PRODUCT INFORMATION

NAME OF THE MEDICINE

Daonil® and Semi-Daonil®

AUSTRALIAN APPROVED NAME

Glibenclamide.

CHEMICAL STRUCTURE

Glibenclamide belongs to the sulphonylurea group of oral antidiabetics. Earlier members of this group are carbutamide, tolbutamide, acetohexamide and chlorpropamide. Chemically, it is 1-\{4 - [2 - (5 -chloro-2-methoxy-benzamido) ethyl] benzenesulphonyl\} - 3 - cyclohexylurea. It has a molecular weight of 494 and an empirical formula of C\textsubscript{23}H\textsubscript{28}ClN\textsubscript{3}O\textsubscript{5}S. It is a white odourless, crystalline powder, practically insoluble in water and in ether, slightly soluble in alcohol and sparingly soluble in chloroform.

CAS REGISTRY NUMBER

10238-21-8

DESCRIPTION

Daonil

Each tablet contains 5 mg of glibenclamide. Excipients are lactose monohydrate, maize starch, pregelatinised maize starch, purified talc, colloidal anhydrous silica and magnesium stearate.

Semi-Daonil®

Each tablet contains 2.5 mg of glibenclamide. Excipients are lactose monohydrate, maize starch, purified talc, colloidal anhydrous silica and magnesium stearate.
Daonil and Semi-Daonil – Glibenclamide

PHARMACOLOGY

Oral hypoglycaemia.

PHARMACODYNAMICS

Mechanism of Action

Daonil appears to lower the blood glucose acutely in healthy individuals and patients with type 2 diabetes by stimulating the release of insulin from the pancreas, an effect dependent upon functioning beta cells. It acts in concert with glucose (improved sensitivity of beta cells to physiological glucose stimulus) and leads to an insulin secretion in the rhythm of meals. Other mechanisms of the hypoglycaemic action associated with short term therapy appear to include reduction of basal hepatic glucose production and enhancement of peripheral insulin action at post-receptor (probably intracellular) sites.

With chronic administration of Daonil and Semi-Daonil in patients with type 2 diabetes, the improvement in glucose tolerance persists despite a gradual decline in glucose- or meal-stimulated secretion of insulin towards pretreatment levels. Extrapancreatic effects appear to contribute substantially to the hypoglycaemic action of the drug during long term administration. The effects appear to include enhanced peripheral sensitivity to insulin and reduction of basal hepatic glucose production. There is evidence that glibenclamide enhances the peripheral action of insulin at post-receptor (probably intracellular) sites and increases insulin binding and/or the number of insulin receptors.

Glibenclamide also exerts a direct inhibitory effect on glucagon-producing alpha cells of the pancreas and increases the release of somatostatin. However, these two pancreatic extra-beta cell actions may play only a minor clinical role.

In addition to its blood glucose lowering effect, glibenclamide has a mild diuretic action and increases free water clearance.

PHARMACOKINETICS

Absorption

Glibenclamide is nearly completely absorbed (84 + 9%) after oral administration and is extensively bound (99%) to serum proteins. The peak serum concentration is reached in 2-6 hours after taking a 5 mg tablet of Daonil and falls within 24 hours to less than 5% of the peak value. The area under the serum concentration time curve (AUC) increases in proportion to increasing doses. Food apparently does not affect the rate or extent of absorption of glibenclamide.

Distribution

Multiple-dose studies with glibenclamide in diabetic patients demonstrate drug level concentration-time curves similar to single-dose studies, indicating no build-up of drug in tissue depots. In non-fasting diabetic patients, the hypoglycaemic action of a single morning dose of glibenclamide persists for 24 hours.
Serum concentrations of glibenclamide appear to decline in a biphasic manner. The elimination half-life of glibenclamide after intravenous dosage is approximately 2 hours, and 2 to 5 hours after oral administration. Some reports indicate a longer half-life of 8 to 10 hours in patients with diabetes.

