A wide spectrum of conditions may compress the spinal cord: degenerative disease, disc herniation and neoplasms are the most common causes; other conditions include trauma, epidural abscess and hematoma. The role of imaging is to establish a radiological diagnosis, to distinguish intrinsic spinal cord disease from extrinsic compression, to define mechanical spine stability and to evaluate the integrity of neural tissue. Clinical presentation, variable in severity and patterns of progression, influences the imaging approach. Imaging is required to assess osseous structures and soft tissues. The need to image ‘across anatomical compartments’ usually requires more than one imaging technique. Multidetector CT is the first-line imaging modality in acute spine trauma. MRI is the best imaging modality to assess soft tissues, image the extradural, intradural and spinal cord compartments. MRI needs to be implemented with fat-suppression techniques applied to T2- and contrast-enhanced T1-weighted sequences. MRI helps predict neurological recovery, providing macroscopic information including both reversible and irreversible histologic changes. Diffusion tensor imaging enables depiction of microstructural changes in the compressed spinal cord, which could further predict tissue viability, and functional prognosis. The role of CT myelography is reserved to selected cases, when MRI is not feasible or results are unclear but, intrathecal injection of contrast from below and above the compression level might be necessary to overcome the diagnostic limitations imposed by a myelographic block. This article summarizes the capabilities and limits of different imaging modalities in the spinal cord compression scenario, suggesting technical imaging protocols and diagnostic pathways, and discusses imaging aspects of various causes of spinal cord compression, following an organization based on their anatomical compartment of origin: osseous, epidural or intradural.

**Keywords:** compression • CT • differential diagnosis • imaging • magnetic resonance • mielopathy • myelo-CT • neoplasms • spinal cord injury

**Background**
Spinal cord compression (SCC) is a medical emergency that requires immediate imaging assessment and appropriate treatment to prevent or minimize neurologic sequelae that could potentially be devastating.

**Clinical presentation**
SCC can manifest with a wide range of clinical presentations, course and degrees of severity. Standardized classification of neurologic impairment is stated by the American Spinal Injury Association; such worldwide standards facilitate clinical patient care, as well as research in spinal cord injury medicine [1].

Generally, patients with SCC have a combination of motor and sensory dysfunction with a segmental distribution referable to one or few contiguous spinal levels [2]. Quadriplegia represents the most serious presentation and, depending on the level of the spinal lesion, there may also be respiratory failure. Acute or severe SCC may also
cause paraparesis or quadriparesis, hyporeflexia followed by hyperreflexia, extensor plantar responses, loss of sphincter tone and sensory deficits. The sensory deficit, with its most cranial involved dermatome, often marks the SCC level (sensory level). Progressive, generally nontraumatic SCC, may initially be associated with different and less specific symptoms, such as back pain, usually followed by the onset of radicular pain symptoms, weakness, sensory loss and changes in sphincter function. Gait disturbance and stumbling can be early symptoms of impending neurologic loss [3]. Urinary retention, secondary to autonomic dysfunction, is the most common bladder complaint, along with saddle distribution sensory loss.

Figure 1. Soft tissues abnormality visualization on CT images. (A & B) Sagittal reformatted and axial CT images show a right posterior paramedian C4–C5 disk herniation with encroachment on the central canal; (C & D) sagittal reformatted and axial CT images demonstrate hyperdense thickening of periodontoideum soft tissue consistent with a post-traumatic epidural hematoma.
for patients with evolving conus medullaris or cauda equina syndrome.

Why imaging?
To confirm a clinically suspected SCC, it is mandatory to perform an imaging work-up. The role of imaging is fundamental to establish a radiological diagnosis, to distinguish intrinsic spinal cord disease from extrinsic cord compression, to define mechanical spine stability, to evaluate the integrity of neural tissue, to provide further radiological information, and to guide treatment if necessary [4].

Imaging techniques
Imaging approaches rely on clinical features and anamnestic criteria. The appropriate use of each available imaging modality is based on the clinical setting, the complexity of spine anatomy and the relative capabilities/limits of each technique. The spine is a complex anatomical environment and, in the setting of SCC, imaging is requested to assess osseous structures and soft tissues, as muscles, discs, ligaments, epidural fat planes, meninges, cerebrospinal fluid (CSF), vessels and spinal cord. The need to image ‘across anatomical compartments’ usually requires more than one imaging technique to establish a final diagnosis [4].

Plain film radiography
Plain films represent the most accessible and less expensive imaging modality in spinal disorders. In general, plain films are acquired in a minimum of two

Figure 2. Spondylotic compressive changes with myelomalacia. (A) Sagittal T2-weighted turbo spin echo image shows degenerative cervical spondylotic changes causing spinal cord compression at two adjacent levels, with intramedullary focal well-defined hyperintense signal in the cord (arrow in A), indicative of chronic compressive myelopathy with gliosis and myelomalacia; (B & C) axial gradient-recalled echo T2*-weighted images confirm the abnormal intramedullary signal (arrowhead in B) and show the degree of central canal stenosis and cord compression at the two levels, due to right paracentral disk herniations.
views: anteroposterior and lateral; additional views are generally required in specific conditions (e.g., open mouth for dens evaluation, flexion/extension for spine stability, and oblique views for peduncles or vertebral foramina assessment). Plain films evaluate the osseous structures and the vertebral alignment but, especially in situations where a SCC is suspected, have severe sensitivity limits, and from a first-line diagnostic imaging tool for spine-related conditions in the past, are nowadays progressively abandoned in favor of more advanced and accurate imaging modalities.

Multidetector CT
Multidetector CT (MDCT) is unparalleled in its capability to detect bone abnormalities, it ensures fast acquisition times, and is progressively widely available on the territory. With an isotropic volumetric acquisition, it guarantees high-quality thin-section multi-planar reformatted images, with views for bone and soft tissues. Complex lesions can be more clearly depicted also using 3D capabilities of MDCT and, in the case of polytrauma patients, the images on the spine can be extracted from the data set of the patient’s total-body CT [5,6]. MDCT can also depict significant soft tissue abnormalities (e.g., disk herniations, ligamentous degenerative hypertrophy, epidural lesions or collections; Figure 1) [7–10]. The most important limitation of this technique remains the inability to provide accurate information on spinal cord lesions and ligamentous injury. Clinicians and radiologists should also consider the significant radiation exposure of patients undergoing MDCT, and therefore, especially in the young population, limit this examination to necessary clinical indications, use available radiation-reduction tools and protocols, and generally apply the ‘as low as reasonably achievable’ imaging principles [11–13].

