Advanced Quantitative Spine Imaging

James F. Griffith, MB, BCh, BAO, MD (CUHK), MRCP (UK), FRCP (Edin), FRCR, FHKCR, FHKAM (Radiology)¹⁰

¹Department of Imaging and Interventional Radiology, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong

Semin Musculoskelet Radiol 2020;24:413–427.

Address for correspondence James F. Griffith, MB, BCh, BAO, MD (CUHK), MRCP (UK), FRCP (Edin), FRCR, FHKCR, FHKAM (Radiology), LG12 Cancer Centre, Prince of Wales Hospital, Shatin, NT, Hong Kong (e-mail: griffith@cuhk.edu.hk).

Abstract	Although advanced quantitative imaging may not be currently used to any degree in
	the routine reporting of spinal examinations, this situation will change in the not too
	distant future. Advanced quantitative imaging has already allowed us to understand a
	great deal more regarding spinal development, marrow physiology, and disease
Keywords	pathogenesis. Radiologists are ideally suited to drive this research forward. To speed
► spine	up this process and optimize the impact of studies reporting spine quantitative data,
► imaging	we should work toward universal standards on the acquisition of spine data that will
 guantification 	allow guantitative studies to be more easily compared, contrasted, and amalgamated.

The spine is the part of the musculoskeletal system most conducive to, and most likely to benefit from, quantitative imaging. Computed tomography (CT) and, more particularly, magnetic resonance imaging (MRI) allow bone and soft tissues to be evaluated free from the superimposition, magnification, and obliquity errors of radiography. This, and new imaging interrogation techniques, has enabled a profusion of new spine imaging data to be realized. This review looks broadly at the current issues, reported data, and potential benefits of advanced quantitative spine imaging, specifically addressing those areas considered to be of highest clinical relevance.

Uses of Quantitative Spine Imaging

Quantitative imaging data can potentially be used in a variety of ways.¹ On an individual patient basis, quantitative spine imaging data can be used in these arenas:

- Diagnosis by having a normal or reference population reference interval and comparing suspicious imaging features with these reference intervals, for example to diagnose developmental canal stenosis or cord atrophy.
- Stratifying disease severity according to predefined reference ranges as in stratifying bone loss by quantitative computed tomography (QCT) or stratifying sarcopenia or the degree of disk degeneration. Providing the patient with easily appreciable objective data as to, for example,

their degree of disk degeneration or muscle mass relative to a matched population cohort.

- *Monitoring disease progress* or *response to therapy* as in, for example, new treatments for spinal cord injury.
- *Predicting disease progress* in the absence of intervention as in, for example, the likelihood of progressive canal stenosis following input of multiple quantitative data into a predictive model.
- *Predicting outcome* following intervention as in patients who have spinal cord decompression.
- *Predicting risk* in, for example, the risk of developing spinal cord injury if engaging in high-energy contact sports.

On a larger scale, quantifiable image data can be used in these situations:

- Define reliable normal and population reference intervals for relevant covariables such as sex, height, and age.
- Use investigative studies and big data research to define disease patterns, compare different populations, and provide valuable data on disease pathogenesis including early detection. This seems to be the main benefit of quantitative spine imaging.

When defining reference intervals for parameters such as developmental spinal canal stenosis, we should appreciate the difference between a *normal sample* and a *population reference interval*.² For a disease with progressive deterioration such as disk degeneration or bone density, reference

Issue Theme Advanced Quantitative MSK Imaging; Guest Editors, Eric Y. Chang, MD and Edwin H.G. Oei, MD, PhD Copyright © 2020 by Thieme Medical Publishers, Inc., 333 Seventh Avenue, New York, NY 10001, USA. Tel: +1(212) 760-0888. DOI https://doi.org/ 10.1055/s-0040-1713608. ISSN 1089-7860. limits are usually obtained in normal young skeletally mature individuals, usually aged 20 to 30 years. "Abnormality" is then rather arbitrarily defined as a deviation > 1 standard deviation (SD) or > 2 SDs from this young Gaussian-distributed population mean as, for example, for the bone mineral density (BMD) T-score. Alternatively, higher than the 90th percentile or > 97% cutoffs can be used.³ The symptomatic and asymptomatic matched populations can be compared to help validate these cutoff values.

For diseases in which there is no known deterioration over time, such as developmental spinal canal size or facet joint orientation, a large cross section of the population of any age can be referenced before determining optimal arbitrary cutoff values. To include only asymptomatic subjects in this population, reference interval seems incorrect because this would, for example, if determining developmental spinal canal size, tend to favor patients with a developmentally normal or capacious spinal canal while conversely including only symptomatic patients would favor patients with a developmental spinal canal stenosis.

Main Issues Limiting the More Widespread Use of Quantitative Spine Imaging

Currently, three overriding issues limit the more widespread use of quantitative spine image data: data reliability, time utilization, and clinical relevance.

Data Reliability

The objective data provided should be precise, reproducible, and clinically relevant.¹ One of the frustrating features currently of quantitative spine imaging is the variability of data reported by different, seemingly rigorously conducted studies. For example, comparing studies reporting cervical cord cross-sectional area (CSA), two studies on asymptomatic European and Japanese subjects, respectively, reported values of ~ 72 to 79 mm² for cervical cord CSA.^{4,5} Another study on asymptomatic Canadians reported values of ~ 93 to 97 mm².⁶

Because objective analysis of imaging-based research databases is normally undertaken by supervised research assistants, close supervision on how (e.g., slice selection, digital zooming, contrast adjustment) and where to obtain measures is critical. This is particularly pertinent for boundary demarcation in CSA measurements as, for example, whether spinal cord measurements are made at the outermost margin of the spinal cord or just outside the cord. Given πr^2 , if the radius changes by $r \pm x$, then CSA will change by $(r \pm x)$.² Because the structures involved are often small, important before data point placement would seem helpful. Even after thorough training, though, readers still tend to produce different readings because there is a tendency to draw lines between gray levels of anatomical structures differently.⁴ Also, small but statistically significant differences can exist between different MR imaging systems due to factors such as corrections of geographic distortion or systematic different placement of the patient.⁴

Given the level of high-resolution imaging data now achievable on all standard body CT and MR systems, the main reason for interstudy variability seems to be differences in measurement techniques such as using sagittal versus axial planes for anteroposterior (AP) diameters, true axial versus oblique axial planes, and variable placement of measurement points. Scientific publications showing precisely where measurements points were placed are very helpful, although articles often do not include such images.⁷ Ideally, like guidelines available on who should acquire dual-energy X-ray absorptiometry (DXA) or QCT measurements, internationally accepted standardized protocols should be devised for the acquisition and measurement of advanced spine imaging quantitative data.⁸ These guidelines could then be adopted into protocols of studies involving acquisition of quantitative spine imaging data. Until this is done, considerable interstudy variability is likely to persist.

Time Utilization

For quantitative imaging data to be integrated into clinical practice, such data need to be obtained in an automated fashion because manual measurements are too time consuming for everyday use. Objective analytical data will someday be produced automatically by computer analysis of spine imaging raw data and be available for review when constructing the standard radiologic report. There is little doubt this will be available in the future with a digital readout of quantifiable data referenced to population or normal intervals like serology laboratory reports. That said, in our experience, current automated image segmentation/quantification techniques in the spine are not up to speed, requiring considerable adjustment following initial automatic segmentation.

Clinical Relevance

Currently, little objective data are used in the day-to-day radiologic reporting of spine examinations because it is time consuming to obtain and of limited clinical relevance in the individual patient setting. Most of the measured variables can be readily reported in daily practice using semiquantitative descriptors such as minimal, mild, moderate, or severe or mild to moderate, for example. There is no doubt, however, that image quantification of the spine will become more widespread, more reliable, and more automated in the not too distant future. Radiologic reports will continue to be composed mainly of descriptive terminology as they are now with quantitative data providing additional backup information.

The remainder of this review looks at 13 specific aspects of quantitative spine imaging focusing mainly on age-related physiologic changes rather than disease-related changes.

Developmental Cervical Spinal Canal Size

To date, no disadvantage has been found in having a large spinal canal. All problems arise when the canal is too small to accommodate the spinal contents (**~Fig. 1**). There are two components to spinal canal stenosis. The first is how large the spinal canal is when it is fully developed (i.e., developmental spinal canal size).⁹ Developmental canal narrowing is measured at the pedicle level removed from acquired degenerative narrowing that occurs at the diskovertebral level. The second is the degree of acquired spinal canal stenosis that occurs due to superimposed narrowing from, for example, disk herniation,



Fig. 1 (a) T2-weighted sagittal and (b) axial images of patient with developmentally normal caliber cervical spine compared with (c) T2-weighted sagittal and (d) axial images of a patient with a developmentally small caliber cervical spinal canal. The patient with a developmentally small caliber cervical spinal canal. The patient with a developmentally small caliber cervical spinal canal. The patient with a developmentally small caliber cervical spinal canal. The patient with a developmentally small caliber cervical spinal canal has developed cord compression with only moderate disk bulging at C4–C5, C5–C6, and C6–C7.

osteophytosis, facet joint and capsular hypertrophy, ligamentum flavum hypertrophy, and spondylolisthesis.

Cervical myelopathy is usually due to acquired degenerative change superimposed on a developmentally small spinal canal.¹⁰ Patients with developmental canal stenosis tend to present 5.5 years earlier with degenerative cervical myelopathy and with worse baseline neurologic function.¹¹ Regarding treatment, surgeons may be more inclined to undertake posterior decompression in patients with significant developmental spinal canal narrowing because it alleviates cord compression by canal expansion rather than by removal of the anterior compressive elements.

