APPLICATION NUMBER:
22-425

CHEMISTRY REVIEW(S)
DATE: April 28, 2009

FROM: Donghao (Robert) Lu, Ph.D.
Division of Pre-Marketing Assessment - 1
Office of New Drug Quality Assessment

TO: File NDA 22-425

SUBJECT: OC recommendation

RECOMMENDATION: The drug product MULTAQ (Dronedarone) film-coated tablet, 400 mg, is recommended as APPROVAL from a CMC perspective.

REVIEW NOTE:

The NDA 22-425 CMC review #1 was completed on 2/18/2008. All other CMC issues have been resolved, except one pending issue on the overall recommendation from the Office of Compliance (OC) on manufacturing facilities. We have now received the OC recommendation indicating no inspection concerns (issued on 4/27/09 - an overall recommendation of acceptable). The Establishment Evaluation Report summary is shown below.
Establishment Evaluation Request

Summary Report

Application: NDA 22425/000
Org Code: 110
Priority: 3P

Sponsor: SANOFI AVENTIS US
55 CORPORATE DR
BRIDGEWATER, NJ 08807

Stamp Date: 27-JUN-2008
PDUFA Date: 30-APR-2009
Action Goal:
District Goal: 26-FEB-2009

Brand Name: DROXERHIST HCL
Estab. Name:
Generic Name: DROXERHIST HCL
Dosage Form: (TABLET)
Strength: 400 MG

FDA Contacts: R. FORTNEY
Project Manager
301-796-1068
D. LU
Review Chemist (HFD-150)
301-796-2059
K. SRINIVASACHAR
Team Leader
301-796-1760

Overall Recommendation: ACCEPTABLE on 27-APR-2009 by R. JOHNSON (HFD-320) 301-796-3334

Establishment: CPN: (b)(4) FEI: (b)(4)
(b)(4)
(b)(4)

DMF No: AADA:

Responsibilities: (b)(4)

Profile: (b)
OAI Status: NONE

Last Milestone: QC RECOMMENDATION
Milestone Date: 15-DEC-08
Decision: ACCEPTABLE
Reason: DISTRICT RECOMMENDATION
Establishment: CFN: (b)(4) FEI: (b)(4)

(b)(4)

(b)(4)

DMF No: AADA:

Responsibilities: (b)(4)

Profile: (b) OAI Status: NONE

Last Milestone: OC RECOMMENDATION

Milestone Date: 20-APR-09

Decision: ACCEPTABLE

Reason: DISTRICT RECOMMENDATION

Establishment: CFN: 9610672 FEI: 3002807193

SANOFI AVENTIS PHARMA SA
69583
NEUVILLE-SUR-SAONE 69583, LYON, FR

DMF No: AADA:

Responsibilities: DRUG SUBSTANCE LABELER

DRUG SUBSTANCE MANUFACTURER
ESTABLISHMENT EVALUATION REQUEST

SUMMARY REPORT

DRUG SUBSTANCE PACKAGER
DRUG SUBSTANCE RELEASE TESTER
DRUG SUBSTANCE STABILITY TESTER

Profile : CSN
OAI Status: NONE

Last Milestone: OC RECOMMENDATION
Milestone Date: 12- Aug-08
Decision : ACCEPTABLE
Reason : BASED ON PROFILE

Establishment : CFN : 9610721  FEI : 1463
SANOFI AVENTIS PHARMA SA
63480
VERTOLAYE, FR

DMF No: AADA:

Responsibilities: DRUG SUBSTANCE MICRONIZER
DRUG SUBSTANCE PACKAGER

Profile : CRU
OAI Status: NONE

Last Milestone: OC RECOMMENDATION
Milestone Date: 04-FEB-09
Decision : ACCEPTABLE
Reason : DISTRICT RECOMMENDATION

Establishment : CFN : 1931809  FEI : 1000117506
SANOFI AVENTIS US LLC
6239 LEMAY FERRY RD
SAINT LOUIS, MO 631292805

DMF No: AADA:
Responsibilities: FINISHED DOSAGE OTHER TESTER
FINISHED DOSAGE PACKAGER

Profile : TCM OAI Status: NONE
Last Milestone: OC RECOMMENDATION
Milestone Date: 02-FEB-09
Decision : ACCEPTABLE
Reason : DISTRICT RECOMMENDATION