Figure 3. Postdecompression visibility of compressive myelopathy effects. (A) Sagittal T2 preoperative image shows degenerative central canal stenosis and cord compression, with no clear high T2 signal in the cord. (B) After surgical decompression, the (C) postoperative sagittal T2-weighted image shows a focus of myelomalacia in the cord (arrow) at the level of previous compression.
MRI

MRI is the best imaging modality to assess soft tissues, ligaments, disks and spinal cord. Any patient with presumed SCC should undergo a MRI examination as soon as possible to reveal the location and severity of spinal cord lesion [8,14–15]. High magnetic field apparels (1.5–3.0 T) should be used to image the spinal cord. Accurate determination of intrinsic or extrinsic spinal cord lesions routinely requires the acquisition of two different pulse sequences, typically unenhanced T1- and T2-weighted images, in two orthogonal planes, usually axial and sagittal. Sagittal T2-weighted images are almost invariably acquired with fast spin echo (FSE) pulse sequences. Several T2-weighted pulse sequences are employed for axial imaging, FSE, gradient-recalled echo (GRE) and 3D constructive interference in the steady state myelographic images, each with specific advantages and limits. The images with the highest anatomical resolution, such as myelographic MRI suffer from intrinsic poor sensitivity for spinal cord parenchymal signal abnormalities; FSE images, with good sensitivity for spinal cord signal abnormalities are often disturbed by CSF flow artifacts, especially along the ventral aspect of the spinal cord. GRE images are insensitive to flow artifacts, and provide good contrast between spinal cord and CSF, yet maintain good sensitivity for spinal cord parenchyma signal abnormalities, and are able to differentiate ‘hard’ calcified or ossified spondylotic spurs (hypointense) from ‘soft’ disc herniations (hyperintense), which may have importance for surgical decisions [16,17]. GRE T1*-weighted images, with an echo time of 20–25 ms are employed, in case of trauma, to detect hemorrhagic spinal cord injuries. Contrast-enhanced T1-weighted images can be necessary if a neoplastic, infectious, or inflammatory condition is suspected, clinically or on the basis of unenhanced images, or when imaging a postoperative spine [18]. The presence of adipose tissue in spinal osseous structures, epidural space, peri- and para-spinal soft tissues implies the need for rigorous fat-suppression techniques to be applied to T2-weighted and contrast-enhanced T1-weighted images [19].

Limits of MRI are the usual contraindications (i.e., pacemaker, metallic foreign bodies in particular locations or claustrophobia), the length of scan, the absolute need for cooperation or of sedation/general anesthesia otherwise,
the difficulty to control vital parameters in critically ill patients, and the relative limited availability of MRI, especially in the emergency situations.

What do we look for on MRI?
The presence of SCC needs to be distinguished from noncompressive spinal cord abnormalities. Therefore, we look for masses or other pathological processes that exert extrinsic mass effect on the spinal cord. In a normal condition, the spinal cord is circumferentially surrounded by CSF, which offers a high contrast background for the visualization of the cord on T2-weighted images. Sagittal T2-weighted images usually offer the most panoramic overview of the spinal segment in examination. This set of images offers immediate evidence of the vertebral body alignment, the CSF surrounding the spinal cord, the morphology, course, size and signal of the spinal cord, and the presence of masses or collections compressing it. If a mass causing dislocation, deformation or compression of the cord is noted, attention to the presence of high T2 signal abnormality is necessary. Notably the presence of an ill-defined high T2 signal and swelling of the spinal cord parenchyma above and below the compression can suggest the presence of edema, suggesting acute compressive myelopathy (Supplementary Figure 1; see online at www.futuremedicine.com/doi/suppl.iim.13.71). Studies in animals [20,21] have shown that acute anterior SCC results in stenosis and occlusion of the epidural venous plexus, which causes breakdown of the blood–spinal cord barrier and results in vasogenic edema. The venous outflow impairment is more severe when compression occurs at two adjacent levels. Subsequently, compression causes impairment of arterial blood flow with cord ischemia and irreversible changes. In this chronic, terminal stage, MRI show a more well-defined high signal
on T2-weighted images, with volume loss indicative of gliosis and myelomalacia (Figure 2). Despite the high sensitivity of sagittal T2-weighted images, axial T2-weighted images are necessary to confirm the spinal cord signal abnormality, the true degree of a SCC and the severity of central canal stenosis (Figure 2B & 2C). It should be noted that in a severely compressed spinal cord, high T2 signal edema may not be visible at the level of maximum compression, but be evident cranially and caudally in the cord. In such cases, edema may become clearly evident once decompression of the cord is achieved, thereby displaying the MRI signs of compressive myelopathy (Figure 3). Contrast-enhanced images are not necessary in the trauma setting, but usually acquired when an infectious or neoplastic process is suspected. In the setting of SCC, on T1-weighted postcontrast images, a dilated and congested enhancing epidural venous plexus can be noted above and below the compressed level, more frequently in the ventral epidural space, but sometimes circumferentially; this represents a potential pitfall, mimicking abnormal enhancing epidural tissue or phlegmon.

In selected cases, to unmask a condition of clinically suspected cervical SCC, but with no clear evidence at standard MRI, MRI with neck flexion and extension can reveal dynamic stenosis (Figure 4) or Hirayama flexion myelopathy (Figure 5) [22,23].

Advanced MRI techniques: future perspective
Standard morphological T1- and T2-weighted images are not always accurate in grading the severity of the myelopathy, and the degree of SCC does not always correlate with clinical severity. Recent technological advances (e.g., diffusion-weighted imaging [DWI], diffusion tensor imaging [DTI] and tractography) have recently been applied to the study of the spinal cord, with the intent to characterize neural tissue at a microstructural level (Figure 6) [24]. Future applications of these techniques that actually are not routinely performed in the clinical ground, may be the assessment of the functional integrity of the axons within the white matter tracts of the compressed spinal cord, and the prognostic prediction upon its treatment [25–28].

