The Immature Pediatric Appendicular Skeleton

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Abstract

Growth and maturation occur in a predictable pattern throughout the body and within each individual bone. In the appendicular skeleton, endochondral ossification predominates in long bones and growth plates. The ends of these long bones are sites of relative weakness in the immature skeleton and prone to injury from acute insult and overuse. We present the normal histoanatomy and physiology of the growth plate complex, highlighting the unique contribution of each component and shared similarities between primary and secondary complexes. Components of the growth plate complex include the physis proper, subjacent vascularity within the growth cartilage, and the ossification front. The second section describes imaging considerations and features of normal and abnormal growth. Finally, we review the Salter-Harris classification for acute fractures and offer examples of characteristic overuse injury patterns involving the epiphyseal (proximal humerus and distal radius), apophyseal (medial epicondyle and tibial tubercle), and secondary growth plate complexes (medial femoral condyle and capitellar osteochondritis dissecans). This article provides a foundation and basic framework to better understand and anticipate potential complications and growth disturbances and to ensure optimal follow-up and early intervention when treatment can be less invasive.

Keywords
► growth disturbance
► athlete
► fracture
► endochondral ossification
► growth plate

Growth and maturation from an osteochondral immature skeleton in children to the predominantly osseous mature skeleton in adults occurs in a predictable pattern throughout the body and within each bone. In flat bones, such as the calvarium, bone formation occurs directly on a mesenchymal scaffold, and is known as intramembranous ossification. In long bones that make up most of the appendicular skeleton, bone formation occurs on a cartilaginous framework called the growth plate (or physis), and this is known as endochondral ossification.1,2 In children and pediatric athletes, traumatic and overuse injuries involving the appendicular skeleton are common, and the latter predominates. These injuries can lead to endochondral ossification dysfunction, producing a spectrum of findings that vary in severity and reversibility, impacting clinical symptomology and long-term outcomes.

During early embryonic development, around week 6 of gestation, the mesenchymal cells differentiate into chondrocytes, responsible for the formation of a cartilaginous model of the future skeleton. Within long bones, chondrocytes within the central portion of this cartilaginous anlage (future diaphysis) undergo cellular hypertrophy and the surrounding matrix calcifies. Around week 7 of gestation, the periosteal sleeve forms, and at the end of week 8, vascular invasion brings mesenchymal cells that differentiate into osteoblasts and osteoclasts.3 Osteoblasts produce an osteoid matrix on the cartilaginous framework that later becomes the future primary ossification center. In contrast, the osteoclasts are
responsible for bone remodeling and formation of the medullary cavity. These primary ossification centers grow and expand bidirectionally toward the ends of the long bone with the two leading edges becoming the proximal and distal primary growth plates\(^4,5\) (\(\text{Fig. 1}\)).

At the ends of these long bones, endochondral growth and ossification occur not only at the primary “discoid” growth plates that underlie epiphyses and apophyses, but also at the secondary “spherical” growth plates that surround epiphyses, apophyses, and small round bones, such as carpal, tarsal, and sesamoid bones. The pressure epiphyses (and the underlying epiphyseal growth plates) are subject to compressive forces, whereas the traction epiphyses or apophyses (and the underlying apophyseal growth plates) serve as attachment sites for tendons to bone and are primarily subject to tensile forces.

The smaller secondary growth plates (or acrophyses) are responsible for the centripetal growth of the underlying secondary ossification centers, which, depending on anatomical location, can be single or multiple and later coalesce as they slowly and progressively replace the unossified growth cartilage\(^1,83\) (\(\text{Fig. 2}\)). These primary and secondary growth plates are sites of relative weakness in the skeletally immature skeleton, especially during periods of rapid growth. Injury to the growth plate may produce a spectrum of clinical findings depending on the type of growth plate involved, anatomical location, which component or components are predominantly injured, the severity of the insult, and the patients’ remaining growth potential.\(^6\)

This article reviews the histoanatomy and unique contribution of each component of the growth plate complex, highlighting shared similarities between primary and secondary growth plates. Imaging considerations and imaging appearances of the growth plates at different stages of skeletal development and maturation are reviewed. Salter-Harris patterns of physeal fractures are then discussed briefly, followed by a more detailed review on our emerging understanding of the pathophysiology that underlies physeal stress injury, divided into those that involve the epiphyseal, apophyseal, and secondary growth plate complexes.

Representative examples with characteristic clinical presentations and key imaging findings that impact treatment

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**Fig. 1** Fetal bone anatomy. Sagittal TRUFI (True Fast Imaging with Steady-State Free Precession) magnetic resonance image from a 37-week gestational fetus shows ossified femoral shaft (arrowhead), cartilaginous epiphyses (arrows, femoral head and condyles), and apophysis (dashed arrow, greater trochanter) underlying the primary growth plates or physes (triangles). Wavy arrow denotes the cartilaginous patella.

