1. Name of the medicinal product

Zaponex 25 mg Tablets
Clozapine

Zaponex 100 mg Tablets
Clozapine

UK Zaponex® Official Recommendations

As a consequence of a recent European regulatory initiative, the Zaponex Summary of Product Characteristics (SmPC) has been harmonised across Europe. The SmPC states that blood monitoring should be carried out in accordance with national-specific official recommendations. These are reproduced below.

The Zaponex Treatment Access System (ZTAS) was developed in order to manage the risk of agranulocytosis associated with clozapine. It is available 24 hours a day. When a monitoring service is not used, evidence suggests a mortality rate from agranulocytosis of 0.3%.\[1\]

This is compared to a mortality rate when Zaponex is used in conjunction with the Zaponex Treatment Access System, of 0.01\%.\[2\]

The Zaponex Treatment Access System provides for the centralised monitoring of leucocyte and neutrophil counts which is a mandatory requirement for all patients in the UK who are treated with Zaponex. The use of Zaponex is restricted to patients who are registered with the Zaponex Treatment Access System. In addition to registering their patients, prescribing physicians must register themselves and a nominated pharmacist with the Zaponex Treatment Access System. All Zaponex-treated patients must be under the supervision of an appropriate specialist and supply of Zaponex is restricted to hospital and retail pharmacies registered with the Zaponex Treatment Access System. Zaponex is not sold to, or distributed through wholesalers.

In the UK, a white cell count with a differential count must be monitored:
- At least weekly for the first 18 weeks of treatment
- At least at 2 week intervals between weeks 18 and 52
- After 1 year of treatment with stable neutrophil counts, patients may be monitored at least at 4 week intervals
- Monitoring must continue throughout treatment and for at least 4 weeks after discontinuation

The Zaponex Treatment Access System maintains a database which includes all patients who have developed abnormal leucocyte or neutrophil findings and who should not be re-exposed to Zaponex or any other brand of clozapine.

Prescribers and pharmacists should adhere to brand prescribing and dispensing of clozapine in order to prevent the disruption to effective monitoring that may be caused if patients switch brands.

Furthermore, in order to protect patient safety, at any one time patients should only be prescribed one brand of clozapine and only registered with the monitoring service connected to that brand.

For further information regarding Zaponex and the Zaponex Treatment Access System please call 0207 365 58 42.


2. Qualitative and quantitative composition

Each tablet contains 25 or 100 mg clozapine.

Excipients with known effect also include lactose monohydrate 48 mg per tablet (Zaponex 25 mg Tablet) or 192 mg per tablet (Zaponex 100 mg Tablet).

For a full list of excipients, see section 6.1.

3. Pharmaceutical form

Tablet.

Zaponex 25 mg: Yellow, round tablet, scored with a division mark on both sides and embossed with “CPN 25” on one side.

Zaponex 100 mg: Yellow, round tablet, scored with a division mark on both sides and embossed with “CPN 100” on one side.

The tablets can be divided into equal doses.

4. Clinical particulars

4.1 Therapeutic indications

Treatment-resistant schizophrenia
Clozapine is indicated in treatment-resistant schizophrenic patients and in schizophrenia patients who have severe, untreatable neurological adverse reactions to other antipsychotic agents, including atypical antipsychotics.

Treatment resistance is defined as a lack of satisfactory clinical improvement despite the use of adequate doses of at least two different antipsychotic agents, including an atypical antipsychotic agent, prescribed for adequate duration.

Psychosis during the course of Parkinson’s disease
Clozapine is also indicated in psychotic disorders occurring during the course of Parkinson’s disease, in cases where standard treatment has failed.

4.2 Posology and method of administration

Posology
The dosage must be adjusted individually. For each patient the lowest effective dose should be used. For doses not realisable/practicable with one strength, other strengths of this medicinal product are available. Cautious titration and a divided dosage schedule are necessary to minimise the risks of hypotension, seizure and sedation.

Initiation of clozapine treatment must be restricted to those patients with a WBC count ≥ 3500/mm³ (3.5x10⁹/L) and an ANC ≥ 2000/mm³ (2.0x10⁹/L) within standardised normal limits.

Dose adjustment is indicated in patients who are also receiving medicinal products that have pharmacodynamic and pharmacokinetic interactions with clozapine, such as benzodiazepines or selective serotonin re-uptake inhibitors (see section 4.5).

Switching from a previous antipsychotic therapy to clozapine
It is generally recommended that clozapine should not be used in combination with other antipsychotics. When clozapine therapy is to be initiated in a patient undergoing oral antipsychotic therapy, it is recommended that the other antipsychotic should first be discontinued by tapering the dosage downwards.

The following dosages are recommended:

Treatment-resistant schizophrenic patients

Starting therapy
12.5 mg once or twice on the first day, followed by 25 mg once or twice on the second day. If well tolerated, the daily dose may then be increased slowly in increments of 25 to 50 mg in order to achieve a dose level of up to 300 mg/day within 2 to 3 weeks. Thereafter, if required, the daily dose may be further increased in increments of 50 to 100 mg at half-weekly or, preferably, weekly intervals.

Therapeutic dose range
In most patients, antipsychotic efficacy can be expected with 200 to 450 mg/day given in divided doses. The total daily dose may be divided unevenly, with the larger portion at bedtime.

Maximum dose
To obtain full therapeutic benefit, a few patients may require larger doses, in which case judicious increments (not exceeding 100 mg) are permissible up to 900 mg/day. However, the possibility of increased adverse reactions (in particular seizures) occurring at doses over 450 mg/day must be borne in mind.

Maintenance dose
After achieving maximum therapeutic benefit, many patients can be maintained effectively on lower doses. Careful downward titration is therefore recommended. Treatment should be maintained for at least 6 months. If the daily dose does not exceed 200 mg, once daily administration in the evening may be appropriate.

Ending therapy
In the event of planned termination of clozapine therapy, a gradual reduction in dose over a 1- to 2-week period is recommended. If abrupt discontinuation is necessary, the patient should be carefully observed for the occurrence of withdrawal reactions (see section 4.4).

Re-starting therapy
In patients in whom the interval since the last dose of clozapine exceeds 2 days, treatment should be re-initiated with 12.5 mg given once or twice on the first day. If this dose is well tolerated, it may be feasible to titrate the dose to the therapeutic level more quickly than is recommended for initial treatment. However, in any patient who has previously experienced respiratory or cardiac arrest with initial dosing (see section 4.4), but was then able to be successfully titrated to a therapeutic dose, re-titration should be carried out with extreme caution.

