C1–C2 Instability Associated with Periodontoid Inflammatory Tissue Leading to Subarachnoid Hemorrhage: A Case Report and Review of the Literature

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Abstract

The authors present a case of atlantoaxial instability associated with C1–C2 inflammatory tissue leading to subarachnoid hemorrhage. A 65-year-old male patient arrived in June 2011 to the emergency unit for cervical pain and fever. Imaging studies documented periodontoid pseudotumor at C1–C2 level. Infective disease was suspected; the patient was therefore hospitalized and treated with antibiotics. Subsequent computed tomographic (CT) scans revealed C1–C2 instability. In August, the patient showed acute neurological deterioration and coma. Urgent brain CT revealed a hemorrhagic lesion which caused compression on the medulla oblongata, subarachnoid hemorrhage, and ventricular dilatation. An external ventricular drainage was positioned. Angio-CT and angiography did not show any vascular abnormalities. Cervical magnetic resonance imaging documented a solid tissue lesion between the atlas arch and axis. The lesion was associated with an epidural and subdural hematoma, exerting compression on brainstem. The patient underwent posterior decompression and C1–C2 fusion according to Harms technique in October, with significant clinical improvement. The authors present a case of atlantoaxial instability associated with a periodontoid pseudotumor at C1–C2 level determining dural sac compression. The patient showed an acute neurological deterioration caused by bleeding of the solid component of the cervical lesion. Hemorrhage of the solid component of periodontoid masses linked to atlantoaxial instability has not yet been reported in literature. To the best of our knowledge, this is the first case of C1–C2 instability with periodontoid pseudotumor leading to subarachnoid hemorrhage.

Keywords
► periodontoid pseudotumor
► atlantoaxial instability
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Introduction

The authors present a singular case of atlantoaxial instability associated with C1–C2 inflammatory bulk tissue leading to intradural hemorrhage involving the foramen magnum and the subarachnoid space. To the best of our knowledge, this is the first case described in literature of atlantoaxial instability associated with subarachnoid bleeding.

Case Report

A 65-year-old male patient presented to the emergency department with cervical pain and fever. Standard sagittal and coronal cervical spine X-rays did not show atlantoaxial instability but underlined diffuse arthesis and inversion of the normal lordosis, thus requiring further radiological examination. Because the patient had a cardiac pacemaker, he underwent cervical spine computed tomography (CT) scan instead of magnetic resonance imaging (MRI). Imaging documented a swollen material located anteriorly to the dens at the C1–C2 transverse ligament level, which exerted initial dural sac compression. An infective etiology was suspected, and the patient was therefore hospitalized and treated with antibiotic therapy. Subsequent CT scans revealed C1–C2 instability (atlas–axis distance: 9.8 mm in July vs. 1.5 mm in May, ►Fig. 1). Indication to C1–C2 posterior arthrodesis was given.

After 1 month, the patient complained of worsening cervical pain and confusion, and was again hospitalized. A few days later, gaze conjugate deviation and severe tetraparesis appeared. Urgent brain CT scan revealed a hemorrhagic hyperdensity involving the lesion, the foramen magnum, and the subarachnoid space (►Fig. 2). Compression on the medulla by the subdural hematoma and ventricular dilatation were also documented. Angio-CT and angiography excluded the presence of any vascular abnormalities. Despite the pacemaker and the risks associated, a cervical magnetic resonance imaging (MRI) was performed. The examination showed a solid hypointense tissue lesion located between atlas arch and axis, next to a fluid component related to a right C1–C2 facet joint synovial cyst. The whole lesion was associated with recent hemorrhage, which exerted compression on brainstem. The medulla and upper spinal cord showed a hyperintense signal in T2-weighted images (►Fig. 3). An external ventricular drainage (EVD) was positioned to control intracranial hypertension. Progressive improvement of the patient’s clinical condition with recovery of consciousness was subsequently observed. The EVD was removed after 5 days and, 1 month after the acute clinical worsening, the patient underwent posterior decompression and C1–C2 fusion according to the Harms technique (►Fig. 4a). Biopsy of the lesion was not feasible due to the unfavorable surgical accessibility.

The patient was then transferred to a rehabilitation unit, showing progressive improvement of the neurological status. At discharge, the patient was able to walk unassisted. A cervical CT scan performed 1 year after surgery showed signs of C1–C2 fusion, an atlantoaxial distance of 3 mm and reduction of periodontoid tissue (►Fig. 4b). Brain CT documented a slight enlargement of the ventricular system.

Discussion

Atlantoaxial instability syndrome includes a wide group of pathological conditions. In fact instability can result from congenital disease, as it occurs in genomic pathologies (Morquio, Down, and Conradi syndromes) and in idiopathic dysplasias (e.g., os odontoideum, fetal warfarin syndrome) or it can be an acquired condition. In the latter group, well-known causes are infections, inflammatory

![Fig. 1](a) Sagittal CT scan (May), showing increased C1–C2 distance (1.5 mm). (b) New cervical CT (July) documented further increase of atlantoaxis distance (9.8 mm). CT, computed tomography.
diseases (in particular in rheumatoid arthritis), tumors, and traumas.¹

The development of an inflammatory mass or pannus at the craniocervical junction due to an atlantoaxial instability is a relatively common condition.² Some authors have called these masses periodontoid pseudotumor³ or retroodontoid pseudotumor according to their position.⁴

However, there are some questions about the nature of this tissue. For some authors, this is an inflammatory granulation tissue, whereas for others, it is a reactive fibrous tissue due to mechanical overload. This last hypothesis would be supported by the fact that pannus is found in patients with no systemic inflammatory process and with only chronic atlantoaxial instability.²

Our clinical case represents an evident C1–C2 instability, related to a significant increase in atlantoaxial distance, as demonstrated in follow-up imaging studies (►Fig. 1). This instability was associated with a focal mass lesion, which exerted initial compression on the dural sac, whose histological definition was not determined preoperatively because of the difficult surgical approachability.

