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Dual diagnosis of epilepsy and psychogenic nonepileptic seizures: Systematic review and meta-analysis of frequency, correlates, and outcomes

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ABSTRACT

Comorbid epilepsy and psychogenic nonepileptic seizures (PNES) represent a serious challenge for the clinicians. However, the frequency, associations, and outcomes of dual diagnosis of epilepsy and PNES are unclear. The aim of the review was to determine the frequency, correlates, and outcomes of a dual diagnosis. A systematic review of all published observational studies (from inception to Dec. 2016) was conducted to determine the frequency, correlates, and outcomes of dual diagnosis. We included studies of individuals of any age reporting a dual diagnosis of epilepsy and PNES. All observational study designs were included with the exception of case reports and case series with fewer than 10 participants.

The mean frequency of epilepsy in patients with PNES across all studies was 22% (95% confidence intervals [CI] 20 to 25%, range: 0% to 90%) while the mean frequency of PNES in patients with epilepsy was 12% (95% CI 10 to 14%, range: 1% to 62%). High heterogeneity means that these pooled estimates should be viewed with caution. A number of correlates of dual diagnosis were reported. Some studies delineated differences in semiology of seizures in patients with dual diagnosis vs. PNES or epilepsy only. However, most of the correlates were inconclusive. Only a few studies examined outcome in patients with dual diagnosis.

Dual diagnosis is common in clinical practice, especially among patients referred to specialized services, and requires careful diagnosis and management.

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1. Introduction

Psychogenic nonepileptic seizures (PNES) are genuinely experienced events resembling epilepsy but without the concomitant electrophysiological electrical activity [1]. The clinical presentation of PNES and epilepsy can be similar. However, approaches to their management are radically different. Epilepsy requires anticonvulsant therapy, in certain cases, surgery and other nonpharmacological methods, while modern etiological models of PNES find that it has much in common with panic disorder and can benefit in a similar way from explanation and psychotherapeutic interventions [2].

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In some cases, PNES and epilepsy coexist. According to different estimates, between 8% and 60% of patients with PNES also have epilepsy [3]. A number of factors could contribute to the development of PNES in epilepsy such as psychiatric comorbidities, cognitive dysfunction, the experience of unpredictable seizures, and problems with social adaptation [4].

To our knowledge, there is no systematic review exploring the frequency, associations, and outcomes of comorbid epilepsy in patients with PNES and vice versa. These data are important for early identification of those who are at risk of the development of comorbid epilepsy and PNES and planning treatment. The aim of the review was to determine the frequency, correlates, and outcomes of dual diagnosis of epilepsy and PNES.

2. Methods

The systematic review was undertaken following Meta-analyses Of Observational Studies in Epidemiology (MOOSE) guidelines [5] for



Review



[☆] Author contributions: MK searched reference lists, extracted the data, and wrote first draft. YX searched databases and performed statistical analysis. All authors take part in the planning of the paper, critically reviewed, revised, and approved the final draft.

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meta-analysis of observational studies and reported following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines [6].

2.1. Search strategy and data extraction

Five databases were searched: Medical Literature Analysis and Retrieval System Online (MEDLINE), Excerpta Medica dataBASE (EMBASE), Psychological Information Database (PsycINFO), Cumulative Index to Nursing and Allied Health Literature (CINAHL), and Allied and Alternative Medicine (AMED) (from inception to 11 Dec. 2016). The following search terms were used as free text or controlled vocabulary (i.e., medical subject headings, EMTREE) as appropriate for each database: 'epilepsy' AND 'seizure', 'attack', 'non-epileptic', 'psychogenic', 'dissociative', 'conversion', 'functional' (full details available in the supplementary file). Titles and abstracts of all references were screened by one author (YX), and full text articles were examined by another (MK) to determine whether they met the inclusion criteria. Further literature was sought through the reference lists of eligible studies (MK). Data extraction included region/country, recruitment site, study period, age, sample size, frequency of dual diagnosis, method of diagnosis, and who made the diagnosis. One researcher (MK) extracted data. When abstracts from conferences were identified, we sought corresponding published journal articles. We reported data from the abstracts only if corresponding journal articles could not be identified. We judged articles to be from the same cohort if there was evidence of overlapping recruitment sites, study dates, authorship, and similar patient characteristics. We included all published observational studies reporting the frequency of people with dual diagnosis of epilepsy and PNES or associations of comorbid epilepsy and PNES as a primary or secondary outcome regardless of duration of the disease. All observational study designs were accepted with the exception of case report and case series of fewer than 10 participants.

All studies were divided into high- and low-quality groups. The studies were qualified as low-quality if they had either specific participant characteristic limits such as one sex, disabled, but not age or convenience, selective, or random sampling. Risk of bias was assessed using a 10-item assessment, which reflected quality criteria for such studies [7].

2.1.1. Statistical analysis

Studies' reported frequency of comorbid epilepsy and PNES was pooled. We conducted quantitative synthesis and produced forest plots in Stata 13 using random effects analysis. Subgroup analysis was conducted based on the recruitment site (i.e., specialized centers, neuropsychiatry units, psychiatry departments, neuropsychology units, hospitals, population-based, databases) and study population (i.e., adults, adults in epilepsy surgery series, children, adults, veterans, intractable seizures, elderly). Statistical heterogeneity and consistency were assessed using the standard Q statistic, with p < 0.05 and I^2 .

3. Results

The search results and selection processes are summarized in Fig. 1. A total of 2773 references were identified, of which 175 full text articles were retrieved to assess for inclusion/exclusion, and a total of 117 studies (122 reports) were considered eligible. Included papers contained data obtained from 17,478 people. Two studies were population-based [8,9] while the rest were hospital-based [1,3,4,10–126]. In the latter, patients were recruited mostly from highly specialized epilepsy centers (tertiary hospitals, academic departments, comprehensive epilepsy programs – 112 of 118 studies). Sixty-nine studies were retrospective, and 49 were prospective. Children were recruited in seven studies,



Fig. 1. PRISMA flow diagram for the systematic review process.

three of which specifically included those with intractable seizures. In one study, patients with juvenile myoclonic epilepsy of different ages were included. One hundred and nine studies were restricted to adults, of which some were exclusively of specific subpopulations: elderly or veterans (n = 7), surgical series (n = 6), treatment-resistant epilepsy (n = 5), "mental retardation" (n = 1), and traumatic brain injury (n = 1). In 72 studies, patients with dual diagnosis were compared with those with PNES. In 21 studies, patients with dual diagnosis were compared with those with PNES and with epilepsy. The method of diagnosis of epilepsy/PNES was video-electroencephalography monitoring (VEM) in 102 of 118 studies.

3.1. Frequency of dual diagnosis

The frequency of epilepsy in patients with PNES varied from 0% to 90% while the frequency of PNES in patients with epilepsy varied from 1% to 62%.

