"I agree that, in practice, most patients have

# significant

'comorbidities' that complicate treatment, as well as research. I believe that not only biological psychiatric research, but also clinical psychiatry would benefit greatly from incorporating the functional psychopathology he describes into our

diagnostic framework."

## Dear Dr. Hollander:

I read with great interest the recent article by Herman M. van Praag, MD.<sup>1</sup> I found enlightening his discussion of the need for revision or supplementation of our standard psychiatric diagnoses in light of the awkward predicaments of biological researchers who attempt to correlate neurobiology with psychopathology. I agree that, in practice, most patients have significant "comorbidities" that complicate treatment, as well as research. I believe that not only biological psychiatric research, but also clinical psychiatry would benefit greatly from incorporating the functional psychopathology he describes into our diagnostic framework.

I write to highlight that the functional psychopathology he describes might be very familiar to anyone with psychoanalytic education. Freud's description of the ego of the structural model<sup>2</sup> included its roles in impulse control, in modulating affect (by mobilization of defense mechanisms), in intellectual functioning, in perception, and in other parameters that can be assessed independently and potentially in a quantitative manner. Heinz Hartmann<sup>3</sup> elaborated on ego functions, adding to its roles dimensions such as adaptability and motoric control. Hartmann<sup>4</sup> also reoriented the psychoanalytic understanding of schizophrenia by suggesting that these patients suffer from ego defects that impair reality testing. Various psychoanalytic researchers such as Greenspan<sup>5</sup> continue to refine our phenomenological understanding of psychological variables and their "functional psychopathology."

These examples illustrate a small sample of the long tradition in psychoanalytic thought of diagnosing ego distortions, deviations, and regressions in individual patients. I believe this approach is very similar to the functional psychopathology described by Dr. van Praag, and that differences and changes in various dimensions of the ego might be correlated with variables of interest in neurobiological research in the way suggested by Dr. van Praag.

In addition, his observation of symptom diversity and variability over time is a fundamental concept in psychoanalytic thought. The manifest symptom is viewed as less enduring and as secondary to the ever-present workings of the ego and its "functional psychopathology."

I am not suggesting that psychoanalytic theory be the basis for revising or supplementing current nosology, but it appears Dr. van Praag's conclusions about the utility of diagnosing apart from nosology are consonant with many decades of careful observations by generations of psychoanalysts. The brilliantly advancing field of neurobiology and the evolving field of psychoanalysis have much to share.

In an article entitled "The Future of Biological Psychiatry," it is interesting to see so much of the history of psychoanalysis. I am heartened on the one hand to see the return of the clinically based observations and perspectives of psychoanalysis, and saddened on the other hand that this important heritage for psychiatry is not recognized immediately in its reincarnation.

# **Burton Hutto, MD** Chapel Hill, NC

## REFERENCES

- Van Praag HM. The future of biological psychiatry. CNS 1 Spectrums. 1997;2:18-25.
- 2. Freud S. The ego and the id. In: Strachey J, ed. The Standard Edition of the Complete Psychological Works of Sigmund Freud. London, England: Hogarth Press; 1953.
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- 4. Hartmann H. Contribution to the metapsychology of schizophrenia. In: The Psychoanalytic Study of the Child. New York: International Universities Press; 1953:8;177-197.
- 5. Greenspan S. Developmentally Based Psychotherapy. New York: International Universities Press; 1996.

## Dear Dr. Hollander:

Dr. Hutto's remarks are well taken. In psychoanalytic theory, there is much that seems to be valuable for further progress and sophistication in psychiatric diagnosing. I underlined this point of view in my inaugural address at the University of Groningen delivered in 1968 and entitled: "The complementary relationship between biological and psychodynamic psychiatry" (published in: Psychiat. Clin. 2, 307-318, 1969).

Unfortunately, the rapprochement has not occurred yet, for I believe in two principle reasons: First, the ever increasing nosological orientation of biological psychiatry, and second, the resistance of psychoanalytically oriented clinicians to put their theses to the test of empirical research. Both obstacles, however, are removable. I would enjoy a merging of forces.