**Fig. 2** Growth plate (physeal) anatomy. (a) Growth plate illustration shows arrangement of the primary and secondary physes, intact vascularity within the growth cartilage, and terminal vascular loops within the ossification front. Chondrocytes undergo apoptosis at and within the zone of provisional calcification (ZPC; shown in purple). (b) Distal femur illustration and (c) coronal intermediate-weighted fat-suppressed magnetic resonance image of the knee from a 10-year-old boy show primary (triangles) and secondary physes (chevrons). (a, b) Adapted and modified from Nguyen et al\(^6\) with permission. Note: Illustrations are not drawn to scale.
Growth Plate Proper
On histology, the growth plate contains chondrocytes that mature as they move from the growth cartilage side toward the ossification front (Fig. 4). Three distinct zones of chondrocytes are recognized, based on their microscopic cellular morphology, organization, and function: reserve, proliferative, and hypertrophic zones. The growth plate is surrounded at its periphery by the perichondrium (Fig. 2).

Reserve Zone
In the reserve (or germinal) zone, chondrocytes are randomly distributed and only divide sporadically. An abundant surrounding extracellular matrix is responsible for storing nutrients, including glycogen and lipids. In this zone, collagen fibrils are disorganized.

Proliferative Zone
In the proliferative zone, chondrocytes are organized into columns that divide rapidly. The top cell of each column (adjacent to the reserve zone) is the progenitor cell, and the rate of division correlates with the rate of bone growth. Less abundant extracellular matrix contains collagen fibrils organized into longitudinal septa. Oxygen and nutrients are derived from arterial branches that arise from the juxaphysyeal growth cartilage and travel within vascular channels and through the reserve zone to nourish the upper proliferative zone. Oxygen tension is the highest in this zone, facilitating robust aerobic metabolism and mitochondrial uptake of calcium.

Hypertrophic Zone
In the hypertrophic zone, chondrocytes mature and undergo internal vacuolation and apoptosis (programmed cell death). This vacuolation process leads to the production of cytoplasmic “holes” and induces nuclear fragmentation and loss of cell membrane integrity. Extracellular glycogen is progressively depleted, and oxygen tension is substantially reduced, leading to anaerobic metabolism and mitochondrial release of calcium, facilitating matrix calcification.
Initially, calcification occurs along the longitudinal septa. Further accumulation of calcium salts creates amorphous deposits of calcium phosphate. These give way to hydroxyapatite crystals that subsequently coalesce, forming the zone of provisional calcification (ZPC). Progressive matrix calcification hinders the diffusion of nutrients and oxygen, further ensuring chondrocyte apoptosis. The ZPC demarcates the transition between the growth plate proper and the subjacent ossification front. In contrast to the arteries within the growth cartilage, the articular loops within the ossification front terminate just short of the ZPC and play no role in nourishing the hypertrophic chondrocytes.

Perichondrium
The perichondrium surrounds the periphery of the growth plate and contains the groove of Ranvier and the ring of LaCroix. The groove of Ranvier is a wedge-shaped structure responsible for the transverse growth of the growth plate. The ring of LaCroix is a dense fibrous band that provides regional mechanical support to this osteochondral junction and is continuous with the periosteum. Like the periosteum, the perichondrium is hypointense on all magnetic resonance imaging (MRI) pulse sequences. But in contrast to the periosteum, the perichondrium is tightly attached to the underlying growth plate. The perichondrial vessels supply the perichondrium, which also nourishes the most peripheral aspects of the growth plate, leading to a relatively more vulnerable central growth plate that can be prone to ischemic insult and can produce a “cupping” deformity.

Growth Cartilage: Epiphyses and Apophyses
With few exceptions, epiphyses and apophyses were initially entirely cartilaginous during fetal and early postnatal development. Nutrient vessels within growth cartilage travel within vascular channels, and small vascular branches extend into the growth plate proper to nourish chondrocytes within the reserve and upper proliferative zones. These vessels produce a gradient of decreasing oxygen tension that extend from the growth cartilage side toward the ossification front side of the growth plate.

Disruption of these nutrient vessels leads not only to endochondral ossification dysfunction but also to epiphyseal and apophyseal deformity. Scattered transphyseal vascular channels are present among infants <12 to 18 months of age that allows vascular communication and spread of disease to cross the growth plate. But these vessels involute with progressive growth, and vascular communication is reestablished only after physeal closure at skeletal maturation. This accounts for the relatively higher frequency of epiphyseal osteomyelitis among infants when compared with older children.

Epiphyses
The immature osteochondral epiphysis contains a superficial layer of avascular articular cartilage, a middle layer of vascularized unossified growth cartilage, and a centrally located SOC. This middle growth cartilage layer is progressively replaced over time by the enlarging SOC, so that at skeletal maturation only the superficial layer of articular cartilage remains.

Apophyses
The immature osteochondral apophysis contains a peripheral layer of vascularized unossified growth cartilage and a centrally located SOC. Analogous to the epiphysis, this growth cartilage layer is progressively replaced over time by the enlarged SOC, so that at skeletal maturation no cartilage remains.

