THE ROLE OF IMAGING IN ADVANCING THE UNDERSTANDING OF THE PATHOGENESIS, DIAGNOSIS AND STAGING OF CENTRAL CHONDROID BONE TUMOURS

By

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There are several other people whose support is much appreciated, and I hope that this sentence gives due reference of my gratitude to them.

Finally, I would like to thank the patients, without them none of this research would have been possible. To them I owe a lifelong debt of gratitude.

Dedication

I would like to dedicate this work to my beloved wife Mariam Jafri and my two sons Yusuf and Idris Douis who despite the little time I devoted to them are always on my mind and in my heart.
Declaration of Authorship

This thesis is submitted to the University of Warwick in support of my application for the degree of Doctor of Philosophy. It has been composed by myself and has not been submitted in any previous application for any degree.

The work presented (including data generated and data analysis) was carried out by the author.

Signed:                                      Date: 02.11.2017

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6) Revision of manuscript

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2) Data review
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Contribution:
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Chapter 7: Is bone scintigraphy necessary in the initial surgical staging of chondrosarcoma of bone?

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Signature of authors:
Abstract

Central chondroid bone tumours are one of the most common primary bone tumours. Benign central chondroid tumours are termed enchondromas and its malignant counterpart are called chondrosarcomas. Enchondromas are frequently observed on routine imaging. Similarly, chondrosarcomas are the second most common primary bone tumour after osteosarcoma. Imaging is crucial in the diagnosis of central chondroid tumours and in the differentiation of enchondromas from chondrosarcomas. Furthermore, imaging plays a vital role in the staging of chondrosarcomas. In this thesis, the published scientific literature on the role of imaging in the diagnosis of benign chondroid tumours and chondrosarcomas and the role of imaging in the staging of chondrosarcomas is reviewed and summarised. Furthermore, the contribution of the authors’ published work is highlighted in the thesis.

The first two articles are review articles which discuss the clinical and imaging features of benign and malignant chondrogenic tumours and the significance of imaging in the diagnosis of these tumours.

The third article is an original article which investigates the theory of the pathogenesis of enchondromas. It is widely believed that enchondromas arise from cartilage islands which are displaced from the growth plate during the process of skeletal maturation. However, this theory is unproven, and the origin of this theory was forgotten prior to the authors’ study. Based on the incidental prevalence of enchondromas of the knee in the adult population of 2.9%, the study assesses the prevalence of cartilage islands/ enchondromas in skeletally immature patients. In this
study, no cartilage islands/enchondromas in skeletally immature patients were identified. The study therefore shows the rarity of enchondromas in skeletally immature individuals which is in contrast to the adult population. Furthermore, in view of the absence of cartilage islands in this study, the study raises doubts about the validity of the unproven theory. Lastly, the very origin of this theory is rediscovered in this thesis which has been forgotten in modern medicine.

The fourth article is an original article which evaluates the role of diffusion-weighted MRI (DWI) in the diagnosis of central cartilage tumours. Prior to the authors’ study the role of DWI in the diagnosis of central cartilage tumours was uncertain. The authors’ study demonstrates that DWI cannot be used to differentiate between enchondromas and chondrosarcomas and that DWI does not aid in the distinction of low-grade chondroid tumours from high-grade chondrosarcomas. This is a finding which was not known prior to the study.

The fifth article is an original article which assesses the utility of conventional MRI in the differentiation of low-grade from high-grade chondrosarcomas of long bone. Prior to the authors’ study the role of conventional MRI in the differentiation of low-grade from high-grade chondrosarcomas of long bone was unknown. The authors’ study shows that bone expansion, active periostitis, soft tissue mass and tumour length can be used to differentiate high-grade from low-grade chondral lesions of long bone on conventional MRI. Furthermore, the presence of these four MRI features shows a diagnostic accuracy of 95.6%. These findings were not known prior to the study and have significantly furthered the knowledge about the role of conventional MRI in the grading of chondrosarcoma of long bone.
The sixth article is an original article which evaluates the role of bone scintigraphy and Computed Tomography of the chest in the staging of chondrosarcoma of bone. Whilst guidelines regarding the staging of bone sarcomas state that bone scintigraphy should be performed to assess for the presence of skeletal metastases and that Computed Tomography (CT) of the chest should be performed to evaluate for possible pulmonary metastases, there has been no research on the utility of bone scintigraphy in chondrosarcoma of bone and on the role of CT-chest in the staging of chondrosarcomas. Furthermore, the prevalence of skeletal and pulmonary metastases of chondrosarcoma at presentation was unknown prior to this study. The authors’ study demonstrated no skeletal metastases on bone scintigraphy in chondrosarcoma of bone at presentation. In contrast, pulmonary metastases were observed in approximately 5% of all patients with chondrosarcoma at presentation on CT-chest. The finding therefore demonstrates the rarity of skeletal metastases in chondrosarcoma of bone at presentation which is in contrast to osteosarcoma and Ewing sarcoma. The study therefore concludes that there is little role for skeletal scintigraphy in the surgical staging of chondrosarcoma. In contrast, the study shows that there is a role for CT-chest in the staging of chondrosarcoma. These above described findings are important new findings and represent a significant contribution to the knowledge base regarding metastatic behaviour of chondrosarcomas at presentation and regarding the staging of chondrosarcoma of bone.
In summary, the authors’ publications have significantly enhanced and furthered the understanding of the pathogenesis of enchondromas, the role of functional MRI in the differentiation of enchondromas from chondrosarcomas, the utility of MRI in the grading of chondrosarcomas and the role of skeletal scintigraphy in the staging of chondrosarcomas.
Abbreviations

ADC = Apparent diffusion coefficient

AP radiograph = Antero-posterior radiograph

CLUMP = cartilaginous lesion of unknown malignant potential

CT = Computed Tomography

DCE-MRI = Dynamic contrast-enhanced Magnetic Resonance Imaging

DWI = Diffusion-weighted Imaging

FDG-PET/CT = Fluorodeoxyglucose-Positron Emission Tomography/Computed Tomography

IDH = Isocitrate dehydrogenase

MRI = Magnetic Resonance Imaging

PDWI = Proton density weighted Imaging

SLICED group = Skeletal Lesions Interobserver Correlation among Expert Diagnosticians Study Group

SUV = Standardized uptake value

T1WI = T1-weighted Imaging

T2WI = T2-weighted Imaging

WHO = World Health Organization

2-HG = 2-Hydroxyglutarate

99m-Tc MDP bone scintigraphy = 99 metastable-Technetium Methylene diphosphonate
Foreword

Primary bone tumours are a heterogeneous group of tumours, which originate in bone or from bone-derived cells. Whilst benign primary bone tumours are relatively common, malignant primary bone tumours (bone sarcomas) represent a rare disease entity. Primary bone tumours are classified according to the World Health Organization Classification of Tumours dependent on their histological composition. Chondrogenic tumours represent a subgroup of bone tumours, which demonstrate cartilaginous differentiation and are amongst the most common primary bone tumours. Enchondroma is a benign bone tumour of hyaline cartilage which represents the second most common benign bone tumour after osteochondroma whilst its malignant counterpart, the chondrosarcoma is a malignant cartilaginous matrix-producing neoplasm which represents the second most common sarcoma of bone after osteosarcoma. Imaging plays a pivotal role in the detection, diagnosis and staging of bone tumours in general and in chondrogenic bone tumours in particular. This is highlighted by the fact that unlike in carcinomas where histopathological evaluation is the gold standard in the diagnosis of tumours, the diagnosis of bone tumours is reliant on a consensus between histopathology and imaging. This is of particular importance in chondrogenic tumours where the differentiation of benign chondrogenic tumours (enchondromas) from malignant chondrogenic tumours (chondrosarcomas) is one of the most challenging diagnoses in orthopaedic oncology and is based on a consensus between clinical findings, imaging and histopathology. Imaging, may it be Radiography, Computed Tomography, Magnetic Resonance Imaging, Skeletal Scintigraphy and more recently Positron Emission Tomography has
revolutionized oncology including the care of patients with sarcoma as it facilitates early detection, diagnosis and staging of cancers thereby significantly contributing to the marked improvement in survival of oncology patients which has been observed over the last 40 years.

Although there is a plethora of evidence supporting the role of imaging in the diagnosis, grading and staging in a wide variety of cancers, the evidence in chondrosarcomas has been very limited. In particular, there has been a paucity of evidence to support the use of functional imaging techniques in the diagnosis of chondrosarcomas.

This thesis discusses four original studies and two review articles published by the author of this thesis which evaluate;

1) the significance of imaging in benign and malignant chondrogenic tumours in general
2) the contribution of MR Imaging in understanding the pathogenesis of enchondromas
3) the role of functional MRI in the diagnosis of chondrosarcomas
4) the significance of conventional MRI in the grading of chondrosarcomas
5) the utility of skeletal scintigraphy in the staging of chondrosarcomas in particular

All original articles and review articles have been published in peer-reviewed journals and all original articles represent significant contributions to the knowledge regarding the pathogenesis of enchondromas, the diagnosis, grading and staging of chondrosarcomas. Furthermore, the hereby presented original articles about the role
of imaging in the diagnosis, grading and staging of chondrosarcomas have resulted in significant changes in patient management. Lastly, the authors’ work, which is being presented for this thesis, has been quoted in multiple national and international conferences and has been cited in multiple peer-reviewed publications and books. The publications form part of the research portfolio of the applicant and are closely related in that they are all imaging research studies on primary chondrogenic bone tumours. The introduction aims to provide both an overview of the topic and a summary of the publications presented describing the role of imaging in the diagnosis, grading and staging of chondrogenic bone tumours prior to the authors’ published work. Furthermore, the limitations of imaging in the diagnosis and in the work-up of patients with chondrogenic tumours are being discussed. Subsequently, the presented work of the author demonstrates the significance and contribution of the authors’ published work in understanding the pathogenesis of enchondromas, the role of imaging in the diagnosis, grading and staging of chondrosarcomas. In the conclusion, the author discusses the potential future role of other advanced imaging techniques in the diagnosis of enchondromas and chondrosarcomas and in the potential role of treatment response in chondrosarcomas.
Chapter 1: Introduction

1.1. Primary bone tumours

Primary bone tumours are divided into benign primary bone tumours, intermediate and malignant primary bone tumours. Whilst benign primary bone tumours are relatively common, intermediate and malignant primary bone tumours are rare disease entities. The true incidence of benign primary bone tumours is unknown because many benign bone neoplasms are clinically indolent and are frequently only depicted as incidental findings on imaging performed for unrelated causes. In contrast, primary malignant bone neoplasms (sarcomas) are rare and account for approximately 0.2% of all neoplasms with an annual incidence of approximately 0.8 per 100000 population.\textsuperscript{1-3}

According to the World Health Organization, primary bone tumours are classified dependent on histopathology. Hence, we differentiate chondrogenic, osteogenic, fibrogenic, fibrohistiocytic, haematopoietic, osteoclastic giant cell rich, notochordal, vascular, myogenic, lipogenic tumours, tumours of undefined neoplastic nature and miscellaneous tumours. The various histological subtypes of primary bone tumours are then further classified dependent on their biological behaviour as benign, intermediate and malignant (Table 1).\textsuperscript{4}
Table 1: WHO classification of bone tumours (adapted from World Health Organization Classification of Tumours of Soft Tissue and Bone, 2013)

<table>
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<th>CHONDROGENIC TUMOURS</th>
<th>FIBROGENIC TUMOURS</th>
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<tr>
<td>Benign</td>
<td>Intermediate (locally aggressive)</td>
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<tr>
<td>Osteochondroma</td>
<td>Desmoplastic fibroma of bone</td>
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<td>Chondroma</td>
<td>Malignant</td>
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<td>Enchondroma</td>
<td>Fibrosarcoma of bone</td>
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<td>Periosteal chondroma</td>
<td>FIBROHISTIOCYTIC TUMOURS</td>
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<td>Osteochondromyxoma</td>
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<td>Subungual exostosis</td>
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<td>Bizarre parosteal osteochondromatous proliferation</td>
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<td>Synovial chondromatosis</td>
<td>Plasma cell myeloma</td>
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<td>Intermediate (locally aggressive)</td>
<td>Solitary plasmacytoma of bone</td>
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<td>Chondromyxoid fibroma</td>
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<td>Atypical cartilaginous tumour/Chondrosarcoma grade 1</td>
<td>GIANT CELL RICH TUMOURS</td>
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<tr>
<td>Intermediate (rarely metasatizing)</td>
<td>Benign</td>
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<tr>
<td>Chondroblastoma</td>
<td>Giant cell lesion of small bones</td>
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<tr>
<td>Malignant</td>
<td>Intermediate (locally aggressive, rarely metasatizing)</td>
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<td>Chondrosarcoma grade 2 and 3</td>
<td>Giant cell tumour of bone</td>
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<td>Dedifferentiated chondrosarcoma</td>
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<td>Low-grade central osteosarcoma</td>
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<td>MYOGENIC</td>
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<td>Intermediate (locally aggressive)</td>
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<td>Aneurysmal bone cyst</td>
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<td>Malignant</td>
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<td>Leiomyosarcoma of bone</td>
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<td>Undifferentiated high-grade pleomorphic sarcoma</td>
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Benign bone tumours are characterized by a limited capacity of local recurrence whilst intermediate tumours often recur and are associated with an infiltrative and locally destructive growth pattern but do not metastasize or rarely metastasize. In contrast, malignant bone tumours (termed bone sarcomas) in addition to demonstrating a locally destructive growth and recurrence also have a significant risk to metastasize which ranges from approximately 20% to 100%.\textsuperscript{1,4} Chondrosarcomas, osteosarcomas, leiomyosarcomas and fibrosarcomas of bone are further graded based on their histopathological features such as relative proportion of cells to matrix, nuclear atypia of tumour cells, irregularity of nuclear contour, enlargement and hyperchromasia of nuclei, mitotic figures and necrosis. The two histopathological grading systems which are most widely adopted are the two-tier system which divides malignant tumours as low-grade and high-grade and the three-tier system which divides malignant bone tumours into grade 1, grade 2 and grade 3 malignant bone tumours.\textsuperscript{1}

1.2 Chondrogenic tumours

Chondrogenic bone tumours are tumours which consist of cartilage and they represent the second largest group of bone tumours after osteogenic bone tumours. Benign chondrogenic bone tumours include osteochondroma, enchondroma, periosteal chondroma, chondromyxoid fibroma, osteochondromyxoma, bizarre parosteal osteochondromatous proliferation, synovial chondromatosis and the chondroblastoma. In contrast, malignant chondrogenic tumours include chondrosarcomas (grade 1-3), periosteal chondrosarcoma, dedifferentiated
chondrosarcoma, mesenchymal chondrosarcoma and clear cell chondrosarcoma (see Table 1). In the following sections, I will focus on one of the most common benign cartilage tumours, the enchondroma and its malignant counterpart, the central chondrosarcoma.

1.2.1 Enchondroma

Enchondroma is a benign hyaline cartilage neoplasm which arises in the medullary cavity.

1.2.1.1 Epidemiology of enchondromas

Enchondromas are relatively common and account for approximately 10-25% of all surgically removed benign bone tumours. The true incidence is significantly higher because many enchondromas are asymptomatic and are frequently only discovered incidentally. This has been highlighted in two previous MRI-studies which discovered that the incidental prevalence of enchondromas on routine MRI-examinations of the knee is 2.9% and the incidental prevalence of enchondromas on routine MRI-examinations of the shoulder is 2.1%. In contrast, enchondromatosis, which is a group of skeletal dysplasias characterized by multiple enchondromas and which includes Ollier’s disease and Maffucci syndrome are rare disease entities. An in-depth discussion of enchondromatosis is beyond the remit of this thesis. I will therefore focus on solitary enchondromas in this thesis unless particularly stated.
Enchondromas demonstrate a wide age distribution which ranges from 5 years to 80 years although the average age at time of diagnosis is 40 years.\textsuperscript{9,10}

### 1.2.1.2 Location and clinical presentation of enchondromas

The most common location of enchondromas are the long tubular bones of the hand (40-65\% of all enchondromas) followed by the major long bones (approximately 25\% of all enchondromas) with the femur being the most common location within the long bones followed by the humerus and tibia. Furthermore, 7\% of all enchondromas are located within the feet. In contrast, enchondromas are rare in the flat bones such as the pelvis, spine, ribs, scapula and sternum.\textsuperscript{9} Whilst enchondromas in the hands and feet may present clinically with palpable swelling or a pathological fracture, enchondromas of the long tubular bones are often asymptomatic and are therefore frequently detected incidentally.\textsuperscript{6,7,9,10}

### 1.2.1.3 Histology of enchondromas

On histopathology, enchondromas are hypocellular, avascular hyaline tumours, which demonstrate abundant hyaline cartilage without cellular atypia (Figure 1). Endosteal erosion may be present in some cases however an important differentiating feature from chondrosarcomas is the lack of entrapment of the host bone by tumour cells. In contrast to enchondromas of long bone, enchondromas of the hands and feet can be more cellular and may demonstrate cellular atypia.\textsuperscript{11}
1.2.1.4 Pathogenesis of enchondromas

The pathogenesis of enchondromas is poorly understood. Although it is widely believed that enchondromas arise from cartilage remnants, which have been displaced from the growth plate during the process of skeletal maturation, this widely held belief is unproven and the origin of this theory is unknown in modern medicine. In chapter 4 of this thesis, I investigate the origin of this theory unravelling unexpected and surprising findings into the very origin of this theory and
critically appraise the origin of this theory. Furthermore, we investigate the theory using MRI.