Herman M. van Praag Maastricht University Maastricht The Netherlands

# SPECTRUN

The International Journal of Neuropsychiatric Medicine Volume 3 - Number 1

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Psychiatrist

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(pp 20-57)					

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Neurologist

## **NEURONTIN®**

(Gabapentin Capsules) Before prescribing, please see full prescribing information. A Brief Summary follows.

## INDICATIONS AND USAGE

in) is indicated as adjunctive therapy in the treatment of particl seizures with and without secondary generalization in adults with epilepsy. CONTRAINDICATIONS

urantin® is contraindicated in patients who have demonstrated hypersensitivity to the drug or its ingredients.

## WARNINGS

## Withdrawal Precipitated Seizure, Status Epilepticus

Virtuardwal receptioned success, standard people Intelligitation of the control document because of the processing scizure frequency. In the plocebe-controlled studies, the incidence of status epilepticus is patients receiving blocebe (2 of 378). Among the 2014 pointers thereit with Neuronith\* course of studies (controlled and uncontrolled) 31(13/5) had status epilepticus. Of these, 14 pointers had no prior history of status epilepticus data between tertament or while on other medications because deducet historical data are not an oblight. The impossible to any whether on an theotment with Neuronith\*'s essociated with a higher or lower rate of status epilepticus than would be expected to occur in a similar population nat therefuel Milescontin\*.

### Tumorigenic Potential

Tumorogenic Fortennica In structor periodic in viro Minter conclopencity studies, on unexpectedy high incidence of prozenic ocian adexactrinous was identified in mole, but not lenale, rats. (See RECAUTORS: Concingenesis, Nutrogenesis, Impairment of Fertility.) The direct significance of this finding is unknown. Circial experience during galagenthis' prematering development provide no direct means to assess its potential for inducing tumoss in humans. In direct studies comprising 2005 potentypers of exposure, new tumos were reported in 10 partients (2 breast, 3 brain, 2 lung, 1 advend). I non-Hodgin's lymphoma, 1 endonetrial conci-tion on it adult, and presenting tumos versions were reported in 10 partients (2 breast, 3 brain, 2 lung). Todenol, 1 non-Hodgin's lymphoma, 1 endonetrial conci-tion on its adult, and presenting tumos versions were reported in 10 partients (2 breast, 3 brain, 2 lung). Todenol, 1 non-Hodgin's lymphoma, 1 endonetrial conci-tion on its adult, and presenting tumos versions were the series of the series in a structure provide for the background incidence and an experience on a structure galageneties of the series of the series of a series of the ser Sudden and Unexplained D

unexplained deaths in patients with epilepsy not receiving nilensy) (ors , whether these fig res are reassuring or raise further concern depends on comp archility of the populations reported upon to the Neurontin® coho

## PRECAUTIONS

Information for Patients

Patients should be instructed to take Neurontin® only as prescribed.

Patients should be advised that Neuronim® may cause dizciness, sommalence and other symptoms and signs of OKS depression. Accordingly, they should be advised neither to drive a car nor to aperate other complex machinery until they have gained softicient experience on Neuronim® to gauge whether or not it affects their mentil and/or mator performance adversely.

## Laboratory Tests

Clinical hids statu do not indicate that routine monitoring of clinical loboratory parameters is necessary for the safe use of Neurontin®. The value of monitoring Neurontin® blood concentrations are abalasted. Neurontin® may be used in combination with other antiepilepit drugs without concern for alternation of the blood concentrations of galaxysets are abalasted. ontiecileatic druas

## **Drug Interactions**

Bocquents is an appreciably, metabolized nor does it interfere with the metabolism of commonly coordineistered antispleptic drugs. The drug interaction data described in this section were obtained from studies involving healthy adults and actients with epileroy.

Preservation: In a single and multiple does study of Neuroniane (400 mg 11.0) in opiectic patients (N = 8) maintained on phenytain monotherapy for at least 2 months, galagentin had no effect on the steady-state trough plasma concentrations of phenytain and phenytain had no effect on galagaentin phanmocokinerics.

