has been previously reported in the past. Detection of a high rate and variability of ETAs associated with CH necessitates a vigilant clinical and diagnostic approach to all babies with CH.

**P286 SPECIFICITY OF MEDULLAR THYROID CANCER IN SIPPP’S AND GORLIN’S MEN SYNDROMES IN PEDIATRIC PATIENTS (8 CASES)**

Anatoly Romanchichen*, Alexandre Gostimskii, Zoya Matveeva, Kristina Vabalayte, Sergei Penedereev, St. Petersburg State Pediatric Medical University, St. Petersburg, Russian Federation

Methods 31325 thyroid patients were operated in the St.-Petersburg Center of Endocrine Surgery and Oncology during 1974 – 2015. Among them 18 (0.06%) cases of MEN were detected, including 8 (0.02%) children. The diagnosis of MEN2 was confirmed by genetic investigation in all 8 children (3 girls, 5 boys) 4–18 years old. Genetic research confirmed syndrome in 7 children, syndrome – in 1 girl 10 years old.

**Results and discussion** The reason for examination of 6 children was MEN identified in adult family members operated on for medullary carcinoma. In 2 cases genetic study was conducted after detection of medullary cancer in pediatric patients. Three patients operated on in childhood belonged to the same family, 5 members of which had syndrome MEH-2a in three generations. Multifocal medullary thyroid cancer with cervical metastases was detected in a girl with MEN-2b. Before surgery, 4 (50%) children did not have manifestations of medullary cancer. Microcarcinoma identified intraoperatively in 1 boy. Increase of the calcitonin levels and thyroid nodules were diagnosed in 4 cases. All 8 patients underwent thyroidectomy, in 7 cases – with central lymphadenectomy. Lateral lymphadenectomy was performed in 2 cases, including 1 Gorlin’s syndrome (RET p.M918T). Histological examination showed normal thyroid in 2 children, C-cell hyperplasia – in 1, medullary cancer – in 5. Multifocal medullary thyroid cancer T4N1M1 was revealed in the patient with MEN-2b. She was operated on 3 times for relapses lymphatic node metastases removing. Medullary thyroid carcinoma and Parathyroid adenoma together were removed in 1 child.

Follow-up results were search in all our pediatric patients operated on during 1989 – 2018 years. All patients are alive. Only in one case we observed hypercalcitoninemia up to 1000 pg/ml after 3 repeated operation on the neck and bayside adrenalectomies.

**Conclusion** Thyroidectomy had prophylactic character only in 3 children 4–10 years old. Early thyroidectomy (under 5 years old) has prevented the development of medullary thyroid cancer in children and adolescent.

**Introduction** Monoallelic pathogenic variants in HNF1A are a well-established cause of maturity-onset diabetes of the young 3 (MODY3). These variants are rarely reported as a cause of neonatal hyperinsulinaemic hypoglycaemia (NHH).

**Case Description** A term infant was delivered by spontaneous vaginal delivery with a birth weight of 3.53 kg (50th centile). He was born to a Gravida 2 Para 1 mother who had manifested gestational diabetes that necessitated insulin therapy in the first trimester of both of her pregnancies. His older sibling also had transient neonatal hyperinsulinism that fully resolved at 1 month of age.

He was admitted to the NICU for the management of transient tachypnoea of the newborn. The infant developed symptoms of hyperglycaemia at 1 hour of life and blood glucose level was measured at 0.4 mmol/L. He continued to become hyperglycaemic despite high glucose infusion rates (GIR). Critical samples were collected whilst hyperglycaemic (plasma glucose 1.7 mmol/L) on day 3 of life confirming the diagnosis of non-ketoic hyperinsulinaemic hypoglycaemia. Given the family history samples from the infant and his parents were sent to Exeter Clinical Laboratory for hyperinsulinism genetic testing.

