Psychogenic nonepileptic seizures and psychogenic movement disorders: two sides of the same coin?

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ABSTRACT
Psychogenic nonepileptic seizures (PNES) and psychogenic movement disorders (PMD) are commonly seen in Neurology practice and are categorized in the DSM-5 as functional neurological disorders/conversion disorders. This review encompasses historical and epidemiological data, clinical aspects, diagnostic criteria, treatment and prognosis of these rather challenging and often neglected patients. As a group they have puzzled generations of neurologists and psychiatrists and in some ways continue to do so, perhaps embodying and justifying the ultimate and necessary link between these specialties.

Keywords: psychogenic nonepileptic seizures, psychogenic movement disorders, conversion disorder, hysteria.

RESUMO
Crises não-epilépticas psicogênicas (CNEP) e distúrbios do movimento psicogênicos (DMP) são comuns na prática e na atualidade são melhor categorizados no DSM-V como distúrbios neurológicos funcionais/desordens de conversão. Esta revisão enfatiza os principais dados históricos, epidemiológicos, clínicos, critérios diagnósticos, tratamento e o prognóstico destes pacientes, frequentemente negligenciados e desafiadores, os quais, como um grupo, tem intrigado gerações de neurologistas e psiquiatras, caracterizando, de forma justificada o elo definitivo entre estas especialidades.

Palavras-chave: crises não-epilépticas psicogênicas, distúrbios do movimento psicogênicos, distúrbio conversivo, histeria.

In the medical sciences, a diagnosis is established when a level of certainty regarding the nature of a condition is achieved. Psychogenic nonepileptic seizures (PNES) and psychogenic movement disorders (PMD) share a slightly less linear diagnostic route. Historically, making a conversion diagnosis required ruling out the originally considered medical diagnosis, i.e., it is neither epilepsy, nor a neurogenic movement disorder. Only then a non-neurologic (in the sense of not resulting from epileptogenic or known motor circuitry imbalance) condition is suspected and raised to the level of diagnosis1. This kind of diagnosis has intrigued generations of physicians and spared no culture, gender or age. A diversity of symptoms, including motor, sensory, cognitive, and/or behavior symptoms are present in these presentations. Patients with PNES or PMD frequently are misdiagnosed as having their initial suspected neurological diseases, and ultimately pose a diagnostic and treatment challenge to many clinicians. Interestingly, it was perhaps this varied presentation that has captured the attention of many professionals through the centuries and lead to the construction and recognition of a very particular diagnosis. Two rather advanced historic civilizations, Egyptians first, and then the Greeks, posited that a displaced hystera (womb), and the consequent suffocation that it produced, was the source of choking, mutism, paralysis and fainting1. Hysteria, a natural development on this concept, was claimed as an explanation for similar phenomena in women and reached almost epidemic proportions in the late 19th century. By then, French
and British women displayed a rather elevated prevalence of hysteria, which was associated with the work of Jean Martin Charcot (Figure 1). His harshest critics proposed that he created this disease and gained profit from it. From a contemporary perspective, it may be tempting to scrutinize the dramatic performances of some of Charcot’s young female patients, who may have been prone to suggestibility of the famous professor’s hypnosis techniques. It was due to Charcot’s astute clinical observations, however, that hysteria moved from an almost gynecological disease to a neurologically conceived condition. Regarding male hysteria, Charcot followed after Galen (who suggested that retained sperm could lead to male hysteria) and later Charles LePois, Briquet and Savill, to offer a full description of hysteria in men (Charcot even mentioned epidemiological data, suggesting that 5% of his hysterical patients were male). The “shell-shock” syndrome described in the World War I trenches substantiated the concept of “male hysteria”. Hysteria, in all fairness, respects no gender, i.e. somatoform symptoms are present in women and men. In spite of the controversy produced by historical reflections and opinions on Charcot’s motives and deep involvement with the field of hysteria, these patients finally came to light in a somewhat “medically organized” fashion and certainly inspired very prestigious minds eager to take this diagnosis to a different level of understanding. Sigmund Freud spent four months at the Salpêtrière between October, 1885 and February, 1886 and his enthusiasm for neurosis and psychology was certainly the product of this stay. His attention is observed in that over 100 out of the 3000 books in Freud’s private library were dedicated to hysteria and hypnosis, and all were authored by either Charcot himself or one of the French professor’s direct pupils. Just prior to Charcot’s death, Charcot acknowledged that he had been mistaken and that hysteria was truthfully a psychiatric disease. By then, Alienism was in the process of becoming the modern psychiatry. Hysteria fell into academic disgrace, raising derogatory connotations and the word itself was removed to the lay domain. The condition was renamed by Babinski to pithiatism; the number of publications on hysteria decreased significantly and at a given point it has been considered an excuse for lack of medical knowledge or poor diagnostic skills. Renowned publications such as the British Medical Journal released papers in the 60’s reflecting the thoughts of authors calling hysteria the “disguise for ignorance and fertile source for clinical errors”. This assumption of diagnostic ignorance was proven incorrect with the advent of advanced diagnostic techniques. In the 1970’s and 1980’s, the birth of epilepsy monitoring units and the technology to document patients and their events for long periods of time surfaced literally all sorts of epilepsy “imitators”, including PNES. Some of them, such as paroxysmal, transitory, tremor-like or tic-like episodes without loss of consciousness can be interpreted as either PNES or PMD, depending on the setting and context that they are disclosed. What was “hysteria” in the past, today has different nomenclatures (e.g. medically unexplained symptoms, somatoform disorders, etc.). As any practicing clinician knows, the disorder clearly still presents itself in the 21st century in the symptoms and semiology of a substantial percentage of patients in both epilepsy and movement disorders clinics. Conversion symptoms very frequently can be a sole phenomenon, however, the vast majority of patients have a psychiatric comorbidity and/or a significant stressor, either recently or remotely. In this paper, the authors intend to review and explore both commonalities, as well as, unique expressions that are present in PNES and PMD. We will systematically discuss PNES and PMD as independent conditions that might share similar natures and etiologies, and hopefully, offer a systematic approach to general management and treatment.