Psychotic disorders occurring during the course of Parkinson’s disease, in cases where standard treatment has failed

Starting therapy
The starting dose must not exceed 12.5 mg/day, taken in the evening. Subsequent dose increases must be by 12.5 mg increments, with a maximum of two increments a week up to a maximum of 50 mg, a dose that cannot be reached until the end of the second week. The total daily amount should preferably be given as a single dose in the evening.

Therapeutic dose range
The mean effective dose is usually between 25 and 37.5 mg/day. In the event that treatment for at least one week with a dose of 50 mg fails to provide a satisfactory therapeutic response, dosage may be cautiously increased by increments of 12.5 mg/week.
Active liver disease associated with nausea, anorexia or jaundice; progressive liver disease, hepatic failure.

Severe renal or cardiac disorders (e.g. myocarditis).

Circulatory collapse and/or CNS depression of any cause.

Alcoholic and other toxic psychoses, drug intoxication, comatose condition.

Uncontrolled epilepsy.

Impaired bone marrow function.

History of clozapine-induced agranulocytosis.

Patients unable to undergo regular blood tests.

Hypersensitivity to the active substance or to any of the excipients.

Zaponex is administered orally.

Method of administration
Zaponex is administered orally.

4.3 Contraindications

- Hypersensitivity to the active substance or to any of the excipients, listed in section 6.1.
- Patients unable to undergo regular blood tests.
- History of toxic or idiosyncratic granulocytopenia/agranulocytosis (with the exception of granulocytopenia/agranulocytosis from previous chemotherapy).
- History of clozapine-induced agranulocytosis.
- Impaired bone marrow function.
- Uncontrolled epilepsy.
- Alcoholic and other toxic psychoses, drug intoxication, comatose conditions.
- Circulatory collapse and/or CNS depression of any cause.
- Severe renal or cardiac disorders (e.g. myocarditis).
- Active liver disease associated with nausea, anorexia or jaundice; progressive liver disease, hepatic failure.
- Paralytic ileus.
- Clozapine treatment must not be started concurrently with substances known to have a substantial potential for causing agranulocytosis; concomitant use of depot antipsychotics is to be discouraged.

Maximum dose
The dose of 50 mg/day should only be exceeded in exceptional cases, and the maximum dose of 100 mg/day must never be exceeded.

Dose increases should be limited or deferred if orthostatic hypotension, excessive sedation or confusion occurs. Blood pressure should be monitored during the first weeks of treatment.

Maintenance dose
When there has been complete remission of psychotic symptoms for at least 2 weeks, an increase in anti-parkinsonian medication is possible if indicated on the basis of motor status. If this approach results in the recurrence of psychotic symptoms, clozapine dosage may be increased by increments of 12.5 mg/week up to a maximum of 100 mg/day, taken in one or two divided doses (see above).

Ending therapy
A gradual reduction in dose by steps of 12.5 mg over a period of at least one week (preferably two) is recommended.

Treatment must be discontinued immediately in the event of neutropenia or agranulocytosis [see section 4.4]. In this situation, careful psychiatric monitoring of the patient is essential since symptoms may recur quickly.

Special populations

Hepatic impairment
Patients with hepatic impairment should receive Zaponex with caution along with regular monitoring of liver function tests [see section 4.4].

Paediatric population
No paediatric studies have been performed. The safety and efficacy of Zaponex in children and adolescents under the age of 16 years have not yet been established. It should not be used in this group until further data become available.

Patients 60 years of age and older
Initiation of treatment is recommended at a particularly low dose [12.5 mg given once on the first day] with subsequent dose increments restricted to 25 mg/day.

Method of administration
Zaponex is administered orally.

4.4 Special warnings and precautions for use

Agranulocytosis
Clozapine can cause agranulocytosis. The incidence of agranulocytosis and the fatality rate in those developing agranulocytosis have decreased markedly since the institution of white blood cell (WBC) counts and absolute neutrophil count (ANC) monitoring. The following precautionary measures are therefore mandatory and should be carried out in accordance with official recommendations.

Because of the risks associated with clozapine, its use is limited to patients in whom therapy is indicated as set out in section 4.1 and:
- who have initially normal leukocyte findings [WBC count ≥ 3500/mm³ (3.5x10⁹/L)] and ANC ≥ 2000/mm³ (2.0x10⁹/L), and
- in whom regular WBC counts and ANC can be performed weekly for the first 18 weeks and at least 4-week intervals thereafter. Monitoring methods must continue throughout treatment and for 4 weeks after complete discontinuation of clozapine.

Before initiating clozapine therapy patients should have a blood test [see “agranulocytosis”) and a history and physical examination. Patients with history of cardiac illness or abnormal cardiac findings on physical examination should be referred to a specialist for further examinations that might include an ECG, and the patient treated only if the expected benefits clearly outweigh the risks [see section 4.3]. The treating physician should consider performing a pre-treatment ECG.

Prescribing physicians must comply fully with the required safety measures.

Prior to treatment initiation, physicians must ensure, to the best of their knowledge, that the patient has not previously experienced an adverse haematological reaction to clozapine that necessitated its discontinuation. Prescriptions should not be issued for periods longer than the interval between two blood counts.

Immediate discontinuation of clozapine is mandatory if either the WBC count is less than 3000/mm³ (3.0x10⁹/L) or the ANC is less than 1500/mm³ (1.5x10⁹/L) at any time during clozapine treatment. Patients in whom clozapine has been discontinued as a result of either WBC or ANC deficiencies must not be re-exposed to clozapine.

At each consultation, a patient receiving clozapine should be reminded to contact the treating physician immediately if any kind of infection begins to develop. Particular attention should be paid to flu-like complaints such as fever or sore throat and to other evidence of infection, which may be indicative of neutropenia. Patients and their caregivers must be informed that, in the event of any of these symptoms, they must have a blood cell count performed immediately. Prescribers are encouraged to keep a record of all patients’ blood results and to take any steps necessary to prevent these patients from accidentally being rechallenged in the future.

Patients with a history of primary bone marrow disorders may be treated only if the benefit outweighs the risk. They should be carefully reviewed by a haematologist prior to starting clozapine.

