

# The Canadian Journal of Neurological Sciences

# Le Journal Canadien des Sciences Neurologiques



## SPECIAL FEATURES

- **Novel Uses of EMG to Study Normal and Disordered Motor Control**  
*Richard B. Stein* ..... 95
- **Mechanisms of Action of Neural Grafts in the Limbic System**  
*Cyörgy Buzsaki and Fred H. Gage* ..... 99
- **Idiopathic Parkinson's disease: Revised Concepts of Cognitive and Affective Status**  
*Ann E. Taylor, J.A. Saint-Cyr and A.E. Lang* ..... 106

## ORIGINAL ARTICLES

- **Immunocytochemical Analysis of Intermediate Filaments in Human Ependymal Tumors**  
*D. Izukawa and B. Lach* ..... 114
- **Cerebral Metastases from Malignant Melanoma**  
*Ivar M. Mendez and Rolando F. Del Maestro* ..... 119
- **Hypomelanosis of Ito. Neurological Complications in 34 Cases**  
*Ignacio Pascual-Castroviejo, Luisa López-Rodríguez, María de la Cruz Medina, Cipriano Salamanca-Maesso and Carmen Roche Herrero* ..... 124
- **Cerebral Arteriovenous Malformations with Associated Arterial Aneurysms: Hemodynamic and Therapeutic Considerations**  
*Douglas Kondziolka, Bruce J. Nixon, Pierre Lasjaunias, William S. Tucker, Karel TerBrugge and Sanford M. Spiegel* ..... 130
- **Late Progression of Post-Encephalitic Parkinson's Syndrome**  
*Donald B. Calne and Andrew J. Lees* ..... 135
- **Parental Age and Birth Order in Alzheimer's disease: A Case-Control Study in the Saguenay-Lac-St-Jean Area (Quebec, Canada)**  
*M. De Braekeleer, S. Froda, D. Gauthrin, H. Tetreault and D. Gauvreau* ..... 139
- **Familial Alzheimer's Disease**  
*A.D. Sadnovick, H. Tuokko, A. Horton, P.A. Baird and B.L. Beattie* ..... 142
- **The Effects of Myoinositol on the Autonomic Neuropathy in the Streptozotocin Diabetic Rat - A Freeze Fracture Study**  
*G. Monckton and H. Marusyk* ..... 147
- **Calcification in a Recent Cerebral Infarct - Radiologic and Pathologic Correlation**  
*J. Parisi, C. Place and S. Nag* ..... 152
- **CANADIAN ASSOCIATION OF NEUROPATHOLOGISTS - Abstracts** ..... 172
- **XXIIIrd CANADIAN CONGRESS OF NEUROLOGICAL SCIENCES - Program and Abstracts** ..... 179

(complete Table of Contents page iii)

**XXIIIrd Canadian Congress of  
Neurological Sciences**

**June 14-18, 1988**

**Quebec City, P.Q.**

**Program and Abstracts ..... page 179**

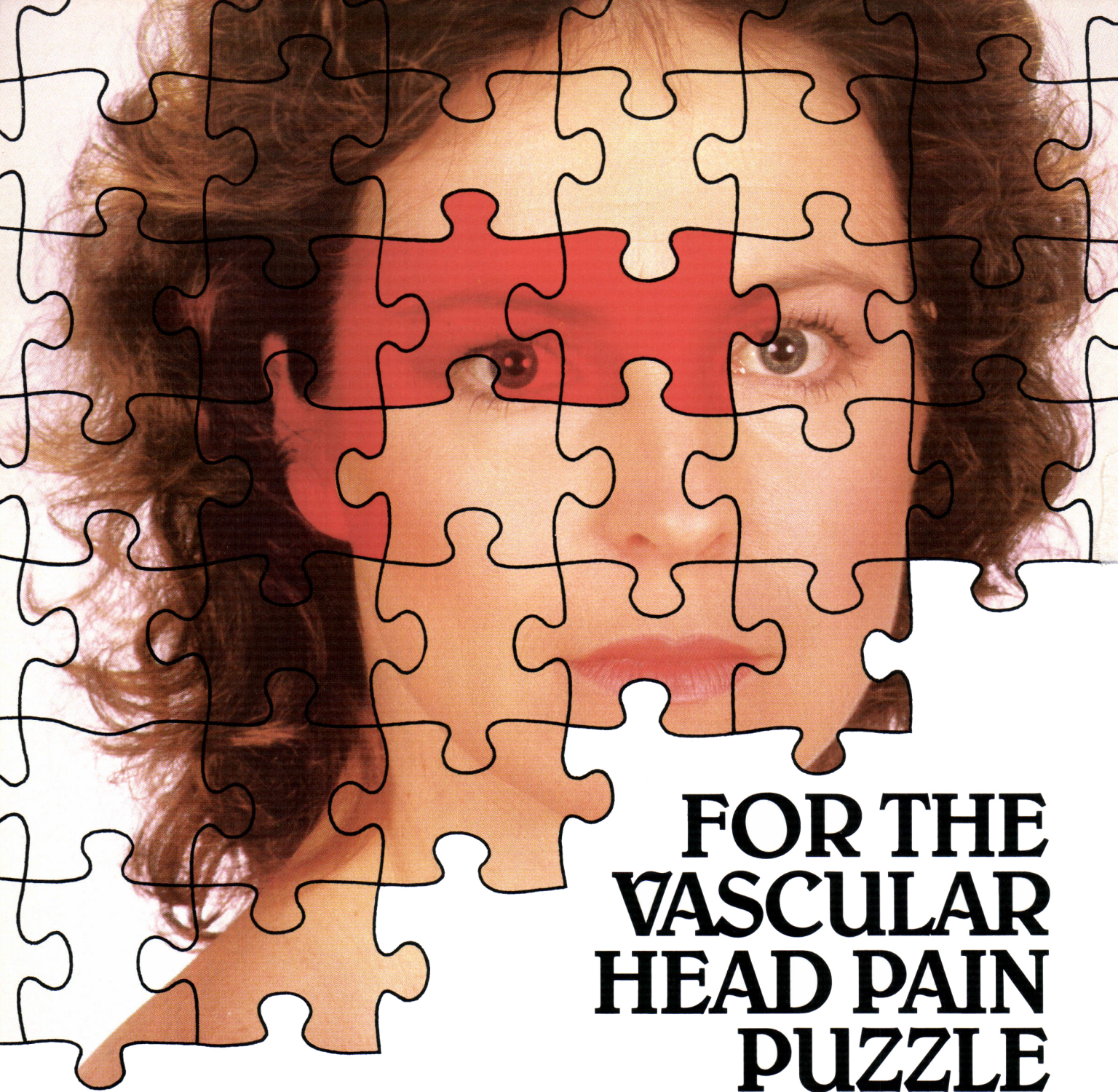
## The Official Journal of

The Canadian Neurological Society

The Canadian Neurosurgical Society

The Canadian Society of Clinical Neurophysiologists

The Canadian Association for Child Neurology





# FOR THE VASCULAR HEAD PAIN PUZZLE

**CAFERGOT<sup>®</sup>**  
To ABORT acute  
vascular headache

**SANDOMIGRAN<sup>®</sup> DS**  
PROPHYLAXIS for chronic  
recurring vascular headache

**SANDOZ<sup>®</sup>**  
▼▼▼

Complete headache therapy  
Sandoz Canada Inc., Dorval, Quebec H9R 4P5

Cafergot contains: ergotamine tartrate/caffeine <sup>®</sup> TM  
Sandomigran DS contains: pizotyline  
Full prescribing information available on request.  



**Editor/Rédacteur en chef** Robert G. Lee *Calgary*  
**Associate Editors/Rédacteurs associés** Yves Lamarre *Montreal* Harvey B. Sarnat *Calgary*  
**Founding Editor/Fondateur-rédacteur** Robert T. Ross *Winnipeg*  
**Book Review Editor/Rédacteur de critiques de livres** T. Peter Seland *Calgary*  
**Managing Editor/Adjoint administratif** Sally A. Gregg *Calgary*

**Editorial Board/Conseil Scientifique**

Albert J. Aguayo <i>Montreal</i>	André Olivier <i>Montreal</i>
Henry J.M. Barnett <i>London</i>	Donald Paty <i>Vancouver</i>
Larry Becker <i>Toronto</i>	Sidney J. Peerless <i>London</i>
Paul Bédard <i>Quebec</i>	Terry Picton <i>Ottawa</i>
George Ebers <i>London</i>	Jean Reiher <i>Sherbrooke</i>
Guy Geoffroy <i>Montreal</i>	Leo P. Renaud <i>Montreal</i>
William J. Logan <i>Toronto</i>	Matthew W. Spence <i>Halifax</i>
Morton Low <i>Vancouver</i>	John Stewart <i>Montreal</i>
John Murphy <i>Toronto</i>	Charles Tator <i>Toronto</i>
Thomas J. Murray <i>Halifax</i>	Simon Verret <i>Quebec</i>

Bryce Weir *Edmonton*

**Publications Committee/Comité de Rédaction**

John Wherrett <i>Toronto</i>	Warren Blume <i>London</i>
Terry Myles <i>Calgary</i>	John Tibbles <i>Halifax</i>

**The Official Journal of:/La Revue Officielle de:**

**The Canadian Neurological Society**

**La Société Canadienne de Neurologie**

President/Président — T. Peter Seland  
Secretary-Treasurer/ — William McCormick,  
Secrétaire-Trésorier Box 2148, Dickson Centre,  
Victoria General Hospital,  
Halifax, Nova Scotia  
B3H 2Y9

**The Canadian Society of Clinical Neurophysiologists**

**La Société Canadienne de Neurophysiologues Clinics**

President/Président — R. Gordon Blair  
Secretary-Treasurer/ — Werner J. Becker,  
Secrétaire-Trésorier 841 Centre Avenue East,  
Calgary, Alberta  
T2E 0A1

**The Canadian Neurosurgical Society**

**La Société Canadienne de Neurochirurgie**

President/Président — Ian Turnbull  
Secretary-Treasurer/ — Harold Hoffman  
Secrétaire-Trésorier Hospital for Sick Children  
555 University Avenue  
Suite 1504  
Toronto, Ontario  
M5G 1X1

**The Canadian Association for Child Neurology**

**L'Association Canadienne de Neurologie Pédiatrique**

President/Président — Kevin Farrell  
Secretary-Treasurer/ — Daniel Keene,  
Secrétaire-Trésorier Suite 208, 150 Montreal Road,  
Vanier, Ontario  
K1L 8H2

The Canadian Journal of Neurological Sciences is published quarterly by University of Calgary Press. The annual subscription rate is \$48.00 for Canada and the U.S.A.; \$48US elsewhere. Interns, Residents Pre- and Post-Doctoral Students \$20.00 per annum. Single copies \$15.00 each. All communications and subscriptions should be sent to the Editor, Canadian Journal of Neurological Sciences, Room 1496, Faculty of Medicine, University of Calgary, 3330 Hospital Drive N.W., Calgary, Alberta, Canada T2N 4N1. Telephone: (403) 220-3062. COPYRIGHT © 1988 by THE CANADIAN JOURNAL OF NEUROLOGICAL SCIENCES INC. No part of this journal may be reproduced in any form without the prior permission of The Canadian Journal of Neurological Sciences. Mailed under second class registration number 3307. Postage paid at Calgary, Alberta. This journal is indexed by *Index Medicus*, *Excerpta Medica* and *Current Contents* — *Clinical Practice and Life Sciences*.

Le Journal Canadien des Sciences Neurologiques est publié trimestriellement par les Presses de l'Université de Calgary. L'abonnement annuel est de \$48.00 pour le Canada et les États-Unis; \$48US ailleurs. Internes, résidents, fellows pré et post-doctoral: \$20.00 par an. Toutes les communications et les abonnements doivent être adressés à l'Éditeur, Journal des Sciences Neurologiques, chambre 1496, Faculté de Médecine, Université de Calgary, 3330 Hospital Drive N.W., Calgary, Alberta, T2N 4N1. Téléphone (403) 220-3062.

DROITS D'AUTEUR © 1988: THE CANADIAN JOURNAL OF NEUROLOGICAL SCIENCES INC. Aucune partie de ce Journal ne peut être reproduite, sous quelque forme que ce soit, sans la permission préalable du Journal Canadien des Sciences Neurologiques. Posté sous permis de second classe no 3307. Port payé à Calgary, Alberta. Le Journal est cité et indexé dans *Index Medicus*, *Excerpta Medica* et *Current Contents* — *Clinical Practice et Life Sciences*.

**Advertising representative/Représentant de publicité** Keith Health Care Communications,  
4953 Dundas St. W., Toronto, Ontario, Canada M9A 1B6 — (416) 239-1233

**Printer/Imprimeur** McAra Printing Limited, 105, 2507 - 12th Street N.E., Calgary, Alberta T2E 7L5

ISSN 0317 - 1671

# This news deserves the front page.

The first ever controlled release form of carbamazepine has been announced by Geigy Pharmaceuticals.