FUNCTIONAL NEUROLOGICAL DISORDERS (CONVERSION DISORDERS)

Psychogenic symptoms, or, functional symptoms, are commonly seen in clinical practice and are estimated to represent 10% of all medical complaints. They are common in neurology, accounting for 10 to 33% of patient visits to a neurologist and 9% of inpatient neurology admissions. The importance of the exam in conversion disorder (CD) has been underscored with the inclusion of the presence of these signs in the diagnostic criteria of CD in DSM-5. Several neuro-
logical symptoms can have psychogenic mechanisms underlying their presentation, including paralysis, sensory loss, blindness, astasia-abasia, amnesia, PNES and PMD\cite{26, 28}. Daum et al. performed a systematic and narrative review about the value of “positive” clinical signs for weakness, sensory and gait disorders in conversion disorder\cite{11}. The authors did not study signs for PNES. They concluded that clinical signs for motor, sensory and gait functional neurological symptoms are numerous, and 14 have been validated (7 motor, 5 sensory, and 2 gait related)\cite{11}. Among positive signs of functional motor, sensory and gait disorders, Hoover sign, abductor sign, abductor finger sign, co-contraction, midline splitting, non-anatomical sensory loss, dragging monoplegic gait, and chair test have been validated\cite{11}. Daum et al. also reviewed non-validated, positive signs of functional disorders, which included non-pyramidal weakness, absent pronator drift, arm drop test, Barré test, platysma sign, Babinski trunk-thigh test, Bowlus-Currier test, excessive slowness, fluctuation, psychogenic Romberg test, walking on ice, sudden knee buckling, astasia-abasia, and expressive behavior\cite{11}. Lombardi et al. published on a sign to detect unilateral upper extremity non-organic paresis, the elbow flex-ex\cite{12}. This test is useful in differentiating between functional and non-neurologic arm paresis\cite{12}.

Functional (psychogenic) neurological disorders/CD have been described as hysteria, somatization disorder, non-organic disorders and medically unexplained symptoms\cite{7, 11, 20}. Stone et al., evaluated 1144 new neurology outpatients with symptoms “unexplained by organic disease”\cite{14}. The most common diagnoses were neurological disease but with symptoms unexplained by this condition (26%), headache disorders (26%) and conversion symptoms (motor, sensory or non-epileptic seizures) (18%)\cite{14}. At follow-up, only 0.4% of 1030 patients had an organic disease diagnosis confirmed. The authors concluded that one-third of new neurology outpatients were diagnosed as having symptoms “unexplained by organic disease”\cite{14}.

