Mutation spectrum of MLL2 in a cohort of kabuki syndrome patients

Lucia Micale1, Bartolomeo Augello1†, Carmela Fusco1, Angelo Selicorni2, Maria N Loviglio1, Margherita Cirillo Silengo3, Alexandre Reymond4, Barbara Gumiero2, Federica Zucchetti2, Ester V D’Addetta1, Elga Belligni5, Alessia Calcagni1, Maria C Digilio5, Bruno Dallapiccola5, Francesca Faravellis, Francesca Forzano6, Maria Accadia7, Aldo Bonfante8, Maurizio Clementi9, Cecilia Daolio9, Sofia Douzgou10, Paola Ferrari11, Rita Fischetto12, Livia Garavelli13, Elisabetta Lapili4, Teresa Mattina15, Daniela Mels16, Maria G Patricelli17, Manuela Priolo18, Paolo Prontera19, Alessandra Renieri20, Maria A Mencarelli20, Gioacchino Scarano21, Matteo della Monica21, Benedetta Toschi22, Licia Turolla23, Alessandra Vancini24, Adriana Zatterale25, Orazio Gabrielli26, Leopoldo Zelante17 and Giuseppe Merla1*†

Abstract

Background: Kabuki syndrome (Niikawa-Kuroki syndrome) is a rare, multiple congenital anomalies/mental retardation syndrome characterized by a peculiar face, short stature, skeletal, visceral and dermatoglyphic abnormalities, cardiac anomalies, and immunological defects. Recently mutations in the histone methyl transferase MLL2 gene have been identified as its underlying cause.

Methods: Genomic DNAs were extracted from 62 index patients clinically diagnosed as affected by Kabuki syndrome. Sanger sequencing was performed to analyze the whole coding region of the MLL2 gene including intron-exon junctions. The putative causal and possible functional effect of each nucleotide variant identified was estimated by in silico prediction tools.

Results: We identified 45 patients with MLL2 nucleotide variants. 38 out of the 42 variants were never described before. Consistently with previous reports, the majority are nonsense or frameshift mutations predicted to generate a truncated polypeptide. We also identified 3 indel, 7 missense and 3 splice site.

Conclusions: This study emphasizes the relevance of mutational screening of the MLL2 gene among patients diagnosed with Kabuki syndrome. The identification of a large spectrum of MLL2 mutations possibly offers the opportunity to improve the actual knowledge on the clinical basis of this multiple congenital anomalies/mental retardation syndrome, design functional studies to understand the molecular mechanisms underlying this disease, establish genotype-phenotype correlations and improve clinical management.

Background

Kabuki syndrome (KS, MIM #147920), also known as Niikawa-Kuroki syndrome, is a rare, multiple congenital anomalies/mental retardation syndrome characterized by a peculiar face, which is defined by long palpebral fissures with eversion of the lateral third of the lower eyelids, short columella with a broad and depressed nasal tip, prominent ears, and a cleft or high-arched palate. Additional features include short stature, skeletal, visceral and dermatoglyphic abnormalities, cardiac anomalies, and immunological defects [1,2]. Kabuki syndrome has an incidence of 1 in 32,000, likely largely underestimated [3]. The vast majority of reported cases are sporadic. After initial and controversial data that associated this condition to chromosomal rearrangement [4,5], mutations in the MLL2 gene identified the underlying cause of Kabuki syndrome in approximately 72% of affected individuals [6,7]. The encoded MLL2 protein is a member of the Mixed Lineage Leukemia (MLL)
family of histone methyltransferases (HMT). The MLL proteins (MLLs) are part of the SET (Su(var)-3-9, Enhancer-of-zeste, Trithorax) family of proteins [8]. The highly conserved SET domain of MLLs confers histone methyltransferase activity, which is the core function of HMTs. MLLs are important in the epigenetic control of active chromatin states [9]. They act as transcriptional co-activators and are involved in embryogenesis and development through, for example, regulation of the expression of the HOX genes and their interaction with nuclear receptors [10,11].

The MLL2 gene encodes a multi-domain-containing protein of 5,537 amino acid residues that can methylate the Lys-4 position of histone H3 (H3K4), an epigenetic mark correlated with transcriptional activity [12,13]. MLL2 is involved in estrogen receptor α (ERα)-mediated signal transduction, acting as a coactivator of a complex that includes ASH2, RBQ3, and WDR5 [14].

In the present study, by direct sequencing of DNA samples from 62 Kabuki patients we identified 42 MLL2 variants, 38 of which are novel.