Establishment : CFN : 9611687 FEI : 3002808341
SANOFI CHIMIE
30390
ARAMON, , FR

DMF No: AADA:

Responsibilities: DRUG SUBSTANCE LABELER
DRUG SUBSTANCE MANUFACTURER
DRUG SUBSTANCE PACKAGER
DRUG SUBSTANCE RELEASE TESTER
DRUG SUBSTANCE STABILITY TESTER
Profile : CSN  OAI Status: NONE

Last Milestone: OC RECOMMENDATION
Milestone Date: 18-AUG-08
Decision: ACCEPTABLE
Reason: BASED ON PROFILE

Establishment: CPN: 9612650  FEI: 3002808206
SANOFI CHIMIE
45 CHEMIN DE METELINE
SISTERON, FR

DMF No: AADA:

Responsibilities: DRUG SUBSTANCE LABELLER
DRUG SUBSTANCE MANUFACTURER
DRUG SUBSTANCE PACKAGER
DRUG SUBSTANCE RELEASE TESTER
DRUG SUBSTANCE STABILITY TESTER

Profile : CSN  OAI Status: NONE

Last Milestone: OC RECOMMENDATION
Milestone Date: 21-APR-09
Decision: ACCEPTABLE
Reason: DISTRICT RECOMMENDATION

Establishment: CPN: 9611342  FEI: 3002808208
SANOFI WINTHROP INDUSTRIE
1, RUE DE LA VIERGE
AMBARES ET LAGRAVE, FR

DMF No: AADA:
Responsibilities: FINISHED DOSAGE MANUFACTURER
FINISHED DOSAGE OTHER TESTER
FINISHED DOSAGE PACKAGER
FINISHED DOSAGE RELEASE TESTER
FINISHED DOSAGE STABILITY TESTER

Profile: TCM
OAI Status: NONE

Last Milestone: OC RECOMMENDATION
Milestone Date: 27-APR-09
Decision: ACCEPTABLE
Reason: DISTRICT RECOMMENDATION

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This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/
Donghao Lu
4/28/2009 09:35:14 AM
CHEMIST

Ramesh Sood
CHEMIST
NDA 22-425

MULTAQ (Dronedarone)
Tablet
400 mg

Sanofi Aventis

Division of Cardio-Renal Drug Products

Donghao (Robert) Lu, Ph.D.

Division of Pre-Marketing Assessment - I
Office of New Drug Quality Assessment
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   A. APPENDICES .............................................................. N/A
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   B. Environmental Assessment Or Claim Of Categorical Exclusion ........................................... N/A

III. Establishment Evaluation Report ................................................................................. N/A

IV. List Of Deficiencies ........................................................................... N/A
Chemistry Review Data Sheet

1. NDA 22-425

2. REVIEW NUMBER: 1

3. REVIEW DATE: 15 AUGUST 2008

4. REVIEWER: Donghao (Robert) Lu, Ph.D.

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<td>NDA 22-425 (Amendment: Revised Proposed Labeling)</td>
<td>13-FEB-09</td>
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7. NAME & ADDRESS OF APPLICANT:

NAME: Sanofi-Aventis U.S. LLC
ADDRESS: 55 Corporate Drive, Bridgewater, NJ 08807
REPRESENTATIVE: Jon Villaume, Ph.D. Vice President, Regulatory Development, Corporate Regulatory Affairs
TELEPHONE: 610-889-6852
8. DRUG PRODUCT NAME/CODE/TYPE:

   PROPRIETARY NAME: MULTAQ (Dronedarone)
   NON-proprietary NAME (USAN): Dronedarone Hydrochloride
   CODE NAME/NUMBER (ONDC ONLY): SR33589B
   CHEMISTRY TYPE / SUBMISSION PRIORITY: 1P

9. LEGAL BASIS FOR SUBMISSION: 505(b)1

10. PHARMACOL. CATEGORY: Anti-Arrhythmic

11. DOSAGE FORM: Tablet

12. STRENGTH/POTENCY: 400 mg

13. ROUTE OF ADMINISTRATION: Oral

14. Rx/OTC DISPENSED: x Rx ___ OTC

15. SPOTS (SPECIAL PRODUCTS ON-LINE TRACKING SYSTEM):
    ___ SPOTS product – Form Completed
    x Not a SPOTS product

16. CHEMICAL NAME, STRUCTURAL FORMULA, MOLECULAR FORMULA, MOLECULAR WEIGHT:

   Name (USAN): Dronedarone hydrochloride
   Name (CAS): Methanesulfonamide, N-[2-butyl-3-[4-[3-(dibutylamino)
               propoxy]benzoyl]-5-benzofuranyl]-, monohydrochloride
   (CAS) Registry Num:
   Structural Formula:

   ![Chemical Structure Image]

   Mol. Formula: C₃₁H₄₅ClN₂O₅S
   Mol. Wt.: 593.22 g/mol
17. RELATED/SUPPORTING DOCUMENTS:

A. DMFs:

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¹ Action codes for DMF Table:
  1 – DMF Reviewed.