The pooled frequency of epilepsy among those with PNES was 22% (95% confidence interval [CI] 20 to 25%) (Fig. 2). The pooled frequency of PNES among those with epilepsy was 12% (95% CI 10 to 14%) (Fig. 3). In both cases, the level of heterogeneity (I^2) of included studies was high i.e., 96.5 and 92.7% correspondingly, p = 0.0001.

Meta-analysis of the data according to the site of recruitment yielded similar results. The pooled frequency of dual diagnosis varied from 2% (95% Cl 1 to 4%) among those recruited from epilepsy surgery programs (3 studies, n = 2872) up to 42% (95% Cl 11 to 72%) among patients from (neuro)psychiatric departments and neuropsychology units (3 studies, n = 161). In the two population-based studies, the pooled frequency of epilepsy among those with PNES (2 studies, n = 833,527) was 14% (95% Cl 4 to 23%) [8,9], and the frequency of PNES among those with epilepsy was 17% in a single study [9].

3.2. Demographic features

There were no differences between those with dual diagnosis, PNES, and epilepsy in the aspects of age, age at disease onset, disease duration, sex, marital, or educational status (see Table 1).

3.3. History details

Patients' history of physical or sexual abuse, abuse in childhood, significant head trauma, and substance abuse, was similar among patients with dual diagnosis and PNES in all five studies [12,17,47,56,72] that reported this. It did not differ among patients with dual diagnosis and epilepsy in two [17,72] out of three studies [17,26,72].

3.4. Seizure characteristics and medication

Nonepileptic seizure characteristics were assessed in ten studies with eight finding differences in semiology of seizures between those with PNES and dual diagnosis. Stiffening of the body (60% vs. 37%, p > 0.05) [12], opisthotonus (18.5% vs. 0%, p > 0.03) [56], convulsive behavior (85% vs. 55.5%, p > 0.05) [47], right-hemibody PNES events (7% vs. 23%, p = 0.054) [77], autonomic symptoms/signs during the seizure (51.6% vs. 23.7%, p = 0.03) [27,99], and postictal state (84% vs. 58%, p = 0.02) [12] were more common in patients with PNES than those with dual diagnosis. Total lack of responsiveness (63% vs. 25%, p < 0.05) [18] and myoclonic seizure semiology (10% vs. 2%, p = 0.073) [77] were observed more often in patients with dual diagnosis than in those with PNES. Absence/staring seizures were less common in dual diagnosis in comparison with patients with epilepsy (9% vs. 41%, p = 0.003) [77].

The median seizure frequency was significantly higher in patients with dual diagnosis than in those with PNES (for instance 30 (range: 2 to 500) vs. 125 (range: 1 to 1000) seizures per year [47]) in two [31,



Fig. 2. The frequency of epilepsy in patients with PNES.

47] of three studies, but, in one study, this held only before the diagnosis of PNES [47].

As expected, patients with dual diagnosis had higher antiepileptic drug (AED) use than patients with PNES (5 studies [17,33,47,40,126]) and in one study, lower AED use than in those with epilepsy [17]. Patients with PNES took more psychotropic drugs than AED in one study [33].

3.5. Miscellaneous clinical features

Two studies explored possible association between dual diagnosis and medical comorbidities. One study (n = 689) showed that patients

Subgroups and author year	Sample si	re Frequency (95% Cl), %		Weight, %
Elderly				
Kipervasser 2007	6 —	- ; 	16 (-13 to 45)	0.37
Subtotal (I-squared = .%, p = .)			16 (-13 to 45)	0.37
			. ,	
Adults		1		
Bayly 2013	9 -	-}- ≇	11 (-9 to 31)	0.69
Preiss 2004	17		29 (7 to 51)	0.63
Pak 2012	20	-+∎-+	5 (-5 to 15)	1.82
Shibaddin 1999	23		13 (-1 to 27)	1.22
Ristic 2004	24	_ · _	38 (18 to 57)	0.75
Marchetti 2009	26	· ·	35 (16 to 53)	0.82
Turner 2011	31		32 (16 to 48)	0.96
Neill 1986	31		35 (19 to 52)	0.93
Benbadis 1996	40	_ ₩ _!	3 (-2 to 7)	2.77
King 1982	40		10 (1 to 19)	1 87
Parra 1999	41		49 (34 to 64)	1.06
Baslet 2010	51		22 (10 to 33)	1 54
Tatum 2016	55		4(-1 to 8)	2.76
Osman 2016	60		62 (50 to 74)	1 40
Acton 2012	71		4 (0 to 9)	2 80
Chu 1988	76		3(-1 to 6)	3 01
Szaflarski 2000	101		16 (0 to 22)	2.01
Walczak 199/	101		10 (5 to 25)	2.20
Such 2000	102		14(7 to 21)	2.57
Hisano 1992	127		13(71019)	2.30
Pabati 2011	134		3(3(012))	2.04
Dalleti 2011	154		14(0(020))	2.54
Cervenka 2013	160		5(2108)	3.04
Andersen 2010	1//		24(18(0.31))	2.40
	194		7 (4 to 11)	3.00
Massout-Tarrus 2016	216		12 (8 to 16)	2.87
Hara 2015	225		14 (9 to 19)	2.83
Bett 1992	227		5 (2 to 8)	3.12
van der Peet 2001	229		11 (7 to 16)	2.91
Devinsky1996	268		8 (4 to 11)	3.08
Elliott 2014	365		23 (19 to 27)	2.87
Caller 2014	816		6 (5 to 8)	3.28
Konnikara 2015	1600		7 (6 to 8)	3.32
Subtotal (I-squared = 88.9%, p = 0.0	000)	\mathbf{O}	13 (11 to 16)	70.43
Intractable seizures				
Rotge 2009	36		8 (-1 to 17)	1.92
Sutula 1981	37		14 (2 to 25)	1.58
Slater 1995	42		2 (-2 to 7)	2.81
Cragar 2003	77	┤_」━╋━	25 (15 to 34)	1.81
Henry 1997	145		8 (4 to 12)	2.85
Rotge 2009	175		13 (8 to 18)	2.74
Subtotal (I-squared = 76.6%, p = 0.0) 01)		11 (5 to 16)	13.70
		1		
Veterans		1		
Hegermiller-Smith 2012	43		14 (4 to 24)	1.68
Subtotal (I-squared = .%, p = .)		$\langle \diamond \rangle$	14 (4 to 24)	1.68
Children				
Dhiman 2014	56		16 (6 to 26)	1.81
Desai 2016	73		17 (8,to 26)	1.99
Subtotal (I-squared = 0.0%, p = 0.87	79)		17 (10 to 23)	3.81
(),		-		
Adults underwent epilepsy surgery				
Whitehead 2015	725		3 (1 to 4)	3.33
Markoula 2013	805		3 (2 to 4)	3.32
Reuber 2002 (a)	1342		1 (0 to 2)	3.36
Subtotal (I-squared = 86.2% , n = 0.0	001)		2 (1 to 4)	10.01
		1 I	<u>,</u> ,	
Overall (I-squared = 92.8%, p = 0.00	00)	•	12 (10 to 14)	100.00
NOTE: Weighte are from any days offers	lucio	Ī	,	
ino i ε: weights are from random eπects ana	iysis			
		1 1 1 1	I	
		0% 25% 50% 75%	100%	

Fig. 3. The frequency of PNES in patients with epilepsy.

