

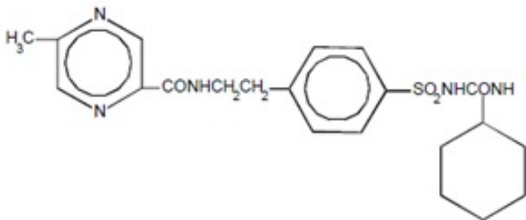
GLIPIZIDE- glipizide tablet
Advagen Pharma Limited

Glipizide Tablets, USP
For Oral Use

DESCRIPTION

Glipizide is an oral blood-glucose-lowering drug of the sulfonylurea class.

The Chemical Abstracts name of glipizide is 1-cyclohexyl-3-[[p-[2-(5-methylpyrazine-carboxamido)ethyl]phenyl]sulfonyl]urea. The molecular formula is $C_{21}H_{27}N_5O_4S$; the molecular weight is 445.55; the structural formula is shown below:



Glipizide is a whitish, odorless powder with a pKa of 5.9. It is insoluble in water and alcohols, but soluble in 0.1 N NaOH; it is freely soluble in dimethylformamide. Glipizide tablets for oral use are available in 2.5 mg, 5 mg, 10 mg and 15 mg strengths.

Inactive ingredients are: colloidal silicon dioxide; croscarmellose sodium, lactose anhydrous; microcrystalline cellulose and stearic acid.

Meets USP Dissolution test 2.

CLINICAL PHARMACOLOGY

Mechanism of Action: The primary mode of action of glipizide in experimental animals appears to be the stimulation of insulin secretion from the beta cells of pancreatic islet tissue and is thus dependent on functioning beta cells in the pancreatic islets. In humans, glipizide appears to lower the blood glucose acutely by stimulating the release of insulin from the pancreas, an effect dependent upon functioning beta cells in the pancreatic islets. The mechanism by which glipizide lowers blood glucose during long-term administration has not been clearly established. In man, stimulation of insulin secretion by glipizide in response to a meal is undoubtedly of major importance. Fasting insulin levels are not elevated even on long-term glipizide administration, but the postprandial insulin response continues to be enhanced after at least 6 months of treatment. The insulinotropic response to a meal occurs within 30 minutes after an oral dose of glipizide in diabetic patients, but elevated insulin levels do not persist beyond the time of the meal challenge. Extrapancreatic effects may play a part in the mechanism of action of oral sulfonylurea hypoglycemic drugs.

Blood sugar control persists in some patients for up to 24 hours after a single dose of glipizide, even though plasma levels have declined to a small fraction of peak levels by that time (see **Pharmacokinetics** below).

Some patients fail to respond initially, or gradually lose their responsiveness to sulfonylurea drugs, including glipizide. Alternatively, glipizide may be effective in some

patients who have not responded or have ceased to respond to other sulfonylureas.

Other Effects: It has been shown that glipizide therapy was effective in controlling blood sugar without deleterious changes in the plasma lipoprotein profiles of patients treated for NIDDM.

In a placebo-controlled, crossover study in normal volunteers, glipizide had no antidiuretic activity and, in fact, led to a slight increase in free water clearance.

Pharmacokinetics:

Gastrointestinal absorption of glipizide in man is uniform, rapid, and essentially complete. Peak plasma concentrations occur 1–3 hours after a single oral dose. The half-life of elimination ranges from 2–4 hours in normal subjects, whether given intravenously or orally. The metabolic and excretory patterns are similar with the two routes of administration, indicating that first-pass metabolism is not significant. Glipizide does not accumulate in plasma on repeated oral administration. Total absorption and disposition of an oral dose was unaffected by food in normal volunteers, but absorption was delayed by about 40 minutes. Thus, glipizide was more effective when administered about 30 minutes before, rather than with, a test meal in diabetic patients. Protein binding was studied in serum from volunteers who received either oral or intravenous glipizide and found to be 98–99% one hour after either route of administration. The apparent volume of distribution of glipizide after intravenous administration was 11 liters, indicative of localization within the extracellular fluid compartment. In mice, no glipizide or metabolites were detectable autoradiographically in the brain or spinal cord of males or females, nor in the fetuses of pregnant females. In another study, however, very small amounts of radioactivity were detected in the fetuses of rats given labelled drug.

The metabolism of glipizide is extensive and occurs mainly in the liver. The primary metabolites are inactive hydroxylation products and polar conjugates and are excreted mainly in the urine. Less than 10% unchanged glipizide is found in the urine.

