Case Report

Grisel syndrome, acute otitis media, and temporo-mandibular reactive arthritis: A rare association

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A B S T R A C T

We present a case report of a four-year-old boy with torticollis and trismus after acute otitis media. Grisel Syndrome diagnosis in association with temporo-mandibular reactive arthritis was admitted, leading to early conservative treatment. GS should be suspected in a child presenting with torticollis after an upper respiratory tract infection or an ENT surgical procedure. The association with temporo-mandibular reactive findings is somehow rarer but not impossible, due to the close vascular communication between retropharyngeal and pterigoid spaces.

1. Background

By definition, Grisel’s syndrome (GS) is a rare, nontraumatic subluxation of C1-C2 joint. Its name is due to the description by Pierre Grisel of three patients with pharyngitis and torticollis due to C1-C2 joint dislocation [1,2].

Although its pathogenesis remains elusive, it is plausible that those with baseline hyperlaxity are at higher risk. That would explain why GS primarily affects children under the age of 12 years and patients with Down syndrome.

Karkos et al. [3] reported 96 cases with non-traumatic atlantoaxial rotary subluxation. Forty-eight percent occurred following infections and 40% after Ear Nose and Throat (ENT) surgery (adenotonsillectomy in 78%). Within the infectious etiology, the main responsible is upper respiratory tract viral infection (83%), followed by retropharyngeal abscess (11%), otitis media (4%) and mumps (2%).

The correct diagnosis is challenging and requires a high index of clinical suspicion and appropriate radiographic imaging. Computerized tomography and/or magnetic resonance imaging of the head and neck are considered the gold standard for GS diagnosis.

The length of time until reduction has been directly related to the failure of medical treatment and to an increased risk of recurrence or permanent neck deformity [4]. Up to 15% of the untreated GS patients develop severe neurological complications: nerve damage, paralysis and even death. Early recognition is of utmost importance to avoid complications.

2. Case presentation

A previously healthy four-year-old boy was admitted to our hospital having neck pain and stiffness, with a rotational misalignment of the mental region to the right side of the neck for six days. There was no history of trauma, but the patient had rhinopharyngitis and acute otitis media diagnosed in the day prior to the beginning of the cervical complaints. He received antibiotic treatment and had remained apyretic.

The physical examination showed: torticollis, associated with spasm and tenderness over right sternocleidomastoid muscle and limited cervical range of motion (Fig. 1); trismus; right temporo-mandibular joint (TMJ) tumorfaction; bilateral tympanic hyperemia; there were no asymmetry or inflammation within the tonsillar area. No neurological, ophthalmological or other ostearticular signs were found.

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3. Investigation

The C-reactive protein was 2.6 mg/dL and the white blood cell count was not altered. Ultrasound examination of the neck revealed multiple enlarged lymph nodes, the biggest one measuring 22 mm × 11 mm in size and was located in the anterior jugular chain.

Contrast-enhanced CT-scan of the cervical region, showed multiple lymphadenopathies in the anterior-lateral cervical region, with no abscess detection (Fig. 2). There was a significant asymmetry of the tissues surrounding the right TMJ due to joint effusion with peripheral halo of contrast intake, suggesting of inflammatory/infectious process (Fig. 3).

4. Treatment and follow-up

First diagnostic impression lead us to assume that this was the case of a secondary torticollis due to cervical lymphadenopathies' inflammatory process. However, initial treatment with diazepam (muscle relaxant), ibuprofen (non-steroid anti-inflammatory) and intravenous (IV) antibiotic (amoxicillin-clavulanate) lead to no further improvement.

This lead to the review of previous CT-scan images. After obtaining adequate multiplanar reconstructions for the cervical spine, additional signs of rotational dislocation of the atlantoaxial joint (Figs. 4 and 5) were identified.

After confirmation of atlantoaxial rotatory subluxation (AARS) – Type 2 dislocation according to Fielding and Hawkins classification [5], neurosurgery consulting was obtained.

In the presence of an AARS with no prior cervical trauma and only acute otitis media as the trigger factor, lead us to establish Grisel Syndrome diagnosis (Deichmueller and Welkoborsky diagnostic algorithm [6] – Table 1).

Being a type 2 AARS with less than 2 weeks of evolution, conservative management was chosen. The patient was submitted to manual subluxation closed reduction under conscious sedation by the neurosurgery team and cervical mobilization restraint was adopted using a Philadelphia collar.

The patient was kept on treatment with ibuprofen and IV amoxicillin-clavulanate for 11 days.

Follow-up contrast enhanced CT-scan performed after 2 weeks, and before hospital release, showed normal alignment of C1-C2, with mild edema of the atlantodental ligament (Figs. 6 and 7). No
Joint effusion was detected in the right TMJ. The patient was then discharged, and neurosurgical follow-up was provided. He kept the cervical collar for 6 weeks, after which there was no recurrence of the AARS.

5. Discussion

As proposed by Welinder et al [7], GS starts as a typical torticollis from the spasm of irritated neck muscles, however, in the prevalence of preexisting lax C1-C2 ligaments, this spasm leads to subluxation of the atlas and axis. This theory can in fact explain the prevalence of GS in children and Down syndrome patients. Additionally, children have wider atlantodental intervals than the general population.

Battiata et al. [8] advanced the two-hit hypothesis to the GS development: the first hit would be the existing of this baseline laxity and a wider anatomical atlantodental space. Followed by the second hit which would be the induction of spasm caused by the inflammatory mediators carried to the cervical muscles by pharyngovertebral venous plexus. The final phase would be the C1-C2 luxation establishment, with secondary inflammatory changes in the transversal ligament and eventually the reabsortive decalcification of C1 and C2 seen in cases with greater disease length.

In this case, the simultaneous reactive joint involvement, with a rare development of TMJ effusion, shows that the hematogenous spread of inflammatory mediators through pharyngovertebral venous plexus may in fact reach the pterygoid space, linking the
weeks, C1-C2 region and TMJ. That would explain trismus and facial asymmetry in the absence of peritonsillar findings in some late detected cases described in literature.

After the AARS diagnosis is established, conservative treatment can be ideally applied if the disease length is shorter than 2 weeks [9]. Conservative or medical treatment consists of C1-C2 subluxation reduction, treatment of the inflammatory and/or infectious process (antibiotics, muscle relaxants and anti-inflammatory therapy) and cervical mobilization restraint using a Sternal-Occipital-Mandibular immobilizer (SOMI) as the first choice of orthosis or simply the Philadelphia collar as a substitute [10].

The majority of GS patients, as our child, do well if diagnosed early [3]. Intractable cases, with a disease length greater than 2 weeks, can be managed with manual repositioning under general anesthesia followed by Minerva orthesis or halo fixation.

Learning points

- Although rare, GS should be suspected in a child presenting with torticollis after an upper respiratory tract infection or an ENT surgical procedure.
- Prompt diagnosis and treatment usually avoid serious complications.
- Previous ligament hyperlaxity may predispose for the development of GS. Muscular spasm induction through the haematogenous spread of inflammatory mediators must be an important trigger for the subluxation.

References