PRODUCT MONOGRAPH

PrMULTAQ®

Dronedarone Tablets

400 mg dronedarone (as dronedarone hydrochloride)

Antiarrhythmic Agent

ATC code: C01BD07

sanofi-aventis Canada Inc.
1755 Steeles Avenue West
Toronto ON,
M2R 3T4

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PART I: HEALTH PROFESSIONAL INFORMATION

SUMMARY PRODUCT INFORMATION

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<tr>
<th>Route of Administration</th>
<th>Dosage Form / Strength</th>
<th>Nonmedicinal Ingredients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral</td>
<td>tablet 400 mg</td>
<td>carnauba wax, colloidal anhydrous silica, crospovidone, hypromellose, lactose monohydrate, macrogol 6000, magnesium stearate, maize starch, poloxamer 407, titanium dioxide.</td>
</tr>
</tbody>
</table>

INDICATIONS AND CLINICAL USE

MULTAQ is indicated for the treatment of patients with paroxysmal or persistent atrial fibrillation (AF) who are in sinus rhythm or who are intended to be cardioverted, to reduce the risk of cardiovascular hospitalization due to atrial fibrillation (see CLINICAL TRIALS). Multaq should only be prescribed after alternative treatment options have been considered.

Pediatrics (< 18 years of age)

No data are available.

CONTRAINDICATIONS

MULTAQ is contraindicated in patients with:

- **Permanent** atrial fibrillation of any duration in which sinus rhythm cannot be restored and attempts to restore it are no longer considered by the attending physician.
- History of, or current heart failure, regardless of NYHA functional class
- Left ventricular systolic dysfunction
- Second- or third-degree atrio-ventricular (AV) block, complete bundle branch block, distal block, sinus node dysfunction, atrial conduction defects or sick sinus syndrome, (except when used in conjunction with a functioning pacemaker)
- Unstable hemodynamic condition(s)
- Bradycardia < 50 bpm
- Patients with liver or lung toxicity related to the previous use of amiodarone
- Co-administration with strong CYP3A4 inhibitors, such as ketoconazole, itraconazole, voriconazole, cyclosporin, clarithromycin, and ritonavir (see DRUG INTERACTIONS)
- Co-administration with drugs inducing torsades de pointes such as phenothiazines, tricyclic antidepressants, and certain oral macrolides, class I and III antiarrhythmics
- QTc Bazett interval ≥ 500 msec
- Severe hepatic impairment
- Pregnancy (see WARNINGS AND PRECAUTIONS, Special populations)
- Lactation (see WARNINGS AND PRECAUTIONS, Special populations)
- History of hypersensitivity reactions to dronedarone or any of its excipients or component of the container. For a complete listing, see DOSAGE FORMS, COMPOSITION AND PACKAGING.

WARNINGS AND PRECAUTIONS

Anticoagulation Therapy

Patients should be appropriately anticoagulated. Where applicable, International Normalized Ratio (INR) should be closely monitored after initiating dronedarone in patients taking vitamin K antagonists as per their label.

Dronedarone should not be used concomitantly with either dabigatran or rivaroxaban since it may increase exposure of dabigatran through P-gp inhibition or rivaroxaban through P-gp and CYP3A4 inhibition (see DRUG INTERACTIONS).

Cardiovascular

Congestive Heart Failure (CHF) or Left Ventricular Systolic Dysfunction

Dronedarone is contraindicated for use in patients in unstable hemodynamic conditions, history of, or current heart failure or left ventricular systolic dysfunction. (see CONTRAINDICATIONS).

Patients with new or worsening heart failure during treatment

Patients should be advised to consult a physician if they develop signs or symptoms of heart failure, such as weight gain, dependent edema, or increasing shortness of breath. If heart failure develops treatment with MULTAQ should be discontinued (see CONTRAINDICATIONS).

Patients should be followed for the development of left ventricular systolic dysfunction during treatment. If left ventricular systolic dysfunction develops, treatment with MULTAQ should be discontinued.
Patients Developing Permanent AF During Treatment

A clinical study in patients with permanent AF (AF duration for at least 6 months) and cardiovascular risk factors was stopped early due to an excess of cardiovascular death, stroke and unplanned cardiovascular hospitalization. It is recommended to perform an ECG at least every 6 months while patients are receiving MULTAQ. If patients treated with MULTAQ develop permanent AF, treatment with MULTAQ should be discontinued.

Patients with Coronary Artery Disease

Caution is needed in patients with coronary artery disease.

QT prolongation

The pharmacological action of dronedarone may induce a moderate (about 10 msec) QTc Bazett prolongation, related to prolonged repolarization. These changes are linked to the therapeutic effect of dronedarone and do not reflect toxicity. Follow up, including ECG, is recommended during treatment. If QTc Bazett interval is ≥ 500 msec, dronedarone should be stopped (see CONTRAINDICATIONS).

Based on clinical experience, dronedarone has a low pro-arrhythmic effect. However, proarrhythmic effects may occur in particular situations such as concomitant use with drugs favoring arrhythmia and/or electrolytic disorders (see WARNINGS AND PRECAUTIONS and DRUG INTERACTIONS).

Endocrine and Metabolism

Electrolyte imbalance
Since antiarrhythmic drugs may be ineffective or may be arrhythmogenic in patients with hypokalemia, any potassium or magnesium deficiency should be corrected before initiation and during dronedarone therapy.

Hepatic

Liver Injury
Hepatocellular liver injury, including acute liver failure requiring transplant, has been reported in patients treated with MULTAQ in the post-marketing setting. Healthcare professionals should advise patients treated with MULTAQ to immediately report symptoms suggesting hepatic injury (such as anorexia, nausea, vomiting, fever, malaise, fatigue, right upper quadrant pain, jaundice, dark urine, or itching). It is not known whether routine periodic monitoring of serum enzymes will detect the development of severe liver injury. However, healthcare professionals should consider obtaining periodic hepatic serum enzymes, especially during the first 6 months of treatment. If hepatic injury is suspected, MULTAQ should be discontinued immediately followed by appropriate blood tests. If liver injury is found, appropriate treatment should be instituted, and investigations should be performed to establish the cause. The safety of restarting MULTAQ in patients who have sustained liver injury from any cause is unknown; accordingly, its use in such patients is not recommended.
Renal

Plasma Creatinine Increase
An increase in plasma creatinine has been observed with dronedarone 400 mg twice daily in healthy subjects and in patients. This increase occurs early after treatment initiation and reaches a plateau after 7 days. Mean increase in atrial fibrillation patients is about 10 µmol/L. Values return to baseline within one week after treatment discontinuation. In a specific study in healthy subjects, this increase was shown to be related to inhibition of creatinine secretion at the tubular level, with no effect on glomerular filtration or on renal blood flow. The same mechanism has also been described with other drugs such as cimetidine, trimethoprim or amiodarone. Increased plasma creatinine could be misinterpreted and lead to inappropriate discontinuation of ACE inhibitors or AIIRAs in patients requiring this treatment (see WARNINGS AND PRECAUTIONS, Monitoring and Laboratory Tests, Management of plasma creatinine increase).

Larger increases in creatinine after dronedarone initiation have been reported in the post-marketing setting. Some cases also reported increases in blood urea nitrogen, possibly due to hypoperfusion secondary to developing CHF (pre-renal azotaemia). In such cases, dronedarone should be stopped. It is recommended to monitor renal function periodically and to consider further investigations as needed (see Post-Market Adverse Drug Reactions, Renal).

Respiratory Disorders
Cases of interstitial lung disease including pneumonitis and pulmonary fibrosis have been reported in post-marketing experience (see ADVERSE REACTIONS, Post-Market Adverse Drug Reactions). Onset of dyspnoea or non-productive cough may be related to pulmonary toxicity and patients should be carefully evaluated clinically. If pulmonary toxicity is confirmed treatment should be discontinued.

Special Populations

Pregnant Women (see CONTRAINDICATIONS)
MULTAQ has been shown to be teratogenic in the rat and is contraindicated in women who are or may become pregnant. Women of childbearing potential should use effective methods of contraception during treatment with MULTAQ. If this drug is used during pregnancy or if the patient becomes pregnant while taking this drug, the patient should be apprised of the potential hazard to the fetus.

MULTAQ can cause fetal harm when administered to pregnant women. In rats, dronedarone caused marked effects on embryo-fetal development at 100 mg/kg/day such as increased post-implantation losses, reduced fetal and placental weights and various external, visceral and skeletal malformations in most fetuses. At lower dosages, up to 50 mg/kg/day (corresponding to 4.5 times the recommended human therapeutic dose), dronedarone had no effects on the litters (with the exception of a transient minor effect on the bodyweight gain of the pups from D1 to D4 post-partum). Dronedarone had no adverse effects on the mothers and their litters up to 30 mg/kg/day. In rabbits, the high dose level (200 mg/kg/day) did not induce any effects to fetuses.
Labor and Delivery
It is not known whether the use of dronedarone during labor or delivery has any immediate or delayed adverse effects. In pre- and post-natal development studies in the rat, at doses up to 50 mg/kg/day, dronedarone induced no effect on the duration of gestation or on parturition (see TOXICOLOGY, Reproduction).

Nursing Women
Dronedarone and its metabolites are excreted in rat milk. Nursing in lactating rats administered dronedarone was associated with minor reduced body-weight gain in the offspring. It is not known whether this drug is excreted in human milk. Therefore, when dronedarone therapy is indicated, the mother should be advised to discontinue nursing (see CONTRAINDICATIONS).

Pediatrics (< 18 years of age)
Safety and efficacy in children below the age of 18 years has not been established. Therefore, use in these patients is not recommended.

Geriatrics (> 65 years of age)
A large number of elderly patients with AF or atrial flutter (AFL) have been included in the MULTAQ clinical program (more than 4500 patients aged 65 years or above, of which more than 2000 patients were 75 years or above). Efficacy and safety were comparable in both elderly and younger patients. Caution is needed in elderly patients≥ 75 years with multiple co-morbidities.

Renal impairment
Patients with renal impairment were included in clinical studies. Consistent with the minimal renal excretion of dronedarone, no pharmacokinetic modification was observed in patients with renal impairment in particular in patients with severe renal impairment though patients with severe renal impairment were present only in small numbers in these studies. From a clinical standpoint, limited experience is available in patients with severe renal impairment (see Post-Market Adverse Drug Reactions, Renal).