Tegretol CR (controlled release), a new and special form of tricyclic carbamazepine, was developed to provide better tolerability plus all the proven efficacy of conventional Tegretol. In fact new <sup>®</sup> Tegretol<sup>®</sup> CR can significantly reduce plasma concentration peaks in a B.I.D. dosage regimen.<sup>1,2</sup> This can result in a more stable pattern of cognitive functioning.<sup>3</sup> New Tegretol CR. Available in breakable 200 mg and 400 mg tablets for easier titration. For further information contact your Geigy representative.

News to act on. For initiating therapy or switching from conventional carbamazepine.

**Tegretol<sup>®</sup> CR.**  
carbamazepine.

MAAP  
LIFE

G 88010

**Geigy**  
Mississauga, Ontario  
L5N 2W5



Table of Contents

**SPECIAL FEATURES**

Novel Uses of EMG to Study Normal and Disordered Motor Control <i>Richard B. Stein</i> .....	95
Mechanisms of Action of Neural Grafts in the Limbic System <i>György Buzsaki and Fred H. Gage</i> .....	99
Idiopathic Parkinson's disease: Revised Concepts of Cognitive and Affective Status <i>Ann E. Taylor, J.A. Saint-Cyr and A.E. Lang</i> .....	106

**ORIGINAL ARTICLES**

Immunocytochemical Analysis of Intermediate Filaments in Human Ependymal Tumors <i>D. Izukawa and B. Lach</i> .....	114
Cerebral Metastases from Malignant Melanoma <i>Ivar M. Mendez and Rolando F. Del Maestro</i> .....	119
Hypomelanosis of Ito. Neurological Complications in 34 Cases <i>Ignacio Pascual-Castroviejo, Luisa López-Rodriguez, María de la Cruz Medina, Cipriano Salamanca-Maesso and Carmen Roche Herrero</i> .....	124
Cerebral Arteriovenous Malformations with Associated Arterial Aneurysms: Hemodynamic and Therapeutic Considerations <i>Douglas Kondziolka, Bruce J. Nixon, Pierre Lasjaunias, William S. Tucker, Karel TerBrugge and Sanford M. Spiegel</i> .....	130
Late Progression of Post-Encephalitic Parkinson's Syndrome <i>Donald B. Calne and Andrew J. Lees</i> .....	135
Parental Age and Birth Order in Alzheimer's disease: A Case-Control Study in the Saguenay-Lac-St-Jean Area (Quebec, Canada) <i>M. De Braekeleer, S. Froda, D. Gauthrin, H. Tetreault and D. Gauvreau</i> .....	139
Familial Alzheimer's Disease <i>A.D. Sadnovick, H. Tuokko, A. Horton, P.A. Baird and B.L. Beattie</i> .....	142
The Effects of Myoinositol on the Autonomic Neuropathy in the Streptozotocin Diabetic Rat - A Freeze Fracture Study <i>G. Monckton and H. Marusyk</i> .....	147
Calcification in a Recent Cerebral Infarct - Radiologic and Pathologic Correlation <i>J. Parisi, C. Place and S. Nag</i> .....	152
Jacob-Creutzfeldt Disease Associated with Wernicke Encephalopathy <i>S. Gaytan-Garcia, J.J. Gilbert, J.H.N. Deck and J.C.E. Kaufmann</i> .....	156
Magnetic Resonance Imaging of Meningio-Angiomatosis <i>Ruben Kuzniecky, Denis Melanson, Yves Robitaille and André Olivier</i> .....	161
In Memoriam: William Strathearn Keith (1902-1987) <i>Ross Fleming</i> .....	165

<b>CORRESPONDENCE</b> .....	167
-----------------------------	-----

<b>BOOK REVIEWS</b> .....	169
---------------------------	-----

<b>CALENDAR OF EVENTS</b> .....	171
---------------------------------	-----

<b>ADVERTISERS INDEX</b> .....	xxv
--------------------------------	-----


<b>CANADIAN ASSOCIATION OF NEUROPATHOLOGISTS - Abstracts</b> .....	172
--	-----

<b>XXIIIrd CANADIAN CONGRESS OF NEUROLOGICAL SCIENCES - Program and Abstracts</b> .....	179
---	-----



*To the parkinsonian patient,  
the little things in life make all the difference*



*Add*  
**PARLODEL<sup>®</sup>**   
(bromocriptine mesylate)  
*Because quality  
of life is the issue*

For brief prescribing information see page xx



# A bright outlook for more epileptic patients

**Depakene**<sup>\*</sup>  
valproic acid

## to control seizures in more patients than ever...

Depakene, a broad-spectrum anticonvulsant, is the drug of choice in absence seizures *with or without tonic-clonic manifestations*.<sup>1,2</sup> Depakene is highly effective in *generalized tonic-clonic seizures*.<sup>3,4</sup>

## ...with fewer side effects than most anticonvulsants

Unlike phenobarbital or phenytoin, Depakene is rarely associated with impaired performance or behavior problems.<sup>5,6</sup> In a year-long survey of adverse reactions to anticonvulsants,<sup>7</sup> sodium valproate, alone or in combination, had one of the lowest incidences of side effects.

**Epival**<sup>\*</sup>  
divalproex sodium

## the same effectiveness as Depakene, with less G.I. upset

Epival, a new form of valproate, is an enteric-coated tablet which is just as effective as Depakene, while minimizing G.I. upset.<sup>8</sup> Thus, Epival helps reduce the risk of poor compliance caused by gastric irritation.





Quality Health Care  
Worldwide  
1888-1988

PHARMACEUTICAL PRODUCTS DIVISION  
ABBOTT LABORATORIES, LIMITED  
MONTREAL, CANADA



\*TM  
©Abbott Laboratories, Limited

For brief prescribing information see page xviii

# The Added Value Of

## Benefit from the Added Value of Nicolet Systems

“Tough buyers” realize that getting the lowest price does not guarantee the best value. The added value of our unique Nicolet Systems provides you with the security of:

- Expert training for all Nicolet systems
- Customer service response in two hours or less
- Applications support from our staff of Ph.D.s
- Seminar series featuring world-renowned speakers
- Complete financial packages and alternatives
- Our 18 years of experience in problem-solving instrumentation

## Nicolet Pathfinder MEGA

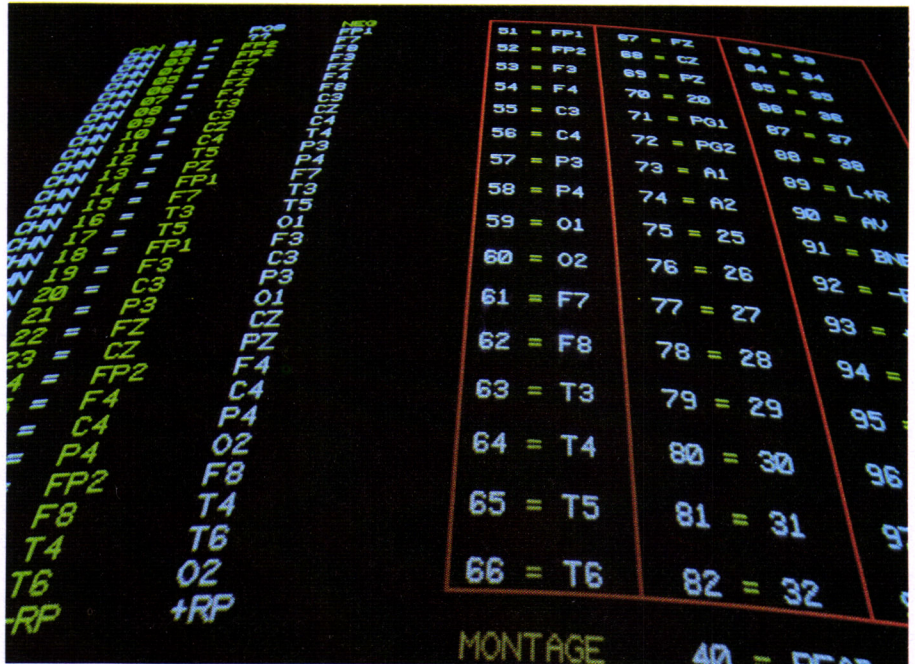
- Internal 8, 16 or 32 channel EP/EEG amplifiers
- Flexible test protocols
- 1.28 megabytes of memory for powerful data analysis
- Topographic Mapping for all clinical/research investigations
- O.R./I.C.U. monitoring for a variety of surgical procedures
- Fully programmable system via MECOL and/or FORTRAN 77



## Nicolet BEAM®

- Benchmark data base from childhood to geriatric

© Nicolet Instrument Corporation



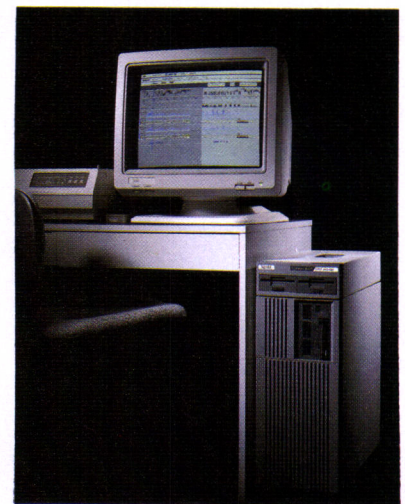
## Introducing the NEW Nicolet Pathfinder MEGA

- Accurate quantification of patient abnormalities
- Complete physician and technologist training
- Data analysis that matches your pace



## Nicolet Electroencephalographs

- 18 or 21 channels
- “Short Menu” allows rapid selection of filters and sensitivity
- Programmable measurement sequences allow automated recordings
- Tailor each montage to your unique requirements



## Nicolet Expert Sleep/Wake™ Analyzer

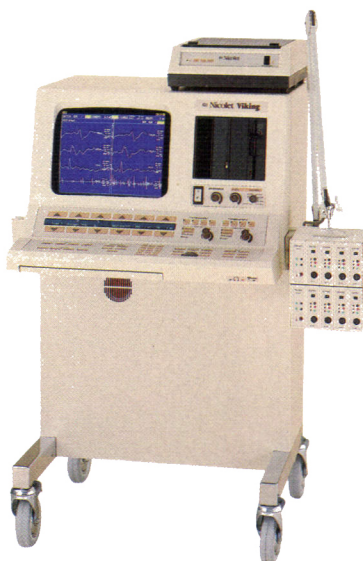
- Cost-effectively analyzes up to four patients simultaneously
- Permits direct validation of analyzed results with raw data
- Performs adaptable analysis of patient's data to accommodate individual variability
- Facilitates detailed analysis of less frequent or short duration events (such as micro-arousals)

# Unique Nicolet Systems



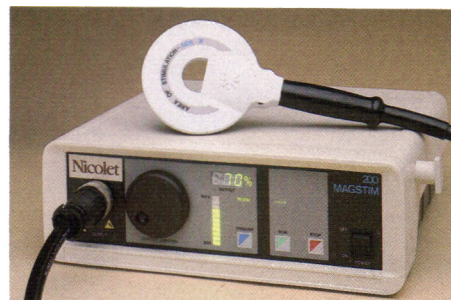
## Nicolet NIM-2™

- Provides the surgeon with an intraoperative tool to locate and identify a nerve directly in the surgical field
- Continuously monitors EMG activity from muscles innervated by the particular nerve
- Acoustically alerts the surgeon when the particular nerve has been activated
- Contains unique technology to minimize the effects of electrocautery interference



## Nicolet Viking

- Unique quantitative capabilities
- Expandable to 8 channels of simultaneous EMG/EP
- Comprehensive, high-quality, integrated reports
- Protocols easily customized by individual users



## Nicolet Magnetic Stimulator

- Fast—no electrodes or skin preparation required
- Painless—ideal for children
- Easily stimulates deep structures
- Extensive safety features for both operator and patient
- Easily interfaced to other Nicolet instruments

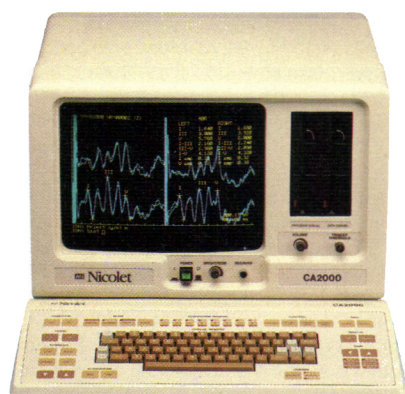


## Electrodes and Supplies

In addition to our complete line of quality systems, Nicolet offers a full line of EEG/EMG/EP electrodes and accessories. For our latest Supplies Catalog and specific brochures on any of our unique Nicolet Systems simply call **TOLL FREE**

Ontario: 1-800-268-2058  
 Eastern/Western Canada: 1-800-387-3385  
 In Quebec Call: 514-678-2134

*Nicolet Instrument Canada Inc.*  
 1-1200 Aerowood Drive  
 Mississauga, Ontario  
 L4W 2S7



## Nicolet CA2000

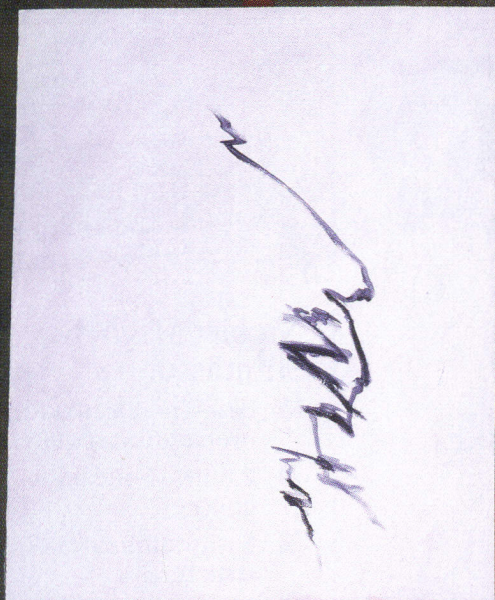
- Comprehensive Evoked Potential and EMG testing
- User-friendly menu operation with 72 sets of user-defined parameters
- Large color screen with optional remote monitors
- Nicolet-engineered to the highest O.R./I.C.U. standards



## Nicolet Compact Four

- All Auditory, Visual and Somatosensory Evoked Potentials
- Complete range of EMG protocols
- Full-feature ENG and ENOG testing
- ERG and EOG for total visual testing

# Productive Antispastic



Without Therapy.