1.2.1.5 Imaging of enchondromas

Imaging is crucial in the identification and diagnosis of enchondromas. Enchondromas of long bones are usually located centrally or eccentrically in a metadiaphyseal location whilst epiphyseal involvement is only observed in 2-5% of all enchondromas of long bones. On radiography, enchondromas usually appear as well-defined lytic lesions within the medullary cavity which demonstrate a geographic pattern of bone destruction. Approximately 95% of all enchondromas in the long bones demonstrate some degree of matrix calcification which is typically referred to as ring-and-arc or popcorn calcification. Enchondromas are usually less than 5cm in maximum length, demonstrate only minimal endosteal scalloping and minimal cortical thickening (Figure 2). Periosteal reaction, bone expansion and cortical destruction are not features of central intramedullary enchondromas of long bones and if present are suspicious for the development of a chondrosarcoma. In contrast to enchondromas of long bones, enchondromas in the hands and feet frequently demonstrate extensive endosteal scalloping, bony expansion and cortical thinning (Figure 3). These findings should therefore not be used to raise the suspicion for the development of a chondrosarcoma in the hands and feet. Furthermore, it may be difficult to demonstrate a calcified matrix in enchondromas of the hand unlike in enchondromas of the long bones. In the absence of trauma however, cortical disruption and periosteal reaction are unusual in enchondromas of the hands and
feet and therefore raise the suspicion of malignant transformation into a chondrosarcoma.\textsuperscript{10,13}

Figure 2: Radiograph of enchondroma of the proximal tibia: Anteroposterior radiograph of the knee demonstrates a 4.3cm in the maximum cranio-caudal dimension measuring enchondroma within the proximal tibia which shows chondroid matrix calcification (arrow) but no endosteal scalloping, cortical destruction or periosteal reaction.
Figure 3: Radiograph of enchondroma of the hand: Anteroposterior radiograph of the little finger demonstrates an expansile radiolucent lesion of the middle phalanx which shows marked endosteal scalloping, cortical thinning and bony expansion with subtle foci of matrix calcification within it (arrow). The imaging findings are in keeping with an enchondroma.
Computed Tomography is useful in demonstrating radiographically occult matrix calcification in enchondromas (Figure 4) and is the imaging modality of choice in the assessment of the degree of endosteal scalloping if present.10,13

Figure 4:

Figure 4: Computed Tomography of enchondroma of the femur. Sagittal CT of an enchondroma of the distal femur demonstrates extensive chondroid matrix calcification (arrow) but no endosteal scalloping.
On MRI, enchondromas of long bones typically demonstrate a well-defined lobular contour, with the lesion appearing as intermediate signal intensity on the T1-weighted images and of increased signal intensity on the T2-weighted images in keeping with hyaline cartilage (Figure 5a,b). Small foci of high signal intensity on the T1-weighted images may be observed within the lesion (Figure 5a) and are thought to represent engulfed normal yellow marrow. On MRI, matrix calcification if present appears as punctate or curvilinear foci of signal void both on T1-weighted images and fluid-sensitive sequences (Figure 6a,b). Thin septa which are of low-signal intensity on the T2-weighted images, are often seen between lobules of cartilage and demonstrate fibrovascular septa which enhance after gadolinium administration (Figure 6c).
Figure 5: MRI of enchondroma of the distal femur: Coronal T1W SE MRI (a) and coronal T2W FS MRI (b) of an enchondroma of the distal femur shows a well-defined lobulated lesion which is of low to intermediate signal intensity on the T1-weighted images (a, small arrow) and of increased signal intensity on the T2 fat-suppressed images (b, small arrow). There are small foci of high signal intensity on the T1-weighted images, (a, large arrow) which represent foci of engulfed normal yellow marrow.
Figure 6:

Figure 6a:  

Figure 6b:
Figure 6: MRI of enchondroma of the distal femur. Coronal T1W SE MRI (a) and coronal T2W FS MRI (b) of an enchondroma of the distal femur demonstrates a 4.9cm chondroid lesion which shows foci of signal void both on the T1-weighted image (a, arrow) and on the T2 fat-suppressed images (b, arrow) in keeping with matrix calcification. Gadolinium-enhanced fat-suppressed T1-weighted MRI examination of the same lesion shows avid peripheral and septal enhancement (c, arrow) as well as nodular enhancement.
On 99m-Technitium methylene diphosphonate (MDP) bone scintigraphy, enchondromas usually demonstrate mild to moderate homogeneous radio-isotope uptake which is equal or less than the radio-isotope uptake observed within the anterior iliac crest (Figure 7a,b). However, approximately 30% of all enchondromas demonstrate avid heterogeneous radio-isotope uptake on 99m-Technitium methylene diphosphonate (MDP) bone scintigraphy which is greater than the uptake observed in the anterior iliac crest (Figure 8).9,10,13,15 Hence, this finding cannot be reliably utilized to differentiate enchondromas from low-grade chondrosarcomas.
Figure 7: MRI and bone scintigraphy of enchondroma of the left distal femur. Coronal T1W SE MRI of the left femur (a) demonstrates a 4.5cm in the maximum cranio-caudal dimension measuring enchondroma in the distal femur. On 99m-Technitium methylene diphosphonate (MDP) bone scintigraphy, the lesion demonstrates only mild homogeneous radio-isotope uptake (b, black arrow) which is less than the radio-isotope uptake observed within the anterior iliac crest.
Figure 8: Bone scintigraphy of enchondroma of the left distal femur. 99m-Technitium methylene diphosphonate (MDP) bone scintigraphy of the enchondroma of the distal femur seen in figure 6 demonstrates avid radio-isotope uptake (black arrow).
On fluoro-deoxyglucose-positron emission tomography (FDG-PET), enchondromas demonstrate mildly increased mean standardized uptake value. However, there is considerable overlap in the SUVmax values between enchondromas and low-grade chondrosarcomas resulting in a poor specificity when SUVmax values fall between 2 and 4.5 which is observed in approximately 46% of all benign and malignant chondroid lesions.\textsuperscript{16,17}

1.2.1.6 Management of enchondromas

Unlike chondrosarcomas which always require surgical intervention unless surgery is contraindicated, the management of enchondromas is dependent on patient symptoms. Asymptomatic enchondromas do not require treatment. In contrast, symptomatic enchondromas can be safely treated with curettage.\textsuperscript{18}

1.2.2 Chondrosarcoma

1.2.2.1 Epidemiology of chondrosarcomas

Chondrosarcomas are the second most common malignant primary bone tumour after osteosarcoma accounting for approximately 20% of all malignant bone tumours and for 3.5% of all biopsied primary bone tumours.\textsuperscript{14} According to the WHO Classification of bone tumours, chondrosarcomas are defined as “a locally aggressive or malignant group of cartilaginous matrix-producing neoplasms with diverse morphological features and clinical behaviour”.\textsuperscript{19} Chondrosarcomas are divided into
conventional chondrosarcomas, dedifferentiated chondrosarcomas, mesenchymal chondrosarcomas and clear cell chondrosarcomas. Conventional chondrosarcomas are further divided dependent on location and origin into primary central chondrosarcomas, secondary central chondrosarcomas, secondary peripheral chondrosarcomas and periosteal chondrosarcomas. Primary central chondrosarcomas are chondrosarcomas, which arise centrally in bone without a benign precursor whilst secondary central chondrosarcomas are central chondrosarcomas, which arise in a pre-existing enchondroma. In contrast, secondary peripheral chondrosarcomas are chondrosarcomas which arise from an osteochondroma whilst periosteal chondrosarcomas originate from the periosteum and hence occur on the surface of bone. Although the above described classification reveals a significant variety of chondrosarcomas, primary central chondrosarcoma is the most common subtype of chondrosarcoma accounting for approximately 85% of all chondrosarcomas. In this thesis, I will therefore focus on “primary central chondrosarcoma” and will refer to it as “chondrosarcoma” unless otherwise stated.

### 1.2.2.2 Location and clinical presentation of chondrosarcomas

Most patients diagnosed with chondrosarcoma are in their 6th decade of life. Although the lesion can arise in any bone which is derived from endosteal ossification, the most common sites in the skeleton are the pelvis, followed by the proximal femur, proximal humerus, distal femur and ribs. In contrast,
chondrosarcomas of the hands and feet are rare. Similarly, chondrosarcoma in the spine and the craniofacial bones are a rare occurrence.

The most frequent clinical presentations are pain and local swelling which are usually present for months prior to diagnosis. Pathological fracture is observed in 3-17% of all chondrosarcomas.

1.2.2.3 Histology of chondrosarcomas

On macroscopy, chondrosarcomas have a translucent, lobular blue-gray or white surface which corresponds to hyaline cartilage. Yellow-white foci are frequently identified within the lesion and represent areas of mineralization. Cortical erosion and destruction, bone expansion and an associated soft tissue mass may be seen (Figure 9).

On histology, the tumour is composed of irregularly shaped lobules of hyaline cartilage which vary in size and shape. The chondrocytes demonstrate atypia, show enlarged hyperchromatic nuclei and frequently demonstrate myxoid change. The presence of host bone entrapment (= permeation of cortical and/or medullary bone) is diagnostic of a chondrosarcoma (Figure 10). In cases where permeation is not identified but cellular atypia is present, the diagnosis of a low-grade chondrosarcoma cannot be reliably made however a low-grade chondrosarcoma cannot be excluded. The classification of these low-grade chondroid tumours remains controversial. Whilst the latest WHO classification of bone tumours does not mention these lesions as a separate disease entity, they invariably are being classified in some centres as enchondromas, in others as grade 1 chondrosarcomas despite the
lack of a permeative growth pattern whilst other institutions classify them as atypical enchondromas, borderline cartilage tumours, grade 0 chondrosarcomas or as cartilaginous tumours of unknown malignant potential (CLUMPs) (Figure 11).\textsuperscript{19,23-25}

The histological criteria for the diagnosis of a chondrosarcoma of the phalanges are different and reliant on the presence of cortical destruction, soft tissue extension or the presence of mitosis.\textsuperscript{19,20}
Figure 9: Macroscopic pathology of a chondrosarcoma of the humerus. Surgical specimen of a coronally sectioned humerus demonstrates lobular white or blue-grey foci of hyaline cartilage within the medullary cavity of the humerus (long, white arrow). Yellow-white foci (green arrow), which are seen within the lesion represent areas of mineralisation. The lesion results in bone expansion, cortical remodelling (curved white arrow) and a soft tissue mass (small, thick arrow). (Image courtesy of Dr S. Vaiyapuri, Department of Pathology, Royal Orthopaedic Hospital, Birmingham, UK).
Figure 10: Histopathology of a grade 1 chondrosarcoma: The specimen demonstrates the hallmark of a grade 1 chondrosarcoma: the presence of host bone entrapment or permeation (black arrows). (Image courtesy of Dr F. Puls, Department of Pathology, Royal Orthopaedic Hospital, Birmingham, UK).
Figure 11: Histopathology of a cartilaginous lesion of unknown malignant potential (CLUMP). The chondrocytes demonstrate atypia and myxoid change. There is however no host bone entrapment. (Image courtesy of Dr S. Vaiyapuri, Department of Pathology, Royal Orthopaedic Hospital, Birmingham, UK).
Conventional intramedullary chondrosarcomas are graded on a scale from I to III. The grading is based on nuclear size, hyperchromasia, cellularity and mitosis. Grade I chondrosarcomas are moderately cellular and contain hyperchromatic nuclei (Figure 10). Grade II chondrosarcomas are more cellular demonstrating more nuclear atypia, hyperchromasia and nuclear size (Figure 12). Grade III chondrosarcomas show increased cellularity, are more pleomorphic, demonstrate increased mitoses and show spindle shape at the periphery of the cartilage lobules (Figure 13). Grading of chondrosarcomas is important because the grade of the lesion correlates with prognosis and overall survival. This fact has been highlighted in multiple studies, most recently in a large retrospective study which demonstrated that the 10-year survival of grade I chondrosarcomas was 95%, the 10-year survival for grade 2 chondrosarcomas was 86% whilst the 10-year survival for grade 3 chondrosarcomas was 55%. A large study of chondrosarcomas identified that 61% of all chondrosarcomas were grade I chondrosarcomas, 36% were grade II chondrosarcomas whilst only 3% were grade III chondrosarcomas.
Figure 12: Histopathology of a grade 2 chondrosarcoma: The chondrocytes demonstrate more nuclear atypia and hyperchromasia. (Image courtesy of Dr F. Puls, Department of Pathology, Royal Orthopaedic Hospital, Birmingham, UK).
Figure 13: Histopathology of a grade 3 chondrosarcoma: The tumour cells are more pleomorphic and increasingly spindle shaped. (Image courtesy of Dr F. Puls, Department of Pathology, Royal Orthopaedic Hospital, Birmingham, UK).
1.2.2.4 Pathogenesis of chondrosarcomas

The pathogenesis of chondrosarcomas remains poorly understood. However, a previous study has demonstrated that somatic mutations in isocitrate dehydrogenase (IDH) 1 and 2 are frequent events in central chondrosarcomas, central and periosteal chondromas occurring in at least 56% of these tumours. In contrast, osteochondromas, peripheral chondrosarcomas and mesenchymal tumours other than central chondrosarcomas, central and periosteal chondromas do not show this mutation.\textsuperscript{33} Mutations in IDH 1 and 2 result in production of the oncometabolite 2-Hydroxyglutarate (2-HG) which in turn drives tumour progression. The exact mechanism by which accumulation of 2-HG may lead to tumourigenesis remains uncertain. However, increasing evidence suggests a possible epigenetic mechanism. The above described findings therefore suggest a causal role of IDH1 and 2 mutations in tumourigenesis of central chondrosarcomas.\textsuperscript{33} In the future, this discovery may therefore potentially result in the development of drugs which target and effectively treat chondrosarcomas showing IDH 1 and 2 mutations.

1.2.2.5 Imaging of chondrosarcomas

Imaging is crucial in the diagnosis of central chondrosarcomas. On radiography, chondrosarcomas typically demonstrate as mixed lytic and sclerotic lesions with a variable degree of chondroid matrix mineralization which is seen in 60-78% of all chondrosarcomas (Figure 14).\textsuperscript{14,15,19,21}
Figure 14:

Radiograph of low-grade chondrosarcoma of distal femur. Anteroposterior radiograph of the distal femur demonstrates a well-demarcated, geographical pattern of bone destruction (large arrow) within the distal femur with extensive chondroid matrix calcification (small arrow).
The amount of calcification is variable, however high-grade chondrosarcomas demonstrate less chondroid matrix mineralization than low-grade chondrosarcomas. Low-grade chondrosarcomas frequently show a geographical pattern of bone destruction (Figure 14) whilst a moth-eaten, permeative pattern of bone destruction favours a high-grade chondrosarcoma or a dedifferentiated chondrosarcoma (Figure 15).\textsuperscript{10,14}. Erosion of the cortex is termed “endosteal scalloping”. Endosteal scalloping of more than two thirds of the depth of the cortex is a hallmark of chondrosarcoma reflecting its increased biological activity (Figure 15).
Figure 15: Radiograph of a high-grade chondrosarcoma (grade 2 chondrosarcoma) of the femur. Lateral radiograph of the femur shows a moth-eaten, permeative pattern of bone destruction within the mid-diaphysis with marked endosteal scalloping (large arrow) and bone expansion. A relatively small focus of chondroid-matrix calcification (small arrow) is observed within the lesion.
Continued growth of the lesion may result in cortical remodeling, cortical thickening, cortical destruction, periosteal reaction, bony expansion (Figure 16) and the development of a soft tissue mass (Figure 17). Within the long bones, a central chondroid lesion which is larger than 5cm is regarded as suspicious for a chondrosarcoma (Figure 18).

