European Guideline on Chronic Pruritus

In cooperation with the European Dermatology Forum (EDF) and the European Academy of Dermatology and Venereology (EADV)

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Abbreviations and Explanations

AD: Atopic dermatitis
AEP: Atopic eruption of pregnancy
CGRP: Calcitonin gene-related peptide
CKD: Chronic kidney disease
CP: Chronic pruritus (longer than 6 weeks)
DIF: Direct immunofluorescence
ICP: Intrahepatic cholestasis of pregnancy
IFSI: International Forum on the Study of Itch
IIF: Indirect immunofluorescence
IL: Interleukin
Itch: Synonymous with pruritus
NSAID: Non-steroidal anti-inflammatory drugs
PAR: Proteinase-activated receptor

PBC: Primary biliary cirrhosis
PEP: Polymorphic eruption of pregnancy
PG: Pemphigoid gestationis
PN: Prurigo nodularis
Pruritus: A skin sensation which elicits the urge to scratch
PUO: Pruritus of unknown origin
PTH: Parathyroid hormone
PV: Polycythaemia vera
RCT: Randomised controlled trials
SSRI: Selective serotonin re-uptake inhibitors
TRP: Transient receptor potential
UV: Ultraviolet
VIP: Vasoactive intestinal peptide
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1. THE CHALLENGE OF WRITING THIS GUIDELINE

Chronic pruritus (CP) is a frequent symptom in the general population and in many skin and systemic diseases (1). Its frequency demonstrates a high burden and an impaired quality of life. This guideline addresses a symptom and not a disease. As a consequence of the diversity of possible underlying diseases, no single therapy concept can be recommended. Each form of pruritus has to be considered individually. There is still a significant lack of randomised controlled trials (RCT), that can be explained by the diversity and complexity of this symptom, multifactorial aetiologies of pruritus and the lack of well-defined outcome measures. To complicate matters, RCT exist for some types of pruritus, but with conflicting results. However, new therapies for improved medical care have been suggested. In addition, many expert recommendations are provided. The health care system in many countries and their social economic situation with constantly reducing financial resources increases the need for guidelines. These recommendations are based on a consensus of participating countries, while also allowing for country-specific treatment modalities, and health care structures. Furthermore, it should be appreciated that some topical and systemic therapies can only be prescribed “off-label” and require informed consent. If such “off-label” therapies cannot be initiated in the physician’s office, cooperation with a specialised centre for pruritus might be helpful.

This guideline addresses all medical disciplines that work with patients suffering from CP. This includes also entities defined by chronic scratch lesions such as prurigo nodularis and lichen simplex. The guidelines are not only focussed on dermatology.

2. DEFINITIONS AND CLINICAL CLASSIFICATION

The definitions presented in this guideline are based on a consensus among the European participants; however, some of them have provoked controversy. Most of the contributors accept pruritus and itch to be synonymous. A practical distinction is that between acute pruritus and chronic forms (lasting six weeks or longer). Pruritus/itch is a sensation that provokes the desire to scratch. According to the International Forum on the Study of Itch (IFSI), CP is defined as pruritus lasting 6 weeks or longer (2). Following the IFSI, the term “pruritus sine materia” will not be used in this guideline (3). In patients with no identified underlying disease, the term “pruritus of unknown origin” or “pruritus of undetermined origin” (PUO) is used. The term “pruritus of unknown aetiology” should be avoided as in most clinically well-defined forms of pruritus the mechanism is unknown (e.g. chronic kidney disease (CKD) associated pruritus). This guideline addresses patients presenting with CP of different, including unknown, origin. If the underlying cause is detected, disease-specific guidelines should be consulted (e.g. atopic dermatitis (AD), cholestatic pruritus) (4–6).

According to the IFSI classification, the aetiology of CP is classified as I “dermatological”, II “systemic”, III “neurological”, IV “somatoform”, V “mixed origin” and VI “others” (2). The IFSI classification comprises a clinical distinction of patients with pruritus on primarily diseased/inflamed skin, pruritus on normal skin and pruritus with chronic secondary scratch lesions.

Somatoform pruritus is defined as pruritus where psychiatric and psychosomatic factors play a critical role in the initiation, intensity, aggravation or persistence of the pruritus. It is best diagnosed using positive and negative diagnostic criteria (5).

3. EPIDEMIOLOGY OF CHRONIC PRURITUS

Data on the prevalence of CP is very limited. The prevalence of CP seems to increase with age (7), but epidemiological studies are missing. It is estimated that about 60% of the elderly (≥65 years of age) suffer from mild to severe occasional pruritus each week (8), entitled senile pruritus or pruritus in the elderly. A population-based cross-sectional study in 19,000 adults showed that about 8–9% of the general population experienced acute pruritus, which was a dominant symptom across all age groups (9). Moreover, it was revealed that pruritus is strongly associated with chronic pain (10). Recent surveys indicate a point-prevalence of CP to be around 13.5% in the general adult population (11) and 16.8% in employees seeking detection cancer screenings (12). The 12-month prevalence of CP was 16.4% and its lifetime prevalence 22.0% in a German population-based cross-sectional study (11). All these data suggest a higher prevalence of CP in the general population than previously reported (11).

CP may be due to both dermatological and systemic diseases. However, the origin of pruritus is unknown in 8–15% of affected patients (1). The frequency of pruritus among patients with a primary rash depends on the skin disease. For example, pruritus is present in all patients with AD and urticaria (13), and about 80% of psoriatic patients (14, 15). Systemic diseases such as primary biliary cirrhosis (PBC) and CKD are associated with CP in 80–100% and 40–70%, respectively (16). In patients with Hodgkin’s lymphoma, pruritus is a frequent symptom, occurring in more than 30% of patients with Hodgkin’s disease.

Only few studies have addressed the frequency of pruritus in primary care. According to the Australian BEACH Program, a continuous national study of general practice activity, pruritus was the presenting complaint for 0.6% of consultations, excluding perianal, periorbital or auricular pruritus (17). In Britain, the
fourth national study of morbidity statistics from general practice (18) was conducted in 1991/1992 with 502,493 patients (1% sample of England and Wales), resulting in 468,042 person-years-at-risk. Pruritus and related conditions was present in 1.04% of consultations (male 0.73%, female 1.33%). On Crete, where patients with cutaneous disorders mostly present to hospitals rather than to primary care, PUO was diagnosed in 6.3% of 3,715 patients in 2003 (19).

4. THE CLINICAL PICTURE OF CHRONIC PRURITUS

4.1. Pruritus in diseased conditions

4.1.1. Pruritus in inflamed and non-inflamed skin. CP may occur as a common symptom in patients with dermatoses with primary skin lesions and systemic diseases without primary skin lesions. In systemic diseases, the skin may appear normal or have skin lesions induced by scratching or rubbing. In this case, a diagnosis might be difficult to establish. Systemic diseases frequently accompanied by pruritus are summarized in Table I. In some cases, pruritus may precede the diagnosis of the underlying disease by years. In the past years, several mechanisms of pruritus on inflamed and normal skin have been identified (see original version of EDF guideline, www.euroderm.org). In the following paragraphs some frequent patient populations and systemic diseases inducing CP are presented.

4.1.2. Pruritus in kidney disease. The pathophysiology of CKD-associated pruritus is unknown. Implicated mechanisms have included direct metabolic factors like increased concentrations of divalent ions (calcium, magnesium), parathyroid hormone (PTH), histamine and tryptase, dysfunction of peripheral or central nerves, the involvement of opioid receptors (μ- and κ-receptors) and xerosis cutis (dry skin) have been suggested as likely candidates (20–28). New data point to a possible role of microinflammation that is quite frequent in uraemia (20, 29).

4.1.3. Pruritus in hepatic diseases. In patients with cholestasis due to mechanical obstruction, metabolic disorders or inflammatory diseases, CP is a frequent symptom (30). It may be quite severe and can even precede the diagnosis of e.g. PBC by years (31). In patients with infective liver disease (hepatitis B or C) or toxic liver disease (e.g. alcohol-induced), pruritus is less frequent. Hepatic pruritus is often generalised, affecting palms and soles in a characteristic way (32). One hypothesis for the mechanism of hepatic pruritus suggests that high opioid tone influences neurotransmission (30). Successful treatment with μ-receptor opioid antagonists such as nalmefene supports this hypothesis (33). It has recently been shown that increased serum autotaxin levels (enzyme that metabolizes lysophosphatidylcholine (LPC) into lysophosphatidic acid (LPA)) and thereby increased LPA levels are specific for pruritus of cholestasis, but not for other forms of systemic pruritus (34). Rifampicin significantly reduced itch intensity and ATX activity in pruritic patients. The beneficial antipruritic action of rifampicin may be explained partly by pregnant X receptor (PXR)-dependent transcription inhibition of ATX expression (34).