Myelo-CT
Myelo-CT is reserved to selected cases when MRI is not feasible or results are unclear, and can render

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**Figure 6. Diffusion tensor tractography.** Diffusion tensor imaging can be performed on the spinal cord where it reveals microstructural arrangements of white matter fibers. In this case of degenerative cervical central canal stenosis and moderate cord compression, (A) MRI sagittal T2-weighted imaging does not reveal a definite signal abnormality in the cord, but (B) tractography shows abnormal arrangement and loss of continuity of the cord’s white matter fibers. (C & D) The control postsurgical decompression reveals reconstitution of normal white matter tracts. Images courtesy of Dr Meng Law, USC Keck Medical Center (CA, USA).
additional information compared with MDCT, with an accurate visualization of the intradural extramedullary compartment (Figure 7) [29]. Myelo-CT is less sensitive than MRI to metallic artifacts, especially in the case of implanted hardware, is not affected by flow artifacts, and offers high contrast and spatial resolution, but owing to complete lack of sensitivity in detecting intrinsic spinal cord abnormalities, can only partially replace MRI. In addition to its diagnostic limits, myelo-CT suffers from a series of disadvantages versus MRI, such as invasiveness of lumbar puncture and intrathecal contrast agent injection, ionizing radiation exposition, need of mobilization of the patient to allow for injected contrast to diffuse from the point of injection to the spine level of interest and the need of injection of contrast from below and above the compression level in case of myelographic block.

**How to image the patient with SCC: diagnostic pathways**

The SCC radiological work-up is generally designed upon the clinical setting. There is a general consensus that spine trauma patients need to be evaluated by emergency MDCT to assess presence of luxations or fractures [30,31]. While a negative MDCT exam is believed to be able to virtually rule out an unstable spine lesion by itself [32,33], patients with neurological symptoms, especially those suggesting SCC, absolutely need diagnostic complement by an emergency MRI, regardless of the positive or negative results of MDCT [34–38]. In all other nontraumatic spinal cord syndromes, in which SCC needs to be ruled out, with an acute, subacute or chronic presentation, MRI can be considered as first-line imaging method. MDCT should be considered as a complementary exam to further characterize osseous abnormalities revealed by MRI. Myelo-CT might be
useful in selected cases, when MRI is not possible or of unsatisfactory diagnostic quality, due to motion or metallic hardware artifacts, or when there is a poor correlation between symptoms and MRI, especially in the chronic degenerative setting [4].

**Lesion’s site of origin**

SCC may be caused by different pathologies arising from or involving different anatomical compartments. The differential diagnosis is strictly influenced by the recognition of the lesion’s compartment of origin. Once the anatomic site of lesion’s origin has been determined, a narrowed differential diagnosis requires consideration of both clinical and specific imaging data.

Table 1 shows an anatomically based differential diagnosis of the most common causes of SCC, depending on the site of origin of the lesion causing SCC.

To evaluate the complexity of SCC in a logical concise manner, we suggest to schematically divide the spinal environment into osseous, epidural and intradural compartments.

In the osseous compartment, we include the osseous spinal column, the discs and the spinal ligaments, bordering the central canal and the neuroforamina.

The epidural compartment is contained between the inner aspect of the bony central canal and the dura mater. Differently from the intracranial compartment, in the spine, the dura mater is not attached to the bone, therefore the epidural space is not virtual but a real space, and contains adipose tissue and a rich vascular network, forming the epidural venous plexus, which is more largely represented in the ventral epidural space. The intradural extramedullary compartment is outlined by the dura mater, and

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**Table 1. Spinal cord compression: site of origin for differential diagnosis.**

<table>
<thead>
<tr>
<th>From osseous compartment</th>
<th>From epidural space</th>
<th>From intradural space</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fractures with posterior wall retropulsion/facet dislocations</td>
<td>Epidural abscess</td>
<td>Primary neoplasm and metastasis</td>
</tr>
<tr>
<td>Metastasis and osseous spine tumor</td>
<td>Epidural hematoma</td>
<td>Subdural hematoma</td>
</tr>
<tr>
<td>Aggressive hemangioma</td>
<td>Disc herniation</td>
<td>Arachnoid cyst</td>
</tr>
<tr>
<td>Degenerative disease</td>
<td>Synovial cyst</td>
<td>–</td>
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</tbody>
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**Figure 8. Burst fracture.** (A) Axial, (B) sagittal reformatted and (C) 3D volume rendering CT images demonstrate the vertebral fracture due to high energy axial load compression force. The vertebral body shows comminution and outward displacement of fragments; the posterior wall is retropulsed and occupies the central canal (arrow in A & B). (D) Axial and (E) sagittal turbo spin echo T₂-weighted and (F) T₁-weighted images allow visualization of soft tissue and neural structures; the spinal cord is severely compressed, with focal high T₂ signal suggestive of edema (arrowhead in D).
includes the subdural and the subarachnoid spaces. While the subdural is a virtual space, between dura and arachnoid, the subarachnoid space is filled with CSF and contains leptomeninges, nerve roots, and arterial and venous intradural spinal vessels.

Finally the intradural intramedullary compartment represents the spinal cord itself, bordered by the pia mater.

**Cause of SCC from the osseous compartment**

**Traumatic fractures & dislocations**

Following a vertebral fracture or dislocation, an osseous fragment can violate the central canal and impinge on the spinal cord, or an entire vertebra can shift axially, disrupt and narrow the integrity of the central canal.

Axial load burst fractures, flexion tear-drop fractures and fractures–dislocations are the most common

**Figure 9. Fracture luxation.** (A & B) Sagittal reformatted multidetector CT images show facet fracture and dislocation (arrow in B) causing severe anterior spondylolisthesis and canal narrowing. After joint dislocation reduction, (C) MRI with sagittal T₂-weighted turbo spin echo and (D) T₂*-weighted gradient-recalled echo is obtained, showing satisfactory partial realignment, but also depicts other crucial findings: a large disk herniation caused by extensive disco–ligamentous complex injury, is still compressing the cord (arrowhead on C), and the cord shows signs of edema and low signal hemorrhage (arrowhead on D).

**Figure 10. Traumatic cord injury.** Four different examples of traumatic cord injuries of different severity. (A) Edematous nonhemorrhagic cord contusion revealed by high T₂ signal and cord swelling. (B) Hemorrhagic cord contusion revealed by low T₂* signal on gradient-recalled echo T₂*-weighted image. (C) Cord compression, with a deformed, angulated, edematous, but apparently intact cord. (D) Complete cord transection, as revealed by a cerebrospinal fluid-like signal gap in the cord at the level of fracture–dislocation and kyphosis (arrow).
traumatic causes of SCC with different underlying mechanism of injury.

