European League Against Rheumatism recommendations for calcium pyrophosphate deposition. Part I: terminology and diagnosis


ABSTRACT

Objectives To agree terminology and to develop recommendations for the diagnosis of calcium pyrophosphate deposition (CPPD).

Methods The European League Against Rheumatism (EULAR) CPPD Task Force, comprising 15 experts from 10 countries, agreed the terms and recommendations for diagnosis of CPPD using a Delphi consensus approach. Evidence was systematically reviewed and presented in terms of sensitivity, specificity and positive likelihood ratio (LR) to support diagnosis; ORs were used for association. Strength of recommendation (SOR) was assessed by the EULAR visual analogue scale.

Results It was agreed that 'CPPD' should be the umbrella term that includes acute calcium pyrophosphate (CPP) crystal arthritis, osteoarthritis (OA) with CPPD and chronic CPP crystal inflammatory arthritis. Chondrocalcinosis (CC) defines cartilage calcification, most commonly due to CPPD and detected by imaging or historological examination. A total of 11 key recommendations were generated on the topics of clinical features, synovial fluid (SF) examination, imaging, comorbidities and risk factors. Definitive diagnosis of CPPD relies on identification of SF CPP crystals. Rapid onset inflammatory symptoms and signs are suggestive but not definitive for acute CPP crystal arthritis. Radiographic CC is not highly sensitive or specific, whereas ultrasonography appears more useful (LR=24.2, 95% CI 3.51 to 168.01) for peripheral joints. Recognised risk factors for CPPD include ageing, OA and metabolic conditions such as primary hyperparathyroidism, haemochromatosis and hypomagnesaemia; familial forms are rare. SORs varied from 53 to 99 (maximum 100).

Conclusion New terms for CPPD were agreed and 11 key recommendations for diagnosis of CPPD were developed using research evidence and expert consensus.

INTRODUCTION

Calcium pyrophosphate deposition (CPPD) occurs almost exclusively in articular tissues, most commonly fibrocartilage and hyaline cartilage, and is the most common cause of chondrocalcinosis (CC). Calcium pyrophosphate (CPP) associated arthritis is the third most common inflammatory arthritis. Recognised risk factors are ageing, osteoarthritis (OA), previous joint trauma/injury, metabolic disease and familial predisposition. The complexity of CPPD in terms of variable phenotypes is compounded by use of different terminologies and classification. In 1961 McCarty and colleagues first identified CPP crystals in synovial fluid (SF) from knees of patients with acute synovitis and CC, introducing the term ‘CPPD’ for ‘calcium pyrophosphate dihydrate’ crystals. Similarity to gout prompted the term ‘pseudogout’ for this ‘crystal-induced arthropathy’. Subsequently other presentations were recognised, many appearing to mimic other forms of arthritis, encouraging proliferation of ‘pseudo’ syndromes and a complex clinical classification of ‘pseudogout’ (type A), ‘pseudo-rheumatoid arthritis’ (type B), ‘pseudo-osteoarthritis’ (with acute attacks, type C; without inflammation, type D), ‘lanthanic or asymptomatic’ (type E) and ‘pseudoneuropathic’ (type F), to which other forms were later added. The term ‘CPPD crystal deposition disease’ was introduced to incorporate all instances of CPP deposition, even though CPPD does not always appear injurious or causal in ‘disease’. The term ‘pyrophosphate arthropathy’ (PA) was later used, particularly in Europe, for CPPD with accompanying structural arthritis. However, some clinicians use the terms CC or pseudogout for any phenotype, whereas others restrict CC for radiographic calcification, pseudogout for acute synovitis and PA for CPPD plus OA.

The group that developed European League Against Rheumatism (EULAR) recommendations for gout considered it desirable to agree a uniform terminology for CPPD and to address issues relating to diagnosis and management. Therefore, the EULAR CPPD Task Force was formed to produce evidence-based recommendations using a combined systematic review and expert consensus approach. Part I, terminology and diagnosis, is presented here.

