NAME OF THE MEDICINE

Active ingredient : Carbamazepine
Chemical name : 5H-dibenz(b,f)azepine-5-carboxamide
Structural formula :

![Structural formula](image)

Molecular formula : $C_{15}H_{12}N_{2}O$  
Molecular weight : 236.3
CAS Registry no. : 298-46-4

DESCRIPTION

Carbamazepine is a white or yellowish-white, crystalline powder; almost odourless; tasteless or with a slightly bitter taste. It has a melting point of 189°-193°C. Carbamazepine is practically insoluble in water and in ether, sparingly soluble in ethanol (96%), and soluble in 10 parts of chloroform.

Each Teril tablet contains 200 mg of carbamazepine and the following inactive excipients: microcrystalline cellulose, pregelatinised maize starch, colloidal anhydrous silica, purified talc, sodium starch glycollate and magnesium stearate.

PHARMACOLOGY

Pharmacodynamics

Pharmacotherapeutic groups: antiepileptic, neurotropic and psychotropic agent.

ATC code: N03 AF01

Antiepileptic agent

Carbamazepine as an antiepileptic agent has been shown to be effective in the treatment of partial seizures (simple and complex) with and without secondary generalisation, generalised tonic-clonic seizures and combinations of these seizure types.

In some clinical studies carbamazepine, given as monotherapy to patients with epilepsy, including children and adolescents, has been reported to exert a mild psychotropic action, including a beneficial effect on attentiveness and cognitive performance and on symptoms of anxiety and depression, as well as a decrease in irritability and aggressiveness. Other studies have not confirmed these findings.
Neurotropic agent

As a neurotropic agent, carbamazepine is clinically effective in relieving paroxysmal attacks of pain in idiopathic trigeminal neuralgia.

Psychotropic agent

As a psychotropic agent, carbamazepine has shown clinical efficacy as treatment for mania as well as for the maintenance treatment of bipolar affective disorders, when given either as monotherapy or in combination with neuroleptics, antidepressants or lithium.

Mechanism of action

The mechanism of action of carbamazepine has been partially elucidated. Carbamazepine stabilises hyperexcited nerve membranes, inhibits repetitive neuronal discharges and reduces synaptic propagation of excitatory impulses. It is conceivable that prevention of repetitive firing of sodium-dependent action potentials in depolarised neurons via use- and voltage-dependent blockade of sodium channels may be its main mechanism of action.

Whereas reduction of glutamate release and stabilisation of neuronal membranes may account mainly for the antiepileptic effects, it is speculated that the depressant effect on dopamine and noradrenaline turnover could be responsible for the antimanic properties of carbamazepine.

Carbamazepine possesses anticholinergic and antidiuretic activity and may suppress ventricular automaticity through its membrane depressant effect.

Pharmacokinetics

Absorption

Absorption from the gastrointestinal tract is relatively slow yet almost complete.

Plasma concentrations.

When taken as a single oral dose, a peak concentration of unchanged carbamazepine is reached within 4 to 24 hours (majority within 12 hours). During one study, the peak following the oral administration of 400 mg was approximately 4.5 microgram/mL.

Steady state plasma concentrations of carbamazepine are attained within about 1 to 2 weeks, depending individually upon auto-induction by carbamazepine and hetero-induction by other enzyme-inducing drugs, as well as on pre-treatment status, dosage and duration of treatment. The therapeutic range at steady state is 4 to 12 microgram/mL (or 17 to 50 micromol/L) carbamazepine. The main metabolite, carbamazepine -10, 11-epoxide, possesses anticonvulsant activity and reaches concentrations approximately 30% of those of carbamazepine.

Bioavailability.

Absolute bioavailability could not be determined, as an intravenous formulation was not developed. Nevertheless, it appears that systemic availability approaches 100% and is unaffected by food.

Serum protein binding.

70 to 80%. The concentration of unchanged substance in saliva and cerebrospinal fluid (CSF) reflects the non-protein bound fraction present in plasma.

Distribution

The concentration of unchanged drug in the CSF and saliva is approximately 20 to 30% of that attained in plasma. Milk concentration ranges from 25 to 60% of the plasma concentration. Carbamazepine readily crosses the placenta. The apparent volume of distribution was found to be 0.8 to 1.9 L/kg.
Metabolism

Carbamazepine is metabolised in the liver via the epoxide-diol pathway, the main metabolite (carbamazepine-10, 11-epoxide) being pharmacologically active. Cytochrome P450 3A4 has been identified as the major isoform responsible for the formation of carbamazepine-10, 11-epoxide. Human microsomal epoxide hydrolase has been identified as the enzyme responsible for the formation of the 10,11-transdiol derivative from carbamazepine-10,11 epoxide. Carbamazepine is capable of inducing its own metabolism by the hepatic mono-oxygenase system.

Elimination

The elimination half-life of unchanged carbamazepine following a single oral dose averaged 36 hours whereas, after repeated administration which leads to hepatic enzyme induction, it averaged 16 to 24 hours, depending on the duration of treatment. In patients receiving concomitant treatment with other liver-enzyme inducing drugs (e.g. phenytoin, phenobarbitone), half-life values averaging 9 to 10 hours have been found. The mean elimination half-life of the 10, 11-epoxide metabolite in the plasma is about 6 hours following single oral doses of the epoxide itself.

Excretion:

Following a single 400 mg dose, 72% was excreted in the urine mainly in the form of epoxidated, hydroxylated and conjugated metabolites. Some 28% of the dose was excreted in the faeces.

Note. Pharmacokinetics were not altered in the elderly. There are no data on patients with impaired hepatic or renal function.

INDICATIONS

Epilepsy

- Complex or simple partial seizures (with or without loss of consciousness), with or without secondary generalisation
- Generalised tonic-clonic seizures
- Mixed seizure patterns incorporating the above.

Teril is suitable for monotherapy and combination therapy. Teril is usually not effective in absence seizures, atonic seizures and myoclonic seizures and should not be used for status epilepticus (see Precautions).

Trigeminal neuralgia

For relief of pain in idiopathic trigeminal neuralgia and trigeminal neuralgia due to multiple sclerosis; and in idiopathic glossopharyngeal neuralgia. (Carbamazepine is not a simple analgesic and is not intended for trivial facial pain or headache).

