BRIFF PRESCRIBING INFORMATION CONSULT BULL PRODUCT MONOGRAPHEOR COMPLETE PRESCRIPING INFORMATION

> **SReminyl** galantamine hydrobromide tablets 4 mg, 8 mg, 12 mg galantamine base

GReminyl Ex galantamine hydrobromide extended release capsules 8 mg, 16 mg, 24 mg galantamine base

Cholinesterase Inhibitor

INDICATIONS AND CLINICAL USE REMINY (nalantamine hydrohromide) and REMINY ER are NUMBER AND CANNEAL USE FEMANT, glasterance hydrocronicle and FERMANIC ET are included for the synchronic between of glasters with most to mostere secretific of the Achieves's type. REMANY, and REMANY, ET sould only be prescribed by the following constalation with clinicism shot are operational for displayers and management of Advance's disease. Secretification is the constrained in the displayers and management of Advance's disease. Secretification with advanced to make a size of the constant of the advanced to the advanced to the constant of the advanced to the advan e our years of agoy, in reals a militar salely information for inconvert, and technivit, but it this plateful opulation (see **WARNINGS AND PRECAUTIONS**). **Pediatrics:** No data are available in childrer erefore, the use of REMINYL and REMINYL ER are not recommended in children under 18 years of age CONTRAINDICATIONS REMIND and REMIND FR are contraindicated in nations with known sitivity to galantamine hydrobromide, other tertiary alkaloid derivatives or to any excipients

WARNINGS AND PRECAUTIONS Carcinogenesis and Mutagenesis See Product Monograph Part II: TOXICOLOGY- Carcinogenicity, Mutagenicity for discussion on animal data Cardiovascular, Because of their pharmacological action, cholinesterase inhibitors have vagotonic effects on the sinoatrial and atrioventricular nodes, leading to bradycardia and heart block. These election in the shaddler of authoristication, seeingly of conjugation and indicate cardiac snay be particularly important to patients with "six's kinus sportioner" or other supresenting cardiac conduction disorders, or to patients taking other drugs concomitantly which significantly slow heart rate. In clinical trials, patients with serious cardiovascular disease were excluded. Caution should be exercised in treating patients with active coronary artery disease or congestive heart failure. It is recommended that REMINML and REMINML ER not be used in patients with cardiac conduction recommended user informer, and inchemits, but not lessed in placents we include, coloration discommillates percept for right burndle transch blockly including "sick situs, syndrome" and those with unexplained synocyal episodes, in anotomized controlled trials, brankpradra was reported at 2°-38 ran syndromine doese up to Almydips composer with CFS for placed by users arrely severe and service led to treatment discontinuation. No increased incidence of heart block was observed at the led to Hardmand discontinuation. We increased incidence of heart block was observed at the recommend dotice. Particle hearted with patients up to 3 mill part in the encommend dotice. The encommend dotice is the encommend dotice in the encommend dotice in schedule of bound at dotie -related increase in risk of grocope placehol 0.7% [2766]. An 9-west conflorational scale of block 3 mill particle. 3 mill particle is scheduled by the scheduled of particle and scheduled of block 3 mill particle. 3 mill particle is scheduled by the scheduled particle and scheduled wheeks 3 mad 4, and 2 mill particle with scheduled particle is the scheduled wheeks 3 mad 4, and 2 mill particle with scheduled particle is the scheduled by more common registrations—bread patients than in placebot headed patients. It should be noted that a toward 1-week does exclusion was used in the solid, which is not normoreald. Whether these cardiac efficies are attenuated by solver thankon rates is not known. Particular causino is varietted charten through a mill particle in the water that schedules. The placetose was the problem of the schedules are attenuated by solver thankon rates is not known. Particular causino is varietted whether through solver the million of the schedules. The schedules in Policiated and the contract of the schedules. The schedules in the schedules in the schedules. contact effects are attentiately assert thation rates is not roome. Perficiels could not swarmed using thation where the majority of passes occurred in the above study. Metabolism, Chroinesterase inhibitors as well as Atherium's disease can be associated with significant weight loss. In controlled clinical thisis, the use of PERMIN, was sociated with weight itss. Weight diseases occurred early during treatment and was related to does. Weight so at 27% occurred me Requestive presents treated with PERMIN and in female patients than in patients receiving placebox. Where weight loss may be of clinical concern, body weight should be monitored. Seathwistediscall: Through their primary action, chilentisesses analisms may be expected in horizoned concern, body weight should be monitored closely for symptoms of active and concerns the controlled and in-Hammatory days. PASISD, it is notified in chilential studies with galantamies, patients with symptomic paper, clicarsion were excelled. Clinical studies with galantamies, patients with symptomic paper, clicarion were controlled contract studies with glastratines, puerfes with symptomatic pulps controlled Christial distribution from microrress, refettle bylaschoi, in the notices of either poptic stoor disease or gestrointestinal bleeding lose ANYERSE REACTIONS;. Galantamine, as a producible consequence of lis pharmacological properties, has been shown to produce trauses, unraining and deather, amonels and welget the These effects appeared now beyongst a reflect of these effects appeared one beyongst a reflect of doses (see ANYERSE REACTIONS), with reases and ventring being more prevailent in women and does give AUMISE PEACHORS, with races and noming being more precised in nome and prefers with one to viewish and compositionly help plasm day operations. The same more sendie to the challenge device effects associated with challenstesse inhibitors and in general are more likely the operations hasses and vorning than are makes, in most case, these effects were of mild to moderate intensity and trassent and time resident during continued PEAMINI treatment or your treatment discontinuation. <u>Reinflatentum Plancing not devered in clinical table of pattername, chromoments may cause belonger on office volume discontinuation pattername, chromoments may cause belonger on office volume discontinuation pattername, chromoments may cause belonger of solution were reported, there was no increase in pattername, chromoments are leading to a particulation of the contraction of cause existent, solution activity particulation of the manufactorium of Athomism's disease. The included Cause existent, solution activity particulation of the contraction of Athomism's disease. The included in the contraction of Athomism's disease. The included Athomism's disease. The included in the contraction of Athomism's disease. The included Athomism's disease. The included in the contraction of Athomism's disease. The included Athomism's disease of the contraction of the contraction of the contraction of Athomism's disease. The included in the contraction of Athomism's disease. The included in the contraction of Athomism's disease and the contraction of Athomism's disease. The contraction of Athomism's disease and the contraction of Athomism's di</u> REMINYL and REMINYL ER treatment for patients with a history of seizure disorder must therefore be carefully evaluated. REMINYL and REMINYL ER have not been studied in patients with moderately careful y evaluate. FEMINNI, and REMINNI, ET have not been studied in patients with moderably senier or severe Arbemer's disease, non-Arbemer dementace or individuals with Parkinsonian leutures. The efficacy and sately of FEMINNI, and FEMINNI, ET in these patient populations are unknown. Per disease's Considerations Aventhesia: Galantamine, as a chrisestesses inhibitor, is filed by beaugrapties. Appropriations Aventhesia: Galantamine, as a chrisestesses inhibitor, is skelly beaugrapties. Applications of the expension during versibles. Registration (see Arbeit christopist of activities of the patients of the patients with a history of activities of control produces and the patients of patients with a history of activities of patients with a history of activities of patients with a history of activities of patients with a history of activities. The patients with a history of activities of patients with a history of activities of patients with a history of activities. The patients with a history of activities of activities. are letter to continued out used sectionals with respect to continue to contin soil of a 11-51, Northinst, and mellinist characteristic data of the polarization in the implainment. There is inhabit information on the pharmacolinetists of paintainnes in result in regarder potents. It is therefore encommended that does excisation with REMINN, or REMINN, EAR Alberiens's doeses potents with resilician implainment presentine elegence of 9 to 80 million the understate with a casion and under conditions of olses monitoring for adverse effects (see ODAGE AND ARMINISTRATION, Secular Inputations). For no data are available on the use of REMINN, FRI publishes with a creatinite observance of less than 9 mL/min, REMINNI, and HEMINNI, ER returned to impose with order population. **Serfatiniss** (2-85 years of age): no controlled clinical studies, the *number of polletes* aged 65 years or over who received PENIMVI. At the represent Cessel to 67 v4 mg/day was 123. Of these patients, 70 received the maximum recommended does of 24 mg/day, There is limited safely information for REMINVI. In this patient population. Since cholinomimetics as well as Alzheimer's disease can be associated with significant weight loss, caution is advised regarding the use of REMINYL and REMINYL ER in elderly patients with low body weigh is advised regarding the use of ROMM, and REMINIT, this individual prices with the tody regist, especially in those 255 years of U.Ber. In Edithy Patients with Serving Connorthal Desizes. There is limited information on the safety of galantamine treatment in patients with mild to moderate Authenies's disease and services/galantime treatment in patients with mild to moderate Authenies's disease and services/galantime treatment and moderate of ROMM, and ROMM, the fin Authenies's disease patient with prices in common ramp of generating publication, solid consistent of high patient production solid process when cannot Patients with MC Cognitive impairment MCDL infesting in messaginaria faits in LOLD fine restroments, double-lind, packaging common and and advised publication of 2 years' contains were completed in non-determent adults with ADL Medical and IAC describes incolor armone more consistent on non-determent adults with ADL Medical and IAC describes incolor armone more countered in non-determent adults with ADL Medical and IAC describes incolor armone more countered in non-determent adults which ADL Medical and IAC describes incolor armone more countered in non-determent adults which armone and the consistence of a more control and a more consistence of the connection of a more control of the control and and a more control of the connection and a more connection and with MCI. Individuals with MCI demonstrate isolated memory impairment greater than expected fo with Mic. Intribution with or electrications assume removing present size electrication. The deposits of ordinar for Alzheimer's Disease. In these trials, REMINYL was not shown to be effective in patients with MIC. In the double-blind portion of these two trials, a total of 1.3 deaths in subjects on REMINYL (in=1026) were recorded and 1 death in subjects. on placebo (n=1022); the reason for this difference is currently unknown. This difference in mortalit has not been observed in PENIMY's studies in Althorner's Disease. Approximately half of the PENIMY's deaths appeared to have resulted from various vascular cases (improcradia infaction, stoke, and souther observed from the resulted from reflocus souther appeared to been souther appeared to been resulted from reflocus, oxide and cascord. There is no evidence of an increased risk of mortality when PENIMY's used in patients with mild to moderate evalence of an increases for sof manifest when Hallman. Such of patients with mich of indicates Affairmen's Disease. Programal Mismane: In a lessings sidely in which save were doed from Dieg Hallman (Hallman) and the side of organizations as a disease of a mighylady of inverseed indicates and fine fine Hallman and produced sideled windows associations and of mighylady of mich the HAPM on a might basis and it is mighylady in a study in which program risk were doesed from the leginizing of organizations froming that it is might be sometiment of the might be seen and it is mighylady, but no adverse efficies on other postated developmental parameters were seen. The doese causing above effects in rats produced slight maternal toxicity. No major malformations were caused in rats given up to 16 mg/kg/day. No drug related teratogenic effects were observed in rabbits given up to 40 mg/kg/day.G2 times the MRHD on a mg/m² bassij during the period of organogenesis. The safety of REMINYL and REMINYL FR in prepnant women has not been established. REMINYL and REMINYL FR nchammt, and an examinar, and in pegical in values as to use or escalable. Including a should not be used in women of childbearing potential unless, in the opinion of the physican, the potential benefit to the patient justifies the potential risk to the fetus. **Nursing Women:** It is not known whether galantamine is excreted in human breast milk and therefore REMINM. and REMINM. En should not be used in nursing mothers. Pediatrics: The safety and effectiveness of REMINYL and REMINYL rring in pediatric patients have not been established

eth many mises occurring in poleuric parents raye not open essousiered. ADVERSE REACTIONS <u>Clinical Trial Adverse Drug Reactions</u> Because clinical trials are conducted under very specific conditions, the adverse drug reaction rates observed in the clinical trials may not reflect the rates observed in practice and should not be compared to the rates in the clinical trials. trials of another drug. Adverse drug reaction information from clinical trials is useful for identifying drugtride of another drug. Adverse drug reaction information from district break sused for defirthly orbup-related actives event and of an aproximation great. A build or 250 professive will multi-blook case. Authorities of disease were treated with PEBMM in Places III controlled crinical studies sing other a 1-week or 4-week dose-excellation period, and 761 patients received PEBMM. 24 register, the maximum recommended maintenance does. The number of patients who compiled the studies was 1666 (75%). The mean duration of treatment for all PEBMM, groups was 150 days range 1-214 days; Adverse treats paid to 10 biocomfination Decent 195, 144(1252) of patients based with PEBMM; disconfinance from Places III controlled crinical trials due to adverse events compared to 5% 699(1159) in the packing organic for patients breadwish with PEBMM in the of deconfination due at adverse events was 14% for makes and 25% for females. In the 4-week dose-excellation freed-dose child Leif Lail 15, 11.19, 65, 6550 of deciration freed with patients with the 15 miles and 150 miles. advises events was 14 % foreigns and 22 % for termines. In the 4-week code-escalation freed-critics shortly EAL LSA-10, 96 CROSEQ of patients freed with REDMM, without one but advises event compared by 7% (2028) in the jackob group. During the dose-escalation phase of this study the inconference of decomfarings due to adverse time was 45 for patients for Foreigns 16 for REDMM. To glittly and 6% for REDMM 24 moyloting. During the maintenance phase. 4% of patients who received REDMM. 18 mg/day and 4% of patients who received REDMM. 18 mg/day and 4% of patients who received REDMM. 24 mg/day influent whom the study due to advise sevents. Table 1.1 shows the most impart adverse. events leading to discontinuation for study GAL-USA-10, in which the recommended 4-week dose

Table 1.1: Most frequent adverse events leading to discontinuation in a placebo-controlled, double-blind trial with a 4-week dose-escalation schedule (GAL-USA-10)

	Recommended 4-week dose escalation					
Adverse Events	Placebo n=286 %	16 mg/day n=279 %	24 mg/day n=273 9			
Nausea	<1	2	4			
Vomiting	0	1	3			
Anorexia	<1	1	<1			
Dizziness	<1	2	1			
Syncope	0	0	1			

Most Frequent Adverse Clinical Events Seen in Association with the Use of REMINYL. The most fro adverse events, defined as those occurring at a frequency of at least 5% and at least twice the rate of placebo in study GAL-USA-10, in which the recommended 4-week dose-escalation schedule was used are shown in Table 1.2. These events were primarily gastrointestinal and tended to occur at a lower rate with 16 mg/day, the initial recommended maintenance dose. Administration of REMMY1 with food, the use of anti-metic medication and ensuring adequate fluid intake may reduce the impact of these

with a 4-week dose increment during dose-escalation and maintenance phases (GAL-USA-10)

		Week 1-12*	Week 1-12'		Week 13-21		
Adverse Events	Placebo n=286 %		24 mg/day n=273 %		16 mg/day n=243 %		
Nausea	5	11	13	<1	4	6	
Vomiting	<1	5	6	<1	2	6	
Diarrhea	5	9	4	2	5	2	
Anorexia	2	5	5	1	2	5	

Dose escalation occurred with 4 weeks per dose increme

Tobe seculation occurred with 4 weeks per dose incoment.

The majority of these adverse events occurred during the dose-escalation period. Nausse and normling, the most inspent adverse events, courtend more frequently all higher doses, lested 57 days in most cases, and the majority of patients had one epocite, the incidence of weight case in his study was charging one second for Weight 12 (21); placebox, <1%; 16 mg/day, 3%; 24 mg/day, 3%; and orm the maintenance phase (Weise 13-21); placebox, <1%; 16 mg/day, 3%; 24 mg/day, 3%. Dose-escalation should be caudious and maintenance objects with emaintenance phase (Weise 13-21); placebox, <1%; 16 mg/day, 3%; 24 mg/day, 3%. Dose-escalation should be caudious and maintenance objects, which emails in Readile and the adjusted according to individue mess. Adverse carefasts Resourde in Controlled Tillus The reported adverse events in FIRMIM*, trais reflect experience gained under closely monitored conditions in a highlyselected patient population. In actual practice or in other clinical trials, these frequency estimates may security private in product in change placed with one can always been expected permanent in private in any one of apply, as the conditions of use, reporting behaviour and the hypes of patients treated in mig diffic. Table 1.3 lists the most common adverse events (adverse events occurring with an incidence of 24 with the following register than individence that present any other the treatment of any incident to indicence uses greates the multi-placebot treatment for four placebo controlled trials for patients treated with 16 or 24 mg/day of REMINYL. The combined values presented in Table 1.3 were derived from trials using a 1-week or the recom

Table 1.3: Adverse events reported in at least 2% of patients with Alzheimer's disease administered REMINYL and at a frequency greater than with placebo (combined 1- and

Body System/ Adverse Events	Placebo (n=801) %	REMINYL* (n=1040) %
Body as a whole - general disorders Fatigue Syncope	3 1	5 2
Central & peripheral nervous system disorders Dizziness Headache Tremor	6 5 2	9 8 3
Gastrointestinal system disorders Nausea Vomitrop Diarrhea Abdominal pain Dyspepsia	9 4 7 4 2	24 13 9 5
Heart rate and rhythm disorders Bradycardia	1	2
Metabolic and nutritional disorders Weight decrease	2	7
Psychiatric disorders Anorexia Depression Insomnia Somnolence	3 5 4 3	9 7 5 4
Red blood cell disorders Anemia	2	3
Respiratory system disorders Rhinits	3	4
Urinary system disorders Urinary tract infection Hematuria	7 2	8 3

Adverse events in patients treated with 16 or 24 mg/day of REMINYL in three placebo-controlled trials with a 1-week dose-escalation period and a 26-week fixed-dose REMINYL treatment, and one placebo-controlled trial with the recommodose REMINYL treatment are included.

No clinically relevant abnormalities in laboratory values were observed. In a cardiovascular safety Inclination present automation in accounty values ever covered in colorisation service. Inclinicating (GL-L2FF, Dipusses greater than the seconds were more common in guidantamin-treater patients than in placeb-releated patients during the case-escalation period see MARNINGS. AND PRECAUTIONS, Mack Frequent Adverse Christia Events Seen in Association with the bloom CE-TEMMELET Adverse reactions in chiract tribids of once-daily treatment with PEMMELET Revented The Committee of sules were similar to those seen with REMINYI, immediate release tablets (see Table 1.4)

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Table 1.4: Adverse events reported in at least 2% of patients with Alzheims administered REMINYL or REMINYL ER and at a frequency greater than placebo

System Organ Class Preferred Term	Placebo (n=320) %	REMINYL (n=326) %	REMINYLER (n=319) %
Body as a whole - general disorders Injury 6 Edema peripheral Fatigue Synoope Fever 1 Leg pain	3 1 1	4 2 4 1 2 2	8 4 4 2 1 <1
Central & peripheral nervous system disorders Dizziness Headache Tremor	4 6 0	7 6 1	10 8 2
Gastrointestinal system disorders Nausea Vomiting Abdominal pain Dyspepsia	5 2 2 2	14 9 3 3	17 7 2 2
Heart rate and rhythm disorders Bradycardia	2	2	3
Metabolic and nutritional disorders Weight decrease Hyperglycemia	1	5 2	4 2
Musculoskeletal system disorders Arthraigia Skeletal pain Arthritis Myalgia	2 1 1	2 3 1	3 2 2 2
Psychiatric disorders Anorexia Depression Anorexi Somnolence Depression aggravated Aggressive reaction Nervousness	3 3 3 2 1 1	7 5 1 2 2 2 2	6 6 4 3 2 2
Respiratory system disorders Rhinitis Pneumonia	3 1	4 2	4 2
Secondary terms Abrasion nost	1	1	2
Skin and appendages disorders Fash	1	<1	3
Urinary system disorders Hematuria Micturition frequency	1 1	1 2	2
Vision disorders Cataract	1	1	2

not otherwise specifie

Other Adverse Events Observed During Clinical Trials FEMINY. has been administered to 3055 patients with Alzheimer's disease during clinical trials workwide. A total of 2357 gallents received galantamine in placebo-controlled trials and 761 patients with Alzheimer's disease received galantamine 24 mg/day, the maximum recommended maintenance dose. About 1000 gatients guardiamine 24 mg/day, use incontinum comminence maintenance dose; reconstructive deservations of personal received galantiamie for at less of one year and approximately 200 patients received galantiamine for two years. To establish the rate of adverse events, data from all patients for any dose of REMINVL in 8 placebo-controlled trials and 6 open-label extension trials were pooled. The methodology to gather and paces commercial sea and open-sea execution may be pool. In the reconsol justice at an oodly here adverse exist was standarded arooms trials, using With Demmolog, All events counting in approximately (1.1% of patients are noticed, except for those already listed desember in stalling, WIO terms for govern to the information, or relately minure execution sea desember. One stalling, WIO terms for govern the information, or relately minure execution sea to counting in all seast 1/100 patients, infequent adverse events - those occurring in 1/100 to 1/1000 patients, rever-tions occurring in 1/1000 to 1/1000 patients, reviews - those occurring in these than 1/1000 to these occurring in microtio in microtio parents; lety rate - more occurring in letter in microtion parents. These address events are not recossive yeards of PSEMM. The terment and in most cases were observed at a similar frequency in placebo-freated patients in the controlled studies. <u>Body as a Whole: General Disorders: Frequency or thest pain, softwaris, here, makes, <u>Cardiospatial System Disorders: Frequent Prepriessor, infraquent pouch in placetisms, professor, depender deviand, cardiac failure, impocardal schemia or infraction. <u>Cardinal & Perpheral Nemous. System (Disorders</u></u></u> cardia claur, mocardia schema or infaction. Cardia & Feighera Herrous Selem Disordes-rintegenet verilon, protectio, consisions, motalinity musice contralisco, presellesia, attacia, phosionesia, paparia, sobiasa, lei crames, trimias, traneient scheme citato' or condrivoracioni acutivori. Gischiomisticali System Disorderis. Propient: fitalerica, infraquent: gestioni entris, fichicip, Fere escopiaga pertration integrita. Bella Rifferin Disorderis, Propient. National protection integrita. Propient Rifferin and superintical techniques, disorderistical techniques, disorderistical techniques, disorderistical status del filtrificial and superintical techniques. Or prolonge, burde branch block. T-area mension, venticular tarbupardia, Para severe tradycardia. Metabolica, Bullitaring Disorderis infraquent. Propiosiona, alianie propiosiana considera Michael Research (Propiosiana silanie) propiosiana considerational propiosi Psychiatric Disorders: Infre uent: anathy, paroniria, paranoid reaction. Ilbido increased, delirium: Rare suicidal ideation, suicide attempt. <u>Urinary System Disorders:</u> Frequent: incontinence; Infrequent hematuria, micturition frequency, cystitis, urinary retention, nocturia, renal calculi. **Post-Marke** hematura, micturition frequency, cystitis, urinary retention, nocturia, renal calculi. Post-h Adverse Drug Reactions Other adverse events from post-approval controlled and uncon clinical trials and post-marketing experience observed in patients treated with REMINYL include: Body as a Whole - General Disorders: dehydration (including rare, severe cases leading to renal insufficiency sammer interest possesses programming company in committee in the committee of the committe exacerbations of the underlying disease processes common in the elderly population

DRUG INTERACTIONS Overview Multiple metabolic pathways and renal excretion are involved in the elimination of glastratimes on o single pathway appears predominant. Based on in vitro studies, CMP206 and CMP344 were the major enzymes notwiden in the metabosism of glastratime. CMP206 was notwiden the brandant of 0 desembly plastratime, whereous PGP344 metablet the brandant of desembly plastratime, vitro of the propriate of glastratime. No could be propriate the control of glastratime. No could be propriate the control of glastratime in vitro of the propriate the control of glastratime. No could be propriate the control of glastratime in vitro of the control of glastratime in vitrol of the control of glastratime in vitrol of the control of glastratime in vitrol o erase inhibitors have the potential to interfere with the activity of anticholineroic medications oblinesterae inhibits see the potential to interies with the activity of amoroming or measures. Its will Distributions and other Considerate inhibits of sensitive self-or the operation when orbinesterae inhibits are given concurrently with successful, seinal re-unmusualer abouting agents or drollinergic agents such as between the lags, and transaction and prefers in the chain it has revised requirings, and individuals and section and prefers in the chain that sensitive interies and individuals or MEMINM. All With these longs time, limited information covering the interaction of REMINM and REMINM CRI with these longs. Distributions of covering the interaction of REMINM and REMINM CRI with these longs. Distributions of Covering the Commissions and the contraction of the contraction inhet information concerning the intension of RSMMML and RSMMML of the United States and Design Intensional Classifications (Elect of Other Dugs on the Metabolism of Galacterians Phemacokinets states to assess the optional of galacterians for intensication with crimediate, restrictine, lectocoracie, enforcement, parametric warfarm and disposit were limited to start term, mostly single-does studies in young healthy volunties. Smalls studies in reflect plantist were not love. In vigit OFFAM modellass the common of parameters and experiment of contractions of parameters device, whereas OFFAS in shoulder in the motion of or desembly opiatronisme. Because galacterinise is also glucorrolidated and excreted unchanged in urine, no single pathway appeass predictional or 1,00 metabolism and Ramifolias Calacterianis essain insistent as a single does of 4 may office 2 of 2-3 may be intensified with the contraction of 100 may play, in-6 makes and 6 females or rankforce (200 mg polity, in-6 makes and 6 females). Circelidate in crossed the bioseabilish for destrictine by accommodate 1,00 mg polity, in-6 make and 6 females of not the permacokinets. to heritases for antitudes (sout mig quary, filen) makes and or filentiases, Limitation included the bloodsability of plastinetine by approximately file. Randforder land not bell on the pharmacovinistics of gelantamine. Reflocorazoie Reflocorazoie, a strong nitribility of CP7344 and an inhibitor of CP7256, at a close of 200 mig b.d. for 4 days, forceased the ALU of algoritamine by 30% when subjects were treated with guidaminet — Aligo b.d. for 6 days in 1-8 miles and 6 filensess. Epithymorphic plantaminetis of CP7344 at a dose of 500 mig pl.d. for 4 days increased the ALU.

of calantamine by 10% when subjects received calantamine 4 mo b.i.d. for 6 days in=8 males and of paintament by 10% when subjects received guaratement and public for Gags region areas and Bermaled, Parender Poursonier, a stoom political of OPCAGE crosses APACT AND public, and bild, and 12 mg bild, guaratemine by 47%, 45%, and 46%, respectively, in 16 healthy volunteers of males and 8 femilies with received guaratement bygether with 20 mg/day parenders (Peter of Scatternine on the Metabolisms of Other Course), and political received on on the Metabolisms of Other Course (Scatternine on the Metabolisms of Other Course). paintings catagracially of or the CHTPAN, CHTP compliance to the province plant of the province plant of the province plant of the province plant of the pla Chronic in witro or in vivo studies on nicotinic receptor modulation have not been conducted. It is unknown whether galantamine has an effect on the pharmacodynamic action of other drugs that act on