Secondary Ossification Centers
Within epiphyses and apophyses, the SOC initially forms following the coalescence of vascular channels centrally, which leads to the release of metalloproteinases (e.g., gelatinase B and collagenase 3) that break down the cartilaginous matrix, triggering chondrocyte hypertrophy, localized calcification, and ossification. The site of transient change within the regional cartilage, occurring just before the appearance of the SOC, is viewed as a pre-ossification center and should not be mistaken for pathology (Fig. 5a).

The newly formed SOC generally has a round morphology and is surrounded by a secondary growth plate that has the same zonal arrangement as the primary growth plate but is smaller in size. As growth continues, portions of the secondary growth plate close at sites where the overlying growth cartilage has been depleted. This results in the SOC losing its initial spherical morphology and conforming to its underlying cartilaginous framework. For example, in the femoral condyles, the portion of the secondary growth plate that faces the primary growth plate closes first, producing a semicircular morphology of the SOC (Fig. 2).

It is also common for some epiphyses and apophyses to have multiple smaller SOCs that coalesce over time, commonly observed in the immature proximal humeral head, trochlea of the distal humerus, and tibial tuberosity of the

Fig. 5 Spectrum of normal ossification involving the immature trochlea. (a) Sagittal T2-weighted fat-suppressed magnetic resonance image from a 6-year-old boy shows the normal appearance of a hyperintense pre-ossification center (arrowhead). (b) Anteroposterior radiograph from a 13-year-old boy shows fragmented and heterogeneously sclerotic appearance of the trochlear ossification (dashed arrow).
proximal tibia. In the humeral head, for example, three (medial head, greater and lesser tuberosity) SOCs coalesce at early childhood to form an irregularly shaped SOC that ultimately become the humeral head.14

Trochlear ossification is well recognized as one of the most widely variable and irregular, and is occasionally mistaken for pathology (► Fig. 5b). However, the bilateral appearance of this finding and the lack of localized symptoms are helpful signs to avoid unintentional false-positive diagnoses.15 Similarly, intra-cartilaginous trochlear tuberosity ossification must be distinguished from Osgood-Schlatter disease and pathologic avulsion fracture. These are described further in the other anatomically designated articles within the current issue of Seminars.

Ossification Front: Metaphyses, Metaphyseal-Equivalents, and SOCs
At the primary spongiosa of the juxaphyseal metaphysis and metaphyseal-equivalent location, and at the peripheral margin of SOCs that abut the ZPC, osteoblasts lay down bone on the calcified cartilaginous bars. Regional vasculature forms terminal vascular loops and capillary tufts that terminate just proximal to the ZPC, ensuring anaerobic metabolism within the adjacent hypertrophic zone of the growth plate proper, facilitating and indirectly ensuring chondrocyte apoptosis3,4 (► Fig. 2). Disruption of these vessels hinders chondrocyte removal and regional bone formation, leading to physeal widening.

Metaphyses and Metaphyseal-Equivalents
Newly formed bone is remodeled within the secondary spongiosa. Internal remodeling entails the removal of cartilaginous bars and the replacement of lamellar bone, whereas external remodeling involves the peripheral reshaping of the metaphysis into its final funnel-shaped morphology.3,4 Regional blood supply is derived from a combination of central-to-skeletal veins and periosteal arteries that penetrate the metaphysis and contribute ~80% and ~20% to the regional blood supply, respectively.16

Secondary Ossification Centers
Newly formed SOC contains hematopoietic red marrow that converts to fatty yellow marrow at ~6 months after its formation. As the SOC grows along its peripheral margin (ossification front), marrow conversion proceeds in the same central-to-peripheral direction.17 This difference in cellular composition is reflected in the change in signal intensity on MRI that must be distinguished from superimposed insult. Latter pathologic marrow edema is more focal and centered over the site of pathology. Normal physiology marrow conversion and reconversion is further discussed in the section, under growth-related imaging features.

Imaging Considerations and Features
At birth, the growth plates are relatively smooth and flat. With progressive growth, undulation of the growth plate can be observed and is believed to be in response to regional anatomy and location-dependent physiologic dynamic biomechanical forces.1,2,13 During early childhood, the thickness of the growth plate remains relatively constant because of an intricate balance between chondrocyte proliferation and apoptosis. Narrowing across the growth plate signals its impending closure and upcoming regional skeletal maturational closure. Closure starts at sites of major undulation that differs between growth plates depending on anatomical location. The final conversion of chondrocytes into bone leaves behind a physeal scar that represents residual ZPC or leftover horizontal trabeculae.18 This physeal scar is slowly remodeled and finally disappears by early adulthood.19

Imaging Modalities
Different imaging modalities can be used to evaluate and monitor for growth disturbance, but the precise approach depends on the patient’s age, skeletal maturity, and clinical question. Imaging is routinely used to confirm and characterize a clinically suspected diagnosis, and to exclude alternative diagnoses. It generally begins with screening radiographs, followed by advanced cross-sectional imaging to further characterize regional anatomy and pathology, and to estimate growth potential, as well as help guide treatment planning. This section provides an overview of the technical considerations for each of the commonly clinically used imaging modalities: radiographs, MRI, computed tomography (CT), and musculoskeletal ultrasonography (US).