Mania and bipolar affective disorders

Treatment of mania and maintenance treatment of bipolar affective disorders to prevent or attenuate recurrence.

CONTRAINDICATIONS

- Known hypersensitivity to carbamazepine or structurally related drugs (e.g. tricyclic antidepressants) or to any other component of the formulation
- Atrioventricular block
• Systemic lupus erythematosus

• History of hepatic porphyrias (e.g. acute intermittent porphyria, variegate porphyria, porphyria cutanea tarda)

• History of bone marrow depression

• Because it is structurally related to tricyclic antidepressants, the use of Teril is contraindicated in combination with monoamine oxidase inhibitors (MAOIs). Before administering Teril, MAOIs should be discontinued for a minimum of 2 weeks, or longer if the clinical situation permits.

• Hepatic failure (as metabolism occurs in the liver, it is recommended that carbamazepine not be given to patients with significant hepatic dysfunction).

PRECAUTIONS

Teril should be given only under medical supervision.

Teril should be prescribed only after a critical benefit-risk appraisal and under close monitoring in patients with a history of cardiac, hepatic or renal damage, adverse haematological reactions to other drugs, or interrupted courses of therapy with Teril.

Serious dermatological reactions

Serious dermatologic reactions, including toxic epidermal necrolysis (TEN: also known as Lyell’s syndrome) and Stevens-Johnson syndrome (SJS), have been reported very rarely with carbamazepine. Patients with serious dermatological reactions may require hospitalization, as these conditions may be life-threatening and may be fatal. Most of the SJS/TEN cases appear in the first few months of treatment with Teril. If signs and symptoms suggestive of severe skin reactions (e.g. SJS/TEN) appear, Teril should be withdrawn at once and alternative therapy should be considered.

Pharmacogenomics

There is growing evidence of the role of different HLA alleles in predisposing patients to immune-mediated adverse reactions.

Association with HLA-A*3101

Human Leukocyte Antigen (HLA)-A*3101 may be a risk factor for the development of hypersensitivity syndrome and cutaneous adverse drug reactions such as SJS, TEN, DRESS, AGEP and maculopapular rash. Retrospective genome-wide studies in Japanese and Northern European populations reported association between severe skin reactions (SJS, TEN, DRESS, AGEP and maculopapular rash) associated with carbamazepine use and the presence of the HLA-A*3101 allele in these patients.

Testing for the presence of HLA-A*3101 allele should be considered in patients with ancestry in genetically at-risk populations (for example, patients of the Japanese and Caucasian populations, patients who belong to the indigenous populations of the Americas, Hispanic populations, people of southern India, and people of Arabic descent), prior to initiating treatment with carbamazepine. The use of carbamazepine should be avoided in patients who are found to be positive for HLA-A*3101, unless the benefits clearly outweigh the risks. Screening is generally not recommended for any current carbamazepine users, as the risk of SJS/TEN, AGEP, DRESS and maculopapular rash is largely confined to the first few months of therapy, regardless of HLA-A*3101 status.

Association with HLA-B*1502

Retrospective studies in patients of Han Chinese and Thai origin found a strong correlation between SJS/TEN skin reactions associated with carbamazepine and the presence in these patients of the Human Leukocyte Antigene (HLA)-B*1502 allele. Higher reporting rates of SJS (rare rather than very rare) are reported in some
countries in Asia (e.g. Taiwan, Malaysia and the Philippines) in which there is a higher prevalence of the HLA-B*1502 allele in the population. The prevalence of carriers of this allele in Asian populations is above 15% in the Philippines, Thailand, Hong Kong and Malaysia, around 10% in Taiwan, around 4% in North China, around 2 to 4% in South Asia including Indians, and less than 1% in Japan and Korea. The prevalence of the HLA-B*1502 allele is negligible in Caucasian, African, indigenous peoples of the Americas, and Hispanic populations sampled.

Testing for the presence of HLA-B*1502 allele should be considered in patients with ancestry in genetically at-risk populations, prior to initiating treatment with carbamazepine. If testing for the presence of the HLA-B*1502 allele should be performed, high-resolution “HLA-B*1502 genotyping” is recommended. The test is positive if either one or two HLA-B*1502 alleles are detected and negative if no HLA-B*1502 alleles are detected.

The use of carbamazepine should be avoided in tested patients who are found to be positive for HLA-B*1502 unless the benefits clearly outweigh the risks. Screening is not generally recommended in patients from populations in which the prevalence of HLA-B*1502 is low. Screening is generally not recommended for any current carbamazepine users, as the risk of SJS/TEN is largely confined to the first few months of therapy, regardless of HLA-B*1502 status.

The identification of subjects carrying the HLA-B*1502 allele and the avoidance of carbamazepine therapy in these subjects has been shown to decrease the incidence of carbamazepine-induced SJS/TEN.

Limitation of genetic screening

Genetic screening results must never substitute for appropriate clinical vigilance and patient management. Many Asian patients positive for HLA-B*1502 and treated with carbamazepine will not develop SJS/TEN and patients negative for HLA-B*1502 of any ethnicity can still develop SJS/TEN. Similarly many patients positive for HLA-A*3101 and treated with carbamazepine will not develop SJS, TEN, DRESS, AGEP or maculopapular rash and patients negative for HLA-A*3101 of any ethnicity can still develop these severe cutaneous adverse reactions. The role of other possible factors in the development of, and morbidity from these severe cutaneous adverse reactions, such as AED dose, compliance, concomitant medications, co-morbidities, and the level of dermatologic monitoring have not been studied.

Other dermatologic reactions

Mild skin reactions (e.g. isolated macular or maculopapular exanthemata) can also occur and are mostly transient and not hazardous, and they usually disappear within a few days or weeks, either during the course of treatment or following a decrease in dosage. However, since it may be difficult to differentiate the early signs of more serious skin reactions from mild transient reactions, the patient should be kept under close surveillance with consideration given to immediately withdrawing the drug should the reaction worsen with continued use.

The HLA-A*3101 allele has been found to be associated with less severe adverse cutaneous reactions from carbamazepine and may predict the risk of these reactions from carbamazepine, such as anticonvulsant hypersensitivity syndrome or non-serious rash (maculopapular eruption). However, the HLA-B*1502 allele has not been found to predict the risk of less severe adverse cutaneous reactions from carbamazepine, such as anticonvulsant hypersensitivity syndrome or non-serious rash (maculopapular eruption).