A developmentally small cervical spinal canal also predisposes to traumatic cord injury because it associated with smaller neural foramina, degenerative foraminal stenosis, and radiculopathy. A developmentally narrow canal may affect cervical kinematics and mechanical loading, potentially accelerating degenerative disease.¹²

Given the frequency of degenerative cervical myelopathy, canal measurements from C3 to C7 are the most useful. Reference intervals should be related to sex and height because these are the two variables that most influence cervical spinal canal size.^{4,13} Radiographic measures (i.e., the Pavlov-Torg ratio) do not correlate well with MR and CT, so radiography should not be relied on to determine cervical spinal canal stenosis.^{14–16} Spinal canal CSA can only be measured on axial images, but AP diameter can be measured on both sagittal and axial images. Sagittal and axial images yield comparable measures for spinal canal AP diameter with axial measurements possibly being very marginally (~ 1.5%) larger.⁴

From an MRI-based study of 140 white asymptomatic volunteers (mean age: 38 years; range: 18–78 years), we can appreciate the considerable variability that exists for developmental cervical spinal canal size.⁴ For example, the AP diameter of the spinal canal, measured on axial images at the midvertebral body level, for a 1.7-m tall woman was 12.5 mm (range: 9.5–16.5 mm) at C3 and 12.2 mm (range: 9.3–16.1 mm) at C6.⁴ This represents a ~ 70% spread for AP diameter across the cohort.

An even greater variability in developmental canal size was seen for spinal CSA than AP diameter.⁴ For 1.7-m tall women, spinal canal CSA was 189 mm² (125–287mm²) at C3 and 177 mm² (117–268 mm²) at C6. This represents ~ 120% for CSA across the cohort. In other words, developmental CSA varies more than AP diameter. Height has a much greater influence on canal AP diameter and CSA (both < 0.001) than age (0.042 and 0.031, respectively).⁴

The normal values reported by Ulbrich et al would have been considered abnormal based on other study criteria. For example, in a study of 1,211 asymptomatic subjects, the mean AP diameter of the spinal canal at the C5 mid-vertebral level was 15.8 ± 1.5 for all subjects, 16.2 ± 1.5 for men and 15.4 ± 1.4 for women.¹⁷ A developmentally narrow canal was defined as an AP diameter < 14 mm,¹⁷ which was present in 10% of subjects. In 0.5%, the canal was < 12 mm, in 2.4%, it was < 13 mm, and in 7.3%, it was < 14 mm. Elsewhere, developmental cervical canal stenosis was defined as a cervical canal diameter < 12 to 13 mm.^{18–20} Asymptomatic subjects with spinal cord compression or cord signal changes had significantly smaller cervical spinal canals than those without cord compression or cord signal change.¹³ The critical AP diameter below which cord compression was likely was < 14.8 mm in men and < 13.9 mm in women.¹³ We can appreciate even from these results the variability seen in reported data. These is a real need for population reference intervals to be established for different heights and sex and move away from the one-measurement-fits-all scenario.

Symptomatic patients show a much higher prevalence of developmental canal narrowing. For example, 50% of symptomatic men and 78% of symptomatic women had an AP diameter of < 15 mm at the C5 mid-vertebral body level.¹³ A total of 68% of those patients with a narrow spinal canal (< 14 mm AP diameter) were female.¹³ This sex difference reflects using the same criteria for men and women.

A study of CT imaging comparing cervical spine injured patients with control subjects concluded that the shape of the spinal canal and not the area put the patient at risk for spinal cord injury.⁷ The values for spinal canal size on this CT

study were generally larger than the values reported from the MRI study.⁷ No comparative study comparing CT versus MRI in cervical spinal measurements has been performed.

Cervical Spinal Cord Dimensions

The spinal cord is malleable and changes shape to conform with the surrounding spinal canal configuration, from circular in the upper cervical region to ellipsoid coronally in the lower cervical region.⁴ Although sagittal images provide an indication of the likelihood of cord injury or compression, cord size should only be measured as cord CSA on axial images because cord AP diameter will give a false underestimation of cord size due to the change in cord shape.

When measuring the spinal cord, cadaveric studies can underestimate cord size measurements due to the effects to fixation and shrinkage.⁶ Also, for in vivo imaging, cord area tends to be slightly larger on T1-weighted rather than T2-weighted axial images.²¹

Men have slightly larger (~ 3.5%) cervical cords than women.^{4,5,21-24} Taller people also have larger spinal cords.^{4,22} The cervical spinal cord size reduced slightly in size with age²⁵ in the order of 1% reduction in cord size per decade.⁶ Spinal cord size seems to remain relatively constant in size during flexion and extension.²⁶ The largest CSA of the spinal cord seems to be at C4–C5 or C5–C6; the CSA of the spinal cord CSA at C3 and C6 is comparable.^{4–6}

Comparing two separate studies that measured the spinal cord CSA in Japanese⁵ and in European⁴ volunteers, cervical spinal cord measurements were consistently \sim 7% smaller for Japanese volunteers, which may be related to differences in height between both cohorts. So, for example, at C6, the difference in average spinal cord CSA for women was 68 mm² versus 75 mm²; for men, it was 71 mm² versus 77 mm².^{4,5}

Within the same population cohort, the spinal cord does vary in size, but this variation is much less than that seen in spinal canal measurements. For 1.7-m tall European women, average spinal cord CSA was 75 mm² (55–101 mm²) at C3 and 75 mm² (56–102 mm²) at C6.⁴ This represents an approximate 40% spread in spinal canal CSA for C3 and C6 across the studied cohort. Because the variation in spinal canal CSA (~ 120%) is much greater than spinal cord CSA (~ 40%), spinal canal size has a much greater bearing on spinal cord to canal mismatch than spinal cord CSA.

The spinal cord occupation ratio (SCOR), which reflects the amount of spinal canal space occupied by the spinal cord, has been used as a measure of spinal cord to canal mismatch.^{5,22,27,28} Patients with spinal cord to canal mismatch are at risk of spinal cord injury, degenerative cervical myelopathy, and neurapraxia. This may be due to less cerebrospinal fluid around the cord, (1) making the cord more prone to degenerative encroachment, (2) reducing the ability to absorb kinetic forces that are transferred to the cord, (3) increasing the likelihood of venous congestive myelopathy, and finally (4) smaller cervical vertebrae may biomechanically be more prone to degeneration.^{12,29} Patients with spinal cord mismatch have worse baseline neurologic function. Because lower baseline neurologic function is a negative predictor of outcome, identification of a significant spinal cord mismatch may alter patient prognosis.¹¹ Comparing SCORs in degenerative cervical myelopathy patients, the prevalence of mismatch was lower in Europe at 2.3% compared with ~ 10% in Asia, Latin America, and North America.³⁰ Routine assessment of developmental spinal canal stenosis may be indicated in those engaged in extreme and high-impact contact sports such as rugby, skiing, basketball, waterskiing, and football. Because there is relatively more space around the cord in taller patients, this probably puts a shorter person at higher risk of cervical cord compression.⁴

The three approaches to assessing spinal cord objectively are canal mismatch quantifying the (1) AP space around the cord as a ratio, (2) AP space around the cord as an absolute measure, or (3) CSA around the cord as a ratio. It is not known which of these approaches is best.

Using the first of these measures, spinal cord AP divided by spinal canal AP diameter, it was recommended that an SCOR > 75% (i.e., a cord occupying > 75% of the canal) at C5, which was close to 2 SDs above the mean, should be used as an indicator of spinal canal to cord mismatch.⁵ Alternatively, averaging SCOR at the two most adjacent noncompressed sites in patients with degenerative cervical myelopathy, a SCOR $\ge 70\%$ was used as an indicator of significant mismatch.³¹

The second approach is to subtract the spinal cord AP diameter from the canal AP diameter.^{32,33} In a retrospective analysis of cervical spine MRI examinations in athletes with and without neuropraxia, a value < 5 mm increased the likelihood of chronic neurapraxia risk, particularly if it was < 4.3 mm.³²

The third approach is to compare CSAs on axial imaging. In a review of patients with minor cervical spine trauma, a SCOR > 80% on axial MRI could identify patients at risk for spinal cord injury.²⁷

Therefore, to date, a spinal cord to canal mismatch can be categorized as a SCOR \geq 75% for an AP ratio, < 5 mm for an AP absolute measure, and \geq 80% for a CSA ratio.^{27,28,32} Clearly, SCOR at the moment can only be measured on MRI, whereas spinal canal dimensions can be measured on both MRI and CT. Spinal canal developmental size varies much more than spinal cord size. Whether spinal cord area is related to or independent of spinal canal area is not known. If the former is the case, the focus should be on spinal canal size rather than SCOR. If the latter is the case, it would seem reasonable to focus on SCOR.

Cervical Vertebral Body Dimensions

The volume of the cervical vertebrae is relevant to study of factors governing cervical vertebral body size as well as corpectomy and cage placement, among other things. Establishing reference standards for normal vertebrae and spinal dimensions can be achieved only after controlling for sex, age, and ethnicity due to the influence of these factors on vertebral anatomy.³⁴ In a study of cervical vertebrae of 277 human skeletons born between 1825 and 1910, a detailed analysis of vertebral body dimensions showed that African Americans had significantly greater vertebral body width and depth in the C3–C5 region than European Americans.³⁴ The heights of the C3 and C4 vertebral bodies were significantly smaller in the African American population. Cervical vertebral bodies became

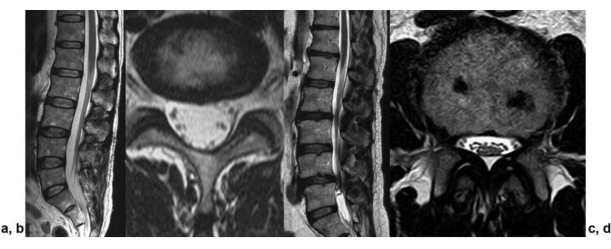


Fig. 2 (a) T2-weighted sagittal and (b) axial images of patient with developmentally normal caliber lumbar spinal canal compared with (c) T2-weighted sagittal and (d) axial images of patient with developmentally small caliber lumbar spinal canal. The patient with a developmentally small caliber lumbar spinal canal has developed lumbar spinal canal stenosis due to only mild anterolisthesis and disk bulging at L4–L5.

wider and deeper with age with the changes in depth much more pronounced than the changes in width.³⁴ There was a significant reduction in vertebral body height with age.³⁴

That males have larger cervical vertebral bodies than females regardless of ethnic origin is well documented.³⁵ While the height of the vertebral body decreases with age mainly between C3 and C6, the vertebral body expands. For example, in C3, between young and old patients, vertebral body width increased by 6% and vertebral body depth increased by 9%, whereas at C7, width increased by $\sim 6\%$ while depth increased by $\sim 15\%$.³⁵ With advancing age, reduction in vertebral body height tends to be greater in women while the increase in vertebral body depth is similar for men and women.³⁶ Because there is no change in the two measurements of the exit foramen with age, it can be concluded that most of the vertebral body expansion in depth seems to occur anteriorly. Another possible explanation for the changing vertebral body shape with age is the charge in cervical spine curvature that occurs to compensate for the increased thoracic kyphosis.³⁷

Developmental Lumbar Spine Canal Size

Lumbar spinal canal stenosis is a common clinical entity. Similar to the cervical spine, two features govern the development of lumbar spinal canal stenosis: developmental lumbar spinal canal size and acquired narrowing of the lumbar spinal canal (**- Fig. 2**). It is important to separate these two elements when quantifying lumbar spinal canal size.