Computed Tomography is the imaging modality of choice in the assessment of endosteal scalloping of chondroid tumours (Figure 19). Furthermore, Computed Tomography is particularly useful in the identification of occult matrix mineralization which is present in 90%-94% of all chondrosarcomas on CT.
Figure 16: Radiograph of a high-grade chondrosarcoma of the proximal femur. Anteroposterior radiograph of the proximal femur shows a mixed lytic, sclerotic lesion (large arrow) which demonstrates cortical remodelling, cortical thickening and bony expansion (small arrow).
Figure 17: Radiograph of a high-grade chondrosarcoma of the proximal femur. Anteroposterior radiograph of the proximal femur shows an ill-defined lesion within the right proximal femur which demonstrates extensive chondroid matrix calcification and which results in cortical destruction and a pathological fracture of the greater trochanter. There is extension of chondroid tissue into the surrounding soft tissue in keeping with an associated large soft tissue mass (arrow).
Figure 18: Radiograph of a low-grade chondrosarcoma of the left proximal humerus. Anteroposterior radiograph of the left proximal humerus shows an 10cm in the maximum cranio-caudal dimension measuring chondroid lesion in the left proximal humerus (arrow) which was a histologically confirmed grade 1 chondrosarcoma. Note the absence of other aggressive features such as bone expansion, cortical destruction or periosteal reaction.
Figure 19: Computed Tomography of a chondrosarcoma of the tibia. Axial CT of the tibia exquisitely demonstrates extensive (more than 2/3) endosteal scalloping (arrow).
Magnetic Resonance Imaging (MRI) most accurately depicts the intraosseous tumour extent, the presence and extent of a soft tissue mass. On MRI, chondrosarcomas typically demonstrate a lobulated appearance. The lesion is of low to intermediate signal intensity on the T1-weighted images, frequently demonstrates punctate foci of signal void due to matrix mineralization and may demonstrate foci of high signal intensity on the T1-weighted images which are due to entrapped areas of yellow marrow. On the fluid-sensitive sequences, the tumour demonstrates very high signal intensity due to the high water content within the chondrocytes (Figure 20a,b). The high signal intensity lobules of cartilage cells are frequently separated by septa, which are hypointense on the fluid-sensitive sequences and demonstrate enhancement after gadolinium administration (Figure 21a,b). Endosteal scalloping, soft tissue extension, cortical changes and periosteal reaction are well appreciated on MRI (Figure 22a,b). Whilst perilesional bone marrow-like oedema is uncommon, its presence favours the diagnosis of a chondrosarcoma (Figure 23a,b). Accurate preoperative grading of chondrosarcomas is important because the treatment of low-grade chondrosarcomas (grade 1 chondrosarcomas) significantly differs from that of high-grade chondrosarcomas (grade 2 and grade 3 chondrosarcomas) in many centres. In many bone tumour centres, low-grade chondrosarcomas are treated with curettage whilst high-grade chondrosarcomas are treated with en-bloc excision or amputation.
Figure 20: MRI of a chondrosarcoma of the proximal humerus. Coronal T1W SE MRI (a) and coronal T2W FS MRI (b) of the humerus shows an 8.9cm in the maximum cranio-caudal dimension measuring lesion in the proximal humeral diaphysis which is of low to intermediate signal intensity on the T1-weighted images (a), of high signal intensity on the T2 fat-suppressed images (b) and which demonstrates foci of signal void (arrow) due to matrix mineralisation.
Figure 21: MRI of a chondrosarcoma of the proximal tibia. Coronal T1W SE MRI (a) and gadolinium-enhanced T1W MRI (b) of the tibia shows a chondrosarcoma of the proximal tibia (a) which demonstrates avid septal enhancement after gadolinium administration (b, arrow).
Figure 22: MRI of a chondrosarcoma of the proximal femur. Coronal T2W FS MRI (a) and axial T2W FS MRI of the left femur shows a 17.1cm in the maximum cranio-caudal dimension measuring chondroid lesion with bony expansion (a, arrow), marked endosteal scalloping (b, small arrow), cortical thickening and periosteal reaction (b, large arrow).
Figure 23: MRI of a grade 1 chondrosarcoma of the proximal tibia. Sagittal (a) and axial (b) PD FS MRI-study of the knee shows an only 2.1cm chondroid lesion within the proximal tibial epiphysis. Despite the small size of the lesion, there is extensive peritumoural bone marrow-like oedema (long arrow), soft tissue oedema and cortical destruction (short arrow). Biopsy confirmed a grade 1 chondrosarcoma.
Although biopsy is frequently performed prior to surgery, only a small area of the tumour is sampled which may result in erroneous down-grading of the tumour. Unpublished data from the Royal National Orthopaedic Hospital, Stanmore performed by the author of this PhD has shown that CT-guided biopsy of chondrosarcomas results in erroneous down-grading of the tumour in approximately 10% of all cases. More worryingly, a recent study published by Roitman et al., demonstrated that concordance between preoperative CT-guided bone biopsy and the final pathological grading in chondrosarcomas was 83% in long bones and only 36% in the pelvis.\textsuperscript{44}

This significant discrepancy between preoperative histological grading and final histological grading in chondrosarcomas may result in inadequate treatment of a high-grade chondrosarcoma. Although a few studies have attempted to evaluate the role of MRI in the differentiation of low-grade from high-grade chondrosarcomas, the results of these studies were hampered by small sample size and by the fact that these studies only evaluated very few MRI-criteria associated with chondrosarcomas leading to conflicting results.\textsuperscript{45-48} Therefore, there remained uncertainty about the role of MRI in the differentiation of low-grade chondrosarcomas from high-grade chondrosarcomas. However, the clinical significance of accurate preoperative grading of chondrosarcomas can not be overemphasized as erroneous downgrading of the tumour due to sampling errors on biopsy may lead to curettage of high-grade chondrosarcomas which is inadequate and hence may result in further surgery. In chapter 6 of this thesis, I therefore present a study which forms part of this PhD-
thesis evaluating the utility of MRI in the differentiation of low-grade from high-grade chondrosarcomas.

Fluorodeoxyglucose-positron emission tomography (FDG-PET) is widely used in oncological imaging in the detection, diagnosis, staging, treatment response assessment and evaluation of recurrence of tumours due to its ability to detect glucose uptake in cells with high metabolic activity. PET/CT has therefore been utilized in an attempt to differentiate benign cartilage tumours from malignant cartilage tumours and in the grading of chondrosarcomas. Whilst there is conflicting evidence about the role of PET/CT in the differentiation of benign from malignant cartilage tumours (Figures 24,25), PET/CT can be used to differentiate benign and low-grade malignant cartilage tumours from high-grade malignant cartilage tumours as it has been demonstrated that high-grade chondrosarcomas have a higher SUV than low-grade chondroid tumours and that the SUV increases with higher histological grade of the tumour. However, this differentiation can be reliably made on MRI as demonstrated by our group and hence the role of PET/CT in the diagnosis and characterization of chondrosarcomas is of doubtful clinical significance especially when one considers that MRI forms part of the routine practice in the loco-regional staging of chondrosarcomas whilst PET/CT is not usually used in the staging of chondrosarcomas. Furthermore, in contrast to MRI, PET/CT is associated with a radiation burden. Hence, PET/CT plays no significant role in the diagnostic workup of patients with chondrosarcoma.
Figure 24: Eccentric enchondroma of the humeral head. Axial T2W FS MRI-study of the left humerus (a) shows imaging features in keeping with an eccentric enchondroma of the...
humeral head (a, arrow). On FDG-PET/CT (b), the lesion demonstrates increased FDG-uptake with an SUVmax of 3.6. The lesion was a histologically confirmed enchondroma.

Figure 25:
Figure 25a: Figure 25b:

Figure 25: Grade 1 chondrosarcoma of the left proximal humerus. Coronal CT (a) shows a 19cm in the maximum craniocaudal dimension measuring grade 1 chondrosarcoma of the left proximal humerus. On FDG-PET/CT (b), the lesion demonstrates mild increased FDG-uptake with an SUVmax of 3.4. The lesion was a histologically confirmed grade 1 chondrosarcoma.
On 99m-Technitium methylene diphosphonate (MDP) bone scintigraphy, 82% of all chondrosarcomas demonstrate marked radio-isotope uptake which is greater than the uptake of the anterior iliac crest (Figure 26a,b) whilst avid isotope uptake is only observed in 21% of all enchondromas. Similarly, a heterogeneous radionuclide uptake pattern is observed in 63% of chondrosarcomas but only in 30% of enchondromas. As the above figures however demonstrate, there is significant overlap in the intensity of radio-isotope uptake and the pattern of uptake between enchondromas and chondrosarcomas. Therefore, skeletal scintigraphy cannot be used to differentiate between the two disease entities.
Figure 26: Chondrosarcoma of the proximal femur. Anteroposterior radiograph of the femur (a) shows a heavily calcified ill-defined lesion (arrow) in the right proximal femur. On 99m-Technitium methylene diphosphonate (MDP) bone scintigraphy (b), the lesion demonstrates a heterogeneous and avid radio-isotope uptake which is greater than the uptake of the anterior iliac crest. The lesion was a histologically confirmed grade 2 chondrosarcoma.
1.2.2.6 Staging of chondrosarcomas

In contrast, skeletal scintigraphy is recommended and therefore widely used in the staging of chondrosarcomas at time of presentation to exclude/detect skeletal metastases. Similarly, Computed Tomography of the chest is widely recommended in the staging of chondrosarcomas to evaluate the presence of pulmonary metastases. An in depth review of the literature however reveals that this recommendation is largely based on evidence acquired from osteosarcomas and Ewing sarcomas where skeletal and pulmonary metastases at presentation are observed in 4%-19% of all patients. However, the incidence of skeletal and pulmonary metastases in chondrosarcomas at presentation, the role of skeletal scintigraphy and Computed Tomography of the chest in the staging of chondrosarcomas have not been evaluated previously. In chapter 7, I present a study which forms part of the submitted thesis evaluating the role of bone scintigraphy and Computed Tomography of the chest in the staging of chondrosarcomas showing some intriguing findings.

1.2.2.7 Management of chondrosarcomas

In contrast to enchondromas which are only treated if found to be symptomatic, chondrosarcomas are always treated with surgery. This is of particular importance in view of the fact that chondrosarcomas are insensitive to radiotherapy or chemotherapy and therefore the only hope for cure at present is surgery. The lack of
non-surgical treatment options is therefore associated with a poor prognosis in patients with metastatic chondrosarcoma.

There has been a change in the surgical treatment technique of low-grade and high-grade chondrosarcomas in recent years. Whilst previously, both low-and high-grade chondrosarcomas were treated with limb-salvage surgery and endoprosthetic reconstruction or amputation, low-grade chondrosarcomas are treated in many centres with intralesional curettage and local adjuvant therapy. In contrast, high-grade chondrosarcomas continue to be treated with endoprosthetic reconstruction or amputation. Therefore, accurate preoperative differentiation of low-grade chondrosarcomas from high-grade chondrosarcomas is crucial.

1.2.3 Enchondroma versus low-grade chondrosarcoma

The differentiation of enchondroma from low-grade chondrosarcoma is one of the most difficult topics in musculoskeletal oncology and is challenging for pathologists, radiologists and clinicians alike. In the following section, the scientific evidence regarding the role and limitation of both histopathology and the various imaging modalities in the differentiation of enchondromas from low-grade chondrosarcomas is discussed in-depth thereby emphasizing the difficulties in the differentiation of the two disease entities.
1.2.3.1 Histology

The difficulties that pathologists face in the distinction of the two disease entities has been highlighted in two previous studies which evaluated the interobserver variability in the differentiation of enchondromas from low-grade chondrosarcomas and in the grading of chondrosarcomas even by experienced musculoskeletal pathologists.\textsuperscript{67,68}

The “Skeletal Lesions Interobserver Correlation among Expert Diagnosticians (SLICED) Study Group” quantified the interobserver reliability in the differentiation of enchondromas from low-grade chondrosarcomas and in the grading of chondrosarcomas among a group of experienced musculoskeletal pathologists and radiologists. Nine musculoskeletal pathologists and eight musculoskeletal radiologists who were experts in the interpretation of chondroid tumours reviewed forty-six cartilaginous tumours of long bone which underwent open biopsy, curettage or excision. Both pathologists and radiologists were asked to classify the chondroid tumours as benign, low-grade malignant or high-grade-malignant using whatever criteria they use in their daily clinical practice. This study demonstrated that the interobserver reliability in the grading of cartilaginous neoplasms in long bones was only moderate for pathologists (kappa value=0.443) and only fair for radiologists (kappa value=0.345) although the inclusion of Magnetic Resonance Imaging resulted in a slightly improved agreement between radiologists (kappa value=0.437).\textsuperscript{68}

Similarly, Eefting and co-workers evaluated the interobserver variability in the histological diagnosis of enchondromas from grade 1 chondrosarcomas and in the grading of chondrosarcomas. In their study, the authors included 16 chondroid tumours which were evaluated by 18 musculoskeletal pathologists. The authors
discovered that the distinction between enchondroma and grade 1 chondrosarcoma was most discordant demonstrating only moderate agreement (kappa coefficient=0.54) whilst the differentiation between grade 1 and grade 2 chondrosarcomas demonstrated substantial agreement (kappa coefficient=0.8).\textsuperscript{67,69} This discrepancy is of significance particularly when comparing these findings with the interobserver reliability in the grading observed in other tumours. In contrast to this finding for example the interobserver reliability in the histopathological grading of soft tissue sarcomas demonstrates kappa coefficients of 0.68 and 0.78.\textsuperscript{70,71} Similarly, histopathological grading of breast carcinoma has shown similar kappa coefficients ranging from 0.69 to 0.73.\textsuperscript{72-74}

The diagnostic dilemma in the differentiation of enchondromas from grade 1 chondrosarcomas has also been highlighted in multiple publications. In these publications central cartilage tumours which on histology are moderately cellular, may show focal myxoid change and mild nuclear atypia however fail to demonstrate a permeative growth have been variably classified as “CLUMPs” (= cartilaginous lesions of unknown malignant potential),\textsuperscript{24} “grade 0 chondrosarcomas”, “atypical enchondromas”\textsuperscript{25} or as “borderline lesions”. Whilst the diagnostic uncertainty in the differentiation of enchondromas from grade 1 chondrosarcomas is widely accepted, proven and highlighted in the latest World Health Organization Classification of bone tumours published in 2013, the above described terms for low-grade chondroid tumours which are difficult to classify havenot been adpoted in the most recent WHO Classification of bone tumours.\textsuperscript{75} These terms continue to be used in many bone tumour centres highlighting the diagnostic dilemma in the differentiation of the two disease entities.
The reported interobserver variability and diagnostic uncertainty in the distinction of enchondromas from grade 1 chondrosarcomas is of clinical relevance as the lack of a gold standard in the diagnosis of the two disease entities may result in inappropriate surgical treatment of an enchondroma or in lack of treatment of a grade 1 chondrosarcoma with potentially adverse consequences. Furthermore, there is controversy regarding the biological behaviour and hence treatment approach of these borderline cartilage tumours with some institutions opting for a watch-and-wait approach whilst other centres decide to perform intralesional curettage for these lesions.

Therefore, in the absence of a permeative growth pattern on histology, consensus between imaging findings, histopathology and clinical findings is most prudent in the distinction of enchondromas from grade 1 chondrosarcomas. Hence, in the next section, the scientific evidence of the role and the limitations of the various imaging modalities in the differentiation of enchondromas from low-grade chondrosarcomas is discussed.

The differentiation of enchondromas and grade 1 chondrosarcomas is particularly challenging in the long tubular bones such as the femur, tibia and humerus where enchondromas are very commonly observed. Similarly, the long tubular bones are the most common location for chondrosarcomas accounting for approximately 45% of all cases.
1.2.3.2 Imaging

As stated previously, imaging is vital in the differentiation of enchondromas from grade 1 chondrosarcomas. In the following section, the role of the various imaging modalities in the differentiation of enchondromas from grade 1 chondrosarcomas will be discussed in more detail. The imaging modalities discussed below include: Radiography, Computed Tomography, Skeletal scintigraphy, Positron Emission Tomography and Magnetic Resonance Imaging with a particular emphasis on advanced MRI-techniques which include Diffusion-weighted MRI and Dynamic-contrast enhanced MRI.