Age, sex, age,	disease duration.	and education level in	patients with dual diagnosis in	comparison with those w	th PNES and/or epilepsy.
· · · · · · · · · · · · · · · · · · ·			P		

	D'Alessio et al. [18]	Mari et al. [40]	Kuyk et al. [58]	Asadi-Pooya and Emami [12]	Hara et al. [31]	Hoepner et al. [33]	O'Sullivan et al. [47]	Sadan et al. [56]	Cragar et al. [17]	Eliott and Charyton [26]	Galimberti et al. [27]	Jovik ^b [87]	Reuber et al. [62]	Markoula et al. ^a [89]	Konnikara et al. [88]	De Timary et al. [20]	Duncan and Oto [25]	Turner et al. [72]	Wissel et al. [77]	Glosser et al. ^a [28]
Total N in the study	43	110	85	188	225	114	38	51	106	689	138	71	180	805	2735	103	288	53	138	250
N PNES	24	85	60	156		73	20	27	29	324	31	17			1488	50	257	22	46	228
N Epilepsy	-	-	-	-	194				58	281	69	30	90	779	1135			21	46	
N Dual	19	25	25	32	31	41	18	24	19	84	38	24	90	26	112	53	31	10	46	22
Age	<	<	>	=	= ES	=	=	=	=		=	=								
									= ES	= ES	= ES	= ES	= ES							
Age at onset		=	=		= ES	<	=	=	<		>	>				<	<	=		>ES
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Male	>	=	=	=		=	=	=	=			=			<	=				<es< td=""></es<>
									= ES	< ES		=ES	< ES	< ES						
Disease duration	<pnes< td=""><td>>PNES</td><td></td><td></td><td>= ES</td><td><pnes< td=""><td></td><td></td><td><es< td=""><td></td><td>=</td><td>=</td><td></td><td></td><td></td><td></td><td></td><td>=</td><td>>ES</td><td></td></es<></td></pnes<></td></pnes<>	>PNES			= ES	<pnes< td=""><td></td><td></td><td><es< td=""><td></td><td>=</td><td>=</td><td></td><td></td><td></td><td></td><td></td><td>=</td><td>>ES</td><td></td></es<></td></pnes<>			<es< td=""><td></td><td>=</td><td>=</td><td></td><td></td><td></td><td></td><td></td><td>=</td><td>>ES</td><td></td></es<>		=	=						=	>ES	
									>PNES	= ES	= ES	= ES								
Education level		=	=	=	<es< td=""><td></td><td></td><td></td><td>=</td><td></td><td></td><td>></td><td></td><td></td><td></td><td></td><td></td><td>=</td><td></td><td>=</td></es<>				=			>						=		=
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"<" Denotes those with dual diagnosis are younger than those with PNES or lower proportion of males among those with dual diagnosis than those with PNES.

"=" Those with dual diagnosis are at same age/gender balance/disease duration/education level as those with PNES.

">" Those with dual diagnosis older/more males than those with PNES.

"< **ES**" Those with dual diagnosis younger/fewer males/had shorter disease duration than those with epilepsy.

"= ES" Those with dual diagnosis same age/gender balance/disease duration/education level was lower as those with epilepsy.

">ES" older/disease duration longer in those with dual diagnosis than in epilepsy.

">PNES" education level was higher/disease duration was shorter in patients with PNES than with dual diagnosis.

"<PNES" disease duration was shorter/education level was lower in patients with dual diagnosis than with PNES.

^a Included patients who had undergone resective epilepsy surgery.

^b Studies recruiting children.

Table 2

Comparative analysis of epilepsy location in patients with dual diagnosis and epilepsy.

Study	N/N ^a	Dual diagnosis	Epilepsy
Pillali and Haut [59] Konnikara et al. [88] Wissel et al. [77] Reuber et al. [62]	38/78 112/1488 46/46 90/90	Frontal lobe No difference ^b Right hemisphere No difference ^c	Temporal lobe –

^a Number of patients with dual diagnosis/epilepsy.

^b Temporal lobe epilepsy was diagnosed in 79% (dual) and 75% (epilepsy) respectively.
^c More generalized epileptiform interictal changes were registered in patients with dual diagnosis.

with dual diagnosis in comparison with those with epilepsy more often smoked, suffered from pain, asthma, gastroesophageal reflux disease, and migraine while another one (n = 188) [12] reported no association.

In one small study (n = 20), the authors suggested an interesting way of classifying patients with dual diagnosis into three groups: i) resistant epilepsy with anxiety/depression and normal cognition, ii) learning difficulties and dependent personality, and iii) comorbid cluster B personality disorders, anxiety disorders, psychic trauma, and normal cognition. However, the small numbers meant this was only a hypothesis-generating study [4].

3.6. EEG and neuroimaging

As expected, patients with dual diagnosis, in comparison with those with PNES, more often had abnormalities on EEG [33,61,91,109]. They were less susceptible to EEG induction procedures in one study [40]. Temporal lobe epilepsy was equally common in patients with epilepsy and dual diagnosis according to the biggest study of its kind (n = 1488) [88] while smaller studies yielded conflicting results (Table 2).

3.7. Psychiatric comorbidity and neuropsychological features

Most researchers used DSM-IV criteria for diagnosing psychiatric comorbidities [18,47,72], however, there was no consistent approach across researchers to the assessment or categorization of neuropsychological features. At most, specific neuropsychological features were explored in no more than three studies, making an overall summary difficult.

Generally, researchers did not find a difference between PNES and dual diagnosis in psychiatric morbidity [47,56,72] or suicide attempts [18]. In three studies, patients with PNES and dual diagnosis experienced more behavioral problems than patients with epilepsy [27,49,84].

Compared with patients with PNES, patients with dual diagnosis more often experienced affective and personality disorders [26,58] but less often had posttraumatic stress disorder (PTSD) and dissociative disorder [18,58] (Table 3). Using the Minnesota Multiphasic Personality Inventory, patients with dual diagnosis, compared with patients with epilepsy, scored more highly on the 'hysteria' [17,86,122], 'lassitude– malaise', and 'mental dullness' scales [17]. On 'hypochondriasis' and 'depression' scales, they scored lower than patients with PNES in one study [86].

Five studies [25,62,72,87,91] assessed cognitive function in patients with PNES and/or epilepsy and dual diagnosis and yielded conflicting results. Two out of four studies reported significant differences in cognitive functions between patients with PNES and dual diagnosis [87,91]; two found that cognitive changes could predict coexistence of PNES in epilepsy [25,62] while one reported similar cognitive status for patients with dual diagnosis, PNES, and epilepsy [72].