Developmental lumbar canal size relates to the size of the osseous spinal canal when spinal maturation is complete by the age of 17 years. It is unclear what factors govern lumbar spinal canal size, although it may be related to shorter gestation age, lower birth weight, primiparity, and lower socioeconomic class.³⁸ Developmental lumbar spinal canal size is measured at the level of the pedicles removed from the acquired changes that predominate at the diskovertebral level (**Fig. 3**). To develop population reference intervals, a large cross section of the population irrespective of symptoms must be studied. Using data obtained from abdominal pelvic CT examinations performed for reasons not related to the lumbar spine provides a good method of obtaining a

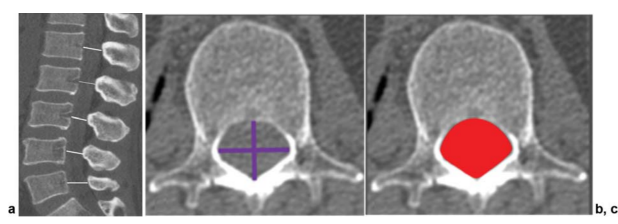


Fig. 3 (a) Developmental canal size is measured at the pedicular level removed from acquired degenerative narrowing that occurs at the diskovertebral level. (b) Anteroposterior and lateral diameters for lumbar developmental canal size. (c) Cross-sectional area for lumbar developmental canal size.

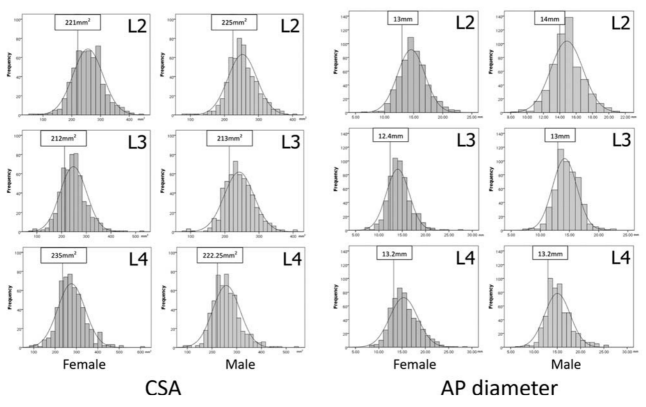


Fig. 4 Population reference intervals for lumbar developmental canal size for (**a**) cross-sectional area (CSA) and (**b**) anteroposterior (AP) diameter of the L2, L3, and L4 lumbar levels for females and males. The demarcation of the smaller quartile (i.e., smaller 25%) at each level is indicated (modified from Griffith et al⁴⁰).

population reference range for spinal dimensions with no need for additional specific imaging^{39,40} (**~Fig. 4**).

Such an approach could be undertaken for each region, race, and sex because considerable cross-population variation does seem to exist in patients with smaller lumbar spinal canals who are more likely to develop symptoms and require surgical intervention.^{39–41} Using this approach, a large (at least 34% between largest and smallest quartiles) variation in lumbar developmental spinal canal CSA can be appreciated.⁴⁰ Spinal canal CSA and depth is consistently smallest at L3, enlarging cranially and caudally within the lumbar region.⁴⁰ There is no accepted definition of what defines a normal or abnormal spinal canal size.⁴² An L3 anteroposterior diameter < 13 mm for males and < 12.4 mm for females represented the smallest 25% for a southern Chinese population.⁴⁰ Similarly, an osseous spinal canal < 212 mm² for males and < 213 mm² for females represented the smallest 25% of the population.⁴⁰

Lumbar Vertebral Body Dimensions

The volume of the thoracic and lumbar vertebral bodies is relevant to vertebroplasty and other procedures. Using CTbased technique, verified by analysis of saw bone vertebral bodies, vertebral body volume increased from T1 to L4 for both female and male subjects.⁴³ Mean thoracic vertebral body volume was ~ 15.0 cm³; mean lumbar vertebral body volume was much higher at ~ 35 cm³.^{43,44} Considerable individual variation in vertebral body size was present ranging from 5.2 cm³ for a T1 vertebral body to 60.6 cm³ for a L4 vertebral body.⁴³ Vertebral body volume in men was larger than that of women only in the lumbar spine, mainly accountable by an increase in vertebral body CSA.⁴⁵ Although thoracic and lumbar vertebral body height is, as expected, closely related to patient height, lumbar vertebral body CSA in males is related to patient weight.⁴⁵ The L5 vertebral body volume is normally up to 5% smaller than the L3 or L4 vertebral bodies.^{43,44} Patients with L5 pars defects tend to have even smaller L5 vertebral bodies, ~ 10% smaller than L4, with greater posterior wedging.⁴⁴

Bone Mineral Density

BMD is the most important surrogate marker of bone strength. It is most widely measured by DXA, although this is an areal rather than volumetric assessment measuring both cortical and trabecular BMD.⁴⁶ Trabecular volumetric BMD (vBMD) by single-energy QCT is a more accurate, although less practical, reference standard for noninvasive vBMD assessment.⁴⁷ With QCT, osteopenia is defined as a trabecular BMD < 120 mg/cm³; osteoporosis is defined as a trabecular BMD < 80 mg/cm³.⁸ Radiation dose for lumbar QCT is ~ 1 mSv for men and 1.6 mSv for women, many times higher than lumbar DXA (1–6 µSV) but comparable with lumbar spine radiography (0.7–2 MSv).^{47,48}

Using spectral detection CT as a feature of dual-energy CT, vBMD can be quantified without using a calibration phantom ("phantomless") yielding high correlation with standard QCT assessment (r = 0.987).⁴⁹

Partial volume averaging due to bone voxels that contain fat will reduce trabecular vBMD measurement by singleenergy QCT by ~ 15%.⁵⁰ A method to correct single-energy QCT for marrow fat content was also recently developed.⁵¹ After first adjusting manufacturer-calibrated values for the European Spine Phantom (ESP-145), density measurements can be adjusted to account for bone marrow adipose tissue (BMAT) content as measured by MRI as follows:

vBMD $_{BMATcorr} = vBMD _{ESPcorr} + 0.7576 \times BMAT$ (%) - 12.96 (mg/cm³).

After correcting for marrow fat content, mean L2–L4 vertebral body vBMD decreased in women from 20 to 80 years by -2.57 ± 0.11 mg/cm³/year, $\sim15\%$ less than the uncorrected value $(-3.00\pm0.13$ mg/cm³/year). Mean vBMD annual decrease for men was -1.70 ± 0.14 mg/cm³/year, $\sim10\%$ less than the uncorrected value $(1.92\pm0.15$ mg/cm³/year).^{51} Dual-energy CT can also accurately measure marrow fat content and enable correction of QCT vBMD data accordingly.^{50,52}

Bone islands are much more common than sclerotic metastases and tend to have higher bone density.^{53,54} This density difference can be used to distinguish bone island and sclerotic metastasis with a high level of certainty. This is done by drawing a region of interest (ROI) as large as possible on the lesion without extending beyond the lesion's margins, irrespective of whether intravenous contrast was administered or not. Either a mean attenuation \geq 885 HU or a maximum attenuation \geq 1,060 HU for the lesion has a sensitivity of 95% and specificity of 96% for diagnosing a bone island.⁵³

Marrow Fat Content and Composition

Magnetic resonance spectroscopy (MRS) can measure bone marrow fat composition as well as content but is limited in allowing assessment of only one vertebral body. Single-voxel MRS-based marrow fat quantification should ideally be based on a multi-TE MRS measurement to minimize confounding effects on proton-density fat fraction and water fractions.⁵⁵

Chemical shift encoded separation techniques (fat-water imaging) can also accurately measure marrow fat content and has a shorter acquisition time of 3 to 4 minutes across a wider scan plane, although it cannot assess marrow fat composition.^{56–58} The reproducibility of ¹H MR spectroscopy and chemical-shift imaging is high, tending to be best in those areas with the highest inherent fat fraction.^{59–61} A standardized T1-signal intensity can also be measured, that is, the vertebral marrow to subcutaneous fat ratio.⁶² Al-though straightforward, accuracy is compromised because T1 signal intensity depends on factors such as coil position, saturation band positioning, and postprocessing homogenization of B1 inhomogeneity, in addition to marrow fat.⁶²

Marrow fat content increases from the cervical to the lumbar spine.^{61,62} When comparing fat measurements between studies, we need to consider from which vertebrae were taken. The L3 vertebral body is the most usually measured, although the T11 and L1 vertebral bodies may be more representative.^{62,63}

There is a gradual physiologic increase in vertebral marrow fat content with age, from $\sim 25\%$ marrow fat content at 25 years to 65% at 65 years.^{64,65} This increase is different in

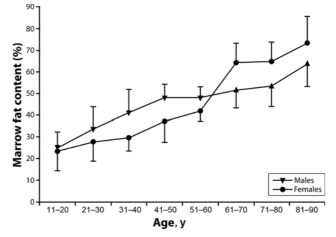


Fig. 5 Marrow fat content, % of lumbar vertebral body stratified for age and sex. Throughout life, marrow fat increases slowly and steadily for both sexes with a dramatic increase in marrow fat content between the ages of 55 and 65 years for women. This mirrors the reduction in bone mineral density that occurs at that time in women.

males and females. Males show a gradual increase of 7% per decade in lumbar vertebral marrow fat content; females show an increase of ~ 5% per decade up to 55 years followed by a dramatic increase in marrow fat content between 55 and 65 years of ~ $25\%^{64,66}$ (**~ Fig. 4**). Male lumbar vertebrae have ~ 10% more fat than females up to 50 years.⁶⁴ Between 50 and 60 years, this sex difference narrows and disappears.⁶⁴ After 60 years, females have ~ 10% more marrow fat in their vertebrae than males⁶⁶ (**~ Fig. 5**).