With Oversedation.

## Information for Authors

The Canadian Journal of Neurological Sciences publishes original articles in the clinical and basic neurosciences. Manuscripts are considered for publication with the understanding that, except for identified review articles, they have not been published elsewhere except in abstract form and are not under simultaneous consideration by another publication. Manuscripts should be submitted to:

The Editor  
Canadian Journal of Neurological Sciences  
Faculty of Medicine,  
University of Calgary  
3330 Hospital Drive N.W.  
Calgary, Alberta T2N 4N1

Manuscripts and all illustrations should be submitted in triplicate. Papers will be accepted in English or French. All papers should be accompanied by an abstract or a résumé of approximately 150 words on a separate page, preferably in both languages, although the Journal will provide the translation if requested. All manuscripts should be double spaced throughout, including references and legends for illustrations. Margins of at least 25 mm should be left on all sides.

For detailed instructions regarding style and layout, authors should refer to "Uniform requirements for manuscripts submitted to biomedical journals". Copies of this document may be obtained by writing to the Journal office, but the main points will be summarized here. Articles should be subdivided under conventional headings of "introduction", "methods and materials", "results" and "discussion" but other headings and subheadings will be considered if more suitable for a particular manuscript. A title page should identify the title of the article, authors, name of institution(s) from which the work originated, and the address and telephone number of the author to whom communications should be addressed. Pages of text should be numbered consecutively. Acknowledgements, including recognition of financial support, should be typed on a separate page at the end of the text.

**References** are to be numbered in the order of citation in the text. Those cited only in tables or in legends for illustrations are numbered in accordance with a sequence established by the first identification in the text of a particular table or illustration. Titles of journals should be abbreviated according to the style used in Index Medicus. References should be complete including the names of the first three authors followed by "et al"

if there are more than three authors, full title, year of publication, volume number, and inclusive pagination for journal articles. Book or chapter references should also include the place of publication and name of the publisher. Examples of correct forms of references follow:

### Journals

Poirier LJ, Filion M, Larochelle L, et al. Physiopathology of experimental parkinsonism in the monkey. *Can J Neurol Sci* 1975; 2: 255-263

### Chapter in a book

McGeer PL, McGeer EG. Amino acid neurotransmitters. In: Siegel GJ, Albers RW, Agranoff BW, Katzman R, eds. *Basic Neurochemistry*. Boston: Little, Brown & Co, 1981: 233-254

**Illustrations** should be high quality glossy black-and-white photographic prints, preferably 127 x 173 mm (5 x 7"). Original artwork and radiographs should not be submitted. The additional cost of colour illustration must be borne by the author; quotations are available upon request from the Journal office. All figures should be identified on the back with the author's name and figure number. Letters and arrows applied to the figures to identify particular findings should be professional appliques suitable for publication. Photomicrographs should include a calibration bar with the scale indicated on the figure or in the legend. Legends for illustrations should be typed on a separate page from the illustrations themselves.

**Tables** should each be on a separate page and be identified with the title or heading. Particular care should be taken in the preparation of tables to ensure that the data are presented in the most clear and precise format. Each column should have a short or abbreviated heading. Place explanatory matter in footnotes, not in the heading. Do not submit tables as photographs.

The SI system (système international d'unités) should be used in reporting all laboratory data, even if originally reported in another system. Temperatures are reported in degrees Celsius. Other measurements should be reported in the metric system. English language text may use either British or American spelling, but should be consistent throughout.

**Review articles** on selected topics also are published by the Journal. These are usually invited, but unsolicited reviews will be considered. It is suggested that authors intending to submit reviews contact the Editor in advance.

**Letters to the Editor** are welcome. These should be limited to two double-spaced pages and may include one illustration and a maximum of four references.

# Therapy Is a Fine Art.

Successful rehabilitation requires both spinal spasticity control and alertness. And that's why Lioresal® stands out.

Lioresal can control painful flexor spasms, but unlike diazepam, oversedation is rarely a problem (1,2,3).

And it effectively stops most flexor spasms overnight, so patients can sleep naturally.

Now there's two strengths:

Lioresal 10mg to initiate therapy and Lioresal D.S. 20mg for maintenance.

Considering that Lioresal has an excellent long-term safety profile, it may be your most effective choice for long-term control of spinal spasticity.

(baclofen)  
**Lioresal**

Lioresal 10mg (initiation therapy)  
Lioresal D.S. 20mg (maintenance therapy)

**Geigy**  
Mississauga, Ontario  
LSN 2WS

PAR  
0377  
G-5139



**With Lioresal.**

For brief prescribing information see page xxiv

## Information aux Auteurs

Le Journal Canadien des Sciences Neurologiques publie des articles originaux dans les sciences neurologiques, cliniques et fondamentales. Les manuscrits ne sont considérés pour publication qu'à la condition expresse, à l'exception des articles de revue clairement identifiés comme tel, qu'ils n'aient pas été publiés ailleurs, sauf sous forme de résumé et qu'ils ne soient pas sous considération simultanée par un autre journal. Les manuscrits doivent être soumis à:

L'Éditeur  
Journal Canadien des Sciences Neurologiques,  
Faculté de Médecine,  
Université de Calgary,  
3330 Hospital Drive, N.W.,  
Calgary, Alberta T2N 4N1

Les manuscrits et toutes les illustrations doivent être soumis en triplicata. Les articles seront acceptés en français ou en anglais. Tous les articles doivent être accompagnés d'un résumé d'environ 150 mots, sur page séparée, préférablement dans les deux langues, quoique le Journal puisse fournir cette traduction sur requête. Les manuscrits doivent être dactylographiés complètement à double interligne y compris les références et les légendes pour illustrations. Des marges d'au moins 25 mm doivent être laissées de tous les côtés.

Pour les conseils plus détaillés sur le style et la présentation du texte, les auteurs doivent se référer au texte intitulé "Règlements uniformes pour les manuscrits soumis aux journaux biomédicaux". On peut obtenir une copie de ce document en écrivant au bureau du Journal, mais en voici les principaux points: Les articles doivent être présentés selon le plan habituel: "Introduction", "Matériel et méthodes", "Résultats" et "Discussion", mais il est possible d'employer d'autres titres ou sous-titres si nécessaire pour un manuscrit en particulier. Sur une page titre séparée on doit identifier le titre de l'article, les auteurs, les institutions d'origine le travail, ainsi que l'adresse et le numéro de téléphone de l'auteur à qui devront être adressées les communications. Les remerciements, incluant ceux pour l'appui financier, doivent être dactylographiés sur page séparée à la fin du texte. Les références doivent être numérotées dans l'ordre où elles sont citées dans le texte. Celles qui sont citées seulement dans les tableaux ou légendes d'illustrations sont numérotées selon la séquence établie par la première identification dans le texte de ces tableaux ou illustrations particulières. Les titres des Journaux doivent être abrégés selon le style utilisé dans Index Medicus. Les références doivent être complètes, incluant le nom des trois premiers auteurs suivis de "et al", s'il y a plus de trois auteurs, le titre complet, l'année de publication, le

numéro du volume et les premières et dernières pages de l'article. Les références aux livres et chapitres de livres doivent aussi inclure le lieu de la publication et le nom de la maison d'édition. Les exemples corrects suivants peuvent être utilisés:

### Journaux

Poirier LJ, Filion M, Larochelle L, et al. Physiopathology of experimental parkinsonism in the monkey. *Can J Neurol Sci* 1975; 2: 255-263

### Chapitre de livre

McGeer PL, McGeer EG, Amino acid neurotransmitters. In: Siegel GJ, Albers RW, Agranoff BW, Katzman R, eds. *Basic Neurochemistry*. Boston: Little, Brown & Co, 1981: 233-254

Les **illustrations** doivent être sur papier brillant de haute qualité et imprimés en blanc et noir, préférablement dans 127 x 173 mm (5 x 7"). Les illustrations et photographies originales ne doivent pas être soumises. Le coût supplémentaire des illustrations en couleur revient entièrement à l'auteur; les coûts détaillés peuvent être obtenus directement au bureau du Journal. Il faut identifier toutes illustrations en inscrivant au dos le nom de l'auteur et le numéro. Toutes lettres ou flèches appliquées aux illustrations pour identifier un aspect particulier doivent être de qualité professionnelle. Les photomicrographies doivent inclure une barre de calibration dont l'échelle est mentionnée dans la légende. Les légendes des illustrations doivent être dactylographiées sur une page séparée de celles-ci.

Les **tableaux** doivent être sur des pages séparées et être identifiés avec titre. On doit prendre un soin particulier dans la préparation de ces tableaux afin d'assurer que les données soient présentées avec le format le plus clair et le plus précis possible. Chaque colonne doit avoir un court titre. Les explications doivent être placées en dessous du tableau et non en sous-titre. Un tableau ne doit pas être soumis sous forme de photographie.

On doit employer le système international d'unités (SI) pour toutes données de laboratoire, même si celles-ci sont originellement présentées dans un autre système. Les températures doivent être citées en degrés Celcius. Les autres données doivent utiliser le système métrique. Les textes en anglais peuvent utiliser l'orthographe anglais ou américain, mais cet usage doit être constant.

Le Journal publie également des **articles de revue** sur des sujets sélectionnées. Ces articles sont généralement sur invitation, mais, à l'occasion, une revue non sollicitée peut être acceptée. Il serait préférable que les auteurs ayant l'intention de soumettre une telle revue contactent d'abord l'Éditeur.

Nous accueillons les **lettres à l'Éditeur**. Celles-ci doivent se limiter à deux pages, double interligne et peuvent contenir une seule illustration et ne citer qu'un maximum de quatre références.

# Une nouvelle à publier à la une.

mpor-  
bab-  
tive:

## Geigy annonce le lancement de la première formulation de carbamazépine à libération contrôlée.

Tegretol CR (libération contrôlée) une nouvelle formulation de carbamazépine tricyclique, a été mis au point pour procurer une meilleure tolérance en plus de toute l'efficacité qu'on reconnaît au Tegretol conventionnel. De fait, le nouveau Tegretol<sup>®</sup> CR diminue significativement les pics de concentration plasmatique avec un régime posologique b.i.d.<sup>1,2</sup> ce qui peut produire un modèle plus stable de fonction cognitive.<sup>3</sup> Nouveau Tegretol CR disponible en comprimés sécables à 200 mg et 400 mg pour faciliter le titrage. Contactez votre représentant Geigy pour de plus amples renseignements.

com  
lais  
tés e

ête

ob

si

pa

vo

vo

dre

on so

Une nouvelle dont il faut tenir compte.  
Pour amorcer le traitement ou le substituer à la  
carbamazépine conventionnelle.

**Tegretol<sup>®</sup> CR.**  
carbamazépine.

**Geigy**

Mississauga, Ontario  
L5N 2W5

PAR  
L'ÉDIT  
G-88010




*A different quality of calm*

*for the child with  
Attention Deficit Disorder*

**Cylert\***  
PEMOLINE

Cylert has helped hyperactive children concentrate, become more attentive and work better<sup>1,2</sup> – but did not suppress their spontaneity.<sup>2</sup> Unlike methylphenidate, Cylert has not shown any significant effect on blood pressure.<sup>3</sup> And if there is a concern about possible abuse, Cylert is a wise choice. There have been no published reports of abuse or dependence and, unlike other stimulants, Cylert is not on the list of controlled drugs.<sup>4</sup> With fewer adverse effects and less risk of addiction, Cylert can offer the hyperactive child a different quality of calm.

 PHARMACEUTICAL PRODUCTS DIVISION  
ABBOTT LABORATORIES, LIMITED  
MONTREAL, CANADA

\*TM

PAAB

For brief prescribing information see page xxiii

# ANOTHER UNEVENTFUL DAY.

## DILANTIN

Extended Phenytoin Sodium Capsules, U.S.P.  
100 mg  
ANTICONSULSANT

### INDICATIONS

Dilantin is indicated for the control of generalized tonic-clonic (grand mal) seizures and complex partial (psychomotor) seizures.

### CONTRAINDICATIONS

Dilantin is contraindicated in those patients with a history of hypersensitivity to hydantoin products.

### WARNINGS

Abrupt withdrawal of phenytoin in epileptic patients may precipitate status epilepticus.