Control in all not in the studies on installine receptor modulation have not been concluded it is unknown whether glostramine has an effect on the pharmacolyramic action of other drugs that act including in control in exposition. Bross-Food International Internations with food them not been established. <u>Drug-Herd Internations</u> Titlescalines in West and products have not been established. <u>Drug-Laboratory International Internations</u> Micropal state of the established of <u>DRUGHER AND ADMINISTRATION REMINITY</u>, (galantamine by pharborounistly and PERMINITY, Eleva-tor indicated for use in patients with mild cognitive impairment (see "MARNINESS AND PRECAUTIONS, Special Tropolations, Patients with Mild Cognitive Impairment (MCI), Martality in investigational Tables in Mild, PERMINIT and PERMINIT Eleva-tion of Indiany Correlation with Chinaco when are experienced in the disposition of the Control of Authorities of Season Elevality (Laboratory) and programment (MCI), Authorities of Season Elevality. Laboratory in the propriet of the disposition of the Control of Authorities of Season Elevality Laboratory and inclination of commission of the commission of the control of evening meals. REMINYI. ER extended release cansules should be administered once daily in the evening lines inclination, and make the control resized qualities about the authention of user our lines in morning, preliability with flood. Patients and caregivers should be advised to enture adequate fluid intake during treatment. **Desiring Considerations =** Opcomitient Tieschmert, in patients treated with potent CMP200 or CMP344 inhibitors, done reductions can be considered. • <u>Special Provisions</u>: Discage adjustments may be required for elderly patients (>55 years old) with low tody, velopit control of the control o (especially females), and patients with hepatic and/or renal impairment. • <u>Missed Dose:</u> The mis dose should be taken at the next scheduled dose. Doses should not be doubled. If therapy has b once show to elabora at the least scheduled uses. Doess show in the bounded, in heady has been interrupted for several days or forger, the patient should be restarted at the lovest does and the does excalated to the current does. Recommended Does and Dosage Adjustment, The disage of REMINYL shown to be effective in controlled clinical trials is 16-32 mg/day given as hince daily dosing. re-barrer, soom to be enterior the control contract has a Co. "Oppling years at the case of 25 mg/sep; sets well breaded than lover does and once of the problem of problem and enterior and of the control co ouse is om jungs, the close should be included in the minimal material cut out of in highly and weeker. If this fillial minimaters does it a seed bleated, a further increase by 24 mg/by, my per considered only after a minimum of 4 weeks at 16 mg/day. The abrupt withdrawed of FEMINYL or FEMINYL Ent in those patients with half observationing does in the effective range was not associated with an nonessed frequency of adverse weeth in comparison with those continuity to excess the same does of that drug. The beneficial effects of FEMINYL and FEMINYL Eraw lost, however, when the drug ooses or und op, in exercise custo in elevitim and relamini, that ever, in even the origin is effectively a few facilities (lose escalation) and even place (in elevitim particular caution, relapid, patricular caution, relapid, integrated (in patricular caution, relapid, integrated (in patricular caution). In elegatic impatricular caution and in elevation (in elevation) and in elevation elevation (in patricular caution). Elegatic impatricular in patricular caution (in elevation) and in elevation (in elevation) and in elevation elevation (in elevation) and in elevation used on the manifest private that the control of th with tood, or at least 1 mess, frein the costage should be incleased to thing office any for at least 4 meters. In these placests, ship does should not exceed a total of 16 mg/day, Sincon not at an analysis on the buse of PEAMM's or PEAMM's ERI in patients with severe hepotic impairment ("Disk" Puch" source of 10-19; PEAMM's and PEAMM's ERI in patients with severe impartite position (see all PEAMM's SAMP PEEAMM's SAMP Impairment if position releases with a real impartite position releases and the maintenance does should be not maintenance to see should be not maintenance to the seed and the notation of the seed and the notation of the seed and the notation of the notation o erally not exceed 16 moviday. Since no data are available on the use of REMINYL or REMINYL ER in nationts with a creatinine clearance loss than 9 ml/min. REMINYL and REMINYL FR are no mended for this population (see **WARNINGS AND PRECAUTIONS**). In a population ively-impaired individuals, sale use of this and all other medications may require supervision

OVERDOCAGE Symptoms: Overdosage with chalmesterase inhibitors can result in cholinergic crisis characterized by severe nausea, vomiting, salvelan, sweating, bradycardia, hypotension, respiratory depression, collapse and convolsions. Nicreasing muscle weakness is a possibility and may result in death if respiratory muscles are involved. In a postmarketing report, one patient who had been taking A my of galantamine deally instantiently in gasted eight 4 mg ballest 52 mg held on he helm day of treatment. Subsequently, she developed transpractia, 07 protongation, vertricolar barbycarda and treatment. Edis debianes de compression by a relief to sold consciousness or brinks the required regular treatment. Edis debianes just prior to inflation of galantamine treatment was normal. Two additional cases of accidental ingestion of 32 mg (nausea, vomiting, and dry mouth; nausea, vomiting, and cases or account in regional or 3.0 mg (installing, or or of 10 mg (installing, or or of 10 mg) excluded in their Insightalizations for other entering with full recovery. One patient, who was prescribed 24 mg/da and heat a history of hallucirations over the previous two years, mistakenly received 24 mg twice daily for 34 days and developed heat of the previous two years, mistakenly received 24 mg twice daily for 34 days and developed heat of the previous two years, mistaken years are of 16 mg/day, inadverteral viruspied. 160 mg and experienced sweating, vomiting, bradycardia, and near-syncope one hour later, which coming and experiences sensoring, interinging, complication, and intern symposite or city does, minimized necessitation frogish treatment. His symptoms revolved within 24-thours. <u>The adminimal fo</u>lialisation in less a plasma half-life of approximatably 7-8 hours. It is recommended that, in case of approximant sensoring, no hardward proximation and approximation and inclination, 4 in any case or necrouse, general supportive inseases smoot or unless Joyle and supportion of significant emolishing diplastination are proficed to be similar to frose of necrosing of other cholinomimetics. These effects generally involve the central nervious system, the parasymptathet nervious system, and the renormalization profit on the addition to muscle weekness or decolations, some or all the following signs of otheringer costs may develop seven masses, vomiting, gastromitestrial cramping, salivation, lacrimation, unitration, defecation, sweating, bradycardia, hypotension, respiratory depression, collapse and convulsions, Increasing muscl uradycaus, imporesson, vieropianov generosov, condepe and commissions in recessing insce-weakness is a possibility and may result in death if respirationy massles are involved. Teriary anticholinergics such as atroprie may be used as an artificite for galantamine overdosage, intraverous atroprine sulphate thratled to effect is recommended at an initial dose of 0.5 to 1.0 mg i.v. with subsequent doses based upon clinical response. Atypical responses in blood pressure and heart rate have been reported with other cholinomimetrs when co-administered with quaternal anticholinergics. It is not known whether galantamine and/or its metabolites can be removed by dialysis (hemodialysis, pertioneal dialysis, or hemofiltration). Dose related signs of toxicity in animats included hypoactivity, tremors, clonic convulsions, salivation, lacrimation, chromodacryorthea, muccid feces

DOSAGE FORMS REMINYL (galantamine hydrobromide), expressed as galantamine base, is available as film-coated tablets in the following strengths: 4 mg galantamine as off-white, circular, biconvex tablets with the inscription "JANSSEN" on one side and "G4" on the other side: 8 mg galantamine as multi-consular biomere tablets with the inscription TAMSSEM" on one side and "69" on the other side. 12 mg galantamine as orange-brown, circular, bicconvex tablets with the inscription "LAMSSEM" on one side and "612" on the other side. REMINYL ER (galantamine hydrobromide) extended release capsules contain white to off-white pellets. The following strengths are available: 6 mg galantamine as white opeque capsules imprinted with "G.8"; 16 mg galantamine as prix opaque capsules imprinted with "G.16"; 24 mg galantamine as caramel opaque capsules imprinted with "G.24".

Product Monograph available upon request

JANSSEN-ORTHO

Janssen-Ortho Inc., Toronto, Ontario M3C 1L9 Last revised: April 2005



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See OBC



PHARMACOLOGIC CLASSIFICATION:

ACTION AND CLINICAL PHARMACOLOGY ALTACE (ramipril) is an angiotensin converting enzyme (ACE) inhibitor

Following oral administration, ALTACE is rapidly hydrolyzed to ramiprilat, its principal

MDICATIONS AND CLINICAL USE: <u>Essential Hypertension</u>, ALTACE (ramipril) is indicated in the treatment of essential hypertension. It may be used alone or in association with thazide diuretics. ALTACE should normally be used in patients in whom treatment with a diuretic or a beta-blocker was found ineffective or has been associated with unacceptable adverse effects. ALTACE can also be tried as an initial agent in those patients in whom use of diuretics and/or beta-blockers are agent in those parents in whom use of underest abund the advokens are contraindicated or in patients with medical conditions in which these drugs frequently cause serious adverse effects. The safety and efficacy of ALTACE in renovascular hypertension have not been established and therefore, its use in this condition is not recommended. The safety and efficacy of concurrent use of ALTACE with antihypertensive agents other than thiazide diuretics have not been established.

antinypercensive agents other than funzable united states in a laver his deep issabilished. Treatment Following Acute Myocardial Infarction ALTAC's is indicated following acute myocardial infarction in clinically stable patients with signs of left ventricular dysfunction to improve survival and reduce hospitalizations for heart failure. Sufficient experience in the treatment of patients with severe (NYTA class IV) heart failure immediately after myocardial infarction is not yet available. (See WARNINGS – Hypotension.)

not yet available. (See WARNINGS – Hypotension.)

MANAGEMENT OF PATIENTS AT INCREASED RISK OF CARDIOVASCULAR

EVENTS: ALTACE may be used to reduce the risk of myocardial infarction, stroke or
cardiovascular death in patients over 55 years of age who are at high risk of
cardiovascular events because of a history of coronary artery disease, stroke,
peripheral artery disease, or diabetes that is accompanied by at least one other
cardiovascular risk factor such as hypertension, elevated total cholesterol levels, low
high density lipoprotein levels, cigarrette smoking, or documented microalbuminuria.
The incidence of the primary outcome (composite of myocardial infarction, stroke and
death from cardiovascular causes) was reduced from 17.8% in the placebo-treated
group to 14.0% in the ramipril-treated group.

group to 14.0% in the ramipri-treated group.

GENERAL: In using ALTACE consideration should be given to the risk of angioedema (see WARNINGS). When used in pregnancy during the second and third trimesters, ACE inhibitors can cause injury or even death of the developing fetus. When pregnancy is detected ALTACE should be discontinued as soon as possible (see WARNINGS — Use in Pregnancy, and INFORMATION FOR THE PATIENT).

CONTRAINDICATIONS: ALTACE (ramipril) is contraindicated in patients who are hypersensitive to this drug, or to any ingredient in the formulation, or in those patients who have a history of angioedema.

who have a history of angioedema. Wannings: Angioedema has been reported in patients with ACE inhibitors, including ALTACE (ramipril). Angioedema associated with laryngeal involvement may be fatal. If laryngeal stridor or angioedema of the face, tongue, or glottis occurs, ALTACE should be discontinued immediately, the patient treated appropriately in accordance with accepted medical care, and carefully observed until the swelling disappears. In instances where swelling is confined to the face and lips, the condition generally resolves without treatment, although antihistamines may be useful in relieving symptoms. Where there is involvement of tongue, glottis, or larynx, likely to cause airway obstruction, appropriate therapy (including, but not limited to 0.3 to 0.5 mL of subcutaneous epinephrine solution 1.1000) should be administered promptly (see ADVERSE REACTIONS).

The incidence of angioedema during ACE inhibitor therapy has been reported to be higher in black than in non-black patients. Patients with a history of angioedema unrelated to ACE inhibitor therapy may be at increased risk of angioedema while receiving an ACE inhibitor (see CONTRAINDICATIONS).

Hypotension: Symptomatic hypotension has occurred after administration of ALTACE Hypotension: Symptomatic hypotension has occurred after administration of ALTACE. usually after the first or second dose or when the dose was increased, it is more likely to occur in patients who are volume depleted by diuretic therapy, dietary salt restriction, dialysis, darrhea, or vomiting, in patients with ischemic heard disease or erebrovascular disease, an excessive fall in blood pressure could result in a myocardial infarction or cerebrovascular accident (see ADVERSE REACTIONS). Because of the potential fall in blood pressure in these patients, therapy with ALTACE should be started under close medical supervision. Such patients should be followed closely for the first weeks of treatment and whenever the dose of ALTACE is increased. In patients with severe congestive heart failure, with or without associated ornal insufficiency, ACE inhibitor therapy may cause excessive hypotension and has been associated with oliguria, and/or progressive azotemia, and rarely, with acute renal failure and/or death.

If hypotension occurs, the patient should be placed in a supine position and, if necessary If hypotension occurs, the patient should be placed in a supine position and, if necessary, receive an intravenous infusion of 0.9% sodium chloride. A transient hypotensive response may not be a contraindication to further doses which usually can be given without difficulty once the blood pressure has increased after volume expansion in hypertensive patients. However, lower doses of ALTACE and/or reduced concomitant durretic therapy should be considered. In patients receiving treatment following acute myocardial infarction, consideration should be given to discontinuation of ALTACE (see ADVERSE REACTIONS — Treatment Following Acute Myocardial Infarction, DOSAGE AND ADMINISTRATION — Treatment Following Acute Myocardial Infarction, Disages and the properties of the patients of the patients

ANU Jannist First III — Treatment Foliowing Acute Myocardial infarction). Neutropenia/faranulocytosis: Agranulocytosis and bone marrow depression have been caused by ACE inhibitors. Several cases of agranulocytosis, neutropenia or leukopenia have been reported in which a causal relationship to ALTACE cannot be excluded. Current experience with the drug shows the incidence to be rare. Periodic monitoring of white blood cell counts should be considered, especially in patients with collagen vascular disease and/or renal disease; <u>Use in Pregnancy</u>, ACE inhibitors can cause fetal and neonatal morbidity and mortality when administered to pregnant women. Several dozen cases have been reported in the world literature. When pregnancy is detected, ALTACE should be discontinued as soon as possible.

PRECAUTIONS: Renal Impairment: As a consequence of inhibiting the renin angiotensin-aldosterone system, changes in renal function have been seen in susceptible individuals. In patients whose renal function may depend on the activity of the renin-angiotensin-aldosterone system, such as patients with bilateral rena of the renin-angiotensin-aldosterone system, such as patients with bilateral renal artery stenosis, unlateral renal artery stenosis to a solitary kidney, or severe congestive heart failure, treatment with agents that inhibit this system has been associated with oliguria, progressive azotemia, and rarely, acute renal failure and/or death. In susceptible patients, concomitant diuretic use may further increase risk. Use of ALTACE should include appropriate assessment of renal function. ALTACE should be used with caution in patients with renal insufficiency as they may require reduced or less frequent doses (see DOSAGE AND ADMINISTRATION). Close monitoring of renal function during therapy should be performed as deemed appropriate in patients with renal insufficiency.

Anaphylactoid Reactions during Membrane Exposure; Anaphylactoid reactions have been reported in patients dialyzed with high-flux membranes (e.g. polyacrylonitrile [PAN]) and treated concomitantly with an ACE inhibitor. Dialysis should be stopped immediately if symptoms such as nausea, abdominal cramps, burning, angloedema, shortness of breath and severe hypotension occur. Symptoms are not relieved by antihistamines, in these patients consideration should be given to using a different type of dialysis membrane or a different class of antihypertensive agents.

Anaphylactoid Reactions during Desensitization: There have been isolated reports of

patients experiencing sustained life threatening anaphylactoid reactions while receiving ACE inhibitors during desensitization treatment with hymenoptera (bees, wasps) venom. In the same patients, these reactions have been avoided when ACE inhibitors were temporarily withheld for at least 24 hours, but they have reappeared upon inadvertent rechallenge

upon inadvertent rechallenge.

Hyperkalemia and Potassium-Sparing Diuretics; Elevated serum potassium (greater than 5.7 mEq.L) was observed in approximately 1% of hypertensive patients in clinical trials treated with ALTACE. In most cases these were isolated values which resolved despite continued therapy, Hyperkalemia was not a cause of discontinuation of therapy in any hypertensive patient. Risk factors for the development of hyperkalemia may include renal insufficiency, diabetes mellitus, and the concomitant use of agents to treat hypokalemia or other drugs associated with increases in serum potassium (see PRECAUTIONS – Drug Interactions).

<u>Surgery/Anesthesia</u>; In patients undergoing surgery or anesthesia with agents producing hypotension, ALTACE may block angiotensin If formation secondary to compensatory renin release. If hypotension occurs and is considered to be due to this mechanism, it may be corrected by volume repletion.

Aortic Stenosis: There is concern, on theoretical grounds, that patients with aortic stenosis might be at particular risk of decreased coronary perfusion when treated with vasodilators because they do not develop as much afterload reduction.

Patients with Impaired Liver Function: Hepatitis (hepatocellular and/or cholestatic), elevations of liver enzymes and/or serum bilirubin have occurred during therapy with ACE inhibitors in patients with or without pre-existing liver abnormalities. In most cases the changes were reversed on discontinuation of the drug.

cases the changes were reversed on discontinuation of the origi.

Elevations of liver enzymes and/or serum bilirubin have been reported with ALTACE (see ADVERSE REACTIONS). Should the patient receiving ALTACE experience any unexplained symptoms particularly during the first weeks or months of treatment, it is recommended that a full set of liver function tests and any other necessary investigations be carried out. Discontinuation of ALTACE should be considered when investigations be care out inscontinuation in Autor. Should be chiscolered winer appropriate. There are no adequate studies in patients with cirrhosis and/or liver dysfunction. ALTACE should be used with particular caution in patients with pre-existing liver abnormalities. In such patients baseline liver function tests should be obtained before administration of the drug and close monitoring of response and metabolic effects should apply.

Nursing Mothers: Ingestion of a single 10 mg oral dose of ALTACE resulted in undetectable amounts of ramipril and its metabolities in breast milk. However, because multiple doses may produce low milk concentrations that are not predictable from single doses, ALTACE should not be administered to nursing mothers.

<u>Pediatric Use</u>: The safety and effectiveness of ALTACE in children have not been established; therefore use in this age group is not recommended.

Use in Elderly: Although clinical experience has not identified differences in response between the elderly (>65 years) and younger patients, greater sensitivity of some older individuals cannot be ruled out.

<u>Patient Alertness</u>: ALTACE may lower the state of patient alertness and/or reactivity, particularly at the start of treatment (see ADVERSE REACTIONS).

particularly at the start of treatment (see ADVERSE REACTIONS).

Cough: A dry, persistent cough, which usually disappears only after withdrawal or lowering of the dose of ALTACE, has been reported. Such possibility should be considered as part of the differential diagnosis of cough.

Purg Interactions: Concomitant Diuretic Therapy: Hypotension may result but can be minimized by discontinuing diuretic or increasing salt intake prior to ramipril treatment and/or reducing initial dose. Agents increasing serum potassium: Use potassium sparing diuretics with caution and monitor frequently. Agents causing renin release; ALTACE arthlypertensive effect increased. Lithium: Lithium levels may be increased. Administer tithium with caution and monitor levels frequently. Antacids: The bioavailability of ALTACE and the pharmacokinetics of ramiprilat were not affected. Digoxin: No change in ramipril; ramiprilat or digoxin serum levels. Warfarin: affected. Digogin; No change in ramipril, ramiprilar of digoxin serum levels. Warfarin; The co-administration of ALTACE with warfarin did not after the anticoagulant effects. Acenocoumarol; No significant changes. Non-steroidal anti-inflammatory agents (NSAID): The antihypertensive effects of ACE inhibitors may be reduced with concomitant administration of NSAIDs (e.g. indomethacin).

NON-BOLD. The administration of NSAIDs (e.g., indomethacin).

ADVERSE REACTIONS: Essential Hypertension, Serious adverse events occurring in North American placebo-controlled clinical trials with ramipril monotherapy in hypertension (n=972) were: hypotension (0.1%); myocardial infarction (0.3%); cerebrovascular accident (0.1%), edema (0.2%); syncope (0.1%), Among all North American ramipri platients [n=1,244], angiodelma occurred in patients treated with ramipril and a diuretic (0.1%). The most frequent adverse events occurring in these trials with ALTACE monotherapy in hypertensive patients (n=651) were: headache (15.1%); disziness (3.7%); asthenia (3.7%); chest pain (2.0%); nausea (1.8%); peripheral edema (1.8%); somelone (1.7%); impotence (1.5%); rash (1.4%); arthritis (1.1%); dispense (1.1%). Discontinuation of therapy due to clinical adverse events was required in 5 patients (0.8%), in placebo-controlled trials, an excess of upper respiratory infection and flu syndrome was seen in the ramipril group. As these studies were carried out before the relationship of cough to ACE inhibitors was recognized, some of these events may represent ramipril-induced cough. In a later 1-year study, increased cough was seen in almost 12% of ALTACE patients, with of patients treated with ALTACE monotherapy in North American controlled clinical rials (n=972) have required discontinuation because of cough.

Treatment Following Acute Myocardial Infarction

trials (n=972) nave required under the street of the stree Treatment Following Acute Myocardial Infarction
Adverse events (except laboratory abnormalities) in a controlled clinical trial of post
Adverse events (except laboratory abnormalities) in a controlled clinical trial of post
Adverse events (except laboratory abnormalities) in a controlled clinical trial of post
Adverse events (except laboratory abnormalities) in a controlled clinical trial of post
Advised patients (n=1,004) were:
hypotension (1,0%); increased cough (7,6%); dizziness/vertigo (5,6%);
nausea/vomiting (3,6%); angina pectoris (2,9%); postural hypotension (2,2%);
syncope (2,1%); heart failure (2,0); severo/resistant heart failure (2,0%; myocardial
infarction (1,7%); vomiting (1,6%); headache (1,2%); abnormal kidney function
(1,2%); abnormal kidney
(1, increases in serum potassium.

DOSAGE AND ADMINISTRATION

Essential Hypertension: Dosage of ALTACE (ramipril) must be individualized. Initiation of therapy requires consideration of recent antihypertensive drug treatment, the extent of blood pressure elevation and salt restriction. The dosage of other antihypertensive agents being used with ALTACE may need to be adjusted.

Monotherapy. The recommended initial dosage of ALTACE in patients not on diuretics is 2.5 mg once daily. Dosage should be adjusted according to blood pressure response, generally, at intervals of at least two weeks. The usual dose range is 2.5 to 10 mg once daily. A daily dose of 20 mg should not be exceeded.

In some patients treated once daily, the antihypertensive effect may diminish towards the end of the dosing interval. This can be evaluated by measuring blood pressure just prior to dosing to determine whether satisfactory control is being maintained for 24 hours. If it is not, either twice daily administration with the same total daily dose, 24 hours. In its 10x, while twice dualy administration with the same total dualy of or an increase in dose should be considered, if blood pressure is not controlled w ALTACE alone, a diuretic may be added. After the addition of a diuretic, it may possible to reduce the dose of ALTACE.

Concomitant Diuretic Therapy: Symptomatic hypotension occasionally may occur following the initial dose of ALTACE and is more likely in patients who are currently being treated with a diuretic. The diuretic should, if possible, be discontinued for two

to three days before beginning therapy with ALTACE to reduce the likelihood of hypotension (see WARNINGS). If the diuretic cannot be discontinued, an initial dose of 1.25 mg of ALTACE should be used with careful medical supervision for several hours and until blood pressure has stabilized. The dosage of ALTACE should subsequently be titrated (as described above) to the optimal response.

subsequently be titrated (as described above) to the optimal response.

Nes in Renal Impairment. For patients with a creatinine clearance below 40 mL/min/

1.73 m² (serum creatinine above 2.5 mg/dL), the recommended initial dose is

1.25 mg of ALTACE once daily. Dosage may be titrated upward until blood pressure is controlled or to a maximum total daily dose of 5 mg. In patients with severe renal impairment (creatinine clearance below 10 mL/min/1.73 m²) the maximum total daily dose of 2.5 mg of ALTACE should not be exceeded.

Treatment Following Acute Myocardial Infarction:

Treatment Following Acute Myocardial Infarction: Initiation of therapy requires consideration of concomitant medication and baseline blood pressure and should be instituted under close medical supervision, usually in a hospital, three to ten days following an acute myocardial infarction in haemodynamically stable patients with clinical signs of heart failure. The recommended initial dosage of ALTACE is 2.5 mg given twice a day (b.i.d.), one in the morning and one in the evening, if tolerated, and depending on the patient's response, dosage may be increased by doubling at intervals of one to three days. The maximum daily dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed 5 mg twice daily (b.i.d). After the initial dose of ALTACE should not exceed

Patients who have been fluid or salt depleted, or treated with diuretics are at an increased risk of hypotension; see WARNINGS – Hypotension). An excessive fall in blood pressure may occur particularly in the following: after the initial dose of ALTACE; after every first increase of dose of ALTACE; after the first dose of a concomitant direction and when increasing the dose of the concentrant direction. It accommiss that direction and when increasing the dose of the concentrant direction. It appropriate, the dose of any concomitant direction should be reduced which may diminish the likelihood of hypotension (see PRECAUTIONS – Drug Interactions). Consideration should be given to reducing the initial dose to 1.25 mg of ALTACE in these patients. <u>Use in Renal Impairment:</u> In patients with impaired renal function (creatinine clearance of 20-50 mL/min/1.73 m* body surface area), the initial recommended dosage is generally 1.25 mg of ALTACE once daily. This dosage may be increased with caution up to 1.25 mg of ALTACE twice daily, depending upon clinical response and

Insufficient data is available concerning the use of ramipril following acute myocardial infarction in patients with heart failure and severe renal failure, (see ACTION AND CLINICAL PHARMACOLOGY – Pharmacokinetics and Metabolism, PRECAUTIONS – Renal Impairment).

Renal impairment). Use in Hepatic Impairment: Insufficient data is available concerning the use of ramipril following acute myocardial infarction in patients with heart failure and hepatic dysfunction. Dose reduction and careful monitoring of these patients is required (see ACTIONS AND CLINICAL PHARMACOLOGY – Pharmacokinetics and Metabolism, PRECAUTIONS — Patients with Impaired Liver Function).

PRECAUTIONS – Patients with Impaired Liver Function).

Management of Patients at Increased Risk of Cardiovascular Events:
Recommended initial dose: 2.5 mg of ALTACE once daily. Depending on the
tolerability, the dose is gradually increased. It is recommended to double the dose
after one week of treatment and – after another three weeks – to increase it to
10 mg. Usual maintenance dose: 10 mg of ALTACE daily (see ACTION AND CLINICAL
PHARMACOLOGY, WARNINGS and PRECAUTIONS). Dosage recommendation or special risk groups such as patients with renal or hepatic impairment, or at an
increased risk of trypotension (fluid or sat depletion, treated with discretics) are to be
followed as previously described (see WARNINGS and PRECAUTIONS).

DOSAGE FORM

a) Composition

a) Composition ALTACE (ramjoril) capsules 1.25 mg, 2.5 mg, 5.0 mg, and 10.0 mg contain the medicinal ingredient ramjoril in quantities of 1.25 mg, 2.5 mg, 5.0 mg, and 10.0 mg respectively. The qualitative formulation for all potencies of ALTACE is: ramjoril, pre-gelatinized starch NF (as filler, gliding agent and disintegration agent) and empty gelatin capsules. Empty gelatin capsules for all potencies of ALTACE are composed of gelatin NF and coloring agents specific to each potency (see below)

POTENCY	CAP	BODY
1.25 mg	Yellow iron oxide Titanium dioxide	Titanium dioxide
2.5 mg	Yellow iron oxide FD & C red no. 3 Titanium dioxide	Titanium dioxide
5.0 mg	FD & C blue no. 2 FD & C red no. 3 Titanium dioxide	Titanium dioxide
10.0 mg	FD & C blue no. 2 FD & C red no. 3 Black iron oxide Titanium dioxide	Titanium dioxide

b) Stability and storage recommendations Store ALTACE (ramipril) in original container at room temperature, below 25°C and not beyond the date indicated on the container.

AVAILABILITY: No. 4 hard gelatin capsules:

- 1.25 mg (white/yellow);2.5 mg (white/orange);5.0 mg (white/red);
- . 10.0 mg (white/blue).

ALTACE capsules 1.25 mg, 2.5 mg, 5.0 mg and 10.0 mg are packaged in cartons of 30 (2 x 15 blister-packed) capsules. Bottles of 100 capsules and 500 capsules also

Product monograph available upon request.

Neterences:

1. ALTACE Product Monograph. 2. The Heart Outcomes Prevention Evaluation Study Investigators (HOPE) Trial. Effects of an angiotensin-converting-enzyme inhibitor, ramipril, on cardiovascular events in high-risk patients. N Engl J Med 2000:342(3):145-53

Distributed by sanofi-aventis Canada Inc. Laval, Quebec H7L 4A8





0.5 mg and 1 mg Capsules (Nabilone)

ACTION

"CESAMET" (nabilone) is a synthetic cannabinoid with antiemetic properties which have been found to be of value in the management of some patients with nausea and vomiting associated with cancer chemotherapy. It also has sedative and psychotropic effects.

After oral administration, comparable peak plasma levels of nabilone and of its carbinol metabolite were attained within 2 hours. The combined plasma concentrations of nabilone and of its carbinol metabolite accounted for, at most, 10 to 20% of the total radiocarbon concentration in plasma. The plasma half-life of nabilone was approximately 2 hours, while that of the total radiocarbon was of the order of 35 hours.

Of the two major possible metabolic pathways, stereo-specific enzymatic reduction and direct enzymatic oxidation, the latter appears to be the more important in man.

The drug and its metabolites are eliminated mainly in the feces (approximately 65%) and to a lesser extent in the urine (approximately 20%). The major excretory pathway is the bilary system.

INDICATIONS

"CESAMET® (nabilane) is indicated in adults for the management of severe nausea and vomiting associated with concer chemotherapy.

CONTRAINDICATIONS

"CESAMET® (nabilione) is contraindicated in patients with known sensitivity to marijuana or other cannabinoid agents, and in those with a history of psychotic reactions.

WARNINGS

"CESAMET" (nabilione) should be used with extreme caution in patients with severe liver dysfunction and in those with a history of non-psychotic emotional disorders.

"CESAMET® should not be taken with alcohol, sedatives, hypnotics, or other psychotomimetic substances.

"CESAMET® should not be used during pregnancy, in nursing mothers, or pediatric patients since its safety under these conditions has not been established.

PRECAUTIONS

Since "CESAMET" (nabilione) will often impair the mental and/or physical abilities required for the performance of potentially hazardous tasks, such as driving a car and operating machinery, the patient should be warned accordingly and should not be permitted to drive or engage in dangerous tasks until the effects of nabilione are no longer greent.

