Constitutional chromosomal events at 22q11 and 15q26 in a child with a pilocytic astrocytoma of the spinal cord

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Abstract

We report on a 9-years-old patient with mild intellectual disability, facial dimorphisms, bilateral semicircular canal dysplasia, periventricular nodular heterotopias, bilateral hippocampal malrotation and abnormal cerebellar foliation, who developed mild motor impairment and gait disorder due to a pilocytic astrocytoma of the spinal cord. Array-CGH analysis revealed two paternal inherited chromosomal events: a 484.3 Kb duplication on chromosome 15q26.3 and a 247 Kb deletion on 22q11.23. Further, a second de novo 1.5 Mb deletion on 22q11.21 occurred. Chromosome 22 at q11.2 and chromosome 15 at q24q26 are considered unstable regions subjected to copy number variations, i.e. structural alterations of genome, mediated by low copy repeat sequences or segmental duplications. The link between some structural CNVs, which compromise fundamental processes controlling DNA stability, and genomic disorders suggest a plausible scenario for cancer predisposition. Evaluation of the genes at the breakpoints cannot account simultaneously for the phenotype and tumour development in this patient. The two paternal inherited CNVs arguably are not pathogenic and do not contribute to the clinical manifestations. Similarly, although the de novo large deletion at 22q11.21 overlaps with the Di George (DGS) critical region and results in haploinsufficiency of genes compromising critical processes for DNA stability, this case lacks several hallmarks of DGS.

Keywords: Pilocytic astrocytoma, Spinal cord, Semicircular canal dysplasia, 15q duplication, 22q11.2 deletion

Background

Patients harbouring deletions within chromosome 22q11 present a wide array of clinical phenotypes, including DiGeorge, Velo-Cardio-Facial (DGS/VCFS) or 22q11.2 deletion syndrome [1-3]. Few reports of malignancy in patients with 22q11.2 deletion syndrome exist, including kidney tumours, hepatoblastomas, neuroblastomas, lymphomas, and, very rarely, central nervous system (CNS) tumours [4-7]. About 97% of DGS/VCFS patients have either a ~3 Mb deletion or, less commonly, a smaller ~1.5 Mb nested deletion [MIM 188400] [1]. Indeed, the long arm of chromosome 22 at band q11.2 is one of the most unstable regions with eight different specific low-copy repeats, designated LCR22s (LCR22-A-LCR22-H), which can yield recurrent Copy Number Variants (CNVs), such as microdeletions and microduplications [8]. A similar unstable region is present even on chromosome 15 at band q24q26 [9]. While the significance of CNVs to human pathology is being recognized, it is unclear how common or functionally relevant CNVs are to the process of carcinogenesis [10].

We present the clinical features of a 9-years-old male harbouring two paternally inherited events – a short 22q11.23 deletion and a 15q26.3 duplication – and a de novo 1.5 Mb deletion at 22q11.2 band. The child developed a spinal cord pilocytic astrocytoma (PA) [11-13]. We previously reported the cytogenetic characterization of the child and her family, suggesting that parental microdeletions/duplications could predispose to the formation of de novo pathogenetically distinct CNVs in offspring [14]. Herein, we attempt to correlate the multiple...
CNVs with both development of the spinal cord PA and clinical phenotype.

Case presentation
Clinicopathological description
Patient's clinical history, including information on family pedigree, and cytogenetic findings were previously reported (Figure 1) [14].

The child came to our attention at the age of 9 years, after a 3-year history of gait disturbance and right leg pain initially responding to anti-inflammatory therapy. Neurological examination revealed pes cavus, motor clumsiness, gait abnormalities with toe-walking and positive heel walk test. Somato-sensory evoked potentials showed low-amplitude potentials in lower limbs. Psychometrics confirmed mental retardation (Raven's Matrices, PM47 = 20th percentile). Electroencephalography was unremarkable. Mild obesity with increased abdominal fat distribution was present. Ophthalmologic examination was normal, while audiometric tests revealed left conductive hearing loss. Cardiac or urologic malformations were absent, as well as immune deficiency or congenital hypocalcaemia.