Phenytoin is not indicated in seizures due to hypoglycemia or other causes which may be immediately identified and corrected.

Phenytoin metabolism may be significantly altered by the concomitant use of other drugs such as:

**A.** Barbiturates may enhance the rate of metabolism of phenytoin. This effect, however, is variable and unpredictable. It has been reported that in some patients the concomitant administration of carbamazepine resulted in an increased rate of phenytoin metabolism.

**B.** Coumarin anticoagulants, disulfiram, phenylbutazone, and sulfaphenazole may inhibit the metabolism of phenytoin, resulting in increased serum levels of the drug. This may lead to an increased incidence of nystagmus, ataxia, or other toxic signs.

**C.** Isoniazid inhibits the metabolism of phenytoin so that with combined therapy, patients who are slow acetylators may suffer from phenytoin intoxication.

**D.** Tricyclic antidepressants in high doses may precipitate seizures, and the dosage of phenytoin may have to be adjusted accordingly.

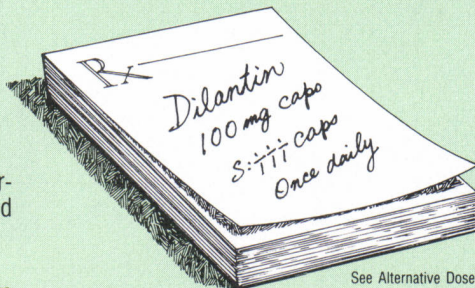
**Usage in Pregnancy:** The effects of Dilantin in human pregnancy and nursing infants are unknown.

The prescribing physician will have to determine the risk/benefit in treating or counselling epileptic women of childbearing potential.

### PRECAUTIONS

The liver is the chief site of biotransformation of phenytoin, patients with impaired liver function may show early signs of toxicity. Elderly patients or those who are gravely ill may show early signs of toxicity.

A small percentage of individuals who have



See Alternative Dose

been treated with phenytoin have been shown to metabolize the drug slowly. Slow metabolism may be due to limited enzyme availability and lack of induction; it appears to be genetically determined.

Phenytoin has been associated with reversible lymph node hyperplasia. If lymph node enlargement occurs in patients on phenytoin, every effort should be made to substitute another anticonvulsant drug or drug combination.

Drugs that control generalized tonic-clonic (grand mal) seizures are not effective for absence (petit mal) seizures. Therefore, if both conditions are present, combined drug therapy is needed.

Hyperglycemia, resulting from the drug's inhibitory effect on insulin release, has been reported. Phenytoin may also raise the blood sugar level in persons already suffering from hyperglycemia.

### ADVERSE REACTIONS

**Central Nervous System:** The most common manifestations encountered with phenytoin therapy include nystagmus, ataxia, slurred speech, and mental confusion. Dizziness, insomnia, transient nervousness, motor twitchings, and headache have also been observed. These side effects may disappear with continuing therapy at a reduced dosage level.

**Gastrointestinal System:** Phenytoin may cause nausea, vomiting, and constipation. Administration of the drug with or immediately after meals may help prevent gastrointestinal discomfort.

**Integumentary System:** Dermatological manifestations sometimes accompanied by fever have included scarlatiniform or morbilliform rashes.

**Hemopoietic System:** Hemopoietic complications, some fatal, have occasionally been reported in association with administration of phenytoin. These have included thrombocytopenia, leukopenia, granulocytopenia, agranulocytosis, and pancytopenia.

**Other:** Gingival hyperplasia occurs frequently; this incidence may be reduced by good oral hygiene including gum massage, frequent brushing and appropriate dental care. Polyarthropathy and hirsutism occur occasionally. Hyperglycemia has been reported. Toxic hepatitis, liver damage, and periarteritis nodosa may occur and can be fatal.

### MANAGEMENT OF OVERDOSE

The mean lethal dose in adults is estimated to be 2 to 5 grams. The cardinal initial symptoms are nystagmus, ataxia and dysarthria. The patient then becomes comatose, the pupils are unresponsive and hypotension occurs. Death is due to respiratory depression and apnea. Treatment is nonspecific since there is no known antidote. First, the stomach should be emptied. If the gag reflex is absent, the airway should be supported. Oxygen, vasopressors and assisted ventilation may be necessary for central nervous system, respiratory and cardiovascular depression. Finally, hemodialysis can be considered since phenytoin is not completely bound to plasma proteins.

### DOSAGE AND ADMINISTRATION

Dosage should be individualized to provide maximum benefit. In some cases, serum blood level determinations may be necessary for optimal dosage adjustments — the clinically effective serum level is usually 10-20 mcg/mL.

**Adult Dose:** Patients who have received no previous treatment may be started on one 100 mg Dilantin Capsule three times daily and the dose then adjusted to suit individual requirements.

**Pediatric Dose:** Initially, 5 mg/kg/day in two or three equally divided doses, with subsequent dosage individualized to a maximum of 300 mg daily. A recommended daily maintenance dosage is usually 4 to 8 mg/kg. Children over 6 years old may require the minimum adult dose (300 mg/day). Pediatric dosage forms available include a 30 mg Capsule, a 50 mg palatably flavoured Infatab, or an oral suspension form containing 30 or 125 mg of Dilantin in each 5 mL.

**Alternative Dose:** Once-a-day dosage for adults with 300 mg of Dilantin may be considered if seizure control is established with divided doses of three 100 mg Capsules daily.

### HOW SUPPLIED

Dilantin 100 mg Capsules; in bottles of 100 & 1000.

Complete prescribing information available upon request.

*Dilantin. Start with it. Stay with it.*

**PARKE-DAVIS**

Parke-Davis Canada Inc., Scarborough, Ontario



\*Reg. T.M. Parke, Davis & Company, Parke-Davis Canada Inc., auth. user.



See IBC



# Counterpoint

100...  
to intrigue against or  
counte...  
counte...  
coun-ter-point \ 'kaunt-er-'póint\ n 1 a : one  
or more independent melodies added above or  
below a given melody 2 a : a complementing  
or contrasting item ; OPPOSITE b : use of contrast  
or interplay of elements in a work of art  
coun-ter-poise \ -,póiz\ vt [ME *countrepe*...  
... more at ...

## The next generation of EMG technology, today.

Counterpoint represents the start of a new era in electromyograph technology.

A system so powerful and flexible it challenges the very definition of what EMG technology can accomplish.

Imagine. One instrument that virtually does it all. And does it fast and accurately. Thanks in large part to the quality of our electrodes and amplifiers. And because at the heart of the system we've incorporated the most advanced signal processor in the industry.

Combine that power with an impressive range of software applications, and you've got a system ready to handle the most demanding jobs, from spontaneous activity and motor nerve conduction velocity to single fiber EMG and power spectrum analysis. Plus more.

And because we know how important it is for you to concentrate on your patient - we've done everything in our power to make Counterpoint as easy to use as possible. The software is user-friendly, footswitch allows you to conduct investigations - hands free - and the electrode arm can be connected to either side of the control panel for comfortable use for everyone.

But the most surprising news is that Counterpoint is totally open to your needs. Data storage and filing can take place in an IBM Personal System/2 or AT compatible environment, so you'll never be left behind by the advent of new technologies or software developments.

Counterpoint masterfully orchestrates all the possibilities in one system. It promises to satisfy the most demanding EMG specialists in the field, and at the same time, makes your job a whole lot easier. And after all, isn't that the point?

Counterpoint from Dantec. Advanced technology in harmony with man.

Please call or write for more information.

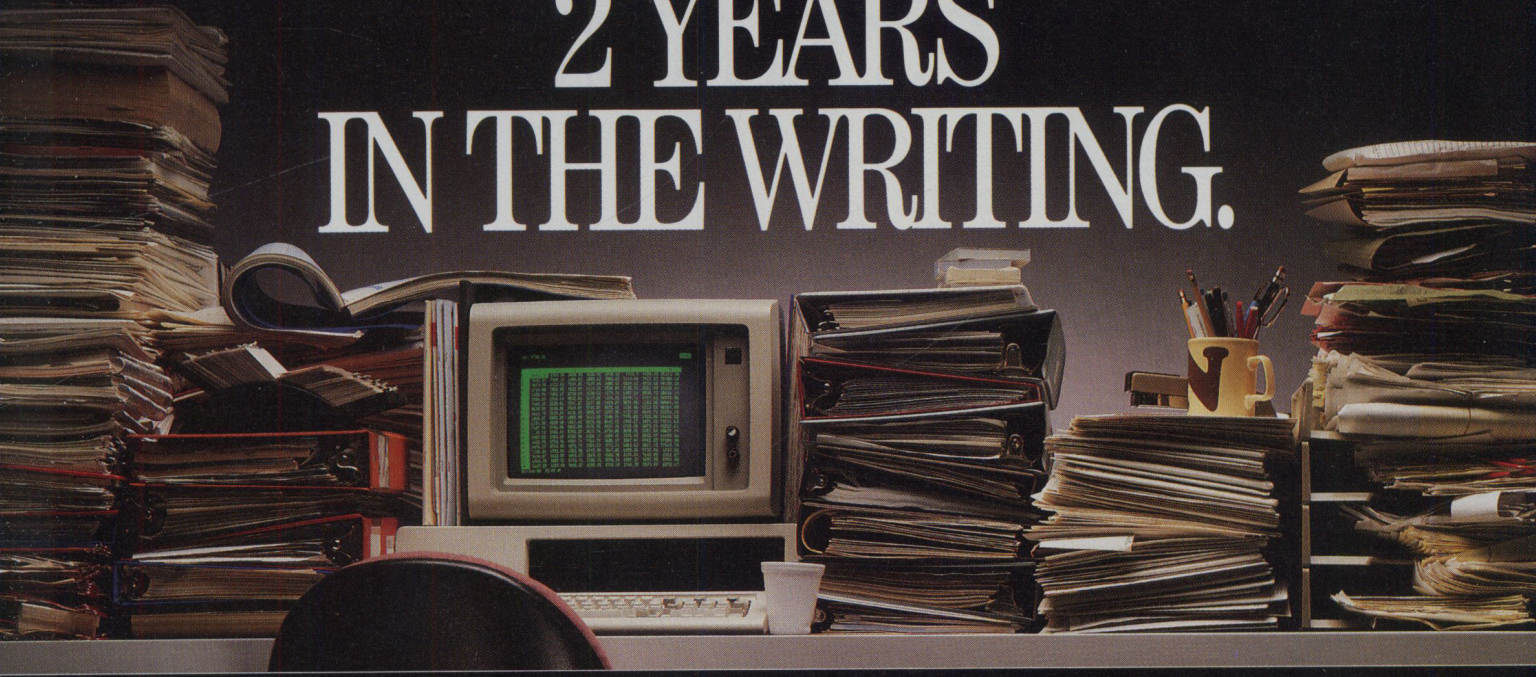


## DANTEC

Electromedical and  
Scientific Equipment Ltd.  
140 Shorting Road  
Scarborough, Ontario, M1S 3S6  
(416) 298-2091



# 2 YEARS IN THE WRITING.



# 2 SECONDS IN THE READING.

"Mersyndol was significantly more effective in decreasing pain, than either acetaminophen and codeine or placebo."<sup>1</sup>

After 2 years and hundreds of headaches there's now clear clinical evidence of the outstanding relief patients have been telling us about for years: "The analgesic superiority of Mersyndol

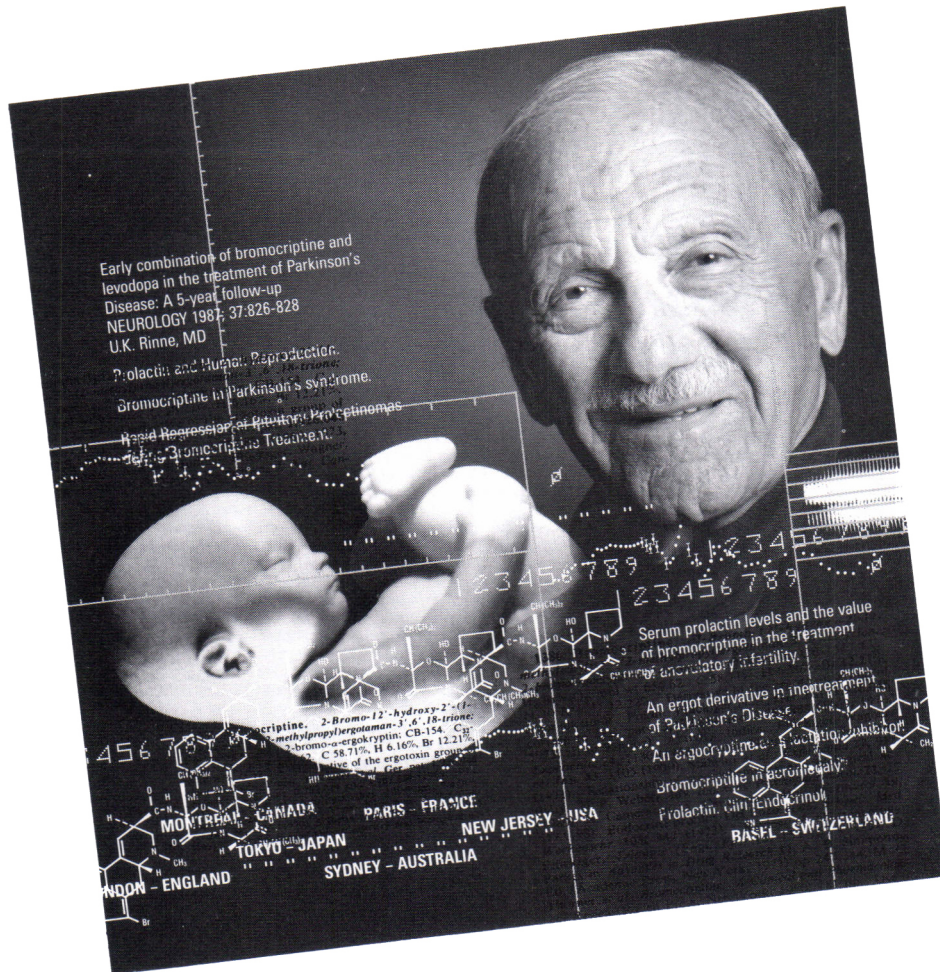
was observed in all headache classes..."<sup>2</sup> tested. You could review the entire report for yourself.