Structural neuroimaging does not reveal a “conversion lesion”. Unexplained neurological symptoms, however, are being studied using functional MRI. In general, studies using this technique have demonstrated that active inhibition in the orbitofrontal cortex and cingulate gyrus may be implicated in functional disorders\cite{20}. Different subtypes of FNDs are shown in Figure 2. The most prevalent psychogenic conditions in practical clinical neurology are generally PNES and PMD\cite{6, 8}.

**PSYCHOGENIC NON-EPILEPTIC SEIZURES**

Non-epileptic seizures (NES) are characterized by paroxysmal, involuntary, usually time-limited alterations in motor and/or sensory function, level of consciousness and behavior that may resemble epileptic seizures (ES), but are not caused by epileptic discharges\cite{16}. They can be of physiologic or psychogenic (PNES) origin. The most common causes of physiologic non-epileptic events, such as syncope\cite{16}, do not pose a significant problem to differential diagnosis when video-electroencephalography (VEEG) monitoring is used\cite{16}. A long list of mental disorders may present as PNES\cite{16}. Patients typically present with episodes characterized by disrupted consciousness or motor/sensory manifestations. Although deficits and symptoms can be produced voluntarily, as in factitious disorders and malingering (found in a small percentage of individuals with neurologic presentations), psychogenic disorder presentations are involuntary (unconsciously produced), as in patients with somatoform disorders and dissociative disorders\cite{16}. Several putative psychological mechanisms of PNES have been conceptualized and described\cite{17}. Some of the mechanisms include a psychodynamic model of primary gain (intended to solve a dilemma, escape an intolerable situation or reduce anxiety), secondary gain (directed to obtain affective or social benefits), a behavioral model of reinforcement of illness behavior (by attention given to symptoms, for instance), and a psychosocial model of maladaptive coping responses to stress (due to deficiencies of coping style, for example)\cite{17}. Dissociation, suggestion and hypnotizability mechanisms also act anchored in pathogenic beliefs that suggest to the patient a severe disease. Suggestion and dissociation also is found in the unconscious modeling of symptoms by the patient, based in his personal past experience or in his conceptualization of the medical problem\cite{18}.

**Figure 2.** Conversion Disorders Functional Neurological Disorders – Subtypes & Comorbidities.
EPIDEMIOLOGY OF PNES

Studies on the prevalence of PNES show variable but clinically significant results, from five to 33% of outpatients receiving treatment for epilepsy, and from 10 to 58% of inpatients treated for refractory epilepsy present PNES. According to Gates such a significant difference in results may be explained by differences in diagnostic criteria for PNES. A female preponderance of up to 80% has been observed in studies of patients with PNES. PNES is present in children and elderly people, but many patients’ age range between the 20’s and 30’s.

IMPACT OF PNES

PNES is as disabling as epilepsy and may lead to severe social and psychological impairments. Patients are also exposed to iatrogenic procedures, such as high doses of AEDs, intravenous AED use, and orotracheal intubation. Quality of life (QoL) measures in patients with PNES reveal that QoL is worse than that of patients with refractory epilepsy and that QoL is related to symptoms and depression. The diagnostic delay in PNES patients has been reported in the past an average of 7.2 years, however, with increasing awareness of PNES by providers and patients with seizures, and the increased access to seizure monitoring units, the time to correct diagnosis is decreasing, considerably.

CLINICAL PATH TO DIAGNOSIS OF PNES

The possibility of PNES being present is usually considered when there is a complete absence of therapeutic response to AED, loss of response (therapeutic failure), or paradoxical responses (worsening or spontaneous and unexpected remissions). Likewise, PNES may be considered because of atypical, multiple, inconsistent or changing seizure patterns, or when the seizures are provoked by evident and specific emotional stress, with a narrow temporal relation to seizure occurrence. Table 1 summarizes important clinical semiologic features of PNES that help distinguish it from epileptic seizures. These elements are considered particularly when the patient demonstrates normal ancillary exams, (interictal routine EEGs, and neuroimaging studies, such as brain CT, MRI and SPECT). The suspected diagnosis raised by a neurologist has a positive predictive value of 84.6% for PNES. Patients with PNES tend to have a greater frequency of seizures than do epilepsy patients. They also have a greater frequency of hospital admissions due to prolonged seizures, or nonepileptic status. The occurrence of seizures in the physician’s office or in the waiting room is very suggestive of PNES, as well as a history of unexplained “chronic pain” or “fibromyalgia.” Antecedent trauma is reported by up to 70% of PNES patients, and sexual abuse in 40%, which may be an underestimate. They also have other psychogenic disorders in approximately 70% of cases.