Other codes indicate why the DMF was not reviewed, as follows:
  2 – Type 1 DMF
  3 – Reviewed previously and no revision since last review
  4 – Sufficient information in application
  5 – Authority to reference not granted
  6 – DMF not available
  7 – Other (explain under "Comments")

² Adequate, Inadequate, or N/A: There is enough data in the application, therefore the DMF did not need to be reviewed.

B. Other Documents:

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<td>21-913</td>
<td>Dronedarone Hydrochloride</td>
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Note: NDA 21-913 (submitted 10 June 2005) was not approved (action: Not Approvable). NDA 22-425 was submitted to replace NDA 21-913 resubmission. The sponsor incorporated by reference NDA 21-913 and the Complete Response (submitted 27 June 2008) to the 29 August 2006 Action Letter to this New Drug Application 22-425.
18. **STATUS:**

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* see CMC review on NDA 21-913
The Chemistry Review for NDA 22-425

The Executive Summary

I. Recommendations

A. Recommendation and Conclusion on Approvability

The drug product MULTAQ (Dronedarone) film-coated tablet, 400 mg, is pending for approval from a CMC perspective. All CMC related issues had been resolved for this application. At the time of writing this review, the office of compliance had not provided a final recommendation for the manufacturing sites in EES. Because of this pending issue, a final recommendation from CMC perspective is not made. The CMC reviewer will file a final memorandum in DFS once the recommendation from the office of compliance is received.

The sponsor should also make changes on the labels:
(1) The established name should be “Dronedarone” to match the 400 mg strength.
(2) Add word “Tablets” in association with the established name, out side of the parenthesis.

B. Recommendation on Phase 4 (Post-Marketing) Commitments, Agreements, and/or Risk Management Steps, if Approvable

II. Summary of Chemistry Assessments

A. Description of the Drug Substance and Drug Product

1. Drug Substance

The drug substance is dronedarone hydrochloride.

Updated information on the drug substance was provided in this NDA and in DMF No. 18409. The DMF was reviewed and found to be adequate (review dated 10-Dec-2008). It is noted that the drug substance section of the dronedarone hydrochloride was originally provided for NDA 21-913. Additional information can be found in the original CMC reviews for NDA 21-913 and DMF No. 18409.

The drug substance section of the dronedarone hydrochloride in NDA 21-913, initially submitted on June 10, 2005 and subsequently amended with updated stability on December 6, 2005, was further amended with updated CMC information. Only those portions with modifications were provided in this submission and they are evaluated as follows.
Dronedarone hydrochloride has a molecular formula of $C_{21}H_{23}ClN_{2}O_{5}S$ and a relative molecular mass of 593.22 g/mol. The salt/base ratio is 1.065. The synthesis route for dronedarone hydrochloride drug substance was presented in NDA 21-913 in two flow charts: (1) synthesis route to [from] main starting materials; and (2) synthesis route to the drug substance from [from]. The structure of dronedarone hydrochloride was elucidated by elemental analysis, ultraviolet spectrometry, infrared spectrometry, nuclear magnetic resonance spectrometry and mass spectrometry. Information for the impurities was provided early with some updated information in this submission.

Dronedarone hydrochloride was subjected to heat, heat and moisture, light stress, and chemical stress. The drug substance was physically and chemically stable based on evaluation of the testing data. Updated information on drug substance stability (tested for 36 months at 30°C/65%RH and 6 months at 40°C/75%RH) was provided. The data showed that dronedarone hydrochloride drug substance was stable during the test period and further supported the retest period of [from].

2. Drug Product

The drug product is dronedarone hydrochloride 400 mg (base equivalent) immediate release film-coated tablets. The drug product section of the dronedarone hydrochloride was originally provided in NDA 21-913. In this submission, only those portions with modifications were provided and they are evaluated as follows. Additional information can be found in CMC reviews for NDA 21-913.

