Imaging of cranial nerves: a pictorial overview

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Abstract

The human body has 12 pairs of cranial nerves that control motor and sensory functions of the head and neck. The anatomy of cranial nerves is complex and its knowledge is crucial to detect pathological alterations in case of nervous disorders. Therefore, it is necessary to know the most frequent pathologies that may involve cranial nerves and recognize their typical characteristics of imaging. Cranial nerve dysfunctions may be the result of pathological processes of the cranial nerve itself or be related to tumors, inflammation, infectious processes, or traumatic injuries of adjacent structures. Magnetic resonance imaging (MRI) is considered the gold standard in the study of the cranial nerves. Computed tomography (CT) allows, usually, an indirect view of the nerve and is useful to demonstrate the intraosseous segments of cranial nerves, the foramina through which they exit skull base and their pathologic changes. The article is a complete pictorial overview of the imaging of cranial nerves, with anatomic and pathologic descriptions and great attention to illustrative depiction. We believe that it could be a useful guide for radiologists and neuroradiologists to review the anatomy and the most important pathologies that involve cranial nerves and their differential diagnosis.

Keywords: Cranial nerves, Magnetic resonance imaging, Computed tomography, Anatomy, Pathology

Key points

- Anatomy of cranial nerves is complex and its knowledge is crucial to detect pathological alterations in case of nervous disorders
- Magnetic resonance imaging (MRI) is the gold standard technique in the study of the cranial nerves
- Steady-state free procession (SSFP) images are the best sequences for the visualization of the cisternal segments showing dark cranial nerves against a background of bright cerebrospinal fluid (CSF)
- Computed tomography (CT) can be useful to evaluate intraosseous segments of cranial nerves, skull base foramina, and bony traumatic lesions
- Cranial nerve dysfunctions may be the result of pathological processes of the cranial nerve itself or may be associated with tumor, inflammation, infectious processes, or traumatic injuries of adjacent structures

Introduction

The human body has 12 pairs of cranial nerves that control motor and sensory functions of the head and neck. This article provides a pictorial overview of the imaging of cranial nerves, with a special focus on their anatomy and pathology. Radiologists and, specially, neuroradiologists should be familiar with the anatomy of each cranial nerve and should know clinical and imaging findings of their dysfunctions.

Imaging

MRI is considered the gold standard in the study of cranial nerves. Table 1 summarizes the most important sequences and features in their study. Several scientific articles have underlined the importance of SSFP sequences for the visualization of the cisternal spaces of cranial nerves thanks to their sub-millimetric spatial and high contrast resolution [1–4]. These sequences typically show dark cranial nerves against a background of bright CSF [2, 5] (Fig. 1). The enhancement of the nerve, after gadolinium administration, is associated to disruption of the blood-nerve barrier and may be secondary to neoplasm, inflammation, demyelination, ischemia, trauma, radiation treatment, and axonal degeneration. Three-dimensional...
T1-weighted gradient echo sequences can provide a better visualization of the nerve enhancement, surrounded by CSF. MRI could also be useful to assess denervation changes of “end-organ muscles” (Table 2) [5]. CT is inferior to MRI in the visualization of the cranial nerves themselves, due to its low contrast resolution. Therefore, it can be useful to evaluate the intrasosseous segments of cranial nerves and the possible associated bony changes. Particularly, CT is optimal to study the bony foramina of the skull base (Table 3, Fig. 2) and bony traumatic lesions [6].

Anatomy

Nuclei of cranial nerves, except I and II, are located in the brainstem (Fig. 3); from their origin, nerves are divided in cisternal, intracranial, and extracranial segments.

**Table 1** Utility of MRI sequences for the study of cranial nerves

| MRI sequences                              | Features                                      |
|--------------------------------------------|-----------------------------------------------|
| T1-weighted sequences                      | - Anatomical definition                       |
|                                            | - Invasion of fat planes                      |
|                                            | - Denervation changes                         |
| T2-weighted sequences                      | - Lesion characterization                     |
|                                            | - Patency of CSF spaces                       |
|                                            | - Denervation changes                         |
| Fast spin echo; steady-state free procession| - Cisternal course of the nerve               |
|                                            | - Neurovascular conflict                      |
| DWI–FLAIR sequences                        | - Ischemic lesion                             |
| Post-gadolinium T1-weighted sequences      | - Enhancement of the nerve                    |
| (eventually with fat-suppression)          | - Perineural spread                           |
|                                            | - Meningeal infiltration                      |

**Fig. 1** MRI of the cisternal tracts of normal cranial nerves. Steady-state free procession (SSFP), axial planes. **a** Olfactory (circles); **b** optic (arrows); **c** oculomotor (arrows); **d** trochlear (arrows); **e** trigeminal (arrows); **f** abducens (circles); **g** facial (thin and dotted arrows) and vestibulocochlear (thick arrows); **h** glossopharyngeal and vagus nerve (circles); **i** spinal accessory (circles); **j** hypoglossal (arrows)
Microscopically, cranial nerves are surrounded by a series of a connective tissue sheaths: endoneurium, perineurium, and epineurium. Axons of each nerve, except I and II, are myelinated by Schwann cells.

