Consensus recommendations on the classification, definition and diagnostic criteria of hip-related pain in young and middle-aged active adults from the International Hip-related Pain Research Network, Zurich 2018

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ABSTRACT

There is no agreement on how to classify, define or diagnose hip-related pain—a common cause of hip and groin pain in young and middle-aged active adults. This complicates the work of clinicians and researchers. The International Hip-related Pain Research Network consensus group met in November 2018 in Zurich aiming to make recommendations on how to classify, define and diagnose hip disease in young and middle-aged active adults with hip-related pain as the main symptom. Prior to the meeting we performed a scoping review of electronic databases in June 2018 to determine the definition, epidemiology and diagnosis of hip conditions in young and middle-aged active adults presenting with hip-related pain. We developed and presented evidence-based statements for these to a panel of 37 experts for discussion and consensus agreement. Both non-musculoskeletal and serious hip pathological conditions (eg, tumours, infections, stress fractures, slipped capital femoral epiphysis), as well as competing musculoskeletal conditions (eg, lumbar spine) should be excluded when diagnosing hip-related pain in young and middle-aged active adults. The most common hip conditions in young and middle-aged active adults presenting with hip-related pain are: (1) femoroacetabular impingement (FAI) syndrome, (2) acetabular dysplasia and/or hip instability and (3) other conditions without a distinct osseous morphology including labral, chondral and/or ligamentum teres pathology. The last category should be limited. Anteroposterior pelvis and lateral femoral head–neck radiographs are the initial diagnostic imaging of choice—advanced imaging should be performed only when requiring additional detail of bony or soft-tissue morphology (eg, for definitive diagnosis, research setting or when planning surgery). We recommend clear, detailed and consistent methodology of bony morphology outcome measures (definition, measurement and statistical reporting) in research. Future research on conditions with hip-related pain as the main symptom should include high-quality prospective studies on aetiology and prognosis. The most common hip conditions in active adults presenting with hip-related pain are: (1) FAI syndrome, (2) acetabular dysplasia and/or hip instability and (3) other conditions without distinct osseous morphology including labral, chondral and/or ligamentum teres conditions. The last category should not be confused with the incidental imaging findings of labral, chondral and/or ligamentum teres pathology in asymptomatic people. Future research should refine our current recommendations by determining the clinical utility of clinical examination and diagnostic imaging in prospective studies.

INTRODUCTION

Young and middle-aged active adults with hip and groin pain often present with a confusing overlap of signs and symptoms.1 There is no consensus on how to define or classify hip disease with different and overlapping intra-articular and extra-articular contributors to symptoms, in an anatomically complex region. Several researchers and consensus
groups have previously attempted to define aspects of hip and groin pain. Among these are consensus agreements on the terminology and definitions for groin pain in athletes and on femoroacetabular impingement (FAI) syndrome.

The Doha Agreement Meeting on terminology and definitions in groin pain in athletes divided groin pain into three major categories: clinical entities for groin pain (adductor-related, iliopsoas-related, inguinal-related and pubic-related), hip-related and other causes of groin pain in athletes. The Doha Agreement concluded that “hip-related pain can be difficult to distinguish from other causes and that it may coexist with other types of groin pain.” A detailed classification of possible causes of hip-related pain in athletes was considered to be outside the scope of the Doha Agreement process. This difficulty was related to the inability for hip examinations to assist in differential diagnosis, and probably only being useful for screening—as most clinical tests have good sensitivity but poor specificity.

There is no consensus on how to classify hip-related pain in young and middle-aged active adults, with prior studies focusing on classifying individual conditions, such as FAI syndrome, acetabular dysplasia, labral tear, chondral lesions and ligamentum teres tears. Each of these conditions is described in detail in online supplementary appendix 1. Briefly, in FAI syndrome, the repetitive mechanical loading of cartilage between the femoral head and acetabular rim is suggested to contribute to acetabular labral and chondrolabral findings, as well as early osteoarthritis. As such, FAI syndrome was defined as a motion-related clinical disorder of the hip with a triad of symptoms, clinical signs and imaging findings (typically cam, pincer or mixed morphology) that represents symptomatic premature contact between the proximal femur and the acetabulum. The primary symptom of FAI syndrome has traditionally been described as groin pain, although pain may also be felt in the back, buttock or thigh. Not having groin pain has been suggested to help exclude FAI syndrome and/or labral tear findings in patients with hip pain. Acetabular dysplasia refers to misalignment between the femoral head and the acetabulum secondary to changes in their shape, size and orientation, which may result in instability and overload of the acetabular rim during normal activities.