Radiographs
Although the cartilaginous growth plates are radiolucent, radiography remains the initial imaging modality of choice because it is readily available and can exclude alternative diagnoses. Radiographs are generally obtained in two (anteneposterior or posteroanterior and lateral) or three (additional oblique) views, depending on the anatomical location and clinical concerns. Additional oblique and stressed views, as well as single or multiple views of the contralateral asymptomatic side, may be helpful in ambiguous cases to detect subtle asymmetric physeal widening, early remodeling, and/or premature physeal closure (► Fig. 6). Full-length long bone or limb radiographs provide an overview of the regional osseous anatomy, and serial examinations are often used to monitor for asymmetric growth, development of angulation deformity, and response to intervention.

Computed Tomography
As with radiographs, the cartilaginous growth plate is hypointensifying, and its presence is only indirectly inferred from the subjacent ossified elements. In the very young, the abundance of unossified growth cartilage cannot be reliably distinguished from the cartilaginous physis. CT is less routinely used among children due to concerns about the potential adverse cumulative effects of ionizing radiation on growing tissue and the abundance of radiolucent cartilage. The main indications for using CT to assess regional physeal anatomy and pathology include a suspected small

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transphyseal osseous bar or tethering, physeal fractures with intra-articular extension, preoperative planning and/or need for intraoperative navigation, and for those patients who are contraindicated for MRI or sedation.

Magnetic Resonance Imaging
MRI is the preferred imaging modality for the comprehensive evaluation of hyaline cartilage, bone marrow signal intensity, and surrounding soft tissues without the use of ionizing radiation (►Fig. 7). The main drawbacks are possible need for patient sedation, limited immediate availability, and higher cost. Hyaline cartilage is mainly composed of water and a solid matrix that consists of primarily type II collagen and large aggregating proteoglycan macromolecules. Therefore, hyaline cartilage is isointense on T1-weighted images and hyperintense on fluid-sensitive images (e.g., T2-weighted fat-saturated, short tau inversion recovery, STIR) when compared with the signal intensity of skeletal muscle.

Articular, growth, and growth plate cartilage are different types of hyaline cartilage and distinguished based on their anatomical location and signal intensities on fluid-sensitive and contrast-enhanced MRI pulse sequences. Cartilage-sensitive pulse sequences such as gradient-echo (GRE) and double-echo steady-state (DESS) images facilitate excellent contrast between the bright cartilage and the dark bone. They can identify and estimate percentage of growth plate closure that directly impacts treatment planning. Although small and centrally located transphyseal bridges in children near skeletal maturity can be treated conservatively, larger, multifocal, and/or eccentrically located bridges in younger skeletally immature children will require serial imaging and potentially surgical intervention (►Fig. 8).

Musculoskeletal Ultrasonography
Cartilage is hypoechogenic on US, but distinction between articular, growth, and growth plate cartilage is not possible and only inferred based on their expected anatomical location. At birth, the diaphyses and metaphyses are ossified, preventing further sonographic penetration. However, the epiphyses and apophyses vary in composition, depending on anatomical location, ranging from completely cartilaginous to partially ossified. Partially ossified epiphyses and apophyses allow variable sonographic penetration that is greater among younger children with smaller body parts and more unossified cartilage. Contralateral comparison, multifocal assessment, and dynamic assessment are possible with US and without the need for patient sedation, making it an ideal complementary imaging modality of choice among younger children. The main drawbacks of US are operator dependence and the current lack of consensus protocols for imaging.
Growth-Related Imaging Features

Unique features of the immature skeleton commonly encountered during clinical practice are laminar growth plate appearance, growth recovery lines, transphyseal tethering, normal patterns of physiologic marrow conversion, and stress-related reconversion.

Laminar Growth Plate Appearance

The bilaminar appearance of the growth plate proper on fluid-sensitive MRI includes the hyperintense cartilaginous portion and the hypointense thinner band of ZPC. On higher spatial resolution MRI, a trilaminar appearance of the growth plate can be observed, typically underlying the larger epiphyses. This is due to a third, less well-defined hyperintense layer, located at the ossification front of the primary spongiosa, abutting the ZPC and containing a few residual chondrocytes, just before complete replacement by bone. Due to the relatively smaller size of the secondary growth plates, only a bilaminar appearance is typically observed (Figs. 2 and 4).