**Methods**

**Subjects and Clinical Data**

Our cohort comprised 62 index patients clinically diagnosed as affected by Kabuki syndrome (Figure 1 and Table 1). Patients were enrolled after obtaining appropriate informed consent by the physicians in charge and approval by the respective local ethics committees. Patients were included in this study whether at least four of the following inclusion criteria were present: 1) long palpebral fissures with eversion of the lateral portion of lower eyelid; 2) broad, arched eyebrows with sparseness; 3) short nasal columella with depressed nasal tip; 4) large, prominent or cupped ears; 5) developmental delay-mental retardation [15].

**Samples preparation**

Genomic DNAs were extracted from fresh and/or frozen peripheral blood leukocytes of the probands and their available family members using an automated DNA extractor and commercial DNA extraction Kits (EZ1, Qiagen, Hilden, Germany).

**PCR-based sequencing of MLL2**

Primers were designed using the Primer 3 Output program (http://frodo.wi.mit.edu/primer3/) to amplify the 54 coding exons of MLL2 (RefSeq NM_003482.3) gene including the intronic flanking sequences. Amplicons and primers were checked both by BLAST and BLAT against the human genome to ensure specificity. A complete list of primers is reported in Additional file 1, Table S1. The amplified products were subsequently purified and sequenced with a ready reaction kit (BigDye Terminator v1.1 Cycle, Applied Biosystems). The fragments obtained were purified using DyeEx plates (Qiagen) and resolved on an automated sequencer (3130xl Genetic analyzer DNA Analyzer, ABI Prism). Sequences were analyzed using the Sequencer software (Gen Code, Ann Arbor, Michigan). Whenever possible the mutations identified were confirmed on a second independent blood sample. The issue of whether the novel MLL2 missense alterations were causative mutations or neutral polymorphisms was addressed by searching dbSNP (http://www.ncbi.nlm.nih.gov/SNP) for their presence; the screening of 100 alleles from healthy unrelated control subjects and from the 1000 Genomes database [16] were used to assess their presence/absence in the general population. All existing and new mutations were described following the recommendations of the Human Genome Variation Society (http://www.hgvs.org/mutnomen).

**In silico analysis**

The putative causal and functional effect of each identified nucleotide variant was estimated by using the following in silico prediction tools: Polyphen http://genetics.bwh.harvard.edu/pph, Align GVG D http://agvgd.iarc.fr/agvgd_input.php, and MutPred http://mutdb.org/profile. Splice sites variants were evaluated for putative alteration of regulatory process at the transcriptional or splicing level with NetGene2 http://www.cbs.dtu.dk/services/NetGene2 and NNSPLICE http://www.fruitfly.org/seq_tools/splice.html. RESCUE-ESE http://genes.mit.edu/burgelab/rescue-ese/ and Fas-ESS http://genes.mit.edu/fas-ess/ online tools were used to predict exon-splicing enhancer and silencer, respectively. RepeatMasker http://www.repeatmasker.org/ was used to screen DNA sequences for the presence of direct repeats. The Coils program http://www.ch.embnet.org/software/COILS_form.html was employed to calculate the probability that the variant induces a conformational change in the coiled-coil domains. Finally multiple species alignment of MLL2 protein was made with the ClustalW software http://www.ebi.ac.uk/Tools/clustalw2/index.html using the following orthologs sequences obtained through the Ensembl genome browser http://www.ensembl.org: P. troglodytes (ENSPTRP00000041051), M. musculus (ENSMUSP0000023741), C. familiaris (gENSCAPF 00000012833), B. taurus (ENSBTAP00000019193), X. tropicalis (ENSTETP00000024427), and D. rerio (ENSDARP00000053862).

**Results and Discussion**

Exome sequencing recently revealed that mutations in the histone methyltransferase MLL2 gene are a major cause of Kabuki syndrome [6]. In a collaborative effort
that involved Italian Institutes, except one, we enrolled 62 individuals with a clinical diagnosis of sporadic Kabuki syndrome (Table 1). We detected nucleotide variants in 73% of the patients (45/62) by direct sequencing of all 54 exons of the *MLL2* gene; the vast majority of which are novel (90%, 38/42 different variants) (Figure 2, Additional file 1, Table S2, Additional file 2, Figure S1 and below) [6,7].

**Nonsense and frameshift mutations**

In agreement with previous reports we identified a majority of truncating mutations (70%, 29/42), three of
which were reported previously (Figure 2, Additional file 1, Table S2) [6,7]. Most of the variants are predicted, if translated, to encode shorter MLL2 proteins either by loss of the entire C-terminal region or parts of it (Figure 2). This region harbors highly conserved domains that are found in a variety of chromatin-associated proteins [17-19]: (i) the helical LXXLL regions involved in the recruitment of the MLL2 complex to the promoters of ERα target genes (Figure 2); (ii) FYRN and FYRC sequence motifs, two poorly characterized phenylalanine/tyrosine-rich regions of around 50 and 100 amino acids [20], respectively; and (iii) a catalytic SET motif that confers histone methyltransferases activity.

