

## SUMMARY OF PRODUCT CHARACTERISTICS

### 1 NAME OF THE MEDICINAL PRODUCT

Paracetamol / Caffeine 500mg/65mg Tablets

### 2 QUALITATIVE AND QUANTITATIVE COMPOSITION

Each tablet contains Paracetamol 500mg and Caffeine 65mg  
For full list of excipients, see section 6.1

### 3 PHARMACEUTICAL FORM

Capsule shaped tablets

White to off white, capsule shaped, biconvex tablets plain on both sides approximately 17.46mm X 7.14mm.

### 4 CLINICAL PARTICULARS

#### 4.1 Therapeutic indications

Symptomatic treatment of mild to moderate pain and/or fever in adults and children aged 16 years or over.

#### 4.2 Posology and method of administration

Oral use.

Adults (16 years and over):

Two tablets up to four times daily. The dose should not be repeated more frequently than every 4 hours. Do not exceed 8 tablets in 24 hours.

Elderly:

As for adults.

Not recommended for children under 16 years.

*Impaired Renal Function:*

In case of renal insufficiency dose adjustment is necessary:

*Impaired Hepatic Function:*

In patients with impaired hepatic function or Gilbert's syndrome, the dose must be reduced or the dosing interval prolonged.

The daily effective dose of paracetamol should not exceed 60 mg/kg/day (up to maximum 2 g paracetamol /day) in the following situations:

- \_ Adults or adolescents weighing less than 50 kg
- \_ Mild to moderate hepatic insufficiency, Gilbert's syndrome (familial non-heamolytic jaundice)
- \_ Dehydration
- \_ Chronic malnutrition
- \_ Chronic alcoholism

Method of administration

Route of administration: Oral

#### **4.3 Contraindications**

Hypersensitivity to paracetamol, caffeine or to any of the excipients listed in section 6.1.

#### **4.4 Special warnings and precautions for use**

Immediate medical advice should be sought in the event of an overdose, even if you feel well, because of the risk of delayed, serious liver damage.

Prolonged or frequent use is discouraged. Patients should be advised not to take other Paracetamol containing products concurrently. Taking multiple daily doses in one administration can severely damage the liver; in such case unconsciousness does not occur. However, medical assistance should be sought immediately. Prolonged use except

under medical supervision may be harmful. In adolescents treated with 60mg/kg daily of Paracetamol, the combination with another antipyretic is not justified except in the case of ineffectiveness.

Caution is advised in the administration of Paracetamol to patients with mild and severe renal insufficiency, mild to moderate hepatocellular insufficiency (including Gilbert's syndrome), severe hepatic insufficiency (Child-Pugh>9), acute hepatitis, concomitant treatment with medicinal products affecting hepatic functions, glucose-6-phosphatedehydrogenase deficiency, haemolytic anaemia, dehydration, alcohol abuse and chronic malnutrition (see section 4.2).

The hazards of overdose are greater in those with non-cirrhotic alcoholic liver disease. Caution should be exercised in cases of chronic alcoholism. The daily dose should not exceed 2 grams in such cases.

Alcohol should not be used during the treatment with paracetamol.

Caution is advised in asthmatic patients sensitive to aspirin, because light reaction bronchospasm with paracetamol (cross-reaction) has been reported in less than 5% of the patients tested.

Paracetamol & Caffeine 500mg/65mg Tablets should be given with care to patients with gout, hyperthyroidism and arrhythmia.

The patient should limit the use of caffeine containing products when taking Paracetamol & Caffeine 500mg/65mg Tablets, as excess caffeine may cause nervousness, irritability, sleeplessness and occasionally rapid heart beat.

Glomerular filtration Dose

10-50 ml/min 1 tablet every 6 hours

< 10 ml/min 1 tablet every 8 hours

As caffeine is found naturally in tea, coffee and chocolate, and in some carbonated drinks there is the potential for users to take more than the recommended 390 mg/day of caffeine (6 tablets) per day. Patients should take account of dietary and other medicinal sources of caffeine and ensure that they do not exceed the stated dose.