Patients who have low WBC counts because of benign ethnic neutropenia should be given special consideration and may only be started on clozapine with the agreement of a haematologist.
White Blood Cell (WBC) counts and Absolute Neutrophil Count (ANC) monitoring

WBC and differential blood counts must be performed within 10 days prior to initiating clozapine treatment to ensure that only patients with normal WBC counts and ANC (WBC count ≥ 3500/mm³ [3.5x10⁹/L] and ANC ≥ 2000/mm³ [2.0x10⁹/L]) will receive Zaponex. After the start of clozapine treatment regular WBC count and ANC must be assessed and monitored weekly for the first 18 weeks, and at least at four-week intervals thereafter.

Monitoring must continue throughout treatment and for 4 weeks after complete discontinuation of clozapine or until haematological recovery has occurred (see “Low WBC count/ANC” below). At each consultation, the patient must be reminded to contact the treating physician immediately if any kind of infection, fever, sore throat or other flu-like symptoms develop. WBC and differential blood counts must be performed immediately if any symptoms or signs of an infection occur.

Low WBC count/ANC

If, during clozapine therapy, either the WBC count falls to between 3500/mm³ [3.5x10⁹/L] and 3000/mm³ [3.0x10⁹/L] or the ANC falls to between 2000/mm³ [2.0x10⁹/L] and 1500/mm³ [1.5x10⁹/L], haematological evaluations must be performed at least twice weekly until the patient’s WBC count and ANC stabilise within the range 3000-3500/mm³ [3.0-3.5x10⁹/L] and 1500-2000/mm³ [1.5-2.0x10⁹/L], respectively, or higher.

Immediate discontinuation of clozapine treatment is mandatory if either the WBC count is less than 3000/mm³ [3.0x10⁹/L] or the ANC is less than 1500/mm³ [1.5x10⁹/L] during clozapine treatment. WBC counts and differential blood counts should then be performed daily and patients should be carefully monitored for flu-like symptoms or other symptoms suggestive of infection. Confirmation of the haematological values is recommended by performing two blood counts on two consecutive days; however, clozapine should be discontinued after the first blood count.

Following discontinuation of clozapine, haematological evaluation is required until haematological recovery has occurred.

<table>
<thead>
<tr>
<th>Blood cell count /mm³ (x10⁹/L)</th>
<th>Action required</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC</td>
<td>ANC</td>
</tr>
<tr>
<td>≥ 3500 (≥ 3.5)</td>
<td>≥ 2000 (≥ 2.0)</td>
</tr>
<tr>
<td>3000-3500 (3.0-3.5)</td>
<td>1500-2000 (1.5-2.0)</td>
</tr>
<tr>
<td>&lt; 3000 (&lt; 3.0)</td>
<td>&lt; 1500 (&lt; 1.5)</td>
</tr>
</tbody>
</table>

If clozapine has been withdrawn and either a further drop in the WBC count below 2000/mm³ [2.0x10⁹/L] occurs or the ANC falls below 1000/mm³ [1.0x10⁹/L], the management of this condition must be guided by an experienced haematologist.

Discontinuation of therapy for haematological reasons

Patients in whom clozapine has been discontinued as a result of either WBC or ANC deficiencies (see table 1) must not be re-exposed to clozapine.

Prescribers are encouraged to keep a record of all patients’ blood results and to take any steps necessary to prevent the patient being accidentally rechallenged in the future.

Discontinuation of therapy for other reasons

Patients who have been on clozapine for more than 18 weeks and have had their treatment interrupted for more than 3 days but less than 4 weeks should have their WBC count and ANC monitored weekly for an additional 6 weeks. If no haematological abnormality occurs, monitoring at intervals not exceeding 4 weeks may be resumed. If clozapine treatment has been interrupted for 4 weeks or longer, weekly monitoring is required for the next 18 weeks of treatment and the dose should be re-titrated (see section 4.2).

Other precautions

This medicinal product contains lactose monohydrate.

Patients with rare hereditary problems of galactose intolerance, the Lapp lactase deficiency or glucose galactose malabsorption should not take this medicine.

Eosinophilia

In the event of eosinophilia, discontinuation of clozapine is recommended if the eosinophil count rises above 3000/mm³ [3.0x10⁹/L]; therapy should be restarted only after the eosinophil count has fallen below 1000/mm³ [1.0x10⁹/L].

Thrombocytopenia

In the event of thrombocytopenia, discontination of clozapine therapy is recommended if the platelet count falls below 50 000/mm³ [50x10⁹/L].

Cardiovascular disorders

Orthostatic hypotension, with or without syncope, can occur during clozapine treatment. Rarely, collapse can be profound and may be accompanied by cardiac and/or respiratory arrest. Such events are more likely to occur with concurrent use of benzodiazepine or any other psychotropic agent [see section 4.5] and during initial titration in association with rapid dose escalation; on very rare occasions they may occur even after the first dose. Therefore, patients commencing clozapine treatment require close medical supervision. Monitoring of standing and supine blood pressure is necessary during the first weeks of treatment in patients with Parkinson’s disease.

Analysis of safety databases suggests that the use of clozapine is associated with an increased risk of myocarditis especially during, but not limited to, the first two months of treatment. Some cases of myocarditis have been fatal. Pericarditis/pericardial effusion and cardiomyopathy have also been reported in association with clozapine use; these reports also include fatalities. Myocarditis or cardiomyopathy should be suspected in patients who experience persistent tachycardia at rest, especially in the first two months of treatment, and/or palpitations, arrhythmias, chest pain and other signs and symptoms of heart failure (e.g., unexplained fatigue, dyspnoea, tachynoea), or symptoms that mimic myocardial infarction. Other symptoms which may be present in addition to the above include flu-like symptoms. If myocarditis or cardiomyopathy is suspected, clozapine treatment should be promptly stopped and the patient immediately referred to a cardiologist.

Patients with clozapine-induced myocarditis or cardiomyopathy should not be re-exposed to clozapine.
Myocardial infarction
In addition, there have been post marketing reports of myocardial infarction which may be fatal. Causality assessment was difficult in the majority of these cases because of serious pre-existing cardiac disease and plausible alternative causes.

QT interval prolongation
As with other antipsychotics, caution is advised in patients with known cardiovascular disease or family history of QT prolongation. As with other antipsychotics, caution should be exercised when clozapine is prescribed with medicines known to increase QTc interval.