Burst fractures consist of a vertically oriented fracture of the vertebral body with lateral dispersion of the fracture fragments: the posterior wall of the vertebral body is disrupted, the distance between the pedicles is widened, and there are associated fractures in the posterior elements. With burst fractures, the energy from the axial load is transferred to the vertebral bodies and intervertebral discs and consequently to neurologic structures, that experience the kinetic energy from the retropulsed vertebral bone fragments. MDCT allows an accurate and clear demonstration of the degree of vertebral comminution, as well as canal compromise by the retropulsed bony fragment (Figure 8) [39]. MRI allows visualization of neural structures and soft tissues [40], detects spinal cord lesions, disk herniations and epidural hematomas (EDHs), possibly contributing to the cord compression.

Flexion teardrop fractures are due to hyperflexion forces of great magnitude that cause a wedge deformity and a fracture of a large triangular portion of the antero–inferior portion of the vertebral body, associated with posterior ligament disruption. This type of spinal injury is usually complicated by middle columns fractures and retropulsion of bony fragments into the spinal canal (Supplementary Figure 2).

Fractures/dislocations consist of different degrees of anterior luxation of the upper vertebra above the lower vertebra consequent to uni- or bi-lateral facet dislocations associated with facet capsules, posterior

Figure 11. Pathologic fracture with spinal cord compression. (A) Sagittal short tau inversion recovery image reveals a collapsed vertebral body with a large epidural mass compressing the spinal cord; (B) sagittal and (C) axial contrast-enhanced fat-suppressed turbo spin echo T1-weighted images highlight the neoplastic tissue that involves the anterior epidural space. Axial image depicts the ‘draw-curtain sign’ (arrows), a sign indicative of neoplastic epidural involvement. (D) Sagittal reformatted CT image demonstrates the lytic pattern of neoplastic growth also at other vertebral levels.
longitudinal ligament, annulus fibrosus, and inter-laminar and interspinous ligament disruption (Figure 9). The spinal cord compromise results from kyphotic and translation deformities with shearing and stretching injuries [41]. MRI plays a fundamental role in the assessment of traumatic cord injury showing a large spectrum of abnormalities, such as cord compression, cord edema, hemorrhagic contusion and transection (Figure 10). Cord edema is detected as high T₂ signal and normal–low T₁ signal within the cord, with a variable grade of swelling [42]. Cord edema is not static and changes significantly within the first 2 weeks following injury; it can gradually resolve or it can evolve in a progressive ascend-

**Figure 12. Spinal cord compression from vertebral tumor.** Extensive myelomatous bony involvement with endocanalar epidural extension: (A) the mass is hypointense on the turbo spin echo T₁-weighted image, (B) slightly hyperintense on T₂-weighted image, with (C & D) intense and homogeneous enhancement on postcontrast fat suppressed images. The spinal cord is laterally displaced (arrow in B) and moderately compressed.

**Figure 13. Vertebral hemangioma with extra-osseous extension.** (A) Sagittal turbo spin echo T₁-weighted and (B) T₂-weighted images show typical MRI features of T₁- and T₂-hyperintense locally aggressive vertebral hemangioma extending into the spinal canal causing compression of the spinal cord. The vertebral body component shows minimal enhancement, while the epidural mass shows contrast enhancement, as depicted on (C) fat-suppressed enhanced T₁-weighted image. (D) Axial T₁-weighted image depicts typical 'draw-curtain' sign and stippled appearance of the vertebral body, representing thickened osseous trabeculae; the cord appears compressed and posteriorly dislocated.
In some cases the swollen and enlarged cord appears compressed in a central canal that would otherwise be of normal width; surgical decompression in such instances should be considered to reduce the risk of vascular cord compromise and further spinal cord damage. The presence of hemorrhage within the cord (hemorrhagic contusion) has been associated with unfavorable prognostic outcome. In these cases, a GRE $T_2^*$-weighted sequence, which we suggest to acquire in the sagittal plane, should be used and may reveal foci of signal loss indicative of presence of blood products (Figure 10B). Cord transection is the most severe cord injury characterized by loss of continuity of the cord, with a CSF or blood-filled gap indicative of complete disruption of the cord (Figure 10D).

**Osteoporotic fractures**

Insufficiency fractures, spontaneous or precipitated by minor trauma, are usually characterized by wedge, concave, biconcave or crush deformity. These are often subclinical, and when occurring at multiple levels can result in spinal deformity, but are almost invariably stable fractures. When posterior wall involvement is seen, this is typically seen at its superior corner, that can be retropulsed and impinge upon the central canal. Since this morphology of the fracture occurs more frequently at the lumbar level, when the retropulsion is significant, it is more frequently cause of radiculopathy.

**Pathologic fractures**

Neoplastic overgrowth inside the vertebra may be responsible for destruction of osseous architecture and

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**Figure 14. Aggressive vertebral hemangioma with atypical MRI features.** The lesion involves two adjacent thoracic vertebral levels that appear hyperintense on (A & D) turbo spin echo $T_2$-weighted images, but hypointense on (B) $T_1$-weighted image, with intense but somewhat inhomogeneous contrast enhancement on (C) enhanced fat-suppressed $T_1$-weighted image. (D) The neoplasm extends into the epidural space compressing the spinal cord. (E) Axial and (F) coronal reformatted CT images show the typical coarse and thickened appearance of the osseous trabeculae, spaced by large vascular spaces.
weakening of the vertebral body. This may result in an insufficiency fracture, but differently from an osteoporotic fracture, the neoplastic soft tissue invading the vertebral body can be displaced posteriorly in the central canal, and more frequently causes SCC. While signal abnormality or contrast-enhancement of the vertebral body cannot reliably distinguish a recent osteoporotic from a malignant fracture, the convex aspect of the posterior wall, a soft tissue epidural mass on MRI, a frankly lytic lesions on CT, or more rarely sclerotic changes, are rather specific signs, when present, of a pathologic fracture (Figure 11 & Supplementary Figure 3). It should be noted that nonfat-suppressed T2-weighted and enhanced T1-weighted images have low sensitivity for neoplastic osseous lesions, due to the confounding effect of an intrinsically high T1 and T2 signal of the normal yellow bone marrow. Fat-suppressed axial T2 and enhanced T1 images, with fat saturation, depict the extent of involvement of the epidural space, the degree of SCC versus dislocation and the presence of an abnormal T2 signal in the cord, the imaging hallmark of compressive myelopathy (Supplementary Figure 5).

