

Spinal Syndromes (GENERAL)

Updated: May 1, 2010

SPINAL STRUCTURES AFFECTED IN VARIOUS DISORDERS 1

ANATOMIC SPINAL SYNDROMES 1

CERVICAL CORD 2

THORACIC CORD 2

LUMBOSACRAL CORD 2

SPINAL CORD TRANSECTION, SPINAL SHOCK 2

SPINAL CORD HEMISECTION 3

 Brown-Sequard syndrome 3

 Dorsal (Posterior) Hemisection 3

 Ventral (Anterior) Hemisection 3

ANTERIOR 2/3 TRANSECTION 3

EXTRAMEDULLARY CORD COMPRESSION 3

CENTRAL CORD SYNDROME 4

CAUDA EQUINA VS. CONUS MEDULLARIS SYNDROME 5

SPINAL COMPLICATIONS 5

 Paresthesias 5

 Decubitus Ulcers 5

 Bladder dysfunction 5

 GI complications 6

 Sexual dysfunction 6

 Malnutrition 6

 Respiratory Failure 6

 Venous Thrombosis & Pulmonary Embolism 6

 Spasticity 6

 Autonomic Dysfunction 6

 Psychiatric Dysfunction 7

SPINAL PROGNOSIS 7

SPINAL REHABILITATION 7

Anatomic localization of MOTOR SYMPTOMS, UMN & LMN lesions → see p. Mov3 >>

Anatomic localization of SENSORY SYMPTOMS → see p. S22 >>

RADICULOPATHY → see p. PN1 >>

Spinal IMAGING → see p. D70 >>

INFECTIONS of spinal cord and vertebrae → see p. Inf3 >>, p. Inf5 >>, p. Inf7 >>

TUMORS of spinal cord and vertebrae → see p. Onc50 >>, p. Onc54 >>

How **vertebra** corresponds to **spinal segment**:

- C1: C1
- C2-7: +1
- T1-6: +2
- T7-9: +3
- T10: L1-2
- T11: L3-4
- T12-L1: other

SPINAL STRUCTURES AFFECTED IN VARIOUS DISORDERS

	Dorsal funiculi (fasc. gracilis & cuneatus)	Lateral funiculi (lateral pyramidal tract, UMN)	Anterior horn (LMN)	Anterolateral system	Intermediolateral column (central autonomic motoneuron)
Subacute Combined Degeneration (vit.B12 def.)	+	+			
ALS		+	+		
Primary Lateral Sclerosis		+			
Familial Spastic Paraplegia	±	+			
Spinal Muscular Atrophy (SMA), Progressive Bulbar Palsy			+		
Syringomyelia	±	±	+	+ (decussating fibers)	±
Tabes dorsalis	+				
Multiple sclerosis	+	+		+	
Poliomyelitis			+		
Shy-Drager syndrome		±	±		+
Tropical spastic paraparesis (HTLV)	±	+			
HIV vacuolar myelopathy	+	+			

Anatomic Spinal Syndromes

- spinal cord contains, in small cross-sectional area, almost entire **motor output** and **sensory input** of trunk and limbs - *spinal cord disorders are frequently devastating.*

LEVEL

(below which sensory / motor / autonomic function is disturbed) – *hallmark of spinal cord damage!* (reflects spinal cord's segmental functional organization)

Sensory level

- use **painful** (sharp pinprick) / **temperature** (dry tuning fork after immersion in cold water) stimulus applied to low back and sequentially moved up toward neck on each side.
- such sensory level (damage to **spinothalamic tract**) is located 1-2 segments* below actual level of **unilateral** spinal cord lesion (but it may be at level of lesion when **bilateral**).
*sensory fibers after synapse in dorsal horn, ascend ipsilaterally for several segments before crossing just anterior to central canal to join opposite spinothalamic tract.

Sweating level – determined by drawing spoon up torso. see p. D1 >>

SEGMENTAL SIGNS

- indicate *upper and lower levels of spinal cord lesion*:

- 1) band of altered **sensation** (hyperalgesia, hyperpathia).
- 2) flaccid paralysis, fasciculations, atrophy in **muscles** innervated by damaged segments.
- 3) absent deep tendon **reflex**.

N.B. with acute transverse lesions, SPINAL SHOCK may be mistaken for extensive damage to many cord segments or polyneuropathy (e.g. Guillain-Barré).

CERVICAL CORD

Best localized by **WEAKNESS pattern** (sensory deficits have less localizing value):

1) **cervicomedullary junction**:

- *extensive lesions* involve adjacent **medullary centers** → vasomotor and respiratory collapse → neurogenic hypotension, apnea → unresponsiveness (difficult diagnosis) → death (in absence of ventilatory support).
- *partial lesions* interrupt **decussating pyramidal tract fibers destined for legs** (cross below those of arms) → "crural paresis" of lower limbs.
- *compressive lesions* produce weakness of ipsilateral shoulder & arm → ipsilateral leg → contralateral leg → contralateral arm.

2) **high cervical cord lesions - life-threatening** (quadriplegia and respiratory paralysis*).

*breathing possible only by accessory muscles of respiration.

3) C₄₋₅ - quadriplegia with **preserved respiratory function** (functional diaphragm)

4) C₅₋₆ - **sparing shoulder muscles** (loss of biceps and brachioradialis reflexes).

5) C₇ - **sparing biceps** (loss of triceps reflex).

6) C₈ - **sparing triceps** (paralyzed fingers and wrist flexion); effort to close hand → extension of wrist and slight flexion of fingers ("preacher's hand").