METHODS

Expert consensus

The Task Force comprised 15 experts from 10 countries. A meeting was organised to agree terminology; subsequently, preagreed terms were circulated for voting. Each participant independently submitted up to 10 propositions on key aspects of diagnosis; consensus was reached using the Delphi technique. A second meeting was organised to discuss recommendations and supporting evidence and to score strength of recommendations (SORs).
Systematic review of research evidence
As with previous projects, research evidence for each proposition was systematically searched (January 1950 to January 2009). Search terms included: calcium pyrophosphate dihydrate and/or deposition, CC, PA, pseudogout, crystal associated diseases/arthritis, crystal deposition diseases and calcium crystals. Studies informing diagnosis of CPPD were included. Case reports, reviews, editorials and commentaries were excluded.

Wherever possible, sensitivity, specificity, likelihood ratios (LR), proportion of CPPD and reliability (κ or intraclass correlation coefficient) were calculated for diagnostic tests; RR or ORs were estimated for association.9 Statistical pooling was undertaken as appropriate and a random effects model was used for heterogenous results.15 The EULAR level of evidence (LOE) for diagnosis,9 and EULAR 0–100 mm visual analogue scale (VAS)13 were used to rank LOE and SOR.

Future research agenda
After the second meeting each participant submitted independently up to 10 future research propositions; consensus was obtained by the Delphi technique.

RESULTS
Terminology of CPPD and its related conditions
The following terms and definitions were agreed:

1. CPP crystals: the simplified term for calcium pyrophosphate dihydrate crystals (similar to ‘sodium urate’ for monosodium urate monohydrate crystals)
2. CPPD: the umbrella term for all instances of CPP crystal occurrence
3. CC: cartilage calcification, identified by imaging or histological examination. This is not always due to CPPD and may occur as an isolated finding in an apparently otherwise normal joint or coexist with structural changes resembling OA.
4. Clinical presentations associated with CPPD:
   - Asymptomatic CPPD: CPPD with no apparent clinical consequence. This may be isolated CC, or OA with CC. Often this is identified incidentally following imaging for other reasons.
   - OA with CPPD: CPPD in a joint that also shows changes of OA, on imaging or histological examination.
   - Acute CPP crystal arthritis: acute onset, self-limiting synovitis with CPPD (replacing the term ‘pseudogout’).
   - Chronic CPP crystal inflammatory arthritis: chronic inflammatory arthritis associated with CPPD.
5. Risk factors that, if present, may be noted in phenotype characterisation:
   - Previous joint injury.
   - Hereditary/familial predisposition to CPPD.
   - Specific diseases (eg, haemochromatosis, primary hyperparathyroidism, hypophosphatasia, hypomagnesaemia).

EULAR recommendations
Of 102 initial propositions, 11 were agreed after 2 Delphi rounds. Recommendations covered four domains: clinical features, SF, imaging and risk factors/associations (see table 1 for the full

Table 1  Propositions and strength of recommendation (SOR), ordered according to topic (clinical features, synovial fluid, imaging, comorbidities and risk factors)