Growth Recovery Lines

Growth recovery lines (also known as growth arrest lines) represent thickened, horizontally oriented trabeculae that form because of a temporary slowdown of longitudinal growth during a time of stress. These lines are surrogate markers that exclude premature physeal closure. They are thickest and most obvious when they are newly formed and within more rapidly growing ossification fronts, such as those found in the bones of the distal forearm and around the knee joint. Over time, these lines are displaced toward the diaphysis and finally disappear with progressive bony remodeling.

Symmetrical growth results in lines that parallel the contour of their respective growth plates. Angled or obliquely oriented growth recovery lines indicate underlying endochondral ossification dysfunction, potential tethering, and/or progressive acquired angulation deformity (Fig. 9). Isolated growth recovery lines in the fibula or the ulna indicate an increased possibility of diffuse premature growth plate closure of the ipsilateral tibia and radius, respectively. Growth recovery lines are typically hypointense on all MRI images and best visualized against a background of hyperintense yellow marrow on non–fat-saturated images (Fig. 8). MRI outperforms radiographs in the earlier detection of these lines by 6 to 7 weeks.

Transphyseal Tethering

Heterogeneous signal intensity and irregular morphology of the growth plate proper can suggest endochondral ossification dysfunction and precede premature physeal closure that can be focal or diffuse. Transphyseal tethering can be osseous or fibrocartilaginous (nonosseous), and the latter can be missed on radiographs and CT. Thus MRI is the preferred imaging modality for the diagnosis and estimation of clinically-suspected transphyseal bridge, helping to guide treatment decision making. Large osseous sites of tethering often contain fatty marrow signal intensity and thus are best depicted on T1-weighted images. In contrast, small bony and nonosseous sites of tethering can have variable signal intensity and may be uncovered owing to the presence of adjacent bone marrow edema produced in response to relatively increased regional tissue rigidity.

Although radiographs are helpful in the initial screening of premature physeal closure and serial monitoring of asymmetric growth, they should not be used to estimate the underlying severity or extent of transphyseal tethering, which can be overestimated on radiographs in more than a third of cases when compared with MRI. Finally, these pathologic sites of tethering should not be confused with a focal periphyseal edema (FOPE) zone that occurs at stereotypical sites of early physiologic growth plate closure. These FOPE zones are best studied and further described in the article on the pediatric knee in this issue.

Bone Marrow Conversion and Reconversion

Normal marrow signal intensity changes with growth, maturation, and in response to stress. A comprehensive review of the pediatric marrow is beyond the scope of this article but has been previously described. Briefly, on MRI, signal intensities of the normal marrow are best visualized against a background of hyperintense yellow marrow on non–fat-saturated images. MRI outperforms radiographs in the earlier detection of these lines by 6 to 7 weeks.
logic marrow must be suspected. Than that observed within the subjacent metaphyses, pathologic signal intensity within the epiphyses and apophyses is lower. At C24 exact and opposite direction from normal physiologic conversion, from yellow back to red marrow occur in the long bone (femur). In the pre- and perinatal period, red marrow (red) predominates within the ossified diaphyses and metaphyses while the epiphyses and apophyses remain completely or predominantly cartilaginous (blue). During early childhood, marrow conversion to yellow marrow (yellow) starts within the mid-diaphysis and progresses bidirectionally toward the metaphyses, slightly faster distally than proximally, so that in late adolescence and throughout adulthood, red marrow is mostly found clustered within the proximal metaphyses. Illustration adapted and modified from Chan et al29 with permission.

Intensity of normal marrow reflects a combination of the distribution of marrow elements (hematopoietic red and fatty yellow marrow) and regional marrow edema. Physiologic conversion from red to yellow marrow occurs throughout life but at a more rapid pace during childhood. In long bones, marrow conversion starts in the mid-diaphysis and bidirectional conversion extends toward the metaphyses, slightly faster in the distal than the proximal metaphyses. By skeletal maturity and early adulthood, residual red marrow is mostly observed within the axial skeleton and proximal metaphyses of the appendicular skeleton (→ Fig. 10).

In children who experience systemic stress, marrow conversion from yellow back to red marrow occur in the exact and opposite direction from normal physiologic conversion. This reversion starts within the metaphyses and progresses toward the diaphysis, epiphyses, and apophyses. At ~ 6 months after the appearance of the SOC, marrow signal intensity follows that of yellow marrow. Thus when signal intensity within the epiphyses and apophyses is lower than that observed within the subjacent metaphyses, physiologic marrow must be suspected.17 Imaging features that favor red marrow over pathologic marrow replacement or infiltration include higher signal intensity relative to skeletal muscle (due to intrinsic fatty elements), feathery margins that interdigitate with adjacent yellow marrow, relative bilaterality, and absence of mass effect.29