- ipsilateral **HORNER'S SYNDROME** may occur at any cervical level lesion.
- damage to **spinal tract of trigeminal nerve** in high cervical region → characteristic **ONION-SKIN PATTERN FACE ANESTHESIA**.

THORACIC CORD

Best localized by **SENSORY LEVEL on trunk** - nipples (T₄), umbilicus (T₁₀), etc. see p. D1 >>

- observe **abdominal wall musculature** and **umbilicus** by asking patient to interlock fingers behind head in supine position and attempt to sit up:
 - **lesions below T₉** paralyze lower abdominal muscles → upward movement of umbilicus (**BEEVOR sign**) + loss of lower superficial abdominal reflexes.
 - **unilateral lesions** → movement of umbilicus to normal side; absent superficial abdominal reflexes on involved side.
- **midline back pain** is useful localizing sign.

LUMBOSACRAL CORD

- lumbar-sacral segments progressively decrease in size - focal lesions are less easily localized.
- L₁₋₂ - cremasteric reflex.
- L₂₋₄ - thigh flexion and adduction, knee extension / patellar reflex.
- L_{5-S₁} - thigh extension, knee flexion, foot and ankle movements / ankle jerk.
- S₂₋₄ - anal sphincter tone / anal wink reflex.

SPINAL CORD TRANSECTION, SPINAL SHOCK

In all vertebrates, acute **spinal cord concussion** or **complete cord transection** is followed by **SPINAL SHOCK** - transient profound depression of all SPINAL REFLEXES below level of injury (in addition to complete PARALYSIS and ANESTHESIA below level):

1. **Flaccid paralysis**
2. **Absence of reflexes** (muscle stretch, plantar, abdominal & cremasteric)
3. **Hypotonic paralysis of bowel & bladder** (ileus, gastroparesis, urinary and bowel retention) ± **priapism**.
4. **Hypotension*** (not present if lesion is below lower thoracic level) with **anhidrosis** and **flushed warm peripheral skin** (→ **poikilothermy**).

*without compensatory tachycardia (if high cervical lesion), i.e. **NEUROGENIC SHOCK** (interrupted sympathetic outflow → vasodilation & bradycardia)

N.B. it is possible to diagnose only **UPPER LEVEL OF INJURY** – sensory loss & flaccid paralysis level.

- **ascending myelitis** - ascending spinal cord edema may rise upper level – may reach dangerous levels (C₄ and above); descending edema is asymptomatic.
- **CAUSE of spinal shock** is uncertain (cessation of tonic bombardment of spinal neurons by excitatory impulses in descending pathways undoubtedly plays role).
 - resting membrane potential of spinal motoneurons is 2-6 mV greater than normal.
- **spinal shock DURATION** is proportionate to *degree of encephalization of motor function* in various species:
 - in frogs & rats it lasts for minutes;
 - in dogs & cats it lasts for 1-2 hours;
 - in monkeys it lasts for days;
 - in humans it lasts for **minimum of 2 weeks** (if complications* are present - it is much longer!)
 - *e.g. infection, malnutrition, anemia, bedsores
- spinal shock may superficially resemble Guillain-Barré syndrome.

RECOVERY FROM SPINAL SHOCK

- SPINAL REFLEXES below level return and become hyperactive (chronic stage of UMN lesion - flaccid plegia changes to spastic paralysis).

- when reflex activity below level returns (i.e. spinal shock is over), check again for sensation / voluntary motor control **below level** – if any is returned, cord transection is incomplete!
- **at lesion level, segmental LMN signs persist** (injury to anterior horns or ventral roots); level where peripheral (LMN) and central (UMN) paralysis abut is reliable indicator of lower level of spinal cord injury!

Now it becomes possible to delineate **UPPER & LOWER LEVELS OF INJURY**.

- recovery of reflex excitability may be due to:
 - 1) **denervation hypersensitivity** to mediators released by remaining spinal excitatory endings.
 - 2) **sprouting of collaterals** from existing neurons → additional excitatory endings on interneurons and motoneurons.

First reflexes to reappear:

- a) sacral reflexes (bulbocavernosus, anal wink)!!! – may return within 24 hours of injury!
- b) slight contraction of leg flexors and adductors in response to noxious stimulus.
- c) knee jerks.

Once spinal reflexes begin to reappear, their threshold steadily drops.

- various different stimuli may evoke **REFLEX SPASMS** (flexor or extensor) that involve many or all of paralyzed muscles;
 - if cord section is incomplete, spasms can be associated with particularly bothersome **pain bursts** (H: **BACLOFEN**).
 - repeated flexor spasms may occur for prolonged periods → **contractures of flexor muscles**.
- afferent stimuli irradiate from one spinal reflex center to another:
 - 1) **threshold of withdrawal reflex is especially low** (minor noxious stimuli → prolonged extremity withdrawal + marked flexion-extension patterns in other three limbs).
 - 2) **withdrawal reflex generalization** may cause **mass reflex** (bladder and rectum evacuation, sweating, piloerection, pallor, BP swings).
 - **mass reflex** can be used to give paraplegic patients degree of bladder and bowel control (initiate urination and defecation by stroking or pinching thighs - **intentional mass reflex**).
- **hyperactive stretch reflexes** can cause **magnet reaction (positive supporting reaction)** (at least in spinal animals). see p. A61 >>
- in incomplete spinal cord transections, **spinal locomotion generators** can be turned on by tonic discharge of discrete area in midbrain (**mesencephalic locomotor region**) → spinal patient can be made to *stand*, and even to produce *walking movements* (e.g. on treadmill). see p. A61 >>
- genital manipulation in spinal male produces **erection** and even **ejaculation**; in spinal female dogs, vaginal stimulation causes tail deviation and movement of pelvis into copulatory position.
- bladder becomes **automatic spastic**; about **bladder and bowel dysfunction** – see below (SPINAL COMPLICATIONS).

