A 6 year old girl followed for severe mental and neuromotor retardation was examined for genetic abnormalities. The daughter of non-consanguinous parents, and sister of a normal brother, the child had important visual problems and significant facial dysmorphism. Her cranial MRI showed mesial temporal sclerosis with hypopompal sulci dilation, hypotrophy and cornual deformity. She had a cytogenomic microarray analysis performed. Her results showed a loss involving chromosome 1p13.3p13.2 (7.32 Mb), indicating a deletion in this region, region that is held important for cases that were evaluated for neurological impairment, congenital anomalies and facial dysmorphism. The cytogenomic microarray analysis also showed a loss involving chromosome 2q23.3q24.1 (4.04 Mb), indicating a deletion in this region. A de novo deletion overlapping this region has been reported to be disease causing in one patient (Malbin J et al), a deletion that has been classified as Variant of Uncertain Significance.

RapidSureDeepDive is a cytogenetic method which detects copy number variants with a much higher resolution than conventional cytogenetic analysis. This technology, which does not require staining or cell culture, uses specially designed chips to study the DNA from various sample types. The deletions found in our patient are quite rare, and maybe the variant of the second deletion will be reclassified as disease causing if proved to be denovo.

A de novo deletion overlapping this region has been reported to be disease causing in one patient (Malbin J et al), the deleted region does not include MBD5, a gene commonly implicated in intellectual disability and hence the deletion has been classified as Variant of Uncertain Significance. The variant will be reclassified as disease causing if proved to be denovo.

Williams-Beuren syndrome (WBS) is developmental disorder caused by microdeletion of genes from chromosome 7. It is estimated in 1 of 10 000 people. WBS could be autosomal-dominant inherited, but usually, it is caused by de novo microdeletion of 26-28 genes on chromosome 7. Usually genes such as CLIP2, NCF1, ELN, GTF2I, GTF2IRD1 and LIMK1 are deleted. ELN gene is detected as a cause of supravalvular aortic stenosis, while absence of NCF1 gene is related to hypertension. Patients with WBS are characterized by cardiovascular diseases, facial dysmorphic features, intellectual disability, unique personality character and endocrine abnormalities.

Case Report We present the case of three patients, two girls at the ages of one and three years and a boy at the age of two years. All three patients have healthy parents and brothers and sisters without known chronic or genetic diseases. All three were presented with facial dysmorphic features which include broad nasal bridge, microretrognathia, large mouth, upper lip is thin while lower lip is thicker. All three patients are cognitively deficient and they have speech problems. The female patient at the age of three years also has hypertension, small teeth, larger neurocranium, atrial and ventricular (muscular) septal defect, gastroesophageal reflux disease (GERD), hypothyroidism and endocrine disorders; hypercalcemia, hypercalciuria, hypothyroidism and early puberty. The male patient also has prominent ears, palmar crease on left palm, pulmonary stenosis, supravalvular aortic stenosis with mitral regurgitation and bilateral inguinal hernia. All three patients were suspects for WBS. Karyograms of all patients were normal, but FISH analysis performed with ELN primer for detection of WBS proved microdeletion on chromosome 7 in region 11 (7q11).

Conclusion WBS is rare disorder characterised by wide range of symptoms and signs. Clinical diagnostic criteria are available for WBS, but confirmation of diagnosis requires detection of microdeletion on chromosome 7q11.2. Early recognition of specific patterns of the disorder, such as facial dysmorphic features together with heart abnormalities, cognitive deficiency and vision problems, is crucial because these patients need to take speech therapist and psychologist therapy as soon as possible because it improves their integration in social environment. Management is focused on treatment of symptoms (eg. hypertension, hypercalcemia), psychological and psychiatric evaluation and speech therapy.