INDICATIONS AND USAGE

Glipizide tablets are indicated as an adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus.

CONTRAINDICATIONS

Glipizide tablets are contraindicated in patients with:

1. Known hypersensitivity to the drug.
2. Type 1 diabetes mellitus, diabetic ketoacidosis, with or without coma. This condition should be treated with insulin.

WARNINGS

SPECIAL WARNING ON INCREASED RISK OF CARDIOVASCULAR MORTALITY

The administration of oral hypoglycemic drugs has been reported to be associated with increased cardiovascular mortality as compared to treatment with diet alone or diet plus insulin. This warning is based on the study conducted by the University Group Diabetes Program (UGDP), a long-term prospective clinical trial designed to evaluate the effectiveness of glucose-lowering drugs in preventing or delaying vascular complications in patients with non-insulin-dependent diabetes. The study involved 823 patients who were randomly assigned to one of four treatment groups (*Diabetes*, 19,

supp. 2: 747-830, 1970).

UGDP reported that patients treated for 5 to 8 years with diet plus a fixed dose of tolbutamide (1.5 grams per day) had a rate of cardiovascular mortality approximately 2½ times that of patients treated with diet alone. A significant increase in total mortality was not observed, but the use of tolbutamide was discontinued based on the increase in cardiovascular mortality, thus limiting the opportunity for the study to show an increase in overall mortality. Despite controversy regarding the interpretation of these results, the findings of the UGDP study provide an adequate basis for this warning. The patient should be informed of the potential risks and advantages of glipizide and of alternative modes of therapy.

Although only one drug in the sulfonylurea class (tolbutamide) was included in this study, it is prudent from a safety standpoint to consider that this warning may also apply to other oral hypoglycemic drugs in this class, in view of their close similarities in mode of action and chemical structure.

PRECAUTIONS

General

Macrovascular Outcomes: There have been no clinical studies establishing conclusive evidence of macrovascular risk reduction with glipizide or any other anti-diabetic drug.

Renal and Hepatic Disease: The metabolism and excretion of glipizide may be slowed in patients with impaired renal and/or hepatic function. If hypoglycemia should occur in such patients, it may be prolonged and appropriate management should be instituted.

Hypoglycemia: All sulfonylurea drugs are capable of producing severe hypoglycemia. Proper patient selection, dosage, and instructions are important to avoid hypoglycemic episodes. Renal or hepatic insufficiency may cause elevated blood levels of glipizide and the latter may also diminish gluconeogenic capacity, both of which increase the risk of serious hypoglycemic reactions. Elderly, debilitated or malnourished patients, and those with adrenal or pituitary insufficiency, are particularly susceptible to the hypoglycemic action of glucose-lowering drugs. Hypoglycemia may be difficult to recognize in the elderly, and in people who are taking beta-adrenergic blocking drugs. Hypoglycemia is more likely to occur when caloric intake is deficient, after severe or prolonged exercise, when alcohol is ingested, or when more than one glucose-lowering drug is used.

Loss of Control of Blood Glucose: When a patient stabilized on any diabetic regimen is exposed to stress such as fever, trauma, infection, or surgery, a loss of control may occur. At such times, it may be necessary to discontinue glipizide and administer insulin.

The effectiveness of any oral hypoglycemic drug, including glipizide, in lowering blood glucose to a desired level decreases in many patients over a period of time, which may be due to progression of the severity of the diabetes or to diminished responsiveness to the drug. This phenomenon is known as secondary failure, to distinguish it from primary failure in which the drug is ineffective in an individual patient when first given.

Hemolytic Anemia: Treatment of patients with glucose 6-phosphate dehydrogenase (G6PD) deficiency with sulfonylurea agents can lead to hemolytic anemia. Because glipizide belongs to the class of sulfonylurea agents, caution should be used in patients with G6PD deficiency and a non-sulfonylurea alternative should be considered. In post-marketing reports, hemolytic anemia has also been reported in patients who did not have known G6PD deficiency.

Laboratory Tests

Blood and urine glucose should be monitored periodically. Measurement of glycosylated hemoglobin may be useful.

Information for Patients

Patients should be informed of the potential risks and advantages of glipizide and of alternative modes of therapy. They should also be informed about the importance of adhering to dietary instructions, of a regular exercise program, and of regular testing of urine and/or blood glucose.

