Cordarone® (amiodarone HCI) **TABLETS** 

DESCRIPTION

Cordarone is a member of a new class of antiarrhythmic drugs with predominantly Class III (Vaughan Williams' classification) effects, available for oral administration as

> Cordarone® (amiodarone HCI) Tablets CI 8050-

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pink, scored tablets containing 200 mg of amiodarone hydrochloride. The inactive ining solvet daules containing 20 mig of announce in environment in the indice. The integrates in ingredients present are colloidal silicon dioxide, lactose, magnesium stearate, povidone starch, and FD&C Red 40. Cordarone is a benzofuran derivative: 2-butyl-3-benzofuranyl 4-[2-(diethylamino)-ethoxy]-3,5-diiodophenyl ketone hydrochloride. It is not chemically related to any other available antiarrhythmic drug.

Amiodarone HCl is a white to cream-colored crystalline powder. It is slightly soluble in water, soluble in alcohol, and freely soluble in chloroform. It contains 37.3% iodine by

CLINICAL PHARMACOLOGY

Electrophysiology/Mechanisms of Action

Electrophysiologymecialanisms of Action
In animals, Cordarone is effective in the prevention or suppression of experimentally induced arrhythmias. The antiarrhythmic effect of Cordarone may be due to at least two major properties: 1) a prolongation of the myocardial cell-action potential duration and refractory period and 2) noncompetitive α- and β-adrenergic inhibition.

Cordarone prolongs the duration of the action potential of all cardiac fibers while caus ing minimal reduction of d'Vid (maximal upstroke velocity of the action potential). The refractory period is prolonged in all cardiac tissues. Cordarone increases the cardiac refractory period without influencing resting membrane potential, except in automatic cells where the slope of the prepotential is reduced, generally reducing automaticity. These electrophysiologic effects are reflected in a decreased sinus rate of 15 to 20% These electrophysiologic energs are reflected in a decreased sinks face or 19 to 20%, increased PR and OT intervals of bout 10%, the development of U-waves, and changes in T-wave contour. These changes should not require discontinuation of Container as they are evidence of its pharmacological action, although Condainer an cause marked sinus bradycardia or sinus arrest and heart block. On rare occasions, OT prolongation has been associated with worsening of arrhythmia (see "WARNINGS").

Hemodynamics
In animal studies and after intravenous administration in man. Cordarone relaxes vas cular smooth muscle, reduces peripheral vascular resistance (afterload), and slightly increases cardiac index. After oral dosing, however, Cordarone produces no significan change in left ventricular ejection fraction (LVEF), even in patients with depressed LVEF After acute intravenous dosing in man. Cordarone may have a mild negative inotropic

**Pharmacokinetics** 

Filluming oral administration in man, Cordarone is slowly and variably absorbed. The bloavailability of Cordarone is approximately 50%, but has varied between 35 and 65% in various studies. Maximum plasma concentrations are attained 3 to 7 hours after a single dose. Despite this, the onset of action may occur in 2 to 3 days, but more commonly takes 1 to 3 weeks, even with loading doses. Plasma concentrations with chronic dosing at 100 to 600 mg/day are approximately dose proportional, with a mean 0.5 mg/L ousning at 100 to ob unjevaly are approximately obes proportional, win a mean J.S. m increase for each 100 mg/dg. These means, however, include considerable indi-vidual variability. Food increases the rate and extent of absorption of Cordarons effects of food upon the bioavailability of Cordarone have been studied in 30 healthy subjects who received a single 600-mg dose immediately after consuming a high-lat meal and following an overnight flast. The area under the plasma concentration-time curve (ALIC) and the peak plasma concentration (Conv.) of amindarone increased by 2.3 conve (not) allow the peak plasma collection and on toward or manufacture microsists of years (range 1.7 to 3.6) and 3.8 (range 1.7 to 3.6) and 3.8 (range 1.7 to 4.4) times, to specifively, in the presence of food. Food also increased the rate of aboration of amidoanne, decreasing the time peak plasma connectration (r<sub>max</sub>) of y5.7%. The mean AUC and mean Cr<sub>max</sub> of desethylamiodarone increased by 5.5% (range 5.0 to 10%) and 32% (range 4.to 40.4%), respectively, but there was no change in the T<sub>max</sub> in the presence of food. Cordarone has a very large but variable volume of distribution, averaging about 60 L/kg

because of extensive accumulation in various sites, especially adjoose tissue and highly perfused organs, such as the liver, lung, and spleen. One major metabolite of Cordarone, desethylamiodarone (DEA), has been identified in man; it accumulates to an even greater extent in almost all tissues. No data are available on the activity of DFA in humans, but in animals, it has significant electrophysiologic and antiarrhythmic effects generally similar to amiodarone itself. DEA's precise role and contribution to the antiar generally similar of amilotation research 20 per better the dividence of manufacturing representations and amilotation are not certain. The dividence of maximal ventricular Class III effects after oral Cordarone administration in humans correlates more closely with DEA accumulation over time than with amilodarone accumulation.

Amiodarone is eliminated primarily by hepatic metabolism and biliary excretion and there is negligible excretion of amiodarone or DEA in urine. Neither amiodarone or DEA in urine. Neither amiodarone or DEA is dialyzable.

In clinical studies of 2 to 7 days, clearance of amiodarone after intravenous administrain clinical studies of a 7 bd/s, cleanate or annovation after finite ventions commission to nin patients with VT and VF ranged between 220 and 440 mth/Rig. Age, exe, renal disease, and hepatic disease (cirrhosis) do not have marked effects on the disposition of amiodarone or DEA Renal impairment does not influence the pharmacokinetiss of amiodarone. After a single dose of intravenous amiodarone in cirrhotic patients, significantly lower Constant average concentration values are seen for DFA, but mean amin cainty lower large, and unchanged. Mornal subjects over 65 years of age 50-year of mental almod-darone levels are unchanged. Mornal subjects over 65 years of age 50-year of mental almod-darone levels and to 10 mlm/kbg) than young explicit subjects (about 150 mlm/kbg) and an increase in 11,0 mlm about 20 to 479. In patients with severe left ventricular dys-function, the pharmacokin dependence of amiddarone are not significantly altered but the ter-minal disposition in 2,0 dt DEA is prolonged. Although no dosage adjustment for patients with renal, hepatic, or cardiac abnormalities has been defined during chronic treatment with Cordarone (amiodarone HCI) Tablets, close clinical monitoring is prudent for elderly natients and those with severe left ventricular dysfunction

Following single dose administration in 12 healthy subjects, Cordarone exhibited multicompartmental pharmacokinetics with a mean apparent plasma terminal elimination Configuration rich primarization 12 days) for amount makes a paper layer particular the many and the state of 107 days, with a mean of approximately 53 days and most patients in the 40- to 55-day range. In the absence of a loading-dose period, steady-state plasma concentrations, at constant oral dosing, would therefore be reached between 130 and 535 days, with an average of 265 days. For the metabolite, the mean plasma-elimination half-life was average or 265 days. For the metabolite, the mean plasma-elimination nair-life was approximately 61 days. These data probably reflect an initial elimination of drug from well-perfused tissue (the 2.5- to 10-day half-life phase), followed by a terminal phase representing extremely slow elimination from poorly perfused tissue compartments such

The considerable intersubject variation in both phases of elimination, as well as unce tainty as to what compartment is critical to drug effect, requires attention to individual responses once arrhythmia control is achieved with loading doses because the correct maintenance dose is determined, in part, by the elimination rates. Daily maintenance doses of Cordarone should be based on individual patient requirements (see "DOSAGE AND ADMINISTRATION")

Cordarone and its metabolite have a limited transplacental transfer of approximately 10 to 50%. The parent drug and its metabolite have been detected in breast milk Cordarone is highly protein-bound (approximately 96%).