Differential diagnosis of any condition requires exclusion of more serious musculoskeletal and non-musculoskeletal pathology, as well screening appropriate body regions. For hip-related pain in young active adults, this could for example include Perthes’ disease, slipped capital femoral epiphysis (SCFE), infection, osteoarthritis, tumours and fractures. While it is beyond the scope and purpose of detailing the process of helping to exclude these pathological concerns, the reader is referred to other recent works detailing this process.

A proper classification of non-arthritic hip disease in active adults into well-defined hip conditions will provide more clarity for clinicians and researchers. This is critically important due to the significant increase in the diagnosis of hip conditions and incidental imaging findings in athletes, as well as arthroscopic hip surgery in general. There is a lack of current best evidence on the classification of hip-related pain, partly due to the heterogeneity in the definition of hip-related pain. Consensus, informed by current best evidence, is needed to provide clarity for clinicians interpreting the findings of research papers, particularly with the increasing knowledge of incidental imaging findings in young and middle-aged active adults.

The International Hip-related Pain Research Network (IHiPRN) was established in 2017 to facilitate collaboration across research groups and disciplines and to improve knowledge dissemination of hip-related pain to clinicians. The specific focus of the IHiPRN is ‘non-red flag pain originating from the hip joint in young and middle-aged active adults’. We identified four topics on hip-related pain for consensus agreement: classification of hip pain, patient-reported outcomes, standardised measurement of physical capacity and physiotherapist-led treatment.

In this paper, we first describe the general consensus process applied to all topics in this series. We then make recommendations on the first topic on how to classify, define and diagnose hip disease in young and middle-aged active adults, with hip-related pain as the main symptom. Other papers in this series will use this classification of hip-related pain in the assessment of (1) patient-reported outcome measures, (2) standardised measurement of physical capacity and (3) physiotherapist-led treatment.

METHODS

Literature review

For the first key area on the classification, definition and diagnostic criteria of hip-related pain in young and middle-aged active adults, working group 1 performed a literature review to identify gaps in the literature to formulate consensus recommendations and provide research recommendations for future study. A summary of the literature review is contained in online supplementary appendix 1. Using components of a scoping review framework, our inclusion and exclusion criteria were the following:

- Study design: Systematic reviews, intervention and observational studies (prospective or retrospective) with a study population of at least 10 young and middle-aged active adults and published in English language, peer-reviewed journals. We excluded conference abstracts, case studies, narrative reviews and non-peer-reviewed studies.
- Population: Research participants diagnosed with hip-related pain or related condition. Participants with primary osteoarthritis, SCFE, Legg-Calve-Perthes or extra-articular condition were excluded as these are not the conditions identified as being of primary prevalence in participants of this age range and pain description.
- Outcome: Definition, epidemiology, clinical and/or imaging diagnosis of any hip-related pain condition.

Consensus meeting

The consensus process used is shown in figure 1.

Selection of expert group members

The IHiPRN leadership group (JLK, KMC, MB, ABM, CLL and KMM) met in January 2017 to identify potential expert group members. Experts were selected based on their track record of publications, and being current active researchers in the field of hip-related pain in young and middle-aged active adults. Many researchers were also clinicians in the field. Potential expert group members were invited to the first IHiPRN Consensus Meeting in Zurich in November 2018 (online supplementary appendix 2), and asked to identify additional experts for invitation.