4.1.4. Pruritus in metabolic and endocrine diseases. In endocrine disorders as hyperthyroidism and diabetes mellitus, less than 10% of patients report pruritus (35, 36). In patients with hypothyroidism, pruritus is most probably driven by xerosis of the skin. Patients with primary hyperparathyroidism do complain about itch in a substantial number of cases (37). The pathophysiology of pruritus in primary hyperparathyroidism is not known. These patients often experience a lack of vitamin D and minerals (e.g. zinc, etc.) which probably contributes to CP. Iron deficiency is frequently associated with pruritus (38). The mechanism for this is unknown. Iron overload as in hemochromatosis may lead to CP (39, 40).

4.1.5. Pruritus in malignancy. Several malignant disorders including tumours, bone marrow diseases and lymphoproliferative disorders may be accompanied by pruritus. In addition to toxic products generated by the tumour itself, allergic reactions to compounds released, and a direct affection on the brain or nerves (in brain tumours) may be the underlying mechanism (8, 41). In polycythemia vera (PV), more than 50% of patients suffer from pruritus (42, 43). Aquagenic pruritus with pinching sensations after contact with water is a characteristic but not necessary feature. It has been suggested that high levels of histamine released by the augmented numbers of basophilic granulocytes might trigger the itch (44). For PV this seems to be most pronounced in patients showing the JAK2 617V mutation (45).
Pruritus in Hodgkin’s disease often starts on the legs and is most severe at night, but generalised pruritus soon ensues. Several factors such as secretion of leukopeptidases and bradykinine, histamine release and high IgE levels with cutaneous depositions may contribute to pruritus in lymphoma (46). Patients with carcinoid syndrome may experience pruritus in addition to flushing, diarrhoea and cardiac symptoms (47).

4.1.6. Pruritus in infectious diseases. Some generalised infections are accompanied by pruritus. Above all, patients infected with HIV may develop a pruritic papular eruption or eosinophilic folliculitis. These entities are easily diagnosed by inspection and histology of the skin and have a high positive predictive value (48, 49). Whether toxocara infections lead to pruritus in a substantial number of patients remains to be confirmed (50).

4.1.7. Pruritus in neurological diseases. Multiple sclerosis, brain infarction and brain tumours are rarely accompanied by pruritus (51, 52). Localised pruritus suggests a neurological origin such as compression of the peripheral or central afferences. This neuropathic origin of localised CP can be found e.g. in postzosteric pruritus, nostalgia paraesthetica and brachioradial pruritus, where an underlying spinal damage is likely (53–56).

4.1.8. Drug induced chronic pruritus. Almost every drug may induce pruritus by various pathomechanisms (Table II) (57). Some may cause urticarial or morbilliform rashes presenting with acute pruritus. Furthermore, drug-induced hepatotoxicity or cholestasis as well as drugs leading to xerosis or phototoxicity may produce CP on normal skin (58). Hydroxyethyl starch, a compound used for fluid restoration, can induce chronic generalised or localised pruritus (59).

### Table II. Drugs that may induce or maintain chronic pruritus (without a rash)

<table>
<thead>
<tr>
<th>Class of drug</th>
<th>Substance (examples)</th>
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<tbody>
<tr>
<td>ACE inhibitors</td>
<td>Captopril, enalapril, lisinopril</td>
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<tr>
<td>Antiarrhythmic agents</td>
<td>Amiodarone, disopyramide, flecaïnine</td>
</tr>
<tr>
<td>Antibiotics</td>
<td>Amoxicillin, ampicillin, cefotaxime, ceftriaxone, chloramphenicol, ciprofloxacin, clarithromycin, clindamycin, cotrimoxazole, erythromycin, gentamycin, metronidazole, minocycline, ofloxacin, penicillin, tetracycline</td>
</tr>
<tr>
<td>Antidepressants</td>
<td>Amitriptylin, citalopram, clomipramin, desipramine, doxepin, fluoxetine, fluvoxamine, imipramine, lithium, maprotiline, mirtazapine, nortriptylne, paroxetine, sertraline</td>
</tr>
<tr>
<td>Antidiabetic drugs</td>
<td>Glimepiride, metformin, tolbutamide</td>
</tr>
<tr>
<td>Antihypertensive drugs</td>
<td>Clonidine, doxazosin, hydralazine, methyl dopa, minoxidil, prazosin, reserpine</td>
</tr>
<tr>
<td>Anticonvulsants</td>
<td>Carbamazepine, clonazepam, gabapentin, lamotrigine, phenobarbital, phenytoin, topiramate, valproic acid</td>
</tr>
<tr>
<td>Anti-inflammatory drugs</td>
<td>Acetylsalicic acid, celecoxib, diclofenac, ibuprofen, indometacin, ketoprofen, naproxen, piroxicam</td>
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<tr>
<td>AT II antagonists</td>
<td>Irbesartan, telmisartan, valsartan</td>
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<tr>
<td>Beta blockers</td>
<td>Acebutolol, atenolol, bisoprolol, metoprolol, nadolol, pindolol, propanolol</td>
</tr>
<tr>
<td>Bronchodilators, mucolytic agents, respiratory stimulants</td>
<td>Aminophylline, doxapram, ipratropium bromide, salmeterol, terbutaline</td>
</tr>
<tr>
<td>Calcium antagonists</td>
<td>Amlodipine, diltiazem, felodipine, isradipine, nifedipine, nimodipine, nisoldipine, verapamil</td>
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<tr>
<td>Diuretics</td>
<td>Amiloride, furosemide, hydrochlorothiazide, spironolactone, triamterene</td>
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<tr>
<td>Hormones</td>
<td>Clomifene, danazol, oral contraceptives, estrogens, progesterone, steroids, testosterone and derivatives, tamoxifen</td>
</tr>
<tr>
<td>Immunosuppressive drugs</td>
<td>Cyclophosphamide, cyclosporine, methotrexate, mycophenolate mofetil, tacrolimus (up to 36%), thalidomide</td>
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<tr>
<td>Antilipids</td>
<td>Clofibrate, fenofibrate, fluvastatin, lovastatin, pravastatin, simvastatin</td>
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<tr>
<td>Neuroleptics</td>
<td>Chlorpromazine, haloperidol, risperidone</td>
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<tr>
<td>Plasma expanders, blood supplying drugs</td>
<td>Hydroxyethyl starch, pentoxifylline</td>
</tr>
<tr>
<td>Tranquilizers</td>
<td>Alprazolam, chloridazepoxide, lorazepam, oxazepam, prazepam</td>
</tr>
<tr>
<td>Uricostatics</td>
<td>Allopurinol, colchicine, probenecid, tiopronin</td>
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</table>
4.2.2. Chronic pruritus in pregnancy. There are no epidemiological studies assessing the prevalence of CP in pregnancy. Pruritus is the leading dermatological symptom in pregnancy estimated to occur in about 18% of pregnancies (64). Pruritus is the leading symptom of the specific dermatoses of pregnancy such as polymorphic eruption of pregnancy (PEP), pemphigoid gestationis (PG), intrahepatic cholestasis of pregnancy (ICP), atopic eruption of pregnancy (AEP), but may also occur in other dermatoses coinciding by chance with pregnancy or in pre-existing dermatoses (64–67). PEP is one the most common gestational dermatoses, affecting about one in 160 pregnancies. While PG, PEP and ICP characteristically present in late pregnancy, AEP starts in 75% of cases before the third trimester (1, 65).

ICP is characterised by severe pruritus without any primary skin lesions, but secondary skin lesions occur due to scratching. It is more prevalent among native Indians in Chile (27.6%) and Bolivia (13.8%) depending on ethnic predisposition and dietary factors (68, 69). ICP has decreased in both countries, e.g. to 14% in Chile. ICP is more common in women of advanced maternal age, multiple gestations, personal history of cholestasis on oral contraceptives and during winter months. Scandinavian and Baltic countries are also more affected (1–2%). In Western Europe and North America, ICP is observed in in 0.4–1% of pregnancies (68–70).

The use of topical and systemic treatments depends on the underlying aetiology of pruritus and the stage and status of the skin. Because of potential effects on the foetus, the treatment of pruritus in pregnancy requires prudent consideration of whether the severity of the underlying disease warrants treatment and selection of the safest treatments available. Systemic treatments such as systemic glucocorticosteroids, a restricted number of antihistamines and ultraviolet phototherapy, e.g. UV A, may be necessary in severe and generalised forms of CP in pregnancy.