He was commenced on diazoxide therapy at 3 mg/kg/day on day 17 of life and following this therapy his blood glucose levels normalised on 4 hourly oral feeds (GIR = 8.3 mg/kg/ min). He successfully tolerated an 8 hour safety fast and was discharged home on day 31 of life. At 7 weeks of life he presented unwell, febrile and hyperglycaemic (blood glucose range 18–22 mmol/l). He had a urinary tract infection and was commenced on antibiotic therapy. His diazoxide dose was discontinued. Close blood glucose observation over a 7 day period and a repeat safety fast demonstrated normoglycaemia.

Genetic testing subsequently demonstrated that both the infant and his mother were heterozygous for a pathogenic HNF1A missense variant (NM_000545.6:c.788G>A:p.Arg263His).

**Discussion** NHH causes significant morbidities including adverse neurodevelopmental sequelae and mortality if untreated. Transient NHH is commonly observed in infants of mothers who have gestational diabetes. HNF1A gene mutation however is a rare cause of transient NHH. Identification of this genetic diagnosis has had major implications for this infant and his family. His mother is currently being appropriately treated for MODY3 and has been counselled about the 50% recurrence risk in subsequent pregnancies. Both children will be observed closely from late childhood due to their risk of developing MODY3.

**P287 TRANSIENT NEONATAL HYPERINSULINAEMIC HYPOGLYCAEMIA SECONDARY TO A HNF1A GENE MUTATION**

1Sabrina Sheridan*, 1Laura McCarron, 1Gillian O’Donnell, 1Eithin Carolan, 1Nuala P. Murphy. 1Department of Paediatric Endocrinology, Children’s University Hospital, Temple St, Dublin, Ireland; 2School of Medicine, University College Dublin, Dublin, Ireland

**Background** Atopic dermatitis (Eczema) is a common chronic remittent inflammatory skin condition. Emollients and topical corticosteroids are the mainstay of treatment and are widely prescribed. This report underlines the danger of combining prescribed and homeopathic treatments and the importance of monitoring steroid prescribing.
Case Report The index is a 2.5 year old girl with moderate eczema associated with nocturnal itch. She was referred to local services and was trialled on topical Clobetasone and Betamethasone cream. After little improvement this was changed to topical Hydrocortisone (1%) and Betamethasone cream. Her parents applied the steroid creams on a continuous basis for at least four months. They also sourced different homeopathic ingredients from Pakistan. Upon analysis of their ingredients in the laboratory, Betamethasone and Dexamethasone were identified in one of them. She required a 5 day course of oral Prednisolone for an allergic reaction to nuts. Over time the eczema remitted but all treatments were continued. After six months, she presented with rapid weight gain over a 2 month period [weight >98th centile]. This was associated with increased skin pigmentation over ankles and knees, striae over her thighs and marked hypertrichosis.

Biochemical testing confirmed absence of diurnal variation with suppressed 8am cortisol levels. She was switched to maintenance eczema care with emollients and topical Tacrolimus. An ACTH stimulation test confirmed adrenal suppression [peak Cortisol 45nmol/L]. Adjunctive investigations for other causes of hypoadrenalism were reported normal. We educated the family on emergency management of cortisol deficiency and prescribed steroid replacement for emergency use only. After four weeks off all steroid containing medications, a repeat ACTH stimulation test confirmed normalisation of adrenal function [peak Cortisol 530nmol/L]. Four months later, her weight has normalised [75–91st centile] and her eczema has remained under control. The pigmentation and striae have faded and her hypertrichosis is no longer evident.

Discussion The role of corticosteroids in the management of inflammation is well established but has resulted in widespread, prolonged and often unsupervised use. There is interindividual variation in steroid sensitivity, small doses can result in dramatic adverse effects. We did suspect that the homeopathic medicine was steroid based. This does not rule-out the possibility that the other ingredients facilitated steroid increased absorption or reduced metabolism.

Conclusion Adrenal suppression is a rare side effect of topical steroid use. Children on courses of steroids for longer than one week require medical monitoring to ensure that the treatment is appropriate, safe and used correctly.