Although electrophysiologic effects, such as prolongation of QTc, can be seen within hours after a parenteral dose of Cordarone effects on abnormal rhythms are not seen nours arer a parenterial dose of Cordarone, effects on annormal mynnins are not seen before 2 to 3 days and usually require 1 to 3 weeks, even when a loading dose is used. There may be a continued increase in effect for longer periods still. There is evidence that the time to effect is shorter when a loading-dose regimen is used. Consistent with the slow rate of elimination, antiarrhythmic effects persist for weeks or consistent with the source of enimination, antiarrhyminic electics persist of weeks or months after Cordarone is discontinued, but the time of recurrence is variable and unpredictable. In general, when the drug is resumed after recurrence of the arrhythmia, control is established relatively rapidly compared to the initial response, presumably because tissue stores were not wholly depleted at the time of recurrence.

Pharmacodynamics

There is no well-established relationship of plasma concentration to effectiveness, but it does appear that concentrations much below 1 mg/L are often ineffective and that levels above 2.5 mg/L are generally not needed. Within individuals dose reductions and ensuing decreased plasma concentrations can result in loss of arrhythmia control ensuing decreased plasma concentrations can resuit in loss or arrhythmia control. Plasma-concentration measurements can be used to identify patients whose levels are unusually low, and who might benefit from a dose increase, or unusually high, and who might have dosage reduction in the hope of minimizing side effects. Some observations have suggested a plasma concentration, dose, or dose/duration relationship for side effects such as pulmonary fibrosis, liver-enzyme elevations, corneal deposits and facial pigmentation, peripheral neuropathy, gastrointestinal and central nervous system

Monitoring Effectiveness

monitoring cleatments of any antiarrhythmic agent in long-term prevention of recur-rent ventricular tachycardia and ventricular fibrillation is difficult and controversial, with highly qualified investigators recommending use of ambulatory monitoring, programmed electrical stimulation with various stimulation regimens, or a combination of these, to assess response. There is no present consensus on many aspects of how best to assess effectiveness, but there is a reasonable consensus on some aspects:

 If a natient with a history of cardiac arrest does not manifest a hemodynamically If a patient with a history of cardiac arrest does not manifest a hemodynamically unstable arriythmia during electrocardiorgaphic monitoring prior to treatment, assess-ment of the effectiveness of Cordarone requires some provocative approach, either sercises or programmed electrical stimulation (PES).
 Whether provocation is also needed in patients who to manifest their life-threatening arriythmia spontaneously is not settled, but there are reasons to consider PES or other

provocation in such natients. In the fraction of natients whose PES-inducible arrhythmia can be made noninducible by Cordarone (a fraction that has varied widely in vari-ous series from less than 10% to almost 40%, perhaps due to different stimulation criteria), the prognosis has been almost uniformly excellent, with very low recurrence (ventricular tachycardia or sudden death) rates. More controversial is the meaning of continued inducibility. There has been an impression that continued inducibility in commune inductionity. There has been an impression that commune inductionity in Cordarone patients may not foretell a poor prognosis but, in fact, many observers have found greater recurrence rates in patients who remain inducible than in those who do not. A number of criteria have been proposed, however, for identifying patients who remain inducible but who seem likely nonetheless to do well on Cordarone. These criteria include increased difficulty of induction (more stimuli or more rapid stimuli), which has been reported to predict a lower rate of recurrence, and ability to tolerate the

induced ventricular tachycardia without severe symptoms, a finding that has been reported to correlate with better survival but not with lower recurrence rates. While these criteria require confirmation and further study in general, easier inducibility or poorer tolerance of the induced arrhythmia should suggest consideration of a need

Several predictors of success not based on PES have also been suggested, including complete elimination of all nonsustained ventricular tachycardia on ambulatory monitoring and very low premature ventricular-beat rates (less than 1 VPB/1,000 normal beats). While these issues remain unsettled for Cordarone, as for other agents, the prescriber of Cordarone should have access to (direct or through referral), and familiarity with, the full range of evaluatory procedures used in the care of patients with life-threatening

It is difficult to describe the effectiveness rates of Cordanne, as these depend on the specific arrhythmia treated, the success criteria used, the underlying cardiac disease of specinic arrityminal readers, the success clients used, in the interpring claudiac baseser of the patient, the number of drugs tride before resorting to Cordarone, the duration of follow-up, the dose of Cordarone, the use of additional antiarrhythmic agents, and many other factors. As Cordarone has been studied principally in patients with refractory life-threatening ventricular arrhythmias, in whom drug therapy must be selected on the basis of response and cannot be assigned arbitrarily, randomized comparisons with other agents or placeho have not been possible. Reports of series of treated patients with a his agents or piaceon have not been possible. Reports or series or treated patients with a nis-tory of cardiac arrest and mean follow-up of one year or more have given mortality (due to arrhythmia) rates that were highly variable, ranging from less than 5% to over 30%, with most series in the range of 10 to 15%. Overall arrhythmia-recurrence rates (fatal and nonfatal) also were highly variable (and, as noted above, depended on response to PES and other measures), and depend on whether natients who do not seem to respond in tally are included. In most cases, considering only patients who seemed to respond well enough to be placed on long-term treatment, recurrence rates have ranged from 20 to 40% in series with a mean follow-up of a year or more

INDICATIONS AND USAGE

Because of its life-threatening side effects and the substantial management difficulties associated with its use (see "WARNINGS" below), Cordarone is indicated only for the treatment of the following documented, life-threatening recurrent ventricular arrhyth mias when these have not responded to documented adequate doses of other available antiarrhythmics or when alternative agents could not be tolerated

Recurrent ventricular fibrillation

2. Recurrent hemodynamically unstable ventricular tachycardia. As is the case for other antiarrhythmic agents, there is no evidence from controlled trials that the use of Cordarone (amiodarone HCI) Tablets favorably affects survival. Cordarone should be used only by physicians familiar with and with access to (directly or through referral) the use of all available modalities for treating recurrent life threatening ventricular arrhythmias, and who have access to appropriate monitoring facilities, including in-hospital and ambulatory continuous electrocardiographic mon toring and electrophysiologic techniques. Because of the life-threatening nature of the arrhythmias treated, potential interactions with prior therapy, and potential exacerbatic of the arrhythmia, initiation of therapy with Cordarone should be carried out in the

CONTRAINDICATIONS

rdarone is contraindicated in severe sinus-node dysfunction, causing marked sinus bradycardia; second- or third-degree atrioventricular block; and when episodes of brady-cardia have caused syncope (except when used in conjunction with a pacemaker). Cordarone is contraindicated in patients with a known hypersensitivity to the drug or to any of its components, including jodine.

WARNINGS

Cordarone is intended for use only in natients with the indicated life-threatening arrhythmias because its use is accompanied by substantial toxicity.

Cordarone has several potentially fatal toxicities, the most important of which is Loffrazine has severa potentially fast lotacities, the most important or winer is proposed to the property of the proposed to the property of only by abnormal liver enzymes. Overt liver disease can occur, however, and has only by adnormal liver enzymes. Over liver disease can occur, nowever, and not been fallal in a few cases. Like other antiarrythmics, Cordarone can exacerbate the arrhythmia, e.g., by making the arrhythmia less well tolerated or more difficult to reverse. This has occurred in Z to 5% of pallents in various series, and significant heart block or sinus bradycardia has been seen in Z to 5%. All of these events should be manageable in the proper clinical settling in most cases. Although the frequency of such proarrhythmic events does not appear greater with Cordarone than with many other agents used in this population, the effects are prolonged

Even in patients at high risk of arrhythmic death, in whom the toxicity of Cordarone is an acceptable risk, Cordarone poses major management problems Cordarone is an adeeptacle risk, order to provide major inanagement problems that could be life-threatening in a population at risk of sudden death, so that every effort should be made to utilize alternative agents first.