The risks of hypoglycemia, its symptoms and treatment, and conditions that predispose to its development should be explained to patients and responsible family members. Primary and secondary failure should also be explained.

Physician Counseling Information for Patients

In initiating treatment for type 2 diabetes, diet should be emphasized 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 hyperglycemia. The importance of regular physical activity should also be stressed, and cardiovascular risk factors should be identified and corrective measures taken where possible. Use of glipizide or other antidiabetic medications must be viewed by both the physician and patient as a treatment in addition to diet and not as a substitution or as a convenient mechanism for avoiding dietary restraint. Furthermore, loss of blood glucose control on diet alone may be transient, thus requiring only short-term administration of glipizide tablets or other antidiabetic medications. Maintenance or discontinuation of glipizide tablets or other antidiabetic medications should be based on clinical judgment using regular clinical and laboratory evaluations.

Drug Interactions:The hypoglycemic action of sulfonylureas may be potentiated by certain drugs including nonsteroidal anti-inflammatory agents, some azoles, and other drugs that are highly protein bound, salicylates, sulfonamides, chloramphenicol, probenecid, coumarins, monoamine oxidase inhibitors, quinolones and beta adrenergic blocking agents. When such drugs are administered to a patient receiving glipizide, the patient should be observed closely for hypoglycemia. When such drugs are withdrawn from a patient receiving glipizide, the patient should be observed closely for loss of control. *In vitro* binding studies with human serum proteins indicate that glipizide binds differently than tolbutamide and does not interact with salicylate or dicumarol. However, caution must be exercised in extrapolating these findings to the clinical situation and in the use of glipizide with these drugs.

Certain drugs tend to produce hyperglycemia and may lead to loss of control. These drugs include the thiazides and other diuretics, corticosteroids, phenothiazines, thyroid products, estrogens, oral contraceptives, phenytoin, nicotinic acid, sympathomimetics, calcium channel blocking drugs, and isoniazid. When such drugs are administered to a patient receiving glipizide, the patient should be closely observed for loss of control. When such drugs are withdrawn from a patient receiving glipizide, the patient should be observed closely for hypoglycemia.

A potential interaction between oral miconazole and oral hypoglycemic agents leading to severe hypoglycemia has been reported. Whether this interaction also occurs with the intravenous, topical, or vaginal preparations of miconazole is not known. The effect of concomitant administration of DIFLUCAN[®] (fluconazole) and glipizide has been demonstrated in a placebo-controlled crossover study in normal volunteers. All subjects received glipizide alone and following treatment with 100 mg of DIFLUCAN as a single daily oral dose for 7 days. The mean percentage increase in the glipizide AUC after fluconazole administration was 56.9% (range: 35 to 81).

In studies assessing the effect of colesevelam on the pharmacokinetics of glipizide ER in healthy volunteers, reductions in glipizide AUC_{0-∞} and C_{max} of 12% and 13%, respectively were observed when colesevelam was coadministered with glipizide ER. When glipizide ER was administered 4 hours prior to colesevelam, there was no significant change in glipizide AUC_{0-∞} or C_{max}, -4% and 0%, respectively. Therefore, glipizide should be administered at least 4 hours prior to colesevelam to ensure that colesevelam does not reduce the absorption of glipizide.

Carcinogenesis, Mutagenesis, Impairment of Fertility: A twenty month study in rats and an eighteen month study in mice at doses up to 75 times the maximum human dose revealed no evidence of drug-related carcinogenicity. Bacterial and *in vivo* mutagenicity tests were uniformly negative. Studies in rats of both sexes at doses up to 75 times the human dose showed no effects on fertility.

Pregnancy:

Glipizide was found to be mildly fetotoxic in rat reproductive studies at all dose levels (5–50 mg/kg). This fetotoxicity has been similarly noted with other sulfonylureas, such as tolbutamide and tolazamide. The effect is perinatal and believed to be directly related to the pharmacologic (hypoglycemic) action of glipizide. In studies in rats and rabbits, no teratogenic effects were found. There are no adequate and well controlled studies in pregnant women. Glipizide should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

Because recent information suggests that abnormal blood glucose levels during pregnancy are associated with a higher incidence of congenital abnormalities, many experts recommend that insulin be used during pregnancy to maintain blood glucose levels as close to normal as possible.

Nonteratogenic Effects: Prolonged severe hypoglycemia (4 to 10 days) has been reported in neonates born to mothers who were receiving a sulfonylurea drug at the time of delivery. This has been reported more frequently with the use of agents with prolonged half-lives. If glipizide is used during pregnancy, it should be discontinued at least one month before the expected delivery date.