Adverse psychotropic reactions can persist for 48 to 72 hours following cessation of treatment.

Since "CESAMET" elevates supine and standing heart rates and causes postural hypotension, it should be used with caution in the elderly and in patients with hypertension or heart disease.

Drug Interactions: Potential interactions between "CESAMET", and diazepam; sodium secobarbital; alcohol; or codeine, were evaluated. The depressant effects of the combinations were additive. Psychomotor function was particularly impaired with concurrent use of diazepam.

Pediatric Use: The safety and efficacy in children under the age of 18 has not been established. Therefore the use of "CESAMET" in this patient population is not recommended.

ADVERSE REACTIONS

The most frequently observed adverse reactions to nabilone and their incidences reported in the course of clinical trials were as follows: drowsiness (66.0%), vertigo (58.8%), psychological high (38.8%), dry mouth (21.6%), depression (14.0%), atoxia (12.8%), blurred vision (12.8%), sensation disturbance (12.4%), anaroexia (7.6%), asthenia (7.6%), headache (7.2%), orthostatic hypotension (5.2%), euphoria (4.0%) and hallucinations (2.0%).

The following adverse reactions were observed in less than 1% of the patients who were administered nabilone in the course of the clinical trials: tachycardia, tremors, syncope, nightmares, distortion in the perception of time, confusion, dissociation, dysphoria, psychotic reactions and seizures.

Spontaneously Reported Adverse Reactions: The following adverse reactions listed in order of decreasing frequency by body system have been reported since *CESAMET* has been marketed. All events are listed regardless of causality assessment.

Blood and Hematopoetic: Leukopenia

Cardiovascular: Hypotension and tachycardia

Eye and Ear: Visual disturbances

Gastrointestinal: Dry mouth, nausea, vomiting, and constipation

Nervous System: Hallucinations, CNS depression, CNS stimulation, ataxia, stupor, vertigo, convulsion, and circumoral paresthesia

Psychiatric: Somnolence, confusion, euphoria, depression, dysphoria, depersonalization, anxiety, psychosis, and emotional lability

Miscellaneous and Ill-Defined Conditions: Dizziness, headache, insomnia, abnormal thinking, chest pain, lack of effect, and face edema

SYMPTOMS AND TREATMENT OF OVERDOSE

Signs and Symptoms: Signs and symptoms which might be expected to occur are psychotic episodes including hallucinations, anxiety reactions, respiratory depression and coma (experience with cases of overdosage of more than 10 mg/day has not yet been reported).

Treatment: Overdosage may be considered to have occurred, even at prescribed dosages, if disturbing psychiatric symptoms are present. In these cases, the patient should be observed in a quiet environment and supportive measures, including reassurance, should be used. Subsequent doses should be withheld until patients have

returned to their baseline mental status; routine dosing may then be resumed if clinically indicated. In such instances, a lower initiating dose is suggested.

If psychotic episodes occur, the patient should be managed conservatively, if possible. For moderate psychotic episodes and anxiety reactions, verbal support and comforting may be sufficient. In more severe cases, antipsychotic drugs may be useful; however, the utility of antipsychotic drugs in cannobinoid psychosis has not been systematically evaluated. Support for their use is drawn from limited experience using antipsychotic agents to manage cannobis overdoses. Because of the potential for drug-drug interactions (eg., additive CNS depressant effects due to nabilone and chlorpromazine), such patients should be closely monitored.

Protect the patient's airway and support ventilation and perfusion. Meticulously monitor and maintain, within acceptable limits, the patient's vital signs, blood gases, serum electrolytes, etc. Absorption of drugs from the gastrointestinal tract may be decreased by giving activated charcool, which, in many cases, is more effective than emesis or lavage; consider charcoal instead of or in addition to gastric emptying. Repeated doses of charcoal over time may hosten elimination of some drugs that have been absorbed. Safeguard the patient's airway when employing gastric emptying or charcoal.

The use of forced diuresis, peritoneal dialysis, hemodialysis, charcoal hemoperfusion, or cholestyramine has not been reported. In the presence of normal renal function, most of a dose of nabilone is eliminated through the biliary system.

Treatment for respiratory depression and comatose state consists in symptomatic and supportive therapy. Particular attention should be paid to the occurrence of hypothermia. If the patient becomes hypotensive, consider fluids, inotropes, and/or vasopressors.

DOSAGE AND ADMINISTRATION Adults:

The usual dosage of "CESAMET" (nabilone) is 1 mg or 2 mg twice a day. The first dose should be given the night before initiating administration of chemotherapeutic medication. The second dose is usually administrated 1 to 3 hours before chemotherapy. If required, administration of "CESAMET" can be continued up to 24 hours after the chemotherapeutic agent is given. The maximum recommended daily dose is 6 mg in divided doses.

"CESAMET" is available in a 0.5 mg strength for dose adjustment within the therapeutic range. Dose adjustment may be required for the purposes of response and tolerance in individual patients. Overdosage may occur even at prescribed dosages, if disturbing psychiatric symptoms are present. In these cases, the patient should be observed in a quiet environment and supportive measures, including reassurance, should be used. Subsequent doses should be withheld until patients have returned to their baseline mental status; routine dosing may then be resumed if clinically indicated. In such instances, a lower initiating dose is suggested.

"CESAMET® contains nabilione in a capsule dosage form and is intended only for oral administration.

STRUCTURAL FORMULA AND CHEMISTRY

Molecular Formula: C₂₄H₃₆O₃ Molecular Weight: 372 U.S.A.N: Nabilone

Chemical Name: $trans(\pm)-3-(1,1-dimethylheptyl)-6,6a,7,8,10,10a-hexahydro-1-hydroxy-6,$

6-dimethyl-9H-dibenzo(b,d),pyran-9-one.

Description: White crystalline powder

Composition

Each 1 mg "CESAMET" capsule contains 1 mg of nabilone, starch, povidone, gelatin, FD&C blue #2 (indigo carmine), red iron oxide and titanium dioxide.

Each 0.5 mg "CESAMET" capsule contains: 0.5 mg of nabilone, starch, povidone, gelatin, titanium dioxyde, D&C red # 33, D&C yellow # 10, FD&C red # 40.

Stability and storage Recommendations

Store at controlled room temperature at 15-30°C.

AVAILABILITY

"CESAMET® 1 mg capsule: each No. 2 hard gelatin capsule, opaque blue cap and white body, imprinted ICN logo on the cap and 3101 on the body, contains 1 mg of nabilone and are available in bottles of 20 capsules.

"CESAMET" 0.5 mg capsule: each 10.4 hard gelatin capsule, opaque red cap and white body, imprinted ICN logo on the cap and 10.2 on the body, contains 0.5 mg of nabilione and are available in bottles of 50 capsules.

"CESAMET® (nabilone) legally is considered to be a narcotic and is subject to the controls which apply to those drugs.

References

- 1. Cesamet (nabilone) Product monograph. Valeant Canada Limited.
- 2. Grotenhermen F and Russo E. Cannabis and Cannabinoids: Pharmacology, Toxicology and Therapeutic Potential. The Haworth Press, Inc. 2002: xxviii.

Product Monograph available upon request



A-15 See page A-11



BETASERON'

THERAPEUTIC CLASSIFICATION

ACTION AND CLINICAL PHARMACOLOGY

Description: BETASERON® (interferon beta-1b) is a purified, sterile, lyophilized protein product produced by recombinant DNA techniques and formulated for use by injection. Interferon beta-1b is manufactured by bacterial fermentation of a strain of Escherichia coli that bears a genetically engineered plasmid containing the gene for human interferon beta_{ser17}. The native gene was obtained from human fibroblasts and altered in a way that substitutes serine for the cysteine residue found at position 17. Interferon beta-1b is a highly purified protein that has 165 amino acids and an approximate molecular weight of 18,500 daltons. It does not include the carbohydrate side chains found in the natural material

General: Interferons are a family of naturally occurring proteins, which have molecular weights ranging from 15,000 to 21,000 daltons. Three major classes of interferons have been identified: alpha, beta, and gamma. Interferon beta-1b, interferon alpha, and interferon gamma have overlapping yet distinct biologic activities. The activities of interferon beta are species-restricted and, therefore, the most pertinent pharma cological information on BETASERON (interferon beta-1b) is derived from studies of human cells in culture and in vivo.

Biologic Activities: Interferon beta-1b has been shown to possess both antiviral and immunomodulatory activities. The mechanisms by which BETASERON exerts its actions in multiple sclerosis (MS) are not clearly understood. However it is known that the biologic response-modifying properties of interferon beta-1b are mediated through its interactions with specific cell receptors found on the surface of human cells. The binding of interferon beta-1b to these receptors induces the expression of a number of interferon-induced gene products (e.g., 2',5'-oligoadenylate synthetase, protein kinase and indoleamine 2,3-dioxygenase) that are believed to be the mediators of the biological actions of interferon beta-1b. A number of these interferon-induced products have been readily measured in the serum and cellular fractions of blood collected from patients treated with interferon beta-1b.

INDICATIONS AND CLINICAL USE

BETASERON (interferon beta-1b) is indicated for:

• the reduction of the frequency of clinical exacerbations in

- ambulatory patients with relapsing-remitting multiple sclerosis. Relapsing-remitting MS is characterized by recurrent attacks of neurologic dysfunction followed by complete or incomplete recovery.
- the slowing of progression in disability and the reduction of the frequency of clinical exacerbations in patients with secondary-progressive multiple sclerosis.

The safety and efficacy of BETASERON in primary progressive MS have not been evaluated.

CONTRAINDICATIONS

BETASERON (interferon beta-1b) is contraindicated in patients with a history of hypersensitivity to natural or recombinant interferon beta, Albumin Human USP, or any other component of the formulation.

WARNINGS

The administration of cytokines to patients with a pre-existing monoclonal gammopathy has been associated with the development of systemic capillary leak syndrome with shock like symptoms and fatal outcome.

In the RR-MS clinical trial, one suicide and four attempted suicides were observed among 372 study patients during a 3-year period. All five patients received BETASERON (interferon beta-1b) (three in the 0.05 mg [1.6 MIU] group and two in the 0.25 mg [8.0 MIU] group). There were no attempted suicides in patients on study who did not receive BETASERON. In the SP-MS study there were 5 suicide attempts in the placebo group and 3 in the BETASERON group including one patient in each group who committed suicide. Depression and suicide have been reported to occur in patients receiving interferon alpha, a related compound. Patients treated with BETASERON should be informed that depression and suicidal ideation may be a side effect of the treatment and should report these symptoms immediately to the prescribing physician. Patients exhibiting depression should be monitored closely and cessation of therapy should be considered.

PRECAUTIONS

General: Rare cases of cardiomyopathy have been reported. If this occurs, and a relationship to BETASERON (interferon beta-1b) is suspected, treatment should be discontinued

Rare cases of thyroid dysfunction (hyper- as well as hypothyroidism) associated with the use of BETASERON have

been reported.

Symptoms of flu syndrome observed with BETASERON therapy may prove stressful to patients with severe cardiac conditions. Patients with cardiac disease such as angina, congestive heart failure or arrhythmia should be monitored osely for worsening of their clinical conditions.

Information to be Provided to the Patient: Patients

should be instructed in injection techniques to assure the safe self-administration of BETASERON (See below and the

BETASERON® INFORMATION FOR THE PATIENT section. Instruction on Self-injection Technique and Procedure.

It is recommended that the first injection be administered or under the direct supervision of, a physician. Appropriate instructions for reconstitution of RETASERON and self-injection using aseptic techniques, should be given to the patient.

A careful review of the **BETASERON® INFORMATION FOR**

THE PATIENT section is also recommended.

Patients should be cautioned against the re-use of needles or syringes and instructed in safe disposal procedures. Information on how to acquire a puncture-resistant container for disposal of used needles and syringes should be given to the

patient along with instructions for safe disposal of full containers. Overall, 80% of patients in the two controlled clinical trials reported injection site reactions at one or more times during therapy. Post-marketing experience has been consistent with

this finding, with infrequent reports of injection site necrosis.

The onset of injection site necrosis usually appears early in therapy with most cases reported to have occurred in the first two to three months of therapy. The number of sites where necrosis has been observed was variable.

Rarely, the area of necrosis has extended to subcutaneous fat or fascia. Response to treatment of injection site necrosis with antibiotics and/or steroids has been variable. In some of these patients elective debridement and, less frequently, skin grafting took place to facilitate healing which could take from three to six months.

Some patients experienced healing of necrotic skin lesions while BETASERON therapy continued. In other cases new necrotic lesions developed even after therapy was discontinued

The nature and severity of all reported reactions should be carefully assessed. Patient understanding and use of aseptic self-injection technique and procedures should be periodically

Flu-like symptoms are not uncommon following initiation of therapy with BETASERON. In the controlled MS clinical trials acetaminophen was permitted for relief of fever or myalgia.

Patients should be cautioned not to change the dosage or the schedule of administration without medical consultation. **Awareness of Adverse Reactions:** Patients should be advised about the common adverse events associated with the use of BETASERON,particularly, injection site reactions and the

flu-like symptom complex (see **ADVERSE REACTIONS**).

Patients should be cautioned to report depression or suicidal ideation (see WARNINGS).

Patients should be advised about the abortifacient potentia of BETASERON (see **PRECAUTIONS**, **Use in Pregnancy**).

Laboratory Tests: The following laboratory tests are recommended prior to initiating BETASERON therapy and at periodic intervals thereafter: thyroid function test, hemoglobin, complete and differential white blood cell counts, platelet counts and blood chemistries including liver function tests A pregnancy test, chest roentgenogram and ECG should also be performed prior to initiating BETASERON therapy. In the controlled MS trials, patients were monitored every 3 months. The study protocol stipulated that BETASERON therapy be discontinued in the event the absolute neutrophil count fell below 750/mm³. When the absolute neutrophil count had returned to a value greater than 750/mm³, therapy could be restarted at a 50% reduced dose. No patients were withdrawn or dose-reduced for neutropenia or lymphopenia. Similarly, if AST/ALT (SGOT/SGPT) levels exceeded 10 times

the upper limit of normal, or if the serum bilirubin exceeded 5 times the upper limit of normal, therapy was discontinued. In each instance during the controlled MS trial, hepatic enzyme abnormalities returned to normal following discontinuation of therapy. When measurements had decreased to below these levels, therapy could be restarted at a 50% dose reduction, if clinically appropriate. Dose was reduced in two patients due to increased liver enzymes; one continued on treatment and one was ultimately withdrawn.

Drug Interactions: Interactions between BETASERON and other drugs have not been evaluated. Although studies designed to examine drug interactions have not been done, it was noted that BETASERON patients (n=180) have received corticosteroid

or ACTH treatment of relapses for periods of up to 28 days.

BETASERON administered in three cancer patients over a dose range of 0.025 mg (0.8 MIU) to 2.2 mg (71 MIU) led to a dose-dependent inhibition of antipyrine elimination. The effect of alternate-day administration of 0.25 mg (8 MIU) BETASERON on drug metabolism in MS patients is unknown

Interferons have been reported to reduce the activity of hepatic cytochrome P450-dependent enzymes in humans and animals. Caution should be exercised when BETASERON is administered in combination with agents that have a narrow therapeutic index and are largely dependent on the hepatic cytochrome P450 system for clearance.

Impairment of Fertility: Studies in female rhesus monkeys with normal menstrual cycles, at doses up to 0.33 mg (10.7 MILI)/kg/day (equivalent to 32 times the recommended human dose based on body surface area comparison) short no apparent adverse effects on the menstrual cycle or on associated hormonal profiles (progesterone and estradiol) when administered over 3 consecutive menstrual cycles. The extrapolability of animal doses to human doses is not known Effects of BETASERON on women with normal menstrua cycles are not known

Use in Pregnancy: BETASERON was not teratogenic at doses up to 0.42 mg (13.3 MIU)/kg/day in rhesus monkeys, but

demonstrated dose-related abortifacient activity when administered at doses ranging from 0.028 mg (0.89 MIL)/kg/day (2.8 times the recommended human dose based on body surface area comparison) to 0.42 mg (13.3 MIL)/kg/day (40 times the recommended human dose based on body surface area comparison)
The extrapolability of animal doses to human doses is not known. Lower doses were not studied in monkeys. Spontaneous abortions while on treatment were reported in 4 patients who participated in the BETASERON RR-MS clinical trial, whereas there was one induced abortion in each of the placebo and BETASERON groups in the SP-MS trial. BETASERON given to rhesus monkeys on gestation days 20 to 70 did not cause tera-togenic effects; however, it is not known if teratogenic effects exist in humans. There are no adequate and well-controlled studies in pregnant women. Women of childbearing potential should take reliable contraceptive measures. If the nationt becomes pregnant or plans to become pregnant while taking BETASERON, the patient should discontinue therapy. It is not nown if interferons alter the efficacy of oral contraceptives.

Nursing Mothers: It is not known whether BETASERON is

excreted in human milk. Given that many drugs are excreted in human milk, there is a potential for serious adverse reactions in nursing infants, therefore a decision should be made whether to

scontinue nursing or discontinue BETASERON treatment.

Pediatric Use: Safety and efficacy in children under

18 years of age have not been established.

Dependence Liability: No evidence or experience suggests that abuse or dependence occurs with BETASERON therapy; however, the risk of dependence has not been syst

ADVERSE REACTIONS

The following adverse events were observed in placebo-controlled clinical studies of BETASERON (interferon beta-1b), at the recommended dose of 0.25 mg (8 MIU), in patients with elapsing-remitting MS (n=124) and secondary-progressive MS (n=360):

1. Relapsing-remitting MS: Injection site reactions (85%) and injection site necrosis (5%) occurred after administration of BETASERON. Inflammation, pain, hypersensitivity, necrosis, and non-specific reactions were significantly associated (p<0.05) with the 0.25 mg (8 MtJ) BETASERON-treated group, compared to placebo. Only inflammation, pain, and necrosis were reported as severe events. The incidence rate for injection site reactions was calculated over the course of 3 years. This incidence rate decreased over time, with 79% of patients experiencing the event during the first 3 months of treatment ared to 47% during the last 6 months. The median time to the first occurrence of an injection site reaction was 7 days Patients with injection site reactions reported these events 183.7days per year. Three patients with drew from the 0.25 mg (8 MIU) BETASERON-treated group for injection site pain

Flu-like symptom complex was reported in 76% of the patients treated with 0.25 mg (8 MILI) BETASERON, A patient was defined as having a flu-like symptom complex if flu-like syndrome or at least two of the following symptoms were concurrently reported: fever, chills, myalgia, malaise or sweating.

Only myalgia, fever, and chills were reported as severe in more than 5% of the patients. The incidence rate for flu-like symptom complex was also calculated over the course of 3 years. The incidence rate of these events decreased ove time, with 60% of patients experiencing the event during the first 3 months of treatment compared to 10% during the last 6 months. The median time to the first occurrence of flu-like symptom complex was 3.5 days and the median duration per patient was 7.5 days per year.

- Laboratory abnormalities included: lymphocyte count < 1500/mm³ (82%),
- ALT (SGPT) > 5 times baseline value (19%), absolute neutrophil count < 1500/mm³ (18%)
- (no patients had absolute neutrophil counts < 500/mm³),

 WBC < 3000/mm³ (16%), and

 total bilirubin > 2.5 times baseline value (6%).

Three patients were withdrawn from treatment with 0.25 mg

(8 MIU) BETASERON for abnormal liver enzymes including of following dose reduction (see **PRECAUTIONS**, **Laboratory** Tests).

one (28%) of the 76 females of childbearing age treated at 0.25 mg (8 MIU) BETASERON and 10 (13%) of the 76 females of childbearing age treated with placebo reported menstrual disorders. All reports were of mild to moderate severity and included: intermenstrual bleeding and spotting early or delayed menses, decreased days of menstrual flow, and clotting and spotting during menstruation.

Mental disorders such as depression, anxiety, emotiona lability, depersonalization, suicide attempts and confusion were observed in this study. Two patients withdrew for confusion. One suicide and four attempted suicides were also reported. It is not known whether these symptoms may be related to the underlying neurological basis of MS, to BETASERON treatment, or to a combination of both. Some similar symptoms have been noted in patients receiving interferon alpha and both interferons are thought to act through the same receptor. Patients who experience the symptoms should be monitored closely and cessation of therapy should be considered.

Additional common clinical and laboratory adverse events associated with the use of BETASERON are listed in the following paragraphs. These events occurred at an incidence of 5% or more in the 124 MS patients treated with 0.25 mg (8 MIU) BETASERON every other day for periods of up to 3 years in the controlled trial, and at an incidence that was at least twice that observed in the 123 placebo patients Common adverse clinical and laboratory events associated with the use of BETASERON were:

- injection site reaction (85%).
- lymphocyte count < 1500/mm³ (82%)
- ALT (SGPT) > 5 times baseline value (19%),
- absolute neutrophil count < 1500/mm3 (18%).
- menstrual disorder (17%), WBC < 3000/mm³ (16%),
- palpitation (8%)
- dyspnea (8%),
- cystitis (8%).
- hypertension (7%),
- breast pain (7%)
- tachycardia (6%), gastrointestinal disorders (6%),
- total bilirubin > 2.5 times baseline value (6%).
- somnolence (6%),
- Jaryngitis (6%)
- pelvic pain (6%
- menorrhagia (6%)
- injection site necrosis (5%), and peripheral vascular disorders (5%).

A total of 277 MS patients have been treated with BETASERON in doses ranging from 0.025 mg (0.8 MILI) to 0.5 mg (16 MIU). During the first 3 years of treatment, withdrawals due to clinical adverse events or laboratory abnormalities not mentioned above included:

• fatigue (2%, 6 patients),

- cardiac arrhythmia (< 1%, 1 patient)
- allergic urticarial skin reaction to injections (< 1%, 1 patient),
- headache (< 1%, 1 patient).
- unspecified adverse events (< 1%, 1 patient), and
- "felt sick" (< 1%, 1 patient)

The table that follows enumerates adverse events and laboratory abnormalities that occurred at an incidence of 2% or more among the 124 MS patients treated with 0.25 mg (8 MIU) BETASERON every other day for periods of up to 3 years in the controlled trial and at an incidence that was at least 2% more than that observed in the 123 placebo patients. Reported adverse events have been re-classified using the standard COSTART glossary to reduce the total number of terms employed in Table 1. In the following table, terms so general as to be uninformative, and those events where a drug cause was remote have been excluded.

le 1: Adverse Events and Laho

Adverse Event	Placebo n=123	0.25 mg (8 MIU) n=124
Body as a Whole		11-124
Injection site reaction*	37%	85%
Headache	77%	84%
Fever*	41%	59%
Flu-like symptom complex*	56%	76%
Pain	48%	52%
Asthenia*	35%	49%
Chills*	19%	46%
Abdominal pain	24%	32%
Malaise*	3%	15%
Generalized edema	6%	8%
Pelvic pain	3%	6%
Injection site necrosis*	0%	5%
Cyst	2%	4%
Necrosis	0%	2%
Suicide attempt	0%	2%
Cardiovascular System		
Migraine	7%	12%
Palpitation*	2%	8%
Hypertension	2%	7%
Tachycardia	3%	6%
Peripheral vascular disorder	2%	5%
Hemorrhage	1%	3%
Digestive System		
Diarrhea	29%	35%
Constipation	18%	24%
Vomiting	19%	21%
Gastrointestinal disorder	3%	6%
Endocrine System		
Goiter	0%	2%
Hemic and Lymphatic System		
Lymphocytes < 1500/mm ³	67%	82%
ANC < 1500/mm ³ *	6%	18%
WBC < 3000/mm ³ *	5%	16%
Lymphadenopathy	11%	14%
Metabolic and Nutritional Disord	ers	
ALT (SGPT) > 5 times baseline*	6%	19%
Glucose < 55 mg/dL	13%	15%
Total bilirubin > 2.5 times baselin	e 2%	6%
Urine protein > 1+	3%	5%
AST (SGOT) > 5 times baseline*	0%	4%
Weight gain	0%	4%
Weight loss	2%	4%
Musculoskeletal System		
Myalgia*	28%	44%
Myasthenia	10%	13%



Nervous System		
Dizziness	28%	35%
Hypertonia	24%	26%
Depression	24%	25%
Anxiety	13%	15%
Nervousness	5%	8%
Somnolence	3%	6%
Confusion	2%	4%
Speech disorder	1%	3%
Convulsion	0%	2%
Hyperkinesia	0%	2%
Amnesia	0%	2%
Respiratory System		
Sinusitis	26%	36%
Dyspnea*	2%	8%
Laryngitis	2%	6%
Skin and Appendages		
Sweating*	11%	23%
Alopecia	- 2%	4%
Special Senses		
Conjunctivitis	10%	12%
Abnormal vision	4%	7%
Urogenital System		
Dysmenorrhea	11%	18%
Menstrual disorder*	8%	17%
Metrorrhagia	8%	15%
Cystitis	4%	8%
Breast pain	3%	7%
Menorrhagia	3%	6%
Urinary urgency	2%	4%
Fibrocystic breast	1%	3%
Breast neoplasm	0%	2%

^{*} significantly associated with BETASERON treatment (p<0.05)

It should be noted that the figures cited in Table 1 cannot be used to predict the incidence of side effects in the course of usual medical practice where patient characteristics and other factors differ from those that prevailed in the clinical trials. The cited figures do provide the prescribing physician with some basis for estimating the relative contribution of drug and nondrug factors to the side effect incidence rate in the population studied.

 Secondary-progressive MS: The incidence of adverse events that occurred in at least 2% of patients treated with 8 MIU BETASERON or placebo for up to three years, orwhere an adverse event was reported at a frequency at least 2% higher with BETASERON than that observed for placebotreated patients in the secondary-progressive study, is presented in Table 2. Adverse events significantly associated with BETASERON compared to placebo (p<0.05) are also indicated in Table 2

Table 2: Incidence of Adverse Events \geq 2% or > 2% Difference (BETASERON vs. Placebo) in the Secondary

riugiessive ma suuy		
Adverse Event	Placebo n=358	0.25 mg (8 MIU) n=360
Body as a Whole		
Asthenia	58%	63%
Flu syndrome*	40%	61%
Pain	25%	31%
Fever*	13%	40%
Back pain	24%	26%
Accidental injury	17%	14%
Chills*	7%	23%
Pain in Extremity	12%	14%
Infection	11%	13%
Abdominal pain*	6%	11%
Malaise	5%	8%
Neck pain	6%	5%
Abscess*	2%	4%
Laboratory test abnormal	1%	3%
Allergic reaction	3%	2%
Chills and fever*	0%	3%
Thorax pain	2%	1%

Cardiovascular System	4%	COV
Vasodilatation Peripheral vascular disorder	4% 5%	6% 5%
Chest pain	4%	5%
Migraine	3%	4%
Hypotension	4%	2%
Hypertension*	2%	4%
Palpitation	3%	2%
Syncope	3%	2%
Hemorrhage	2%	2%
Tachycardia	1%	2%
Digestive System		
Nausea	13%	13%
Constipation	12%	12%
Diarrhea	10%	7%
Gastroenteritis Vomiting	5% 6%	6% 4%
Dysphagia	5%	4%
Gastrointestinal disorder	5%	4%
Tooth disorder	4%	4%
Dyspepsia	4%	4%
Anorexia	2%	4%
Fecal incontinence	3%	2%
Liver function test abnormal	1%	3%
Gastritis	2%	2%
Flatulence	1%	3%
Sore throat	1%	2%
Colitis	2%	0%
Gastrointestinal pain	0%	2%
Gingivitis	0%	2%
lemic and Lymphatic System		
Leukopenia*	5%	10%
Anemia	5%	2%
Ecchymosis	2%	1%
Lymphadenopathy	1%	3%
njection Site	100/	400/
Injection site reaction* Injection site inflammation*	10%	46% 48%
Injection site inflammation	4% 5%	9%
Injection site pain	0%	5%
Injection site hemorrhage	2%	2%
Metabolic and Nutritional Disord		2.70
Peripheral edema	7%	7%
Weight loss	3%	2%
SGPT increased	2%	2%
Hypercholesteremia	2%	1%
Musculoskeletal System		
Myasthenia	40%	39%
Arthralgia	20%	20%
Myalgia*	9%	23%
Bone fracture (not spontaneous)	5%	3%
Muscle cramps	3%	3%
Spontaneous bone fracture	3%	3%
Arthritis	1%	2%
Joint disorder	1%	2%
Nervous System	4404	47000
Headache	41%	47%
Neuropathy	41%	38%
Paresthesia	39%	35%
Hypertonia*	31% 34%	41% 34%
Abnormal gait Depression	34%	27%
Ataxia	23%	19%
Dizziness	14%	14%
Incoordination	120/	110/

Vertigo

Tremor

Neuralgia Movement disorde

Anxiety

Sleep disorde

Hypesthesia

Nervousness

Incoordination

Emotional lability Paralysis

Sweating increased

Somnolence

Vaginal moniliasis Kidney pain Pyelonephritis Prostatic disorde *significantly associated with BETASERON treatment (p<0.05) Seventy-four (74) patients discontinued treatment due to adverse events (23 on placebo and 51 on BETASERON). Injection site reactions were significantly associated with early termination of treatment in the BETASERON group compared to placebo (p<0.05). The highest frequency of adverse events leading to discontinuation involved the nervous system, of which depression (7 on placebo and 11 on BETASERON) was the most common. Significantly more patients on active therapy (14.4% vs. 4.7% on placebo) had elevated ALT (SGPT) values (>5 times

Menorrhagia

Nocturia

14% 13% 8% 12% 11% 10% 8% 9% 6% 7% 6% 5% 5% 4%

11%

8% 8%

8%

8%

6%

6% 5% 5% 6%

6%

Speech disorder Dysarthria 4% 1% 2% 3% 2% 2% 1% 1% 1% 1% Spastic paralysis Convulsi 2% 2% 3% 2% Hyperesthesia Dry mouth 2% 2% 2% Heminlegia Thinking abnormal Myoclonus Respiratory System 32% 28% Rhinitis 20% 12% 16% 9% Pharyngitis 10% 6% 5% Cough increased Sinusitis 5% 6% 5% 3% Pneumonia Upper respiratory tract infection 3% 1% 1% 2% Voice alteration Skin and Append 12% 20% Pruritus 6% 4% 4% 6% Skin disorde 4% 2% 3% 2% Eczema Hernes simple 2% 2% Alopecia 2% 3% 3% Acne 2% 1% 1% 1% Dry skin Subcutaneous hematoma Breast pain 1% 1% Herpes zoster 2% 2% Seborrhea 15% 11% Abnormal vision Amblyopia 10% Diplopia 9% 5% 3% 3% 2% 7% 4% 2% 2% 3% 1% Eye pain Otitis media Conjunctivitis Deafness 3% Optic neuritis Ear disorder 2% 2% 2% 2% 2% 1% 1% Tinnitus Urogenital System 25% 22% Urinary tract infection 15% 8% 7% Urinary tract disorder 10% 9% 7% Cystitis 7% 8% Urinary urgency 13% 5% 6% Menstrual disorder Increased urinary frequency 9% 6% 12% 4% 3% 2% 2% 2% 2% 2% 0% Metrorrhagia Urinary retention 6% 4% Vaginitis Amenorrhea 4% 4% 4% 4% 1% 2% 0% Dysuria Impotence Menopause

and gamma-GT values in the BETASERON group throughout the study. In the BETASERON group, most ALT (SGPT) abnormalities resolved spontaneously with continued treatment whereas some resolved upon dose reduction or temporary discontinuation of treatment.

