EXCLUSIVITY SUMMARY

NDA # 200890 SUPPL # HFD # 520

Trade Name Isopto Carpine

Generic Name pilocarpine hydrochloride ophthalmic solution 1%, 2% & 4%

Applicant Name Alcon

Approval Date, If Known June 22, 2010

PART I IS AN EXCLUSIVITY DETERMINATION NEEDED?

1. An exclusivity determination will be made for all original applications, and all efficacy supplements. Complete PARTS II and III of this Exclusivity Summary only if you answer "yes" to one or more of the following questions about the submission.

   a) Is it a 505(b)(1), 505(b)(2) or efficacy supplement?  

      YES ☑  NO

      If yes, what type? Specify 505(b)(1), 505(b)(2), SE1, SE2, SE3, SE4, SE5, SE6, SE7, SE8  

      505(b)(2)

   c) Did it require the review of clinical data other than to support a safety claim or change in labeling related to safety? (If it required review only of bioavailability or bioequivalence data, answer "no.")

      YES ☑  NO

      If your answer is "no" because you believe the study is a bioavailability study and, therefore, not eligible for exclusivity, EXPLAIN why it is a bioavailability study, including your reasons for disagreeing with any arguments made by the applicant that the study was not simply a bioavailability study.

      If it is a supplement requiring the review of clinical data but it is not an effectiveness supplement, describe the change or claim that is supported by the clinical data:
d) Did the applicant request exclusivity?  
YES ☐ NO ☒

If the answer to (d) is "yes," how many years of exclusivity did the applicant request?

e) Has pediatric exclusivity been granted for this Active Moiety?  
YES ☐ NO ☒

If the answer to the above question in YES, is this approval a result of the studies submitted in response to the Pediatric Written Request?

IF YOU HAVE ANSWERED "NO" TO ALL OF THE ABOVE QUESTIONS, GO DIRECTLY TO THE SIGNATURE BLOCKS AT THE END OF THIS DOCUMENT.

2. Is this drug product or indication a DESI upgrade?  
YES ☒ NO ☐

IF THE ANSWER TO QUESTION 2 IS "YES," GO DIRECTLY TO THE SIGNATURE BLOCKS ON PAGE 8 (even if a study was required for the upgrade).

PART II  FIVE-YEAR EXCLUSIVITY FOR NEW CHEMICAL ENTITIES  
(Answer either #1 or #2 as appropriate)

1. Single active ingredient product.

Has FDA previously approved under section 505 of the Act any drug product containing the same active moiety as the drug under consideration? Answer "yes" if the active moiety (including other esterified forms, salts, complexes, chelates or clathrates) has been previously approved, but this particular form of the active moiety, e.g., this particular ester or salt (including salts with hydrogen or coordination bonding) or other non-covalent derivative (such as a complex, chelate, or clathrate) has not been approved. Answer "no" if the compound requires metabolic conversion (other than deesterification of an esterified form of the drug) to produce an already approved active moiety.  

YES ☒ NO ☐

If "yes," identify the approved drug product(s) containing the active moiety, and, if known, the NDA #(#s).
NDA# 20-237 Salagen Tablets (pilocarpine hydrochloride)
NDA# 18-796 Pilopine HS (pilocarpine hydrochloride ophthalmic gel) 4%
NDA# 20-619 BetopticPilo

2. Combination product.

If the product contains more than one active moiety (as defined in Part II, #1), has FDA previously approved an application under section 505 containing any one of the active moieties in the drug product? If, for example, the combination contains one never-before-approved active moiety and one previously approved active moiety, answer "yes." (An active moiety that is marketed under an OTC monograph, but that was never approved under an NDA, is considered not previously approved.)

YES ☐ NO ☒

If "yes," identify the approved drug product(s) containing the active moiety, and, if known, the NDA #(s).

NDA#
NDA#
NDA#

IF THE ANSWER TO QUESTION 1 OR 2 UNDER PART II IS "NO," GO DIRECTLY TO THE SIGNATURE BLOCKS ON PAGE 8. (Caution: The questions in part II of the summary should only be answered “NO” for original approvals of new molecular entities.) IF “YES,” GO TO PART III.

PART III THREE-YEAR EXCLUSIVITY FOR NDAs AND SUPPLEMENTS

To qualify for three years of exclusivity, an application or supplement must contain "reports of new clinical investigations (other than bioavailability studies) essential to the approval of the application and conducted or sponsored by the applicant." This section should be completed only if the answer to PART II, Question 1 or 2 was "yes."

1. Does the application contain reports of clinical investigations? (The Agency interprets "clinical investigations" to mean investigations conducted on humans other than bioavailability studies.) If the application contains clinical investigations only by virtue of a right of reference to clinical investigations in another application, answer "yes," then skip to question 3(a). If the answer to 3(a) is "yes" for any investigation referred to in another application, do not complete remainder of
summary for that investigation.  

YES ☒ NO ☐

IF "NO," GO DIRECTLY TO THE SIGNATURE BLOCKS ON PAGE 8.

2. A clinical investigation is "essential to the approval" if the Agency could not have approved the application or supplement without relying on that investigation. Thus, the investigation is not essential to the approval if 1) no clinical investigation is necessary to support the supplement or application in light of previously approved applications (i.e., information other than clinical trials, such as bioavailability data, would be sufficient to provide a basis for approval as an ANDA or 505(b)(2) application because of what is already known about a previously approved product), or 2) there are published reports of studies (other than those conducted or sponsored by the applicant) or other publicly available data that independently would have been sufficient to support approval of the application, without reference to the clinical investigation submitted in the application.

(a) In light of previously approved applications, is a clinical investigation (either conducted by the applicant or available from some other source, including the published literature) necessary to support approval of the application or supplement?

YES ☒ NO ☐

If "no," state the basis for your conclusion that a clinical trial is not necessary for approval AND GO DIRECTLY TO SIGNATURE BLOCK ON PAGE 8:

(b) Did the applicant submit a list of published studies relevant to the safety and effectiveness of this drug product and a statement that the publicly available data would not independently support approval of the application?

YES ☐ NO ☒

(1) If the answer to 2(b) is "yes," do you personally know of any reason to disagree with the applicant's conclusion? If not applicable, answer NO.

YES ☐ NO ☐

If yes, explain:

(2) If the answer to 2(b) is "no," are you aware of published studies not conducted or sponsored by the applicant or other publicly available data that could independently demonstrate the safety and effectiveness of this drug product?

YES ☒ NO ☐
If yes, explain:

Published clinical trials support the safety and efficacy of the drug product.

(c) If the answers to (b)(1) and (b)(2) were both "no," identify the clinical investigations submitted in the application that are essential to the approval:

Studies comparing two products with the same ingredient(s) are considered to be bioavailability studies for the purpose of this section.

3. In addition to being essential, investigations must be "new" to support exclusivity. The agency interprets "new clinical investigation" to mean an investigation that 1) has not been relied on by the agency to demonstrate the effectiveness of a previously approved drug for any indication and 2) does not duplicate the results of another investigation that was relied on by the agency to demonstrate the effectiveness of a previously approved drug product, i.e., does not redemonstrate something the agency considers to have been demonstrated in an already approved application.

a) For each investigation identified as "essential to the approval," has the investigation been relied on by the agency to demonstrate the effectiveness of a previously approved drug product? (If the investigation was relied on only to support the safety of a previously approved drug, answer "no.")

<table>
<thead>
<tr>
<th>Investigation #1</th>
<th>YES □</th>
<th>NO □</th>
</tr>
</thead>
<tbody>
<tr>
<td>Investigation #2</td>
<td>YES □</td>
<td>NO □</td>
</tr>
</tbody>
</table>

If you have answered "yes" for one or more investigations, identify each such investigation and the NDA in which each was relied upon:

b) For each investigation identified as "essential to the approval", does the investigation duplicate the results of another investigation that was relied on by the agency to support the effectiveness of a previously approved drug product?

<table>
<thead>
<tr>
<th>Investigation #1</th>
<th>YES □</th>
<th>NO □</th>
</tr>
</thead>
<tbody>
<tr>
<td>Investigation #2</td>
<td>YES □</td>
<td>NO □</td>
</tr>
</tbody>
</table>
If you have answered "yes" for one or more investigation, identify the NDA in which a similar investigation was relied on:

c) If the answers to 3(a) and 3(b) are no, identify each "new" investigation in the application or supplement that is essential to the approval (i.e., the investigations listed in #2(c), less any that are not "new"):

4. To be eligible for exclusivity, a new investigation that is essential to approval must also have been conducted or sponsored by the applicant. An investigation was "conducted or sponsored by" the applicant if, before or during the conduct of the investigation, 1) the applicant was the sponsor of the IND named in the form FDA 1571 filed with the Agency, or 2) the applicant (or its predecessor in interest) provided substantial support for the study. Ordinarily, substantial support will mean providing 50 percent or more of the cost of the study.

a) For each investigation identified in response to question 3(c): if the investigation was carried out under an IND, was the applicant identified on the FDA 1571 as the sponsor?

Investigation #1

IND #

YES ☐ NO ☐

! Explain:

Investigation #2

IND #

YES ☐ NO ☐

! Explain:

(b) For each investigation not carried out under an IND or for which the applicant was not identified as the sponsor, did the applicant certify that it or the applicant's predecessor in interest provided substantial support for the study?
Investigation #1

YES ☐ NO ☐

Explain:

Investigation #2

YES ☐ NO ☐

Explain:

(c) Notwithstanding an answer of "yes" to (a) or (b), are there other reasons to believe that the applicant should not be credited with having "conducted or sponsored" the study? (Purchased studies may not be used as the basis for exclusivity. However, if all rights to the drug are purchased (not just studies on the drug), the applicant may be considered to have sponsored or conducted the studies sponsored or conducted by its predecessor in interest.)

YES ☐ NO ☐

If yes, explain:

Literature based NDA.

Name of person completing form: Lori Marie Gorski.
Title: Regulatory Health Project Manager
Date: June 28, 2010

Name of Office/Division Director signing form: Wiley A. Chambers, M.D.
Title: Acting Director

Form OGD-011347; Revised 05/10/2004; formatted 2/15/05
<table>
<thead>
<tr>
<th>Application Type/Number</th>
<th>Submission Type/Number</th>
<th>Submitter Name</th>
<th>Product Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>NDA-200890</td>
<td>ORIG-1</td>
<td>ALCON INC</td>
<td>PILOCARPINE HYDROCHLORIDE OPTHALMIC SOLUTION, 1%, 2% AND 4%</td>
</tr>
</tbody>
</table>

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

/s/

LORI M GORSKI
06/28/2010
Original NDA Exclusivity Summary

WILEY A CHAMBERS
06/28/2010
PEDIATRIC PAGE

NDA/BLA#: 200890  Supplement Number:  NDA Supplement Type (e.g. SE5): ______
Division Name: Division of Anti-Infective and Ophthalmology  PDUFA Goal Date: June 22, 2010  Stamp Date: December 22, 2009
Proprietary Name: ISOPTO Carpine
Established/Generic Name: pilocarpine hydrochloride ophthalmic solution 1, 2, 4%
Dosage Form: topical ophthalmic solution
Applicant/Sponsor: Alcon Research, Ltd.

Indication(s) previously approved (please complete this question for supplements and Type 6 NDAs only):
(1) ______
(2) ______
(3) ______
(4) ______

Pediatric use for each pediatric subpopulation must be addressed for each indication covered by current application under review. A Pediatric Page must be completed for each indication.

Number of indications for this pending application(s): 4
(Attach a completed Pediatric Page for each indication in current application.)

Indication 1: The reduction of elevated intraocular pressure (IOP) in patients with open-angle glaucoma or ocular hypertension

Q1: Is this application in response to a PREA PMR? Yes □ Continue
No ☑ Please proceed to Question 2.

If Yes, NDA/BLA#: ______  Supplement #:______  PMR #:______

Does the division agree that this is a complete response to the PMR?
□ Yes. Please proceed to Section D.
□ No. Please proceed to Question 2 and complete the Pediatric Page, as applicable.

Q2: Does this application provide for (If yes, please check all categories that apply and proceed to the next question):
(a) NEW ☐ active ingredient(s) (includes new combination); ☑ indication(s); ☐ dosage form; ☐ dosing regimen; or ☐ route of administration?*
(b) ☐ No. PREA does not apply. Skip to signature block.
* Note for CDER: SE5, SE6, and SE7 submissions may also trigger PREA.

Q3: Does this indication have orphan designation?
□ Yes. PREA does not apply. Skip to signature block.
☑ No. Please proceed to the next question.

Q4: Is there a full waiver for all pediatric age groups for this indication (check one)?
□ Yes: (Complete Section A.)
☑ No: Please check all that apply:
□ Partial Waiver for selected pediatric subpopulations (Complete Sections B)
□ Deferred for some or all pediatric subpopulations (Complete Sections C)
☑ Completed for some or all pediatric subpopulations (Complete Sections D)
□ Appropriately Labeled for some or all pediatric subpopulations (Complete Sections E)
□ Extrapolation in One or More Pediatric Age Groups (Complete Section F)
(Please note that Section F may be used alone or in addition to Sections C, D, and/or E.)