Overall, there is at least a 40 to 50% increase in fat cell content with increasing age. This increase in fat content happens primarily due to an increase in fat cell number rather than fat cell size,⁶⁷ and it seems to be associated with little, if any, change in marrow fatty acid profile.⁶⁸ This increase in fat cell volume occurs primarily at the expense of red marrow volume. Although the trabecular volume decreases with age, the overall percentage decrease in trabecular volume is small at ~ 5% in volume. Because the marrow cavity is a defined space and vascular sinusoids and other marrow structures do not seem to expand with age, we can infer that for any increase in fat cell content, there is almost a corresponding decrease in functioning or red marrow content. In other words, marrow fat content should be a good surrogate marker for hematopoietic marrow content (**– Fig. 6**).⁶⁹

The inverse relationship between increasing marrow fat and trabecular bone loss in osteoporosis has been recognized histologically for 50 years.⁷⁰ MRI, and more recently dualenergy CT, have enabled marrow fat content to be quantified noninvasively on a large scale and at different anatomical sites.⁵² Over and above the physiologic increase in marrow fat content with age, osteoporosis is associated with an even greater increase in marrow fat content in older men with normal BMD, osteopenia, and osteoporosis was $50.1 \pm 8.7\%$, $55.7 \pm 10.2\%$, and $58.2 \pm 7.8\%$, respectively (p = 0.002).⁷¹ Similarly, for older women, marrow fat content in those with normal BMD, osteopenia, and osteoporosis was $59.2 \pm 10.0\%$,

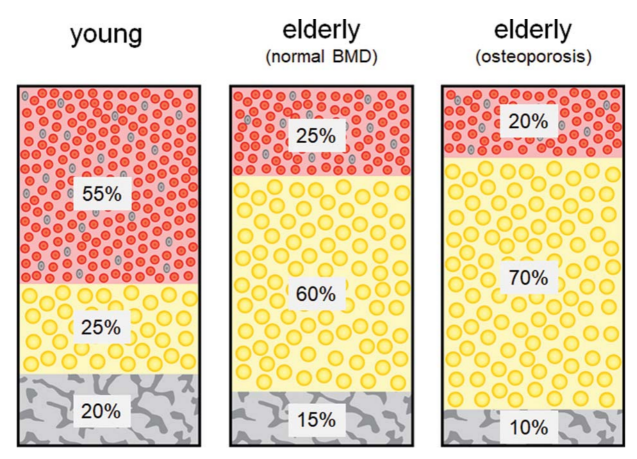


Fig. 6 The bone marrow is confined by the rigid bone cortex. As marrow fat content increases with age, red marrow content decreases because any age-related with or without osteoporotic-related change in trabecular bone volume is relatively small. In other words, marrow fat content is a surrogate marker of red marrow fat content. Physiologic changes in bone marrow content are exaggerated in patients with osteoporosis. BMD, bone mineral density.

 $63.3 \pm 9.5\%$, and $67.7 \pm 8.5\%$, respectively (p = 0.002).⁷² Similar results were shown using three-dimensional gradient-echo chemical-shift-encoded fat-water imaging.⁵⁶ In a study of 51 patients (28 females; mean age: 69.7 ± 9.0 years), mean fat content of normal BMD vertebrae ($56.3 \pm 14.8\%$) was higher than that of osteoporotic vertebrae ($62.4 \pm 11.0\%$; p = 0.007).⁵⁶ Trabecular bone loss, which can be specifically measured by QCT, as opposed to DXA that usually measures both trabecular and cortical bone loss, seems to be much more associated with increased marrow fat content than cortical bone loss.⁷³

Higher prevalent marrow fat predicts future increased osteoporotic bone loss. Postmenopausal women with a marrow fat content above the median experienced average bone loss of 4.7% over 4 years, whereas those with a marrow fat content below the median only experienced bone loss of 1.6%.⁷⁴ Similarly, for each 1 SD (\sim 8%) increase in baseline marrow fat content, trabecular vBMD tended to increase by \sim 0.9% more over a 3-year period.⁷⁵

Increased marrow fat may also increase vertebral fracture risk. Mean vertebral marrow fat was 55% in those with a prevalent vertebral fracture compared with 45% (p < 0.001) in those without a prevalent vertebral fracture even after adjusting for the effect of marrow fat on DXA measurements.⁷⁶ To date, no longitudinal studies have investigated whether marrow fat content is related to incident vertebral fracture.⁷³ In addition, osteoporosis marrow fat is also increased in other conditions associated with reduced BMD such as anorexia nervosa, alcohol abuse, spinal cord injury, and prolonged bed rest.⁵²

Although chronic osteoporotic-type fractures can be readily recognized on MRI, quantitative MR imaging can be helpful in distinguishing between acute/subacute vertebral fracture and malignant fracture. The presence of fat within the fractured vertebral body on T1-weighted images is very helpful sign in identifying the fracture as osteoporotic.⁷⁷ In equivocal cases, chemical shift fat to water imaging can augment standard imaging in quantifying fat content within the fractured vertebral body. Either the absolute fat content or a fat fraction ratio can be used where the fat content of the fractured vertebral body is compared with the fat content of an adjacent normal-appearing vertebral body. A fat fraction < 9% within the fractured vertebral body or a fat fraction ratio < 0.2 indicates malignancy with a 96% sensitivity and 89% specificity.^{78,79}

MRS can also evaluate fat composition. The portion of unsaturated vertebral marrow fat in healthy young subjects was higher than postmenopausal osteoporotic women (0.127 versus 0.091),⁸⁰ a finding confirmed by spectroscopy of iliac crest aspirates.⁷³ Patients with prevalent vertebral fracture patients have 1.7% lower unsaturation levels and 2.9% higher vertebral marrow fat saturation levels than those with no fracture.⁸¹ People with diabetes also had 1.3% lower unsaturation levels, higher vertebral marrow fat saturation levels,

Bone Marrow Perfusion

MRI can measure marrow perfusion with high reproducibly using empirical time-intensity curve measures such as maximal enhancement (Emax) and enhancement slope (Eslope).⁶⁰ Pharmacokinetic modeling can also be applied, although these measures are less robust and more theoretical. Within the rigid confines of the bone cortex, because the two main constituents of bone marrow are red and fatty marrow, fatty marrow content is a surrogate marker for red marrow content. In other words, the higher the fatty marrow content, the lower the red marrow content. Red marrow, being much more metabolically active than fatty marrow, is the main driver of bone marrow perfusion. Therefore, as expected, changes in bone marrow perfusion coincide well with changes in red marrow content.⁶⁷ Bone marrow perfusion is also related to endothelial dysfunction and atherosclerosis with a weak negative correlation (r = -0.33; p = 0.0018) between vertebral body perfusion and carotid artery intima media thickness.⁸²

Vertebral marrow perfusion reduces with aging.^{83–87} Overall Emax was > 60% greater in subjects aged < 50 years (58.2 ± 44.6) than subjects aged > 50 years (21.9 ± 14.8) .⁸³ For females only, an even greater difference was apparent with Eslope in those < 50 years (87.2 ± 54.1) being 80% greater than those > 50 years (18.0 ± 13.8) .⁸³ Although vertebral bone marrow perfusion is higher in young females than young males, the rate of perfusion decline is also higher in females, such that vertebral bone marrow perfusion is higher in older men than older women.⁸³ This pattern closely matches changes that occur in red to fatty marrow composition with age. Also, in line with changing marrow fat content, the upper (L1–L2) lumbar vertebral bodies are better perfused than the lower (L3–L5) vertebral bodies.^{85,86}

Bone perfusion is also closely related to BMD. As BMD decreases, marrow fat increases. Patients with osteoporosis

have lower bone marrow perfusion than those with osteopenia, whereas patients with osteopenia have lower bone marrow perfusion than those with normal BMD⁸⁸ (**-Fig. 7**). This reduction in perfusion affects the marrow but not the surrounding muscle, indicating it is an integral marrow effect rather than part of a more generalized circulatory impairment.⁷² Bone perfusion is a critical element in bone fracture healing including microfracture healing. Wee can appreciate how compromised perfusion in osteoporotic bone could lead to impaired microfracture healing and thus to microfracture propagation until a spontaneous macroscopic insufficiency fracture occurs, occasionally followed by fracture nonunion.^{88,89}

Bone Marrow Diffusion

Free movement of extracellular water molecules in the bone marrow is affected by the cellular compaction and the amount of interstitial fluid. Extracellular water motion also depends on factors such as blood flow, capillary permeability, interstitial pressure, temperature, and the viscosity of interstitial fluid. The standard measure of water diffusivity in tissues is the apparent diffusion coefficient (ADC) acquired from single-shot echo planar imaging. The ADC of water is 3.0×10^{-3} mm²/s. The ADC of bone marrow is the lowest of nearly all tissues at 0.20 to 0.60×10^{-3} mm²/s, and particularly fatty marrow that has an ADC about half that of red marrow.^{90–92}