Following this expression of interest, four key areas were identified as priorities for consensus. These four key areas were the following:

1. Classification of hip pain (including use of clinical tests and imaging).
2. Patient-reported outcome measures for hip pain (including hip-related measures and maybe others including pain/coping/fear/utility measures).
Need for international consensus established January 2017

IHIPRN leadership group (JLK, KMC, MB, ABM, CLL, KMK) selected Expert Group members

Four key priority areas identified and four working groups established

- Leadership group identified experts to lead each of four working groups
- Working group members determined by discussion between leadership group and working group leaders

Pre-agreement meeting process

- Four working group prepared evidence summary and developed draft consensus recommendations
- Evidence summary and draft consensus recommendations presented to entire expert group prior to Zurich meeting

Agreement meeting process (November 2018)

- Working groups revised evidence summaries and draft consensus recommendations
- Working group leaders presented evidence summaries to all expert group members for discussion/further revision.
- 38 expert group members rated each recommendation on 10-point Likert scale
- Scores pooled and median for each recommendation determined

Recommendations from four working groups form international consensus

3. Standardised measurement of physical capacity in hip-related pain (including clinical measures, biomechanics, electromyography, physical activity, functional performance and return to sport).


The leadership group then identified experts to lead each of the four working groups. These were MPR and RA (group 1), ABM and CLL (group 2), FI and JLK (group 3), JLK and MB (group 4). Members of the working groups were then determined following discussion between the leadership group and the working group leaders. The working group members for this paper were MPR, RA, JLK, HPD, JJH, AW, PVK, AK, ABM, EA, PH and KMW.

Expert group demographics

Areas of expertise among the expert group participants included physiotherapy, orthopaedic surgery, sport and exercise medicine, biomechanics, diagnostics, imaging and radiology, patient-reported outcome measures and exercise science. In addition, many participants were also expert clinicians who regularly treat young and middle-aged active adults with hip-related pain.

Premeeting process

Prior to the meeting, the four working groups prepared evidence summaries (based on literature searches and syntheses) to help inform recommendations and underpin discussions at the consensus meeting. These evidence summaries and draft consensus recommendations were presented to all meeting participants via email at least 1 week prior to the Zurich meeting.

Consensus meeting process

At the beginning of the consensus meeting in Zurich, the working groups discussed and revised the draft consensus recommendations. Evidence summaries and consensus recommendations
were then presented to, and discussed by the whole expert group, and further revisions were made. Following discussion, each participant was asked to vote on the recommendation on a 10-point Likert scale, where 0 was considered to be ‘inappropriate’ and 9 ‘appropriate’, as described previously.31 32 Participants were blinded to each other’s scoring. Scores were pooled and the median (IQR) for each recommendation was determined. Recommendations that scored 0–3 were considered inappropriate, scored 4–6 were considered uncertain and scored 7–9 were considered appropriate.31 32 The level of available evidence was charted, and summaries of this for each condition are contained in figures 2–9 in a 2×2 quadrant (magnitude × precision) accordingly:

- Red quadrant: values for both magnitude and precision are low, not typically recommended for clinical decision-making. These measures are Not Recommended.
- Yellow quadrant: values for either magnitude or precision are low. Caution is recommended for clinical decision-making. These measures are Cautiously Recommended.

**Determining diagnostic utility for various clinical and radiological measures**

Diagnostic values (eg, sensitivity, specificity, likelihood ratios) for the various clinical and radiological measures are reported in the appendix. Since determination of clinical utility of these values is often poorly done or misunderstood, we have attempted to provide the reader with a graphical representation of how to potentially use and incorporate these values in clinical practice (eg, what clinical value should they be given). To do this requires examination of not only diagnostic accuracy but also magnitude of diagnostic probability shifts,34 precision and the quality of the study examining such measures.

Values used in this paper were stratified accordingly:

- **Magnitude**: size of the shift in probability as previously defined.34
- **Precision**: how repeatable are the results. This is defined as highest number divided by lowest number for the CI of the post-test probability value for each measure (eg, 95% CI for a post-test probability value of 49 to 93 has a precision value of 93/49=1.9). A lower value indicates higher precision. This was described previously as the upper-to-lower confidence limit ratio.35 A probability shift with a wide CI is imprecise and unstable.35

**Study Quality**: high or low study quality based on previous systematic reviews that have quantified study quality.36 37 Charted as:

- **High-quality study**:●
- **Low-quality study**:●

Diagnostic utility was charted, and summaries of this for each condition are contained in figures 2–9 in a 2×2 quadrant (magnitude × precision) accordingly:

- **Red quadrant**: values for both magnitude and precision are low, not typically recommended for clinical decision-making. These measures are Not Recommended.
- **Yellow quadrant**: values for either magnitude or precision are low. Caution is recommended for clinical decision-making. These measures are Cautiously Recommended.