Monitoring and Laboratory Tests
It is recommended to closely monitor patients taking MULTAQ including ECG at least every 6 months. If patients treated with MULTAQ develop permanent AF, treatment with MULTAQ should be discontinued.

Management of plasma creatinine increase
It is recommended that baseline values of plasma creatinine be established 7 days after initiation of treatment with dronedarone.

If the result obtained for the plasma creatinine is above the upper limit of normal as provided by the laboratory, this value should be used as the new reference baseline taking into account that this may be expected with dronedarone, since the drug can affect baseline values. An increase in creatinemia should not necessarily lead to the discontinuation of treatment with dronedarone or
discontinuation of treatment with ACE-inhibitors or angiotensin receptor blockers (ARBs) (see WARNINGS AND PRECAUTIONS, Renal).

Liver function testing

Healthcare professionals should advise patients treated with MULTAQ to immediately report symptoms suggesting hepatic injury (such as anorexia, nausea, vomiting, fever, malaise, fatigue, right upper quadrant pain, jaundice, dark urine, or itching). It is not known whether routine periodic monitoring of serum enzymes will detect the development of severe liver injury. However, healthcare professionals should consider obtaining periodic hepatic serum enzymes, especially during the first 6 months of treatment. If hepatic injury is suspected, MULTAQ should be discontinued immediately followed by appropriate blood tests. If liver injury is found, appropriate treatment should be instituted, and investigations should be performed to establish the cause. The safety of restarting MULTAQ in patients who have sustained liver injury from any cause is unknown; accordingly, its use in such patients is not recommended.

Further laboratory testing is at the discretion of the attending physician.

ADVERSE REACTIONS

Adverse Drug Reaction Overview

The safety profile of dronedarone 400mg twice daily in patients with AF or AFL is based on 5 placebo controlled studies, ATHENA, EURIDIS, ADONIS, ERATO and DAFNE. In these studies, a total of 6285 patients were randomized and treated. Of these, 3282 patients were treated with dronedarone 400 mg twice daily, and 2875 received placebo.

The mean exposure across studies was 12 months. In ATHENA, the maximum follow-up was 30 months.

Assessment of intrinsic factors such as race, gender or age on the incidence of any treatment emergent adverse events did not suggest any excess of adverse events in a particular sub-group.

In pooled clinical trials, premature discontinuation due to adverse reactions occurred in 11.8% of the dronedarone-treated patients and in 7.7% in the placebo-treated group. The most common reasons for discontinuation of therapy with MULTAQ were gastrointestinal disorders (3.2 % of patients versus 1.8% in the placebo group).

The most frequent adverse reactions observed with dronedarone 400 mg twice daily in the 5 studies were diarrhea, nausea and vomiting, fatigue and asthenia

Clinical Trial Adverse Drug Reactions

Because clinical trials are conducted under very specific conditions the adverse reaction rates observed in the clinical trials may not reflect the rates observed in practice and should not be compared to the rates in the clinical trials of another drug. Adverse drug
reaction information from clinical trials is useful for identifying drug-related adverse events and for approximating rates.

Table 1 displays adverse reactions ≥ 1% associated with dronedarone 400 mg twice daily in AF or AFL patients, presented by system organ class and by decreasing order of frequency. Adverse reactions included in the system organ class “Investigation” are presented separately in Table 2.

| Table 1 - Adverse reactions occurring ≥ than 1% of patients taking dronedarone in placebo-controlled clinical trials |
|---------------------------------------------------------------|---------------|---------------|
|                                                              | Placebo       | Dronedarone 400 mg twice daily |
|                                                              | (N=2875)      | (N=3282)      |
| Gastrointestinal disorders                                   |               |               |
| Diarrhoea                                                     | 5.8%          | 9.0%          |
| Nausea                                                        | 3.1%          | 4.9%          |
| Abdominal pains                                               | 2.8%          | 3.5%          |
| Vomiting                                                      | 1.1%          | 2.0%          |
| Dyspepsia                                                     | 1.0%          | 1.5%          |
| General disorders and administration site conditions         |               |               |
| Fatigue                                                       | 3.6%          | 4.3%          |
| Asthenia                                                      | 1.7%          | 2.3%          |
| Cardiac disorders                                             |               |               |
| Bradycardia                                                   | 1.3%          | 3.3%          |
| Skin and subcutaneous tissue disorders                       |               |               |
| Rashes (including generalised, macular, maculopapular)        | 1.6%          | 2.7%          |
| Pruritus                                                       | 0.9%          | 1.3%          |

Less Common Clinical Trial Adverse Drug Reactions (<1%)

Nervous system disorders: Ageusia and dysgeusia.

Skin and subcutaneous tissue disorders: Dermatitis, dermatitis allergic, eczema, erythemas (including erythema and rash erythematous and photosensitivity reaction.

In addition, the following laboratory data/ECG parameters were reported with dronedarone 400 mg twice daily.

| Table 2 - : Laboratory data/ECG parameters |
|--------------------------------------------|-----------------|-----------------|
|                                             | Placebo         | Dronedarone 400 mg twice daily |
|                                             | (N=2875)        | (N=3282)        |
| Blood creatinine increased ≥ 10% five days after treatment initiation | 20.6%          | 50.9%          |
| QTc Bazett prolonged (> 450 msec in male > 470 msec in female) | 18.7%          | 27.6%          |
Post-Market Adverse Drug Reactions

The following adverse reactions have been identified during post-approval use of MULTAQ. Because these reactions are reported voluntarily from a population of an unknown size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

Cardiac disorders:
Congestive heart failure
A few cases of atrial flutter with 1:1 atrioventricular conduction have been reported.

Hepatic:
Serum hepatic enzymes and serum bilirubin increase: Hepatocellular liver injury, including acute liver failure requiring transplant, has been reported in a few patients (see WARNINGS AND PRECAUTIONS).

Immune system disorders:
Anaphylactic reactions including angioedema

Renal:
Marked increase in serum creatinine, pre-renal azotemia and acute renal failure, often in the setting of heart failure (see WARNINGS AND PRECAUTIONS, Cardiovascular) or hypovolemia, have been reported in patients taking MULTAQ. In most cases, these effects appear to be reversible upon drug discontinuation and with appropriate medical treatment. However, rare cases of acute renal failure resulting in death have been reported postmarketing. Although all such cases were complex and confounded by multiple illnesses and concomitant medications, a contributive role of dronedarone in the development of renal failure could not be excluded. Renal function should be monitored periodically (see WARNINGS AND PRECAUTIONS, Renal).

Respiratory disorders:
Interstitial lung disease including pneumonitis and pulmonary fibrosis (a number of patients had been previously exposed to amiodarone).

Vascular disorders:
Vasculitis, including leukocytoclastic vasculitis (Schönlein-Henoch purpura)

DRUG INTERACTIONS

Overview

Dronedarone is primarily metabolized by CYP 3A4 (see ACTION AND CLINICAL PHARMACOLOGY, Pharmacokinetics) and is a moderate inhibitor of CYP 3A4 and a mild inhibitor of CYP 2D6. Therefore, inhibitors and inducers of CYP 3A4 have the potential to interact with dronedarone, and dronedarone has the potential to interact with medicinal products
that are substrates of CYP 3A4 and CYP 2D6. It also has the potential to inhibit P-glycoprotein (P-gP) transport. Dronedarone and/or its metabolites also have the potential to inhibit Organic Anion Transporter (OAT), Organic Anion Transporting Polypeptide (OATP) and Organic Cation Transporter (OCT) transport in vitro. Dronedarone has no significant potential to inhibit CYP1A2, CYP2C9, CYP2C19, CYP2C8 and CYP 2B6.

A potential pharmacodynamic interaction can also be expected with beta-blockers, calcium antagonists and digitalis.

Co-administration of drugs prolonging the QT interval (such as phenothiazines, bepridil, tricyclic antidepressants, certain oral macrolides, Class I and III antiarrhythmics, which may induce torsades de pointes) is contraindicated because of the potential risk of proarrhythmia (see CONTRAINDICATIONS).

In clinical trials, patients treated with dronedarone received a variety of concomitant medications including beta-blockers, digitalis, calcium antagonists (including those with heart rate-lowering effects), statins and oral anticoagulants.