Although it has not been yet experimentally verified for the MLL2 gene, the prevalence of premature termination mutations may result in the partial transcripts degradation through nonsense-mediated mRNA decay (NMD). NMD is an evolutionarily conserved process that typically degrades transcripts containing premature termination codons (PTCs) to prevent translation of unnecessary or aberrant and possible transcripts [21]. The NMD process takes place when PTCs are located more than 50-55 nucleotides upstream of an exon-exon junction [22]. As 86% (25/29) of such detected MLL2 mutations follow this rule it is likely that the consequent MLL2 haploinsufficiency could be the driving force for the onset of the Kabuki syndrome.

Indel variants

Our screen revealed three not yet described indel variants located in the C-terminal region of the protein (see samples KB71, KB77, and KB53 in Additional file 1, Table S2). They might have resulted from slipped mispairing between direct repeats or through the insertion or deletion of a single base within a mononucleotide tract (Additional file 1, Table S3), as already reported [23].

COILS algorithm predicted the amplification of one of the five coiled-coil putative domains for the c.11819_11836dupTTCAACAACAGCAGCAGC (p. Lys3940_Gln3945dup) (Figure 2 and data not shown), a domain involved in protein-protein interaction. This variant was inherited from the apparently asymptomatic mother. It is thus impossible to conclusively determine the pathogenic nature of the resulting protein.

Splice site variants

We detected 3 variants located at the splice site junctions, two of which are novel [7] (Additional file 1, Table S2); the in silico modeling predicted complete or partial abrogation of the junction formation with a pathogenic impact. The c.400+1G>A, occurring within the invariant GT donor splice site in intron 3-4, results in the disruption of the canonical splice site and it is expected to produce an aberrant protein of only 135 residues. The c.401-3A>G, occurring 3 bp away from the next intron-exon junction, is predicted to create a new acceptor splice site at position -3 within intron 3-4 that could lead to a frameshift encoding a mutant protein with a premature stop at 84 codons downstream. Finally, c.13999+5G>A decreases the donor site score prediction, possibly resulting in a less efficient intron splicing. Unfortunately, RNA from these patients was unavailable preventing further investigation of the effect of these variants.

| Table 1 Clinical features of our cohort of Kabuki syndrome patients |
|---------------------------------|-----------------|-----------------|-----------------|
| **Gender** | 36/62 (58.1%) Male | 26/62 (41.9%) Female |
| **General features** | | |
| Short stature | 38/62 (61.2%) | |
| Microcephaly | 18/62 (29%) | |
| Neonatal problems | 42/62 (67.7%) | |
| **Facial** | | |
| Long palpebral fissures | 59/62 (95.2%) | |
| Everted lower eyelids | 53/62 (85.5%) | |
| Large dysplastic ears | 56/62 (90.3%) | |
| Arched eyebrows, sparse lateral one third | 51/62 (82.2%) | |
| Flat nasal tip | 43/62 (69.3%) | |
| Abnormal dentition | 33/62 (53.2%) | |
| High/cleft palate | 37/62 (59.7%) | |
| Strabismus | 26/62 (41.9%) | |
| Blue sclerae | 11/62 (17.7%) | |
| Micrognathia | 20/62 (32.2%) | |
| Ptosis | 32/62 (51.6%) | |
| Broad nasal root | 39/62 (62.9%) | |
| Oligodontie | 23/62 (37.1%) | |
| Thin upper and full lower lip | 44/62 (71%) | |
| **Limb/skeletal** | | |
| Persistent fetal pads | 47/62 (75.8%) | |
| Brachy/clinodactyly | 39/62 (62.9%) | |
| Lax joints | 30/62 (48.4%) | |
| Hip dislocation | 8/62 (12.9%) | |
| **Visceral anomalies** | | |
| Cardiac anomalies | 37/62 (59.7%) | |
| Urogenital anomalies | 24/62 (38.7%) | |
| **Neurologic** | | |
| MR | 52/62 (83.9%) | |
| Hypotonia | 37/62 (59.7%) | |
| Seizures | 13/62 (21.4%) | |
| **Other clinical features (most recurrent)** | | |
| frequent infections | 26/62 (41.9%) | |
| leftearly breast development | 10/62 (16.1%) | |
| lefthypacusia | 7/62 (11.3%) | |
| skeletal anomalies | 6/62 (9.7%) | |
| leftthyroid anomalies | 4/62 (6.9%) | |
| leftagenesis/dysgenesis corpus callosum | 3/62 (5.2%) | |