Lymphopenia (<1500/mm³) was observed in 90.9% of ETASERON patients compared to 74.3% of placebo patients and neutropenia (<1400/mm³) was noted in 18.0% BETASERON and 5.1% placebo patients.

baseline value). Flevations were also observed in AST (SGOT)

DOSAGE AND ADMINISTRATION FOR SUBCUTANEOUS USE ONLY

BETASERON (interferon beta-1b) should only be prescribed by (or following consultation with) dinicians who are experienced in the diagnosis and management of multiple sclerosis.

The recommended dose of BETASERON for both relapsing-

remitting and secondary-progressive MS patients is 0.25 mg (8 MIU) injected subcutaneously every other day. Limited data regarding the activity of a lower dose in relapsing-remitting MS patients are presented above (see **ACTION AND CLINICAL PHARMACOLOGY, Clinical Trials**).

In the secondary-progressive MS study, patients initiated treatment with half the dose (4 MIU s.c. every other day) for a period of 2 weeks prior to escalating to the recommended dose of 8 MIU (s.c. every other day).

Efficacy of treatment for longer than 2 years has not been substantially demonstrated in relapsing-remitting multiple sclerosis. For secondary-progressive multiple sclerosis, safety and efficacy data beyond 3 years are not available.

To reconstitute lyophilized BETASERON for injection, use a

sterile syringe and needle to inject 1.2 mL of the diuent supplied, Sodium Chloride, 0.54% Solution, into the BETASERON vial. Gently swirl the vial of BETASERON to dissolve the drug com-pletely; do not shake. Inspect the reconstituted product visually and discard the product before use if it contains particulate matter or is discolored. After reconstitution with accompanying diluent, each mL of solution contains 0.25 mg (8 MIU) interferon

beta-1b, 13 mg Albumin Human USP and 13 mg Mannitol USP.
Withdraw 1 mL of reconstituted solution from the vial into a sterile syringe fitted with a 27-gauge 1/2-inch needle and inject the solution subcutaneously. Sites for self-injection include abdomen, buttocks and thighs, A vial is suitable for single use only; unused portions should be discarded (See BETASERON® [interferon beta-1b] INFORMATION FOR THE PATIENT section for SELF-INJECTION PROCEDURE.)

AVAILABILITY OF DOSAGE FORMS

BETASERON (interferon beta-1b) is presented in single-use vials of lyophilized powder containing 0.3 mg (9.6 MlU) interferon beta-1b, 15 mg Albumin Human USP, and 15 mg
Mannitol, USP. BETASERON is supplied in cartons containing 15 vials of medication and 15 vials of diluent (2 mL of Sodium Chloride 0.54% solution, per vial).

Product Monograph available upon request.

REFERENCES:

- Data on file. Berlex Canada Inc., 1999
- Product Monograph of PBETASERON® (Interferon beta-1b), Berlex Canada, June 1999.
- Series varieties, Julie 1999.

 3. The IFNB Multiple Sclerosis Study Group and the University of British Columbia MG/MRI Analysis Group, Interferor beta-to in the treatment of multiple sclerosis: Final outcome of the randomised controlled trial. Neurology 1995; 45:1227-1285.

2260 32nd Avenue, Lachine, Québec H8T 3H4







SUMMARY PRODUCT

Classification

Route of	Dosage Form /	Clinically Relevant		
Administration	Strength	Nonmedicinal Ingredients		
Oral	Capsules, 25 mg, 50 mg, 75 mg, 150 mg, 300 mg	Lactose monohydrate For a complete listing, see Dosage Forms, Composition and Packaging section.		

INDICATIONS AND CLINICAL USE

Adults: LYRICA (pregabalin) is indicated for the management of neuropathic pain associated with:

- · Diabetic peripheral neuropathy and
- · Postherpetic neuralgia

Geriatrics (>65 years of age): Pregabalin oral clearance tended to decrease with increasing age. This decrease in pregabalin oral clearance is consistent with age-related decreases in creatinine clearance. Reduction of pregabalin dose may be required in patients who have age-related compromised renal function. (see **WARNINGS AND PRECAUTIONS**, Geriatrics [>65 years of age])

Pediatrics (<18 years of age): The safety and efficacy of pregabalin in pediatric patients (<18 years of age) have not been established and its use in this patient population is not recommended (see **WARNINGS AND PRECAUTIONS**, Pediatrics).

CONTRAINDICATIONS

Patients who are hypersensitive to pregabalin or to any ingredient in the formulation or component of the container.

WARNINGS AND PRECAUTIONS

Tumorigenic Potential

In standard preclinical in vivo lifetime carcinogenicity studies of pregabalin, a high incidence of hemangiosarcoma was identified in two different strains of mice (see **Preclinical Toxicology**). The clinical significance of this finding is uncertain. Clinical experience during pregabalin's premarketing development provides no direct

means to assess its potential for inducing tumors in humans.

In clinical studies across various patient populations, comprising 6396 patientyears of exposure in 8666 patients ranging in age from 12 to 100 years, new or worsening-preexisting tumors were reported in 57 patients. The most common malignant tumor diagnosed was skin carcinoma [17 patients] followed by breast manignant tumor diagnosed was skill carcinoma (1) patients) provided by treast carcinoma (8) patients), prostatic carcinoma (6) patients), carcinoma not otherwise specified (6) patients) and bladder carcinoma (4) patients). Without knowledge of the background incidence and recurrence in similar populations not treated with LYRICA (pregabalin), it is impossible to know whether the incidence seen in these cohorts is or is not affected by treatment.

Ophthalmological Effects

opiniamilological recess
In controlled studies, pregabalin treatment was associated with vision-related adverse events such as blurred vision (amblyopia) (6% pregabalin and 2% placebo) and diplopia (2% pregabalin and 0.5% placebo). Approximately 1% of pregabalin-treated patients discontinued treatment due to vision-related adverse events (primarily blurred vision). Of the patients who did not withdraw, the blurred vision resolved with continued dosing in approximately half of the cases (see Post-

Marketing Adverse Drug Reactions)

Marketing Adverse Drug Reactions:

Prospectively planned ophthalmologic testing, including visual acuity testing, formal visual field testing and dilated funduscopic examination, was performed in over 3600 patients. In these patients, visual acuity was reduced in 7% of patients treated with pregabalin, and 5% of placebo-treated patients. Visual field changes were detected in 13% of pregabalin-treated, and 12% of placebo-treated patients. Funduscopic changes were observed in 2% of pregabalin-treated, and 2% of placebo-treated patients. At this time, clinical significance of the ophthalmologic findings is unknown.

Patients should be informed that if changes in vision occur, they should notify their publicing in the visual disturbance partiests further assessment including discontinuation.

physician. If visual disturbance persists, further assessment, including discontinuation of pregabalin, should be considered. More frequent assessments should be considered for patients who are already routinely monitored for ocular conditions.

Peripheral Edema

In controlled clinical trials pregabatin treatment caused peripheral edema in 6% of patients (336/5509) compared with 2% of patients (42/2384) in the placebo group. In these studies 0.5% (28/5508) of pregabalin patients and 0.2% (4/2384) of placebo patients withdrew due to peripheral edema (see **ADVERSE REACTIONS**, Peripheral Edema).

under to peripheral eveniral size ADVCHAS. ARCHOTONS, Tempheral couries in controlled clinical trials of up to 13 weeks in duration of patients without clinically significant heart or peripheral vascular disease, there was no apparent association between peripheral edema and cardiovascular complications such as hypertension or congestive heart failure. In the same trials, peripheral edema was not associated with laboratory changes suggestive of deterioration in renal or hepatic function.

Higher frequencies of weight gain and peripheral edema were observed in patients taking both LYRICA (pregabalin) and a thiazolidinedione antidiabetic agent compared to patients taking either drug alone. The majority of patients using thiazoldimedione antidiabetic agents in the overall safety database were participants in studies of pain associated with diabetic peripheral neuropathy. In this population, peripheral edema was reported in 3% (2/60) of patients who were using thiazoldimedione antidiabetic was reported in 3% (Z/60) of patients who were using hiszolidinedione antidiabetic agents only, 8% (68/859) of patients who were treated with pregabalin only, and 19% (23/120) of patients who were on both pregabalin and thiazolidinedione antidiabetic agents. Similarly, weight gain was reported in 0% (0/60) of patients on thiazolidinediones only, 4% (58/859) of patients on pregabalin only, and 7.5% (9/120) of patients on both drugs.

As the thiazolidinedione class of antidiabetic drugs can cause weight gain and/or fluid referition, possibly exacerbating or leading to heart failure, care should be taken when co-administering LYRICA and these agents.

Because there are limited data on congestive heart failure patients with New York Heart Association (NYHA) Class III or IV cardiac status, LYRICA should be used with caution in these patients

Weight Gain

Pregabalin treatment was associated with weight gain. In pregabalin controlled clinical trials of up to 13 weeks, a gain of 7% or more over baseline weight was observed in 8% of pregabalin-treated patients and 2% of placebo-treated patients. Few patients treated with pregabalin (0.2%) withdrew from controlled trials due to weight gain (see **ADVERSE REACTIONS**, Weight Gain). Pregabalin associated weight gain was related to dose and duration of exposure, but did not appear to be associated with baseline BMI, gender or age. Weight gain was not limited to patients with edema (see WARNINGS AND PRECAUTIONS, Peripheral Edema)

Although weight gain was not associated with clinically important changes in blood pressure in short-term controlled studies, the long-term cardiovascular effects of pregabalin-associated weight gain are unknown.

Among diabetic patients, pregabalin-treated patients gained an average of 1.6 kg (range: 16 to 16 kg), compared to an average 0.3 kg (range: -10 to 9 kg) weight gair in placebo patients. In a cohort of 333 diabetic patients who received pregabalin for at least 2 years, the average weight gain was 5.2 kg.

While the effects of pregabalin-associated weight gain on glycemic control have not been systematically assessed, in controlled and longer-term open label clinical trials with diabetic patients, pregabalin treatment did not appear to be associated with loss of glycemic control (as measured by $HbA_{\rm IC}$).

Dizziness and Somnolence

In controlled neuropathic pain studies, pregabalin caused dizziness in 23% of patients (424/1831) compared to 7% in placebo (58/857). Somnolence was experienced by 14% (256/1831) and 4% (33/857) of the patients treated with pregabalin and placebo, respectively. These events begin shortly after the initiation of therapy and generally occur more frequently at higher doses. In these studies, dizziness and somnolence led to withdrawal of 3.5% and 2.6% of the pregabalin-treated patients, respectively. For the remaining patients (359 and 208, respectively) who experienced these events, dizziness and somnolence persisted until the last dose of pregabalin in 43% and 58% of the patients, respectively (see **ADVERSE REACTIONS**, Tables 2 and 4, and Post-Marketing Adverse Drug Reactions).

Accordingly, patients should be advised not to drive or operate complex machinery or engage in other hazardous activities until they have gained sufficient experience on pregabalin to gauge whether or not it affects their mental and/or motor performance

adversely (see CONSTIMER INFORMATION) **Abrupt or Rapid Discontinuation**

llowing abrupt or rapid discontinuation of pregabalin, some patients reported symptoms including insomnia, nausea, headache and diarrhea. Pregabalin should be tapered gradually over a minimum of one week rather than discontinued abruptly (see **ADYERSE REACTIONS**, Adverse Events Following Abrupt or Rapid be tapered gradua

Sexual Function/Reproduction

Impairment of Male Fertility

Preclinical Data

In fertility studies in which male rats were orally administered pregabalin (50 to 2500 mg/kg) prior to and during mating with untreated females, a number of adverse reproductive and developmental effects were observed. These included decreased sperm counts and sperm motility, increased sperm abnormalities, reduced fertility, increased preimplantation embryo loss, decreased litter size, decreased fetal body weights and an increased incidence of fetal abnormalities. Effects on sperm and fertility parameters were reversible in studies of this duration (3-4 months). The no-effect dose for male reproductive toxicity in these studies (100 mg/kg) was associated with a plasma pregabalin exposure (AUC) approximately 3 times human exposure at the maximum recommended dose (MRD) of 600 mg/day.

leaposite at the indumini recommendeur observiming to obtoing use. In addition, adverse effects on reproductive organ (testes, epididymides) histopathology were observed in male rats exposed to pregabalin (500 to 1250 mg/kg) in general toxicology studies of four weeks or greater duration. The no-effect does for male reproductive organ histopathology in rats (250 mg/kg) was associated with a plasma exposure approximately 8 times human exposure at the MRD.

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In a fertility study in which female rats were given pregabalin (500, 1250 or 2500 mg/kg) orally prior to and during matting and early gestation, disrupted estrous cyclicity and an increased number of days to mating were seen at all doses and embryolethality occurred at the highest dose. The low dose in this study produced a plasma exposure approximately 9 times that in humans receiving the MRD. A noeffect dose for female reproductive toxicity in rats was not established. The clinical reprofusers were finding in a programmed to the productive toxicity in rats was not established. The clinical reprofusers are finding in a programmed to the productive toxicity in rats was not established. significance of female fertility findings in animals is unknown

Human Data

In a double-blind, placebo-controlled clinical trial to assess the effect of pregabalin on sperm motility, 30 healthy male subjects were exposed to pregabalin 600 mg/day for 3 months (one complete sperm cycle). Pregabalin did not exhibit significant day ion similaris (one complete symmetric preparation of healthy male subjects, as measured by semen analysis, when compared with placebo (n=16). However, due to the small sample size and short-term exposure to pregabalin (only one complete sperm cycle), no conclusions can be made regarding possible reproductive effects of pregabalin during long-term exposure. Effects on other male reproductive parameters in humans have not been adequately studied.

Special Populations

Renal

Because pregabalin is eliminated primarily by renal excretion, the dose of pregabalin

a related for olderly nations with renal impairment (see ACTION) AND CLINICAL PHARMACOLOGY and DOSAGE AND ADMINISTRATION)

Adjustment of Dose in Renally-Impaired Patients

In patients with a medical history of significant renal insufficiency, daily dosages should be reduced accordingly (see Table in **DOSAGE AND ADMINISTRATION, Dosing Considerations**).

Preclinical Data

Pregabalin was not teratogenic in mice, rats or rabbits. Pregabalin induced fetal toxicity in rats and rabbits at ≥39 times the mean human exposure at the maximum recommended clinical dose of 600 mg/day |AUC_{0.24} of 123 µg *hr/mL]. In the prenatal-postnatal toxicity study, pregabalin induced offspring developmental toxicity in rats at ≥5 times the maximum recommended human exposure. No developmental effects occurred at 2 times the maximum recommended human exposure (see **PRODUCT**

MONOGRAPH

Human Data

Pregnant Women

There are no adequate and well-controlled studies in pregnant women. Pregabalin should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Labour and Delivery

The effects of pregabalin on labour and delivery in pregnant women are unknown. In the prenatal-postnatal study in rats, pregabalin prolonged gestation and induced dystocia at exposures ≥47 times the mean human exposure [AUC_{I0.24)} of im recommended clinical dose of 600 mg/day (see

PRODUCT MONOGRAPH)

It is not known if pregabalin is excreted in human breast milk; however, it is present in the milk of rats. Because of the potential for adverse reactions in nursing infants from pregabalin, a decision should be made whether to discontinue nursing or to , taking into account the importance of the drug to the mother (see PRODUCT MONOGRAPH)

Pediatrics (<18 years of age)

The safety and efficacy of pregabalin in pediatric patients (<18 years of age) have not been established

Geriatrics (>65 years of age)

Of the 1831 patients who received pregabalin in neuropathic pain studies, 528 were 65 to 74 years of age, and 452 were 75 years of age or older. No significant differences in efficacy were observed between these patients and younger patients. Pregabalin oral clearance tended to decrease with increasing age. This decrease in pregabalin oral clearance is consistent with age-related decre

clearance. Reduction of pregabalin dose may be required in patients who have age related compromised renal function. In general, the incidence of adverse events did not increase with age.

Creatine Kinase Elevations

Pregabalin treatment was associated with creatine kinase elevations. Mean changes Pregabalin treatment was associated with creatine kinase elevations. Mean changes in creatine kinase from baseline to the maximum value were 60 U/I, for pregabalin-treated patients and 28 U/L for the placebo patients. In all controlled trials across multiple patient populations, 2% of patients on pregabalin and 1% of placebo patients had a value of creatine kinase at least three times the upper limit of normal. Three pregabalin-treated subjects had events reported as rhabdomyolysis in premarketing clinical trials. The relationship between these myopathy events and pregabalin is not completely understood because the cases had documented factors that may have caused or contributed to these events. Prescribers should instruct patients to promptly report unexplained muscle pain, tenderness or weakness, particularly if these muscle symptoms are accompanied by malaise or fever. Pregabalin treatment should be discontinued if myopathy is diagnosed or suspected or if markedly elevated creatine

Laboratory Changes, Decreased Platelet Count

Pregabalin treatment was associated with a decrease in platelet count. Pregabalintreated subjects experienced a mean maximal decrease in platelet count of 20×10^{3} /µL, compared to 11×10^{3} /µL in placebo patients. In the overall database of controlled trials, 2% of placebo patients and 3% of pregabalin patients experienced a potentially clinically significant decrease in platelets, defined as 20% below baseline value and $<150 \times 10^3/\mu L$.

In randomized controlled trials, pregabalin was not associated with an increase in ding related adverse eve

ECG Changes, PR Interval Prolongation

Pregabalin treatment was associated with mild PR interval prolongation. In analyses of clinical trial ECG data, the mean PR interval increase was 3-6 msec at pregabalin doses a 200 mg/day. This mean change difference was not associated with an increased risk of PR increase a 25% from baseline, an increased percentage of subjects with on-treatment PR >200 msec, or an increased risk of adverse events of nd or third degree AV block.

Information for Patients

Dizziness and Somnolence

Patients should be counseled that LYRICA (pregabalin) may cause dizziness, somnolence, blurred vision and other CNS signs and symptoms. Accordingly, they should be advised not to drive, operate complex machinery or engage in other hazardous activities until they have gained sufficient experience on pregabalin to gauge whether or not it affects their mental, visual and/or motor performance

Visual Disturbances

Patients should be counseled that LYRICA may cause visual disturbances. Patients should be informed that if changes in vision occur, they should notify their physician (see WARNINGS AND PRECAUTIONS, Ophthalmologic Effects).

Abrupt or Rapid Discontinuation

Patients should be advised to take LYRICA as prescribed. Abrupt or rapid discontinuation may result in insomnia, nausea, headache or diarrhea.

Edema and Weight Gain

Patients should be counseled that LYRICA may cause edema and weight gain

Recents should be advised that LYRICA may cause edema and weight gain. Patients should be advised that concomitant treatment with LYRICA and a thiazolidinedione antidiabetic agent may lead to an additive effect on edema and weight gain. For patients with preexisting cardiac conditions, this may increase the risk of heart failure.

Muscle Pain, Tenderness or Weakness

Patients should be instructed to promptly report unexplained muscle pain, tenderness or weakness, particularly if accompanied by malaise or fever.

Concomitant Treatment with CNS Depressants, Alcohol

Patients who require concomitant treatment with central nervous system depressants such as opiates or benzodiazepines should be informed that they may experience additive CNS side effects, such as somnolence.

Patients should be told to avoid consuming alcohol while taking LYRICA, as LYRICA may potentiate the impairment of motor skills and sedation of alcohol.

Pregnant Woman

Patients should be instructed to notify their physician if they become pregnant or ntend to become pregnant during therapy, and to notify their physician if they are preast-feeding or intend to breast-feed during therapy.

Animal Studies in Male Reproduction

In preclinical studies in rats, pregabalin was asso male-mediated teatogenicity (see WARNINGS AND PRECAUTIONS, Sexual Function/Reproduction). The clinical significance of this finding is uncertain; however, men being treated with LYRICA who plan to father a child should be informed of the potential risk of male-mediated teratogenicity.

Diabetic patients should be instructed to pay particular attention to skin integrity while being treated with LYRICA. Some animals treated with pregabalin developed skin ulcerations, although no increased incidence of skin lesions associated with LYRICA was observed in clinical trials (see **PRODUCT MONOGRAPH**).

Patients should be informed of the availability of a patient information leaflet, and they should be instructed to read the leaflet prior to taking LYRICA.

Preclinical Toxicology Carcinogenesis

A dose-dependent increase in the incidence of malignant vascular tumors (hemangiosarcomas) was observed in two strains of mice (B6C3F1 and CD-1) given Internatinguisatromasy was ususerved in two starties of the mice bocks? The arms to Co-1 given pregabalin (200, 1000 or 5000 mg/kg) in the diet for two years. Plasma pregabalin exposure (AUC1) in mice receiving the lowest dose that increased hemangiosarcomas was approximately equal to the human exposure at the maximum recommended dose (MRD) of 600 mg/day. A no-effect dose for induction of hemangiosarcomas in mice was not established. In an investigative study in female B6C3F1 mice, chronic treatment (24 months) with prepabalin at 1000 mg/kg caused an increased incidence of hemangiosarcoma, consistent with previous studies, but not at 50 or 200 mg/kg. on retainguistancome, consistent with previous studies, but hold at 90 at 200 mg/kg. Discontinuation of treatment after 12 months at 1000 mg/kg did not significantly reduce the incidence of hemangiosarcoma at 24 months. Evidence of carcinogenicity was not seen in two studies in Wistar rats following dietary administration of pregabalin for two years at doses (50, 150 or 450 mg/kg in males and 100, 300 or 900 mg/kg in females) that were associated with plasma exposures in males and females up to approximately 14 and 24 times, respectively, human exposure at the MQD. The disease isonificance is humans of this folionic princip undersum. MRD. The clinical significance in humans of this finding in mice is unknown

Mutagenesis

Pregabalin is not genotoxic based on results of a battery of in vitro and in vivo tests. Pregabalin was not mutagenic in bacteria or in mammalian cells in vitro, was not clastogenic in mammalian systems in vitro and in vivo, and did not induce nscheduled DNA synthesis in mouse or rat hepatocytes.

Dermatopathy

Skin lesions ranging from erythema to necrosis were seen in repeated-dose toxicology studies in both rats and monkeys. The etiology of these skin lesions is unknown. At the maximum recommended human dose (MRD) of 600 mg/day, there is a 2-fold safety margin for the dermatological lesions. The more severe dermatopathies involving necrosis were associated with pregabalin exposures (as expressed by plasma AUCs) of approximately 3 to 8 times those achieved in humans given the MRD. No increase in incidence of skin lesions was observed in clinical studies.

Ocular lesions

Ocular lesions (characterized by retinal atrophy [including loss of photoreceptor cells] and/or corneal inflammation/mineralization) were observed in two lifetime carcinogenicity studies in Wistar rats. These findings were observed at plasma pregabalin exposures (AUC) ≥2 times those achieved in humans given the maximum recommended dose of 600 mg/day. A no-effect dose for ocular lesions was not established. Similar lesions were not observed in lifetime carcinogenicity studies in two strains of mice or in monkeys treated for 1 year. The clinical significance of

Monitoring and Laboratory Tests

Routine therapeutic drug monitoring or clinical laboratory testing is not required for patients treated with LYRICA (pregabalin) (see **ADVERSE REACTIONS**).

ADVERSE REACTIONS

Adverse Drug Reaction Overview Clinical Trial Adverse Drug Reactions

Clinical Intel Audies Dring Reactions in all controlled and uncontrolled trials, more than 8666 patients have received LYRICA (pregabalin), with 83% of exposure at dosages of 300 mg/day or above and 32% at dosages of 600 mg/day or higher. Approximately 4010 patients had at least 6 months of exposure, ad 15 had at least 1 year of exposure, and 39\$ had at least 2 years of exposure to pregabalin. In controlled trials, 1831 patients with neuropathic

ain received pregabaling

Most Common Adverse Events in All Controlled Clinical Studies of Neuropathic Pain

The most commonly observed adverse events (≥5% and twice the rate of that seen in placebo) in pregabalin-treated patients were: dizziness, somnolence, peripheral edema and dry mouth. Adverse events were usually mild to moderate in intensity. Discontinuation Due to Adverse Events

In all controlled studies, the discontinuation rate due to adverse events was 14% for patients receiving pregabalin and 7% for patients receiving placebo. The most common reasons for discontinuation due to adverse events (\$2%) in the pregabalin treatment groups were dizziness and somolence. Other adverse events that led to withdrawal more frequently in the pregabalin group than the placebo group were ataxia [1%] and asthenia, confusion, headache and nausea (<1% each).

In controlled neuropathic pain studies, the discontinuation rate due to adverse events was 11% for pregabalin and 5% for placebo. The most common reasons for discontinuation due to adverse events (≥2%) in the pregabalin treatment groups were dizziness and somnolence. Other adverse events that led to withdrawal more frequently in the pregabalin group than the placebo group were confusion (1%) and asthenia, peripheral edema and ataxia (<1% each).

Incidence of Adverse Events in Controlled Clinical Studies of Neuropathic Pain Insummanes of adverse events, investigator's terms for individual adverse events have been grouped into a smaller number of standardized categories using the COSTART IV dictionary. The prescriber should be aware that the percentages in Table 1 through Table 6 cannot be used to predict the frequency of adverse events in the course of usual medical practice where patient characteristics and other factors may differ from those prevailing during clinical studies. Similarly, the cited frequencies cannot be directly compared with figures obtained from other clinical investigations involving different treatments, uses or investigators. An inspection of these frequencies, however, does provide the prescriber with one basis to estimate the relative contribution of drug and non-drug factors to the adverse event incidences in the population studied.