1.2.3.2.1 Radiography

Geirnaert and co-workers have previously evaluated the role of radiography in the differentiation of enchondromas from grade 1 chondrosarcomas. The authors included 35 enchondromas and 43 central grade 1 chondrosarcomas. The diagnosis of chondroid tumours was based on histology and long-term follow-up. In their study, 51% of all enchondromas were located in the hands and feet whilst only 5% of all grade 1 chondrosarcomas were located in the hands and feet. In contrast, within the axial skeleton, grade 1 chondrosarcomas were statistically more commonly observed than enchondromas whilst there was no statistically significant difference in the number of enchondromas and grade 1 chondrosarcomas identified in the femora and humeri. Furthermore, in their study, grade 1 chondrosarcomas were significantly larger (median size: 5cm) than enchondromas (median size: 2cm) although there was
significant overlap in size when enchondromas of the hands and feet were excluded. The authors also discovered that only the presence of ill-defined margins and lobulated contours on radiography were statistically significant differentiating features between enchondromas and grade 1 chondrosarcoma. The authors however reported that radiographic features such as endosteal scalloping, cortical thinning, destruction, periosteal reaction and soft tissue extension were not statistically significant differentiating features. The major limitation of this publication is however the study design which did not differentiate between the radiographic appearances of low-grade chondroid tumours of the hands and feet, low-grade chondral tumours of the axial skeleton and low-grade chondroid tumours of the long tubular bones. Furthermore, 51% of enchondromas in this study were located in the hands and feet whilst 35% of grade 1 chondrosarcomas were located in the axial skeleton. The diagnostic challenge in the differentiation of enchondromas and grade 1 chondrosarcomas is however largely one centered around the long tubular bones (such as the femur, humerus, tibia) because both enchondromas and chondrosarcomas are frequently observed in this location particularly as 45% of all chondrosarcomas are diagnosed in the long tubular bones. Therefore, Murphey and co-workers evaluated the role of radiographs in the distinction of enchondromas from chondrosarcomas of the appendicular skeleton. The authors included 92 enchondromas and 95 chondrosarcomas which included 35 grade 1 chondrosarcomas (37% of all chondrosarcomas), 29 grade 2 chondrosarcomas (31% of all chondrosarcomas) and 31 grade 3 chondrosarcomas (33% of all chondrosarcomas). They found that a depth of endosteal scalloping of more than two-third of the cortical thickness as well as the extent of endosteal
scalloping were statistically significant differentiating features on radiography. Furthermore, increased tumour size, the presence of cortical destruction, cortical thickening, pathological fracture, periosteal reaction and soft tissue extension were statistically significant differentiating features favouring the diagnosis of a chondrosarcoma. A major limitation of this study is that the study cohort included chondrosarcomas in general and did not differentiate between low-grade and high-grade chondrosarcomas in their analysis. It is well established that both the histological and imaging differentiation of enchondromas from high-grade chondrosarcomas does not represent a diagnostic challenge. In contrast, the difficulty lies in the differentiation of enchondromas from low-grade chondrosarcomas. Therefore, the results published by Murphey et al. have to be interpreted in the context of this significant limitation and it cannot be assumed that the described differentiating features are also applicable in the differentiation of enchondromas from low-grade chondrosarcomas of the appendicular skeleton.15

1.2.3.2.2 Computed Tomography

In the above quoted publication by Murphey et al., the authors also evaluated the role of CT in the differentiation of enchondromas and chondrosarcomas of the appendicular skeleton. In their study, the authors analyzed the CT-examinations of 88 lesions of which 39 were enchondromas and 49 lesions were chondrosarcomas. On CT, the size of the lesion, depth and extent of endosteal scalloping, cortical destruction, the presence of a pathological fracture, periosteal reaction, and soft tissue extension were statistically significant differentiating features. In contrast,
cortical remodeling, cortical thickening, the presence and extent of matrix
calcification as well as the attenuation values of the non-mineralized component on
CT were not statistically significant. As stated above, the major limitation of this study
is the lack of differentiation of enchondromas from low-grade chondrosarcomas.15

1.2.3.2.3 Skeletal Scintigraphy

The role of skeletal scintigraphy in the differentiation of enchondromas from
chondrosarcomas in the appendicular skeleton has also been assessed by Murphey
and co-workers. The authors retrospectively reviewed the skeletal scintigraphs of 67
enchondromas and 51 chondrosarcomas. Isotope uptake in the lesions was graded
on a scale of 1-3. Grade 1 was classified as uptake within the lesion less than that of
the anterior iliac crest, grade 2 was similar to the uptake in the anterior iliac crest
whilst grade 3 was defined as uptake greater than in the anterior iliac crest.
Furthermore, the authors assessed if the radio-isotope in the lesions was
homogeneous or heterogeneous. The authors found that bone scintigraphy showed
greater isotope uptake when compared to the anterior iliac crest in 82% of
chondrosarcomas (figure 26) whilst this finding was only observed in 21% of all
enchondromas (figure 8). In contrast, 79% of all enchondromas demonstrated equal
or lower activity than the anterior iliac crest (figure 7b). Similarly, heterogeneous
uptake was observed in 63% of all chondrosarcomas but only in 30% of all
enchondromas. This finding was thought to be due to increased biological activity of
chondrosarcomas.15 Although the difference in radio-isotope uptake between
enchondromas and chondrosarcomas of the appendicular skeleton was statistically
significant in this study, the authors as stated above did not differentiate between enchondromas and grade 1 chondrosarcomas and therefore these findings cannot be used to differentiate between the two disease entities.

1.2.3.2.4 Positron emission tomography

The potential role of FDG PET in the differentiation of benign cartilaginous tumours and chondrosarcomas has been assessed in three studies.

Aoki et al. studied the SUV values in four enchondromas, one osteochondroma and six chondrosarcomas which included one grade 1 chondrosarcoma, four grade 2 chondrosarcomas and one grade 3 chondrosarcoma. They found that the mean SUV in benign cartilage tumours was 0.96 whilst the mean SUV of chondrosarcomas was 2.23. However, the sample size of the study was very small and included only one grade 1 chondrosarcoma. Therefore, the findings cannot be used to differentiate enchondromas from low-grade chondrosarcomas.

Similarly, Feldman et al. also studied the role of PET in 29 chondroid tumours. In this study, the benign chondroid tumour group included 11 enchondromas and 7 osteochondromas whilst the chondrosarcoma-group included 5 grade 1 chondrosarcomas, 2 grade 2 chondrosarcomas, 1 grade 3 chondrosarcoma and 2 dedifferentiated chondrosarcomas. The authors found that a cut-off maximum SUV of 2.0 resulted in a sensitivity of 90.9%, a specificity of 100% and a diagnostic accuracy of 96.6% in the differentiation of benign cartilage tumours from chondrosarcomas.

In view of the heterogeneity of benign cartilage tumours which were included in this study and in view of the inclusion of low-grade and high-grade chondrosarcomas, no
conclusions about the role of PET in the differentiation of enchondromas from low-grade chondrosarcomas can be made based on this study.\textsuperscript{50}

In contrast, a study by Lee and co-workers retrospectively assessed PET in 35 cartilaginous tumours which included ten enchondromas, three osteochondromas, twelve grade 1 chondrosarcomas, five grade 2 chondrosarcomas and five grade 3 chondrosarcomas. In this study, the mean SUV of benign chondral tumours was 1.147, the mean SUV of grade 1 chondrosarcomas was 0.898 whilst the mean SUV of high-grade chondrosarcomas was 6.903. The authors therefore concluded that although PET could be used to differentiate low-grade chondrosarcomas from high-grade chondrosarcomas, it could not distinguish between benign cartilage tumours and grade 1 chondrosarcomas (Figures 24, 25). Hence, there is no role for PET in the differentiation of enchondromas from low-grade chondrosarcomas.\textsuperscript{16}

\textbf{1.2.3.2.5 Magnetic Resonance Imaging}

The utility of MRI in the differentiation of enchondromas from chondrosarcomas has been investigated in a few studies. In the following section, the role of anatomical MRI in the differentiation of the two disease entities will be discussed followed by a review of the literature on the role of functional MRI (dynamic-contrast enhanced MRI and diffusion-weighted MRI) in the distinction of enchondromas from low-grade chondrosarcomas.
1.2.3.2.5.1 Anatomical Magnetic Resonance Imaging

Murphey et al. retrospectively analyzed the MR Imaging of 35 enchondromas and 33 chondrosarcomas of the appendicular skeleton. They evaluated lesion size, endosteal scalloping, cortical remodeling, cortical destruction, pathological fracture, periosteal reaction, cortical thickening and soft tissue extension. The group found that the mean length of enchondromas was 5cm whilst the mean length of chondrosarcomas was 8cm. Similarly, the depth and extent of endosteal scalloping was significantly different in enchondromas and chondrosarcomas with endosteal scalloping of more than two-thirds being observed in 67% of chondrosarcomas but only in 11% of enchondromas. Furthermore, cortical remodeling, cortical destruction, soft tissue extension and pathological fracture were statistically significant differentiating features between enchondromas and chondrosarcomas whilst cortical thickening and periosteal reaction could not differentiate between the two disease entities. However, a significant limitation of this study is that the authors did not perform a subgroup analysis to evaluate potential differentiating MRI-features between enchondromas and low-grade chondrosarcomas.\(^\text{15}\)

There have been two studies which evaluated the role of MRI in the differentiation of enchondromas from low-grade chondrosarcomas. Ferrer-Santacreu et al. assessed the role of MRI in the differentiation of enchondromas and low-grade chondrosarcomas of the appendicular skeleton in 82 patients. The authors found no statistically significant differentiating imaging features between enchondromas and low-grade chondrosarcomas.\(^\text{78}\)
Another study by Choi et al. assessed the MRI-features in 16 enchondromas and in 18 low-grade chondrosarcomas. In contrast to Ferrer-Santacreu, the authors found that predominantly intermediate signal intensity on T1WI, a multilobular appearance on contrast-enhanced T1WI, cortical destruction, soft tissue extension, surrounding bone marrow oedema, abnormal soft tissue signal, epiphysial involvement and location in the flat bone were more commonly observed in low-grade chondrosarcomas than in enchondromas. A significant limitation of this publication is that the authors included chondrosarcomas of the flat bones and enchondromas of the hand in their study. As previously stated, the differentiation of enchondromas from low-grade chondrosarcomas does not represent a diagnostic challenge in these locations. Similarly, the imaging findings of chondroid lesions of the hands and pelvis are very different to the imaging findings of chondroid lesions of the appendicular system. Therefore, the role of conventional MRI in the differentiation of enchondromas from low-grade chondrosarcomas remained uncertain until very recently. A recent study by the author of this thesis specifically evaluated the role of conventional MRI, DCE-MRI and clinical findings in the differentiation of enchondromas from low-grade chondrosarcomas of long bones (see Appendix 1). In this study, we retrospectively analyzed the MRI-findings in 60 central chondroid tumours which included 27 enchondromas, 10 cartilaginous tumours of unknown malignant potential, 15 grade 1 chondrosarcomas and 8 high-grade chondrosarcomas. The subgroup analysis which evaluated the conventional MRI-findings of enchondromas and grade 1 chondrosarcomas, revealed that more than 2/3 endosteal scalloping demonstrated a sensitivity of 71.4% (95% confidence intervals: 41.9%-91.6%) and a specificity of 92.9% (95% confidence interval: 76.5%-
bone expansion showed a sensitivity of 50% (95% confidence interval: 23%-77%) and a specificity of 89.3% (95% confidence interval: 71.8%-97.7%), cortical destruction showed a sensitivity of 35.7% (95% confidence interval: 12.8%-64.9%) and a specificity of 100% (95% confidence interval: 87.7%-100%) whilst the presence of a soft tissue mass revealed a sensitivity of 40% (95% confidence interval: 16.3%-67.7%) and a specificity of 100% (95% confidence interval: 87.2%-100%). Of note is that the presence of pain attributed to the lesion resulted in a sensitivity of 60% (95% confidence interval: 32.3%-83.7%) and a specificity of 88.9% (95% confidence interval: 70.8%-97.7%) in the differentiation of enchondromas from grade 1 chondrosarcomas. Subsequent calculation of Odds ratios showed an Odds ratio for pain attributable to the lesion of 12 (95% confidence interval: 2.5-58.5), an Odds ratio for more than 2/3 endosteal scalloping of 25 (95% confidence interval: 4.2-15.7), and an Odds ratio for bone expansion of 14.3 (95% confidence interval: 2.5-83.1) whilst cortical destruction and the presence of a soft tissue mass were diagnostic for a grade 1 chondrosarcoma. Of note is that the 95% confidence intervals for the sensitivity, specificity and the Odds ratios were wide. This finding is most likely due to the small sample size of the study. Despite the wide confidence intervals of the results, the study demonstrated a positive relationship between pain attributable to the lesion, more than 2/3 endosteal scalloping, bone expansion and the diagnosis of a grade 1 chondrosarcoma. Although the study confirmed that cortical destruction and the presence of a soft tissue mass are diagnostic for a grade 1 chondrosarcoma, these findings are not commonly observed in grade 1 chondrosarcomas. However, the study demonstrated that more than 2/3 of endosteal scalloping is the most sensitive conventional MRI sign of grade 1 chondrosarcomas. Similarly, pain attributed to the
lesion was an important clinical sign of grade 1 chondrosarcomas highlighting the importance of a close collaboration between radiologists, clinicians and pathologists in order to differentiate between enchondromas from grade 1 chondrosarcomas. 

1.2.3.2.5.2 Dynamic Contrast-Enhanced Magnetic Resonance Imaging

Dynamic contrast-enhanced MRI (DCE-MRI) is a functional MRI-technique which allows the non-invasive assessment of tumour vascularity. DCE-MRI has previously been utilized in an attempt to differentiate benign bone tumours from malignant bone tumours. More specifically, there have been three studies which attempted to differentiate benign chondroid tumours from chondrosarcomas using DCE-MRI. Geirnaerdt and co-workers assessed eight enchondromas, eleven osteochondromas and eighteen chondrosarcomas which included seven grade 1 chondrosarcomas, nine grade 2 chondrosarcomas and two grade 3 chondrosarcomas using DCE-MRI in an attempt to differentiate benign cartilage tumours from chondrosarcomas. The authors evaluated the start of contrast uptake (early versus delayed) and the progression of contrast uptake (exponential versus gradual) within chondroid tumours. They found that early enhancement was observed in 89% of all chondrosarcomas but not in enchondromas whilst delayed enhancement was seen in 11% of chondrosarcomas and in 38% of enchondromas. In contrast, no enhancement was described in 63% of all enchondromas but in no case of a chondrosarcoma. Furthermore, exponential uptake was seen in 61% of all chondrosarcomas but not in enchondromas whilst gradual enhancement was
observed in 38% of enchondromas and in 39% of chondrosarcomas. The authors therefore concluded that lack of enhancement within a cartilage tumour excluded a chondrosarcoma. Furthermore, they concluded that the differentiation of benign cartilage tumours from chondrosarcomas was possible with a sensitivity of 61%, a specificity of 95%, a positive predictive value of 92% and a negative predictive value of 72%. A significant limitation of this study is that the authors assessed the contrast uptake pattern in enchondromas versus chondrosarcomas in general and did not perform a subgroup analysis evaluating the contrast uptake pattern in enchondromas versus low-grade chondrosarcomas. As stated previously, the distinction between enchondromas and high-grade chondrosarcomas does not represent a diagnostic challenge. A further limitation of this study is the small sample size which most likely would have hampered a subgroup analysis.