4.2.3. Chronic pruritus in children. There are no epidemiological studies assessing the prevalence of CP in children (1, 64). Differential diagnosis of CP in children has a wide spectrum (64) but is dominated by AD. The cumulative prevalence of AD is between 5 to 22% in developed countries. The German Atopic Dermatitis Intervention Study (GADIS) showed a significant correlation between the pruritus intensity and severity of AD and sleeplessness (71, 72). A Norwegian cross-sectional questionnaire-based population study in adolescents revealed a pruritus prevalence of 8.8%. Pruritus was associated with mental distress, gender, sociodemographic factors, asthma, rhinoconjunctivitis and eczema (73). Itching of mild to moderate severity may occur in acne (74, 75).

There are no studies about systemic causes of CP among children. It must be assumed that systemic causes in children are mostly based on genetic diseases or systemic diseases, e.g. biliary atresia or hypoplasia, familial hyperbilirubinemia syndromes, polycystic kidney disease. Drug-induced pruritus without any specific skin symptoms appears to be rare in children (1). Common medications associated with CP in adults play a minor role in children due to limited use at that age.

When considering treatment, the physician must remember that topically applied drugs may cause intoxication due to the different body volume/body surface area rate. In addition, the licensed age for the drug must be taken into account. Low- (class 1, 2) to medium-strength (class 3) glucocorticosteroids may be applied in children. Topical immunomodulators are used for AD and pruritus in children ≥ 2 years, but in some European countries e.g. pimecrolimus is licensed for use in children > 3 months. Topical capsaicin is not used in children < 10 years. The dosages of systemic drugs need to be adapted in children. Ultraviolet phototherapy should be performed with caution due to possible long-term photodamage of the skin.

5. DIAGNOSTIC MANAGEMENT

5.1. Patient’s history, examination and clinical characteristics of pruritus

The collection of the patient’s history and a thorough clinical examination are crucial at the first visit, as it forms an assessment of their pruritus including intensity, onset, time course, quality, localisation, triggering factors and the patient’s theory of causality. Attention should be paid to incidents preceding or accompanying the onset of pruritus (e.g. pruritus following bathing). It is also important to consider the methods used to relieve pruritus, e.g. brushes. This helps with the interpretation of clinical findings such as the absence of secondary skin lesions in the mid-back known as the “butterfly sign” that indicates that the patient cannot reach this area by hand and is thus unable to scratch it. It is also important to ask about preexisting diseases, allergies, atopic diathesis and drug intake (Table II). A great deal of helpful information can be obtained using questionnaires. There are no definite clinical findings related to specific pruritic diseases (76), but awareness of the following anamnestic aspects and clinical findings may help with the diagnosis of the cause of pruritus:

- When several family members are affected, scabies or other parasites should be considered.
- The relationship between pruritus and special activities is important: Pruritus during physical activity is suggestive of cholinergic pruritus. It is common in patients with atopic eczema and mild forms of cholinergic pruritus. Pruritus provoked by skin cooling after bathing should prompt consideration of aquagenic pruritus. It may be associated with or precede PV or myelodysplastic syndrome, and
screening for these diseases should be performed intermittently.
• Nocturnal generalised pruritus associated with chills, fatigue, tiredness and “B” symptoms (weight loss, fever and nocturnal sweating) raises the possibility of Hodgkin’s disease.
• Somatoform pruritus rarely disturbs sleep; most other pruritic diseases cause nocturnal wakening.
• Seasonal pruritus frequently presents as “winter itch”, which may also be the manifestation of pruritus in the elderly due to xerosis cutis and asteatotic eczema.

A patient’s history should always include all current and recent medications, infusions, and blood transfusions. Severe pruritus can lead to considerable psychological distress. This should not be underestimated by the physician and should be addressed directly. CP can be accompanied by behavioural/adjustment disorder and a withdrawal from social and work life (77). In these cases, psychosomatic counselling is required. CP with excoriations sometimes progressing to self-mutilation can be caused by psychiatric disease such as delusional parasitosis. Such patients need psychiatric examination and if necessary treatment. A solely psychological cause of pruritus should not be diagnosed without psychiatric examination.

Examination of patients with CP includes a thorough inspection of the entire skin including mucous membranes, scalp, hair, nails, and anogenital region. The distribution of primary and secondary skin lesions should be recorded together with skin signs of systemic disease. General physical examination should include palpation of the liver, kidneys, spleen, and lymph nodes.

There is no standardised method of documenting pruritus. The sensation of pruritus is subject to much inter- and intra-individual variation due to tiredness, anxiety, stress. Questionnaires deliver self-reported information regarding various aspects of CP. So far, no structured questionnaire exists, but the questionnaire should consider the patients’ perspective, the medical doctors’ perspective and needs of various measurements of clinical trials. Several different questionnaires in different languages for different pruritic diseases have been developed, but so far no definite questionnaires exist. Additional tools are needed to better assess the different dimensions of CP and better tailor management. With this goal in mind, a special interest group (SIG) was initiated by members of the IFSI to determine which of the various psychometric properties of CP questionnaires offer the greatest utility in the evaluation of CP (78). The intensity of pruritus is usually assessed by scales such as the visual analogue scale (VAS) or the numeric rating scale (79, 80). When using a VAS, the scale ranges from 0–10 and is graphically presented as a bar chart. However, these methods often fail to consider the frequency of itch attacks over the course of a day. For patients with severe PUO, it can be helpful to keep a diary in order to allow for clearer attribution of the symptoms.

5.2. Diagnostic algorithm and diagnostics
Laboratory screening, clinical and technical approaches and investigations are summarised in Table III and IV. All this helps to follow a diagnostic algorithm (Fig. 1).

6. THERAPY
6.1. General principles
In the patient with CP it is important to establish an individual therapy regimen according to their age, pre-existing diseases, medications, quality and intensity of pruritus. Most importantly, elderly patients, pregnant women and children need special attention. As the care of patients with CP often extends over a long period, with initial uncertainty about the origin of their pruritus, frustration regarding the failure of past therapies and general psychological stress frequently occurs. The diagnostic procedures and therapy should be discussed with the patient in order to achieve best possible concordance and compliance. It must be remembered that some therapies are not licensed for CP and can only be prescribed “off-label”. This requires separate informed consent.

<table>
<thead>
<tr>
<th>Table III. Diagnostics: laboratory screening, diverse approaches and investigations</th>
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<td>Chronic pruritus:</td>
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<td>First-step lab screening</td>
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<td>Chronic pruritus:</td>
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<td></td>
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</tbody>
</table>

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**Table IV. Laboratory and technical investigations in chronic pruritus due to systemic diseases**