**RESULTS**

**Results of the consensus meeting: consensus meeting participants**

The consensus meeting in Zurich, Switzerland (17–18 November 2018) was attended in person by 37 out of the 44 HHPRN participants; one additional participant attended via videoconferencing.

- **Green quadrant**: high magnitude and high precision. These measures are Recommended if study quality is high. Cautious recommendation for low-quality studies.

  We have drawn these figures in quadrants to help the reader, but just as with the quadrants in a cost-effectiveness curve, we appreciate the axis are continuous and there are no discrete cutoffs from ‘low to high’. 

![Figure 2](image.png)

**Figure 2** Magnitude and precision of clinical tests for diagnosis and screening of FAI syndrome and labral pathology. (+), positive test; (-), negative test; FADIR, flexion adduction internal rotation; FAI, femoroacetabular impingement; IR, internal rotation; MRA, magnetic resonance arthrogram. Reference standards are listed in ()

![Figure 3](image.png)

**Figure 3** Magnitude and precision of imaging tests for diagnosis and screening of FAI syndrome. (+), positive test; (-), negative test; FAI, femoroacetabular impingement; MRA, magnetic resonance arthrogram; T, Tesla.
Consensus statement

**Figure 4** Magnitude and precision of clinical tests for diagnosis and screening of acetabular dysplasia and hip instability. (+), positive test; (-), negative test; AB-HEER, abduction–hyperextension–external rotation; FABER, flexion abduction external rotation; FPAW, foot progression angle walking; HEER, hyperextension–external rotation; PIT, prone instability test; (+) positive test, (-) negative test.

**Figure 5** Magnitude and precision of imaging tests for diagnosis and screening of labral conditions. (+), positive test; (-), negative test; CTA, CT arthrogram; MRA, magnetic resonance arthrogram; T, Tesla; US, ultrasound. Reference standards are listed in (1).

**Figure 6** Magnitude and precision of imaging tests for diagnosis and screening of chondral conditions. (+), positive test; (-), negative test; MRA, magnetic resonance arthrogram.

Expert group demographics
The expert group of 27 physiotherapists, 7 orthopaedic surgeons, 4 sport and exercise medicine physicians, 4 exercise scientists, 1 radiologist and 1 general medical doctor included 12 women and 32 men. Participants had a mean (SD) age of 41 (11) years and 20 (10) years of experience. Twenty-one participants were from Europe, 13 from Australia/New Zealand, 8 from North America and 2 from the Middle East. All participants were fluent English speakers.

Literature search and evidence synthesis
Considerations
While synthesising available evidence to inform the consensus process, a number of important aspects require consideration: (1) Many of the included studies were performed in high volume hip surgeon practice settings. The patients in these studies have a high a priori probability of having a hip condition, leading to a limited ability of an examination method or imaging modality to substantially alter the post-test probability. Future studies in general practice or physiotherapy settings might improve our understanding of the value of these tests. (2) Some conditions are defined by certain radiological appearances or cut-off values (eg, cam morphology where alpha angle is >60, pincer morphology where lateral centre edge angle (LCEA) is >40, acetabular dysplasia where LCEA is <20), but there is no consensus on specific imaging parameters to diagnose different hip conditions. As such, it is inappropriate to examine the value of imaging to assess the presence of a condition. (3) The third condition ‘other’ includes isolated or combined labral, chondral and ligamentum teres conditions as the most likely cause of hip-related nociception in the absence of distinct osseous morphology. This category should not be confused with the incidental imaging findings in asymptomatic people. Labral, chondral and/or ligamentum teres findings are common in asymptomatic individuals and can coexist in patients with FAI syndrome or acetabular dysplasia and/or hip instability.
The IHiPRN consensus group met in Zurich in November 2018, and considered how to classify, define and diagnose hip disease in young and middle-aged active adults with hip-related pain as the main symptom. The working group conducted a scoping review, a useful framework to collate and summarise information on a broad topic, presented three clinical, one combined clinical and research and two research recommendations for consensus discussion and agreement.

