Chronic prostatitis: current treatment options

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Abstract: Male chronic prostatitis/chronic pelvic pain syndrome (CP/CPPS) is represented by a heterogeneous group of symptoms that can cause an important impairment of daily quality of life for patients. Diagnosis of CP/CPPS is often not clear and treatment can be challenging, as it varies according to the different causative factors and derived symptoms. Differently from approaches used in the past, the diagnosis and subsequent treatment rely on separating this entity from chronic bacterial prostatitis and considering it as a multifactorial disease. Autoimmunity and inflammation, myofascial tenderness, neuroinflammation, and psychological causes have been clearly related to this disease, and therefore CPPS should not only be considered as related to benign prostatic enlargement. A multitude of different symptoms related to urinary, genital, rectal, and perineal areas can be attributed to this condition and therefore should be routinely investigated in patients, as well as possible differential diagnoses which can cause the same symptoms, such as pudendal nerve entrapment syndrome. The aim of this narrative review is to focus on CPPS after an infectious cause has been excluded.

Keywords: chronic prostatitis, chronic pelvic pain syndrome, pudendal neuralgia, physical therapy, pharmacological treatment

Introduction

The term “chronic prostatitis” is used to define and include many different symptomatological patterns, and its understanding is still enigmatic for many physicians and patients. Overall, it is estimated that the prevalence of chronic prostatitis among the male population is about 4.5–9%, with recurrence rates increasing up to 50% with increasing age.\(^1\)\(^-\)\(^3\) Therefore, it has a similar prevalence to that of ischemic heart disease and higher than diabetes.\(^4\) Although idiopathic urogenital and anorectal pain syndromes are not uncommon, effective treatments remain elusive for this patient group. Pain and functional disorders in these parts of the body can be embarrassing, limiting the desire to discuss the symptoms with the physician; similarly, clinicians may not be familiar enough with these syndromes, leading to misdiagnosis. Moreover, most of the involved patients usually complain of many different symptoms, not limited to “pure” prostatodynia but also presenting with: lower urinary tract symptoms (LUTS) with pollakiuria, dysuria, nocturia, urinary dribbling, or weak urinary stream; symptoms related to the anorectal area, such as constipation, sensation of foreign body in the rectus, and rectal pain during and after defecation; symptoms related to the external genitalia, represented by genital pain or burning and premature ejaculation, spontaneous sexual stimulation, or alteration of orgasms;\(^5\)\(^,\)\(^6\) patients can also refer to an associated low back pain, worsened in the sitting position. These symptoms can appear simultaneously or progressively, generating an increasing sensation of discomfort and anxiety for the
patients, who often do not feel completely understood by their referring physician, with a progressive impairment of their quality of life (QoL).

Following the definition provided by the US National Institutes of Health (NIH) classification, chronic prostatitis is divided into different categories: a combination of chronic bacterial prostatitis (CBP), chronic pelvic pain syndrome (CPPS), or asymptomatic prostatitis. The overall NIH classification of prostatitis syndromes includes:

- **Category I:** Acute bacterial prostatitis (ABP), due to acute bacterial infection determining prostatitis symptoms, systemic infection, and acute bacterial UTI.
- **Category II:** Chronic bacterial prostatitis (CBP), with a demonstrated chronic bacterial prostatic infection with or without prostatitis symptoms.
- **Category III:** Chronic prostatitis/chronic pelvic pain syndrome (CP/CPPS), where an infective agent is absent, and the disease is led by chronic pelvic pain symptoms and voiding symptoms in the absence of UTI. Those patients present with urological pain or discomfort in the pelvic region, associated with urinary symptoms and/or sexual dysfunction, lasting for at least 3 of the previous 6 months. This category is further divided into IIIA, inflammatory, and IIIB, non-inflammatory.
- **Category IV:** Asymptomatic inflammatory prostatitis (AIP) due to prostate inflammation in the absence of genitourinary tract symptoms, always associated with CPPS.