See p. TrS5 >> for American Spinal Injury Association (ASIA) system for examination and classification of spinal cord injury

TREATMENT (huge doses of glucocorticoids, etc) → see p. TrS5 >>

SPINAL CORD HEMISECTION

BROWN-SEQUARD SYNDROME

ETIOLOGY

- 1) traumatic hemisections (e.g. stab wound, lateral mass fracture in cervical spine)
- 2) extramedullary tumors
- 3) extramedullary abscesses
- 4) vasculitis (as in SLE).

CLINICAL FEATURES

- caudal to hemisection:

I. **Contralateral effects** – loss of pain-temperature sensation (tr. spinothalamicus).

N.B. sensory level is located 1-2 segments below level of lesion!!!

II. **Ipsilateral effects**:

- 1) **UMN paralysis** (tr. corticospinalis lat.);
 - if high cervical – hemidiaphragm paralysis.
- 2) **loss of discriminative touch-proprioception** (dorsal funiculus)
- 3) **loss of sweating** (descending autonomic fibers in ventral funiculus)
 - if high cervical – Horner syndrome.
- 4) **SEGMENTAL*** – **anesthesia / radicular pain** (dorsal root), **LMN paralysis** (ventral horn)

*i.e. hemisection segment

N.B. bowel and bladder control is usually intact!

DORSAL (POSTERIOR) HEMISECTION

- 1) **dorsal funiculus** – loss of vibration and position sense.
- 2) **tr. corticospinalis lat.** – paralysis.

VENTRAL (ANTERIOR) HEMISECTION

- 1) **tr. spinothalamicus** – loss of pain & temperature sense, loss of urge to urinate.
- 2) **tr. reticulospinalis** – anhidrosis, vasodilation-hypotension, loss of voluntary* bladder-bowel control; if rostral to C₃ – paralysis of automatic breathing.

*reflex emptying intact

ANTERIOR 2/3 TRANSECTION

- **anterior spinal artery** occlusion (supplies whole spinal cord, except dorsal funiculi).

Clinically – VENTRAL HEMISECTION + **paralysis**.

EXTRAMEDULLARY CORD COMPRESSION

ETIOLOGY

1. Spinal or epidural **abscess / hematoma**
2. **Tumor** (85% - vertebral metastases) - may present acutely even though tumor has been present for weeks or longer.
3. Epidural **granuloma** (e.g. neurocysticercus).
4. Cervical or thoracic **herniated intervertebral disk** (central herniation may cause acute compression without local pain).
5. Trauma
6. Atlantoaxial subluxation.

CLINICAL FEATURES

SEGMENTAL features - most reliable indication of lesion level (longitudinal location)!

- 1) **LMN paralysis** (ventral horn)
- 2) **anesthesia / prominent radicular pain** (dorsal root)

vs. intramedullary lesions - tend to produce poorly localized burning pain rather than radicular pain

- other strongly localizing symptoms – **local back pain, tenderness over spine** (N.B. some lesions are painless!).
- radicular pain may be exacerbated by Valsalva maneuvers, straight-leg raising test.
- site of compression in *transverse plane* may determine clinical symptoms (e.g. laterally located lesion → Brown-Sequard syndrome).

N.B. because most lesions twist cord and also interfere with vascular supply to sites beyond compression, neurological signs may not demarcate exact transverse site!

- certain spinal tracts are more vulnerable to compression than others:
corticospinal tracts > posterior column > spinothalamic & descending autonomic fibers.

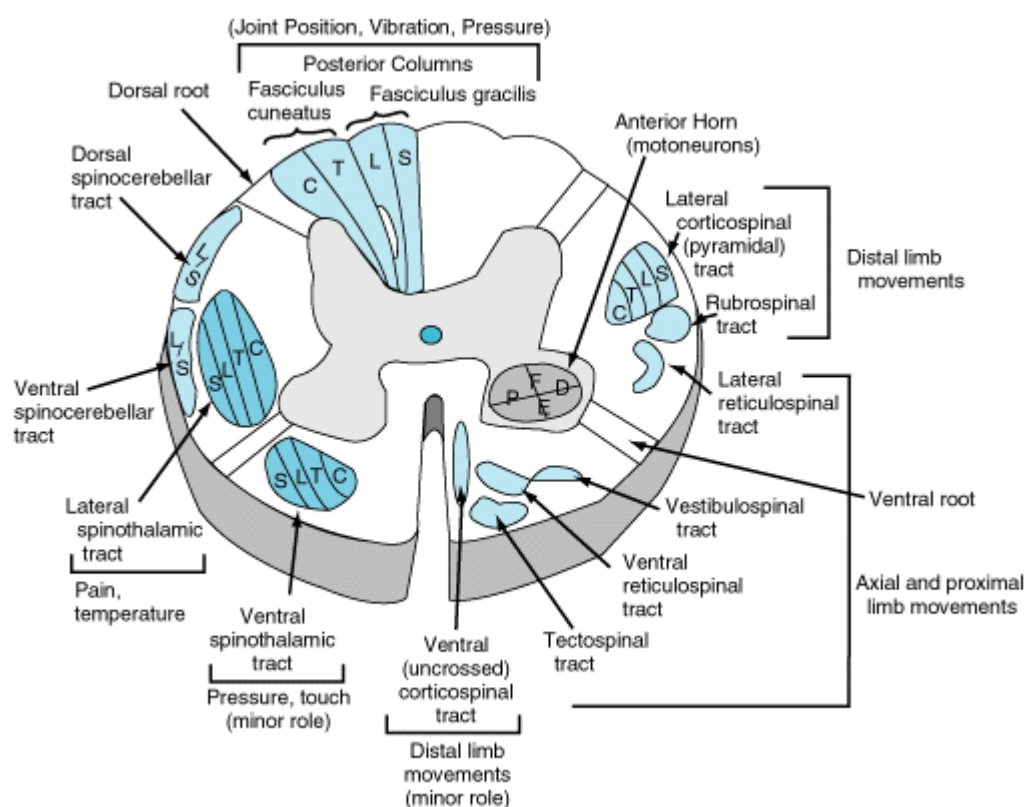
Earliest manifestations in **lower body parts** – due to **Flatau law** (superficial location of lumbosacral fibers in lateral spinal cord - susceptible to external compression):