Functionally, they may have motor efferents and/or sensory afferents (Table 4).

I cranial nerve - Olfactory nerve

The olfactory nerve is a sensitive nerve that conveys olfactory stimuli from the nasal cavity to the brain [7]. It consists of white matter tracts not surrounded by Schwann cells. Olfactory receptors are located in the mucosa of nasal cavity and olfactory filaie represent their axons. They enter the anterior cranial fossa through the cribriform plate and terminate in the olfactory bulb [6]. Successively, the nerve courses between the gyrus rectus and the medial orbital gyrus. The secondary axons terminate in the inferomedial temporal lobe, uncus, and entorhinal cortex [3] (Fig. 4).

II cranial nerve - Optic nerve

Optic nerve represents an extension of central nervous system and is myelinated by oligodendrocytes [8].

| Nerve | Muscles to evaluate for denervation |
|-------|-----------------------------------|
| III   | Superior, inferior, medial rectus, and inferior oblique muscles |
| IV    | Superior oblique muscle |
| V     | Muscles of mastication: masseter, temporalis, and pterygoid (V3) |
| VI    | Lateral rectus muscle |
| X     | Vocal cords |
| XI    | Sternocleidomastoid, trapezius muscles |
| XII   | Tongue muscles |

III cranial nerve - Oculomotor nerve

The oculomotor nerve has a somatic motor function of most ocular extrinsic muscles (inferior, superior, middle rectus, inferior oblique, and levator palpebrae superior muscles) and a parasympathetic function (ciliaris and sphincter pupillae muscles) [6]. Somatic motor fibers of oculomotor originate from the nuclear complex located in the midbrain at the level of the superior colliculi, ventral to the cerebral aqueduct and the periaqueductal gray matter. The parasympathetic fibers arise in the Edinger-Westphal nucleus, situated dorsally to the oculomotor nuclear complex. The nerve emerges from the interpeduncular cistern running through the perimesencephalic cistern, superiorly to the posterior cerebral artery (PCA) and inferiorly to the superior cerebellar artery (SCA) [6]. Successively, the nerve enters in the cavernous sinus where is the most superior nerve, to reach the orbit through the superior orbital fissure [6] (Fig. 5).

IV cranial nerve - Trochlear nerve

The trochlear nerve innervates only the superior oblique muscle. It is the unique nerve with a root zone arising from the posterior brainstem where its nucleus lies [3]. After exiting the pons, the nerve curves over the superior cerebellar peduncle and then runs between the SCA and the PCA. Successively, it runs along the free margin of the tentorium and then enters the cavernous sinus, inferior to the III nerve and superior to the ophthalmic division of the V nerve [6]. Passing the cavernous sinus, the trochlear nerve enters the orbit through the superior orbital fissure [3] (Fig. 5).

V cranial nerve - Trigeminal nerve

The trigeminal nerve is the largest cranial nerve, composed of sensory and motor roots [3]. Four trigeminal nuclei (spinal nucleus of V, principal sensory nucleus of V, motor nucleus of V, and mesencephalic nucleus) extend from the midbrain down to the upper cervical medulla [6]. The nerve emerges straight forward from the lateral pons and pass in the Meckel’s cave, where is located the trigeminal ganglion (Gasserian ganglion), from which the nerve splits into three subdivisions. The
ophthalmic (V1) and the maxillary (V2) divisions pass in the lateral wall of cavernous sinus, inferior to the VI cranial nerve. The ophthalmic division exits the skull via the superior orbital fissure (with the III, IV, VI cranial nerves and the superior ophthalmic vein). The maxillary division exits the skull through the foramen rotundum and crosses the pterygopalatine fossa. Successively, it enters the orbits via the inferior orbital fissure, passes within the infraorbital groove, and reaches the face through the infraorbital foramen. The mandibular division (V3) exits the skull via the foramen ovale and passes in the infratemporal fossa, where it divides into some branches (meningeal branch, medial and lateral pterygoid nerves, masseteric nerve, deep temporal nerve, buccal nerve, auriculotemporal nerve, lingual nerve, and inferior alveolar nerve) [3, 6] (Fig. 6).