Categories III and IV prostatitis cover a wide range of multifaceted chronic pain syndromes with varied clinical patterns, diagnostic pathways, and treatments. This field of research has rapidly evolved during the past few years, aiming at a wider understanding of the involved pathophysiological patterns. The physician should not only be focused on “usual” urological symptoms and complaints but also investigate all possible related symptoms in the anorectal, urinary, and genital area, as well as neurological, immunological, and psychological factors that may be related and influence the disease process.

Moreover, symptoms may also be due to other pathophysiology not related to prostatic inflammation, such as pudendal neuralgia due to pudendal nerve compression, which is often misdiagnosed and treated as a CPPS, with limited results.8,9

Recent studies suggest that many chronic pain conditions, particularly fibromyalgia (FM), chronic fatigue syndrome (CFS), and irritable bowel syndrome (IBS), share several demographic, clinical, and psychosocial aspects, as well as pain-related subjective and objective features, with CPPS, perhaps reflecting a mutual primary pathophysiology.10,11 Temporomandibular joint dysfunction, postural anomalies, asymmetry of body axis, and muscle disorders contribute to the long-term development of severe pain syndromes that can also reflect in the urogenital area.12–14

Given the diverse range of symptoms and possible etiologies, it is evident that a “one size fits all” diagnostic investigation and therapeutic approach is not possible, and the treatment can represent a real challenge for the involved physician, who has to evaluate the patient as a whole, taking into account not only aspects directly related to the reported symptoms, as stated in a recent paper on the European Association of Urology guidelines.15 The aim of this narrative review is to focus on CPPS after an infectious/CPPS after an infectious cause has been excluded. The principal studies considered in this work and discussed below are listed in Table 1.

### The UPOINT clinical phenotyping system for CPPS

The most widely adopted questionnaire for clinical evaluation of CPPS is the National Institutes of Health Chronic Prostatitis Symptom Index (NIH-CPSI).16 This tool, validated in 1999, is composed of nine different questions investigating pain, urinary symptoms, and QoL related to CPPS. In 2009, Shoskes et al17 proposed a dedicated clinical classification of CPPS, to separately identify the different possible reported symptoms. It considers: Urinary symptoms (mostly regarding the storage phase but also the voiding phase); Psychosocial dysfunction (depression or catastrophizing thoughts); Organ-specific findings (prostate tenderness or swelling, leukocytosis in prostatic fluid, hematospermia, prostatic calcifications); Infection (exclusion of infective etiology or bowel contamination); Neurological/systemic (presence of abdominal and/or pelvic pain, IBS, FM, CFS); and Tenderness of muscles (presence of palpable muscle spasm or trigger points in abdomen and pelvic floor).

Classifying patients according to the clinical phenotype can allow for exploration of the relative contribution of each domain to the severity of symptoms and, ultimately, to the treatment response by focusing on the main complaints from the individual patient. The
Table 1 Summary of the most relevant studies included in this review