**Drug-Drug Interactions**

<table>
<thead>
<tr>
<th>Agent</th>
<th>Ref</th>
<th>Effect</th>
<th>Clinical comment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antidepressants</strong></td>
<td>T</td>
<td>No effect</td>
<td>Dronedarone is a weak inhibitor of CYP2D6 in humans, so it is predicted to have limited interaction on antidepressants that are metabolized by CYP2D6.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Proarrhythmic effect</td>
<td>Co-administration of dronedarone with tricyclic antidepressants and other medicinal products inducing torsades de pointes is contraindicated (see CONTRAINDICATIONS).</td>
</tr>
<tr>
<td><strong>Beta-blockers</strong></td>
<td>CT</td>
<td>↑ metoprolol and propranolol exposure</td>
<td>Beta-blockers that are metabolized by CYP 2D6 can have their exposure increased by dronedarone. Moreover, beta-blockers have the potential to interact with dronedarone from a pharmacodynamic point of view. Dronedarone 800 mg daily increased metoprolol exposure by 1.6-fold and propranolol exposure by 1.3-fold (i.e. much below the 6-fold differences observed between poor and extensive CYP 2D6 metabolizers). In clinical trials, bradycardia was more frequently observed when dronedarone was given in combination with beta-blockers. Due to the pharmacokinetic interaction and possible pharmacodynamic interaction, beta-blockers should be used with caution if taken concomitantly with dronedarone.</td>
</tr>
<tr>
<td>(e.g. metoprolol and propranolol)</td>
<td></td>
<td>Bradycardia</td>
<td></td>
</tr>
<tr>
<td>Agent</td>
<td>Ref</td>
<td>Effect</td>
<td>Clinical comment</td>
</tr>
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</tr>
<tr>
<td>Calcium antagonists (e.g. verapamil,</td>
<td>CT</td>
<td>↑ dronedarone exposure</td>
<td>Calcium antagonists are substrates and/or moderate inhibitors of CYP3A4. Moreover, calcium antagonists with heart-rate lowering properties have the potential to interact with dronedarone from a pharmacodynamic point of view.</td>
</tr>
<tr>
<td>diltiazem, nifedipine)</td>
<td></td>
<td></td>
<td>Repeated doses of diltiazem (240 mg twice daily), verapamil (240 mg once daily) and nifedipine (20 mg twice daily) resulted in an increase in dronedarone exposure of 1.7-, 1.4-, and 1.2-fold, respectively.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>↑ verapamil and nisoldipine</td>
<td>Calcium antagonists also have their exposure increased by dronedarone (400 mg twice daily) (verapamil by 1.4-fold, and nisoldipine by 1.5-fold). In clinical trials, there was no evidence of safety concerns when dronedarone was co-administered with calcium antagonists with heart rate-lowering effects.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>exposure</td>
<td>Overall, due to the pharmacokinetic interaction and possible pharmacodynamic interaction, calcium antagonists with depressant effects on sinus and atrio-ventricular node such as verapamil and diltiazem should be used with caution when associated with dronedarone.</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td>CT</td>
<td>No effect</td>
<td>No interaction was observed between dronedarone and clopidogrel.</td>
</tr>
<tr>
<td>Dabigatran</td>
<td>CT</td>
<td>↑ dabigatran exposure</td>
<td>Dronedarone should not be used concomitantly with dabigatran since it may increase exposure through P-gp inhibition, and thereby the risk of bleeding.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>When dabigatran etexilate 150 mg once daily was co-administered with dronedarone 400 mg twice daily, the dabigatran exposures (C&lt;sub&gt;max&lt;/sub&gt; and AUC&lt;sub&gt;0-24&lt;/sub&gt;) were increased by 1.7- and 2-fold, respectively.</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>In a retrospective, claims database cohort study in the United States (US), no statistically significant increased risk of bleeding diagnoses leading to hospitalization or emergency department visit was observed with the concomitant use of dabigatran and dronedarone compared to those taking dabigatran alone in patients with non-valvular atrial fibrillation (NVAF). Among those concomitantly taking dabigatran and dronedarone, there was a statistically significant increased risk of ICD-diagnoses of gastrointestinal bleeding.</td>
</tr>
<tr>
<td>Agent</td>
<td>Ref</td>
<td>Effect</td>
<td>Clinical comment</td>
</tr>
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<td>---------------------------</td>
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<td>----------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Digoxin</td>
<td>CT</td>
<td>↑ digoxin exposure</td>
<td>Dronedarone (400 mg twice daily) increased digoxin exposure by 2.5-fold by inhibiting P-gP transporter. Moreover, digitalis has the potential to interact pharmacodynamically with dronedarone. In clinical trials, increased levels of digitalis were observed when dronedarone was co-administered with digitalis. The concomitant use of digitalis and dronedarone is generally not recommended. Patients should be treated with digitalis and dronedarone only if there is a specific therapeutic need and no alternative treatment available. These patients should be closely monitored for serum digoxin levels, especially during the first week of co-administration. Clinical and ECG monitoring are also recommended, and the digoxin dose should be adjusted as appropriate.</td>
</tr>
<tr>
<td>Drugs prolonging the QT interval (e.g. phenothiazines, bepridil, tricyclic antidepressants, certain oral macrolides, Class I and III antiarrhythmics)</td>
<td>T</td>
<td>Proarrhythmic effect</td>
<td>Co-administration of drugs prolonging the QT interval (inducing torsades de pointes) is contraindicated because of the potential risk of proarrhythmia. (see CONTRAINDICATIONS).</td>
</tr>
<tr>
<td>Erythromycin (moderate CYP 3A4 inhibitor)</td>
<td>CT</td>
<td>↑ dronedarone exposure</td>
<td>Repeated doses of erythromycin (500 mg three times a day for 10 days) resulted in an increase in steady-state dronedarone exposure of 3.8-fold.</td>
</tr>
<tr>
<td>Agent</td>
<td>Ref</td>
<td>Effect</td>
<td>Clinical comment</td>
</tr>
<tr>
<td>-------</td>
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<td>--------</td>
<td>------------------</td>
</tr>
<tr>
<td>Factor Xa inhibitors (e.g. rivaroxaban, edoxaban, apixaban)</td>
<td>CT</td>
<td>↑ factor Xa inhibitor exposure</td>
<td>As dronedarone may increase exposure of factor Xa inhibitors through P-gp and/or CYP3A4 inhibition, an assessment of the risk for thromboembolic events and bleeding should be made when concomitant use is recommended. Consider dose reduction of the respective factor Xa inhibitors according to label recommendation. Dronedarone should not be used with rivaroxaban since it may increase exposure and thereby the risk of bleeding. In a retrospective cohort study using the Truven Health MarketScan database in the US, a significantly increased risk of ICD-diagnoses of bleeding leading to hospitalization or emergency department visit was observed, driven by gastrointestinal bleeding, in NVAF patients with the concomitant use of rivaroxaban and dronedarone compared to those taking rivaroxaban alone. Administration of dronedarone 400 mg twice daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 5 increased edoxaban AUC and C&lt;sub&gt;max&lt;/sub&gt; by 85% and 46%, respectively. Dose reduction of edoxaban is required for concomitant use with dronedarone. In a retrospective cohort study using the Truven Health MarketScan database in the US, no increased risk of bleeding diagnoses leading to hospitalization or emergency department visit was observed in NVAF patients with the concomitant use of apixaban with dronedarone compared to those taking apixaban alone.</td>
</tr>
<tr>
<td>Losartan (CYP 2C9 substrate)</td>
<td>CT</td>
<td>No effect</td>
<td>Although not considered to be a clinically significant interaction, co-administration of losartan 100 mg once daily for 14 days with dronedarone 400 mg bid decreased the C&lt;sub&gt;max&lt;/sub&gt; of losartan by 18% and the C&lt;sub&gt;max&lt;/sub&gt; and AUC&lt;sub&gt;0-24&lt;/sub&gt; of the active metabolite by 25% and 21% respectively.</td>
</tr>
<tr>
<td>Metformin (OCT1 and OCT2 substrate)</td>
<td>CT</td>
<td>No effect</td>
<td>No interaction was observed between dronedarone and metformin.</td>
</tr>
<tr>
<td>Omeprazole (CYP 2C19 substrate)</td>
<td>CT</td>
<td>No effect</td>
<td>No interaction was observed between dronedarone and omeprazole.</td>
</tr>
<tr>
<td>Oral contraceptives (e.g. ethinylestradiol and levonorgestrel)</td>
<td>CT</td>
<td>No effect</td>
<td>No decreases in ethinylestradiol and levonorgestrel were observed in healthy subjects receiving dronedarone (800 mg twice daily) concomitantly with oral contraceptives. However, an increase of 28% in ethinylestradiol exposure and a 19% increase in levonorgestrel exposure were observed.</td>
</tr>
</tbody>
</table>
Table 3 - Established or Potential Drug-Drug Interactions

<table>
<thead>
<tr>
<th>Agent</th>
<th>Ref</th>
<th>Effect</th>
<th>Clinical comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pantoprazole</td>
<td>CT</td>
<td>No effect</td>
<td>Pantoprazole (40 mg once daily), a drug increasing gastric pH without any effect on cytochrome P450, had no significant pharmacokinetic interaction with dronedarone.</td>
</tr>
<tr>
<td>P-glycoprotein (P-gp) substrates (e.g. doxorubicin, fexofenadine and talinolol)</td>
<td>T</td>
<td>↑ P-gp plasma concentration</td>
<td>Dronedarone inhibits P-gp, and interactions may therefore occur with doxorubicin, fexofenadine and talinolol.</td>
</tr>
<tr>
<td>Potent CYP 3A4 inducers (e.g. rifampicin, phenobarbital, carbamazepine, phenytoin)</td>
<td>CT</td>
<td>↓ dronedarone exposure</td>
<td>Rifampicin (600mg once daily) decreased dronedarone exposure by 5-fold with no major change in its active metabolite exposure. Co-administration of rifampicin or other potent CYP3A4 inducers are not recommended with dronedarone as they decrease its exposure.</td>
</tr>
<tr>
<td>Potent CYP 3A4 inhibitors (e.g. ketoconazole, itraconazole, voriconazole, ritonavir, cyclosporin, clarithromycin)</td>
<td>CT</td>
<td>↑ dronedarone exposure</td>
<td>Repeated doses of ketoconazole (200 mg daily), a strong CYP 3A4 inhibitor, resulted in a 17-fold increase in dronedarone exposure. Concomitant use of ketoconazole as well as other potent CYP 3A4 inhibitors is contraindicated (see CONTRAINDICATIONS).</td>
</tr>
<tr>
<td>Sirolimus and Tacrolimus (CYP 3A4 substrates)</td>
<td>T</td>
<td>↑ sirolimus and tacrolimus exposure</td>
<td>Dronedarone can increase plasma concentrations of tacrolimus and sirolimus (CYP3A4 substrates) when given orally. Monitoring of their plasma concentrations and appropriate dosage adjustment is recommended when these drugs are co-administered with dronedarone.</td>
</tr>
</tbody>
</table>
### Table 3 - Established or Potential Drug-Drug Interactions

<table>
<thead>
<tr>
<th>Agent</th>
<th>Ref</th>
<th>Effect</th>
<th>Clinical comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Statins (Substrates of CYP3A4 and/or P-gP substrates e.g. simvastatin, lovastatin, atorvastatin and pravastatin)</td>
<td>CT</td>
<td>↑ simvastatin and simvastatin acid exposure</td>
<td>Dronedarone can increase exposure of statins that are substrates of CYP3A4 and/or P-gP substrates. Dronedarone (400 mg twice daily) increased simvastatin and simvastatin acid exposure by 4-fold and 2-fold respectively.</td>
</tr>
<tr>
<td></td>
<td>CT</td>
<td>↑ atorvastatin, rosvastatin</td>
<td>There was a weak interaction of dronedarone on atorvastatin (1.7-fold) and on statins transported by OATP, such as rosvastatin (1.4-fold).</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>↑ lovastatin, and pravastatin</td>
<td>It is predicted that dronedarone could also increase the exposures of lovastatin, and pravastatin within the same range as simvastatin acid. In clinical trials, there was no evidence of safety concerns when dronedarone was co-administered with statins metabolized by CYP3A4. As high doses of statins increase the risk of myopathy, concomitant use of statins which are CYP3A4 substrates and/or P-gP substrates should be undertaken with caution. Lower starting dose and maintenance doses of statins should be considered according to the statin label recommendations and patients monitored for clinical signs of muscular toxicity. A significant interaction of dronedarone on statins which are not CYP3A4/P-gP substrates such as fluvastatin is unlikely.</td>
</tr>
<tr>
<td>Theophylline (CYP 1A2 substrate)</td>
<td>CT</td>
<td>↓ theophylline exposure</td>
<td>Concomitant administration of dronedarone 400 mg bid with theophylline 400 mg bid resulted in decreased theophylline exposure (18% decrease in AUC_{0-12} and a 17% decrease in C_{max} of theophylline).</td>
</tr>
</tbody>
</table>
Table 3 - Established or Potential Drug-Drug Interactions