Cerebrovascular adverse events
An approximately 3-fold increased risk of cerebrovascular adverse events has been seen in randomised placebo controlled clinical trials in the dementia population with some atypical antipsychotics. The mechanism for this increased risk is not known. An increased risk cannot be excluded for other antipsychotics or other patient populations. Clozapine should be used with caution in patients with risk factors for stroke.

Risk of thromboembolism
Since clozapine may be associated with thromboembolism, immobilisation of patients should be avoided. Cases of venous thromboembolism (VTE) have been reported with antipsychotic drugs. Since patients treated with antipsychotics often present with acquired risk factors for VTE, all possible risk factors for VTE should be identified before and during treatment with clozapine and preventive measures undertaken.

Seizures
Patients with a history of epilepsy should be closely observed during clozapine therapy since dose-related convulsions have been reported. In such cases, the dose should be reduced (see section 4.2) and, if necessary, an anti-convulsant treatment should be initiated.

Anticholinergic effects
Clozapine exerts anticholinergic activity, which may produce undesirable effects throughout the body. Careful supervision is indicated in the presence of prostatic enlargement and narrow-angle glaucoma. Probably on account of its anticholinergic properties, clozapine has been associated with varying degrees of impairment of intestinal peristalsis, ranging from constipation to intestinal obstruction, faecal impaction and paralytic ileus (see section 4.8). On rare occasions these cases have been fatal. Particular care is necessary in patients who are receiving concomitant medications known to cause constipation (especially those with anticholinergic properties such as some antipsychotics, antidepressants and anti-parkinsonian treatments), have a history of colonic disease or a history of lower abdominal surgery as these may exacerbate the situation. It is vital that constipation is recognised and actively treated.

Fever
During clozapine therapy, patients may experience transient temperature elevations above 38°C, with the peak incidence within the first 3 weeks of treatment. This fever is generally benign. Occasionally, it may be associated with an increase or decrease in the WBC count. Patients with fever should be carefully evaluated to rule out the possibility of an underlying infection or the development of agranulocytosis. In the presence of high fever, the possibility of neuroleptic malignant syndrome (NMS) must be considered. If the diagnosis of NMS is confirmed, clozapine should be discontinued immediately and appropriate medical measures should be administered.

Metabolic changes
Atypical antipsychotic drugs, including clozapine, have been associated with metabolic changes that may increase cardiovascular/ cerebrovascular risk. These metabolic changes may include hyperglycaemia, dyslipidemia, and body weight gain. While atypical antipsychotic drugs may produce some metabolic changes, each drug in the class has its own specific profile.

Hyperglycaemia
Impaired glucose tolerance and/or development or exacerbation of diabetes mellitus has been reported rarely during treatment with clozapine. A mechanism for this possible association has not yet been determined. Cases of severe hyperglycaemia with ketoacidosis or hyperosmolar coma have been reported very rarely in patients with no prior history of hyperglycaemia, some of which have been fatal. When follow-up data were available, discontinuation of clozapine resulted most often in resolution of the impaired glucose tolerance, and reinstitution of clozapine resulted in its reoccurrence. Patients with an established diagnosis of diabetes mellitus who are started on atypical antipsychotics should be monitored regularly for worsening of glucose control. Patients with risk factors for diabetes mellitus (e.g. obesity, family history of diabetes) who are starting treatment with atypical antipsychotics should undergo fasting blood glucose testing at the beginning of treatment and periodically during treatment. Patients who develop symptoms of hyperglycaemia during treatment with atypical antipsychotics should undergo fasting blood glucose testing. In some cases hyperglycaemia has resolved when the atypical antipsychotic was discontinued; however, some patients required continuation of antidiabetic treatment despite discontinuation of the suspect drug. The discontinuation of clozapine should be considered in patients where active medical management of their hyperglycaemia has failed.

Dyslipidemia
Undesirable alterations in lipids have been observed in patients treated with atypical antipsychotics including clozapine. Clinical monitoring, including baseline and periodic follow-up lipid evaluations in patients using clozapine is recommended.

Weight gain
Weight gain has been observed with atypical antipsychotic use, including clozapine. Clinical monitoring of weight is recommended.

Rebound withdrawal effects
Acute withdrawal reactions have been reported following abrupt cessation of clozapine therefore gradual withdrawal is recommended. If abrupt discontinuation is necessary (e.g. because of leukopenia), the patient should be carefully observed for the recurrence of psychotic symptoms and symptoms related to cholinergic rebound, such as profuse sweating, headache, nausea, vomiting and diarrhoea.

Special populations
Hepatic impairment
Patients with stable pre-existing liver disorders may receive clozapine, but need regular liver function tests. Liver function tests should be performed in patients in whom symptoms of possible liver dysfunction, such as nausea, vomiting and/or anorexia, develop during clozapine therapy. If the elevation of the values is clinically relevant (more than 3 times the UNL) or if symptoms of jaundice occur, treatment with clozapine must be discontinued. It may be resumed (see "Re-starting therapy" under section 4.2) only when the results of liver function tests are normal. In such cases, liver function should be closely monitored after re-introduction of the drug.
Patients aged 60 years and older
Initiation of treatment in patients aged 60 years and older is recommended at a lower dose (see section 4.2).
Orthostatic hypotension can occur with clozapine treatment and there have been reports of tachycardia, which may be sustained. Patients aged 60 years and older, particularly those with compromised cardiovascular function, may be more susceptible to these effects.

Increased mortality in elderly people with dementia
Data from two large observational studies showed that elderly people with dementia who are treated with antipsychotics are at a small increased risk of death compared with those who are not treated. There are insufficient data to give a firm estimate of the precise magnitude of the risk and the cause of the increased risk is not known.

Zaponex is not approved for the treatment of dementia-related behavioural disturbances.

4.5 Interaction with other medicinal products
and other forms of interaction

Contraindication of concomitant use
Substances known to have a substantial potential to depress bone marrow function must not be used concurrently with clozapine (see section 4.3).

Long-acting depot antipsychotics (which have myelosuppressive potential) must not be used concurrently with clozapine because these cannot be rapidly removed from the body in situations where this may be required, e.g. neutropenia (see section 4.3).

Alcohol should not be used concomitantly with clozapine due to possible potentiation of sedation.