<table>
<thead>
<tr>
<th>Studies</th>
<th>Study type</th>
<th>Domain analyzed</th>
<th>Treatment(s)</th>
<th>No. of patients</th>
<th>p</th>
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<tbody>
<tr>
<td>Zhao et al(^{29})</td>
<td>RCT</td>
<td>• Pain</td>
<td>Celecoxib vs placebo</td>
<td>32 vs 32</td>
<td>&lt;0.006</td>
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<tr>
<td></td>
<td></td>
<td>• Quality of life</td>
<td></td>
<td></td>
<td>&lt;0.032</td>
</tr>
<tr>
<td>Bates et al(^{10})</td>
<td>RCT</td>
<td>• NIH-CPSI</td>
<td>Oral prednisolone vs placebo</td>
<td>18</td>
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</tr>
<tr>
<td>Wagenlehner et al(^{32})</td>
<td>RCT</td>
<td>• NIH-CPSI</td>
<td>Pollen extract vs placebo</td>
<td>70 vs 69</td>
<td>0.0126</td>
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<tr>
<td></td>
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<td>• Pain</td>
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<td></td>
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<tr>
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<td></td>
<td>• Quality of life</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Cai et al(^{35})</td>
<td>RCT</td>
<td>• NIH-CPSI</td>
<td>Pollen extract vs ibuprofen</td>
<td>41 vs 46</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>• Pain</td>
<td></td>
<td></td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Quality of life</td>
<td></td>
<td></td>
<td>0.002</td>
</tr>
<tr>
<td>Pontari et al(^{36})</td>
<td>RCT</td>
<td>• NIH-CPSI</td>
<td>Pregabalin vs placebo</td>
<td>218 vs 106</td>
<td>&lt;0.05</td>
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<tr>
<td></td>
<td></td>
<td>• ≥6 points NIH-CPSI score decrease</td>
<td></td>
<td></td>
<td>0.07</td>
</tr>
<tr>
<td>Nickel et al(^{38})</td>
<td>RCT</td>
<td>• NIH-CPSI</td>
<td>Alfuzosin vs placebo</td>
<td>136 vs 136</td>
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<tr>
<td>Kaplan et al(^{34})</td>
<td>RCT</td>
<td>• NIH-CPSI group 1</td>
<td>Saw palmetto vs finasteride</td>
<td>32 vs 32</td>
<td>0.41</td>
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<td></td>
<td></td>
<td>• NIH-CPSI group 2</td>
<td></td>
<td></td>
<td>&lt;0.003</td>
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<td>Fitzgerald et al(^{39})</td>
<td>RCT</td>
<td>Patient global response assessment</td>
<td>Myofascial physical therapy vs global massage</td>
<td>24 vs 23</td>
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<td>Lee et al(^{42})</td>
<td>RCT</td>
<td>Predefined clinical response criterion</td>
<td>Acupuncture vs sham procedures</td>
<td>44 vs 45</td>
<td>0.017</td>
</tr>
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<td>Schneider et al(^{63})</td>
<td>Prospective series</td>
<td>• Pain</td>
<td>TENS</td>
<td>60</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Quality of life</td>
<td></td>
<td></td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

**Note:** Significant values (p<0.05) are shown in bold.

**Abbreviations:** RCT, randomized controlled trial; NIH-CPSI, National Institutes of Health Chronic Prostatitis Symptom Index; TENS, posterior tibial nerve stimulation/transcutaneous electrical nerve stimulation.
Figure 1 Different phenotypes of chronic prostatitis/chronic pelvic pain syndrome (CP/CPPS) according to the UPOINT classification system,\textsuperscript{17,20} with the most relevant research work cited in the text.

authors found that patients identify their complaints in three main areas and only 22% of them reported a single positive domain. Several works in the literature confirm and validate the clinical utility of this approach.\textsuperscript{18,19}

This evidence demonstrates that CPPS originates from different patterns of pathology and that the therapeutic approach is wide and differs in relation to the reported symptoms. Moreover, the resolution of one “aspect” can lead to the resolution of others, even if not strictly related. Magri et al,\textsuperscript{20} applying the UPOINT classification system on 1,227 patients in a multicentric European series, noticed that many of these patients also complained about sexual dysfunction. Ejaculatory pain is also specifically reported in the NIH-CPSI symptom index (question 2b). Therefore, an implementation of the UPOINT classification has been proposed, adding the domain of sexual dysfunction (erectile dysfunction, impaired sexual desire, and orgasmic dysfunction) as a clinical pattern related to CPPS. Consistent with the study by Magri et al,\textsuperscript{20} in a US study on 162 men, those with a positive sexual dysfunction domain were found to report a lower QoL.\textsuperscript{21} A 2016 meta-analysis estimated the overall prevalence of sexual dysfunction in men with CP/CPPS as up to 0.62 (95% CI 0.48–0.75), while the prevalence of erectile dysfunction and premature ejaculation was 0.29 (95% CI 0.24–0.33) and 0.40 (95% CI 0.30–0.50), respectively.\textsuperscript{22}

The understanding of this clinical phenotyping system for CP/CPPS (Figure 1) can explain why this disease has a wide multifactorial genesis, potentially different in each patient, therefore generating an individual multifaceted complex of symptoms for every patient diagnosed with CP/CPPS. A review by Magistro et al\textsuperscript{23} analyzed 28 randomized controlled trials (RCTs) evaluating various treatments for CP/CPPS, and underlined that monotherapy is never enough to achieve symptom relief and that the therapeutic approach should focus on the different symptomatic pattern presented by the patient in a multimodal setting.