<table>
<thead>
<tr>
<th>Agent</th>
<th>Ref</th>
<th>Effect</th>
<th>Clinical comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Warfarin and other vitamin K</td>
<td>CT</td>
<td>↑ S-warfarin exposure</td>
<td>Dronedarone (600 mg twice daily) increased S-warfarin by 1.2-fold with no change in R-warfarin and increased INR by only 1.07-fold.</td>
</tr>
<tr>
<td>antagonists (CYP 2C9 substrate)</td>
<td></td>
<td></td>
<td>In the ATHENA study, more patients taking oral anticoagulants experienced clinically significant INR elevations (≥5) within 1 week of starting dronedarone (4.7%) versus placebo (1.7%). The dose of oral anticoagulant at baseline and any change in dose were not monitored, but after 1 month, there were no further differences between the treatment groups. Furthermore, no excess risk of bleeding was observed in the dronedarone group.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Increased INR (≥5) values with or without bleeding events were reported in post-marketing cases in patients treated concomitantly with oral vitamin K antagonists and dronedarone. In 70% of these cases, this increase occurred within 2 weeks of starting dronedarone. In 80% of the cases where the dose of the anticoagulant was adjusted, INR recovered to therapeutic values. INR should be monitored closely for the first 6 weeks after initiating dronedarone treatment in patients taking vitamin K antagonists.</td>
</tr>
</tbody>
</table>

Legend: C = Case Study; CT = Clinical Trial; T = Theoretical

**Drug-Food Interactions**

**Grapefruit juice**
Repeated doses of 300 mL of grapefruit juice (inhibitor of the CYP3A4) three times daily resulted in a 3-fold increase in dronedarone exposure. Therefore, patients should be warned to avoid grapefruit juice beverages while taking dronedarone.

**Drug-Herb Interactions**

Co-administration of St John’s Wort, a potent CYP3A4 inducer, is not recommended with dronedarone as this herb decreases its exposure.

**DOSAGE AND ADMINISTRATION**

There is limited information on the optimal timing to switch from amiodarone to MULTAQ. It should be considered that amiodarone may have a long duration of action after discontinuation due to its long half-life. If a switch is considered, this should be done with caution under the supervision of a specialist.

**Recommended Dose and Dosage Adjustment**

The recommended dosage is 400 mg twice daily in adults. MULTAQ should be taken as one tablet with the morning meal and one tablet with the evening meal.
Treatment with class I or III antiarrhythmics (such as flecainide, propafenone, quinidine, disopyramide, dofetilide, sotalol, amiodarone) must be stopped before starting MULTAQ.

Treatment with MULTAQ can be initiated in an outpatient setting.

**Renal Impairment**
No dose adjustment is required in patients with renal impairment (see WARNINGS AND PRECAUTIONS, Special Populations and ACTION AND CLINICAL PHARMACOLOGY, Pharmacokinetics).

**Severe Hepatic impairment**
MULTAQ is contraindicated in patients with severe hepatic impairment (see CONTRAINDICATIONS).

**Missed Dose**
If a dose of this medication has been missed, the missed dose should be skipped and the patient should go back to the regular dosing schedule. The dose should not be doubled.

**OVERDOSAGE**
In the event of overdosage, the patient’s cardiac rhythm and blood pressure should be monitored in addition to general supportive measures. Treatment should be supportive and based on symptoms.
It is not known whether dronedarone and/or its metabolites can be removed by dialysis (hemodialysis, peritoneal dialysis or hemofiltration). There is no specific antidote available.

For the management of a suspected drug overdose, contact your regional Poison Control Centre.

**ACTION AND CLINICAL PHARMACOLOGY**

**Mechanism of Action**
In animals, dronedarone prevents atrial fibrillation or restores normal sinus rhythm depending on the model used. It also prevents ventricular tachycardia and ventricular fibrillation in several animal models. These effects most likely result from its electrophysiological properties belonging to all four Vaughan-Williams classes. Dronedarone is a multichannel blocker inhibiting the potassium currents (including \( I_{K(Ach)} \), \( I_{Kur} \), \( I_{Kr} \), \( I_{Ks} \)) and thus prolonging cardiac action potential and refractory periods (Class III). It also inhibits the sodium currents (Class Ib) and the calcium currents (Class IV). It non-competitively antagonizes adrenergic activities (Class II).

**Pharmacodynamics:**
In animal models, dronedarone reduces heart rate. It prolongs Wenckebach cycle length and AH-, PQ-, QT-intervals; with no marked effect on QTc-, HV- and QRS-intervals. It increases effective refractory periods of the atrium, atrio-ventricular node and ventricle with minimal degree of reverse-use dependency.

Dronedarone decreases arterial blood pressure and myocardial contractility (dP/dt\textsubscript{max}) with no change in left ventricular ejection fraction and reduces myocardial oxygen consumption.

Dronedarone has vasodilatory properties, more pronounced in coronary arteries (related to the activation of the nitric oxide pathway) than in peripheral arteries.

Dronedarone displays indirect antiadrenergic effects; it reduces alpha-adrenergic blood pressure response to epinephrine and beta1 and beta2 responses to isoproterenol.

**Pharmacokinetics:**

<table>
<thead>
<tr>
<th>Single dose mean</th>
<th>C\textsubscript{max}</th>
<th>t\textsubscript{1/2} (h)</th>
<th>AUC\textsubscript{0-\infty}</th>
<th>Clearance</th>
<th>Volume of distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>400 mg</td>
<td>67.2 (36)</td>
<td>17.6 (56)</td>
<td>474 (33)</td>
<td>937 (35)</td>
<td>21000 (31)</td>
</tr>
</tbody>
</table>

**Absorption:**
Dronedarone should be taken with food. Following oral administration in fed conditions, dronedarone is well absorbed (at least 70 %) from intestines to blood. However due to presystemic first pass metabolism, the absolute bioavailability of dronedarone (given with food) is 15 %. Concomitant intake of food increases dronedarone bioavailability by on average 2- to 4-fold. After oral administration in fed conditions, peak plasma concentrations of dronedarone and the main circulating active metabolite (N-debutyl metabolite) are reached within 3 to 6 hours.

After repeated administration of 400 mg twice daily, steady state is reached within 4 to 8 days of treatment and the mean accumulation ratio for dronedarone ranges from 2.6 to 4.5. The steady state mean dronedarone C\textsubscript{max} is 84-147 ng/mL and the exposure of the main N-debutyl metabolite is similar to that of the parent compound.

The pharmacokinetics of dronedarone and its N-debutyl metabolite both deviate moderately from dose proportionality: a 2-fold increase in dose results in an approximate 2.5- to 3.0- fold increase with respect to C\textsubscript{max} and AUC.

**Distribution:**
The in vitro plasma protein binding of dronedarone and its N-debutyl metabolite is > 98 % and not saturable. Both compounds bind mainly to albumin. After intravenous (IV) administration the volume of distribution at steady state (V\textsubscript{ss}) ranges from 1200 to 1400 L

**Metabolism:**
Dronedarone is extensively metabolized, mainly by CYP3A4. The major metabolic pathway includes N-debutylation to form the main circulating active metabolite followed by oxidation,
oxidative deamination to form the inactive propanoic acid metabolite, followed by oxidation, and direct oxidation. Monoamine oxidases contribute partially to the metabolism of the active metabolite of dronedarone. The N-debutyl metabolite exhibits pharmacodynamic activity but is 3 to 10-times less potent than dronedarone.

**Excretion:**
After oral administration, approximately 6 % of the labeled dose is excreted in urine mainly as metabolites (no unchanged compound excreted in urine) and 84 % are excreted in feces mainly as metabolites. After IV administration the plasma clearance of dronedarone ranges from 130 to 150 L/h. The terminal elimination half-life of dronedarone is around 25 - 30 hours and that of its N-debutyl metabolite around 20 - 25 hours. In patients, dronedarone and its metabolite are completely eliminated from the plasma within 2 weeks after the end of a 400 mg twice daily-treatment.

**Special Populations and Conditions**

The pharmacokinetics of dronedarone in patients with atrial fibrillation is consistent with that in healthy subjects. The main sources of variability in dronedarone exposure (age, gender, bodyweight, concomitant treatment with weak to moderate CYP3A4 inhibitors) remain modest in their magnitude (less than 2-fold).

Assessment of intrinsic factors such as race, gender or age on the incidence of any treatment emergent adverse events did not suggest any excess of adverse events in a particular sub-group.

**Gender**
In female patients, dronedarone exposures are on average 30% higher as compared to male patients (see ADVERSE EVENTS, Overview).

**Race**
Pharmacokinetic differences due to race were not assessed.

**Elderly**
Of the total number of subjects in clinical studies of dronedarone, 73% were 65 years of age and over and 34% were 75 and over. In patients aged 65 years old and above, dronedarone exposures are 23% higher in comparison with patients aged below 65 years (see WARNINGS AND PRECAUTIONS, Special Populations).

**Pediatric population**
The pharmacokinetics of dronedarone were not assessed in pediatric patients.

**Liver Impairment**
In subjects with moderate hepatic impairment, dronedarone total and unbound exposures are increased by 1.3-fold and by 2-fold respectively. Those of the active metabolite are decreased by 1.6-fold to 1.9-fold. The effect of severe hepatic impairment on the pharmacokinetics of dronedarone was not assessed (see CONTRAINDICATIONS).
Renal Impairment
Consistent with the very weak renal excretion of dronedarone, no pharmacokinetic modification was observed in patients with renal impairment in particular in patients with severe renal impairment (see WARNINGS AND PRECAUTIONS, Special Populations).

STORAGE AND STABILITY

Store at 25°C, with excursion permitted between 15 and 30°C.

Store in the original package.

DOSAGE FORMS, COMPOSITION AND PACKAGING

MULTAQ is provided as tablets for oral administration.