Vertebral neoplasms

Tumors arising from the vertebral osseous structures can cause SCC, even in the absence of a pathologic fracture, in the case of extra-osseous neoplastic overgrowth and invasion of the epidural space. Multiple myeloma and osseous metastasis from solid tumors represent the most common causes of SCC from osseous spine tumors. MRI usually shows a low T1 signal, a high T2 signal and contrast enhancement in the affected bone and in the epidural mass (Figure 12 & Supplementary Figure 4). It should be noted that nonfat-suppressed T2-weighted and enhanced T1-weighted images have low sensitivity for neoplastic osseous lesions, due to the confounding effect of an intrinsically high T1 and T2 signal of the normal yellow bone marrow. Fat-suppressed axial T2 and enhanced T1 images, with fat saturation, depict the extent of involvement of the epidural space, the degree of SCC versus dislocation and the presence of an abnormal T2 signal in the cord, the imaging hallmark of compressive myelopathy (Supplementary Figure 5).

Figure 15. Chronic compressive myelopathy. (A) Sagittal turbo spin echo T1-weighted and (B) T2-weighted images show severe cervical spondylotic changes with central canal stenosis at C3–C4. The cord is compressed and displays a focal well defined area of high T2 signal (arrow on B), which is confirmed in the left side of the cord by (C) the axial gradient-recalled echo T2-weighted image (arrow on C). This focal cord lesion with sharp borders is supposed to represent an area of chronic gliosis and myelomalacia.
which further direct the diagnosis toward a neoplastic etiology. Clinical information and patient’s history are obviously essential data to reduce the spectrum of differential diagnosis.

Nonenhanced CT with a bone reconstruction algorithm is extremely helpful in defining both lytic and blastic lesions and in visualizing the presence of cortical destruction.

**Aggressive hemangioma**

Vertebral hemangioma is a postcapillary cavernous vascular malformation, usually characterized by extremely slow flow, which may expand the periosteum and extend into the soft tissues. Vertebral hemangioma may be symptomatic when progressive spinal cord or nerve–root compression occurs due to growth of the lesion beyond the confines of the vertebral body, or because of compression fracture or epidural hemorrhage [51]. MRI can show the typical hyperintense T1 and T2 signal (Figure 13). Presence of highly vascular or fibrotic components may render imaging characteristics atypical, with a low T1 signal (Figure 14), posing a more difficult differential diagnosis with a neoplastic lesion. Contrast-enhancement of such lesions is variable, can be intense or inhomogeneous (Figures 13 & 14). CT in such cases might be useful depicting the coarse and stippled appearance of the osseous trabeculae of the vertebral body and the bony expansion and remodeling (Figure 14). MRI is the method of choice to depict the extent to the posterior elements, to the epidural space, and the SCC [52].

![Figure 16. Epidural hematoma. (A & B) Sagittal reformatted and axial CT images show a L1 burst fracture with posterior wall retropulsion. (C) Sagittal turbo spin echo images evidence an extensive epidural collection hyperintense to the spinal cord on T2-weighted and (D) T1-weighted images. The collection appears more relatively hyperintense on fat-suppressed (E) T1-weighted and (F) T2-weighted images. (G & H) Axial turbo spin echo T2-weighted images evidence the severe mass effect on the spinal cord caused by the fractured posterior wall and the circumferential epidural hematoma.](image-url)
Degenerative osseous & nonosseous causes of SCC

Vertebral posterior osteophytes, disc-osteophyte complexes, ligamentous calcification and ossification, articular masses hypertrophy, and degenerative antero- and retro-listhesis are the most common cause of spinal stenosis, nerve root compression and compressive changes of the spinal cord, especially in the cervical region. These spondylotic changes, present with high prevalence in the elderly population, with different degrees of severity and with variable clinical expression, can be responsible of neurological symptoms that develop slowly, over months to years, generally related to vascular compromise of neural tissue. An accurate imaging diagnosis, and strict clinical correlation are important to guide treatment and to predict prognosis. MRI is the most commonly used tool for delineating the extent of the degenerative changes, the degree of narrowing of the spinal canal and the spinal cord alterations, secondary to edema, ischemia or myelomalacia. A standard imaging protocol for such conditions include sagittal T2- and T1-weighted sequences, completed with an axial T2-weighted set of images. CT can be of complementary use to precisely depict the anatomy of calcific and osseous structures, measuring precise diameters of central and foramina bony canals. Plain films, usually performed with dynamic views, can provide valuable information (e.g., spine stability) to guide treatment. CT myelography is used when MRI is not feasible, or results are incongruent with clinical data.

Typical MRI findings of compressive myelopathy is an intramedullary high signal on T2-weighted images at the level of cord compression, which can be fuzzy and vague, or intensely bright and well defined (Figure 15 & Supplementary Figure 1) [53,54]. It has been reported that high signal intensity of the spinal cord reflects pathological changes, both reversible and irreversible. Increased T2 signal intensity in the

Figure 17. Epidural abscess with spinal cord compression. (A) Sagittal fat-suppressed turbo spin echo T2-weighted image shows two possible foci of spondylodiscitis, at C3–C4 and C5–C6, with prevertebral soft tissues thickening and edema, likely a phlegmon, and a ventral epidural fluid collection from C2 to C5 compressing the spinal cord, likely an abscess. No contrast agent was injected due to renal insufficiency. The spinal cord shows high T2 signal (arrow on A) suggesting edema and myelopathy. (B) shows the CT-guided C3–C4 disc aspiration, which retrieved purulent fluid, probably also partially draining the abscess. Microbiology analysis revealed Staphylococcus aureus.
The spinal cord usually corresponds to water content or occasionally to the presence of blood or serum [55–58]. In a cadaveric study, Ohshio et al. [59] related high signal intensity on T2-weighted images of the cord to histologic changes, such as gliosis, edema, necrosis, myelomalacia or spongiform changes in the gray matter with edema, Wallerian degeneration, demyelination and necrosis of the white matter. There is controversy about the prognostic role of high signal intensity on T2-weighted images: several authors have reported intramedullary high signal as a sign of poor prognosis, while others reported no correlation with prognosis [60–62]. A study [60] found that a predominantly well-defined area of intramedullary high signal intensity on T2-weighted has a poor prognostic meaning towards recovery after surgical decompression, whereas patients with predominantly ‘faint and fuzzy’ borders area of intramedullary high signal intensity have the same outcome as those without signal intensity changes. Another study reported that patients with multisegmental areas of high T2 signal intensity have a poorer prognosis than patients with unisegmental areas of abnormal signal intensity [63]. Intramedullary contrast enhancement in compressive myelopathy has also been described [64,65], and some authors have reported a negative prognostic factor for recovery in patients undergoing decompressive surgery (Supplementary Figure 6) [66].