Nursing Mothers: Although it is not known whether glipizide is excreted in human milk, some sulfonylurea drugs are known to be excreted in human milk. Because the potential for hypoglycemia in nursing infants may exist, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother. If the drug is discontinued and if diet alone is inadequate for controlling blood glucose, insulin therapy should be considered.

Pediatric Use: Safety and effectiveness in children have not been established.

Geriatric Use: A determination has not been made whether controlled clinical studies of glipizide included sufficient numbers of subjects aged 65 and over to define a difference in response from younger subjects. Other reported clinical experience has not identified differences in responses between the elderly and younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy.

ADVERSE REACTIONS

In U.S. and foreign controlled studies, the frequency of serious adverse reactions reported was very low. Of 702 patients, 11.8% reported adverse reactions and in only 1.5% was glipizide discontinued.

Hypoglycemia: See **PRECAUTIONS** and **OVERDOSAGE** sections.

Gastrointestinal: Gastrointestinal disturbances are the most common reactions. Gastrointestinal complaints were reported with the following approximate incidence: nausea and diarrhea, one in seventy; constipation and gastralgia, one in one hundred. They appear to be dose-related and may disappear on division or reduction of dosage. Cholestatic jaundice may occur rarely with sulfonylureas: glipizide should be discontinued if this occurs.

Dermatologic: Allergic skin reactions including erythema, morbilliform or maculopapular eruptions, urticaria, pruritus, and eczema have been reported in about one in seventy patients. These may be transient and may disappear despite continued use of glipizide; if skin reactions persist, the drug should be discontinued. Porphyria cutanea tarda and photosensitivity reactions have been reported with sulfonylureas.

Hematologic: Leukopenia, agranulocytosis, thrombocytopenia, hemolytic anemia (see **PRECAUTIONS**), aplastic anemia, and pancytopenia have been reported with sulfonylureas.

Metabolic: Hepatic porphyria and disulfiram-like reactions have been reported with sulfonylureas. In the mouse, glipizide pretreatment did not cause an accumulation of acetaldehyde after ethanol administration. Clinical experience to date has shown that glipizide has an extremely low incidence of disulfiram-like alcohol reactions.

Endocrine Reactions: Cases of hyponatremia and the syndrome of inappropriate antidiuretic hormone (SIADH) secretion have been reported with this and other sulfonylureas.

Miscellaneous: Dizziness, drowsiness, and headache have each been reported in about one in fifty patients treated with glipizide. They are usually transient and seldom require discontinuance of therapy.

Laboratory Tests: The pattern of laboratory test abnormalities observed with glipizide was similar to that for other sulfonylureas. Occasional mild to moderate elevations of SGOT, LDH, alkaline phosphatase, BUN, and creatinine were noted. One case of jaundice was reported. The relationship of these abnormalities to glipizide is uncertain, and they have rarely been associated with clinical symptoms.

Post-Marketing Experience The following adverse events have been reported in post-marketing surveillance:

Hepatobiliary: Cholestatic and hepatocellular forms of liver injury accompanied by jaundice have been reported rarely in association with glipizide; glipizide should be discontinued if this occurs.

OVERDOSAGE

There is no well documented experience with glipizide overdosage. The acute oral toxicity was extremely low in all species tested (LD₅₀ greater than 4 g/kg).

Overdosage of sulfonylureas, including glipizide, can produce hypoglycemia. Mild hypoglycemic symptoms without loss of consciousness or neurologic findings should be treated aggressively with oral glucose and adjustments in drug dosage and/or meal patterns. Close monitoring should continue until the physician is assured that the patient is out of danger. Severe hypoglycemic reactions with coma, seizure, or other neurological impairment occur infrequently, but constitute medical emergencies

requiring immediate hospitalization. If hypoglycemic coma is diagnosed or suspected, the patient should be given a rapid intravenous injection of concentrated (50%) glucose solution. This should be followed by a continuous infusion of a more dilute (10%) glucose solution at a rate that will maintain the blood glucose at a level above 100 mg/dL. Patients should be closely monitored for a minimum of 24 to 48 hours since hypoglycemia may recur after apparent clinical recovery. Clearance of glipizide from plasma would be prolonged in persons with liver disease. Because of the extensive protein binding of glipizide, dialysis is unlikely to be of benefit.