Precautions including dose adjustment
Clozapine may enhance the central effects of CNS depressants such as narcotics, antihistamines, and benzodiazepines. Particular caution is advised when clozapine therapy is initiated in patients who are receiving a benzodiazepine or any other psychotropic substance. These patients may have an increased risk of circulatory collapse, which, on rare occasions, can be profound and may lead to cardiac arrest and/or respiratory arrest. It is not clear whether cardiac or respiratory collapse can be prevented by dose adjustment.

Because of the possibility of additive effects, caution is essential in the concomitant administration of substances possessing anticholinergic, hypotensive, or respiratory depressant effects.

Owing to its anti-alpha-adrenergic properties, clozapine may reduce the blood-pressure-increasing effect of norepinephrine or other predominantly alpha-adrenergic agents and reverse the pressor effect of epinephrine.

Concomitant administration of substances known to inhibit the activity of some cytochrome P450 isozymes may increase the levels of clozapine, and the dose of clozapine may need to be reduced to prevent undesirable effects. This is more important for CYP 3A4 inhibitors such as azole antifungicals, cimetidine, erythromycin, and protease inhibitors are unlikely, although some have been reported. Because the plasma concentration of clozapine is increased by caffeine intake and decreased by nearly 50% following a 5-day caffeine-free period, dosage changes of clozapine may be necessary when there is a change in caffeine-drinking habit. In cases of sudden cessation of smoking, the plasma clozapine concentration may be increased, thus leading to an increase in adverse effects.

Cases have been reported of an interaction between citalopram and clozapine, which may increase the risk of adverse events associated with clozapine. The nature of this interaction has not been fully elucidated.

Concomitant administration of substances known to induce cytochrome P450 enzymes may decrease the plasma levels of clozapine, leading to reduced efficacy. Substances known to induce the activity of cytochrome P450 enzymes and with reported interactions with clozapine include, for instance, carbamazepine (not to be used concomitantly with clozapine, due to its myelosuppressive potential), phenytoin and rifampicin. Known inducers of CYP1A2, such as omeprazole, may lead to decreased clozapine levels. The potential for reduced efficacy of clozapine should be considered when it is used in combination with these substances.

Other
Concomitant use of lithium or other CNS-active agents may increase the risk of development of neuroleptic malignant syndrome (NMS).

Rare but serious reports of seizures, including onset of seizures in non-epileptic patients, and isolated cases of delirium where clozapine was co-administered with valproic acid have been reported. These effects are possibly due to a pharmacodynamic interaction, the mechanism of which has not been determined.

Caution is called for in patients receiving concomitant treatment with other substances which are either inhibitors or inducers of the cytochrome P450 isozymes. With tricyclic antidepressants, phenothiazines and type 1 anti-arrhythmics, which are known to bind to cytochrome P450 2D6, no clinically relevant interactions have been observed thus far.

As with other antipsychotics, caution should be exercised when clozapine is prescribed with medicines known to increase QTc interval, or causing electrolyte imbalance.

An outline of drug interactions believed to be most important with clozapine is given in Table 2 (See next page). The list is not exhaustive.
Table 2: Reference to the most common drug interactions with clozapine

<table>
<thead>
<tr>
<th>Drug</th>
<th>Interactions</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone marrow suppressants (e.g. carbamazepine, chloramphenicol, sulphonamides (e.g. co trimoxazole), pyrazolone analgesics (e.g. phenylbutazone), penicillamine, cytotoxic agents and long-acting depot injections of antipsychotics)</td>
<td>Interact to increase the risk and/or severity of bone marrow suppression</td>
<td>Clozapine <strong>must not be used</strong> concomitantly with other agents having a well known potential to suppress bone marrow function (see section 4.3)</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>Concomitant use may increase risk of circulatory collapse, which may lead to cardiac and/or respiratory arrest</td>
<td>Whilst the occurrence is rare, caution is advised when using these agents together. Reports suggest that respiratory depression and collapse are more likely to occur at the start of this combination or when clozapine is added to an established benzodiazepine regimen</td>
</tr>
<tr>
<td>Anticholinergics</td>
<td>Clozapine potentiates the action of these agents through additive anticholinergic activity</td>
<td>Observe patients for anticholinergic side-effects, e.g. constipation, especially when using to help control hypersalivation</td>
</tr>
<tr>
<td>Antihypertensives</td>
<td>Clozapine can potentiate the hypotensive effects of these agents due to its sympathomimetic antagonistic effects</td>
<td>Caution is advised if clozapine is used concomitantly with antihypertensive agents. Patients should be advised of the risk of hypotension, especially during the period of initial dose titration</td>
</tr>
<tr>
<td>Alcohol, MAOIs, CNS depressants, including narcotics and benzodiazepines</td>
<td>Enhanced central effects. Additive CNS depression and cognitive and motor performance interference when used in combination with these substances</td>
<td>Caution is advised if clozapine is used concomitantly with other CNS active agents. Advise patients of the possible additive sedative effects and caution them not to drive or operate machinery</td>
</tr>
<tr>
<td>Highly protein bound substances (e.g. warfarin and digoxin)</td>
<td>Clozapine may cause an increase in plasma concentration of these substances due to displacement from plasma proteins</td>
<td>Patients should be monitored for the occurrence of side effects associated with these substances, and doses of the protein bound substance adjusted, if necessary</td>
</tr>
<tr>
<td>Phenytoin</td>
<td>Addition of phenytoin to clozapine regimen may cause a decrease in the clozapine plasma concentrations</td>
<td>If phenytoin must be used, the patient should be monitored closely for a worsening or recurrence of psychotic symptoms</td>
</tr>
<tr>
<td>Lithium</td>
<td>Concomitant use can increase the risk of development of neuroleptic malignant syndrome (NMS)</td>
<td>Observe for signs and symptoms of NMS</td>
</tr>
<tr>
<td>CYP1A2 inducing substances (e.g. omeprazole)</td>
<td>Concomitant use may decrease clozapine levels</td>
<td>Potential for reduced efficacy of clozapine should be considered</td>
</tr>
<tr>
<td>CYP1A2 inhibiting substances (e.g. fluvoxamine, caffeine, ciprofloxacin)</td>
<td>Concomitant use may increase clozapine levels</td>
<td>Potential for increase in adverse Care is also required upon cessation of concomitant CYP1A2 inhibiting medications as there will be a decrease in clozapine levels</td>
</tr>
</tbody>
</table>
4.6 Fertility, pregnancy and lactation

Pregnancy
For clozapine, there are only limited clinical data on exposed pregnancies. Animal studies do not indicate direct or indirect harmful effects with respect to pregnancy, embryonal/foetal development, parturition or postnatal development (see section 5.3). Caution should be exercised when prescribing to pregnant women.