This consensus paper builds on previous hip and groin pain consensus papers. It classifies hip disease in young and middle-aged active adults (with hip-related pain as the main symptom) into three hip conditions: (1) FAI syndrome, (2) acetabular dysplasia and/or hip instability and (3) other conditions without distinct osseous morphology. This final category includes patients with isolated or combined labral, chondral and ligamentum teres findings (including a range of distinct and overlapping pathologies including tears, cysts, erosions or hyper trophy). It may also include patients with no imaging findings. We also report the clinical utility of diagnostic findings for each of these conditions. It is important to acknowledge that these conditions often coexist. We also acknowledge (but did not include here) other hip conditions with hip-related pain as the main symptom including osteoarthritis, infections, transient osteoporosis, stress fractures and tumours. We also acknowledge that currently diagnostic accuracy is limited to lower quality cohort studies with high pretest probability of disease. To what extent diagnostic accuracy is altered in studies of lower pretest probability is not understood, although one high-quality study with a lower pretest probability suggests the probability shift is similarly low. Other studies in settings of low pretest probability are necessary to determine the diagnostic accuracy of these various assessments in primary care setting.

Our evidence-informed classification of hip disease in young and middle-aged active adults classifies hip-related pain into three condition categories (once non-musculoskeletal and musculoskeletal serious hip pathology and competing body regions are excluded): FAI syndrome, acetabular dysplasia and/or hip instability, and other conditions, such as labral, chondral and/or ligamentum teres conditions. It is important, however, to appreciate that these conditions often coexist in patients with hip-related pain. These recommendations are not intended to be absolute for all young and middle-aged active adults with hip-related pain—a multifactorial condition. The clinical approach to hip disease in active adults with hip-related pain is highly contextual and should be integrated with clinical expertise using evidence-based practice and clinical decision-making models. Each of the consensus recommendations are discussed below and represented in the clinical case example (Box).

Clinical recommendation 1 (C1). A negative flexion adduction internal rotation (FADIR) test helps to rule out hip disease in young and middle-aged active adults presenting with hip-related pain.

The diagnostic accuracy of the clinical assessment is mostly limited to clinical special tests and we cautiously recommended the FADIR test as the only clinical test for utilisation in attempt to help rule out potential hip-related pain. While the FADIR test is represented in our ‘cautiously recommend’ quadrant and the Thomas test is in the ‘recommend’ quadrant, the specifics of consensus recommendations and consensus voting results can be found in table 1 and figures 10–12. The consensus group considered all recommendations to be appropriate.

**Discussions**

**Findings**

Both non-musculoskeletal and serious hip pathological conditions (eg, tumours, infections, stress fractures, SCFE), as well as competing musculoskeletal conditions (eg, lumbar spine) should be excluded when diagnosing hip-related pain in young and middle-aged active adults. Five hip conditions in young and middle-aged active adults presenting with hip-related pain met the inclusion criteria of this review: FAI syndrome; acetabular dysplasia and/or hip instability; labral, chondral and ligamentum teres conditions. We classified the last three conditions as ‘other conditions’. It is important to note that labral, chondral and ligamentum teres pathology are common on imaging in asymptomatic people (and best-labelled ‘incidental findings’ in this context).

**Results of the consensus on topic 1: classification of hip-related pain**

The IHiPRN delegates at the November 2018 Zurich consensus meeting discussed and voted on six recommendations: three recommendations for clinicians, one recommendation for clinicians and researchers and two for researchers. The final
the studies examining these clinical tests is worthy of consideration. The Thomas test was examined in one high-quality study not specifically designed to determine diagnostic accuracy. The FADIR test has been examined in several studies; they are low-quality studies, most not designed to determine diagnostic accuracy. We therefore feel it is important to provide the clinician with the most appropriate clinical measure to potentially screen for all hip-related pain, based on the best current evidence, and recommend the FADIR test despite its noted limitations. Other clinical special tests (eg, Thomas test, prone instability test, ligamentum teres tear test) were considered by the expert group, and these tests have a:

- Very limited ability to confirm FAI syndrome by increasing post-test probability.
- Limited to substantial ability to help confirm acetabular dysplasia and/or hip instability in low-quality studies.
- Substantial ability to help rule in labral conditions in only one high-quality study.
- Unknown ability to rule in chondral conditions.
- Moderate post-test probability for confirming diagnosis of ligamentum teres conditions.