<table>
<thead>
<tr>
<th>No.</th>
<th>Proposition</th>
<th>LoE</th>
<th>SOR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Although often asymptomatic, CPPD can present variable clinical phenotypes, most commonly OA with CPPD, acute CPP crystal arthritis and chronic inflammatory arthritis.</td>
<td>Ib</td>
<td>90 (86 to 94)</td>
</tr>
<tr>
<td>2</td>
<td>The rapid development of severe joint pain, swelling and tenderness that reaches its maximum within 6–24 h, especially with overlying erythema, is highly suggestive of acute crystal inflammation though not specific for acute CPP crystal arthritis.</td>
<td>IV</td>
<td>88 (84 to 93)</td>
</tr>
<tr>
<td>3</td>
<td>Presentation with features suggesting crystal inflammation involving the knee, wrist or shoulder of a patient over age 65 years is likely to be acute CPP crystal arthritis. The presence of radiographic CC and advanced age increases this likelihood, but definitive diagnosis needs to be crystal proven.</td>
<td>Ib/IIb</td>
<td>81 (74 to 89)</td>
</tr>
<tr>
<td>4</td>
<td>OA with CPPD particularly targets knees with chronic symptoms and/or acute attacks of crystal-induced inflammation. Compared to OA without CPPD, it may associate with more inflammatory symptoms and signs, an atypical distribution (eg, radiocarpal or midcarpal, glenohumeral, hindfoot or midfoot involvement) and prominent cyst and osteophyte formation on radiographs.</td>
<td>Ib/IIb</td>
<td>53 (38 to 68)</td>
</tr>
<tr>
<td>5</td>
<td>Chronic CPP crystal inflammatory arthritis presents as chronic oligoarthritis or polyarthritis with inflammatory symptoms and signs and occasional systemic upset (with elevation of CRP and ESR); superimposed flares with characteristics of crystal inflammation support this diagnosis. It should be considered in the differential diagnosis of rheumatoid arthritis and other chronic inflammatory joint diseases in older adults. Radiographs may assist diagnosis, but the diagnosis should be crystal proven.</td>
<td>Ib</td>
<td>83 (72 to 93)</td>
</tr>
<tr>
<td>6</td>
<td>Definitive diagnosis of CPPD is by identification of characteristic CPP crystals (parallelepipedic, predominantly intracellular crystals with absent or weak positive birefringence) in synovial fluid, or occasionally biopsied tissue.</td>
<td>Ib</td>
<td>94 (90 to 97)</td>
</tr>
<tr>
<td>7</td>
<td>A routine search for CPP (and urate) crystals is recommended in all synovial fluid samples obtained from undiagnosed inflamed joints, especially from knees or wrists of older patients.</td>
<td>IV</td>
<td>99 (97 to 100)</td>
</tr>
<tr>
<td>8</td>
<td>Radiographic CC supports the diagnosis of CPPD, but its absence does not exclude it.</td>
<td>Ib</td>
<td>97 (92 to 102)</td>
</tr>
<tr>
<td>9</td>
<td>Ultrasonography can demonstrate CPPD in peripheral joints, appearing typically as thin hyperechoic bands within hyaline cartilage and hyperechoic sparkling spots in fibrocartilage. Sensitivity and specificity appear excellent and possibly better than those of conventional x-rays.</td>
<td>Ib</td>
<td>78 (70 to 87)</td>
</tr>
<tr>
<td>10</td>
<td>Acute CPP crystal arthritis and sepsis may coexist, so when infection is suspected microbiological investigation should be performed even if CPP crystals and/or CC are identified.</td>
<td>III</td>
<td>96 (93 to 100)</td>
</tr>
<tr>
<td>11</td>
<td>In patients with CPPD, risk factors and associated comorbidities should be assessed, including OA, prior joint injury, predisposing metabolic disease (including haemochromatosis, primary hyperparathyroidism, hypomagnesaemia) and rare familial predisposition. Metabolic or familial predisposition should particularly be considered in younger patients (&lt; 55) and if there is florid polyarticular CC.</td>
<td>Ib/IIb</td>
<td>94 (89 to 99)</td>
</tr>
</tbody>
</table>
Acute attacks Knee 1 (100) 36–80 Radiographic CC 0.50 (0.28 to 0.72) 0.90 (0.63 to 0.97) 1.18 (0.82 to 1.71) Doherty et al\textsuperscript{42}

Chronic CPP crystal inflammatory arthritis

Pain Knee, wrist, elbow, shoulder and hip 4 (429) 36–97 SF crystals and/or radiographic CC 0.21 (0.12 to 0.30) 0.80 (0.72 to 0.89) 1.96 (1.12 to 3.44) Gordon et al\textsuperscript{1}

Stiffness Knee 2 (227) 36–80 Radiographic CC 0.35 (0.19 to 0.50) 0.81 (0.74 to 0.88) 1.56 (1.16 to 2.11) Doherty et al\textsuperscript{42}

Swelling/effusion Knee, wrist, elbow, shoulder and hand 5 (432) 36–97 Radiographic CC 0.40 (0.23 to 0.56) 0.77 (0.68 to 0.87) 1.56 (1.16 to 2.11) Doherty et al\textsuperscript{42}

Tenderness Knee 2 (227) 36–80 Radiographic CC 0.13 (0.02 to 0.24) 0.85 (0.79 to 0.91) 0.91 (0.37 to 2.24) Doherty et al\textsuperscript{42}