The tablet is white film-coated oblong tablet with "4142" engraved on one side and a double wave marking on the other. All of the excipients used in the formulation are pharmacopoeial materials, and, as such, are tested against the relevant pharmacopoeial monographs. The drug product manufacturing process, [from], was evaluated early in NDA 21-913. The in-process control tests, which ultimately determine the quality of the drug product, are performed at the following steps in the manufacturing program: [from].

The primary packaging components for dronedarone hydrochloride tablets are [from] Aluminum blister packs and [from] bottles. CMC information to support the 100, 200 and 500 mL [from] bottles, replacing the [from] bottles described in the original NDA, were provided. The stability studies of dronedarone hydrochloride 400 mg film-coated tablets, packaged in the [from] bottles, were provided in this submission, including both primary stability study results and stability results supporting the new container closure system. The stability data supported the proposed expiration period (shelf life) of 36 months.
B. Description of How the Drug Product is Intended to be Used

The drug product MULTAQ (dronedarone hydrochloride) Tablets, 400 mg is indicated for rhythm and rate control in patients with atrial fibrillation (AF) or atrial flutter, in order to maintain normal sinus rhythm or to decrease ventricular rate. AF is a disorder found in about 2.2 million Americans. In it the heart's two small upper chambers (the atria) quiver instead of beating effectively. As a result, blood may pool and clot. Translocation of the clot to an artery in the brain results in a stroke. Stroke occurs in about 15 percent of the individuals that have AF. The likelihood of developing AF increases with age; three to five percent of people over 65 have atrial fibrillation. Multaq is a prescription medicine used in adults to help control these illnesses. Multaq can lower the risk for having to go into the hospital for heart problems or the risk for death.

C. Basis for Approvability or Not-Approval Recommendation

From a CMC perspective, Sanofi Aventis has submitted sufficient and appropriate information to support the approval of the drug product. It should be noted that most of the CMC issues were resolved in NDA 21-913, except one CMC concern that an acceptance criterion for should be listed in addition to the current criterion. In this submission, the resolution to this CMC concern was provided. In addition, other updated information were also provided. Based on this review, all CMC related issues had been resolved.

III. Administrative

A. Reviewer’s Signature

\s\ Donghao (Robert) Lu, Ph.D.

B. Endorsement Block

\s\ Ramesh Sood, Ph.D.

C. CC Block
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/s/
Donghao Lu
2/18/2009 01:56:12 PM
CHEMIST

Ramesh Sood
2/18/2009 03:32:57 PM
CHEMIST
Initial Quality Assessment
Branch I

OND Division: Division of Cardiovascular and Renal Products
NDA: 21-913
Applicant: Sanofi-Aventis
Letter Date: 27 June 2008
Stamp Date: 27 June 2008
PDUFA Date: 27 Dec. 2008
Tradename: Multaq
Established Name: Dronedarone
Dosage Form: Tablets, 400 mg
Route of Administration: Oral
Indication: Reduction in risk of cardiovascular hospitalization or death in patients with atrial fibrillation or flutter
Assessed by: Kasturi Srinivasachar
ONDQA Fileability: Yes

Summary
This is a complete response resubmission of the original NDA which received a ‘Not Approvable’ action on 29 Aug. 2006 for clinical reasons. There were relatively minor CMC issues listed in the action letter concerning the acceptance criteria for drug substance particle size and drug product dissolution. Sanofi has responded to these and, in addition, has provided updated drug substance and drug product information. The DMF 18409 has also been amended. The container closure system for the drug product has been changed based on a comparability protocol submitted in the original NDA. These updates as well as the responses to the CMC questions in the action letter will need an in-depth review.

Labeling
The established name should be ‘dronedarone’ and not ‘dronedarone hydrochloride’ to match the dosage form strength.

Comments and Recommendations
The application is fileable. Facilities, original as well as new ones identified in the resubmission, will be entered into EES and the reviewer should verify the accuracy and completeness of the entries. The CMC reviewer who reviewed the original NDA is recommended for this application.