De Coninck et al. also evaluated the role of DCE in the differentiation of enchondromas from chondrosarcomas. In their retrospective study, the authors included 75 enchondromas and 31 chondrosarcomas which included 18 low-grade chondrosarcomas, 10 intermediate grade chondrosarcomas and 3 high grade chondrosarcomas. The authors found that enhancement within the tumour which was two times more when compared to muscle combined with a 76 degree slope of the uptake curve resulted in a 100% sensitivity and 63.3% specificity in the detection of chondrosarcomas. A significant limitation of this study is the inclusion of low-grade and high-grade chondrosarcomas and the fact that the authors did not differentiate between low-grade chondrosarcomas from high-grade chondrosarcomas in this study. Therefore, the obtained results cannot be utilized in the differentiation of enchondromas from low-grade chondrosarcomas. Hence, the role of DCE-MRI in
the differentiation of the two disease entities remained unclear until very recently. As stated above, the author of this thesis re-evaluated the role of conventional MRI, DCE-MRI and clinical assessment in the differentiation of enchondromas from low-grade chondrosarcomas and high-grade chondrosarcomas of long bone (see Appendix 1). In this study, we evaluated the median angle of the curve, the median absolute enhancement of the curve and the relative enhancement of the curve for enchondroma, CLUMPs, grade 1 chondrosarcoma and high-grade chondrosarcomas. Whilst the study demonstrated a statistically significant difference in the median angle of the curve this was due to differences in the angle of the curve between enchondromas versus high-grade chondrosarcomas, due to differences in the angle of the curve between the CLUMP-group and grade 1 chondrosarcomas and due to differences in the group between the CLUMP-group and the high-grade chondrosarcoma group. Similarly, the statistically significant difference in the median absolute enhancement of the various curves was due to differences in the absolute enhancement of enchondromas versus high-grade chondrosarcomas and due to differences between the CLUMP-group and high-grade chondrosarcoma group. In contrast, there was no statistically significant difference in the pairwise comparison between the various chondroid groups. Similarly, there was no statistically significant difference of the various DCE-MRI parameters between enchondromas and grade 1 chondrosarcomas. This study therefore demonstrates that the previously reported potential role of DCE-MRI in the differentiation of enchondromas from chondrosarcomas is most likely due to the inclusion of high-grade chondrosarcomas. In contrast, according to our findings, DCE-MRI cannot be utilized to differentiate enchondromas from low-grade chondrosarcomas.\textsuperscript{80}
Another functional MRI-technique which has gained increased popularity and is a promising tool in the diagnosis and treatment response assessment in oncological imaging is diffusion-weighted MRI (DWI). DWI exploits the principle of Brownian motion of water molecules. Within biological tissues, the motion of water molecules is restricted by barriers such as large molecules and cell membranes. The degree of diffusion restriction of water molecules in biological tissue is inversely correlated to tissue cellularity and the integrity of cell membranes. Hence, high cellular tissue may demonstrate more restricted diffusion than low cellular tissue.\(^{85,86}\) DWI has therefore been utilized to differentiate benign skeletal lesions from malignant skeletal lesions.\(^ {87,88}\) As one differentiating feature between enchondromas from low-grade chondrosarcomas on histology is cellularity, DWI appears to be an attractive functional imaging technique in the differentiation of the two disease entities. There has been a paucity of evidence in the literature about the potential role of DWI in the differentiation of enchondromas from low-grade chondrosarcomas.

A recent study assessed the utility of DWI in the differentiation of enchondromas from chondrosarcomas. The authors found that the ADC-values between enchondromas, grade 1, grade 2 and grade 3 chondrosarcomas were statistically significant, however the differences were only small and the authors therefore concluded that a larger sample size was required to assess the utility of DWI in the differentiation of enchondromas from grade 1 chondrosarcomas.\(^ {89}\) In the fifth chapter of this thesis, the author therefore presents a study evaluating the role of
DWI in the differentiation of enchondromas from low-grade chondrosarcomas which showed some interesting findings.

Whilst it has become evident from the extensive review of the literature, that imaging plays a role in the diagnosis of central chondroid bone tumours, there remains uncertainty about the role that imaging may play in advancing the understanding of the pathogenesis of enchondromas and in the precise role that imaging may play in the diagnosis, grading and staging of chondrosarcomas of bone. In the following chapters, the authors’ published work submitted for consideration of a PhD therefore investigates the theory that imaging may aid in the understanding of the pathogenesis of enchondromas and may aid in the diagnosis, grading and staging of chondrosarcomas of bone. The first two chapters are review articles which summarize the role of imaging in the diagnosis of enchondromas and chondrosarcomas. The third article investigates the role that MRI may play in advancing the understanding of the pathogenesis of enchondromas. The fourth article assesses the role of DWI-MRI as a functional MRI-technique in the differentiation of enchondromas from low-grade chondrosarcomas. The fifth article evaluates the utility of conventional MRI in the grading of chondrosarcomas of long bone. Finally, the sixth article examines the role of bone scintigraphy and Computed Tomography of the chest in the staging of chondrosarcomas. The presented thesis therefore investigates the role of imaging in the diagnosis, grading and staging of chondrosarcomas.
Chapter 8: Discussion

8.1 Summary

Enchondromas are the second most common benign cartilage tumour whilst chondrosarcomas are the second most common malignant primary bone tumour after osteosarcoma. Despite this, there has been a paucity of evidence about the role of imaging in the diagnosis, grading and staging of chondrosarcomas. The hereby presented portfolio of peer-reviewed published work submitted for consideration of a PhD thesis significantly enhances the understanding of the pathogenesis of enchondromas and significantly contributes to the knowledge about the role of imaging in the diagnosis, grading and staging of chondrosarcomas.

The first two articles, which form part of the thesis are review articles providing an overview of imaging of benign chondroid tumours and chondrosarcomas. In these articles, the role of imaging in the diagnosis of benign chondroid tumours and chondrosarcomas, in the grading and staging of chondrosarcomas is discussed.

The third article, which is an original article, critically appraises the commonly held belief about the pathogenesis of enchondromas. Whilst it is widely believed that enchondromas arise from cartilage remnants which have been displaced from the growth plate during the process of skeletal maturation, the origin of this very theory has been forgotten in modern medicine. Furthermore, this theory remains unproven. In the original article entitled “Can MR imaging challenge the commonly accepted theory of the pathogenesis of solitary enchondroma of long bone?”, we have
investigated the origin of the above stated theory and have surprisingly discovered that this theory was first postulated by the “father of modern pathology” Rudolf Virchow in 1863 in his 3-volume book “Die krankhaften Geschwülste” and later expanded upon by Virchow in the article “Über die Entstehung des Enchondroma und seine Beziehungen zu der Ecchondrosis und der Exostosis cartilaginea” which he published in “Monatsberichte der Königlich Preussischen Akademie der Wissenschaften” in 1875. Furthermore, we discovered after studying the original manuscripts in German, that Virchow’s theory was based on observations he made in patients suffering from rickets (Figure 27).\textsuperscript{90,91} Until our publication, both the origin and historic background of this theory were unknown in modern medicine. The re-discovery of the very origin, background and reasoning of this theory in our article is therefore a significant contribution to the history of medicine in general and the understanding of the history of the pathogenesis of enchondromas in particular.

We subsequently challenged this theory using MR Imaging in skeletally immature patients. The article which forms part of the thesis, raises doubts about the widely believed but unproven theory that enchondromas arise from displaced cartilage islands during the process of skeletal maturation as we did not identify any macroscopically detectable cartilage islands on MRI. The doubt about the validity of this theory is also supported by the discovery of mutations in the genes encoding Isocitrate dehydrogenase 1 and 2 (IDH1 and IDH2) in enchondromas, periosteal chondromas and central chondrosarcomas therefore highlighting their role in the tumourigenesis of enchondromas.\textsuperscript{33} There are limitations to our study. Our investigations and study do not refute the theory that enchondromas arise from
cartilage remnants which have been displaced from the growth plate during the process of skeletal maturation. The study raises doubts about the validity of this theory for the following reasons:

Firstly, Virchow based his theory on observations he made in pathology specimen of children with rickets where he observed proliferations and islands of cartilage extending from the growth plate into the metadiaphysis. He therefore concluded that such foci of cartilage proliferation could have persisted into adulthood and would have subsequently formed into enchondromas. Whilst rickets was endemic in Europe of the 19th century, its incidence although slightly increasing lately, has dramatically reduced since the 19th century and can therefore not be an important cause for the development of enchondromas. Other causes, which could result in disruption of the growth plate, cannot be ruled out as potential causative agents in the pathogenesis of enchondromas. In particular, trauma in the skeletally immature individual, continues to be frequently observed and remains a possible cause for the development of enchondromas. This is of particular importance when one considers that repetitive injuries as seen in elite athletes or single event insults such as fractures, may lead to injury and disruption of the growth plate and on MRI may manifest themselves as extension of physeal cartilage into the metaphysis. Similar extensions of cartilage from the growth plate into the metaphysis were observed in seven MRI-examinations of six patients in our studied cohort (2.9% of all cases). In all seven MRI-studies, the cartilage extensions into the growth plate were in continuity with the growth plate. The aetiology of these cartilage extensions in our study remains unknown due to the retrospective nature of the study. We can
therefore not exclude that such cartilage extensions may become separated from the growth plate at a later stage during the process of skeletal maturation and could potentially develop into enchondromas. This is of importance especially when one considers that growth plate closure is usually observed at the age of 16 years and the median age of patients in our study was 13 years. However, as stated above, similar cartilage extensions have been described previously and have been attributed to single event or multi event injuries such as previous trauma which either regress after cessation of the repetitive injury or in the case of severe injuries may lead to significant clinically evident growth disturbances.

Furthermore, we did not identify any enchondromas in our cohort which evaluated 240 MRI-studies of the knee in 209 skeletally immature patients. The absence of enchondromas in skeletally immature patients in our study is however in contrast to the prevalence of incidental enchondromas on MRI-examinations of the knee in adults which has been quoted to be 2.8-2.9%. Our study therefore also highlights the rarity of enchondromas in skeletally immature individuals which is in contrast to adults where enchondromas are frequently observed.

We cannot exclude that some of the patients in our study could potentially develop displacement of cartilage islands into the growth plate in the future or may have developed displacement of cartilage islands not depicted on conventional MRI. Hence, the presented study does not refute the above stated theory. Our study findings and the investigation and subsequent rediscovery of the very origin of the theory justifies raising doubts about the validity of the widely believed but unproven theory. Furthermore, the discovery that a majority of enchondromas are associated
with mutations in the genes IDH1 and IDH2 strongly favours that enchondromas are neoplastic lesions rather than developmental abnormalities.³³
Figure 27:
Figure 27 (Illustrations taken from the article: “Über die Entstehung des Enchondroma und seine Beziehungen zu der Ecchondrosis und der Exostosis cartilaginea” published in “Monatsberichte der Königlich Preussischen Akademie der Wissenschaften” in 1875 by R. Virchow)

In figure 27.1 Virchow uses an image of pathology specimen in a child with rickets of the distal femur to demonstrate proliferation of cartilage of the growth plate with islands of cartilage which have “separated completely and lie backwards within the fully developed spongiosa” (translated from German into English by me).

In figure 27.2 Virchow shows an image of a pathology specimen in a child with rickets of the distal femur to demonstrate “a large and a smaller completely separated island of proliferating cartilage which lies within the spongiosa” (translated from German into English by me).

In figure 27.3 Virchow demonstrates an image of a pathology specimen of a skeletally mature patient with an enchondroma of the distal femur. In his publication, he states that the location of the enchondroma “correlates so precisely with the location of the islands of cartilage observed in the skeletally immature bones (of figure 27.1 and 27.2) that one cannot have any doubt, that such a remnant of cartilage really persisted (into adulthood)” (translated from German into English by me).
The diagnosis and grading of chondrosarcomas represent major challenges for pathologists, clinicians and radiologists alike. Particularly, the differentiation of enchondromas from low-grade chondrosarcomas is one of the most difficult and controversial diagnostic dilemmas in musculoskeletal oncology. Whilst anatomical MRI has been used in attempt to differentiate between the two disease entities, there remains considerable uncertainty in the diagnosis.

In the fourth article (original article), we have attempted to use diffusion-weighted MRI (DWI), - a non-invasive functional MRI-technique which indirectly assesses the cellularity of biologic tissues - in an attempt to differentiate enchondromas from chondrosarcomas. A previous retrospective study stated that DWI might be promising in the differentiation of enchondromas from low-grade chondrosarcomas. However, the results of this study were hampered by a small sample size which included only 14 central chondroid tumours. In this study, the authors measured mean ADC-values in three consecutive slices and correlated the mean ADC-values with the histological grading of the chondroid tumours. Whilst the authors found a statistically significant difference in the ADC-values of enchondromas from chondrosarcomas, they concluded that there was only a slight difference in the ADC-values of enchondromas from chondrosarcomas and therefore a significantly larger patient cohort was required to evaluate the role that DWI may play in the differentiation of enchondromas from chondrosarcomas and low-grade chondrosarcomas in particular.89

The aim of our study was therefore to investigate the potential role that DWI could play in the differentiation of enchondromas from chondrosarcomas in general and in
the differentiation of enchondromas from low-grade chondrosarcomas in a larger patient cohort. In the fourth article, which forms part of this PhD-thesis, we therefore re-assessed the role of DWI in the differentiation of enchondromas from chondrosarcomas in a larger patient cohort. We included 52 central chondroid bone tumours of which there 24 enchondromas, 5 cartilaginous lesions of unknown malignant potential, 15 grade 1 chondrosarcomas, three grade 2 chondrosarcomas, two grade 3 chondrosarcomas and three dedifferentiated chondrosarcomas. We found that there was a statistically significant difference in the mean and minimum ADC values in all groups. Post hoc analysis demonstrated that this difference in mean and minimum ADC values was due to the ADC values in the dedifferentiated chondrosarcoma group. In contrast, the mean and minimum ADC-values of low-grade chondroid tumours and high-grade chondrosarcomas were not statistically significantly different. We therefore concluded that DWI could not differentiate between enchondromas and chondrosarcomas and that it does not aid in the distinction of low-grade chondroid tumours from high-grade chondrosarcomas. This study which forms part of the portfolio of publications submitted for consideration of a PhD therefore significantly advances the understanding about the role of Diffusion-weighted MRI in the differentiation of enchondromas from chondrosarcomas.

There are several limitations to our study.

Firstly, the retrospective nature of the study and the fact that the study was performed in a bone tumour centre most likely has resulted in a selection bias as many enchondromas observed on routine MRI-examinations are not referred to a
bone tumour centre where the study was performed. Secondly, whilst 81% of all central cartilage tumours were histologically confirmed, ten enchondromas did not undergo histological confirmation representing 19% of all central cartilaginous tumours and 42% of all enchondromas in our study. We have applied stringent criteria for designating a lesion as an enchondroma based on widely accepted imaging features.

Thirdly, minimum and mean ADC values were measured in one slice. This slice was carefully selected to correspond with the slice on conventional MR imaging, which showed the most aggressive imaging features. Furthermore, there is no standardised technique in the literature which has been widely accepted for analysis of DWI-data. Whilst some authors have analysed DWI-data using a whole-tumour volume approach, analysis of DWI data using the region of interest approach continues to be most widely performed technique similar to the analysis method used in our publication. Although analysis of skeletal lesions using a whole tumour volume approach is likely to be more accurate taking into consideration heterogeneity of ADC-values within skeletal lesions, this technique is very time-consuming, not widely performed and very difficult to implement in routine clinical practice due to its impracticality and time-consuming nature. Thus, the aim of our data analysis was to investigate an analysis method which could be easily implemented in routine clinical practice and hence if found to be useful could be widely adopted. Whilst there have been no studies which compared the different analysis methods of DWI in bone tumours, there has been a very recent study which evaluated the interobserver variability of three selective region-of-interest
measurement protocols for apparent diffusion coefficient quantifications in soft
tissue tumours and compared them with whole tumour volume ADC measurements.
The authors found that whilst all selective and whole tumour volume measurements
offered good to excellent interobserver agreement, the selective observer based
method of ADC measurement resulted in the closest values to whole tumour volume
measurements. The authors therefore concluded that a simplified observer-based
manual method of ADC quantification in the evaluation of soft tissue tumours was
comparable to whole tumour volume measurements of the minimum and mean
ADCs, and required significantly less analysis time. Further research is required to
evaluate which analysis method in bone tumours and skeletal lesions are most
feasible and accurate in the interpretation of ADC-values in daily clinical practice. This
is of importance if DWI is to be implemented as a routine MRI-sequence in daily
clinical practice.

Furthermore, the inter-observer reliability of measurement of ADC-values has not
been tested in this article because both the slice selection and the measurement of
ADC-values was performed by one reader only. Therefore, the reliability of the study
remains untested. Lastly, we did not calculate 95% confidence intervals for the
minimum and mean ADC-values. However, the measured sample mean values in the
study may not reflect the population mean values. Therefore, the range of ADC
values which contain the true population mean with 95% certainty cannot be
quantified based on the study results provided.