Carbomazepine: Steadystate trough plasma carbamazepine and carbamazepine 10, 11 epoxide concentrations were not affected by concomitant gabapentin (400 mg TLD.; N = 12) macokinetics were unaltered by carbamazepine ac . Likewise, gabapentin p

Valproix Acids: The mean steady-state trough serum valproix acid concentrations prior to and during concomitant gabapentin administration (400 mg 1.1.0.; N = 17) were not different and neither were gologentin phormackinetic parameters offected by valgravic acid.
Phonobarhited: Estimates of steady-state phormacokinetic parameters for phenobarhited or gologentin (300 mg T.D.; N =12) are identical whether the drugs are administered clone

or together

Constitute: In the presence of cirreticine or 300 mg Q.D.: (N = 12) the mean apparent and clearance of galagentin fell by 14% and creatinine clearance fell by 10%. Thus cirreticine appared to after the renal excention of both galagentin and creatinine, an endogenous marker of renal function. This small decrease in excention of galagentin by cirreticine is not expected to

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Pregnancy Carport C Goopentin his been shown to be letituix in indents, causing deliged costicution of several hores in the skull, vertebrue, torsimite, and indiniss. These effects occurad when pregnant mice severed and does of 1000 on 3000 mg/kg/dxy intrips feed of organopases, or oppositionelly in 14 times the maximum does of 36000 mg/dby given to explexite patients on on g/m bass. The readingt a letituity or oppositionelly in the human does on on g/m bass, there not verse does prior to and during mating, and hanguing patrition, pars from all does groups (500), 1000 mg/kg/dxy letituity and the letituity of the base of the control of the set of the

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It is not known if gebopertin is societed in human mik and the effect on the runsing infant is unknown. However, because many drugs are excreted in human mik, Neurontin® should be used in normal who examines the society of the beautits clearly converging the tisks.

Sofety and effectiveness in pediatric patients below the one of 12 years have not been established

Geriatric Use

to systematic studies in genatric patients have been conducted. Adverse clickal events reported among 59 Neuronnin \* exposed patients over ope 65 did not affier in kind from those reported for younger individuals. The small number of older individuals evaluated, however, limits the strength of any conclusions reached about the influence, if any, of age on the kind and incidence of adverse events or laboratory abcornality associated with the use of Neuronia e. Because Neurontia e alminated primarity by renal excretion, the dose of Neuronia e should be adjusted as nated in DOSAGE AND ADMINISTRATION (Table 2) for elderly patients with

componiest event buchton. Creatinine dearnore is difficult to messure in outpatients and serum creatinine may be reduced in the elderly because of decreased muscle mass. Creatinine dearners (Cr.2) can be reasonably well estimated using the exaction of (ackardth and Gouth.

for females	$C_{r_{*}} = (0.85)(140 \text{age})(\text{wt})/[(72)(S_{r_{*}})]$	

for moles

 $C_{C_{f}} = (140 \text{ toge})(\text{wt})/[(12)(S_{C_{f}})]$ whis in kilograms and  $S_{C_{f}}$  is serum creatining ine in mg/dl ADVERSE REACTIONS

The most commonly observed adverse events associated with the use of Neurontin® in combination with other antiepleptic drugs, not seen at an equivalent frequency among placebo-treated

The max cannot be covered service events associated with the USE of reactions " in combination with other antippleptic drugs, not seen at an equivalent frequency among pixele-tected potents, were somehered, dozines, atouts, thinge, and pastognus. Approximately 7% of the 2014 forkholds who exceed Reaconding in periodical into discontinued testment because of an adverse event. The objects events most commonly costored with withdowed were somehered (12%), atouto (0.8%), taigue (0.6%), naceo and/or vorniting (0.6%), and dizziness (0.6%). Incidence in Controlled Chinical Trials

lable 1 lass treatmentenergent signs and symptons that ocurred in at least 1% of Neuronin "heated patients with epileosy participating in placebocontrolled trials and were numerically more common in the Neurontin" group. In these studies, either Neuronin" or placebo was added to the patient's current antiepilepit drug theory. Atherse events were usually mide to moderate in intensity