Each tablet of MULTAQ contains dronedarone hydrochloride equivalent to 400 mg dronedarone”.

The inactive ingredients are:

Core of the tablets: colloidal anhydrous silica, crospovidone, hypromellose, lactose monohydrate, magnesium stearate, maize starch, poloxamer 407.

Coating / Polishing of the tablets: carnauba wax, hypromellose, macrogol 6000, titanium dioxide.

MULTAQ 400 mg tablets are provided as white film-coated tablets for oral administration, oblong-shaped, engraved with a double wave marking on one side and “4142” code on the other side in: bottles of 60 tablets, 180 tablets, 500 tablets, and in boxes of 4 blisters (15 tablets per blister).
PART II: SCIENTIFIC INFORMATION

PHARMACEUTICAL INFORMATION

Drug Substance

Common name: Dronedarone hydrochloride

Chemical name:

N-{2-butyl-3-[4-(3-dibutylaminopropoxy)benzoyl]benzofuran-5-yl} methanesulfonamide, hydrochloride.

Molecular formula and molecular mass: Its empirical formula is C_{31}H_{44}N_{2}O_{5}S, HCl with a relative molecular mass of 593.2 (556.8 for dronedarone base).

Structural formula:

![Structural formula of Dronedarone hydrochloride]

Physicochemical properties: Dronedarone HCl is a white to practically white fine powder that is practically insoluble in water and freely soluble in methylene chloride and methanol.
CLINICAL TRIALS

The efficacy of dronedarone in the reduction of risk of cardiovascular hospitalization or death from any cause was demonstrated in patients with AF (atrial fibrillation) or a history of AF/AFL and additional risk factors in the ATHENA multicenter, multinational, double blind, and randomized placebo-controlled study. Patients were to have at least one risk factor (including age, hypertension, diabetes, prior cerebrovascular accident, left atrium diameter ≥ 50 mm or LVEF<0.40) together with AF/AFL and sinus rhythm both documented within the last 6 months. Patients could be in AF/AFL or in sinus rhythm after spontaneous conversion or following any procedures. Four thousand six hundred and twenty eight (4628) patients were randomized and treated for up to 30 months maximum (median follow-up: 22 months) with either dronedarone 400 mg twice daily (2301 patients) or placebo (2327 patients), in addition to conventional therapy including beta-blockers (71%), ACE inhibitors or angiotensin receptor blockers (ARBs) (69%) digitalis (14%), calcium antagonists (14%), statins (39%), oral anticoagulants (60%), chronic antiplatelet therapy (5%) and/or diuretics (54%).

The primary endpoint of the study was the time to first hospitalization for ‘cardiovascular reasons’ or death from any cause. With few exceptions, the 17 components comprising ‘cardiovascular hospitalization’ included many not generally accepted as being causally related to atrial fibrillation. Secondary endpoints evaluated were time to death from any cause, time to first hospitalization for ‘cardiovascular reasons’. Time to cardiovascular death and time to sudden death were also assessed.

Patients ranged in age from 23 to 97 years and 42% were over 75 years old. Forty seven percent (47%) patients were female and a majority Caucasian (89%). The majority had hypertension (86%) and structural heart disease (60% - which included coronary artery disease, congestive heart failure and left ventricular ejection fraction <45%. Twenty five percent (25%) had AF at baseline.

Dronedarone reduced the incidence of cardiovascular hospitalization or death from any cause by 24.2% when compared to placebo.

The curves showing the overall event rate are displayed in Figure 1.
Figure 1 – Kaplan-Meier cumulative incidence curves from randomization to first cardiovascular hospitalization or death from any cause truncated at 24 months

As can be seen from Table 6, it was a reduction in the recurrence of atrial fibrillation and other supraventricular rhythm disorders that resulted in the reduction in hospitalization for
‘cardiovascular reasons’. Hospitalizations for all other reasons were very similar in both dronedarone and placebo groups.

| Table 6-Number (%) of patients with a cardiovascular hospitalization according to pre specified main reason per the Investigator during on-study period – all randomized patients (EFC5555/ATHENA) |
|-------------------------------------------------|----------------|----------------|
| Placebo (N=2327) | Dronedarone 400 mg BID (N=2301) |
| Any cardiovascular hospitalization | 859 (36.9%) | 675 (29.3%) |
| Atrial fibrillation and other supraventricular rhythm disorders | 457 (19.6%) | 296 (12.9%) |
| Worsening CHF, including pulmonary edema or dyspnea of cardiac origin | 92 (4.0%) | 78 (3.4%) |
| Myocardial infarction or unstable angina | 61 (2.6%) | 48 (2.1%) |
| Stable angina pectoris or atypical chest pain | 41 (1.8%) | 45 (2.0%) |
| TIA or stroke (except intracranial hemorrhage) | 35 (1.5%) | 28 (1.2%) |
| Transcutaneous coronary, cerebrovascular or peripheral procedure | 31 (1.3%) | 27 (1.2%) |
| Implantation of a pacemaker, ICD or any other cardiac device | 29 (1.2%) | 32 (1.4%) |
| Major bleeding (requiring two or more units of blood or any intracranial hemorrhage) | 24 (1.0%) | 21 (0.9%) |
| Syncope | 24 (1.0%) | 21 (0.9%) |
| Cardiovascular surgery except cardiac transplantation | 23 (1.0%) | 21 (0.9%) |
| Blood pressure related (hypotension, hypertension; except syncope) | 21 (0.9%) | 21 (0.9%) |
| Atherosclerosis related (if not otherwise specified) | 8 (0.3%) | 11 (0.5%) |
| Ventricular tachycardia (non-sustained and sustained VT) | 6 (0.3%) | 6 (0.3%) |
| Pulmonary embolism or deep vein thrombosis | 3 (0.1%) | 10 (0.4%) |
| Non-fatal cardiac arrest | 2 (<0.1%) | 3 (0.1%) |
| Ventricular extrasystoles | 1 (<0.1%) | 1 (<0.1%) |
| Ventricular fibrillation | 1 (<0.1%) | 1 (<0.1%) |
| Cardiovascular infection | 0 (0%) | 4 (0.2%) |
| Other ventricular arrhythmia | 0 (0%) | 1 (<0.1%) |

The reduction in ‘cardiovascular hospitalization' or death from any cause was consistent in all subgroups, irrespective of baseline characteristics or concomitant medications (see Figure 2).
**Figure 2**– Relative risk (dronedarone 400 mg BID versus placebo) estimates with 95% confidence intervals according to selected baseline characteristics-first cardiovascular hospitalization or death from any cause

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>N</th>
<th>RR [95% CI] (a)</th>
<th>P-value (b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;65</td>
<td>873</td>
<td>0.89 [0.71;1.11]</td>
<td></td>
</tr>
<tr>
<td>[65-75[</td>
<td>1830</td>
<td>0.71 [0.60;0.83]</td>
<td></td>
</tr>
<tr>
<td>&gt;=75</td>
<td>1925</td>
<td>0.75 [0.65;0.87]</td>
<td>0.27</td>
</tr>
<tr>
<td>Gender</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>2459</td>
<td>0.74 [0.64;0.85]</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>2169</td>
<td>0.77 [0.67;0.89]</td>
<td>0.65</td>
</tr>
<tr>
<td>Presence of AF/AFL</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>1155</td>
<td>0.74 [0.61;0.91]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>3473</td>
<td>0.76 [0.68;0.85]</td>
<td>0.85</td>
</tr>
<tr>
<td>Structural Heart Disease</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>2732</td>
<td>0.76 [0.67;0.85]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>1853</td>
<td>0.77 [0.65;0.92]</td>
<td>0.85</td>
</tr>
<tr>
<td>LVEF&lt;35% or NYHA&gt;=class I</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>1417</td>
<td>0.74 [0.63;0.87]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>3146</td>
<td>0.77 [0.68;0.87]</td>
<td>0.71</td>
</tr>
<tr>
<td>LVEF(%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;35</td>
<td>179</td>
<td>0.68 [0.44;1.03]</td>
<td></td>
</tr>
<tr>
<td>&gt;=35</td>
<td>4365</td>
<td>0.76 [0.69;0.84]</td>
<td>0.58</td>
</tr>
<tr>
<td>Beta blocking agents</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>3269</td>
<td>0.78 [0.69;0.87]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>1359</td>
<td>0.71 [0.58;0.86]</td>
<td>0.41</td>
</tr>
<tr>
<td>ACE or All receptor antagonists</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>3216</td>
<td>0.74 [0.66;0.83]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>1412</td>
<td>0.79 [0.66;0.95]</td>
<td>0.59</td>
</tr>
<tr>
<td>Digitalis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>629</td>
<td>0.76 [0.59;0.98]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>3999</td>
<td>0.76 [0.68;0.84]</td>
<td>0.96</td>
</tr>
<tr>
<td>Calcium antagonists (c)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>638</td>
<td>0.63 [0.48;0.82]</td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>3990</td>
<td>0.78 [0.70;0.87]</td>
<td>0.15</td>
</tr>
</tbody>
</table>

- a Determined from Cox regression model
- b P-value of interaction between baseline characteristics and treatment based on Cox regression model
- c Calcium antagonists with heart rate lowering effects restricted to diltiazem, verapamil and bepridil

With respect to the secondary endpoint of ‘death from any cause’, the difference between the number of deaths in the two treatment groups did not reach statistical significance (p=0.1758).
DETAILED PHARMACOLOGY

Animal

Pharmacodynamics

_in vitro_ pharmacology

The _in vitro_ pharmacology activities of dronedarone were similar to those of amiodarone but with a higher potency of dronedarone in most studies. The _in vitro_ electrophysiological studies demonstrated that dronedarone (0.01 to 30 μmol/L), as well as amiodarone, is a multi-channel blocker since it inhibits, through a direct interaction, several native ionic channels of cardiomyocytes: rapid sodium channel and L-type and T-type calcium currents, outward potassium currents: I_K1, I_Kr, I_Ks, I_M and I_K(ACH) and the pacemaker current If, and recombinant human channels human Ether-a-go-go Related Gene (hERG), (I_Kr) and Kv1.5 (I_Kur) stably transfected in CHO/human embryonic kidney (HEK) cells.