As the evidence for the clinical utility of these other tests is very limited or insufficient for ruling non-arthritis hip disease in or out in young and middle-aged active adults, we do not recommend their use by clinicians to diagnose the source of hip-related pain. In addition, we do not know the diagnostic accuracy of symptoms for conditions characterised by hip-related pain as mentioned earlier. The value of range-of-motion (ROM), muscle performance and other clinical tests for the diagnosis of a condition characterised by hip-related pain is unknown, as such tests currently lack diagnostic accuracy.

Clinical recommendation 2 (C2). Diagnostic utility of imaging for hip disease in young and middle-aged active adults presenting with hip-related pain is limited; we recommend that imaging should never be used in isolation but combined with the patient’s symptoms and clinical signs (when making treatment and other relevant decisions).

In young and middle-aged active adults with hip disease, this recommendation on diagnostic utility of imaging should underpin any agreement on diagnostic criteria. Diagnostic imaging demonstrates post-test probability shifts ranging from very small to substantial for the diagnosis of FAI syndrome, and acetabular dysplasia and/or hip instability, and small to moderate shifts at best for labral, chondral and ligamentum teres pathology. Diagnostic imaging has a very limited ability to confirm the diagnosis of a particular condition as the cause of pain. As such, we agreed with previous recommendations that the diagnostic criteria should include appropriate diagnostic imaging, but also consideration of symptoms and clinical signs.\(^1\)\(^2\)\(^3\) Diagnosis of the primary cause of hip-related pain should never be made using imaging alone. The next step in the research process is to evaluate the performance of these diagnostic criteria, which primarily includes clinical signs and diagnostic imaging for the various hip conditions with hip-related pain as the primary symptom.

Clinical recommendation 3 (C3). We recommend anteroposterior (AP) pelvis and lateral femoral head–neck radiographs to assist in diagnosing the hip conditions characterised by hip-related pain, and MRI/magnetic resonance arthrogram (MRA) or CT scan where three-dimensional morphological assessment is indicated.

Our recommendation is consistent with prior recommendations\(^3\) that the initial diagnostic imaging should at least include AP pelvic and lateral femoral head–neck view radiographs. The latter can, for example, include a Dunn view, a frog-leg (also known as Lauenstein) view or a cross-table view. This will provide the clinician with an understanding of the underlying morphology of the hip, which may be relevant for the patient presenting with hip-related pain. When indicated (clinically or in research), advanced imaging can be used for more clarity on
three-dimensional morphology or to assess intra-articular structures. For intra-articular structures such as the labrum, cartilage and ligamentum teres, we recommend MRI/MRA or CT arthrogram scan. Incidental intra-articular findings are common in asymptomatic individuals and should be managed with the appropriate clinical sensitivity. Again, caution is suggested as the clinical utility of diagnostic imaging can have only small shifts in post-test probability as a result of either positive or negative findings. As such, a comprehensive clinical examination, including screening for other conditions that might present with hip-related pain (eg, of the spine and pelvis), is required, and the diagnosis of the cause of hip-related pain should never be made on imaging alone.

Clinical and research recommendation 1 (CR1). Hip-related pain may be further categorised after imaging into: (1) FAI syndrome; (2) acetabular dysplasia and/or hip instability and (3) other conditions causing hip-related pain. This last category could include soft-tissue conditions without specific bony morphology.

For both clinical and research purposes, imaging can help to categorise hip disease in young and middle-aged active adults presenting with hip-related pain into the following conditions: (1) FAI syndrome, (2) acetabular dysplasia and/or hip instability and (3) other conditions causing hip-related pain without specific bony morphology, such as labral, chondral and/or ligamentum teres conditions. Many hip-related pain conditions—especially labral and chondral conditions—coexist with FAI syndrome and acetabular dysplasia and/or hip instability. The third category (similar to previous recommendations) is less distinct with variable clinical presentations and includes intra-articular conditions without a distinct bony morphology. We acknowledge that special tests and imaging have high rates of false-positive findings and that patients can present with symptoms and signs similar to that of FAI syndrome, but have normal bony morphology. The third category of ‘other conditions’ will include these patients. Finally, we acknowledge that patients may present with pain, but normal morphology or no findings consistent with other pathology on imaging can explain nociception. Standardised and well-reported methods in research, such as those suggested in this recommendation, will reduce research heterogeneity, while improving the accuracy of summary estimates in meta-analyses.