**VI cranial nerve - Abducens nerve**

The abducens nerve is a motor nerve that emerges from the abducens nucleus, located just beneath the floor of the IV ventricle in the dorsal pons [3]. The nerve courses anteriorly to exit the brainstem at the pontomedullary junction and crosses preoptic cistern in a dorsal-to-ventral direction. Successively, it descends near to the posterior aspect of the clivus, passing through the Dorello’s canal, and enters in the cavernous sinus [9]. It is the unique cranial nerve that courses through the central venous portion of the sinus. Finally, it enters the orbit via the superior orbital fissure to innervate the lateral rectus muscle [3, 6] (Fig. 5).

**VII cranial nerve - Facial nerve**

The facial nerve is a mixed nerve and consists of two portions: the proper VII nerve (with motor function) and the intermediate nerve (with sensory and parasympathetic motor fibers) [6]. The motor nucleus is located in the lower portion of the pons and motor fibers exit from the lateral portion of the pontomedullary sulcus. Successively, the nerve traverses the cerebellopontine cistern and enters the temporal bone through the internal acoustic meatus. In the petrous bone, it passes through the facial canal, giving three segments (labyrinthine, tympanic, and mastoid). The nerve exits the skull base at the stylomastoid foramen and reaches the
parotid gland [6]. From the mastoid portion, the nerve of the stapedius muscle and corda tympani emerge [6]. The afferent sensory fibers that originate from receptors of the first two-thirds of the tongue arrive at the geniculate ganglion, via the chorda tympani. They run with the intermediate nerve in the internal acoustic meatus and the cerebellopontine cistern to terminate in the nucleus of tractus solitarius. The parasympathetic fibers of the intermediate nerve arise from the superior salivatory nucleus. At the geniculate ganglion, the great petrosal nerve provides parasympathetic fibers for the lacrimal gland and the mucosa of mouth, nose, and pharynx [6] (Fig. 7).

Table 4 Summary of efferents and afferents fibers of cranial nerves

| Motor efferents                  | General somatic efferents | General visceral efferents | Special visceral efferents |
|----------------------------------|---------------------------|---------------------------|---------------------------|
|                                  | III, IV, VI, XII          | III, VII, IX, X           | V, VII, IX, X, XI         |
| Sensory afferents                |                           |                           |                           |
| General somatic afferents        | V, VII, X                 |                           |                           |
| General visceral afferents       | IX, X                     |                           |                           |
| Special afferents                | I, II, VII, IX, X         |                           |                           |

VIII cranial nerve - Vestibulocochlear nerve

The vestibulocochlear nerve is formed by two nerves: the cochlear nerve that transmits sound and originates in the cochlear membrane, and the vestibular nerve (divided in superior and inferior) that controls balance and originates from the ciliary sensory cells in the membranous labyrinth [10]. Both nerves traverse the internal acoustic meatus, together with the facial nerve, and pass in the cerebellopontine cistern. In the internal acoustic meatus, the nerve
Fig. 4 Schematic drawing of the olfactory nerve

Fig. 5 Schematic drawing of the optic and oculomotor nerves
is so placed: the facial nerve is anterosuperior, the vestibular superior nerve is posterosuperior, the cochlear nerve is anteroinferior, and the inferior vestibular nerve is posteroinferior (for this, remember the acronym “Seven Up Coke Down”). Successively, vestibulocochlear nerve enters the brainstem and reaches the dorsal and ventral cochlear nucleus and the four vestibular nuclei that are located in the dorsolateral pons [6] (Fig. 8).

IX cranial nerve - Glossopharyngeal nerve
The glossopharyngeal nerve is a mixed sensory, motor, and secretory nerve. It emerges from the lateral medulla into the lateral cerebellomedullary cistern, where it is closely associated with the flocculus of cerebellum [3]. Successively, the nerve exits the skull through the jugular foramen (pars nervosa), and it enters in the carotid space; successively, it turns lateral to the carotid artery and the stylopharyngeal muscle and continues lateral to the palatine tonsil area. Its lingual branch ends in the posterior sublingual space, to innervate the posterior third of the tongue (taste and sensory). Other nervous branches of the extracranial segment of IX nerve are pharyngeal (sensation from posterior oropharynx and soft palate), sinus nerve (parasympathetic supply to carotid body and sinus), stylopharyngeus (motor innervation to the stylopharyngeus muscle), and tympanic branch or Jacobson nerve (sensory information from the middle ear) [6] (Fig. 9).