Table 1. Clinical features of psychogenic nonepileptic seizures (PNES).

<table>
<thead>
<tr>
<th>Feature</th>
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<tr>
<td>Long duration</td>
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<tr>
<td>Fluctuating course</td>
</tr>
<tr>
<td>Asynchronous movements</td>
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<tr>
<td>Pelvic thrusting</td>
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<tr>
<td>Side-to-side head or body movement</td>
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<tr>
<td>Closed eyes during episode</td>
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<tr>
<td>Ictal crying</td>
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<tr>
<td>Memory recall</td>
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DIAGNOSIS DURING VIDEO EEG

Video EEG (VEEG) is the “gold standard” for proper diagnosis of PNES. During VEEG monitoring, behavior and electroencephalographic activity are simultaneously registered. A spontaneous or elicited event is defined as a PNES when there is no ictal EEG evidence of epileptic discharges before, during or after the ictus, and semiology is consistent with PNES and not epilepsy. The event must be critically investigated in the context of clinical data because simple partial epileptic seizures, parietal lobe and hemispheric seizures may occur without evident epileptic discharges. The International League Against Epilepsy (ILAE) NES Task Force published minimal requirements for NES diagnosis, which utilizes history, semiology and EEG (preferably video) to establish diagnosis. As is done with PMD diagnosis, levels of diagnostic certainty are ranked based on what data are available from history, witnessed event and diagnostic testing, with levels of Possible, Probable, Clinically Established, and Documented diagnosis (see Table 2).

Provocative procedures, such as saline provocation, hypnosis, simple suggestions, suggestive interview or a mixture of them have been used to obtain a typical event, however, the ethics of provocative procedures has been raised. The average sensitivity of saline provocation across studies with VEEG monitoring is approximately 74%. Routine activation procedures (hyperventilation and photic stimulation) can be used during EEG for seizure induction and do not pose the risk of compromising the physician-patient alliance. Approximately 10% of patients with PNES have epilepsy, when studied using the most stringent criteria. Other reports show a range from 5.3 to 73% of patients with PNES with mixed epilepsy. This variability in different studies may reflect several methodological aspects. A study carried in a Brazilian tertiary center found this association occurring in 50% of PNES patients, a higher association.
level than found in studies in the US. Comorbid psychiatric disorders are the rule in patients with PNES, however, brief psychiatric consults do not always find clear abnormalities during the mental status examination that are found with more comprehensive anamnesis. Comorbidity with depressive, anxiety and borderline and obsessive compulsive personality disorders is high. Patients who receive no feedback or intervention after VEEG have no improvement or worsening of PNES. In contrast, a correct diagnosis or a therapeutic communication can reduce or even abolish PNES. Diagnosis and treatment might result in a significant reduction of utilization and cost of health programs.

More recent epidemiological data show that PMD represents 2% to 3% of all cases seen in movement disorders clinics and occur more commonly in women (ratio of men to women about 1:5) between 37 and 50 years of age. PMD is not consistent over time and are not congruent with the classical definitions of neurological movement disorders. The association of changes in abnormal movements with distractibility is very common and is an important observation used during examination.