**Metabolism**

Glibenclamide is completely metabolised in the liver. The drug is metabolised at the cyclohexyl ring principally to a 4-trans-hydroxy derivative. A second metabolite, the 3-cis-hydroxy derivative, also occurs. These metabolites contribute some hypoglycaemic action; they are weakly active (0.25% and 2.5%, respectively, as glibenclamide) in rabbits.

**Excretion**

Glibenclamide is excreted as metabolites in the bile and urine, approximately 50% by each route. In patients with renal insufficiency, depending on the degree of the renal excretion disorder, there is increased elimination of the metabolites via the bile. This dual excretory pathway is qualitatively different from that of other sulphonylureas, which are excreted primarily in the urine.

Glibenclamide appears to be only minimally removed by haemodialysis.

**INDICATIONS**

Daonil and Semi-Daonil are indicated as an adjunct to diet to lower the blood glucose in patients with non-insulin-dependent diabetes mellitus (type 2) whose hyperglycaemia cannot be controlled by diet alone. Because of its broad and predictable action, Daonil and Semi-Daonil are often suitable for the management of patients who have failed to respond to other oral antidiabetics.

In initiating treatment for non-insulin-dependent diabetes, diet should be emphasised as the primary form of treatment. Caloric restriction and weight loss are essential in the obese diabetic patient. Proper dietary management alone may be effective in controlling the blood glucose and symptoms of hyperglycaemia. The importance of regular physical activity should also be stressed, and cardiovascular risk factors should be identified and corrective measures taken where possible. If this treatment programme fails to reduce symptoms and/or blood glucose the use of an oral sulphonylurea should be considered. Use of Daonil and Semi-Daonil must be viewed by both the physician and patient as a treatment in addition to diet, and not as a substitute for diet or as a convenient mechanism for avoiding dietary restraint.

**CONTRAINDICATIONS**

1. Known hypersensitivity or allergy to glibenclamide or any of the excipients.
2. Insulin-dependent diabetes (type 1 or juvenile onset diabetes) or in those with diabetes complicated by ketosis.
3. Treatment of diabetic ketoacidosis.
4. Serious metabolic decompensation with acidosis, in particular precoma and coma.
5. Severe impairment of renal function.
6. Severe hepatic dysfunction.
PRECAUTIONS

The treatment of diabetes requires regular checks. Until optimal control is achieved, or when changing from one product to another, or when tablets are not taken regularly, the patient's alertness and capacity to react may be impaired to such an extent that he or she may not be fit to drive, or to operate machinery.

When situations of unusual stress arise (e.g. trauma, emergency or elective surgery, febrile infections), blood glucose regulation may deteriorate and a temporary change to insulin may become necessary to maintain good metabolic control.

It should be borne in mind that there is a possibility of cross-sensitivity to sulphonamides and their derivatives. Persons allergic to other sulphonamide derivatives may develop an allergic reaction to glibenclamide as well.

Epidemiological studies suggest that the administration of glibenclamide is associated with an increased risk of cardiovascular mortality, when compared to treatment with metformin or gliclazide. This risk was especially observed in patients with diagnosed coronary diseases.

HYPOGLYCAEMIC REACTIONS

Severe hypoglycaemia, which may be prolonged and is potentially lethal, can be induced by all sulphonylureas.

Debilitated, malnourished, or geriatric patients and patients with mild disease or impaired hepatic or renal function should be carefully monitored and dosage of glibenclamide should be carefully adjusted in these patients, since they may be predisposed to developing hypoglycaemia. Renal or hepatic insufficiency may cause increased serum concentrations of glibenclamide and hepatic insufficiency may also diminish gluconeogenic capacity, both of which increase the risk of severe hypoglycaemic reactions.

Alcohol ingestion (see Interactions With Other Medicines) intense or prolonged exercise, deficient caloric intake, use of more than one antidiabetic agent, severe endocrine disorders and adrenal or pituitary insufficiency may also predispose patients to the development of hypoglycaemia.

