APO-Clonidine

NAME OF THE MEDICINE

Active ingredient: Clonidine hydrochloride
Chemical name: 2,6-dichloro-N-2-imidazolidinylidene-benzenamine hydrochloride
Molecular formula: C₉H₉N₃Cl₂.HCl
Molecular weight: 266.56
CAS number: 4205-91-8
Laboratory designation: ST 155anaesthesia

Structural formula:

![Structural formula of Clonidine hydrochloride]

DESCRIPTION

Clonidine hydrochloride is a white or almost white, crystalline powder. It is soluble in ethanol, slightly soluble in chloroform and practically insoluble in ether. One gram is soluble in 13 mL of water (20°C)

APO-Clonidine tablets contain 100 micrograms of clonidine hydrochloride.

APO-Clonidine tablets contain the excipients Allura Red AC, hyprolose, microcrystalline cellulose, magnesium stearate, maize starch, lactose monohydrate, calcium hydrogen phosphate, colloidal anhydrous silica.

PHARMACOLOGY

Mode of Action

Antihypertensive effect

Clonidine hydrochloride is predominantly an antihypertensive agent whose mechanism of action appears to be central alpha₂ adrenergic stimulation, as demonstrated in animal studies. This results in the inhibition of bulbar sympathetic cardioaccelerator and sympathetic vasoconstrictor centres, thereby causing a decrease in sympathetic outflow from the brain. There is an increase in vagal activity which produces a decrease in heart rate. There is also an increase in baroreceptor activity. Additionally clonidine hydrochloride stimulates peripheral alpha₁ adrenergic receptors. This is reflected by a small transient pressor effect (5-10 mmHg systolic blood pressure) following parenteral use. A transient rise in blood sugar may also occur following large doses of clonidine hydrochloride. The peripheral effects of clonidine
hydrochloride generally require isolated organ type preparations for their demonstration, as in the intact animal or man, the central action predominates.

Use in migraine prophylaxis and menopausal flushing

Smaller doses of clonidine hydrochloride may be used for migraine prophylaxis and the alleviation of symptoms in menopausal flushing. The mechanism of action appears to be modification of the response of peripheral blood vessels to vasoconstrictor and vasodilator stimuli including noradrenaline, isoprenaline and angiotensin.

Pharmacokinetic Studies

Absorption and distribution

The pharmacokinetics of clonidine is dose-proportional in the range of 75-300 micrograms. Clonidine, the active ingredient of clonidine hydrochloride, is well absorbed from the gastrointestinal tract and undergoes a minor first pass effect. Peak plasma concentrations are reached within 1-3 hours after oral administration. The duration of action varies from 6-12 hours, the duration of action being longer in the milder hypertensives. The plasma protein binding is 30-40%.

Metabolism and excretion

The terminal elimination half-life of clonidine has been found to range from 9-26 hours in patients with normal renal function. With impaired renal function it has been reported to increase to 18-48 hours.

The metabolic pathway of clonidine involves cleavage of the imidazolidine ring and the hydroxylation of the phenyl ring. Five metabolites have been identified in man and include para-hydroxy-clonidine and dichlorophenylguanidine.

Two-thirds of an administered dose is excreted in the urine (about half of which is unchanged Clonidine hydrochloride) and the remainder is excreted in the faeces.

The antihypertensive effect is reached at plasma concentrations between about 0.2 and 2.0 ng/mL in patients with normal renal function. The hypotensive effect is attenuated or decreases with plasma concentrations above 2.0 ng/mL.

Given intravenously, clonidine hydrochloride is effective within five minutes, has a maximum hypotensive action within 20 to 30 minutes, and the effect lasts for several hours. Following intramuscular administration, clonidine hydrochloride is effective within 5 to 10 minutes. The maximum hypotensive effect is reached after 75 minutes and the duration of action is approximately 5 hours.

In a study designed to evaluate the pharmacokinetics of clonidine following administration of clonidine hydrochloride controlled release tablets (formulation not registered in Australia) in 30 patients (13 white patients, 6 black patients and 11 Hispanic patients), the pharmacokinetics was found to be similar between subjects from different racial groups.

The pharmacokinetics of clonidine is not influenced by food.
INDICATIONS

Oral: All grades of essential hypertension.
Renal hypertension.

The prophylactic management of migraine or recurrent vascular headaches which occur in adult patients with a frequency of more than once a month and are not adequately relieved by appropriate therapy for the acute attack. Alleviation of symptoms due to localised vasodilatation in menopausal flushing.

