Complete this log if the subject experienced any clinical adverse events (including intercurrent events) since the last visit. Check the “None” box and instruct the subject to initial and date the source documentation box if the subject has not experienced any clinical adverse events since the last visit.

0 None

* Please complete a Serious Adverse Event Reporting (SERIOUS) form.
** Please complete the appropriate Change in Medications form.
*** Please complete the Concomitant Medications (CMED) form.

<table>
<thead>
<tr>
<th>DESCRIPTION OF ADVERSE EVENT (1000)</th>
<th>1. ICD9 CODE (1010)</th>
<th>2. DATE STARTED (Top Line) (1020)</th>
<th>3. DATE STOPPED (Bottom Line) (1030)</th>
<th>4. ONGOING at current visit (1040)</th>
<th>5. TYPE (1050)</th>
<th>6. SEVERITY (1060)</th>
<th>7. SERIOUS (1070)</th>
<th>8. LIKELIHOOD OF RELATIONSHIP TO STUDY DRUG(S) (1080)</th>
<th>9. CHANGE IN STUDY DRUG(S) (1090)</th>
<th>10. OUTCOME (Skip if #3 is missing) (1100)</th>
<th>11. TREATMENT REQUIRED (1110)</th>
<th>12. ONGOING at final visit (1120)</th>
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</tr>
</tbody>
</table>
Coordinator Instructions: Before administering this questionnaire, give the subject his or her diary cards from the two weeks immediately preceding the visit. The subject should refer to the diary cards when answering each question. The AEQ must be the last asthma questionnaire completed at a given visit.

Subject Instructions: Please consider your last two weeks of asthma control in answering these questions. Check the box next to the response that best describes your average weekly asthma symptoms, rescue use and nighttime awakenings in the past two weeks.

1. In the past two weeks, how often have you experienced asthma symptoms?
   - 0  Less than or equal to 2 days a week
   - 1  3 to 5 days per week
   - 2  6 or more days per week, but not continual
   - 3  Continual (multiple times every day) (1000)

2. In the past two weeks, how often have you used your rescue beta-agonist medicine (e.g., albuterol (Proventil, Ventolin)), aside from preventative use prior to exercise?
   - 0  Less than or equal to 2 days per week
   - 1  3 to 5 days per week
   - 2  6 days per week
   - 3  At least once per day (daily) (1010)

3. In the past two weeks, how often have you awakened at night due to asthma symptoms?
   - 0  No awakenings or awakened 1 night during the 2 weeks
   - 1  1 night per week
   - 2  2 or 3 nights per week
   - 3  4 or more nights per week (1020)
AM1® QUALITY CONTROL

(Technician completed)

1. Serial Number of AM1® being tested

2. Serial Number of turbine being tested

3. Test date

4. Is a new AM1® device being tested for this subject?
   If YES, indicate the primary reason.
   - 1 First issuing
   - 2 “Old” device failed QC testing
   - 3 “Old” device had display problems
   - 4 “Old” device experienced battery failure
   - 5 “Old” device was recalled
   - 6 “Old” device was lost
   - 7 Other

5. Trial 1 (1060/1070)

6. Trial 2 (1080/1090)

7. Trial 3 (1100/1110)

8. Trial 4 (1120/1130)

9. Trial 5 (1140/1150)

Clinic Use Only
Median Relative Bias __ __ __ __ __ %
Inter-quartile Range __ __ __ __ __ %

The Median Relative Bias is the third largest value of the 5 measures of relative bias. The Inter-quartile Range is determined by subtracting the relative bias of rank 2 from the relative bias of rank 4.

When a subject receives a new AM1® or turbine for the first time, the median relative bias must be between -15% and +15%, and the inter-quartile range must be less than 10%.

When a subject returns to the clinic with a used AM1®, (i) subtract the original median relative bias (the median relative bias when the AM1® or turbine was first dispensed) from the current median relative bias, and (ii) subtract the original inter-quartile range (the inter-quartile range when the AM1® or turbine was first dispensed) from the current inter-quartile range. The difference for (i) must be between -5% and +5% and the difference for (ii) must be less than +5% for the AM1® to be reissued to the subject.