Neuroimaging and pathological examination
Spine magnetic resonance imaging (MRI) revealed enlargement of the inferior dorsal spinal cord due to a solid tumour, hypointense on T1-weighted images and hyperintense on T2-weighted images, with mild, inhomogeneous contrast enhancement. Leptomeningeal invasion and dissemination were absent (Figure 2a, b). Brain MRI showed small nodules of subependymal gray matter heterotopia, bilateral hippocampal malrotation, and abnormal cerebellar foliation (Figure 2c, d). Temporal bone computerized tomography revealed bilateral digenesis of the lateral semicircular canals and vestibules forming a single cavity.

![Figure 1 Clinical features of the proband and family pedigree. (a) In the family pedigree the propositus (IV-1) is indicated by a full-black square and an arrow - bold cross within square (II-7) indicates the second degree uncle with pes cavus - black corner of square (II-4) and circle (II-1) indicate second degree relatives who died for tumour under 55 years - black corner of circle (I-6) indicate a second degree aunt informative for the patient - NA indicates no chromosomal abnormalities. (b, c) The patient presented mild facial dimorphisms.](image-url)
with normal appearing cochleae (Figure 2e). Signs of left chronic otitis media were found.

After a period of surveillance, the patient underwent neurosurgical intervention. The absence of a cleavage plane between the tumour and the surrounding tissue hampered total or subtotal resection, due to high risk of functional damage. Therefore, only a biopsy was performed. Histological findings were consistent with PA. The labelling index, as evaluated by the monoclonal antibody Ki67-MIB-1, was 1%. No expression of p53 protein was found.

Adjuvant therapies were planned to be introduced just in case of clinical and/or imaging worsening. After four years, the child is neurologically stable with no remarkable change in tumour volume.

**CNV analysis**

CNVs were analyzed using UCSC Genome Browser (http://www.genome.ucsc.edu) and Decipher Database (http://decipher.sanger.ac.uk/, version 6.0) website tools. The 15q26.3 duplication, with proximal breakpoints between base pairs 100,051,182 and 100,996,155, involved myocyte enhancer factor 2A (MEF2A) [MIM 600660], a target of ERK/MAPK pathway. Defects in this gene were reported in autosomal dominant coronary artery disease 1 with myocardial infarction (ADCAD1) [MIM 608320]. The duplication also involved putative peptidoglycan-binding domain containing 4 (LYSMD4) [XM_005254863], ADAM metalloprotease with thrombospondin type 1 motif, 17 (ADAMTS17) [DQ_217943] and interrupted ceramide synthase 3 (CERS3) [MIM 615276]. Mutations in ADAMTS17 can cause autosomal recessive Weill-Marchesani-like syndrome [MIM 613195] while alteration of both ADAMTS17 and CERS3 have been associated with autosomal recessive congenital ichthyosis 9 (ARCI9) [MIM 615023].

The 22q11.23 deletion (chr22:25,664,618-25,911,651), involving low density lipoprotein receptor-related protein 5-like (LRP5L) [NM_182492], has never been associated with any disease.

The de novo 22q11.21 deletion (chr22:18,894,835-20,311,763) involved 24 genes located within the DGS critical region (DGCR). Besides T-box 1 (TBX1), a transcription factor regulating developmental processes, several genes are involved, as clathrin, heavy chain-like 1 (CLTCL1) [MIM 601273], RAN binding protein 1 (RANBP1) [MIM 601180], histone cell cycle regulator (HIRA) [MIM 600237], cell division cycle 45 (CDC45) [MIM 603465], and catechol-O-methyltransferase (COMT) [MIM 116790] partially overlapping thioredoxin reductase 2 (TXNRD2) [MIM 606448]. These genes are active in many cellular processes, including cell cycle progression, senescence, signal transduction, apoptosis and gene regulation.

**Discussion**

Locus-specific rates of CNV formation underlie not just rare complex disorders involving multiple congenital
abnormalities but also sporadic diseases, and perhaps cancer predisposition. The principal models currently invoked to explain genomic instability and susceptibility to cancer, conjecture the deleterious consequences of some structural CNVs on fundamental processes, including DNA replication and damage response (DDR) processes [8,15].