which were reported previously (Figure 2, Additional file 1, Table S2) [6,7]. Most of the variants are predicted, if translated, to encode shorter MLL2 proteins either by loss of the entire C-terminal region or parts of it (Figure 2). This region harbors highly conserved domains that are found in a variety of chromatin-associated proteins [17-19]: (i) the helical LXXLL regions involved in the recruitment of the MLL2 complex to the promoters of ERα target genes (Figure 2); (ii) FYRN and FYRC sequence motifs, two poorly characterized phenylalanine/tyrosine-rich regions of around 50 and 100 amino acids [20], respectively; and (iii) a catalytic SET motif that confers histone methyltransferases activity.

Although it has not been yet experimentally verified for the MLL2 gene, the prevalence of premature termination mutations may result in the partial transcripts degradation through nonsense-mediated mRNA decay (NMD). NMD is an evolutionarily conserved process that typically degrades transcripts containing premature termination codons (PTCs) to prevent translation of unnecessary or aberrant and possible transcripts [21]. The NMD process takes place when PTCs are located more than 50-55 nucleotides upstream of an exon-exon junction [22]. As 86% (25/29) of such detected MLL2 mutations follow this rule it is likely that the consequent MLL2 haploinsufficiency could be the driving force for the onset of the Kabuki syndrome.

Indel variants

Our screen revealed three not yet described indel variants located in the C-terminal region of the protein (see samples KB71, KB77, and KB53 in Additional file 1, Table S2). They might have resulted from slipped mispairing between direct repeats or through the insertion or deletion of a single base within a mononucleotide tract (Additional file 1, Table S3), as already reported [23].

COILS algorithm predicted the amplification of one of the five coiled-coil putative domains for the c.11819_11836dupTTCAACAACAGCAGCAGC (p. Lys3940_Gln3945dup) (Figure 2 and data not shown), a domain involved in protein-protein interaction. This variant was inherited from the apparently asymptomatic mother. It is thus impossible to conclusively determine the pathogenic nature of the resulting protein.

Splice site variants

We detected 3 variants located at the splice site junctions, two of which are novel [7] (Additional file 1, Table S2); the in silico modeling predicted complete or partial abrogation of the junction formation with a pathogenic impact. The c.400+1G>A, occurring within the invariant GT donor splice site in intron 3-4, results in the disruption of the canonical splice site and it is expected to produce an aberrant protein of only 135 residues. The c.401-3A>G, occurring 3 bp away from the next intron-exon junction, is predicted to create a new acceptor splice site at position -3 within intron 3-4 that could lead to a frameshift encoding a mutant protein with a premature stop at 84 codons downstream. Finally, c.13999+5G>A decreases the donor site score prediction, possibly resulting in a less efficient intron splicing. Unfortunately, RNA from these patients was unavailable preventing further investigation of the effect of these variants.
**Missense variants**

Missense *de novo* variants have already been found in Kabuki patients. Ng and colleagues reported 8 pathogenic missense variants, two of which were recurrent in affected patients. As these were all mapping within the last exon of *MLL2* that encodes the different conserved C-terminal domains of the protein (see above), the authors suggested that such mutations are tolerated, while mutations elsewhere are lethal. By *in silico* analysis, Paulussen *et al.* proposed the pathogenicity of two missense variants located within that C-terminal region [6,7]. We detected seven patients with a single or two missense variants (KB28 and KB38 patients; Additional file 1, Table S2). PCR amplification, cloning and sequencing showed that both sets of the two sequence changes in KB28 and KB38 patients are located on one allele. From a phenotypic point of view, the two patients with pairs of missense variants do not appear to be more severely affected than affected individuals with single variants. Sequencing of the corresponding exons in the KB38 parents demonstrated that both variants arose *de novo*, while the KB28 patient inherited the variant from the apparently asymptomatic mother. We had also accessed to DNA of parents of carriers of missense variants (Additional file 1, Table S2). Yet, in both cases the *MLL2* variant was inherited from the apparently asymptomatic father.