CONTRAINDICATIONS

Clonidine hydrochloride should not be used in patients with known hypersensitivity to the active ingredient, clonidine hydrochloride, and in patients with severe bradyarrhythmia resulting from either sick sinus syndrome or AV block of second or third degree.

In case of rare hereditary conditions that may be incompatible with an excipient of the product (please refer to PRECAUTIONS) the use of the product is contraindicated.

PRECAUTIONS

Special care should be exercised in treating patients who have a history of depression or who have advanced cerebrovascular disease. Reduction of blood pressure in the latter circumstances may itself cause mental changes. Concurrent administration of tricyclic antidepressants may require adjustment of clonidine hydrochloride dosage.

Although a transient rise in blood sugar has been noted occasionally in humans treated with clonidine hydrochloride, which may be due to a pharmacologic alpha-adrenomimetic effect of the drug, no case of induced diabetes mellitus due to clonidine hydrochloride has been reported. Patients with clinical diabetes mellitus should be watched for a possible increase in their requirements of anti-diabetic therapy.

Clonidine hydrochloride should be used with caution in patients with mild to moderate bradyarrhythmia such as low sinus rhythm, with disorders of cerebral or peripheral perfusion, polyneuropathy, and constipation.

No therapeutic effect of clonidine hydrochloride can be expected in the treatment of hypertension caused by phaeochromocytoma.

Since clonidine hydrochloride and its metabolites are extensively excreted in the urine, careful adjustment of dosage is required in patients with renal insufficiency (see Dosage and Administration, Renal insufficiency).

As with other anti-hypertensives, treatment with clonidine hydrochloride should be monitored particularly carefully in patients with heart failure or severe coronary heart disease.

Termination of oral therapy should be gradual (e.g. over more than 7 days).

Sudden cessation of antihypertensive therapy is known to be associated in some instances with rebound hypertension which in some cases may be severe. This may occur with clonidine hydrochloride particularly in patients receiving more than the maximum recommended dose of 900 micrograms per day.

Following sudden discontinuation of clonidine hydrochloride after prolonged treatment with high doses, restlessness, palpitations, rapid rise in blood pressure, nervousness, tremor,
headache or nausea have been reported.

An excessive rise in blood pressure following discontinuation of clonidine hydrochloride therapy can be reversed by intravenous phentolamine (see Interactions with other medicines).

If long-term treatment with a β-blocker needs to be interrupted, the β-blocker should be gradually phased out first, then clonidine.

Patients who wear contact lenses should be warned that treatment with clonidine hydrochloride may cause decreased lacrimation.

APO-Clonidine tablets contain 209.8 mg of Lactose per maximum recommended daily dose. Patients with the rare hereditary conditions of galactose intolerance e.g. galactosaemia should not take this medicine.

**Anaesthesia**

Abrupt withdrawal of clonidine hydrochloride is undesirable. Limited evidence suggests that it is unnecessary to withdraw clonidine hydrochloride before anaesthesia and that maintenance of therapy is preferable to abrupt withdrawal. In the peri-operative period clonidine hydrochloride can, where necessary, be administered parenterally until oral therapy is resumed.

Where therapy with clonidine hydrochloride is to be suspended before operation, withdrawal should be gradual (i.e. over more than 7 days) and monitored by regular observation of blood pressure.

**Menopausal Flushing**

The efficacy of clonidine in the treatment of menopausal flushing has only been demonstrated in the first year after onset of symptoms.

**Effects on Fertility**

Clinical studies on the effect of clonidine on human fertility have not been conducted.

Clonidine had no effect on fertility in male or female rats when administered orally at doses up to 0.15 mg/kg/day (35% higher than the maximum recommended total daily dose of clonidine in humans, based on body surface area).

**Use in Pregnancy (Category B3)**

Clonidine hydrochloride has not shown teratogenic potential when tested in rats, but in some circumstances the incidence of embryonic and perinatal deaths was increased with doses comparable to those used clinically for antihypertensive therapy.

There are limited data from the use of clonidine in pregnant women, but the experience with clonidine hydrochloride since marketing does not include any positive evidence of adverse effect on the foetus. Since this experience cannot exclude such an effect, clonidine hydrochloride should be used during pregnancy only when the benefit clearly justifies the possible risk to the foetus.

Clonidine passes the placental barrier, and may lower the heart rate of the foetus. There is no adequate experience regarding the long-term effects of prenatal exposure.

