the possible prohibition of performing sports, to the results of surgery as scars, a possible source of shame.

Parents of children with heart disease may experience higher stress levels than normal parental function and may feel very stressed about issues related to accountability and social integration.

Methods This research aims to investigate the quality of life in pediatric age both in terms of purely cardiological aspects (CardioPeds) and of general quality of life (PedsQL), also evaluating, starting from 12 years, the possible presence of depressive symptoms (PHQ-9) or anxious symptoms (GAD-7). These tests were administered both to the children and the parents, with the addition for these last ones of the compilation of the PSI, to investigate the parental distress.

The research allowed to divide the sample of 500 patients into 6 predominant pathologies, in order to compare the quality of life of children in different diseases and to understand if a specific pathology is associated or not with a lower quality of life.

Results Through an adequate statistical analysis it was found that the quality of life perceived by the subjects included in the study is significantly better than that perceived by the parents about the same children and adolescents.

Conclusions This study was one of the first to investigate the quality of life in congenital and/or acquired pediatric heart disease. However, the results obtained require further studies, in order to deepen what has emerged.

**GP35 STUDY OF THE STIFFNESS OF THE VASCULAR WALL IN CHILDREN FROM FAMILIES WITH A BURDENED HISTORY OF CARDIOVASCULAR DISEASES**

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Medical emergency center in Kazan conducted targeted screening to select patients with ischemic strokes, myocardial infarctions, angina pectories, widespread atherosclerosis of peripheral vessels, angiographically confirmed coronary atherosclerosis, ischemic heart disease and dislipidemia. 230 patients were selected (men ≤55 years, women ≤ 60 years). Their children, grandchildren (144 people) (Group 1) aged 5–17 years were examined. Control group of conditionally healthy children (112 people) was formed (Group 2), their family history was collected with the aim of excluding heredity burdened by CVD. A biochemical blood test was performed to determine the lipid profile; thus, a group with dyslipidemia (Group 1a) was selected among children with burdened family history, which included 76 people (53%), lipidogram indicators (Group 1b) that were normal (68 people, 47%). Rigidity indices of main vessels were determined: PTT (pulse wave propagation time), PWVao (pulse wave propagation speed in the aorta), AIx (augmentation index), (dP/dt) max (max blood pressure rise rate). An ambulatory daily monitor, BPLab® software using Vastosyn® technology was used.

Results In general, in Group 1, the rigidity indicators of the vascular wall were similar (average daily PTT 145.3 ± 15.9 ms, PWVao 6.7 ± 0.9 m/s, AIx 4.9 ± 8.1%, (dP/dt) max 667.9 ± 119.8 mm Hg/s) with those of Group 2 (average daily PTT 140.5 ± 15.2 ms, PWVao 6.4 ± 1.1 m/s, AIx 5.1 ± 8.2%, (dP/dt) max 668.1 ± 121.3 mm Hg/s) and were statistically significant (p < 0.05). However, it was found that if we consider non-maximum daily indicators, then in Group 1 they were significantly higher (p < 0.05) (max daily PTT 199.9 ± 10.2 ms, PWVao 16.3 ± 1.4 m/s, AIx 35.3 ± 3.4%, (dP/dt) max 1098.1 ± 120.5 mm Hg/s) than in Group 2 (max daily PTT 181.7 ± 9.6 ms, PWVao 13.1 ± 1.9 m/s, AIx 28.3 ± 1.4%, (dP/dt) max 1005.1 ± 110.3 mm Hg/s). Comparing the parameters between Group 1a and 1b, no statistically significant difference was found between the mean values, but statistically significant difference was found between the maximum values.

Conclusions Direct relationship has been revealed between the presence of a burdened family history of CVDs, changes in the lipid profile of parents, relatives of the 1st and 2nd lines, and changes in the rigidity of the wall of the great vessels in their children. The rigidity of the vascular wall was significantly higher in Group 1 than in Group 2, differed between children from Group 1a and Group 1b.
or inadequate intake. Children with feeding problems also tended to eat less than children without feeding problems. There was a trend towards more feeding problems in patients with chromosomal abnormalities or other associated anomalies.

**Conclusion** Feeding disorder is often and frequent long-term sequel in children after neonatal or early infancy heart surgery. Patients with chromosomal and associated anomalies and those who underwent multiple cardiac surgeries are at risk of developing feeding difficulties and later feeding disorders. These patients need to be selected for preventive strategies and nutritional intervention should be offered in order to increase the caloric intake of the child and to develop a sound feeding relationship in the family.

**GP38** SPECKLE TRACKING AS A MODERN ECHOCARDIOGRAPHIC Technique IN ASSESSING CARDIAC FUNCTION IN CHILDREN

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**Introduction** Speckle tracking is a new method of assessing cardiac function in adults, but used only at a research level in children. The aim is to implement this method in children with congenital and acquired cardiac pathology and oncologic disease as a new method of detecting cardiac function deterioration.

**Methods and material** We selected patients with congenital or acquired heart disease and oncologic pathology then performed echocardiography, cardiac biomarkers and speckle tracking and compared the results.

**Results** Currently we have 31 patients out of which 55% had a cardiac pathology, 26% had an oncological disease and 19% were collagenases or vasculitides. However, the study is ongoing and we hope to achieve a larger patient pool.

When considering the patients with congenital heart disease 47% had modified GLS(global longitudinal strain) and cardiac biomarkers, but a normal EF(ejection fraction). Aortic coarctation, aortic regurgitation, complex cyanogenic malformations, ventricular septal defect, complete AV block, hypertrophic cardiomyopathy and dilated cardiomyopathy were the maladies with altered GLS but with normal ejection fraction.

In the case of the complete AV block, she had a new dual chamber pacemaker installed, after which she developed symptoms of cardiac failure, GLS was under normal values, EF was 54% and had a septal dysynchrony, however after modifying the device’s parameters the GLS was normal and there was no septal dysynchrony.

Patients with chemotherapy frequently develop cardiac toxicity, 44% of patients were diagnosed with cardiotoxicity with the help of speckle tracking and cardiac biomarkers.

One patient with Rhabdomyosarcoma with an initial normal GLS, chemotherapy was initiated, after which he accused palpitations. The GLS was at the lower normal limit, thus confirming cardiotoxicity due to Doxorubicin.

One patient with LES was monitored by speckle tracking. Initial normal GLS, was altered during a disease flare after which pericarditis developed.

We had two patients with Kawasaki disease complicated with severe coronary aneurysms which were followed-up by speckle tracking.

**Conclusions** Speckle tracking can be used as a fast and safe method to determine cardiac function in children, being well correlated with cardiac biomarkers, even when EF is normal.