Modulation of inflammatory process

Since the early 2000s, the immunological mechanisms responsible for chronic inflammation in CP/CPPS have been exhaustively explored. Not surprisingly, many different factors seem to be involved. In vivo and in vitro CP/CPPS studies\textsuperscript{24,25} showed a sort of autoimmunity against prostate cells induced by the inflammation, with the recruiting leukocytes including Th1 cells and mast cells, which enhance and trigger the development of CPPS,\textsuperscript{26} in a similar way to that described for rheumatoid arthritis, multiple sclerosis, and inflammatory bowel diseases.\textsuperscript{27,28}

Thus, medications able to interrupt this molecular mechanism can have a primary therapeutic role.

An RCT by Zhao et al\textsuperscript{29} demonstrated the efficacy of celecoxib (dosage 200 mg four times per day, administered for 6 weeks) over placebo for pain modulation; however, symptoms suddenly re-presented after treatment discontinuation. Another trial failed to demonstrate a benefit from oral prednisolone administration over 4 weeks.\textsuperscript{30} Similarly, studies involving monoclonal antibodies did not show significant clinical CPPS improvement.\textsuperscript{31} Common anti-inflammatory drugs (ie, NSAIDs) seem to give rapid relief of symptoms but only for a short period (2–4 weeks). This limits the application of these drugs to acute phases of the disease, without the possibility for long-term administration. Moreover, their well-known side effects further limit their applicability for pain reduction in CP/CPPS.

A multicentric RCT by Wagenlehner et al\textsuperscript{32} investigated the therapeutic benefit of 12 weeks of treatment with pollen extract (Cernilton) in men with CP/CPPS. Participants were evaluated using the NIH-CPSI questionnaire, the number of leukocytes in post-prostatic massage urine, the International Prostate Symptom Score (IPSS), and the sexuality domain of a life satisfaction questionnaire at baseline and after 6 and 12 weeks of treatment. Overall, a decrease in the NIH-CPSI total score by at least 25% or at least 6 points was noticed in the 70.6% of patients undergoing this treatment compared to placebo. Therefore, several studies demonstrated the anti-inflammatory and anti-proliferative role of pollen extract in CP/CPPS due to the inhibition of prostaglandin and leukotriene synthesis in both molecular and clinical trials. Moreover,
flower pollen extracts inhibit 5α-reductase after prolonged therapy and this can also contribute to the reported amelioration of LUTS reported in clinical trials. No major adverse events have been reported in the literature and this treatment is active part of daily urological practice.\textsuperscript{33,34} An RCT by Cai et al\textsuperscript{35} investigated the anti-inflammatory properties of oral administration of pollen extract in association with vitamins in CPPS. Patients were equally divided into two homogeneous groups, receiving oral capsules of DEPROX 500\textsuperscript{®} (two capsules every 24 hours) or ibuprofen (600 mg, one tablet three times a day) for 4 weeks. At the final evaluation, a greater and statistically significant improvement in urinary function and QoL was noticed in the DEPROX 500 group compared with the ibuprofen group. Therefore, these therapeutic agents can improve total symptoms, pain, and QoL in CP/CPPS patients.

**Neuromodulatory therapies**

Neuromodulating medications have a widespread application in the treatment of neuropathic pain, and studies have begun to define their role in the treatment of CP/CPPS.