The fifth article (original article) presented as part of the submitted PhD-thesis
represents an original article which evaluates the role of MRI in the grading of
chondrosarcomas. Whilst a previous publication showed that the presence of a soft tissue mass favoured the diagnosis of a high-grade chondrosarcoma, the utility of this study was hampered by the fact that the authors only assessed a few MRI-features which are associated with chondrosarcomas and by the small sample size of the study.\(^4\) Therefore, there remained uncertainty about the role of other MR Imaging features in the differentiation of high-grade from low-grade appendicular chondrosarcomas. Hence, we evaluated a wide range of MRI-features, which are used in the characterization of chondrosarcomas representing the largest study published on this topic. In this retrospective study, we analysed the MRI-examinations of 179 chondrosarcomas of long bones and divided them into low-grade chondroid lesions and high-grade chondrosarcomas. We found that on multivariate analysis, bone expansion, active periostitis, the presence of a soft tissue mass, and tumour length are statistically significant differentiating factors between low-grade and high-grade chondroid tumours. On logistic regression analysis, the Odds ratio for bone expansion, active periostitis, soft tissue mass and tumour length were: 8.8 (95% confidence interval 2.4-32.4), 52.8 (5-562.2), 21.1 (4-111.1) and 1.4 (1.2-1.7) for the diagnosis of a high-grade chondrosarcoma. The above stated figures in the study demonstrate wide confidence intervals for all four MRI-features which could be due to the variation of MRI-features in the high-grade chondrosarcoma group although the sample size albeit large in our study could also be a contributing factor. However, it is important to note that the study demonstrated a positive relationship between the above described MRI features and the diagnosis of a high-grade chondrosarcoma. Our study also demonstrated that the presence of the above described four MRI features showed a diagnostic accuracy of 95.6%. We therefore
concluded that these imaging features accurately predicted the presence of a high-grade chondrosarcoma. Thus, MRI can be used accurately to differentiate low-grade chondroid tumours from high-grade chondrosarcomas of the major long bones which in turn may avoid inadequate surgery. This is of particular clinical importance when one considers that biopsy may result in sampling errors of chondrosarcoma of long bone of up to 17% and hence may result in inappropriate treatment.\textsuperscript{44}

There are nevertheless limitations to our study.

Firstly, due to the retrospective nature of our study, the MRI protocol was not standardised. In view of the rarity of chondrosarcomas, there have been no prospective studies evaluating the role of MRI in the diagnosis and grading of chondrosarcomas. Furthermore, our article continues to be one of the largest studies evaluating the role of MRI in chondrosarcoma hence highlighting challenges faced in the development and conduct of potential prospective studies which would require inclusion of a large patient cohort in order to evaluate MRI-features in the differentiation of low-grade from high-grade chondrosarcomas.

Secondly, our study included a higher number of high-grade chondrosarcomas than is usually observed in other series which was most likely due to the tertiary nature of the institution in which the study was conducted. This may therefore have led to a selection bias.

Thirdly, the differentiation of enchondromas from low-grade chondrosarcomas demonstrate a low reliability even among subspecialised histopathologists due to the lack of a gold standard in the distinction of the two disease entities.\textsuperscript{67,68} The difficulty
that radiologists, pathologists and clinicians face in the differentiation of enchondromas from low-grade chondrosarcomas was also reflected in our study by the fact that our study included 28 “borderline malignant chondroid lesions” which were classified as grade 0.5 chondrosarcomas. All chondral tumours at the institution in which the study was performed were reviewed by two musculoskeletal pathologists and the diagnosis of benign versus malignant chondral tumours was made in consensus with radiologists, pathologists and clinicians. Furthermore, the author of this thesis performed an unpublished analysis at the time when the original analysis was performed which excluded the above stated 28 “borderline malignant chondroid lesions” or CLUMPs. The results of this analysis demonstrated that on multivariate analysis, bone expansion, active periostitis, soft tissue mass and tumour length continued to be statistically significant differentiating factors whilst the other MRI-features continued to be not statistically significant. Therefore, the inclusion of 28 indeterminate chondroid lesions has not resulted in a significant change in the findings of the publication.

Lastly, the inter-observer reliability of the above described conventional MRI-findings remains untested as the analysis of the MRI-features was performed in consensus between the two readers and in case of disagreement the opinion of the most senior reviewer was accepted. However, this approach to the data analysis precludes assessment of inter-observer reliability of the above described MRI-features.

In conclusion, the above described study identified several MRI features which allow accurate differentiation of low-grade chondroid tumours from high-grade chondrosarcomas of long bones. Therefore, this study significantly enhances the
understanding and knowledge that MRI plays in the accurate preoperative grading of chondrosarcomas of long bones. Furthermore, the findings of this study may result in a more accurate diagnosis prior to surgery and may significantly improve patient care.\textsuperscript{51}

The sixth article (original article) included in the submitted PhD-thesis evaluates the role of staging in chondrosarcomas. Prior to this publication, it has been widely assumed that skeletal scintigraphy and Computed Tomography of the chest are mandatory in the initial staging of chondrosarcomas to assess for the presence of skeletal and pulmonary metastases. This management approach however has been based on evidence obtained from other sarcomas such as osteosarcoma and Ewing sarcoma which demonstrate skeletal and pulmonary metastases in 4-19\% at initial presentation. The biological behaviour of chondrosarcomas is very different to osteosarcomas and Ewing sarcomas which tend to be more aggressive. Hence, this management approach was not evidence-based.

We have therefore investigated the role of whole-body skeletal scintigraphy and Computed Tomography of the chest in the initial surgical staging of chondrosarcomas. In this retrospective study, we evaluated the bone scintigraphy reports and bone scintigraphs in 188 patients with chondrosarcomas. We discovered 195 chondrosarcomas in 188 patients. In our study, there were no patients with skeletal metastases at presentation. However, we discovered 3 patients with multifocal chondrosarcomas, 2 of whom had known Ollier’s disease and who would have been adequately imaged on the staging MRI-study. A third patient with multifocal chondrosarcoma was found to have a low-grade central chondrosarcoma
in the opposite femur. This finding did not alter surgical management as the lesion was observed and remained unchanged. We therefore concluded that there is little role of skeletal scintigraphy in the initial surgical staging of chondrosarcomas. The importance of this publication also lies in demonstrating that unlike in osteosarcoma and Ewing sarcoma, skeletal metastases in chondrosarcomas at presentation are very rare. This is of significance because previous studies have demonstrated that incidental enchondromas are commonly observed on routine MRI-examinations of the knee and shoulder.\textsuperscript{6,7} Since publication of our findings, the rarity of skeletal metastases in chondrosarcoma has been corroborated by Gulia et al. who evaluated 69 patients with chondrosarcoma and identified only one patient with skeletal metastases (1.6\% of all cases). Of note is, that in their study the single patient with skeletal metastases was also found to have concomitant pulmonary metastases. In contrast, Gulia et al. confirm that the prevalence of pulmonary metastases in chondrosarcoma of bone is significant being observed in 9.8\% of all cases (6 patients) in their patient cohort.\textsuperscript{113}

Similarly, a multicentre retrospective study published by Andreou et al. recently, evaluated the metastatic potential of grade 1 chondrosarcomas. The authors included 225 patients with grade 1 chondrosarcoma and found that 14 patients developed metastases after a median time of 49 months with a range of 4 to 125 months. In their study, only two patients (0.9\% of all patients) developed skeletal metastases whilst 12 patients (5.3\% of all patients) developed pulmonary metastases. This recent study further supports our findings that skeletal metastases
are a rare occurrence in chondrosarcoma of bone in contrast to pulmonary metastases.\textsuperscript{114}

We also assessed the CT-chest reports of all patients with chondrosarcoma at the time of diagnosis. In our study, we identified pulmonary metastases in approximately 5\% of chondrosarcomas at initial presentation. In contrast to skeletal scintigraphy, we found therefore evidence supporting the added value of Computed Tomography of the chest in the detection of pulmonary metastases in the initial staging of chondrosarcomas.

There are limitations to our study.

Firstly, of the 352 patients identified on the database with a chondrosarcoma, 154 did not have bone scintigraphy images or reports available for review. Whilst review of clinical letters in these patients made no reference about the presence of skeletal metastases, we were therefore not in a position to definitely rule out that some of these patients may have presented with occult skeletal metastases.

Secondly, of the 195 chondrosarcomas included in the study, only bone scintigrams, which demonstrated increased uptake outside the site of the chondrosarcoma according to the bone scintigraphy report were re-reported by the first author of the article. Whilst review of the clinical notes made no reference to the presence of skeletal metastases, we are again unable to rule out that some of these patients may have had occult skeletal metastases. Furthermore, the inter-observer reliability of the findings of the re-reported bone scintigrams has not been tested because bone scintigraphy studies were re-reported by the author of this thesis only.
Thirdly, in our patient cohort, three patients were diagnosed with synchronous multifocal chondrosarcomas. Whilst the presence of multifocal synchronous chondrosarcomas could be interpreted as skeletal metastases, two of these patients suffered from Ollier’s disease which is characterised by multifocal enchondromas in which malignant transformation is a well-recognised complication therefore strongly favouring that the presence of further chondrosarcomas is due to multiple chondrosarcomas rather than skeletal metastases. In contrast to the other two cases, the third patient with synchronous multifocal chondrosarcomas was not known to suffer from Ollier’s disease as there were no further manifestations on other imaging to suggest multiple enchondromas. The existence of multicentric primary chondrosarcomas in the absence of enchondromatosis or hereditary multiple exostosis remains uncertain. The third patient in our cohort was found to have a grade 2 chondrosarcoma of the left pubis, a grade 2 chondrosarcoma of the left proximal femur and a grade 1 chondrosarcoma of the right proximal femur. We therefore cannot exclude that one of the lesions in the left femur or pelvis represented a skip metastasis. The preoperative staging MRI-study accurately depicted both lesions. In contrast, the grade 1 chondrosarcoma of the contralateral right proximal femur was only depicted on the bone scintigraphy study. However, this lesion in view of its lower grade most likely represented a primary chondrosarcoma rather than a skeletal metastasis.

Although some clinicians may wish to continue performing skeletal scintigraphy or may even consider whole-body MRI in the initial staging of chondrosarcomas, our findings highlight the rarity of skeletal metastases in this disease entity. Hence, the
presence of other adjacent chondroid lesions on skeletal scintigraphy or whole-body MRI should not be misinterpreted as skip metastases. This is of particular importance when one considers that incidental enchondromas are identified in approximately 2% of all MRI-scans of the shoulder and in approximately 3% of all MRI-scans of the knee.\textsuperscript{6,7}

In conclusion, this study, which forms part of the PhD-thesis demonstrates the rarity of skeletal metastases in chondrosarcoma of bone at presentation and highlights the significant prevalence of pulmonary metastases. This study therefore significantly advances the knowledge about the role of skeletal scintigraphy and Computed Tomography of the chest in the initial staging of chondrosarcoma of bone.

In summary, the authors’ published work submitted for consideration of a PhD, investigates the theory that imaging may aid in the understanding of the pathogenesis of enchondromas and may aid in the diagnosis, grading and staging of chondrosarcomas of bone.
8.2 Future research

Whilst the author’s work has significantly advanced the understanding about the pathogenesis of enchondromas, the role of imaging in the diagnosis, grading and staging of chondrosarcomas, there remains diagnostic uncertainty in the differentiation of enchondromas from low-grade chondrosarcomas.

Although the author’s most recent publication demonstrated that the presence of more than 2/3 of endosteal scalloping on conventional MR shows a sensitivity of 71.4% and a specificity of 92.9% in the differentiation of enchondromas from low-grade chondrosarcomas and hence is the most sensitive feature on conventional MRI, the differentiation of the two disease entities remains challenging. Similarly, diffusion-weighted MRI cannot be used to differentiate between enchondromas and low-grade chondrosarcomas as demonstrated by the author. Furthermore, whilst dynamic-contrast enhanced MRI aids in the detection of high-grade chondrosarcomas, DCE-MRI crucially cannot be utilized to differentiate enchondromas from low-grade chondrosarcomas as published by the author of this thesis very recently (see Appendix 1)."}

Other imaging techniques such as radiography, CT and PET/CT are of limited value in the distinction of the two disease entities. Other functional imaging modalities such as conventional MR-spectroscopy and hyperpolarized MRI have not been evaluated in the diagnosis of enchondromas and low-grade chondrosarcomas. These functional imaging techniques have the potential to allow unique and fascinating insights into the tumour biology, into the molecular and genetic make-up of tumours in general.
and of chondroid tumours in particular. It has been shown relatively recently that a
majority of enchondromas and chondrosarcomas have somatic mutations in
isocitrate dehydrogenase 1 and 2 (IDH 1 and 2) which produce the oncometabolite
2-hydroxyglutarate. This oncometabolite could therefore potentially be identified
and the concentration of this oncometabolite could be quantified non-invasively
using conventional MR-spectroscopy or hyperpolarized MRI. Quantification of this
oncometabolite could therefore possibly result in a cut-off value which could
differentiate enchondromas from low-grade chondrosarcomas more reliably. These
imaging techniques may therefore potentially hold the key to unravel the diagnostic
uncertainty in the distinction of enchondromas from low-grade chondrosarcomas.

Whilst molecular MRI-techniques warrant further research, at present, the
differentiation of enchondromas from low-grade chondrosarcomas continues to be
challenging. However, a very recent publication by the author of this thesis
demonstrated that certain clinical and conventional MRI-features aid in the
distinction of the two disease entities. In particular, the presence of more than 2/3
of endosteal scalloping on conventional MRI and pain attributed to the lesion
demonstrate a moderate sensitivity in the differentiation of enchondromas from
low-grade chondrosarcomas whilst the presence of pain attributed to the lesion and
more than 2/3 of endosteal scalloping are highly specific for a grade 1
chondrosarcoma. The authors’ findings however require validation in a larger
prospective patient cohort. Hence, further research is required to evaluate the role
of conventional MRI in the differentiation of the two disease entities.
At present, there remains considerable diagnostic uncertainty in the differentiation of enchondromas from low-grade chondrosarcomas. In particular, the malignant potential of central chondroid lesions which do not demonstrate permeation on histopathology remains unknown. Hence, a major recommendation of this thesis is that future studies should focus on the malignant potential of low-grade central chondroid lesions which are not treated. Therefore, prospective long-term follow-up studies should be performed to assess the malignant potential of such low-grade chondroid lesions and to assess potential predictive imaging biomarkers. Surgery remains the only hope for cure in chondrosarcoma and the presence of skeletal metastases is associated with a bleak outcome. Advances in the molecular understanding of chondrosarcomas are likely to result in the development of non-surgical therapies such as agents which specifically target mutations in chondrosarcoma cells. Therefore, the above described molecular imaging techniques may in future also allow the development of a personalized, molecular fingerprint of chondrosarcomas which in turn may allow non-invasive monitoring of treatment response in patients with chondrosarcoma beyond the current concept of conventional, anatomical imaging thus potentially resulting in a paradigm shift in imaging. Furthermore, molecular imaging may facilitate the development of new drugs which combine diagnostic and therapeutic capabilities (so-called theranostic drugs). In particular, the efficacy of drugs targeting IDH 1 and 2 mutations in chondrosarcomas could be combined with in-vivo and real-time assessment of treatment response in these patients using the above described functional imaging techniques.
Therefore, future imaging research in chondrosarcomas should focus on the role of conventional MRI in the differentiation of enchondromas from low-grade chondrosarcomas. However, imaging research should also investigate the potential role that the non-invasive detection and quantification of oncometabolites produced by genetic mutations may play in the diagnosis and treatment response assessment of chondrosarcomas.


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Appendix 1: What are the differentiating clinical and MRI-features of enchondromas from low-grade chondrosarcomas?

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Abstract
Objectives To evaluate the role of clinical assessment, conventional and dynamic contrast-enhanced MRI in differentiating enchondromas from chondrosarcomas of long bone.
Methods The following clinical and MRI findings were assessed: age, gender, pain, pain attributable to lesion, tumour location, tumour length, presence, depth of endosteal scalloping, bone marrow oedema, soft tissue oedema, cortical destruction, periosteal reaction, bone expansion, macroscopic fat, calcification, soft tissue mass, haemorrhage, dynamic contrast-enhanced MRI. Clinical and MRI findings were compared with histopathological grading.
Results Sixty patients with central chondroid tumours were included (27 enchondromas, 10 cartilaginous lesions of unknown malignant potential, 15 grade 1 chondrosarcomas, 8 high-grade chondrosarcomas). Pain attributed to lesion, tumour length, endosteal scalloping > 2/3, cortical destruction, bone expansion and soft tissue mass were differentiating features between enchondromas and grade 1 chondrosarcomas. Dynamic contrast-enhanced MRI could not differentiate enchondromas from grade 1 chondrosarcomas.
Conclusions Previously reported imaging signs of chondrosarcomas are useful in the diagnosis of grade 1 lesions but have lower sensitivity than in higher grade lesions. Deep endosteal scalloping is the most sensitive imaging sign of grade 1 chondrosarcomas. Pain due to the lesion is an important clinical sign of grade 1 chondrosarcomas. Dynamic contrast-enhanced MRI is not useful in differentiating enchondromas from grade 1 chondrosarcomas.