**Table 2.** Overview of proposed diagnostic levels of certainty for psychogenic nonepileptic seizures.

<table>
<thead>
<tr>
<th>Diagnostic level</th>
<th>History</th>
<th>Witnessed event (semiology)</th>
<th>EEG</th>
</tr>
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<tbody>
<tr>
<td>Possible</td>
<td>+</td>
<td>By witness or self-report/description</td>
<td>No epileptiform activity in routine or sleep deprived interictal EEG</td>
</tr>
<tr>
<td>Probable</td>
<td>+</td>
<td>By clinician who reviewed video recording or in person, showing typical semiology of PNES</td>
<td>No epileptiform activity in routine or sleep deprived interictal EEG</td>
</tr>
<tr>
<td>Clinically established</td>
<td>+</td>
<td>By clinician experienced in diagnosis of seizure disorders (on video or in person), showing typical semiology of PNES, while not on EEG</td>
<td>No epileptiform activity in routine or ambulatory ictal EEG during a typical ictus/event in which the semiology would make ictal epileptiform EEG activity expectable during equivalent epileptic seizures</td>
</tr>
<tr>
<td>Documented</td>
<td>+</td>
<td>By clinician experienced in diagnosis of seizure disorders, showing typical semiology of PNES, while on video EEG</td>
<td>No epileptiform activity immediately before, during or after ictus captured on ictal Video EEG with typical PNES semiology</td>
</tr>
</tbody>
</table>

**Key:** “+”: History characteristics consistent with PNES. EEG: Electroencephalogram. (As noted in the text, additional tests may affect the certainty of the diagnosis – for instance, self-protective maneuvers or forced eye closure during unresponsiveness or normal postictal prolactin levels with convulsive seizures).

**PROGNOSIS OF PNES**

Adults with PNES having risk factors that include comorbid depression, a personality disorder and a history of abuse were more likely to have persistence of seizures, compared to those without these comorbidities. Several studies show that early and correct diagnosis of PNES, followed by adequate treatment, could lead either to remission in 19 to 25%, or to improvement in 75 to 95%. Therefore, correct diagnosis and treatment might result in a significant reduction of utilization and cost of health programs.

**PSYCHOCGENIC MOVEMENT DISORDERS**

Identifying psychogenic movement disorders can be a challenging task in neurological practice. Historically, Charcot was known for his great interest in psychogenic cases and for paying special attention in its definition and treatment. Charcot considered that besides women, men employed in labor, effeminate men and children were prone to the development of hysterical neurological deficits, including PNES (then referred to as hysteroepilepsy) and PMD.

**DIAGNOSIS OF PMD**

The semiology and characteristics of PMD are important for the diagnosis. The onset of PMD is usually abrupt and occurs in the context of a precipitating event. Disabilities may be selective, and signs may improve with distraction. In general, PMD are not consistent over time and are not congruent with the classical definitions of neurological movement disorders. The association of changes in abnormal movements with distractibility is very common and is an important observation used during examination. Fatigability, particularly in patients presenting with tremor, is also common. In some cases, abnormal movements stop when fatigue appears. PMD can present with a wide variety of manifestations, including tremor, dystonia, chorea, gait disorders, parkinsonism, tics and ataxia, usually in combination and affecting multiple body parts. The diagnosis of PMD has been classically viewed as one of exclusion. However, recent clinical advances and advances in neurophysiological examinations have led to the establishment of diagnostic criteria that help clinicians make a more accurate diagnosis and manage the disease better.

There are some clinical characteristics of PMD that are more common in clinical neurological practice. Table 3 summarizes the signs and characteristics associated with PMD.
are traditionally classified into the following four categories according to basic definitions proposed originally by Fahn and Williams in 1988 and subsequently modified by Fahn in 1994. The same diagnostic level category names are used for PNES, however, the criteria for the diagnostic designation differ significantly for PMD:

(i) Documented PMD: Documented PMD includes those patients who have complete resolution of PMD following psychotherapy, psychological suggestion by the physician, physiotherapy or administration of a placebo with suggestion or are witnessed as being free of symptoms when left alone and supposedly unobserved;

(ii) Clinically established PMD: Clinically established PMD is inconsistent over time or incongruent with the typical presentation of a classical movement disorder. In the presence of either of the above, the patient must have any of the additional manifestations, including other neurological signs, multiple somatizations, obvious psychiatric disturbance, disappearance of the PMD with distraction and excessive (almost deliberate) slowing;

(iii) Probable PMD: Probable PMD includes patients with incongruous and inconsistent movements in the absence of any of the other features listed in category 2 (Clinically established) to support the diagnosis of PMD and patients with a movement disorder that is consistent and congruent with a classical neurological movement disorder but who have other features, such as disappearance of the movement with distraction or other psychogenic neurologic disorders and multiple somatizations;

(iv) Possible PMD: Possible PMD is characterized by clinical features of PMD occurring in the presence of an emotional disturbance.