3.8. Employment and disability status

Employment status did not differ among patients with dual diagnosis and PNES in two studies [47,58]. Disability status and length of medical care for the seizure disorder were analyzed in four studies, which yielded conflicting results [17,26,77,126].

3.9. Outcomes in patients with dual diagnosis

Unfavorable outcomes in the group with dual diagnosis were reflected in a greater number of emergency department visits in comparison with epilepsy [15] and more frequent hospital visits [56] and rarer achievement of remission than in those with PNES [78].

Two studies demonstrated that detection of dual diagnosis could improve subsequent outcomes in those patients [13,56]. After diagnosis of PNES, psychogenic events more often ceased in the group with dual diagnosis as opposed to the group with PNES in one study (22% vs. 58%) [56].

4. Discussion

Dual diagnosis is more frequent in this systematic review than previously reported [3]. This could be explained partly by the recruitment of patients in most studies from specialized epilepsy centers where complex and unusual cases concentrate. However, this relatively high frequency of dual diagnosis was also shown in two population-based studies suggesting that it is also common in overall populations of patients primarily diagnosed with epilepsy or PNES.

The lowest frequency of dual diagnosis was registered in surgical series (2%) which is probably explained by careful presurgical examination and exclusion of most of the patients with comorbid PNES. In contrast, the highest frequency of dual diagnosis was observed among patients referred to (neuro)psychiatry/neuropsychology units (42%), which probably reflects factors leading to referral to those services. This finding emphasizes the importance of considering the recruitment setting when looking at comorbidities.

Dual diagnosis was almost twice as frequent in the studies recruiting people with PNES than in the studies recruiting those with epilepsy. It could be that a certain proportion of the patients with epilepsy developed PNES, which then predominated in the clinical picture; such a pattern is common in our experience.

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Psychiatric comorbidities in patients with dual diagnosis, PNES, and epilepsy.

Study	Dual	PNES	Epilepsy	Findings
Sadan et al. [56]	24	27		PNES = dual diagnosis
Turner et al. [72]	10	22	21	ES = PNES = dual diagnosis
O'Sullivan et al. [47]	18	20		PNES = dual diagnosis
Wissel et al. [77]	46	46	46	ES < dual diagnosis; PNES < dual diagnosis depression, anxiety and stressor as a trigger of a seizure
Helmstaedter [84]	335			Dual diagnosis > ES general behavioral problems
Owzcarek [49]	152			Dual diagnosis > ES anxiety and neuroticism
Ito et al. [57]	165			Dual diagnosis > ES dissociation
Eliott and Charyton [26]	84	32	281	Dual diagnosis > PNES as depression, anxiety, bipolar disorder, and personality disorder
Kuyk et al. [58]	25	60		Dual diagnosis > PNES personality disorders;
				Dual diagnosis < PNES somatoform disorder
D'Alessio et al. [18]	19	24		PNES = dual diagnosis

Demographic features of patients with dual diagnosis, PNES, and epilepsy were similar. Despite some differences in individual studies, no consistent or specific semiological signs of dual diagnosis were found which may differentiate it from patients who had PNES or epilepsy alone.

Varying findings on psychiatric comorbidities and neuropsychological correlates in patients with dual diagnosis reflect the clinical heterogeneity of this phenomenon. On the other hand, only some authors performed formal psychiatric evaluation to assess psychiatric comorbidities while the rest used only psychometric scales, which could be considered as a limitation of those studies. There are several potential overlapping mechanisms of the development of PNES in epilepsy. These include a) anxiety and other psychiatric comorbidity arising from the experience of epilepsy; b) the way in which epilepsy may act as a 'symptom scaffold' on which is built a recurrent conditioned response to arousal [2]; c) as an involuntary substitute symptom especially in a population with intellectual disability and after successful surgery for epilepsy. In the latter, case reduction in epileptic seizure frequency leads to the development of PNES driven by secondary gains such as caregiver attention or activity avoidance [4]. The development of PNES after epilepsy surgery could be not only "compensatory" but also a result of psychological stress associated with the operation [60, 63].

Very few studies assessed outcomes in patients with dual diagnosis. Some data showed that dual diagnosis predisposes patients to worse outcomes, but once the correct diagnosis is made, the number of events and number of AED is likely to decrease. This emphasizes the importance of timely diagnosis of PNES in patients with epilepsy.

In clinical practice, the data suggest that patient with treatment-resistant epilepsy is at higher risk of developing PNES, and vice versa. The dual diagnosis should be considered in the cases of the unexpected development of new seizure types or increase in their frequency. Correctly identifying dual diagnosis is typically harder than isolated PNES and especially recording both types of events on VEM. In some cases, details from the patient's history can be misleading. For instance, mild traumatic brain injury is a risk factor for the development of not only epilepsy but also posttraumatic stress disorder and PNES [127]. On the other hand, although stress and adverse experience are considered established risk factors for the development of PNES in some individuals, they can also contribute to the development/exacerbation of epilepsy [128,129].

This systematic review had several limitations. We included studies regardless how PNES, epilepsy, and dual diagnosis were identified. In the majority of cases, the diagnosis was confirmed by VEM while some authors used more relaxed diagnostic criteria. There was a high degree of heterogeneity between the studies that even a random effects meta-analysis may not have compensated for. Nonetheless, we think the meta-analysis has some face validity in describing the published literature although summary values should be interpreted with caution. We also did not evaluate publication bias. There are likely to be other series of patients with ES, where PNES was not a focus of the study title or abstract but it is recorded as a comorbidity. We were not able to analyze studies in relation to the frequency of intellectual disability within individual studies, as these data were rarely available. Clinical experience suggests that patients with intellectual disability have a particularly high rate of dual diagnosis. Methodological limitations of some studies could also affect the dual diagnosis. For instance, provocative tests (verbal suggestion, saline injection) were used in 2 studies during VEM, and in some studies, the 'criteria for epileptiform discharges' were not clearly described. None of the studies presented a priori power calculations for the sample size.

Future research could attempt to build on the categorization of patients with dual diagnosis of PNES and epilepsy under different subtypes depending on mechanisms of development and clinical features. Thus far, there is no work studying effective treatment of PNES in those with dual diagnosis as they tend to be excluded from clinical trials of AEDs. Clinicians working in this area tend to adopt a model of treatment, which focuses on helping patients and their family identify individual seizure types and then treat accordingly. An additional element of therapy in those patients can involve a focus on understanding the hypothesized mechanisms of association described above. Psychogenic nonepileptic seizures are so common in certain patients with epilepsy (those with cognitive decline, affective disorders etc.) that it begs the question of whether preemptive education or psychological interventions are warranted and may be helpful in those high-risk groups?

5. Conclusion

Dual diagnosis is relatively common among those diagnosed with PNES or epilepsy, especially in those who referred to specialized epilepsy centers. This indicates the importance of considering this comorbidity, not only in patients with PNES but also in a population with epilepsy. Future research should pursue potential mechanisms of the development of PNES in epilepsy, describe individual risk factors and test possible interventions for the treatment and possibly early detection and prevention of the development of PNES in patients with epilepsy.