Instability Knee 2 (227) 36–80 Radiographic CC 0.18 (0.06 to 0.31) 0.82 (0.51 to 1.14) 0.95 (0.44 to 2.04) Doherty et al\textsuperscript{42}

Synovial fluid

CPP crystals Not specified 194 Expert diagnosis 0.95 (0.92 to 1.00) 0.86 (0.80 to 0.93) 7.09 (4.27 to 11.76) Lumberas et al\textsuperscript{59}

Radiograph

CC Wrist 18 71 (41–95) CPP crystals 0.29 (−0.05 to 0.62) 0.20 (−0.15 to 0.55) 0.36 (0.10 to 1.25) Utsinger et al\textsuperscript{44}

Ultrasound

Cartilage calcification Knee 43 68 (40–92) CPP crystals 0.87 (0.69 to 1.04) 0.96 (0.90 to 1.03) 24.2 (3.51 to 168.01) Filippou et al\textsuperscript{62}

Achilles tendon calcification Heel 107 65 (42–92) Radiographic CC plus CPP crystals 0.56 (0.45 to 0.71) 0.98 (0.94 to 1.02) 29 (4.11 to 204.05) Falsetti et al\textsuperscript{68}

Plantar facia calcification Heel 107 65 (42–92) Radiographic CC plus CPP crystals 0.16 (0.06 to 0.25) 0.98 (0.94 to 1.02) 7.89 (1.04 to 60.16) Falsetti et al\textsuperscript{68}

CC, chondrocalcinosis; CPP, calcium pyrophosphate; LR, likelihood ratio.

**Table 2** Validity of diagnostic tests for calcium pyrophosphate deposition (CPPD)

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### Clinical features

**Proposition 1** (also see detailed proposition in table 1)

CC may be an apparently isolated phenomenon or occur with structural changes of OA.\textsuperscript{16–18} Hospital-based series suggest that OA with CPPD may differ from OA without CPPD in showing more osteophytosis,\textsuperscript{19–21} different joint involvement\textsuperscript{22–26} and more inflammatory features (table 3). However, whether OA with CPPD is a distinct OA ‘subset’ remains unclear. Isolated CC and OA with CPPD may be clinically occult or associate with acute CPP crystal arthritis; OA with CPPD may also associate with pain, stiffness and functional limitation. Much less commonly, atypical or periarticular CPPD may associate with tendinitis,\textsuperscript{27} tenosynovitis,\textsuperscript{28} bursitis,\textsuperscript{29} tumorous CPPD\textsuperscript{30} or syndromes relating to spinal involvement\textsuperscript{31}

**Proposition 2**

Direct literature evidence to support this recommendation was not found. However, one nested case-control study of postmeniscectomy knees showed those with CC had five times more risk (LR=5.00, 95% CI 2.27 to 11.02) of self-limiting acute attacks compared to those without CC.\textsuperscript{32} One hospital series observed that in acute CPP crystal arthritis symptoms and signs usually resolve within 3–4 days.\textsuperscript{33} Rapid development of acute synovitis with pain, stiffness, swelling/effusion and marked tenderness (erythema) is highly characteristic of crystal synovitis but these features are not specific to one crystal\textsuperscript{3} so crystal identification is required for precise diagnosis.

**Proposition 3**

A community cross-sectional survey in Sweden found that CC is most common in knees (8.5%), then wrists (5.1%) and hands (1.7%).\textsuperscript{34} Prevalence of knee CC in this study was similar to that in the USA (8.1%)\textsuperscript{17} and UK (7.0%)\textsuperscript{18} studies so this distribution may be generalisable to other populations. Several hospital series report similar distribution but greater prevalence for each joint\textsuperscript{35} (figure 1). The glenohumeral joint appears less common affected.\textsuperscript{36} Ageing is a major risk factor. CPPD is rare under age 50,\textsuperscript{18} but increases dramatically afterwards\textsuperscript{2} (OR=2.27 to 11.02) (figure 2). The risk doubles every decade between 45 and 85 years (OR=2.25, 95% CI 1.79 to 2.82) independently of other risk factors.\textsuperscript{34} CPPD under age 45 should raise the possibility of familial\textsuperscript{37} or CPP crystal-induced synovitis according to site and age has not been calculated. Radiographic CC is often taken as a surrogate for CPPD but does not necessarily predict SF CPP crystal identification according to limited evidence for the wrist\textsuperscript{34} (table 2) and the knee.\textsuperscript{35} Definitive diagnosis of CPPD has to
be crystal proven usually by examination of SF (occasionally histological).