The child paternally inherited the 15q26.3 duplication. This distal duplication involved four genes active in several cellular processes encompassing MEF2A which is a target of ERK/MAPK pathway. Although the pathway plays an important role in PA [16], defects in MEF2A have never been associated to this tumour. On the other hand, paternally inherited 22q11.23 short deletion arguably contribute neither to clinical phenotype nor to tumour development.

The present case had a de novo 1.5 Mb deletion at chromosome 22q11.21 (Figure 1a), one of the most rearrangement-prone regions in the human genome associated with genetic disorders including DGS/VCFS.

Breakpoints analysis revealed that the heterozygous deletion contains several genes of the critical region for DGS/VCFS, i.e. several DGCRs, encompassing TBX1, who is responsible for most of malformations of this syndrome [17]. Twenty-four genes resulted haploinsufficient including cell division cycle 45 (CDC45), catechol-O-methyltransferase (COMT) and thioredoxin reductase 2 (TXNRD2) which could increase risk of malignancy in patients with 22q11.2 DS [4,8,18].

Although 22q11.21 deletion overlaps with the DGS/VCFS region [2], the clinical phenotype was only partially related with this syndrome making untangling genotype-phenotype relationship difficult. In fact, several hallmarks of 22q11 deletion syndrome lacked, such as palatal abnormalities, immuno-deficiency, cardiac and genitourinary malformations, and hypocalcaemia [2,3]. However, the patient presented motor and language delay with behavioural immaturity consistent with the syndrome and cerebellar foliation abnormalities. Brain malformations are frequent in DGS/VCFS patients, including polymicrogyria, callosal agenesis and cerebellar vermal hypoplasia (Figure 2c, d) [20]. Intriguingly, bilateral inner ear malformations were found, consisting in bilateral semicircular canal dysplasia (Figure 2e). This abnormality has been occasionally described in DGS/VCFS [21], but its clinical significance is unknown due to the lack of correlation with sensor-neural hearing loss.

This child developed a spinal cord PA (WHO grade I) at the age of 9 years. This rare neoplasm accounts for 2-5% of all CNS tumours in children and young adults [12,13]. PA may be encountered in tumour syndromes like Neurofibromatosis type 1 [22] and, rarely, Frasier and Noonan syndromes [23,24]. Only sporadic cases of brain tumour in children with 22q11 deletion syndrome were reported [5-7]. Indeed, cytogenetic studies demonstrated the association between CNS rhabdoid tumors and chromosome 22q11.2 alterations, with SMARCB1 [MIM 601607] deletions and/or mutations [7], while the first case of pleomorphic xanthoastrocytoma in a 16 year-old patient with DGS/VCFS, harbouring V600E BRAF mutation has been recently reported [6]. The identified 22q11.2 deletion was proximal and did not encompass SMARCB1 locus.

The combination of 22q11.1 deletion with a proximal 15q11.2-q13.3 duplication was described, in a case with psychomotor delay, mild facial dysmorphisms, learning difficulties, but without tumour [25]. This spinal PA likely represent a sporadic event and relationship with DGS/VCFS is uncertain. Thus, despite extent of the de novo deletion at 22q11.2 and little contribution of the paternal inherited CNVs, no features apparently correlated with either tumour development or phenotypic defects. Accordingly, the possibility of the occurrence of the spinal tumour merely by chance should be considered.

Conclusion
Paternal inherited CNVs did not lead to pathological consequences. Even no consistent genotype-phenotype correlation has been shown for the 22q11.21 deletion: a definite link between tumour and chromosomal event does not exist. Perhaps the de novo, large 22q11.21 deletion, apparently together with the inherited benign variants, may contribute to development of both spinal PA and complex phenotype.

These novel observations warrant ongoing reporting of similar cases to untangle the complex genotype-phenotype relationship with constitutional 22q11.2 deletion. The biological consequences and the contribution to cancer predisposition of chromosomal rearrangements involving CNVs of some DDR components deserve in-depth investigation.