The proposal of an analgesic neuromodulation of CP/CPPS derives from some studies concerning the effects of a neuromodulatory drug, pregabalin, in increasing dosages (from 150 mg to 600 mg daily) over 6 weeks of treatment, which showed an amelioration of pain symptoms. This is probably due to a chronic activation of sensitive terminations in CP/CPPS that lead to the transmission of nociceptive signals. However, patients did not report symptom resolution but mostly a modest improvement in pain symptoms. Therefore, this effect has not been clearly demonstrated and pregabalin should only be prescribed as a second-line treatment.\textsuperscript{36}

A modification of prostate innervation during age and with the development of benign prostatic hyperplasia (BPH) has been demonstrated,\textsuperscript{37} with an increase in α1-adrenoceptor density leading to an augmented basal tone of the urethral sphincter, and a subsequent increase in urine pressure inside the bladder that can contribute towards increasing the inflammatory response.\textsuperscript{38}

Therefore, neuromodulator therapy can have a consistent role in this pathophysiological pattern in combination with other drugs (ie, α-blockers).

Cordaro et al\textsuperscript{39} investigated the role of the fatty acid amide-signaling molecule palmitoylethanolamide (PEA) in combination with a biological precursor of resveratrol with antioxidant properties named polydatin, for the reduction of BPH-induced neural-mediated inflammation and prostate growth as well as for CPPS symptom modulation. PEA is an endogenous fatty acid amide, belonging to the class of nuclear factor agonists with affinity to cannabinoid-like G-coupled receptors GPR55 and GPR119, with neuromodulatory and anti-inflammatory properties.\textsuperscript{40,41} PEA’s mechanism of action, depicted for the first time in 1993 by the Nobel Prize winner Rita Levi-Montalcini, is described as autacoid local injury antagonism,\textsuperscript{42,43} and acts as a regulator of mast cell-induced inflammation at the level of sensory nerve endings. Therefore, PEA can reduce neuropathic pain and disorders based on glial cell overactivation, such as in diabetes and glaucoma.\textsuperscript{44} Histological findings in CP/CPPS patients' prostates often show infiltrating lymphocytes and macrophages around glandular elements, and therefore PEA could have similar neuronal and immune-modulating effects on prostatic cells. Although not yet validated for CP/CPPS through RCTs, such medications are already in clinical use and have shown a beneficial effect in other clinical applications, for example in reducing the vascular inflammation and oxidative stress responsible for atherosclerosis.\textsuperscript{45}

**Alpha-blockers, phosphodiesterase type 5 inhibitors and 5α-reductase inhibitors**

The pharmacological influence of selective α-blocking agents (such as silodosine or terazosine) is directed against the smooth muscle in the prostatic gland and capsule. Moreover, α-lytic agents are active on the region of the bladder neck in patients with BPH, which is frequently associated with CP. A reduction of voiding pressures and of voiding flow patterns can significantly reduce the discomfort of the patient with CP/CPPS and could be considered in a multimodal therapeutic regimen.\textsuperscript{46,47}

A multicenter, randomized, double-blind, placebo-controlled trial involving 292 patients evaluated the possible efficacy of the α-blocker alfuzosin in reducing overall symptoms in CP/CPPS patients, with the primary outcome being a reduction of at least 4 points in the NIH-CPSI score after 12 weeks of therapy compared to placebo.\textsuperscript{48} The results were discouraging, with similar response rates in the two groups (34.8% vs 36%, \(p=0.9\)). Other studies reported the relief of symptoms by α-blockers, but always limited to patients with bladder outlet obstruction.\textsuperscript{49–52} Evidently, those drugs provide relief from voiding symptoms rather than from pain. So, α-blockers should not be prescribed for patients without voiding issues and attention must be paid when prescribing them to young patients owing to the known negative effect on ejaculation.
Another effective type of drug with the potential to reduce LUTS, erectile dysfunction, and symptoms of CP/CPPS is the phosphodiesterase-5 inhibitor (PDE5-I). A study by Oelke et al.53 demonstrated a similar efficacy of PDE5-I compared to common \( \alpha \)-blockers in terms of amelioration of LUTS and restoration of urinary flow rate, with the known additional positive effect on erectile function.