Breastfeeding
Animal studies suggest that clozapine is excreted in breast milk and has an effect in the nursing infant; therefore, mothers receiving clozapine should not breast-feed.

Women of child-bearing potential
A return to normal menstruation may occur as a result of switching from other antipsychotics to clozapine. Adequate contraceptive measures must therefore be ensured in women of childbearing potential.

4.7 Effects on ability to drive and use machines

Owing to the ability of clozapine to cause sedation and lower the seizure threshold, activities such as driving or operating machinery should be avoided, especially during the initial weeks of treatment.

4.8 Undesirable effects

Summary of the safety profile
For the most part, the adverse event profile of clozapine is predictable from its pharmacological properties. An important exception is its propensity to cause agranulocytosis (see section 4.4). Because of this risk, its use is restricted to treatment-resistant schizophrenia and psychosis occurring during the course of Parkinson’s disease in cases where standard treatment has failed. While blood monitoring is an essential part of the care of patients receiving clozapine, the physician should be aware of other rare but serious adverse reactions, which may be diagnosed in the early stages only by careful observation and questioning of the patient in order to prevent morbidity and mortality.

The most serious adverse reactions experienced with clozapine are agranulocytosis, seizures, cardiovascular effects and fever (see section 4.4). The most common side effects are drowsiness-sedation, and dizziness. The very common adverse reactions observed include drowsiness/sedation, and dizziness.

Blood and lymphatic system
Development of granulocytopenia and agranulocytosis is a risk inherent to clozapine treatment. Although generally reversible on withdrawal of treatment, agranulocytosis may result in sepsis and can prove fatal. Because immediate withdrawal of treatment is required to prevent the development of life-threatening agranulocytosis, monitoring of the WBC count is mandatory (see section 4.4). Table 3 below summarises the estimated incidence of agranulocytosis for each clozapine treatment period.

<table>
<thead>
<tr>
<th>Treatment period</th>
<th>Incidence of agranulocytosis per 100,000 person-weeks$^2$ of observation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weeks 0-18</td>
<td>32.0</td>
</tr>
<tr>
<td>Weeks 19-52</td>
<td>2.3</td>
</tr>
<tr>
<td>Weeks 53 and higher</td>
<td>1.8</td>
</tr>
</tbody>
</table>

(2) Person-time is the sum of individual units of time that the patients in the registry were exposed to clozapine before experiencing agranulocytosis. For example, 100,000 person-weeks could be observed in 1,000 patients who were in the registry for 100 weeks [100*1000=100,000], or in 200 patients who were in the registry for 500 weeks [200*500=100,000] before experiencing agranulocytosis.

The cumulative incidence of agranulocytosis in the UK Clozaril Patient Monitoring Service lifetime registry experience (0 - 11.6 years between 1989 and 2001) is 0.78%. The majority of cases (approximately 70%) occur within the first 18 weeks of treatment.

Metabolic and Nutritional Disorders
Impaired glucose tolerance and/or development or exacerbation of diabetes mellitus has been reported rarely during treatment with clozapine. On very rare occasions, severe hyperglycaemia, sometimes leading to ketoacidosis/hyperosmolar coma, has been reported in patients on clozapine treatment with no prior history of hyperglycaemia. Glucose levels normalised in most patients after discontinuation of clozapine and in a few cases hyperglycaemia recurred when treatment was reinitiated. Although most patients had risk factors for non-insulin-dependent diabetes mellitus, hyperglycaemia has also been documented in patients with no known risk factors (see section 4.4).

Nervous System Disorders
The very common adverse reactions observed include drowsiness/sedation, and dizziness.

Clozapine can cause EEG changes, including the occurrence of spike and wave complexes. It lowers the seizure threshold in a dose-dependent manner and may induce myoclonic jerks or generalised seizures. These symptoms are more likely to occur with rapid dose increases and in patients with pre-existing epilepsy. In such cases the dose should be reduced and, if necessary, anticonvulsant treatment initiated. Carbamazepine should be avoided because of its potential to depress bone marrow function, and with other anticonvulsant agents the possibility of a pharmacokinetic interaction should be considered. In rare cases, patients treated with clozapine may experience delirium.

Very rarely, tardive dyskinesia has been reported in patients on clozapine who had been treated with other antipsychotic agents. Patients in whom tardive dyskinesia developed with other antipsychotics have improved on clozapine.

Cardiac Disorders
Tachycardia and postural hypotension with or without syncope may occur, especially in the initial weeks of treatment. The prevalence and severity of hypotension is influenced by the rate and magnitude of dose titration. Circulatory collapse as a result of profound hypotension, in particular related to aggressive titration, with the possible serious consequences of cardiac or pulmonary arrest, has been reported with clozapine.
A minority of clozapine-treated patients experience ECG changes similar to those seen with other antipsychotics, including S-T segment depression and flattening or inversion of T waves, which normalise after discontinuation of clozapine. The clinical significance of these changes is unclear. However, such abnormalities have been observed in patients with myocarditis, which should therefore be considered.

Isolated cases of cardiac arrhythmias, pericarditis/pericardial effusion and myocarditis have been reported, some of which have been fatal. The majority of the cases of myocarditis occurred within the first 2 months of initiation of therapy with clozapine. Cardiomyopathy generally occurred later in the treatment.

Eosinophilia has been co-reported with some cases of myocarditis (approximately 14%) and pericarditis/pericardial effusion; it is not known, however, whether eosinophilia is a reliable predictor of carditis.

Signs and symptoms of myocarditis or cardiomyopathy include persistent tachycardia at rest, palpitations, arrhythmias, chest pain and other signs and symptoms of heart failure (e.g. unexplained fatigue, dyspnoea, tachypnoea), or symptoms that mimic myocardial infarction. Other symptoms which may be present in addition to the above include flu-like symptoms.

Sudden, unexplained deaths are known to occur among psychiatric patients who receive conventional antipsychotic medication but also among untreated psychiatric patients. Such deaths have been reported very rarely in patients receiving clozapine.

**Vascular Disorders**
Rare cases of thromboembolism have been reported.

**Respiratory System**
Respiratory depression or arrest has occurred very rarely, with or without circulatory collapse [see sections 4.4 and 4.5].