We confirm that we have read the Journal's position on issues involved in ethical publication and affirm that this report is consistent with those guidelines.

Supplementary data to this article can be found online at https://doi. org/10.1016/j.yebeh.2018.10.010.

Conflict of interest

None of the authors have any conflict of interest to disclose.

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References

- Chen JJ, Caller TA, Mecchella JN, Thakur DS, Homa K, Finn CT, et al. Reducing severity of comorbid psychiatric symptoms in an epilepsy clinic using a colocation model: results of a pilot intervention. Epilepsy Behav 2014;39:92–6.
- [2] Brown RJ, Reuber M. Towards an integrative theory of psychogenic non-epileptic seizures (PNES). Clin Psychol Rev 2016;47:55–70.
- [3] Benbadis SR, Agrawal V, Tatum WO. How many patients with psychogenic nonepileptic seizures also have epilepsy? Neurology 2001;57:915–7.
- [4] Magaudda A, Gugliotta SC, Tallarico R, Buccheri T, Alfa R, Laganà A. Identification of three distinct groups of patients with both epilepsy and psychogenic nonepileptic seizures. Epilepsy Behav 2011;22:318–23.
- [5] Stroup DF, Berlin JA, Morton SC, Olkin I, Williamson GD, Rennie D, et al. Meta-analysis of observational studies in epidemiology: a proposal for reporting. JAMA 2000; 283:2008–12.
- [6] Moher D, Liberati A, Tetzlaff J, Altman DG, PRISMA Group. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. PLoS Med 2009; 6:e1000097.
- [7] Quality assessment tool for observational cohort and cross-sectional studies. Available at: https://www.nhlbi.nih.gov/health-topics/study-quality-assessment-tools, Accessed date: 1 March 2018.
- [8] Szaflarski JP, Ficker DM, Cahill WT, Privitera MD. Four-year incidence of nonepileptic seizures in Hamilton County, OH. Neurology 2000;55:1561–3.
- [9] Sigurdardottir KR, Olafsson E. Incidence of psychogenic seizures in adults: a population-based study in Iceland. Epilepsia 1998;39:749–52.

- [10] Alessi R, Vincentiis S, Rzezak P, Valente KD. Semiology of psychogenic nonepileptic seizures: age-related differences. Epilepsy Behav 2013;27:292–5.
- [11] Anderson J, Hill J, Alford M, Oto M, Russell A, Razvi S. Healthcare resource utilization after medium-term residential assessment for epilepsy and psychogenic nonepileptic seizures. Epilepsy Behav 2016;62:147–52.
- [12] Asadi-Pooya AA, Emami M. Demographic and clinical manifestations of psychogenic non-epileptic seizures: the impact of co-existing epilepsy in patients or their family members. Epilepsy Behav 2013;27:1–3.
- [13] Baheti NN, Radhakrishnan A, Radhakrishnan K. A critical appraisal on the utility of long-term video-EEG monitoring in older adults. Epilepsy Res 2011;97:12–9.
- [14] Bayly J, Carino J, Petrovski S, Smit M, Fernando DA, Vinton A, et al. Time-frequency mapping of the rhythmic limb movements distinguishes convulsive epileptic from psychogenic nonepileptic seizures. Epilepsia 2013;54:1402–8.
- [15] Caller TA, Chen JJ, Harrington JJ, Bujarski KA, Jobst BC. Predictors for readmissions after video-EEG monitoring. Neurology 2014;83:450–5.
- [16] Cervenka MC, Lesser R, Tran TT, Fortuné T, Muthugovindan D, Miglioretti DL. Does the teddy bear sign predict psychogenic nonepileptic seizures? Epilepsy Behav 2013;28:217–20.
- [17] Cragar DE, Schmitt FA, Berry DTR, Cibula JE, Dearth CM, Fakhoury TA. A comparison of MMPI-2 decision rules in the diagnosis of nonepileptic seizures. J Clin Exp Neuropsychol 2003;25:793–804.
- [18] D'Alessio L, Giagante B, Oddo S, Silva WW, Solís P, Consalvo D, et al. Psychiatric disorders in patients with psychogenic non-epileptic seizures, with and without comorbid epilepsy. Seizure 2006;15:333–9.
- [19] Davies KG, Blumer DP, Lobo S, Hermann BP, Phillips BL, Montouris GD. De novo nonepileptic seizures after cranial surgery for epilepsy: incidence and risk factors. Epilepsy Behav 2000;1:436–43.
- [20] De Timary P, Fouchet P, Sylin M, Indriets JP, de Barsy T, Lefèbvre A, et al. Non-epileptic seizures: delayed diagnosis in patients presenting with electroencephalographic (EEG) or clinical signs of epileptic seizures. Seizure 2002;11:193–7.
- [21] Desai D, Desai S, Jani T. Juvenile myoclonic epilepsy in rural Western India: not yet a benign syndrome. Epilepsy Res Treat 2016;2016:1435150.
- [22] Devinsky O, Sanchez-Villaseñor F, Vazquez B, Kothari M, Alper K, Luciano D. Clinical profile of patients with epileptic and nonepileptic seizures. Neurology 1996;46: 1530–3.
- [23] Dhiman V, Sinha S, Rawat VS, Harish T, Chaturvedi SK, Satishchandra P. Semiological characteristics of adults with psychogenic nonepileptic seizures (PNESs): an attempt towards a new classification. Epilepsy Behav 2013;27:427–32.
- [24] Dhiman V, Sinha S, Rawat VS, Vijaysagar KJ, Thippeswamy H, Srinath S, et al. Children with psychogenic non-epileptic seizures (PNES): a detailed semiologic analysis and modified new classification. Brain Dev 2014;36:287–93.
- [25] Duncan R, Oto M. Predictors of antecedent factors in psychogenic nonepileptic attacks: multivariate analysis. Neurology 2008;71:1000–5.
- [26] Elliott JO, Charyton C. Biopsychosocial predictors of psychogenic non-epileptic seizures. Epilepsy Res 2014;108:1543–53.
- [27] Galimberti CA, Teresa Ratti M, Murelli R, Marchioni E, Manni R, Tartara A. Patients with psychogenic nonepileptic seizures, alone or epilepsy-associated, share a psychological profile distinct from that of epilepsy patients. J Neurol 2003;250:338–46.
- [28] Glosser G, Roberts D, Glosser DS. Nonepileptic seizures after resective epilepsy surgery. Epilepsia 1999;40:1750–4.
- [29] Gordon PC, Valiengo LDCL, Proença ICGF, Kurcgant D, Jorge CL, Castro LH, et al. Comorbid epilepsy and psychogenic non-epileptic seizures: how well do patients and caregivers distinguish between the two. Seizure 2014;23:537–41.
- [30] Hamid H, Fodeh SJ, Lizama AG, Czlapinski R, Pugh MJ, WC Jr LaFrance, et al. Validating a natural language processing tool to exclude psychogenic nonepileptic seizures in electronic medical record-based epilepsy research. Epilepsy Behav 2013; 29:578–80.
- [31] Hara K, Adachi N, Akanuma N, Ito M, Okazaki M, Matsubara R, et al. Dissociative experiences in epilepsy: effects of epilepsy-related factors on pathological dissociation. Epilepsy Behav 2015;44:185–91.
- [32] Herskovitz M. Psychogenic nonepileptic seizure patterns in patients with epilepsy. Psychosomatics 2015;56:78–84.
- [33] Hoepner R, Labudda K, May TW, Schöndienst M, Bien CG, Brandt C. Distinguishing between patients with pure psychogenic nonepileptic seizures and those with comorbid epilepsy by means of clinical data. Epilepsy Behav 2014;35:54–8.
- [34] Jones SG, O'Brien TJ, Adams SJ, Mocellin R, Kilpatrick CJ, Yerra R, et al. Clinical characteristics and outcome in patients with psychogenic nonepileptic seizures. Psychosom Med 2010;72:487–97.
- [35] Kalogjera-Sackellares D, Sackellares JC. Intellectual and neuropsychological features of patients with psychogenic pseudoseizures. Psychiatry Res 1999;86:73–84.
- [36] Kellinghaus C, Loddenkemper T, Dinner D, Lachhwani D, Lüders HO. Non-epileptic seizures of the elderly. J Neurol 2004;251:704–9.
- [37] Kipervasser S, Neufeld MY. Video-EEG monitoring of paroxysmal events in the elderly. Acta Neurol Scand 2007;116:221–5.
- [38] Marchetti RL, Kurcgant D, Neto JG, Von Bismark MA, Fiore LA. Evaluating patients with suspected nonepileptic psychogenic seizures. J Neuropsychiatry Clin Neurosci 2009;21:292–8.
- [39] Martin R, Burneo J, Prasad A, Powell T, Faught E, Knowlton R, et al. Frequency of epilepsy in patients with psychogenic seizures monitored by video-EEG. Neurology 2003;61:1791–2.
- [40] Mari F, Di Bonaventura C, Vanacore N, Fattouch J, Vaudano AE, Egeo G, et al. Video-EEG study of psychogenic nonepileptic seizures: differential characteristics in patients with and without epilepsy. Epilepsia 2006;47(Suppl. 5):64–7.
- [41] Massot-Tarrús A, McLachlan RS. Marijuana use in adults admitted to a Canadian epilepsy monitoring unit. Epilepsy Behav 2016;63:73–8.