The pescible should be notee that these liques, obtained when Neuronin® was odded to concurrent antipolopic drug thempy, current be used to pescit the troppency of adverse events in the coarse of cond maked process where potent documentations and the latters may all the time from providing damage that documents, however, because control to detext yoursel while the time and from the distinct intergrations and one damage that the state state is a state of the state state. Nonexel, the event begaves across the detext yoursel while the state of the one theorem is however, because of the state states, however, because state the restoring physician with one basis to estimate the relative contribution of drug and nondrug factors to the adverse event incidences in the population studied

TABLE 1. Treatment-Emergent Adverse Event Incidence in Controlled Add-On Trials (Events in at least 1% of Neurontin\* (Gabagentin Capsules) patients and numerically more frequent than in the placebo group

	Neurontin <sup>®0</sup>	Placebo <sup>0</sup>		Neurontin <sup>®0</sup>	Placebo <sup>a</sup>
Body System/	N = 543	N = 378	Body System/	N = 543	N = 378
Adverse Event	%	%	Adverse Event	%	%
Body As A Whole			Nervous System (continued)		
Fotigue	11.0	5.0	Tremor	6.8	3.2
Weight Increase	2.9	1.6	Nervousness	2.4	1.9
Back Pain	1.8	0.5	Dysarthria	2.4	0.5
Peripheral Edema	17	0.5	Amnesia	2.2	0.0
ardiovas <u>cular</u>			Depression	1.8	1.1
Vasadilatation	1.1	0.3	Thinking Abnormal	1.7	1.3
Digestive <u>System</u>			Twitching	1.3	0.5
Dyspepsia	2.2	0.5	Coordination Abnormal	1.1	0.3
Mouth or Throat Dry	1.7	0.5	Respiratory System		
Constipation	1.5	0.8	Rhinitis	4.1	3.7
Dental Abnormalities	1.5	0.3	Pharyngitis	2.8	1.6
Increased Appetite	1.1	0.8	Coughing	1.8	1.3
lematologic and Lymphatic Systems			Skin and Appendages		
Leukopenia	1.1	0.5	Abrasion	1.3	0.0
Ausculoskeletal System			Pruritus	1.3	0.5
Myalgia	2.0	1.9	Urogenital System		
Frocture	1.1	8.0	Impotence	1.5	1.1
ervous System			Special Senses		
Somolence	19.3	8.7	Diplopia ,	5.9	1.9
Dizziness	17.1	6.9	Amblycpio <sup>D</sup>	4.2	1.1
Ataxia	12.5	5.6	Laboratory Deviations		
Nystagmus	8.3	4.0	WBC Decreased	1.1	0.5

<sup>0</sup> Plus background antiepileptic drug therapy <sup>b</sup> Amblyopia was often described as blurred visior

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Other events in more than 1% of patients but equally or more frequent in the placebo group included; headache, viral infection, fever, nausea and/or vamiting, abdaminal pain, diarrhea, as, confusion, insomnia, emotional lability, rash, acne

Among the treatment-emergent adverse events occurring at an incidence of at lenst 10% of Neurontin-treated patients, somviolence and atoxin appeared to exhibit a positive doseresmonse relatio

The overall incidence of adverse events and the types of adverse events seen were similar among men and women treated with Neurontin®. The incidence of adverse events increased slightly with increasing ope in portents treated with after Mexamini's calculate access on the same increase interact man measurement in a measurement or overset events histoped and there are insufficient due to support a statement regarding the distribution of oderse events by roce.

### Other Adverse Events Observed During All Clinical Trials

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cleady lated in the previous bable, those too general to be informative, and those not reasonably associated with the use of the dag. Events as further abceller within loady system categories and enumerated in order of decreasing frequency using the following definitions: frequent adverse events are defined as those occurs in in in at loss 11/100 pointers, interest anterese events and the control of the date of the date of the set of Body As A Whole: Frequent astheria, malaise, face edema; Infrequent: allergy, generalized edema, weight devease, chill; Rare: stronge feelings, lassitude, alcohol intolerance,

rer effec nager control Cardiovescular System: Frequent Inventoria, Infraquent hypotension, angina pectoria, peripheral vasolar disorder, polytatica, tachyandia, migrarie, murrur, Rare atrial Minderian, heart failure, fromtoghlealtis, deep throntoghlealtis, myscardial infraction, centroversular accident, pulmarary thrombosis, venticular extraspatiles, backcardia, premature atrial contraction,

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Endocrine System: Rave hyperthyroid, hypothyroid, goiter, hypoestogen, ovarian failure, epididymitis, swollen testicle, cushingoid appearance.