Typical amounts of caffeine available from dietary sources are

Brewed coffee; 50-100mg/100ml\*

Instant coffee and tea: 20-73mg/100ml\*

Carbonated drinks (cola) 9-19mg/100ml\*

Chocolate 5-20mg/100ml

(\*100ml is equivalent to about 1 small cup of fluid)

#### **4.5 Interaction with other medicinal products and other forms of interaction**

Hepatotoxic substances may increase the possibility of Paracetamol accumulation and overdose. The risk of hepatotoxicity of paracetamol may be increased by drugs which induce liver microsomal enzymes such as barbiturates, tricyclic antidepressants, and alcohol.

Probenecid causes an almost two-fold reduction in clearance of Paracetamol by inhibiting its conjugation with glucuronid acid. A reduction of the Paracetamol dose should be considered for concomitant treatment with probenecid.

Salicylamide may prolong the elimination half-life of Paracetamol.

Metoclopramide accelerate absorption of Paracetamol.

Cholestyramine reduces absorption of Paracetamol.

Concomitant use of Paracetamol (4 g per day for at least 4 days) with oral anticoagulants may lead to slight variations of INR values. In this case, increased monitoring of INR values should be done during the duration of the combination and after its discontinuation.

Isoniazid reduces paracetamol clearance by 20%, with possible potentiation of its action and/or toxicity, by inhibiting its metabolism in the liver. The clinical relevance is unknown.

Paracetamol decreases the bioavailability of lamotrigine with possible reduction of its effect due to possible induction of its metabolism in the liver.

Co-administration of paracetamol with zidovudine may result in neutopenia or hepatotoxicity. However, these effects have not been consistently reported. The chronic / multiple dose paracetamol use in patients on zidovudine therapy should be avoided, however, if chronic paracetamol and zidovudine are to be given concurrently white blood count and

liver function tests should be monitored particularly in malnourished patients.

Paracetamol may affect the pharmacokinetics of chloramphenicol. Monitoring of chloramphenicol plasma levels is recommended if combining paracetamol with chloramphenicol injection treatment.

Interference with laboratory tests: Paracetamol may affect uric acid tests by wolframato phosphoric acid, and blood sugar tests by glucose-oxidase-peroxidase.

### **Caffeine**

Phenylpropanolamine increases caffeine plasma concentrations four-fold. There is a risk of additive CNS adverse events. Isolated reports describe the development of acute psychosis when caffeine was given with phenylpropanolamine.

Fluvoxamine, a potent inhibitor of CYP 1 A2, markedly reduces the clearance of caffeine. Concomitant administration may lead to caffeine intoxication.

Ciprofloxacin reduces caffeine metabolism, leading to two-fold increases in caffeine plasma concentrations.

Caffeine, a CNS stimulant, has an antagonistic effect towards the action of sedatives and tranquilizers. Caffeine may enhance the tachycardic effect of phenylpropanolamine and other sympathomimetic drugs.

Caffeine can increase blood pressure and counters the hypotensive action of Beta blockers such as atenolol, metoprolol, oxprenolol and propranolol. This medicine should not be used at the same time as beta blockers.

Disulfiram decreases caffeine clearance by up to 50%. Concomitant use of disulfiram and Paracetamol & Caffeine 500mg/65mg Tablets should be avoided.

Dipyridamole: injectable dipyrimidole: decrease of the vasodilating effect of dipyrimidole.

Treatment with caffeine should be discontinued at least 5 days before myocardial imaging. Coffee, tea and chocolate consumption should be avoided in the 24 hours preceding the test. Use with caution.

Enoxacin: increase of caffeine plasmatic concentrations due to a decrease of its hepatic metabolism, which can lead to excitement or hallucinations. Concomitant use is therefore not recommended.

Mexiletine: increase of caffeine plasmatic concentration due to inhibition of its hepatic metabolism with mexiletine. To be taken into account.

Norfloxacin: increase of caffeine plasmatic concentration due to inhibition of its hepatic metabolism with norfloxacin. To be taken into account.

Stiripentol: possible increase of caffeine plasmatic concentration with risk of overdose, due to its hepatic metabolism inhibition. Use with caution.

Caffeine exerts a competitive inhibition of the metabolism of clozapine. Therefore clozapine and caffeine must not be used concurrently.