The difficulty of using Cordarone effectively and safely itself poses a significant risk

to patients. Patients with the indicated arrhythmias must be hospitalized while the loading dose of Cordarone is given, and a response generally requires at least one week, usually two or more. Because absorption and elimination are variable. week, issuary two indice. Because assorption and entimation are variable, maintenance-dose selection is difficult, and it is not unusual to require dosage decrease or discontinuation of treatment. In a retrospective survey of 192 patients with ventricular tachyarrhythmias, 84 required dose reduction and 18 required at with Hemitoter decipality minimals, Se Frequent duse relations and to require a have reported 15 to 20% overall frequencies of discontinuation due to adverse reac-tions. The time at which a previously controlled life-threatening arrhythmia will recur after discontinuation or does adjustment is unpredictable, ranging from weeks to months. The patient is obviously at great risk during this time and may need prolonged hospitalization. Altempts to substitute other antiarrythmic agents when Cordarone must be stopped will be made difficult by the gradually, but unpre-dictably, changing amiodarone body burden. A similar problem exists when Cordarone is not effective; it still poses the risk of an interaction with whatever sub-

Morfailly in the National Heart, Lung and Blood Institute's Cardiac Arrhythmia Suppression Trial (CAST), a long-term, multi-centered, randomized, double-blind study in Marchael and Cast (CAST), a long-term, multi-centered, randomized, double-blind study in Marchael Institution, more reliable to the State of the Stat

Cordarone therapy was evaluated in two multi-centered, randomized, double-blind, placebo-controlled trials involving 1202 (Canadian Amidodarone Myocardial Infarction Arrhythmia Trial; CAMIAT) and 1486 (European Myocardial Infarction Internation on Implicit Life Control of the Lead to the Update Implementation of the Control of the Lead to the Update Implementation of the Control of the Lead to the Control of the Con

	Placebo		Amiodarone		Relative Risk	
	N	Deaths	N	Deaths		95%CI
EMIAT	743	102	743	103	0.99	0.76-1.31
CAMIAT	596	68	606	57	0.88	0.58-1.16

These data are consistent with the results of a pooled analysis of smaller, controlled studies involving patients with structural heart disease (including myocardia

**Pulmonary Toxicity** 

Pullinulary Loucity
There have been postmarketing reports of acute-onset (days to weeks) pulmonary injury in patients treated with oral Cordarone with or without initial I.V. therapy. Findings have included pulmonary infilitates or N-ray, bronchospasm, wheezing, fever, dyspnea, cough, hemoptysis, and hypoxia. Some cases have progressed to respiratory failure and/or death. Cordarone (amiodarone HCI) Tablets may cause a clinical syndrome of cough and pro-

Cordatine (anniodatine Hot) Tables may clause a clinical syndrome or cough and pro-gressive dyspinea accompanied by functional, radiographic, gallium-scan, and patholog ical data consistent with pulmonary toxicity, the frequency of which varies from 2 to 7% in most published reports, but is as high as 10 to 17% in some reports. Therefore, when Cordarone therapy is initiated, a baseline chest X-ray and pulmonary-function tests, including diffusion capacity, should be performed. The patient should return for a history, physical exam, and chest X-ray every 3 to 6 months.

Pulmonary toxicity secondary to Cordarone seems to result from either indirect or direct toxicity as represented by hypersensitivity pneumonitis or interstitial/alveola pneumonitis, respectively.

Patients with preexisting pulmonary disease have a poorer prognosis if pulmonary

Hypersensitivity pneumonitis usually appears earlier in the course of therapy, and rechallenging these patients with Cordarone results in a more rapid recurre

Bronchoalveolar lavage is the procedure of choice to confirm this diagnosis, which can be made when a T suppressor/cytotoxic (CD8-positive) lymphocytosis is noted. Steroid therapy should be instituted and Cordarone therapy discontinued in these natients. Interstitial/alveolar pneumonitis may result from the release of oxygen radicals and/or phospholipidosis and is characterized by findings of diffuse alveolar damage, interstitial pneumonitis or fibrosis in lung biopsy specimens. Phospholipidosis (foamy cells. preunionals or initiosis in multiplicity specimens. Prospicitions of a company and program that of the present in most cases of Cordarone-induced pulmonary toxicity, however, these changes also are present in approximately 50% of all palents on Cordarone therapy. These cells should be used as markers of therapy, but not as evidence of toxicity. A diagnosis of Cordarone-induced interstitial/alveolar pneumonitis should lead, at a minimum, to dose reduction or, preferably, to withdrawal of the Cordarone to establish reversibility, especially if other acceptable antiarrhythmic therapies are available. Where these measures have been acceptance animal minum. uneaples are available. Write insert insert instances have been instituted, a reduction in symptoms of amiodarone-induced pulmonary toxicity was usually noted within the first week, and a clinical improvement was greatest in the first two to three weeks. Chest X-ray changes usually resolve within two 16 our months. According to some experts, steroids may prove beneficial. Prednisone in doses of 40 to 60 mg/day or equivalent doses of other steroids have been given and tapered over the course of several weeks depending upon the condition of the patient. In some cases rechallenge with Cordarone at a lower dose has not resulted in return of toxicity. Reports suggest that the use of lower loading and maintenance doses of Cordarone are associated with a decreased incidence of Cordarone-induced pulmonary toxicity. In a patient receiving Cordarone, any new respiratory symptoms should suggest the possibility of pulmonary toxicity, and the history, physical exam, chest X-ray, and pulmonary-function tests (with diffusion capacity) should be repeated and evaluated. A 15% decrease in diffusion capacity has a high sensitivity but only a moderate specificity for pulmonary toxicity; as the decrease in diffusion capacity approaches 30%, the sen-sitivity decreases but the specificity increases. A gallium-scan also may be performed as part of the diagnostic workup.

Fatalities, secondary to pulmonary toxicity, have occurred in approximately 10% of cases. However, in patients with life-threatening arrhythmias, discontinuation of Cordarone therapy due to suspected drug-induced pulmonary toxicity should be under-taken with caution, as the most common cause of death in these patients is sudden cardiac death. Therefore, every effort should be made to rule out other causes of respirationy impairment (i.e., congestive heart failure with Swan-Ganz catheterization if nec-estation, respiratory infection, pulmonary embolism, malignancy, etc.) before discontinu-ing Cordarone in these patients. In addition, bronchoalveolar lavage, transbronchial lung biopsy and/or open lung biopsy may be necessary to confirm the diagnosis, especially in those cases where no acceptable alternative therapy is available If a diagnosis of Cordarone-induced hypersensitivity pneumonitis is made, Cordarone should be discontinued, and treatment with steroids should be instituted. If a diagnosis of Cordarone-induced interstitial/alveolar pneumonitis is made, steroid therapy should be instituted and, preferably, Cordarone discontinued or, at a minimum, reduced in dosage. Some cases of Cordarone-induced interstitial/alyeolar pneumonitis may res olar pneumonitis may resolve following a reduction in Cordarone dosage in conjunction with the administration of steroids. In some patients, rechallenge at a lower dose has not resulted in return of interstitial/alveolar pneumonitis; however, in some patients (perhaps because of severe alveolar damage) the pulmonary lesions have not been reversible.

Worsened Arrhythmia Cordarone, like other antiarrhythmics, can cause serious exacerbation of the presenting arrhythmia, a risk that may be enhanced by the presence of concomitant antiarrhyth-mics. Exacerbation has been reported in about 2 to 5% in most series, and has included new ventricular fibrillation, incessant ventricular tachycardia, increased resistance to cardioversion, and polymorphic ventricular tachycardia associated with QTc prolonga-tion (torsades de pointes [TdP]). In addition, Cordarone has caused symptomatic bradycardia or sinus arrest with suppression of escape foci in 2 to 4% of patients Fluoroquinolones, macrolide antibiotics, and azoles are known to cause QTc prolonga-tion. There have been reports of QTc prolongation, with or without TdP, in patients taking amindarone when fluoroquinolones, macrolide antihiotics, or azoles were administered concomitantly. (See "Drug Interactions, Other reported interactions with amioda-

The need to co-administer amiodarone with any other drug known to prolong the QTc interval must be based on a careful assessment of the potential risks and benefits of doing so for each patient.