DOSAGE AND ADMINISTRATION

There is no fixed dosage regimen for the management of diabetes mellitus with glipizide or any other hypoglycemic agent. In addition to the usual monitoring of urinary glucose, the patient's blood glucose must also be monitored periodically to determine the minimum effective dose for the patient; to detect primary failure, i.e., inadequate lowering of blood glucose at the maximum recommended dose of medication; and to detect secondary failure, i.e., loss of an adequate blood-glucose-lowering response after an initial period of effectiveness. Glycosylated hemoglobin levels may also be of value in monitoring the patient's response to therapy.

Short-term administration of glipizide tablets may be sufficient during periods of transient loss of control in patients usually controlled well on diet.

In general, glipizide tablets should be given approximately 30 minutes before a meal to achieve the greatest reduction in postprandial hyperglycemia.

Initial Dose: The recommended starting dose is 5 mg, given before breakfast. Geriatric patients or those with liver disease may be started on 2.5 mg.

Titration: Dosage adjustments should ordinarily be in increments of 2.5–5 mg, as determined by blood glucose response. At least several days should elapse between titration steps. If response to a single dose is not satisfactory, dividing that dose may prove effective. The maximum recommended once daily dose is 15 mg. Doses above 15 mg should ordinarily be divided and given before meals of adequate caloric content. The maximum recommended total daily dose is 40 mg.

Maintenance: Some patients may be effectively controlled on a once-a-day regimen, while others show better response with divided dosing. Total daily doses above 15 mg should ordinarily be divided. Total daily doses above 30 mg have been safely given on a b.i.d. basis to long-term patients.

In elderly patients, debilitated or malnourished patients, and patients with impaired renal or hepatic function, the initial and maintenance dosing should be conservative to avoid hypoglycemic reactions (see **PRECAUTIONS** section).

Patients Receiving Insulin: As with other sulfonylurea-class hypoglycemics, many stable non-insulin-dependent diabetic patients receiving insulin may be safely placed on Glipizide tablets. When transferring patients from insulin to glipizide tablets, the following general guidelines should be considered:

- For patients whose daily insulin requirement is 20 units or less, insulin may be discontinued and glipizide therapy may begin at usual dosages. Several days should elapse between glipizide titration steps.
- For patients whose daily insulin requirement is greater than 20 units, the insulin dose should be reduced by 50% and glipizide therapy may begin at usual dosages. Subsequent reductions in insulin dosage should depend on individual patient response. Several days should elapse between glipizide titration steps.

During the insulin withdrawal period, the patient should test urine samples for sugar and

ketone bodies at least three times daily. Patients should be instructed to contact the prescriber immediately if these tests are abnormal. In some cases, especially when patient has been receiving greater than 40 units of insulin daily, it may be advisable to consider hospitalization during the transition period.

Patients Receiving Other Oral Hypoglycemic Agents:As with other sulfonylurea-class hypoglycemics, no transition period is necessary when transferring patients to glipizide. Patients should be observed carefully (1–2 weeks) for hypoglycemia when being transferred from longer half-life sulfonylureas (e.g., chlorpropamide) to glipizide due to potential overlapping of drug effect.

When colesevelam is coadministered with glipizide ER, maximum plasma concentration and total exposure to glipizide is reduced. Therefore, glipizide tablets should be administered at least 4 hours prior to colesevelam.

HOW SUPPLIED

Glipizide tablets 2.5 mg are round, white to off-white, uncoated tablets, debossed on one side and Λ on other side. They are supplied as follows:

NDC 72888-115-30, bottles of 30.

NDC 72888-115-01, bottles of 100.

NDC 72888-115-05, bottles of 500.

NDC 72888-115-00, bottles of 1000.

Glipizide tablets 5 mg are Round, white to off-white, uncoated tablets, debossed Λ on one side and break line on other side. They are supplied as follows:

11

12

NDC 72888-116-30, bottles of 30.

NDC 72888-116-01, bottles of 100.

NDC 72888-116-05, bottles of 500.

NDC 72888-116-00, bottles of 1000.

Glipizide tablets 10 mg are Round, white to off-white, uncoated tablets, debossed Λ on one side and break line on other side. They are supplied as follows:

14

NDC 72888-117-30, bottles of 30.

NDC 72888-117-01, bottles of 100.

NDC 72888-117-05, bottles of 500.

NDC 72888-117-00, bottles of 1000.