The missense variants are distributed across the entire length of the *MLL2* gene (Figure 2). They were not found in 50 healthy unrelated control samples and were absent from the 1000 Genomes database [16]. The putative functional relevance and pathogenicity of these *MLL2* missense variants were predicted by *in silico* software. The PolyPhen program, which predicts possible impact of an amino acid substitution on the structure and function of a human protein, identified only the p.Pro2841Thr variant as possibly damaging. Accordingly to the criteria of Align-GVGD all missense variants were predicted to be deleterious (Table 2). Finally, we used the computational model MutPred, designed to classify an amino acid substitution as disease-associated or neutral in human. MutPred predicted that four of the identified missense variants have a high probability (≥0.5) of being deleterious and generated *in silico* hypothesis for the possible pathological mechanism for three of them (Table 2). The analysis of the mutated residues in 7 *MLL2* proteins orthologs showed that all the missense variants occurred at amino acid residues evolutionarily conserved (Figure 2).
### Table 2 In silico prediction of pathogenic effect of MLL2 missense and splice site variants

| ID   | Exon | Mutation                  | AA change   | Polyphen | Align | GVGD | Predicted Probability of deleterious mutation | Confident in silico hypothesis |
|------|------|---------------------------|-------------|----------|-------|------|---------------------------------------------|-------------------------------|
| KB32 | 11   | c.3773G>A                 | p.Arg1258Gln| benign   | deleterious | 0.16 | loss of loop (p = 0.0288), loss of catalytic residue at R1258 (p = 0.0301); gain of helix (p = 0.0349) | no change  
|      |      |                           |             |          |       |     |                                             |                               |
| KB28 | 15   | c.[4249A>G;4252C>A]       | p.[Met1471Val; Leu1418Met] | benign; benign | deleterious; deleterious | 0.46; 0.47 | none; none | no change | gain of one ESS  
|      |      |                           |             |          |       |     |                                             |                               |
| KB34 | 16   | c.4565A>G                 | p.Gln1522Arg | benign   | deleterious | 0.50 | none | no change | loss of one ESE no change  
|      |      |                           |             |          |       |     |                                             |                               |
| KB27 | 34   | c.8521C>A                 | p.Pro2841Thr | possibly damaging | deleterious | 0.24 | gain of phosphorylation P2841 (p = 0.028) | gain of two ESEs loss of one ESS  
|      |      |                           |             |          |       |     |                                             |                               |
| KB38 | 48   | c.[15084C>G;15100T>G]     | p.[Asp 5028Glu; Phe5034Val] | benign; benign | deleterious; deleterious | 0.42; 0.56 | none; none | gain of three ESEs no change | gain of two ESSs no change  
|      |      |                           |             |          |       |     |                                             |                               |
| KB76 | 48   | c.1517A>C                 | p.His5059pro | benign   | deleterious | 0.71 | none | no change | gain of ubiquitination at KS544 (p = 0.0396)  
|      |      |                           |             |          |       |     |                                             |                               |
| KB17 | 50   | c.16019G>A                | p.Arg5340Gln | benign   | deleterious | 0.53 | none | no change | no change  
|      |      |                           |             |          |       |     |                                             |                               |

### Splice site modification prediction

| ID   | Intron | Mutation                  | NetGene | NNSplice | Predicted Protein | Conclusion |
|------|--------|---------------------------|---------|----------|-------------------|------------|
| KB31 | Intron 3-4 | c.400+1G>A   | Loss of DS | Loss of DS | 135 AA with novel 2 AA | deleterious |
| KB20 | Intron 3-4 | c.401-3A>G   | New AS at-3 | New AS at-3 | 217 AA with novel 84 AA | deleterious |
| KB29 | Intron 42-43 | c. 13999+5 G>A | Lower confidence of DS prediction | Lower confidence of DS prediction | | unpredictable |
Finally, we employed RESCUE-ESE and Fas-ESS tools on missense variants and frameshift mutations to predict associated splicing phenotypes by identifying sequence changes that disrupt or alter predicted Exonic Splicing Enhancers (ESE) and Exonic Splicing Silencers (ESS). ESE and ESS are short oligonucleotides that can enhance or inhibit pre-mRNA splicing when present in exons, playing important roles in constitutive and alternative splicing. A variation that disrupts an ESE, for instance, could cause exon skipping which would result in the exclusion of an entire exon from the mRNA transcript. Conversely, a substitution in the ESS sequence promotes the use of adjacent splice sites, often contributing to alternative splicing. As reported in Table 2 and Additional file 1, Table S3, we found that some of the MLL2 mutations lead to creation of new ESEs and/or to disruption of predicted wild type ESEs/ESSs. As secondary structures or adjacent negative elements also participate to the modulation of the splicing event mediated by ESE and ESS, we retain that association to functional studies will enable to better understand the role of the reported cases of ESEs/ESSs disruption or alteration in the complex phenotypic spectrum observed in the Kabuki patients.