Use of lithium carbonate and caffeine may cause a small decrease in serum lithium levels. Therefore concomitant ingestion of caffeine should be avoided. In case of concomitant use, the risk of an increase in serum lithium on abrupt cessation of caffeine should be taken into account.

Monoamine oxidase inhibitors may increase the stimulant effects of caffeine.

Methoxsalen reduces clearance of caffeine and may increase the effects of caffeine.

Phenytoin doubles caffeine clearance, although caffeine does not affect the metabolism of phenytoin.

Pipemidic acid reduces caffeine clearance, enhancing the effects of caffeine.

Theophylline and caffeine share the same metabolic pathway, leading to decreased clearance times for theophylline when used concurrently with caffeine. Concomitant use should be avoided.

Levothyroxine, like caffeine can increase blood pressure, and therefore these two active ingredients should not be used concurrently.

Ephedrine and caffeine interact to produce significant cardiovascular effects. Therefore caffeine should be avoided when ephedrine is being taken.

#### **4.6 Fertility, pregnancy and lactation**

##### Pregnancy

Epidemiological studies in human pregnancy have shown no ill effects due to paracetamol used in the recommended dosage, but patients should follow the advice of their doctor regarding its use.

There is a potential increased risk of lower birth weight and spontaneous abortion associated with caffeine consumption during pregnancy.

##### Lactation

Paracetamol is excreted in breast milk but not in a clinically significant amount. Available published data do not contraindicate breast feeding.

Caffeine in breast milk may have a stimulating effect on breast-fed infants. Irritability and poor sleeping pattern in the infant have been reported.

##### Fertility

There is insufficient information available on the effects of Paracetamol and Caffeine on human fertility.

#### **4.7 Effects on ability to drive and use machines**

Paracetamol & Caffeine mg/65mg Tablets has no or negligible influence on the ability to drive and use machines.

#### 4.8 Undesirable effects

Adverse events from historical clinical trial data are both infrequent and from minimal patient exposure. Accordingly, adverse events reported from extensive post-marketing experience at therapeutic/labelled dose are listed below by system organ class and frequency.

Frequencies are defined as: very common ( $>1/10$ ), common ( $>1/100$ ,  $<1/10$ ), uncommon ( $>1/1000$ ,  $<1/100$ ), rare

( $>1/10,000$ ,  $<1/1000$ ), very rare ( $<1/10,000$  including isolated reports) and not known (cannot be estimated from available data).

Frequency	System	Symptoms
Common $>1/100$ , $<1/10$	Psychiatric disorders:	Insomnia, restlessness, anxiety
	Gastrointestinal disorders	Gastrointestinal disorder
	Nervous system disorders	Dizziness, Nervousness
Rare $>1/10000$ - $<1/1000$	Blood and lymphatic system disorders	Platelet disorders, stem cell disorders.
	Immune system disorders	Anaphylactic reactions, allergic dermatitis, Allergies (excluding angioedema).
	Psychiatric disorders	Depression NOS, confusion, hallucinations.
	Nervous system disorders	Tremor NOS, headache NOS
	Eye disorders	Abnormal vision.
	Cardiac disorders	Oedema.
	Gastrointestinal disorders	Haemorrhage NOS, abdominal pain NOS, diarrhoea NOS, nausea, vomiting.
	Hepato-biliary disorders	Hepatic function abnormal, hepatic failure, hepatic necrosis, jaundice.
Skin and subcutaneous tissue disorders	Pruritus, rash, sweating, purpura, angioedema, urticaria	

	General disorders and administration site conditions	Dizziness (excluding vertigo), malaise, pyrexia, sedation, drug interaction NOS.
	Injury, poisoning and procedural complications	Overdose and poisoning
Very Rare (< 10 000)	Hepato-biliary disorders	Hepatotoxicity, hepatic function abnormal, increased transaminases
	General disorders and administration site conditions	hypersensitivity reaction (requiring discontinuation of treatment)
	Blood and lymphatic system disorders	thrombocytopenia leukopenia neutropenia hemolytic anemia agranulocytosis
	Metabolism and nutrition disorders	Hypoglycaemia
	Renal and urinary disorders	Sterile pyuria (cloudy urine) and renal side effects
	Respiratory, thoracic and mediastinal disorder	Bronchospasm
	Skin and subcutaneous disorders	Serious skin reactions have been reported.