As known for BPH treatment, \( \alpha \)-reductase inhibitors (5-ARIs) reduce prostate volume by inhibiting the conversion of testosterone to dihydrotestosterone, the metabolically active form responsible for prostatic growth. An RCT by Kaplan et al.54 provided better amelioration of symptoms compared to saw palmetto over a 1-year period. In another study considering a group of men enrolled in a prostate cancer risk reduction study, the NIH-CPSI scores decreased significantly for those under dutasteride compared to placebo.55 Therefore, although a clear action on CP/CPPS has not been demonstrated, the reported amelioration of CPSI scores could encourage 5-ARI therapy in selected patients.

**Physiotherapy and pelvic floor muscle relaxation**
Prostatic massage, perineal or pelvic floor massage, and myofascial trigger-point release have been proposed as a beneficial treatment modality for patients who complain of perineal soreness and difficulty in bladder/rectal evacuation. Muscle tenderness is clearly an important cause of chronic pain and the derived increase in intrapelvic pressure can also lead to worsening of bladder/prostatic symptoms in addition to possible effects in developing chronic orchialgia or low back pain.56,57

Anderson et al, in a series of 74 patients, demonstrated the relationship between specific areas of pelvic pain and specific myofascial trigger points, showing a clear relationship of visceral pain to particular muscular areas.58 Many studies have reported pain relief after muscular massage or physical therapy. One of the most relevant is a study by Fitzgerald et al.59 which demonstrated better symptom relief for CPPS patients undergoing specific myofascial therapy over global massage, with global response assessment response rates of 57% vs 21% (\( p=0.03 \)). It is thus important to refer patients to highly specialized physiotherapy centers.60

Other physical therapy interventions include electromagnetic therapy, microwave thermotherapy, extracorporeal shockwave therapy, acupuncture, and posterior tibial nerve stimulation/transcutaneous electrical nerve stimulation (TENS). Some publications have shown a clear beneficial effect of acupuncture, probably by reducing the neuropathic pain.60 Lee et al, in a controlled trial, demonstrated its efficacy for CPPS symptom reduction compared to placebo.62 Attention has also been focused on TENS, owing to its proven efficacy on musculoskeletal pain. The advantages of this treatment are the possibility to deliver it at the patient's home and the absence of side effects. Schneider et al,63 in a series of 60 patients with CPPS refractory to \( \alpha \)-blockers and common analgesics, described symptom relief in 50% of subjects after 12 weeks of TENS treatment, with perceived amelioration in 70%. Even if TENS is effective, there are no RCTs proving its efficacy on chronic pain relief.64

**Psychological aspects**
An important aspect of CP/CPPS is the long-term duration and evolution of the symptoms. These can be frustrating and disabling, with consequent limitations on the patient’s daily activities and QoL. Psychological support has been proposed as an integrative therapy in the vision of a multimodal approach.55 This is focused on perceiving patients’ internal beliefs in relation to chronic pain, which are often causative of depression or catastrophic thinking. Moreover, there is often a variable perception of pain, which can increase owing to the person focusing on the symptoms. Therefore, patients should be investigated regarding their social support and interactions, daily and family life, and relationship and sexual issues, to orient them towards healthy coping strategies.66

The importance of multidisciplinary treatment is emphasized by several reviews, where the need for high-quality psychological treatment evaluation is underlined.67,68 Therefore, a mental distress evaluation should always be carried out by a dedicated psychologist or psychiatric team, with the aim of addressing susceptible patients.

**Differential diagnosis with pudendal nerve entrapment syndrome**
Originating from the sacral branches S2, S3, and S4, the pudendal nerve (PN) carries sensation from the external genitalia and the skin of the perineum and anus regions, as well as the motor supply to various pelvic muscles, including the male or female external urethral sphincter and the external anal sphincter. Its main branches are the superficial branch, with a sensitive innervation from the perineum and scrotum/labia region, and the deep branch, with a motor
supply for bulbocavernous, ischiocavernous, superficial and deep transverse perineus, and sphincter urethrae muscles. The terminal branch of the PN is the dorsal nerve of the penis or clitoris. Therefore, this nerve is related to the entire pelvic and anogenital area, and its compression or irritation can lead to a wide variety of symptoms.