Conclusions

Our study increases the number of identified MLL2 mutations and variants, and emphasizes some characteristics of the spectrum of MLL2 mutations associated with this pathology, further providing insight into its etiology. The in silico analysis predicts that the identified MLL2 missense, splice-site and indel variants might be pathogenic. Other studies reported the presence of such MLL2 variants predicted to be associated with the disease. However, their biological significance and pathogenicity were not unambiguously demonstrated; therefore further and more functionally oriented studies are needed to understand the nature of these variants and their possible role in the disease. Solving these issues is relevant to avoid any incorrect interpretation and diagnosis of other Kabuki cases carrying such MLL2 variants.

We were unable to found any detectable point mutation and/or small del/dup in the coding region of MLL2 gene in 27% (17/62) of the Kabuki syndrome patients. Mutations in MLL2 regulatory regions, exon microduplications and/or microdeletions, as well as genetic heterogeneity of the syndrome may account for these negative results. Alternatively, some of these patients might have been misdiagnosed as a result of the complex clinical spectrum covered by this pathology, thus possibly highlighting the need to more accurately select Kabuki cases before conducting the analysis.

In summary, this study underlines the relevance of mutational screening of the MLL2 gene among patients with Kabuki syndrome. The identification of a large spectrum of MLL2 mutations will offer the opportunity to improve the actual knowledge on the clinical basis of this multiple congenital mental retardation syndrome, to design functional studies to understand the molecular mechanisms underlying the disease, to establish genotype-phenotype correlations, to improve the clinical management, and to identify potential targets for therapy.

Additional material

Additional file 1: Table S1. Oligos used in this study. Table S2. MLL2 mutations identified in our cohort of KS patients and as reported in the literature. Table S3. Repeats (underlined and highlighted in red) that might mediate micro-deletions, micro-insertion/deletions (indel), and micro-duplications in the MLL2 gene.

Acknowledgements and funding

We thank all the families that agree to participate and get possible this study. This work was in part supported by grants from the Italian Ministry of Health (Ricerca Corrente 2008-10), the Fondazione Banca del Monte di Foggia “Domenico Siniscalco Ceci”, the Jerome Lejeune Foundation, and with the contribution of Ministero degli Affari Esteri, Direzione Generale per la Promozione e la Cooperazione Culturale (2009-2010) to QM. We thank Mariani Foundation, Milan for the support to the activity of Ambulatorio di Genetica Clinica Pediatrica Fondazione MMBA to AS. AR is supported by grants from the Swiss National Science Foundation. We thank the Galliera Genetic Bank - Network of Telethon Genetic Biobanks (project GTB07001) for providing us with some Kabuki samples. The founders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Written consent for publication was obtained from the patients or their relatives.

Author details

1Medical Genetics Unit, IRCCS Casa Sollievo della Sofferenza Hospital, 71013 San Giovanni Rotondo, Italy. 2Ambulatorio Genetica Clinica Pediatrica, Clinica Pediatrica Università Milano Bicocca, Fondazione MMBA AOS Gerardo Monza, Italy. 3Dipartimento di Scienze Pediatriche, Università di Torino, Italy. 4Center for Integrative Genomics, University of Lausanne, Lausanne, Switzerland. 5Medical Genetics, Bambino Gesù Paediatric Hospital, IRCCS, Rome, Italy. 6Division of Medical Genetics, Galleria Hospital, Genova, Italy. 7Laboratory of Medical Genetics, “V.Fazzà” Hospital, Lecce, Italy. 8Medical Genetics Unit, St. Bassiano Hospital, Bassano del Grappa, Italy. 9Dipartimento Pediatra, Genetica Clinica, Padova, Italy. 10Department of Genetics, Institute of Child Health, “Aghia Sophia” Children’s Hospital, Athens, Greece. 11Division Materno Infantile, Università degli studi Modena, Italy. 12U.O. Malattie Metaboliche PO Giovanni XXIII, AOU Policlinico Consorziale, Bari, Italy. 13Clinical Genetics Unit, S.Maria Nuova Hospital Reggio Emilia, Italy. 14Medical Genetics Unit, Children’s Hospital Anna Meyer, Firenze, Italy. 15Genetica Medica, Università di Catania, Catania, Italy. 16Area Funzionale di Genetica Clinica Pediatrica, Dipartimento di Pediatría, Università degli Studi di Napoli “Federico II”, Italy. 17Biologia Molecolare e Citogenetica, Diagnostica e Ricerca San Raffaele, Milano, Italy. 18Unita’ Operativa di Genetica Medica, Azienda Ospedaliera Bianchi-Melacrinio-Morelli, Reggio Calabria, Italy. 19Medical Genetics Unit, University of Perugia, “S. Maria della Misericordia” Hospital, Perugia, Italy. 20Medical Genetics Section, Biotechnology Department, University of Siena, Italy. UOC Genetica Medica, Dipartimento di Emergenza
Urgenza e dei Servizi Diagnostici, Azienda Ospedaliera Universitaria Senese, Siena, Italy. 21 UOC Genetica Medica, Azienda Ospedaliera RN “G.Rummo”, Benevento, Italy. 22 Medical Genetics Section, Cytogenetics and Molecular Genetics Unit, Santa Chiara University Hospital, Pisa, Italy. 23Ambulatorio di Genetica Medica, Azienda ULSS 9, Treviso, Italy. 24Newborn Intensive Care Unit, Maggiori Hospital, Bologna, Italy. 25Servizio di Genetica, ASL NAPOLI 1 P.S.I. Elena d’Aosta Napoli, Italy. 26Istituto di Scienze Materno-Infantili, Universita Politecnica delle Marche, Ancona, Italy.