Key Points
- Differentiation of enchondroma from low-grade chondrosarcoma is challenging for radiologists and pathologists.
- The utility of clinical assessment, conventional and dynamic contrast-enhanced MRI was uncertain.
- Clinical assessment and conventional MRI aid in differentiating enchondromas from low-grade chondrosarcoma.
- Dynamic contrast-enhanced MRI cannot differentiate enchondromas from grade 1 chondrosarcoma.

Keywords Conventional MRI · Dynamic contrast-enhanced MRI · Enchondroma · Chondrosarcoma · Pain

Abbreviations and acronyms
CS Chondrosarcoma
CLUMP Cartilaginous lesion of unknown malignant potential
DCE MRI Dynamic contrast-enhanced MRI
Introduction

Chondrosarcoma (CS) of bone is the second most common malignant primary bone tumour after osteosarcoma [1]. Similarly, its benign counterpart, the enchondroma, represents the second most commonly observed primary benign bone tumour after osteochondroma, and is frequently seen on routine MRI-examinations of the knee and shoulder [2-4]. The differentiation of enchondromas from low-grade chondrosarcomas is one of the most difficult distinctions for radiologists, pathologists and clinicians, alike. The reason for this diagnostic uncertainty in the differentiation of the two disease entities on histopathology is the lack of a gold standard for the diagnosis of chondrosarcomas [5, 6]. Whilst the presence of permeation on histopathology is diagnostic for a chondrosarcoma, it is the lack of permeation in cartilaginous tumours with imaging findings, atypical for an enchondroma (e.g. size > 5 cm), which results in the diagnostic conundrum [6-8]. In the absence of clear cut diagnostic criteria on histopathology, it is frequently stated that the differentiation between the two disease entities is based on a consensus decision between radiology, histopathology and clinical findings [7]. However, this approach may result in potential over- or under-diagnosis of chondrosarcomas based on the different diagnostic criteria which may significantly differ from centre to centre. This lack of clarity in the differentiation of enchondromas from low-grade chondrosarcomas may therefore be a confounding factor in previous studies published on this topic and may explain the discrepancy in findings between various centres. Similarly, there have been various, sometimes conflicting, reports about the role of MRI in the differentiation of enchondromas from low-grade chondrosarcomas [8-12]. This again may be due to lack of clarity in the histopathological diagnosis of chondrosarcomas, as previous studies did not state how the diagnosis of a chondrosarcoma was established.

The aim of this article is therefore to evaluate the utility of conventional and dynamic contrast-enhanced MRI (DCE-MRI) as well as the role of clinical assessment in the differentiation of enchondromas from chondrosarcomas of the long bones with particular emphasis in the distinction of enchondromas from low-grade chondrosarcomas of long bones, using the most stringent inclusion criterion for the diagnosis of a chondrosarcoma: the presence of a permeative growth pattern on histopathology.

Material and methods

Patients

Institutional Review Board approval was obtained before commencement of the study to perform this retrospective study (ROH 16-021)) as a service evaluation. All patients with central chondroid tumours of long bone who underwent conventional MRI and dynamic contrast-enhanced MRI at our institution were identified in this retrospective study. Data were collected on patient age, gender, tumour location, MRI-findings, patient symptoms and histological grading of the chondral lesions. Patients were included if the MR-findings were pathognomonic for an enchondroma based on widely accepted MRI-criteria [8] or if a biopsy, curettage or resection was performed confirming the diagnosis of either an enchondroma, a CLUMP or chondrosarcoma.

In total, there were 60 patients with central chondroid tumours of long bone. There were 27 patients with an enchondroma, 10 patients with a CLUMP, 15 patients with a grade 1 CS and 8 patients with high-grade CS. The HG-CS group consisted of 1 grade 3 CS, 4 dedifferentiated CS and 3 grade 2 CS.

A detailed description of the materials and methods including the MRI-protocol, the image data analysis, clinical findings, histopathology findings, the correlation of histopathology with MRI and clinical findings as well as the methods of the statistical analysis can be found in the supplementary document.

Results

The demographics, presence of pain, pain attributed to the lesion, lesion location and lesion length of enchondromas, CLUMPs, grade 1 CS and high-grade CS are summarised in Table 1. Only pain attributed to the lesion and tumour length were statistically differentiating factors between enchondromas and grade 1 CS.

The presence and depth of endosteal scalloping, bone marrow oedema, soft tissue oedema, cortical destruction, periosteal reaction, bone expansion, soft tissue mass, macroscopic fat, haemorrhage, calcification and the severity of calcification in the enchondroma-, CLUMP-, grade 1 CS and high-grade CS are illustrated in Table 2. Only depth of endosteal scalloping of more than 2/3 (= grade 3 endosteal scalloping), cortical destruction, bone expansion and the presence of a soft tissue mass were differentiating features between enchondromas and grade 1 CS.

The various DCE-MRI parameters (angle of the DCE-MRI curve, absolute enhancement and relative enhancement on DCE-MRI) in patients with enchondromas, CLUMPs, grade 1 CS and high-grade CS are illustrated in Table 3. None of the above mentioned DCE-MRI parameters was statistically
<table>
<thead>
<tr>
<th>Demographics</th>
<th>Total (60)</th>
<th>Enchondroma (27)</th>
<th>CLUMPs (10)</th>
<th>Grade 1 CS (15)</th>
<th>HG-CS (8)</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Male</td>
<td>41</td>
<td>5</td>
<td>6</td>
<td>6</td>
<td>2</td>
<td>0.091</td>
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<tr>
<td>Female</td>
<td>19</td>
<td>22</td>
<td>4</td>
<td>9</td>
<td>6</td>
<td></td>
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<tr>
<td>Median age (y)</td>
<td>50.5</td>
<td>49</td>
<td>54</td>
<td>49</td>
<td>64</td>
<td>0.067</td>
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<tr>
<td>Presence of pain (%)</td>
<td>40 (66.7%)</td>
<td>16 (59.3%)</td>
<td>6 (60%)</td>
<td>11 (73.3%)</td>
<td>7 (87.5%)</td>
<td>0.46</td>
</tr>
<tr>
<td>Pain thought to be due to the lesion (%)</td>
<td>20 (33.3%)</td>
<td>3 (11.1%)</td>
<td>1 (10%)</td>
<td>9 (60%)</td>
<td>7 (87.5%)</td>
<td>0.000**</td>
</tr>
<tr>
<td>Scoleral distribution</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humerus</td>
<td>29</td>
<td>15</td>
<td>4</td>
<td>8</td>
<td>2</td>
<td>0.41</td>
</tr>
<tr>
<td>Femur</td>
<td>20</td>
<td>6</td>
<td>5</td>
<td>4</td>
<td>5</td>
<td></td>
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<tr>
<td>Tibia</td>
<td>10</td>
<td>6</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td></td>
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<tr>
<td>Fibula</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
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<tr>
<td>Osseous location (1)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Proximal</td>
<td>34</td>
<td>19</td>
<td>5</td>
<td>8</td>
<td>2</td>
<td>0.306</td>
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<td>Distal</td>
<td>21</td>
<td>8</td>
<td>5</td>
<td>4</td>
<td>4</td>
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<tr>
<td>Proximal and distal</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>2</td>
<td></td>
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<tr>
<td>Osseous location (2)</td>
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<td></td>
<td></td>
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<tr>
<td>Diaphysis</td>
<td>22</td>
<td>10</td>
<td>8</td>
<td>3</td>
<td>1</td>
<td>0.024*</td>
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<tr>
<td>Diaphysis and Metaphysis</td>
<td>24</td>
<td>12</td>
<td>2</td>
<td>7</td>
<td>3</td>
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<tr>
<td>Diaphysis, Metaphysis and Epiphysis</td>
<td>8</td>
<td>1</td>
<td>0</td>
<td>4</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Metaphysis and Epiphysis</td>
<td>6</td>
<td>4</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Median lesion length (cm) [range]</td>
<td>4.7 [2.4-14.3]</td>
<td>7.9 [6.6-12]</td>
<td>10.5 [2.1-26.2]</td>
<td>12.5 [4.3-29.6]</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Due to differences in lesion length between CLUMPS group and HG-CS group (p = 0.048).

**Subgroup analysis showed that this was due to differences in tumour length between enchondroma vs grade 1 CS-group (p = 0.006).
Table 2  Presence of endosteal scalloping, depth of endosteal scalloping, presence of bone marrow oedema, soft tissue oedema, cortical destruction, periosteal reaction, bone expansion, soft tissue mass, macroscopic fat, haemorrhage, calcification and the severity of calcification in patients with enchondromas, CLUMPs and grade 1 CS and high-grade CS

<table>
<thead>
<tr>
<th>MRJ Features</th>
<th>Total (60)</th>
<th>Enchondroma (27)</th>
<th>CLUMP (10)</th>
<th>Grade 1 CS (15)</th>
<th>HG-CS (8)</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endosteal scalloping present (%)</td>
<td>50 (83.3%)</td>
<td>20 (74%)</td>
<td>7 (70%)</td>
<td>15 (100%)</td>
<td>8 (100%)</td>
<td>p = 0.000*</td>
</tr>
<tr>
<td>Depth of endosteal scalloping (%)</td>
<td>10 (16.7%)</td>
<td>7 (25.9%)</td>
<td>3 (30%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>p = 0.015, due to differences in the enchondroma vs grade 1 CS group (p = 0.001), due to differences in the CLUMP-group vs grade 1 CS group (p = 0.037) and due to differences in CLUMP-group vs HG-CS group (p = 0.002)</td>
</tr>
<tr>
<td>Grade 1</td>
<td>20 (33.3%)</td>
<td>10 (37%)</td>
<td>5 (50%)</td>
<td>5 (33.3%)</td>
<td>0 (0%)</td>
<td></td>
</tr>
<tr>
<td>Grade 2</td>
<td>9 (15%)</td>
<td>8 (29.6%)</td>
<td>1 (1%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td></td>
</tr>
<tr>
<td>Grade 3</td>
<td>21 (35%)</td>
<td>2 (7.4%)</td>
<td>1 (1%)</td>
<td>10 (66.7%)</td>
<td>8 (100%)</td>
<td></td>
</tr>
<tr>
<td>Bone marrow oedema (%)</td>
<td>21 (35%)</td>
<td>8 (29.6%)</td>
<td>0 (0%)</td>
<td>7 (46.7%)</td>
<td>6 (72%)</td>
<td>p = 0.04*, due to differences in the CLUMP-group vs HG-CS group (p = 0.012)</td>
</tr>
<tr>
<td>Soft tissue oedema (%)</td>
<td>20 (33.3%)</td>
<td>1 (4.8%)</td>
<td>0 (0%)</td>
<td>5 (33.3%)</td>
<td>8 (100%)</td>
<td>p = 0.000*</td>
</tr>
<tr>
<td>Cortical destruction</td>
<td>13 (21.7%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>5 (33.3%)</td>
<td>8 (100%)</td>
<td>p = 0.024, due to differences in the enchondroma vs grade 1 CS group (p = 0.006), due to differences in the CLUMP-group vs grade 1 CS group (p = 0.036) and due to differences in the CLUMP-group vs HG-CS group (p = 0.000)</td>
</tr>
<tr>
<td>Periosteal reaction</td>
<td>14 (23.3%)</td>
<td>1 (3.7%)</td>
<td>0 (0%)</td>
<td>5 (33.3%)</td>
<td>8 (100%)</td>
<td>p = 0.000*</td>
</tr>
<tr>
<td>Bone expansion</td>
<td>15 (25%)</td>
<td>2 (7.4%)</td>
<td>0 (0%)</td>
<td>7 (46.7%)</td>
<td>6 (75%)</td>
<td>p = 0.012, due to differences in the enchondroma vs grade 1 CS group (p = 0.000), due to differences in the CLUMP-group vs HG-CS group (p = 0.000)</td>
</tr>
<tr>
<td>Soft tissue mass</td>
<td>14 (23.3%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>6 (40%)</td>
<td>8 (100%)</td>
<td>p = 0.000*</td>
</tr>
<tr>
<td>Macroscopic fat</td>
<td>44 (73.3%)</td>
<td>24 (89%)</td>
<td>9 (90%)</td>
<td>9 (60%)</td>
<td>2 (33.3%)</td>
<td>p = 0.000*</td>
</tr>
<tr>
<td>Haemorrhage</td>
<td>2 (3.3%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>2 (25%)</td>
<td>p = 0.47, due to differences in the enchondroma vs HG-CS group (p = 0.006)</td>
</tr>
<tr>
<td>Calcification</td>
<td>55 (91.7%)</td>
<td>26 (96.3%)</td>
<td>9 (90%)</td>
<td>13 (86.7%)</td>
<td>7 (87.5%)</td>
<td></td>
</tr>
<tr>
<td>Severity of calcification</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mild</td>
<td>16 (26.7%)</td>
<td>6 (22.2%)</td>
<td>2 (20%)</td>
<td>6 (40%)</td>
<td>2 (25%)</td>
<td>p = 0.44</td>
</tr>
<tr>
<td>Moderate</td>
<td>28 (46.7%)</td>
<td>13 (48.2%)</td>
<td>4 (40%)</td>
<td>7 (46.7%)</td>
<td>4 (50%)</td>
<td></td>
</tr>
<tr>
<td>Severity</td>
<td>11 (18.3%)</td>
<td>7 (25.9%)</td>
<td>3 (30%)</td>
<td>0 (0%)</td>
<td>1 (12.5%)</td>
<td></td>
</tr>
</tbody>
</table>
significant in the differentiation of enchondromas from grade 1 CS.

Figures 1, 2, 3 and 4 demonstrate typical conventional and DCE MR-images of enchondroma (Fig. 1), a CLUMP (Fig. 2), grade 1 CS (Fig. 3) and high-grade CS (Fig. 4) in our study.

Subsequently, we calculated the sensitivity, specificity, positive predictive value and negative predictive value (see Table 4) and the Odds ratio (see Table 5) for grade 3 endosteal scalloping, cortical destruction, bone expansion, soft tissue mass and pain attributed to the lesion in the differentiation of enchondromas from grade 1 CS.

### Discussion

The differentiation of enchondroma from low-grade CS is one of the most challenging and difficult diagnoses in bone sarcoma imaging and pathology alike. This has been highlighted in previous studies which demonstrated a moderate interobserver reliability in the distinction of the two disease entities, even amongst experienced sarcoma radiologists and experienced sarcoma pathologists [5, 6]. The differentiation of enchondromas and grade 1 chordomas is particularly challenging in the long tubular bones such as the femur, tibia and humerus, where enchondromas are very commonly observed. Similarly, the long tubular bones are the most common location for chordosarcomas, accounting for approximately 45% of all cases [7].

A lack of clear cut diagnostic criteria on histopathology is the cause for this diagnostic uncertainty [5, 6]. Hence it is frequently stated that the differentiation between the two disease entities is based on a consensus decision between radiology, histopathology and clinical findings [7, 8]. However, this approach may result in potential over- or underdiagnosis of chordosarcomas based on the different diagnostic criteria in the various sarcoma centres. This is reflected in the fact that in the absence of a permissive growth pattern on histopathology, these lesions are variably termed as large enchondromas, CLUMPS (cartilaginous lesion of unknown malignant potential) or grade 0 CS [13, 14]. This lack of clarity in the differentiation of enchondromas from low-grade CS may therefore be a confounding factor in previous studies published on this topic and may explain the discrepancy in findings between various centres. In contrast, the presence of a permissive growth pattern on histopathology is diagnostic for a chordosarcoma [1, 6].

Distinction of the two disease entities is crucial, however, as enchondromas do not require surgical intervention unless symptomatic, whilst grade 1 CS are invariably treated with surgery, which may either be resection or, increasingly, curettage [15–18].