- 1) early **sacral** sensory loss (**tr. spinothalamicus**) - EXTRAMEDULLARY lesions cause **ascending pain & temperature loss**
- 2) early spastic weakness in **legs** (**tr. corticospinalis lat.**).

vs. INTRAMEDULLARY lesions – descending pain & temperature loss with long spare of perineal-sacral sensation; corticospinal signs may appear late.

- 3) urinary retention (**tr. reticulospinalis**).
- 4) gait ataxia (**tr. spinocerebellaris**).

FLATAU law – **topographic fiber lamination** – greater distance nerve fibers (of long tracts) run lengthwise in cord, more they tend to be situated toward its periphery.



Source of picture: William F. Ganong "LANGE Review of Medical Physiology", 21st ed. (2003); McGraw-Hill / Appleton & Lange; ISBN-13: 978-0071402361 >>

- lesion is above highest dermatome involved in deficit (radiographic studies should be tailored to visualize cord at and above level of sensory deficit).
- distinction between EXTRADURAL (generally malignant) and INTRADURAL (generally benign) masses is important; long duration of symptoms favors intradural origin.

DIAGNOSIS

Proper treatment requires expeditious diagnosis! - therapy will not reverse fixed paralysis of > 48 h duration (acute spinal cord compression is neurologic emergency!)

Neuroimaging (MRI is method of choice)

- **acute postmyelography decompression** may occur with compressive lesions → emergency decompressive laminectomy.

Lumbar puncture is contraindicated in compressive lesion!

TREATMENT

Spinal cord compression is emergency! see p. Onc56 >>

- 1) **steroids** – give immediately!
- 2) **antibiotics** (if indicated).
- 3) immediate **radiotherapy** (for cord compression due to malignancy).
- 4) **surgical decompression** (where radiotherapy is not effective)

CENTRAL CORD SYNDROME

- pathological process **starts centrally and proceeds centrifugally** → characteristically evolving motor and sensory signs.

ETIOLOGY

1. **Syringomyelia**
2. **Intramedullary cord tumors** (esp. central canal ependymoma)
3. **AVM**
4. **Anterior spinal artery ischemia.**
5. **Spinal cord trauma:** see p. TrS5 >>
 - a) **neck hyperextension** in presence of narrow spinal canal → cord compression between bony bars anteriorly and thickened ligamentum flavum posteriorly → cord hypoperfusion in central watershed distribution.
 - b) **hematomyelia** (usually confined to central gray matter)

CLINICAL FEATURES

Characteristic initial presentation - combination of **SEGMENTAL** (at level of lesion) features:

1. **Loss of pain and temperature sensation** – due to lesion to central cord portion where **spinothalamic fibers decussate**.
 - because only decussating spinothalamic tract fibers are affected, loss of pain and temperature is **bilateral** but affects only those **segments** of spinal cord involved in pathological process (**suspended sensory loss** with normal sensation above and below lesion).
 - may produce poorly localized **burning pain**.
vs. extramedullary cord compression – radicular pain
 - **posterior column** sensation is preserved (**disassociated sensory loss**).
2. **LMN signs** (SYRINGOMYELIA or TUMOR usually invade **anterior horns** early);
 - in SYRINGOMYELIA, segmental pattern characteristically begins in upper cervical segments (**distal arms** suffer first!).
 - in CERVICAL TRAUMA, initial quadriplegia is replaced over minutes by leg recovery.

If lesion expands centrifugally, it may compromise other spinal structures:

- 1) **lateral corticospinal tracts** - late involvement!

vs. extramedullary cord compression – early, with legs affected first

- 2) **ascending** (vs. decussating) **spinothalamic tract** fibers
N.B. because spinothalamic tracts are topographically laminated (**FLATAU law** - sacral fibers in most ventral-lateral position), sacral dermatomes are long preserved (**sacral sparing**) – INTRAMEDULLARY lesions cause **descending loss of pain and temperature sensation**.

vs. EXTRAMEDULLARY cord compression - **ascending loss of pain and temperature sensation** with early **sacral** involvement

- 3) **posterior columns**
- 4) **intermediolateral columns** → autonomic manifestations (Horner's syndrome, sudomotor and vasomotor dysfunction, trophic changes [esp. hands]).