**Section A: Fully Waived Studies (for all pediatric age groups)**

Reason(s) for full waiver: (check, and attach a brief justification for the reason(s) selected)
- [ ] Necessary studies would be impossible or highly impractical because:
  - [ ] Disease/condition does not exist in children
  - [ ] Too few children with disease/condition to study
  - [ ] Other (e.g., patients geographically dispersed): ______
- [ ] Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients AND is not likely to be used in a substantial number of pediatric patients.
- [ ] Evidence strongly suggests that product would be unsafe in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)
- [ ] Evidence strongly suggests that product would be ineffective in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)
- [ ] Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)
- [ ] Justification attached.

If studies are fully waived, then pediatric information is complete for this indication. If there is another indication, please complete another Pediatric Page for each indication. Otherwise, this Pediatric Page is complete and should be signed.

**Section B: Partially Waived Studies (for selected pediatric subpopulations)**

Check subpopulation(s) and reason for which studies are being partially waived (fill in applicable criteria below):

*Note: If Neonate includes premature infants, list minimum and maximum age in “gestational age” (in weeks).*

<table>
<thead>
<tr>
<th>Reason (see below for further detail):</th>
</tr>
</thead>
<tbody>
<tr>
<td>minimum</td>
</tr>
<tr>
<td>Neonate</td>
</tr>
<tr>
<td>Other</td>
</tr>
<tr>
<td>Other</td>
</tr>
<tr>
<td>Other</td>
</tr>
<tr>
<td>Other</td>
</tr>
<tr>
<td>Other</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? [ ] No; [ ] Yes.

Are the indicated age ranges (above) based on Tanner Stage? [ ] No; [ ] Yes.

Reason(s) for partial waiver (check reason corresponding to the category checked above, and attach a brief justification):
- # Not feasible:
  - [ ] Necessary studies would be impossible or highly impractical because:
    - [ ] Disease/condition does not exist in children
    - [ ] Too few children with disease/condition to study
    - [ ] Other (e.g., patients geographically dispersed): ______
- * Not meaningful therapeutic benefit:
  - [ ] Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric...
patients in this/these pediatric subpopulation(s) AND is not likely to be used in a substantial number of pediatric patients in this/these pediatric subpopulation(s).

† Ineffective or unsafe:

☐ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

△ Formulation failed:

☐ Applicant can demonstrate that reasonable attempts to produce a pediatric formulation necessary for this/these pediatric subpopulation(s) have failed. (Note: A partial waiver on this ground may only cover the pediatric subpopulation(s) requiring that formulation. An applicant seeking a partial waiver on this ground must submit documentation detailing why a pediatric formulation cannot be developed. This submission will be posted on FDA’s website if waiver is granted.)

☐ Justification attached.

For those pediatric subpopulations for which studies have not been waived, there must be (1) corresponding study plans that have been deferred (if so, proceed to Sections C and complete the PeRC Pediatric Plan Template); (2) submitted studies that have been completed (if so, proceed to Section D and complete the PeRC Pediatric Assessment form); (3) additional studies in other age groups that are not needed because the drug is appropriately labeled in one or more pediatric subpopulations (if so, proceed to Section E); and/or (4) additional studies in other age groups that are not needed because efficacy is being extrapolated (if so, proceed to Section F). Note that more than one of these options may apply for this indication to cover all of the pediatric subpopulations.
**Section C: Deferred Studies (for selected pediatric subpopulations).**

Check pediatric subpopulation(s) for which pediatric studies are being deferred (and fill in applicable reason below):

<table>
<thead>
<tr>
<th>Deferrals (for each or all age groups):</th>
<th>Reason for Deferral</th>
<th>Applicant Certification †</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ready for Approval in Adults</td>
<td>Need Additional Adult Safety or Efficacy Data</td>
</tr>
<tr>
<td>Population</td>
<td>minimum</td>
<td>maximum</td>
</tr>
<tr>
<td>Neocate</td>
<td>_ wk. _ mo.</td>
<td>_ wk. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>All Pediatric Populations</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>Date studies are due (mm/dd/yy):</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☐ No; ☐ Yes.

* Other Reason: _____

† Note: Studies may only be deferred if an applicant submits a certification of grounds for deferring the studies, a description of the planned or ongoing studies, evidence that the studies are being conducted or will be conducted with due diligence and at the earliest possible time, and a timeline for the completion of the studies. If studies are deferred, on an annual basis applicant must submit information detailing the progress made in conducting the studies or, if no progress has been made, evidence and documentation that such studies will be conducted with due diligence and at the earliest possible time. This requirement should be communicated to the applicant in an appropriate manner (e.g., in an approval letter that specifies a required study as a post-marketing commitment.)

If all of the pediatric subpopulations have been covered through partial waivers and deferrals, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.
Section D: Completed Studies (for some or all pediatric subpopulations):

Pediatric subpopulation(s) in which studies have been completed (check below):

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
<th>PeRC Pediatric Assessment form attached?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>___ wk. ___ mo.</td>
<td>___ wk. ___ mo.</td>
<td>Yes □</td>
</tr>
<tr>
<td>Other</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
<td>Yes □</td>
</tr>
<tr>
<td>Other</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
<td>Yes □</td>
</tr>
<tr>
<td>Other</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
<td>Yes □</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>0 yr. 0 mo.</td>
<td>16 yr. 11 mo.</td>
<td>Yes ☑</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☑ No; □ Yes.
Are the indicated age ranges (above) based on Tanner Stage? ☑ No; □ Yes.

Note: If there are no further pediatric subpopulations to cover based on partial waivers, deferrals and/or completed studies, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.

Section E: Drug Appropriately Labeled (for some or all pediatric subpopulations):

Additional pediatric studies are not necessary in the following pediatric subpopulation(s) because product is appropriately labeled for the indication being reviewed:

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>___ wk. ___ mo.</td>
<td>___ wk. ___ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>___ yr. ___ mo.</td>
<td>___ yr. ___ mo.</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? □ No; ☑ Yes.
Are the indicated age ranges (above) based on Tanner Stage? □ No; ☑ Yes.

If all pediatric subpopulations have been covered based on partial waivers, deferrals, completed studies, and/or existing appropriate labeling, this Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.

Section F: Extrapolation from Other Adult and/or Pediatric Studies (for deferred and/or completed studies)

Note: Pediatric efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations if (and only if) (1) the course of the disease/condition AND (2) the effects of the product are sufficiently similar between the reference population and the pediatric subpopulation for which information will be extrapolated. Extrapolation of efficacy from studies in adults and/or other children usually requires supplementation with other information obtained from the target pediatric subpopulation, such as

IF THERE ARE QUESTIONS, PLEASE CONTACT THE CDER PMHS VIA EMAIL (cederpmhs@fda.hhs.gov) OR AT 301-796-0700.
pharmacokinetic and safety studies. Under the statute, safety cannot be extrapolated.

Pediatric studies are not necessary in the following pediatric subpopulation(s) because efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations:

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
<th>Extrapolated from:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Adult Studies</td>
<td>Other Studies</td>
<td></td>
</tr>
<tr>
<td>Neonate</td>
<td>__ wk. __ mo.</td>
<td>__ wk. __ mo.</td>
<td>☐</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☑ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☐ No; ☑ Yes.

Note: If extrapolating data from either adult or pediatric studies, a description of the scientific data supporting the extrapolation must be included in any pertinent reviews for the application.

**Indication 2**: The management of acute angle-closure glaucoma

Q1: Is this application in response to a PREA PMR? ☑ Continue

Yes ☐ Continue

No ☑ Please proceed to Question 2.

If Yes, NDA/BLA#: _______ Supplement #:_______ PMR #:_____

Does the division agree that this is a complete response to the PMR?

☐ Yes. Please proceed to Section D.

☐ No. Please proceed to Question 2 and complete the Pediatric Page, as applicable.

Q2: Does this application provide for (If yes, please check all categories that apply and proceed to the next question):

(a) NEW ☑ active ingredient(s) (includes new combination); ☑ indication(s); ☑ dosage form; ☑ dosing regimen; or ☐ route of administration?*

(b) ☐ No. PREA does not apply. **Skip to signature block.**

* Note for CDER: SE5, SE6, and SE7 submissions may also trigger PREA.

Q3: Does this indication have orphan designation?

☐ Yes. PREA does not apply. **Skip to signature block.**

☒ No. Please proceed to the next question.
Q4: Is there a full waiver for all pediatric age groups for this indication (check one)?

☐ Yes: (Complete Section A.)
☒ No: Please check all that apply:

☐ Partial Waiver for selected pediatric subpopulations (Complete Sections B)
☐ Deferred for some or all pediatric subpopulations (Complete Sections C)
☒ Completed for some or all pediatric subpopulations (Complete Sections D)
☐ Appropriately Labeled for some or all pediatric subpopulations (Complete Sections E)
☐ Extrapolation in One or More Pediatric Age Groups (Complete Section F)

(Please note that Section F may be used alone or in addition to Sections C, D, and/or E.)

<table>
<thead>
<tr>
<th>Section A: Fully Waived Studies (for all pediatric age groups)</th>
</tr>
</thead>
</table>

Reason(s) for full waiver: (check, and attach a brief justification for the reason(s) selected)

☐ Necessary studies would be impossible or highly impracticable because:
- Disease/condition does not exist in children
- Too few children with disease/condition to study
- Other (e.g., patients geographically dispersed): ______

☐ Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients AND is not likely to be used in a substantial number of pediatric patients.

☐ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)

☐ Justification attached.

If studies are fully waived, then pediatric information is complete for this indication. If there is another indication, please complete another Pediatric Page for each indication. Otherwise, this Pediatric Page is complete and should be signed.
### Section B: Partially Waived Studies (for selected pediatric subpopulations)

Check subpopulation(s) and reason for which studies are being partially waived (fill in applicable criteria below):

*Note: If Neonate includes premature infants, list minimum and maximum age in “gestational age” (in weeks).*

<table>
<thead>
<tr>
<th>Reason (see below for further detail):</th>
<th>minimum</th>
<th>maximum</th>
<th>Not feasible #</th>
<th>Not meaningful therapeutic benefit *</th>
<th>Ineffective or unsafe †</th>
<th>Formulation failed ∆</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>wk. __ mo.</td>
<td>wk. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Other</td>
<td>yr. __ mo.</td>
<td>yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Other</td>
<td>yr. __ mo.</td>
<td>yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>Other</td>
<td>yr. __ mo.</td>
<td>yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☐ No; ☐ Yes.

Reason(s) for partial waiver (check reason corresponding to the category checked above, and attach a brief justification):

# Not feasible:
- ☐ Necessary studies would be impossible or highly impracticable because:
  - ☐ Disease/condition does not exist in children
  - ☐ Too few children with disease/condition to study
  - ☐ Other (e.g., patients geographically dispersed): _____

* Not meaningful therapeutic benefit:
- ☐ Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients in this/these pediatric subpopulation(s) AND is not likely to be used in a substantial number of pediatric patients in this/these pediatric subpopulation(s).

† Ineffective or unsafe:
- ☐ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations *(Note: if studies are partially waived on this ground, this information must be included in the labeling.)*
- ☐ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations *(Note: if studies are partially waived on this ground, this information must be included in the labeling.)*
- ☐ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations *(Note: if studies are partially waived on this ground, this information must be included in the labeling.)*

∆ Formulation failed:
- ☐ Applicant can demonstrate that reasonable attempts to produce a pediatric formulation necessary for this/these pediatric subpopulation(s) have failed. *(Note: A partial waiver on this ground may only cover the pediatric subpopulation(s) requiring that formulation. An applicant seeking a partial waiver on this ground must submit documentation detailing why a pediatric formulation cannot be developed. This submission will be posted on FDA’s website if waiver is granted.)*

☐ Justification attached.

For those pediatric subpopulations for which studies have not been waived, there must be (1) corresponding study plans that have been deferred (if so, proceed to Sections C and complete the PeRC Pediatric Plan Template); (2) submitted studies that have been completed (if so, proceed to Section D and complete the PeRC Pediatric Assessment form); (3) additional studies in other age groups that are not needed because the drug is appropriately labeled in one or more pediatric subpopulations (if so, proceed to Section E); and/or (4) additional studies in other age groups that are not needed because efficacy is being extrapolated (if so, proceed to Section F). Note that more than one of these options may apply for this indication to cover all of the

IF THERE ARE QUESTIONS, PLEASE CONTACT THE CDER PMHS VIA EMAIL (cderpmhs@fda.hhs.gov) OR AT 301-796-0700.
Section C: Deferred Studies (for selected pediatric subpopulations).

Check pediatric subpopulation(s) for which pediatric studies are being deferred (and fill in applicable reason below):

<table>
<thead>
<tr>
<th>Deferrals (for each or all age groups):</th>
<th>Reason for Deferral</th>
<th>Applicant Certification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Population</td>
<td>Ready for Approval in Adults</td>
<td>Need Additional Adult Safety or Efficacy Data</td>
</tr>
<tr>
<td>Neonate</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>All Pediatric Populations</td>
<td>□</td>
<td>□</td>
</tr>
</tbody>
</table>

Date studies are due (mm/dd/yy): _____

Are the indicated age ranges (above) based on weight (kg)? □ No; □ Yes.

Are the indicated age ranges (above) based on Tanner Stage? □ No; □ Yes.

