MUL TIPLE INTRACRANIAL SCHWANNOMAS: CASE REPORT

Mario Mihalj1, Krešimir Dolić2, Pavao Jurinović1, Nikolina Ivica Miše1, Marina Titlić1 and Irena Pintarić1

1Clinical Department of Neurology, 2Clinical Department of Diagnostic and Interventional Radiology, Split University Hospital Center, Split, Croatia

SUMMARY – Schwannomas are benign encapsulated tumors arising from the sheaths of peripheral nerves. They present as slowly enlarging solitary lumps, which may cause neurological defects. Multiple schwannomas in non-neurofibromatosis type 2 patients are extremely rare. We report a case of a 60-year-old female patient, without any family history of neurofibromatosis or schwannomatosis, presented with trigeminal neuralgia and progressive facial nerve palsy. Magnetic resonance imaging revealed the presence of acoustic schwannoma involving facial nerve and trigeminal schwannoma of the cisternal part of the nerve involving gasserian ganglion (Meckel’s cave). After gamma knife radiosurgery, trigeminal neuralgia was relieved completely with improvement of facial nerve palsy.

Key words: Neurilemmoma – diagnosis; Neurilemmoma – surgery; Neurroma, acoustic – diagnosis; Neurroma, acoustic – surgery; Trigeminal nerve diseases – surgery; Case reports

Introduction

Schwannomas (neurilemmomas) are benign, slow-growing, encapsulated tumors originating from Schwann cells that form myelin sheath around peripheral, cranial, and autonomic nervous systems. They develop on the outside of the nerve, but may push it aside or against adjacent structures causing damage1. Schwannomas account for 7% of all intracranial tumors2, and among them vestibular schwannomas are most common. However, those originating from the trigeminal ganglion or nerve root, or other cranial nerves are considered rare1.

We report a case of schwannomas of the vestibular and trigeminal nerves in a non-neurofibromatosis type 2 (NF2) female patient who presented with trigeminal neuralgia and progressive facial nerve palsy.

Case Report

A 60-year-old female was admitted to the Clinical Department of Neurology, Split University Hospital Center, because of acute right-side facial palsy (House-Brackmann grade IV). She did not complain of any episode of vertigo, dizziness, or tinnitus. The basic laboratory tests (peripheral blood and blood chemistry findings) were within the normal limits.

Her medical history showed that she also had suffered from right-sided infranuclear facial palsy (House-Brackmann grade III) a year before. Otherwise, she was healthy and no other focal neurological signs or symptoms including hearing disturbances were elicited at that time. She was administered 20 mg prednisone per day for 5 days, including gastric and local eye protection therapy. Prednisone was discontinued by gradual reduction of dosage. After physiotherapy, facial function improved gradually to grade II. Six months later, she presented with right trigeminal neuralgia. The pain affected initially the maxillary and then the ophthalmic divisions of the trigeminal nerve, characterized as sharp and lancinating pain of moder-
ate to severe intensity. Systemic and neurological examinations were normal. In particular, trigeminal sensation was intact bilaterally and trigeminal reflexes were normal. Conventional analgesia in high dosage relieved pain. Based on her positive medical history of neuralgia, magnetic resonance imaging (MRI) of the brain was performed to reveal the presence of right-side trigeminal and vestibular nerve schwannomas also involving the ganglion segment of the facial nerve (Fig. 1).

Ophthalmologic and general physical examinations did not reveal any findings suggestive of neurofibromatosis. There was no family history of neurologic disturbances, café-au-lait spots, tumors of the skin or central nervous system.

The patient underwent gamma knife radiosurgery for trigeminal schwannoma and vestibular schwannoma. An obvious decrease in tumor volumes was observed. Trigeminal neuralgia was relieved completely with improvement of facial nerve palsy (House-Brackmann grade II) during 9-month clinical follow up.

Discussion

Vestibular schwannomas are the most common intracranial site of schwannoma occurrence. They comprise 90% of all cranial nerve schwannomas, followed by the second most common but much rarer trigeminal schwannomas.

Cranial nerve schwannomas usually manifest sporadically as a single neoplasm. The presence of multiple schwannomas in the same patient suggests tumorigen-
control, preservation of multiple nerve functions, and good quality of life in patients with both vestibular schwannoma and trigeminal schwannoma\textsuperscript{10,11}.

The presented case is unique because it shows a rare phenotype of multiple schwannomas (unilateral vestibular schwannoma and trigeminal schwannoma) in a non-NF2 patient with no other stigmata of neurofibromatosis.

It also shows that progressive facial palsy without any disturbances of hearing is probably not Bell's palsy. In patients with such symptoms, especially if they have recurrent facial palsy and other neurological signs, it is important to examine other rare causes.

References


Sažetak

MULTIPLI INTRAKRANIJSKI ŠVANOMI: PRIKAZ SLUČAJA

M. Mihalj, K. Dolić, P. Jurinović, N. Ivica Miše, M. Titlić i I. Pintarić

Švanomi su dobroćudni inkapsulirani tumori koji potječu iz ovojnica perifernih živaca. Najčešće se javljaju kao pojedinačni sporo rastući tumori koji mogu uzrokovati neurološke defekte. Mnogostruki švanomi su iznimno rijetki u bolesnika koji nemaju neurofibromatouzu tip 2. Prikazujemo slučaj 60-godišnje bolesnice s negativnom obiteljskom anamnezom za neurofibromatozu ili švanomatozu, koja se prezentirala s neuralgijom trigeminusa i progresivnom paralizom facijalnog živca. Magnetska rezonanca prikazala je prisutnost akustičnog švanoma sa zahvaćanjem i facijalnog živca te švanoma cisternalnog dijela trigeminalnog živca sa širenjem u područje gangliona istog (Meckelova šupljina). Nakon “gamma knife” radiokirurgije simptomi trigeminalne neuralgije su se u potpunosti povukli uz značajno kliničko poboljšanje paralize facijalnog živca.

Ključne riječi: Neurilemom – dijagnostika; Neurilemom – kirurgija; Neumom, akustični – dijagnostika; Neumom, akustični – kirurgija; Trigeminalni živac, bolesti – kirurgija; Prikazi slučaja