Overall, ADC values tend to decrease with increasing age; the correlation (r = -0.3 to -0.4; p = 0.001) is weak.^{93,94} Mean vertebral ADC values in subjects aged < 30 years ($0.54 \pm 0.07 \times 10^{-3} \text{ mm}^2/\text{s}$) is higher than that in subjects aged > 30 years ($0.47 \pm 0.08 \times 10^{-3} \text{ mm}^2/\text{s}$).⁹⁴ The correlation between decreasing ADC values and increasing age is higher in women (r = -0.581; p < 0.001) than men,^{93,95,96} most likely due to age-related increase in marrow fat and decrease in marrow perfusion. Marrow ADC values also decrease from L1 to L5 in line with increasing fat and decreasing perfusion from the upper to the lower lumbar region.⁹⁷

ADC values in middle-aged to elderly patients with normal BMD were 0.47 ± 0.03 ; those with osteopenia were

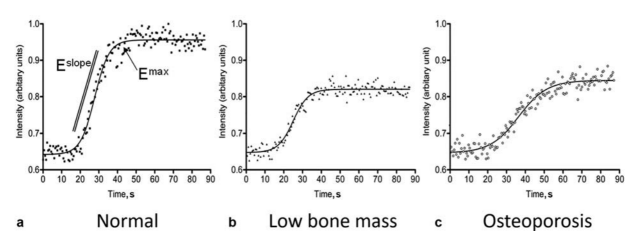


Fig. 7 Typical time-intensity perfusion curves of vertebral body marrow perfusion from dynamic contrast-enhanced magnetic resonance imaging examination in subjects with (a) normal bone mineral density (BMD), (b) low bone mass (osteopenia), and (c) osteoporosis. The rate and degree of intravenous contrast in subjects with osteoporosis is less than in normal BMD subjects. Emax, maximal enhancement; Eslope, enhancement slope.

 0.42 ± 0.02 and with osteoporosis were $0.39\pm0.03.^{98}$ Similar differences were found by some 72,99 but not all 100 other studies, probably related to different examination protocols, analytical methods, and cohorts. 73

Bone Marrow Metabolism

Based on the degree of glucose utilization, fluorodeoxyglucose accumulation provides a measure of tissue metabolic activity and, as such, can be used as a useful measure of bone marrow metabolic activity.¹⁰¹ Combining volumetric MRI data and quantitative metabolic data from positron emission tomography (PET) to calculate the relative metabolic activity of red and yellow marrow in the L1, L3, and L5 vertebral bodies, the mean maximum standardized uptake (SUV_{max}) of fat and red marrow was 0.38 ± 0.1 and 2.6 ± 0.6 , respectively. In other words, metabolic activity of red marrow was seven times that of fatty marrow.¹⁰² Bone marrow metabolic activity tended to decrease with age more in the proximal femora and humeri (r = -0.60 to -0.67; p < 0.01) than the axial skeletal (sternal manubrium, 12th thoracic, 5th lumbar vertebra, and iliac crests) correlation coefficient -0.28 to -0.48; p < 0.05).¹⁰³

A ¹⁸F-NaF PET/CT provides a specific measure of mineralized bone, rather than bone marrow metabolism and can also quantify arterial calcification.^{104,105} Analyzing this tracer uptake in patients undergoing ¹⁸F-NaF PET/CT for suspected bone metastases, SUV_{mean} in the lumbar spine was 6.9 ± 1.9 (2.2–14.4), which was several times higher than the femoral neck, $2.7 \pm 1.0 (0.9-8.0)$.¹⁰⁰ Mean regional bone metabolism at the lumbar spine and femoral neck decreased with increasing age (r = -0.44; p < 0.001) and overall was lower in female patients.¹⁰⁰ Bone mineral metabolism was also negatively correlated with hypertension (p = 0.003), hypercholesterolemia (p = 0.01), and prior cardiovascular events, which is not surprising given the well-known association between osteoporosis and arteriosclerosis, particularly arterial calcification.^{100,106}

Disk Volume and Degeneration

Disk volume is relevant to the study of disk degeneration and other aspects of spinal pathology.¹⁰⁷ The volume of a normal lumbar intervertebral disk is $\sim 10~{\rm cm}^3$ when measured in the evening and $\sim 11~{\rm cm}^3$ when measuring in the morning after a night's rest. In other words, the disk, through the effect of its hydrophilic proteoglycan molecules, attaches $\sim 0.9~{\rm cm}^3$ of water into the disk overnight. Hence people are slightly taller in the morning and astronauts in space, who lack this cyclical fluid flow as a result of microgravity, lose height and have back pain. In osteoporotic and low bone mass patients, while central disk height is increased, giving rise to the biconcave appearance, overall lumbar disk volume is actually reduced compared with normal BMD patients.¹⁰⁸ This may be due to compromised nutritional supply from the neighboring vertebrae as BMD decreases.¹⁰⁸

To improve the quantification of disk degeneration, surrogate MR measures of tissue hydration, such as T1-, T2-, and T2* relaxation times, and T1 ρ have been investigated. Diskal T2 relaxation times decrease with disk hydration and, to a lesser extent, with reduced proteoglycan and collagen content.^{109–112} T2 relaxation provides a continuous measure of small changes in disk composition over time such as diurnal variation in disk water content¹¹² and the effect of normal aging.^{112,113} T2 relaxation times of the nucleus pulposus decrease ~ 10% per decade with physiologic aging.¹¹³ Overall disk T2 relaxation time tends to be lower in chronic low back pain patients than in symptomatic controls, especially in the posterior annulus fibrosus.¹¹⁴ This heightens the importance of addressing different areas of the disk when comparing different cohorts. On T2-weighted images, T2 or T1 ρ maps, both the nucleus pulposus and the inner fibers of the annulus fibrosus have comparable signal intensity because the boundary between these two components of the disk is indistinct.

Several methods have been described to demarcate the nucleus and annulus on MR images, either based on visual assessment or quantitative measures based on equal areas of five or seven ROIs placed horizontally across the disk, with or without an intervening gap. The central three ROIs are usually defined as the nucleus pulposus. Visual assessment seems to produce a more reliable delineation of the nucleus and annulus than quantitative delineation.¹¹⁵ T1p relaxation measurement, which probes the interaction between water molecules and their macromolecular environment, has the potential to identify early biochemical changes in the disk, although the specific determinants of T1p relaxation are not well understood. T1p mapping may be slightly more sensitive than T2 mapping for disk degeneration, although it is more time consuming. Both T2 and T1p have a floor effect such that with more severe degeneration neither T2 or T1p measurements will change. Differences in T1p values between healthy and degenerative disks can be small, whereas errors in T1p quantification are many including B1 RF inhomogeneity and BO field inhomogeneity as well as data acquisition. This limits its clinical application as does the current lack of validation with a reference standard.¹¹⁶

Ligamentum Flavum Thickness

The ligamentum flavum extends from C2 to S1, bridging the anteroinferior part of the more cephalad lamina to the posterosuperior aspect of the more caudad lamina. It prevents separation of the posterior elements during spinal flexion and restores an erect posture after flexion. With increasing age, there is reduced elasticity of the ligament with an increase in the collagen and a decrease in the elastic fiber content. Most ligamentum flavum hypertrophy is caused by reduced disk height bulking the ligament rather than true hypertrophy of the ligament. 107,117 Hypertrophy of the ligamentum flavum reduces the size of the central canal, lateral recesses, and exit foramina, compressing the descending and exiting nerve roots, and leading to radiculopathy. Average thickness of the ligamentum flavum in the cervical spine is 2.2 ± 0.4 mm, thinnest at C2–C3 and thickest in the lower cervical region.¹¹⁸

Average thickness of the ligamentum flavum in the thoracic region is 2.6 ± 0.7 mm, thinnest in the upper and thickest in the mid- to lower thoracic region.¹¹⁹ No age variation was apparent.¹¹⁹ Maximum thickness of the ligamentum flavum in the thoracic spine is at the T10–T11 level (3.3 ± 0.9 mm),

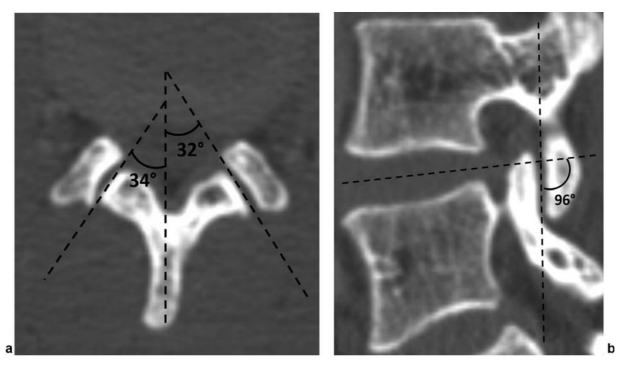


Fig. 8 The axial facet joint angle is typically measured between the mid-vertebral body sagittal line and a line drawn between the anteromedial and posterolateral edges of the superior articular facet at the disk level. In this example, the angles are 32 degrees and 34 degrees. There is no facet joint tropism. The sagittal angle is measured as the angle between a line bisecting the intervertebral disk and one drawn between the superior and inferior aspects of the facet joint. In this example, the angle is 96 degrees.

possibly due to larger tensile forces and mobility at this level.¹¹⁹ In the lumbar spine, ligamentum flavum thickness > 4 mm is considered hypertrophy.^{102,120} The ligamentum flavum tends to be thickest at L4–L5 followed by L5–S1 and is ~ 5 to 10% thicker on the left side.¹²¹

Facet Joint Orientation

Facet joint orientation/asymmetry is relevant to spinal biomechanics and seems to be related to the development of spondylolisthesis.¹²²