* Other Reason: _____

† Note: Studies may only be deferred if an applicant submits a certification of grounds for deferring the studies, a description of the planned or ongoing studies, evidence that the studies are being conducted or will be conducted with due diligence and at the earliest possible time, and a timeline for the completion of the studies. If studies are deferred, on an annual basis applicant must submit information detailing the progress made in conducting the studies or, if no progress has been made, evidence and documentation that such studies will be conducted with due diligence and at the earliest possible time. This requirement should be communicated to the applicant in an appropriate manner (e.g., in an approval letter that specifies a required study as a post-marketing commitment.)

If all of the pediatric subpopulations have been covered through partial waivers and deferrals, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.
Section D: Completed Studies (for some or all pediatric subpopulations).

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
<th>PeRC Pediatric Assessment form attached?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>__ wk. __ mo.</td>
<td>__ wk. __ mo.</td>
<td>☐ Yes ☐ No</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐ Yes ☐ No</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐ Yes ☐ No</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>☐ Yes ☐ No</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>0 yr. 0 mo.</td>
<td>16 yr. 11 mo.</td>
<td>☒ Yes ☐ No</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☒ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☐ No; ☒ Yes.

Note: If there are no further pediatric subpopulations to cover based on partial waivers, deferrals and/or completed studies, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.

Section E: Drug Appropriately Labeled (for some or all pediatric subpopulations):

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>__ wk. __ mo.</td>
<td>__ wk. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☒ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☒ No; ☐ Yes.

If all pediatric subpopulations have been covered based on partial waivers, deferrals, completed studies, and/or existing appropriate labeling, this Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.

Section F: Extrapolation from Other Adult and/or Pediatric Studies (for deferred and/or completed studies)

Note: Pediatric efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations if (and only if) (1) the course of the disease/condition AND (2) the effects of the product are sufficiently similar between the reference population and the pediatric subpopulation for which information will be extrapolated. Extrapolation of efficacy from studies in adults and/or other children usually requires supplementation with other information obtained from the target pediatric subpopulation, such as

IF THERE ARE QUESTIONS, PLEASE CONTACT THE CDER PMHS VIA EMAIL (cederpms@fda.hhs.gov) OR AT 301-796-0700.
Pediatric studies are not necessary in the following pediatric subpopulation(s) because efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations:

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
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<th>Extrapolated from:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neocate</td>
<td>_ wk. _ mo.</td>
<td>_ wk. _ mo.</td>
<td>Adult Studies?</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td></td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td></td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)?  □ No; □ Yes.
Are the indicated age ranges (above) based on Tanner Stage?  □ No; □ Yes.

Note: If extrapolating data from either adult or pediatric studies, a description of the scientific data supporting the extrapolation must be included in any pertinent reviews for the application.

Indication 3: The prevention of postoperative elevated IOP associated with laser surgery

Q1: Is this application in response to a PREA PMR?  □ Yes  □ Continue
          □ No  ✗ Please proceed to Question 2.

If Yes, NDA/BLA#: _____  Supplement #:_____  PMR #: ____

Does the division agree that this is a complete response to the PMR?
  □ Yes. Please proceed to Section D.
  □ No. Please proceed to Question 2 and complete the Pediatric Page, as applicable.

Q2: Does this application provide for (If yes, please check all categories that apply and proceed to the next question):
(a) □ NEW active ingredient(s) (includes new combination); □ indication(s); □ dosage form; □ dosing regimen; or □ route of administration?*

(b) □ No. PREA does not apply. Skip to signature block.

* Note for CDER: SE5, SE6, and SE7 submissions may also trigger PREA.

Q3: Does this indication have orphan designation?
  □ Yes. PREA does not apply. Skip to signature block.
  ✗ No. Please proceed to the next question.
Q4: Is there a full waiver for all pediatric age groups for this indication (check one)?

☐ Yes: (Complete Section A.)
☒ No: Please check all that apply:
  ☐ Partial Waiver for selected pediatric subpopulations (Complete Sections B)
  ☐ Deferred for some or all pediatric subpopulations (Complete Sections C)
  ☒ Completed for some or all pediatric subpopulations (Complete Sections D)
  ☐ Appropriately Labeled for some or all pediatric subpopulations (Complete Sections E)
  ☐ Extrapolation in One or More Pediatric Age Groups (Complete Section F)

(Please note that Section F may be used alone or in addition to Sections C, D, and/or E.)

Section A: Fully Waived Studies (for all pediatric age groups)

Reason(s) for full waiver: (check, and attach a brief justification for the reason(s) selected)

☐ Necessary studies would be impossible or highly impracticable because:
  ☉ Disease/condition does not exist in children
  ☐ Too few children with disease/condition to study
  ☐ Other (e.g., patients geographically dispersed): ______

☐ Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients AND is not likely to be used in a substantial number of pediatric patients.

☐ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations (Note: if studies are fully waived on this ground, this information must be included in the labeling.)

☐ Justification attached.

If studies are fully waived, then pediatric information is complete for this indication. If there is another indication, please complete another Pediatric Page for each indication. Otherwise, this Pediatric Page is complete and should be signed.
Section B: Partially Waived Studies (for selected pediatric subpopulations)

Check subpopulation(s) and reason for which studies are being partially waived (fill in applicable criteria below):

Note: If Neonate includes premature infants, list minimum and maximum age in “gestational age” (in weeks).

<table>
<thead>
<tr>
<th>Reason (see below for further detail):</th>
<th>minimum</th>
<th>maximum</th>
<th>Not feasible#</th>
<th>Not meaningful therapeutic benefit*</th>
<th>Ineffective or unsafe†</th>
<th>Formulation failed△</th>
</tr>
</thead>
<tbody>
<tr>
<td>□ Neonate</td>
<td>_ wk. __ mo.</td>
<td>_ wk. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>□ Other</td>
<td>_ yr. __ mo.</td>
<td>_ yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>□ Other</td>
<td>_ yr. __ mo.</td>
<td>_ yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>□ Other</td>
<td>_ yr. __ mo.</td>
<td>_ yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
<tr>
<td>□ Other</td>
<td>_ yr. __ mo.</td>
<td>_ yr. __ mo.</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
<td>☐</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☑ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☐ No; ☑ Yes.

Reason(s) for partial waiver (check reason corresponding to the category checked above, and attach a brief justification):

# Not feasible:

☐ Necessary studies would be impossible or highly impracticable because:

☐ Disease/condition does not exist in children

☐ Too few children with disease/condition to study

☐ Other (e.g., patients geographically dispersed): ______

* Not meaningful therapeutic benefit:

☐ Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients in this/these pediatric subpopulation(s) AND is not likely to be used in a substantial number of pediatric patients in this/these pediatric subpopulation(s).

† Ineffective or unsafe:

☐ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

☐ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

△ Formulation failed:

☐ Applicant can demonstrate that reasonable attempts to produce a pediatric formulation necessary for this/these pediatric subpopulation(s) have failed. (Note: A partial waiver on this ground may only cover the pediatric subpopulation(s) requiring that formulation. An applicant seeking a partial waiver on this ground must submit documentation detailing why a pediatric formulation cannot be developed. This submission will be posted on FDA’s website if waiver is granted.)

☐ Justification attached.

For those pediatric subpopulations for which studies have not been waived, there must be (1) corresponding study plans that have been deferred (if so, proceed to Sections C and complete the PeRC Pediatric Plan Template); (2) submitted studies that have been completed (if so, proceed to Section D and complete the PeRC Pediatric Assessment form); (3) additional studies in other age groups that are not needed because the drug is appropriately labeled in one or more pediatric subpopulations (if so, proceed to Section E); and/or (4) additional studies in other age groups that are not needed because efficacy is being extrapolated (if so, proceed to Section F). Note that more than one of these options may apply for this indication to cover all of the

IF THERE ARE QUESTIONS, PLEASE CONTACT THE CDER PMHS VIA EMAIL (cderpmhs@fda.hhs.gov) OR AT 301-796-0700.
### Section C: Deferred Studies (for selected pediatric subpopulations)

Check pediatric subpopulation(s) for which pediatric studies are being deferred (and fill in applicable reason below):

<table>
<thead>
<tr>
<th>Deferrals (for each or all age groups):</th>
<th>Reason for Deferral</th>
<th>Applicant Certification</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ready for Approval in Adults</td>
<td>Need Additional Adult Safety or Efficacy Data</td>
</tr>
<tr>
<td>Neonate</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>Other</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>All Pediatric Populations</td>
<td>□</td>
<td>□</td>
</tr>
</tbody>
</table>

Date studies are due (mm/dd/yy): _____

Are the indicated age ranges (above) based on weight (kg)? □ No; □ Yes.

Are the indicated age ranges (above) based on Tanner Stage? □ No; □ Yes.

* Other Reason: _____

† Note: Studies may only be deferred if an applicant submits a certification of grounds for deferring the studies, a description of the planned or ongoing studies, evidence that the studies are being conducted or will be conducted with due diligence and at the earliest possible time, and a timeline for the completion of the studies. If studies are deferred, on an annual basis applicant must submit information detailing the progress made in conducting the studies or, if no progress has been made, evidence and documentation that such studies will be conducted with due diligence and at the earliest possible time. This requirement should be communicated to the applicant in an appropriate manner (e.g., in an approval letter that specifies a required study as a post-marketing commitment.)

If all of the pediatric subpopulations have been covered through partial waivers and deferrals, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.
### Section D: Completed Studies (for some or all pediatric subpopulations)

Pediatric subpopulation(s) in which studies have been completed (check below):

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
<th>PeRC Pediatric Assessment form attached?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>_ wk. _ mo.</td>
<td>_ wk. _ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>0 yr. 0 mo.</td>
<td>16 yr. 11 mo.</td>
<td>Yes ☒ No ☐</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☒ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☒ No; ☐ Yes.

**Note:** If there are no further pediatric subpopulations to cover based on partial waivers, deferrals and/or completed studies, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.

### Section E: Drug Appropriately Labeled (for some or all pediatric subpopulations):

Additional pediatric studies are not necessary in the following pediatric subpopulation(s) because product is appropriately labeled for the indication being reviewed:

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>_ wk. _ mo.</td>
<td>_ wk. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>_ yr. _ mo.</td>
<td>_ yr. _ mo.</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☒ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☒ No; ☐ Yes.

If all pediatric subpopulations have been covered based on partial waivers, deferrals, completed studies, and/or existing appropriate labeling, this Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.

### Section F: Extrapolation from Other Adult and/or Pediatric Studies (for deferred and/or completed studies)

**Note:** Pediatric efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations if (and only if) (1) the course of the disease/condition AND (2) the effects of the product are sufficiently similar between the reference population and the pediatric subpopulation for which information will be extrapolated. Extrapolation of efficacy from studies in adults and/or other children usually requires supplementation with other information obtained from the target pediatric subpopulation, such as

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Pediatric studies are not necessary in the following pediatric subpopulation(s) because efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations:

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
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<td></td>
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<td>Adult Studies?</td>
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<td></td>
<td></td>
<td></td>
<td>Other Pediatric Studies?</td>
</tr>
<tr>
<td>Neonate</td>
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<td>__ wk. __ mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
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<td>__ yr. __ mo.</td>
<td></td>
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<tr>
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<td></td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td></td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)?    ☐ No; ☐ Yes.
Are the indicated age ranges (above) based on Tanner Stage?    ☐ No; ☐ Yes.

Note: If extrapolating data from either adult or pediatric studies, a description of the scientific data supporting the extrapolation must be included in any pertinent reviews for the application.

**Indication 4:** Induction of miosis

**Q1:** Is this application in response to a PREA PMR?    ☐ Yes ☐ Continue
☐ No ☒ Please proceed to Question 2.

If Yes, NDA/BLA#: _______ Supplement #:_______ PMR #:______

Does the division agree that this is a complete response to the PMR?
☐ Yes. Please proceed to Section D.
☐ No. Please proceed to Question 2 and complete the Pediatric Page, as applicable.

**Q2:** Does this application provide for (If yes, please check all categories that apply and proceed to the next question):
(a) NEW ☐ active ingredient(s) (includes new combination); ☒ indication(s); ☐ dosage form; ☐ dosing regimen; or ☐ route of administration?*
(b) ☐ No. PREA does not apply. **Skip to signature block.**

* **Note for CDER:** SE5, SE6, and SE7 submissions may also trigger PREA.

**Q3:** Does this indication have orphan designation?
☐ Yes. PREA does not apply. **Skip to signature block.**
☒ No. Please proceed to the next question.
Q4: Is there a full waiver for all pediatric age groups for this indication (check one)?

☐ Yes: (Complete Section A.)
☒ No: Please check all that apply:
  ☐ Partial Waiver for selected pediatric subpopulations (Complete Sections B)
  ☐ Deferred for some or all pediatric subpopulations (Complete Sections C)
  ☒ Completed for some or all pediatric subpopulations (Complete Sections D)
  ☐ Appropriately Labeled for some or all pediatric subpopulations (Complete Sections E)
  ☐ Extrapolation in One or More Pediatric Age Groups (Complete Section F)

(Please note that Section F may be used alone or in addition to Sections C, D, and/or E.)

<table>
<thead>
<tr>
<th>Section A: Fully Waived Studies (for all pediatric age groups)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reason(s) for full waiver: <strong>(check, and attach a brief justification for the reason(s) selected)</strong></td>
</tr>
</tbody>
</table>

☐ Necessary studies would be impossible or highly impracticable because:
  ☐ Disease/condition does not exist in children
  ☐ Too few children with disease/condition to study
  ☐ Other (e.g., patients geographically dispersed): ______

☐ Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients AND is not likely to be used in a substantial number of pediatric patients.