**Proposition 4**
The knee is a target site for OA \(^46\)–\(^48\) and the most common site for CPPD \(^1\)\(^34\)–\(^36\) (figure 1). A total of 4 cross-sectional and 5 case-control studies (4517 subjects in all) provided quantitative data for analysis of the association between OA and CPPD. \(^1\)\(^17\)\(^18\)\(^20\)\(^38\)\(^49\)–\(^52\). The pooled OR was 2.66 (95% CI 2.00 to 3.54). Results were consistent between cross-sectional (2.52, 95% CI 1.86 to 3.44) and case-control (1.10 (0.58 to 2.08) studies, suggesting people with OA are three times more likely to have CPPD.

CPPD may associate with more inflammatory features (eg, pain, stiffness, effusion, more severe pain and disability) and more rapid progression\(^53\) than in knees without CPPD, but the associations are marginal and not useful for diagnosis (table 2).

CPPD may associate with more inflammatory features (eg, pain, stiffness, effusion, more severe pain and disability) and more rapid progression\(^53\) than in knees without CPPD, but the associations are marginal and not useful for diagnosis (table 2).

### Table 3 Risk factors and comorbidities associated with calcium pyrophosphate deposition (CPPD)

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>No. studies</th>
<th>No. subjects</th>
<th>OR (95% CI)*</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (every 10 years from 40 to 90)</td>
<td>1</td>
<td>1851</td>
<td>2.25 (1.79 to 2.82)</td>
<td>Zhang et al(^41)</td>
</tr>
<tr>
<td>Female gender</td>
<td>8</td>
<td>5042</td>
<td>0.89 (0.58 to 1.38)</td>
<td>Gordon et al(^1) Felson et al(^3) Neame et al(^18) Cruz et al(^34) Doherty et al(^38) Vinyayejujul et al(^37) Ellman et al(^39) Sanmarti et al(^50)</td>
</tr>
<tr>
<td>BMI (WHO grade)</td>
<td>1</td>
<td>1851</td>
<td>0.90 (0.70 to 1.14)</td>
<td>Zhang et al(^41)</td>
</tr>
<tr>
<td>Familial aggregation</td>
<td>2</td>
<td>2000</td>
<td>1.10 (0.58 to 2.08)</td>
<td>Zhang et al(^41) Fernández Dapica and Gómez-Reino(^16)</td>
</tr>
<tr>
<td>OA</td>
<td>9</td>
<td>4517</td>
<td>2.66 (2.00 to 3.54)</td>
<td>Gordon et al(^1) Felson et al(^3) Neame et al(^18) Riestra et al(^20) Doherty et al(^38) Al-Arfaj(^49) Menkes et al(^50) Sanmarti et al(^50) Stucki et al(^52)</td>
</tr>
<tr>
<td>OST</td>
<td>3</td>
<td>1906</td>
<td>1.26 (0.76 to 2.09)</td>
<td>Neame et al(^18) Bourqui et al(^34) Hansen et al(^55)</td>
</tr>
<tr>
<td>JSN</td>
<td>4</td>
<td>2043</td>
<td>1.24 (0.91 to 1.69)</td>
<td>Neame et al(^18) Riestra et al(^20) Schouten et al(^54) Bourqui et al(^54)</td>
</tr>
<tr>
<td>Cysts</td>
<td>3</td>
<td>367</td>
<td>2.94 (0.92 to 4.96)</td>
<td>Ledingham et al(^53) Bourqui et al(^34) Hansen et al(^55)</td>
</tr>
<tr>
<td>Trauma/injury</td>
<td>1</td>
<td>100</td>
<td>5.00 (1.77 to 14.11)</td>
<td>Doherty et al(^32)</td>
</tr>
<tr>
<td>RA</td>
<td>2</td>
<td>818</td>
<td>0.18 (0.08 to 0.41)</td>
<td>Doherty et al(^32) Brasseur et al(^36) Yashiro et al(^38) Alexander et al(^38) Haux et al(^37) Pritchard and Jessop(^18) Rynes and Merzig(^69)</td>
</tr>
<tr>
<td>Hyperparathyroidism</td>
<td>5</td>
<td>976</td>
<td>3.03 (1.15 to 8.02)</td>
<td></td>
</tr>
<tr>
<td>Hypomagnesaemia</td>
<td>1</td>
<td>144</td>
<td>13.5 (2.76 to 127.3)</td>
<td></td>
</tr>
<tr>
<td>Diuretics</td>
<td>1</td>
<td>1277</td>
<td>2.17 (1.02 to 4.19)</td>
<td></td>
</tr>
</tbody>
</table>