The axial facet joint angle is typically measured either between the mid-vertebral body sagittal line and a line drawn between the anteromedial and posterolateral edges of the superior articular facet at the disk level¹²² (**Fig. 8**). Normally, the facet joint angle moves away from the mid-sagittal line, from an average of \sim 42 to 58 degrees while descending the lumbar spine.¹⁰²

In a population study, using data deemed from routine abdominopelvic CT examinations, and defining spondylolisthesis as an intervertebral body shift > 7% of the AP diameter of the inferior vertebral body, spondylolisthesis was present in 8.7% of males and 16.4% of females, and most prevalent at L4–L5.¹²² Facet joint angles in the axial plane decreased, on average, with age in women but not in men, from ~ 48 to 43 degrees at L4–L5 and from ~ 53 to 47 degrees at L5–S1.¹²² Facet joint angles in the sagittal plane decreased in men more than women¹²² (**¬Fig. 8**). This sagittal decrease in men was from ~ 83 to 80 degrees at L3–L4 and from ~ 81 to 79 degrees at L4–L5.¹²² This sagittalization and axialization of the facet joint with age may help explain why older people are more prone to degenerative spondylolisthesis.¹²²

Facet joint tropism refers to facet joint angle symmetry. Facet joint trophism is considered acceptable if the side-to side difference in facet joint angles in either the axial or the sagittal plane is < 7 degrees.¹²² Facet joint tropism in the axial plane is considered moderate if the difference is 7 to 15 degrees and severe if the difference is > 15 degrees.¹⁰² Facet joint trophism is most severe at L4–L5, where a 20% prevalence of moderate and a 6% prevalence of severe trophism can exist.¹⁰² Also, patients with degenerative spondylolisthesis tend to have more facet joint tropism.¹²²

In conclusion, we can appreciate how the many aspects of spine anatomy and physiology make it particularly conducive to quantitative analysis. Quantitative analysis of spine imaging data has begun to provide us with a much better understanding of spinal disease. More standardized analytics methods will allow more ready comparison of research studies with radiologists in a great position to drive this research forward.

Conflict of Interest None declared.

References

- 1 Weaver O, Leung JWT. Biomarkers and imaging of breast cancer. AJR Am J Roentgenol 2018;210(02):271–278
- 2 Whyte MB, Kelly P. The normal range: it is not normal and it is not a range. Postgrad Med J 2018;94(1117):613–616

- ³ Flechtner-Mors M, Schwab KO, Fröhlich-Reiterer EE, et al. Overweight and obesity based on four reference systems in 18,382 paediatric patients with type 1 diabetes from Germany and Austria. J Diabetes Res 2015;2015:370753
- 4 Ulbrich EJ, Schraner C, Boesch C, et al. Normative MR cervical spinal canal dimensions. Radiology 2014;271(01):172–182
- 5 Kato F, Yukawa Y, Suda K, Yamagata M, Ueta T. Normal morphology, age-related changes and abnormal findings of the cervical spine. Part II: Magnetic resonance imaging of over 1,200 asymptomatic subjects. Eur Spine J 2012;21(08):1499–1507
- 6 Fradet L, Arnoux PJ, Ranjeva JP, Petit Y, Callot V. Morphometrics of the entire human spinal cord and spinal canal measured from in vivo high-resolution anatomical magnetic resonance imaging. Spine 2014;39(04):E262–E269
- 7 Matsuura P, Waters RL, Adkins RH, Rothman S, Gurbani N, Sie I. Comparison of computerized tomography parameters of the cervical spine in normal control subjects and spinal cord-injured patients. J Bone Joint Surg Am 1989;71(02):183–188
- 8 American College of Radiology. Practice parameter for the performance of quantitative computed tomography (QCT) bone densitometry. Available at: https://www.acr.org/Quality-Safety/Standards-Guidelines/Practice-Guidelines-by-Modality/ CT. Accessed April 12, 2020
- 9 Zhang JT, Wang LF, Liu YJ, et al. Relationship between developmental canal stenosis and surgical results of anterior decompression and fusion in patients with cervical spondylotic myelopathy. BMC Musculoskelet Disord 2015;16:267
- 10 Kadoya S, Nakamura T, Kwak R, Hirose G. Anterior osteophytectomy for cervical spondylotic myelopathy in developmentally narrow canal. J Neurosurg 1985;63(06):845–850
- 11 Nouri A, Martin AR, Nater A, et al. Influence of magnetic resonance imaging features on surgical decision-making in degenerative cervical myelopathy: results from a global survey of AOSpine international members. World Neurosurg 2017;105:864–874
- 12 Morishita Y, Naito M, Hymanson H, Miyazaki M, Wu G, Wang JC. The relationship between the cervical spinal canal diameter and the pathological changes in the cervical spine. Eur Spine J 2009; 18(06):877–883
- 13 Nagata K, Yoshimura N, Hashizume H, et al. The prevalence of cervical myelopathy among subjects with narrow cervical spinal canal in a population-based magnetic resonance imaging study: the Wakayama Spine Study. Spine J 2014;14(12):2811–2817
- 14 Blackley HR, Plank LD, Robertson PA. Determining the sagittal dimensions of the canal of the cervical spine. The reliability of ratios of anatomical measurements. J Bone Joint Surg Br 1999;81 (01):110–112
- 15 Herzog RJ, Wiens JJ, Dillingham MF, Sontag MJ. Normal cervical spine morphometry and cervical spinal stenosis in asymptomatic professional football players. Plain film radiography, multiplanar computed tomography, and magnetic resonance imaging. Spine 1991;16(06):S178–S186
- 16 Prasad SS, O'Malley M, Caplan M, Shackleford IM, Pydisetty RK. MRI measurements of the cervical spine and their correlation to Pavlov's ratio. Spine 2003;28(12):1263–1268
- 17 Nakashima H, Yukawa Y, Suda K, Yamagata M, Ueta T, Kato F. Narrow cervical canal in 1211 asymptomatic healthy subjects: the relationship with spinal cord compression on MRI. Eur Spine J 2016;25(07):2149–2154
- 18 Hayashi H, Okada K, Hamada M, Tada K, Ueno R. Etiologic factors of myelopathy. A radiographic evaluation of the aging changes in the cervical spine. Clin Orthop Relat Res 1987;(214):200–209
- 19 Inoue H, Ohmori K, Takatsu T, Teramoto T, Ishida Y, Suzuki K. Morphological analysis of the cervical spinal canal, dural tube and spinal cord in normal individuals using CT myelography. Neuroradiology 1996;38(02):148–151
- 20 Kasai Y, Akeda K, Uchida A. Physical characteristics of patients with developmental cervical spinal canal stenosis. Eur Spine J 2007;16(07):901–903

- 21 Kim G, Khalid F, Oommen VV, et al. T1- vs. T2-based MRI measures of spinal cord volume in healthy subjects and patients with multiple sclerosis. BMC Neurol 2015;15:124
- 22 Ishikawa M, Matsumoto M, Fujimura Y, Chiba K, Toyama Y. Changes of cervical spinal cord and cervical spinal canal with age in asymptomatic subjects. Spinal Cord 2003;41(03):159–163
- 23 Kameyama T, Hashizume Y, Ando T, Takahashi A. Morphometry of the normal cadaveric cervical spinal cord. Spine 1994;19(18):2077–2081
- 24 Lee MJ, Cassinelli EH, Riew KD. Prevalence of cervical spine stenosis. Anatomic study in cadavers. J Bone Joint Surg Am 2007;89(02):376–380
- 25 Yanase M, Matsuyama Y, Hirose K, et al. Measurement of the cervical spinal cord volume on MRI. J Spinal Disord Tech 2006;19 (02):125–129
- 26 Sayıt E, Aghdasi B, Daubs MD, Wang JC. The occupancy of the components in the cervical spine and their changes with extension and flexion. Global Spine J 2015;5(05):396–405
- 27 Rüegg TB, Wicki AG, Aebli N, Wisianowsky C, Krebs J. The diagnostic value of magnetic resonance imaging measurements for assessing cervical spinal canal stenosis. J Neurosurg Spine 2015;22(03):230–236
- 28 Nouri A, Montejo J, Sun X, et al. Cervical cord-canal mismatch: a new method for identifying predisposition to spinal cord injury. World Neurosurg 2017;108:112–117
- 29 Countee RW, Vijayanathan T. Congenital stenosis of the cervical spine: diagnosis and management. J Natl Med Assoc 1979;71 (03):257–264
- 30 Nouri A, Martin AR, Tetreault L, et al. MRI analysis of the combined prospectively collected AOSpine North America and international data: the prevalence and spectrum of pathologies in a global cohort of patients with degenerative cervical myelopathy. Spine 2017;42(14):1058–1067
- 31 Nouri A, Tetreault L, Nori S, Martin AR, Nater A, Fehlings MG. Congenital cervical spine stenosis in a multicenter global cohort of patients with degenerative cervical myelopathy: an ambispective report based on a magnetic resonance imaging diagnostic criterion. Neurosurgery 2018;83(03):521–528
- 32 Presciutti SM, DeLuca P, Marchetto P, Wilsey JT, Shaffrey C, Vaccaro AR. Mean subaxial space available for the cord index as a novel method of measuring cervical spine geometry to predict the chronic stinger syndrome in American football players. J Neurosurg Spine 2009;11(03):264–271
- 33 Tierney RT, Maldjian C, Mattacola CG, Straub SJ, Sitler MR. Cervical spine stenosis measures in normal subjects. J Athl Train 2002;37(02):190–193
- 34 Ezra D, Masharawi Y, Salame K, Slon V, Alperovitch-Najenson D, Hershkovitz I. Demographic aspects in cervical vertebral bodies' size and shape (C3-C7): a skeletal study. Spine J 2017;17(01): 135–142
- 35 Stemper BD, Yoganandan N, Pintar FA, et al. Anatomical gender differences in cervical vertebrae of size-matched volunteers. Spine 2008;33(02):E44–E49
- 36 Pashkova IG, Kosourov AK. Age dependent changes in cervical region of the spine according to data of nuclear magnetic tomography. [in Russian]. Morfologiia 2004;125(01):80–82
- 37 Boyle JW, Nikitov SA, Boardman AD, Booth JG, Booth K. Nonlinear self-channeling and beam shaping of magnetostatic waves in ferromagnetic films. Phys Rev B Condens Matter 1996;53(18): 12173–12181
- 38 Papp T, Porter RW, Craig CE, Aspden RM, Campbell DM. Significant antenatal factors in the development of lumbar spinal stenosis. Spine 1997;22(16):1805–1810
- 39 Schizas C, Schmit A, Schizas A, Becce F, Kulik G, Pierzchała K. Secular changes of spinal canal dimensions in western Switzerland: a narrowing epidemic? Spine 2014;39(17):1339–1344
- 40 Griffith JF, Huang J, Law SW, et al. Population reference range for developmental lumbar spinal canal size. Quant Imaging Med Surg 2016;6(06):671–679