The most recent consensuses seem to point to a rather poor specificity of conventional MRI findings, such as a high T2 signal in the cord, in discriminating potentially reversible edema and ischemia from irreversible myelomalacia and gliosis; several researches on DWI and DTI on spinal cord have been carried out in the past few years and have showed superior correlation of DWI and DTI measurements with clinical and electrophysiological signs of myelopathy, compared with conventional MRI [67–72]. Similarly, fractional anisotropy measurements seem to be able to predict recovery upon surgical decompression [28]. Nevertheless, these techniques have not yet been implemented in routine imaging, and require further research.

**Spinal cord injury without radiographic abnormality**

Spinal cord injury without radiographic abnormality is defined as the presence of spinal cord injury in the absence of a fracture or dislocation on plain radiography or CT scan. An MRI rules out SCC in the patient with neurological deficit following spinal trauma. Spinal cord injury without radiographic abnormality is usually due to contusion or cord stretching. It occurs in the pediatric population, as a consequence of children’s vertebral column elasticity and vulnerability to deforming forces, and in elderly patients with pre-existing cervical degenerative changes, such as spondylosis.

![Figure 18. Epidural abscess with spinal cord compression.](image-url)

(A) Sagittal turbo spin echo images show an extensive posterior epidural collection, hyperintense on T2-weighted image, (B) hypointense on T1-weighted image with (C) peripheral enhancement on fat-suppressed enhanced T1-weighted image. (D) Axial turbo spin echo fat-suppressed T2-weighted image depicts the extradural location of the fluid collection well, with anterior displacement of the dura (arrow on D), and the mass effect on the spinal cord. (E) Axial diffusion-weighted image shows high signal intensity within the extradural abscess. (F & G) display the cord swelling and edema, well visible after surgical decompression of the abscess, despite resolution of the spinal cord compression (also see Figure 4).
or disc herniation, resulting in narrowing of the sagittal diameter of the spinal canal. Impingement of the cervical cord by these degenerative changes may explain the mechanism of spinal cord injury in these cases [73–75].

Causes of SCC from the epidural compartment
Epidural hematoma
EDHs are frequently encountered in the setting of spinal trauma or after spinal surgery [76,77]; spontaneous EDHs are rare, generally occurring in older patients (aged 50–80 years) with risk factors such as coagulopathy [78]. Since the dura is not firmly attached to the bones in the spine, EDHs can also derive from low pressure bleeding phenomena, such as venous hemorrhages. The imaging characteristics of EDH vary with the oxidative state of the hemorrhage and the clot retraction. MDCT, providing the first imaging findings in the trauma setting, with the soft tissue window setting, is often helpful in detecting the hyperdense, frequently crescent-shaped acute hematoma; however, MDCT might suffer from low sensitivity in regions with artifacts or when the EDH is isodense. MRI is regarded as the imaging modality of choice to evaluate EDH; depending on the status of the hemoglobin breakdown products, EDHs have variable signal intensity, from isointense to hyperintense on T1-weighted sequences, and from hypointense to hyperintense on T2-weighted sequences (Supplementary Figure 7).

For EDHs that appear hyperintense on T1- and T2-weighted sequences, fat suppression, also applied to the nonenhanced T1- and T2-weighted images, allows better depiction of the extension of the hematoma against the dark background of the suppressed fat signal, and better appreciation of the relationship with surrounding structures (Figure 16) [79].

Hematomas generally do not show significant enhancement (Supplementary Figure 7), especially in the early stages; this can be an important diagnostic finding in differentiating these lesions from abscess or tumor.

Figure 19. Disc herniations. (A–C) Case 1 with (A) sagittal and (B) axial turbo spin echo T2-weighted images showing multilevel cervical spondylotic changes with disc/osteophytes complexes, central canal stenosis, cord compression and apparently chronic myelopathic changes. Note that on turbo spin echo, soft disc herniations and disc–osteophyte complexes are not easily differentiated because both appear T2-hypointense (arrows on A & B). In such cases, (C) the corresponding axial gradient-recalled echo T2-weighted image depicts the soft disc component as hyperintense (arrow on C), while an osteophyte would have appeared markedly hypointense. (D & E) Case 2 displays a large acute traumatic disc herniation compressing the spinal cord. Note the very high T2 signal in the disc fragment due to high water content, and the ill-defined high T1 signal in the cord, meaning cord edema, on the (E) axial gradient-recalled echo T1-weighted image.
Epidural abscess

Epidural abscess is a dreadful complication of spondylodiscitis. The phlegmon is usually isointense or hypointense on T₁-weighted images, hyperintense on T₂-weighted images, with peripheral enhancement after gadolinium injection [80]. Fat-suppressed T₁-weighted and contrast-enhanced T₁-weighted sequences are helpful to highlight the infectious site of origin which is often the disc space and subchondral areas along the disc endplates (Figure 17), and to depict the typical peripheral enhancement and central fluid component of the collection (Figure 18) [19]. The contrast-enhancing area often extends in the adjacent soft tissues, beyond the peripheral collection rim, indicating the phlegmonous component, characterized by inflammatory response and edema (Supplementary Figure 8). Recently, DWI has been used to study epidural abscesses; the purulent fluid collection, when the causative agent is a pyogenic germ, appears markedly hyperintense on DWI (Figure 18), and hypointense on the corresponding apparent diffusion coefficient map [81].

CT is performed as a complementary imaging study, generally when MRI is not available or contraindicated. It can render additional information on erosive changes of the bones, especially along the disc endplates in case of spondylodiscitis, and it is also a valuable guide for percutaneous biopsies (Figure 17) [82].