- [42] Metrick ME, Ritter FJ, Gates JR, Jacobs MP, Skare SS, Loewenson RB. Nonepileptic events in childhood. Epilepsia 1991;32:322–8.
- [43] Müller T, Merschhemke M, Dehnicke C, Sanders M, Meencke H-J. Improving diagnostic procedure and treatment in patients with non-epileptic seizures (NES). Seizure 2002;11:85–9.
- [44] Neill JC, Alvarez N. Differential diagnosis of epileptic versus pseudoepileptic seizures in developmentally disabled persons. Appl Res Ment Retard 1986;7:285–98.
- [45] O'Brien FM, Delanty N, Dineen C, Murphy KC. Psychogenic non-epileptic seizures in an Irish tertiary referral centre for epilepsy. Ir J Psychol Med 2009;26:174–8.
 [46] Osman A, Seri S, Cavanna AE. Clinical characteristics of patients with epilepsy in a
- [46] Osman A, Seri S, Cavanna AE. Clinical characteristics of patients with epilepsy in a specialist neuropsychiatry service. Epilepsy Behav 2016;58:44–7.
- [47] O'Sullivan SS, Spillane JE, McMahon EM, Sweeney BJ, Galvin RJ, McNamara B, et al. Clinical characteristics and outcome of patients diagnosed with psychogenic nonepileptic seizures: a 5-year review. Epilepsy Behav 2007;11:77–84.
- [48] Owczarek K, Jędrzejczak J. Patients with coexistent psychogenic pseudoepileptic and epileptic seizures: a psychological profile. Seizure 2001;10:566–9.
- [49] Owczarek K. Anxiety as a differential factor in epileptic versus psychogenic pseudoepileptic seizures. Epilepsy Res 2003;52:227–32.
- [50] Parra J, Iriarte J, Kanner AM. Are we overusing the diagnosis of psychogenic nonepileptic events? Seizure 1999;8:223–7.
- [51] Patidar Y, Gupta M, Khwaja G, Chowdhury D, Batra A, Dasgupta A. Clinical profile of psychogenic non-epileptic seizures in adults: a study of 63 cases. Ann Indian Acad Neurol 2013;16:157.
- [52] Plioplys S, Doss J, Siddarth P, Bursch B, Falcone T, Forgey M, et al. A multisite controlled study of risk factors in pediatric psychogenic nonepileptic seizures. Epilepsia 2014;55:1739–47.
- [53] Rawat VS, Dhiman V, Sinha S, Vijay Sagar KJ, Thippeswamy H, Chaturvedi SK, et al. Co-morbidities and outcome of childhood psychogenic non-epileptic seizures – an observational study. Seizure 2015;25:95–8.
- [54] Ribaï P, Tugendhaft P, Legros B. Usefulness of prolonged video-EEG monitoring and provocative procedure with saline injection for the diagnosis of non epileptic seizures of psychogenic origin. J Neurol 2006;253:328–32.
- [55] Rotge JY, Lambrecq V, Marchal C, Pedespan JM, Burbaud P, Rougier A, et al. Conversion disorder and coexisting nonepileptic seizures in patients with refractory seizures. Epilepsy Behav 2009;16:350–2.
- [56] Sadan O, Neufeld MY, Parmet Y, Rozenberg A, Kipervasser S. Psychogenic seizures: long-term outcome in patients with and without epilepsy. Acta Neurol Scand 2016; 133:145–51.
- [57] Ito M, Adachi N, Okazaki M, Kato M, Onuma T. Evaluation of dissociative experiences and the clinical utility of the dissociative experience scale in patients with coexisting epilepsy and psychogenic nonepileptic seizures. Epilepsy Behav 2009; 16:491–4.
- [58] Kuyk J, Swinkels WAM, Spinhoven P. Psychopathologies in patients with nonepileptic seizures with and without comorbid epilepsy: how different are they? Epilepsy Behav 2003;4:13–8.
- [59] Pillai JA, Haut SR. Patients with epilepsy and psychogenic non-epileptic seizures: an inpatient video-EEG monitoring study. Seizure 2012;21:24–7.
- [60] Reuber M, Kral T, Kurthen M, Elger CE. New-onset psychogenic seizures after intracranial neurosurgery. Acta Neurochir 2002;144:901–7.
- [61] Reuber M, Fernandez G, Bauer J, Singh DD, Elger CE. Interictal EEG abnormalities in patients with psychogenic nonepileptic seizures. Epilepsia 2002;43:1013–20.
- [62] Reuber M, Qurishi A, Bauer J, Helmstaedter C, Fernandez G, Widman G, et al. Are there physical risk factors for psychogenic non-epileptic seizures in patients with epilepsy? Seizure 2003;12:561–7.
- [63] Reuber M, Kurthen M, Fernández G, Schramm J, Elger CE. Epilepsy surgery in patients with additional psychogenic seizures. Arch Neurol 2002;59:82–6.
- [64] Shihabuddin B, Abou-Khalil B, Fakhoury T. The value of combined ambulatory cassette-EEG and video monitoring in the differential diagnosis of intractable seizures. Clin Neurophysiol 1999;110:1452–7.
- [65] Silva W, Giagante B, Saizar R, D'Alessio L, Oddo S, Consalvo D, et al. Clinical features and prognosis of nonepileptic seizures in a developing country. Epilepsia 2001;42: 398–401.
- [66] Slater JD, Brown MC, Jacobs W, Ramsay RE. Induction of pseudoseizures with intravenous saline placebo. Epilepsia 1995;36:580–5.
- [67] Syed TU, Arozullah AM, Suciu GP, Toub J, Kim H, Dougherty ML, et al. Do observer and self-reports of ictal eye closure predict psychogenic nonepileptic seizures? Epilepsia 2008;49:898–904.
- [68] Syed TU, Arozullah AM, Loparo KL, Jamasebi R, Suciu GP, Griffin C, et al. A self-administered screening instrument for psychogenic nonepileptic seizures. Neurology 2009;72:1646–52.
- [69] Szabõ L, Siegler Z, Zubek L, Liptai Z, Körhegyi I, Bánsági B, et al. A detailed semiologic analysis of childhood psychogenic nonepileptic seizures. Epilepsia 2012;53:565–70.
- [70] Tamer SK. The pediatric non-epileptic seizure. Indian J Pediatr 1997;64:671-6.
- [71] Tatum WO, Diciaccio B, Yelvington KH. Cortical processing during smartphone text messaging. Epilepsy Behav 2016;59:117–21.
- [72] Turner K, Piazzini A, Chiesa V, Barbieri V, Vignoli A, Gardella E, et al. Patients with epilepsy and patients with psychogenic non-epileptic seizures: video-EEG, clinical and neuropsychological evaluation. Seizure 2011;20:706–10.
- [73] Vincentiis S, Valente KD, Thomé-Souza S, Kuczinsky E, Fiore LA, Negrão N. Risk factors for psychogenic nonepileptic seizures in children and adolescents with epilepsy. Epilepsy Behav 2006;8:294–8.
- [74] Walczak TS, Williams DT, Berten W. Utility and reliability of placebo infusion in the evaluation of patients with seizures. Neurology 1994;44:394–9.
- [75] Westbrook LE, Devinsky O, Geocadin R. Nonepileptic seizures after head injury. Epilepsia 1998;39:978–82.