CAUDA EQUINA vs. CONUS MEDULLARIS syndrome

CONUS MEDULLARIS – tapered caudal termination of spinal cord (*lower sacral & coccygeal* segments).

CAUDA EQUINA – collection of intradural elongated roots of *lumbar & sacral* spinal nerves.

Feature	CAUDA EQUINA	CONUS MEDULLARIS
Pain	Severe radicular pain (sciatica) & low back pain	Back pain (less severe than radicular pain)
Sensory loss	Asymmetric saddle anesthesia* – all modalities (radicular sensory loss)	Bilateral saddle anesthesia* (usually restricted to perianal region) – all modalities or touch preservation.
Motor deficits	Asymmetrical areflexic para- / monoplegia	Absent!!! (or mild distal leg paresis)
Evacuation disorder	Late and mild – hypotonic bladder (urinary retention)**	Early - atonic bladder (urinary retention with overflow incontinence), atonic anal sphincter (constipation with incontinence)
Impotence	±	+
Bulbocavernosus (S ₂₋₄) & anal wink (S ₄₋₅) reflexes	+	ABSENT

***saddle anesthesia** - sensory loss confined to S₃₋₅ dermatomes.

**may be limited to asymptomatic bladder retention noted only on postvoid catheterization (> 100 mL)

Patient may not feel urge to urinate! (ask every patient with back pain about difficulty with urination and defecation)

Nerve roots in cauda equina:

- **poorly developed epineurium** - particularly susceptible to injury (in peripheral nerves well developed epineurium protects against compressive and tensile stresses).
- **relative hypovascularity** in proximal third of root (nutritional supply is supplemented with increased vascular permeability* and diffusion from surrounding CSF).
*may result in edema compounding initial and sometimes seemingly slight injury.

Causes of CAUDA EQUINA syndrome:

- 1) tumor
- 2) abscess
- 3) lumbar spinal stenosis
- 4) lumbar disk disease
- 5) arachnoiditis
- 6) spinal anesthesia
- 7) trauma.

MRI is criterion standard for initial evaluation.

TREATMENT - directed at underlying cause.

- in *acute* or *traumatic* syndrome, some suggest **METHYLPREDNISOLONE** (similar to traumatic spinal cord injury); steroids have not shown significant benefit in penetrating trauma.
- **surgical decompression**, e.g. lumbar laminectomy (timing is controversial - immediate, early, and late surgery shows varying results).
cauda equina injuries (involving peripheral nerves rather than spinal cord) are surgically remediable for longer periods than conus medullaris injuries

SPINAL COMPLICATIONS

- higher anatomical level of injury, greater risk of complications.
- **usual symptoms** associated with medical illnesses **may be lacking** (because of *destruction of afferent pain pathways*).
- **unexplained fever, spasticity worsening, neurologic function deterioration** should prompt search for underlying cause (infection, thrombophlebitis, intraabdominal pathology).
N.B. *loss of normal thermoregulation* can produce recurrent fever (**quadruplegic fever**)!

PARESTHESIAS

- **burning / shooting pains** below level of spinal cord lesion.

- causes:
 - a) selective deafferentation of spinothalamic pathway with preservation of function of dorsal columns.
 - b) abnormal discharge of thalamic neurons.
 - c) another occult lesion in conus or cauda equina.
- treat as neuropathic pain; avoid narcotics (bowel and bladder adverse effects).

DECUBITUS ULCERS

see p. 2217 >>

- predisposed by *immobility* and *lack of sensation*!

BLADDER DYSFUNCTION

see p. 2590a >>

SPINAL SHOCK – **atonic bladder**; H: intermittent catheterization to prevent urinary retention → permanent bladder atony.

CHRONIC STAGE – bladder dysfunction depends on level of lesion:

- a) **lesions above sacral parasympathetic nucleus** – within several days of injury, automatic **spastic bladder with detrusor-sphincter dyssynergia** develops (bladder re-education should begin promptly!);

b) *lesions of conus medullaris or cauda equina* – **atonic bladder**.

- bone matrix protein breakdown + immobilization → osteoporosis, large Ca^{2+} release → hypercalcemia and hypercalciuria → **urinary calcium stones** (H: urine acidification).
- stones + **dysfunctional bladder** → urinary stasis → **UTI** (most common complication of spinal cord injury!!!), **hydronephrosis**, **autonomic dysreflexia**.

GU tract is primary source of infection after cord trauma!

- prophylactic antibiotics are not indicated.

GI COMPLICATIONS

- **paralytic ileus** almost universally occurs after cord trauma; H: nasogastric suctioning.
 - in few days, small bowel function returns to normal, but large bowel and rectal function may be lost permanently.
- *for several weeks* after acute spinal injury (anal sphincter is atonic) **laxatives** and **digital disimpaction** are necessary in most patients to ensure at least biweekly evacuation;
 - **GLYCERIN** suppositories are useful (insert \approx 20 min before desired time of evacuation); also stool softeners (e.g. **DOCUSATE**), stool bulking agents (e.g. **PSYLLIUM**).
 - avoid anus stretching!
 - **flatus tube** may be helpful.
- *later*, start training for **REGULAR DEFECATION** - **GLYCERIN** suppositories on alternate days.
 - Both bowel and bladder sphincter reflexes can be trained to provide reflex emptying if lesions spare lower motor neurons.*
- consider **prophylaxis for GI stress ulcers**.