P289 METABOLIC COMPLICATIONS OF OBESITY IN CHILDREN

Irina Osokina*. Science Centre of the Siberian Branch of Russian Academy of Science, Institute for Medical Studies of the North, Krasnoyarsk, Russian Federation

Introduction Children with obesity have high risk of metabolic complications.

Aim To reveal frequency of metabolic complications in children with obesity.

Methods We examined 91 obese children, aged 7–17 (BMI > 95 percentile). Mean age of children was 12.46 ± 3.5 yrs. Obesity in children manifested at 5.3±0.5 yrs. Family anamnesis characterized by obesity in 71%, diabetes type 2 - in 25%, hypertension - in 53% in the first degree relatives. All patients were examined by clinical, biochemical, ultrasound methods. An oral glucose tolerance test (OGTT) accompanied by four point of insulinema was performed. HOMA index was calculated according to the standard formula. In children with fatty liver chronic hepatitis were excluded. Metabolic syndrome (MS) was diagnosed according to a classical definition (Weiss’s criteria).

Results In obese children metabolic complications were found in 39.6%. The prevalence of the metabolic complications was as follows: hypertriglyceridemia 38.2%, glucose intolerance 17.6%, hypertension 52.6%. Increase of serum cholesterol was revealed in 24%, low density lipoproteins - in 14%, decrease of high-density lipoproteins – in 32% children. Metabolic syndrome was found in 18 (19.8%) patients with BMI 30.9 ± 3.4 kg/m². Insulin resistance revealed in 25% children. HOMA index was 4.6 ± 3.3 mU/L. Ultrasound signs of fatty liver were shown in 40 patients.

Conclusions This study showed a high prevalence of metabolic complications among obese schoolchildren.

P290 POSSIBILITIES OF GENETIC RESEARCH IN THE PREVENTION OF THE DEVELOPMENT OF EXCESS MASS OF THE BODY AND OBESITY IN CHILDREN

1Elena Bulatova, 1Olga Lagno, 1Alexander Shabalov*, 2Valeriy Polunovskiy, 1Ludmila Selenstsova. 1St. Petersburg State Pediatric Medical University, St. Petersburg, Russian Federation; 2LLC Natsionalnyi Tsentr Geneticheskikh Issledovaniy, Novosibirsk, Russian Federation

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Relevance Conducting genetic research in pediatric practice can help to create personalized dietary and therapeutic recommendations, as well as the implementation of appropriate physical activity.

Objective To determine the presence of polymorphisms of genes that contribute to the development of excess fat mass, increase the speed of carbohydrate absorption in patients with overweight and obesity to create further personalized recommendations on diet therapy and physical activity.

Patients and methods Surveyed 9 children (3 boys and 6 girls) aged from 6 to 10 years. Anamnesis was collected, an objective examination of patients, a biochemical examination of blood (lipidogram, fasting glucose), and a hormonal examination (insulin, TSH) were performed. All children underwent genetic screening based on DNA analysis (material - buccal epithelium) with the definition of gene polymorphisms: the FABP2 gene, which regulates the rate of fat absorption; PPARG gene, affecting the growth rate of fat cells; gene ADRB2, affecting the rate of carbohydrate consumption in the blood during exercise; The TCF7L2 gene, which regulates insulin secretion rate when glucose is released.

Results 4 patients were overweight, grade 1 obesity - 2; 2 degrees - 2; Grade 3 obesity - 1 child. In all nine patients, unfavorable gene polymorphisms were found, contributing to impaired fat metabolism. In 5 children, according to DNA analysis, a reduced rate of consumption of carbohydrate reserves in response to an increase in the level of adrenaline in the blood was noted. 4 children showed a genetic predisposition to a decrease in the rate of insulin secretion in response to glucose intake. In a 3rd degree obese patient, all the studied gene polymorphisms that adversely affect carbohydrate and fat metabolism were found.

Conclusion All patients showed a genetic predisposition to impaired metabolism of fats and/or carbohydrates. Grade 3 obesity is associated with all modifications of genes that adversely affect both carbohydrate and fat metabolism, and