If risk factors for hypoglycaemia are present, it may be necessary to adjust the dosage of glibenclamide or the entire therapy. This also applies whenever illness occurs during therapy or the patient's lifestyle changes.

Elderly patients are particularly susceptible to hypoglycaemic action of glucose-lowering drugs. Hypoglycaemia may be difficult to recognise in the elderly. The initial and maintenance dosing should be conservative to avoid hypoglycaemic reactions.

Hypoglycaemia can, almost always, be promptly controlled by immediate intake of carbohydrates (glucose or sugar, e.g. in the form of sugar lumps, sugar-sweetened fruit juice or tea).
Despite initially successful countermeasures, hypoglycaemia may recur. Patients must, therefore, remain under close observation.

Severe hypoglycaemia, or a protracted episode, which can only be temporarily controlled by usual amounts of sugar, further requires immediate treatment and follow-up by a physician and, in some circumstances, in-patient hospital care.

Patients receiving glibenclamide should be monitored with regular clinical and laboratory evaluations, including blood and urine glucose determinations, to determine the minimum effective dosage and to detect primary failure (inadequate lowering of blood glucose concentration at the maximum recommended dosage) or secondary failure (loss of control of blood glucose concentration following an initial period of effectiveness) to the drug. Glycosylated haemoglobin measurements may also be useful for monitoring the patient's response to glibenclamide therapy. During the withdrawal period in patients in whom glibenclamide is replacing insulin, patients should be instructed to test their urine for glucose and ketones at least 3 times daily, and to report the results to their physician; when feasible, patient or laboratory monitoring of blood glucose concentration is preferable. Care should be taken to avoid ketosis, acidosis and coma during the withdrawal period in patients being switched from insulin to glibenclamide. If adequate lowering of blood glucose concentration is no longer achieved during maintenance therapy with glibenclamide, the drug should be discontinued.

As is necessary during treatment with any blood-glucose-lowering drug, the patient and the physician must be aware of the risk of hypoglycaemia. Patients and responsible family members should be made aware of the signs and symptoms of hyperglycaemia and hypoglycaemia and the prompt action required in the event of such occurrences. Symptoms of hyperglycaemia include severe thirst, dry mouth, frequent micturition and dry skin. Possible symptoms of hypoglycaemia include intense hunger, nausea, vomiting, sweating, tremor, pareses, sensory disturbances, restlessness, irritability, aggressiveness, depression, confusion, speech disorders, aphasia, visual disorders, impaired concentration, impaired alertness and reactions, headaches, dizziness, disturbed sleep, helplessness, loss of self-control, delirium, transient neurological disorders such as cerebral convulsions, lassitude, sleepiness, somnolence, loss of consciousness up to and including coma, shallow respiration and bradycardia. In addition, signs of adrenergic counter-regulation may be present, such as sweating, clammy skin, anxiety, tachycardia, hypertension, palpitations, angina pectoris and cardiac arrhythmias.

The clinical picture of a severe hypoglycaemic attack may resemble that of a stroke. The symptoms of hypoglycaemia nearly always subside when hypoglycaemia is corrected.

In the presence of a genetic defect in metabolism, the elimination half-life may be prolonged.

Because of its broad and predictable hypoglycaemic effect, Daonil and Semi-Daonil should be taken immediately before breakfast. Patients who eat only a light breakfast should defer the first dose of the day until lunch time.

Some improvement in glucose tolerance may take place after a few weeks' treatment with Daonil and Semi-Daonil. The clinical status should be checked within the first 4-8 weeks and at regular intervals thereafter so as to ascertain whether it is possible to cut down the dose or cease glibenclamide therapy. Correction of dosage must also be considered whenever the patient’s weight changes, the patient’s life-style changes or other factors arise that cause an increased susceptibility to hypoglycaemia or hyperglycaemia.
HAEMOLYTIC ANAEMIA

Treatment of patients with glucose-6-phosphate dehydrogenase (G6PD) deficiency with sulphonylurea agents can lead to haemolytic anaemia. Since glibenclamide belongs to the class of sulphonylurea agents, caution should be used in patients with G6PD-deficiency and a non-sulphonylurea alternative should be considered.