Most of the information used to delineate the currently available diagnostic criteria for PMD come from case series. The three most significant of these series are those described by Lang, including the case series of PMD evaluated by Fahn et al., Jankovic et al., and Lang et al. in which tremor, dystonia, myoclonus and mixed (often bizarre) movement disorders were the most common PMD.

Additionally, in a Brazilian series published in 2010, tremor was the most frequent PMD (55.6%). The second most common PMD semiology was pure dystonia (33.3%), agreeing with data in the medical literature in which the frequency of this presentation ranges from 15% to 53% in Parkinsonism and myoclonus PMD were uncommon, each occurring in four patients. Factor et al. studied 28 patients with PMD and found that the most common semiology was tremor (50%), followed by dystonia, myoclonus and Parkinsonism. Distractibility (86%) and abrupt onset (54%) were the most common clinical characteristics in this series of patients. Twenty-five percent presented with combined PMD and neurological movement disorder.

In general psychogenic or functional tremor (PT) is the most common psychogenic movement disorder. It can occur in any body part, although the hands and arms are the most frequently involved. More rarely, PT can occur in the head and legs. Of the different clues suggesting tremor may be psychogenic, the most common are distractibility, entrainment (a change in the original tremor frequency to match the frequency of a repetitive task performed in another limb) and the presence of coactivation (the co-contraction sign). In general, PT is thought never to affect the fingers, tongue or face. The dystonia semiology represents the second most common form of PMD.

In neurological clinical practice the border between functional or psychogenic, dystonia (FD) and neurological dystonia is not clear-cut. Historically, several forms of dystonia have been considered to be of psychogenic origin, such as blepharospasm and writer’s cramp. In the last 30 years, however, with the great advances in genetics, neurophysiology and neuroimaging, the phenotypes of primary idiopathic dystonia have been clearly defined. Some forms of fixed dystonia associated with previous peripheral trauma and with pain similar to chronic regional pain syndrome (“causalgia-dystonia”), have been considered FD. The most
common forms of FD are blepharospasm, limb focal dystonia and abductor laryngeal dystonia or paradoxical vocal cord dysfunction^35^-^38^.

The next most common form of PMD is myoclonus, now defined as functional myoclonus (FM)^39^. In this setting, electrophysiological tests are needed to ensure correct diagnosis, particularly those using electroencephalogram-electromyography back-averaging^39^. Functional parkinsonism, chorea, tics and ataxia are rarely reported^40^.

**Prognosis OF PMD**

Patients with PMD usually have a poor prognosis^41^-^42^. Cases with at least six months of symptoms at the time of diagnosis had little or no response to proposed treatments^41^-^42^, Feinstein et al. evaluated psychiatric outcomes of their patients with a mean follow up of 3.2 years, showed that PMD persisted in more than 90% of them, especially in cases with comorbid major depression, anxiety and personality disorders^41^, In contrast, Thomas et al., evaluated 228 patients with PMD with a mean follow-up of 3.4 years (6 months to 12 years), and concluded that symptoms improved in 56.6% of patients, worsened in 22.1% and remained the same in 21.3%^41^, Ertan et al. studied 49 patients with PMD in a tertiary clinic in Turkey and concluded that the response to treatment was poor, with a high rate of drop out of these patients in the follow-up^41^.

**PSYCHOGENIC MOVEMENT DISORDERS AND NON-EPILEPTIC SEIZURES**

An intriguing question is whether PMD and PNES share the same psychopathological comorbidities, particularly anxiety, depression and other conditions, such as personality disorders and trauma/abuse histories^45^-^46,^47^-^48^. Against this background, Grimaldi et al. performed a prospective comparative study in which they investigated the presence of anxiety and depression in 17 patients, nine of whom had PMD and eight PNES^45^. They concluded that all the patients had the same demographic and psychopathological profile, although in the group with PNES there was a greater incidence of anxiety disorders and a family history of epilepsy^45^. Driver-Dunckley et al. studied 172 patients in a retrospective chart review, comparing 116 patients with PNES with 56 with PMD^46^. They found that 82% of the patients were female and that 70% had chronic pain, 55% subjective cognitive complaints, 47% fatigue and 45% a history of childhood abuse. The patients with PNES had coexisting epilepsy in 17% of cases, and those with PMD had coexisting neurologic movement disorders in 9%. The authors concluded that PNES and PMD had the same psychopathology, with more similarities than differences^46^. Hopp et al. studied 104 patients with PMD and 35 with PNES using different cognitive, psychological and social function measures^47^. They demonstrated that patients with PNES and PMD, despite differences in their phenomenology and demographics, shared the same psychiatric symptoms, suggesting that PNES and PMD represent different presentations of a single disorder^47^. Mula published an editorial about PNES and PMD discussing the commonalities between these two conditions^48^. He suggested that these disorders occupy a gray area between neurology and psychiatry and commented on the poor level of integration between neurologists and psychiatrists^48^.