**Gastrointestinal System**
Constipation and hypersalivation have been observed very frequently, and nausea and vomiting frequently. Very rarely ileus may occur [see section 4.4]. Rarely clozapine treatment may be associated with dysphagia. Aspiration of ingested food may occur in patients presenting with dysphagia or as a consequence of acute overdose.

**Hepatobiliary Disorders**
Transient, asymptomatic elevations of liver enzymes and rarely, hepatitis and cholestatic jaundice may occur. Very rarely, fulminant hepatic necrosis has been reported. If jaundice develops, clozapine should be discontinued [see section 4.4]. In rare cases, acute pancreatitis has been reported.

**Renal Disorders**
Isolated cases of acute interstitial nephritis have been reported in association with clozapine therapy.

**Reproductive and Breast Disorders**
Very rare reports of priapism have been received.

**General Disorders**
Cases of neuroleptic malignant syndrome (NMS) have been reported in patients receiving clozapine either alone or in combination with lithium or other CNS-active agents.

Acute withdrawal reactions have been reported [see section 4.4].

**Tabulated list of adverse reactions**
The table below [Table 4] summarises the adverse reactions accumulated from reports made spontaneously and during clinical studies.

<table>
<thead>
<tr>
<th>Blood and lymphatic system disorders</th>
<th>Common</th>
<th>Leukopenia/decreased WBC/neutropenia, eosinophilia, leukocytosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncommon</td>
<td></td>
<td>Agranulocytosis</td>
</tr>
<tr>
<td>Rare</td>
<td></td>
<td>Anaemia</td>
</tr>
<tr>
<td>Very rare</td>
<td></td>
<td>Thrombocytopenia, thrombocythaemia</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Metabolism and nutrition disorders</th>
<th>Common</th>
<th>Weight gain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rare</td>
<td></td>
<td>Impaired glucose tolerance, diabetes mellitus</td>
</tr>
<tr>
<td>Very rare</td>
<td></td>
<td>Ketoacidosis, hyperosmolar coma, severe hyperglycaemia, hypertriglyceridaemia, hypercholesterolaemia</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Psychiatric disorders</th>
<th>Common</th>
<th>Dysarthria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncommon</td>
<td></td>
<td>Dysphemia</td>
</tr>
<tr>
<td>Rare</td>
<td></td>
<td>Restlessness, agitation</td>
</tr>
<tr>
<td><strong>Nervous system disorders</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------------------------------</td>
<td></td>
</tr>
<tr>
<td><strong>Very common</strong></td>
<td>Drowsiness/sedation, dizziness</td>
<td></td>
</tr>
<tr>
<td><strong>Common</strong></td>
<td>Headache, tremor, rigidity, akathisia, extrapyramidal symptoms, seizures/convulsions/myoclonic jerks</td>
<td></td>
</tr>
<tr>
<td><strong>Uncommon</strong></td>
<td>Neuroleptic malignant syndrome</td>
<td></td>
</tr>
<tr>
<td><strong>Rare</strong></td>
<td>Confusion, delirium</td>
<td></td>
</tr>
<tr>
<td><strong>Very rare</strong></td>
<td>Tardive dyskinesia, obsessive compulsive symptoms</td>
<td></td>
</tr>
<tr>
<td><strong>Not known</strong></td>
<td>Cholinergic syndrome (after abrupt withdrawal)<em>, EEG changes</em></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Eye disorders</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Common</strong></td>
<td>Blurred vision</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Cardiac disorders</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Very common</strong></td>
<td>Tachycardia</td>
</tr>
<tr>
<td><strong>Common</strong></td>
<td>ECG changes</td>
</tr>
<tr>
<td><strong>Rare</strong></td>
<td>Circulatory collapse, arrhythmias, myocarditis, pericarditis/pericardial effusion</td>
</tr>
<tr>
<td><strong>Very rare</strong></td>
<td>Cardiomyopathy, cardiac arrest</td>
</tr>
<tr>
<td><strong>Not known</strong></td>
<td>Myocardial infarction which may be fatal*, chest pain/angina pectoris*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Vascular disorders</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Common</strong></td>
<td>Hypertension, postural hypotension, syncope</td>
</tr>
<tr>
<td><strong>Rare</strong></td>
<td>Thromboembolism</td>
</tr>
<tr>
<td><strong>Not known</strong></td>
<td>Venous thromboembolism, including cases of pulmonary embolism and cases of deep vein thrombosis</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Respiratory, thoracic and mediastinal disorders</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Rare</strong></td>
<td>Aspiration of ingested food, pneumonia and lower respiratory tract infection which may be fatal</td>
</tr>
<tr>
<td><strong>Very rare</strong></td>
<td>Respiratory depression/arrest</td>
</tr>
<tr>
<td><strong>Not known</strong></td>
<td>Nasal congestion*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Gastrointestinal disorders</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Very common</strong></td>
<td>Constipation, hypersalivation</td>
</tr>
<tr>
<td><strong>Common</strong></td>
<td>Nausea, vomiting, anorexia, dry mouth</td>
</tr>
<tr>
<td><strong>Rare</strong></td>
<td>Dysphagia</td>
</tr>
<tr>
<td><strong>Very rare</strong></td>
<td>Parotid gland enlargement, intestinal obstruction/paralytic ileus/faecal impaction</td>
</tr>
<tr>
<td><strong>Not known</strong></td>
<td>Diarrhoea*, abdominal discomfort/heartburn/dyspepsia*</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Hepatobiliary disorders</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Common</strong></td>
<td>Elevated liver enzymes</td>
</tr>
<tr>
<td><strong>Rare</strong></td>
<td>Hepatitis, cholestatic jaundice, pancreatitis</td>
</tr>
<tr>
<td><strong>Very rare</strong></td>
<td>Fulminant hepatic necrosis</td>
</tr>
</tbody>
</table>
Very rare events of ventricular tachycardia and QT prolongation which may be associated with Torsades De Pointes have been observed although there is no conclusive causal relationship to the use of this medicine.

**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.

### 4.9 Overdose

In cases of acute intentional or accidental clozapine overdose for which information on the outcome is available, mortality to date is about 12%. Most of the fatalities were associated with cardiac failure or pneumonia caused by aspiration and occurred at doses above 2000 mg. There have been reports of patients recovering from an overdose in excess of 10,000 mg. However, in a few adult individuals, primarily those not previously exposed to clozapine, the ingestion of doses as low as 400 mg led to life-threatening comatose conditions and, in one case, to death. In young children, the intake of 50 to 200 mg resulted in strong sedation or coma without being lethal.