X cranial nerve - Vagus nerve
The vagus nerve emerges from the lateral medulla, from the base of the nucleus ambiguus and the dorsal nucleus of the vagus [6]. The nerve enters the lateral cerebellomedullary cistern and exits the skull through the jugular foramen (pars vascularis) between the glossopharyngeal nerve and the accessory nerve. Into and below the jugular foramen, the nerve forms the superior ganglion, and just below the jugular foramen, the nerve forms the inferior vagal ganglion. Successively, it descends vertically in the retrostyloid space together with the carotid artery, to join the upper mediastinum where it gives off the recurrent laryngeal nerve. Successively, the vagus nerve enters the abdominal cavity through the esophageal hiatus [6] (Fig. 9).
XI cranial nerve - Accessory nerve

The accessory nerve has both cranial and spinal roots [3]. Cranial roots emerge into the lateral cerebellomedullary cistern, while spinal roots emerge from upper cervical segment of the spinal cord (from C0 to C5) and pass superiorly through the foramen magnum into the cisterna magna. Successively, conjoined cranial and spinal roots enter the jugular foramen, posterior to the glossopharyngeal and vagus nerve. Within the jugular foramen, cranial roots mix with vagus nerve at the level of the superior vagal ganglion, while spinal roots descend laterally to the

Fig. 7 Schematic drawing of the facial nerve

Fig. 8 Schematic drawing of the vestibulocochlear nerve
Fig. 9 Schematic drawing of the glossopharyngeal and vagus nerves.

Fig. 10 Schematic drawing of the accessory and hypoglossal nerves.
internal carotid artery and internal jugular vein, and achieve the sternocleidomastoid and trapezius muscle [3, 6] (Fig. 10).

XII cranial nerve - Hypoglossal nerve
The hypoglossal nerve emerges from the medulla oblongata as a series of rootlets extending from the ventrolateral sulcus of the medulla into the lateral cerebellomedullary cistern, near to the vertebral artery and the posterior inferior cerebellar artery (PICA). The nerve exits the skull through the hypoglossal canal and then runs medial to the glossopharyngeal, vagus, and accessory nerves, deeply to the digastric muscle, to innervate a large part of extrinsic and intrinsic tongue muscles [3] (Fig. 10).

Pathology
Pathology of cranial nerves involves a wide spectrum of etiologies that includes primary or secondary neoplastic involvement, inflammation, infections, traumatic injuries, ischemic lesions, compression by adjacent structures, and congenital pathologies (Table 5).
Here, we review and illustrate the most important and frequent pathologies that may involve cranial nerves.
All images and cases were collected in our Institute.

Table 5 Summary of pathologies that may involve cranial nerves

| Neoplasm |
|-----------------|
| Primary neural tumor |
| Compressive mass (sellar, paracavernous, bone tumor) |
| Leptomeningeal or perineural spread of tumor |

| Inflammation |
|-----------------|
| Optic neuritis |
| Multiple sclerosis, neuromyelitis optica spectrum disorders |
| Pseudotumor (ToIosa-Hunt syndrome) |
| Sarcoidosis |
| Gradengio syndrome |
| Vestibular neuritis, labyrinthitis |

| Infection |
|-----------------|
| Abscess, encephalitis, meningitis, thrombophlebitis, viral neuritis (Ramsey-Hunt syndrome) |
| Skull base osteomyelitis |

| Trauma |
|-----------------|
| Vascular |
| Neurovascular compression, aneurysm |

| Ischemia |
|-----------------|
| Hemorrhage |
| Superficial siderosis |

| Congenital |
|-----------------|
| Neurofibromatosis (I and II), Kallman syndrome, aplasia/hypoplasia of the optic nerve, Duane syndrome, Moebius syndrome |

Neoplasms
Schwannoma and neurofibroma
Schwannoma is a benign nerve sheath tumor that originates from Schwann cells. Approximately 90% of intracranial schwannomas arise from the VIII cranial nerve with trigeminal involvement as the second most common (0.8–5%) [11, 12]. Intracranial schwannomas can be associated with neurofibromatosis 2 (NF2), a genetic condition characterized by the presence of multiple schwannomas, meningiomas, and ependymomas. Patient with NF2 may have bilateral vestibular schwannomas or non-vestibular schwannomas [13]. Imaging shows the slow growth of the tumor, with bone remodeling of neural foramen and deformation of the adjacent brain tissue. Features of end-organ compromise can be present. On CT, schwannomas have low-intermediate attenuation with variable enhancement. MR imaging findings include iso-hypointensity on T1-weighted sequences, high signal on T2-weighted sequences, and marked enhancement on post-contrast images, eventually with unenhanced cystic spaces [13] (Figs. 11, 12, 13, and 14).

Neurofibromas rarely involve cranial nerves, and are more commonly found extracranially. They are typically associated with neurofibromatosis 1 (NF1) and can show malignant transformation. NF1 is a genetic condition characterized by the presence of café au lait spots, neurofibromas, optic nerve glioma, osseous lesions, iris hamartomas, and axillary or inguinal freckling (Figs. 15 and 16).