Hypersensitivity

Teril may trigger hypersensitivity reactions, including Drug Rash with Eosinophilia and Systemic Symptoms (DRESS), a delayed multi-organ hypersensitivity disorder with fever, rash, vasculitis, lymphadenopathy, pseudo lymphoma, arthralgia, leukopenia, eosinophilia, hepato-splenomegaly, abnormal liver function tests and vanishing bile duct syndrome (destruction and disappearance of intrahepatic bile ducts), that may occur in various combinations. Other organs may also be affected (e.g. lungs, kidneys, pancreas, myocardium, colon) (see ADVERSE EFFECTS).

Patients who have exhibited hypersensitivity reactions to carbamazepine should be informed that approximately 25 to 30% of these patients may experience hypersensitivity reactions with oxcarbazepine (Trileptal®). Cross-hypersensitivity can occur between carbamazepine and phenytoin (see INTERACTIONS).
In general, if signs and symptoms suggestive of hypersensitivity reactions occur, Teril should be withdrawn immediately.

**Seizures**

Teril should be used with caution in patients with mixed seizures which include absences, either typical or atypical. In all of these conditions, carbamazepine may exacerbate seizures. In case of exacerbation of seizures, Teril should be discontinued.

**Special Populations**

**Hepatic Function**

Baseline and periodic evaluations of hepatic function, particularly in patients with a history of liver disease and in elderly patients, must be performed during treatment with Teril. The drug should be withdrawn immediately in cases of aggravated liver dysfunction or active liver disease.

**Renal Function**

Baseline and periodic complete urinalysis and BUN determinations are recommended.

**Geriatrics**

Due to drug interactions and different antiepileptic drug pharmacokinetics, the dosage of Teril should be selected with caution in elderly patients.

**Hyponatremia**

Hyponatremia is known to occur with carbamazepine. In patients with pre-existing renal conditions associated with low sodium or in patients treated concomitantly with sodium lowering medicinal products (e.g. diuretics, medicinal products associated with inappropriate ADH secretion), serum sodium levels should be measured prior to initiating carbamazepine therapy. Thereafter, serum sodium levels should be measured after approximately two weeks and then at monthly intervals for the first three months during therapy, or according to clinical need. These risk factors may apply especially to elderly patients.

**Hypothyroidism**

Carbamazepine may reduce serum concentrations of thyroid hormones through enzyme induction requiring an increase in dose of thyroid replacement therapy in patients with hypothyroidism. Hence thyroid function monitoring is suggested to adjust the dosage of thyroid replacement therapy.

**Haematological effects**

Aplastic anaemia and agranulocytosis (in some cases fatal) have been reported in association with the use of carbamazepine. However, due to the very low incidence of these conditions, meaningful risk estimates for carbamazepine are difficult to obtain. The overall risk in the general untreated population has been estimated at 4.7 persons per million per year for agranulocytosis and 2 persons per million per year for aplastic anaemia.

Although reports of transient or persistent reductions in platelet count or white cell count are not uncommon in association with the use of carbamazepine, data are not available to estimate accurately their incidence or outcome. Nevertheless, the vast majority of leukopenia cases have not progressed to aplastic anaemia or agranulocytosis. Nonetheless, complete blood counts, including platelets and possibly reticulocytes and serum iron, should be obtained before treatment, as a baseline and periodically thereafter.

If during treatment definitely low or decreased white blood cell or platelet counts are observed, the patient and the complete blood count should be monitored closely. Teril should be discontinued if any evidence of significant bone marrow depression appears.
Because the onset of potentially serious blood dyscrasias may be rapid, patients should be made aware of early toxic signs and symptoms of a potential haematological problem, as well as symptoms of dermatological or hepatic reactions. If reactions such as fever, sore throat, and rash, ulcers in the mouth, easy bruising, petechial or purpuric haemorrhage appear, the patient should be advised to consult their physician immediately.

**Anticholinergic effects**

Teril has shown mild anticholinergic activity. Patients with increased intraocular pressure, urinary retention or prostatism should, therefore, be observed closely during therapy (see Adverse Effects).

**Ophthalmological effects**

Carbamazepine therapy has been associated with punctate cortical lens opacities and conjunctivitis, although a direct causal relationship has not been established. Baseline and periodic ophthalmological examinations are recommended.

**Psychiatric effects**

The possibility of activation of a latent psychosis and, in elderly patients, of confusion or agitation should be borne in mind.

**Suicidal Behaviour and Ideation**

Antiepileptic drugs (AEDs), including carbamazepine, increase the risk of suicidal thoughts or behaviour in patients taking these drugs for any indication. Patients treated with any AED for any indication should be monitored for the emergence or worsening of depression, suicidal thoughts or behaviour, and/or any unusual changes in mood or behaviour.

Pooled analyses of 199 placebo-controlled clinical trials (mono- and adjunctive therapy) of 11 different AEDs showed that patients randomised to one of the AEDs had approximately twice the risk (adjusted Relative Risk 1.8, 95% CI:1.2, 2.7) of suicidal thinking or behaviour compared to patients randomised to placebo. Based on the subgroup analysis, the adjusted relative risks (estimated odds ratio) of suicidal behaviour or ideation was 0.65 (95% CI: 0.08, 4.42) for carbamazepine compared to placebo. In these trials, which had a median treatment duration of 12 weeks, the estimated incidence rate of suicidal behaviour or ideation among 27,863 AED-treated patients was 0.43%, compared to 0.24% among 16,029 placebo-treated patients, representing an increase of approximately one case of suicidal thinking or behaviour for every 530 patients treated. There were four suicides in drug-treated patients in the trials and none in placebo-treated patients, but the number is too small to allow any conclusion about drug effect on suicide.

The increased risk of suicidal thoughts or behaviour with AEDs was observed as early as one week after starting drug treatment with AEDs and persisted for the duration of treatment assessed. Because most trials included in the analysis did not extend beyond 24 weeks, the risk of suicidal thoughts or behaviour beyond 24 weeks could not be assessed.