*Meta-analysis was undertaken to pool results from multiple studies.*

BMI, body mass index; JSN, joint space narrowing; OA, osteoarthritis; OST, osteophyte; RA, rheumatoid arthritis.

**Proposition 5**
Although the population prevalence is unknown, hospital series of chronic CPP crystal inflammatory arthritis show that most are mono/oligoarthritis (9%) though some (11%) are polyarticular.\(^55\) There may be non-specific elevation of C reactive protein and erythrocyte sedimentation rate\(^53\) but diagnostic identification of CC is more useful since this mainly results from CPPD and is less likely to occur with rheumatoid arthritis (OR=0.18, 95% CI 0.08 to 0.41)\(^50\)\(^56\) (table 3). Nevertheless, definitive diagnosis relies on CPP crystal identification, usually in SF.
**Recommendations**

**SF examination**

Proposition 6

Identification of SF CPP crystals (usually by light, compensated polarised light or phase contrast microscopy) is the recommended reference standard for diagnosis of CPP crystal-associated arthritis (figure 3). First suggested by McCarty in 1962, this has been used repeatedly as a major diagnostic feature for CPP syndromes. Its validity and reliability have been systematically reviewed. \(^5\) \(^7\) Training in SF CPP crystal identification results in better sensitivity (0.95, 95% CI 0.92 to 1.02) and reasonable specificity (0.86, 95% CI 0.80 to 0.93) compared to expert (‘gold standard’) identification \(^5\) \(^9\) (table 2). Trained observer reliability is very good (\(\kappa=0.79\), 95% CI not reported) \(^5\) \(^9\). No quantitative cut-off of crystals is recommended; even one or a few crystals are clinically significant. The more characteristic the crystal morphology, the more confident the diagnosis.

Proposition 7

Although there are no specific studies, examination of SF for CPP crystals should be undertaken for any undiagnosed inflammatory arthritis since CPPD is a common cause of joint inflammation \(^2\) and may present atypically.

**Imaging**

Proposition 8

Radiographic CC is a useful imaging marker, often taken as a surrogate for CPPD (figure 4), but common discordance with positive SF crystal identification reduces its diagnostic usefulness. \(^3\) \(^5\) \(^4\) \(^4\) \(^5\) \(^6\) Detection of CC in proven cases of CPP crystal arthritis varies from 29% to 95% depending on population and joint examined. \(^3\) \(^5\) \(^4\) \(^4\) \(^5\) \(^6\) The sensitivity and specificity of CC against the ‘gold’ standard remain unknown. However, one small case control study (n=18) \(^4\) suggested that at the wrist CC is neither sensitive (0.29) nor specific (0.20) for diagnosis (table 2); the likelihood of a patient with wrist CC having SF CPP crystals identified was only 3%. However, results of this
small study may not be generalisable and further studies are required.

There are several possible reasons for discordance between CC and SF crystal positivity including: lack of specificity of CC for CPPD (basic calcium phosphates may also cause this); low sensitivity of radiographs for detecting CC; possible greater difficulty in identifying small numbers of SF CPP crystals in situations other than acute CPP crystal synovitis (especially when there is isolated CC and no cartilage fibration to encourage crystal shedding); and reduced ability to identify CC when there is significant cartilage loss. Also being small and weakly, or non-birefringent, CPP crystals may be underdetected.