Pudendal neuralgia, also known as Alcock’s canal syndrome, was first described by Amarenco et al in 1987, in a French paper. It may be related to or be secondary to childbirth, pelvic surgery, intense cycling, sacroiliac skeletal abnormalities, or age-related changes. The most well-known diagnostic criteria for PN are the Nantes criteria, first published in 2008. The diagnosis is mainly clinical, and its five main signs according to these criteria are: 1) pain in the anatomical territory of the PN (ie, between the anus and the penis/clitoris area), which is 2) worsened by sitting; 3) symptoms do not wake the patient during the night; 4) no objective sensory loss on clinical examination; and 5) positive anesthetic PN block.

As can be seen from the Nantes criteria, the diagnosis is mainly clinical. Pereira et al related the possible different symptoms to the site of PN compression, with four different sites: sacral root emergency; infrapiriform below the sacroischiatic ligament; Alcock’s canal; and terminal branches (ie, perineal nerve, dorsal clitoris, and inferior anal nerve).

Symptoms are extremely variable. Pain is probably the last symptom experienced and patients often report a long history of functional problems. The symptoms are often increased in the seated position and became worse during the daytime, with relief during the night; a cutaneous allodynia is typical and due to PN irritation, the symptoms can increase after sexual intercourse, with a delay of 24–48 hours, or after defecation. Women describe worsening of the symptoms when they have sexual intercourse associated with an orgasm. Similarly, men may report changes in the quality of orgasms and premature ejaculation.

The coexistence of pain in the perineum and lower limbs/pseudosciatica is indicative of a disorder of the second segment of the PN. The PN can be entrapped under the sacroischiatic ligament simultaneously with the sciatic and inferior gluteal nerves. PN compression in this area is often the cause of pudendal neuralgia and can be treated with conservative measures or with surgical decompression.

Conservative measures include muscle stretches and exercises to correct spasms and imbalances, as well as electrical stimulation and biofeedback. Postural anomalies also have to be investigated and corrected.

Anesthetic infiltration of the PN provides the definitive diagnostic test for PN entrapment, providing a temporary resolution of reported symptoms, and should be performed before any attempt at surgical decompression.

Owing to the highly variable differences in symptom presentation, pudendal neuralgia is often underestimated and probably often misdiagnosed and confused with CPPS. This condition should always be ruled out in patients presenting with apparent CPPS, particularly at a young age.

**Conclusions**

CP/CPPS is a multivariate and complex disease, often presenting a difficult diagnostic framework. This pathology has a consistent impact on patients’ QoL as it can last for years or a lifetime if not correctly identified and treated. Moreover, owing to its high prevalence, it imposes a significant economic burden on the healthcare system. Practitioners need to move away from old-fashioned habits such as endless courses of empiric antibiotics or underestimation of the patients’ reported problems in the absence of an objective finding, and embrace a multimodal approach to CPPS. Moreover, differential diagnoses such as PN entrapment syndrome should always be considered and investigated.

**Abbreviations list**

ABP, acute bacterial prostatitis; AIP, asymptomatic inflammatory prostatitis; 5-ARI, 5α-reductase inhibitor; BPH, benign prostatic hyperplasia; CBP, chronic bacterial prostatitis; CFS, chronic fatigue syndrome; CP/CPPS, chronic prostatitis/chronic pelvic pain syndrome; FM, fibromyalgia; IBS, irritable bowel syndrome; IPSS, International Prostate Symptom Score; LUTS, lower urinary tract symptoms; NIH, National Institutes of Health; NIH-CPSI, National Institutes of Health Chronic Prostatitis Symptom Index; PDE5-L, phosphodiesterase-5 inhibitor; PEA, palmitoylethanolamide; PN, pudendal nerve; QoL, quality of life; TENS, transcutaneous electrical nerve stimulation.

**Disclosure**

The authors report no conflicts of interest in this work.

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