Esthesioneuroblastoma
Esthesioneuroblastoma (ENB) or olfactory neuroblastoma is a rare malignant tumor of neural crest origin, arising from the olfactory epithelium of the nasal vault. It presents typically in the second and sixth decades of life, without gender predilection [14]. Clinical presentation may include, in a various combination, epistaxis, nasal obstruction, decreased olfactory function, diplopia, and proptosis. ENB may invade paranasal sinuses, orbits, and anterior cranial fossa, and, if metastatic, involve local lymph nodes, with distant metastasis to lungs, liver, and bone. At CT, ENB does not have specific features appearing as a homogeneous soft tissue mass in the nasal cavity. However, CT is useful to evaluate bone involvement. On MRI, ENB usually appears hypointense on T1-weighted sequences and intermediate to hyperintense on T2-weighted sequences. Contrast enhancement is avid and homogeneous, except for areas of necrosis or hemorrhage. MRI may be useful to detect dural involvement [14] (Fig. 17).

Optic nerve glioma
Optic nerve glioma is a relatively rare tumor that typically occurs in children. If associated with NF1, glioma can be multifocal and bilateral. In NF1 patient, optic nerve is the most common site of the tumor, whereas in non-NF1
Fig. 11 Small intracanalicular vestibulocochlear schwannoma. MRI steady-state free procession (SSFP) (a) and T1-weighted post-gadolinium (b) axial images show a small hypointense lesion on steady-state sequence with avid contrast-enhancement visible in the intracanalicular segment of internal auditory canal (arrows).

Fig. 12 Small trigeminal schwannoma. MRI T1-weighted post-gadolinium axial (a) and coronal (b) images show a homogeneous enhancing mass in the right Meckel's cave (dotted circles).

Fig. 13 Large trigeminal schwannoma. CT axial image (soft tissues and bone windows, a, b, respectively) demonstrates a mild iso-attenuated mass (empty arrow) with erosion of the left petrous apex (arrow); MRI T1-weighted post-gadolinium axial image (c) shows an avid enhancing dumbbell-shaped mass (asterisk), that extends in the Meckel's cave, along the cisternal course of the left trigeminal nerve.
patients, chiasma involvement is the most frequent. On MRI, the tumor typically appears isointense on T1-weighted images and iso-hyperintense on T2-weighted images, with variable contrast enhancement; in non-NF1 patients, cystic components may be detected (Fig. 18). If present, perineural arachnoidal gliomatosis shows hyperintense signal on T2-weighted sequences and may have contrast enhancement [8].

Perineural spread of tumor
Two types of perineural growth of tumor are described: perineural invasion (PNI), a microscopical process that affect small nerves, and perineural spread of tumor (PNS) that extends along the sheaths covering larger central nerves [15]. The incidence of perineural spread is about 2.5–5% and may occur with any head and neck malignancy [16, 17]. It is typically associated with carcinoma arising from minor or major salivary glands (e.g., adenoid cystic carcinoma), mucosal or cutaneous squamous cell carcinoma, basal cell carcinoma, melanoma, lymphoma, and sarcoma. In most cases, the peripheral branches of second (V2) and third (V3) divisions of the trigeminal nerve, and the descending branch of the facial nerve serve as conduit for perineural spread. Also, the ophthalmic nerve and the hypoglossal nerve may be interested [17]. Clinical findings include neural dysfunction
symptoms, as pain, dysesthesias, or muscle denervation atrophy [5]. For radiologist, the knowledge of the anatomy of cranial nerves is crucial to evaluate the perineural spread. On MRI, it can be visualized as segmental nerve thickening and enhancement on post-gadolinium sequences. Typically, imaging techniques with fat suppression are used to better increase the conspicuity of the enhancement [17]. Skull base invasion may result as replacement of normal fatty marrow signal [5] (Figs. 19, 20, and 21). Tumoral growth may cause enlargement of foramina or canals through which nerves exit the skull.

**Leptomeningeal spread of tumor**

Primary (medulloblastoma, ependymoma, oligodendroglioma, and glioblastoma) and secondary (leukemia/lymphoma, breast cancer, lung cancer, and melanoma) tumors may spread through subarachnoid spaces with extension along the cranial nerves. VII and VIII cranial seem to be the most affected nerves [18]. Clinically, symptoms of irritation and neural compression are present with multiple cranial neuropathies and mental status changes, secondary to meningeal irritation and hydrocephalus. The nerve may enhance on post-gadolinium sequences and appear enlarged (Fig. 22). Neural enhancement can be very subtle and better demonstrated on post-Gadolinium fluid-attenuated inversion recovery (FLAIR) sequences [5, 19–21].