**CONCLUDING REMARKS**

Moving the field forward for better understanding of PMD, PNES and other somatoform disorders will require increased collaboration between neurology and psychiatry. A renaissance of neuropsychiatry is being driven by advances in functional neuroimaging, neuroscience and the treatment needs of the aging population with neurodegenerative neuropsychiatric disorders. Viewing somatoform/conversion disorders from a combined lens of neurology/psychiatry provides a comprehensive assessment approach and opens avenues of collaborative management.

“**Lumpers or Splitters**”

From prior research the question arises, should PMD and PNES be “lumped or split”, that is, are they “variations on a theme”, or independent populations? Larger samples for fully powered designed studies may require incorporating both PMD and PNES. On the other hand, discrete samples may be needed to identify regions of interest and putative networks in biomarker focused studies. Also of note is that a number of patients with PNES also have other movement symptoms without change in level of consciousness, apart from their ictus, possibly generating a third group of “mixed PNES/PMD”. These issues can be addressed by linking research centers to reach target sample sizes.

**Treatment of PNES and PMD**

Great strides have been made in validating treatment for patients with PNES in the past decade. Prior to the NINDS/NIMH/AES supported NES Workshop in 2005, only class III and IV level treatment data existed^49^-^50^. The NES workshop set the NES research benchmarks, and PNES treatment is a target of the NINDS Epilepsy Research Benchmarks^50^. Our group and others have been systematically developing treatments for patients with PNES and studying biomarkers of PNES. This is being accomplished through multi-disciplinary, multi-modal, multi-site work. Examples of advances in PNES treatment over the last decade include examining clinical trial methodology in PNES in an open label study of sertraline for PNES^50^, followed by a pilot placebo-controlled
RCT with sertraline. Standard medical care (SMC) or treatment as usual (TAU) is in the US, and in a cross-cultural comparison with Chile has been described. A psychotherapy initially used in epilepsy has been modified for PNES and was used to conduct an open label trial for PNES. With pharmacologic, psychotherapeutic and SMC data, a multi-centered, RCT comparing PNES-CBT, sertraline, PNES-CBT and sertraline, and SMC/TAU was conducted. Based on the successful reduction in seizures, improvement in comorbidities, QoL and functioning in the two groups treated with the manualized therapy, the treatment workbook is being published along with a therapist’s guide. Providers at sites across the US are being trained in administering the manualized treatment. These and other modalities studied in PNES treatments are reviewed in the IALAE NES management article. Advances in PMD treatment include an open label trial of sertraline for PMD. In another study, psychodynamic psychotherapy was used in a single-blind treatment trial for patients with PMD. Non-psychotherapy modalities including physiotherapy also have been used to treat patients with PMD. A summary of cognitive behavioral approaches used in patients with functional neurological disorders provides a review of studies in a variety of somatoform presentations.

With the momentum of recognition by the Movement Disorders Society (MDS) and the International League Against Epilepsy (ILAE) designating these populations’ symptoms as significant disorders found in patients with movement disorders and seizures, task forces are setting standards for diagnosis and treatment. Examples include the ILAE NES Task Force establishing the minimum requirements for making the diagnosis of PNES, and summarized the literature on PNES management. Text books are published summarizing the NES, and PMD literature. National research funding sources (Institutes, Medical Societies and voluntaries) now are providing support for studies examining treatment and mechanisms for patients with conversion disorders. With greater identification of the somatoform disorders by neurologists, psychiatrists, primary care physicians, opportunities for treatment are opened. With a neuropsychiatric conceptualization and formulation and with more management options now available, this population is being demystified and providers are empowered to effectively treat patients. Cross-disciplinary and cross-cultural collaboration will continue to facilitate advances in this common and challenging neuropsychiatric disorder.

References


