and development, hypotonia, intellectual disability and seizures. Its frequency is estimated at 1/20000 to 1/50000 births with a female predilection. It’s due to a genetic material deletion near the short arm of chromosome 4 (4p-). Through this case report we aim to remind the mains and associated features of this rare disorder as well as its management.

**Methods** We report the case of a male newborn with a Wolf-Hirschhorn syndrome.

**Results** A full term newborn was born by c-section indicated for a severe growth restriction. Birth weight was 1850 g, length was 45 cm and head circumference was 30.5 cm. Clinical examination at birth noted an important hypotonia, cyanosis with no associated signs of respiratory distress, distinctive facial features including a broad and flat nasal bridge, a high forehead, widely spaced eyes with iridian coloboma, poorly provided eyebrows, poorly formed ears with a pre-tragic fistula, micrognathia, thin lips and posterior cleft palate associated to a spina bifida occulta, posterior hypospadias, testicular ectopia and mottled skin. Echocardiography showed a type interruption of the aortic arch. The ultrasound examination of brain and abdomen was normal. The genetic tests concluded to a Wolf-Hirschhorn syndrome with a (4p) deletion. The boy died at the age of one month due to a severe bronchiolitis. Parental genetic tests are planned.

**Conclusion** Wolf-Hirschhorn is a rare genetic disorder. It may be inherited but generally it occurs as a random event. Its outcome depends on associated malformations especially congenital heart disease. We insist on the importance of genetic counseling and antenatal diagnosis when index cases are registered.

**P120 HYPTERTENSION AND ACCESSORY RENAL ARTERIES**

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**Introduction** Arterial hypertension is a major risk factor for cardiovascular, cerebrovascular and renal morbidity and mortality. Renovascular disease is the third most common cause of hypertension in children after coarctation of the aorta and parenchymal renal disease. The spectrum of renovascular disorders in children is very broad compared to adults. Besides fibromuscular dysplasia and renal artery stenosis as the main causes, narrow/hypoplastic accessory renal artery or multiple accessory renal artery or multiple renal arteries could also be involved in ischemic nephropathy.

**Case report** We describe the case of a 15-year-old girl who presented with palpitations, chest pain and headache. The physical examination revealed a high blood pressure (over the 99th percentile for age, gender and height). She had insignificant personal and family medical history regarding the cardiovascular risk factors. Laboratory analysis and relevant radiological evaluation were obtained in order to determine the etiology for suspected secondary hypertension. Direct plastic renal, plastic aldosterone, urinary catecholamines, plastic metanephrines, vanilmandelic acid, thyroid function tests were all within normal limits. In order to rule out possible renal artery stenosis, CT angiography of the renal artery was performed, surprisingly demonstrating a small caliber (hypoplastic) accessory right superior renal artery.

**Conclusion** Physicians should be alert to the possibility of the causal role of accessory renal arteries in patients with severe and difficult to manage hypertension. For several reasons this possibility may be greater in children.

**P121 TO SCAN OR NOT TO SCAN ...AGAIN**

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A 13 year old male was referred to our regional centre for review of a left sided periorbital cellulitis by ENT, ophthalmology and paediatric teams. Left eye pain began 4 days previously and was initially felt to be secondary to allergies. However, one day prior to transfer to our unit intravenous cefotaxime was commenced at referring hospital and neuroimaging (CT Brain) was performed; this revealed a preseptal periorbital cellulitis with no intracranial extension. Upon arrival he was unwell and lethargic and still spiking temperatures whereupon his antimicrobial cover was broadened to include i.v fluoxacillin and metronidazole.

Inflammatory markers were elevated with a CRP of 227 mg/L (0–5) while white cell count was slightly elevated at 13x10^9/L (4–10). Initial blood cultures were negative but repeat blood cultures taken after 3 days of antibiotics were positive with a pathogenic gram negative organism. Following discussion with microbiology colleagues antibiotic coverage was broadened with an initial improvement over 24 hours but subsequently fevers recurred with increasing lethargy but no focal neurological signs. Inflammatory markers were slowly falling. In view of the clinical deterioration repeat imaging was requested, despite a previously normal CT Brain. MRI revealed a subdural abscess anterior to the left frontal lobe and extensive left sided sinusits. He was emergently transferred for neurosurgical review and urgent surgery to drain the abscess and nasal sinusits. He recovered well following 6 weeks of intravenous antibiotics.

Subdural abscess is a rare but recognised complication of periorbital cellulitis, particularly secondary to sinusits. A systematic review of the intracranial complications of rhinosinusitis revealed that the vast majority (70%) occurred in young adolescent males with a morbidity rate of 27% and a mortality rate of 3%. In our case repeat neuroimaging was undertaken due to a clinical deterioration and allowed for timely surgical intervention with a good outcome.

**P122 CELIAC DISEASE ASSOCIATED WITH DEPRESSION IN ADOLESCENTS**

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**Introduction** Celiac disease is an immune-mediated systemic disorder elicited by gluten which causes malabsorption and small-bowel mucosal inflammation in genetically susceptible
individuals. Although the gastrointestinal tract is the target of autoimmune insult, celiac disease is also associated with extraintestinal problems: autoimmune disorders, malignancies, dermatologic conditions, rheumatologic conditions, neurologic and psychiatric disorders. The psychological symptoms are one of the most interesting and unexpected presentations, which may be the result of nutrient malabsorption or increased levels of proinflammatory cytokines.

Case report We present the case of a 15 year old adolescent girl with a history of autoimmune thyroiditis on replacement therapy with levothyroxine, who was hospitalised because of behaviour disorders and oligobradimenorrhea. On clinical examination the patient presented drowsiness, sudden behavioural changes, learning problems, bradydalia and a tendency to isolate herself. Blood tests showed positive anti-TG antibodies (>10x normal) and EMA antibodies. HLA typing was also performed and was positive for HLA-DQ2 cis: DQA1*05 - DQB1*02 - DRB1*03. On the psychiatric consult she was diagnosed with a minor depressive syndrome. She was immediately started on a gluten free diet. Although she showed low compliance to the diet on the follow-up examinations, removing gluten resulted in positive effects on the psychiatric symptomatology. We would also like to add that the patient refused the oesophagogastroduodenoscopy and psychological counselling.

Conclusion Recent studies have shown an association between celiac disease and psychiatric conditions, especially depression. Medical personnel should be aware of this atypical presentation of the disease in order to correctly diagnose and treat celiac disease. Also, psychiatric patients who are resistant to traditional therapy should be investigated for celiac disease in case they also present symptoms of CD or if a positive family history exists.