The risk of suicidal thoughts or behaviour was generally consistent among drugs in the data analysed. The finding of increased risk with AEDs of varying mechanisms of action and across a range of indications suggests that the risk applies to all AEDs used for any indication. The risk did not vary substantially by age (5-100 years) in the clinical trials analysed. Table 1 shows absolute and relative risk by indication for all evaluated AEDs.

**Table 1. Risk by indication for antiepileptic drugs in the pooled analysis**

<table>
<thead>
<tr>
<th>Indication</th>
<th>Placebo patients with events / 1000 patients</th>
<th>Drug patients with events / 1000 patients</th>
<th>Relative Risk: Incidence of events in Drug patients/ Incidence in Placebo patients</th>
<th>Risk Difference: Additional Drug patients with events per 1000 patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epilepsy</td>
<td>1.0</td>
<td>3.4</td>
<td>3.5</td>
<td>2.4</td>
</tr>
<tr>
<td>Psychiatric</td>
<td>5.7</td>
<td>8.5</td>
<td>1.5</td>
<td>2.9</td>
</tr>
<tr>
<td>Other</td>
<td>1.0</td>
<td>1.8</td>
<td>1.9</td>
<td>0.9</td>
</tr>
<tr>
<td>Total</td>
<td>2.4</td>
<td>4.3</td>
<td>1.8</td>
<td>1.9</td>
</tr>
</tbody>
</table>
The relative risk for suicidal thoughts or behaviour was higher in clinical trials for epilepsy than in clinical trials for psychiatric or other conditions, but the absolute risk differences were similar for the epilepsy and psychiatric indications.

Anyone considering prescribing Trileptal or any other AED must balance this risk with the risk of untreated illness. Epilepsy and many other illnesses for which AEDs are prescribed are themselves associated with morbidity and mortality and an increased risk of suicidal thoughts and behaviour. Should suicidal thoughts and behaviour emerge during treatment, the prescriber needs to consider whether the emergence of these symptoms in any given patient may be related to the illness being treated.

Patients, their caregivers, and families should be informed that AEDs increase the risk of suicidal thoughts and behaviour and should be advised of the need to be alert for the emergence or worsening of the signs and symptoms of depression, any unusual changes in mood or behaviour, or the emergence of suicidal thoughts, behaviour, or thoughts about self-harm. Behaviours of concern should be reported immediately to healthcare providers.

**Endocrinological effects**

Breakthrough bleeding has been reported in women taking carbamazepine while using hormonal contraceptives. The reliability of hormonal contraceptives may be adversely affected by Teril (see **INTERACTIONS WITH OTHER MEDICINES**). Women of childbearing potential should be advised to consider using alternative forms of birth control while taking Teril. Due to enzyme induction, Teril may cause failure of the therapeutic effect of any drugs containing oestrogen and/or progesterone (e.g., failure of contraception) (see **INTERACTIONS WITH OTHER MEDICINES**).

There have very rare been reports of impaired male fertility and/or abnormal spermatogenesis.

**Monitoring of plasma concentrations**

Although correlations between dosage and plasma concentrations of carbamazepine, and between plasma concentrations and clinical efficacy or tolerability are rather tenuous, monitoring of the plasma concentrations may be useful in the following circumstances: dramatic increase in seizure frequency; verification of patient compliance; during pregnancy; when treating children or adolescents; in suspected absorption disorders; in suspected toxicity when more than one drug is being used (see **INTERACTIONS WITH OTHER MEDICINES**).

**Dose reduction and withdrawal effects**

Abrupt dose reduction or withdrawal may precipitate convulsions or even status epilepticus, therefore carbamazepine should be withdrawn gradually. If treatment with Teril has to be withdrawn abruptly in a patient with epilepsy, the changeover to the new antiepileptic compound should be made under cover of a suitable drug (e.g. intravenous diazepam or intravenous phenytoin).

**Interference with serological testing**

Carbamazepine may result in false positive perphenazine concentrations in HPLC analysis due to interference.

Carbamazepine and the 10,11-epoxide metabolite may result in false positive tricyclic antidepressant concentration in fluorescence polarized immunoassay method.

**Effects on ability to drive or use machines**

The patient's ability to react may be impaired by the medical condition resulting in seizures and adverse reactions including dizziness, drowsiness, ataxia, diplopia, impaired accommodation and blurred vision reported with carbamazepine, especially at the start of treatment or in association with dose adjustments. Patients should, therefore, exercise due caution when driving a vehicle or operating machinery.
Carcinogenicity, Mutagenesis, Impairment of Fertility

In rats treated with oral carbamazepine at doses of 25, 75 and 250 mg/kg/day for 2 years, the incidence of hepatocellular tumours was dose-dependently increased in females, and aspermatogenesis and testicular atrophy were observed at all doses. This dose range is 0.2 to 2 times the maximum recommended clinical dose of 1200 mg/day, on a surface area basis. The significance of the carcinogenicity findings relative to the use of carbamazepine in humans is not known. There have been very rare reports of impaired male fertility and/or abnormal spermatogenesis.

Genotoxicity

Bacterial and mammalian mutagenicity studies yielded negative results.

Use in Pregnancy (Category D)

In animals (mice, rats, rabbits), oral administration of carbamazepine during organogenesis led to increased embryonic mortality at daily doses which caused maternal toxicity (above 200 mg/kg/day, or about 1.5 times the maximum recommended clinical dose on a surface area basis). In rats, there was also some indication of abortion at 300 mg/kg/day. Near-term rat foetuses showed growth retardation at maternally toxic doses. In mice, oral administration of carbamazepine at doses of 40 to 240 mg/kg/day (less than the maximum recommended clinical dose on a surface area basis) caused defects (mainly dilatation of cerebral ventricles) in 4.7% of exposed foetuses compared with 1.3% of controls. In rats, a small number of congenital abnormalities occurred following oral administration of carbamazepine at doses of 250 and 650 mg/kg (respectively, 2 and 4 times the maximum recommended clinical dose on a surface area basis).

The risk of a mother with epilepsy giving birth to a baby with an abnormality is about three times that of the general population. Some of this risk is due to the anticonvulsant drugs taken. Although carbamazepine has been known to produce malformations in one animal species (the rat), the significance of this in humans is not known. Mothers taking more than one anticonvulsant drug might have a higher risk of having a baby with a malformation than mothers taking one drug. Overall, the risk of having an abnormal child as a result of medication is far outweighed by the dangers to the mother and foetus of uncontrolled epilepsy.