10. Did the AM1® pass?
   - 1 Yes
   - 0 No

11. If NO, is this the second test with this turbine at this visit?
   - 1 Yes
   - 0 No

   ➔ If NO, retest the AM1® with the same turbine and complete another AM1® Quality Control form.
   ➔ If YES, issue a new turbine and complete another AM1® Quality Control form. If 2 turbines have been tested with this device, issue a new device and turbine and complete another AM1® Quality Control form.
ASTHMA HISTORY

1. Approximately how old were you when chest symptoms suggesting asthma first appeared? (Enter '00' if subject was under 1 year.)
   ■ 1 Yes  ■ 0 No  ■ 8 Don’t Recall
   ___ ___ years

2. How old were you when a doctor first diagnosed you with asthma?
   ___ ___ years

2a. What caused you to seek medical care?
   ■ 1 Acute Symptoms
   ■ 2 Chronic Symptoms
   ■ 3 Other

3. Have any of your immediate blood relatives been told by a physician that they have asthma? (Check the 'N/A' box if the subject does not have biological siblings or children.)
   3a. Mother
   ■ 1 Yes  ■ 0 No  ■ 8 Don’t Know
   3b. Father
   ■ 1 Yes  ■ 0 No  ■ 8 Don’t Know
   3c. Brother(s) or Sister(s)
   ■ 1 Yes  ■ 0 No  ■ 8 Don’t Know  ■ 9 N/A
   3d. Child(ren)
   ■ 1 Yes  ■ 0 No  ■ 8 Don’t Know  ■ 9 N/A

ASTHMA SYMPTOMS

4. On average, over the last 4 weeks, how often have you experienced each of the following asthma symptoms:

4a. Cough (deep chest)
   ■ 0 Never
   ■ 1 Less than or equal to twice a week
   ■ 2 More than twice a week but not daily
   ■ 3 Daily
   ■ 4 Continuously

4b. Sputum (phlegm or mucous from the lungs)
   ■ 0 Never
   ■ 1 Less than or equal to twice a week
   ■ 2 More than twice a week but not daily
   ■ 3 Daily
   ■ 4 Continuously
4c. Chest tightness (difficulty taking a deep breath, pressure in the chest)

- 0 Never
- 1 Less than or equal to twice a week
- 2 More than twice a week but not daily
- 3 Daily
- 4 Continuously (1100)

4d. Wheezing (whistling or musical sound in the chest)

- 0 Never
- 1 Less than or equal to twice a week
- 2 More than twice a week but not daily
- 3 Daily
- 4 Continuously (1110)

4e. Shortness of breath

- 0 Never
- 1 Less than or equal to twice a week
- 2 More than twice a week but not daily
- 3 Daily
- 4 Continuously (1120)

4f. Nighttime symptoms (waking due to asthma)

- 0 Never
- 1 One or two nights a month
- 2 More than two nights per month but at most one night a week
- 3 More than one night a week but not frequently
- 4 Frequently (1130)

5. How often do you use your rescue beta-agonist medicine (e.g., Albuterol (Proventil, Ventolin)) other than to pretreat prior to exercise?

- 0 Less than or equal to twice a week
- 1 Greater than twice a week but not daily
- 2 Daily but not four times a day
- 3 Greater than or equal to four times a day (1140)

6. How do you categorize your asthma symptoms throughout the course of the year?

- 1 Relatively the same all year
- 2 Vary by season(s) (1150)

If 'Vary by season(s)', do your asthma symptoms worsen during the...

6ai. Winter?

- 0 No

6a(ii). Spring?

- 0 No

6a(iii). Summer?

- 0 No

6a(iv). Fall?

- 0 No
7. In the last 12 months, how many... (Enter ‘00’ if none)

7a. Asthma episodes have you had that required emergency care or an unscheduled office visit? ____ ____ (1200)

7b. Hospitalizations have you had due to asthma? ____ ____ (1210)

7c. Courses of oral corticosteroid therapy (e.g., Prednisone) for asthma have you taken? ____ ____ (1220)

7d. Days of work, school or housework have you missed due to asthma? ____ ____ ____ days (1230)

→ If Question #7d > 0, complete Question #7di.

7di. In the past 3 months, how many days of work, school, or housework have you missed due to asthma? ____ ____ ____ days (1240)

ASTHMA TRIGGERS

8. Do any of the following currently provoke your asthma?

8a. Exercise/Sports [ ] Yes [ ] No [ ] Don’t Know (1250)

8b. Menstrual cycle
   (If subject is male or postmenopausal, check N/A) [ ] Yes [ ] No [ ] Don’t Know [ ] N/A (1260)

8c. Aspirin or non-steroidal anti-inflammatory drugs (e.g., Aleve, Motrin) [ ] Yes [ ] No [ ] Don’t Know (1270)

8d. Colds, upper respiratory infections, sinus infections [ ] Yes [ ] No [ ] Don’t Know (1280)

8e. Irritants (e.g., smoke, pollution, odors, perfumes, chemicals) [ ] Yes [ ] No [ ] Don’t Know (1290)

8f. Weather conditions (e.g., cold, humidity) [ ] Yes [ ] No [ ] Don’t Know (1300)

8g. Emotional stress [ ] Yes [ ] No [ ] Don’t Know (1310)