☐ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations *(Note: if studies are fully waived on this ground, this information must be included in the labeling.)*

☐ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations *(Note: if studies are fully waived on this ground, this information must be included in the labeling.)*

☐ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations *(Note: if studies are fully waived on this ground, this information must be included in the labeling.)*

☐ Justification attached.

*If studies are fully waived, then pediatric information is complete for this indication. If there is another indication, please complete another Pediatric Page for each indication. Otherwise, this Pediatric Page is complete and should be signed.*
## Section B: Partially Waived Studies (for selected pediatric subpopulations)

Check subpopulation(s) and reason for which studies are being partially waived (fill in applicable criteria below):

Note: If Neonate includes premature infants, list minimum and maximum age in “gestational age” (in weeks).

<table>
<thead>
<tr>
<th>Reason (see below for further detail):</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Not feasible*</td>
<td>Not meaningful therapeutic benefit†</td>
<td>Ineffective or unsafe‡</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>minimum</th>
<th>maximum</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>wk. mo.</td>
<td>wk. mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. mo.</td>
<td>yr. mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. mo.</td>
<td>yr. mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. mo.</td>
<td>yr. mo.</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. mo.</td>
<td>yr. mo.</td>
<td></td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? □ No; □ Yes.

Are the indicated age ranges (above) based on Tanner Stage? □ No; □ Yes.

Reason(s) for partial waiver (check reason corresponding to the category checked above, and attach a brief justification):

### # Not feasible:
- □ Necessary studies would be impossible or highly impracticable because:
  - □ Disease/condition does not exist in children
  - □ Too few children with disease/condition to study
  - □ Other (e.g., patients geographically dispersed): _____

### * Not meaningful therapeutic benefit:
- □ Product does not represent a meaningful therapeutic benefit over existing therapies for pediatric patients in this/these pediatric subpopulation(s) AND is not likely to be used in a substantial number of pediatric patients in this/these pediatric subpopulation(s).

### † Ineffective or unsafe:
- □ Evidence strongly suggests that product would be unsafe in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)
- □ Evidence strongly suggests that product would be ineffective in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)
- □ Evidence strongly suggests that product would be ineffective and unsafe in all pediatric subpopulations (Note: if studies are partially waived on this ground, this information must be included in the labeling.)

### ∆ Formulation failed:
- □ Applicant can demonstrate that reasonable attempts to produce a pediatric formulation necessary for this/these pediatric subpopulation(s) have failed. (Note: A partial waiver on this ground may only cover the pediatric subpopulation(s) requiring that formulation. An applicant seeking a partial waiver on this ground must submit documentation detailing why a pediatric formulation cannot be developed. This submission will be posted on FDA’s website if waiver is granted.)

### Notice

For those pediatric subpopulations for which studies have not been waived, there must be (1) corresponding study plans that have been deferred (if so, proceed to Sections C and complete the PeRC Pediatric Plan Template); (2) submitted studies that have been completed (if so, proceed to Section D and complete the PeRC Pediatric Assessment form); (3) additional studies in other age groups that are not needed because the drug is appropriately labeled in one or more pediatric subpopulations (if so, proceed to Section E); and/or (4) additional studies in other age groups that are not needed because efficacy is being extrapolated (if so, proceed to Section F). Note that more than one of these options may apply for this indication to cover all of the

IF THERE ARE QUESTIONS, PLEASE CONTACT THE CDER PMHS VIA EMAIL (cedermhbs@fda.hhs.gov) OR AT 301-796-0700.
pediatric subpopulations.

Section C: Deferred Studies (for selected pediatric subpopulations).

Check pediatric subpopulation(s) for which pediatric studies are being deferred (and fill in applicable reason below):

<table>
<thead>
<tr>
<th>Deferrals (for each or all age groups):</th>
<th>Reason for Deferral</th>
<th>Applicant Certification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Population</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neonate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All Pediatric Populations</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Date studies are due (mm/dd/yy):</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)?  
☐ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage?  
☐ No; ☐ Yes.

* Other Reason: ______

† Note: Studies may only be deferred if an applicant submits a certification of grounds for deferring the studies, a description of the planned or ongoing studies, evidence that the studies are being conducted or will be conducted with due diligence and at the earliest possible time, and a timeline for the completion of the studies. If studies are deferred, on an annual basis applicant must submit information detailing the progress made in conducting the studies or, if no progress has been made, evidence and documentation that such studies will be conducted with due diligence and at the earliest possible time. This requirement should be communicated to the applicant in an appropriate manner (e.g., in an approval letter that specifies a required study as a post-marketing commitment.)

If all of the pediatric subpopulations have been covered through partial waivers and deferrals, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.
### Section D: Completed Studies (for some or all pediatric subpopulations).

**Pediatric subpopulation(s) in which studies have been completed (check below):**

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
<th>PeRC Pediatric Assessment form attached?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>__ wk. __ mo.</td>
<td>__ wk. __ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
<td>Yes ☐ No ☐</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>0 yr. 0 mo.</td>
<td>16 yr. 11 mo.</td>
<td>Yes ☑ No ☐</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☒ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☒ Yes; ☐ No

*Note: If there are no further pediatric subpopulations to cover based on partial waivers, deferrals and/or completed studies, Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.*

### Section E: Drug Appropriately Labeled (for some or all pediatric subpopulations):

**Additional pediatric studies are not necessary in the following pediatric subpopulation(s) because product is appropriately labeled for the indication being reviewed:**

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>__ wk. __ mo.</td>
<td>__ wk. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>Other</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>__ yr. __ mo.</td>
<td>__ yr. __ mo.</td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? ☐ No; ☐ Yes.

Are the indicated age ranges (above) based on Tanner Stage? ☑ Yes; ☐ No.

*If all pediatric subpopulations have been covered based on partial waivers, deferrals, completed studies, and/or existing appropriate labeling, this Pediatric Page is complete and should be signed. If not, complete the rest of the Pediatric Page as applicable.*

### Section F: Extrapolation from Other Adult and/or Pediatric Studies (for deferred and/or completed studies)

*Note: Pediatric efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations if (and only if) (1) the course of the disease/condition AND (2) the effects of the product are sufficiently similar between the reference population and the pediatric subpopulation for which information will be extrapolated. Extrapolation of efficacy from studies in adults and/or other children usually requires supplementation with other information obtained from the target pediatric subpopulation, such as*

*IF THERE ARE QUESTIONS, PLEASE CONTACT THE CDER PMHS VIA EMAIL (cderpmhs@fda.hhs.gov) OR AT 301-796-0700.*
pharmacokinetic and safety studies. Under the statute, safety cannot be extrapolated.

Pediatric studies are not necessary in the following pediatric subpopulation(s) because efficacy can be extrapolated from adequate and well-controlled studies in adults and/or other pediatric subpopulations:

<table>
<thead>
<tr>
<th>Population</th>
<th>minimum</th>
<th>maximum</th>
<th>Extrapolated from:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Adult Studies?</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Other Pediatric Studies?</td>
</tr>
<tr>
<td>Neonate</td>
<td>wk. ___</td>
<td>wk. ___</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. ___</td>
<td>yr. ___</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. ___</td>
<td>yr. ___</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>yr. ___</td>
<td>yr. ___</td>
<td></td>
</tr>
<tr>
<td>All Pediatric Subpopulations</td>
<td>yr. ___</td>
<td>yr. ___</td>
<td></td>
</tr>
</tbody>
</table>

Are the indicated age ranges (above) based on weight (kg)? □ No; □ Yes.
Are the indicated age ranges (above) based on Tanner Stage? □ No; □ Yes.

Note: If extrapolating data from either adult or pediatric studies, a description of the scientific data supporting the extrapolation must be included in any pertinent reviews for the application.

If there are additional indications, please complete the attachment for each one of those indications. Otherwise, this Pediatric Page is complete and should be signed and entered into DFS or DARRTS as appropriate after clearance by PeRC.

This page was completed by:

{See appended electronic signature page}

Lori Marie Gorski
Regulatory Project Manager

(Revised: 6/2008)
<table>
<thead>
<tr>
<th>Application Type/Number</th>
<th>Submission Type/Number</th>
<th>Submitter Name</th>
<th>Product Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>NDA-200890</td>
<td>ORIG-1</td>
<td>ALCON INC</td>
<td>PILOCARPINE HYDROCHLORIDE OPTHALMIC SOLUTION, 1%, 2% AND 4%</td>
</tr>
</tbody>
</table>

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

/s/

-----------------------------------------------
LORI M GORSKI
07/02/2010
peds page for original NDA approval
**ACTION PACKAGE CHECKLIST**

### APPLICATION INFORMATION

<table>
<thead>
<tr>
<th>NDA #</th>
<th>200890</th>
</tr>
</thead>
<tbody>
<tr>
<td>BLA #</td>
<td></td>
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<tr>
<td>NDA Supplement #</td>
<td></td>
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<tr>
<td>BLA STN #</td>
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<tr>
<td>If NDA, Efficacy Supplement Type:</td>
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</table>

<table>
<thead>
<tr>
<th>Proprietary Name:</th>
<th>Isopto Carpine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Established/Proper Name:</td>
<td>pilocarpine hydrochloride</td>
</tr>
<tr>
<td>Dosage Form:</td>
<td>ophthalmic solution 1, 2 &amp; 4 %</td>
</tr>
</tbody>
</table>

| Applicant: | Alcon Research Ltd |
| Agent for Applicant (if applicable): | Division: Division of Anti-Infective and Ophthalmology Products |

<table>
<thead>
<tr>
<th>RPM:</th>
<th>Lori Marie Gorski</th>
</tr>
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</table>

**NDAs:**

<table>
<thead>
<tr>
<th>NDA Application Type:</th>
<th>☒ 505(b)(1)</th>
<th>☒ 505(b)(2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Efficacy Supplement:</td>
<td>☒ 505(b)(1)</td>
<td>☒ 505(b)(2)</td>
</tr>
</tbody>
</table>

(A supplement can be either a (b)(1) or a (b)(2) regardless of whether the original NDA was a (b)(1) or a (b)(2). Consult page 1 of the 505(b)(2) Assessment or the Appendix to this Action Package Checklist.)

**505(b)(2) Original NDAs and 505(b)(2) NDA supplements:**

Listed drug(s) relied upon for approval (include NDA #(/s) and drug name(/s)):

Provide a brief explanation of how this product is different from the listed drug.

☒ If no listed drug, check box and explain: application is a literature based NDA

**Two months prior to each action, review the information in the 505(b)(2) Assessment and submit the draft to CDER OND IO for clearance. Finalize the 505(b)(2) Assessment at the time of the approval action.**

On the day of approval, check the Orange Book again for any new patents or pediatric exclusivity.

☒ No changes  ☐ Updated  Date of check: June 18, 2010

If pediatric exclusivity has been granted or the pediatric information in the labeling of the listed drug changed, determine whether pediatric information needs to be added to or deleted from the labeling of this drug.

<table>
<thead>
<tr>
<th>Proposed action</th>
</tr>
</thead>
<tbody>
<tr>
<td>User Fee Goal Date is June 22, 2010</td>
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</table>

<table>
<thead>
<tr>
<th>Previous actions (specify type and date for each action taken)</th>
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</thead>
<tbody>
<tr>
<td>☒ None</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>If accelerated approval or approval based on efficacy studies in animals, were promotional materials received?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes: Promotional materials to be used within 120 days after approval must have been submitted (for exceptions, see <a href="http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/ucm069965.pdf">http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/ucm069965.pdf</a>). If not submitted, explain not accelerated</td>
</tr>
</tbody>
</table>

| ☐ Received |

---

1 The **Application Information** section is (only) a checklist. The **Contents of Action Package** section (beginning on page 5) lists the documents to be included in the Action Package.