Adverse Events From Controlled Clinical Studies of Neuropathic Pain

Diabetic Peripheral Neuropathy

Table 1 lists all adverse events, regardless of causality, occurring in ≥2% of patients lable I lists all adverse events, legardies or causainty, occurring in ≥₹% or patients with heuropathic pain associated with diabetic peripheral neuropathy receiving pregabalin for at least one of the pregabalin groups, and for which the incidence was greater than in the placebo group. A majority of pregabalin-treated patients in these studies had adverse events with a maximum intensity of mild or moderate. In these studies, 979 patients received pregabalin and 459 patients received placebo for up to 13 weeks

Table 1. Incidence (%) of Treatment-Emergent Adverse Events in Placebo-Controlled Studies in Neuropathic Pain Associated with Diabetic Peripheral Neuropathy (Events in at Least 2% of Patients Receiving Pregabalin and More Frequent Than in Placebo-Treated Patients)

		Pregabalin (mg/day)			
Body System Preferred Term	Placebo (n = 459) %	75 (n = 77) %	150 (n = 212) %	300 (n = 321) %	600 (n = 369) %
Body as a whole					
Infection	6.1	3.9	7.5	8.4	4.6
Asthenia	2.4	3.9	1.9	4.4	7.3
Pain	3.9	5.2	4.2	2.5	4.9
Accidental injury	2.8	5.2	2.4	2.2	5.7
Back pain	0.4	0.0	2.4	1.2	1.9
Chest pain	1.1	3.9	1.4	1.2	1.6
Face edema	0.4	0.0	0.9	0.9	2.2
Digestive system		- Carlotte Company			
Dry mouth	1.1	2.6	1.9	4.7	6.5
Constipation	1.5	0.0	2.4	3.7	6.0
Diarrhea	4.8	5.2	2.8	1.9	3.0
Flatulence	1.3	2.6	0.0	2.2	2.7
Vomiting	1.5	1.3	0.9	2.2	1.1
Hemic and lymph	natic system	1			
Ecchymosis	0.2	2.6	0.5	0.6	0.3
Metabolic and no	utritional di	sorders			
Peripheral edema	2.4	3.9	6.1	9.3	12.5
Weight gain	0.4	0.0	4.2	3.7	6.2
Edema	0.0	0.0	1.9	4.0	1.9
Hypoglycemia	1.1	1.3	3.3	1.6	1.1
Nervous system					
Dizziness	4.6	7.8	9.0	23.1	29.0
Somnolence	2.6	3.9	6.1	13.1	16.3
Neuropathy	3.5	9.1	1.9	2.2	5.4
Ataxia	1.3	6.5	0.9	2.2	4.3
Vertigo	1.1	1.3	1.9	2.5	3.5
Confusion	0.7	0.0	1.4	2.2	3.3
Euphoria	0.0	0.0	0.5	3.4	1.6
Thinking abnormal ^a	0.0	1.3	0.0	0.9	3.0

		Pregabalin (mg/day)			
Body System Preferred Term	Placebo (n = 459) %	75 (n = 77) %	150 (n = 212) %	300 (n = 321) %	600 (n = 369) %
Abnormal gait	0.0	1.3	0.0	0.6	2.7
Reflexes decreased	1.7	3.9	0.5	1.2	1.4
Amnesia	0.2	2.6	0.9	0.0	2.2
Hypesthesia	0.7	2.6	0.0	0.0	0.8
Hyperalgesia	0.2	2.6	0.0	0.0	0.3
Respiratory syst	em				
Dyspnea	0.7	2.6	0.0	1.9	1.9
Skin and append	dages				
Pruritus	1.3	2.6	0.0	0.9	0.0
Special senses					
Blurred vision ^b	1.5	2.6	1.4	2.8	1.5
Conjunctivitis	0.2	2.6	1.4	0.6	0.3

- a Thinking abnormal primarily consists of events related to difficulty with concentration/attention but also includes events related to cognition and language problems and slow thinking.
- b Investigator term: summary level term is amblyopia

Discontinuation in Controlled Clinical Studies of Diabetic Peripheral Neuropathy

Approximately 9% of patients receiving pregabalin and 4% receiving placebo discontinued from controlled diabetic peripheral neuropathy studies due to adverse events. The adverse events most commonly leading to discontinuation are presented

Table 2. Adverse Events Most Frequently (≥2% of patients) Leading to Discontinuation in Placebo-Controlled Studies in Patients with ociated with Diabetic Perinhe

	Nu	ımber (%) o	f Patients			
		Pregabalin (mg/day)				
COSTART Preferred Term	Placebo (n = 459)	75 (n = 77)	150 (n = 212)	300 (n = 321)	600 (n = 369)	
Dizziness	2 (0.4)	0 (0.0)	3 (1.4)	6 (1.9)	21 (5.7)	
Somnolence	0 (0.0)	0 (0.0)	0 (0.0)	5 (1.6)	15 (4.1)	

Postherpetic Neuralgia

Table 3 lists all adverse events, regardless of causality, occurring in ≥2% of patients with neuropathic pain associated with postherpetic neuralgia receiving pregabilin for at least one of the pregabalin groups, and for which the incidence was greater than in the placebo group. A majority of pregabalin-treated patients in these studies had adverse events with a maximum intensity of mild or moderate. In these studies, 852 patients received pregabalin and 398 patients received placebo for up to 13 weeks.

Table 3. Incidence (%) of Treatment-Emergent Adverse Events in Placebo-Controlled Studies in Neuropathic Pain Associated with Postherpetic Neuralgia (Events in at Least 2% of Patients Receiving Pregabalin and More Frequent Than in Placebo-Treated Patients)

		Pregabalin (mg/day)						
Body System Preferred Term	Placebo (n = 398)	75 (n = 84) %	150 (n = 302) %	300 (n = 312) %	600 (n = 154) %			
Body as a who	le							
Infection	3.5	14.3	8.3	6.4	2.6			
Headache	5.3	4.8	8.9	4.5	8.4			
Pain	3.8	4.8	4.3	5.4	4.5			
Asthenia	4.0	3.6	5.0	2.6	5.2			
Accidental injury	1.5	3.6	2.6	3.2	5.2			
Flu syndrome	1.3	1.2	1.7	2.2	1.3			
Face edema	0.8	0.0	1.7	1.3	3.2			
Malaise	1.0	2.4	0.3	0.6	0.0			
Cardiovascula	r system							
Vasodilatation	1.3	2.4	1.0	0.6	0.0			
Digestive syste	em							
Dry mouth	2.8	7.1	7.0	6.1	14.9			
Constipation	2.3	3.6	4.6	5.4	5.2			
Diarrhea	4.0	2.4	4.3	3.5	4.5			
Flatulence	1.0	2.4	1.3	1.6	3.2			
Vomiting	0.8	1.2	0.7	2.9	2.6			
Metabolic and	nutritional o	disorders						
Peripheral	3.5	0.0	7.9	15.7	16.2			
edema		0.500						
Weight gain	0.3	1.2	1.7	5.4	6.5			
Edema	1.3	0.0	1.0	2.2	5.8			
Hyperglycemia	0.8	2.4	0.3	0.0	0.0			
Nervous system								
Dizziness	9.3	10.7	17.9	31.4	37.0			
Somnolence	5.3	8.3	12.3	17.9	24.7			
Ataxia	0.5	1.2	2.0	5.4	9.1			
Abnormal gait	0.5	0.0	2.0	3.8	7.8			
Confusion	0.3	1.2	2.3	2.9	6.5			
Thinking abnormal ^a	1.5	0.0	1.7	1.3	5.8			
Incoordination	0.0	2.4	1.7	1.3	2.6			
Amnesia	0.0	0.0	1.0	1.3	3.9			
Speech disorder	0.0	0.0	0.3	1.3	3.2			
Insomnia	1.8	0.0	0.7	2.2	0.0			
Euphoria	0.0	2.4	0.0	1.3	1.3			
Nervousness	0.5	0.0	1.0	0.3	2.6			
Tremor	1.5	1.2	0.0	1.0	2.6			
Hallucinations	0.0	0.0	0.3	0.3	3.2			
Hyperesthesia	0.3	2.4	0.3	0.0	1.3			
Respiratory sy								
Bronchitis	0.8	0.0	1.3	1.0	2.6			
Pharyngitis	0.8	0.0	2.6	0.6	0.6			

		Pregabalin (mg/day)						
Body System Preferred Term	Placebo (n = 398) %	75 (n = 84) %	150 (n = 302) %	300 (n = 312) %	600 (n = 154) %			
Rhinitis	1.8	1.2	0.7	0.6	3.2			
Skin and apper	ndages	•	•					
Rash	3.0	2.4	2.0	2.9	5.2			
Special senses	3							
Blurred vision ^b	2.5	1.2	5.0	5.1	9.1			
Diplopia	0.0	0.0	1.7	1.9	3.9			
Abnormal vision	0.3	0.0	1.0	1.6	5.2			
Urogenital sys	tem							
Urinary tract infection	1.5	0.0	2.3	1.6	3.2			

a. Thinking abnormal primarily consists of events related to difficulty with a Trilling automina primary consists of events related to unificulty with concentration/attention but also includes events related to cognition and language problems and slow thinking.

b Investigator term; summary level term is amblyopia.

Discontinuation in Controlled Clinical Studies of Posthernetic Neuralgia Approximately 14% of patients receiving pregabalin and 7% receiving placebo discontinued from controlled postherpetic neuralgia studies due to adverse eve adverse events most commonly leading to discontinuation are presented in Tab

Table 4. Adverse Events Most Frequently (>2% of patients) Leading to Discontinuation in Placebo-Controlled Studies in Patients with Neuropathic Pain Associated with Postherpetic Neuralgia

	1	Number (%)	of Patients						
COSTART		Pregabalin (mg/day)							
Preferred Term	Placebo (n = 398)	75 (n = 84)	150 (n = 302)	300 (n = 312)	600 (n = 154)				
Dizziness	3 (0.8)	0 (0.0)	11 (3.6)	12 (3.8)	12 (7.8)				
Somnolence	1 (0.3)	0 (0.0)	6 (2.0)	12 (3.8)	10 (6.5)				
Confusion	1 (0.3)	0 (0.0)	2 (0.7)	5 (1.6)	8 (5.2)				
Peripheral edema	1 (0.3)	0 (0.0)	2 (0.7)	5 (1.6)	5 (3.2)				
Ataxia	0 (0.0)	0 (0.0)	1 (0.3)	5 (1.6)	4 (2.6)				
Abnormal gait	0 (0.0)	0 (0.0)	0 (0.0)	4 (1.3)	4 (2.6)				
Hallucinations	0 (0.0)	0 (0.0)	0 (0.0)	1 (0.3)	4 (2.6)				
Dry mouth	1 (0.3)	0 (0.0)	0 (0.0)	0 (0.0)	4 (2.6)				

Incidence of Most Common Dose-Related Treatment-Emergent Adverse

Most common dose-related treatment-emergent adverse events are presented in Table 5 (diabetic peripheral neuropathy) and Table 6 (postherpetic neuralgia).

Table 5. Incidence (%) of Most Common Dose-Related Treatment-Emergent Adverse Events in Placebo-Controlled Studies in Neuropathic Pain Associated with Diabetic Peripheral Neuropathy

Adverse Event Preferred Term		Pregabalin (mg/day)						
	Placebo (n = 459) %	75 (n = 77) %	150 (n = 212) %	300 (n = 321) %	600 (n = 369) %			
Dizziness	4.6	7.8	9.0	23.1	29.0			
Somnolence	2.6	3.9	6.1	13.1	16.3			
Peripheral edema	2.4	3.9	6.1	9.3	12.5			
Asthenia	2.4	3.9	1.9	4.4	7.3			
Dry mouth	1.1	2.6	1.9	4.7	6.5			
Weight gain	0.4	0.0	4.2	3.7	6.2			
Constipation	1.5	0.0	2.4	3.7	6.0			
Blurred vision ^a	1.5	2.6	1.4	2.8	5.7			

a Investigator term; summary level term is amblyopia.

Table 6. Incidence (%) of Most Common Dose-Related Treatment-Emergent Adverse Events in Placebo-Controlled Studies in Neuropathic Pain Associated with Posthernetic Neuralgia

Adverse		Pregabalin (mg/day)						
Event Preferred Term	Placebo (n = 398) %	75 (n = 84) %	150 (n = 302) %	300 (n = 312) %	600 (n = 154) %			
Dizziness	9.3	10.7	17.9	31.4	37.0			
Somnolence	5.3	8.3	12.3	17.9	24.7			
Peripheral edema	3.5	0.0	7.9	15.7	16.2			
Dry mouth	2.8	7.1	7.0	6.1	14.9			
Blurred vision ^a	2.5	1.2	5.0	5.1	9.1			
Ataxia	0.5	1.2	2.0	5.4	9.1			
Weight gain	0.3	1.2	1.7	5.4	6.5			
Abnormal gait	0.5	0.0	2.0	3.8	7.8			

a Investigator term; summary level term is amblyopia.

Adverse Events Following Abrupt or Rapid Discontinuation

Following abrupt or rapid discontinuation of pregabalin, some patients reported symptoms including insomnia, nausea, headache and diarrhea. Pregabalin should he tapered gradually over a mir mum of one week rather than discontinued abruptly (see WARNINGS AND PRECAUTIONS, Abrupt or Rapid Discontinuation)

Drug Abuse and Dependence/Liability

In a study of recreational users (n=15) of sedative/hypnotic drugs, including alcohol, a single dose of LYRICA (pregabalin) 450 mg received subjective ratings of "good drug effect", "high", and "liking" to a degree that was similar to a single dose of diazepam 30 mg, in controlled clinical studies in over 5500 patients, 4% of LYRICAtreated patients and 1% of placebo-treated patients overall reported euphoria as an adverse event. However, in clinical trials of diabetic peripheral neuropathy, euphoria was reported as an adverse event by 1.8% of LYRICA-treated patients and 0% of placebo-treated patients, and in clinical trials of postherpetic neuralgia, euphoria was reported as an adverse event by 0.9% of LYRICA-treated patients and 0% of placebo-treated patients. In clinical studies, following abrupt or rapid discontinuation of pregabalin, some patients reported symptoms including insomnia, nausea, headache or diarrhea suggestive of physical dependence (see WARNINGS AND PRECAUTIONS, Abrupt or Rapid Discontinuation).

Pregabalin is not known to be active at receptor sites associated with drugs of abuse. As with any CNS active drug, physicians should carefully evaluate patients for history of drug abuse and observe them for signs of LYRICA misuse or abuse (e.g., development of tolerance, dose escalation, drug-seeking behaviour).

Other Events Observed During the Premarketing Evaluation of LYRICA

Following is a list of treatment-emergent adverse events reported during Following is a list or treatment-emergent adverse events reported ourning premarketing assessment of LYRICA in clinical trials (over 8600 adult subjects) except those already listed in the previous tables or elsewhere in labeling. In the tabulations that follow, a COSTART-based dictionary of terminology has been used tabulations that follow, a CUSTAH-loased dictionary of terminology has been used to classify reported adverse events. The frequencies presented, therefore, represent the proportion of the over 8600 adult individuals exposed to multiple doses of LYRICA who experienced an event of the type cited on at least 1 occasion while receiving LYRICA. It is important to emphasize that although the events reported occurred during treatment with LYRICA, they were not necessarily caused by it.

Less Common Clinical Trial Adverse Drug Reactions (<2%)

Events are further categorized by body system and listed in order of decreasing frequency according to the following definitions: frequent adverse events are those occurring on 1 or more occasions in at least 1/100 patients; infrequent adverse events are those occurring in 1/100 to 1/1000 patients; rare events are those occurring in fewer than 1/1000 patients.

Body System	Adverse Events
Body as a v	whole
Frequent	Flu syndrome, back pain, allergic reaction, fever, generalized edema
Infrequent	Neck pain, neoplasm, cellulitis, cyst, chills, malaise, overdose, moniliasis, hernia, viral infection, photosensitivity reaction, pelvic pain, abdomen enlarged, abscess, neck rigidity, lab test abnormal, drug level increased, carcinoma, sepsis, suicide attempt, reaction unevaluable
Rare	Infection fungal, unexpected benefit, chills and fever, body odor, drug level decreased, halitosis, hangover effect, injection site reaction, hormone level altered, hypothermia, infection bacterial, injection site hemorrhage, intentional overdose, mucous membrane disorder, accidental overdose, adenoma, anaphylactoid reaction, ascites, chest pain substernal, death, sarcoidosis, sudden death, immune system disorder, increased drug effect, injection site pain, Lupus Erythematosus syndrome, medication error, sarcoma, shock, tolerance decreased
Cardiovaso	cular
Frequent	Hypertension, vasodilatation
Infrequent	Palpitation, migraine, tachycardia, peripheral vascular disorder, electrocardiogram abnormal, cardiovascular disorder, angina pectoris, congestive heart failure, hemorrhage, myocardial infarct, hypotension, postural hypotension, ventricular extrasystoles, atrial fibrillation, coronary atrey disorder, bradycardia, cerebovascular disorder, sinus bradycardia, myocardial ischemia, bundle branch block, AV block first degree, arteriosclerosis, deep thrombophlebtis, phiebitis, arterial anomaly, heart failure, pulmonary embolus, retinal vascular disorder, soncose vein
Rare	Heart arrest, vascular anomaly, occlusion, supraventricular tachycardia, atrial arrhythmia, atrial flutter, cerebral infarct, coronary occlusion, thrombophlebitis, thrombosis, cardiomegaly, extrasystoles, pallor, AV block, AV block second degree, cardiomyopathy, peripheral gangrene, CII interval prolonged, retinal artery occlusion, supraventricular extrasystoles, cerebral hemorrhage, digitalis intoxication, ventricular arrhythmia, aortic stenosis, bigeminy, cerebrovascular disorder, left heart failure, ventricular tarchyacrdia, AV block complete, carotid occlusion, carotid thrombosis, cor pulmonale, embolus lower extremity, endocarditis, heart block, increased capillary fragility, intracranial aneurysm, nodal tarchycardia, CII interval shortened, retinal vein thrombosis, ST elevated, T inverted, vascular headache, vasculitis
Digestive s	system
Frequent	Nausea, diarrhea, anorexia, gastrointestinal disorder
Infrequent	Gastroenteritis, tooth disorder, periodontal abscess, colitis, gastriis, liver function tests abnormal, increased salivation, thirst, nausea and vomiting, rectal disorder, gingivitis, dysphagia, stomatitis, mouth ulceration, cholelithiasis, rectal hemorrhage, gastrointestinal hemorrhage, glossitis, tooth caries, abnormal stools, cholecystitis, melena, oral moniliasis, esophagitis, tongue disorder, chellitis, tongue edema
Rare	Eructation, pancreatitis, stomach ulcer, ulcerative stomatitis, esophageal stenosis, fecal incontinence, gum hemorrhage, intestinal obstruction, enteritis, peptic ulcer, enterocolitis, gum hyperplasia, hepatomegaly, liver fatty deposit, tenesmus, biliary pain, fecal impaction, jaundice, periodontitis, ulcerative colitis, aphthous stomatitis, cholestatic jaundice, gastrointestinal carcinoma, hemorrhagic gastritis, hepatitis, liver tenderness, anusea, voniting and diarrhes, salivary gland enlargement, stomach atony, bloody diarrhea, cardiospasm, duodenal ulcer, gamma glutamyl transpeptidase increased, hematemesis, hepatoma, intestinal stenosis, intestinal ulcer, leukoplakia of mouth, necrotizing pancreatitis, pancreas disorder, pseudomembranous colitis, sialadenitis, stomach ulcer hemorrhage, tongue discoloration
Endocrine	system
Infrequent	Diabetes mellitus, hypothyroidism
Rare	Goiter, prolactin increased, thyroid disorder, gonadotropic follicle stim hormone increase, hyperthyroidism, thyroiditis, adrenal insufficiency, parathyroid disorder, thyroid carcinoma, thyroid neoplasia, virilism
Hemic and	lymphatic
Infrequent	Anemia, leukopenia, thrombocytopenia, lymphadenopathy, hypochromic anemia, leukocytosis, eosinophilia
Rare	hypochromia alemae, leukocyosis, eosinipinia Lymphocytosis, petechia, iron deficiency anemia, cyanosis, lymphodema, polycythemia, lymphoma like reaction, megalobiastic anemia, splenomegaly, purpura, thrombocythemia, thrombocytopenic purpura, chronic leukemia, coagulation disorder, erythrocytes abnormal, leukemoid reaction, lymphangitis, macrocytic anemia, pancytopenia, prothrombin decreased, rupture of spleen, sedimentation rate increased

Body System	Adverse Events
	and nutritional
nfrequent	Hyperglycemia, SGPT increased, hypoglycemia, hypokalemia, hypercholesteremia, SGOT increased, weight loss, hyperlipemia, amylase increased, hyperuricemia, alkaline phosphatase increased, creatinine increased, hyponatremia, gout, dehydration, BUN increased, healing abnormal
Rare	Hypercalcemia, hyperkalemia, hypocalcemia, bilirubinemia, alcohol intolerance, hypoglycemic reaction, ketosis, calcium disorder, hypochloremia, hypomagnesemia, hypoproteinemia, NPN increased, uremia, acidosis, avitaminosis, enzymatic abnormality, gamma globulins increased, hypernatremia, hypophosphatemia, lactic acidosis, obesity
Musculosk	celetal system
Frequent	Arthralgia, myalgia, arthritis, leg cramps, myasthenia
Infrequent	Tendon disorder, arthrosis, joint disorder, bone disorder, tenosynovitis, bursitis, tendinous contracture, osteoporosis, tendon rupture, bone pain
Rare	Rheumatoid arthritis, osteomyelitis, rhabdomyolysis, myopathy, muscle atrophy, myositis, pyogenic arthritis, bone neoplasm, musculoskeletal congenital anomaly, pathological fracture
Nervous sy	
Frequent	Insomnia, anxiety, libido decreased, depersonalization, hypertonia, neuropathy
Infrequent	Reflexes decreased, sleep disorder, abnormal dreams, hostility, hallucinations, hyperkinesia, personality disorder, dysarthria, hyperesthesia, hypokinesia, circumoral parethesia, libido increased, neuralgia, vestibular disorder, aphasia, movement disorder, hyperalgesia, apathy, hypotonia, convulsion, facial paralysis, psychosis
Rare	Drug dependence, neuritis, paranoid reaction, CNS depression, CNS neoplasia, manic reaction, neurosis, extrapyramidal syndrome, meningitis, hemiplegia, reflexes increased, akathisia, delirium, paralysis, withdrawal syndrome, brain edema, CNS stimulation, dyskinesia, encephalopathy, foot drop, grand mal convulsion, hypalgesia, enripheral neuritis, psychotic depression, addiction, arachnoiditis, cerebellar syndrome, cogwheel rigidity, dementia, dystonia, Guillain-Barre syndrome, intractanial hemorrhage, multiple sclerosis, myelitis, schizophrenic reaction, subarachnoid hemorrhage, torticollis
Respirator	y system
Frequent	Sinusitis, rhinitis, dyspnea, cough increased, pneumonia, lung disorder
Infrequent	Asthma, epistaxis, laryngitis, voice alteration, respiratory disorder, sputum increased
Rare	Apnea, emphysema, aspiration pneumonia, hyperventilation, lung edema, pleural disorder, atelectasis, hemoptysis, hiccup, hypoxal, laryngismus, lung fibrosis, pleural effusion, lung function decreased, pulmonary hypertension, yawn, bronchiedtasis, bronchiolitis, carcinoma of lung, hypoventilation, laryngeal neoplasia, nasal septum disorder, pneumothorax
Skin and a	ppendages
Infrequent	Pruritus, sweating, skin disorder, acne, dry skin, alopecia, skin ulcep, herpes simplex, urticaria, nail disorder, eczema, herpes zoster, skin benign neoplasm, fungal dermatitis, maculopapular rash, vesiculobullous rash, skin cartinoma, furunculosis, skin discoloration, skin hypertrophy, psoriasis, seborrhea, hirsutism
Rare	Skin nodule, angioedema, cutaneous moniliasis, skin atrophy, exfoliative dermatitis, pustular rash, ichthyosis, skin melanoma, subcutaneous nodule, sweating decreased, hair disorder, lichenoid dermatitis, melanosis, miliaria, purpuric rash, skin necrosis, Stevens Johnson syndrome
Special se	ense
Frequent	Eye disorder, conjunctivitis, otitis media
Infrequent	Retinal disorder, tinnitus, eye pain, cataract specified, dry eyes, taste perversion, ear pain, lacrimation disorder, ear disorder, deafness, eye hemorrhage, photophobia, glaucoma, vitreous disorder, corneal lesion, ottis externa, refraction disorder, blepharitis, retinal edema, taste loss, abnormality of
Rare	accommodation Hyperacusis, keratitis, mydriasis, parosmia, ptosis, retinal hemorrhage, color blindness, retinal depignment, retinal detachment, retinal description, retinal description, retinal description, retinal
	detachment, corneal opacity, corneal ulcer, iritis, niight blindness, optic atrophy, retinal degeneration, cataract NOS, scleritis, strabismus, anisocoria, blindness, exophthalmos, keratoconjunctivitis, ophthalmoplegia, papilledema
Urogenital	
Frequent	Anorgasmia
Infrequent	Urinary frequency, urinary incontinence, cystitis, abnormal ejaculation, urination impaired, dysuria, metrorrhagia, hematuria, vaginal moniliasis, prostatic disorder, vaginitis, dysmenorrhea, urinary urgency, kidney calculus, breast pain, menstrual disorder, amenorrhea, menorrhagia, kidney function abnormal, nephritis, urine abnormality, vaginal hemorrhage, urinary retention, urinary tract disorder, leukorrhea, preast neoplasm, menopause, oliguria, polyuria, albuminuria, pyuria
Rare	Breast carcinoma, penis disorder, papanicolau smear suspicious, fibrooystic breast, prostatic carcinoma, uterine fibroide slarlaged, acute kidney failure, creatinine clearance decreased, nephrosis, nocturia, polycystic kidney, bladder carcinoma, breast enlargement, cervicitis, cervic disorder, female leatation, glyosuria, gynecomastia, hypomenorrhea, kidney pain, mastitis, pyelonephritis, kidney failure, breast abscess, epididymitis, orchitis, prostate neoplasia, prostatic specific antigen increase, salpingitis, urogenital disorder, unlithiasis, utenne disorder, vulvovaginal disorder, balanitis, bladder calculus, calcium crystalluria, cervix neoplasm, dysparemia, endometrial

Comparison of Gender and Race

The overall adverse event profile of pregabalin was similar between women and men. There are insufficient data to support a statement regarding the distribution of adverse experience reports by race.

Peripheral Edema

Incidence of peripheral edema in controlled neuropathic pain studies was 10.4% in the pregabalin group compared with 2.9% in the placebo group. In clinical trials, these events of peripheral edema were dose-related, mostly mild to moderate in intensity and rarely led to withdrawal. Perinheral edema was not associated with intensity and lately led to withdrawar. Peripheral events was to associated with cardiovascular complications such as hypertension or congestive heart failure and there was no evidence of hemodilution or changes in any laboratory parameters indicative of underlying organ dysfunction (see **WARNINGS AND PRECAUTIONS**, Peripheral Edema

In the controlled neuropathic pain studies, patients on pregabalin had a higher incidence (5.9%) of weight gain as defined by a ≥7% increase from baseline weight as compared with the placebo group (1.6%). The mean change in the pregabalin group was an increase of 1.5 kg compared with 0.2 kg in the placebo group; few patients (0.1%) withdrew due to weight gain. This weight gain was dose-related, and not associated with clinically important changes in blood pressure or cardiovascular adverse events. There was no relationship between baseline body mass index and the incidence of $\geq 7\%$ weight gain in the controlled trials.

Based on the results of a controlled study of reproductive function in healthy male volunteers, the \$27% weight gain on pregabalin appeared to be reversible. In this study, there were no reports of peripheral edema (see **WARNINGS AND** PRECAUTIONS, Weight Gain

Abnormal Hematologic and Clinical Chemistry Findings

In all controlled trials, 1,0% of patients on pregabalin and 0,5% of placebo patients had an increase in creatine kinase of -3x upper limit of normal. Renal dysfunction was generally not associated with the elevated creatine kinase in these patients. Mean changes in creatine kinase in pregabatin-treated patients and 4.8 U/L for the placebo patients (see **DOSAGE AND ADMINISTRATION**, Patients with Renal Impairment). Routine therapeutic
drug monitoring or clinical laboratory testing is not required for patients treated with
LYRICA (see **WARNINGS AND PRECAUTIONS**).

Post-Marketing Adverse Drug Reactions

POST-marketing Aurores or Jug neacuous

The worldwide post-marketing experience to date with LYRICA is consistent with the clinical program. The most frequently reported adverse events from spontaneous post-marketing reports for LYRICA are shown below. There are insufficient data to support an estimate of their incidence or to establish causation.

See disorders: diplopia, vision blurred, visual disturbance. There have also been rare reports of accommodation disorder, eyelid edema and eye redness (see WARNINGS AND PRECAUTIONS, Ophthalmological Effects).

Gastrointestinal disorders: diarrhea, dry mouth, nausea, vomiting

General disorders and administration site conditions: fatigue, feeling

Nervous system disorders: ataxia, coordination abnormal, dizziness, dysarthria, headache, memory impairment, paresthesia, somnolence, speech disorder, tremor (see WARNINGS AND PRECAUTIONS, <u>Dizziness and Somnolence</u>).