Among the inward currents, dronedarone had binding affinity for and inhibited the rapid sodium channel. Thus, it reduced the rate of phase 0 of the Action Potential (AP) (dV/dt_max) in atrium and ventricle, since dV/dt_max is a reasonable indicator of changes in the sodium current. The decrease in dV/dt_max by the drug was frequency-dependent, particularly at high rates (100 and 200 beats/min) and with rapid onset kinetics; an effect characteristic of class IB antiarrhythmic agents like lidocaine and also amiodarone.

Dronedarone also had binding affinity for and blocked L-type calcium channels use dependently with rapid onset kinetics and thus displayed calcium antagonistic activity and class IV antiarrhythmic properties. This finding was in accordance with a reduction in the calcium transient (intracellular calcium concentration) and a decrease in shortening (contraction parameter) of isolated ventricular cells or with a reduction of contraction amplitude of isolated atrium, ventricle and heart.

The main potassium currents involved in cardiac repolarization are I_to, I_K1, I_Kr, I_Ks and I_M at the ventricular level, plus I_K(ACH) and I_Kur in the atrium. Dronedarone weakly increased I_to, weakly decreased I_K1 but concentration-dependently reduced I_Kr, I_Ks, I_K(ACH) and I_Kur. At the atrial level, the APD lengthening observed in rabbit atria might be explained by the inhibition of one of these currents involved in repolarization phase. Dronedarone also blocked the carbachol induced I_K(ACH) in atrial cells. These results suggested that dronedarone could effectively antagonize the AP shortening induced by vagal stimulation, as has been demonstrated with amiodarone which inhibits the carbachol-induced APD shortening; dronedarone is thus likely to have antiarrhythmic activity in pathologies where an increase in vagal tone might play a role such as, for example, atrial fibrillation.

Dronedarone showed vasodilatory properties: it caused concentration-dependent decreases in K+ depolarization- and norepinephrine-induced contraction of aortic strips and decreased
coronary resistance of isolated hearts perfused at constant pressure. All these effects of dronedarone might be related, at least in part, to its calcium antagonistic or class IV antiarrhythmic property. However in coronary arteries, as its vasodilating effect was inhibited by a nitric oxide synthase inhibitor, L-NOARG, the coronary-dilator effect of dronedarone was probably related to activation of the nitric oxide pathway.

**In vivo pharmacology**

The *in vivo* electrophysiological and hemodynamic profiles as well as anti-adrenergic activities of dronedarone were similar to those of amiodarone.

Dronedarone decreased heart rate (HR) in the majority of experiments carried out in anesthetized or conscious rats, dogs and pigs. Dronedarone increased both Atrial effective refractory period (ERP) and Atrio-ventricular (AV) nodal ERP whereas ventricular ERP was slightly prolonged with a minimal degree of reverse frequency-dependency. With the reduction in HR, ventricular interval duration (QT) on the electrocardiogram (ECG) was often prolonged but corrected QT (QTc) was rarely prolonged. These acute electrophysiological effects of dronedarone tended to be enhanced after chronic oral dosing.

The multifactorial mechanism of action of dronedarone contributed to its hemodynamic effects. Dronedarone had α₁-, β₁- and β₂-anti adrenergic effects (although there was weak affinity for these receptors) and calcium antagonist properties that might contribute to the vasodilating and, possibly, to the negative inotropic effects. Also, dronedarone transiently increased coronary blood flow in dogs. Main hemodynamic effects of dronedarone were a decrease in contractility (LV dP/dt max) and an increase in LV end diastolic pressure observed at relatively high concentrations and after intravenous administration.

Chronic oral treatment with dronedarone also decreased plasma concentrations of noradrenaline and these changes correlated with the reduction of HR. This observation suggested that the reduction in the release of catecholamines might participate in the antiadrenergic action of dronedarone as well as the non-competitive β-adrenoceptor antagonistic activity, the down-regulation of the β-adrenoceptor and calcium antagonistic property. A direct activation of α₂ presynaptic adrenoceptors seems unlikely since dronedarone has a weak affinity for α₂ receptors (IC₅₀ = 28 μmol/L) and the peak plasma concentration (1.3 to 1.9 μmol/L) was much lower.

**Pharmacokinetics**

**Absorption and Bioavailability**

Dronedarone appeared to be well absorbed (64 to 95% of the administered dose) following single oral administration in both rats and dogs, based on combined urinary and biliary excretion of radioactivity, and on percentage of parent drug excreted in feces that is likely non-absorbed. Dronedarone underwent an enterohepatic circulation involving about 11% of the dose over 24 hours post dosing as measured in rats intraduodenally infused with the bile of rats pretreated with dronedarone. As a result of the first pass extraction, the absolute oral bioavailability of dronedarone was low (14 to 22%) in rats and dogs and was similar to that found in humans (15%).
SR35021 exposure represented around 22 to 28% of that of dronedarone in rats and around 11% in dogs. In mice, SR35021 achieved plasma levels roughly similar to those of dronedarone.

In toxicokinetic studies, AUC and $C_{\text{max}}$ generally increased more than expected by dose proportionality in rats, rabbits and dogs while they increased roughly in proportion to dose in mice. No significant accumulation of parent drug and SR35021 was observed following repeated oral administration in mice and dogs, while a 2- to 3-fold increase in dronedarone and SR35021 exposure was observed in rats. Steady state was generally already achieved on day 14 of the multiple dose and toxicology studies in mice, rats and dogs whatever the dose, except in rats for doses above and including 17.5 mg/kg/day (later achievement between 1 and 3 months of treatment). No major gender effect was observed.

**Distribution and Protein Binding**

Following both single and multiple oral administrations to rats and dogs, circulating radioactivity was mainly distributed in plasma rather than in blood cells, independent of the route or duration of administration.

Dronedarone and SR35021 were very highly bound to plasma proteins (>99.5% for dronedarone and >98% for SR35021) in all animal species tested (mice, rats, rabbits, dogs, macaques) and humans, ranging from 99.7 to 99.9%. No saturation of protein binding was observed within the range of concentrations tested (1 to 10 000 ng/mL for dronedarone and 50 to 10 000 ng/mL for SR35021) encompassing largely those reached in all animal species and in human.

In rats, dogs and macaques, the pharmacokinetics of dronedarone was characterized by a large volume of distribution. Tissue distribution was investigated in male and female albino rats and in pigmented male rats following single dose and in male albino rats following repeated doses. Both qualitative and quantitative distribution studies were performed. Radioactivity was rapidly and extensively distributed in various tissues. In addition to the gastro-intestinal tract, high concentrations of total radioactivity were observed in the liver and kidney at each timepoint. Radioactivity was also distributed in pituitary gland, lung, adrenals, pancreas, spleen, thyroid, salivary glands, brown fat, Harder’s glands, pineal body and heart. Actual concentrations of total radioactivity were higher following multiple doses, but radioactivity was rapidly eliminated such that the concentrations decreased markedly by 24 hours post dose and were below the limit of quantification between 96 and 336 hours after the last dose. A similar pattern of distribution and elimination of total radioactivity was observed following both single and multiple oral doses of 10 mg/kg $[^{14}\text{C}]$-dronedarone to male rats. At higher doses of 25 and 100 mg/kg affinity of radioactivity for mesenteric lymph nodes was assessed to support carcinogenicity studies. Radioactivity concentration in mesenteric lymph nodes was 2- to 8-fold higher than in mandibular lymph nodes and 5- to 38-fold higher than in blood after single or 28-day repeated administrations. Radioactivity showed an affinity for melanin containing tissues (uveal tract, eye and skin) in male-pigmented rats.

Radioactivity was shown to cross the placental barrier in the rat and in the rabbit, and was detected in fetuses in addition to the fetal area (trophoblast and placenta).
**Metabolism**

Overall, at least 30 metabolites have been detected in the different studied species. The main routes of biotransformation consisted in:

- N-debutylation leading to SR35021 with further oxidation;
- Oxidative deamination leading to the O-propanoic acid derivative (SR90154) with further oxidation;
- Oxidation of the parent compound.

SR35021 has antiarrhythmic, electrophysiological and hemodynamic activities similar to those of dronedarone but is 3 to 10 times less potent than dronedarone. Hence, SR35021 may contribute to the pharmacological activity of dronedarone but not to a major extent. SR90154 has very little or no activity.

Dronedarone undergoes extensive metabolism after oral administration with less than 15% excreted unchanged. In plasma, following single oral administration, dronedarone accounted for approximately 4 to 45% of total radioactivity across species. Numerous metabolites were identified. The main circulating metabolites were SR35021 (N-debutylated metabolite) in mice, SR90154 (propanoic acid derivative) in rats and dogs and oxidated derivatives of dronedarone, SR35021 and SR90154 in rabbits. In human, the main circulating metabolites were SR90154 and SR35021, with plasmatic exposures similar or higher to dronedarone. Exposures estimated in animals for those two metabolites during long term toxicology studies (muse, rat dog) at NOEL or at 1st dose with effects, showed levels in the range or higher than exposure expected in human at the therapeutic dose.

Various oxidative derivates from dronedarone, SR35021 and SR90154 were also circulating in human, accounting for 1.1 to 5.8% of total radioactivity. These metabolites were generally quantified in plasma from animal species for 1.8 to 21% of total radioactivity. Three circulating human metabolites from secondary oxidation pathways (Trihydroxy derivative of dronedarone and N-ethanol derivatives of SR35021) could not be quantified in animal plasma but were observed, or their direct precursor, in excreta of at least one animal species.

The different animal species were representative of the main metabolite pathways observed in human. This support that animals in toxicological studies were exposed to the main human metabolites.

**Excretion and Elimination**

In rats, dogs and macaques, the pharmacokinetics of dronedarone were characterized by a high plasma clearance and a moderate terminal half-life. Dronedarone is rapidly eliminated by metabolic clearance with no excretion of unchanged drug in urine (mouse, rat, rabbit, dog, macaque), and less than 0.5% of unchanged drug in rat bile. In most of the studied species, following oral administration, 72% to 97% of total radioactivity was excreted in feces. Around 44% and 69% of the dose was recovered in the 0-48 hour rat bile collection period following oral and intravenous administrations, respectively. Urinary excretion of radioactivity was less than 9% of the dose, whatever the species. In feces, dronedarone represented 5% to 12% of the administered dose whatever the species examined and could represent non-absorbed drug. These
data demonstrate that metabolism followed by biliary excretion is the main pathway for dronedarone elimination.