**Signs and symptoms**
Drowsiness, lethargy, areflexia, coma, confusion, hallucinations, agitation, delirium, extrapyramidal symptoms, hyperreflexia, convulsions; hypersalivation, mydriasis, blurred vision, thermolability; hypotension, collapse, tachycardia, cardiac arrhythmias; aspiration pneumonia, dyspnoea, respiratory depression or failure.

**Treatment**
There are no specific antidotes for clozapine. Gastric lavage and/or administration of activated charcoal within the first 6 hours after the ingestion of the drug. Peritoneal dialysis and haemodialysis are unlikely to be effective. Symptomatic treatment under continuous cardiac monitoring, surveillance of respiration, monitoring of electrolytes and acid-base balance. The use of epinephrine should be avoided in the treatment of hypotension because of the possibility of a ‘reverse epinephrine’ effect.

Close medical supervision is necessary for at least 5 days because of the possibility of delayed reactions.
5. Pharmacological Properties

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Antipsychotics, diazepines, oxazepines and thiapropenes [ATC code N05A H02]

Clozapine has been shown to be an antipsychotic agent that is different from classic antipsychotics.

In pharmacological experiments, the compound does not induce catalepsy or inhibit apomorphine- or amphetamine-induced stereotyped behaviour. It has only weak dopamine-receptor-blocking activity at D₁, D₂, D₃ and D₄ receptors, but shows high potency for the D₂ receptor, in addition to potent anti-alpha-adrenergic, anticholinergic, antihistaminic, and arousal-reaction-inhibiting effects. It has also been shown to possess antiserotonergic properties.

Clinically, clozapine produces rapid and marked sedation and exerts antipsychotic effects in schizophrenic patients resistant to other drug treatment. In such cases, clozapine has proven effective in relieving both positive and negative schizophrenic symptoms mainly in short-term trials. In an open clinical trial performed in 319 treatment-resistant patients treated for 12 months, a clinically relevant improvement was observed in 37% of patients within the first week of treatment and in an additional 44% by the end of 12 months. The improvement was defined as about 20% reduction from baseline in Brief Psychiatric Rating Scale Score. In addition, improvement in some aspects of cognitive dysfunction has been described.

Compared to classic antipsychotics, clozapine produces fewer major extrapyramidal reactions such as acute dystonia, parkinsonian-like side effects and akathisia. In contrast to classic antipsychotics, clozapine produces little or no prolactin elevation, thus avoiding adverse effects such as gynaecomastia, amenorrhoea, galactorrhoea, and impotence.

A potentially serious adverse reaction caused by clozapine therapy is granulocytopenia and agranulocytosis occurring at an estimated incidence of 3% and 0.7%, respectively. In view of this risk, the use of clozapine should be limited to patients who are treatment-resistant or patients with psychosis in Parkinson's disease when other treatment strategies have failed (see section 4.1) and in whom regular haematological examinations can be performed (see sections 4.4 and 4.8).

5.2 Pharmacokinetic properties

Absorption

The absorption of orally administered clozapine is 90 to 95%; neither the rate nor the extent of absorption is influenced by food.

Clozapine is subject to moderate first-pass metabolism, resulting in an absolute bioavailability of 50 to 60%.

Distribution

In steady-state conditions, when given twice daily, peak blood levels occur on an average at 2.1 hours (range: 0.4 to 4.2 hours), and the volume of distribution is 1.6 l/kg. Clozapine is approximately 95% bound to plasma proteins.

Biotransformation/metabolism

Clozapine is almost completely metabolised before excretion. Of the main metabolites only the dimethyl metabolite was found to be active. Its pharmacological actions resemble those of clozapine, but are considerably weaker and of short duration.

Elimination

Its elimination is biphasic, with a mean terminal half-life of 12 hours (range: 6 to 26 hours). After single doses of 75 mg the mean terminal half-life was 7.9 hours; it increased to 14.2 hours when steady-state conditions were reached by administering daily doses of 75 mg for at least 7 days.

Only trace amounts of unchanged drug are detected in the urine and faeces, approximately 50% of the administered dose being excreted as metabolites in the urine and 30% in the faeces.

Linearity/non-linearity

Dosage increases from 37.5 mg to 75 mg and 150 mg given twice daily were found to result during steady state in linearly dose-proportional increases in the area under the plasma concentration/time curve (AUC), and in the peak and minimum plasma concentrations.

5.3 Preclinical safety data

Preclinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity and carcinogenic potential (for reproductive toxicity, see section 4.6).

6. Pharmaceutical particulars

6.1 List of excipients

Lactose monohydrate
Povidone
Pregelatinised starch
Maize starch
Talc
Colloidal anhydrous silica
Magnesium stearate

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

Zaponex 25 mg Tablets in blister packaging: 4 years
Zaponex 100 mg Tablets in blister packaging: 5 years
Zaponex 25 and 100 mg Tablets in HDPE containers: 2 years

6.4 Special precautions for storage

For the HDPE container:
Do not store above 25 °C. Keep the plastic container tightly closed. Store in the original packaging.

For the blister packs:
Do not store above 25 °C. Store in the original packaging. Keep blister in the outer carton.

6.5 Nature and contents of container

Zaponex Tablets are packaged in PVC/PVDC/aluminium blisters containing 10 or 14 tablets and per 500 tablets in a HDPE container. The blisters containing 10 tablets are available in pack sizes of 30, 60, 90 and 300 tablets, the blisters containing 14 tablets in pack sizes of 28 and 84 tablets. The blisters are packaged in a lithographed carton box, with a patient information leaflet enclosed.

Not all pack sizes may be marketed.
6.6 Special precautions for disposal and other handling

No special requirements.

7. Marketing Authorisation Holder

Leyden Delta BV
Neerbosscheweg 620
6544 LL Nijmegen
The Netherlands

8. Marketing Authorisation Number(s)

Zaponex 25 mg Tablets: PL 32553/0001
Zaponex 100 mg Tablets: PL 32553/0002

9. Date of first authorisation/renewal of the authorisation

Date of Licence Granted: 18 June 2004
Date of Last Renewal: 18 June 2009

10. Date of revision of the text

July 2015