**Meningioma**

Meningioma is the commonest extra-axial tumor of central nervous system and arises from the meningothelial...
cells of arachnoid layer. Typical locations include parasagittal and lateral aspect of the cerebral convexity, sphenoid wing, middle cranial fossa, and olfactory groove. The tumor may show transforaminal extension, for example into the orbit or following the trigeminal course. After the acoustic schwannoma, it represents the second most common mass in the cerebellopontine angle. On imaging, meningioma appears as a lobular, extra-axial mass with well-circumscribed margins and a broad-based dural attachment. On MRI, it is isointense/hypointense relative to gray matter on the T1-weighted sequences and isointense/hyperintense relative to gray matter on the T2-weighted sequences. A rim of high T2 signal that separates the mass from the adjacent brain is often visible (“cleft signs”). Post-contrast sequences show marked and homogeneous enhancement with rare central nonehancing areas, corresponding to necrosis or calcification phenomena. In up to 72% of cases, a “dural tail” is evident, as

![Fig. 18 Optic nerve gliomas. MRI T1-weighted sagittal (a), axial (b), and coronal (c) images show bilateral intraorbital hypointense masses (dotted circles) centered on the optic nerve. FLAIR images (d-f) well depict the isointense masses causing enlargement of the optic nerves (dotted circles).](image1)

![Fig. 19 Perineural spread of tumor. Patient with squamous carcinoma of the retromolar trigone. CT sagittal images at 0 (a), 6 (b), and 12 (c) months follow-up. Note the progressive infiltration (arrows) of the pterygopalatine fossa (asterisks) with obliteration of the normal fatty tissue. In (c), it is evident the invasion of Meckel’s cave (empty arrow).](image2)
enhancement of the neighboring dura mater. Possibly associated bony changes can be osteolysis, hyperostosis, and enlargement of the adjacent foramina [21] (Fig. 23). Meningioma may also affect the optic nerve, appearing as homogeneous enhancing lesion around the optic nerve, with intermediate T1 and T2 signal [22].

Inflammation
Optic neuritis
The term optic neuritis (ON) usually refers to inflammatory process that involve optic nerve. Clinically, it presents with painful eye movements and visual loss. On MRI, the nerve appears swollen and hyperintense on T2-weighted sequences, with contrast enhancement best seen on fat-suppressed T1-weighted sequences [8] (Fig. 24). ON may be observed in autoimmune (for example multiple sclerosis and neuromyelitis optica spectrum disorders) and systemic diseases (sarcoidosis, lupus erythematosus, Wegener disease, Sicca syndrome, Behcet disease) [23]. Particularly, in neuromyelitis optica spectrum disorder (NMOSD), nerve involvement may be bilateral and more severe than in multiple sclerosis (MS). NMOSD is also characterized by involvement of spinal cord, with brain MRI features not suggestive for MS [8].

Tolosa-Hunt syndrome
Tolosa-Hunt is an uncommon disease related to a retro-orbital pseudotumor extending in the cavernous sinus [24, 25]. It is an idiopathic disorder that can manifest with a unilateral painful ophthalmoplegia, diplopia, and deficit of the V1 branch of trigeminal nerve. Bilateral pseudotumor is described in 5% of cases [24]. Tolosa-Hunt consists in a granulomatous inflammation of the lateral wall of the cavernous sinus or superior orbital fissure. Imaging techniques can show an infiltrative soft tissue mass within the cavernous sinus that appears enlarged. The tissue is hypo-isointense on T2-weighted sequences, with avid contrast enhancement on post-gadolinium images (Fig. 25). Differential diagnosis may include lymphoma, sarcoidosis, metastatic disease, meningioma, infection, granulomatous pachimeningitis, Meckel’s cave schwannoma, aneurysm, and pituitary macroadenoma [25]. Clinical findings may overestimate this condition and MRI is critical to exclude other similar conditions, allowing for precise management and therapeutic planning. After diagnosis, a good response to steroid therapy can be achieved.

Bell’s palsy
Bell’s palsy (idiopathic facial palsy) is the most common cause of unilateral peripheral nerve neuropathy. Clinical presentation is variable, with dysgeusia, mastoid pain, impaired salivation, and lacrimation. A Herpes-Simplex-Virus reactivation seems to be the most probable etiology [5]. Prognosis is often good and the treatment is based on corticosteroid therapy, often with the addition of acyclovir. Typical Bell’s palsy does not require imaging studies. However, MRI may be useful to exclude other lesions responsible of atypical Bell’s palsy (gradual-onset palsy, failure to improve with time) or recurrent palsy [26]. On MRI, the facial nerve shows increased enhancement on post-contrast sequences that may involve one or more segments, without nodularity (Fig. 26). Enhancement of distal intrameatal and labyrinthine segments is typical [26]. On T2-weighted sequences, the nerve may be hyperintense. In case of irregular or nodular enhancement, other causes of pathology (such as perineural spread of tumor) should be evaluated. Bilateral Bell’s palsy is unusual, often associated with HIV or other systemic pathologies like sarcoidosis, Lyme, and syphilis [5, 26–28].