USE IN PREGNANCY (CATEGORY C)

It is important to achieve strict normoglycaemia during pregnancy. Glibenclamide must not be taken during pregnancy. The patient must change over to insulin during pregnancy. The sulphonylureas may enter the foetal circulation and cause neonatal hypoglycaemia. In animal studies, embryotoxicity and/or birth defects have been demonstrated.

Patients planning a pregnancy must inform their physician. It is recommended that such patients change to insulin.

Category C: Drugs which, owing to their pharmacological effects, have caused or may be suspected of causing, harmful effects on the human foetus or neonate without causing malformations. These effects may be reversible. Relevant texts should be consulted for further details.

USE IN LACTATION

It is not known whether glibenclamide is excreted in milk or whether it has a harmful effect on the newborn. To prevent possible ingestion with breast milk, glibenclamide must not be taken by breastfeeding women. If necessary, the patient must change over to insulin, or must stop breastfeeding.

PAEDIATRIC USE

The safety and efficacy of glibenclamide in children have not been established. Glibenclamide is not recommended for use in this age group.

INTERACTIONS WITH OTHER MEDICINES

An increased incidence of elevated liver enzymes was observed in patients receiving glibenclamide concomitantly with bosentan. Both bosentan and glibenclamide inhibit the bile salt export pump, leading to intracellular accumulation of cytotoxic bile salts. Therefore, this combination should not be used (see CONTRAINDICATIONS).

Other drugs given at the same time as sulphonylureas may cause undesirable depression or elevation of the blood sugar level.

Glibenclamide is mainly metabolized by CYP2C9 and to a lesser extent by CYP3A4. This should be taken into account when glibenclamide is coadministered with inducers or inhibitors of CYP2C9.

Drugs which may potentiate the hypoglycaemic action of Daonil and Semi-Daonil include insulin, other oral antidiabetic agents, alcohol, ACE inhibitors, aminosalicylic acid, anabolic steroids and male sex hormones, azapropazone, beta-receptor blockers, bezafibrate, biguanides,
chloramphenicol, clarithromycin, clofibrate, clonidine, co-trimoxazole, coumarin derivatives, cyclophosphamide, disopyramide, fenfluramine, fenyardiol, fibrates, fluoxetine, gemfibrozil, guanethidine, heparin, ifosfamide, MAO-inhibitors, miconazole, oxpentifylline (parenteral, in high doses), oxyphenbutazone, para-aminosalicylic acid, phenylbutazone, phenyramidol, phosphamides, probenecid, quinolone antibiotics, ranitidine, reserpine, salicylates, sulphipyrazoline, certain long-acting sulphonamides, tetracycline compounds, tritoqualine and trophysamide. Highly protein-bound drugs which may also potentiate the hypoglycaemic action of Daonil and Semi-Daonil due to glibenclamide displacement from plasma proteins, include oral anticoagulants, hydantoins, salicylates and other non-steroidal anti-inflammatory agents.

Drugs which may cause an attenuation of the hypoglycaemic action of Daonil and Semi-Daonil include adrenaline (epinephrine) and other sympathomimetic agents, alcohol, acetazolamide, barbiturates, calcium channel blockers, cimetidine, clonidine, corticosteroids, diazoxide, diuretics, glucagon, isoniazid, large doses of laxatives, nicotinic acid (high dosage), oestrogens, progestogens, phenothiazine derivatives, phenytoin, ranitidine, rifampicin, ritodrine and thyroid hormones.