Proposition 9
Case-control studies have examined the usefulness of ultrasonography (US) in detecting calcification in knees (figure 5),

wrist and shoulders.

US of the knee appears sensitive (0.87, 95% CI 0.69 to 1.04) and specific (0.96, 95% CI 0.90 to 1.08) for detection of SF CPP crystals. Positive US findings may strongly suggest the diagnosis of CPPD (LR=24.2, 95% CI 3.51 to 168.01). Given a positive US result, the likelihood of a patient having SF CPP crystals in the same joint (eg, knee) in the UK population, for example, would be 65%. Suitability of US to detect calcification varies between sites (table 2) and it is insensitive for deep structures (eg, spine). One direct comparison showed US to be more sensitive (100%) than x-rays (82%) in identifying CPPD. However, although US seems a promising technique for crystal identification the few published studies emanate from just a few centres and further studies are required.

Comorbidities and risk factors
Proposition 10
One study in a tertiary centre retrospectively reviewed all cases with crystal positive SF samples identified over a 7-year period. 265 positive samples, 183 (69.0%) contained MSU crystals, 81 (30.6%) contained CPP crystals and 1 (0.4%) contained both. Four (1.5%) also had positive cultures; of these, three were from joints with CPP crystals.

Proposition 11
As discussed, OA and ageing are major risk factors for CPPD. There are common risk factors for OA and CPPD (eg, ageing, joint injury) but OA cartilage also directly encourages deposition of calcium crystals (basic calcium phosphates as well as CPP). Conversely, CPPD could be a primary factor that causes, or amplifies joint damage in OA. Whether gender influences CPPD remains controversial, but pooling of eight cross sectional/case-control studies yields a non-significant association (OR=0.89, 95% CI 0.58 to 1.38). Although body mass index is a major risk factor for OA, there is no evidence of association with CPPD (table 3).

Future research agenda
After four Delphi rounds five propositions were agreed.

1. Whether there are significant differences (eg, in symptoms, joint distribution, clinical outcomes) between OA with and without CPPD requires further study.
2. Clinical studies (using crystal identification as the gold standard for diagnosis) are required to better define the spectrum of CPPD and its possible clinical subsets.
3. An optimal protocol (including training) and agreed European standard for the identification of CPP crystals in SF needs to be established.
4. The validity of different imaging techniques for the diagnosis of CPPD should be evaluated, leading to recommendations about their application in routine care.
5. Further studies are required to determine whether correction of associated metabolic disease (eg, primary hyperparathyroidism, haemochromatosis) influences progression and outcome of CPPD associated arthropathy.
DISCUSSION
A large number of terms are used for CPPD and its clinical phenotypes. However, the plethora of ‘pseudo’ syndromes and inconsistent use of terms such as pseudogout and CC often cause confusion. The EULAR CPPD Task Force therefore suggests ‘calcium pyrophosphate deposition’ (CPPD) as the umbrella term for all instances of CPP crystal deposition. Under this umbrella, asymptomatic CPPD, OA with CPPD, acute CPP crystal arthritis and chronic CPP crystal inflammatory arthritis are included. It was felt that terms prefixed by ‘pseudo’ should be abandoned because they do not specify the causative crystal, are probably not discrete clinical subsets, are a source of potential confusion for patients, and intimate that CPP crystals are of secondary importance and interest compared to sodium urate. The term chondrocalcinosis is retained for cartilage calcification, which is most commonly due to CPPD. It is appreciated that introduction of new terminology may prove inconvenient in the short term, but would be beneficial in the long term for research, education and clinical practice.

There are several limitations to this project. Firstly, we established expert consensus prior to systematic review of research evidence. While this approach addresses clinically relevant questions, it is not comprehensive. Secondly, the recommendations are not intended to classify CPPD for research purposes but to provide recommendations for clinical diagnosis. Thirdly, clinical studies used different gold standards for diagnosis, making comparison between diagnostic tests difficult. Furthermore, research evidence for diagnosis of CPPD is generally sparse and of poor quality. We therefore provided a suggested future research agenda to encourage expansion of the evidence base.

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None.

REFERENCES


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