Version: 6/8/10
### Application Characteristics

<table>
<thead>
<tr>
<th>Review priority:</th>
<th>Standard</th>
<th>Priority</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical classification (new NDAs only):</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Fast Track</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rolling Review</td>
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<td></td>
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<tr>
<td>Orphan drug designation</td>
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<td></td>
</tr>
<tr>
<td>Rx-to-OTC full switch</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rx-to-OTC partial switch</td>
<td></td>
<td></td>
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<tr>
<td>Direct-to-OTC</td>
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<td></td>
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<tr>
<td>NDAs: Subpart H</td>
<td>BLAs: Subpart E</td>
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<tr>
<td>Accelerated approval (21 CFR 314.510)</td>
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<td>Restricted distribution (21 CFR 314.520)</td>
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<td>Subpart I</td>
<td></td>
<td></td>
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<tr>
<td>Approval based on animal studies</td>
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<td></td>
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<tr>
<td>Submitted in response to a PMR</td>
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<td></td>
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<td>Submitted in response to a PMC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Submitted in response to a Pediatric Written Request</td>
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<td></td>
</tr>
</tbody>
</table>

**Comments:** Currently the product is on the FDA compliance list of Medically Necessary Unapproved Marketed Drugs.

### BLAs only

- Ensure **RMS-BLA Product Information Sheet for TBP** and **RMS-BLA Facility Information Sheet for TBP** have been completed and forwarded to OPI/OBI/DRM (Vicky Carter)
  - Yes, dates

### BLAs only: Is the product subject to official FDA lot release per 21 CFR 610.2 (approvals only)

- Yes | No

### Public communications (approvals only)

- Office of Executive Programs (OEP) liaison has been notified of action
  - Yes | No
- Press Office notified of action (by OEP)
  - Yes | No

- Indicate what types (if any) of information dissemination are anticipated
  - None
  - HHS Press Release
  - FDA Talk Paper
  - CDER Q&As
  - Other

---

2 Answer all questions in all sections in relation to the pending application, i.e., if the pending application is an NDA or BLA supplement, then the questions should be answered in relation to that supplement, not in relation to the original NDA or BLA. For example, if the application is a pending BLA supplement, then a new **RMS-BLA Product Information Sheet for TBP** must be completed.
## Exclusivity

- **Is approval of this application blocked by any type of exclusivity?**
  - No ☒ Yes ☐

  - **NDAs and BLAs:** Is there existing orphan drug exclusivity for the “same” drug or biologic for the proposed indication(s)? Refer to 21 CFR 316.3(b)(13) for the definition of “same drug” for an orphan drug (i.e., active moiety). This definition is NOT the same as that used for NDA chemical classification.
  - No ☒ Yes ☐
    - If, yes, NDA/BLA # and date exclusivity expires:

  - **(b)(2) NDAs only:** Is there remaining 5-year exclusivity that would bar effective approval of a 505(b)(2) application? *(Note that, even if exclusivity remains, the application may be tentatively approved if it is otherwise ready for approval.)*
  - No ☒ Yes ☐
    - If yes, NDA # and date exclusivity expires:

  - **(b)(2) NDAs only:** Is there remaining 3-year exclusivity that would bar effective approval of a 505(b)(2) application? *(Note that, even if exclusivity remains, the application may be tentatively approved if it is otherwise ready for approval.)*
  - No ☒ Yes ☐
    - If yes, NDA # and date exclusivity expires:

  - **(b)(2) NDAs only:** Is there remaining 6-month pediatric exclusivity that would bar effective approval of a 505(b)(2) application? *(Note that, even if exclusivity remains, the application may be tentatively approved if it is otherwise ready for approval.)*
  - No ☒ Yes ☐
    - If yes, NDA # and date exclusivity expires:

  - **NDAs only:** Is this a single enantiomer that falls under the 10-year approval limitation of 505(u)? *(Note that, even if the 10-year approval limitation period has not expired, the application may be tentatively approved if it is otherwise ready for approval.)*
  - No ☒ Yes ☐
    - If yes, NDA # and date 10-year limitation expires:

## Patent Information (NDAs only)

- **Patent Information:** Verify that form FDA-3542a was submitted for patents that claim the drug for which approval is sought. If the drug is an old antibiotic, skip the Patent Certification questions.
  - Verified ☒ Not applicable because drug is an old antibiotic.

  - **Patent Certification [505(b)(2) applications]:** Verify that a certification was submitted for each patent for the listed drug(s) in the Orange Book and identify the type of certification submitted for each patent.
  - 21 CFR 314.50(i)(1)(i)(A) ☒
  - 21 CFR 314.50(i)(1)(ii) ☐ (iii)

- **[505(b)(2) applications] If the application includes a paragraph III certification,** it cannot be approved until the date that the patent to which the certification pertains expires (but may be tentatively approved if it is otherwise ready for approval).
  - No paragraph III certification ☒
    - Date patent will expire

- **[505(b)(2) applications] For each paragraph IV certification,** verify that the applicant notified the NDA holder and patent owner(s) of its certification that the patent(s) is invalid, unenforceable, or will not be infringed (review documentation of notification by applicant and documentation of receipt of notice by patent owner and NDA holder). *(If the application does not include any paragraph IV certifications, mark “N/A” and skip to the next section below (Summary Reviews)).*
  - N/A (no paragraph IV certification) ☒
    - Verified

Version: 6/8/10
[505(b)(2) applications] For each paragraph IV certification, based on the questions below, determine whether a 30-month stay of approval is in effect due to patent infringement litigation.

Answer the following questions for each paragraph IV certification:

(1) Have 45 days passed since the patent owner’s receipt of the applicant’s notice of certification?

(Note: The date that the patent owner received the applicant’s notice of certification can be determined by checking the application. The applicant is required to amend its 505(b)(2) application to include documentation of this date (e.g., copy of return receipt or letter from recipient acknowledging its receipt of the notice) (see 21 CFR 314.52(e)).)

If “Yes,” skip to question (4) below. If “No,” continue with question (2).

(2) Has the patent owner (or NDA holder, if it is an exclusive patent licensee) submitted a written waiver of its right to file a legal action for patent infringement after receiving the applicant’s notice of certification, as provided for by 21 CFR 314.107(f)(3)?

If “Yes,” there is no stay of approval based on this certification. Analyze the next paragraph IV certification in the application, if any. If there are no other paragraph IV certifications, skip the rest of the patent questions.

If “No,” continue with question (3).

(3) Has the patent owner, its representative, or the exclusive patent licensee filed a lawsuit for patent infringement against the applicant?

(Note: This can be determined by confirming whether the Division has received a written notice from the (b)(2) applicant (or the patent owner or its representative) stating that a legal action was filed within 45 days of receipt of its notice of certification. The applicant is required to notify the Division in writing whenever an action has been filed within this 45-day period (see 21 CFR 314.107(f)(2))).

If “No,” the patent owner (or NDA holder, if it is an exclusive patent licensee) has until the expiration of the 45-day period described in question (1) to waive its right to bring a patent infringement action or to bring such an action. After the 45-day period expires, continue with question (4) below.

(4) Did the patent owner (or NDA holder, if it is an exclusive patent licensee) submit a written waiver of its right to file a legal action for patent infringement within the 45-day period described in question (1), as provided for by 21 CFR 314.107(f)(3)?

If “Yes,” there is no stay of approval based on this certification. Analyze the next paragraph IV certification in the application, if any. If there are no other paragraph IV certifications, skip to the next section below (Summary Reviews).

If “No,” continue with question (5).
(5) Did the patent owner, its representative, or the exclusive patent licensee bring suit against the (b)(2) applicant for patent infringement within 45 days of the patent owner’s receipt of the applicant’s notice of certification?

(Note: This can be determined by confirming whether the Division has received a written notice from the (b)(2) applicant (or the patent owner or its representative) stating that a legal action was filed within 45 days of receipt of its notice of certification. The applicant is required to notify the Division in writing whenever an action has been filed within this 45-day period (see 21 CFR 314.107(f)(2)). If no written notice appears in the NDA file, confirm with the applicant whether a lawsuit was commenced within the 45-day period).

If “No,” there is no stay of approval based on this certification. Analyze the next paragraph IV certification in the application, if any. If there are no other paragraph IV certifications, skip to the next section below (Summary Reviews).

If “Yes,” a stay of approval may be in effect. To determine if a 30-month stay is in effect, consult with the OND ADRA and attach a summary of the response.

---

**CONTENTS OF ACTION PACKAGE**

- **Copy of this Action Package Checklist**
  - included

  **Officer/Employee List**
  - List of officers/employees who participated in the decision to approve this application and consented to be identified on this list (approvals only)
    - Included
  - Documentation of consent/non-consent by officers/employees
    - Included

- **Action Letters**
  - Copies of all action letters (including approval letter with final labeling)
    - Action(s) and date(s) included

- **Labeling**
  - Package Insert (write submission/communication date at upper right of first page of PI)
    - Most recent draft labeling. If it is division-proposed labeling, it should be in track-changes format.
    - Approved PI attached to approval letter
    - November 24, 2009
    - Original applicant-proposed labeling
    - none
    - Example of class labeling, if applicable

---

3 Fill in blanks with dates of reviews, letters, etc.

Version: 6/8/10
- Medication Guide/Patient Package Insert/Instructions for Use (write submission/communication date at upper right of first page of each piece)

- Most-recent draft labeling. If it is division-proposed labeling, it should be in track-changes format.

- Original applicant-proposed labeling

- Example of class labeling, if applicable

- Labels (full color carton and immediate-container labels) (write submission/communication date on upper right of first page of each submission)

- Most-recent draft labeling

- Proprietary Name
  - Acceptability/non-acceptability letter(s) (indicate date(s))
  - Review(s) (indicate date(s))

- Labeling reviews (indicate dates of reviews and meetings)

<table>
<thead>
<tr>
<th>Administrative / Regulatory Documents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Administrative Reviews (e.g., RPM Filing Review/Memo of Filing Meeting) (indicate date of each review)</td>
</tr>
<tr>
<td>505(b)(2) Assessment (indicate date)</td>
</tr>
<tr>
<td>NDAs only: Exclusivity Summary (signed by Division Director)</td>
</tr>
<tr>
<td>Application Integrity Policy (AIP) Status and Related Documents [<a href="http://www.fda.gov/ICECI/EnforcementActions/ApplicationIntegrityPolicy/default.htm">http://www.fda.gov/ICECI/EnforcementActions/ApplicationIntegrityPolicy/default.htm</a>]</td>
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<tr>
<td>This application is on the AIP</td>
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<tr>
<td>â Yes â No</td>
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<tr>
<td>If yes, Center Director’s Exception for Review memo (indicate date)</td>
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<tr>
<td>If yes, OC clearance for approval (indicate date of clearance communication)</td>
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<tr>
<td>Pediatrics (approvals only)</td>
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<tr>
<td>Date reviewed by PeRC March 24, 2010</td>
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<tr>
<td>If PeRC review not necessary, explain:</td>
</tr>
<tr>
<td>Pediatric Page (approvals only, must be reviewed by PERC before finalized)</td>
</tr>
<tr>
<td>Debarment certification (original applications only): verified that qualifying language was not used in certification and that certifications from foreign applicants are cosigned by U.S. agent (include certification)</td>
</tr>
<tr>
<td>Outgoing communications (letters (except action letters), emails, faxes, telecons)</td>
</tr>
<tr>
<td>Internal memoranda, telecons, etc.</td>
</tr>
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</table>

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4 Filing reviews for scientific disciplines should be filed behind the respective discipline tab.
## Minutes of Meetings

<table>
<thead>
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<th>Details</th>
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</thead>
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<td>N/A or no mtg</td>
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<tr>
<td>Pre-NDA/BLA meeting (indicate date of mtg)</td>
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<tr>
<td>EOP2 meeting (indicate date of mtg)</td>
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<tr>
<td>Other milestone meetings (e.g., EOP2a, CMC pilots) (indicate dates of mtgs)</td>
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## Advisory Committee Meeting(s)

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<td></td>
<td>No AC meeting</td>
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</table>

### Decisional and Summary Memos

- **Office Director Decisional Memo (indicate date for each review)**: None
- **Division Director Summary Review (indicate date for each review)**: None; June 21, 2010
- **Cross-Discipline Team Leader Review (indicate date for each review)**: None; June 22, 2010
- **PMR/PMC Development Templates (indicate total number)**: None

### Clinical Information

#### Clinical Reviews

- **Clinical Team Leader Review(s) (indicate date for each review)**
- **Clinical review(s) (indicate date for each review)**: June 21, 2010
- **Social scientist review(s) (if OTC drug) (indicate date for each review)**: None

#### Financial Disclosure reviews(s) or location/date if addressed in another review

- **Financial Disclosure review(s)** or location/date if addressed in another review
  - included in clinical review
  - If no financial disclosure information was required, check here □ and include a review/memo explaining why not (indicate date of review/memo)

#### Clinical reviews from immunology and other clinical areas/divisions/centers (indicate date of each review)

- **None**

#### Controlled Substance Staff review(s) and Scheduling Recommendation (indicate date of each review)

- **None**

#### Risk Management

- **REMS Documents and Supporting Statement (indicate date(s) of submission(s))**
- **REMS Memo(s) and letter(s) (indicate date(s))**
- **Risk management review(s) and recommendations (including those by OSE and CSS) (indicate date of each review and indicate location/date if incorporated into another review)**: None

#### DSI Clinical Inspection Review Summary(ies) (include copies of DSI letters to investigators)

- **None requested** included

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5 Filing reviews should be filed with the discipline reviews.