Acute Physeal Fracture
Growth plates are two to five times weaker than the surrounding soft tissues, making them the “weak link” and prone to injury.30 The primary growth plates underlying pressure epiphyses are the best studied in the published literature, and ~ 18% of all pediatric fractures involve the growth plate with the peak incidence of these fractures overlapping with the adolescent growth spurt.31,32 This overlap is postulated to be the result of a combination of factors, such as temporary bone fragility where longitudinal growth outpaces bone mineralization, the ability to generate greater force, and the involution of the perichondral sleeve that stabilizes the osteochondral interface.14,33–35 Moreover, the relatively lower stiffness of the hypertrophic chondrocytes, in combination with their large size, renders them particularly vulnerable to injury from excessive mechanical loading.36

The Salter-Harris classification of physeal fractures remains the system most widely used clinically for describing these fractures. Type I fractures are uncommon, accounting for < 10% of growth plate fractures, and transverse fracture cleavage planes are centered within the growth plate. Type II fractures are most common, accounting for about three quarters of growth plate fractures, and fracture cleavage planes involve the metaphysis. Types III and IV fractures are uncommon intra-articular fractures that involve the epiphysis, or both the epiphysis and metaphysis, respectively. These latter fractures often require preoperative CT to determine precise fracture alignment and the need for surgical reduction. Relatively rare type V fractures, postulated to involve compression injury, are only diagnosed retrospectively after the development of growth disturbance.6,19,26

Physeal Overuse Injury
In contrast to high-energy acute insults that often lead to fractures and/or dislocations, the growing participation in pediatric sports with higher intensity training and earlier specialization has led to increased rates of overuse injuries that result from repetitive microtrauma at submaximal stress.37 It is postulated that in these skeletally immature athletes, this stress initially and predominantly temporally compromises vascular perfusion within the ossification front, leading to prolonged survival of hypertrophic chondrocytes and lengthening of the fragile chondrocyte columns, producing growth plate widening.38,39

Laor and colleagues38 originally proposed two distinct patterns of chondrocyte extension into the ossification front: a focal “tongue” of persistent cartilage from focal vascular insult versus a “broad band” from a more diffuse insult. The latter diffuse pattern is more commonly encountered among these pediatric athletes.41 This pattern carries an overall favorable prognosis and is often reversible if diagnosed and treated early, allowing restoration of regional perfusion.8 However, if undiagnosed or if the patient is noncompliant with conservative treatment, the biomechanically relatively weaker osteochondral junction is vulnerable to additional insults and stresses, increasing the risk for progressive injury that can affect the growth plate proper and adjacent growth cartilage, which is less reversible and can lead to permanent growth disturbance.8,41 (→ Fig. 3).

The following section discusses our emerging understanding of the pathophysiology that underlies physeal
stress injury, divided into those injuries that involve the epiphyseal, apophyseal, and secondary growth plate complexes.

**Epiphyseal Growth Plate Complex**

The epiphyseal growth plate complexes, including the primary growth plate proper and subjacent epiphysis and metaphysis, are found at one or both ends of a long bone. For the smaller long bones of the hands and feet (e.g., metacarpals, metatarsals, and phalanges), this complex is found at only one end, whereas for the larger long bones of the appendicular skeleton (e.g., humeri and femurs), this complex is found at both ends of the bone. In the latter larger long bones, the proximal and distal epiphyseal physis often have differential growth rates and thus do not contribute equally to the final length of the underlying bone.

For example, ~ 71% of femoral and 57% of tibial lengthening occur at the distal femoral and proximal tibial primary physis, which account for ~ 37% and 25% of the overall total length of the lower extremity, respectively. This information is critically important because it impacts the potential for regional remodeling and severity of the resulting growth disturbance (Fig. 2). The best studied examples of overuse injury that involve the epiphyseal growth plate complex include the proximal humerus among overhead throwing athletes and the distal radius among gymnasts and platform divers.

**Proximal Humeral Epiphyseal Physeal Complex**

Among baseball pitchers, a high level of external rotational torque is placed on the proximal humerus during the late cocking phase of a pitch. This repetitive microtraumatic shear, torque, or traction on the proximal humeral physeal complex is postulated to damage the most susceptible vertically oriented collagen fibers within the hypertrophic zone and subjacent vascularity within the newly formed bone of the metaphyseal ossification front. Suboptimal throwing technique, excessive throwing volume, high throwing speeds, and playing breaking pitches (e.g., curveballs) can all predispose this physeal complex to cumulative stress and injury. These athletes present with pain during overhead activities that worsen during throws and on physical examination; tenderness is centered over the proximal humerus.

Imaging assessment starts with radiographs, which should include an external rotation view that profiles the anterolateral aspect of the humeral physis, the most common site of early injury. Radiographic findings include growth plate widening, indistinctness or loss of ZPC (reflecting active disease), irregularity and sclerosis within the subjacent metaphysis (reflecting chronicity). Physiologic growth plate closure starts centrally and progresses in a centripetal pattern. When early intervention is successful, healing occurs with normalization of the growth plate width and reconstitution of the ZPC.