But your patients are the best proof. And that's where Mersyndol headache relief speaks volumes.

ANALGESIC  
**Mersyndol**<sup>®</sup>  
WITH CODEINE TABLETS  
(Acetaminophen - Codeine - Doxylamine)

AN EXTRA INGREDIENT. FOR EXTRA STRONG HEADACHE RELIEF.

For brief prescribing information see page xxvi



## A CELEBRATION OF EXPERIENCE

Parlodel celebrates a milestone publication. Combined L-Dopa and Bromocriptine Therapy for Parkinson's Disease\* is study number 5,000 for Parlodel. That's more than one original publication per day for over twelve years.

Why this unprecedented long-term interest? From its initial indication for suppression of postpartum lactation, to its current use in treating hyperprolactinemic infertility and Parkinson's disease, the wide therapeutic potential of Parlodel continues to fire the curiosity of physicians and medical researchers.

In turn, their experience has made Parlodel one of the best documented products available world-wide.

**PARLODEL**<sup>®</sup>  
(bromocriptine mesylate)

\*Robertson HA, Robertson GS  
Combined L-Dopa and Bromocriptine Therapy for Parkinson's Disease: A Proposed Mechanism of Action.  
Clinical Neuropharmacology 1987; Vol. 10, No. 4:384-7.

Availability: Tablets each containing 2.5 mg bromocriptine, as mesylate, available in bottles of 100.  
Capsules each containing 5 mg bromocriptine, as mesylate, available in bottles of 100.

Product monograph available to physicians and pharmacists upon request.

Sandoz Canada Inc., Dorval, Quebec H9R 4P5





# Depakene<sup>®</sup> Epival<sup>®</sup>

**ACTION** Valproic acid and divalproex sodium are chemically-related anticonvulsants. Although their mechanism of action has not yet been established, it has been suggested that their activity is related to increased brain levels of gamma-aminobutyric acid (GABA). The effect on the neuronal membrane is unknown. Epival (divalproex sodium) dissociates into valproic acid in the gastrointestinal tract.

Peak serum levels of valproic acid occur in 3 to 4 hours. The serum half-life ( $t_{1/2}$ ) of valproic acid is typically in the range of 6 to 16 hours. Half-lives in the lower part of the above range are usually found in patients taking other anti-epileptic drugs. A slight delay in absorption occurs when the drug is administered with meals but this does not affect the total absorption. Valproic acid is rapidly distributed throughout the body and the drug is strongly bound (90%) to human plasma proteins. Increases in dose may result in decreases in the extent of protein-binding and variable changes in valproic acid clearance and elimination.

The therapeutic plasma concentration range is believed to be from 50 to 100  $\mu\text{g/mL}$ . Occasional patients may be controlled with serum levels lower or higher than this range. A good correlation has not been established between daily dose, serum level and therapeutic effect.

Elimination of valproic acid and its metabolites occurs principally in the urine, with minor amounts in the feces and expired air. Very little unmetabolized parent drug is excreted in the urine. The principal metabolite formed in the liver is the glucuronide conjugate.

See **WARNINGS** section regarding statement on fatal hepatic dysfunction.

**INDICATIONS AND CLINICAL USE** Sole or adjunctive therapy in the treatment of simple or complex absence seizures, including petit mal; useful in primary generalized seizures with tonic-clonic manifestations. May also be used adjunctively in patients with multiple seizure types which include either absence or tonic-clonic seizures.

In accordance with the International Classification of Seizures, simple absence is defined as a very brief clouding of the sensorium or loss of consciousness (lasting usually 2-15 seconds) accompanied by certain generalized epileptic discharges without other detectable clinical signs. Complex absence is the term used when other signs are also present.

**CONTRAINDICATIONS** Should not be administered to patients with hepatic disease or significant dysfunction. Contraindicated in patients with known hypersensitivity to the drug.

**WARNINGS** Hepatic failures resulting in fatalities has occurred in patients receiving DEPAKENE<sup>®</sup> (valproic acid). These incidences usually have occurred during the first six months of treatment with DEPAKENE<sup>®</sup> (valproic acid). A recent survey study of valproate use in the United States in nearly 400,000 patients between 1978 and 1984, has shown that children under two years of age who received the drug as part of multiple anticonvulsant therapy were at greatest risk (nearly 20-fold increase) of developing fatal hepatotoxicity. These patients typically had other medical conditions such as congenital metabolic disorders, mental retardation or organic brain disease, in addition to severe seizure disorders. The risk in this age group decreased considerably in patients receiving valproate as monotherapy. Similarly, patients aged 3 to 10 years were at somewhat greater risk if they received multiple anticonvulsants than those who received only valproate. Risk generally declined with increasing age. No deaths have been reported in patients over 10 years of age who received valproate alone.

If DEPAKENE<sup>®</sup> (valproic acid) is to be used in children two years old or younger, it should be used with extreme caution and as a sole agent. The benefits of seizure control should be weighed against the risk. Serious or fatal hepatotoxicity may be preceded by non-specific symptoms such as loss of seizure control, malaise, weakness, lethargy, anorexia, and vomiting. Patients and parents should be instructed to report such symptoms. Because of the non-specific nature of some of the early signs, hepatotoxicity should be suspected in patients who become unwell, other than through obvious cause, while taking Epival or Depakene.

Liver function tests should be performed prior to therapy and at frequent intervals thereafter especially during the first 6 months. However, physicians should not rely totally on serum biochemistry since these tests may not be abnormal in all instances, but should also consider the results of careful interim medical history and physical examination. Caution should be observed in patients with a prior history of hepatic disease. Patients with various unusual congenital disorders, those with severe seizure disorders accompanied by mental retardation, and those with organic brain disease may be at particular risk.

In high-risk patients, it might also be useful to monitor serum fibrinogen and albumin for decrease in concentrations and serum ammonia for increases in concentration. If changes occur, the drug should be discontinued. Dosage should be titrated to and maintained at the lowest dose consistent with optimal seizure control.

The drug should be discontinued immediately in the presence of significant hepatic dysfunction, suspected or apparent. In some cases, hepatic dysfunction has progressed in spite of discontinuation of the drug. The frequency of adverse effects particularly elevated liver enzymes may increase with increasing dose. Therefore, the benefit gained by improved seizure control by increasing the dosage must be weighed against the increased incidence of adverse effects sometimes seen at higher dosages.

**Use in Pregnancy:** According to recent reports in the medical literature, valproic acid may produce teratogenicity in the offspring of women receiving the drug during pregnancy. The incidence of neural tube defects in the

fetus may be increased in mothers receiving valproic acid during the first trimester of pregnancy. Based upon a single report, it was estimated that the risk of valproic acid exposed women having children with spina bifida is approximately 1.2%. This risk is similar to that which applies to non-epileptic women who have had children with neural tube defects (encephaly and spina bifida). Animal studies have demonstrated valproic acid induced teratogenicity, and studies in human females have demonstrated placental transfer of the drug.

Multiple reports in the clinical literature indicate an association between the use of anti-epileptic drugs and an increased incidence of birth defects in children born to epileptic women taking such medication during pregnancy. The incidence of congenital malformations in the general population is regarded to be approximately 2%; in children of treated epileptic women, this incidence may be increased 2- to 3-fold. The increase is largely due to specific defects, eg. congenital malformations of the heart, cleft lip or palate, and neural tube defects. Nevertheless, the great majority of mothers receiving anti-epileptic medications deliver normal infants.

Data are more extensive with respect to diphenylhydantoin and phenobarbital, but these drugs are also the most commonly prescribed anti-epileptics. Some reports indicate a possible similar association with the use of other anti-epileptic drugs, including trimethadione, paramethadione, and valproic acid. However, the possibility also exists that other factors, eg. genetic predisposition or the epileptic condition itself may contribute to or may be mainly responsible for the higher incidence of birth defects.

Anti-epileptic drugs should not be discontinued in patients to whom the drug is administered to prevent major seizures, because of the strong possibility of precipitating status epilepticus with attendant hypoxia and risks to both the mother and the unborn child. With regard to drugs given for minor seizures, the risks of discontinuing medication prior to or during pregnancy should be weighed against the risk of congenital defects in the particular case and with the particular family history.

Epileptic women of child-bearing age should be encouraged to seek the counsel of their physician and should report the onset of pregnancy promptly to him. Where the necessity for continued use of anti-epileptic medication is in doubt, appropriate consultation is indicated.

**Nursing Mothers:** Valproic acid is excreted in breast milk. Concentrations in breast milk have been reported to be 1 to 10% of serum concentrations. As a general rule, nursing should not be undertaken while a patient is receiving Epival (divalproex sodium) or Depakene (valproic acid).

**Fertility:** Chronic toxicity studies in juvenile and adult rats and dogs demonstrated reduced spermatogenesis and testicular atrophy at doses of valproic acid greater than 200 mg/kg/day in rats and 90 mg/kg/day in dogs. Segment 1 fertility studies in rats have shown that doses up to 350 mg/kg/day for 60 days have no effect on fertility. The effect of divalproex sodium and valproic acid on the development of the testes and on sperm production and fertility in humans is unknown.

**LONG-TERM TOXICITY STUDIES IN RATS AND MICE INDICATED A POTENTIAL CARCINOGENIC RISK.**

**PRECAUTIONS: Hepatic dysfunction:** See **CONTRAINDICATIONS** and **WARNINGS**.

**General:** Because of reports of thrombocytopenia and inhibition of platelet aggregation, platelet counts and bleeding-time determination are recommended before instituting therapy and at periodic intervals. It is recommended that patients be monitored for platelet count prior to planned surgery. Clinical evidence of hemorrhage, bruising or a disorder of hemostasis/coagulation is an indication for reduction of dosage or withdrawal of therapy pending investigation.

Hyperammonemia with or without lethargy or coma has been reported and may be present in the absence of abnormal liver function tests; if elevation occurs the drug should be discontinued.

Because Depakene or Epival may interact with other anti-epileptic drugs, periodic serum level determinations of concurrently administered anti-epileptics are recommended during the early part of therapy. (See **DRUG INTERACTIONS**.) There have been reports of breakthrough seizures occurring with the combination of valproic acid and phenytoin.

Depakene and Epival are partially eliminated in the urine as a ketone-containing metabolite which may lead to a false interpretation of the urine ketone test.

There have been reports of altered thyroid function tests associated with valproic acid; the clinical significance of these is unknown.

**Driving and Hazardous Occupations:** May produce CNS depression, especially when combined with another CNS depressant, such as alcohol. Therefore, patients should be advised not to engage in hazardous occupations, such as driving a car or operating dangerous machinery, until it is known that they do not become drowsy from the drug.

**Drug Interactions:** May potentiate the CNS depressant action of alcohol.

There is evidence that valproic acid may cause an increase in serum phenobarbital levels, by impairment of non-renal clearance. This phenomenon can result in severe CNS depression. The combination of valproic acid and phenobarbital has also been reported to produce CNS depression without significant elevations of barbiturate or valproic acid serum levels. Patients receiving concomitant barbiturate therapy should be closely monitored for neurological toxicity. Serum barbiturate drug levels should be obtained, if possible, and the barbiturate dosage decreased, if indicated.

Primidone is metabolized into a barbiturate, and therefore, may also be involved in a similar or identical interaction.

There is conflicting evidence regarding the interaction of valproic acid with phenytoin (See **PRECAUTIONS - General**). It is not known if there is a change in unbound (free) phenytoin serum levels. The dosage of phenytoin should be adjusted as required by the clinical situation.

The concomitant use of valproic acid and clonazepam may produce absence status.

Caution is recommended when valproic acid or divalproex sodium is administered with drugs affecting coagulation, eg. acetylsalicylic acid and warfarin (See **ADVERSE REACTIONS**).

**ADVERSE REACTIONS** The most commonly reported adverse reactions are nausea, vomiting and indigestion. Since valproic acid has usually been used with other anti-epileptics, it is not possible in most cases to determine whether the adverse reactions mentioned in this section are due to valproic acid alone or to the combination of drugs.