A careful assessment of the notential risks and benefits of administering Cordarone must made in patients with thyroid dysfunction due to the possibility of arrhythmia break bugh or exacerbation of arrhythmia in these patients. Liver Injury

Elevations of hepatic enzyme levels are seen frequently in patients exposed to Cordarone and in most cases are asymptomatic. If the increase exceeds three times normal, or doubles in a patient with an elevated baseline, discontinuation of Cordarone or dosage reduction should be considered. In a few cases in which biopsy has been done, the histology has resembled that of alcoholic hepatitis or cirrhosis. Hepatic failure has been a rare cause of death in patients treated with Cordarone

Loss of Vision

Cases of optic neuropathy and/or optic neuritis, usually resulting in visual impairment, have been reported in patients treated with amiodarone. In some cases, visual impairment has progressed to permanent blindness. Optic neuropathy and/or neuritis may occur at any time following initiation of therapy. A causal relationship to the drug has not been clearly established. If symptoms of visual impairment appear, such as changes in visual acuity and decreases in peripheral vision, prompt ophthalmic examination is recommended. Appearance of ontic neuropathy and/or neuritis calls for reevaluation of Cordarone therapy. The risks and complications of antiarrhythmic therapy with Cordarone must be weighed against its benefits in patients whose lives are threatened by cardiac arrhythmias. Regular ophthalmic examination, including funduscopy and slit-lamp examination, is recommended during administration of Cordarone. (See "ADVERSE REACTIONS")

Neonatal Hypo- or Hyperthyroidism

Cordarone can cause fetal harm when administered to a pregnant woman. Although Cordarone use during pregnancy is uncommon, there have been a small number of published reports of congenital golter/hypothyroidism and hyperthyroidism. If Cordarone is used during pregnancy, or if the patient becomes pregnant while taking Cordarone, the patient should be apprised of the potential hazard to the fetus. In general, Cordarone (amiodarone HCI) Tablets should be used during pregnancy only if the notential benefit to the mother justifies the unknown risk to the fetus. In pregnant rats and rabbits, amiodarone HCI in doses of 25 mg/kg/day (approximately 0.4 and 0.9 times, respectively, the maximum recommended human maintenance dose\*) had no adverse effects on the fetus. In the rabbit, 75 mg/kg/day (approximately 2.7 times the maximum recommended human maintenance dose\*) caused abortions in 2.7 times the maximum recommenden unimal mannenance dose\*) caused abortions i greater than 90% of the animals. In the rat, doses of 50 mg/kg/day or more were asso-ciated with slight displacement of the testes and an increased incidence of incomplete ossification of some skull and digital bones; at 100 mg/kg/day or more, fetal body weights were reduced; at 200 mg/kg/day, there was an increased incidence of fetal resorption. (These doses in the rat are approximately 0.8, 1.6 and 3.2 times the maximum recommended human maintenance dose.\*) Adverse effects on fetal growth and survival also were noted in one of two strains of mice at a dose of 5 mg/kg/day (approximately 0.04 times the maximum recommended human maintenance dose\*).

PRECAUTIONS

Impairment of Vision

Optic Neuropathy and/or Neuritis

Cases of optic neuropathy and optic neuritis have been reported (see "WARNINGS"). Corneal Microdeposits

\*600 mg in a 50 kg patient (doses compared on a body surface area basis)

Corneal microdeposits appear in the majority of adults treated with Cordarone. They are usually discernible only by slit-lamp examination, but give rise to symptoms such as visual halos or blurred vision in as many as 10% of patients. Corneal microdeposits are reversible upon reduction of dose or termination of treatment. Asymptomatic microde e are not a reason to reduce dose or discontinue treatment (see "ADVERSE

Neurologic
Chronic administration of oral amiodarone in rare instances may lead to the development of peripheral neuropathy that may resolve when amiodarone is discontinued, but

Photosensitivity
Cordarone has induced photosensitization in about 10% of patients; some protection may be afforded by the use of sun-barrier creams or protective clothing. During longterm treatment, a blue-gray discoloration of the exposed skin may occur. The risk may he increased in patients of fair complexion or those with excessive sun exposure, and may be related to cumulative dose and duration of therapy.

Thyroid Abnormalities Condarone inhibits peripheral conversion of thyroxine  $(T_a)$  to triiodothyronine  $(T_3)$  and and contract ministry per pineta conversion of the youth  $(1_3)$  for inducting of the  $(1_3)$  fand increased levels of inactive reverse  $T_3$  (r $T_3$ ) in clinically euthyroid patients. It is also a potential source of large amounts of inorganic iodine. Because of its release of inorganic iodine, or perhaps for other reasons, Cordarone can cause either hypothyroidism or hyperthyroidism. Thyroid function should be monitored prior to treatment and periodically thereafter, particularly tunction studin be immunited print or treatment and performance in elderly patients, and in any patient with a history of thyroid nodules, goiter, or other thyroid dysfunction. Because of the slow elimination of Cordarone and its metabolites, high plasma iodide levels, altered thyroid function, and abnormal thyroid-function tests may persist for several weeks or even months following Cordarone withdrawal.

thypothyroidism has been reported in 2 to 4% of patients in most series, but in 8 to 10% in some series. This condition may be identified by relevant clinical symptoms and particularly by elevated serum TSH levels. In some clinically hypothyroid amiodarone-treated patients, free thyroxine index values may be normal. Hypothyroidism is best managed by Cordarone dose reduction and/or thyroid hormone supplement. However therapy must be individualized, and it may be necessary to discontinue Cordarone (amiodarone HCl) Tablets in some patients.

Hyperthyroidism occurs in about 2% of patients receiving Cordarone, but the incidence may be higher among patients with prior inadequate dietary iodine intake. Cordarone-induced hyperthyroidism usually poses a greater hazard to the patient than hypothyinduced hyperhyrolism disaday boses a greater nazar to the patient and hypothyr-roldism because of the possibility of arrhythmia breakthrough or aggravation, which may result in death. In fact, IF ANY NEW SIGNS OF ARRHYTHMIA APPEAR, THE POS-SIBILITY OF HYPERTHYROLDISM SHOULD BE CONSIDERED. Hyperthyroidism is best identified by relevant clinical symptoms and signs, accompanied usually by abnormally elevated levels of serum T<sub>3</sub>, RIA, and further elevations of serum T<sub>4</sub>, and a subnormal serum TSH level (using a sufficiently sensitive TSH assay). The finding of a flat TSH response to TRH is confirmatory of hyperthyroidism and may be sought in equivocal response to first sometiment of injecting local many accompany fordarone-induced hyperthy-roldism, aggressive medical treatment is indicated, including, if possible, dose reduc-tion or withdrawal of Cordarone. The institution of antithyroid drugs, \(\theta\)-adrenergic blockers and/or temporary corticosteroid therapy may be necessary. The action of antithyroid drugs may be especially delayed in amindarone-induced thyrotoxicosis anunyinut drugs may be especially delayed in animodatine-moused informations because of substantial quantities of preformed thyroid hormones stored in the gland. Radioactive iodine therapy is contraindicated because of the low radioidoine uptake associated with amiodarone-induced hyperthyroidism. Experience with thyroid surgen, in this setting is extremely limited, and this form of therapy runs the theoretical risk of inducing thyroid storm. Cordarone-induced hyperthyroidism may be followed by a transient period of hypothyroidism.