Research recommendation 1. Measures of bony morphology should be reported in detail. We recommend that bony morphology outcome measures (such as the alpha angle or centre-edge angle) should be clearly defined, measured and reported (eg, detailed methodological description; blinding; per hip/per person reporting with statistical correction as appropriate; reliability measures).

We recommend that researchers provide the following as part of the methodology section of research reports when using bony morphology imaging outcome measures: (1) define the outcome measure; (2) report a reliable and complete description of imaging methodology and details on how the outcome is measured and (3) provide detail of the statistical analysis used.

Research recommendation 2. Future research recommendations: We recommend large-scale, interdisciplinary research on aetiology and prognosis for each of the listed three hip conditions (characterised by hip-related pain).

Studies that examine the aetiology and prognosis of the three hip conditions are critical if we are to effectively classify, define and diagnose hip-related pain in young and middle-aged active adults. To classify and define each condition, we must identify the aetiology of the condition. Understanding the aetiology of the conditions will eventually enable effective primary, secondary and tertiary prevention strategies to be developed. We must conduct studies that examine the prognosis of young and middle-aged people with hip-related pain. We can determine prognosis by conducting research that explores the relationship of each of the three conditions with other factors, such as pain, other symptoms, psychological factors, socioeconomic factors and physical impairments. We will then be able to begin to develop accurate assessment processes and effective, targeted treatments.

A hypothetical clinical case example (not requiring patient consent statement) describing the implementation of recommendations C1, C2, C3 and CR1 into clinical practice.

Patient described is a 25-year-old male football player with a 6-month history of anterior hip and groin pain, aggravated by kicking and changing direction activities.

C1: A negative flexion-adduction-internal rotation (FADIR) test may help to rule out hip in young and middle-aged active adults presenting with hip-related pain. The FADIR test should be performed as part of a comprehensive physical examination. If the test is negative, the clinician could consider that the source of pain is possibly not the hip joint. If the test is positive, the hip may be a source of nociception, and further assessment is required.

The patient has a positive FADIR test.

C2: Diagnostic utility of imaging for hip disease in young and middle-aged active adults presenting with hip-related pain is limited; it should be interpreted together with the patient’s symptoms and clinical signs. His anteroposterior (AP) pelvis and lateral radiographs have been obtained. The report states the presence of cam morphology. While these findings may be contributing to his pain and match the clinical presentation, the clinician should be aware of the prevalence of cam morphology (66%) in athletes.

C3: We recommend AP pelvis and lateral femoral head-neck radiographs to assist in diagnosing the hip conditions characterised by hip-related pain, and MRI or CT scan where three-dimensional morphological assessment is indicated. He has a positive FADIR test and has cam morphology on radiographs. In consultation with his medical team, he has decided to pursue surgical options to address his continued hip pain. Referral for MRI or CT scan may assist in identifying labral, chondral or other hip conditions when clinically indicated; especially if non-surgical treatment such as physiotherapy has not succeeded and surgery is being considered, MRI may be necessary to better plan surgery.

CR1: Hip-related pain may be further categorised after imaging into: (1) femoroacetabular impingement (FAI) syndrome, (2) acetabular dysplasia and/or hip instability and (3) other conditions causing hip-related pain. This last category could include soft-tissue conditions without specific bony morphology. The comprehensive subjective assessment of the patient’s complaints/pain, physical examination and imaging undertaken suggest that he may have FAI syndrome (positive FADIR test, cam morphology and labral lesions on MRI scan). Categorising his condition as FAI syndrome may assist him in understanding his condition and may aid the shared decision-making process regarding appropriate management decisions.
A. Agreeing on the classification of hip disease presenting with hip-related pain (ie, the different conditions).
B. Agreeing on the diagnostic criteria for each hip condition.