<table>
<thead>
<tr>
<th>Laboratory and technical screening-basic</th>
<th>Creatinine, AST, ALT, alkaline phosphatase, bilirubin, TSH, complete blood count, glucose, chest X-ray, (Ca, y-GT, stool test for parasites in genito-anal pruritus)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Metabolic and endocrine diseases</strong></td>
<td></td>
</tr>
<tr>
<td>Renal insufficiency</td>
<td>Lab I: Creatinine, (and urea for elderly)</td>
</tr>
<tr>
<td></td>
<td>Lab II: phosphate, PTH, HCO3-, urinalysis, urine protein concentration. ANA, anti-ds-DNS-Ab, ANCA, Anti-GBM-Ab etc.</td>
</tr>
<tr>
<td></td>
<td>Tech: sonography of the kidneys, CT or MRI</td>
</tr>
<tr>
<td>Liver diseases with or without cholestasis</td>
<td>Lab I: y-GT, AP, bilirubin, AST, ALT, (and HB-, HC-antibodies, if a risk-patient)</td>
</tr>
<tr>
<td></td>
<td>Lab II: LDH, AMA, ANA, Anti-HBc-Ab, HBs-Ag, Anti-HCV-Ab, anti-smooth muscle Ab, antiaactin Ab</td>
</tr>
<tr>
<td></td>
<td>Tech: sonography of the liver, CT or MRT, (Magnetic resonance cholangiogram (MRC) or endoscopic retrograde cholangiogram (ERC) to rule out primary sclerosing cholangitis)</td>
</tr>
<tr>
<td><strong>Hyperparathyroidism</strong></td>
<td>Lab I: PTH, Calcium (only, if symptoms or signs of hyperparathyroidism (“stones, bones, moans and abdominal groans and psychiatric overtones”))</td>
</tr>
<tr>
<td></td>
<td>Lab II: phosphate, Vit D (1,25-Vit D, 25 Vit-D)</td>
</tr>
<tr>
<td></td>
<td>Tech: sonography of the parathyroid glands, scintigraphy, MRI</td>
</tr>
<tr>
<td><strong>Hyper- and hypothyroidism</strong></td>
<td>Lab I: TSH,</td>
</tr>
<tr>
<td></td>
<td>Lab II: T3, T4, MAKs and TRAKs</td>
</tr>
<tr>
<td></td>
<td>Tech: sonography of the thyroid glands, Iodine-scintigraphy</td>
</tr>
<tr>
<td><strong>Anemia</strong></td>
<td>Lab I: complete blood count including MCV and MCHC, LDH</td>
</tr>
<tr>
<td></td>
<td>Lab II: ferritin, transferrin saturation (TSAT) – optionally:</td>
</tr>
<tr>
<td></td>
<td>Lab III: Bone marrow aspiration with iron staining</td>
</tr>
<tr>
<td><strong>Iron deficiency</strong></td>
<td>Lab I: ferritin</td>
</tr>
<tr>
<td></td>
<td>Lab II: transferrin saturation (TSAT)</td>
</tr>
<tr>
<td><strong>Malabsorption</strong></td>
<td>(Lab-tests only in case of a typical history (pancreas disease, intestinal resection) or symptoms like chronic diarrhea or steatorrhea and weight loss.)</td>
</tr>
<tr>
<td></td>
<td>Lab I: Serum protein, serum albumine, calcium, blood count, gliadin-antibody</td>
</tr>
<tr>
<td></td>
<td>Lab II: Vitamin A (hyperkeratosis by Vitamin A deficiency), Vitamin B12 (neuropathy by Vitamin B deficiency)</td>
</tr>
<tr>
<td></td>
<td>Tech: endoscopy with biopsy</td>
</tr>
<tr>
<td><strong>Other diseases</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Pruritus of the elderly</strong></td>
<td>Lab I: Lab screening: creatinine, ALT, AST, alkaline phosphatase, bilirubin, TSH, full blood count, + BUN, (+ estimated creatinine clearance)</td>
</tr>
<tr>
<td><strong>Infective diseases</strong></td>
<td>HIV:</td>
</tr>
<tr>
<td></td>
<td>HIV-antibodies, Westernblot</td>
</tr>
<tr>
<td></td>
<td>Parasitoses including Helminthosis, Giardia lambia (rare): stool culture and microscopic examination</td>
</tr>
<tr>
<td><strong>Haematological disorders</strong></td>
<td>Polycythemia vera:</td>
</tr>
<tr>
<td></td>
<td>Lab I: blood count, thrombocytes, sedimentation rate</td>
</tr>
<tr>
<td></td>
<td>Lab II: to rule out secondary erythrocytosis: O₂ saturation, erythropoietin (EPO) level (renal cell carcinoma or polycystic kidneys)</td>
</tr>
<tr>
<td></td>
<td>Lab III: bone marrow with chromosomal aberrations, Tech: sonography, CT or MRI of the spleen, Lymphoma:</td>
</tr>
<tr>
<td><strong>Neurological diseases</strong></td>
<td>Lab I: blood count, blood smear, thrombocytes, sedimentation rate,</td>
</tr>
<tr>
<td></td>
<td>Lab II: Bone marrow with chromosomal aberrations, Tech: sonography, CT or MRI of the abdomen, thorax and additional affected areas, (PET)</td>
</tr>
<tr>
<td><strong>Psychiatric or psychosomatic diseases</strong></td>
<td>Psychiatric and psychosomatic exploration, psychiatric short questionnaire for depressive and anxiety disorder</td>
</tr>
<tr>
<td><strong>Pregnancy with or without cholestasis</strong></td>
<td>Lab I: y-GT, AP, bilirubin, AST, ALT, bile acids</td>
</tr>
<tr>
<td></td>
<td>Lab II: Virus screen: hepatitis A, B, C, Epstein Barr and cytomegalovirus, a liver autoimmune screen for chronic active hepatitis and primary biliary cirrhosis (anti-smooth muscle and antimitochondrial antibodies) (67)</td>
</tr>
<tr>
<td></td>
<td>Tech: liver ultrasound</td>
</tr>
<tr>
<td><strong>Drug induced pruritus</strong></td>
<td>Lab I: y-GT, AP, bilirubin, AST, ALT, LDH</td>
</tr>
<tr>
<td></td>
<td>Skin biopsy in case of HES exposition (electron microscopy)</td>
</tr>
</tbody>
</table>
First, the patient should be informed about general pruritus-relieving measures (Table V). They include simple and helpful measures such as wet and cold wraps, application of lotio alba, etc. Application of short-time localised heat has shown promising itch-relieving results in case reports and an experimental study (81). Prior to further symptomatic therapy, the patient should be subject to a careful diagnostic evaluation and therapy given for any underlying disease (see Tables III, IV). If pruritus still persists, combined or consecutive step-by-step symptomatic treatment is necessary (Table VI). Pharmacologic interventions for specific pruritic diseases, e.g. urticaria should be performed according to the guideline of the specific disease and the field’s Cochrane Group (82, 83).

6.2. Causative therapy and aetiology specific treatment

CP can be addressed by treating the underlying disease. Therapeutic measures include specific treatments of underlying dermatoses, avoidance of contact allergens, discontinuation of implicated drugs, specific internal, neurological and psychiatric therapies, surgical treatment of an underlying tumour or transplantation of organs. Normally, there is sudden relief of pruritus when the underlying disease improves, e.g. when Hodgkin’s disease responds to chemotherapy or when a patient with PBC has been transplanted. For some underlying diseases, specific treatments have proven to be successful in relieving pruritus, even if the underlying disease is

Table V. General measures for treating chronic pruritus

<table>
<thead>
<tr>
<th>Avoidance of</th>
<th>Factors that foster dryness of the skin, e.g. dry climate, heat (e.g. sauna), alcoholic compresses, ice packs, frequent washing and bathing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contact with irritant substances</td>
<td>Contact with irritant substances (e.g. compresses with rivanol, chamomile, tea-tree oil)</td>
</tr>
<tr>
<td>Very hot and spicy food</td>
<td>Very hot and spicy food, large amounts of hot drinks and alcohol</td>
</tr>
<tr>
<td>Excitement, strain, negative stress</td>
<td>Excitement, strain, negative stress</td>
</tr>
<tr>
<td>In atopic patients</td>
<td>In atopic patients: avoidance of aerogen allergens (e.g. house dust and house dust mites) which may aggravate pruritus</td>
</tr>
<tr>
<td>Application of</td>
<td>Mild, non-alcaline soaps, moisturizing syndets and shower/bathing oils</td>
</tr>
<tr>
<td>L Hartwater, bathing time not exceeding 20 min</td>
<td>L Hartwater, bathing time not exceeding 20 min</td>
</tr>
<tr>
<td>Soft clothing permeable to air</td>
<td>Soft clothing permeable to air, e.g. cotton, silver-based textiles</td>
</tr>
<tr>
<td>Skin moisturizer on a daily basis</td>
<td>Skin moisturizer on a daily basis especially after showering and bathing</td>
</tr>
<tr>
<td>with symptomatic relief</td>
<td>Topicals with symptomatic relief especially for pruritus at night: creams/lotions/sprays with e.g. urea, campher, menthol, polidocanol, tannin preparations</td>
</tr>
<tr>
<td>Wet, cooling or fat-moist-wrappings</td>
<td>Wet, cooling or fat-moist-wrappings, wrappings with black tea, short and lukewarm showers</td>
</tr>
<tr>
<td>Relaxation techniques</td>
<td>Autogenic training, relaxation therapy, psychosocial education</td>
</tr>
<tr>
<td>Education</td>
<td>Coping with the vicious circle of itch–scratch–itch</td>
</tr>
<tr>
<td>Educational training programs</td>
<td>Educational training programs e.g. for children suffering from atopic dermatitis or chronic pruritus (71, 72, 254)</td>
</tr>
</tbody>
</table>

Fig. 1. Diagnostic algorithm.
not treated. Aetiology specific treatments act on a known or hypothetically assumed pathogenesis of pruritus in underlying diseases. For only a few of these treatments evidence of efficacy can be found in controlled studies. Treatments for CP in specific diseases are presented in Tables VII–XI. When deciding the choice of treatment, consideration should be given to the level of evidence, side-effects, practicability, costs, availability of a treatment and individual factors such as patient’s age.

6.3. Symptomatic therapy: topical

6.3.1. Local anaesthetics. Local anaesthetics act via different groups of skin receptors. They can be used for pain, dysaesthesia and pruritus. Benzocaine, lidocaine, pramoxine as well as a mixture of prilocaine and lidocaine are widely used topically, but have only a short-term effect. In experimental studies, the antipruritic effect of local anaesthetics is limited in diseased skin, e.g. AD (84, 85). Successful application in the treatment of local anaesthetics is presented in Tables VII–XI. When deciding the choice of treatment, consideration should be given to the level of evidence, side-effects, practicability, costs, availability of a treatment and individual factors such as patient’s age.