SEXUAL DYSFUNCTION

1. Mechanical and pharmacologic interventions
2. Psychosocial counseling

Men:

- in men, **priapism** is seen early (esp. after high cord lesions) → reflex but no psychogenic erection.
- **semen** quality and motility is reduced because of repeated UTIs.

Women:

- paraplegia and tetraplegia result in **menstrual cycle interruption** for months, but this returns with time - **conception** and **pregnancy** are possible.
- women may experience life-threatening **autonomic hyperreflexia** during delivery.

MALNUTRITION

- **anorexia** → early loss of weight occurs in many spinal patients.
- patients (like all immobilized patients) *catabolize large amounts of body protein* → develop **negative nitrogen balance**.
- protein may be lost through **bedsores**.
- prophylaxis / treatment - diet high in protein, calories, and vitamins (incl. parenteral hyperalimentation).
- **calcium & vitamin D supplementation** - to avoid **osteoporosis**.

RESPIRATORY FAILURE

- respiratory failure is caused by:
 - a) **neurological compromise**
 - b) **pain**
 - c) **retropharyngeal hematoma** (from cervical trauma)
- **acute pulmonary edema** has occurred after cervical spine injuries unassociated with significant head injury.
- respiratory failure is exacerbated by **CNS depressants**, **immobilization in recumbency**, **abdominal distention** (from paralytic ileus).
- atelectasis → pneumonia.
- check at regular intervals - vital capacity, arterial blood gases / pulse oximetry.
- for **cervical cord lesions**:
 - 1) **artificial ventilation** (tracheal intubation → tracheostomy)
 - 2) **phrenic nerve pacemakers** - for lesions at C₅ or above.
 - NeuRx DPS RA/4 Respiratory Stimulation System (FDA approved) - implantable electronic device that stimulates diaphragm - allows to breathe for at least 4 hours a day without a mechanical ventilator.
 - 3) chest **physical therapy**
 - 4) **negative-pressure cuirass** (to alleviate atelectasis, particularly if lesion is below C₄).

N.B. in lesions above T₁₀, there is no effective coughing!

H: regular nasotracheal suctioning, chest physiotherapy, use of rotating beds or frames

VENOUS THROMBOSIS & PULMONARY EMBOLISM

- high risk in **acute cord injury**.
 - 1) **calf-compression devices** (for first two weeks)
 - 2) **anticoagulation**: **HEPARIN** (5000 U SC every 12 h) → **WARFARIN** (INR 2-3) for 3 months in persistent paralysis.

SPASTICITY

- major late complication of spinal cord disease (weeks ÷ months after initial insult).
- most severe spasticity - incomplete traumatic injury, multiple sclerosis.
- if lesion involves upper cervical cord, spasms may involve all four extremities, trunk, and bladder.
- spasms of extremities are usually **flexor** (but may also be extensor).
- severe spasticity may lead **contractures**.
- treatment (if spasms are painful, interfere with rehabilitation, or delay healing of bedsores)
 - see p. Mov3 >>

AUTONOMIC DYSFUNCTION

- descending pathways from brain normally coordinate sympathetic activity and modulate segmental autonomic reflexes; spinal cord transection may be attended by **autonomic hyperreflexia** (affecting bowel, bladder, sexual, temperature-regulation, and cardiovascular functions).
- **BLOOD PRESSURE is generally normal at rest**, but precise feedback regulation normally supplied by baroreceptor reflexes is absent:

- **wide swings in BP are common** (quadriparetic patients exhibit both orthostatic hypotension and supine hypertension after upward tilting).
- **vasopressin & renin-angiotensin-aldosterone system** have enhanced role in maintenance of orthostatic arterial pressure.
- patients are at risk of **bradycardia & cardiac arrest** during **tracheal suction** (or other maneuvers that activate **vagovagal reflexes**).
- inability to sense heat or cold exposure below level of injury → **dangerous increases / decreases in body temperature**.

PAROXYSMAL AUTONOMIC HYPERREFLEXIA (S. AUTONOMIC DYSREFLEXIA)

- in lesions above major splanchnic sympathetic outflow (i.e. lesions above T5-6; e.g. affects 85% patients with lesion above C6).
- **trigger - noxious stimulus below level of cord lesion** (e.g. fecal impaction, bladder distention, catheter insertion, UTI, decubitus ulcer).
- sensory inputs activate sympathetic neurons of intermediolateral nuclei in thoracic spinal cord → **massive reflex activation of sympathetic outflow below lesion** → **vasoconstriction** (below level of lesion), **tachycardia, systemic hypertension** (up to 300 mmHg!!!*)
*may lead to life-threatening hypertensive encephalopathy, stroke, retinal hemorrhage!
- reflex pathways (via carotid and aortic baroreceptors) then **inhibit sympathetic activity above cord lesion** → **vasodilation** (**flushing, nasopharyngeal congestion, headache**), **diaphoresis** above level of lesion, **bradycardia**.
- N.B. descending pathways are blocked - sympathetic hyperactivity below lesion continues.
- **prophylaxis-treatment:**
 - 1) removal of offending stimuli.
 - 2) BP can often be lowered by **tilting head upward**.
 - 3) **ganglionic blockers** (**MECAMYLAMINE**, 2.5-5 mg)
 - 4) short-acting **antihypertensives** (e.g. **CLONIDINE** prophylactically to reduce hypertension resulting from bladder stimulation; **NIFEDIPINE**).