Henerologic and Lymphanic Systems: Frequer processing described is builds reading from physical insura, infrequent anemia, thrombosytopenia, lymphoderopathy, Rave WBC court increased, lymphonytosis, ron-Hodylin's lymphono, bleeding time increased.

sculoskeletal System: Frequent antimology, Infrequent, tendinitis, arthritis, joint stiffness, joint swelling, positive Romberg test, Rore: costochondritis, osteoporosis, busitis, contracture. 

Respiratory System: Frequent, pneumonia; Infrequent, epistaxis, dyspnea, conea, Rare, mucasitis, aspiration pneumonia, hyperventilation, hiccup, laryngitis, nasal obstruction, snoring,

oventilation, lung edema. Dementological: Infrazont, objecta, eczema, dry skin, increased sweating, unicania, hirsutism, seborthea, cyst, herces simplex; Rare; herces zoster, skin discolor, skin papules,

Vermitergenet integration of the second part of urgency, vaginal pain, breast pain, testicle pain.

Special Senses: Frequent chromal vision; Anfequent culturat, conjunctivitis, eyes dry, eye pain, visual held delect, photopholia, biotecti or uniated patosis, eye hemandruge, hordealum, henring lass, cantole, finituita, intere ar Frichica, olita, justa lass, unada labe, eye mithing, and hilans, Sie eyes hitting, automati accountabilito, patomative eyes, entrophydi, automati, and constrainti, la toras explosation paties errore, entrophydi, automa, intis, constrainti, disclassificate durings, binthese latitud adperationi, micios, disclassificita, dispensitive for a disclassification disclassification disclassification dispensitive for advective disclassification disclassification dispensitive for advective disclassification disclassification dispensitive for advective disclassification disclassification disclassification disclassification dispensitive for advective disclassification strabismus, eustachian tube dysfunction, labyrinthitis, atitis externa, add smell.

## Postintroduction Reports

Adverse events social with Neuronia's that have been received since market intraduction, that are not listed above, and that may have no causal relationship to drug, include the following: erythema multiforme, Stevens kolmson syndrome and elevated liver function tests. DRUG ABUSE AND DEPENDENCE

endence potential of Neurontin® has not been evaluated in human studies. The abuse and dea OVERDOSAGE

A lethal dase of gabapentin was not identified in mice and rats receiving single and dases as kigh as 8000 mg/kg. Signs of acute taxicity in animals included ataxia, labored breathing, grosis, sedation, hypoactivity, or excitatio

Anne and overdoses of Neurontin® up to 49 crams have been reported. In these cases, double vision, shared speech, drawsiness, lethoray and domines were observed. All notients recovered with supportive core.

Gabacentin can be removed by hemodialysis. Although hemodialysis has not been performed in the few averalose cases reported, it may be indicated by the patient's clinical state or in patients with size from transferred

### DOSAGE AND ADMINISTRATION

Records P is service mended for coldon therapy in patients over 12 years of age. Evidence beging on its safety and effectiveness in pediatric patients below the age of 12 is not available. Neuronin \* is given analy with an without food.

Heardman<sup>+</sup> is given andly winn a winnut hoad. The effective does of the summan<sup>+</sup> is 900 to 1600 mg/day and given in divided does (these times o day) using 300 nor 400mg capacies. Thinkin to an effective does can take place apply, over a few doys, giving 300 mg on Day 1, 300 mg twice a day on Day 2, and 300 mg these times a day on Day 3. To minimize potential side effects, especially samediance, duziness, faitigue, and chands, the find does on Day. I mgo be chrinicated or beathman. If executing 400 me 400mg capacies Themese aday us to 1800 mg/day, Decayses up to 2400 mg/day hore beare will belanded in largered ministra date. Decay 6 300 mg/day hore bear datinised to a small number of patients for a relatively short duration, and have been well betende. The maximum time between doces in the LLD, schedule should not exceed 12 hours.