Psychiatric disorders: confusional state, depression, insomnia, psychotic disorder There have been rare reports of psychotic disorders in patients receiving pregabalin.

Renal and urinary disorders: urinary retention Respiratory, thoracic and mediastinal disorders: dyspnea

Skin and subcutaneous tissue disorders: pruritus

DRUG INTERACTIONS

Since pregabalin is predominately excreted unchanged in the urine, undergoes negligible metabolism in humans (-2% of a dose recovered in urine as metabolites), does not inhibit drug metabolism in vitro, and is not bound to plasma proteins, LYRICA (pregabalin) is unlikely to produce, or be subject to, pharmacokinetic interactions

In Vitro Studies: In vitro drug metabolism studies revealed that pregabalin at concentrations which were, in general, 10-fold greater than observed in Phase 2/3 clinical trials, does not inhibit human CYP1A2, CYP2A6, CYP2C9, CYP2C19, CYP2D6, CYP2E1, and CYP3A4 enzyme systems.

In Vivo Studies: The drug interaction data described in this section were obtained from studies involving healthy adults, patients with epilepsy, and patients with chronic pain disorde

Carbamazepine, valproic acid, lamotrigine, phenytoin, phenobarbital, and

In vitro and in vivo studies showed that LYRICA is unlikely to be involved in significant pharmacokinetic drug interactions. Specifically, there are no clinically significant pharmacokinetic interactions between pregabalin and the following antiepileptic pharmacoxinetic interactions between programma and the fortunary anti-propriate drugs: carbamazepine, valiptoic acid, lamotrigine, phenytoin, phenobarbital, and topiramate. Important pharmacokinetic interactions would also not be expected to occur between pregabalin and commonly used antiepileptic drugs.

Tiagabine: The results of a population pharmacokinetic analysis indicated that in patients with partial seizures tiagabine had no clinically significant effect on egabalin clearance

pregabalin clearance.

Gabapentin: The pharmacokinetics of pregabalin and gabapentin were investigated in 12 healthy subjects following concomitant single dose administration of 100 mg pregabalin and 300 mg gabapentin, and in 18 healthy subjects following concomitant multiple dose administration of 200 mg pregabalin ghan and 200 mg gabapentin open down of the gabapentin pharmacokinetics following single and multiple dose administration were unaltered by pregabalin cadeministration. The rate of pregabalin absorption was reduced by approximately 25% (single dose administration) and 18% (multiple dose administration) based on lower C_{ess} values, however, the extent of pregabalin absorption was reduced by approximately approximately 25% (single dose administration) and 18% (multiple dose administration) based on lower C_{ess} values, however, the extent of pregabalin absorption was reaffected by acknowledged to the control of absorption was unaffected by gabapentin coadministration

Oral Contraceptives: Pregabalin coadministration (200 mg TID) had no effect on the steady state pharmacokinetics of norethindrone and ethinyl estradiol (1 mg/35 µg, espectively) in healthy subjects.

Lorazepam: Multiple dose administration of pregabalin (300 mg BID) in healthy subjects had no effect on the rate and extent of lorazepam single dose pharmacokinetics and single dose administration of lorazepam (1 mg) had no clinically significant effect on the steady state pharmacokinetics of pregabalin.

Oxycodone: Multiple dose administration of pregabalin (300 mg BID) in

healthy subjects had no effect on the rate and extent of oxycodone single dose pharmacokinetics. Single dose administration of oxycodone (10 mg) had no clinically significant effect on the steady state pharmacokinetics of pregabalin.

Ethanol: Multiple dose administration of pregabalin (300 mg BID) in healthy subjects had no effect on the rate and extent of ethanol single dose pharmacokinetics and single dose administration of ethanol (0.7 g/kg) had no clinically significant effect on the steady state pharmacokinetics of pregabalin.

Diuretics, Oral Hypoglycemics, and Insulin: A population pharmacokinetic analysis in patients with chronic pain showed no clinically significant effect on pregabalin clearance with the concomitant use of diuretics, oral hypoglycemics,

carcinoma, endometrial disorder, glomerulitis, hydronephrosis, ovarian cancer, unintended pregnancy, urethral pain, urethritis, urogenital anomaly, urogenital neoplasia, uterine hemorrhage

Pharmacodynamic

Multiple oral doses of pregabalin co-administered with oxycodone, lorazepam, or ethanol did not result in clinically important effects on respiration. Pregabalin appears to be additive in the impairment of cognitive and gross motor function caused by oxycodone. Pregabalin may potentiate the effects of ethanol and lorazepam

Drug-Food Interactions

The rate of pregabalin absorption is decreased when given with food resulting in a decrease in C_{max} by approximately 25% to 30% and an increase in T_{max} to approximately 3 hours. However, administration of pregabalin with food has no clinically relevant effect on the total amount of pregabalin absorbed. Therefore, pregabalin can be taken with or without food

Drug-Herb Interactions

(pregabalin) has no known drug/herb interactions

Drug-Laboratory Interactions

LYRICA (pregabalin) has no known drug/laboratory test interactions

DOSAGE AND ADMINISTRATION

Dosing Considerations

Patients with Impaired Renal Function

Pregabalin is primarily eliminated from the systemic circulation by renal excretion as unchanged drug. In patients with a medical history of significant renal insufficiency, daily dosages should be reduced

accordingly (see <u>Dosage Adjustment Based on Renal Function</u>, below). In accordance with current clinical practice, if LYRICA (pregabalin) has to be discontinued, it is recommended this should be done gradually over a minimum of 1 week (see

WARNINGS AND PRECAUTIONS, Abrupt or Rapid Discontinuation

Adults

Neuropathic pain associated with diabetic peripheral neuropathy

Neuropannic pain associated with rateeur peripheral neuropany. The recommended starting dose for LYRIC4 is 150 mg/day, given in two or three divided doses (75 mg BID or 50 mg TID), with or without food in patients with a creatinine clearance rate of at least 60 mL/min. Efficacy of LYRIC4 has been demonstrated within the first week. Based on individual patient response and tolerability, the dose may be increased to 150 mg BID (300 mg/day) after one week. tolerability. The dose may be increased to 15 dm gBill you mg/roay) after one week. For patients who experience significant and ongoing pain and can tolerate pregabalin 300 mg/day well, maximum daily dose of 600 mg (300 mg twice a day, BID) can be used. However, in clinical trials, LYBICA 600 mg/day did not provide additional significant efficacy and patients treated with this dose experienced markedly higher rates of adverse events and discontinued the trial more frequently.

Neuropathic pain associated with postherpetic neuralgia

The recommended starting dose for LYRICA is 150 mg/day, given in two or three divided doses (75 mg BID or 50 mg TID), with or without food in patients with a creatinine clearance rate of at least 60 mL/min. Efficacy of LYRICA has been demonstrated within the first week. Based on individual patient response and tolerability, the dose may be increased to 150 mg BID (300 mg/day) after one week. toleratinity, the use finely be increased to 150 mig for 150 migrually rate one week. For patients who experience significant and ongoing pain and can tolerate pregabalin 300 mg/day well, maximum daily dose of 600 mg (300 mg twice a day, BID) can be used. However, in clinical trials, LYRIGA 600 mg/day did not provide additional significant efficacy and patients treated with this dose experienced markedly higher rates of adverse events and discontinued the trial more frequently.

Dosage Adjustment Based on Renal Function

LYRICA is primarily eliminated by renal excretion. Therefore, the dose should be adjusted for patients with reduced renal function. Pregabalin clearance is directly proportional to creatinine clearance. Therefore, dosing adjustment should be based on creatinine clearance ($C_{\rm lo}$), as indicated in Table 7. To use this desire table as

To use this dosing table, an estimate of the patient's creatinine clearance ($CL_{\mathbb{C}}$) in mL/min is needed. $CL_{\mathbb{C}}$ in mL/min may be estimated from serum creatinine (mg/dL) determination using the Cockcroft and Gault equation:

$$CL_{\odot} = \frac{[140 - age (years)] \times weight (kg)}{72 \times serum creatinine (mg/dL)}$$
 (x 0.85 for female patients)

Pregabalin is effectively removed from plasma by hemodialysis. Over a 4-hour hemodialysis treatment, plasma pregabalin concentrations are reduced by approximately 50%. For patients receiving hemodialysis, pregabalin daily dose should be adjusted based on renal function. In addition to the daily dose adjustment, a supplemental dose should be given immediately following every 4-hour hemodialysis treatment (see Table 7).

Table 7. Pregabalin Dosage Adjustment Based on Renal Function

Creatinine Clearance (CL _C ,) (mL/min)	Total P	regabalin Dai (mg/day)ª	Dose Regimen	
≥60	150	300	600	BID or TID
30-60	75	150	300	BID or TID
15-30	25-50	75	150	QD or BID
<15	25	25-50	75	QD

Patients on the 25 mg QD regimen: take one supplemental dose of 25 mg or 50 mg

Patients on the 25-50 mg QD regimen: take one supplemental dose of 50 mg or 75 mg

ents on the 75 mg QD regimen: take one supplemental dose of 100 mg or 150 mg

TID = Three divided doses: BID = Two divided doses: QD = Single daily dose Total daily dose (mg/day) should be divided as indicated by dose regimen to provide mg/dose.

b Supplementary dose is a single additional dose

Geriatrics (>65 years): Pregabalin oral clearance tended to decrease with increasing age. This decrease in pregabalin oral clearance is consistent with agerelated decreases in creatinine clearance. Reduction of pregabalin dose may be required in patients who have age-related compromised renal function.

Pediatrics (<18 years of age): The safety and efficacy of pregabalin in pediatric patients (<18 years of age) have not been established and its use in this patient population is not recommended.

Administration

en orally with or without food (see ACTION AND CLINICAL PHARMACOLOGY).

Signs, Symptoms and Laboratory Findings of Acute Overdosage in Humans

The highest known dose of pregabalin received in the clinical development program was 15,000 mg in 1 patient. The types of adverse events experienced by patients who received an overdose were not clinically different from other patients receiving recommended doses of pregabalin

Treatment or Management of Overdose

There is no specific antidote for overdose with pregabalin. If indicated, elimination of unabsorbed drug may be attempted by emesis or gastric lavage; usual precautions should be observed to maintain the airway. General supportive care of the patient is indicated including monitoring of vital signs and observation of the clinical status of the patient. A Certified Poison Control Center should be contacted for up-to-date information on the management of overdose with pregabalin.

Hemodialysis

Standard hemodialysis procedures result in significant clearance of pregabaling (approximately 50% in 4 hours) and should be considered in cases of overdose. Although hemodialysis has not been performed in the few known cases of overdose, it may be indicated by the patient's clinical state or in patients with significant renal

ACTION AND CLINICAL PHARMACOLOGY

Mechanism of Action

Pharmacodynamics

LYRICA (pregabalin) binds with high affinity to the alpha₂-delta protein (a calcium channel subunit) of brain tissues and has analgesic, antiepileptic and anxiolytic activity. Pregabalin is known chemically as (S)-3-(aminomethyl)-5-methylhexanoic

Although the mechanism of action of pregabalin is unknown, results with genetically Authough the mechanism of action of pregadant in sunknown, resuns with generically modified mice and with compounds structurally-related to pregabalin indicate that selective binding to the alpha-delta protein is required for analgesic, antiepileptic and anxiolytic action in animal models. In vitro, pregabalin reduces the release of several neurotransmitters, suggesting a modulatory action on calcium channel

Prenabalin does not mimic GABA at GABA, or GABA, recentors, nor does it augment GABA_A responses like benzodiazepines or barbiturates. In contrast to vascular calcium channel blockers, pregabalin does not alter systemic blood pressure or cardiac function. Various in vitro and in vivo results differentiate pregabalin from GABA uptake inhibitors or GABA transaminase inhibitors. In addition, pregabalin does not block sodium channels, it is not active at opiate receptors, it does not alter cyclooxygenase enzyme activity, it is not a serotonin agonist, it is not a dopamine antagonist, and it is not an inhibitor of dopamine, serotonin or noradrenaline

Pregabalin treatment reduces pain-related behavior in neuropathic animal models of diabetes, peripheral nerve damage or chemotherapeutic insult and in a model of musculoskeletal-associated pain. Pregabalin given intrathecally prevents pain-related behaviors and reduces pain-related behavior caused by spinally administered agents, suggesting that it acts directly on tissues of the spinal cord or brain.

Pharmacokinetics

All pharmacological actions following pregabalin administration are due to the All pharmacological actions following pregabatin administration are due to the activity of the parent compound; pregabatin is not appreciably metabolized in humans. Mean steady-state plasma pregabatin concentration-time profiles following 75, 300 and 600 mg/day given in equally divided doses every 8 hours (TID) and 600 mg/day given in equally divided doses every 12 hours (BID) are shown in Table 8. Pregabatin pharmacokinetics are linear over the recommended daily dose range. Inter-subject pharmacokinetic variability for pregabatin is low (<20%).

Table 8. Pregabalin Mean (CV%*) Steady-State Pharmacokinetic Parameter Values in Healthy Volunteers

Dose (mg)	Regimen	Daily Dose (mg/ day)	n	C _{maxss} (µg/ mL)	t _{max} (hr)	C _{minss} (µg/ mL)	AUC _(0-t) (μg•hr/ mL)	t _{1/2} (hr)	C _{L/F} (mL/ min)
25	TIDb	75	8	1.39	0.9	0.45	6.7	5.9	64.1
25	IIU°	/5		-19.5	-34.2	-25	-18.3	-17.3	-16.1
100	TID	300	6	5.03	0.8	1.94	25.2	6.3	68.9
100	טוו	300		-21.3	-31	-33.6	-23	-19.6	-20.9
200	TID	600	11	8.52	0.9	3.28	41.7	6.3	81
				-14.8	-22.2	-29.2	-12.8	-13.6	-11.7
300	BID:	600	8	9.07	1.4	2.6	59	6.7	85.1
				-10.5	-57.1	-15.5	-6.4	-16.2	-6.4

Time of peak plasma concentration at steady state

Steady-state trough plasma concentration

Area under the plasma concentration-time curve during one dosing interval VIIIC at steady state

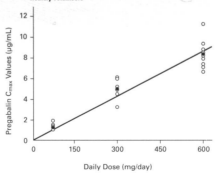
Flimination half-life

Oral clearance reent coefficient of variation

Total daily dose given in equally divided doses every 8 hours Total daily dose given in equally divided doses every 12 hours

Absorption: Pregabalin is rapidly absorbed when administered in the fasted state, with peak plasma concentrations occurring within 1.5 hours following both single- and multiple-dose administration. Pregabalin oral bioavailability is ≥90% and is independent of dose. C_{max} (Figure 1) and AUC values increase proportionally following single- and multiple-dose administration. Following repeated administration, steady state is achieved within 24 to 48 hours. Multiple dose pharmacokinetics are predictable from single-dose data.

Figure 1. Individual and Mean Steady-State Pregabalin C_{max} Values Following 75, 300 and 600 mg/day Given in Equally Divided Doses TID (q8h) to Healthy Volunteers



Solid line is the regression line going through the origin; individual (O) and mean (\spadesuit) values.

Distribution: In preclinical studies, pregabalin has been shown to readily cross the blood brain barrier in mice, rats and monkeys. Pregabalin is a substrate for system L transporter which is responsible for the transport of large amino acids across the blood-brain barrier. Pregabalin has been shown to cross the placenta in rats and is present in the milk of lactating rats. In humans, the apparent volume of distribution of pregabalin following oral administration is approximately 0.5 L/kg. Pregabalin is not bound to plasma proteins. At clinically efficacious doses of 150 and 600 mg/day, the average steady-state plasma pregabalin concentrations were approximately 1.5 and 6.0 µg/ml., respectively.

approximately 1.3 and on bignit, respectively. Metabolism: Pregabalin undergoes negligible metabolism in humans. Following a dose of radiolabeled pregabalin, approximately 98% of the radioactivity recovered in the urine was unchanged pregabalin. The N-methylated derivative of pregabalin, the major metabolite of pregabalin found in urine, accounted for 0.9% of the dose. In preclinical studies, pregabalin (S-enantiomer) did not undergo racemization to the R-enantiomer in mice_rats_rabbits or monkeys.

Excretion: Pregabalin is eliminated from the systemic circulation primarily by renal excretion as unchanged drug. Pregabalin mean $t_{1/2}$ is 6.3 hours. Pregabalin elimination is proportional to creatinine clearance. Pregabalin clearance is reduced in patients with impaired renal function (see **DOSAGE AND ADMINISTRATION**).

Special Populations and Conditions

Pregabalin undergoes negligible metabolism, is not bound to plasma proteins and is eliminated predominately as unchanged drug by renal excretion. Clinically important differences in pregabalin pharmacokinetics due to race and gender have not been observed and are not anticinated

Pediatrics: Pharmacokinetics of pregabalin have not been studied in paediatric

Geriatrics: Pregabalin oral clearance tended to decrease with increasing age. This decrease in pregabalin oral clearance is consistent with age-related decreases in creatinine clearance. Reduction of pregabalin dose may be required in patients who have age-related compromised renal function (see WARNINGS AND PRECAUTIONS and DOSAGE AND ADMINISTRATION).

Gender: A population pharmacokinetic analysis of the Phase 2/3 clinical program showed that the relationship between daily dose and pregabalin drug exposure is similar between genders when adjusted for gender-related differences in creatinine

Race: A population pharmacokinetic analysis of the Phase 2/3 clinical program showed that the relationship between daily dose and pregabalin drug exposure is similar among Caucasians, Blacks and Hispanics.

similar among Jaucasians, blacks and inspanies.

Renal Insufficiency: Because renal elimination is the major elimination pathway, dosage reduction in patients with renal dysfunction is necessary. Pregabalin is effectively removed from plasma by hemodialysis. Following a 4-hour hemodialysis treatment, plasma pregabalini concentrations are reduced by approximately 50%. For patients on hemodialysis, dosing must be modified (see DOSAGE AND

ADMINISTRATION) STORAGE AND STARILITY

Store at 15°C-30°C

DOSAGE FORMS, COMPOSITION AND PACKAGING

Each capsule of LYRICA (pregabalin) contains 25, 50, 75, 150 or 300 mg pregabalin, lactose monohydrate, maize starch and talc. The capsule shells contain gelatin and titanium dioxide. In addition, the orange capsule shells contain red iron oxide and trainion blokke. In author, the origing bassive steins contain the buffer capability and the white capability shells contain sodium lauryl sulfate and colloidal silicon dioxide. Colloidal silicon dioxide is a manufacturing aid, which may not be present. The markings on the capsules are in black ink, which contains shellac, black iron oxide, propylene glycol, potassium hydroxide and water

Capsules are packaged in HDPE bottles containing 60 capsules, and PVC/aluminum

PHARMACEUTICAL INFORMATION

Drug Substance Proper name:

pregabalin

Chemical name

(S)-3-(aminomethyl)-5-methylhexanoic acid

Molecular formula C.H.-NO

Molecular mass: 159.23

Physicochemical properties

Structural formula:

Pregabalin is a white crystalline solid. It is soluble in water and in both basic and acidic aqueous solutions

Product Monograph available upon request

Last revised: June 3, 2005

Neterences:

1. LYRICA Product Monograph, June 2005.

2. Data on file, Pfizer Canada Inc., study 1008-196.

3. Freynhagen R, et al. Efficacy of pregabalin in neuropathic pain evaluated in a 12-week, randomised, double-blind, multicentre, placebo-controlled trial of flexible- and fixed-dose regimens. Pain 2005:115:254-263



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Pfizer Canada Inc., licensee





power you can trust™

LIPITOR (atorvastatin calcium) 10 mg, 20 mg, 40 mg and 80 mg tablets

THERAPEUTIC CLASSIFICATION: Lipid Metabolism Regulator

ACTIONS AND CLINICAL PHARMACOLOGY

Please refer to the Product Monograph for complete ACTIONS AND CLINICAL PHARMACOLOGY information.

INDICATIONS AND CLINICAL LISE

Hypercholesterolemia

LIPITOR (atorivastatin calcium) is indicated as an adjunct to lifestyle changes, including diet (at least equivalent to the Adult Treatment Panel III (ATP III) TLC diet), for the reduction of elevated total cholesterol (total-C), LDL-C, TG and apolipoprotein B (apo B) in hyperlipidemic and dyslipidemic conditions, when response to diet and other nonpharmacological measures alone

Primary hypercholesterolemia (Type Ila); Combined (mixed) hyperlipidemia (Type Ilb), including familial combined hyperlipidemia, regardless of whether cholesterol or triglycerides are the lipid abnormality of concern; Dysbetalipoproteinemia (Type III); Hypertriglyceridemia (Type III); Familial hypercholesterolemia (hypercholesterolemia, LPITOR should be used as an adjunct to treatments such as LDL aphresis, or as monotherapy if such treatments are not available; an adjunct to diet to reduce total-C, LDL-C, and apo B levels in boys and postmenarchal girls, 10 to 17 years of age with heterozygous familial hypercholesterolemia, if after an adequate trial of diet therapy the following findings are still present:

a. LDL-C remains ≥4.9 mmol/L (190 mg/dL) or

- b. LDL-C remains ≥4.1 mmol/L (160 mg/dL) and:

 there is a positive family history of premature cardiovascular disease or
 - two or more other CVD risk factors are present in the pediatric patient

LIPTOR also raises HDL-cholesterol and therefore lowers the LDL-C/HDL-C and total-C/HDL-C ratios in patients with primary hypercholesterolemia and combined (mixed) hyperlipidemia (Fredrickson Type IIa and IIb dyslipidemia). In pooled data from 24 controlled clinical trials, LIPTOR raised HDL-C levels 5%-7% in primary hypercholesterolemic (Type IIa) patients and 10%-15% in mixed (Type IIb) dyslipidemic patients

In clinical trials, LIPTOR (10 to 80 mg/day) significantly improved lipid profiles in patients with a wide variety of hyperlipidemic and dyslipidemic conditions. In 2 dose-response studies in mildly to moderately hyperlipidemic patients (Fredrickson Types IIa and IIb), LIPTOR reduced the levels of total cholesterol (29-45%), LDL-C (39-60%), apo B (32-50%), TG (19-37%), and increased high density lipoprotein cholesterol (HDL-C) levels (5-9%). Comparable responses were achieved in patients with heterozygous familial hypercholesterolemia, non-familial forms of hypercholesterolemia, combined hyperlipidemia, including familial combined hyperlipidemia and patients with non-insulin dependent diabetes mellitus. In patients with hypertriglyceridemia (Type IV), LIPITOR (10 to 80 mg daily) reduced TG (25-56%) and LDL-C levels (23-40%). LIPITOR has not been studied in conditions where the major abnormality is elevation of chylomicrons (TG levels >11 mmol/L), i.e., Types I and V

In an open-label study in patients with dysbetalipoproteinemia (Type III), LIPITOR (10 to 80 mg daily) reduced total-C (40-57%), TG (40-56%) and IDL-C + VLDL-C levels (34-58%).

In an open-label study in patients with homozygous familial hypercholesterolemia (FH), LIPITOR (10 to 80 mg daily) reduced mean LDL-C levels (22%), in a pilot study, LIPITOR 80 mg/day showed a mean LDL-C lowering of 30% for patients not on plasmapheresis and of 31% for patients who continued plasmapheresis. A mean LDL-C lowering of 35% was observed in receptor defective patients and of 19% in receptor negative patients.

Prior to initiating therapy with LIPITOR, secondary causes should be excluded for elevations in plasma lipid levels (e.g., poorly controlled diabetes mellitus, hypothyroidism, nephrotic syndrome, dysproteinemias, obstructive liver disease and alcoholism), and a lipid profile performed to measure total cholesterol, LDL-C, HDL-C and TG. For patients with TG <4.52 mmol/L (<400 mg/dL), LDL-C can be estimated using the following equation

LDL-C (mmol/L) = total-C - [(0.37 x (TG) + HDL-C)] LDL-C (mg/dL) = total-C - [(0.2 x (TG) + HDL-C)]'

For patients with TG levels >4.52 mmol/L (>400 mg/dL), this equation is less accurate and LDL-C concentrations should be measured directly or by ultracentrifugation

Patients with high or very high triglyceride levels, i.e., >2.2 mmol/L (200 mg/dL) or >5.6 mmol/L (500 mg/dL), respectively, may require triglyceride-lowering therapy (fenofibrate, bezafibrate or nicotinic acid) alone or in combination with LIPITOR.

In general, combination therapy with fibrates must be undertaken cautiously and only after risk-benefit analysis (see WARNINGS – Muscle Effects; PRECAUTIONS – Pharmacokinetic Interaction Studies and Potential Drug

Elevated serum triglycerides are most often observed in patients with the metabolic syndrome (abdominal obesity, atherogenic dyslipidemia [elevated triglycerides, small dense LDL particles and low HDL-cholesterol], insulin resistance with or without glucose intolerance, raised blood pressure and prothrombic and proinflammatory states).

(For the treatment of specific dysligidemias, refer to the Report of the Canadian Working Group on Hypercholesterolemia and Other Dyslipidemias or to the US NCEP Expert Panel on Detection, Evaluation and Treatment of High Blood Cholesterol in Adults [Adult Treatment Panel III], under REFERENCES).

When drugs are prescribed, attention to therapeutic lifestyle changes (reduced intake of saturated fats and cholesterol, weight reduction, increased physical activity, ingestion of soluble fibres) should always be maintained and reinforced.

Prevention of Cardiovascular Disease

LIPITOR is indicated to reduce the risk of myocardial infarction in adult hypertensive patients without clinically evident coronary heart disease, but with at least 3 additional risk factors for coronary heart diseases such as: age \$55 years, male sex, smoking, type 2 diabetes, left ventricular hypertrophy, other specified abnormalities on ECG, microalbuminuria or proteinuria, ratio of plasma total cholesterol to HDL-C >6 or premature family history of coronary heart disease.

LIPITOR is also indicated to reduce the risk of myocardial infarction and stroke in adult patients with type 2 diabetes mellitus and hypertension without clinically evident coronary heart disease, but with other risk factors such as age ≥55 years, retinopathy, albuminuria or smoking.

Hypersensitivity to any component of this medication.

Active liver disease or unexplained persistent elevations of serum transaminases exceeding 3 times the upper limit of normal (see WARNINGS)

Pregnancy and nursing women: Cholesterol and other products of cholesterol biosynthesis are essential components for fetal development (including synthesis of steroids and cell membranes). LIPITOR should be administered to women of childbearing age only when such patients are highly unlikely to conceive and have been informed of the possible harm. If the patient becomes pregnant while taking LIPITOR, the drug should be discontinued immediately and the patient apprised of the potential harm to the fetus. Atherosclerosis being a chronic process, discontinuation of lipid metabolism regulating drugs during pregnancy should have little impact on the outcome of long-term therapy of primary hypercholesterolemia (see **PRECAUTIONS – Use in Pregnancy, Use in Nursing Mothers**).

Pharmacokinetic Interactions

The use of HMG-CoA reductase inhibitors has been associated with severe myopathy, including rhabdomyolysis, which may be more frequent when they are co-administered with drugs that inhibit the cytochrome P-450 enzyme system. Atomastatin is metabolized by cytochrome P-450 isoform 3A4 and as such may interact with agents that inhibit this enzyme (see WARNINGS – Muscle Effects; PRECAUTIONS – Pharmacokinetic Interaction Studies and Potential Drug Interactions, Cytochrome P-450-mediated Interactions).

Effects on skeletal muscle such as myalgia, myopathy and very rarely, rhabdomyolysis have been reported in patients treated with LIPITOR. Very rare cases of rhabdomyolysis with acute renal failure secondary to myoglobinuria, have been reported with LIPITOR and other HMG-CoA reductase inhibitors.

Myopathy, defined as muscle pain or muscle weakness in conjunction with increases in creatine kinase (CK) values to >10 times the upper limit of normal, should be considered in any patient with diffuse myalgia, muscle tenderness or weakness, and/or marked elevation of CK. Patients should be advised to report promptly any unexplained muscle pain, tenderness or weakness, particularly if accompanied by malaise or fever. Patients who develop any signs or symptoms suggestive of myopathy should have their CK levels measured. LIPITOR therapy should be discontinued if markedly elevated CK levels are measured or myopathy is diagnosed or suspected.

Predisposing Factors for Myopathy/Rhabdomyolysis: LIPITOR, as with other HMG-CoA reductase inhibitors, should be prescribed with caution in patients with predisposing factors for myopathy/rhabdomyolysis. Such factors include

Personal or family history of hereditary muscular disorders; Previous history of muscle toxicity with another HMG-CoA reductase inhibitor; Concomitant use of a fibrate or niacin; Hypothyroidism; Alcohol abuse; Excessive physical exercise; Age >70 years; Renal impairment; Hepatic impairment; Diabetes with hepatic fatty change; Surgery and trauma; Fraility; Situations where an increase in plasma levels of active ingredient may occur.