Not known: Irritability, Palpitations, tachycardia, Edema of the larynx, anaphylactic shock, anaemia, bronchospasm\*, liver alteration and hepatitis, renal alteration (severe renal impairment, nephrite interstitial, haematuria, anuresis), gastrointestinal effects and vertigo have been reported.

\* There have been cases of bronchospasm with paracetamol, but these are more likely in asthmatics sensitive to aspirin or other NSAIDs.

When the recommended dosing regimen is combined with dietary caffeine intake, the resulting higher dose of caffeine may increase the potential for caffeine-related adverse effects such as nervousness, dizziness, insomnia, restlessness, anxiety, irritability, headache, gastrointestinal disorder and palpitations.

### Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the Yellow Card Scheme; website: [www.mhra.gov.uk/yellowcard](http://www.mhra.gov.uk/yellowcard)

## **4.9 Overdose**

Liver damage is possible in adults who have taken 10 g or more of paracetamol. Ingestion of 5 g or more of paracetamol may lead to liver damage if the patient has risk factors (see below).

### **Risk factors**

If the patient

a) Is on long term treatment with carbamazepine, phenobarbitone, phenytoin, primidone, rifampicin, St John's Wort or other drugs that induce liver enzymes.

Or

b) Regularly consumes ethanol in excess of recommended amounts.

Or

c) Is likely to be glutathione deplete e.g. eating disorders, cystic fibrosis, HIV infection, starvation, cachexia.

### **Symptoms**

Symptoms of paracetamol overdose in the first 24 hours are pallor, nausea, vomiting, anorexia and abdominal pain.

Liver damage may become apparent 12 to 48 hours after ingestion. Abnormalities of glucose metabolism and metabolic acidosis may occur. In severe poisoning, hepatic failure may progress to encephalopathy, haemorrhage, hypoglycaemia, cerebral oedema, and death. Acute renal failure with acute tubular necrosis, strongly suggested by loin pain, haematuria and proteinuria, may develop even in the absence of severe liver damage. Cardiac arrhythmias and pancreatitis have been reported.

### **Management**

Immediate treatment is essential in the management of paracetamol overdose. Despite a lack of significant early symptoms, patients should be referred to hospital urgently for immediate medical attention. Symptoms may be limited to nausea or vomiting and may not reflect the severity of overdose or the risk of organ damage. Management should be in accordance with established treatment guidelines.

Treatment with activated charcoal should be considered if the overdose has been taken within 1 hour. Plasma paracetamol concentration should be measured at 4 hours or later after ingestion (earlier concentrations are unreliable).

Treatment with N-acetylcysteine may be used up to 24 hours after ingestion of paracetamol, however, the maximum protective effect is obtained up to 8 hours post-ingestion. The effectiveness of the antidote declines sharply after this time. If required the patient should be given intravenous N-acetylcysteine, in line with the established dosage schedule. If vomiting is not a problem, oral methionine may be a suitable alternative for remote areas, outside hospital. Management of patients who present

with serious hepatic dysfunction beyond 24h from ingestion should be discussed with the National Poisons Information Unit or a liver unit.

## **Caffeine**

### **Symptoms**

Common symptoms include anxiety, nervousness, restlessness, insomnia, excitement, muscle twitching, confusion, convulsions. For high intake of caffeine, hyperglycemia could also appear. Cardiac Symptoms include tachycardia and cardiac arrhythmia.

It must be noted that for clinically significant symptoms of caffeine overdose to occur with this product, the amount ingested would be associated with serious paracetamol-related toxicity.

### **Management**

The symptoms of caffeine overdose are controlled by reducing or stopping caffeine intake

## **5 PHARMACOLOGICAL PROPERTIES**

### **5.1 Pharmacodynamic properties**

Pharmacotherapeutic Group: Analgesics; Other Analgesics and Antipyretics;  
Analides: Paracetamol, combinations excl.

ATC code: N02B E51

The combination of paracetamol and caffeine is a well-established analgesic combination.