Kasturi Srinivasachar
Pharmaceutical Assessment Lead
Ramesh Sood, Ph.D.
Branch Chief

July 3, 2008
Date

July 3, 2008
Date
This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/
Kasturi Srinivasachar
7/3/2008 12:03:42 PM
CHEMIST

Ramesh Sood
7/7/2008 07:47:44 AM
CHEMIST
NDA 21-913

MULTAQ® (Dronedarone) Tablets, 400 mg

Sanofi Aventis

REVIEW #2

Donghao R. Lu, Ph.D.
(Drug Substance Reviewer)

William C. Timmer, Ph.D.
(Drug Product Reviewer)

Division of Cardio-Renal Drug Products
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Chemistry Review Data Sheet

1. NDA 21-913

2. REVIEW NUMBER: 2

3. REVIEW DATE: 7 April 2006

4. REVIEWER: Donghao R. Lu, Ph.D.  Drug Substance
   William C. Timmer, Ph.D.  Drug Product

5. PREVIOUS DOCUMENTS:

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7. NAME & ADDRESS OF APPLICANT:

   | NAME:             | Sanofi Aventis       |
   | ADDRESS:          | 11 Great Valley Parkway; Malvern, PA, 19355. |
   | REPRESENTATIVE:   | Douglas A. Greene, M.D. |
   | TELEPHONE:        | 610-889-6425        |
8. DRUG PRODUCT NAME/CODE/TYPE:

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9. LEGAL BASIS FOR SUBMISSION: 505(b)1

10. PHARMACOL. CATEGORY: Anti-Arrhythmic

11. DOSAGE FORM: Tablet

12. STRENGTH/POTENCY: 400 mg

13. ROUTE OF ADMINISTRATION: Oral

14. Rx/OTC DISPENSED: \( _x_ \text{Rx} \quad ___\text{OTC} \)

15. \textbf{SPOTS (SPECIAL PRODUCTS ON-LINE TRACKING SYSTEM):}
   
   \( x \) SPOTS product – Form Completed
   
   \( x \) Not a SPOTS product

16. CHEMICAL NAME, STRUCTURAL FORMULA, MOLECULAR FORMULA, MOLECULAR WEIGHT:

   Name (USAN): Dronedarone hydrochloride
   Name (CAS): methanesulfonamide, \( N\)-[2-butyl-3-[4-[3-(dibutylamino) propoxy]benzoyl]-5-benzofuranyl]-, monohydrochloride
   Molc. Formula: \( C_{31}H_{45}ClN_2O_5S \)
   Molc. Wt.: 593.22 g/mol
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1 Action codes for DMF Table:

1 – DMF Reviewed.

Other codes indicate why the DMF was not reviewed, as follows:

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3 – Reviewed previously and no revision since last review
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2 Adequate, Inadequate, or N/A: There is enough data in the application, therefore the DMF did not need to be reviewed.

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<td>Dronedarone HCl</td>
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<th>RECOMMENDATION</th>
<th>DATE</th>
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<td>Acceptable</td>
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<td>J.M. D’Ambrogio</td>
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<tr>
<td>Methods Validation</td>
<td>-- to be initiated --</td>
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<td>W.C. Timmer, Ph.D.</td>
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<td>J. Jahng, Pharm.D.</td>
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<td>EA</td>
<td>FONSI</td>
<td>29-MAR-06</td>
<td>B. Nguyen, Ph.D.</td>
</tr>
</tbody>
</table>
The Chemistry Review for NDA 21-913

The Executive Summary

I. Recommendations

A. Recommendation and Conclusion on Approvability

The drug product Multaq (Dronedarone HCl) Tablets, 400 mg is recommended as APPROVABLE from a CMC perspective.

B. Recommendation on Phase 4 (Post-Marketing) Commitments, Agreements, and/or Risk Management Steps, if Approvable

There are no Phase IV commitments.

The sponsor has included a Comparability Protocol which involves packaging components. This protocol has been reviewed and found to be acceptable.

II. Summary of Chemistry Assessments

A. Description of the Drug Substance and Drug Product

1. Drug Substance

The drug substance is dronedarone hydrochloride.

Detailed information on the drug substance was provided in DMF No. 18409. The DMF was reviewed and found to be adequate (review dated 10-Jan-2006).

Data from the studies of elemental analysis, UV, IR, NMR and MS demonstrated that the structure was adequately defined.

The synthesis routes and the use of reagents appear adequate.

The impurities detected during the synthesis and development of the DS were evaluated. Analytical methods were developed for the control of the impurities listed above. These methods were briefly described in the tables containing the
CHEMISTRY REVIEW

Executive Summary Section

previous methods and the current methods.