LIPTOR therapy should be temporarily withheld or discontinued in any patient with an acute serious condition suggestive of myopathy or having a risk factor predisposing to the development of renal failure secondary to rhabdomyolysis (such as sepsis, severe acute infection, hypotension, major surgery, trauma, severe metabolic, endocrine and electrolyte disorders and uncontrolled seizures).

LIPITOR therapy should be discontinued if markedly elevated CPK levels occur or myopathy is diagnosed or suspected. The risk of myopathy and rhabdomyolysis during treatment with HMG-CoA reductase inhibitors is increased with concurrent administration of cyclosporin, fibric acid derivatives, erythromycin, clarithromycin, niacin (incotinic acid), azole antifungals or netazodone. As there is no experience to date with the use of LIPHOR given concurrently with these drugs, with the exception of pharmacokinetic studies conducted in healthy subjects with erythromycin and clarithromycin, the benefits and risks of such combined therapy should be carefully considered (see PRECAUTIONS - Pharmacokinetic Interaction Studies and Potential Drug Interactions

Hepatic Effects

In clinical trials, persistent increases in serum transaminases >3 times the upper limit of normal occurred in <1% of patients who received LIPITOR. When the dosage of LIPITOR was reduced, or when drug treatment was interrupted or discontinued, serum transaminase levels returned to pretreatment levels. The increases were generally not associated with jaundice or other clinical signs or symptoms. Most patients continued treatment with a reduced dose of LIPITOR without clinical sequelae.

Liver function tests should be performed before the initiation of treatment, and periodically thereafter. Special attention should be paid to patients who develop elevated serum transaminase levels, and in these patients measurements should be repeated promptly and then performed more frequently.

If increases in alanine aminotransferase (ALT) or aspartate aminotransferase (AST) show evidence of progression, particularly if they rise to >3 times the upper limit of normal and are persistent, the dosage should be reduced or the drug discontinued.

LIPITOR, as well as other HMG-CoA reductase inhibitors, should be used with caution in patients who consume substantial quantities of alcohol and/or have a past history of liver disease. Active liver disease or unexplained transaminase elevations are contraindications to the use of LIPITOR; if such a condition should develop during therapy, the drug should be discontinued.

PRECAUTIONS

General

Before instituting therapy with LIPITOR (atorvastatin calcium), an attempt should be made to control elevated serum lipoprotein levels with appropriate diet, exercise, and weight reduction in overweight patients, and to treat other underlying medical problems (see INDICATIONS AND CLINICAL USE). Patients should be advised to inform subsequent physicians of the prior use of LIPITOR or any other lipid-lowering agents

Effect on the Lens

Current long-term data from clinical trials do not indicate an adverse effect of atorvastatin on the human lens

Effect on Ubiquinone (CoQ10) Levels

Significant decreases in circulating ubiquinone levels in patients treated with atorvastatin and other statins have been observed. The clinical significance of a potential long-term statin-induced deficiency of ubiquinone has not been established. It has been reported that a decrease in myocardial ubiquinone levels could lead to impaired cardiac function in patients with borderline congestive heart failure.

Effect on Lipoprotein (a)

In some patients, the beneficial effect of lowered total cholesterol and LDL-C levels may be partly blunted by a concomitant increase in Lp(a) lipoprotein concentrations. Present knowledge suggests the importance of high Lp(a) levels as an emerging risk factor for coronary heart disease. It is thus desirable to maintain and reinforce lifestyle changes in high risk patients placed on atorvastatin therapy.

Hypersensitivity

An apparent hypersensitivity syndrome has been reported with other HMG-CoA reductase inhibitors which has included 1 or more of the following features: anaphylaxis, angloedema, lupus erythematous-like syndrome, polymyalgia rheumatica, vasculitis, purpura, thrombocytopenia, leukopenia, hemolytic anemia, positive ANA, ESR increase, eosinophilia, arthritis, rathralgia, urticaria, asthenia, photosensitivity, fever, chills, flushing, malaise, dyspnea, toxic epidermal necrolysis, erythema multiforme, including Stevens-Johnson syndrome. Although to date hypersensitivity syndrome has not been described as such, LIPITOR should be discontinued if hypersensitivity is suspected.

Use in Pregnancy

LIPITOR is contraindicated during pregnancy (see CONTRAINDICATIONS).

There are no data on the use of LIPITOR during pregnancy, LIPITOR should be administered to women of childbearing age only when such patients are highly unlikely to conceive and have been informed of the potential hazards. If the patient becomes pregnant while taking LIPITOR, the drug should be discontinued and the patient apprised of the potential risk to the fetus.

Use in Nursing Mothers

In rats, milk concentrations of atoryastatin are similar to those in plasma. It is not known whether this drug is excreted in human milk. Because of the potential for adverse reactions in nursing infants, women taking LIPITOR should not breast-feed (see **CONTRAINDICATIONS**).

Pediatric Use

Safety and effectiveness of LIPITOR in patients 10-17 years of age (N=140) with heterozygous familial hypercholesterolemia have been evaluated in a controlled clinical trial of 6 months duration in adolescent boys and postmenarchal girls. Patients treated with LIPITOR had a safety and tolerability profile generally similar to that of placebo. Doses >20 mg have not been studied in this patient population.

LIPITOR had no effect on growth or sexual maturation in boys and in girls. The effects on menstrual cycle were not assessed [see ADVERSE REACTIONS and DOSAGE AND ADMINISTRATION for Heterozygous Familial Hypercholesterolemia in

Pediatric Patients (10-17 years of age)).

Adolescent females should be counselled on appropriate contraceptive methods while on LIPITOR therapy (see CONTRAINDICATIONS; PRECAUTIONS – Use in Pregnancy). LIPITOR has not been studied in controlled clinical trials involving in pre-pubertal patients or patients younger than 10 years of age.

Doses of LIPITOR up to 80 mg/day for 1 year have been evaluated in 8 pediatric patients with homozygous familial hypercholesterolemia

Geriatric Use

Treatment experience in adults 70 years or older (N=221) with doses of LIPITOR up to 80 mg/day has demonstrated that the safety and effectiveness of atorvastatin in this population was similar to that of patients <70 years of age. Pharmacokinetic evaluation of atorvastatin in subjects over the age of 65 years indicates an increased AUC. As a precautionary measure, the lowest dose should be administered initially.

Elderly patients may be more susceptible to myopathy (see WARNINGS - Muscle Effects - Predisposing Factors for Myonathy/Rhabdomyolysis)

Renal Insufficiency

Plasma concentrations and LDL-C lowering efficacy of LIPITOR was shown to be similar in patients with moderate renal insufficiency compared with patients with normal renal function. However, since several cases of mabdomyolysis have been reported in patients with a history of renal insufficiency of unknown severity, as a precautionary measure and pending further experience in renal disease, the lowest dose (10 mg/day) of LIPITOR should be used in these patients. Similar precautions apply in patients with severe renal insufficiency (creatine clearance < 30 mL/min (<0.5 mL/sec)); the lowest dosage should be used and implemented cautiously (see WARNINGS – Muscle Effects; PRECAUTIONS – Pharmacokinetic Interaction Studies and Potential Drug Interactions). Refer also to DOSAGE AND ADMINISTRATION.

Endocrine Function

HMG-CoA reductase inhibitors interfere with cholesterol synthesis and as such might theoretically blunt adrenal and/or gonadal steroid production. Clinical studies with atorvastatin and other HMG-CoA reductase inhibitors have suggested that these agents do not reduce plasma cortisol concentration or impair adrenal reserve and do not reduce basal plasma testosterone concentration. However, the effects of HMG-CoA reductase inhibitors on male fertility have not been studied in adequate numbers of patients. The effects, if any, on the pituitary-gonadal axis in premenopausal women are unknown.

Patients treated with atorvastatin who develop clinical evidence of endocrine dysfunction should be evaluated appropriately. Caution should be exercised if an HMG-CoA reductase inhibitor or other agent used to lower cholesterol levels is administered to patients receiving other drugs (e.g., ketoconazole, spironolactone or cimetidine) that may decrease the levels of endogenous steroid hormones.

Pharmacokinetic Interaction Studies and Potential Drug Interactions

Pharmacokinetic interaction studies conducted with drugs in healthy subjects may not detect the possibility of a potential drug interaction in some patients due to differences in underlying diseases and use of concomitant medications (see PRECAUTIONS - Geriatric Use, Renal Insufficiency; Patients with Severe Hypercholesterolemia).

Concomitant Therapy with Other Lipid Metabolism Regulators: Based on post-marketing surveillance, gemfibrozil, fenofibrate, other fibrates and lipid-lowering doses of niacin (nicotinic acid) may increase the risk of myopathy when given concomitantly with HMG-CoA reductase inhibitors, probably because they can produce myopathy when given alone (see WARNINGS - Muscle Effects). Therefore, combined drug therapy should be approached with caution.

Bile Acid Sequestrants:

Patients with mild to moderate hypercholesterolemia; LDL-C reduction was greater when LIPITOR 10 mg and colestipol 20 g were coadministered (-45%) than when either drug was administered alone (-35% for LIPITOR and -22% for colestipol).

Patients with severe hypercholesterolemia: LDL-C reduction was similar (-53%) when LIPITOR 40 mg and colestipol 20 g were coadministered when compared to LIPTOR 80 mg alone. Plasma concentration of atorvastatin was lower (approximately 26%) when LIPTOR 40 mg plus colestipol 20 g were coadministered compared with LIPTOR 40 mg alone. However, the combination drug therapy was less effective in lowering triglycerides than LIPTOR monotherapy in both types of hypercholesterolemic patients.

When LIPITOR is used concurrently with colestipol or any other resin, an interval of at least 2 hours should be maintained between the two drugs, since the absorption of LIPITOR may be impaired by the resin.

Fibric Acid Derivatives (Gemfibrozil, Fenofibrate, Bezafibrate) and Niacin (nicotinic acid): Although there is limited experience with the use of LIPITOR given concurrently with fibric acid derivatives and niacin, the benefits and risks of such combined therapy should be carefully considered. The risk of myopathy during treatment with drugs in this class, including atorvastatin, is increased with concurrent administration (see WARNINGS – Muscle Effects).

Coumarin Anticoagulants: LIPITOR had no clinically significant effect on prothrombin time when administered to patients receiving chronic warfarin therapy

Digoxin: In healthy subjects, digoxin pharmacokinetics at steady-state were not significantly altered by coadministration of digoxin 0.25 mg and LIPITOR 10 mg daily. However, digoxin steady-state concentrations increased approximately 20% following

coadministration of digoxin 0.25 mg and LIPITOR 80 mg daily. Patients taking digoxin should be monitored appropriately. Antihypertensive agents (amlodipine): In clinical studies, LIPITOR was used concomitantly with antihypertensive agents without evidence to date of clinically significant adverse interactions. In healthy subjects, atorvastatin pharmacokinetics were not altered by the coadministration of LIPITOR 80 mg and amlodipine 10 mg at steady state.

(quinapril): In a randomized, open-label study in healthy subjects, steady-state quinapril dosing (80 mg QD) did not significantly affect the pharmacokinetic profile of atorvastatin tablets (10 mg QD).

Oral Contraceptives and Hormone Replacement Therapy: Coadministration of LIPITOR with an oral contraceptive containing 1 mg norethindrone and 35 µg ethinyl estradiol increased plasma concentrations (AIUC levels) of norethindrone and ethinyl estradiol by approximately 30% and 20%, respectively. These increases should be considered when selecting an oral contraceptive. In clinical studies, LIPITOR was used concomitantly with estrogen replacement therapy without evidence of clinically significant adverse interactions.

Antacids: Administration of aluminum and magnesium based antacids, such as Maalox® TC Suspension, with LIPITOR decreased plasma concentrations of LIPTOR by approximately 35%. LDL-C reduction was not altered but the triglyceride-lowering effect of LIPTOR may be affected.

Cimetidine: Administration of cimetidine with LIPITOR did not alter plasma concentrations or the LDL-C lowering efficacy of LIPITOR, however, the triglyceride-lowering effect of LIPITOR was reduced from 34% to 26%

Cytochrome P-450-mediated Interactions: Atorvastatin is metabolized by the cytochrome P-450 isoenzyme, CYP 3A4 Frythromycin, a CYP 3A4 inhibitor, increased atorvastatin plasma levels by 40%. Coadministration of CYP 3A4 inhibitors, such antifungal agents (i.e., itraconazole, ketoconazole), protease inhibitors, or the antidepressant nefazodone, may have the potential to increase plasma concentrations of HMG-CoA reductase inhibitors, including LIPITOR. Caution should thus be exercised with concomitant use of these agents (see WARNINGS – Pharmacokinetic Interactions, Muscle Effects; PRECAUTIONS – Renal Insufficiency, Endocrine Function; DOSAGE AND ADMINISTRATION).

Terfenadine: In healthy subjects, coadministration of maximum doses of atorvastatin (80 mg) and terfenadine (120 mg) a CYP 3A4 substrate, was shown to produce a modest increase in terfenadine AUC. The QTc interval remained unchanged However, since an interaction between these two drugs cannot be excluded in patients with predisposing factors for arrhythmia, (e.g., pre-existing prolonged QT interval, severe coronary artery disease, hypokalemia), caution should be exercised when these agents are coadministered (see WARNINGS – Pharmacokinetic Interactions; DOSAGE AND ADMINISTRATION)

Antipyrine: Antipyrine was used as a non-specific model for drugs metabolized by the microsomal hepatic enzyme (cytochrome P-450) system. LIPITOR had no effect on the pharmacokinetics of antipyrine, thus interactions with other druos netabolized via the same cytochrome isozymes are not expected.

Macrolide Antibiotics (azithromycin, clarithromycin, erythromycin): In healthy adults, coadministration of LIPITOR (10 mg 0D) and azithromycin (500 mg 0D) did not significantly after the plasma concentrations of atorvastatin. However, coadministration of atorvastatin (10 mg 0D) with erythromycin (500 mg 0ID) or clarithromycin (500 mg BID), which are both CYP 3A4 inhibitors, increased plasma concentrations of atorvastatin by approximately 40% and 80%, respectively (see WARNINGS - Muscle Effects)

Protease Inhibitors (nelfinavir mesylate): In healthy adults, coadministration of nelfinavir mesylate (1250 mg BID), a known CVP 3A4 inhibitor, and atorvastatin (10 mg 0D) resulted in increased plasma concentrations of atorvastatin. AUC and C_{max} of atorvastatin were increased by 74% and 122% respectively.

Patients with Severe Hypercholesterolemia

Higher drug dosages (80 mg/day) required for some patients with severe hypercholesterolemia (including familial hypercholestrolemia) are associated with increased plasma levels of atovastatin. Caution should be exercised in such patients who are also severely renally impaired, elderly, or are concomitantly being administered digoxin or CYP 3A4 inhibitors (see WARNINGS – Pharmacokinetic Interactions, Muscle Effects; PRECAUTIONS – Pharmacokinetic Interactions of the Company o

LIPITOR may elevate serum transaminase and creatine kinase levels (from skeletal muscle). In the differential diagnosis of chest pain in a patient on therapy with LIPITOR, cardiac and noncardiac fractions of these enzymes should be determined.

ADVERSE REACTIONS

LIPITOR is generally well-tolerated. Adverse reactions have usually been mild and transient. In controlled clinical studies (placebo-controlled and active-controlled comparative studies with other lipid-lowering agents) involving 2502 patients, <2% of patients were discontinued due to adverse experiences attributable to LIPITOR. Of these 2502 patients, 1721 were treated for at least 6 months and 1253 for 1 year or more.

Adverse experiences occurring at an incidence ≥1% in patients participating in placebo-controlled clinical studies of LIPITOR and reported to be possibly, probably or definitely drug related include constipation, diarrhea, dyspepsia, flatulence, nausea, headache, pain, myalgia and asthenia.

The following additional adverse events were reported in clinical trials (not all have been associated with a causal relationship to LIPITOR therapy); muscle cramps, myositis, myopathy, paresthesia, peripheral neuropathy, pancreatitis, hepatitis, cholestatic jaundice, anorexia, vomiting, alopecia, pruritus, rash, impotence, hyperglycemia and hypoglycemia.

Heterozygous Familial Hypercholesterolemia in Pediatric Patients (ages 10-17 years)

In a 26-week controlled study in boys and postmenarchal girls (n=187, where 140 patients received LIPITOR), the safety and tolerability profile of LIPITOR 10 to 20 mg daily was similar to that of placebo. The adverse events reported in ≥1% of patients were abdominal pain, depression and headache (see **PRECAUTIONS – Pediatric Use**).

Laboratory Changes and Adverse Events

The criteria for clinically significant laboratory changes were >3 X the upper limit of normal (ULN) for liver enzymes, and >5 X ULN for creatine kinase. A total of 8 unique subjects met one or more of these criteria during the double-blind phase. Hence, the incidence of patients who experienced abnormally high enzymatic levels (AST/ALT and creatine kinase) was >4% (8/187).

Five atorvastatin and one placebo subjects had increases in CK >5 X ULN during the double-blind phase; two of the five atorvastatin-treated subjects had increases in CK >10 X ULN. Two subjects had clinically significant increases in ALT.

Post-Market Adverse Drug Reaction: The following adverse events have also been reported during post-marketing experience with LIPITOR, regardless of causality assessment: Very rare reports: severe myopathy with or without rhabdomyolysis (see WARNINGS – Muscle Effects; PRECAUTIONS – Renal Insufficiency, Pharmacokinetic Interaction Studies and Potential Drug Interactions, Isolated reports: Opencomastia, thromboryopenia, arthraigia and allergic reactions including urticaria, angioneurotic edema, anaphylaxis and bullous rashes (including erythema multiforme, Stevens-Johnson syndrome and toxic epidermal necrolysis) and fatigue. These may have no causal relationship to atorvastatin.

Abnormal Hematologic and Clinical Chemistry Findings

Ophthalmologic observations: see PRECAUTIONS

Laboratory Tests: Increases in serum transaminase levels have been noted in clinical trials (see PRECAUTIONS).

DOSAGE AND ADMINISTRATION

Patients should be placed on a standard cholesterol-lowering diet [at least equivalent to the Adult Treatment Panel III (ATP III) TLC diet] before receiving LIPITOR, and should continue on this diet during treatment with LIPITOR. If appropriate, a program of weight control and physical exercise should be implemented.

Prior to initiating therapy with LIPITOR, secondary causes for elevations in plasma lipid levels should be excluded. A lipid profile should also be performed

Primary Hypercholesterolemia and Combined (Mixed) Dyslipidemia, Including Familial Combined Hyperlipidemia

The recommended starting dose of LIPITOR is 10 or 20 mg once daily, depending on the patient's LDL-C reduction required The recommended starting oose of LIPTOR is 10 or 20 mg once daily, depending on the patients LID-C-creduction requires (see Tables 1 and 2). Patients who require a large reduction in LID-C. (more than 45%) may be started at 40 mg once daily. The dosage range of LIPTOR is 10 to 80 mg once daily. Doses can be given at any time of the day, with or without food, and should preferably be given in the evening. A significant therapeutic response is evident within 2 weeks, and the maximum response is usually achieved within 2-4 weeks. The response is maintained during chronic therapy. Adjustments of dosage, if necessary, should be made at intervals of 2 to 4 weeks. The maximum dose is 80 mg/day.

TABLE 1. Dose-Response in Patients With Mild-to-Moderate Hypercholesterolemia (Mean Percent Change from Baseline)

	LIPITOR Dose (mg/day)					
Lipid Parameter	10 (N=22)	20 (N=20)	40 (N=21)	80 (N=23)		
Total-C: 7.1 mmol/L ^b (273 mg/dL) ^b	-29	-33	-37	-45		
LDL-C: 4.9 mmol/L ^b (190 mg/dL) ^b	-39	-43	-50	-60		

Results are pooled from 2 dose-response studies

The dosage of LIPITOR should be individualized according to the baseline LDL-C, total-C/HDL-C ratio and/or TG levels to achieve the recommended target lipid values at the lowest dose needed to achieve the LDL-C target (see Recommendations or the Management of Dyslipidemia and the Prevention of Cardiovascular Disease (Canada), summarized below in Table 2, and/or the Third Report of the US National Cholesterol Education Program (NCEP Adult Treatment Panel III)), and the patient's response. Lipid levels should be monitored periodically and, if necessary, the dose of LIPITOR adjusted based on target lipid levels recommended by guidelines.

TABLE 2. Canadian Recommendations for the Target Lipid Values Based on Level of Risk

Risk Category	Target Levels				
	LDL-C level (mmol/L)		Total-C/HDL-C ratio		
High¹ (10-year risk of CAD ≥20%, or a history of diabetes mellitus¹¹ or any atherosclerotic disease)	<2.5		<4.0		
Moderate (10-year risk 11%-19%)	<3.5	and	<5.0		
Low ^{†††} (10-year risk ≤10%)	<4.5	and	<6.0		

Note: LDL-C = low-density lipoprotein cholesterol.

Apolipoprotein B can be used as an alternative measurement, particularly for follow-up of patients treated with statins. An potable value of a patient stream as an electromy incasurement, particularly for rollow-up or patients freated with statins. An optimal level of apolipoprotein B in a patient at high risk is <0.9 g/L, in a patient at moderate risk <1.05 g/L and in a patient at low risk <1.2 g/L.

*Includes patients with chronic kidney disease and those undergoing long-term dialysis.

In the 'very low' risk stratum, treatment may be deferred if the 10-year estimate of cardiovascular disease is <5% and the LDL-C level is <5.0 mmol/L.

Severe Dyslipidemias

In patients with severe dyslipidemias, including homozygous and heterozygous familial hypercholesterolemia and dysbetalipoproteinemia (Type III), higher dosages (up to 80 mg/day) may be required (see WARNINGS – Pharmacokinetic Interactions, Muscle Effects; PRECAUTIONS – Pharmacokinetic Interaction Studies and Potential Drug Interactions). Heterozygous Familial Hypercholesterolemia in Pediatric Patients (10-17 years of age)

In this population, the recommended starting dose of LIPITOR is 10 mg/day; the maximum recommended dose is 20 mg/day (doses >20 mg/day have not been studied in this patient population). Doses should be individualized according to the recommended goal of therapy (see NCEP Pediatric Panel Guidelines; INDICATIONS AND CLINICAL USE). Adjustments should be made at intervals of 4 weeks or more.

NCEP (National Cholesterol Education Program) Pediatric Panel Guidelines: Classification of cholesterol levels in pediatric patients with a familial history of hypercholesterolemia or premature cardiovascular disease is summarized below:

Category	Total-C (mmol/L [mg/dL])	LDL-C (mmol/L [mg/dL])	
Acceptable	<4.4 [170]	<2.8 [110]	
Borderline	4.4-5.1 [170-199]	2.8-3.3 [110-129]	
High	≥5.2 [200]	≥3.4 [130]	

Concomitant Therapy

See PRECAUTIONS - Drug/Laboratory Test Interactions.

Dosage in Patients With Renal Insufficiency

AVAILABILITY OF DOSAGE FORMS

LIPITOR (atorvastatin calcium) is available in dosage strengths of 10 mg, 20 mg, 40 mg and 80 mg atorvastatin per tablet. 1. Friedewald WT. et al. Clin Chem 1972:18(6):489-502.

For a copy of the Product Monograph or full Prescribing Information, please contact:



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Ropinirole (as ropinirole hydrochloride)

TABLETS: 0.25 mg, 1.0 mg, 2.0 mg, 5.0 mg

THERAPEUTIC CLASSIFICATION: AntiParkinsonian Agent / Dopamine Agonist INDICATIONS AND CLINICAL USE: RECUIP® (ropinirole hydrochloride) is indicated in the treatment of the signs and symptoms of idiopathic Parkinson's disease. RECUIP® can be used both as early therapy, without concomitant levodopa and as an adjunct to levodopa. Three year and five year active-comparator controlled clinical trials have been conducted.

CONTRAINDICATIONS: REQUIP* (ropinirole hydrochloride) is contraindicated in patients with a known hypersensitivity to ropinirole hydrochloride or the excipients of the drug product.

WARNINGS: Sudden Onset of Sleep - Patients receiving treatment with REQUIP* (ropinirole hydrochloride), and other dopaminergic agents have reported suddenly falling asleep while engaged in activities of daily living, including operating a motor vehicle, which has sometimes resulted in accidents. Although some of the patients reported somnolence while on REQUIP® others perceived that they had no warning signs, such as excessive drowsiness, and believed that they were alert immediately prior to the event. Physicians should alert patients of the reported cases of sudden onset of sleep. bearing in mind that these events are NOT limited to initiation of therapy Patients should also be advised that sudden onset of sleep has occurred without warning signs. If drowsiness or sudden onset of sleep should occur. patients should immediately contact their physician. Until further information is available on the management of this unpredictable and serious adverse event, patients should be warned not to drive or engage in other activities where impaired alertness could put themselves and others at risk of serious injury or death (e.g., operating machines). Episodes of falling asleep while engaged in activities of daily living have also been reported in patients taking other dopaminergic agents, therefore, symptoms may not be alleviated by substituting these products. Presently, the precise cause of this event is unknown, It is known that many Parkinson's disease patients experience alterations in sleep architecture, which results in excessive daytime sleepiness or spontaneous dozing, and that dopaminergic agents can also induce sleepiness. There is insufficient information to determine whether this event is associated with REQUIP®, all dopaminergic agents or Parkinson's disease itself. Orthostatic Symptoms - Dopamine agonists appear to impair the systemic regulation of blood pressure with resulting orthostatic symptoms of dizziness or lightheadedness, with or without documented hypotension. These symptoms appear to occur especially during dose escalation. Therefore patients treated with dopamine agonists should be carefully monitored for signs and symptoms of orthostatic hypotension, especially during dose escalation (see DOSAGE and ADMINISTRATION) and should be informed of this risk. Hallucinations - Early Therapy: In placebo- controlled trials, REQUIP (ropinirole hydrochloride) caused hallucination in 5.1% of patients during early therapy (1.4% in the placebo group). Hallucination was of sufficient severity that it led to discontinuation in 1.3% of patients. The incidence of hallucination was dose-dependent. In a 5-year study comparing REQUIP® with levodopa in early Parkinson's patients, the overall incidence of hallucinations was 17.3% (31/179) for patients treated with REQUIP® and 5.6% (5/89) for levodopa patients. Hallucinations led to discontinuation of the study treatment in 5.0% of REQUIP* and 2.2% of levodopa patients. In a 3-year study comparing REQUIP® with another dopamine agonist, the overall incidence of hallucinations was 9.5% (16/168) for patients treated with REQUIP® and 9.0% (15/167) for patients receiving active comparator. Hallucinations led to discontinuation of the study treatment in 2.4% of REQUIP® patients and 3.0% of comparator patients. Concomitant Selegiline: In a 5-year study, REQUIP® patients receiving concomitant selegiline reported a higher incidence of hallucinations (23.5%) than did those without (12.2%); this subpopulation effect was not seen in the L-dopa arm (hallucinations with concomitant selegiline = 2.0% vs hallucinations without selegiline = 8.0%). Adjunct Therapy: Hallucinations were experienced by 10.1% of patients receiving REQUIP® and levodopa, compared to 4.2% receiving placebo and levodopa Hallucinations were of sufficient severity that it led to discontinuation in 1.9% of patients. The incidence of hallucinations was dose dependent.