AUTONOMIC HYPERREFLEXIA in addition to **SOMATIC HYPERREFLEXIA (SPASTICITY)** may lead to accumulation of contractures, bladder, bowel, and skin disorders, which eventually cause severe wasting and death!

PSYCHIATRIC DYSFUNCTION

- **depression** (following initial period of denial) occurs in almost all patients and may be masked by jocularity.
- **suicide rate** is 5 times higher than in general population (lower for men; 2 times higher in marginally disabled persons compared to more severely affected individuals).
- **narcotic addiction** is also occasionally problem.

SPINAL PROGNOSIS

No effective means to promote repair of injured spinal cord tissue!

- if total loss of motor power & sensation distal to level (feature of complete transection) persist for > 24 hours* - 99% will not have functional recovery.
*ensure that **spinal shock** is not present and **sacral sparing** is carefully excluded
- after **acute spinal cord lesion**, prospects for significant recovery fade after ≈ **4 months** (recovery plateaus between 6 and 12 months);
 - many patients even after complete spinal cord injuries, regain 1-2 levels (or some key muscles) after > 1 year – esp. important in high cervical lesions!
- prognosis in **TRANSECTED SPINAL CORDS** used to be very poor (**LIFE EXPECTANCY** is greatly decreased);
in past, renal failure was leading cause of death after spinal cord trauma.
currently, pulmonary problems (pneumonia, pulmonary emboli, sepsis) are single most common cause of morbidity and mortality after spinal cord trauma.
 - antibiotics and meticulous attention to nutrition, fluid balance, skin care, bladder function, and general nursing care have reduced **mortality to 6%**.
- **SPORT aspects:**
 - any injury that necessitates internal surgical spinal stabilization obviates return to contact sports.
 - minor injuries that heal correctly with bracing may not limit athletic involvement.

Disability and survival associated with spinal cord damage are determined by:

- 1) level of lesion
- 2) completeness of transection
- 3) age (prognosis is better with **younger age**).

Expected Neurologic Function Following Complete Cord Lesions:

Level	Self-Care	Transfers	Maximum Mobility
High quadriplegia (C1-4)	Dependent on others; requires respiratory support (e.g. implantation of diaphragmatic stimulators)	Dependent on others	Motorized wheelchair
Low quadriplegia (C5-8)	Partially independent with adaptive equipment	May be dependent or independent	May use manual wheelchair, drive automobile with adaptive equipment
Paraplegia (below T1)	Independent	Independent	Ambulates short distances with aids

Even complete high cervical cord lesions may be compatible with productive life!

SPINAL REHABILITATION

- best carried out in experienced **spinal centers**.
- best if single physician organizes long-term approach.
- start early (once spine stabilization has been achieved).
 - early range of motion prevents contractures, diminishes risk of venous thrombosis, protects skin, and boosts morale.
 - bed should be fitted with **footboards** to keep ankles and toes in neutral position.
 - **soft braces** to fix lower extremities in neutral position.
 - exercises to **strengthen unaffected muscles**.
 - gradual progression toward **vertical position** (simultaneous monitoring of systemic BP - horizontal position for prolonged period results in sympathetic tone loss)
- major focus of rehabilitation:
 - 1) bowel management

- 2) bladder management
- 3) transfer techniques
- ultimate aim - AMBULATION & ECONOMIC INDEPENDENCE.
- **psychological support** throughout disease course is necessary (severe depression can occur after losing control of body).
- **special adaptive devices** may allow patients to drive.
- recently, role of *CENTRAL PATTERN GENERATORS* and possibility of activating standing and stepping circuits after SCI even in chronic injury phase has been addressed;
 - in 1914, Graham Brown demonstrated existence of central pattern generators for walking in animals (neuronal networks capable of creating rhythmic motor activity in absence of phasic sensory input).
 - theoretically, similar system exists in humans and can be activated by repeated exercise or stimulation of walking pathways; exercise programs have been developed (incl. suspended body weight support system over treadmill to facilitate walking and bicycles designed for SCI).

Spinal Cord Independence Measure (SCIM) - only comprehensive functional rating scale designed specifically for patients with spinal cord injury

- development of SCIM started in 1994.
- third international version (SCIM III) intended to overcome intercultural differences was formulated in 2002

SCIM - SPINAL CORD INDEPENDENCE MEASURE

Loewenstein Rehabilitation Hospital, Department IV

(Version 1, May 1996, Raanana, Israel)

Patient Name: ID: Examiner Name:

(The score attached to the relevant description of each function should be placed in the adjacent square below the relevant date)

Self-Care

DATE

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1. **Feeding** (cutting, opening containers, bringing food to mouth, holding cup with fluid)
- 0. Needs parenteral, gastrostomy or fully assisted oral feeding
 - 1. Eats cut food using several adaptive devices for hand and dishes
 - 2. Eats cut food using only one adaptive device for hand; unable to hold cup
 - 3. Eats cut food with one adaptive device; holds cup
 - 4. Eats cut food without adaptive devices; needs a little assistance (e.g., to open containers)
 - 5. Independent in all tasks without any adaptive device

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2. **Bathing** (soaping, manipulating water tap, washing)
- 0. Requires total assistance
 - 1. Soaps only small part of body with or without adaptive devices
 - 2. Soaps with adaptive devices; cannot reach distant parts of the body or cannot operate a tap
 - 3. Soaps without adaptive devices; needs a little assistance to reach distant parts of body
 - 4. Washes independently with adaptive devices or in specific environmental setting
 - 5. Washes independently without adaptive devices