**Gastrointestinal:** Nausea, vomiting and indigestion are the most commonly reported side effects at the initiation of therapy. These effects are usually transient and rarely require therapy. Diarrhea, abdominal cramps and constipation have also been reported. Anorexia with some weight loss and increased appetite with some weight gain have also been seen.

**CNS Effects:** Sedative effects have been noted in patients receiving valproic acid alone but are found most often in patients on combination therapy. Sedation usually disappears upon reduction of other anti-epileptic medication. Ataxia, headache, myasthenia, diplopia, asterixis, "spots before the eyes", tremor, dysarthria, dizziness, and incoordination have rarely been noted. Rare cases of coma have been reported in patients receiving valproic acid alone or in conjunction with phenobarbital.

**Dermatologic:** Transient increases in hair loss have been observed. Skin rash and petechiae have rarely been noted.

**Endocrine:** There have been reports of irregular menses and secondary amenorrhea in patients receiving valproic acid.

Abnormal thyroid function tests have been reported (See **PRECAUTIONS**). **Psychiatric:** Emotional upset, depression, psychosis, aggression, hyperactivity and behavioural deterioration have been reported.

**Musculoskeletal:** Weakness has been reported.

**Hematopoietic:** Thrombocytopenia has been reported. Valproic acid inhibits the second phase of platelet aggregation (See **PRECAUTIONS**). This may be reflected in altered bleeding time. Bruising, hematoma formation and frank hemorrhage have been reported. Relative lymphocytosis and hypofibrinogenemia have been noted. Leukopenia and eosinophilia have also been reported. Anemia and bone marrow suppression have been reported.

**Hepatic:** Minor elevations of transaminases (eg. SGOT and SGPT) and LDH are frequent and appear to be dose related. Occasionally, laboratory tests also show increases in serum bilirubin and abnormal changes in other liver function tests. These results may reflect potentially serious hepatotoxicity (See **WARNINGS**).

**Metabolic:** Hyperammonemia (See **PRECAUTIONS**). Hyperglycemia has been reported and associated with a fatal outcome in a patient with pre-existing non-ketotic hyperglycemia.

**Pancreatic:** There have been reports of acute pancreatitis occurring in association with therapy with valproic acid.

**SYMPTOMS AND TREATMENT OF OVERDOSAGE** In a reported case of overdose with valproic acid after ingesting 36 g in combination with phenobarbital and phenytoin, the patient presented in deep coma. An EEG recorded diffuse slowing, compatible with the state of consciousness. The patient made an uneventful recovery.

Naloxone has been reported to reverse the CNS-depressant effects of valproic acid overdose.

Because naloxone could theoretically also reverse the anti-epileptic effects of Depakene or Epival, it should be used with caution.

Since Epival tablets are enteric-coated, the benefit of gastric lavage or emesis will vary with the time since ingestion. General supportive measures should be applied with particular attention to the prevention of hypovolemia and the maintenance of adequate urinary output.

**DOSE AND ADMINISTRATION** The recommended initial dosage is 15 mg/kg/day, increasing at one week intervals by 5 to 10 mg/kg/day until seizures are controlled or side effects preclude further increases.

The maximal recommended dosage is 60 mg/kg/day. When the total daily dose exceeds 125 mg, it should be given in a divided regimen (See Table).

The frequency of adverse effects (particularly elevated liver enzymes) may increase with increasing dose. Therefore, the benefit gained by improving seizure control must be weighed against the increased incidence of adverse effects.

As the dosage is raised, blood levels of phenobarbital or phenytoin may be affected (See **PRECAUTIONS**).

Patients who experience G.I. irritation may benefit from administration of the drug with food or by a progressive increase of the dose from an initial low level. The capsules or tablets should be swallowed without chewing.

**AVAILABILITY** Depakene (valproic acid) is available as orange-coloured, soft gelatin capsules of 250 mg in bottles of 100 capsules; pale yellow, oval, soft gelatin enteric-coated capsules of 500 mg in bottles of 100 capsules; and as a red syrup containing the equivalent of 250 mg valproic acid, as the sodium salt, per 5 mL in bottles of 450 mL.

Epival (divalproex sodium) enteric-coated tablets are available as salmon-pink coloured tablets of 125 mg; peach-coloured tablets of 250 mg; lavender-coloured tablets of 500 mg. Supplied in bottles of 100 tablets.

Table of Initial Doses by Weight (based on 15 mg/kg/day)

kg	lb	Dosage		
		Total daily dose (mg)	equivalent to valproic acid Dose 1	Dose 2 Dose 3
10-24.9	22-54.9	250	125	0 125
25-39.9	55-87.9	500	250	0 250
40-59.9	88-131.9	750	250	250 250
60-74.9	132-164.9	1,000	250	250 500
75-89.9	165-197.9	1,250	500	250 500

Product monograph available on request.

- Henrikson O, Johannessen SI. Clinical and pharmacokinetic observations on sodium valproate-A 5-year follow-up study in 100 children with epilepsy. *Acta Neurol Scand* 1982;65:504-523.
- Delgado-Escueta AV et al. *Medical Progress: The Treatable Epilepsies*. New Engl J Med 1983;308(26):1576-1584.
- Turnbull DM et al. A comparison of phenytoin and valproate in previously untreated adult epileptic patients. *J Neuro Neurosurg Psych* 1982; 45:55-59.
- Covanis A et al. Sodium Valproate: Monotherapy and Polytherapy. *Epilepsia* 1982;23:693-720.
- Loiseau P. Rational Use of Valproate: Indications and Drug Regimen in Epilepsy. *Epilepsia* 1984;25(1):S65-S72.
- American Academy of Pediatrics' Committee on Drugs. Behavioural and Cognitive Effects of Anticonvulsant Therapy. *Pediatrics* 1985; 76(4):644-647.
- Beghi E et al. Adverse Reactions to Antiepileptic Drugs: A Multicenter Survey of Clinical Practice. *Epilepsia* 1986;27(4):323-330.
- Wilder BJ et al. Gastrointestinal tolerance of divalproex sodium. *Neurology* 1983;33(6): 808-811.

©Abbott Laboratories, Limited  
• TM  
Printed in Canada

PHARMACEUTICAL PRODUCTS DIVISION  
ABBOTT LABORATORIES, LIMITED  
MONTREAL, CANADA

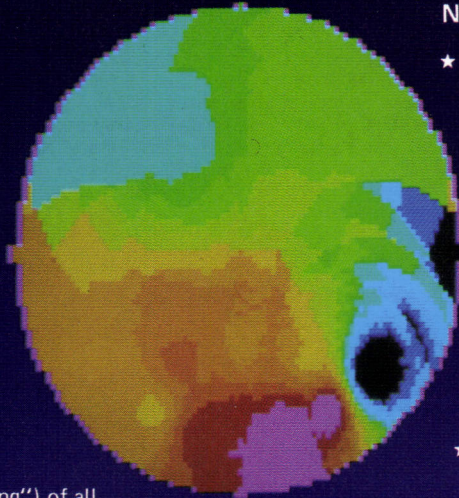
MAAD PAAB

See pages vi, vii

# New Dimension in EEG and Evoked

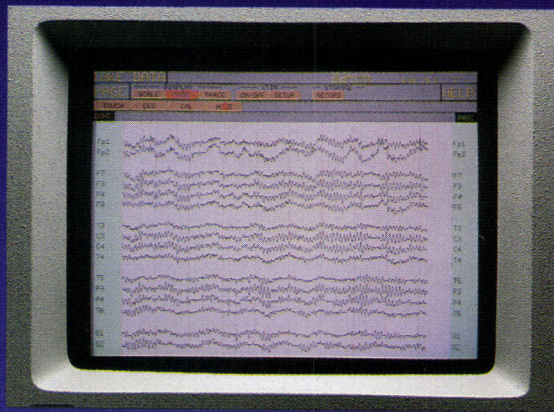
## SEEG

- ★ Advanced electroencephalography and evoked potentials testing
- ★ Up to 32 recording channels
- ★ Software—control using a "mouse"
- ★ Storage of raw and digitally processed EEG on (hard)disk or streaming tape
- ★ Review of EEG in any desired montage
- ★ Software for spectral, spatial and statistical analysis and for specific applications
- ★ Topographic display ("brain—mapping") of all EEG and EP data

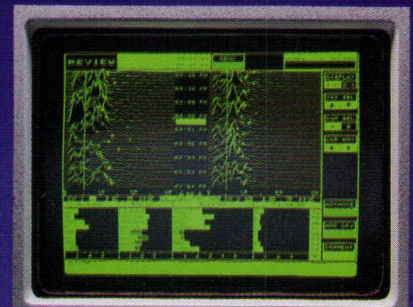


## NeuroScope

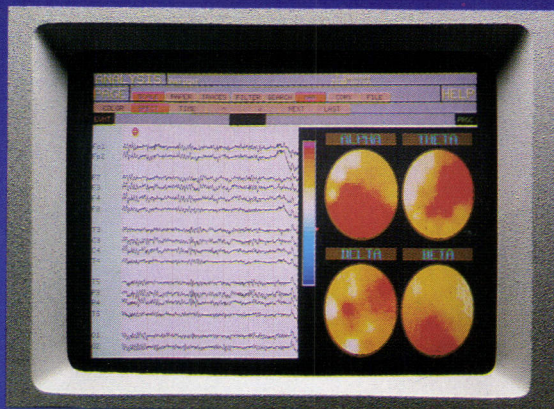
- ★ Unique touch screen control
- ★ Performs up to four—channel auditory, visual and somatosensory evoked potentials
- ★ Can be configured for EEG, CSA and EP Monitoring
- ★ Computer—generated electrode montage including impedance check
- ★ All stimulator functions for VEP, AEP and SEP
- ★ Test results on floppy disk for review and analysis
- ★ Built in thermal printer



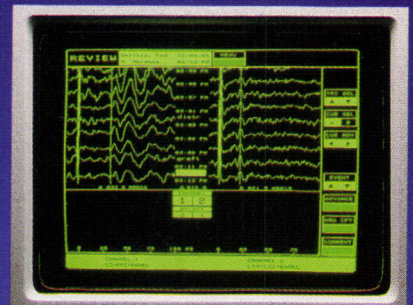
Electroencephalography



EEG and EP Monitoring During Carotid Endarterectomy



Quantitative Analysis



Evoked Potentials in Brain and Spinal Surgery



DANTEC ELECTRONICS Ltd  
140 Shorting Road  
Scarborough  
Ontario M1S 3S6  
Canada  
Telephone: (416) 298-2091  
Telex: 6525137 dantec tor



NeuroScope

SEEG



# Because quality of life is the issue

**ACTIONS** ParloDel (bromocriptine mesylate) is a dopaminomimetic ergot derivative with D<sub>2</sub> type dopamine receptor agonist activity, and has also D<sub>1</sub> dopamine receptor antagonist properties. The dopaminomimetic activity of bromocriptine in the striatum is considered responsible for the clinical benefits seen in selected patients with Parkinson's disease, when low doses of the drug are gradually added to levodopa therapy in patients on long-term treatment who develop late side effects of levodopa or no longer respond to the medication. Excessive dopaminomimetic drive may, however, provoke psychotic and other adverse reactions.

The extreme variability in G.I. tract absorption and the extensive and individually variable first-pass metabolism are responsible for the broad variability in plasma concentrations of bromocriptine and, in part, for the variability in dose response.

**INDICATIONS\*** **Parkinson's Disease:** ParloDel (bromocriptine mesylate) has been found to be clinically useful as an adjunct to levodopa (usually with a decarboxylase inhibitor), in the symptomatic management of selected patients with Parkinson's disease who experience prominent dyskinesia or wearing off reactions on long-term levodopa therapy.

Patients on long-term treatment who are beginning to deteriorate on levodopa therapy may be controlled by reducing the dose of levodopa and adjusting the frequency and schedule of drug administration. Patients maintained on optimal dosages of levodopa who still experience prominent dyskinesia and/or end-of-dose failure may benefit from the concomitant use of ParloDel, by decreasing the occurrence and/or severity of these manifestations. Since rapid escalation of bromocriptine doses causes severe adverse reactions, it is recommended to combine a slow increase of ParloDel, usually with a concomitant, gradual and limited reduction of levodopa dosage. Continued efficacy of bromocriptine for more than two years has not been established and there is some evidence that its efficacy tends to wane. Evidence available indicates that there is no consistent benefit from bromocriptine in patients who have not responded previously to levodopa, and studies have shown significantly more adverse reactions in bromocriptine-treated patients than in patients treated with levodopa. ParloDel is not recommended in the treatment of newly diagnosed patients or as the sole medication in Parkinson's disease.

**CONTRAINDICATIONS** Other than sensitivity to ergot alkaloids, no absolute contraindications to treatment with ParloDel (bromocriptine mesylate) are known. For procedure during pregnancy see "Use in Pregnancy" under Precautions.

**WARNINGS** Long-term treatment (6-36 months) with ParloDel in doses of 20 to 100 mg/day has been associated with pulmonary infiltrates, pleural effusion and thickening of the pleura in a few patients. Where ParloDel was discontinued, these changes slowly reverted to normal.

**PRECAUTIONS** ParloDel (bromocriptine mesylate) may cause hypotension, primarily postural; periodic monitoring of the blood pressure, particularly during the first days of therapy, is advisable. In some patients dizziness (vertigo) may occur with ParloDel; patients should therefore be cautioned against activities requiring rapid and precise responses, such as driving an automobile or operating dangerous machinery, until their response has been determined.

