWSIC1 defects. These findings suggest that the precise BWS phenotype is determined by the loss of function of specific genes in the imprinted cluster. The association of exomphalos with germline CDKN1C mutations but not BWSIC1 defects (these patients in contrast are predicted to have IGF2 overexpression but no alterations in CDKN1C) is consistent with the phenotypic differences between “BWS mice” with CDKN1C inactivation or IGF2 expression.30,31 The overlapping phenotypes of BWS patients with germline CDKN1C mutations and IGF2 overexpression (as assumed in BWSIC1 defect cases) suggest that both gene products may act in the same biochemical pathway. Although the two imprinting domain model for 11p15.5 predicts that maternal CDKN1C expression will be downregulated in BWSCR2 cases, to date it has not been possible to determine CDKN1C expression patterns in patients with LOM at KvDMR1. However, the high frequency of exomphalos in patients with putative BWSIC2 defects is consistent with the hypothesis that LOM at KvDMR1 will lead to reduced CDKN1C expression. A further possibility is that molecular variability in patients with BWSIC2 defects will also be influenced by whether this is associated with LOI of IGF2.

The ability to predict which BWS patient are at risk for embryonal tumours would enhance clinical management. Although a very high incidence of Wilms and other tumours had been suggested in BWS patients with UPD, more recent studies suggest that the risk is lower than feared previously.17,18 It is of interest that although LOI of IGF2 and H19 methylation (consistent with a BWSIC1 defect) are common in Wilms tumour,19,20 loss of imprinting of KCNQ1OT (LIT1) (consistent with a BWSIC2 defect) has not been found in Wilms tumours.19 This might suggest that the risk of embryonal tumours would be less in BWS children with BWSIC2 defects (possibly related to the lack of apparent H19 silencing in these cases32). Consistent with this hypothesis, we found that embryonal tumours were confined to BWS patients with UPD and BWSIC1 defects. However, further studies are required to delineate the exact risks of embryonal tumours in different subgroups of BWS. Although overexpression of IGF2 has long been suspected as a cause for BWS, the precise reason for LOI of IGF2 (if this, as expected, leads to increased IGF2 protein levels) in patients with partial or no features of BWS37,38 suggest that LOI of IGF2 per se is not sufficient to cause BWS and alterations in expression or mutations of other genes (for example, CDKN1C) may also be required for the BWS phenotype.

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Engel, Smallwood, Harper, et al


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Jacqueline R Engel, Alan Smallwood, Antonita Harper, Michael J Higgins, Mitsuo Oshimura, Wolf Reik, Paul N Schofield and Eamonn R Maher

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