Disc herniations & synovial cysts

Disc herniations are among the most commonly encountered findings in MRI of the cervical and lumbar spine, less commonly found in the thoracic spine. A common cause of radicular symptoms and signs, due to direct compressive effects and/or local inflammatory response, they are more rarely cause of SCC. Through the rupture or fissuration of the annulus fibrosus and posterior longitudinal ligament, as a result of trauma or much more frequently in relation to degenerative phenomena, extruded disc material may extend into the epidural space causing cord compression. The herniating disc material can be contained by some residual fibers of the stretched and deformed annulus, or be completely extruded in the epidural space. Large disc herniations or a combination of disc herniation on a pre-existing central canal stenosis can cause significant SCC (Figure 19). Sagittal and axial T₁-weighted images depict the disc herniation, characterize it on the base of the morphology and continuity with the parent disc as contained or extruded, assess the mass effect on the spinal cord, and the presence of cord edema. The signal of the disc fragment can be variable, usually depending on the hydration status of the nuclear material. It is important to note that the herniated disc fragment can display a signal different from the parent disc. The use of gradient echo pulse sequences performed on the axial plane greatly enhances the ability to obtain diagnostic images for the clinical evaluation of cervical disc herniation, offering a sharp delineation of bone/osteophytes, that show low signal, and disc margin, with high signal (Figure 19), an excellent contrast between the spinal cord and surrounding subarachnoid space, and clear visualization of the neural foramina and exiting nerve roots (Figure 19) [83,84]. In case a contrast agent is injected, the disc fragment can have a peripheral rim of enhancement, better visible on fat-suppressed
enhanced T1-weighted images, which might help in the differential diagnosis with a neoplastic mass, such as a meningioma or a schwannoma.

A rarer entity, especially in the cervical and thoracic spine, which can represent a cause of SCC from the epidural space, is the synovial cyst. It most commonly arises in the setting of degenerative changes and originates from the facet joints. If the synovial cyst spurs from the anterior and medial margin of the articular capsule and grows in the central canal, it can compress the spinal cord from the postero-lateral epidural space. The clearest diagnosis of a synovial cyst is provided by MRI, which characteristically shows a well-circumscribed cystic mass, with a fluid-like signal on T1- and T2-weighted images, typically with a thin capsule and no contrast enhancement [85]. Variations to these typical features are possible, with cysts showing calcific walls, hyperproteinaceous fluid content, with high T1- and low T2 signals. Axial T2-weighted images again demonstrate the cyst, its relationship with the facet joint, its epidural location, mass effect on the spinal cord, and imaging signs of myelopathy when present. A diagnostic doubt can sometimes be cleared by CT-guided injection of contrast agent in the facet joint space, resulting in contrast opacification of the cyst.

Extramedullary hematopoiesis

Extramedullary hematopoiesis (EMH) is a physiologic response to chronic anemia observed in various hematologic disorders, such as thalassemias, myelofibrosis and polycythemia [86]. EMH is commonly seen in the abdomen or chest, rarely in the epidural space [87]. MRI is obviously the first selected imaging modality for the diagnosis of EMH, clearly showing the site and extent of the lesion in relation to the displaced spinal cord. Active recent hematopoietic extramedullary lesions have rich vasculature, while inactive older lesions have more fatty tissue and iron deposits [88]. If the patient is treated with blood transfusions, the lesion may decrease in size and

Figure 21. Subdural hematoma. (A) Sagittal turbo spin echo T1- and (B) T2-weighted images show a hyperintense intradural extramedullary collection anterior and posterior to the spinal cord (arrows in A & B). The thin dark dural line is seen at its place (arrowhead in A), defining the intradural location of the collection. (C & D) Axial turbo spin echo fat-suppressed T2-weighted images reveal a circumferential disposition of the collection around the spinal cord with a festooned appearance (arrows in C & D).
appear on MRI with massive iron deposition. Active lesions show intermediate signal intensity in both T₁- and T₂-weighted MRI; enhancement is minimal or absent differentiating it from other epidural lesions such as abscesses or metastases. Older inactive lesions show high signal intensity in both T₁- and T₂-weighted MRI due to fatty infiltration or low signal intensity in both T₁- and T₂-weighted MRI due to iron deposition.

Angiolipoma
Angiolipoma of the spine is a benign neoplasm consisting of both mature fatty tissue and abnormal vascular elements, usually characterized by a slow progressive clinical course [89]. Because angiolipomas are mostly hyperintense on T₁-weighted images and often nearly isointense with epidural fat, fat-suppression MRI is particularly well suited for the investigation of these tumors. The lesion can also present a variable grade of hypointensity, predictive of its degree of vascularity, likely to be encountered at surgery [90]. On T₁-weighted images, the tumors have variable signal intensity attributed to the variable vascular and adipose elements of the tumor and most lesions enhance after gadolinium administration.

Causes of SCC from the intradural compartment
Spinal subdural hematoma
Spinal subdural hematomas (SSDH) are contained within the virtual space existing between the apposed dural and arachnoidal meningeal sheets. They can be caused by abnormalities of coagulation, blood dyscrasias, trauma, underlying neoplasm and arteriovenous malformation, and invasive spinal procedures [91]. MRI is the preferred imaging modality to evaluate subdural hematomas. It provides information on their location, extent, morphology and variable T₁ and T₂ signals related to hemoglobin’s breakdown products. A SSDH is recognized and distinguished by an EDH by the observation that the thin T₁-hypointense line of the dura is not displaced and the T₁-hyperintense epidural fat is regularly seen. SSDH can extend circumferentially inside and along the borders of the thecal sac, by which it possibly causes SCC, and displays a typical appearance on axial images (Figures 20 & 21) [92].

Figure 22. Meningioma. (A) Sagittal turbo spin echo T₂-weighted, (B) coronal 3D constructive interference in the steady state and (C) axial and (D) sagittal turbo spin echo fat-suppressed enhanced T₁-weighted images show an expansile intradural-extramedullary mass with homogeneous enhancement laterally displacing the spinal cord. Small dural tail is apparent on high-resolution 3D constructive interference in the steady state image (arrow on B).
Primary neoplasms
Intradural–extramedullary tumors lie within the dural borders of the thecal sac, but outside of the spinal cord. Meningioma and schwannoma are the most common primary tumors in this location. These are expansile masses contained by the T2-hypointense thin dural line, that determine extrinsic dislocation/compression of the spinal cord, that does not appear enlarged, differently from intramedullary tumors. The epidural fat lining, visible as a T1-hyperintense rim around the thecal sac, is preserved. CT can be helpful in defining mineralization or bone remodeling in these tumors, but MRI with multiplanar T2-weighted and contrast-enhanced T1-weighted sequences, clearly defines tissue characteristics, the soft tissue planes and neural structures surrounding these lesions. In consideration of the intradural location of these masses, the background to mass contrast is assured by the presence of the CSF, therefore, in theory, use of fat-suppression techniques should not be necessary; although, in practical terms, if a mass is large enough to expand the thecal sac, the borders of the mass may be very close to the adipous epidural compartment, and still fat suppression applied to the T2-weighted and to the contrast-enhanced T1-weighted sequences helps to delineate the real borders of the mass. Imaging differentiation of meningioma from schwannoma is not always easy. T1 and T2 signal characteristics often overlap, with schwannomas demonstrating slightly more heterogeneously increased T2 signals. Dural tail and calcifications are predictors of meningioma, whereas an expansile-enhancing mass, extending across the intervertebral foramen, is strongly suggestive of a neural sheet tumor, although meningiomas are also known to display this behavior (Figures 22 & 23) [93,94].