Version: 6/8/10
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<tr>
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<td>Statistical Team Leader Review(s) <em>(indicate date for each review)</em></td>
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<td>Statistical Review(s) <em>(indicate date for each review)</em></td>
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<tr>
<td>DSI Clinical Pharmacology Inspection Review Summary <em>(include copies of DSI letters)</em></td>
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<td>• ADP/T Review(s) <em>(indicate date for each review)</em></td>
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<td>• Supervisory Review(s) <em>(indicate date for each review)</em></td>
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<td>Statistical review(s) of carcinogenicity studies <em>(indicate date for each review)</em></td>
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<tr>
<td>ECAC/CAC report/memo of meeting</td>
<td>None Included in P/T review, page</td>
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<tr>
<td>DSI Nonclinical Inspection Review Summary <em>(include copies of DSI letters)</em></td>
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</table>

<table>
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<td>• Branch Chief/Team Leader Review(s) <em>(indicate date for each review)</em></td>
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<td>• Product quality review(s) including ONDQA biopharmaceutics reviews <em>(indicate date for each review)</em></td>
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<td>Microbiology Reviews</td>
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<td>• NDAs: Microbiology reviews (sterility &amp; pyrogenicity) <em>(OPS/NDMS)</em> <em>(indicate date of each review)</em></td>
<td>Not needed May 10, 2010</td>
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<tr>
<td>• BLAs: Sterility assurance, microbiology, facilities reviews <em>(DMPQ/MAPCB/BMT)</em> <em>(indicate date of each review)</em></td>
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<td>Reviews by other disciplines/divisions/Centers requested by CMC/quality reviewer <em>(indicate date of each review)</em></td>
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<tr>
<td>Environmental Assessment (check one) (original and supplemental applications)</td>
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<tr>
<td>Categorical Exclusion <em>(indicate review date)</em> <em>(all original applications and all efficacy supplements that could increase the patient population)</em></td>
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<tr>
<td>Review &amp; FONSI <em>(indicate date of review)</em></td>
<td>May 14, 2010</td>
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<tr>
<td>Review &amp; Environmental Impact Statement <em>(indicate date of each review)</em></td>
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</tr>
</tbody>
</table>

| Facilities Review/Inspection | |
| NDAs: Facilities inspections *(include EER printout)* *(date completed must be within 2 years of action date)* *(only original NDAs and supplements that include a new facility or a change that affects the manufacturing sites)* | Date completed: June 16, 2010 |
| □ | Acceptable |
| □ | Withhold recommendation |
| □ | Not applicable |
| BLAs: TB-EER *(date of most recent TB-EER must be within 30 days of action date)* *(original and supplemental BLAs)* | Date completed: N/A |
| □ | Acceptable |
| □ | Withhold recommendation |

| NDAs: Methods Validation *(check box only, do not include documents)* | |
| □ | Completed |
| □ | Requested |
| □ | Not yet requested |
| □ | Not needed (per review) |

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6 I.e., a new facility or a change in the facility, or a change in the manufacturing process in a way that impacts the Quality Management Systems of the facility.
Appendix to Action Package Checklist

An NDA or NDA supplemental application is likely to be a 505(b)(2) application if:

1. It relies on published literature to meet any of the approval requirements, and the applicant does not have a written right of reference to the underlying data. If published literature is cited in the NDA but is not necessary for approval, the inclusion of such literature will not, in itself, make the application a 505(b)(2) application.

2. Or it relies for approval on the Agency's previous findings of safety and efficacy for a listed drug product and the applicant does not own or have right to reference the data supporting that approval.

3. Or it relies on what is "generally known" or "scientifically accepted" about a class of products to support the safety or effectiveness of the particular drug for which the applicant is seeking approval. (Note, however, that this does not mean any reference to general information or knowledge (e.g., about disease etiology, support for particular endpoints, methods of analysis) causes the application to be a 505(b)(2) application.)

Types of products for which 505(b)(2) applications are likely to be submitted include: fixed-dose combination drug products (e.g., heart drug and diuretic (hydrochlorothiazide) combinations); OTC monograph deviations (see 21 CFR 330.11); new dosage forms; new indications; and, new salts.

An efficacy supplement can be either a (b)(1) or a (b)(2) regardless of whether the original NDA was a (b)(1) or a (b)(2).

An efficacy supplement is a 505(b)(1) supplement if the supplement contains all of the information needed to support the approval of the change proposed in the supplement. For example, if the supplemental application is for a new indication, the supplement is a 505(b)(1) if:

1. The applicant has conducted its own studies to support the new indication (or otherwise owns or has right of reference to the data/studies).

2. And no additional information beyond what is included in the supplement or was embodied in the finding of safety and effectiveness for the original application or previously approved supplements is needed to support the change. For example, this would likely be the case with respect to safety considerations if the dose(s) was/were the same as (or lower than) the original application.

3. And all other “criteria” are met (e.g., the applicant owns or has right of reference to the data relied upon for approval of the supplement, the application does not rely for approval on published literature based on data to which the applicant does not have a right of reference).

An efficacy supplement is a 505(b)(2) supplement if:

1. Approval of the change proposed in the supplemental application would require data beyond that needed to support our previous finding of safety and efficacy in the approval of the original application (or earlier supplement), and the applicant has not conducted all of its own studies for approval of the change, or obtained a right to reference studies it does not own. For example, if the change were for a new indication AND a higher dose, we would likely require clinical efficacy data and preclinical safety data to approve the higher dose. If the applicant provided the effectiveness data, but had to rely on a different listed drug, or a new aspect of a previously cited listed drug, to support the safety of the new dose, the supplement would be a 505(b)(2).

2. Or the applicant relies for approval of the supplement on published literature that is based on data that the applicant does not own or have a right to reference. If published literature is cited in the supplement but is not necessary for approval, the inclusion of such literature will not, in itself, make the supplement a 505(b)(2) supplement.

3. Or the applicant is relying upon any data they do not own or to which they do not have right of reference.

If you have questions about whether an application is a 505(b)(1) or 505(b)(2) application, consult with your ODE’s ADRA.
<table>
<thead>
<tr>
<th>Application Type/Number</th>
<th>Submission Type/Number</th>
<th>Submitter Name</th>
<th>Product Name</th>
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<tbody>
<tr>
<td>NDA-200890</td>
<td>ORIG-1</td>
<td>ALCON INC</td>
<td>PILOCARPINE HYDROCHLORIDE OPTHALMIC SOLUTION, 1%, 2% AND 4%</td>
</tr>
</tbody>
</table>

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

/s/

LORI M GORSKI
06/25/2010
action package checklist
Memorandum

***Pre-Decisional Agency Information ***

Date: June 14, 2010

To: Lori Gorski, Project Manager
   Division of Anti-Infective and Ophthalmology Products

From: Beth Carr, Pharm.D., Regulatory Review Officer
      Sheila Ryan, Pharm.D., Group Leader
      Division of Drug Marketing, Advertising, and Communications
      (DDMAC)

Subject: Isopto Carpine (pilocarpine hydrochloride ophthalmic solution) 1%, 2%, and 4%
         NDA 200890

DDMAC has reviewed the proposed product labeling for Isopto Carpine (pilocarpine hydrochloride
ophthalmic solution) 1%, 2%, and 4%, dated 6/7/2010, and we offer the following comments. Please
feel free to contact me at (301) 796-3674 with any questions or clarifications.

4 page(s) of Draft Carton and Container Labels have been Withheld in Full immediately following this page as B4 (CCI/TS
<table>
<thead>
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<td>PILOCARPINE HYDROCHLORIDE OPHTHALMIC SOLUTION, 1%, 2% AND 4%</td>
</tr>
</tbody>
</table>

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

/s/

BETH M CARR
06/14/2010
**REQUEST FOR DDMAC LABELING REVIEW CONSULTATION**

**Please send immediately following the Filing/Planning meeting**

TO: Wayne Amchin  
DDMAC Regulatory Project Manager

FROM: Lori Gorski  
Project Manager  
Division of Anti-Infective & Ophthalmology Products  
Phone 301-796-0722  
E-mail lori.gorski@fda.hhs.gov

REQUEST DATE: June 7, 2010  
IND NO.: NDA 200890  
TYPE OF DOCUMENTS: Original NDA

NAME OF DRUG: Isopto Carpine (pilocarpine hydrochloride ophthalmic solution) 1%, 2% and 4%

PRIORITY CONSIDERATION: Priority

CLASSIFICATION OF DRUG:  

DESIRED COMPLETION DATE: One week from your receipt or sooner

NAME OF FIRM: Alcon  
PDUFA Date: June 22, 2020

**TYPE OF LABEL TO REVIEW**

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<td>[ ] INITIAL PROPOSED LABELING</td>
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</table>

EDR link to submission:  
The network location is: `\CDSESUB1\EVSPROD\NDA200890\200890.ENX`

Please Note: There is no need to send labeling at this time. DDMAC reviews substantially complete labeling, which has already been marked up by the CDER Review Team. The DDMAC reviewer will contact you at a later date to obtain the substantially complete labeling for review.

COMMENTS/SPECIAL INSTRUCTIONS:  
Attached is the Division's revised label for Pilocarpine. The PDUFA date is June 22. Please let me know when you can respond with comments to the Division. Thanks – Lori Gorski 796-0722

Labeling Meetings: We may have one more labeling meeting if needed, there is not currently one scheduled.  
Wrap-Up Meeting: May 14, 2010

SIGNATURE OF REQUESTER: Lori Marie Gorski

SIGNATURE OF RECEIVER

METHOD OF DELIVERY (Check one)  
[ ] eMAIL  
[ ] HAND
<table>
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/s/

LORI M GORSKI
06/07/2010
DDMAC consult of divisions revised label for an original NDA. Label sent via email.
NDA 200,890

Alcon Research Ltd.
Attention: Michael C. Son, Ph.D, RAC
Senior Manager, Regulatory Affairs
6201 South Freeway, R3-52
Fort Worth, TX 76134-2099

Dear Dr. Son:

Please refer to your New Drug Application (NDA) submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Isopto® Carpine (pilocarpine hydrochloride ophthalmic solution), 1%, 2%, and 4%.

We refer to our FDA Information Request letter dated May 7, 2010, draft responses from Michael Son, Alcon Research Ltd. sent on May 12, 2010, and teleconferences between Michael Son, Alcon Research Ltd, and Jeannie David, FDA, on May 14, 2010 and May 20, 2010. We request a prompt written response in order to continue our evaluation of your NDA.

Please provide as soon as possible all available quantitative data on pilocarpine hydrochloride drug substance in lots of (b)(4).

To facilitate prompt review of your response, please also provide an electronic courtesy copy of your response to both Jeannie David, Regulatory Project Manager in the Office of New Drug Quality Assessment (Jeannie.David@fda.hhs.gov), and Lori Gorski, Regulatory Project Manager the Office of New Drugs (Lori.Gorski@fda.hhs.gov).

If you have any questions regarding this letter, call Jeannie David, Regulatory Project Manager, at (301) 796-4247.

Sincerely,

Stephen P. Miller, Ph.D.
Acting Chief, Branch V
Division of New Drug Quality Assessment II
Office of New Drug Quality Assessment
Center for Drug Evaluation and Research
<table>
<thead>
<tr>
<th>Application Type/Number</th>
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</table>

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/s/

STEPHEN P MILLER
05/25/2010
MEMORANDUM OF MEETING MINUTES

MEETING DATES: May 14, 2010 and May 20, 2010
TIME: multiple
APPLICATION: NDA 200,890
DRUG NAME: pilocarpine hydrochloride ophthalmic solution, 1%, 2%, and 4%
SPONSOR: Alcon Research Ltd.
TYPE OF MEETING: Impromptu teleconferences with applicant
PHONE NUMBER CALLED: Michael Son, Ph.D., RAC

FDA PARTICIPANTS:

Office of New Drug Quality Assessment (ONDQA)
Rao Kambhampati, Ph.D., Review Chemist (May 14, 2010, teleconference only)
Jeannie David, M.S., Regulatory Project Manager

EXTERNAL PARTICIPANTS:

Alcon Research Ltd.
Michael Son, Ph.D., RAC, Senior Manager, Regulatory Affairs

BACKGROUND:

FDA issued an Information Request letter dated May 7, 2010, regarding chemistry review of the NDA. Alcon provided a draft response by email to Jeannie David, FDA on May 12, 2010 (attached). The following teleconference discussions took place.

POINTS DISCUSSED:

May 14, 2010, and May 20, 2010, discussions of Alcon’s May 12, 2010, email draft response:

- For point 2, Alcon indicated that they were having difficulty in getting additional quantitative data on the from the drug substance manufacturer. Alcon seemed willing to support a limit of but indicated that was unwilling to agree. Alcon stated that had informed them that only of their lots will meet . Alcon stated that they had only limited information in hand on exact; most information is in Certificates of Analyses that report only NMT .
  
  FDA requested that Alcon provide as much batch analysis data as they have available on .

- For point 3, Alcon requested if for drug substance and for drug product would be acceptable. FDA agreed.

- For point 4, Alcon indicated that they agree with NMT for unknown unspecified impurities, and requested if NMT would be ok for unknown specified impurities. FDA indicated this was acceptable, and asked that Alcon attempt to identify these impurities in the future. FDA requested that Alcon submit revised drug substance and drug product specification tables as requested in the May 7, 2010, letter. Alcon agreed to add this to their May 12, 2010 draft response.

Note, only Alcon’s May 12, 2010, email and Isopto Carpine Draft Partial CMC Responses (May 2010).pdf are attached below. The files: 32s44-batch-analyses.pdf, 70198F Chromatograms.pdf, 85917F Chromatograms.pdf are not attached.
Dear Ms. David,

As we discussed earlier today, I am sending you the partial responses to the ISOPTO Carpine NDA (CMC comments received on May 7, 2010). Please note that the draft responses being provided have not been reviewed by all involved functions at Alcon; therefore, the responses may not be final. Specifically, the responses to comments #2, #3 and #7 are still being reviewed and finalized.

Once we hear back from you, we will submit the complete responses electronically through the FDA gateway. In the meantime, please let me know if you have any further questions. Thank you.

Regards,
Michael C. Son, Ph.D., RAC
Senior Manager, Regulatory Affairs
Alcon Laboratories, Inc.
Michael.Son@AlconLabs.com
(817) 551-8120

Dear Michael,

Attached is an electronic courtesy copy of a CMC Information Request letter issued today, May 7, 2010.

We request a response by May 12, preferably early in the day. An email copy of the response you will be submitting to the NDA will be sufficient to meet that timeline. After review of the letter with your team, please notify me if it will be feasible to meet this date.

Please contact me for any questions regarding this letter.