Volatile Anesthetic Agents: Close perioperative monitoring is recommended in patients undergoing general anesthesia who are on amindarone therapy as they may be more sensitive to the myocardial depressant and conduction effects of halogenated inhala-

Hypotension Postbypass: Rare occurrences of hypotension upon discontinuation of cardiopulmonary bypass during open-heart surgery in patients receiving Cordarone have been reported. The relationship of this event to Cordarone therapy is unknown Adult Respiratory Distress Syndrome (ARDS): Postoperatively, occurrences of ARDS have been reported in patients receiving Cordarone therapy who have undergone either cardiac or noncardiac surgery. Although patients usually respond well to vigorous res-piratory therapy, in rare instances the outcome has been fatal. Until further studies have performed, it is recommended that FiO<sub>2</sub> and the determinants of oxygen delivery to the tissues (e.g., SaO<sub>2</sub>, PaO<sub>2</sub>) be closely monitored in patients on Cordarone.

Information for Patients
Patients should be instructed to read the accompanying Medication Guide each tire
they refill their prescription. The complete text of the Medication Guide is reprinted.

Elevations in liver enzymes (SGOT and SGPT) can occur. Liver enzymes in patients or relatively high maintenance doses should be monitored on a regular basis. Persistent significant elevations in the liver enzymes or henatomenaly should alert the physician to significant elevations in the level enzymes on replacements of consider reducing the maintenance dose of Cordarone or discontinuing therapy.

Cordarone (amiodarone HCI) Tablets alters the results of thyroid-function tests, causing an increase in serum T<sub>4</sub> and serum reverse T<sub>3</sub>, and a decline in serum T<sub>3</sub> levels. Despite these biochemical changes, most patients remain clinically euthyroid.

Drug Interactions

Amindarone is metabolized to desethylamindarone by the cytochrome P450 (CYP450) enzyme group, specifically cytochrome P450 3A4 (CVP3A4) and CVP2C8. The CVP3A4 isoenzyme is present in both the liver and intestines (see "CLINICAL PHARMACOLOGY, Pharmacokinetics"). Amiodarone is also known to be an inhibitor of CYP3A4.

Therefore, amiodarone has the potential for interactions with drugs or substances that may be substrates, inhibitors or inducers of CYP3A4. While only a limited number of in vivo drug-drug interactions with amiodarone have been reported, the potential for other interactions should be anticipated. This is especially important for drugs associated with serious toxicity, such as other antiarrhythmics. If such drugs are needed, their dose should be reassessed and, where appropriate, plasma concentration measured. In view of the long and variable half-life of amiodarone, potential for drug interactions exists, not only with concomitant medication, but also with drugs administered after discontinuation of amindarone

Since amiodarone is a substrate for CYP3A4 and CYP2C8, drugs/substances that inhibit CVP3A4 may decrease the metabolism and increase serum concentrations of

### amindarone. Reported examples include the following:

### Protease inhibitors:

Protease inhibitors are known to inhibit CYP3A4 to varying degrees. A case report of one patient taking amiodarone 200 mg and indinavir 800 mg three times a day resulted in increases in amiodarone concentrations from 0.9 mg/L to 1.3 mg/L. DEA concentra-tions were not affected. There was no evidence of toxicity. Monitoring for amiodarone toxicity and serial measurement of amiodarone serum concentration during concomitant protease inhibitor therapy should be considered.

Histamine H<sub>2</sub> antagonists: Cimetidine inhibits CYP3A4 and can increase serum amiodarone levels.

### Other substances: Grapefruit juice given to healthy volunteers increased amiodarone AUC by 50% and Crass by 84%, and decreased DEA to unquantifiable concentrations. Grapefruit juice inhibits CYP3A4-mediated metabolism of oral amiodarone in the intestinal mucosa, resulting in increased plasma levels of amiodarone; therefore, grapefruit juice should be a concentration of the intestinal mucosa, resulting in increased plasma levels of amiodarone; therefore, grapefruit juice should be a concentration of the not be taken during treatment with oral amiodarone. This information should be considered ered when changing from intravenous amiodarone to oral amiodarone (see "DOSAGE

AND ADMINISTRATION") AND ADMINISTRATION ).

Amilodarone may suppress certain CYP450 enzymes, including CYP1A2, CYP2C9, CYP2D6, and CYP3A4. This inhibition can result in unexpectedly high plasma levels of other drugs which are metabolized by those CYP450 enzymes. Reported examples of this interaction include the following:

### Immunosunnressives

Cyclosporine (CYP3A4 substrate) administered in combination with oral amiodarone has been reported to produce persistently elevated plasma concentrations of cyclosporine resulting in elevated creatinine, despite reduction in dose of cyclosporine. HMG-CoA reductase inhibitors:

Simvastatin (CYP3A4 substrate) in combination with amiodarone has been associated with reports of myopathy/rhabdomyolysis.

Cardiovasculars: Cardiac glycosides: In patients receiving digoxin therapy, administration of oral amiodarone regularly results in an increase in the serum digoxin concentration that may reach toxic levels with resultant clinical toxicity. Amiodarone taken concomitantly with digoxin increases the serum digoxin concentration by 70% after one day. On initiation of oral amiodarone, the need for digitalis therapy should be reviewed and the dose reduced by approximately 50% or discontinued. If digitalis treatment is continued, serum levels should be closely monitored and patients observed for clinical evidence of toxicity. These precautions probably should apply to digitoxin administration as well Antiarrhythmics:

Other antiarrhythmic drugs, such as quinidine, procainamide, disopyramide, and

phenytoin, have been used concurrently with oral amiodarone.
There have been case reports of increased steady-state levels of quinidine, procainamide, and phenytoin during concomitant therapy with amiodarone. Phenytoin decreases serum amiodarone levels. Amiodarone taken concomitantly with quindine increases quinidine serum concentration by 33% after two days. Amiodarone taken concomitantly with procalimatile for less than seven days increases plasma concentrations of procainamide and n-acetyl procainamide by 55% and 33%, respectively Quinidine and progainamide doses should be reduced by one-third when either is administered with amiodarone. Plasma levels of **flecainide** have been reported to increase in the presence of oral amiodarone; because of this, the dosage of flecainide should be adjusted when these drugs are administered concomitantly. In general, any added antiarrhythmic drug should be initiated at a lower than usual dose with careful monitoring.

Combination of amiodarone with other antiarrhythmic therapy should be reserved for patients with life-threatening ventricular arrhythmias who are incompletely responsive to a single agent or incompletely responsive to a mindarone. During transfer to amiodarone the dose levels of previously administered agents should be reduced by 30 to 50% several days after the addition of amiodarone, when arrhythmia suppression should be beginning. The continued need for the other antiarrhythmic agent should be reviewed after the effects of amindarone have been established, and discontinuation ordinarily should be attempted. If the treatment is continued, these patients should be particularly carefully monitored for adverse effects, especially conduction disturbances and exacerbation of tachyarrhythmias, as amiodarone is continued. In amiodarone treated patients who require additional antiarrhythmic therapy, the initial dose of such agents should be approximately half of the usual recommend

### Antihynartanciyac

Antinypertensives:

Amindarone should be used with caution in patients receiving 8-receptor blocking agents (e.g., propranolol, a CYP3A4 inhibitor) or calcium channel antagonists (e.g., verapamil, a CYP3A4 substrate, and diltiazem, a CYP3A4 inhibitor) because of the possible potentiation of bradycardia, sinus arrest, and AV block; if necessary, amiodarone can continue to be used after insertion of a pacemaker in patients with severe bradycar

### Anticoagulants:

Antihintics:

Potentiation of warfarin-type (CVP2CQ and CVP3AA substrate) anticognitant reconnect is almost always seen in patients receiving amiodarone and can result in serious or fatal bleeding. Since the concomitant administration of warfarin with amiodarone increases the prothrombin time by 100% after 3 to 4 days, the dose of the anticoaculant should be reduced by one-third to one-half, and prothrombin times should be monitored

Some drugs/substances are known to accelerate the metabolism of amindarone by Stimulating the synthesis of CYP3A4 (enzyme induction). This may lead to low amiodarone serum levels and potential decrease in efficacy. Reported examples of this interaction include the following:

Rifampin is a potent inducer of CYP3A4. Administration of rifampin concomitantly with oral amiodarone has been shown to result in decreases in serum concentrations of amiodarone and desethylamiodarone.