### Table 3

<table>
<thead>
<tr>
<th>DCE-MRI features</th>
<th>Enchondroma</th>
<th>CLUMP (10)</th>
<th>Grade 1 CS (15)</th>
<th>HG-CS (8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median angle of the curve in degrees</td>
<td>57.8 (8.8 – 80)</td>
<td>35 (17)</td>
<td>71 (30.1–97.9)</td>
<td>72.9 (53.8–96.9)</td>
</tr>
<tr>
<td>$p$ Value due to differences in the angle of the curve between enchondroma vs CLUMP group and grade 1 CS group</td>
<td>$p = 0.0009$</td>
<td>$p = 0.0009$</td>
<td>$p = 0.0009$</td>
<td></td>
</tr>
<tr>
<td>Relative enhancement (range in brackets)</td>
<td>0.85 (0.041–1.6)</td>
<td>0.13 (0.01–1.5)</td>
<td>0.98 (0.037–2.62)</td>
<td>1.6 (0.002–2.76)</td>
</tr>
<tr>
<td>$p$ Value due to differences in absolute enhancement between enchondroma vs HG-CS group</td>
<td>$p = 0.0017$</td>
<td>$p = 0.0027$</td>
<td>$p = 0.0027$</td>
<td></td>
</tr>
<tr>
<td>$p$ Value due to differences in absolute enhancement between CLUMP group vs HG-CS group</td>
<td>$p = 0.0279$</td>
<td>$p = 0.0027$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Pairwise comparison not statistically significant.
Fig. 1 A 44-year-old female patient with an enchondroma of the left proximal tibia. a Coronal STIR MR image shows an enchondroma measuring 3.3 cm in the maximum cranio-caudal dimension. b Axial T1W SE MR image demonstrates foci of chondroid matrix mineralization (small arrow) and islands of entrapped fat (large arrow) but no endosteal scalloping. c, d DCE-MRI demonstrates an angle of the curve of 57.4 degrees and a relative enhancement of 286% when compared to muscle. The patient complained of knee pain, which was thought to be due to medial femorotibial joint osteoarthritis as evidenced on the MRI study (see Figure 1a).

There also remains significant uncertainty about the role of conventional and dynamic contrast-enhanced MRI in the differentiation of enchondromas from low-grade chondrosarcomas. Previous studies have stated the limited utility of MRI in the differentiation of enchondromas from low-grade CS, while other studies have described a particularly important role for dynamic contrast-enhanced MRI in the differentiation of enchondromas from low-grade CS [8, 10–12, 19]. This discrepancy in findings can in part be attributed to the lack of clear diagnostic histopathological criteria that may have been applied, or in the inclusion of low-grade and high-grade chondrosarcomas in these studies.

Murphey et al., e.g., retrospectively analysed the MR imaging of 35 enchondromas and 33 chondrosarcomas of the appendicular skeleton. They evaluated lesion size, endosteal scalloping, cortical remodelling, cortical destruction, pathological fracture, periosteal reaction, cortical thickening and soft tissue extension. The group found that the mean length of enchondromas was 5 cm, whilst the mean length of chondrosarcomas was 8 cm. Similarly, the depth and extent of endosteal scalloping was significantly different in enchondromas and chondrosarcomas, with endosteal scalloping of more than two-thirds being observed in 67% of chondrosarcomas but in only 11% of enchondromas. Furthermore, cortical destruction and soft tissue extension were statistically significant differentiating features between enchondromas and chondrosarcomas, whilst cortical remodelling, pathological fracture, cortical thickening and periosteal...
reaction could not differentiate between the two disease entities on MRI. However, this study included both low-grade CS and high-grade CS. Therefore, the results in their study cannot be utilised to differentiate enchondromas from low-grade CS [8].

In contrast, Ferrer-Santacreu et al. assessed the role of MRI in the differentiation of enchondromas and low-grade chondrosarcomas of the appendicular skeleton in 82 patients. The authors found no statistically significant differentiating imaging features between enchondromas and low-grade chondrosarcomas [11].

Another study by Choi et al. assessed the MRI-features in 16 enchondromas and in 18 low-grade chondrosarcomas. In contrast to Ferrer-Santacreu, the authors found that predominantly intermediate signal intensity on T1WI, a multilobular appearance on contrast-enhanced T1WI, cortical destruction, soft tissue extension, surrounding bone marrow oedema, abnormal soft tissue signal, epiphyseal involvement and location in the flat bone were more commonly observed in low-grade chondrosarcomas than in enchondromas. However, the authors included chondrosarcomas of the flat bones and enchondromas of the hand in their study. However, the differentiation of enchondromas from low-grade chondrosarcomas does not represent a diagnostic challenge in these locations [10].

In contrast, Crim et al. evaluated the utility of MRI and radiography in the differentiation of enchondromas from
Fig. 3 A 64-year-old male patient with a grade 1 chondrosarcoma of the right distal femur. 

(a) Coronal STIR MR image shows a grade 1 chondrosarcoma measuring 8.6 cm in the maximum anteroposterior dimension. 

(b) Axial T1W SE MR image demonstrates more than 2/3 of endosteal scalloping of the posterior cortex (arrow).

c, d DCE-MRI shows an angle of the curve of 50.8 degrees and a relative enhancement of 190% when compared to muscle. The patient complained of lower thigh pain in the region of the lesion, which was thought to be due to the chondroid lesion. Biopsy followed by resection confirmed a well-differentiated cartilage tumour with focal areas of permissive growth in keeping with a grade 1 chondrosarcoma.

Low-grade CS in 53 cases. Their study included 32 enchondromas, 6 borderline-malignant lesions and 12 chondrosarcomas. The authors found that bone expansion and cortical thickening were rare but specific features of chondrosarcoma, and that both radiography and MRI had limitations in the evaluation of low-grade chondroid lesions [12].

De Coninck et al. also evaluated the role of DCE in the differentiation of enchondromas from chondrosarcomas. In their retrospective study, the authors included 75 enchondromas and 31 chondrosarcomas, which included 18 low-grade chondrosarcomas, 10 intermediate grade chondrosarcomas and 3 high grade chondrosarcomas. The authors found that enhancement within the tumour, which was two times more when compared to muscle, combined with a 76 degree slope of the uptake curve resulted in a 100% sensitivity and 63.3% specificity in the detection of chondrosarcomas. However, a significant limitation of this study was the inclusion of low-grade and high-grade chondrosarcomas. Therefore, the obtained results cannot be utilised in the differentiation of enchondromas from low-grade chondrosarcomas [9].

Furthermore, there remains uncertainty about the precise diagnosis of low-grade CS, because in many centres the diagnosis is based on a consensus decision between radiology, histopathology and clinical findings, therefore likely resulting in an incorporation bias. Hence, the role of conventional MRI...
Fig. 4  A 49-year-old male patient with a grade 2 chondrosarcoma of the left proximal tibia.  a  Coronal T1W SE MR image shows a grade 2 chondrosarcoma measuring 4.7 cm in the maximum cranio-caudal dimension that shows cortical destruction and a soft tissue mass (arrow).  b  Axial T1W SE MR image demonstrates cortical destruction, a soft tissue mass, soft tissue oedema (long arrow) and extensive bone marrow oedema (short arrow).  c, d  DCE-MRI demonstrates an angle of the curve of 80.4 degrees and a relative enhancement of 50% when compared to muscle.  The patient complained of resting pain and night pain, which was thought to be due to the chondroid lesion.  Biopsy followed by resection revealed a grade 2 chondrosarcoma and DCE-MRI in the differentiation of the enchondromas from low-grade CS remains unclear.

An important aim of our study was therefore to minimise a potential incorporation bias by utilising the histopathological

<table>
<thead>
<tr>
<th>Significant features</th>
<th>Sensitivity (CI)</th>
<th>Specificity (CI)</th>
<th>PPV (CI)</th>
<th>NPV (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain due to lesion</td>
<td>60% (CI: 32.3%-83.7%)</td>
<td>88.9% (CI: 70.8%-97.7%)</td>
<td>75% (CI: 42.8%-94.5%)</td>
<td>80% (CI: 61.4%-92.3%)</td>
</tr>
<tr>
<td>Grade 3 endosteal scalloping</td>
<td>71.4% (CI: 41.9%-91.6%)</td>
<td>92.9% (CI: 76.5%-99.2%)</td>
<td>83.3% (CI: 51.6%-97.9%)</td>
<td>86.7% (CI: 69.3%-96.2%)</td>
</tr>
<tr>
<td>Bone expansion</td>
<td>50% (CI: 23%-77%)</td>
<td>89.3% (CI: 71.8%-97.7%)</td>
<td>79% (CI: 43.8%-93.3%)</td>
<td>78.1% (CI: 60%-90.7%)</td>
</tr>
<tr>
<td>Cortical destruction</td>
<td>35.7% (CI: 12.8%-64.9%)</td>
<td>100% (CI: 87.7%-100%)</td>
<td>100% (CI: 47.8%-100%)</td>
<td>75.7% (CI: 58.8%-88.2%)</td>
</tr>
<tr>
<td>Soft tissue mass</td>
<td>49% (CI: 16.3%-67.7%)</td>
<td>100% (CI: 87.2%-100%)</td>
<td>100% (CI: 54.1%-100%)</td>
<td>75% (CI: 57.8%-87.9%)</td>
</tr>
</tbody>
</table>
Table 5  Odds ratio for pain attributed to the lesion, grade 3 endosteal scalloping and bone expansion in the differentiation of enchondromas from grade 1 CS

<table>
<thead>
<tr>
<th>Significant features</th>
<th>Odds Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain due to lesion</td>
<td>12 (CI: 2.5-58.5)</td>
</tr>
<tr>
<td>Grade 3 endosteal scalloping</td>
<td>25 (CI: 4.2-157)</td>
</tr>
<tr>
<td>Bone expansion</td>
<td>14.3 (CI: 2.5-85.1)</td>
</tr>
<tr>
<td>Cortical destruction</td>
<td>*</td>
</tr>
<tr>
<td>Soft tissue mass</td>
<td>*</td>
</tr>
</tbody>
</table>

*Odds ratios for cortical destruction and soft tissue mass could not be calculated as these findings were not present in enchondromas.

criterion which is diagnostic for a chondrosarcoma – the presence of a permeative growth pattern. To the best of our knowledge, ours is the first study to evaluate the utility of conventional MRI, DCE-MRI and clinical findings in the differentiation of enchondromas from low-grade CS utilising the presence of a permeative growth pattern as a diagnostic criterion.

We found that tumour length, depth of endosteal scalloping, bone expansion, cortical destruction and pain attributed to the lesion were statistically differentiating features between enchondromas and grade 1 CS. Furthermore, grade 3 endosteal scalloping, cortical destruction, bone expansion, soft tissue mass and pain attributed to the lesion demonstrated a high specificity in the differentiation of enchondromas from grade 1 CS, whilst only grade 3 endosteal scalloping and pain attributed to the lesion demonstrated a moderate sensitivity. In contrast, cortical destruction, bone expansion and soft tissue mass showed only a low sensitivity. Grade 3 endosteal scalloping and bone expansion was observed in two enchondromas. The first case was an eccentrically located enchondroma in the humerus measuring 2.8 cm in which the eccentric location of the lesion was most likely the cause for the grade 3 endosteal scalloping and bone expansion (Fig. 5) [20]. In contrast, the second case of an enchondroma with grade 3 endosteal scalloping and bone expansion was a 4.3 cm centrally located enchondroma within the proximal metaphysis of the proximal humerus. In both cases, the diagnosis of an enchondroma was established histologically after curettage of the lesions.

However, age, gender, lesion location, the mere presence of endosteal scalloping, bone marrow oedema, soft tissue oedema, periosteal reaction, macroscopic fat, calcification, severity of calcification and the presence of intratumoural haemorrhage were not statistically differentiating factors. Moreover, various DCE-MRI such as slope of the curve, relative and absolute enhancement were not able to differentiate enchondromas from low-grade CS. This finding is of significance, as previous studies stated that DCE-MRI is an important tool in the differentiation of the two disease entities. This discrepancy in findings can be explained by the fact that the above mentioned studies included both low-grade and high-grade CS and did not differentiate between low-grade and high-grade CS. Therefore the findings in those studies cannot be utilised to differentiate enchondromas from low-grade CS, but only to differentiate enchondromas from CS in general [9, 19]. This is also supported in our study, which demonstrated a statistically significant difference in DCE-MRI parameters in the differentiation of low-grade chondroid tumours and HGS-CS. However, for the radiologist, the challenge does not lie in the differentiation of enchondromas from chondrosarcomas per se but in the distinction of enchondromas from low-grade CS. Furthermore, previous studies demonstrated that conventional MRI could differentiate low-grade CS from high-grade CS [21, 22]. Hence, the above mentioned study is of limited utility for the practising radiologist.

In view of the great difficulties in differentiating enchondromas from low-grade CS, there has been increased interest in utilising other functional MRI-techniques to aid in the distinction of the two disease entities. Douis and co-workers, e.g., have previously investigated the utility of DWI in the differentiation of enchondromas from chondrosarcomas. The authors found that DWI could not differentiate enchondromas only.

Fig. 5  A 39-year-old female patient with an eccentric enchondroma of the right proximal humerus. a, b Coronal T1W SE MR image shows an eccentric enchondroma measuring 2.8 cm in the maximum cranio-caudal dimension that shows grade 3 endosteal scalloping (arrow) and bone expansion due to its location. A subsequent curettage demonstrated an enchondroma only.
from CS and that DWI could not aid in the distinction of low-grade chondroid tumours from high-grade CS. Hence, the distinction of enchondromas from low-grade CS remains challenging and reliant on the close collaboration of radiologists, pathologists and clinicians.

There are a few limitations to our study. Firstly, we specifically differentiated between enchondromas, CLUMPs, low-grade CS and high-grade CS. This is of relevance, as in many articles on this topic, this distinction, particularly with regards to low-grade cartilaginous tumours which are difficult to categorise into a benign or low-grade malignant group, is frequently not made. However, the categorisation into the above mentioned groups in our publication reflects the diagnostic challenge and uncertainty that radiologists, clinicians and pathologists frequently face in clinical practise. The nature of these CLUMPs remains uncertain. Hence, these lesions are termed CLUMPs (cartilaginous lesions of unknown malignant potential) at our institution and our protocol is currently to follow-up these cases with yearly radiographs and clinical assessment.

Secondly, in our study, the diagnosis of 17 of 27 enchondromas was not histologically confirmed and no follow-up in these patients was performed. However, we applied stringent diagnostic criteria for the MR-diagnosis of enchondromas in the absence of a histological confirmation, as described by Murphey and coworkers [8].

A further limitation is that the MRI-findings were reviewed by a single reader (ID). However, the reader of the MRI-studies is a fellowship-trained musculoskeletal radiologist with extensive experience in the interpretation of bone sarcomas.

Furthermore, there is likely a selection bias in our study because only a small proportion of patients who were referred to our institution with chondroid tumours underwent a conventional MRI and DCE-MRI in-house. Many of these cases, however, were selected for DCE-MRI in an attempt to reduce the diagnostic uncertainty, because the diagnosis was not clear on previously obtained conventional imaging.

Finally, in view of the retrospective nature of the study, we had to rely on the electronic patient records with regards to the clinical assessment. Whilst some may argue that such an assessment is subjective, our findings reveal that clinical evaluation demonstrated a high specificity and moderate sensitivity in the differentiation of the two disease entities. This is largely due to the fact that the pain in chondrosarcoma is characterised by a constant resting pain, and may be associated with night pain as observed in our study. In contrast, pain not attributed to the lesion was frequently mechanical in nature and could usually be attributed to degenerative processes (such as meniscal tears or osteoarthritis).

Conclusion

Grade 1 CS remains a difficult imaging diagnosis. Our findings show that DCE-MRI could not differentiate enchondromas from grade 1 CS. In agreement with previous reports, we found that tumour length, depth of endosteal scalloping, cortical destruction, bone expansion, soft tissue mass and the presence of pain attributed to the lesion suggest CS as opposed to enchondroma. However, except for the presence of deep endosteal scalloping and pain attributed to the lesion, the sensitivity of these signs was low in grade 1 CS.

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Compliance with ethical standards

Guarantor The scientific guarantor of this publication is Mark A. Davies.

Conflict of interest The authors of this manuscript declare no relationships with any companies whose products or services may be related to the subject matter of the article.

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Statistics and biometry Peter Nightingale kindly provided statistical advice for this manuscript.

Ethical approval Institutional Review Board approval was obtained.

Informed consent Written informed consent was not required for this study because of the retrospective nature of the study.

Methodology

• retrospective
• observational
• performed at one institution

References


Appendix 2: List of publications by candidate

Published articles


13. Puls F, Arbajian E, Magnusson L, **Douis H**, Kindblom LG, Mertens F


14. Puls F, Niblett A, Marland G, Gaston CL, **Douis H**, Mangham DC, Sumathi VP,


15. Chechlacz M, Terry A, Demeyere N, **Douis H**, Bickerton WL, Rotshtein P,


Book Chapters