Concomitant treatment with beta-receptor blockers, clonidine, reserpine, guanethidine or other sympatholytic drugs may mask the warning symptoms of a hypoglycaemic attack. The symptoms of hypoglycaemia may also be milder or absent where hypoglycaemia develops gradually or where there is autonomic neuropathy. In rare instances, potentiation or attenuation of the blood-sugar-lowering effect of Daonil and Semi-Daonil have been observed during concomitant treatment with H₂ receptor antagonists, clonidine or reserpine.

In very rare cases, an intolerance to alcohol may occur. Both acute and chronic alcohol intake, or excessive alcohol ingestion by people who drink occasionally, may attenuate the hypoglycaemic effect of glibenclamide or dangerously potentiate it by delaying its metabolic inactivation. Disulfiram-like reactions have occurred very rarely following the concomitant use of alcohol and glibenclamide.

Glibenclamide may either potentiate or weaken the effect of coumarin derivatives.

Glibenclamide may increase cyclosporin plasma concentration and potentially lead to its increased toxicity. Monitoring and dosage adjustment of cyclosporin are therefore recommended when both drugs are co-administered.

Colestevlam binds to glibenclamide and reduces glibenclamide absorption from the gastrointestinal tract. No interaction was observed when glibenclamide was taken at least 4 hours before colestevlam. Therefore glibenclamide should be administered at least 4 hours prior to colestevlam.

Food does not alter the bioavailability or other pharmacokinetic parameters of glibenclamide.

**ADVERSE EFFECTS**

Clinical experience in the use of Daonil and Semi-Daonil has shown that side effects serious enough to compel discontinuation of therapy are uncommon, even during long-term therapy. However, if adverse effects persists, the drug should be discontinued.
HYPOGLYCAEMIA

Hypoglycaemia may not only be severe, but also prolonged and fatal (see PRECAUTIONS, Hypoglycaemic Reactions and OVERDOSAGE).

EYE DISORDERS

Especially at the start of treatment, there may be temporary visual impairment due to the change in blood glucose levels. The cause is a temporary alteration in the turgidity and hence the refractive index of the lens, this being dependent on blood glucose level.

GASTROINTESTINAL REACTIONS

Adverse gastrointestinal effects such as nausea, vomiting, epigastric fullness or sensation of pressure, abdominal pain, anorexia, heartburn, dyspepsia and diarrhoea are the most common adverse reactions to glibenclamide, occurring in about 1-2% of patients. Glibenclamide-induced adverse gastrointestinal effects appear to be dose related and may subside following a reduction in dosage. Pancreatitis has been reported rarely.

DERMATOLOGIC REACTIONS

Hypersensitivity reactions, allergic or pseudoallergic reactions may occur. Allergic skin reactions e.g. pruritus, erythema, urticaria, erythematous and maculo-papular and bullous skin eruptions or psoriasiform drug eruptions occur in 1.5% of treated patients. These may be transient and may disappear despite continued use of glibenclamide; if skin reactions persist, the drug should be discontinued. In isolated cases, mild reactions in the form of urticaria may develop into serious and even life-threatening reactions with dyspnoea and fall in blood pressure, sometimes progressing to shock. In the event of urticaria, a physician must therefore be notified immediately.

A hypersensitivity reaction may be directed against glibenclamide itself, but may alternatively be triggered by excipients. Allergy to sulphonamide derivatives may also be responsible for an allergic reaction to glibenclamide.

In isolated cases, allergic vasculitis may arise and, in some circumstances, may be life-threatening. In isolated cases, hypersensitivity of the skin to light may occur, and sodium concentration in the serum may decrease. Porphyria cutanea tarda and pellagra-like changes have been reported with sulphonylureas.