Table VII. Therapeutic options in chronic kidney disease-associated pruritus

<table>
<thead>
<tr>
<th>Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 1</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Step 2</td>
</tr>
<tr>
<td>Step 3</td>
</tr>
</tbody>
</table>

Concomitant treatment in every step
• Diagnostics and treatment of underlying disease
• General therapeutic measures (Table V)
• In sleep disorders: sedative H1-antihistamines, tranquilizers, tricyclical antidepressants or neuroleptics
• Psychosomatic care, behavioural therapy for scratch behaviour
• In erosive scratch lesions: disinfecting measures, topical corticosteroids

*There is no evidence for the following diagnoses: cholestatic pruritus, nephrogenic pruritus

Table VI. Stepwise symptomatic-therapeutic approach in chronic pruritus (> 6 weeks)

<table>
<thead>
<tr>
<th>Therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Step 1</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Step 2</td>
</tr>
<tr>
<td>Step 3</td>
</tr>
</tbody>
</table>

Concomitant treatment in every step
• Diagnostics and treatment of underlying disease
• General therapeutic measures (Table V)
• In sleep disorders: sedative H1-antihistamines, tranquilizers, tricyclical antidepressants or neuroleptics
• Psychosomatic care, behavioural therapy for scratch behaviour
• In erosive scratch lesions: disinfecting measures, topical corticosteroids

*There is no evidence for the following diagnoses: cholestatic pruritus, nephrogenic pruritus

Table VIII. Therapeutic options in hepatic and cholestatic pruritus

<table>
<thead>
<tr>
<th>Antipruritic effects confirmed in controlled studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Activated charcoal 6 g/day (41)</td>
</tr>
<tr>
<td>• Gabapentin 300 mg 3×/week postdischarge (170)</td>
</tr>
<tr>
<td>• Gamma-linolenic acid cream 3×/day (261)</td>
</tr>
<tr>
<td>• Capsaicin 3–5×/day (98, 99)</td>
</tr>
<tr>
<td>• UVB phototherapy (237)</td>
</tr>
<tr>
<td>• Acupuncture at the Quchi (LI11) acupoint (262)</td>
</tr>
<tr>
<td>• Nalfurafine intravenously postdischarge (25)</td>
</tr>
<tr>
<td>• Thalidomide 100 mg/day (211)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Antipruritic effects confirmed in case reports</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Cholestyramine (41)</td>
</tr>
<tr>
<td>• Tacrolimus ointment 2×/day (124, 125)</td>
</tr>
<tr>
<td>• Cream containing structured physiological lipids with endocannabinoids (110)</td>
</tr>
<tr>
<td>• Mirtazapine (179)</td>
</tr>
<tr>
<td>• Cromolyn sodium (151)</td>
</tr>
<tr>
<td>• Erythropoetin 36 IU/kg 3×/week (263)</td>
</tr>
<tr>
<td>• Lidocaine 200 mg i.v./day (41)</td>
</tr>
<tr>
<td>• Ketotifen 1–2 mg/day (150)</td>
</tr>
</tbody>
</table>

Table IX. Antipruritic therapy of atopic dermatitis

<table>
<thead>
<tr>
<th>Antipruritic effects confirmed in controlled studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Glucocorticosteroids (topical and oral)</td>
</tr>
<tr>
<td>• Cyclosporin A</td>
</tr>
<tr>
<td>• Leukotriene antagonists (e.g. zafirlukast)</td>
</tr>
<tr>
<td>• Interferon-gamma, i.c.</td>
</tr>
<tr>
<td>• Antihistamines (topical and systemic)</td>
</tr>
<tr>
<td>• Nalfurafine 30 mg i.v. 1×/day (272)</td>
</tr>
<tr>
<td>• Gabapentin 300 mg i.v. 2×/day (268)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Antipruritic effects confirmed in case reports</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Macrolide antibiotics</td>
</tr>
<tr>
<td>• Immunoglobuline, i.v.</td>
</tr>
<tr>
<td>• UV A1-/UV B 311-Therapie</td>
</tr>
</tbody>
</table>

*We refer to the current guideline for atopic dermatitis and ref. 280.
Table X. Therapeutic options in polycythaemia vera

<table>
<thead>
<tr>
<th>Effects confirmed in case reports</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Paroxetine 20 mg/day (42, 181)</td>
</tr>
<tr>
<td>• Hydroxyzine (42)</td>
</tr>
<tr>
<td>• Fluoxetine 10 mg/day (181)</td>
</tr>
<tr>
<td>• Aspirin (282)</td>
</tr>
<tr>
<td>• Cimetidine 900 mg/day (283, 284)</td>
</tr>
<tr>
<td>• Pizotifen 0.5 mg 3×/day (285)</td>
</tr>
<tr>
<td>• Cholesteramine (286)</td>
</tr>
<tr>
<td>• Ultraviolet B phototherapy (241)</td>
</tr>
<tr>
<td>• Photochemotherapy (PUVA) (287, 288)</td>
</tr>
<tr>
<td>• Transcutaneous electrical nerve stimulation (289)</td>
</tr>
<tr>
<td>• Interferon-alpha (290–293)</td>
</tr>
</tbody>
</table>

Table XI. Therapeutic options in aquagenic pruritus

<table>
<thead>
<tr>
<th>Effects confirmed in case reports (294–296)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Topical capsaicin 0.025–1% twice/d for 4 weeks</td>
</tr>
<tr>
<td>• Glyceryl trinitrate topically 2%</td>
</tr>
<tr>
<td>• Transdermal application of scopulinam, topically 3 or 9%</td>
</tr>
<tr>
<td>• Baths with sodium bicarbonate (0.5–1 kg/bath)</td>
</tr>
<tr>
<td>• Bath and systemic PUVA, UVB (242–245)</td>
</tr>
<tr>
<td>• Propranolol 10 to 80 mg/day</td>
</tr>
<tr>
<td>• Clonidine 0.1 mg twice/day</td>
</tr>
<tr>
<td>• Astemizol 10 mg/day</td>
</tr>
<tr>
<td>• Ibuprofen (prior to bathing)</td>
</tr>
<tr>
<td>• Pregabalin 150–300 mg/day</td>
</tr>
<tr>
<td>• Antihistamines, e. g. hydroxyzine 25 mg/d, chlorpheneramine 8 mg/day, cetirizine, loratadine, fexofenadine, terfenadine</td>
</tr>
<tr>
<td>• H2-blockers: cimetidine 900 mg/day</td>
</tr>
<tr>
<td>• Opioid receptor antagonists, e. g. naltrexone 25–50 mg/day</td>
</tr>
<tr>
<td>• Selective serotonin reuptake inhibitors, e. g. paroxetine 20 mg/day, fluoxetine 10 mg/day</td>
</tr>
<tr>
<td>• Interferon-alpha 2b 5×3 mil IE 1st week, 3×3 mil IE 2nd – 4th week</td>
</tr>
</tbody>
</table>

Effects confirmed in RCT
| Acetyl salicylic acid 300–500 mg/day |

6.3.2. Glucocorticosteroids. Pruritus experimentally induced by histamine was significantly suppressed by topical hydrocortisone when compared to placebo (87). All other clinical studies apply to an underlying inflammatory dermatosis in which “pruritus” was one parameter amongst many. Clinical experience shows that topical glucocorticosteroids can be effective if itch is the consequence of an inflammatory dermatosis. Use of topical glucocorticosteroids to treat the symptom of pruritus is not advised in the absence of an inflammatory dermatosis. Topical glucocorticosteroids with a favourable side-effect profile (e.g. fluticasonepropionate, methylprednisolon-aceponate or mometasonfuorate) are to be preferred (88, 89). In some cases the anti-inflammatory effect of glucocorticosteroids is helpful, but insufficient to completely abolish pruritus (90).

**Expert recommendation**: Initial short-term application of topical glucocorticosteroids can be recommended in CP associated with an inflammatory dermatosis, but should not be used as long-term treatment or in the absence of a primary rash.

6.3.3. Capsaicin. Capsaicin (trans-8-metyl-N-vanillyl-6-nonenamide) is the pungent agent of chilli peppers and is used as a pain-relieving medication (91). Topical application of capsaicin activates sensory C-fibres to release neurotransmitters inducing dose-dependent erythema and burning. After repeated applications of capsaicin, the burning fades due to tachyphylaxis and retraction of epidermal nerve fibres (91). However, pruritus reoccurs some weeks after discontinuation of therapy indicating no permanent degeneration of the nerve fibres (92).

The greater the initial dose of capsaicin and the more frequent the applications, the sooner the desensitization will appear and pruritus disappear. Severe initial burning may be a side-effect of topical application. Cooling of the skin can also reduce the capsaicin-evoked burning. More unusual adverse effects of capsaicin include cough or sneezing due to inhalation of capsaicin from the skin or from the jar and its effect on sensory nerve fibres in the mucous membranes (91). It appears that such adverse effects are less bothersome for patients with severe pruritus compared to patients with slight pruritus (unpublished observations). A lower concentration of capsaicin and less frequent applications will induce tachyphylaxis later but may give a better compliance. The concentration of capsaicin varies in different studies, but 0.025% capsaicin is tolerated well by most patients. If capsaicin is not available in this concentration as a standard drug it can be produced using a lipophilic vehicle. Capsaicin is also well soluble in alcohol; capsaicin 0.025% in *spir dil* can be used to treat itchy scalp (not published). A weaker concentration of 0.006% capsaicin is recommended for intertriginous skin e.g. pruritus ani (93).