2. Prognosis
A. Determining the performance of these criteria (eg, diagnostic accuracy of clinical and radiological examination).
B. Investigating the impact of the diagnostic components of a specific hip condition on diagnostic or prognostic thinking (eg, stratification of patients into high and low risk; determining the prognostic value of the different conditions on long-term outcomes, for example, osteoarthritis).
C. Determining the impact on diagnostic or therapeutic strategies (eg, determining if advanced imaging is required for diagnosis, agreeing on an appropriate treatment strategy).

3. Financial, health-related, and patient and provider satisfaction outcomes (eg, cost-effectiveness studies, implementation of satisfaction outcome scores in the clinic and report in treatment outcome studies).

Research investigating the hip conditions in young and middle-aged active adults with hip-related pain has not completed this logical process. Additionally, while we point out that progress has been made with steps 1 and 3, the remaining steps are largely ignored with current literature focusing on outcomes for treatment strategies. Thus, we recommend large-scale, prospective and interdisciplinary research on aetiology and prognosis for each of the three primary condition categories of hip disease with hip-related pain as the main symptom and their associated bony morphologies and soft-tissue pathology.

Limitations
Several limitations require consideration regarding the clinical and research recommendations discussed in this paper. The group of participants who voted on these recommendations at the first IHiPRN meeting in Zurich included many different professions, and diversity in research and clinical experience. However, due to the invitation process and budgetary and logistical limitations, the group cannot be considered fully inclusive. Therefore, the contributions were limited to the participants who were part of the consensus process. We acknowledge the possibility of bias, and homogeneity of expert opinion, and that future consensus meetings will aim to include a broader range of experts, as well as consumers (patient representatives).

The extremely high level of agreement demonstrated in figures 10–12 may represent a ceiling effect or bias. In previous work using similar RAND Appropriateness Method scales, that bias can be reduced by having an ‘effective group leadership’. The design of IHiPRN, working groups, collaborations premeeting, during and postmeeting all were focused on limiting any potential bias.
This consensus is based on a scoping review of only English language studies with no quality assessment of included studies. It is possible that we missed important studies in other languages and included low-quality studies or reviews. Interpretation of study quality was inferred from the included systematic reviews. Consensus statement voting did not cover all aspects of this paper. For example, studies without quality assessment (eg, studies on condition definition and epidemiology) were not addressed in consensus statements.
We acknowledge that other hip conditions not addressed in this paper can also lead to hip-related pain in young active adults. The conditions described are most prevalent in young and middle-aged active adults with hip-related pain.
Finally, there are a number of concerns with diagnostic accuracy studies, including: (1) typically performed in settings of high pretest probability (eg, surgical populations), (2) the majority of these studies are low-quality cohort studies with a high risk of bias in many quality domains—diagnostic accuracy might therefore be overestimated and (3) several patients with hip-related pain might have more than one hip condition (on imaging or in surgery). It is worth mentioning that caution is warranted in interpretation of these consensus statements knowing that the information available is based on biased high-prevalence, lower-level cohort evidence.

CONCLUSION
The first IHiPRN Consensus Meeting was held in Zurich, Switzerland, in November 2018. The IHiPRN participants developed and gained consensus on three clinical recommendations, one
clinical and research recommendation and two research recommenda-
tions, all of which were an amalgamation of best available
evidence and expert opinion. Both non-musculoskeletal and serious hip pathological conditions (eg, tumours, infections, SCFE), as well as competing musculoskeletal conditions (eg, lumbar spine) should first be excluded before categorising hip disease in young and middle-aged active adults presenting with hip-related pain. Once these are excluded, hip-related pain should be categorised into: (1) FAI syndrome, (2) acetabular dysplasia and/or hip instability and (3) other conditions without labral clinical signs and diagnostic imaging is unknown for patients recognise that the diagnostic capability of matching symptoms, should be further assessment of intra-articular structures or morphology is indicated. Clearly this is an area requiring further research to provide both the clinician and researcher greater clarity regarding imaging utility.

Research studies reporting bony morphology should provide clear definitions and detailed imaging and reporting methods. We recommend large-scale, prospective and interdisciplinary research on aetiology and prognosis of hip-related pain where the over-lapping nature of the various conditions is considered.

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REFERENCES
26 Poole C. Low P-values or narrow confidence intervals: which are more dubious? *Epidemiology* 2001;12:291–4.