The feature that makes this case unusual and surprising is the sudden subdural and subarachnoid hemorrhage (SAH) associated with the lesion, which caused compression of the brainstem and ventricular dilatation (►Fig. 2).

In literature, the association between C1–C2 instability, consequent periodontoid mass, and SAH has not been reported yet. The complete cerebral and cervical vessel

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**Fig. 2** (a) Axial CT demonstrates a hemorrhagic hyperdensity involving the foramen magnum and the subarachnoid space, exerting compression on the medulla oblongata. (b) Axial CT scan reveals evident ventricular dilatation; hematic hyperdensity can be seen in the occipital horns. CT, computed tomography.

**Fig. 3** (a, b) Sagittal T2-weighted MRI documents hemorrhage at C1–C2 level associated to the periodontoid lesion. The hemorrhage exerts compression on brainstem. The medulla oblongata and upper spinal tract showed a hyperintense signal. MRI, magnetic resonance imaging.
angiography excluded the presence of any vascular abnormality and this led to the conclusion that the bleeding originated from the cervical lesion.

Radiological studies, in particular the MRI, documented a probable reactive/flogistic disease at C1–C2 level. The peridontoid tissue was associated with a fluid component at C1–C2 right joints, probably due to a synovial cyst. The presence of a synovial cyst is sign of joint instability, as Jost et al suggested. Even though the pathogenesis of synovial cysts is unclear, Jost et al proposed the release of inflammatory factors by stressed joints as a possible pathological mechanism, resulting in the formation of the cyst or extrusion of synovium through the joint capsule caused by trauma or instability. In literature, cases of hemorrhagic synovial cysts have been reported, but in the case presented, the bleeding involved the solid component of the lesion, as MRI clearly demonstrated (Fig. 5). The synovial cyst can therefore be considered an epiphenomenon of the joint instability and not the origin of the hemorrhage.

Hemorrhages of solid component of peridontoid masses linked to atlantoaxial instability have not, to our knowledge, been reported in the literature so far.

The hematoma determined compression of the brainstem and diffuse SAH, with acute and massive dilatation of the ventricles, which required urgent EVD positioning. The hydrocephalus resolved during the following hospitalization, and so the shunt was removed.

In our case, there was no abnormal fluorodeoxyglucose uptake noted on positron emission tomography scan and serum tumor markers were also negative. The radiological features of the lesion and the progressive cranio cervical instability suggested an inflammatory disease, such as rheumatoid arthritis; however, this hypothesis was not confirmed by the analysis of autoantibodies, markers of common rheumatological diseases.

The histological diagnosis was not achieved since the lesion could not be safely reached through the posterior approach. Cases of atlantoaxial instability with peridontoid mass unrelated to rheumatoid arthritis have been reported in literature, and histological examinations in these cases showed inflammatory or scar tissue, suggesting that atlantoaxial instability induced local inflammation and then the developing of a mass.

The mechanism, which correlated atlantoaxial instability, hemorrhage, and intradural blood diffusion, is not clear. A hypothesis could be that the atlantoaxial instability determined inflammation and frailty of the dura mater, eventually causing it to tear when the lesion bled. This mechanism would also explain blood also diffused in the subarachnoid space.

Surgical treatment of peridontoid pseudotumor can be transoral odontoid resection followed by anterior plating or, more commonly, posterior decompression and fusion. Anterior approach is recommended when atlantoaxial kyphosis is not reducible and could be the best way to control ventral spinal cord compression. The posterior approach aims to pannus regression, due to atlantoaxial arthrosis. Several reports described not only regression of peridontoid lesions...
Pseudotumor after vertebral stabilization but also clinical improvement of myelopathy. Pseudotumor resection allows histopathological examination but has a high risk of severe neurological deficits. In the case presented, a posterior approach with decompression and C1–C2 fusion with instrumentation was performed and led to progressive neurological improvement. After 1 year, the patient was able to walk again. CT cervical scan documented reduction of the periodontoid mass and of the atlas-axis distance.

Conclusions
The growth of an inflammatory mass or pannus in the craniovertebral junction due to atlantoaxial instability is a well-known condition called periodontoid pseudotumor. Even if such a dramatic clinical course is exceptional a significant increase of C1–C2 distance in short-term radiological follow-up should be considered alarming and should induce an expeditious surgical treatment. In literature, there have been no cases of pannus acute bleeding leading to brainstem compression and, in particular, this could be the first case of C1–C2 instability associated with SAH.

Conflict of Interest
The authors do not declare any conflict of interest about the development and management of this case and this article.

Permission Request
The patient, even if no data can be directly linked to him/her, declared his/her permission to publish this case report.

Statement of Authors’ Approval
All the authors approve the content of the article submitted.

Note
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References