HAEMATOLOGIC REACTIONS

Anaemia, leukopaenia, thrombocytopenia, thrombocytopenic purpura, agranulocytosis, pancytopenia, eosinophilia, haemolytic anaemia, aplastic anaemia, bone marrow aplasia, eosinophilia and coagulation disorders have been reported with sulphonylureas. Potentially life-threatening changes in the blood picture may occur. They may include, rarely, mild to severe thrombopenia (e.g. presenting as purpura) and, in isolated cases, haemolytic anaemia, erythrocytopenia, granulocytopenia, agranulocytosis and (for example, due to myelosuppression) pancytopenia. In principle, these reactions are reversible once glibenclamide has been withdrawn.
HEPATIC REACTIONS

Increased liver enzymes (AST, ALT), abnormal liver function, cholestasis, cholestatic hepatitis, granulomatous hepatitis and bilirubinaemia have been reported with sulphonylureas. In isolated cases there may be hepatitis, elevation of liver enzyme levels and/or cholestasis and jaundice which may progress to life-threatening liver failure but can regress after withdrawal of glibenclamide.

MISCELLANEOUS

Although a causal relationship has not been established, the following adverse effects have been reported in patients receiving glibenclamide: paresthesia, blindness, deafness, diplopia, visual disturbances, tremor, convulsions (other than withdrawal), encephalopathy, confusion, acute psychosis, abnormal renal function, acute renal failure, ocular disturbances (accommodation changes, crystalline lens changes), lactic acidosis, alopecia/hipotrichosis, hyponatraemia, syndrome of inappropriate secretion of antidiuretic hormone (SIADH), arthralgia, arthritis, cerebrovascular disorders, headache, facial oedema, angioedema, weight gain, hypersensitivity vasculitis and increased sweating.

DOSAGE AND ADMINISTRATION

Dosage of glibenclamide must be based on regular blood and urine glucose determinations and must be carefully individualised to obtain optimum therapeutic effect. The dosage of glibenclamide must be the lowest possible dose which is effective. If appropriate glibenclamide dosage regimens are not followed, hypoglycaemia may be precipitated. It is very important not to skip meals after the tablets have been taken.

In newly treated patients with diabetes, stabilisation should be commenced with one Semi-Daonil tablet daily, taken immediately before breakfast. Patients who eat only a light breakfast should defer the first dose of the day until lunchtime.

After 3-5 days, the blood sugar and urine sugar should be checked. If good control has been achieved, the daily dose of one Semi-Daonil tablet is continued as maintenance therapy.

If control is unsatisfactory, elevation of the daily dose in steps of 2.5 mg is necessary at intervals of 7 days up to a maximum of 15 mg, or, in exceptional cases, 4 tablets (20 mg) daily.

Daily allotments of up to 10 mg can be taken as a single dose before breakfast; daily dosage in excess of 10 mg should be taken before the evening meal.

As an improvement in control of diabetes is, in itself, associated with higher insulin sensitivity, glibenclamide requirements may fall as treatment proceeds. To avoid hypoglycaemia, timely dose reduction or cessation of Daonil therapy must therefore be considered.

Mistakes, e.g. forgetting to take a dose, must never be corrected by subsequently taking a larger dose. Measures for dealing with such mistakes (in particular forgetting a dose or skipping a meal), or in the event a dose cannot be taken at the prescribed time, must be discussed and agreed between physician and patient beforehand.

In the management of type 2 diabetes mellitus, oral hypoglycaemic administration is not a substitute for appropriate dietary control.
When transferring patients from other oral antidiabetic drugs, it is recommended to begin with the usual starting dose (2.5 to 5 mg) per day. Depending on the pharmacokinetic and pharmacodynamic characteristics of the previous medication, a drug-free transition period may be necessary in order to avoid overlapping drug effects possibly resulting in hypoglycaemia.