Comprehensive information for all the impurities at the starting material level, at
the intermediate level and at the final synthesis level were presented and
described in the review of the DMF.

Dronedarone hydrochloride, was subjected to heat, heat and moisture, light stress,
and chemical stress. The DS was physically and chemically stable based on
evaluation of the testing data. In addition, the DS appears stable under the storage
condition of 30°C/65% RH for 9 months for primary batches and 48 months for
supporting batch.

2. Drug Product

The drug product is dronedarone hydrochloride 400 mg\(^1\) (base equivalent)
immediate release film-coated tablets.

Dronedarone is active by an oral route. Therefore, tablet dosage forms were formulated for use. The \(\text{(b) (4)}\) were used for Phase 1
and 2A clinical trials, while the film-coated tablets were used for Phase 2B and 3
clinical trials. For reference, bioavailability between the film-coated tablets used
for Phase 2B and the film-coated tablets used for Phase 3 studies has been
established; refer to Section 2.7 Biopharmaceutics.

The commercial DP is a finished tablet that has a white film-coat. The tablet is
oblong and engraved with « 4142 » code on one side and with a double wave
marking on the other side.

All of the excipients used in the formulation are pharmacopoeial materials and,
as such, are tested against the relevant pharmacopoeial monographs.

The DP manufacturing process, \(\text{(b) (4)}\), has been evaluated in this
review. The quality of the DP is controlled by verifying the batch number and
weight of each component. All materials used are tested and released as per
standard practices prior to use. The manufacturing rooms are verified and
inspected to be clean and acceptable for use and only materials required for
specific operation are in the production area.

The in-process control tests, which ultimately determine the quality of the DP,
are performed at the following steps in the manufacturing program: \(\text{(b) (4)}\).

---

\(^{1}\) The drug product contains 426 mg of dronedarone hydrochloride which corresponds to 400 mg dronedarone base.
Again, comprehensive details of the process are available in the manufacturing development report in the submission.

All manufacturing operations are performed in compliance with current GMPs. The batch size of the DP is typically film-coated tablets. Appropriate analytical tests and specifications have been developed and appropriately validated. Details of the validation studies are available in the submission.

The results obtained from analysis of the clinical and primary stability batches during the accelerated, long-term, and photostability studies indicate that the impurities or degradation products do not arise from the DP itself.

The primary packaging components for dronedarone hydrochloride tablets are blister packs and bottles. The blister packs are composed of a rigid, transparent, colourless foil of 20 µm thickness. The 75 ml and 200 ml square white opaque bottles are closed with a child-proof tamper-proof screw cap. The 500 ml rectangular white opaque bottle is closed with a tamper-proof push-fit cap.

B. Description of How the Drug Product is Intended to be Used

The drug product MULTAQ (dronedarone hydrochloride) Tablets, 400 mg is indicated for rhythm and rate control in patients with atrial fibrillation (AF) or atrial flutter, in order to maintain normal sinus rhythm or to decrease ventricular rate.

AF is a disorder found in about 2.2 million Americans. In it the heart's two small upper chambers (the atria) quiver instead of beating effectively. As a result, blood may pool and clot. Translocation of the clot to an artery in the brain results in a stroke. Stroke occurs in about 15 percent of the individuals that have AF. The likelihood of developing AF increases with age; three to five percent of people over 65 have atrial fibrillation.

There are three major goals of medical treatment of AF: the restoration of normal sinus rhythm, control of the ventricular rate during AF, and prevention of blood clot formation.

Restoration of Sinus Rhythm: Sinus rhythm is often restored with medications by slowing the conduction of electrical impulses, decreasing the excitability and
automaticity of cardiac cells, or prolonging the refractory period of cardiac tissue. Several medications may be used to terminate AF including procainimide (Pronestyl), quinidine, disopyramide (Nорpace), amiodarone (Cordarone), and dofetilide (Tikosyn).

Control of Ventricular Rate: To effectively reduce the symptoms associated with AF, it is important that the ventricular rate be controlled. The irregular, flopping sensation in the chest is from the irregular ventricular beat in response to AF. Thus, the faster the ventricles go, the more symptomatic patients usually become. The goal of medications such as beta blockers, calcium channel blockers, and digoxin is to slow down the heart rate by decreasing the excitability of the cardiac cells.