Consent
Written informed consent was obtained from the patient’s parents for publication and any accompanying images of this case report. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

Abbreviations
DGS/VCFS: DiGeorge or Velo-cardio-facial syndrome; CNS: Central nervous system; PA: pilocytic astrocytoma; MRI: Magnetic resonance imaging; WHO: World Health Organization; CNV: Copy number variation; DDR: DNA damage response; ADCAD1: Autosomal dominant coronary artery disease 1 with myocardial infarction; ERK/MAPK: Extracellular signal-regulated kinases/Mitogen-activated protein kinases; MEF2A: Myocyte enhancer factor 2A; LYSMD4: putative peptidoglycan-binding, domain containing 4;
ADAMTS17: ADAM metallopeptidase with thrombospondin type 1 motif, 17; CERS3: Ceramide synthase 3; ARCI: Autosomal recessive congenital ichthyosis; LRPSL: Low density lipoprotein receptor-related protein S-like; DgCRC: DiGeorge syndrome critical region genes; TBX1: T-box 1; CLTC1: Clathrin, heavy chain-like 1; HIRA: Histone cell cycle regulator; RANBP1: RAN binding protein 1; CDC45: Cell division cycle 45; COMT: Catechol-O-methyltransferase; TXNRD2: Thioreredox reductase 2.

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

Authors’ contributions
SM and MS provided assistance for the clinical data, participated in the design of the study and wrote the manuscript. AR participated in coordination of the study. GM, CM, AR and PN carried out radiological and pathological data review, respectively, and gave editorial assistance. ET carried out cytogenetic studies. PDM revised the manuscript and with EM and MP gave editorial assistance. AC provided clinical data. MLG participated in the design of the study, provided clinical data and helped to draft the manuscript. VC conceived the study, participated in its design and coordination and drafted the manuscript. All authors read and approved the final manuscript.