Clonidine hydrochloride may also induce transitory elevation of blood glucose and impairment of glucose tolerance. Children born to mothers treated with clonidine hydrochloride during pregnancy should be specifically examined for changes in glucose
metabolism. During pregnancy the oral forms of clonidine are preferred. Intravenous injection of clonidine should be avoided.

Non-clinical studies in rats do not indicate direct or indirect harmful effects with respect to reproductive toxicity (see Effects on Fertility).

Post partum a transient rise in blood pressure in the newborn cannot be excluded.

Use in Lactation

Clonidine is excreted in human milk. As the effect on the newborn is not known, infants born to mothers being treated with clonidine hydrochloride should not be breast fed.

Use in children and adolescents

The use and the safety of clonidine in children and adolescents has little supporting evidence in randomised controlled trials and therefore can not be recommended for use in this population.

In particular, when clonidine is used off-label concomitantly with methylphenidate in children with ADHD, serious adverse reactions, including death, have been observed. Therefore, clonidine in this combination is not recommended.

Genotoxicity

Comprehensive investigations have not been performed to assess the potential genotoxic effects of clonidine. Clonidine showed no activity in the Ames test for mutagenicity or mouse micronucleus test for clastogenicity.

Carcinogenicity

The carcinogenic potential of clonidine has not been assessed in an adequate range of studies. In rats, dietary administration of clonidine at doses up to 1.2 mg/kg/day (males) or 1.5 mg/kg/day (females) did not cause carcinogenic effects. These doses are 10-14 times the maximum recommended human daily dose of clonidine, based on body surface area.

Effects on Ability to Drive and Use Machines

No studies on the effects on the ability to drive and use machines have been performed.

However, patients should be advised that they may experience undesirable effects such as dizziness, sedation and accommodation disorder during treatment with clonidine hydrochloride. Therefore, caution should be recommended when driving a car or operating machinery. If patients experience the above mentioned side effects they should avoid potentially hazardous tasks such as driving or operating machinery.

INTERACTIONS WITH OTHER MEDICINES

If the patient is on antihypertensive therapy, care should be taken as even a small dose of clonidine may further lower blood pressure and necessitate adjustment of the antihypertensive regime.

When clonidine hydrochloride is used as an antihypertensive agent, additional clonidine for the prophylaxis of migraine or the alleviation of symptoms in menopausal flushing should not
be prescribed. Clonidine hydrochloride may potentiate the effects of alcohol, sedatives, hypnotics or other centrally active substances.

Although retinal, lens or corneal damage have not been detected with clonidine therapy, follow up procedures, such as ophthalmoscopy, are recommended.

Substances which raise blood pressure or induce a sodium and water retaining effect such as nonsteroidal anti-inflammatory drugs can reduce the therapeutic effect of clonidine.

Substances with α₂-adrenergic receptor blocking properties, such as phentolamine, may abolish the α₂-adrenergic receptor mediated effects of clonidine in a dose-dependent way.

Concomitant administration of drugs with a negative chronotropic or dromotropic effect such as β-blockers or digitalis glycosides can cause or potentiate bradycardiac rhythm disturbances.

It cannot be ruled out that concomitant administration of a β-blocker will cause or potentiate peripheral vascular disorders.

The antihypertensive effect of clonidine may be reduced or abolished and orthostatic regulation disturbances may be provoked or aggravated by concomitant administration of tricyclic antidepressants or neuroleptics with α-receptor blocking effects.

Based on observations in patients in a state of delirium alcoholicum, it has been suggested that high intravenous doses of clonidine may increase the arrhythmogenic potential (QT-prolongation, ventricular fibrillation) of high intravenous doses of haloperidol.

**ADVERSE EFFECTS**

The following adverse events (regardless of causality) and incidences are based on a review of 22 clinical studies comprising 640 patients treated with clonidine hydrochloride.