The daily excretion of radioactivity in feces remained constant from day 1 to day 14 during the repeated dosing regimen in mice and dogs and slightly increased in rats (80% of a dose on day 1 to 106% on day 14 over the 0-24 hour period). Over a 168-hour period, total recovery after single dosing was more than 90% in all species examined, indicating that drug excretion was almost complete.

Radioactivity was excreted in milk after treatment of lactating dams.

**Human**

**Pharmacodynamics**

**Increase in Serum Creatinine**

ANDROMEDA, a study performed in 627 patients with left ventricular dysfunction and a recent hospitalization for a severe episode of congestive heart failure was stopped prematurely due to an excess of deaths in the dronedarone group[n=25 versus 12 (placebo), p=0.027].

Exploration of the ANDROMEDA findings suggests that increased values of plasma creatinine could be misinterpreted and lead to inappropriate discontinuation of ACE inhibitors or AIIRAs in patients requiring this treatment.

While the proportion of patients treated with ACE inhibitors/AIIRAs at baseline was similar in the placebo and dronedarone groups, 84.2% and 88.4%, respectively, the proportion of patients discontinuing these agents during the study was higher in the dronedarone group 13.2% compared to placebo (5.7%).

Therefore, in the ATHENA trial (see CLINICAL TRIALS) which excluded patients in an unstable hemodynamic condition including congestive heart failure of stage NYHA IV (see CONTRAINDICATIONS) the possibility of a creatinine increase with dronedarone and its appropriate interpretation were explained to investigators, with the recommendation not to inappropriately discontinue ACE inhibitors or AIIRAs in patients requiring this treatment (see WARNINGS AND PRECAUTIONS, Renal and WARNINGS AND PRECAUTIONS, Monitoring and Laboratory Tests).

**TOXICOLOGY**

**Single-Dose Toxicity**

Single dose toxicity studies with oral (1500 and 2000 mg/kg) or intravenous (up to 20 mg/kg) dronedarone administration were performed in mice and rats of both sexes.

Orally, the maximum nonlethal dose in the mouse and the rat was 2000 mg/kg. A single oral administration of 2000 mg/kg of dronedarone caused some clinical signs in rats only (prostration,
The findings from the oral studies suggest that dronedarone has a very limited potential for toxicity in humans in acute overdose situations.

**Repeat-Dose Toxicity**

In the pivotal subacute and chronic toxicity studies, dronedarone was administered from 2 to 160 mg/kg/day in rats from 2 weeks to 6 months and from 5 to 140 mg/kg/day in dogs from 2 weeks to 1 year by the oral route.

Health deterioration changes, subsequent to the other treatment-related effects described below, were observed at doses higher than 60 mg/kg/day. Health deterioration was clearly dose related throughout the studies conducted with dronedarone.

Signs of health deterioration were marked at the high dose of 160 mg/kg/day in rats and led to mortality in 4/20 rats from day 14 to day 16 in the 2 week study after a decline in health status: clinical signs of poor condition were characterized by piloerection, reduced activity, soiled urogenital area, occurring mainly in the second week of the study and was associated with weight loss and decreased food consumption.

The presence of a poor general health condition was supported at histopathological examination by the occurrence with classical health deterioration changes consisting of fat tissue atrophy, decreased hepatocytic margination (indicative of decreased physiological glycogen storage) and reduced size of hepatocytes, renal cortical tubular distensions, atrophy of crural muscle, atrophy (serous dedifferentiation) in the pancreas, parotid and submaxillary glands, atrophy of the small intestine mucosa, lymphoid atrophy of the thymus (involution) or lymph nodes, hyperplasia, necrosis or hemorrhage in the adrenal cortex, fatty involution of the bone marrow, thinning of primary spongiosa in the proximal tibial epiphysis, spermatogenesis changes and atrophy of the male sexual glands.

In the chronic studies in the rat (up to 6 months) and dog (up to 1 year), health deterioration was slight and noted only at the highest dose and mainly shown by weight loss or decreased weight gain associated with decreased food consumption, with a tendency for recovery during the chronic studies.

In subacute and chronic toxicity studies, slight and reversible phospholipidosis (accumulation of foamy macrophages) was observed in mesenteric lymph nodes mainly in the rat. This effect is considered specific to this species and not relevant to humans.

Electrocardiographic changes were noted at 160 mg/kg/day in the 2 week study and at all dose levels in the chronic studies in rats and, in dogs, from 25 mg/kg/day upwards in the 2 week study, at 60 mg/kg/day in the 3 month study and at 45 mg/kg/day in the 1 year study in dogs, which consisted of decreased HR and increased PR and QT (associated with increased QTc only during
the 1 year study), sometimes associated with slightly increased heart weight. These variations were related to the pharmacological properties of the test compound. No ventricular premature complex, no ventricular tachycardia or ventricular fibrillations were observed in any of the studies and there was no death associated with cardiac effects.

No mortality other than that imputed to nonspecific effects in rats at 160 mg/kg/day was attributed to the compound. It should be noted that no sudden death occurred during any of the studies.

Dronedarone was well tolerated up to and including 60 mg/kg/day in the 3 month rat and 3 month dog studies, up to 50 mg/kg/day in the 6 month rat study and up to 45 mg/kg/day in the one year dog study.

Genotoxicity
Dronedarone had no genotoxic effects, based on one in vivo micronucleus test in mice and four in vitro tests: the Ames test with or without metabolic activation, a DNA repair test on rat hepatocytes, a gene mutation assay on hamster fibroblasts and a cytogenetic study of human lymphocytes.

Carcinogenicity
In 2 year-oral carcinogenicity studies, the highest dronedarone dose administered for 24 months was 70 mg/kg/day in rat and 300 mg/kg/day in mice. Observations were increased incidence of mammary gland tumors in female mice, histiocytic sarcomas in mice and hemangiomas at the mesenteric lymph node level in rats, all at the highest tested dose only (corresponding to an exposure of 5 to 10 times that of the human therapeutic dose).

- **Mammary gland adenocarcinomas/adenoacanthomas** were observed in female mice, with statistical significance at 300 mg/kg/day for adenocarcinomas when compared to the control groups. As mammary tumors are known to be influenced by hormones, prolactin levels were measured in two separate studies and were shown to be slightly increased with a statistically significant difference in treated animals compared to controls. The presence of malignant epithelial tumors in the mammary gland of female mice was not considered to represent a significant risk factor for humans for the following reasons:
  - These tumors were limited to one sex and one species and high dose group.
  - These changes were not associated with any target organ toxicity.
  - The morphology of these neoplasms is very species specific and similar neoplasms are not observed in human breast cancer.
  - The limited increase in tumor incidence was proved to be associated with a consistently moderate increase in prolactin, and the human relevance of rodent prolactin induced mammary carcinogenesis is believed to be low.
• **Histiocytic sarcomas (hemolymphoreticular system)** observed in mice at the highest dose were not considered to represent a significant risk factor for humans due to the following reasons:

  - These changes were observed at a low incidence, with no similar effects in male or female rats, and the relevance of such marginal findings remained equivocal since this could be the result of a normal variation or represent a weak carcinogenic response. There is a great variability of background incidence and this may also be partly attributed to the variety of terminology used.
  - The incidences were similar in magnitude to published incidences in untreated male and female mice from a large reference study.
  - No chemical products induce histiocytic sarcoma in rats; in mice, three compounds are known to cause such changes, two of them are clearly genotoxic and the remaining one has not undergone a complete battery of genotoxic tests.
  - There was no increase in the frequency of other lymphoreticular tumors in both carcinogenicity studies.
  - No analogous neoplasm is known to occur in humans.

• **Benign hemangiomas** observed in the mesenteric lymph nodes in the rat, at the highest dose only were not considered to be relevant for human:

  - It is known that hemangiomas are not precancerous changes and do not transform into malignant hemangiosarcomas in either animals or man.
  - Hemangiomas in man are relatively common in many organs and in many cases are present over the lifespan of an individual but hemangiomas and hemangiosarcomas of lymph nodes are extremely rare in humans.
  - Spontaneous proliferative vascular changes similar to those observed in the rat mesenteric lymph node have been described in mesenteric lymph node in man but are considered completely innocuous. In the literature, it is described how venous or lymphatic obstruction leads to vascular transformation of lymph nodes. The increased incidence in the rat is considered to be in relation with phospholipidosis and accumulation of foamy macrophages in the mesenteric nodes seen in this species, particularly susceptible to this type of effect. It was actually shown that dronedarone accumulates preferentially in the mesenteric lymph nodes when compared with other lymph nodes and with plasma. Thus, the proliferative vascular changes seen in the rat are considered to be reactive-proliferative, rare in occurrence and innocuous, therefore not relevant for man.
  - In conclusion, only the high dose group was involved in both species for these neoplastic changes and an acceptable exposure multiple (5 to 10) exists between the plasma exposures at which an increased incidence of tumors is seen, and the exposure in man at the expected human therapeutic dose. Additionally, none of these observations was considered relevant for humans.
Reproductive and Developmental Toxicity

Dronedarone was not shown to alter fertility in animal studies up to 100 mg/kg/day because the treatment did not induce any adverse effect upon mating performance and fertility or male reproductive organ weights and sperm motility and count at any dosage. Dronedarone is teratogenic in the rat at 100 mg/kg/day. The dose level of 50 mg/kg/day, at which a ratio exposure between rat and human at therapeutic dose is around 3, is not teratogenic. Dronedarone is not teratogenic in rabbits at 60 mg/kg/day. Although the number of litters was low at the dose level of 200 mg/kg/day, it was considered as nonteratogenic. In the light of these findings, dronedarone should be contraindicated in pregnant women. The effects of this compound on the development of offspring (decreased viability indices during lactation at high dose levels, i.e. 90 mg/kg/day) were noted however, the most important changes were the marked tissue concentration and the potential effects on the thyroid function of the fetus.

It is not known whether dronedarone is excreted in human milk. In rats, $[^{14}\text{C}]$-radiolabel was detected in maternal milk after oral administration of 30 mg/kg/day and subsequently in gastrointestinal contents of pups. Milk radioactivity/plasma radioactivity ratio ranged between 2 and 4. Because many drugs are excreted in human milk and because the potential for serious adverse reactions in nursing infants from dronedarone is unknown, women should be advised not to breast feed during treatment with dronedarone.

No studies have been performed in juvenile animals as the target population for the claimed indication is a predominantly elderly adult population.