Care should be exercised when administering ParloDel concomitantly with phenothiazines or antihypertensive agents. Due to drug interaction at the receptor site, dosage should be adjusted accordingly.

Alcohol should be avoided during treatment with ParloDel. In some patients, the concomitant use of ParloDel and alcohol has given rise to alcohol intolerance and an increase in the severity and incidence of ParloDel's possible adverse reactions.

ParloDel should always be taken with food. In cases

where severe adverse effects, such as nausea, vomiting, vertigo or headaches are severe or persisting, the therapeutic dosage of ParloDel should be reduced to half of one tablet daily (1.25 mg) and increased gradually to that recommended. The dopamine antagonist domperidone may be useful in the control of severe gastrointestinal side effects in parkinsonian patients receiving ParloDel (see Drug Interactions).

As with all medication, ParloDel should be kept safely out of the reach of children.

**Use in Pregnancy:** If the patient wishes to become pregnant, ParloDel (bromocriptine mesylate) should be stopped as soon as possible after conception is suspected. In this event immunological confirmation should be done immediately. When pregnancy is confirmed, ParloDel, like all other drugs, should be discontinued unless, in the opinion of the treating physician, the possible benefit to the patient outweighs the potential risk to the fetus.

In human studies with ParloDel (reviewed by Turkali, I.), there were 1410 reported pregnancies, which yielded 1236 live and 5 stillborn infants from women who took ParloDel (bromocriptine mesylate) during early pregnancy. Among the 1241 infants, 43 cases (31 minor and 12 major) of congenital anomalies were reported. The incidence (3.46%) and type of congenital malformations and the incidence of spontaneous abortions (11.13%) in this group of pregnancies does not exceed that generally reported for such occurrences in the population at large.

**Use in Parkinson's Disease:** Use of ParloDel (bromocriptine mesylate), particularly in high doses, may be associated with mental confusion and mental disturbances. Since patients with Parkinson's disease may manifest varying degrees of dementia, caution should be exercised when treating such patients with ParloDel.

ParloDel administered alone or concomitantly with levodopa may cause visual or auditory hallucinations. These usually resolve with dosage reduction, but discontinuation of ParloDel may be required in some cases. Rarely, after high doses, hallucinations have persisted for several weeks following discontinuation of ParloDel. Caution should be exercised when administering ParloDel to patients with a history of myocardial infarction, particularly if they have a residual atrial, nodal or ventricular arrhythmia.

Symptomatic hypotension can occur and, therefore, caution should be exercised when administering ParloDel, particularly in patients receiving antihypertensive medication. Periodic evaluation of hepatic, hematopoietic, cardiovascular and renal function is recommended.

**Drug Interactions:** The concomitant use of erythromycin may increase bromocriptine plasma levels.

Domperidone, a dopamine antagonist, may cause increases in serum prolactin. In so doing, domperidone may antagonise the therapeutically relevant prolactin lowering effect of ParloDel. It is possible that the anti-tumorigenic effect of ParloDel in patients with prolactinomas may be partially blocked by domperidone administration.

**ADVERSE REACTIONS** The most frequently observed adverse reactions are nausea, vomiting, headache and gastrointestinal side effects such as abdominal pain, diarrhea and constipation. All these effects may be minimized or even prevented by giving small initial doses of bromocriptine and by taking it with food.

Postural hypotension which can, on rare occasions, lead to fainting and "shock-like" syndromes has been reported in sensitive patients. This is most likely to occur during the first few days of ParloDel treatment.

When bromocriptine is added to levodopa therapy, the incidence of adverse reactions may increase. The most common newly appearing adverse reactions in combination therapy were: nausea, abnormal involuntary movements,

hallucinations, confusion, "on-off" phenomenon, dizziness, drowsiness, faintness, fainting, vomiting, asthenia, abdominal discomfort, visual disturbance, ataxia, insomnia, depression, hypotension, shortness of breath, constipation and vertigo.

Less common adverse reactions include anorexia, anxiety, blepharospasm, dry mouth, dysphagia, edema of the feet and ankles, erythromelalgia, epileptiform seizures, fatigue, headache, lethargia, mottling of skin, nasal stuffiness, nervousness, nightmares, paresthesia, skin rash, urinary frequency, urinary incontinence, urinary retention and rarely signs or symptoms of ergotism such as tingling of fingers, cold feet, numbness, muscle cramps of feet and legs or exacerbation of Raynaud's syndrome.

Abnormalities in laboratory tests may include elevation of blood urea nitrogen, SGOT, SGPT, GGPT, CPK, alkaline phosphatase and uric acid, which are usually transient and not of clinical significance.

The occurrence of adverse reactions may be lessened by temporarily reducing dosage to one-half tablet two or three times daily.

**SYMPTOMS AND TREATMENT OF OVERDOSE** There have been several reports of acute overdosage with ParloDel (bromocriptine mesylate) in children and adults. No life threatening reactions have occurred. Symptoms reported included nausea, vomiting, dizziness, drowsiness, hypotension, sweating and hallucinations. Management is largely symptomatic; the cardiovascular system should be monitored. Metoclopramide can be used to antagonize the emesis and hallucinations in patients who have taken high doses.

**DOSAGE AND ADMINISTRATION** ParloDel (bromocriptine mesylate) should always be taken with food.

Although ParloDel (bromocriptine mesylate) has been found clinically useful in decreasing the severity and frequency of "on-off" fluctuations of late levodopa therapy, the decision to use bromocriptine as adjunctive treatment and the selection of dosage must be individualized in each case. A low dose is recommended. The initial dose of ParloDel is one half of a 2.5 mg tablet (1.25 mg) at bedtime with food to establish initial tolerance. Thereafter, the recommended dosage is 2.5 mg daily in two divided doses, with meals, (half a 2.5 mg tablet twice daily). The dosage may be increased very gradually, if necessary, by adding an additional 2.5 mg per day, once every 2 to 4 weeks, to be taken always in divided doses with meals. Increments should usually not exceed 2.5 mg. Clinical assessments are recommended at two week intervals or less during dosage titration, to ensure that the lowest effective dosage is not exceeded. The usual dosage range is from a few milligrams to 40 mg daily in two or three divided doses with meals. The median dose varies with the experience of individual investigators, but can be around 10 mg daily or higher. During initial titration it is recommended that the dosage of levodopa should be maintained, if possible. Subsequently, it might be desirable to combine a slow increase of bromocriptine with a concomitant, limited and gradual reduction of levodopa.

#### AVAILABILITY

TABLETS each containing 2.5 mg bromocriptine, as mesylate, available in bottles of 100.  
CAPSULES each containing 5 mg bromocriptine, as mesylate, available in bottles of 100.

\*For information on other approved indications, please consult the ParloDel product monograph, available to physicians and pharmacists on request.

**SANDOZ**

Sandoz Canada Inc.  
P.O. Box 385  
Dorval, Quebec H9R 4P5

## Brief Prescribing Information

### **Tegretol**<sup>®</sup> (carbamazepine)

**TEGRETOL**<sup>®</sup> 200 mg  
**TEGRETOL**<sup>®</sup> CHEWTABS<sup>™</sup> 100 mg and 200 mg  
**TEGRETOL**<sup>®</sup> CR 200 mg and 400 mg

#### Action

TEGRETOL (carbamazepine) has anticonvulsant properties which have been found useful in the treatment of psychomotor epilepsy and, as an adjunct in the treatment of partial epilepsies, when administered in conjunction with other anticonvulsant drugs to prevent the possible generalization of the epileptic discharge. A mild psychotropic effect has been observed in some patients, which seems related to the effect of the carbamazepine in psychomotor or temporal lobe epilepsy.

TEGRETOL relieves or diminishes the pain associated with trigeminal neuralgia often within 24 to 48 hours.

Like other tricyclic compounds, TEGRETOL has a moderate anticholinergic action which is responsible for some of its side effects. A tolerance may develop to the action of TEGRETOL after a few months of treatment and should be watched for.

TEGRETOL may suppress ventricular automaticity due to its membrane-depressant effect similar to that of quinidine and procainamide, associated with suppression of phase 4 depolarization of the heart muscle fibre. A number of investigators have reported a deterioration of EEG abnormalities with regard to lead alterations and a higher incidence of records with nil beta activity, during carbamazepine-combined treatment.

The absorption of carbamazepine in man is relatively slow. When taken in a single oral dose, TEGRETOL (carbamazepine tablets) and TEGRETOL CHEWTABS (carbamazepine chewable tablets) yield peak plasma concentrations of unchanged carbamazepine within 4-24 hours. With respect to the quantity of carbamazepine absorbed, there is no clinically relevant difference between the various dosage forms. When TEGRETOL CR (carbamazepine controlled release tablets) are administered repeatedly, they yield a lower average maximal concentration of carbamazepine in the plasma, without a reduction in the average minimal concentration. This tends to result in a lower incidence of intermittent concentration-dependent adverse drug reactions. It also ensures that the plasma concentrations remain largely stable throughout the day, thereby making it possible to manage with a twice-daily dosage.

Carbamazepine becomes bound to serum proteins to the extent of 70-80%. The concentration of unchanged substance in the saliva reflects the non-protein-bound portion present in the serum (20-30%).

The elimination half-life of unchanged carbamazepine in the plasma averages approximately 36 hours following a single oral dose, whereas after repeated administration, which leads to autoinduction of hepatic enzymes, it averages only 16-24 hours, depending on the duration of the medication. In patients receiving concomitant treatment with other enzyme-inducing anti-epileptic agents, half-life values average 9-10 hours have been found. Only 2-3% of the dose, whether given singly or repeatedly, is excreted in the urine in unchanged form. The primary metabolite is the pharmacologically active 10, 11-epoxide.

In man, the main urinary metabolite of carbamazepine is the trans-diol derivative originating from the 10, 11-epoxide; a small portion of the epoxide is converted into 9-hydroxymethyl-10-carbamoyl-acridan. Other important biotransformation products are various monohydroxylated compounds, as well as the N-glucuronide of carbamazepine.

The therapeutic range for the steady-state plasma concentration of carbamazepine generally lies between 4-10 mcg/ml.

#### Indications and Clinical Use

##### A. Trigeminal Neuralgia:

TEGRETOL (carbamazepine) is indicated for the symptomatic relief of pain of trigeminal neuralgia only during periods of exacerbation of true or primary trigeminal neuralgia (tic douloureux). It should not be used preventively during periods of remission. In some patients, TEGRETOL has relieved glossopharyngeal neuralgia. For patients who fail to respond to TEGRETOL, or who are sensitive to the drug, recourse to other accepted measures must be considered.

TEGRETOL is not a simple analgesic and should not be used to relieve trivial facial pains or headaches.

##### B. TEGRETOL has been found useful in:

1. the management of psychomotor (temporal lobe) epilepsy and,
2. as an adjunct, in some patients with secondary or partial epilepsies with complex symptomatology or secondarily generalized seizures, when administered in combination with other antiepileptic medication.
3. as an alternative medication in patients with generalized tonic-clonic seizures who are experiencing marked side effects or fail to respond to other anticonvulsant drugs.

TEGRETOL is not effective in controlling petit mal, minor motor, myoclonic and predominantly unilateral seizures, and does not prevent the generalization of epileptic discharge. Moreover, recent information suggests that exacerbation of seizures may occasionally occur in patients with atypical absences.

#### Contraindications

TEGRETOL (carbamazepine) should not be administered to patients with a history of hepatic disease or serious blood disorder.

TEGRETOL should not be administered immediately before, in conjunction with, or immediately after a monoamine oxidase inhibitor. When it seems desirable to administer TEGRETOL to a patient who has been receiving an MAO inhibitor, there should be as long a drug-free interval as the clinical condition allows, but in no case should this be less than 14 days. Then the dosage of TEGRETOL should be low initially, and increased very gradually.

TEGRETOL should not be administered to patients presenting atrioventricular heart block. (See Sections on Action and Precautions).

Safe use in pregnancy has not been established. Therefore, TEGRETOL should not be administered during the first 3 months of pregnancy. TEGRETOL should not be given to women of child-bearing potential unless, in the opinion of the physician, the expected benefits to the patient outweigh the possible risk to the fetus (See Reproductive Studies). Because of demonstrated toxicity in nursing animals TEGRETOL should not be administered to nursing mothers.

TEGRETOL should not be administered to patients with known hypersensitivity to carbamazepine or to any of the tricyclic compounds, such as amitriptyline, trimipramine, imipramine, or their analogues or metabolites, because of the similarity in chemical structure.

#### Warnings

Although reported infrequently, serious adverse effects have been observed during the use of TEGRETOL (carbamazepine). Agranulocytosis and aplastic

anemia have occurred in a few instances with a fatal outcome. Leucopenia, thrombocytopenia, hepatocellular and cholestatic jaundice, and hepatitis have also been reported. It is, therefore, important that TEGRETOL should be used carefully and close clinical and frequent laboratory supervision should be maintained throughout treatment in order to detect as early as possible signs and symptoms of a possible blood dyscrasia.