Glipizide tablets 15 mg are Round, white to off-white, uncoated tablets, debossed Λ on one side and break line on other side. They are supplied as follows:

111

NDC 72888-231-01, bottles of 100.

NDC 72888-231-05, bottles of 500.

NDC 72888-231-00, bottles of 1000.

RECOMMENDED STORAGE:

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

All trademarks are the property of their respective owners.

Manufactured by:

Rubicon Research Limited
Thane, 421506 India

Distributed by:

Advagen Pharma Ltd
East Windsor, NJ 08520, USA.

Revision:04, 02/2026

PACKAGE LABEL-PRINCIPAL DISPLAY PANEL

Glipizide Tablets 2.5 mg - NDC 72888-115-30 - 30 Tablets Label

Each tablet contains: glipiZIDE, USP.....2.5 mg

DOSAGE AND USE: See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at <https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-115-30

glipiZIDE
Tablets, USP

2.5 mg

30 Tablets Rx Only

ADVAGEN

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA
Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.
Mfg Lic No. : KD-682 Rev.02, 01/26 XXXX

Overprinting Area
43 mm X 15 mm

Glipizide Tablets 2.5 mg - NDC 72888-115-01 - 100 Tablets Label

Each tablet contains: glipiZIDE, USP.....2.5 mg

DOSAGE AND USE: See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at <https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-115-01

glipiZIDE
Tablets, USP

2.5 mg

100 Tablets Rx Only

ADVAGEN

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA
Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.
Mfg Lic No. : KD-682 Rev.02, 01/26 XXXX

Overprinting Area
43 mm X 15 mm

Glipizide Tablets 2.5 mg - NDC 72888-115-05 - 500 Tablets Label


Each tablet contains: glipiZIDE, USP.....2.5 mg
DOSAGE AND USE: See accompanying prescribing information.
Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].
 Dispense in a tightly closed container using child resistant closure.
 Keep this and all medication out of the reach of children.
Package Insert available at <https://www.advagenpharma.com/pil/glipizidetabets>

NDC 72888-115-05
glipiZIDE
Tablets, USP
2.5 mg
 500 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**, East Windsor, NJ 08520, USA
 Manufactured by: **Rubicon Research Ltd.**, Thane 421506, India.
 Mfg Lic No. : KD-682
 Rev.02, 01/26

XXXX

Overprinting Area
 50 mm X 20 mm



Glipizide Tablets 2.5 mg - NDC 72888-115-00 - 1000 Tablets Label


Each tablet contains: glipiZIDE, USP.....2.5 mg
DOSAGE AND USE: See accompanying prescribing information.
Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].
 Dispense in a tightly closed container using child resistant closure.
 Keep this and all medication out of the reach of children.
Package Insert available at <https://www.advagenpharma.com/pil/glipizidetabets>

NDC 72888-115-00
glipiZIDE
Tablets, USP
2.5 mg
 1000 Tablets Rx Only

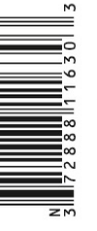
Distributed by: **Advagen Pharma Ltd.**, East Windsor, NJ 08520, USA
 Manufactured by: **Rubicon Research Ltd.**, Thane 421506, India.
 Mfg Lic No. : KD-682
 Rev.02, 01/26

XXXX

Overprinting Area
 50 mm X 20 mm



Glipizide Tablets 5 mg - NDC 72888-116-30 - 30 Tablets Label


Each tablet contains: glipiZIDE, USP.....5 mg
DOSAGE AND USE: See accompanying prescribing information.
Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].
 Dispense in a tightly closed container using child resistant closure.
 Keep this and all medication out of the reach of children.
Package Insert available at <https://www.advagenpharma.com/pil/glipizidetabets>

NDC 72888-116-30
glipiZIDE
Tablets, USP
5 mg
 30 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**, East Windsor, NJ 08520, USA
 Manufactured by: **Rubicon Research Ltd.**, Thane 421506, India.
 Mfg Lic No. : KD-682 Rev.04, 08/24 22439