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References
1. Tan TY, Gordon CT, Amor DJ, Farlie PG: Developmental perspectives on copy number abnormalities of the 22q11.2 region. Clin Genet 2010, 78:201–218.
2. Scamber PJ: The 22q11 deletion syndromes. Hum Mol Genet 2000, 9:2421–2426.
3. Ben-Shachar S, Ou Z, Shaw CA, Ben-Shachar S, Ou Z, Shaw CA, Belmont JW, Patel MS, Hummel M, Amato S, Tartaglia N, Berg J, Sutton VR, Lalani SR, Chinnault AC, Cheung SW, Lupski JR, Patel A: 22q11.21 distal deletion: a recurrent genomic disorder distinct from DiGeorge syndrome and velocardiofacial syndrome. Am J Hum Genet 2008, 82:214–221.
4. McDonald-McGinn DM, Reilly A, Wallgren-Pettersson C, McDonald-McGinn DM, Reilly A, Wallgren-Pettersson C, Hoyne HE, Yang SP, Adam MP, Zackai EH, Sullivan KE: Malignancy in chromosome 22q11.2 deletion syndrome (DiGeorge syndrome/velocardiofacial syndrome). Am J Hum Genet 2006, 140A:906–909.
5. Asamoto H, Fumita M: Di George syndrome associated with glioma and two kinds of viral infection. N Engl J Med 1977, 296:1235.
6. Murray JC, Donahue DJ, Malik SI, Murray JC, Donahue DJ, Malik SI, Dzunk YB, Braly EZ, Dougherty MU, Eaton KW, Biegel JA: Temporal lobe hemispheric xanthoastrocytoma and acquired BRAF mutation in an adolescent with the constitutional 22q11.2 deletion syndrome. J Neurooncol 2011, 102:509–514.
7. Beddow RA, Smith M, Kidd A, Beddow RA, Smith M, Kidd A, Corbett R, Hunter AG: Diagnosis of distal 22q11.2 deletion syndrome in a patient with a teratoid/habdominal tumour. Eur J Med Genet 2011, 54:295–298.
8. Colnaghi R, Carpenter G, Volker M, O’Dricoll M: The consequences of structural genomic alterations in humans: genomic disorders, genomic instability and cancer. Semin Cell Dev Biol 2011, 22:875–885.
9. Stankevicz P, Lupski JR: Genome architecture, rearrangements and genomic disorders. Trends Genet 2002, 18:74–87.
10. Shilen A: Malkin: Copy number variations and cancer susceptibility. Curr Opin Oncol 2010, 22:55–63.
11. Louis DN, Ohgaki H, Louis DN, Ohgaki H, Westler OD, Cavanee WK, Burger PC, Jouvet A, Scheithauer BW, Kleihues P: WHO Classification of tumours of the central nervous system. 2007th edition. Lyon: IARC; 2007.
12. Roопrapat C, Houten JK: Spinal cord astrocytomas: presentation, management, and outcome. Neurolourg Clin N Am 2006, 17:29–36.
13. Huisman TA: Pediatric tumors of the spine. Cancer Imaging 2009, 9:545–548.
14. Capra V, Mascelli S, Garri ML, Nozza P, Vaccari C, Bricco L, Cuoco C, Gimelli G, Taisano E: Parenteral imbalances involving chromosomes 15q and 22q may predispose to the formation of De novo pathogenic microdeletions and microduplications in the offspring. PLoS One 2013, 8(3):e57910.
15. Gu W, Zhang F, Lupski J: Mechanisms for human genomic rearrangements. Pathogens 2008, 1:4.
16. Jones DTW, Gronych J, Lichter P, Witt O, Pilsner SM: MANK pathway activation in pilocytic astrocytoma. Cell Mol Life Sci 2012, 69:1799–1811.
17. Rauch A, Zink S, Zweier C, Rauch A, Zink S, Zweier C, Thiel CT, Koch A, Rauch R, Lasicor J, Hufferweier U, Weyand M, Singer H, Hofbeck M: Systematic stateful of atypical deletions reveals genotype–phenotype correlation in 22q11.2. J Med Genet 2005, 42:871–876.
18. Lindsay EA: Chromosomal microdeletions: dissecting del22q11 syndrome. Nat Rev Genet 2001, 2:558–568.
19. Digilio MC, Pacifico C, Tieri L, Digilio MC, Pacifico C, Tieri L, Marino B, Gannotti A, Dallapiccola B: Audiological findings in patients with microdeletion 22q11 (di George/velocardiofacial syndrome). Br J Audiol 1999, 33:329–333.
20. Mori T, Mori K, Fuji K, Mori T, Mori K, Fuji K, Eoda Y, Miyakazi M, Harada M, Kagami S: Neuroradiological and neurofunctional examinations for patients with 22q11.2 deletion. Neuropediatrics 2011, 42:215–221.
21. Yamashita K, Yoshia H, Hidatsashita I, Tamashina K, Kama I, Inoguchi T, Honda H: Sensonarneud health loss: there is no correlation with isolated dysplasia of the lateral semi-circular canal on temporal bone CT. Acta Radiol 2011, 52(2):229–233.
22. Ohgaki H, Kim YH, Steinach JP: Nervous system tumors associated with familial tumor syndromes. Curr Opin Neurol 2010, 23(6):583–591.
23. Suttibh V, Huff V, Wolfle JE, Ketonen L, Sutibh V, Huff V, Wolfle JE, Ketonen L, Lang FF, Jr, Stewart J, Langford L, Herzog CE: Bilateral gonadalblastoma with dysgerminoma and pilocytic astrocytoma with WT1 GT-IVS9 mutation: A 46 XY phenotypic female with Fraser syndrome. Pediatr Blood Cancer 2009, 53(7):1349–1351.
24. Schuettelz LE, McDonald S, Whitesell K, Schuettelz LE, McDonald S, Whitesell K, Desrusseau DM, Grange DK, Gurnett CA, Wilson BD: Pilocytic astrocytoma in a child with Noonan syndrome. Pediatr Blood Cancer 2009, 53(6):1147–1149.
25. Manolakos E, Sarri C, Vetro A, Manolakos E, Sarri C, Vetro A, Kefalas K, Leze E, Sofocleus C, Kitsos G, Merou K, Kokotas H, Papadopoulo A, Attillas A, Petersen MB, Kitsou-Tselli S: Combined 22q11.1-q11.2 deletion with 15q11.2-q13.3 duplication identified by array-CGH in a 6 years old boy. Mol Cytogenet 2011, 4(11).

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