Authors’ contributions
GM designed the study and obtained the necessary financial support. BA, MNL, CF, AC, and EVD carried out the molecular genetic studies. LM and GM interpreted the results and wrote the manuscript with the help of LZ and AR. All other authors provided samples. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Received: 11 March 2011 Accepted: 9 June 2011 Published: 9 June 2011

References
1. Kuroki Y, Suzuki Y, Chyo H, Hata A, Matsui i: A new malformation syndrome of long palpebral fissures, large ears, depresso nasal tip, and skeletal anomalies associated with postnatal dwarfism and mental retardation. J Pediatr 1981, 99(4):570-573.
2. Niikawa N, Matsuura N, Fukushima Y, Ohsawa T, Kaji T: Kabuki make-up syndrome: a syndrome of mental retardation, unusual facies, large and protruding ears, and postnatal growth deficiency. J Pediatr 1981, 99(4):565-569.
3. Niikawa N, Kuroki Y, Kaji T, Matsuura N, Ishikiriyama S, Tonomi H, Ishikawa N, Yamada Y, Fujita M, Uremoto H, et al: Kabuki make-up (Niikawa-Kuroki) syndrome: a study of 62 patients. Am J Med Genet 1988, 31(3):565-589.
4. Cusco I, del Campo M, Villadrich M, Gonzalez E, Gener B, Galan E, Toledo L, Perez-Jurado LA: Array-CGH in patients with Kabuki-like phenotype: identification of two patients with complex rearrangements including 2q37 deletions and no other recurrent aberration. BMC Med Genet 2008, 9:27.
5. Milunsky JM, Huang XL: Unmasking Kabuki syndrome: chromosome 8p22-8p23 duplication revealed by comparative genomic hybridization and BAC-FISH. Clin Genet 2003, 64(6):509-516.
6. Ng SB, Bigham AW, Buckingham CJ, Hannibal MC, McMillin MJ, Gildersleeve H, Beck AE, Tabor HK, Cooper GM, Mofford HC, Lee C, Turner EH, Smith JD, Reider MJ, Yoshiura K, Matsumoto N, Ohta T, Nikawa N, Nickerson DA, Bamshad MJ, Shendure J: Exome sequencing identifies MLL2 mutations as a cause of Kabuki syndrome. Nat Genet 2010, 42(9):790-793.
7. Paulussen AD, Stegmann AP, Blok MJ, Tserpelis D, Posma-Velter C, Dettich Y, Smeets HJ, Devriendt K, Schrander-Stumpel CT, Mancini GM, Wessels MW, Hennekam RC, Vreeburg M, Geraedts J, de Ravel T, Fryns JP, Smeets HJ, Devriendt K: Identification of novel protein domain families associated with nuclear hormone signaling and mRNA processing. Mol Cell Biol 2009, 29(7):2030-2047.
8. Nilestra O, Bhojraj A, Narte LP, Gilbert P: A map of human genome variation from population-scale sequencing. Nature 2010, 467(7319):1061-1073.
9. Balconi LS, Ronne H: Evidence of domain swapping within the junmorni family of transcription factors. Trends Biochem Sci 2000, 25(6):274-276.
10. Durbin RM, Abecasis GR, Altshuler DL, Auton A, Brooks LD, Gibbs RA, Hurles ME, McVean GA: A map of human genome variation from population-scale sequencing. Nature 2010, 467(7319):1061-1073.
11. Eissenberg JC, Shilatifard A: When cells stop making sense: effects of nonsense codons on RNA metabolism in vertebrate cells. RNA 1995, 1(5):453-465.
12. Nagy E, Maquat LE: A rule for termination-codon position within intron-containing genes: when nonsense affects RNA abundance. Trends Biochem Sci 1998, 23(6):198-199.
13. Prasad R, Zhadanov AB, Sedkov Y, Bullitch F, Druck T, Ralliappali R, Yano T, Alder H: Coreset CM, Huebner K, Mazo A, Canani E: Structure and expression pattern of human ALR, a novel gene with strong homology to ALL-1 involved in acute leukemia and to Drosophilal trithorax. Oncogene 1997, 15(5):549-560.
14. Garcia-Alai MM, Ali MD, Jorgensen RS, Boryczko MS: The structure of the FVR domain of transforming growth factor beta receptor 1. Protein Sci 2010, 19(7):1432-1438.