In general, patients who were previously maintained on insulin dosages up to 40 units daily may be transferred directly to glibenclamide and administration of insulin may be abruptly discontinued; the initial glibenclamide dosage is 2.5-5 mg daily in patients whose insulin dosage was less than 20 units daily and 5 mg daily in patients whose insulin dosage was 20-40 units daily. In patients requiring insulin dosages greater than 40 units daily, an initial glibenclamide dosage of 5 mg daily should be started and insulin dosage reduced by 50%. Subsequently, insulin is withdrawn gradually and dosage of glibenclamide is increased in increments of 1.25-2.5 mg daily every 2-10 days, according to the patient's tolerance and therapeutic response. During the period of insulin withdrawal, patients should test their urine at least 3 times daily for glucose and acetone, and should be instructed to report the results to their physician so that appropriate adjustments in therapy may be made if necessary; when feasible, patient or laboratory monitoring of blood glucose concentration is preferable. The presence of persistent ketonuria with glycosuria, ketosis, and/or inadequate lowering or persistent elevation of blood glucose concentration indicates that the patient requires insulin therapy.

If adequate control is no longer possible with diet and DAONIL and SEMI-DAONIL (maximum 20 mg daily), good results may be obtained by combined administration of Daonil and a biguanide derivative.

**OVERDOSAGE**

For information on the management of overdose, contact the Poisons Information Centre on 131126.

**PATHOGENESIS**

Acute glibenclamide toxicity may result from excessive dosage and numerous conditions may predispose patients to the development of glibenclamide-induced hypoglycaemia (see CONTRAINDICATIONS, PRECAUTIONS). Acute overdose as well as long-term treatment with too high a dose of glibenclamide may lead to severe, protracted, life-threatening hypoglycaemia. Fatal hypoglycaemia has occurred with ingestion of as little as 2.5 to 5 mg of the drug.

**MANIFESTATIONS**

Acute glibenclamide overdosage is manifested principally as hypoglycaemia, which may be severe and has occasionally been fatal. Severe hypoglycaemia may result in loss of consciousness and seizures, with resultant neurologic sequelae.

**TREATMENT**

In case of overdosage with glibenclamide, a doctor has to be called immediately. At the first signs of hypoglycaemia, the patient must immediately take sugar, preferably glucose, unless a doctor has already started care.
Since hypoglycaemia and its clinical symptoms may recur after apparent clinical recovery (even after several days), close and continued medical supervision and possibly referral to a hospital are indicated. In particular, significant overdosage and severe reactions, e.g. with unconsciousness or other neurological dysfunctions, are emergency cases and require immediate care and hospitalisation.

If hypoglycaemic coma is diagnosed or suspected, administration of glucagon (adults: 0.5 - 1 mg) i.v., s.c. or i.m., or i.v. infusion of a 20% glucose solution (adults: 40-100 mL) is indicated, until the patient recovers consciousness. In infants, glucose must be dosed very carefully, accompanied by close monitoring of blood glucose, taking into account the risk of potentially severe hyperglycaemia. Other symptomatic therapy (e.g. anticonvulsants) should be administered as necessary.

In cases of acute intake of large amounts of glibenclamide, detoxification e.g. by medicinal charcoal as an absorbent, is indicated.

After acute glucose replacement has been completed, it is usually necessary to give an intravenous glucose infusion in lower concentration to ensure that the hypoglycaemia does not recur. The patient’s blood glucose level should be carefully monitored for at least 24 hours. In severe cases with a protracted course, hypoglycaemia, or the danger of slipping back into hypoglycaemia, may persist for several days.

**PRESENTATION AND STORAGE CONDITIONS**

**DAONIL**

Tablets, 5 mg (white, biplane oblong tablet with a score-line on both sides. LDI is engraved each side of the score-line and inverted. The other side is plain).

Store below 25°C. Protect from light.

**SEMI-DAONIL**

Tablets, 2.5 mg (white, round, scored, marked LDY with HOECHST logo).

*Not marketed

**POISON SCHEDULE OF THE MEDICINE**

Prescription Medicine (Schedule 4)

**NAME AND ADDRESS OF SPONSOR**

sanofi-aventis australia pty ltd
12-24 Talavera Road
Macquarie Park NSW 2113
DATE OF FIRST INCLUSION IN THE ARTG

Semi-Daonil: 24 November 1993
Daonil: 27 April 2000

DATE OF MOST RECENT AMENDMENT

1 September 2016