- 41 Wiley MR, Hee Jo C, Khaleel MA, McIntosh AL. Size matters: which adolescent patients are most likely to require surgical decompression for lumbar disk herniations? J Pediatr Orthop 2019;39(10):e791–e795
- 42 Steurer J, Roner S, Gnannt R, Hodler J; LumbSten Research Collaboration. Quantitative radiologic criteria for the diagnosis of lumbar spinal stenosis: a systematic literature review. BMC Musculoskelet Disord 2011;12:175
- 43 Limthongkul W, Karaikovic EE, Savage JW, Markovic A. Volumetric analysis of thoracic and lumbar vertebral bodies. Spine J 2010;10(02):153–158
- 44 Wilms GE, Willems E, Demaerel P, De Keyzer F. CT volumetry of lumbar vertebral bodies in patients with hypoplasia L5 and bilateral spondylolysis and in normal controls. Neuroradiology 2012;54(08):839–843
- 45 Caula A, Metmer G, Havet E. Anthropometric approach to lumbar vertebral body volumes. Surg Radiol Anat 2016;38(03):303–308
- 46 Griffith JF, Engelke K, Genant HK. Looking beyond bone mineral density : Imaging assessment of bone quality. Ann N Y Acad Sci 2010;1192:45–56
- 47 Löffler MT, Sollmann N, Mei K, et al. X-ray-based quantitative osteoporosis imaging at the spine. Osteoporos Int 2020;31(02): 233–250
- 48 Adams JE. Quantitative computed tomography. Eur J Radiol 2009;71(03):415–424
- 49 Roski F, Hammel J, Mei K, et al. Bone mineral density measurements derived from dual-layer spectral CT enable opportunistic screening for osteoporosis. Eur Radiol 2019;29(11):6355–6363
- 50 Sfeir JG, Drake MT, Atkinson EJ, et al. Evaluation of crosssectional and longitudinal changes in volumetric bone mineral density in postmenopausal women using single- versus dualenergy quantitative computed tomography. Bone 2018; 112:145–152
- 51 Cheng X, Li K, Zhang Y, et al. The accurate relationship between spine bone density and bone marrow in humans. Bone 2020; 134:115312
- 52 Bredella MA, Daley SM, Kalra MK, Brown JK, Miller KK, Torriani M. Marrow adipose tissue quantification of the lumbar spine by using dual-energy CT and single-voxel (1)H MR spectroscopy: a feasibility study. Radiology 2015;277(01):230–235
- 53 Ulano A, Bredella MA, Burke P, et al. Distinguishing untreated osteoblastic metastases from enostoses using CT attenuation measurements. AJR Am J Roentgenol 2016;207(02):362–368
- 54 Sala F, Dapoto A, Morzenti C, et al. Bone islands incidentally detected on computed tomography: frequency of enostosis and differentiation from untreated osteoblastic metastases based on CT attenuation value. Br J Radiol 2019;92(1103):20190249
- 55 Dieckmeyer M, Ruschke S, Cordes C, et al. The need for T_2 correction on MRS-based vertebral bone marrow fat quantification: implications for bone marrow fat fraction age dependence. NMR Biomed 2015;28(04):432–439
- 56 Kühn JP, Hernando D, Meffert PJ, et al. Proton-density fat fraction and simultaneous R2* estimation as an MRI tool for assessment of osteoporosis. Eur Radiol 2013;23(12):3432–3439
- 57 Zhang Y, Zhou Z, Wang C, et al. Reliability of measuring the fat content of the lumbar vertebral marrow and paraspinal muscles using MRI mDIXON-Quant sequence. Diagn Interv Radiol 2018; 24(05):302–307
- 58 Karampinos DC, Baum T, Nardo L, et al. Characterization of the regional distribution of skeletal muscle adipose tissue in type 2 diabetes using chemical shift-based water/fat separation. J Magn Reson Imaging 2012;35(04):899–907
- 59 Bernard CP, Liney GP, Manton DJ, Turnbull LW, Langton CM. Comparison of fat quantification methods: a phantom study at 3.0T. J Magn Reson Imaging 2008;27(01):192–197
- 60 Griffith JF, Yeung DK, Chow SK, Leung JC, Leung PC. Reproducibility of MR perfusion and (1)H spectroscopy of bone marrow. J Magn Reson Imaging 2009;29(06):1438–1442

- 61 Baum T, Yap SP, Dieckmeyer M, et al. Assessment of whole spine vertebral bone marrow fat using chemical shift-encoding based water-fat MRI. J Magn Reson Imaging 2015;42(04):1018–1023
- 62 Sieron D, Drakopoulos D, Loebelenz LI, et al. Correlation between fat signal ratio on T1-weighted MRI in the lower vertebral bodies and age, comparing 1.5-T and 3-T scanners. Acta Radiol Open 2020;9(01):2058460120901517
- 63 Belavy DL, Quittner MJ, Ridgers ND, Shiekh A, Rantalainen T, Trudel G. Specific modulation of vertebral marrow adipose tissue by physical activity. J Bone Miner Res 2018;33(04):651–657
- 64 Kugel H, Jung C, Schulte O, Heindel W. Age- and sex-specific differences in the 1H-spectrum of vertebral bone marrow. J Magn Reson Imaging 2001;13(02):263–268
- 65 Duda SH, Laniado M, Schick F, Strayle M, Claussen CD. Normal bone marrow in the sacrum of young adults: differences between the sexes seen on chemical-shift MR imaging. AJR Am J Roentgenol 1995;164(04):935–940
- 66 Griffith JF, Yeung DKW, Ma HT, Leung JC, Kwok TCY, Leung PC. Bone marrow fat content in the elderly: a reversal of sex difference seen in younger subjects. J Magn Reson Imaging 2012;36(01):225–230
- 67 Griffith JF, Wang YX, Zhou H, et al. Reduced bone perfusion in osteoporosis: likely causes in an ovariectomy rat model. Radiology 2010;254(03):739–746
- 68 Griffith JF, Yeung DK, Ahuja AT, et al. A study of bone marrow and subcutaneous fatty acid composition in subjects of varying bone mineral density. Bone 2009;44(06):1092–1096
- 69 Griffith JF. Age-related changes in the bone marrow. Curr Radiol Rep 2017;5:24
- 70 Dunnill MS, Anderson JA, Whitehead R. Quantitative histological studies on age changes in bone. J Pathol Bacteriol 1967;94(02): 275–291
- 71 Griffith JF, Yeung DK, Antonio GE, et al. Vertebral bone mineral density, marrow perfusion, and fat content in healthy men and men with osteoporosis: dynamic contrast-enhanced MR imaging and MR spectroscopy. Radiology 2005;236(03):945–951
- 72 Griffith JF, Yeung DK, Antonio GE, et al. Vertebral marrow fat content and diffusion and perfusion indexes in women with varying bone density: MR evaluation. Radiology 2006;241(03): 831–838
- 73 Li X, Shet K, Xu K, Rodríguez JP, Pino AM, Kurhanewicz J, Schwartz A, Rosen CJ. Unsaturation level decreased in bone marrow fat of postmenopausal women with low bone density using high resolution magic angle spinning (HRMAS) 1H NMR spectroscopy. Bone. 2017;105:87–92
- 74 Griffith JF, Yeung DK, Leung JC, Kwok TC, Leung PC. Prediction of bone loss in elderly female subjects by MR perfusion imaging and spectroscopy. Eur Radiol 2011;21(06):1160–1169
- 75 Woods GN, Ewing SK, Sigurdsson S, et al. Greater bone marrow adiposity predicts bone loss in older women. J Bone Miner Res 2020;35(02):326–332
- 76 Wehrli FW, Hopkins JA, Hwang SN, Song HK, Snyder PJ, Haddad JG. Cross-sectional study of osteopenia with quantitative MR imaging and bone densitometry. Radiology 2000;217(02):527–538
- 77 Griffith JF. Identifying osteoporotic vertebral fracture. Quant Imaging Med Surg 2015;5(04):592–602
- 78 Schmeel FC, Luetkens JA, Feißt A, et al. Quantitative evaluation of T2* relaxation times for the differentiation of acute benign and malignant vertebral body fractures. Eur J Radiol 2018;108:59–65
- 79 Suh CH, Yun SJ, Jin W, Park SY, Ryu CW, Lee SH. Diagnostic performance of in-phase and opposed-phase chemical-shift imaging for differentiating benign and malignant vertebral marrow lesions: a meta-analysis. AJR Am J Roentgenol 2018; 211(04):W188–W197
- 80 Yeung DK, Griffith JF, Antonio GE, Lee FK, Woo J, Leung PC. Osteoporosis is associated with increased marrow fat content and decreased marrow fat unsaturation: a proton MR spectroscopy study. J Magn Reson Imaging 2005;22(02):279–285