Vestibular neuritis and labyrinthitis
Vestibular neuritis presents as unilateral acute vertigo without hearing loss. Nausea and vomiting are also associated. It affects young and mid-adults with no sexual preponderance. The etiology is various. Viral, bacterial, and protozoan infections have been reported, but also allergic and autoimmune causes are described. Inflammation of the vestibular nerve
**Fig. 21** Perineural spread of tumor. MRI T1-weighted post-gadolinium axial (a) and coronal (b) images. Note the enlargement and the enhancement of the mandibular branch of trigeminal nerve (V2) (dotted circles).

**Fig. 22** Leptomeningeal spread of tumor. Patient with lung carcinoma. MRI axial image of T1-weighted post-gadolinium sequence demonstrates a large metastasis of the pons (asterisks), bilateral leptomeningeal enhancement of trigeminal nerves (a, circles), and bilateral facial and vestibulocochlear nerves (b, dotted circles).

**Fig. 23** Sphenoid wing meningioma. MRI T1-weighted coronal image (a), T2-weighted (b), and T1-post gadolinium (c) axial images. A large mass with avid enhancement is visible, suggestive for meningioma of the right sphenoid wing (asterisks) with orbital involvement, displacement of the optic nerve (arrows), and the globe (empty arrows). Signs of hyperostosis are also evident (arrowhead).
may be complicated by demyelination, with loss of function, not always reversible [29]. MRI shows hyperintensity of the cisternal tract of the vestibular nerve on T2-weighted and FLAIR sequences [30] with enhancement on post-gadolinium images [31] (Fig. 27). Acute labyrinthitis is similar to vestibular neuritis with associated hearing loss and tinnitus [5].

Infection
Infectious meningitis may result from viral, bacterial, fungal, or parasitic infection. Viral infections, mostly related to *Herpes simplex virus type 1*, *Cytomegalovirus*, and *Varicella zoster*, may manifest with cranial nerve involvement and abnormal enhancement of the nerve on MRI [28]. Bacterial meningitis is typically caused by *Hae-mophilus Influenzae*, *Streptococcus pneumoniae*, and *Neisseria Meningitidis*. Imaging is often normal, although MRI post-contrast sequences can demonstrate leptomeningeal enhancement [5]. Intracranial Tuberculosis, typically in pediatric population, can manifest as leptomeningitis with involvement of cranial nerves (Fig. 28). *Cryptococcus neoformans* is associated with the Cryptococcal meningitis, characterized by optic neuropathy. Necrosis of the optic nerve and chiasm by cryptococcal organism have been described. *Rhinocerebral mucormycosis* is fungal infection in immunocompromised patients, with sinonasal disease that may progress to the orbit and cavernous sinus. It can be complicated by vascular and perineural invasion and local thrombotic infarction. Cranial neuroschistosomiasis, less common than the spinal form, is characterized by a granulomatous reaction that leads to increase of intracranial pressure and focal neurologic signs. Lyme disease, caused by *Borrelia burgdorferi*, may involve any of the cranial nerves, with a predilection for the facial nerve, that appears thickened and enhanced [28].

Trauma
Accidental or iatrogenic trauma, causing edema, hematoma, or disruption of the fibers, may lead to nerve impairment. Traction, stretching, impingement, and transection of the fibers are the typical mechanisms of injury [32]. High-resolution CT is the imaging modality of choice to detect skull base fractures and foraminal involvement. MRI may be useful to demonstrate intraneural edema or hemorrhage, especially on T2*-weighted sequences. Olfactory nerve may be involved in the closed injury of basal frontal lobe, with or without association of cribriform plate fracture (Fig. 29). Optic and oculomotor nerves may be affected in the orbital and optic canal fractures. Lesions of the abducens nerve are reported in the fracture of the clivus and petrous apex. Injuries of the facial nerve are associated to temporal bone fractures, that are classified as transverse or longitudinal based on their relationship to the long axis of the petrous temporal bone. They occur in 7% of temporal bone fractures with predilection for the geniculate ganglion area and the stylomastoid canal with edema,
Fig. 26 Bell’s Palsy. MRI T1-weighted pre- (a) and post-gadolinium (b), coronal images; T1-weighted post-contrast axial image (c). In a, b, note the swelling and enhancement of the left facial nerve (dotted circles); in c, the enhancement around the geniculate ganglion (arrow).