- [76] Whitehead K, O'Sullivan S, Walker M. Impact of psychogenic nonepileptic seizures on epilepsy presurgical investigation and surgical outcomes. Epilepsy Behav 2015; 46:246–8.
- [77] Wissel BD, Dwivedi AK, Gaston TE, Rodriguez-Porcel FJ, Aljaafari D, Hopp JL, et al. Which patients with epilepsy are at risk for psychogenic nonepileptic seizures (PNES)? A multicenter case–control study. Epilepsy Behav 2016;61:180–4.
- [78] Yadav A, Agarwal R, Park J. Outcome of psychogenic nonepileptic seizures (PNES) in children: a 2-year follow-up study. Epilepsy Behav 2015;53:168–73.
- [79] Acton EK, Doll K, Charles J, Shih J, Tatum WO. Respiratory compromise in PNEA patients on the EMU. Epilepsy Curr 2012;12(Suppl. 1):3.173 [Abstract].
- [80] Betts T, Boden S. Diagnosis, management and prognosis of a group of 128 patients with non-epileptic attack disorder. Part I. Seizure 1992;1:19–26.
- [81] Chu NS. Long-term ambulatory EEG evaluation of epileptic seizures and non-epileptic attacks: a study of 100 patients. Chin Med J (Taipei) 1988;42:359–66.
- [82] Farooq O, Agarwal N, Li P, Jani R, Boudreau M, Kerr S, et al. The presence of non-epileptic seizures in an epilepsy monitoring unit (EMU). Epilepsy Curr 2014;14:1.132 [Abstract].
- [83] Hegermiller-Smith B, Schooff DM. Differences in psychiatric co-morbidities between epileptic and non-epileptic seizures among the veteran population. A study conducted at the Durham VA Medical Center. Epilepsy Curr 2012;12(Suppl. 1):3.096 [Abstract].
- [84] Helmstadter C. Behavioural features of patients with psychogenic non-epileptic seizures (PNES). Epilepsy Curr 2015;15:1.283 [Abstract].
- [85] Hernandez AW, Bailey L, Johnson C, Perry MS, Malik SI. Prevalence of non-epileptic events in children with and without epilepsy admitted to a level 4 epilepsy center. Epilepsy Curr 2013;13:3.354 [Abstract].
- [86] Jedrzejczak J, Owczarek K, Majkowski J. Psychogenic pseudoepileptic seizures: clinical and electroencephalogram (EEG) video-tape recordings. Eur J Neurol 1999;6: 473–9.
- [87] Jovic NJ. Cognitive performance of children and adolescents with pseudoepileptic seizures. Psihijatrija Danas 1998;30:423–39.
- [88] Konikkara JJ, Pacheco J, Van Ness P, Agostini M, Hays R, Howe-Martin L, et al. Unique characteristics of patients with comorbid epileptic and psychogenic nonepileptic seizures. Epilepsy Curr 2015;15:426–7.
- [89] Markoula S, De Tisi J, Foong J, Duncan JS. Psychogenic nonepileptic seizures after adult epilepsy surgery. Epilepsy Curr 2013;13:3.271 [Abstract].
- [90] Pak A, Anschel DJ, Zhang S. Using the body outline task in epilepsy and nonepileptic seizures. Epilepsy Curr 2013;13:3.271 [Abstract].
- [91] Reuber M, Fernández G, Helmstaedter C, Qurishi A, Elger CE. Evidence of brain abnormality in patients with psychogenic nonepileptic seizures. Epilepsy Behav 2002;3:249–54.
- [92] Villalobos R, Gonzales E. Non-epileptic seizures in a population of diagnosed pediatric epilepsy patients. Epilepsy Curr 2015;15:12.274 [Abstract].
- [93] Wada JA. Differential diagnosis of epilepsy. EEG Suppl 1985;37:285-311.
- [94] Ziemba KS, Drazkowski JF. Driving safety in people with non-epileptic events. Epilepsy Curr 2015;15:247.
- [95] Preiss J, Vojtech Z, Haas T. Is it possible to diagnose pseudoseizures (non-epileptic psychogenic seizures) by Dissociative Experience Scale (DES)? Ceska Slov Psychiatr 2004;100:197–203.
- [96] Uhlmann C, Eisele F, Flammer E. Diagnostik und Therapie von Patienten mit nichtepileptischen dissoziativen Krampfanfällen in einer Abteilung für Epileptologie. Z Psychosom Med Psychother 2017;57:288–94.
- [97] Gugliotta SC, Alfa R, Lagana AMA. Presa in carico e valutazione psichica di una popolazione di pazienti con Crisi Psicogene (PNES) Psychological evaluation and management of patients with psychogenic non-epileptic seizures (PNES). Bol Lega Ital Epil 2013;145:35–43.
- [98] Turner K, Piazzini A, Barbieri V, Chiesa V, Vignoli A, Gardella E, et al. Psychiatric and cognitive profiles of patients with epilepsy and psychogenic non-epileptic seizures (PNES). Bol Lega Ital Epil 2010;140:40–2.
- [99] Galimberti C, Ratti M, Murelli R, Sartori I, Zanotta N, Manni R, et al. Non-epileptic psychogenic seizures: clinical and psychological findings. Bol Lega Ital Epil 1998: 327–8.
- [100] Hisano T, Adachi N, Onuma T. Clinical characteristics of 10 epileptic patients with pseudoseizures. J Jpn Epilepsy Soc 1992;10:209–14.
- [101] Ristic AJ, Petrovic I, Vojvodic N, Janković S, Sokić D. Phenomenology and psychiatric origins of psychogenic non-epileptic seizures. Srp Arh Celok Lek 2004;132:22–7.