Prevention of Blood Clot Formation: During AF the atria lose their organized pumping action and fibrillate (quiver) in response to the continuous electrical stimulation. In normal sinus rhythm, the atria contract, the valves open and blood fills the ventricles (the lower chambers). The ventricles then contract to complete the organized cycle of contraction that occurs with each heartbeat.

Since the atria don't contract during AF, the blood is not able to empty efficiently from the atria into the ventricles with each heartbeat. Blood can then pool and become stagnant in the atria, creating a site for blood clot formation. Since the left side of the heart pumps the oxygenated blood to all parts of the body, clot formation in the left atrium can become a primary source of stroke in patients with AF.

Other current treatment options for AF include medications, electrical cardioversion, ablations, pacemakers, and surgery. The choice of therapy is quite individualized and is usually based on the degree of disability and symptoms associated with the AF.

C. Basis for Approvability or Not-Approval Recommendation

From a CMC perspective, Sanofi Aventis has submitted sufficient and appropriate information to support an APPROVABLE recommendation for the drug product.

The principal CMC issue involves the particle size distribution. In particular, the drug substance reviewer R. Lu, Ph.D., has submitted the following information request to the sponsor:

In order to better define and control the drug substance particle size distribution please add an acceptance criterion for \[(b) (4)\] in addition to the current \[(b) (4)\] criterion, or justify why it is not necessary to have an acceptance criterion for \[(b) (4)\].
At present, this is the only outstanding CMC issue.

Aside from above, the physical and chemical characteristics, impurity profile, and stability for dronedarone hydrochloride tablets are adequately demonstrated in this submission. The acceptance criteria are appropriate to ensure the identity, strength, quality, potency, and purity of the finished drug product. The criteria are also adequate to assure consistent quality so as to eliminate batch-to-batch variations. In particular, the HPLC assay provides an acceptable degree of separation of dronedarone impurities and degradants. Based on analysis of the stability data, the approved shelf life for Multaq (drodenarone) Tablets, 400 mg is 18 months at room temperature when protected from light.

III. Administrative

A. Reviewer’s Signature

\~ Robert Lu, Ph.D.
\~ William C. Timmer, Ph.D.

B. Endorsement Block

R. Sood, Ph.D. / 7-APR-06
Donghao R. Lu, Ph.D. / 7-APR-06
William C. Timmer, Ph.D. / 7-APR-06

C. CC Block
This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/
---------------------
William Timmer  
4/7/2006 12:41:49 PM  
CHEMIST

Ramesh Sood  
4/7/2006 02:33:41 PM  
CHEMIST
NDA 21-913

MULTAQ® (Dronedarone) Tablets, 400 mg

Sanofi Aventis

Donghao R. Lu, Ph.D.
(Drug Substance Reviewer)

William C. Timmer, Ph.D.
(Drug Product Reviewer)

Division of Cardio-Renal Drug Products
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   B. Endorsement Block ........................................................................................................................... 11
   C. CC Block ......................................................................................................................................... 11

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Chemistry Review Data Sheet

1. NDA 21-913

2. REVIEW NUMBER: 1

3. REVIEW DATE: 1 September 2005

4. REVIEWER: Donghao R. Lu, Ph.D. Drug Substance
             William C. Timmer, Ph.D. Drug Product

5. PREVIOUS DOCUMENTS:

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7. NAME & ADDRESS OF APPLICANT:

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<tr>
<th>NAME:</th>
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<tr>
<td>ADDRESS:</td>
<td>11 Great Valley Parkway; Malvern, PA, 19355.</td>
</tr>
<tr>
<td>REPRESENTATIVE:</td>
<td>Douglas A. Greene, M.D.</td>
</tr>
<tr>
<td>TELEPHONE:</td>
<td>610-889-6425</td>
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8. DRUG PRODUCT NAME/CODE/TYPE:

<table>
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<tr>
<th>PROPRIETARY NAME</th>
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<tr>
<td>NON-PROPRIETARY NAME (USAN)</td>
<td>Dronedarone Hydrochloride</td>
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<tr>
<td>CODE NAME/ NUMBER (ONDC ONLY)</td>
<td>SR33589B</td>
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<td>CHEMISTRY TYPE / SUBMISSION PRIORITY</td>
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9. LEGAL BASIS FOR SUBMISSION: 505(b)1