Topical capsaicin’s effects have been confirmed in controlled clinical trials for different pain syndromes and neuropathy as well as notalgia paraesthesia (94), brachioradial pruritus (95), pruritic psoriasis (96, 97) and haemodialysis-related pruritus (98, 99). Case reports and case series described effects in hydroxyethyl starch-induced pruritus (100, 101), prurigo nodularis (100, 102–104), lichen simplex (100, 103), nummular eczema (100), aquagenic pruritus (105) and PUVA-associated pruritus (106).

**Expert recommendation**: Capsaicin can be effective in localised forms of CP, but patient compliance due to side-effects can restrict usage.
6.3.4. Cannabinoid receptor agonists. Topical cannabinoid receptor agonists are a new development since 2003 and appear to have antipruritic and analgesic properties. Experimentally induced pain, pruritus and erythema could be reduced by application of a topical cannabinoid agonist (107, 108). One cosmetic product containing the cannabinoid receptor and peroxisome proliferator-activated receptor alpha (PPAR-α) agonist, N-palmitoylethanolamine, is currently on the market. In (non-vehicle controlled) clinical trials and case series, it proved to have antipruritic effects in prurigo, AD, CKD-associated pruritus and PUO (109–111) as well as analgetic effects in postzosteric neuralgia (112).

Expert recommendation: Cannabinoid receptor agonists can be effective in the treatment of localised pruritus.

6.3.5. Tacrolimus and pimecrolimus. The effects of tacrolimus and pimecrolimus on pruritus are mediated both through their immunological and neuronal properties (113). Paradoxically, while they can induce transient pruritus at the beginning of treatment, in the medium-term they may provide an alternative treatment for many causes of pruritus. They are very effective against pruritus in AD (114). Furthermore, tacrolimus ointment is more effective at reducing pruritus when compared with vehicle and pimecrolimus cream (114). Clinical trials have shown benefit of both pimecrolimus and tacrolimus in seborrhoeic dermatitis, genital lichen sclerosis, intertriginous psoriasis and cutaneous lupus erythematosus and – only for tacrolimus – in resistant idiopathic pruritus ani (115–122). In other diseases, the available data are limited to small case series, or individual cases e.g. hand eczema (pimecrolimus), rosacea (tacrolimus), graft-versus-host disease (tacrolimus), vulval pruritus (tacrolimus) or Netherton’s syndrome (tacrolimus, pimecrolimus). Topical tacrolimus has been shown anecdotally to be effective in pruritus associated with systemic diseases such as PBC (123) and chronic renal insufficiency (124, 125).

However, these observations have not been confirmed in a controlled study on CKD-associated pruritus (126, 127). Both substances can be used to treat localised forms of CP such as genital pruritus (128).

Expert recommendation: Tacrolimus and pimecrolimus are effective in localised forms of CP.

6.3.6 Acetylsalicylic acid. Topical acetylsalicylic acid (acetylsalicylic acid/dichloromethane solution) has been described to have antipruritic effects in occasional patients with lichen simplex (129). However, this beneficial effect could not be confirmed in experimentally induced itch with histamine (130).

Expert recommendation: Due to the lack of studies, topical acetylsalicylic acid can currently not be recommended for CP.

6.3.7. Doxepin. The tricyclic antidepressant doxepin showed antipruritic effects when applied as a 5% cream in double-blind studies for treatment of AD (131), lichen simplex, nummular dermatitis and contact dermatitis (132). Topical doxepin therapy is not licensed and not used in any European country except for the UK (Xepin©) (133–135).

Expert recommendation: Due to the increased risk of contact allergy, especially when the treatment exceeds 8 days, topical doxepin cannot be recommended.

6.3.8. Zinc, menthol and camphor. Although zinc oxide has been used in dermatology for over 100 years due to its anti-inflammatory, antiseptic and anti-pruritic properties and its safety, there is only scarce literature on its effects. Prescriptions of zinc are frequent, with concentrations varying from 10 to 50% in creams, lotions, lotions, ointments and pastes that are useful in the treatment of pruritus, especially for localised forms of pruritus, in children as well as in adults (136).

Menthol is an alcohol obtained from mint oils, or prepared synthetically. Applied to the skin and mucous membranes, menthol dilates blood vessels, causing a sensation of coldness, followed by an analgesic effect (136). Menthol is used in dusting powders, liniments, lotions and ointments in concentrations from 1–10% (136). Menthol binds to the TRPM8 receptor (137) that belongs to the same TRP family of excitatory ion channels as TRPV1, the capsaicin receptor. These two receptors have been shown to co-exist occasionally in the same primary afferent neurons and promote thermosensations at a wide range of temperatures: 8–28°C and >50°C, respectively (137). Short-term application of such medications in CP in combination with other topical or systemic therapies can be recommended.

Camphor is an essential oil containing terpenes, it is soluble in alcohol (136). Applied to the skin it causes a sensation of warmth that is followed by a mild degree of anaesthesia (136). Camphor has been used in dermatology for decades in liniments, lotions and ointments in concentrations from 2–20%. It has been shown to specifically activate another constituent of the TRP ion channel family, namely TRPV3 (138). Recently, camphor was demonstrated to activate capsaicin receptor, TRPV1, while menthol also activates the camphor receptor, TRPV3. These findings illustrate the complexity of sensory perception and explain the efficacy of ointments containing both menthol and camphor (136).

Expert recommendation: Short term application of camphor, menthol and zinc in CP in combination with other topical or systemic therapies can be recommended.

6.3.9. Mast cell inhibitors. In a multi-center, double-blind, placebo-controlled trial, application of a 3% hydrogel formulation of tiacrilast against vehicle in AD led to no significant improvement of pruritus (139). Pruritus in AD responds to topical sodium cromoglycate (140), that was proved by a recent placebo-controlled study (141).

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**Expert recommendation**: There is limited evidence to recommend the use of topical mast cell inhibitors for CP.

### 6.4. Systemic therapy

#### 6.4.1. Antihistamines

Antihistamines are the most widely used systemic antipruritic drugs in dermatology. Most antihistamines that have been tried in pruritus belong to the H1 type. First generation antihistamines, such as chlorpheniramine, clemastine, cyproheptadine, diphenhydramine, hydroxyzine, and promethazine are known to bind not only to H1-receptors but also to muscarinic, α-adrenergic, dopamine or serotonin receptors and have a central sedative effect. Due to side effects, the application of sedative antihistamines is nowadays limited. Second generation antihistamines like cetirizine, levocetirizine, loratadine, desloratadine, ebastine, fexofenadine and rupafine have minimal activity on non-histaminic receptors, little sedative effect, and a longer duration of action compared to the first generation (142). Non-sedative H1-receptor antagonists offer an effective reduction of pruritus in diseases associated with increased mast cell degranulation like urticaria or mastocytosis (142). However, the doses required to alleviating pruritus in urticaria often amount to up to four times the licensed dose (143). Higher doses of the second generation antihistamines enhance their soporific side effects (142), which may contribute to their efficacy. A recent case series suggest that updosing of antihistamines may also be beneficial in CP (144).

Systemic H1-antihistamines are often employed to combat itch in AD, but only sedative antihistamines have shown some benefit, mainly by improving sleep (145). Hydroxyzine is the most commonly used antihistaminic of the first generation showing sedative, anxiolytic and antipruritic activities. In adult patients it is recommended as an antipruritic agent in the dosage 75–100 mg/day. In children the effective dose is 1–2.5 mg/kg/day. In a controlled study, addition of hydroxyzine resulted in a 750-fold increase in the dose of histamine required to elicit itch. There was a five-fold increase following both cyproheptadine and placebo and a ten-fold increase following diphenhydramine (146). In addition, hydroxyzine was significantly more effective in reducing histamine-induced pruritus than neuroleptics, like thiopixene, chlorpromazine and thoridazine (147).

In addition, antihistamines are widely used as first-line drugs for treatment of CP associated with different systemic diseases such as chronic renal failure, cholestasis, hematopoetic diseases and thyroid disorders. However, conventional doses of antihistamines in the treatment of pruritus in internal diseases have not proven to be effective (142).