Review of a cohort of 1457 women exposed to carbamazepine not combined with valproate revealed a 1% incidence of spina bifida. A smaller cohort of 50 women on carbamazepine monotherapy produced two babies with spina bifida. Other congenital anomalies such as craniofacial defects, cardiovascular malformations, hypospadias and anomalies involving various body systems (e.g. fingernail hypoplasia and developmental disorder) have been reported. There is evidence suggestive of an increased risk of malformations in humans when carbamazepine has been used in combination with other anticonvulsant drugs. The risk of malformations following exposure to carbamazepine as polytherapy may vary depending on the specific drugs used and may be higher in polytherapy combinations that include valproate. Monotherapy is recommended wherever possible. Patients should be counselled regarding the possibility of an increased risk of malformations and specialist prenatal diagnosis including detailed mid-trimester ultrasound should be offered. Minimum effective doses should be given and monitoring of plasma levels is recommended. The plasma concentration could be maintained in the lower side of the therapeutic range provided seizure control is maintained (see PHARMACOLOGY - Plasma concentrations). If pregnancy occurs in a woman receiving Teril, or if the question of initiating treatment with Teril arises during pregnancy, the drug's expected benefits must be carefully weighed against its possible hazards, particularly in the first 3 months of pregnancy.

Folic acid deficiency is known to occur in pregnancy. Antiepileptic drugs have been reported to aggravate folic acid deficiency. This deficiency may contribute to the increased incidence of birth defects in the offspring of treated epileptic women. Folic acid supplementation has, therefore, been recommended before and during pregnancy. Folic acid supplementation (5 mg) should be commenced four weeks prior to and continue for twelve weeks after conception.

In the neonate

In order to prevent bleeding disorders in the offspring, it has also been recommended that vitamin K1, be given to the mother during the last weeks of pregnancy as well as to the neonate.
There have been a few cases of neonatal seizures and/or respiratory depression associated with maternal Teril and other concomitant anticonvulsant drug use. A few cases of neonatal vomiting, diarrhoea and/or decreased feeding have also been reported in association with maternal Teril use. These reactions may represent a neonatal withdrawal syndrome.

**Use in Lactation**

Carbamazepine passes into human milk (about 25 to 60% of plasma concentrations). The benefits of breastfeeding should be weighed against the possibility of adverse effects occurring in the infant. Mothers taking Teril may breastfeed their infants, provided the infant is observed for possible adverse reactions (e.g. excessive somnolence, allergic skin reaction). There have been some reports of cholestatic hepatitis in neonates exposed to carbamazepine during antenatal and/or during breastfeeding. Therefore breastfed infants of mothers treated with carbamazepine should be carefully observed for adverse hepatobiliary effects.

**INTERACTIONS WITH OTHER MEDICINES**

Cytochrome P450 3A4 (CYP 3A4) is the main enzyme catalysing the formation of the active metabolite carbamazepine-10,11-epoxide. Co-administration of inhibitors of CYP 3A4 may result in an increased plasma concentration of carbamazepine which could induce adverse reactions. The dose of Teril may have to be adjusted.

Co-administration of CYP 3A4 inducers might increase the rate of carbamazepine metabolism, thus leading to a decrease in carbamazepine plasma concentration and a potential decrease in the therapeutic effect. Similarly, discontinuation of a CYP 3A4 inducer may decrease the rate of metabolism of carbamazepine, leading to an increase in carbamazepine plasma levels. The dose of Teril may have to be adjusted.

Carbamazepine is a potent inducer of CYP3A4 and other phase I and phase II enzyme systems in the liver, and may therefore reduce plasma concentrations of co-medications mainly metabolised by CYP3A4 by induction of their metabolism (see PRECAUTIONS).

Human microsomal epoxide hydrolase has been identified as the enzyme responsible for the formation of the 10,11-transdiol derivative from carbamazepine-10,11 epoxide. Co-administration of inhibitors of human microsomal epoxide hydrolase may result in increased carbamazepine-10,11 epoxide plasma concentrations.

**Agents that may raise carbamazepine plasma concentrations:**

Since high plasma concentrations of carbamazepine may result in adverse reactions (e.g. dizziness, drowsiness, ataxia, diplopia), the dosage of Teril should be adjusted accordingly and/or the plasma concentrations monitored when used concomitantly with the substances described below.

- **Analgesics, anti-inflammatory drugs:** dextropropoxyphene, ibuprofen.
- **Androgens:** danazol.
- **Antibiotics:** macrolide antibiotics (e.g. erythromycin, troleandomycin, josamycin, clarithromycin), ciprofloxacin.
- **Antidepressants:** possibly desipramine, fluoxetine, fluvoxamine, nefazodone, paroxetine, trazodone.
- **Antiepileptics:** vigabratin.
- **Antifungal:** azoles (e.g. itraconazole, ketoconazole, fluconazole, voriconazole).
- **Antihistamines:** loratadine, terfenadine.
- **Antipsychotics:** olanzapine, quetiapine.
Antituberculosis: isoniazid.

Antivirals: protease inhibitors for HIV treatment (e.g. ritonavir).

Carbonic anhydrase inhibitors: acetazolamide

Cardiovascular drugs: diltiazem, verapramil.

Gastrointestinal drugs: possibly cimetidine, omeprazole.

Muscle relaxants: oxybutynin, dantrolene.

Platelet aggregation inhibitors: ticlopidine.

Other interactions: grapefruit juice, nicotinamide (in adults, only in high doses).

Agents that may raise the active metabolite carbamazepine-10,11-epoxide plasma levels:

Since raised plasma carbamazepine-10,11-epoxide levels may result in adverse reactions (e.g. dizziness, drowsiness, ataxia, diplopia), the dosage of Teril should be adjusted accordingly and/or the plasma levels monitored when used concomitantly with the substances described below:

Quetiapine, valproic acid, valnoctamide, valpromide and primidone.

Agents that may decrease carbamazepine plasma concentrations:

The dose of Teril consequently may have to be adjusted when used concomitantly with the substances described below.