Overprinting Area
 43 mm X 15 mm



Glipizide Tablets 5 mg - NDC 72888-116-01 - 100 Tablets Label

Each tablet contains:
glipiZIDE, USP.....5 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at
<https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-116-01

glipiZIDE
Tablets, USP

5 mg

100 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22440

Overprinting Area
43 mm X 20 mm

Glipizide Tablets 5 mg - NDC 72888-116-05 - 500 Tablets Label

Each tablet contains:
glipiZIDE, USP.....5 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at
<https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-116-05

glipiZIDE
Tablets, USP

5 mg

500 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22441

Overprinting Area
50 mm X 20 mm

Glipizide Tablets 5 mg - NDC 72888-116-00 - 1000 Tablets Label

Each tablet contains:
glipiZIDE, USP.....5 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at
<https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-116-00

glipiZIDE
Tablets, USP

5 mg

1000 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22442

Overprinting Area
50 mm X 20 mm

Glipizide Tablets 10 mg - NDC 72888-117-30 - 30 Tablets Label

Each tablet contains:
glipiZIDE, USP.....10 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at
<https://www.advagenpharma.com/pil/glipizidetabets>

NDC 72888-117-30

glipiZIDE
Tablets, USP

10 mg

30 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

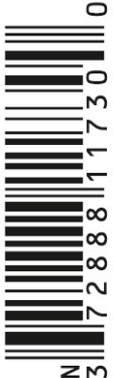
Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22443

Overprinting Area
43 mm X 20 mm





Glipizide Tablets 10 mg - NDC 72888-117-01 - 100 Tablets Label

Each tablet contains:
glipiZIDE, USP.....10 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at
<https://www.advagenpharma.com/pil/glipizidetabets>

NDC 72888-117-01

glipiZIDE
Tablets, USP

10 mg

100 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22444

Overprinting Area
50 mm X 20 mm





Glipizide Tablets 10 mg - NDC 72888-117-05 - 500 Tablets Label

Each tablet contains:
glipiZIDE, USP.....10 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at
<https://www.advagenpharma.com/pil/glipizidetabets>

NDC 72888-117-05

glipiZIDE
Tablets, USP

10 mg

500 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

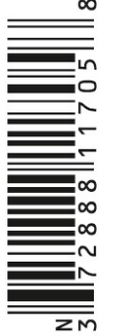
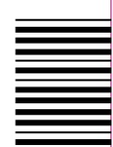
Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22445

Overprinting Area
50 mm X 20 mm


Glipizide Tablets 10 mg - NDC 72888-117-00 - 1000 Tablets Label



Each tablet contains:
glipiZIDE, USP.....10 mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at <https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-117-00

glipiZIDE Tablets, USP

10 mg

1000 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.04, 08/24

22446

Overprinting Area
56 mm X 25 mm



Glipizide Tablets 15 mg - NDC 72888-231-01 - 100 Tablets Label



Each tablet contains:
glipiZIDE, USP.....15mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at <https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-231-01

glipiZIDE Tablets, USP

15 mg

100 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

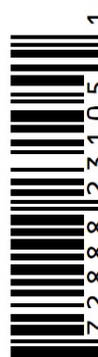
Mfg Lic No. : KD-682

Rev.01, 02/26

Overprinting Area
45 mm X 20 mm



Glipizide Tablets 15 mg - NDC 72888-231-05 - 500 Tablets Label



Each tablet contains:
glipiZIDE, USP.....15mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at <https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-231-05

glipiZIDE Tablets, USP

15 mg

500 Tablets Rx Only

Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.01, 02/26

Overprinting Area
50 mm X 25 mm



Glipizide Tablets 15 mg - NDC 72888-231-00 - 1000 Tablets Label



Each tablet contains:
glipiZIDE, USP....15mg

DOSAGE AND USE:
See accompanying prescribing information.

Store at 20° to 25°C (68° to 77°F); excursions permitted to 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature].

Dispense in a tightly closed container using child resistant closure.

Keep this and all medication out of the reach of children.

Package Insert available at <https://www.advagenpharma.com/pil/glipizidetables>

NDC 72888-231-00

glipiZIDE Tablets, USP

15 mg

1000 Tablets

Rx Only



Distributed by: **Advagen Pharma Ltd.**,
East Windsor, NJ 08520, USA

XXXX

Manufactured by: **Rubicon Research Ltd.**,
Thane 421506, India.