Long-term toxicity studies in rats indicated a potential carcinogenic risk (See Section on Toxicology). Therefore, the possible risk of drug use must be weighed against the potential benefits before prescribing carbamazepine to individual patients.

#### Precautions

##### Monitoring of Hematological and Other Adverse Reactions:

Complete blood studies, including platelet counts, and evaluation of hepatic and renal function and urinalysis should be carried out before treatment is instituted. Careful clinical and laboratory supervision should be maintained throughout treatment, including frequent performance of complete blood counts, in order to detect any early signs or symptoms of blood dyscrasia. Should any signs or symptoms or abnormal laboratory findings suggestive of blood dyscrasia or liver disorder occur, TEGRETOL (carbamazepine) should be immediately discontinued until the case is carefully reassessed. Non-progressive or fluctuating asymptomatic leucopenia, which is encountered, does not generally call for the withdrawal of TEGRETOL. However, treatment with TEGRETOL should be discontinued if the patient develops leucopenia which is progressive or accompanied by clinical manifestations, e.g. fever or sore throat.

##### Urinary Retention and Increased Intraocular Pressure:

Because of its anticholinergic action, TEGRETOL should be given cautiously, if at all, to patients with increased intraocular pressure or urinary retention. Such patients should be followed closely while taking the drug.

##### Occurrence of Behavioural Disorders:

Because it is closely related to the other tricyclic drugs, there is some possibility that TEGRETOL might activate a latent psychosis, or, in elderly patients, produce agitation or confusion, especially when combined with other drugs. Caution should also be exercised in alcoholics.

##### Use in Patients with Cardiovascular Disorders:

TEGRETOL should be used cautiously in patients with a history of coronary artery disease, organic heart disease, or congestive failure. If a defective conductive system is suspected, an ECG should be performed before administering TEGRETOL, in order to exclude patients with atrioventricular block.

##### Driving and Operating Hazardous Machinery:

Because dizziness and drowsiness are possible side effects of TEGRETOL, patients should be warned about the possible hazards of operating machinery or driving automobiles.

##### Drug Interactions:

Induction of hepatic enzymes in response to TEGRETOL may have the effect of diminishing the activity of certain drugs that are metabolized in the liver. This should be considered when administering TEGRETOL concomitantly with other anti-epileptic agents and drugs such as theophylline. Concomitant administration of TEGRETOL with verapamil, diltiazem, erythromycin, troleandomycin, cimetidine, propoxyphene or isoniazid, has been reported to result in elevated plasma levels of carbamazepine. Since an increase in the blood levels of carbamazepine may result in unwanted effects (e.g. dizziness, headache, ataxia, diplopia and nystagmus may occur), the dosage of carbamazepine should be adapted accordingly and blood levels monitored.

The concomitant administration of carbamazepine and lithium may increase the risk of neurotoxic side effects.

In patients receiving oral anticoagulant medication, the dosage of the anticoagulant should be readjusted to clinical requirements whenever treatment with TEGRETOL is initiated or withdrawn.

TEGRETOL, like other anticonvulsants, may adversely affect the reliability of oral contraceptives. Patients should accordingly be advised to use some alternative, non-hormonal method of contraception.

TEGRETOL, like other psycho-active drugs, may reduce the patient's alcohol tolerance; it is therefore advisable to abstain from alcohol consumption during treatment.

TEGRETOL should not be administered in conjunction with an MAO inhibitor. (See Section on Contraindications).

#### Adverse Reactions

The reactions which have been most frequently reported with TEGRETOL (carbamazepine) are drowsiness, unsteadiness on the feet, vertigo, dizziness, gastrointestinal disturbances, and nausea. These reactions usually occur only during the initial phase of therapy. They have rarely necessitated discontinuing TEGRETOL therapy, and can be minimized by initiating treatment at a low dosage.

The more serious adverse reactions observed are the hematologic, hepatic, cardiovascular and dermatologic reactions, which require discontinuation of therapy. If treatment with TEGRETOL has to be withdrawn abruptly, the change-over to another anti-epileptic drug should be effected under cover of diazepam.

The following adverse reactions have been reported:

**Hematologic** - Transitory leucopenia, eosinophilia, hyponatremia, leucocytosis, thrombocytopenic purpura, agranulocytosis, macrocytic anemia and aplastic anemia. In a few instances, deaths have occurred.

**Hepatic** - During the long-term administration of TEGRETOL, abnormalities in liver function tests, cholestatic and hepatocellular jaundice, and hepatitis have been reported.

**Dermatologic** - The following reactions occurred during treatment with TEGRETOL: skin sensitivity reactions and rashes, erythematous rashes, pruritic eruptions, urticaria, photosensitivity, pigmentary changes, neurodermatitis and in rare cases Stevens-Johnson syndrome, toxic epidermal necrolysis, exfoliative dermatitis, alopecia, diaphoresis, erythema multiforme, erythema nodosum, and aggravation of disseminated lupus erythematosus.

**Neurologic** - The reactions reported as occurring during treatment with TEGRETOL include vertigo, somnolence, ataxia, confusion, headache, fatigue, blurred vision, spatial hallucinations, transient diplopia and oculomotor disturbances, speech disturbances, abnormal involuntary movements and increase in motor seizures. In addition, peripheral neuritis and paresthesia, depression with agitation, talkativeness, nystagmus, hyperacusis, and tinnitus have been reported but only very rarely. There have been some reports of paralysis and other symptoms of cerebral arterial insufficiency but no conclusive relationship to the administration of TEGRETOL could be established.

**Cardiovascular** - Thromboembolism, recurrence of thrombophlebitis in patients with a prior history of thrombophlebitis, primary thrombophlebitis, congestive heart failure, aggravation of hypertension, Stokes-Adams in patients with AV block, hypertension, syncope and collapse, edema, aggravation of coronary artery disease. Some of these complications including myocardial infarction and arrhythmia) have been associated with other tricyclic compounds.

**Genitourinary** - Urinary frequency, acute urinary retention, oliguria with elevated blood pressure, azotemia, renal failure, and impotence. Elevation of BUN, albuminuria and glycosuria also have been observed.

**Respiratory** - Pulmonary hypersensitivity characterized by fever, dyspnea, pneumonitis or pneumonia

**Gastrointestinal** - Disturbances associated with TEGRETOL therapy have included nausea, vomiting, gastric or abdominal discomfort, diarrhea or constipation, anorexia and dryness of the mouth and throat, glossitis and stomatitis.

**Ophthalmic** - There is no conclusive evidence that TEGRETOL produces pathological changes in the cornea, lens or retina. However, it should be recognized that many phenothiazines and related drugs have been shown to cause eye changes. By anatomy, periodic eye examinations, including slit-lamp funduscopy and tonometry, are recommended.

Other reactions reported during treatment with TEGRETOL include fever and chills, aching joints and muscles, leg cramps, conjunctivitis, and adenopathy or lymphadenopathy.

#### Symptoms and Treatment of Overdosage

##### Symptoms of Overdosage:

The symptoms of overdosage include dizziness, ataxia, drowsiness, stupor, nausea, vomiting, restlessness, agitation, disorientation, tremor, involuntary movements, opisthotonus, abnormal reflexes (slowed or hyperactive), mydriasis, nystagmus, flushing, cyanosis, and urinary retention. Hypotension or hypertension may develop. Coma may ensue. EEG and ECG changes may occur. The laboratory findings in isolated instances of overdosage have included leukocytosis, reduced leukocyte count, glycosuria and acetonaemia.

##### Treatment of Overdosage:

There is no known specific antidote to TEGRETOL (carbamazepine). Experience with accidental TEGRETOL overdosage is limited. Since TEGRETOL is chemically related to the tricyclic antidepressants, reference to treatment of TOFRANIL (imipramine) overdosage is relevant.

It is recommended that emesis be induced, and that gastric lavage be performed. Vital signs should be watched and symptomatic treatment should be administered as required. Hyperirritability may be controlled by the administration of parenteral diazepam or barbiturates. However, barbiturates should not be used if drugs that inhibit monoamine oxidase have also been taken by the patient, either in overdosage or in recent therapy (within two weeks).

Barbiturates may also induce respiratory depression, particularly in children. It is therefore advisable to have equipment available for artificial ventilation and resuscitation when barbiturates are employed. Paraldehyde may be used to counteract muscular hypertonus without producing respiratory depression.

Shock (circulatory collapse) should be treated with supportive measures, including intravenous fluids, oxygen, and corticosteroids.

It is recommended that the electrocardiogram be monitored, particularly in children, to detect any cardiac arrhythmias or conduction defects.

#### Dosage and Administration

##### Use in Epilepsy (See Indications):

A low initial daily dosage of TEGRETOL (carbamazepine) with a gradual increase in dosage is advised. Dosage should be adjusted to the needs of the individual patient.

TEGRETOL tablets and CHEWTABS should be taken in 2 to 4 divided doses daily, with meals whenever possible.

The controlled-release characteristics of TEGRETOL CR reduce the daily fluctuations of plasma carbamazepine. TEGRETOL CR tablets (either whole or, if so prescribed, only half a tablet) should be swallowed unchewed with a little liquid during or after a meal. These controlled-release tablets should be prescribed as a twice-daily dosage. If necessary, three divided doses may be prescribed.

##### Adults and Children Over 12 Years of Age:

Initially, 100 to 200 mg once or twice a day depending on the severity of the case and previous therapeutic history. The initial dosage is progressively increased, in divided doses, until the best response is obtained. The usual optimal dosage is 800 to 1200 mg daily. In rare instances some adult patients have received 1600 mg. As soon as disappearance of seizures has been obtained and maintained, dosage should be reduced very gradually until a minimum effective dose is reached.

##### Children 6-12 Years of Age:

Initially, 100 mg in divided doses on the first day increase gradually by adding 100 mg per day until the best response is obtained. Dosage should generally not exceed 1000 mg daily. As soon as disappearance of seizures has been obtained and maintained, dosage should be reduced very gradually until a minimum effective dose is reached.

##### Use in Trigeminal Neuralgia:

The initial daily dosage should be small, 200 mg taken in 2 doses of 100 mg each is recommended. The total daily dosage can be increased by 200 mg/day until relief of pain is obtained. This is usually achieved at dosage between 200 and 800 mg daily, but occasionally up to 1200 mg/day may be necessary. As soon as relief of pain has been obtained and maintained, progressive reduction in dosage should be attempted until a minimum effective dosage is reached. Because trigeminal neuralgia is characterized by periods of remission, attempts should be made to reduce or discontinue the use of TEGRETOL at intervals of not more than 3 months, depending upon the individual clinical course.

Prophylactic use of the drug in trigeminal neuralgia is not recommended.

#### Availability

**TEGRETOL tablets 200 mg** Each white, round, flat, bevelled-edge double-scored tablet engraved GEIGY on one side contains 200 mg carbamazepine. Available in bottles of 100 and 500 tablets.

**TEGRETOL CHEWTABS 100 mg** Pale pink, round, flat, bevelled-edge tablets with distinct red spots. GEIGY engraved on one side and MR on the other. Fully bisected between the M and R. Each chewable tablet contains 100 mg carbamazepine. Available in bottles of 100 CHEWTABS.

**TEGRETOL CHEWTABS 200 mg** Pale pink, oval, biconvex tablets with distinct red spots. GEIGY engraved on one side and PU engraved on the other. Fully bisected between the P and U. Each chewable tablet contains 200 mg carbamazepine. Available in bottles of 100 CHEWTABS.

**TEGRETOL CR 200 mg** Beige-orange, capsule-shaped, slightly biconvex tablet, engraved CG/CG on one side and HCH/CH on the other. Fully bisected on both sides. Each controlled-release tablet contains 200 mg carbamazepine. Available in bottles of 100 tablets.

**TEGRETOL CR 400 mg** Brownish-orange, capsule-shaped, slightly biconvex tablet, engraved CG/CG on one side and ENE/ENE on the other. Fully bisected on both sides. Each controlled-release tablet contains 400 mg carbamazepine. Available in bottles of 100 tablets.

Protect from heat and humidity.

#### Geigy


Mississauga, Ontario  
LSN 2W5



G 8800

1. Kramer G, Besser R, Katzmann K, Theisohn M.: Slow release carbamazepine in the treatment of epilepsy. *Akt. Neurol.* 1985; 12: 70-74. 2. Product Monograph: Data On File. 3. Adenkanp AP, Alpheris WJ, Moerland MC, Ohevaner N, Van Parys JAP. Controlled release carbamazepine: Cognitive side effects in patients with epilepsy. *Epilepsia* 1987; 28: 507-514.

See obc



**Consider 'Prolopa'<sup>®</sup>  
if her Parkinson  
therapy isn't  
working anymore!**

Dividing daily dosage of levodopa into more frequent, smaller doses of 'Prolopa' 50-12.5 appears to reduce severity of

- dose-related fluctuations
- abnormal involuntary movements and
- "on-off" phenomena<sup>1</sup>,

thereby improving patient response to therapy.

**Prolopa<sup>®</sup>** Helps return  
the simple pleasures of living

levodopa/benserazide 4:1



© Original Research in Medicine and Chemistry For brief prescribing information see page xxiv G4188



**No "breaking"  
required**



Prolopa<sup>®</sup>  
Capsule 50-12.5