The ran exercising to manifer galapentin plasma concentrations to optimize Neurontin® therapy. Further, because there are no significant plasma;colorient interactions among Neu after commanly used antisplastic drugs, the addition of Neurontin® does not after the plasma levels of these drugs appreciably.

If Neuronin''s discontinued and/or an alternate anticonculsant medication is added to the therapy, this should be done gradually over a minimum of 1 week. Dosage adjustment in patients with compromised renal function or undergoing hematiallysis is recommended as follows:

TABLE 2. Neumatin® Dosone Based on Reput Function

Renal Function		
Creatinine Clearance	Total Daily Dase	Dose Regimen
(mL/min)	(mg/day)	(mg)
>60	1200	400 T.I.D.
30-60	600	300 B.I.D.
15-30	300	300 Q.D.
<15	150	300 Q.O.D. <sup>d</sup>
Hernodiolysis	_	200-300 <sup>b</sup>

<sup>0</sup> Every other day

Loading dose of 300 to 400 mg in patients who have never received Neurontin®, then 200 to 300 mg Neurontin® following each 4 hours of hemodialysic

Caution: Federal law prohibits dispensing without prescription.

PARKE-DAVIS

v of Warner-Lambert Co. ©1 prris Plains, NJ 07950 USA

David G, age 31\* NEURONTIN, 1800 mg a day as adjunctive therapy for partial seizures

# NEURONTIN ADJUNCTIVE THERAPY OFFERS EASY AND RAPID TITRATION FOR IMPROVED INDIVIDUAL CONTROL

■ NEURONTIN can be rapidly titrated to effect, up to 1800 mg/day (600 mg tid).<sup>#8</sup> In clinical studies, doses of 3600 mg/day were well tolerated in a small number of patients during short-term administration

■NEURONTIN has no pharmacokinetic interactions with commonly prescribed first-line AEDs: valproic acid, carbamazepine, phenobarbital, or phenytoin

■ NEURONTIN offers the confidence that comes from experience in over 300,000 patients

\*Hypothetical patient

Please see adjacent page for a brief summary of full prescribing information.

gabapentin capsules 100 mg, 300 mg, 400 mg WELL TOLERATED...EASILY

WELL TOLERATED...EASILY TITRATED...PROVEN EFFICACY

NEURONTIN is indicated as adjunctive therapy in the treatment of partial seizures with and without secondary generalization in adults (>12 years old). NEURONTIN is contraindicated in patients who have demonstrated hypersensitivity to the drug or its ingredients.

In placebo-controlled studies, status epilepticus occurred in 0.6% (3/543) of NEURONTIN-treated patients vs 0.5% (2/378) of placebo-treated patients. Because adequate historical data are not available, it is impossible to say whether treatment with NEURONTIN is associated with a higher or lower rate of status epilepticus.

In placebo- controlled studies (n=543), the most common adverse events associated with NEURONTIN were somnolence (19.3% vs 8.7% with placebo); dizziness (17.1% vs 6.9% with placebo); ataxia (12.5% vs 5.6% with placebo); fatigue (11% vs 5% with placebo); nystagmus (8.3% vs 4% with placebo).

† Because NEURONTIN is eliminated renally, dosage adjustment is recommended in renally compromised patients or those patients undergoing hemodialysis. Please see Dosage and Administration section of full prescribing information for schedule.

+ To minimize potential side effects, especially somnolence, dizziness, fatigue, and ataxia, the first dose on Day 1 may be administered at bedtime.

§ Titration to an effective dose can take place rapidly, over a few days, giving 300 mg on Day 1, 300 mg twice a day on Day 2, and 300 mg three times a day on Day 3. Once titrated to 900 mg/day (300 mg tid), if necessary the dose may be increased using 300-mg or 400-mg capsules three times a day, up to 1800 mg/day.

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