Although identified in human skin, H2-receptors play a minor role in pruritus and H2-receptor antagonists alone have no antipruritic effect (145, 148). A combination of H2-antihistamines and H1-antihistamines has been used in treatment of pruritus in small trials but the results are conflicting (145, 148). A combination of H1-antihistamine with a leukotriene antagonist has been reported to alleviate pruritus in chronic urticaria (149). **Expert recommendation**: Antihistamines are effective in treating CP in urticaria. Antihistamines are of some value for itch in AD and CP of diverse origin. As there is limited evidence of antipruritic effects of non-sedating antihistamines in AD, PV and CP of diverse origin, sedating antihistamines can be recommended to be applied during night time for sleep improvement. Hydroxyzine is the first choice of the majority of physicians trying to control CP but its sedative effect may contraindicate its use in the elderly.

#### 6.4.2. Mast cell inhibitors

Ketotifen, a mast cell stabilizer, showed antipruritic effects in single patients with CKD-associated pruritus (150). Two patients with CKD-associated pruritus (151) and Hodgkin’s lymphoma (152) showed a significant antipruritic effect with the mast cell stabilizer cromoglicic acid. **Expert recommendation**: There is insufficient evidence to recommend the systemic use of mast cell inhibitors for CP.

#### 6.4.3. Glucocorticosteroids

There are no studies investigating the efficacy of the exclusive use of systemic glucocorticosteroids in CP. In clinical experience, pruritus ceases within approximately 30 min of i.v. glucocorticosteroids in the treatment of urticaria or drug-induced exanthema. Likewise, in AD, allergic contact dermatitis, dyshidrosis and bullous pemphigoid, rapid reduction of pruritus is observed, which can be explained by the high anti-inflammatory potency of glucocorticosteroids. Thus, while systemic glucocorticosteroids should not be considered as an antipruritic drug for long-term therapy, short-term use is possible in cases of severe pruritus, but should not be prescribed for a period of more than two weeks (153) because of severe side-effects.

Prednisone is the most commonly selected oral corticosteroid initially at a daily dose that can range from 2.5–100 mg daily or more, usually starting in a dose of 30–40 mg daily. In exceptional cases, i.v. methylprednisolone is used at a dose of 500 mg/day to 1 g/day, because of its high potency and low sodium-retaining activity. It is important to remember that the dosage should be tapered in accordance with the severity of pruritus. Before discontinuing systemic therapy one may change to topical corticosteroid therapy. Corticosteroids should be used with caution in children and the elderly as well as in patients with relevant metabolic disorders such as diabetes. **Expert recommendation**: Systemic corticosteroids can be used as short-term treatment in severe cases of CP, but should not be used for longer than 2 weeks.
6.4.4. Opioid receptor agonists and antagonists. Experimental and clinical observations have demonstrated that pruritus can be evoked or intensified by endogenous or exogenous μ-opioids (154). This phenomenon can be explained by activation of spinal opioid receptors, mainly μ-opioid receptors. Reversing this effect with μ-opioid antagonists thus leads to an inhibition of pruritus (112). The opposite is true for κ-opioids. Their binding to μ-opioid receptors leads to inhibition of pruritus (155).

Several clinical studies have demonstrated that different μ-opioid receptor antagonists may significantly diminish pruritus (30, 33, 156–160). In double-blind RCT, μ-opioid receptor antagonists such as naloxone, naltrexone and naltrexone have exhibited high antipruritic potency. For example, pruritus in chronic urticaria, AD, and cholestatic pruritus has shown therapeutic response to nalmefine (10 mg twice daily) and naltrexone (50–100 mg/day) (161, 162). Controlled studies have also been performed in patients with CKD-associated pruritus (26, 27, 163). Results were variable from significant reduction of pruritus to no response. Case reports have demonstrated efficacy in prurigo nodularis, macular amyloidosis, lichen amyloidosis, pruritus in mycosis fungoides, psoriasis vulgaris, aquagenic pruritus, hydroxylethyl starch-induced pruritus and PUO.

Nalfurafine, a preferential μ-opioid receptor agonist, was investigated in CKD-associated CP in two large RTCs (25, 164). Both trials demonstrated significant clinical benefit of nalfurafine in patients with uremic pruritus (155) within the first seven days of treatment. The drug is currently licensed in Japan only. **Expert recommendation:** Opioid receptor antagonists may be effective in cholestatic pruritus and AD but their side-effect profile needs to be considered. Nalfurafine can be applied in Japanese patients with uremic pruritus.

6.4.5. Gabapentin and pregabalin. Gabapentin is an antiepileptic drug, also used in neuropathic disorders causing pain or pruritus (165). The mechanisms of action of gabapentin, a 1-amino-methyl-cyclo-hexane acetic acid and a structural analogue of the inhibitory neurotransmitter γ-aminobutyric acid (GABA) remain unclear. It is used in postherpetic neuralgia (166), especially with paroxysmal pain or pruritus. Anecdotal indications are brachioradial pruritus (167) and cutaneous T-cell lymphoma (168). Pilot studies have been performed for the treatment of pruritus caused by burns and wound healing in children demonstrating antipruritic effects of gabapentin (169). Double-blind, randomised, placebo-controlled trials were performed for CKD-associated pruritus (170) and cholestatic pruritus (171). Gabapentin was safe and effective for treating CKD-associated pruritus (172, 173). Pregabalin is similar to gabapentin and a more recent drug. Its use has been suggested in a case of cetuximab-related pruritus and aquagenic pruritus (174, 175). A recent controlled trial demonstrated a significant antipruritic effect of pregabalin in patients on haemodialysis within one month (176). **Expert recommendation:** Gabapentin and pregabalin can be recommended in the treatment of CKD-associated pruritus and neuropathic CP.

6.4.6. Antidepressants. Psychoemotional factors are known to modulate the ‘itch threshold.’ Under certain circumstances, they can trigger or enhance CP (177). Itch is a strong stressor and can elicit psychiatric disease and psychological distress. Depressive disorders are present in about 10% of patients with CP (77). Consequently, depressive symptoms are treated in these patients, and some antidepressants also exert an effect on pruritus through their pharmacological action on serotonin and histamine. SSRIs, such as paroxetine, can have an antipruritic effect on patients with PV, psychogenic or paraneoplastic pruritus and other patients with chronic PUO (178). Antidepressants, like mirtazapine (179) and especially doxepin (180) have been effective in urticaria, AD and HIV-related pruritus.

The SSRI paroxetine (20 mg/day) has exhibited antipruritic effects in pruritus due to PV (181), paraneoplastic pruritus (182, 183) and psychiatric disease (184). In two patients, pruritus was induced by discontinuation of paroxetine treatment for depression (185). A RCT in pruritus of non-dermatologic origin confirmed the antipruritic effect of paroxetine (178). In a two-armed proof-of-concept study with paroxetine and fluvoxamine, patients with CP of dermatological origin reported significant antipruritic effect (186). Sertraline proved efficacy in cholestatic pruritus as demonstrated in a RCT (187). As severe cardiac side effects have been described, especially in elderly, this therapy should be used with caution. A psychosomatic/psychiatric examination before starting the treatment is recommended because of its stimulative effects. **Expert recommendation:** SSRIs can be recommended for the treatment of somatoform pruritus, paraneoplastic CP, PUO and cholestatic pruritus. Mirtazapine can be recommended in CP of AD.

6.4.7. Serotonin receptor antagonists. Due to the pathophysiological significance of serotonin in different diseases such as kidney and liver diseases, serotonin receptor antagonists (of the 5-HT3 type) such as ondansetron (8 mg 1–3/day), topisetron (5 mg/day) and granisetron (1 mg/day) have been used anecdotally to treat pruritus (188–194). Contradictory or negative results have been reported in partly controlled studies using ondansetron for cholestatic pruritus (188, 195, 196) and opioid-induced pruritus (197–199). An antipruritic effect was reported for ondansetron in CKD-associated pruritus (200). However, this could not be confirmed in subsequent controlled studies (201–203) later on.
**Expert recommendation**: Due to the lack of convincing evidence, serotonin receptor antagonists cannot be recommended in the treatment of CP.

6.4.8. Thalidomide. A number of mechanisms for the antipruritic action of thalidomide have been proposed including a central depressant effect (204), a local effect on proliferated neural tissue in PN (205), and the antagonism of TNF-α (206).

The best results with thalidomide in CP have been achieved in PN. Several studies have shown a rapid decrease of pruritus on thalidomide (50–300 mg/day) (207, 208). A prospective open trial of thalidomide 100 mg/day, followed by narrow-band UVb (TL-01) showed a high response with minimal side-effects (209). Likewise, good results have been seen in HIV-positive patients with PN (210). There is one randomised double-blind crossover trial of the successful treatment of CKD-associated pruritus with thalidomide (211). Thalidomide is teratogenic and there is a dose-related risk of neuropathy, especially in high daily doses (>100 mg/day) (212).

**Expert recommendation**: Though there is evidence for its antipruritic effect, thalidomide is not recommended for the treatment of CP due to its side effects.