**Distal Forearm Epiphyseal Physeal Complexes**

Among gymnasts, repetitive compressive and rotational forces are placed on the distal forearm that preferentially affects the distal radius over the distal ulna because the former has a greater cross-sectional area and is responsible for the bulk of the load transmission across the wrist joint. These athletes typically present with localized pain and tenderness centered over the distal forearm, often without a history of acute trauma. Symptoms increase with load-bearing activities and are relieved with rest.

Imaging assessment begins with radiographs where findings of overuse injury are observed, and can be limited to the distal radius or involve both the distal radius and ulna. In contrast to baseball pitchers where findings are often limited to the dominant throwing arm, findings among these gymnasts are usually bilateral, but one side may be worse and more symptomatic than the other.

MRI findings of overuse injury are often observed, and again marrow edema suggests an active ongoing insult. When early intervention is successful, healing occurs with centripetal ossification of the metaphyseal cartilage, leading to normalization of the growth plate width and reconstitution of the ZPC. However, ongoing stress and recurrent injury can lead to irreversible damage, resulting in stunted growth and long-term deformity. This latter observation is postulated to account for the higher prevalence of ulnar positive variance among gymnasts (Fig. 3).

**Apophyseal Growth Plate Complex**

Apophyses contribute to the morphology at the distal ends of mature bone and serve as attachment sites for tendons and ligaments. Analogous to the epiphyseal primary physis, apophyseal primary physis are weaker than the attaching soft tissues and thus prone to failure that can present as acute avulsion fractures and overuse injuries. The latter apophyses is underrecognized and underdiagnosed, often incidentally discovered after an acute injury. It is postulated that cumulative submaximal repetitive stresses weaken the
already venerable osteochondral junction, requiring less energy to produce a frank fracture. The best studied examples of apophyseal injuries include the medial epicondyle among overhead throwers and tibial tubercle among athletes who perform repetitive kicking and jumping (quadriiceps contraction) motions.53

Medial Epicondyle Apophyseal Physeal Complex

Repetitive valgus stress across the elbow can produce a spectrum of findings (also known as thrower’s elbow). The most frequent site of injury involves the medial elbow. Among skeletally immature athletes, medial epicondyle apophysitis is common, whereas among skeletally mature athletes, soft tissue (ulnar collateral ligament and flexor-pronator mass) injuries predominate.54,55 Patients typically present with medial elbow pain that worsens with throwing.

Imaging assessment again starts with radiographs, which can be normal, and findings may be relatively subtle. But in those immature athletes with prolonged symptoms, findings of overuse injury can include growth plate widening, fragmentation of the medial epicondyle ossification, and regional sclerosis. Medial epicondyle overgrowth from regional hyperemia can occur, leading to advanced skeletal maturation.56 (►Fig. 6).

Superimposed avulsion fractures can either include the entire medial epicondyle apophysis (centered at the primary physis) or only the distal epicondyle that underlies the attachment of the ulnar collateral ligament (centered at the secondary physis).54 MRI is preferred over CT for its overall more comprehensive assessment of the elbow, including the presence of marrow edema (that suggests ongoing active stress) (►Fig. 12), integrity of ulnar collateral ligament and flexor-pronator mass, and any findings of internal derangement within the elbow, such as osteochondritis dissecans (OCD).57,58 further discussed in the section on the secondary growth plate complex.

Tibial Tubercle Apophyseal Physeal Complex

Tibial tubercle develops in four stages: cartilaginous, apophyseal, epiphyseal, and osseous stages. The first cartilaginous stage (at approximately < 8 years of age in girls and < 9 years in boys) exists before the appearance of the secondary ossification centers. During the second, apophyseal stage (~ 8–12 years of age in girls and 9–14 years in boys), ossification centers appear, enlarge, and coalesce, whereas during the third epiphyseal stage (~ 10–15 years of age in girls and 11–17 years in boys), the maturing apophyseal ossification coalesce with the proximal tibial epiphyseal ossification. The final osseous stage (approximately > 15 years of age in girls and > 17 years in boys) is reached after complete physeal closure at skeletal maturity.

The transition from fibrocartilage to columnar physeal cartilage and ultimate physeal closure occur in a proximal to distal direction, predisposing tissue at the distal end of the tibial tubercle to differential stress and the risk for biomechanical failure, particularly near skeletal maturation.59 Currently, there is no consensus regarding whether or not overuse injury (e.g., Osgood-Schlatter disease) predisposes patients to avulsion fractures, but both processes share the same culprit mechanism of forceful quadriiceps contraction (often associated with kicking, sprinting, or jumping) transmitted to the immature tibial tubercle by the attaching patellar tendon.60–62