PRECAUTIONS: Cardiovascular - Since REQUIP® (ropinirole hydrochloride) has not been studied in patients with a history or evidence of significant cardiovascular disease including myocardial infarction, unstable angina cardiac decompensation, cardiac arrhythmias, vaso-occlusive disease (including cerebral) or cardiomyopathy, it should be used with caution in such patients. There is limited experience with REQUIP® in patients treated with antihypertensive and antiarrhythmic agents. Consequently, in such patients, the dose of REQUIP® should be titrated with caution. Orthostatic Symptoms -Orthostatic symptoms of dizziness or lightheadedness as well as somnolence may occur during REQUIP® therapy. Neuroleptic Malignant Syndrome - A symptom complex resembling the neuroleptic malignant syndrome (characterized by elevated temperature, muscular rigidity, altered consciousness, and autonomic instability), with no other obvious etiology, has been reported in association with rapid dose reduction, withdrawal of, or changes in anti-Parkinsonian therapy. A single spontaneous report of a symptom complex resembling the neuroleptic malignant syndrome has been observed in a 66 year old diabetic male patient with Parkinson's disease, who developed fever, muscle stiffness, and drowsiness 8 days after beginning REQUIP® treatment. The patient also experienced acute bronchitis, which did not respond to antibiotic treatment. REQUIP® was discontinued three days before the patient died. The reporting physician considered these events to be possibly related to REOLIP® treatment (see DOSAGE AND ADMINISTRATION) A single spontaneous report of severe muscle pain has been reported in a 66 year old male patient around his thigh. The reporting physician considered the event to be probably related to REQUIP® treatment. Retinal Pathology in Rats - In a two year carcinogenicity study in albino Sprague-Dawley rats, retinal atrophy was observed at incidences of 0%, 1.4%, 1.4% and 10% of male rats and 0%, 4.4%, 2.9% and 12.9% of female rats dosed at 0, 1.5, 15 and 50 mg/kg/day respectively. The incidence was significantly higher in both male and female animals dosed at 50 mg/kg/day. The 50 mg/kg/day dose represents a 2.8 fold greater exposure (AUC) and a 13.1 fold greater exposure (Consu) to ropinirole in rats than the exposure would be in humans at the maximum recommended dose of 24 mg/day. The relevance of this finding to humans is not known. Pregnancy - The use of REOUIP® during pregnancy is not recommended. REQUIP® given to pregnant rats during organogenesis (gestation days 8 through 15) resulted in decreased fetal body weight at 60 mg/kg/day (approximately 3-4 times the AUC at the maximal human dose of 8 mg t.i.d.), increased fetal death at 90 mg/kg/day (approximately 5 times the AUC at the maximal human dose of 8 mg t.i.d.) and digital malformations at 150 mg/kg/day (approximately 8-9 times the AUC at the maximal human dose of 8 mg t.i.d.). These effects occurred at maternally toxic doses. There was no indication of an effect on development of the conceptus at a maternally toxic dose of 20 mg/kg/day in the rabbit. In a perinatal-postnatal study in rats. 10 mg/kg/day of REQUIP® (approximately 0.5 - 0.6 times the AUC at the maximal human dose of 8 mg t.i.d.) impaired growth and development of nursing offspring and altered neurological development of female offspring. Nursing Mothers - Since REQUIP® suppresses lactation, it should not be administered to mothers who wish to breast-feed infants. Studies in rats have shown that REQUIP® and/or its metabolites cross the placenta and are excreted in breast milk. Consequently, the human fetus and/or peopate may be exposed to dopamine agonist activity. Use in Women Receiving Estrogen Replacement Therapy- In female patients on long-term treatment with conjugated estrogens, oral clearance was reduced and elimination half-life prolonged compared to patients not receiving estrogens. In patients, already receiving estrogen replacement therapy, REQUIP® may be titrated in the recommended manner according to clinical response. However, if estrogen replacement therapy is stopped or introduced during treatment with REQUIP®, adjustment of the REQUIP® dosage may be required. Pediatric Use - Safety and effectiveness in the pediatric population have not been established. Renal and Hepatic Impairment - No dosage adjustment is needed in patients with mild to moderate renal impairment (creatinine clearance of 30 to 50 mL/min). Because the use of REQUIP® in patients with severe renal impairment or hepatic impairment has not been studied, administration of REQUIP® to such patients is not recommended. Drug Interactions - Psychotropic Drugs: Neuroleptics and other centrally active dopamine antagonists may diminish the effectiveness of REQUIP® Therefore, concomitant use of these products is not recommended. Based on population pharmacokinetic assessment, no interaction was seen between REQUIP® and tricyclic antidepressants or benzodiazepines. Anti-Parkinson Drugs: Based on population pharmacokinetic assessment, there were no interactions between REQUIP® and drugs commonly used to treat Parkinson's disease, i.e., selegiline, amantadine, and anticholinergics. Levodopa: The potential pharmacokinetic interaction of levodopa/ carbidopa (100 mg/10 mg b.i.d.) and REQUIP® (2 mg t.i.d.) was assessed in levodopa naive (de novo) male and female patients with Parkinson's disease (n=30, mean age 64 years). The rate and extent of availability of REQUIP® at steady state were essentially the same with or without levodopa. Similarly, the rate and extent of availability of levodopa, as well as its elimination half-life, were essentially the same in the presence and absence of REQUIP®. Inhibitors of CYP1A2: Ciprofloxacin: The effect of ciprofloxacin (500 mg b.i.d.) on the pharmacokinetics of REQUIP® (2 mg t.i.d.) was studied in male and female patients with Parkinson's disease (n=12. mean age 55 years). The extent of systemic availability of REQUIP® was significantly increased when coadministered with ciprofloxacin (AUC increased by 1.84 fold). Thus, in patients already receiving CYP1A2 inhibitors such as ciprofloxacin, REQUIP® therapy may be instituted in the recommended manner and the dose titrated according to clinical response. However, if therapy with a drug known to be an inhibitor of CYP1A2 is stopped or introduced during treatment with REQUIP®, adjustment of the REQUIP® dosage will be required Substrates of CYP1A2: Theophylline: The effect of oral theophylline (300 mg b.i.d.) on the pharmacokinetics of REQUIP® (2 mg t.i.d.) was studied in male and female patients with Parkinson's disease (n=12, mean age 59 years). There was no marked change in the rate or extent of availability of REQUIP® when coadministered with theophylline. Similarly, coadministration of REQUIP® with intravenous theophylline (5 mg/kg) did not result in any marked change in the pharmacokinetics of theophylline. It is therefore unlikely that substrates of CYP1A2 would significantly after the pharmacokinetics of REQUIP®, and vice-versa, **Digoxin**: The effect of REQUIP® (2 mg t.i.d.) on the pharmacokinetics of digoxin (0.125-0.25 mg o.d.) was studied in male and female patients with Parkinson's disease (n=10, mean age 72 years). Coadministration at steady state with REQUIP® resulted in a 10% decrease in digoxin AUC although mean trough digoxin plasma concentrations were unaltered. However, the effect of higher recommended doses of REQUIP® on the pharmacokinetics of digoxin is not known. Alcohol: No information is available on the potential for interaction between REQUIP® and alcohol. As with other centrally active medications, patients should be cautioned against taking REQUIP® with alcohol. Psycho-Motor Performance - (see WARNINGS Sudden Onset of Sleep)

ADVERSE REACTIONS: Adverse Reactions Associated with Discontinuation of Treatment – Of 1599 patients who received REQUIP® (ropinirole hydrochloride) during the premarketing clinical trials, 17.1% in

early-therapy studies and 17.3% in adjunct-therapy studies discontinued treatment due to adverse reactions. The events resulting in discontinuation of RECUIP® in 1% or more of patients were as follows: Early therapy: nausea (6.4%), dizziness (3.8%), aggravated Parkinson's disease (1.3%), hallucination (1.3%), headache (1.3%), somnolence (1.3%) and vomiting (1.3%). Adjunct therapy: dizziness (2.9%), dyskinesia (2.4%), confusion (2.4%), vomiting (2.4%), hallucination (1.9%), nausea (1.9%), anxiety (1.9%), and increased sweating (1.4%). Patients over 75 years of age (n=130) showed slightly higher incidences of withdrawal due to hallucination, confusion and dizziness than patients less than 75 years of age. Most Frequent Adverse Events - Adverse events occurring with an incidence of greater than, or equal to, 10% were as follows: Early therapy: nausea, dizziness, somnolence, headache, peripheral edema, vomiting, syncope, fatigue and viral infection, Adiunct therapy: dyskinesia, nausea, dizziness, somnolence and headache. Donamine agonists with an ergoline chemical structure have been associated with adverse experiences such as retroperitoneal fibrosis, erythromelalgia and pulmonary reactions. REQUIP® has a novel, non-ergoline chemical structure and no reports of such events have been observed in clinical trials. Incidence of Adverse Events in Placebo Controlled Trials - The incidence of postural hypotension, an event commonly associated with initiation of dopamine agonist therapy, was not notably different from placebo in clinical trials. However, decreases in systolic blood pressure to < 90 mmHg have been observed in 13% (<65 years), 16% (65 - 75 years) and 7.6% (>75 years) of patients treated with REQUIP®. Table 2 lists adverse events that occurred at an incidence of 1% or more among REQUIP®-treated patients who participated in placebo-controlled trials for up to one year. Patients were dosed in a range of 0.75 mg to 24 mg/day. Reported adverse events were classified using a standard World Health Organization (WHO)-based dictionary terminology. The prescriber should be aware that these figures can not be used to predict the incidence of adverse events in the course of usual medical practice where patient characteristics and other factors differ from those which prevailed in the clinical trials. Similarly, the cited frequencies can not be compared with figures obtained from other clinical investigations involving different treatments, uses and investigators. The cited figures, however, do provide the prescribing physician with some basis for estimating the relative contribution of drug and non-drug factors to the adverse events incidence rate in the population studied

Adverse events with	incidence :	LE 2 ≥1% from all pla	cebo-controlled		
early and adjunct therapy studies Early Therapy Adjunct Therapy					
	REQUIP®	Placebo	REQUIP*	Placebo	
	N = 157 occurrence	N = 147 % occurrence	N = 208 % occurrence	N = 120 % occurrence	
Autonomic Nervous System	NO CONTENDE	A GCCATCHOL	/# OCCUPTORIOS	/ OCCUPTION	
Sweating Increased	6.4	4.1	7.2	1.7	
Mouth Dry	5.1	3.4	5.3	0.8	
Flushing	3.2	0.7	1.4	0.8	
Body as a Whole General					
Peripheral Edema Fatique	13.4	4.1 4.1	3.9	2.5	
Injury	10.0	4.1	10.6	9.2	
Pain	7.6	4.1	5.3	3.3	
Asthenia	6.4	1.4	-	-	
Drug Level Increased	4.5	2.7	6.7	3.3	
Chest Pain	3.8	2.0	-	-	
Malaise	3.2	0.7	1.4	8.0	
Therapeutic Response	1.0	0.7			
Decreased Cellulitis	1.9	0.7	-	-	
Cellulitis Influenza-like Symptoms	1.3	0.0	1.0	0.0	
innuenza-like Symptoms Fever		_	1.4	0.0	
Cardiovascular General			1.1	0.0	
Syncope	11.5	1.4	2.9	1.7	
Hypotension Postural	6.4	4.8	-	-	
Hypertension	4.5	3.4	3.4	3.3	
Hypotension	1.9	0.0	2.4	0.8	
Cardiac Failure	-	-	1.0	0.0	
Central and Peripheral Nervo					
Dizziness Dyskinesia	40.1	21.8	26.0 33.7	15.8 12.5	
Headache	17.2	17.0	16.8	11.7	
Ataxia (Falls)	-	- 17.0	9.6	6.7	
Tremor	_		6.3	2.5	
Paresthesia	-	_	5.3	2.5	
Hyperesthesia	3.8	2.0	-	-	
Dystonia	-	-	4.3	4.2	
Hypokinesia	-	-	5.3	4.2	
Paresis	_	-	2.9	0.0	
Speech Disorder	-	-	1.0	0.0	
Vertigo Carpal Tunnel Syndrome	1.9	0.0 0.7	_	-	
Gastrointestinal System	1.3	0.7		~	
Nausea	59.9	21.8	29.8	18.3	
Vomiting	12.1	6.8	7.2	4.2	
Dyspepsia	9.6	4.8	-	-	
Constipation	8.3	7.5	5.8	3.3	
Abdominal Pain	6.4	2.7	8.7	7.5	
Diarrhea	-	-	4.8	2.5	
Anorexia	3.8	1.4	-	-	
Flatulence	2.5	1.4	1.9	0.8	
Tooth Disorder Saliva Increased	1.9	0.7	1.0	0.8	
Saliva increased Colitis	1.3	0.0	2.4	0.8	
Dysphagia	1.3	0.0	2.4	0.8	
Periodontitis	1.3	0.0	1.4	0.8	
Eructation	-	1-1	1.4	0.0	
Fecal Incontinence	$(1-\epsilon)^{-1}$	-	1.0	0.0	
Hemorrhoids	-	-	1.0	0.0	
Gastroesophageal Reflux	-	-	1.0	0.0	
Gastrointestinal Disorder (NOS)	-	-	1.0	0.0	
Tooth Ache Hearing and Vestibular	-	-	1.0	0.0	
Hearing and Vestibular Tinnitus	1.3	0.0	-	_	
Heart Rate and Rhythm	1.0	0.0			
Palpitation	3.2	2.0	2.9	2.5	

SKULL-BASE NEUROSURGEON



The **Department of Clinical Neurosciences and the Calgary Health Region** invite applications for a full-time academic position as a neurosurgeon with special expertise in skull-base surgery at the Assistant Professor level or higher to be responsible for teaching, research and patient care.

Salary support and start-up funding will be available through successful application to the Alberta Heritage Foundation for Medical Research (AHFMR) and/or the Canadian Institutes of Health Research (CIHR) and support through an Alternate Relationship Plan in Neurosurgery supported by Alberta Health and Wellness, Calgary Health Region and the Faculty of Medicine.

The Department of Clinical Neurosciences is a progressive clinical and academic Department in the Faculty of Medicine, closely affiliated with the Hotchkiss Brain Institute. Calgary is a vibrant multicultural city situated in close proximity to the Rocky Mountains, Banff National Park and Lake Louise.

Qualifications include an MD, or equivalent, a FRCS or equivalent in Neurosurgery and eligibility for licensure in the Province of Alberta. Clinical fellowship training in skull-base surgery is required in addition to a track record of academic excellence in basic or clinical research, related to pathology that affects the brain. Desired areas include, but are not restricted to, expertise in neurooncology, neuroendoscopy, radiosurgery and skull-base anatomy. Preference will be given to individuals with one or more of these skill-sets and to those competitive for grant support from AHFMR and CIHR.

Please forward curriculum vitae, statement of research interests and the names of three referees by August 31, 2006 to:

Dr. Rajiv Midha, Head

Division of Neurosurgery Department of Clinical Neurosciences University of Calgary 1403 – 29 Street N.W. Calgary, AB, Canada T2N 2T9

In accordance with Canadian Immigration requirements, priority will be given to Canadian citizens and permanent residents of Canada. The University of Calgary respects, appreciates and bonours diversity.

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Not just rewarding work - but the kind of rewards you've been working toward!



Reward yourself by settling in the place
National Geographic calls "One of the
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Known as mother nature's recreational
playground, you can ski perfect powder,
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paths, or ride the trails of the Mountain
Bike Capital of Canada. Our homes are
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Life here is simply...more rewarding.

At the Kootenay Boundary Regional Hospital in Trail, British Columbia, you can create the rewarding practice you are looking for – and the lifestyle rewards you have been working toward.

Serving a catchment area of 85,000, you will provide a broad range of neurological services with access to all required diagnostics and support from a comprehensive team of specialists.

Neurologists at Kootenay Boundary Regional Hospital qualify for recruitment and retention incentives, receive one of the highest levels of remuneration for health professionals in Canada and qualify for BC's highly competitive on-call payment program.

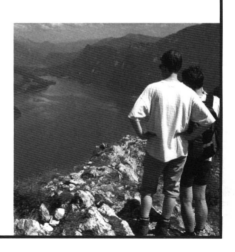
Don't wait. Call us.

Dr. Bob StreichDirector, Medical Services

Kootenay Boundary robert.streich@interiorhealth.ca Tel 250 304-2888 ext 230









KING FAISAL SPECIALIST HOSPITAL AND RESEARCH CENTRE

The King Faisal Specialist Hospital and Research Centre – Riyadh is JCIA accredited and operates 865 beds (159 day-care beds) with approximately 8,000 employees. Located in the capital of Saudi Arabia, the hospital provides a full range of tertiary, secondary and primary health care programs at its main campus facility, as well as the King Fahad National Children's Cancer Centre. The KFSHRC is the national referral center for Oncology; Organ Transplantation; Cardiovascular Diseases; and, Genetic Diseases. Affiliated with the King Saud University (Faculty of Medicine), the hospital's postgraduate education programs support both Residency and Fellowship training.

DEPARTMENT OF NEUROSCIENCES

The Department of Neurosciences provides tertiary care in Neurosurgery, Neurology, Pediatric Neurology and Psychiatry. There are five Consultant Neurosurgeons, five Pediatric Neurologists, three Psychiatrists, seven Neurologists, two Clinical Neurophysiologists and two Clinical Neuropsychologists. Residency training programs in Neurology, Neurosurgery and Psychiatry and Fellowship Programs in Pediatric Neurology and Pediatric Neurosurgery are offered. We have approximately 25 residents and fellows. Annually, 1500 admissions, 800 operations and 21000 outpatient visits occur. The Department has a Comprehensive Epilepsy and Movement Disorder Program. Numerous subspecialty clinics are in place. Neurosurgical theaters have modern equipment and navigational system. The Department is committed to research with approximately 20 papers are published annually.

The King Faisal Specialist Hospital and Research Centre is seeking a Chairman for the Department of Neurosciences.

The successful candidate will assume the overall responsibility for the quality of patient care within the Department of Neurosciences and for all professional and administrative activities of the Department that has approximately 25 consultants, 25 residents, and seven administrative staff.

The applicants must have graduated from an accredited medical school, completed training in a recognized training program and be certified by the appropriate authority in the country where the candidate was trained.

It is expected that the successful candidate will have a minimum of 10 years of clinical experience in specialty or subspecialty activity including 3 years experience in a responsible administrative capacity. Academic, research and teaching background are essential.

The deadline for submission of applications is 60 days from date of publication. Please direct your electronic responses to:

E-Mail: anasser@kfshrc.edu.sa

Abdulaziz Al Nasser, MD, CPE

Co-Chairman, Medical Chairman Search Committee King Faisal Specialist Hospital & Research Centre – Riyadh Post Office Box 3354 (MBC62), Riyadh, 11211, Saudi Arabia

Telephone: 00966 1 4427395 or 4647272 ext 31868
Fax: 00966 1 4427397 or 4423055
All responses to King Faisal Specialist Hospital and Research

Centre – Riyadh are confidential. For more information about the King Faisal Specialist Hospital and Research Centre – Riyadh, please visit our web site at www.kfshrc.edu.sa.

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Employment Opportunity for Neuro-Ophthalmologist

The Division of Ophthalmology, McMaster University, Hamilton, Ontario is currently seeking a neuroophthalmologist or ophtho-neurologist. The successful candidate will participate in teaching (undergraduate and graduate levels), research (clinical or basic science) and will carry out a clinical practice in a tertiary university setting. Successful candidates will be fellowship trained. Salaries at McMaster University are competitive and will be commensurate with the seniority and experience of the candidate. McMaster University is committed to employment equity and encourages applications from all qualified candidates, including aboriginal peoples, persons with disabilities, members of visible minorities, and women. In accordance with Canadian immigration requirements this advertisement is directed to Canadian citizens and permanent residents.

Please reply to:
John Harvey MD, FRCSC
Professor & Head, Division of Ophthalmology
McMaster University Medical Centre, Room 4V2
1200 Main Street West, Hamilton, ON L8N 3Z5
905-521-2100 x76662
jtharvey@mcmaster.ca

NEUROLOGY - NEW YORK



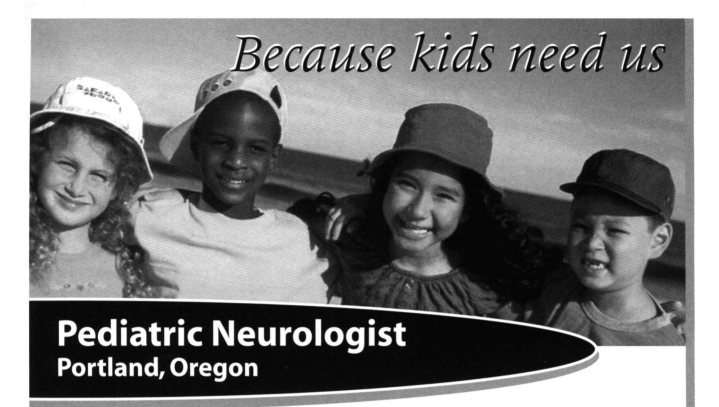
The Guthrie Clinic in Corning, New York is the clinic's largest regional office. There are 60 physicians practicing at two locations in Corning. The clinic is seeking a general neurologist with any subspecialty interest to join another neurologist in practice. Salary guarantee plus full benefits package.

Rated one of the top 50 communities in the country Corning offers a rich tapestry of cultural and recreational opportunities in the beautiful Finger Lakes Region of upstate New York.

Contact Ann Lamb 800-678-7858, x63486 314-726-0026 (fax) alamb@cejkasearch.com

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Legacy Health System is seeking a BE/BC Pediatric Neurologist to join a growing Children's Neurology Program at Legacy Emanuel Children's Hospital. Services are provided by one employed Pediatric Neurologist, as well as one private practice Pediatric Neurologist. The practice consists of inpatient consultation to PICU, NICU and Acute Pediatrics, outpatient consultations, and EEG interpretation without focus on attention deficient disorder. Collaborative interdisciplinary care with other pediatric specialists such as physiatrists, neurosurgeons, intensivists and hospitalists is common. Call is 1 in 3 covering weekdays and 1 in 7 for weekend call.

Legacy Children's Hospital is a 155-bed facility that includes an 18-bed PICU and a 45-bed Level III Neonatal ICU. The hospital has over 90 board-certified pediatric sub-specialists providing care in cardiology, endocrinology, gastroenterology, genetics, infectious disease, intensive care, neonatology, nephrology, neurology, neurosurgery, orthopedics, otolaryngology, pulmonology, pediatric surgery, and urology.

Legacy Health System is a five-hospital system based in Portland, Oregon. It is one of the largest Oregon-based health systems in the state, providing a comprehensive array of specialty services including: a Level I trauma center, an advanced minimally invasive surgery program, a large cancer program, The Oregon Burn Center, one of the largest heart surgery programs in Oregon, and complete women's and children's services. Legacy has also invested heavily in research and education, with over a dozen associated residency programs, plus a freestanding Clinical Research & Technology Center, which supports numerous clinical trials. Academic affiliations are available with Oregon Health Sciences University.

Candidates should be Board Certified or Board Eligible with an interest in general pediatric neurology and a strong interest in Epilespy. Candidates should have an interest in both rare and common neurological conditions and have strong EEG reading capability.

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Legacy is committed to delivering comprehensive and family-centered care with compassion and excellence. We offer a competitive salary and benefits plan, and relocation assistance. Candidates should contact Vicki Owen, Sr. Recruitment Consultant, Legacy Employment Services at 1 (866) 888-4428, ext. 6. Applications are required and can be accessed at our website, www.legacyhealth.org. Please reference position number 091914. In addition, please email your CV to vowen@lhs.org. AA/EOE

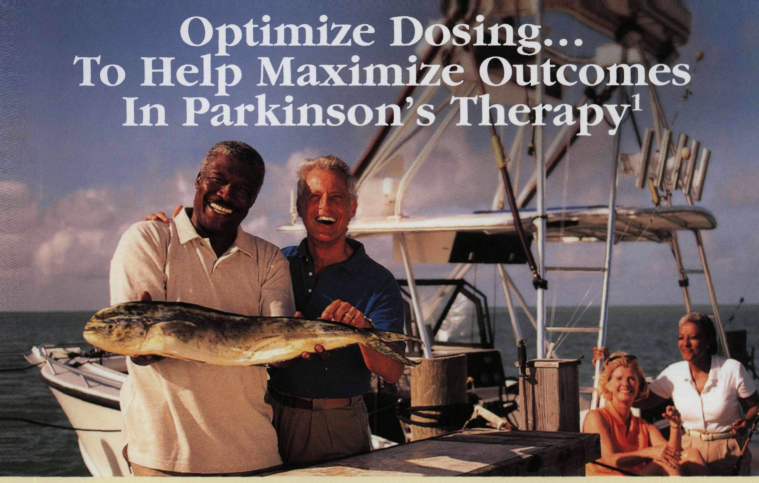
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Titrate to help maximize patient benefit. In at least 75% of the patients who responded to REQUIP®, doses of up to 9 mg/day were necessary to ensure a first therapeutic response.19

Three Reasons to Prescribe REQUIP®

REQUIP® delayed the use of L-dopa

34% (n=29 of 85) of REQUIP® monotherapy patients completed the entire 5-year study without requiring L-dopa supplementation²⁰

Low risk of dyskinesia

Only 5% of REQUIP® monotherapy patients developed dyskinesia compared with 36% of L-dopa patients^{2*}

Low supplementary dose of L-dopa needed

When used with adjunct L-dopa, REQUIP® patients required an average of 43% less L-dopa (427 ± 221 mg) than patients on L-dopa alone $(753 \pm 398 \text{ mg})^2$

- In early treatment of Parkinson's disease over the course of a 5-year multicentre, prospective, double-blind, flexible-dose study, with 268 patients randomized to either REQUIP® (n=179) or L-dopa and benserazide (a decarboxylase inhibitor) (n=89). Open label L-dopa was available as supplementary medication. 2,3 p<0.001
- * Prior to supplementation with L-dopa
- x Data from 3 large phase III double-blind trials of ropinirole monotherapy in early Parkinson's disease were examined: a 5-year L-dopa-controlled trial (n=179), a 3-year bromocriptine-controlled trial (n=168), both with planned interim analysis and a 6-month placebo-controlled
- † Please consult the Warnings section of the Product Monograph.³
- ® REQUIP is a registered trademark, used under license by GlaxoSmthKline Inc.

References: 1. Korczyn AD et al. Dosing with ropinirole in a clinical setting. Acta Neurologica Scandinavica 2002; 106:200-204. 2. Rascol O et al. A five-year study of the incidence of dyskinesia in patients with early Parkinson's disease who were treated with ropinirole or levodopa. N Eng J Med 2000;342(20):1484-1491. 3. Product Monograph of REQUIP® (ropinirole hydrochloride), GlaxoSmithKline, March 2004.

REQUIP® (ropinirole hydrochloride) is indicated in the treatment of the signs and symptoms of idiopathic Parkinson's disease.

REQUIP® can be used both as early therapy, without concomitant levodopa and as an adjunct to levodopa. Patients receiving treatment with REQUIP® and other dopaminergic agents have reported the sudden onset of sleep while engaged in daily activities. Patients should be warned not to drive or engage in other activities where impaired alertness could put themselves or others at risk.^{3†}

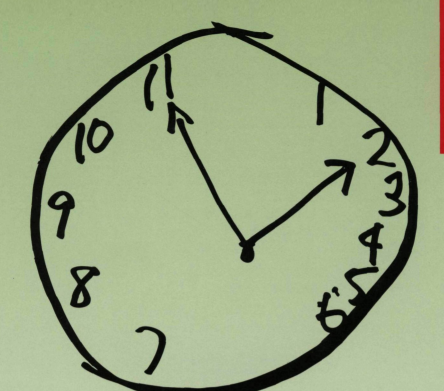
Adverse events occurring with an incidence of greater than, or equal to, 10% were as follows: Early therapy: nausea, dizziness, somnolence, headache, peripheral edema, vomiting, syncope, fatigue and viral infection. Adjunct therapy: dyskinesia, nausea, dizziness, somnolence and headache. REQUIP® is contraindicated in patients with a known hypersensitivity to ropinirole hydrochloride or the excipients of the drug product.











Once-a-Day REMINYL ER

Take the Time to Look at REMINYL* ER.

Consider once-a-day REMINYL* ER as initial treatment in AD.1

"REMINYL and **"REMINYL ER** (galantamine hydrobromide) are indicated for the symptomatic treatment of patients with mild to moderate dementia of the Alzheimer's type. REMINYL ER has not been studied in controlled clinical trials for longer than 6 months.

The most common side effects for REMINYL ER (vs. placebo) in a clinical trial were nausea (17% vs. 5%), dizziness (10% vs. 4%), injury (8% vs. 6%) and headache (8% vs. 6%). For patients who experienced adverse events, the majority occurred during the dose-escalation phase.

There is no evidence that galantamine alters the course of the underlying dementing process.

† Data does not support an indication for either vascular dementia (VaD) or Alzheimer's disease (AD) and concomitant cerebrovascular disease (AD+CVD).

In patients with moderately impaired hepatic function (Child-Pugh score of 7-9), based on pharmacokinetic modelling, dosing with REMINYL should begin with 4 mg once daily for at least 1 week. For REMINYL ER, based on pharmacokinetic modelling, dosing should begin with 8 mg every other day for at least 1 week. Then the dosage should be increased to 4 mg twice a day for REMINYL or 8 mg once daily for REMINYL ER for at least 4 weeks. In these patients, daily doses should not exceed 16 mg/day. REMINYL and REMINYL ER are not recommended in patients with severe hepatic impairment (Child-Pugh score of 10-15).

In patients with renal impairment (creatinine clearance of 9-60 mL/min), dose escalation should proceed cautiously and the maintenance dose should generally not exceed 16 mg/day. REMINYL and REMINYL ER are not recommended in patients with creatinine clearance of less than 9 mL/min.

Dose reductions can be considered in patients treated with potent CYP2D6 or CYP3A4 inhibitors.

REFERENCE: 1. REMINYL* (galantamine hydrobromide tablets), REMINYL* ER (galantamine hydrobromide extended-release capsules) Product Monograph, JANSSEN-ORTHO Inc., September 29, 2005.

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ROJA053700E



AD with Cerebrovascular Disease and VaD data for REMINYL now included in Product Monograph.†