Best regards,
Jeannie

Jeannie David, M.S.
Regulatory Project Manager
ISSUE 1

We noticed that the Certificates for Analysis that were provided in the original NDA submission for pilocarpine hydrochloride batches contained numerical percentages for the following related substances (impurities): any other impurities (individually not identified in Ph. Eur. monograph), and sum of other impurities. Therefore, we recommend that you resubmit the Batch Analyses including the actually observed values.

RESPONSE

The Ph. Eur. Related substances test may be interpreted as a limit test, but in consideration of the Agency’s request, Alcon has calculated numeric values for the batches listed in Table 3.2.S.4.4-1 of Section 3.2.S.4.4. Please also note that Alcon does not currently have a specification for “Any Unspecified Impurity Not Identified in the Ph. Eur. Monograph”, therefore those values were obtained from . Upon approval, Alcon commits to adopt the proposed specifications for drug substance listed in Section 3.2.S.4.1.

Table 3.2.S.4.4-1 of Section 3.2.S.4.4 has been revised to include actually observed values for the following related substances (impurities): , any other impurities (individually not identified in Ph. Eur. monograph), and sum of other impurities.

Revisions to Module 3:

Section 3.2.S.4.4
ISSUE 2

Please tighten the acceptance criterion for content in the drug substance specification to be consistent with the observed range based on the batch analyses currently submitted to the NDA).

RESPONSE

Alcon accepts the content result from the, and the specification is based on experience with the manufacturing process. While the drug substance batches submitted in the NDA would, in fact, meet the tighter limits suggested, other batches produced in the same time-frame would not. Considering that the proposed specification of NMT the ICH limit of 3000 ppm for this Class 2 solvent, the safety margin provided by the current specification seems adequate.

NOTE: Discussions with the drug substance manufacturer, are currently ongoing.
ISSUE 3

With regard to the acceptance criterion in the drug product specification, you stated that the drug substance criterion for content is NMT of Active. However, we noticed that the drug substance Certificates of Analysis do not contain an acceptance criterion for content. Also, the batch analysis and stability data that were provided in the NDA for the drug product batches contained at release and at all the stability test points. Therefore, please tighten the acceptance criterion for content in the drug product specification to NMT.

RESPONSE

The Ph. Eur. Monograph specifies . Therefore, the Ph. Eur. specification of NMT is applicable, since is a specified impurity other than Since is grouped into a sum of “other impurities” for the drug substance, a separate specification for would also need to be included in the drug substance. This requires approval from the drug substance manufacturer.

NOTE: Discussions with the drug substance manufacturer, are currently ongoing.
**ISSUE 4**

With regard to your response to Issue 3(c) in the Amendment dated April 15, 2010, regarding the acceptance criterion for Any Individual Unspecified Impurity content, we recommend the following changes: Revise the acceptance criteria for Any Individual Unknown Unspecified Impurity to NMT \( (b) (4) \) and add a specified limit for the unknown impurity that is being observed, “Unknown Impurity RRT=0.XY” with an acceptance criterion of NMT \( (b) (4) \).

**RESPONSE**

Alcon agrees to revise the acceptance criteria for Any Individual Unknown Unspecified Impurity to NMT \( (b) (4) \) and add specified limits for the unknown impurities being observed.
**ISSUE 5**

Based on your response to Issue 4 in the Amendment dated April 15, 2010, we do not believe that an overage is appropriate because there does not appear to be significant loss of active during manufacture of the product. If you believe that an overage is important for any strength of pilocarpine hydrochloride ophthalmic solution, please provide the following information for each batch of the drug product: calculated drug substance weight (based on the assay value and targeting 100% in the drug product); actually added drug substance weight; overage of drug substance; expected assay value for the drug product; and actually observed assay value for the drug product.

**RESPONSE**

A review of the last 10 manufacturing batches of each concentration with the overage (including the calculated drug substance weight, expected assay value and observed assay value) was performed as shown in Tables 5-1 through 5-3. Based on this analysis, Alcon agrees that the 1% formulation does not require an overage. However, an overage of up to is warranted for the 2% and the 4% formulations.
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/s/

JEANNIE C DAVID
05/25/2010
Information Request

Alcon Research Ltd.
Attention: Michael C. Son, Ph.D, RAC
Senior Manager, Regulatory Affairs
6201 South Freeway, R3-52
Fort Worth, TX 76134-2099

Dear Dr. Son:

Please refer to your New Drug Application (NDA) submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Isopto® Carpine (pilocarpine hydrochloride ophthalmic solution), 1%, 2%, and 4%.

We are reviewing the Chemistry, Manufacturing and Controls sections of your submission and have the following comments and information requests. Please address the following CMC comments and recommendations that are related to your amendment dated April 15, 2010. We would appreciate your response by May 12, 2010.

1. We noticed that the Certificates for Analysis that were provided in the original NDA submission for pilocarpine hydrochloride batches contained numerical percentages for the following related substances (impurities): (b)(4), any other impurities (individually not identified in Ph.Eur. monograph), and sum of other impurities. Therefore, we recommend that you resubmit the Batch Analyses including the actually observed values.

2. Please tighten the acceptance criterion for (b)(4) content in the drug substance specification to be consistent with the observed range (b)(4) based on the batch analyses currently submitted to the NDA).

3. With regard to the (b)(4) acceptance criterion in the drug product specification, you stated that the drug substance criterion for (b)(4) content is NMT (b)(4) of Active. However, we noticed that the drug substance Certificates of Analysis do not contain an acceptance criterion for (b)(4) content. Also, the batch analysis and stability data that were provided in the NDA for the drug product batches contained (b)(4) at release and at all the stability test points. Therefore, please tighten the acceptance criterion for (b)(4) content in the drug product specification to NMT (b)(4)

4. With regard to your response to Issue 3(c) in the amendment dated April 15, 2010, regarding the acceptance criterion for Any Individual Unspecified Impurity content, we recommend the following changes: Revise the acceptance criteria for Any Individual Unknown Unspecified Impurity to NMT (b)(4) and add a specified limit for the unknown impurity that is being observed, “Unknown Impurity RRT= 0.XY” with an acceptance criterion of NMT (b)(4)
5. Based on your response to Issue 4 in the amendment dated April 15, 2010, we do not believe that an overage is appropriate because there does not appear to be significant loss of active during manufacture of the product. If you believe that an overage is important for any strength of pilocarpine hydrochloride ophthalmic solution, please provide the following information for each batch of the drug product: calculated drug substance weight (based on the assay value and targeting 100% in the drug product); actually added drug substance weight; overage of drug substance; expected assay value for the drug product; and actually observed assay value for the drug product.

6. Please provide representative HPLC chromatograms of the drug product stability batches #70198F and #85917F for the initial, 12 month, 24 month, and 36 month time point samples and for the resolution standard.

7. We recommend that for the next three years you include both accelerated conditions (40°C/25%RH) with testing points of 1, 2, 3, and 6 months and long-term conditions (25°C/40%RH) with testing points of 3, 6, 9, 18, 24 and 36 months in your yearly stability commitment. This will establish a baseline of accelerated data on three batches for future post-approval changes.

8. Since the NDA submission does not contain stability data for the drug product stored under refrigerated conditions and no freeze thaw cycling studies were performed on the drug product, please change the storage statement to “Store at 15º to 25ºC (59º to 77ºF) and protect from freezing.”

To facilitate prompt review of your response, please also provide an electronic courtesy copy of your response to both Jeannie David, Regulatory Project Manager in the Office of New Drug Quality Assessment (Jeannie.David@fda.hhs.gov), and Lori Gorski, Regulatory Project Manager the Office of New Drugs (Lori.Gorski@fda.hhs.gov).

If you have any questions regarding this letter, call Jeannie David, Regulatory Project Manager, at (301) 796-4247.

Sincerely,

(See appended electronic signature page)

Stephen P. Miller, Ph.D.
Acting Chief, Branch IV
Division of Pre-Marketing Assessment II
Office of New Drug Quality Assessment
Center for Drug Evaluation and Research
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/s/

STEPHEN P MILLER
05/07/2010
Hi Mike

There are 2 additional requests from the clinical reviewer of NDA 200890, pilocarpine. Please respond with an electronic submission through the gateway. Let me know if you have any questions.

1) Section 5.3 of the proposed label states, (b)(4) The annotated label cites the Salagen label and the submitted literature.

Can Alcon be more specific about the location of the supportive information for this statement?

2) The 120-day safety update should be submitted to the application.

Thanks.
Lori Gorski
Project Manager
Division of Anti-Infective & Ophthalmology Products
Phone 301-796-0722
Fax 301-796-9881
E-mail lori.gorski@fda.hhs.gov
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/s/

LORI M GORSKI
05/03/2010
clinical request for information
Hi Mike

I've had a request for clarification from the pharm tox reviewer. Please respond with a submission through the gateway.

Thanks and call me if you have any questions.

1. Please confirm that the drop size of ISOPTOCarpine is (b)(4)

2. Per NDA Section 2.7.2.3.2 Human PK Studies, please show the calculations used to derive this figure.

3. In the pregnancy and the nonclinical toxicology sections of the labeling, the multiples of animal dose to MROHD (maximum recommended ophthalmic human dose) were shown. Please show the calculations used to derive these figures.

Lori Gorski
Project Manager
Division of Anti-Infective & Ophthalmology Products
Phone 301-796-0722
Fax 301-796-9881
E-mail lori.gorski@fda.hhs.gov
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/s/

LORI M GORSKI
04/22/2010
pharm tox request
NDA 200890

Alcon Research, Ltd.
6201 South Freeway, R3-52
Fort Worth, Texas 76134

ATTENTION: Michael C. Son, Ph.D.
Senior Manager, Regulatory Affairs

Dear Dr. Son:

Please refer to your New Drug Application (NDA) dated December 22, 2009, received December 22, 2009, submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Pilocarpine Hydrochloride Ophthalmic Solution, 1%, 2% and 4%.

We also refer to your December 21, 2009, correspondence, received December 22, 2009, requesting review of your proposed proprietary name, Isopto Carpine. We have completed our review of the proposed proprietary name, Isopto Carpine and have concluded that it is acceptable for this product.

However, we note that you have developed a naming convention for your product line that uses the prefix “Isopto” and, in certain cases, contains the established name within the proprietary name. For future reference, we discourage the incorporation of established names in proprietary names and also the use of the prefix “Isopto” for your other products because it will contribute to name similarity in a drug class with many overlapping product characteristics.

The proposed proprietary name, Isopto Carpine, will be re-reviewed 90 days prior to the approval of the NDA. If we find the name unacceptable following the re-review, we will notify you.

If any of the proposed product characteristics as stated in your December 21, 2009, submission are altered prior to approval of the marketing application, the proprietary name should be resubmitted for review.
If you have any questions regarding the contents of this letter or any other aspects of the proprietary name review process, contact Brantley Dorch, Safety Regulatory Project Manager in the Office of Surveillance and Epidemiology, at (301) 796-0150. For any other information regarding this application contact the Office of New Drugs (OND) Regulatory Project Manager, Lori Gorski, at (301) 796-0722.

Sincerely,

{See appended electronic signature page}

Carol Holquist, RPh
Director
Division of Medication Error Prevention and Analysis
Office of Surveillance and Epidemiology
Center for Drug Evaluation and Research
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/s/

CAROL A HOLQUIST
03/22/2010
NDA 200,890

INFORMATION REQUEST

Alcon Research Ltd.
Attention: Michael C. Son, Ph.D, RAC
Senior Manager, Regulatory Affairs
6201 South Freeway, R3-52
Fort Worth, TX 76134-2099

Dear Dr. Son:

Please refer to your New Drug Application (NDA) submitted under section 505(b) of the Federal Food, Drug, and Cosmetic Act for Isopto® Carpine (pilocarpine hydrochloride ophthalmic solution), 1%, 2%, and 4%.

We are reviewing the Chemistry, Manufacturing and Controls sections of your submission and have the following comments and information requests. We request a prompt written response in order to continue our evaluation of your NDA. We would appreciate your response by April 16, 2010.

1. The Alcon’s Batch Analysis Tables 2.3.S.4-3 and 3.2.S.4.4-1 for the drug substance do not contain actually observed results for Related Substances (impurities) and ordinary impurities instead they were shown as NMT of certain percentage. Please provide actually observed results.

2. Please include microbial limits test in the NDA drug substance acceptance specification. Since this test is included in the drug substance specification, the test results from the Certificate of Analysis can be routinely used to accept the drug substance.

3. In the drug product specification please make the following changes:
   a. Tighten bacterial endotoxins acceptance criterion from to
   b. For 4% strength drug product, tighten Osmolality test acceptance criterion from to 550-600 mOsm/Kg.
   c. For the topical ophthalmic drug products we recommend an acceptance criterion of for Any Individual Unspecified Impurity content, therefore, please tighten the proposed acceptance criterion for Any Individual (Single) Unspecified Impurity content as much as possible.
   d. On the basis of the release and stability study results, we recommend that you tighten the following:
      i. content from NMT of Active to NMT of Active.
      ii. content from NMT of Active to NMT of Active.
iii. Total Impurities content from NMT \( (h) (4) \) of Active to NMT \( (h) (4) \) of Active.

iv. Viscosity from \( (h) (4) \) to \( (h) (4) \)

4. We noticed that most of the decomposition of pilocarpine hydrochloride occurs during storage of the drug product, therefore, we do not recommend 3% overage of pilocarpine hydrochloride in the batch formula of the drug product.