Osgood-Schlatter disease is generally a clinical diagnosis and patients are treated conservatively.63 But in those with atypical symptoms or concern for superimposed avulsions, imaging assessment starts with radiographs. In contrast to Osgood-Schlatter disease with typical radiographic findings of tibial tubercle fragmentation, distal patellar tendinosis, and/or overlying soft tissue edema,59 tibial tubercle fractures are characterized by fracture extent that can be limited to the anterior cortex of the tuberosity or involve the apophyseal growth plate, epiphyseal growth plate, and extend into the knee joint and/or the posterior proximal tibial metaphysis.61,62 (►Fig. 13). MRI is preferred over CT, particularly for those patients with intra-articular extension, to detect internal derangement, soft tissue injury, and/or entrapment.59,62 Rare complications include growth arrest with genu recurvatum and limb-length discrepancy, particularly among those who have sustained injury at a younger age.61,62

Secondary Growth Plate Complex

The secondary growth plates are responsible for the enlargement of the secondary ossification centers. These peripherally located growth plates grow in a centripetal direction, slowly and progressively replacing the overlying unossified

Fig. 12  Examples of apophyseal primary periphyseal stress injury (PPSI) and avulsion fracture of the medial epicondyle. (a) Axial intermediate-weighted (left) and T2-weighted fat-suppressed MR (middle) images from a 9-year-old boy show irregularity and widening of the medial epicondyle physis (triangle) and surrounding reactive marrow edema (circle). (b) Axial T2-weighted fat-suppressed MR (right) image from a different 12-year-old boy shows anterior displacement of the avulsed medial epicondyle (arrow) from its donor site ( bracket) with more intense reactive marrow edema and an elbow effusion (star).
growth cartilage.\textsuperscript{1} It is postulated that injury to the secondary growth plate complex may be the pathophysiologic mechanism that underlies the development of OCD\textsuperscript{64} (\textsuperscript{\textdagger}Fig. 2). OCD is one of the leading causes of chronic knee pain among children and young adults.\textsuperscript{65} The medial femoral condyle is the most common anatomical location throughout the body and within the knee, thus, the best studied. OCD lesions were classically divided into juvenile (JOCD) and adult lesions according to whether the regional growth plates are open or closed at the time of diagnosis. This is important because whereas symptomatic adult lesions rarely heal spontaneously, patients with JOCD are commonly asymptomatic, and more than half can heal with conservative management.\textsuperscript{66–70} This latter observation has been attributed to the presence of vascularized unossified growth cartilage among skeletally immature children, providing an opportunity for self-repair and healing.\textsuperscript{9}

Radiographic findings include focal subchondral lucency, marginal sclerosis, progeny mineralization, fragmentation, and/or intra-articular bodies. MRI is the preferred next imaging modality of choice to characterize lesion stability and to monitor treatment response, helping optimize the decision and timing for surgical intervention. Although several MRI features have been proposed,\textsuperscript{68,71,72} in the knee, the most clinically used criteria are the presence of an osteochondral defect (empty crater), overlying chondrolysis, disruption of the radius of curvature, fluid-like high signal intensity at the interface, cysts (circumferential or a single cyst > 5 mm), and/or multiple breaks in the subchondral bone plate\textsuperscript{72,73} (\textsuperscript{\textdagger}Fig. 14).

Less commonly, OCD lesions are found within the elbow joint that disproportionately involve the capitellum,\textsuperscript{74,75} particularly among baseball pitchers and gymnasts. Repetitive valgus overload with the elbow in flexion is believed to cause these lesions among baseball players, whereas weight bearing with the elbow in extension may be the cause among gymnasts.\textsuperscript{76} Histopathologic analysis previously showed that compressive forces can cause focal endochondral ossification dysfunction at these physiologically active, peripherally located secondary physis.\textsuperscript{64,77–79}

In contrast to medial femoral condyle lesions, capitellar OCD lesions are more likely to be unstable at the time of diagnosis, but it is uncertain whether this is due to either or to a combination of regional differences (more tenuous regional blood supply within the distal humerus)\textsuperscript{80,81} or delayed clinical presentation (\textsuperscript{\textdagger}Fig. 15). On MRI, although the presence of an osteochondral defect, intra-articular body, overlying cartilage changes, subchondral bone plate disruption, and hyperintense rim at the interface were significantly more common among unstable lesions, the number or size of the perilesional cysts did not significantly differ between stable and unstable capitellar OCD when arthroscopy was used as the reference standard.\textsuperscript{82}

Regardless of the anatomical location, treatment for OCD depends on lesion stability. Stable lesions are treated conservatively, but unstable lesions or lesions that have failed a trial of conservative therapy, particularly in those who have reached skeletal maturity, are often treated surgically to prevent further joint damage and reduce the risk for future premature osteoarthritis.\textsuperscript{82}

**Summary**

Growth and maturation occur in a predictable pattern, and endochondral ossification predominates within long bones of the appendicular skeleton, occurring in both primary and secondary growth plate complexes. These growth plate
complexes are sites of relative weakness, especially during pubescence, and they are at risk for acute insult or overuse injuries. A basic knowledge of the normal histoanatomy and physiology of the growth plate complex ensures an improved understanding of the various patterns of injury and shared similarities between different anatomical locations and physeal complexes.

Conflict of Interest
None declared.

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