Bilateral dysfunction of superior laryngeal nerves as a consequence of cervical necrotising fasciitis

Obojestranska disfunkcija zgornjih grlnih živcev kot posledica cervikalnega nekrozantnega fasciitisa

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Introduction

Necrotizing fasciitis (NF) is a life-threatening, rapidly-spreading soft tissue infection\(^1\)\(^2\) that causes progressive necrosis of deep fasciae, leading to necrosis of neighbouring soft tissues.\(^3\) The head and neck are involved in 2.6% of cases.\(^4\)\(^5\) According to Krenk,\(^6\) Wolf\(^2\) and Lanišnik\(^7\), the incidence of cervical NF (CNF) is 0.0001–0.0002. It can be either idiopathic or secondary;\(^4\) in the latter case, it is caused by infections, trauma or surgery.\(^2\)

The infection process begins 48 hours after an initial insult.\(^8\) Early clinical signs and symptoms are nonspecific.\(^4\) Initially, there is
Klinični primer/Case report

Figure 1: Axial CT image of the neck demonstrating fluid and gas collections, thickening of the subcutaneous tissues and cervical muscles, fluid and gas collections and fascial plane blunting or dissection. The presence of gas within the soft tissues in the absence of previous surgery, radiotherapy or trauma is a pathognomonic sign. In edentulous patients, CT scan can also establish eventual odontogenic origin of infection. In dentate patients with dental fillings, however, panoramic radiograph of the mandible is additionally recommended, due to artefacts on the CT scan. Another diagnostic option is surgical approach, i.e. a 2-cm-long neck incision through the platysma. If loose tissue planes, friable muscles, murky dishwater discharge, minimal resistance after finger insertion and no bleeding are found, the CNF is diagnosed.

Intravenous broad-spectrum antibiotics should be given in high doses. Initially, an antibiotic effective against aerobic and anaerobic bacteria infection is prescribed. Later on, the therapy should be targeted against the microorganisms isolated from the microbiological samples. Nutritional, hemodynamic and intensive care supportive measures are vital. Surgical treatment is the key intervention. The excision and debridement must be quick and substantial in order to decrease the bacterial load, halt both the release of inflammatory mediators and the spread of necrotizing process. The excision with fasciotomy should be continued until the tissues bleeding freely on incision are encountered. Two adjunctive treatment modalities that could theoretically reduce necrosis development are intravenous Immunoglobulin Gamma and Hyperbaric Oxygenation.

The prognosis of CNF is grave as dangerous complications may appear in a matter of days leading to descending necrotizing mediastinitis, multi-organ failure, septic shock and death. The lesion of the cranial nerves as a complication of CNF has only occasionally been reported in the literature. Cutilli and Chueng stated CNF can extend into a patchy, smooth, shiny erythema of the cervical skin, without sharp demarcation from uninvolved skin. It is accompanied by edema, crepitation, tenderness, pain on palpation and fever. The pain is more severe than expected, and may be felt deep in the muscles, but the affected skin area may also be insensitive. Later on, the skin becomes dusky, covered with blisters and bullae and finally necrotises. If left untreated, the skin breaks down and copious purulent debris and “dishwater” fluid ensue. The sepsis develops within 48 hours of symptom onset. Laboratory tests show a very rapid rise of CRP values and leukocytosis with a left shift. The CT scan of the neck and chest shows specific signs indicating the CNF and its extent, and offers anatomic information necessary for surgical planning. CT findings predicting CNF include diffuse thickening of the skin, subcutaneous tissues, fasciae and muscles, fluid and gas collections and fascial plane blunting or dissection.
Wolf² reported the neural deficits of the accessory nerve, hypoglossal nerve and of marginal branch of the facial nerve in 35%. Lanišnik⁷ reported partial facial nerve palsy in 33%. To the best of our knowledge, the dysfunction of the superior laryngeal nerve from CNF has not yet been reported.

Case report

A 71-year-old edentulous female with arterial hypertension and diabetes mellitus type II suffered from two-day-long dysphagia, odynophagia and hoarseness without fever. The transnasal flexible endoscopy revealed normal mobility of the larynx with barely notable mucosal swelling of the retrocricoid area and both piriform sinuses, and copious accumulation of the saliva in the hypopharynx. There were erythema, soft swelling and crepitations present over the prelaryngeal skin and the upper chest. It was excruciatingly tender to palpation.

The laboratory blood tests showed leukocytosis \(18.52 \times 10^9/L\) with left shift neutrophilia \(16.12 \times 10^9/L\), elevated CRP \(330\,\text{mg/L}\) and procalcitonine levels \(9.85\,\text{mg/L}\). An urgent CT scan of the neck and chest showed some fluid collections and widely spread collections of gas, thickening of the subcutaneous tissues and cervical muscles, and signs of soft tissue dissection, which is characteristic for CNF (Fig. 1). Moreover, the gas collections and minor streaks of fluid collections were also seen in the anterior mediastinum all the way to the tracheal carina, i.e. signs of descending mediastinitis (Fig. 2). No further diagnostics was performed. Empiric intravenous antibiotic therapy with amoxicillin-clavulanic acid and metronidazole was started immediately.