#### Paracetamol

##### ANALGESIC:

The mechanism of analgesic action has not been fully determined. Paracetamol may act predominantly by inhibiting a prostaglandin synthesis in the central nervous system (CNS) and to a lesser extent through a peripheral action by blocking pain-impulse generation. The peripheral action may also be due to inhibition of prostaglandin synthesis or to inhibition of the synthesis or actions of other substances that sensitise pain receptors to mechanical or chemical stimulation.

##### ANTIPYRETIC:

Paracetamol probably produces antipyresis by acting centrally on the hypothalamic heat-regulating centre to produce peripheral vasodilation resulting in increased blood flow through the skin, sweating, and heat loss. The central action probably involves inhibition of prostaglandin synthesis in the hypothalamus.

#### Caffeine

Central nervous system stimulant – Caffeine stimulates all levels of the CNS, although its cortical effects are milder and of shorter duration than those of amfetamines.

##### ANALGESIA ADJUNCT:

Caffeine constricts cerebral vasculature with an accompanying decrease in cerebral blood flow and in the oxygen tension of the brain. It is believed that caffeine helps to

relieve headache by providing a more rapid onset of action and/or enhanced pain relief with lower doses of analgesic.

## 5.2 Pharmacokinetic properties

### PARACETAMOL

#### Absorption and Fate

Paracetamol is readily absorbed from the gastro-intestinal tract with peak plasma concentrations occurring about 30 minutes to 2 hours after ingestion. It is metabolised in the liver and excreted in the urine mainly as the glucuronide and sulphate conjugates. Less than 5% is excreted as unchanged paracetamol. The elimination half-life varies from about 1 to 4 hours. Plasma-protein binding is negligible at usual therapeutic concentrations but increases with increasing concentrations.

A minor hydroxylated metabolite which is usually produced in very small amounts by mixed-function oxidases in the liver and which is usually detoxified by conjugation with liver glutathione may accumulate following paracetamol overdose and cause liver damage.

Physiopathological Variations: Renal Insufficiency: In cases of severe renal insufficiency (creatinine clearance lower than 10 ml/min) the elimination of paracetamol and its metabolites is delayed

### CAFFEINE

#### Absorption and Fate

Caffeine is absorbed readily after oral administration and is widely distributed throughout the body. Caffeine is metabolised almost completely via oxidation, demethylation, and acetylation, and is excreted in the urine as 1-methyluric acid, 1-methylxanthine, 7-methylxanthine, 1,7-dimethylxanthine (paraxanthine), 5-acetylamino-6-formylamino-3-methyluracil (AFMU), and other metabolites with only about 1% unchanged.

## 5.3 Preclinical safety data

There is no data of relevance to the prescriber that have not already been included in other sections of SPC.

## 6 PHARMACEUTICAL PARTICULARS

### 6.1 List of excipients

Povidone K-30 (E1201)

Povidone K-90 (E1201)

Potato starch

Pregelatinised starch

Purified Talc  
Croscarmellose sodium  
Stearic acid (E570)  
Magnesium stearate

## **6.2 Incompatibilities**

Not applicable

## **6.3 Shelf life**

24 months

## **6.4 Special precautions for storage**

No special storage conditions.

## **6.5 Nature and contents of container**

Paracetamol & Caffeine 500mg/65mg Tablets are packaged in blister packs comprising of white opaque PVC/PVdC (20 micron/40gsm) and with backing of foil, which are placed in an outer carton along with leaflet. These are available in the pack sizes of 4, 6, 12 and 16 tablets.

Not all pack sizes may be marketed

## **6.6 Special precautions for disposal**

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

## **7 MARKETING AUTHORISATION HOLDER**

Bristol Laboratories Limited

Unit 3, Canalside, Northbridge Road,  
Berkhamsted, Hertfordshire,  
HP4 1EG, United Kingdom

**8    MARKETING AUTHORISATION NUMBER(S)**

PL 17907/0305

**9    DATE OF FIRST AUTHORISATION/RENEWAL OF THE  
AUTHORISATION**

21/01/2015

**10   DATE OF REVISION OF THE TEXT**

01/06/2016