6.4.9. Leukotriene receptor antagonist, TNF antagonists.

Leukotriene receptor antagonists (e.g. montelukast) and TNF-α antagonists influence the pathogenesis of AD. They have been used in combination with antihistamines as antipruritic therapy. Montelukast has also been used in several types of urticaria as well as in combination with antihistamines. A combination of H1-antihistamine with a leukotriene antagonist has been reported to alleviate pruritus in chronic urticaria (204).

**Expert recommendation**: Due to the lack of evidence, leukotriene receptor antagonists and TNF antagonists cannot be recommended in the treatment of CP.

6.4.10. Cyclosporin A. Pruritus in AD responds to treatment with cyclosporin A as demonstrated in controlled double-blind studies (213, 214). Cyclosporin A has been administered in PN for 24 to 36 weeks, using doses of 3.0–4.5 mg/kg/day. Improvement was observed in both pruritus and skin lesions after two weeks of treatment (215, 216). It seems likely that in these diseases cyclosporin A acts on pruritus through its immunological effects. However, direct effects on nerve endings are also possible, suggested by successful use in non-immunological diseases as reported in several studies, e.g. 10 patients with senescent pruritus were treated with cyclosporin A 5 mg/kg/day for 8 weeks (217). All patients of this uncontrolled, open study responded. Case reports describe antipruritic effects in dystrophic epidermolysis bullosa associated CP and in CKD-associated pruritus (218, 219).

**Expert recommendation**: Cyclosporin A can be recommended in the treatment of CP in AD or in PN.

6.4.11. Aprepitant. Substance P (SP) has a dominant role in pruritus induction in the skin. Via binding to the neurokinin 1 receptor (NKR1) on keratinocytes, blood vessels and mast cells, SP promotes inflammation and mast cell degranulation. SP is released from sensory neurons. In conditions with hyperplasia of skin nerves (AD, PN), SP levels are increased. Accordingly, inhibition of the pruritogenic effects of SP by blocking the corresponding receptor may have antipruritic effects. Several case reports suggest a positive role of the NKR1 receptor antagonist aprepitant in CP, e.g. cutaneous T-cell lymphoma, solid tumours and drug-induced pruritus (220–223). Recently, a proof-of-concept study in 20 patients showed significant, antipruritic effects in chronic, therapy-refractory pruritus of various origins with a one-week monotherapy of aprepitant (224). The highest response rate was observed in patients with atopic diathesis and PN. RCT are missing.

**Expert recommendation**: NKR1 antagonists, in particular aprepitant, are promising substances in the therapy of CP. Aprepitant might be used as a second-line option in therapy refractory cases, e.g. in patients with AD and PN.

6.5. UV phototherapy

Ultraviolet (UV)-based therapy is well established for treating pruritus and utilizes UVB (290–320 nm) and UVA (320–400 nm). The light sources include broadband UVB (BB-UVB, 290–320 nm, peaks at 313 nm), narrowband UVB (NB-UVB, 311 nm), broadband UVA (320–400 nm, peaks at 355 nm), and UVA1 (340–400 nm, peaks at 365 nm) (225).

Inflammatory dermatoses associated with pruritus respond well to different UV treatments including UVA1 311. For the treatment of AD, early studies demonstrated that UVB was better than placebo (226). In a recent study NB-UVB was better than BB-UVB and both were better than placebo (227). In the treatment of pruritus of AD, BB-UVB and UVA were equally effective in a half-body comparison (228). In a more recent study, NB-UVB was insignificantly better than UVA1 for pruritus (229). In AD, phototherapy seems to act locally rather than systemically: When one half of the body was treated with UVB and the other half was not, only the treated side improved (226).

For the treatment of prurigo PUVA, UVA1 and NB-UVB proved to be effective in a RCT, with PUVA and UVA1 superior to NB-UVB (230).

For many other skin diseases, a number of studies have demonstrated the efficacy of UV treatment, e.g. psoriasis, lichen planus, T-cell lymphoma, solar, chronic, and idiopathic urticaria, and urticaria pigmentosa.

It can be assumed that in cases of pruritic inflammatory dermatoses pruritus is reduced by inhibiting pro-inflammatory mediators and induction of anti-
inflammatory and immunosuppressive factors. UVB mainly affects epidermal keratinocytes and Langerhans’ cells, due to its limited penetration into the skin. UVA1, in contrast, reaches to the dermis and therefore can affect T lymphocytes, mast cells, and dermal dendritic cells, e.g. induces apoptosis of these cells (225). However, UVB-induced apoptosis of mast cells has been argued to explain relief of pruritus (231). Furthermore, phototherapy leads to a reduction of CGRP-immunoreactive nerve fibres in the skin (232).

In conditions with pruritus on primarily non-inflamed skin, UV therapy has been particularly effective in CKD-associated pruritus (233, 234). In a placebo-controlled trial, UVA alone was ineffective for this condition (235). However, an antipruritic effect was seen in CKD-associated pruritus when treated with combined UVA/UVB phototherapy (236). BB-UVB alone was effective in treating CKD-associated pruritus. It was remarkable that in spite of placebo control (only one body half was treated) an improvement of pruritus occurred over the entire body (237), suggesting a systemic antipruritic effect. In an open pilot study using NB-UVB 14/20, CKD-associated pruritus patients responded well to treatment (238). Also in a recent study NB-UVB appeared to be effective in reduction of CKD-associated pruritus (239). However in another case NB-UVB treatment was unsuccessful, but BB-UVB helped (240).

UV therapy has also been reported to be effective in a number of cases of metabolic itch. In PV, 8/10 patients responded to NB-UVB in an open study (241). Aquagenic pruritus has shown response to bath PUVA therapy (242) and systemic PUVA (243) for the duration of therapy. To treat aquagenic pruritus, PUVA was found to be superior to BB-UVB in 5 patients (244). Recently, two patients with aquagenic pruritus have been reported with a good, but ephemeral response to NB-UVB (245).

In HIV patients with pruritus, UVB produced significant relief of pruritus in an open study with 21 patients (33% primary pruritus, 66% eosinophilic folliculitis) (246). In a single case report, a patient with Hodgkin’s disease responded well to BB-UVB (247).

A retrospective analysis of children up to the age of 18 years suffering from AD and psoriasis suggests NB-UVB treatment (248). In children, longer follow-up is essential to determine true carcinogenic risk of UV therapy.

**Expert recommendation:** UV therapy can be applied for CP. The mode of UV phototherapy depends on the underlying disease. UVA as well as UVB (NB-UVB/BB-UVB) as well as a combination of UVA/UVB relieve CP in certain diseases. UV phototherapy can be used in combination with topical and/or systemic treatment except for calcineurin inhibitors and immunosuppressant drugs.

### 6.6. Psychosomatic therapy (relaxation techniques and psychotherapy)

The vicious itch-scratch cycle has to be taken into account when a patient is treated for pruritus. In addition to causal and symptomatic therapy, behavioural therapy to avoid scratching should be considered, e.g. conscious suppression of the reflex by intense concentration, distraction or alternative scratching techniques such as habit reversal (249). This is very important in patients with prurigo nodularis who might show an unconscious automatic scratching behaviour.

Adjuvant psychosocial programmes are most effective in AD (72, 250–252). Such programmes include strategies for breaking the vicious circle of itching and scratching, relaxation and stress management techniques as well as strategies for dealing with relapses. A similar educational programme was developed for patients with CP (253, 254). It is currently established for in-patient hospital treatment of patients with pruritic dermatoses using behavioural therapy in the context of an integrated psychosomatic treatment (255, 256). In patients with coexisting depression, psychotherapy in combination with psychotropic medication can be helpful even to treat pruritus of different aetiology (257). Most publications on psychotherapeutic/psychopharmacologic interventions, however, refer to small groups or single case reports. In neurotic excoriations, combined psychopharmacotherapy is also often indicated (257–260).

**Expert recommendation:** Relaxation techniques and education programmes for CP patients are useful as a complementary treatment for managing CP.

### 7. KEY SUMMARY OF DISCUSSION CONCERNING COUNTRY-SPECIFIC PROCEDURES

- **Antihistamines:** Sedative H1 antihistamines are first-choice therapy in CP to improve night-time sleep. Studies on application of higher doses are yet to be conducted.
- **UV phototherapy** is recommended for generalised pruritus, especially in elderly pruritus patients or in case of contraindications for systemic therapy.
- **Anticonvulsants/pain modulators** are recommended in neuropathic pruritus.
- **Antidepressants** are recommended in forms of CP not responding to other therapies.
- **Systemic glucocorticosteroids** are not recommended for treatment of CP except of very severe and desperate cases.
- **Serotonin receptor antagonists and thalidomide** are not recommended for treatment.

This guideline is in accordance with the EDF guideline on chronic pruritus finished in 2010 (www.euroderm.org). This is an updated version.
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European Guideline on Chronic Pruritus

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