Other substances, including herbal preparations: St. John's Wort (Hypericum perforatum) induces CYP3A4. Since amiodarone is a substrate for CYP3A4, there is the potential that the use of St. John's Wort in patients receiving amiodarone could result in reduced amiodarone levels.

Other reported interactions with amiodarone:
Fentanyl (CYP3A4 substrate) in combination with amiodarone may cause hypotension. bradycardia, and decreased cardiac output.

Sinus bradycardia has been reported with oral amiodarone in combination with lidocaine (CYP3A4 substrate) given for local anesthesia. Seizure, associated with increased lidocaine concentrations, has been reported with concomitant administration

Dextromethorphan is a substrate for both CYP2D6 and CYP3A4. Amiodarone inhibits CYP2D6.

Cholestyramine increases enterohepatic elimination of amiodarone and may reduce its serum levels and t<sub>1/2</sub>.

Disonvramide increases OT prolongation which could cause arrhythmia

Fluoroquinolones, macrolide antibiotics, and azoles are known to cause QTc prolongation. There have been reports of QTc prolongation, with or without TdP, in patients taking amiodarone when fluoroquinolones, macrolide antibiotics, or azoles were administered concomitantly. (See "WARNINGS, Worsened Arrhythmia".)

Hemodynamic and electrophysiologic interactions have also been observed after con-comitant administration with propranolol, diltiazem, and verapamil.

Volatile Anesthetic Agents (See "PRECAUTIONS, Surgery, Volatile Anesthetic Agents,") In addition to the interactions noted above, chronic (>2 weeks) oral Cordarone administration impairs metabolism of phenytoin, dextromethorphan, and methotrexate

### Electrolyte Disturbances

Electroyle disturbances Since antiarry/thmic drugs may be ineffective or may be arrhythmogenic in patients with hypokalemia, any potassium or magnesium deficiency should be corrected before instituting and during Cordarone therapy. Use caution when coadministering Cordarone with drugs which may induce thypokalemia and/or hypomagnesemia.

Carcinogenesis, Mutagenesis, Impairment of Fertility
Amiodarone HCl was associated with a statistically significant, dose-related increase in the incidence of thyroid tumors (follicular adenoma and/or carcinoma) in rats. The incidence of thyroid tumors was greater than control even at the lowest dose level tested, i.e., 5 mg/kg/day (approximately 0.08 times the maximum recommended human

Mutagenicity studies (Ames, micronucleus, and lysogenic tests) with Cordarone were

negative.

In a study in which amiodarone HCl was administered to male and female rats, beginning 9 weeks prior to mating, reduced fertility was observed at a dose level of 90 mg/kg/day (approximately 1.4 times the maximum recommended human maintenance dose\*). \*600 mg in a 50 kg patient (dose compared on a body surface area basis)

# Pregnancy: Pregnancy Category D See "WARNINGS, Neonatal Hypo- or Hyperthyroidism."

Labor and Delivery
It is not known whether the use of Cordarone during labor or delivery has any immediate or delayed adverse effects. Preclinical studies in rodents have not shown any effect of Cordarone on the duration of gestation or on parturition.

### Nursing Mothers

Nursing womers

Cordarone and one of its major metabolites, desethylamiodarone (DEA), are excreted in human milk, suggesting that breast-feeding could expose the nursing infant to a significant dose of the drug. Nursing offspring of lactating rats administered Cordarone have been shown to be less viable and have reduced body-weight gains. Therefore, when Cordarone therapy is indicated, the mother should be advised to discontinue nursing.

The safety and effectiveness of Cordarone® (amiodarone HCI) Tablets in pediatric patients have not been established.

### Geriatric Use

Clinical studies of Cordarone Tablets did not include sufficient numbers of subjects comical soudies of voidance habets and not include softwarent indicates or soughests aged 65 and over to determine whether they respond differently from younger subjects. Other reported clinical experience has not identified differences in responses between the elderly and younger patients. In general, does 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

Adverse reactions have been very common in virtually all series of patients treated with Cordarone for ventricular arrhythmias with relatively large doses of drug (400 mg/day and above), occurring in about three-fourths of all patients and causing discontinuation in 7 to 18%. The most serious reactions are pulmonary toxicity, exacerbation of arrhythmia, and rare serious liver injury (see "WARNINGS"), but other adverse effects arriyalinia, and rate serious well injuly (see Wanhamos ), but ourie adverse eneus occisitute important problems. They are often reversible with dose reduction or cessation of Cordarone treatment. Most of the adverse effects appear to become more frequent with continued treatment beyond six months, although rates appear to remain relatively constant beyond one year. The time and dose relationships of adverse effects are under continued study.

Neurologic problems are extremely common, occurring in 20 to 40% of patients and including malaise and fatigue, tremor and involuntary movements, poor coordination and gait, and peripheral neuropathy; they are rarely a reason to stop therapy and may respond to dose reductions or discontinuation (see "PRECAUTIONS").

Gastrointestinal complaints, most commonly nausea, vomiting, constipation, and anorexia, occur in about 25% of patients but rarely require discontinuation of drug. These commonly occur during high-dose administration (i.e., loading dose) and usually respond to dose reduction or divided doses.

Ophthalmic abnormalities including optic neuropathy and/or optic neuritis, in some cases progressing to permanent blindness, papilledema, corneal degeneration, phot sensitivity, eye discomfort, scotoma, lens opacities, and macular degeneration have been reported. (See "WARNINGS.")

Asymptomatic corneal microdeposits are present in virtually all adult patients who have been on drug for more than 6 months. Some patients develop eye symptoms of halos, photophobia, and dry eyes. Vision is rarely affected and drug discontinuation is rarely

Dermatological adverse reactions occur in about 15% of nationts with photosensitivity being most common (about 10%). Sunscreen and protection from sun exposure may be helpful, and drug discontinuation is not usually necessary. Prolonged exposure to Cordarone occasionally results in a blue-gray pigmentation. This is slowly and occasionally incompletely reversible on discontinuation of drug but is of cosmetic impor-

Cardiovascular adverse reactions, other than exacerbation of the arrhythmias, include the uncommon occurrence of congestive heart failure (3%) and bradycardia. Bradycardia usually responds to dosage reduction but may require a pacemaker for control. CHF rarely requires drug discontinuation. Cardiac conduction abnormalities occur infrequently and are reversible on discontinuation of drug.

The following side-effect rates are based on a retrospective study of 241 patients treated for 2 to 1.515 days (mean 441.3 days).

The following side effects were each reported in 10 to 33% of patients: Gastrointestinal: Nausea and vomiting.

## The following side effects were each reported in 4 to 9% of patients:

Dermatologic: Solar dermatitis/photosensitivity. Neurologic: Malaise and fatigue, tremor/abnormal involuntary movements, lack of coor-

dination, abnormal galt/ataxia, dizziness, paresthesias. Gastrointestinal: Constipation, anorexia. Ophthalmologic: Visual disturbances.

Hepatic: Abnormal liver-function tests Respiratory: Pulmonary inflammation or fibrosis

## The following side effects were each reported in 1 to 3% of patients:

hyroid: Hypothyroidism, hyperthyroidism. Neurologic: Decreased libido, insomnia, headache, sleep disturbances Cardiovascular: Congestive heart failure, cardiac arrhythmias, SA node dysfunction.