Fig. 27 Vestibular neuritis. MRI FLAIR (a) and T1-weighted post-gadolinium (b, c) axial images show mild hyperintensity (a, arrow) and enhancement (b, arrow) of the right vestibular nerve and right labyrinth (a, c, dotted circles).

Fig. 28 Tuberculosis (TBC). MRI T1-weighted post-gadolinium axial image demonstrates pathologic enhancing tissue involving the basal cistern and ventral surface of the brainstem (asterisks), with leptomeningeal enhancement of bilateral trigeminal (a, circles), facial, and vestibulocochlear nerves (b, dotted circles).
hematoma, or partial/complete transection of the nerve [5, 6, 33, 34]. Iatrogenic injuries may complicate a variety of surgical procedures. Facial nerve paralysis may follow temporal bone and parotid surgery, while lower cranial nerves may be involved during neck dissection; in particular, the recurrent laryngeal nerve may be injured during thyroid and parathyroid surgery [32].

Vascular

Neurovascular compression

Neurovascular compression syndromes are caused by vascular structures (usually arteries) that directly contact the cisternal segment of a cranial nerve (Fig. 30). Typically, the transition zone between the central and the peripheral myelin is the most vulnerable region of the nerve [35]. Clinical presentation is variable and includes trigeminal and glossopharyngeal neuralgia, vestibular paroxysm, and hemifacial spasm. However, neurovascular compression should be evaluated only in the presence of specific clinical features [6]. Also, nonhemorrhagic aneurysms may be associated to nerve dysfunction, for example in the case of vision loss due to compression of II nerve or optic chiasm by proximal internal carotid artery, and III, IV, or VI cranial nerves’ palsy secondary to intracavernous aneurysm. Imaging may be useful to detect the displacement and the compression of a nerve by redundant arteries. Steady-state sequences and corresponding Time-of-Flight (TOF) or 3D T1-weighted gadolinium-enhanced images are very useful for diagnosis and preoperative evaluation [35].

Fig. 30 Neurovascular compression of facial nerve. Patient with left facial nerve palsy. MRI axial steady-state free procession (a) and T1-weighted post-gadolinium, MIP reconstruction (b) images show a neurovascular compression of the left facial nerve (dotted circles)
Ischemia
Ischemic lesions of the brainstem may cause isolated nerve palsies, more commonly in elderly population. Cranial nerves between the III and the VI are the most affected. Diffusion-weighted imaging (DWI) sequences are the best to visualize an acute ischemic lesion (Figs. 31 and 32), whereas FLAIR and T2-weighted sequences are useful to visualize subacute or old infarctions [6]. The anatomical knowledge of the nuclei of the cranial nerves is fundamental to interpret ischemic nerve palsies.

Hemorrhage
Superficial siderosis
Superficial siderosis is a rare pathological condition that results from chronic deposition of hemosiderin in the subpial layers of the brain and spinal cord. Classically, it is bilateral and it may present with the typical triad: ataxia, sensorineural hearing loss, and myelopathy. Other clinical presentations include dysarthria, nystagmus, myelopathy, bladder dysfunction, sensory and pyramidal signs. Hemosiderin deposition has predilection for the superior cerebellar vermis, cerebellar folia, frontal lobe, temporal cortex, brainstem, spinal cord, nerve roots and cranial nerve, mainly the VIII. Rarely, extracranial nerve palsies, optic and trigeminal neuropathy have been reported in literature. MRI is useful to identify hemosiderin deposition, based on magnetic susceptibility, better demonstrated as hypointensity on T2*-weighted sequences (blooming effect) (Fig. 33) [36].

Conclusion
Cranial nerve dysfunctions may be the result of pathological processes of the cranial nerve itself or associated with tumor conditions, inflammation, infectious processes, or traumatic injuries. Neuroradiologists play a fundamental role in diagnosis. The knowledge of the anatomy of each nerve is crucial to detect the site of pathological alterations. Furthermore, it is necessary to know the most frequent pathologies of cranial nerves and their typical imaging features.

Abbreviations
CSF: Cerebrospinal fluid; CT: Computed tomography; DWI: Diffusion-weighted imaging; ENB: Esthesioneuroblastoma; FLAIR: Fluid-attenuated inversion recovery; MRI: Magnetic resonance imaging; MS: Multiple sclerosis; NF1: Neurofibromatosis 1; NF2: Neurofibromatosis 2; NMOSD: Neuromyelitis optica spectrum disorder; ON: Optic neuritis; PCA: Posterior cerebral artery; PICA: Posterior inferior cerebellar artery; PNI: Perineural invasion; PSN: Perineural spread; SCA: Superior cerebellar artery; SSFP: Steady state free procession; TOF: Time-of-flight

Guarantor
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