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3. **Dressing** (preparing clothes, dressing upper and lower body, undressing)
- 0. Requires total assistance
 - 1. Dresses upper body partially (e.g., without buttoning) in special setting (e.g., back support)
 - 2. Independent in dressing and undressing upper body. Needs much assistance for lower body
 - 3. Requires little assistance in dressing upper or lower body
 - 4. Dresses and undresses independently, but requires adaptive devices and/or special setting
 - 5. Dresses and undresses independently, without adaptive devices

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4. **Grooming** (washing hands and face, brushing teeth, combing hair, shaving, applying makeup)
- 0. Requires total assistance
 - 1. Performs only one task (e.g., washing hands and face)
 - 2. Performs some tasks using adaptive devices; needs help to put on/take off devices
 - 3. Performs some tasks using adaptive devices; puts on/takes off devices independently
 - 4. Performs all tasks with adaptive devices or most tasks without devices
 - 5. Independent in all tasks without adaptive devices

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Respiration and Sphincter Management

DATE

5. **Respiration**
- 0. Requires assisted ventilation
 - 2. Requires tracheal tube and partially assisted ventilation
 - 4. Breathes independently but requires much assistance in tracheal tube management
 - 6. Breathes independently and requires little assistance in tracheal tube management
 - 8. Breathes without tracheal tube, but sometimes requires mechanical assistance for breathing
 - 10. Breathes independently without any device

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6. **Sphincter management - Bladder**
- 0. Indwelling catheter
 - 5. Assisted intermittent catheterization or no catheterization, residual urine volume > 100cc
 - 10. Intermittent self-catheterization
 - 15. No catheterization required, residual urine volume < 100cc

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7. **Sphincter management - Bowel**
- 0. Irregularity, improper timing or very low frequency (less than once in 3 days) of bowel movements
 - 5. Regular bowel movements, with proper timing, but with assistance (e.g., for applying suppository)
 - 10. Regular bowel movements, with proper timing, without assistance

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8. **Use of toilet** (perineal hygiene, clothes adjustment before/after, use of napkins or diapers)
- 0. Requires total assistance
 - 1. Undresses lower body, needs assistance in all the remaining tasks
 - 2. Undresses lower body and partially cleans self (after); needs assistance in adjusting clothes and/or diapers
 - 3. Undresses and cleans self (after); needs assistance in adjusting clothes and/or diapers
 - 4. Independent in all tasks but needs adaptive devices or special setting (e.g., grab-bars)
 - 5. Independent without adaptive devices or special setting

Mobility (room and toilet)

9. Mobility in bed and action to prevent pressure sores

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- 0. Requires total assistance
- 1. Partial mobility (turns in bed to one side only)
- 2. Turns to both sides in bed but does not fully release pressure
- 3. Releases pressure when lying only
- 4. Turns in bed and sits up without assistance
- 5. Independent in bed mobility; performs push-ups in sitting position without full body elevation
- 6. Performs push-ups in sitting position

10. Transfers: bed-wheelchair (locking wheelchair, lifting footrests, removing and adjusting arm rests, transferring, lifting feet)

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- 0. Requires total assistance
- 1. Needs partial assistance and/or supervision
- 2. Independent

11. Transfers: wheelchair-toilet-tub (if uses toilet wheelchair - transfers to and from; if uses regular wheelchair - locking wheelchair, lifting footrests, removing and adjusting arm rests, transferring, lifting feet)

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- 0. Requires total assistance
- 1. Needs partial assistance and/or supervision, or adaptive device (e.g., grab-bars)
- 2. Independent

Mobility (indoors and outdoors)

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12. Mobility indoors (short distances)

- 0. Requires total assistance
- 1. Needs electric wheelchair or partial assistance to operate manual wheelchair
- 2. Moves independently in manual wheelchair
- 3. Walks with a walking frame
- 4. Walks with crutches
- 5. Walks with two canes
- 6. Walks with one cane
- 7. Needs leg orthosis only
- 8. Walks without aids

13. Mobility for moderate distances (10 - 100 meters)

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- 0. Requires total assistance
- 1. Needs electric wheelchair or partial assistance to operate manual wheelchair
- 2. Moves independently in manual wheelchair
- 3. Walks with a walking frame
- 4. Walks with crutches
- 5. Walks with two canes
- 6. Walks with one cane
- 7. Needs leg orthosis only
- 8. Walks without aids

14. Mobility outdoors (more than 100 meters)

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- 0. Requires total assistance
- 1. Needs electric wheelchair or partial assistance to operate manual wheelchair
- 2. Moves independently in manual wheelchair
- 3. Walks with a walking frame
- 4. Walks with crutches
- 5. Walks with two canes
- 6. Walks with one cane
- 7. Needs leg orthosis only
- 8. Walks without aids

15. Stair management

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- 0. Unable to climb or descend stairs
- 1. Climbs 1 or 2 steps only, in a training setup
- 2. Climbs and descends at least 3 steps with support or supervision of another person
- 3. Climbs and descends at least 3 steps with support of handrail and/or crutch and/or cane
- 4. Climbs and descends at least 3 steps without any support or supervision

16. Transfers: wheelchair-car (approaching car, locking wheelchair, removing arm and foot rests, transferring to and from car, bringing wheelchair into and out of car)

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- 0. Requires total assistance
- 1. Needs partial assistance and/or supervision, and/or adaptive devices
- 2. Independent without adaptive devices

BIBLIOGRAPHY for ch. "Spinal Disorders" → follow this [LINK >>](#)