Gastrointestinal: Abdominal pain. Henatic: Nonspecific henatic disorders Other: Flushing, abnormal taste and smell, edema, abnormal salivation, coagulation

# The following side effects were each reported in less than 1% of patients Blue skin discoloration, rash, spontaneous ecchymosis, alopecia, hypotension. and car

matitis, blue skin discoloration, hyperthyroidism, and hypothyroidism

diac conduction abnormalities In surveys of almost 5,000 patients treated in open U.S. studies and in published reports of treatment with Cordarone, the adverse reactions most frequently requiring discontinuation of Cordarone included pulmonary infiltrates or fibrosis, paroxysmal ventricular tachycardia, congestive heart failure, and elevation of liver enzymes. Other symptoms causing discontinuations less often included visual disturbances, solar der

## Postmarketing Reports

rosiniarketing repuirs in postmarketing surveillance, sinus arrest, hepatitis, cholestatic hepatitis, cirrhosis, epididymtils, impotence, vasculitis, pseudotumor cerebri, syndrome of inappropriate antidiuretic hormone secretion (SIADH), thrombocytopenia, angioedema, bronchiolitis andudureux orimine secureum (slovini, internatory)periana, anguereumia, oriminomo obliterans organizing pneumonia (possibly fatal), bronchospasm, possibly fatal respiratory disorders (including distress, failure, arrest, and ARDS), fever, dyspnea, cough, hempolysis, wheezing, hypoxia, pulmonary inflitrates, pelleuritis, pancreatitis, toxic epidermal necrolysis (sometimes fatal), myopathy, muscle weakness,

rhabdomyolysis, hemolytic anemia, aplastic anemia, pancytopenia, neutropenia, erythema multiforme, Stevens-Johnson syndrome, exfoliative dermatitis, pruritus, hallucination, confusional state, disorientation, and delirium also have been reported in natients receiving Cordarone

There have been cases, some fatal, of Cordarone overdose.

In addition to general supportive measures, the patient's cardiac rhythm and blood pressure should be monitored, and if bradycardia ensues, a 8-adrenergic agonist or a pacemaker may be used. Hypotension with inadequate tissue perfusion should be treated n positive inotropic and/or vasopressor agents. Neither Cordarone nor its metaboliti

The acute oral LD50 of amiodarone HCl in mice and rats is greater than 3,000 mg/kg.

DOSAGE AND ADMINISTRATION
BECAUSE OF THE UNIQUE PHARMACOKINETIC PROPERTIES, DIFFICULT DOSING
SCHEDULE, AND SEVERITY OF THE SIDE EFFECTS IF PATIENTS ARE IMPROPERLY
MONITORED, CORDARONE SHOULD BE ADMINISTERED ONLY BY PHYSICIANIS WHO
ARE EXPERIENCED IN THE TREATMENT OF LIFE THERETENING ARRHYTHMIAS WHO ARE THOROUGHLY FAMILIAR WITH THE RISKS AND BENEFITS OF CORDARONE THERAPY, AND WHO HAVE ACCESS TO LABORATORY FACILITIES CAPABLE OF ADE-QUATELY MONITORING THE FFFECTIVENESS AND SIDE EFFECTS OF TREATMENT. In order to insure that an antiarrhythmic effect will be observed without waiting several months, loading doses are required. A uniform, optimal dosage schedule for administration of Cordarone has not been determined. Because of the food effect on absorption, Cordarone should be administered consistently with regard to meals (see "CLINICAL PHARMACOLOGY"). Individual patient titration is suggested according to

the following guidelines: For life-threatening ventricular arrhythmias, such as ventricular fibrillation or hemody-namically unstable ventricular tachycardia: Close monitoring of the patients is indicated during the loading phase, particularly until risk of recurrent ventricular tachycardia or fibrillation has abated. Because of the serious nature of the arrhythmia and the lack of infinition has abared. Declared in the along should be offer a minylimite allowed that of the predictable time course of effect, of the along should be performed in a hospital setting. Loading doses of 800 it, 600 mg/day are required for 1 to 3 weeks (occasionally longer) until mild the apeutic response occurs. (Administration of Cordarone in divided doses with meaks is suggested for total daily doses of 1,000 mg or higher, or when pastrointestinal intolerance occurs.) If side effects become excessive, the dose should be reduced. Elimination of recurrence of ventricular fibrillation and tachycardia usually occurs within 1 to 3 weeks, along with reduction in complex and total ventricula

Since grapefruit juice is known to inhibit CYP3A4-mediated metabolism of oral amiodarone in the intestinal mucosa, resulting in increased plasma levels of amiodarone. grapefruit juice should not be taken during treatment with oral amiodarone (see "PRECAUTIONS, Drug Interactions").

Upon starting Cordarone therapy, an attempt should be made to gradually discontinue prior antiarrhythmic drugs (see section on "Drug Interactions"). When adequate prior animal royunito: Usig Sees Section (in 'I'rug interactions'); vi witer discription arrivithmic animal royunito (in 'I'rug interactions'); vi witer discription arrivithmic animal royunito (in 'I'rug in 'I'rug in

single daily dose, or in patients with severe gastrointestinal intolerance, as a b.i.d. dose. In each natient, the chronic maintenance dose should be determined according to in each patient, the chronic maintenance does enotion be determined according in antiarrhythmic effect as assessed by symptoms, Holter recordings, and/or progr electrical stimulation and by patient tolerance. Plasma concentrations may be he evaluating nonresponsiveness or unexpectedly severe toxicity (see "CLINICAL PHARMACOLIGY").

# The lowest effective dose should be used to prevent the occurrence of side effects

The lowest enective loses should not less of prevent une occurrence or since enects, in all instances, the physician must be guided by the severity of the individual patient's arrhythmia and response to therapy. When dosage adjustments are necessary, the patient should be closely monitored for an extended period of time because of the long and variable half-life of Cordarone and the difficulty in predicting the time required to attain a new steady-state level of drug.

Dosage suggestions are s	Loading Dose (Daily)	Adjustment and Maintenance Dose (Daily)		
Ventricular Arrhythmias	1 to 3 weeks	~1 month	usual maintenance	
	800 to 1,600 mg	600 to 800 mg	400 mg	

Cordarone® (amiodarone HCI) Tablets are available in bottles of 60 tablets and in Redipak® cartons containing 100 tablets (10 blister strips of 10) as follows: 200 mg, NDC 0008-4188, round, convex-faced, pink tablets with a raised "C" and marked "200" on one side, with reverse side scored and marked "WYETH" and "4188."

Keep tightly closed. Store at room temperature, approximately 25°C (77°F).
Protect from light.
Dispense in a light-resistant, tight container.
Use carlon to protect contents from light.

# **Medication Guide**

# Cordarone® 'KOR-DU-RON Tablets (amindarone HCI)

### Rx only

Read the Medication Guide that comes with Cordarone Tablets before you start taking them and each time you get a refill. There may be new information. This Medication Guide does not take the place of talking with your doctor about your medical condition

# What is the most important information I should know about Cordarone Tablets? Cordarone Tablets can cause serious side effects that can lead to death including:

- lung damage liver damage
- beat problems
- Call your doctor or get medical help right away if you have any symptoms such as the onlowing.

  shortness of breath, wheezing, or any other trouble breathing; coughing, chest pain,
- or spitting up of blood nausea or vomiting; passing brown or dark-colored urine; feel more tired than usual;
- your skin and whites of your eyes get yellow; or have stomach pain heart pounding, skipping a beat, beating very fast or very slowly; feel light-headed or

# Because of these possible side effects, Cordarone Tablets should only be used in adults with life-threatening heartbeat problems called ventricular arrhythmias, for which other treatments did not work or were not tolerated.