Mfg Lic No. : KD-682

Rev.01, 02/26



Overprinting
Area
50 mm X 25 mm



GLIPIZIDE

glipizide tablet

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:72888-115
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
GLIPIZIDE (UNII: X7WDT95N5C) (GLIPIZIDE - UNII:X7WDT95N5C)	GLIPIZIDE	2.5 mg

Inactive Ingredients

Ingredient Name	Strength
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CROSCARMELOSE SODIUM (UNII: M28OL1HH48)	
ANHYDROUS LACTOSE (UNII: 3SY5LH9PMK)	
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)	
STEARIC ACID (UNII: 4ELV7Z65AP)	

Product Characteristics

Color	white (white to off-white)	Score	no score
Shape	ROUND	Size	6mm
Flavor		Imprint Code	UpArrowhead;I1
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72888-115-30	30 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
2	NDC:72888-115-01	100 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
3	NDC:72888-115-05	500 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
4	NDC:72888-115-	1000 in 1 BOTTLE; Type 0: Not a Combination	10/03/2023	

00	Product	10/03/2023	
Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA214874	10/03/2023	

GLIPIZIDE

glipizide tablet

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:72888-116
Route of Administration	ORAL		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
GLIPIZIDE (UNII: X7WDT95N5C) (GLIPIZIDE - UNII:X7WDT95N5C)	GLIPIZIDE	5 mg

Inactive Ingredients	
Ingredient Name	Strength
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CROSCARMELOSE SODIUM (UNII: M28OL1HH48)	
ANHYDROUS LACTOSE (UNII: 3SY5LH9PMK)	
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)	
STEARIC ACID (UNII: 4ELV7Z65AP)	

Product Characteristics			
Color	white (white to off-white)	Score	2 pieces
Shape	ROUND	Size	8mm
Flavor		Imprint Code	UpArrowhead;I2
Contains			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72888-116-30	30 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
2	NDC:72888-116-01	100 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
3	NDC:72888-116-05	500 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
4	NDC:72888-116-00	1000 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date

ANDA	ANDA214874	10/03/2023	
------	------------	------------	--

GLIPIZIDE

glipizide tablet

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:72888-117
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
GLIPIZIDE (UNII: X7WDT95N5C) (GLIPIZIDE - UNII:X7WDT95N5C)	GLIPIZIDE	10 mg

Inactive Ingredients

Ingredient Name	Strength
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CROSCARMELOSE SODIUM (UNII: M28OL1HH48)	
ANHYDROUS LACTOSE (UNII: 3SY5LH9PMK)	
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)	
STEARIC ACID (UNII: 4ELV7Z65AP)	

Product Characteristics

Color	white (white to off-white)	Score	2 pieces
Shape	ROUND	Size	10mm
Flavor		Imprint Code	UpArrowhead;I4
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72888-117-30	30 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
2	NDC:72888-117-01	100 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
3	NDC:72888-117-05	500 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	
4	NDC:72888-117-00	1000 in 1 BOTTLE; Type 0: Not a Combination Product	10/03/2023	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA214874	10/03/2023	

GLIPIZIDE

glipizide tablet

Product Information				
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:72888-231	
Route of Administration	ORAL			
Active Ingredient/Active Moiety				
	Ingredient Name	Basis of Strength	Strength	
	GLIPIZIDE (UNII: X7WDT95N5C) (GLIPIZIDE - UNII:X7WDT95N5C)	GLIPIZIDE	15 mg	
Inactive Ingredients				
	Ingredient Name		Strength	
	SILICON DIOXIDE (UNII: ETJ7Z6XBU4)			
	CROSCARMELOSE SODIUM (UNII: M28OL1HH48)			
	ANHYDROUS LACTOSE (UNII: 3SY5LH9PMK)			
	MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)			
	STEARIC ACID (UNII: 4ELV7Z65AP)			
Product Characteristics				
Color	white (white to off-white)	Score	2 pieces	
Shape	ROUND	Size	12mm	
Flavor		Imprint Code	UpArrowhead;I11	
Contains				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72888-231-01	100 in 1 BOTTLE; Type 0: Not a Combination Product	02/04/2026	
2	NDC:72888-231-05	500 in 1 BOTTLE; Type 0: Not a Combination Product	02/04/2026	
3	NDC:72888-231-00	1000 in 1 BOTTLE; Type 0: Not a Combination Product	02/04/2026	
Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
ANDA	ANDA214874	10/03/2023		

Labeler - Advagen Pharma Limited (051627256)

Establishment

Name	Address	ID/FEI	Business Operations
Rubicon Research Limited		677604197	manufacture(72888-115, 72888-116, 72888-117, 72888-231) , analysis(72888-115, 72888-116, 72888-117, 72888-231) , label(72888-115, 72888-116, 72888-117, 72888-231) , pack(72888-115, 72888-116, 72888-117, 72888-231)