Antiepileptics: oxcarbazepine, phenobarbital, phenytoin, primidone, progabide, and, although the data are partly contradictory, possibly also clonazepam, valproic acid or valpromide

Antineoplasics: cisplatin or doxorubicin

Antituberculosis: rifampicin

Bronchodilators or anti-asthma drugs: theophylline, aminophylline

Dermatological drugs: isotretinoin has been reported to alter the bioavailability and/or clearance of carbamazepine and carbamazepine-10,11-epoxide. Carbamazepine plasma concentrations should be monitored.

Other interactions: herbal preparations containing St John’s wort (Hypericum perforatum)

Effect of carbamazepine on plasma concentrations of concomitant drugs:

Due to induction of the hepatic mono-oxygenase enzyme system, carbamazepine may lower the plasma concentration and or diminish or even abolish the activity of certain drugs that are metabolised by this system. The dosage of the following drugs may have to be adjusted to clinical requirements:

Analgesics, anti-inflammatory agents: buprenorphine, methadone, paracetamol (long term administration of carbamazepine and paracetamol may be associated with hepatotoxicity), tramadol.

Antibiotics: Doxycycline, rifabutin.

Anticoagulants: oral anticoagulants (warfarin).
Antidepressants: bupropion, citalopram, mianserin, nefazodone, sertraline, trazodone, tricyclic antidepressants (e.g. imipramine, amitriptyline, nortriptyline, clomipramine).

Antiemetics: aprepitant.

Antiepileptics: clobazam, clonazepam, ethosuximide, lamotrigine, oxcarbazepine, primidone, tiagabine, topiramate, valproic acid. Plasma phenytoin levels have been reported both to be raised and to be lowered by carbamazepine, and mephenytoin plasma levels have been reported in rare instances to increase.

Antifungals: Itraconazole, voriconazole.

Antihelmintics: praziquantel, albendazole.

Antineoplastics: imatinib, cyclophosphamide, lapatinib, temsirolimus.

Antipsychotics: clozapine, haloperidol, bromperidol, olanzapine, quetiapine, risperidone, ziprasidone, aripiprazole, paliperidone.

Antivirals: protease inhibitors for HIV treatment (e.g. indinavir, ritonavir, saquinavir).

Anxiolytics: alprazolam, midazolam.

Bronchodilators or anti-asthma drugs: theophylline.

Contraceptives: hormonal contraceptives (alternative contraceptive methods should be considered).

Cardiovascular drugs: calcium channel blockers (dihydropyridine group) e.g. felodipine, digoxin, simvastatin, atorvastatin, lovastatin, cerivastatin, ivabradine.

Corticosteroids: corticosteroids (e.g. prednisolone, dexamethasone).

Drugs used in erectile dysfunction: tadalafil.

Immunosuppressants: cyclosporin, everolimus, tacrolimus, sirolimus.

Thyroid agents: levothyroxine.

Other drug interactions: products containing oestrogens and/or progesterones.

**Combinations that require specific consideration:**

Concomitant use of carbamazepine and levetiracetam has been reported to increase carbamazepine-induced toxicity.

Concurrent use of carbamazepine and isoniazid has been reported to increase isoniazid-induced hepatotoxicity.

Combined use of carbamazepine and lithium or metoclopramide on one hand and carbamazepine and neuroleptics (haloperidol, thioridazine) on the other may lead to an increase in neurological adverse reactions (with the latter combination even in the presence of “therapeutic” plasma level concentrations).

The causative role of carbamazepine in inducing or contributing to the development of a serotonin syndrome during concomitant use with selective serotonin reuptake inhibitors is unclear. (Symptoms of serotonin...
Teril – Product Information

syndrome include hyperthermia, tremor, convulsions, sweating, muscle contractions and changes in mental state, including confusion, irritability and extreme agitation).

Concurrent medication with Teril and some diuretics (hydrochlorothiazide, frusemide) may lead to symptomatic hyponatraemia.

Carbamazepine may antagonise the effects of non-depolarising muscle relaxants (e.g. pancuronium). Their dosage may need to be raised and patients should be monitored closely for more rapid recovery from neuromuscular blockade than expected.

Carbamazepine, like other psychoactive drugs, may reduce alcohol tolerance. It is, therefore, advisable for the patient to abstain from alcohol.

Teril should not be administered with monoamine oxidase inhibitors (MAOI) (see CONTRAINDICATIONS).

Concurrent use of Teril with hormonal contraceptives may render this type of contraceptive ineffective (see PRECAUTIONS-Endocrinological effects).

**ADVERSE EFFECTS**

Particularly at the start of treatment with Teril, or if the initial dosage is too high, or when treating elderly patients, certain types of adverse reaction occur very commonly or commonly, e.g. CNS adverse reactions (dizziness, headache, ataxia, drowsiness, fatigue, diplopia), gastrointestinal disturbances (nausea, vomiting), as well as allergic skin reactions. Such reactions can be minimised by starting with a low dose.

The dose-related adverse reactions usually abate within a few days, either spontaneously or after a dosage reduction. The occurrence of CNS adverse reactions may be a manifestation of relative overdosage or significant fluctuation in concentration of the drug in plasma. In such cases it is advisable to monitor the plasma concentrations and possibly to lower the daily dosage and/or divide it into 3 or 4 fractional doses.

Adverse drug reactions from clinical trials are listed by MedDRA system organ class. Within each system organ class, the adverse drug reactions are ranked by frequency, with the most frequent reactions first. Within each frequency grouping, adverse drug reactions are presented in order of decreasing seriousness. In addition, the corresponding frequency category for each adverse drug reaction is based on the following convention (CIOMS III): very common (≥1/10); common (≥1/100, <1/10); uncommon (≥1/1,000, <1/100); rare (≥1/10,000, <1/1,000); very rare (<1/10,000).

**Blood and lymphatic system disorders**

**Very common:** Leucopenia.

**Common:** eosinophilia, thrombocytopenia.

**Rare:** leucocytosis, lymphadenopathy.

**Very rare:** agranulocytosis/aplastic anaemia (with fatal outcome in some cases), pancytopenia, aplasia pure red cell, anaemia, anaemia megaloblastic, reticulocytosis, haemolytic anaemia.