15. Doerks T, Copley RR, Schuh C, Ponting CP, Bork P: Systematic identification of novel protein domain families associated with nuclear functions. Genome Res 2002, 12(1):47-56.
16. Ansari KI, Mandal SS: Topological features of the MLL2 complex as a coactivator for estrogen receptor alpha. J Biol Chem 2006, 281(23):15714-15720.
17. Xie M, Rao SM, Zhu YJ: Identification of the MLL2 complex as a coactivator for estrogen receptor alpha. J Biol Chem 2006, 281(23):15714-15720.
18. Prasad R, Zhadanov AB, Sedkov Y, Bullitch F, Druck T, Ralliappali R, Yano T, Alder H: Coreset CM, Huebner K, Mazo A, Canani E: Structure and expression pattern of human ALR, a novel gene with strong homology to ALL-1 involved in acute leukemia and to Drosophilal trithorax. Oncogene 1997, 15(5):549-560.
19. Garcia-Alai MM, Ali MD, Jorgensen RS, Boryczko MS: The structure of the FVR domain of transforming growth factor beta receptor 1. Protein Sci 2010, 19(7):1432-1438.
20. Doerks T, Copley RR, Schuh C, Ponting C, Bork P: Systematic identification of novel protein domain families associated with nuclear functions. Genome Res 2002, 12(1):47-56.
21. Nagy E, Maquat LE: A rule for termination-codon position within intron-containing genes: when nonsense affects RNA abundance. Trends Biochem Sci 1998, 23(6):198-199.
22. Prasad R, Zhadanov AB, Sedkov Y, Bullitch F, Druck T, Ralliappali R, Yano T, Alder H: Coreset CM, Huebner K, Mazo A, Canani E: Structure and expression pattern of human ALR, a novel gene with strong homology to ALL-1 involved in acute leukemia and to Drosophilal trithorax. Oncogene 1997, 15(5):549-560.
23. Garcia-Alai MM, Ali MD, Jorgensen RS, Boryczko MS: The structure of the FVR domain of transforming growth factor beta receptor 1. Protein Sci 2010, 19(7):1432-1438.
24. Doerks T, Copley RR, Schuh C, Ponting CP, Bork P: Systematic identification of novel protein domain families associated with nuclear functions. Genome Res 2002, 12(1):47-56.
25. Nagy E, Maquat LE: A rule for termination-codon position within intron-containing genes: when nonsense affects RNA abundance. Trends Biochem Sci 1998, 23(6):198-199.
26. Prasad R, Zhadanov AB, Sedkov Y, Bullitch F, Druck T, Ralliappali R, Yano T, Alder H: Coreset CM, Huebner K, Mazo A, Canani E: Structure and expression pattern of human ALR, a novel gene with strong homology to ALL-1 involved in acute leukemia and to Drosophilal trithorax. Oncogene 1997, 15(5):549-560.
27. Garcia-Alai MM, Ali MD, Jorgensen RS, Boryczko MS: The structure of the FVR domain of transforming growth factor beta receptor 1. Protein Sci 2010, 19(7):1432-1438.
28. Doerks T, Copley RR, Schuh C, Ponting CP, Bork P: Systematic identification of novel protein domain families associated with nuclear functions. Genome Res 2002, 12(1):47-56.
29. Nagy E, Maquat LE: A rule for termination-codon position within intron-containing genes: when nonsense affects RNA abundance. Trends Biochem Sci 1998, 23(6):198-199.
30. Prasad R, Zhadanov AB, Sedkov Y, Bullitch F, Druck T, Ralliappali R, Yano T, Alder H: Coreset CM, Huebner K, Mazo A, Canani E: Structure and expression pattern of human ALR, a novel gene with strong homology to ALL-1 involved in acute leukemia and to Drosophilal trithorax. Oncogene 1997, 15(5):549-560.
31. Garcia-Alai MM, Ali MD, Jorgensen RS, Boryczko MS: The structure of the FVR domain of transforming growth factor beta receptor 1. Protein Sci 2010, 19(7):1432-1438.