Nerve injuries are a common complication of upper extremities fractures. Especially the long arm nerves are prone to injuries. After such an injury the muscles are denervated. Fortunately, if nerves are injured but the continuity of the nerve sheath is intact a regeneration by spreading of the axonal fibres starts early after the injury and finally a new neuromuscular connection – a neuromuscular endplate- is reestablished. Till today it is unknown whether the newly formed neuro-muscular endplate is functional equivalent to the original. During the last years a new technic became routinely available in paediatric neurology to study the neuro-muscular endplate: stimulated single-fibre-EMG. This technic allows a precise and objective assessment of the neuro-muscular connection. At our hospital we care together with our colleagues from the paediatric surgery department for children with fractures and nerve injuries. During the last years we have adopted the technique of single fibre EMG and included it into our clinical repertoire for traumatic nerve injuries.

It has been previously hypothesis that the newly formed neuromuscular endplate is not as reliable as the original one. This fact would have major clinical implications for the support and training after nervous injuries. To address this question, we have set up this pilot project to conduct routine measurements of the recovering nerve using single fibre-EMG. Our preliminary data shows, that these measurements can be reliable conducted in the pediatric population and we are now in the process of applying for ethical approval to analyse a larger cohort.

In terms of diagnostic procedures, 24-hour oesophageal pH monitoring was positive in all the cases of Sandifer’s where it was performed, while upper GI endoscopy ± biopsy and barium X-ray were diagnostic only in a subset of cases.

A range of treatment options were applied in the reviewed literature, including dietary changes, pharmacological management, enteral tube feeding, and surgical approach. These treatment options are consistent with the 2015 NICE guidelines on management of GORD in children and young people.

Successful treatment of the underlying gastro-oesophageal pathology led to a complete or near-complete resolution of the neurological symptoms in all of the cases.

**Discussion**

As Sandifer syndrome is driven by the underlying gastro-oesophageal reflux, it is not surprising that its investigations and management showed to be consistent with those of GORD. It is evident from the literature that many patients were originally misdiagnosed with various neuropsychiatric diagnoses that led to unnecessary testing and ineffective medications with significant side effects. Earlier diagnosis of Sandifer’s would have allowed to avoid them.

Rasmussen syndrome is a subacute inflammatory encephalitis and one of the causes of continual partial seizures. The onset of convulsive attacks is usually preceded by an episode of non-specific fever, usually of viral origin (eg, CMV infection). Attacks are usually partial, long-lasting and frequent. It most often occurs in children under 10 years of age. EEG findings show diffuse paroxysmal activity. The disease is progressive, with development of various neurological deficits (hemiplegia, hemianopsia, and aphasia). Here we present a case of a 3-year-old girl, with normal early psychomotor development, who had the first absence-type convulsive attack during her 10th year. Corticosteroids were also administered during the exacerbations. In a further course of the disease, a clinical MR of the endocranium showed the excision of a hypertrophy of the left hemisphere and discreet left-sided Walerian degeneration, when diagnosis of Rasmussen syndrome was suspected. Further diagnostic procedures included immunological examination of the liquor and the presence of oligoclonal IgG strains in the liquor (LGI1, AMPA, GABA-B) have been introduced into the therapy. In March 2017, new MR of the endocranium showed the existence of a hypertrophy of the left hemisphere and discreet left-sided Walerian degeneration, when diagnosis of Rasmussen’s syndrome was suspected. Further diagnostic procedures included immunological examination of the liquor and the presence of oligoclonal IgG strains in the liquor (LGI1, AMPA, GABA-B) was confirmed. Anti-NMDA autoantibodies were negative, and therapy with specific immunoglobulins was administered. Corticosteroids were also administered during the exacerbations. In a further course of the disease, a clinical finding reveals a worsening of neurological status (spastic right-sided hemiparesis, facial nerve paralysis and dysarthria). At the end of 2017, the girl had an average of 4-5 convulsive attacks per month, which led the patient to a decision for hemisperectomy on the left side.