Phototoxicity

The studies evaluating drug photosensitivity were conducted in the guinea-pig, which is widely accepted as a predictive model for man.

Dronedarone was shown to be only slightly phototoxic, i.e. phototoxic at high dose levels (from 100 mg/kg/day upwards), but not phototoxic at lower doses even following repeated administration. Whereas phototoxicity induced by amiodarone was still noted several weeks after treatment was stopped, the only change seen with dronedarone was the presence of a non-specific mild erythema (a reaction also observed in non treated animals during the course of the study) which persisted for at least 11 days after the last dronedarone administration. In addition, dronedarone was shown not to induce photoallergy, unlike amiodarone.

Immunotoxicity

In the specific immune function study, no primary or compound-related immunotoxicity was observed. No effect interpreted as an impairment of immune function was observed in the general toxicology studies; the thymic atrophy observed in some animals in different studies resulted from overall health deterioration and nonspecific stress (as indicated by several other histopathology findings involving other organs), commonly observed when animals are given very high dosages. Some slight increases in IgG and IgM levels were observed in rats only but these were not considered to be toxicologically relevant in the absence of other compound related effects on the immune system.
In conclusion, the extensive examination of immunotoxicity and immune function in animals has not demonstrated any compound related effect.
REFERENCES


PART III: CONSUMER INFORMATION

**MULTAQ®**
Dronedarone hydrochloride

This leaflet is part III of a three-part "Product Monograph" published when MULTAQ® was approved for sale in Canada and is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about MULTAQ. Contact your doctor or pharmacist if you have any questions about the drug.

**ABOUT THIS MEDICATION**

What the medication is used for:
MULTAQ helps control abnormal heart rhythm called atrial (AF) fibrilllation and can lower the risk for having to go into the hospital for heart problems.

MULTAQ has not been studied in children.

What it does:
MULTAQ is a prescription medicine used in adults to help control abnormal heart rate and rhythm called atrial fibrillation.

When it should not be used:
Do not take MULTAQ if:
- You have an allergy to dronedarone, the active ingredient in MULTAQ, or anything else in MULTAQ. See “What the nonmedicinal ingredients are” for a complete list of ingredients.
- You have heart problems such as heart block or very slow heart rate (bradycardia), unless you have a pacemaker.
- Your ECG (electrocardiogram), a record of the electrical activity of your heart shows a heart disorder called “prolonged QT corrected interval” (more than 500 msec).
- You have a severe heart problem with severe shortness of breath, swelling of your feet or legs, trouble breathing while lying down or sleeping, shortness of breath while moving around (signs of heart failure).
- You take certain medicines that can interact with MULTAQ. See the list under the heading “Do not use these medicines with MULTAQ”.
- You have a type of atrial fibrillation called permanent atrial fibrillation. In permanent AF, a decision has been made not to change back your heart rhythm to normal.
- The percentage of blood leaving your heart each time it contracts is too low (condition called left ventricular dysfunction).
- You have severe liver or lung problems.
- You are pregnant. MULTAQ can harm your baby.
- You are breast-feeding. MULTAQ may pass into your milk. Do not take MULTAQ if you are breast-feeding. Do not breast-feed if you take MULTAQ.

Do not use these medicines with MULTAQ and inform your doctor and pharmacist if you are taking these medications:
- NORVIR® (ritonavir) for HIV infection
- NIZORAL® (ketocazole) and SPORANOX® (itraconazole) for fungal infections
- BIAxin® (clarithromycin) for infections
- Cyclosporine for organ transplant
- Some medicines that can affect how your heart beats:
  - Medicines for mental illness called phenothiazines such as chlorpromazine and thioridazine
  - Medicines for depression such as nortryptiline, ANAFRANIL® (chlomipramine)
  - Medicines for abnormal heart rhythm or fast heartbeat such as sotalol, CORDARONE® (amiodarone), dofetilde, TAMBOCOR® (flecainide), RYTHMOL® (propafenone)
  - PRADAXA® (dabigatran) to thin your blood
  - XARELTO® (rivaroxaban) to thin your blood

What the medicinal ingredient is:
dronedarone hydrochloride

What the nonmedicinal ingredients are:
carnauba wax, colloidal anhydrous silica, crospovidone, hypromellose, lactose monohydrate, macrogol 6000, magnesium stearate, maize starch, poloxamer 407 and titanium dioxide

What dosage forms it comes in:
Film coated tablets containing 400 mg dronedarone, as dronedarone hydrochloride.

**WARNINGS AND PRECAUTIONS**

**BEFORE you use MULTAQ talk to your doctor or pharmacist if you are using:**

- Medicine for high blood pressure, chest pain, or other heart conditions, such as:
  - verapamil, diltiazem, nifedipine, metoprolol, propranolol, atenolol, digoxin
  - benazepril and captopril in a therapeutic class called ACE inhibitors
  - losartan, and valsartan in a therapeutic class called ARBs
- Any of these medicines:
  - Fexofenadine, a medicine for allergy or hay fever
  - A medicine to treat cancer (doxorubicin)
  - Statin medicine to lower blood cholesterol such as atorvastatin (LIPITOR®) or simvastatin
  - Rifampicin used for tuberculosis
  - Erythromycin
  - Phenobarbital, carbamazepine, and phenytoin used for seizure disorders
  - St. John’s Wort herbal medicine for depression
Tell your doctor about all your heart problems, including if you have a problem when your heart does not pump the blood through your body as well as it should (heart failure). You should call your doctor right away if any of the following signs appear or if you already had them and they worsen:
- shortness of breath when moving around
- trouble breathing while lying down or sleeping
- swelling of your feet or legs
- weight increase

Tell your doctor about all your liver problems including life threatening liver failure. Your doctor will order blood tests to check your liver function before you start taking MULTAQ and during treatment. In some cases MULTAQ treatment may need to be stopped. You should call your doctor right away if any of the following signs and symptoms of liver problems appear:
- loss of appetite, nausea, vomiting,
- fever, feeling unwell, unusual tiredness,
- itching,
- yellowing of the skin or the whites of the eyes (jaundice),
- unusual darkening of the urine,
- right upper stomach area pain or discomfort.

Tell your doctor about all your other health problems. Including:
- Any health problem that could lead to a low level of potassium or magnesium in your blood.

Tell your doctor if:
- You are pregnant, or planning to become pregnant.
- You are breast-feeding a baby.

Blood test and ECG while taking MULTAQ
While you are taking MULTAQ, your doctor may perform heart examination tests such as an ECG or/and may take blood tests to assess your medical condition and how you tolerate the treatment. The results of one of the blood tests for kidney function (blood creatinine) may be changed by MULTAQ and may lead your doctor to take into account another reference for the “normal” value. Please notify any other doctor you may consult for the first time about your new "normal" value for your blood creatinine.

INTERACTIONS WITH THIS MEDICATION
MULTAQ can interact with certain other medicines and cause serious side effects.

Tell your doctor about all medicines you take. Keep a list of them with you at all times. Show it to your doctor and pharmacist each time you get a new medicine. Do not take any new medicines while you are taking MULTAQ unless you have talked with your doctor. Include all non-prescription medicines, vitamins, and herbal remedies. MULTAQ and certain other medicines can interact with each other, causing serious side effects. Sometimes the dose of other medicines must be changed when they are used with MULTAQ.

Tell your doctor about:
- Water pills for high blood pressure. They could lower your blood potassium and magnesium.
- All heart medicines and blood pressure medicines
- Medicines for hay fever, cancer, or high cholesterol
- Medicines to treat infections, depression, or seizures
- Medicines used in organ transplants
- St. John’s Wort
- Medication for fungal infections

You should avoid:
- Drinking grapefruit juice or eating grapefruit.

PROPER USE OF THIS MEDICATION
Usual dose:
Take MULTAQ as your doctor tells you at the same time each day.
Take 1 MULTAQ tablet (400 mg) 2 times a day with meals.
- Take the first MULTAQ tablet with your morning meal.
- Take the second MULTAQ tablet with your evening meal.

Keep taking MULTAQ even if you feel better. If you stop, your condition may get worse. Keep taking MULTAQ until your doctor tells you to stop. Make sure to observe all appointments with your doctor for check-ups.

Overdose:
In case of drug overdose, contact a health care practitioner, hospital emergency department or Regional Poison Control Centre immediately, even if there are no symptoms.

Taking too much MULTAQ can be dangerous. You may need medical care right away.

Missed Dose:
If you miss a dose, wait and take your next dose at your regular time. Do not take 2 doses at the same time. Do not make up for a missed dose.

SIDE EFFECTS AND WHAT TO DO ABOUT THEM
The common side effects are:
- digestive disturbances such as diarrhea, nausea, vomiting, abdominal pain and indigestion
- feeling tired and weak

Other side effects may also be observed:
• skin problems such as rash and itching redness of skin, eczema, inflammation of the skin, abnormal sensitivity of the skin to sunlight
• change in your sense of taste, loss of taste.

Talk with your doctor about any side effects that bother you. These are not all the side effects of MULTAQ. Your doctor and pharmacist have a more complete list.

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<thead>
<tr>
<th>SERIOUS SIDE EFFECTS, HOW OFTEN THEY HAPPEN AND WHAT TO DO ABOUT THEM</th>
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<td>Symptoms</td>
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This is not a complete list of side effects. For any unexpected effects while taking MULTAQ, contact your doctor or pharmacist.

Reporting Side Effects
You can report any suspected side effects associated with the use of health products to Health Canada by:

- Visiting the Web page on Adverse Reaction Reporting (http://www.hc-sc.gc.ca/dhp-mp's/medeff/report-declaration/index-eng.php) for information on how to report online, by mail or by fax; or
- Calling toll-free at 1-866-234-2345.

NOTE: Contact your health professional if you need information about how to manage your side effects. The Canada Vigilance Program does not provide medical advice.

HOW TO STORE IT

Keep MULTAQ and all other medications out of the reach and sight of children.

Keep the tablets in their original package and store at room temperature (15°-30°C).

MORE INFORMATION

This document plus the full product monograph, prepared for health professionals can be found on Health Canada’s website (http://hc-sc.gc.ca/index-eng.php), at www.sanofi.ca, or by contacting the sponsor, sanofi-aventis Canada Inc., at:
1-800-265-7927

This leaflet was prepared by sanofi-aventis Canada Inc.

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BIAXIN® is a registered trade-mark of Abbott Laboratories
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