- 81 Patsch JM, Li X, Baum T, et al. Bone marrow fat composition as a novel imaging biomarker in postmenopausal women with prevalent fragility fractures. J Bone Miner Res 2013;28(08):1721–1728
- 82 Chen WT, Ting-Fang Shih T, Hu CJ, Chen RC, Tu HY. Relationship between vertebral bone marrow blood perfusion and common carotid intima-media thickness in aging adults. J Magn Reson Imaging 2004;20(05):811–816
- 83 Chen WT, Shih TT, Chen RC, et al. Vertebral bone marrow perfusion evaluated with dynamic contrast-enhanced MR imaging: significance of aging and sex. Radiology 2001;220(01):213–218
- 84 Montazel JL, Divine M, Lepage E, Kobeiter H, Breil S, Rahmouni A. Normal spinal bone marrow in adults: dynamic gadoliniumenhanced MR imaging. Radiology 2003;229(03):703–709
- 85 Savvopoulou V, Maris TG, Vlahos L, Moulopoulos LA. Differences in perfusion parameters between upper and lower lumbar vertebral segments with dynamic contrast-enhanced MRI (DCE MRI). Eur Radiol 2008;18(09):1876–1883
- 86 Hillengass J, Stieltjes B, Bäuerle T, et al. Dynamic contrastenhanced magnetic resonance imaging (DCE-MRI) and diffusion-weighted imaging of bone marrow in healthy individuals. Acta Radiol 2011;52(03):324–330
- 87 Baur A, Stäbler A, Bartl R, Lamerz R, Scheidler J, Reiser M. MRI gadolinium enhancement of bone marrow: age-related changes in normals and in diffuse neoplastic infiltration. Skeletal Radiol 1997;26(07):414–418
- 88 Griffith JF, Yeung DK, Tsang PH, et al. Compromised bone marrow perfusion in osteoporosis. J Bone Miner Res 2008;23(07): 1068–1075
- 89 Wang YX, Griffith JF, Kwok AW, et al. Reduced bone perfusion in proximal femur of subjects with decreased bone mineral density preferentially affects the femoral neck. Bone 2009;45(04): 711–715
- 90 Biffar A, Dietrich O, Sourbron S, Duerr HR, Reiser MF, Baur-Melnyk A. Diffusion and perfusion imaging of bone marrow. Eur J Radiol 2010;76(03):323–328
- 91 Ward R, Caruthers S, Yablon C, Blake M, DiMasi M, Eustace S. Analysis of diffusion changes in posttraumatic bone marrow using navigator-corrected diffusion gradients. AJR Am J Roentgenol 2000;174(03):731–734
- 92 Li X, Schwartz AV. MRI assessment of bone marrow composition in osteoporosis. Curr Osteoporos Rep 2020;18(01):57–66
- 93 Jie H, Hao F, Na LX. Vertebral bone marrow diffusivity in healthy adults at 3T diffusion-weighted imaging. Acta Radiol 2016;57 (10):1238–1243
- 94 Herrmann J, Krstin N, Schoennagel BP, et al. Age-related distribution of vertebral bone-marrow diffusivity. Eur J Radiol 2012; 81(12):4046–4049
- 95 Yeung DK, Wong SY, Griffith JF, Lau EM. Bone marrow diffusion in osteoporosis: evaluation with quantitative MR diffusion imaging. J Magn Reson Imaging 2004;19(02):222–228
- 96 Nonomura Y, Yasumoto M, Yoshimura R, et al. Relationship between bone marrow cellularity and apparent diffusion coefficient. J Magn Reson Imaging 2001;13(05):757–760
- 97 Lasbleiz J, Le Ster C, Guillin R, Saint-Jalmes H, Gambarota G. Measurements of diffusion and perfusion in vertebral bone marrow using intravoxel incoherent motion (IVIM) with multishot, readout-segmented (RESOLVE) echo-planar imaging. J Magn Reson Imaging 2019;49(03):768–776
- 98 Hatipoglu HG, Selvi A, Ciliz D, Yuksel E. Quantitative and diffusion MR imaging as a new method to assess osteoporosis. AJNR Am J Neuroradiol 2007;28(10):1934–1937
- 99 Tang GY, Lv ZW, Tang RB, et al. Evaluation of MR spectroscopy and diffusion-weighted MRI in detecting bone marrow changes in postmenopausal women with osteoporosis. Clin Radiol 2010; 65(05):377–381
- 100 Shih TT, Liu HC, Chang CJ, Wei SY, Shen LC, Yang PC. Correlation of MR lumbar spine bone marrow perfusion with bone mineral density in female subjects. Radiology 2004;233(01):121–128

- 101 Blebea JS, Houseni M, Torigian DA, et al. Structural and functional imaging of normal bone marrow and evaluation of its agerelated changes. Semin Nucl Med 2007;37(03):185–194
- 102 Basu S, Houseni M, Bural G, et al. Magnetic resonance imaging based bone marrow segmentation for quantitative calculation of pure red marrow metabolism using 2-deoxy-2-[F-18]fluoro-Dglucose-positron emission tomography: a novel application with significant implications for combined structure-function approach. Mol Imaging Biol 2007;9(06):361–365
- 103 Fan C, Hernandez-Pampaloni M, Houseni M, et al. Age-related changes in the metabolic activity and distribution of the red marrow as demonstrated by 2-deoxy-2-[F-18]fluoro-D-glucosepositron emission tomography. Mol Imaging Biol 2007;9(05): 300–307
- 104 Derlin T, Janssen T, Salamon J, et al. Age-related differences in the activity of arterial mineral deposition and regional bone metabolism: a 18F-sodium fluoride positron emission tomography study. Osteoporos Int 2015;26(01):199–207
- 105 Kurata S, Shizukuishi K, Tateishi U, et al. Age-related changes in pre- and postmenopausal women investigated with 18F-fluoride PET—a preliminary study. Skeletal Radiol 2012;41(08): 947–953
- 106 Griffith JF, Kumta SM, Huang Y. Hard arteries, weak bones. Skeletal Radiol 2011;40(05):517–521
- 107 Sudhir G, Vignesh Jayabalan S, Gadde S, Venkatesh Kumar G, Karthik Kailash K. Analysis of factors influencing ligamentum flavum thickness in lumbar spine: a radiological study of 1070 disc levels in 214 patients. Clin Neurol Neurosurg 2019; 182:19–24
- 108 Kwok AW, Wang YX, Griffith JF, et al. Morphological changes of lumbar vertebral bodies and intervertebral discs associated with decrease in bone mineral density of the spine: a cross-sectional study in elderly subjects. Spine 2012;37(23):E1415–E1421
- 109 Tertti M, Paajanen H, Laato M, Aho H, Komu M, Kormano M. Disc degeneration in magnetic resonance imaging. A comparative biochemical, histologic, and radiologic study in cadaver spines. Spine 1991;16(06):629–634
- 110 Marinelli NL, Haughton VM, Muñoz A, Anderson PA. T2 relaxation times of intervertebral disc tissue correlated with water content and proteoglycan content. Spine 2009;34(05):520–524
- 111 Kerttula L, Kurunlahti M, Jauhiainen J, Koivula A, Oikarinen J, Tervonen O. Apparent diffusion coefficients and T2 relaxation time measurements to evaluate disc degeneration. A quantitative MR study of young patients with previous vertebral fracture. Acta Radiol 2001;42(06):585–591
- 112 Boos N, Wallin A, Gbedegbegnon T, Aebi M, Boesch C. Quantitative MR imaging of lumbar intervertebral disks and vertebral bodies: influence of diurnal water content variations. Radiology 1993;188(02):351–354
- 113 Marinelli NL, Haughton VM, Anderson PA. T2 relaxation times correlated with stage of lumbar intervertebral disk degeneration and patient age. AJNR Am J Neuroradiol 2010;31(07):1278–1282
- 114 Ogon I, Takebayashi T, Takashima H, et al. Analysis of chronic low back pain with magnetic resonance imaging T2 mapping of lumbar intervertebral disc. J Orthop Sci 2015;20(02):295–301
- 115 Mok GSP, Zhang D, Chen SZ, Yuan J, Griffith JF, Wang YXJ. Comparison of three approaches for defining nucleus pulposus and annulus fibrosus on sagittal magnetic resonance images of the lumbar spine. J Orthop Translat 2016;6:34–41
- 116 Wáng YX. Towards consistency for magnetic resonance (MR) relaxometry of lumbar intervertebral discs. Quant Imaging Med Surg 2016;6(04):474–477
- 117 Altinkaya N, Yildirim T, Demir S, Alkan O, Sarica FB. Factors associated with the thickness of the ligamentum flavum: is ligamentum flavum thickening due to hypertrophy or buckling? Spine 2011;36(16):E1093–E1097
- 118 Rahmani MS, Terai H, Akhgar J, et al. Anatomical analysis of human ligamentum flavum in the cervical spine: Special

Seminars in Musculoskeletal Radiology Vol. 24 No. 4/2020

consideration to the attachments, coverage, and lateral extent. J Orthop Sci 2017;22(06):994–1000

- 119 Chelladurai A, Balasubramaniam S, Anbazhagan SP, Gnanasihamani S, Ramaswami S. Dorsal spinal ligamentum flavum thickening: a magnetic resonance imaging study. Asian Spine J 2018;12(01):47–51
- 120 Munns JJ, Lee JY, Espinoza Orías AA, et al. Ligamentum flavum hypertrophy in asymptomatic and chronic low back pain subjects. PLOS One 2015;10(05):e0128321
- 121 Kolte VS, Khambatta S, Ambiye MV. Thickness of the ligamentum flavum: correlation with age and its asymmetry—an magnetic resonance imaging study. Asian Spine J 2015;9(02): 245–253
- 122 Morimoto M, Higashino K, Manabe H, et al. Age-related changes in axial and sagittal orientation of the facet joints: Comparison with changes in degenerative spondylolisthesis. J Orthop Sci 2019;24(01):50–56