Based on the clinical signs, the radiological findings and the laboratory test results, NF was diagnosed and a surgical procedure under general anaesthesia was planned. The transverse collar incision was carried out and a skin flap in the subplatysmal plane was elevated. Dirty brown-black dishwater and a typical foul odour were noted. The subplatysmal fatty tissue, the prelaryngeal muscles, the superficial part of the thyroid gland, the lymph nodes and fatty tissue of nodal levels III and IV in the left, the parts of the left scaleni muscles had already necrotized (Fig. 3). All necrotic tissues were removed until freshly bleeding tissue was reached. Drains were placed into the parapharyngeal, retropharyngeal and the left pterygoid spaces and a transcervical digital mediastinotomy was performed. Malodourous brown-black dishwater again poured out of all aforementioned neck spaces and the mediastinum. The microbiologic samples and a biopsy of the affected tissue were taken. At the end of the procedure, direct laryngoscopy and tracheostomy were performed and a nasogastric feeding tube was inserted. Even after extensive discussion with the patient and her relatives the origin of CNF could not be found.

The microbiological culture from the wound smear obtained during the surgery revealed the following microorganisms: *Streptococcus constellatus ssp. constellatus*, *Lecleria adecarboxylata*, *Lactobacillus catenaformis*, *Prevotella intermedia*, *Prevotella buccae* and *Parvimonas (Micromonas) micra*. According to this finding and the sensitivity of microorganisms, we changed the antibiotics for intravenous clindamycin and cefotaxime. The histological finding of the biopsy confirmed our clinical suspicion of NF.

After the surgical procedure, the patient’s voice became breathy due to inadequate...
Figure 4: Slow healing of the surgical neck wound on the 12th postoperative day.

Figure 5: The patient’s neck with the surgical wound healed two and a half months after surgery.

glottic closure. In a few weeks, her voice became normal. However, several serious complications, such as slow healing of the neck wound (Fig. 4), pleural effusion, pneumonia, acute renal failure, septic shock, depression and dysphagia occurred. All of them, except the dysphagia, resolved within four weeks of intensive medical treatment, supportive measures and rigorous daily wound care (Fig. 5).

The time from CNF surgery and tracheostomy to decannulation was 40 days. Subsequent fiberendoscopic evaluation of swallowing showed that there was a massive spillage of liquids and semiliquids over the aryepiglottic folds and posterior commissure into the larynx (Fig. 6). The coughing reflex appeared when the bolus passed the larynx and entered the trachea. Rough sensibility testing with the tip of the instruments, e.g. touch of the aryepiglottic folds and the glottis, showed marked hyposensitivity, i.e. no laryngeal adductor reflex was elicited. On gross observation, there were neither laryngeal elevation, nor closure of the laryngeal entrance by the epiglottis or adduction of the vocal folds during swallowing.

Radiographic and repeated fiberendoscopic evaluations of the upper airway tract during swallowing showed massive aspiration (Fig. 6). The patient was discharged from the hospital seven weeks after surgery with a gastrostomy feeding tube but without tracheostomy.

Two-and-a-half months after surgery, the patient spontaneously started to swallow without aspiration, prompting us to remove the gastrostomy tube. The glottic closure was complete on examination.

Discussion

In this report, we have described a unique case of severe dysphagia with aspiration in a patient with CNF. Considering the reasons for dysphagia in our patient, several aspects should be emphasized. Firstly, the prelaryngeal muscle removal during surgery resulted in both the inadequate laryngeal elevation and, consequently, the lack of passive closure of its entrance by the epiglottis. Secondly, there was the sensory deficit of the supraglottic mucosa. It was discovered by touching the supraglottic mucosa bilaterally with the tip of the endoscope, which failed to elicit the reflexive adduction of the vocal folds. Therefore, there was no airway protection during swallowing. Considering both normal movement of the vocal cords during phonation and breathing and evident sensory deficit of the supraglottic mucosa, we concluded that the lack of vocal cords adduction during swallowing with consequent aspiration was attributable to the dysfunction of both internal branches of the superior laryngeal nerves.

The internal branches of the superior laryngeal nerves carry the sensory impulses from the laryngeal mucosa down to the vocal cords. They enter the larynx through the thyrohyoid membrane that is situated posteriorly and in close proximity to the prelaryngeal muscles that were most severely affected by CNF. Therefore, the paralysis of
The dysfunction of other cranial nerves after CNF, i.e. accessory nerves, hypoglossal nerves and facial nerves, have already been reported, but only by few authors. Nevertheless, this supports our thesis that CNF can affect the surrounding cranial nerves and that the involvement of a particular nerve depends on the location and severity of CNF.

**Conclusion**

Our case clearly shows that patients with CNF are prone to toxic neuritis-mediated neurological deficits of the cranial nerves, which are considered to be another complication of CNF in addition to those already known. Unless they are recognized and dealt with immediately, further complications follow. In the case of dysfunction of the superior laryngeal nerves, the principal complication is dysphagia leading to aspiration pneumonia, dehydration and cachexia.

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**References**