Intradural & leptomeningeal metastasis
The subarachnoid space is a frequent localization for metastatic deposits. MRI is the preferred imaging modality [95] able to demonstrate region of bulky disease, represented by intradural and extramedullary nodules, to visualize clumping of nerve root and spinal cord alterations. In a patient being investigated for suspected SCC, the T1-weighted sequence may show a dislocation/compression of the spinal cord and high signal
within it, due to edema adjacent to a pathological extra-
medullary deposit. The demonstration of intradural
metastasis and leptomeningeal lesions is easily obtained
on conventional postcontrast T1-weighted images, but
the use of the postcontrast images with fat suppression
increases the evidence of leptomeningeal linear and
nodular enhancement areas along the spinal cord sur-
face and nerve roots, resulting in better differentiation
from dural thickening/enhancement [19]. In the case of
infectious and inflammatory processes, the pattern of
enhancement is usually linear and uniform, whereas it
tends to be thicker and nodular in cases of tumor spread.
The appearance of advanced leptomeningeal carcinoma-
tosis, characterized by thick perimedullary and perineu-
ral enhancement, has been described as ‘sugar coating’
for its appearance on contrast-enhanced T1-weighted
images (Figure 24) [96].

**Arachnoid cyst**
Spinal arachnoid cysts are rare lesions most often
located in the mid-to-lower thoracic spine. Most spi-
nal arachnoid cysts are asymptomatic and discovered
incidentally by MRI; neuropathic pain, numbness,
weakness and myelopathy can be present due to mass
effect on the spinal cord [97]. MRI is the imaging
modality of choice to assess the size, nature and extent
of cystic lesion, as well as the mass effect on the cord
(Figure 25) [98,99]. On T1- and T2-weighted images, the
signal within a cyst has the same intensity as CSF. On
thin axial T2-weighted images, it is possible to assess
the mass effect upon the spinal cord, and the possibly
associated MRI signs of myelopathy, such as edema
and gliosis. High-resolution thin sections (<1 mm)
myelographic MRI pulse sequences, heavily fluid sen-
titive, can precisely depict the cyst walls. The spinal
cord can appear flattened. Large cysts can display
flow-related artifacts, caused by transmitted pulsations
from the circulating CSF. CT can be helpful to assess
the remodeling of bone canal determined by the cyst.
CT myelography can be used as a complementary and
often confirmatory exam, since it may demonstrate or
exclude a communication between the subarachnoid
space and the cyst, which is important for surgical
planning [100].

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**Figure 24. Two cases of meningeal metastasis.**
(A) A turbo spin echo fat-suppressed enhanced T1-weighted image shows dural nodular-enhancing deposits compressing the spinal cord, in this patient with primary lung cancer.
(B & C) Sagittal turbo spin echo T2-weighted and fat-suppressed enhanced T1-weighted images show extensive bulky subarachnoid leptomeningeal carcinomatosis, compressing the spinal cord, from a myxopapillary ependymoma.
Neuroenteric cyst
Neuroenteric cysts are rare congenital epithelium-lined cysts of the CNS; they are usually located anteriorly to the spinal cord. The most common myelographic and CT myelographic appearance is that of a lobulated intradural-extradural mass. Neuroenteric cysts are usually isointense to hyperintense relative to CSF on long repetition time sequences and isointense or slightly hyperintense to CSF on T1-weighted images. These signal characteristics correlate with the high-protein-content fluid within the cysts, described at surgery as milky or mucinous.

Intramedullary lesion
An enlargement of the spinal cord, due to vascular, neoplastic or inflammatory processes, may determine an obliteration of CSF surrounding the spinal cord, causing a SCC through a vicious circle of congestive myelopathy.

Conclusion
Imaging is essential to confirm and characterize a clinical suspect of SCC. CT is the preferred imaging modality to assess bone structures, especially in the trauma setting, whereas MRI is advantageous in the evaluation of soft tissues and is unique in the visualization of spinal cord, both in acute, sub-acute and chronic settings. For a schematic approach to the imaging differential diagnosis of the various causes of SCC, it is important to determine the site of origin of the lesion, whether from the osseous, epidural, or intradural compartment in the spine. Further research will likely bring to the clinical arena advanced microstructural imaging techniques able to provide diagnostic and prognostic information in various SCC conditions.

Future perspective
Recent technological advances (e.g., DWI, DTI and tractography) will likely be able to further characterize neural tissue at a microstructural level, predicting tissue viability, functional prognosis and patient recovery.

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Imaging in spinal cord compression

- Multidetector CT (MDCT) is the preferred imaging modality to assess bone structures especially in acute trauma patients and absolutely needs diagnostic complemeny by MRI when spinal cord compression (SCC) is suspected. MRI can be considered as the first-line imaging method in all other nontraumatic spinal cord syndromes in which SCC needs to be ruled out.

Strategies to approach SCC

- SCC may be caused by different pathologies arising from or involving different anatomical compartments. The recognition of the lesions' compartment of origin narrows the broad spectrum of differential diagnosis.

Anatomically based causes of SCC

- From the osseous compartment: traumatic fractures and dislocations, osteoporotic and pathologic fractures, vertebral neoplasms, aggressive hemangioma and degenerative changes such as vertebral posterior osteophytes, disc-osteophyte complexes, ligamentous calcification and ossification, articular masses hypertrophy, and antero/retrolisthesis.
- From the epidural compartment: hematoma, abscess, disc herniations and synovial cyst.
- From intradural compartment: subdural hematoma, primary neoplasms, intradural and leptomeningeal metastasis.

References

Papers of special note have been highlighted as:

- of considerable interest
5. Discusses various imaging modalities used in different clinical contexts of myelopathy.
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