**Immune system disorders**

**Rare:** a delayed multi-organ hypersensitivity disorder with fever, rashes, vasculitis, lymphadenopathy, pseudo lymphoma, arthralgia, leucopenia, eosinophilia, hepatosplenomegaly, abnormal liver function tests and vanishing bile duct syndrome (destruction and disappearance of the intrahepatic bile ducts), occurring in various combinations. Other organs may also be affected (e.g. lungs, kidneys, pancreas, myocardium, and colon).
Very rare: anaphylactic reactions, angioedema, hypogammaglobulinaemia.

**Endocrine disorders**

**Common:** oedema, fluid retention, weight increase, hyponatraemia and blood osmolarity decreased due to an antidiuretic hormone (ADH)-like effect, leading in rare cases to water intoxication accompanied by lethargy, vomiting, headache, confusional state, neurological disorders.

**Very rare:** galactorrhoea; gynaecomastia.

**Metabolism and Nutrition disorders**

**Rare:** folate deficiency, decreased appetite.

**Very rare:** porphyria acute (acute intermittent porphyria and variegate porphyria), porphyria non-acute (porphyria cutanea tarda).

**Psychiatric disorders**

**Rare:** hallucinations (visual or auditory), depression, aggression, agitation, anorexia, restlessness, confusional state.

**Very rare:** activation of psychosis.

**Nervous system disorders**

**Very common:** ataxia, dizziness, somnolence.

**Common:** diplopia, headache, increases in seizure frequency in patients with atypical absences.

**Uncommon:** abnormal involuntary movements (e.g. tremor, asterixis, dystonia, tics), nystagmus.

**Rare:** orofacial dyskinesia, choreoathetosis, eye movement disorder, speech disorders (e.g. dysarthria, slurred speech), neuropathy peripheral, paraesthesia, paresis. There have been some reports of paralysis and other symptoms of cerebral arterial insufficiency, but no conclusive relationship to the administration of carbamazepine could be established.

**Very rare:** neuroleptic malignant syndrome (the causative role of carbamazepine in inducing or contributing to the development of neuroleptic malignant syndrome, especially in conjunction with neuroleptics, is unclear), aseptic meningitis with myoclonus and peripheral eosinophilia, dysgeusia.

**Eye disorders**

**Common:** Accommodation disorders (e.g. blurred vision)

**Very rare:** lenticular opacities (see Precautions), conjunctivitis.

**Ear and labyrinth disorders**

**Very rare:** hearing disorders (e.g. tinnitus, hyperacusis, hypoacusis, change in pitch perception).
Cardiac disorders

Rare: cardiac conduction disorders.

Very rare: arrhythmia, Stokes-Adams attacks and syncope associated with atrioventricular block, bradycardia, cardiac failure, aggravation of symptoms of coronary insufficiency.

Vascular disorders

Rare: hypertension or hypotension

Very rare: circulatory collapse, thromboembolism (e.g. pulmonary embolism), thrombophlebitis

Respiratory, thoracic and mediastinal disorders

Very rare: pulmonary hypersensitivity characterised by fever, dyspnoea, pneumonitis or pneumonia.

Gastrointestinal disorders

Very common: vomiting, nausea.

Common: dry mouth.

Uncommon: diarrhoea, constipation

Rare: abdominal pain.

Very rare: pancreatitis, glossitis, stomatitis.

Hepato-biliary disorders

Rare: cholestatic hepatitis, parenchymal (hepatocellular) or mixed type, vanishing bile duct syndrome, jaundice.

Very rare: hepatic failure, granulomatous liver disease.

Skin and subcutaneous tissue disorders

Very common: urticaria, which may be severe, dermatitis allergic.

Uncommon: dermatitis exfoliative and erythroderma.

Rare: systemic lupus erythematosus, pruritus.

Very rare: Stevens-Johnson syndrome, toxic epidermal necrolysis, photosensitivity reaction, erythema multiforme, erythema nodosum, pigmentation disorder, purpura, acne, hyperhidrosis, alopecia, hirsutism.

Musculoskeletal, connective tissue and bone disorders

Rare: muscular weakness

Very rare: bone metabolism disorders (decrease in plasma calcium and blood 25-hydroxy-cholecalciferol) leading to osteomalacia/osteoporosis, arthralgia, myalgia, muscle spasms.
Renal and urinary disorders

Very rare: tubulointerstitial nephritis, renal failure, renal impairment (e.g. albuminuria, haematuria, oliguria and blood urea increased azotaemia), urinary retention, urinary frequency.

Reproductive system

Very rare: sexual dysfunction/erectile dysfunction, spermatogenesis abnormal (with decreased sperm count and/or motility).

General disorders

Very common: fatigue

Investigations

Very common: gamma-glutamyltransferase increased (due to hepatic enzyme induction), usually not clinically relevant.

Common: blood alkaline phosphatase increased.

Uncommon: transaminases increased.

Very rare: intraocular pressure increased, blood cholesterol increased, high density lipoprotein increased, blood triglycerides increased. Thyroid function test abnormal: decreased L-Thyroxin (free thyroxine, thyroxine, triiodothyronine) and increased blood thyroid stimulating hormone, usually without clinical manifestations, blood prolactin increased.

Adverse drug reactions from spontaneous reports (frequency not known):

The following adverse drug reactions have been derived from post-marketing experience with carbamazepine via spontaneous case reports and literature cases. Because these reactions are reported voluntarily from a population of uncertain size, it is not possible to reliably estimate their frequency which is therefore categorized as not known. Adverse drug reactions are listed according to system organ classes in MedDRA. Within each system organ class, ADRs are presented in order of decreasing seriousness.

Infections and infestations: Reactivation of Human herpes virus 6 infection

Blood and lymphatic system disorders: Bone marrow failure.

Nervous system disorders: Sedation, memory impairment.

Gastrointestinal disorders: Colitis.

Musculoskeletal and connective tissue disorders: Fracture.

Immune system disorders: Drug Rash with Eosinophilia and Systemic Symptoms (DRESS).

Skin and subcutaneous tissue disorders: Acute Generalised Exanthematous Pustulosis (AGEP), lichenoid keratosis, onychomadesis.

Investigations: Bone density decreased.
DOSAGE AND ADMINISTRATION

This product is unable to deliver all dosage regimens.

The tablets may be taken during or after meals with a little liquid.

Use in Pregnancy

Women of childbearing age under treatment with Teril should be counselled to inform their medical practitioner immediately if pregnancy is suspected (see Precautions – Use in Pregnancy).