10. PHARMACOL. CATEGORY: Anti-Arrhythmic

11. DOSAGE FORM: Tablet

12. STRENGTH/POTENCY: 400 mg

13. ROUTE OF ADMINISTRATION: Oral

14. Rx/OTC DISPENSED: _x_Rx ___OTC

15. SPOTS (SPECIAL PRODUCTS ON-LINE TRACKING SYSTEM):

   ____SPOTS product – Form Completed

   ___x__Not a SPOTS product

16. CHEMICAL NAME, STRUCTURAL FORMULA, MOLECULAR FORMULA, MOLECULAR WEIGHT:

   ![Chemical Structure](image)

   Name (USAN): Dronedarone hydrochloride
   Name (CAS): methanesulfonamide, N-[2-butyl-3-[4-[3-(dibutylamino)propoxy]benzoyl]-5-benzofuranyl]-, monohydrochloride
   Molc. Formula: C_{31}H_{45}ClN_{2}O_{5}S
   Molc. Wt.: 593.22 g/mol
17. RELATED/SUPPORTING DOCUMENTS:

A. DMFs:

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1 Action codes for DMF Table:

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Executive Summary Section

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<td>F. Zielinski, Ph.D.</td>
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The Chemistry Review for NDA 21-913

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The impurities detected during the synthesis and development of the DS were evaluated. Analytical methods were developed for the control of the impurities listed above. These methods were briefly described in the tables containing the
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The in-process control tests, which ultimately determine the quality of the DP, are performed at the following steps in the manufacturing program:

---

\(^1\) The drug product contains 426 mg of dronedarone hydrochloride which corresponds to 400 mg dronedarone base.
Again, comprehensive details of the process are available in the manufacturing development report in the submission.

All manufacturing operations are performed in compliance with current GMPs.

The batch size of the DP is typically film-coated tablets.

Appropriate analytical tests and specifications have been developed and appropriately validated. Details of the validation studies are available in the submission.

The results obtained from analysis of the clinical and primary stability batches during the accelerated, long-term, and photostability studies indicate that the impurities or degradation products do not arise from the DP itself.

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**Restoration of Sinus Rhythm:** Sinus rhythm is often restored with medications by slowing the conduction of electrical impulses, decreasing the excitability and
automaticity of cardiac cells, or prolonging the refractory period of cardiac tissue. Several medications may be used to terminate AF including procainimide (Pronestyl), quinidine, disopyramide (Nortace), amiodarone (Cordarone), and dofetilide (Tikosyn).

**Control of Ventricular Rate:** To effectively reduce the symptoms associated with AF, it is important that the ventricular rate be controlled. The irregular, flopping sensation in the chest is from the irregular ventricular beat in response to AF. Thus, the faster the ventricles go, the more symptomatic patients usually become. The goal of medications such as beta blockers, calcium channel blockers, and digoxin is to slow down the heart rate by decreasing the excitability of the cardiac cells.

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Other current treatment options for AF include medications, electrical cardioversion, ablations, pacemakers, and surgery. The choice of therapy is quite individualized and is usually based on the degree of disability and symptoms associated with the AF.

**C. Basis for Approvability or Not-Approval Recommendation**

From a CMC perspective, Sanofi Aventis has submitted sufficient and appropriate information to support the approval of the drug product. The physical and chemical characteristics, impurity profile, and stability for dronedarone hydrochloride tablets are adequately demonstrated in this submission. The acceptance criteria are appropriate to ensure the identity, strength, quality, potency, and purity of the finished drug product. The criteria are also adequate to assure consistent quality so as to eliminate batch-to-batch variations. In particular, the HPLC assay provides an acceptable degree of separation of dronedarone impurities and degradants. Based on analysis of the stability data, the approved shelf life for Multaq (drodenarone) Tablets, 400 mg is 18 months at room temperature when protected from light.
III. Administrative

A. Reviewer’s Signature

\$\$\$ Robert Lu, Ph.D.

\$\$\$ William C. Timmer, Ph.D.

B. Endorsement Block

R. Sood, Ph.D. / 14 Mar-06
Donghao R. Lu, Ph.D. / 01-MAR06
William C. Timmer, Ph.D. / 01-MAR-06

C. CC Block

83 pages of B4 (TS/CCI) material has been withheld in full after this page.
This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/
William Timmer
CHEMIST

Ramesh Sood
3/17/2006 01:25:19 PM
CHEMIST