5. Please provide \( (h) (4) \) test results information for a drug product batch that was stored under room temperature stability conditions through the proposed expiration dating period. If the data are not available, please initiate the study and provide the results as they become available.

6. During stability study, for the pilocarpic acid impurity content test, at the 12 month time point all lots contained \( (h) (4) \) except lot #70198F (1% strength) and lot #85917F (4% strength) which contained \( (h) (4) \) and \( (h) (4) \) respectively. Please provide an explanation.

7. During stability study, for the Total Impurities content test, at the 12 month time point all lots contained \( (h) (4) \) of Total Impurities except lot #70198F and lot #85917F which contained \( (h) (4) \) and \( (h) (4) \), respectively. Please provide an explanation.

8. The stability data included 12, 24, and 36 month time point testing results only. Please provide 3, 6, 9, 12, and 18 month test point results for registration batches (if available) or for supportive stability batches.

9. Provide accelerated and stress stability study results for the drug product. If the data are not available, please initiate the study and provide the data as they become available.

To facilitate prompt review of your response, please also provide an electronic courtesy copy of your response to both Jeannie David, Regulatory Project Manager in the Office of New Drug Quality Assessment (Jeannie.David@fda.hhs.gov), and Lori Gorski, Regulatory Project Manager the Office of New Drugs (Lori.Gorski@fda.hhs.gov).

If you have any questions regarding this letter, call Jeannie David, Regulatory Project Manager, at (301) 796-4247.

Sincerely,

{See appended electronic signature page}

Stephen P. Miller, Ph.D.
Acting Chief, Branch IV
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/s/

STEPHEN P MILLER

03/19/2010
Hi Mike

Below is a request for literature articles that have been referenced in the pilo application. Please submit this information to the NDA application.

If you have any questions please give me a call.

Thanks
Lori Gorski
301-796-0722

Source: Allingham


Source: Bartlett


# From Netland and Allen


Mazor Z, Ticho U, Rehany U, Rose L: Piloplex, a new long-acting pilocarpine polymer salt,


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From Roger and Lightman
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/s/

LORI M GORSKI
03/11/2010
request for lit articles
Dear Dr. Son:

Please refer to your new drug application (NDA) dated December 22, 2009, received December 22, 2009, submitted pursuant to section 505(b)(2) of the Federal Food, Drug, and Cosmetic Act for Isopto Carpine (pilocarpine hydrochloride ophthalmic solution) 1%, 2% and 4%.

We have completed our filing review and have determined that your application is sufficiently complete to permit a substantive review. Therefore, this application was considered filed 60 days after the date we received your application in accordance with 21 CFR 314.101(a). The review classification for this application is Priority. Therefore, the user fee goal date is June 22, 2010.

We are reviewing your application according to the processes described in the Guidance for Review Staff and Industry: Good Review Management Principles and Practices for PDUFA Products. Therefore, we have established internal review timelines as described in the guidance, which includes the timeframes for FDA internal milestone meetings (e.g., filing, planning, mid-cycle, team and wrap-up meetings). Please be aware that the timelines described in the guidance are flexible and subject to change based on workload and other potential review issues (e.g., submission of amendments). We will inform you of any necessary information requests or status updates following the milestone meetings or at other times, as needed, during the process. If major deficiencies are not identified during the review, we plan to communicate proposed labeling and, if necessary, any postmarketing requirement/commitment requests by June 1, 2010.

At this time, we are notifying you that, we have not identified any potential review issues. Please note that our filing review is only a preliminary evaluation of the application and is not indicative of deficiencies that may be identified during our review.

Under the Pediatric Research Equity Act (PREA) (21 U.S.C. 355c), all applications for new active ingredients, new indications, new dosage forms, new dosing regimens, or new routes of administration are required to contain an assessment of the safety and effectiveness of the product for the claimed indications in pediatric patients unless this requirement is waived, deferred, or inapplicable.
We acknowledge receipt of your request for a full waiver of pediatric studies for this application. Once we have reviewed your request and the application we will notify you of our decision.

If you have any questions, call Lori Gorski, Regulatory Project Manager, at (301) 796-0722.

Sincerely,

[See appended electronic signature page]

Wiley A. Chambers, M.D.
Acting Director
Division of Anti-Infective and Ophthalmology Products
Office of Antimicrobial Products
Center for Drug Evaluation and Research
Application Type/Number  Submission Type/Number  Submitter Name  Product Name
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/s/

WILEY A CHAMBERS
03/05/2010
NDA 200890

NDA ACKNOWLEDGMENT

Alcon Research Ltd.
Attention: Michael C. Son, Ph.D, RAC
Senior Manager, Regulatory Affairs
6201 South Freeway, R3-52
Fort Worth, TX 76134-2099

Dear Dr. Son:

We have received your new drug application (NDA) submitted pursuant to section 505(b)(2) of the Federal Food, Drug, and Cosmetic Act (FDCA) for the following:

Name of Drug Product: Isopto Carpine (pilocarpine hydrochloride ophthalmic solution) 1%, 2% and 4%

Date of Application: December 22, 2009

Date of Receipt: December 22, 2009

Our Reference Number: NDA 200890

Unless we notify you within 60 days of the receipt date that the application is not sufficiently complete to permit a substantive review, we will file the application on February 19, 2010, in accordance with 21 CFR 314.101(a).

If you have not already done so, promptly submit the content of labeling [21 CFR 314.50(l)(i)] in structured product labeling (SPL) format as described at http://www.fda.gov/ForIndustry/DataStandards/StructuredProductLabeling/default.htm. Failure to submit the content of labeling in SPL format may result in a refusal-to-file action under 21 CFR 314.101(d)(3). The content of labeling must conform to the content and format requirements of revised 21 CFR 201.56-57.

Please note that you are responsible for complying with the applicable provisions of sections 402(i) and 402(j) of the Public Health Service Act (PHS Act) (42 USC §§ 282(i) and (j)), which was amended by Title VIII of the Food and Drug Administration Amendments Act of 2007 (FDAAA) (Public Law No. 110-85, 121 Stat. 904). Title VIII of FDAAA amended the PHS Act by adding new section 402(j) (42 USC § 282(j)), which expanded the current database known as ClinicalTrials.gov to include mandatory registration and reporting of results for applicable
clinical trials of human drugs (including biological products) and devices. FDAAA requires that, at the time of submission of an application under section 505 of the FDCA, the application must be accompanied by a certification that all applicable requirements of 42 USC § 282(j) have been met. Where available, the certification must include the appropriate National Clinical Trial (NCT) control numbers. 42 USC 282(j)(5)(B). You did not include such certification when you submitted this application. You may use Form FDA 3674, Certification of Compliance, under 42 U.S.C. § 282(j)(5)(B), with Requirements of ClinicalTrials.gov Data Bank, to comply with the certification requirement. The form may be found at http://www.fda.gov/opacom/morechoices/fdaforms/default.html.


The NDA number provided above should be cited at the top of the first page of all submissions to this application. Send all submissions, electronic or paper, including those sent by overnight mail or courier, to the following address:

Food and Drug Administration  
Center for Drug Evaluation and Research  
Division of Anti-Infective and Ophthalmology Products  
5901-B Ammendale Road  
Beltsville, MD 20705-1266

All regulatory documents submitted in paper should be three-hole punched on the left side of the page and bound. The left margin should be at least three-fourths of an inch to assure text is not obscured in the fastened area. Standard paper size (8-1/2 by 11 inches) should be used; however, it may occasionally be necessary to use individual pages larger than standard paper size. Non-standard, large pages should be folded and mounted to allow the page to be opened for review without disassembling the jacket and refolded without damage when the volume is shelved. Shipping unbound documents may result in the loss of portions of the submission or an unnecessary delay in processing which could have an adverse impact on the review of the submission. For additional information, please see http://www.fda.gov/Drugs/DevelopmentApprovalProcess/FormsSubmissionRequirements/DrugMasterFil esDMFs/ucm073080.htm
If you have any questions, call Lori Gorski, Regulatory Project Manager, at (301) 796-0722.

Sincerely,

{See appended electronic signature page}

Maureen P. Dillon-Parker
Chief, Project Management Staff
Division of Anti-Infective and Ophthalmology Products
Office of Antimicrobial Products
Center for Drug Evaluation and Research
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/s/

MAUREEN P DILLON PARKER
01/29/2010
Hi Mike - We have the following comment regarding the pediatric section of the pilocarpine application. Please call me if you have any questions. Thanks.

We acknowledge receipt of your request for a full waiver of pediatric studies for this application. The request for a waiver from pediatric studies is inadequate and is therefore denied. There is sufficient information available in the public domain to support various pediatric indications for pilocarpine hydrochloride solution. Please revise the pediatric section of your application to reflect the intended and documented usage for this product.

Lori Gorski
Project Manager
Division of Anti-Infective & Ophthalmology Products
Phone 301-796-0722
Fax 301-796-9881
E-mail lori.gorski@fda.hhs.gov
<table>
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<tr>
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<th>Submission Type/Number</th>
<th>Submitter Name</th>
<th>Product Name</th>
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<tbody>
<tr>
<td>NDA-200890</td>
<td>ORIG-1</td>
<td>ALCON INC</td>
<td>PILOCARPINE HYDROCHLORIDE OPHTHALMIC SOLUTION, 1%, 2% AND 4%</td>
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/s/

LORI M GORSKI
01/28/2010
request update to pediatric section of NDA from waiver to complete
**TEAM MEETING AGENDA**  
January 25, 2010

**NDA**  
200890

**Drug**  
Isopto Carpine (pilocarpine hydrochloride ophthalmic solution) 1%, 2% and 4%

**Indication**  
The reduction of IOP in patients with open angle glaucoma or ocular hypertension, for acute angle-closure glaucoma, postoperative elevated IOP associated with laser surgery and Induction of miosis.

**Sponsor**  
Alcon

**Received**  
December 22, 2009

**Day 60:**  
February 19, 2010

**Day 74:**  
March 5, 2010

**First Reviews**  
May 7, 2010

**User Fee Date**  
June 22, 2010

<table>
<thead>
<tr>
<th>Primary Reviewer</th>
<th>Team Leader</th>
<th>Filable</th>
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<tr>
<td><strong>Project Manager</strong></td>
<td>Lori Gorski</td>
<td>pre-38</td>
<td>505b2</td>
</tr>
<tr>
<td></td>
<td>Maureen Dillon Parker</td>
<td>Medically Necessary Unapproved Marketed Drug</td>
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</table>

**Micro**  
Denise Miller  
Jim McVey

**Stats**  
Rima Izem  
Yan Wang

**Pharm/Tox**  
Conrad Chen  
Wendy Schmidt

**CMC**  
Rao Kambhampati  
Steve Miller

**Clin Pharm**  
Eric Zhang  
Chuck Bonapace

**Clinical**  
Bill Boyd  
No filing issues

**OSE**  
Brantley Dorch  
Proprietary name under review

**DSI**  
Jean Mulinde  
Kassa Ayalew  
Consult to be sent
Hi All

Just a recap of Mondays filing meeting for Alcon's pilocarpine literature NDA. This is a Priority review.

1. If you have any issues or requests to convey to Alcon please email them to me as soon as possible.

2. There were no filing issues. See reviewers filing reviews for specific information.

3. First review target date is May 7th and everyone agreed they could meet that date or sooner.

4. Rima and Bill will provide Lori with a request for additional literature to support the proposed indications.

5. Alcon has been advised they should change their pediatric section from 'complete waiver' of studies to 'completed' and provide available literature for labeling children. This application will go to PERC on March 24, 2009.

6. Lori will follow up with a consult to DSI and with DDMAC once the label has been drafted by the division.

If I’ve missed anything please let me know!

Thanks everyone.

Lori Gorski
Project Manager
Division of Anti-Infective & Ophthalmology Products
Phone 301-796-0722
Fax 301-796-9881
E-mail lori.gorski@fda.hhs.gov
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/s/
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LORI M GORSKI
01/29/2010
RPM filing review original NDA
Hi Mike

Attached is a request from the statistical reviewer for information for NDA 200890, Isopto Carpine. Please submit the response electronically.

We have received your study reports and data for clinical studies in pdf format. If possible, please send all the data submitted for these clinical studies in electronic format (.xpt extension) with documentation (define.pdf file). This will assist us in conducting our review and statistical analyses.

Thanks.

Lori Gorski
Project Manager
Division of Anti-Infective & Ophthalmology Products
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/s/

LORI M GORSKI
01/13/2010
stat request for info
Hi Mike - How are you?

I have a couple regulatory requirements for the pilocarpine NDA.

1. The application was submitted pursuant to section 505(b)2 of the Federal Food, Drug, and Cosmetic Act since the division's decision will rely on the literature articles that have been submitted. Please submit a new Form FDA 356h stating the application description as a 505(b)2.

2. It's a requirement that a Form FDA 3674 be included with every application submitted. If there is one in your NDA please provide the location where the form can be found. Otherwise please submit a Form FDA 3674.

Thanks Mike. If you have any question give me a call.

Lori Gorski
Project Manager
Division of Anti-Infective & Ophthalmology Products
Phone 301-796-0722
Fax 301-796-9881
E-mail lori.gorski@fda.hhs.gov
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/s/

LORI M GORSKI
01/11/2010
redequest for regulatory forms for original NDA