Use in the Elderly

In elderly subjects, the dosage of Teril should be selected with caution.

Dosage Recommendations

Epilepsy:

Wherever possible, Teril should be prescribed as monotherapy. Treatment should be initiated with a low daily dosage, to be slowly increased until an optimal effect is obtained (see PHARMACOLOGY – Plasma Concentrations). After obtaining adequate seizure control, the dosage may be reduced very gradually to the minimum effect level.

Determination of plasma concentrations may help in establishing the optimum dosage (see PHARMACOLOGY – Plasma Concentrations).

When Teril is added to existing anticonvulsant therapy, this should be done gradually while maintaining, or if necessary adjusting, the dosage of the other anticonvulsant(s) (see INTERACTIONS WITH OTHER MEDICINES).

Adults and children over 15 years:

Initially 100 mg to 200 mg once or twice daily. Slowly raise the dosage until an optimum response is obtained - generally at 400 mg two or three times daily. In some patients, 1600 or even 2000 mg daily may be required in rare instances.

Children aged 6 – 15 years:

The initial dose for children 13 to 15 years is as for adults. The recommended maintenance doses are:

- 6 to 10 years: 400 to 600 mg daily.
- 11 to 15 years: 600 to 1,000 mg daily.

Daily dose should generally not exceed 1,000 mg.

Children aged less than 6 years:

Limited data are available concerning the safety and efficacy in children less than 6 years old.

Trigeminal neuralgia:

The recommended initial dose is 200 to 400 mg daily in two divided doses increasing by 200 mg each day in divided doses until pain relief is obtained. This is usually achieved with doses up to 800 mg daily. Larger doses should be given as three to four divided doses. The maximum dose should not exceed 1200 mg daily. As soon as the pain is well controlled, gradually reduce the dosage to the minimal effective level. Because trigeminal neuralgia is characterised by periods of remission, attempts should be made to reduce or discontinue the use of carbamazepine at intervals of not more than three months.
Mania and maintenance treatment of bipolar affective disorder:

The dosage range is 400 to 1,600 mg daily. When used alone in mania the starting dose of carbamazepine should be 200 to 400 mg daily in two divided doses. Dosage should be increased to 800 to 1,000 mg during the first week by daily increments of 200 mg and up to 1,600 mg if no response is found after a second week.

For maintenance treatment, carbamazepine is commenced at a dosage of 200 to 400 mg daily in two divided doses. Dosage should be increased weekly by increments of 100 mg. Due to auto induction, concentrations may fall after two to three weeks and dosage increases may be necessary after this time. The same plasma level range as for anticonvulsant therapy is considered adequate (4 to 12 microgram/mL; 17 to 50 micromol/L). However dose increases should be titrated against the appearance of side effects.

OVERDOSE

Signs and symptoms

The presenting signs and symptoms of overdosage develop within one to three hours of ingestion and usually involve the central nervous, cardiovascular, respiratory systems and the adverse drug reactions mentioned under “ADVERSE EFFECTS”. Relapse and aggravation of symptoms on the second and third day after overdose may occur. This is thought to be due to delayed absorption, possibly due to production of a gastric mass of tablets.

Central nervous system:

CNS depression, disorientation, depressed level of consciousness, somnolence, agitation, hallucination, coma, blurred vision, slurred speech, dysarthria, nystagmus, ataxia, dyskinesia, initially hyperreflexia, later hyporeflexia, convulsions (especially in small children), psychomotor disturbances, myoclonus, hypothermia, mydriasis.

Respiratory system:

Respiratory depression, pulmonary oedema.

Cardiovascular system:

Tachycardia, hypotension, at times hypertension, conduction disturbance with widening of QRS complex, syncope in association with cardiac arrest.

Gastrointestinal system:

Vomiting, delayed gastric emptying, reduced bowel motility.

Musculoskeletal system:

Rhabdomyolysis

Renal function:

Retention of urine, oliguria or anuria; fluid retention, water intoxication due to an ADH-like effect of carbamazepine.

Laboratory findings:

Hyponatraemia (see Management), leukocytosis, leukopenia, hypokalaemia, metabolic acidosis, hyperglycaemia, glycosuria, acetonuria, increased muscle creatine phosphokinase.
Management

There is no specific antidote. Management should be guided initially by the patient's clinical condition. All patients suspected of serious overdose should be admitted to hospital and the plasma carbamazepine concentration measured to confirm carbamazepine poisoning and to ascertain the size of the overdose.

Administration of activated charcoal. If the patient's level of consciousness is impaired, intubation may be necessary to protect the airway. Supportive medical care in an intensive care unit with cardiac monitoring and careful correction of electrolyte imbalance.

Hyponatraemia is not usually a problem in acute overdosage. However, in chronic intoxication it may be managed by fluid restriction and slow and careful intravenous infusion of sodium chloride 0.9%. These measures may be useful in preventing brain damage.

Special recommendations

Hypotension:

Intravenous fluid replacement. If the patient fails to respond, consider intravenous dopamine or dobutamine.

Disturbances of cardiac rhythm:

There are no data regarding drug treatment of carbamazepine induced arrhythmias. These should, therefore, be handled according to the circumstances in each patient.

Convulsions:

Initially, administer a benzodiazepine (e.g. diazepam) if seizures occur. If seizures recur, another anticonvulsant, e.g. phenobarbitone (with caution because of increased respiratory depression), may be considered.

Charcoal haemoperfusion has been recommended. Forced diuresis, haemodialysis, and peritoneal dialysis have been reported not to be effective.

For further information on the management of overdosage, contact the Poisons Information Centre (131126).

PRESENTATION AND STORAGE CONDITIONS

Teril 200 mg tablet
White, uncoated, marked CB on one side and G on the reverse.
200
Available in bottles of 200 tablets.

Store below 30°C.

NAME AND ADDRESS OF THE SPONSOR

Alphapharm Pty Limited

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POISON SCHEDULE OF THE MEDICINE

S4 – Prescription Only Medicine

DATE OF FIRST INCLUSION IN THE AUSTRALIAN REGISTER OF THERAPEUTIC GOODS (THE ARTG)

20/09/1991

DATE OF MOST RECENT AMENDMENT

03/12/2015