*Endocrine disorders:*
  ≥0.01% and <0.1%  gynaecomastia

*Psychiatric disorders:*
  ≥1% and <10%  depression, sleep disorder
  ≥0.1% and <1%  delusional perception, hallucination, nightmare

*Not known*
  confusional state, libido decreased

*Nervous system disorders:*
  ≥10%  dizziness, sedation
  ≥1% and <10%  headache
  ≥0.1% and <1%  paraesthesia

*Eye disorder:*
  ≥0.01% and <0.1%  lacrimation decreased

*Not known*
  accommodation disorder
Cardiac disorders:
≥0.1% and <1% sinus bradycardia
≥0.01% and <0.1% atrioventricular block
Not known bradyarrhythmia

Vascular disorders:
≥10% orthostatic hypotension
≥0.1% and <1% Raynaud's phenomenon

Respiratory, thoracic and mediastinal disorders:
≥0.01% and <0.1% nasal dryness

Gastrointestinal disorders:
≥10% dry mouth
≥1% and <10% constipation, nausea, salivary gland pain, vomiting
≥0.01% and <0.1% colonic pseudo-obstruction

Skin and subcutaneous tissue disorders:
≥0.1% and <1% pruritus, rash, urticaria
≥0.01% and <0.1% alopecia

Reproductive system and breast disorders:
≥1% and <10% erectile dysfunction

General disorders and administration site conditions:
≥1% and <10% fatigue
≥0.1% and <1% malaise

Investigations:
≥0.01% and <0.1% blood glucose increased

Most adverse effects are mild and tend to diminish with continued therapy.

Occasional reports of abnormal liver function tests and cases of hepatitis have also been reported.

DOSAGE AND ADMINISTRATION

The dosage recommendations are as follows:

Antihypertensive - initially 50-100 micrograms two to three times daily adjusted in small increments according to the patient's individual blood pressure response. If adequate control is not achieved with a daily dose of 600 micrograms of clonidine hydrochloride alone, additional therapy should be considered. Since the hypotensive effect of clonidine hydrochloride is dose dependent, it is usual to titrate the dose to satisfy the requirements for each patient. In impaired renal and hepatic function the half-life is prolonged and the dosage regime should be monitored carefully.

Migraine prophylaxis - initially 25 micrograms morning and evening. If necessary, after two
weeks, this may be increased to 50 micrograms twice daily, then to a total daily dose of 150 micrograms. If the frequency of attacks is significantly reduced, consideration may be given to gradually ceasing therapy as remission may be sustained in a proportion of patients. Duration of treatment will depend upon the frequency and severity of attacks.

Menopausal flushing - initially 25 micrograms morning and evening. If after two weeks there has been no remission, increase to 50 micrograms twice daily. If necessary this may be increased to a total daily dose of 150 micrograms. Duration of treatment will depend upon the frequency and severity of attacks but long-term efficacy (longer than 8 weeks) in the treatment of menopausal flushing has not been established.

Renal insufficiency
Dosage must be adjusted:
- according to the individual antihypertensive response which can show high variability in patients with renal insufficiency
- according to the degree of renal impairment.

Careful monitoring is required. Since only a minimal amount of clonidine is removed during routine haemodialysis, there is no need to give supplemental clonidine following dialysis.

OVERDOSAGE
For information on the management of overdose, contact the Poisons Information Centre on 13 11 26 (Australia).

Symptoms
The most important features of clonidine overdose are likely to be bradycardia and sedation, respiratory depression including apnoea and somnolence including coma. Blood pressure response may be variable and may vary from severe hypotension (due to central sympathetic inhibition and vagal stimulation) to severe hypertension (due to direct alpha agonist activity). Treatment must therefore be appropriate to the clinical features (i.v. atropine followed by a pressor amine if necessary in patients with hypotension or an alpha blocker such as phentolamine for patients with hypertension). Other features which may be seen include weakness, vomiting, diminished or absent reflexes, skin pallor, hypothermia, cardiac arrhythmias and constricted pupils with poor reaction to light.

Management
General supportive measures with regular checks of pulse, B.P., ECG, blood sugar and body temperature should be undertaken. The blood pressure should be monitored carefully for 48 hours following the overdose, as a later hypertensive phase may be associated with declining blood levels of clonidine.

PRESENTATION AND STORAGE CONDITIONS
APO-Clonidine tablets are scored, pink, compressed tablets, impressed with the symbol YS 01 on one side and the score line on the reverse side. Each tablet contains 100 micrograms of clonidine hydrochloride.

APO-Clonidine tablets are available in HDPE bottles fitted with PP child resistant closure containing 100 tablets.

APO-Clonidine tablets should be stored below 25°C.
NAME AND ADDRESS OF THE SPONSOR

Southern Cross Pharma Pty Ltd
56 Illabunda Dr
Malua Bay, NSW
2536 Australia

POISON SCHEDULE OF THE MEDICINE

S4 – Prescription Only Medicine

DATE OF FIRST INCLUSION ON THE AUSTRALIAN REGISTER OF THERAPEUTIC GOODS:

24th November 2016