- [102] van der Peet J, Swinkels W, Duijsens I. Personality disorders in patients admitted to an epilepsy clinic. Tijdschr Psychiatr 2001;43:683–91.
- [103] Meierkord H, Will B, Fish D, Shorvon S. The clinical features and prognosis of pseudoseizures diagnosed using video-EEG telemetry. Neurology 1991;41:1643–6.
 [104] Lesser RP, Lueders H, Dinner DS. Evidence for epilepsy is rare in patients with psy-
- chogenic seizures. Neurology 1983;33:502–4.
 [105] Krumholz A. Niedermever E. Psychogenic seizures: a clinical study with follow-up
- [106] Barry E, Krumholz A, Bergey GK, Chatha H, Alemayehu S, Grattan L. Nonepileptic
 [106] Barry E, Krumholz A, Bergey GK, Chatha H, Alemayehu S, Grattan L. Nonepileptic
- posttraumatic seizures. Epilepsia 1998;39:427–31.
- [107] Mayor R, Howlett S, Grünewald R, Reuber M. Long-term outcome of brief augmented psychodynamic interpersonal therapy for psychogenic nonepileptic seizures: seizure control and health care utilization. Epilepsia 2010;51:1169–76.
- [108] Lancman ME, Brotherton TA, Asconapé JJ, Penry JK. Psychogenic seizures in adults: a longitudinal analysis. Seizure 1993;2:281–6.
- [109] Lelliott PT, Fenwick P. Cerebral pathology in pseudoseizures. Acta Neurol Scand 1991;83:129–32.
- [110] Henry TR, Drury I. Non-epileptic seizures in temporal lobectomy candidates with medically refractory seizures. Neurology 1997;48:1374–82.
- [111] Cohen RJ, Suter C. Hysterical seizures: suggestion as a provocative EEG test. Ann Neurol 1982;11:391–5.
- [112] Baslet G, Roiko A, Prensky E. Heterogeneity in psychogenic nonepileptic seizures: understanding the role of psychiatric and neurological factors. Epilepsy Behav 2010;17:236–41.
- [113] Benbadis SR, Lancman ME, King LM, Swanson SJ. Preictal pseudosleep: a new finding in psychogenic seizures. Neurology 1996;47:63–7.
- [114] Kotagal P, Costa M, Wyllie E, Wolgamuth B. Paroxysmal nonepileptic events in children and adolescents. Pediatrics 2002;110:e46.
- [115] Kanner AM, Parra J, Frey M, Stebbins G, Pierre-Louis S, Iriarte J. Psychiatric and neurologic predictors of psychogenic pseudoseizure outcome. Neurology 1999;53: 933–8.
- [116] King DW, Gallagher BB, Murvin AJ, Smith DB, Marcus DJ, Hartlage LC, et al. Pseudoseizures: diagnostic evaluation. Neurology 1982;32:18–23.
- [117] Lempert T, Schmidt D. Natural history and outcome of psychogenic seizures: a clinical study in 50 patients. J Neurol 1990;237:35–8.
- [118] Seneviratne U, Briggs B, Lowenstern D, D'Souza W. The spectrum of psychogenic non-epileptic seizures and comorbidities seen in an epilepsy monitoring unit. J Clin Neurosci 2011;18:361–3.
- [119] Gulick TA, Spinks IP, King DW. Pseudoseizures: ictal phenomena. Neurology 1982; 32:24–30.
- [120] Sutula TP, Sackellares JC, Miller JQ, Dreifuss FE. Intensive monitoring in refractory epilepsy. Neurology 1981;31:243–7.
- [121] Leis AA, Ross MA, Summers AK. Psychogenic seizures: ictal characteristics and diagnostic pitfalls. Neurology 1992;42:95–9.
- [122] Jedrzejczak J, Owczarek K. Psychogenic pseudoepileptic seizures: analysis of the clinical data in 1990–1997. Krankenhauspsychiatrie 1999;10:36–40.
- [123] Paquet JM, Turpin JC, Luaute JP, Gross J-C. Diagnosis of pseudo-seizures using prolonged video-coupled EEG. Ann Med Psychol 1998;156:631–4.
- [124] Fujimoto S, Mizuno K, Takasaka Y, Shibata H, Kanayama M, Ishikawa T. Ictal electroencephalographic recordings of patients with seizure. Rinsho Byori 1995;43: 865–70.
- [125] Holmes GL, Sackellares JC, McKiernan J, Ragland M, Dreifuss FE. Evaluation of childhood pseudoseizures using EEG telemetry and video tape monitoring. J Pediatr 1980;97:554–8.
- [126] Duncan R, Graham CD, Oto M, Russell A, McKernan L, Copstick S. Primary and secondary care attendance, anticonvulsant and antidepressant use and psychiatric contact 5–10 years after diagnosis in 188 patients with psychogenic non-epileptic seizures. J Neurol Neurosurg Psychiatry 2014;85:954–8.
- [127] Berg AT, Altalib HH, Devinsky Ö. Psychiatric and behavioral comorbidities in epilepsy: a critical reappraisal. Epilepsia 2017;58:1123–30.
- [128] Galtrey CM, Mula M, Cock HR. Stress and epilepsy: fact or fiction, and what can we do about it? Pract Neurol 2016;16:270–8.
- [129] Lang JD, Taylor DC, Kasper BS. Stress, seizures, and epilepsy: patient narratives. Epilepsy Behav 2018;80:163–72.