Cordarone Tablets can cause other serious side effects. See "What are the nossible or reasonably likely side effects of Cordarone Tablets?" for more information If you get serious side effects during treatment with Cordarone Tablets you may need to

stop Cordarone Tablets, have your dose changed, or get medical treatment. Talk with your doctor before you stop taking Cordarone Tablets. You may still have side effects after stopping Cordarone Tablets because the mediTell all your healthcare providers that you take or took Cordarone Tablets. This information is very important for other medical treatments or surgeries you may have

### What are Cordarone Tablets?

what are contained abules?

Cordarone is a medicine used in adults to treat life-threatening heartbeat problems called ventricular arrhythmias, for which other treatment did not work or was not tolercalled verification and initialists, or without once treatment out in the work was not ober-bade. Cordarone Tablets have not been shown to help people with life-threatening heart-beat problems live longer. Treatment with Cordarone Tablets should be started in a hospital to monitor your condition. You should have regular cheek-ups, blood tests, chest x-rays, and eye exams before and during treatment with Cordarone Tablets to check for serious side effects.

Cordarone Tablets have not been studied in children.

### Who should not take Cordarone Tablets?

- who snoul not lake Corarone I adiets?

  Do not take Cordrone Tablets if you:

   have certain heart conditions (heart block, very slow heart rate, or slow heart rate with dizziness or lightheadedness)

   have an allergy to amiodarone, iodine, or any of the other ingredients in
- Cordarone Tablets. See the end of this Medication Guide for a complete list of ingre-dients in Cordarone Tablets.

# What should I tell my doctor before starting Cordarone Tablets?

- Tell your doctor about all of your medical conditions including if you:
   have lung or breathing problems
- have liver problems
- have or had thyroid problems
- have blood pressure problems
- nave blood pressure problems
  are pregnant or planning to become pregnant. Cordarone can harm your unborn
  baby. Cordarone can stay in your body for months after treatment is stopped.
  Therefore, talk with your doctor before you plan to get pregnant.
  are breastleeding. Cordarone passes into your milk and can harm your baby. You
- should not breast feed while taking Cordarone, Also, Cordarone can stay in your body for months after treatment is stopped.

Tell your doctor about all the medicines you take including prescription and nonpre-scription medicines, vitamins and herbal supplements. Cordarone Tablets and certain other medicines can interact with each other causing serious side effects. Sometimes the dose of Cordarone Tablets or other medicines must be changed when they are used

- together. Especially, tell your doctor if you are taking:

  antibiotic medicines used to treat infections
- depression medicines
- cimetidine (Tagamet®), a medicine for stomach ulcers or indigestion
- seizure medicines diabetes medicines
- cyclosporine an immunosuppressive medicine
- dextromethorphan, a cough medicine medicines for your heart, circulation, or blood pressure
- water pills (diuretics) high cholesterol or bile medicines
- narcotic pain medicines St. John's Wort

Know the medicines you take. Keep a list of them with you at all times and show it to your doctor and pharmacist each time you get a new medicine. Do not take any new medicines while you are taking Cordarone Tablets unless you have talked with your

### How should I take Cordarone Tablets?

- flow should I take Cordarone Tablets?

  Take Cordarone Tablets exactly as prescribed by your doctor.

  The dose of Cordarone Tablets you take has been specially chosen for you by your doctor and may change during treatment. Keep taking your medicine until your doctor tells you to stop. Do not stop taking it because you teel better. Your condition may get worse. Talk with your doctor if you have side effects.
- Your doctor will tell you to take your dose of Cordarone Tablets with or without
- rour doctor will early us case you undeed ordinatine faules within on willowing meals. Make sure you take Condarone Tablets the same way each time.

  Do not drink grapefruit juice during treatment with Cordarone Tablets. Grapefruit juice affects how Cordarone is absorbed in the stomach.

  Taking too many Cordarone Tablets can be dangerous. If you take too many
- Cordarone Tablets, call your doctor or go to the nearest hospital right away. You may
- need medical care right away.

  If you miss a dose, do not take a double dose to make up for the dose you missed. Continue with your next regularly scheduled dose
- What should I avoid while taking Cordarone Tablets?
- Do not drink grapefruit juice during treatment with Cordarone Tablets. Grapefruit juice affects how Cordarone is absorbed in the stomach. Avoid exposing your skin to the sun or sun lamps. Cordarone Tablets can cause a photosensitive reaction. Wear sun-block cream or protective clothing when out in the
- Avoid pregnancy during treatment with Cordarone Tablets. Cordarone can harm
- your unborn baby.

  Do not breastfeed while taking Cordarone Tablets. Cordarone passes into your milk and can harm your baby.

What are the nossible or reasonably likely side effects of Cordarone Tablets? what are the possible or reasonably likely side effects that lead to death including lung damage, liver damage, and worse heartbeat problems. See "What is the most important information I should know about Cordarone Tablets?"

# Some other serious side effects of Cordarone Tablets include:

- unite dimer serious since elects of curardine l'abilest include:

  vision problems that may lead to permanent blindness. You should have regular
  eye exams before and during treatment with Cordarone Tablets. Call your doctor if
  you have blurred vision, see halos, or your eyes become sensitive to light,
  nerve problems. Cordarone Tablets can cause a feeling of "pins and needles" or numbness in the hands, legs, or feet, muscle weakness, uncontrolled movements
- poor coordination, and trouble walking. thyroid problems. Cordarone Tablets can cause hypothyroidism or hyperthyroidism. Infruit profiles. Cortainer latties can cause injunyiouslin or hyperinfruitism. Your doctor may arrange regular blood tests to check your thyroid function during treatment with Cordarone. Call your doctor if you have weight loss or weight gain, restlessness, weakness, heat or oold intolerance, hair thinning, sweating, changes it your menses, or swelling of your neck (gotter).
- skin problems. Cordarone Tablets can cause your skin to be more sensitive to the sun or to turn a bluish-gray color. In most patients, skin color slowly returns to nor mal after stopping Cordarone Tablets. In some patients, skin color does not return to

Other side effects of Cordarone Tablets include nausea, vomiting, constipation, and loss

Call your doctor about any side effect that bothers you.

These are not all the side effects with Cordarone Tablets. For more information, ask your doctor or pharmacist

## How should I store Cordarone Tablets?

- Store Cordarone Tablets at room temperature. Protect from light. Keep Cordarone Tablets in a tightly closed container.

  Safely dispose of Cordarone Tablets that are out-of-date or no longer needed.

# Keen Cordarone Tablets and all medicines out of the reach of children.

General information about Cordarone Tablets General Information about Cordarone Tablets

Medicates are sometimes prescribed for purposes other than those listed in a
Medication Guide. Do not use Cordarone Tablets for a condition for which it was not
prescribed. Do not share Cordarone with other people, even if they have the same symptoms that you have. It may harm them

If you have any questions or concerns about Cordarone Tablets, ask your doctor or healthcare provider. This Medication Guide summarizes the most important info about Cordarone Tablets. If you would like more information, talk with your doct can ask your doctor or pharmacist for information about Cordarone Tablets that was



This Medication Guide may have been revised after this inis Medication Guide may nave been revised after mis copy was produced. For more information and the most current Medication Guide, please visit www.wyeth.com or call our medical communications department toll-free at 1-800-934-5556.



What are the ingredients in Cordarone Tablets?

Inactive Ingredients: colloidal silicon dioxide, lactose, magnesium stearate, povidone starch, and FD&C Red 40

This Medication Guide has been approved by the U.S. Food and Drug Administration Rx only

# Manufactured for Wveth®

Wyeth Pharmaceuticals Inc. Philadelphia, PA 19101

by Sanofi Winthrop Industrie 1, rue de la Vierge 33440 Ambares, France

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