8h. Food additives/preservatives (e.g., MSG, etc.) [ ] Yes [ ] No [ ] Don’t Know (1320)

8i. Other (please specify)—————————————————————————— [ ] Yes [ ] No (1330)
ALLERGIES
9. Do you have allergies to pets, pollen, dust, etc.?
   - Yes ☐  - No ☐  - Don’t Know ☐

   If YES,
   
   9a. Do your allergies provoke your asthma?
       Yes ☐  - No ☐  - Don’t Know ☐
   
   9b. Were your allergies diagnosed by a doctor?
       Yes ☐  - No ☐  - (1360)
   
   9c. How do you categorize your allergies?
       Relatively the same all year ☐  
       Vary by season(s) ☐  

   ➔ If 'Vary by season(s)', do your allergies worsen during the...

   9ci. Winter?
       Yes ☐  - No ☐  - (1380)
   
   9cii. Spring?
       Yes ☐  - No ☐  - (1390)
   
   9ciii. Summer?
       Yes ☐  - No ☐  - (1400)
   
   9ciii. Fall?
       Yes ☐  - No ☐  - (1410)

10. Have you ever had eczema (i.e., prolonged itchy, scaly, weepy skin rash)?
    - Yes ☐  - No ☐  - Don’t Know ☐

   10a. If YES, was your eczema diagnosed by a doctor?
        Yes ☐  - No ☐  - (1420)

11. Have any of your immediate blood relatives been told by a physician that they have allergies/eczema/hay fever?
    (Check the 'N/A' box if the subject does not have biological siblings or children.)

   11a. Mother
        Yes ☐  - No ☐  - Don’t Know ☐
   
   11b. Father
        Yes ☐  - No ☐  - Don’t Know ☐
   
   11c. Brother(s) or Sister(s)
        Yes ☐  - No ☐  - Don’t Know ☐  
        N/A ☐
   
   11d. Child(ren)
        Yes ☐  - No ☐  - Don’t Know ☐
        N/A ☐

SMOKING HISTORY
12. Were you ever a smoker?
    - Yes ☐  - No ☐  - (1472)

   12a. If YES, how many years did you smoke?
        (total number of years)
        ____ ____ years (1474)

13. Did you grow up in a household where you were exposed to tobacco smoke?
    - Yes ☐  - No ☐  - (1480)

14. In an average week, approximately how many hours are you exposed to other people’s tobacco smoke in all environments?
    ____ ____ hours (1490)

Subject Source Documentation
Subject’s Initials:  ____  ____ (1500)
Date:  ____ / ____ / ____ ____ (1510)
CONCOMITANT MEDICATIONS FOR ASTHMA/ALLERGY AND ADVERSE EVENTS

(Calendar completed)

**Instructions:** Since signing the informed consent or last study visit, list all prescription and over-the-counter (OTC) concomitant medications used to treat asthma/allergy symptoms and adverse events. Do not list routine use of study drugs or rescue medications. Check the “None” box if the subject has not started taking any medications since signing the informed consent or last study visit. If the medication is not related to an adverse or laboratory event, leave the event number missing and check the “N/A” box. If the subject is still taking the medication at the end of the current visit, check the “ongoing at current visit” check box and leave the stop date missing. All ongoing medications should be reviewed at subsequent visits to document the stop date of a medication. At the last study visit or an early termination visit, review all ongoing medication and indicate a stop date or check the “ongoing at final visit” check box on the data collection form and update the medication data in the ACRN data entry application.

At the final study visit or early termination visit, forward all concomitant medications for asthma/allergy-related medications and adverse events forms to the DCC.

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<th>NAME OF MEDICATION (1000)</th>
<th>CODE (1010)</th>
<th>RELATED EVENT (1020)</th>
<th>DOSE (1030)</th>
<th>UNITS (1040)</th>
<th>FREQUENCY (1050)</th>
<th>ROUTE</th>
<th>START DATE (MM/DD/YYYY) (1060)</th>
<th>STOP DATE (MM/DD/YYYY) (1070)</th>
<th>ONGOING AT CURRENT VISIT (1080)</th>
<th>ONGOING AT FINAL VISIT (1090)</th>
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### Units and Frequency Codes for Concomitant Medications

#### Codes for Units

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<td>mcg (μg)</td>
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<tr>
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<td>ml</td>
</tr>
<tr>
<td>4</td>
<td>mg/ml</td>
</tr>
<tr>
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<td>mEq</td>
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<tr>
<td>6</td>
<td>g</td>
</tr>
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<td>U</td>
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<td>teaspoon</td>
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<td>9</td>
<td>patch</td>
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<tr>
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<td>puffs (oral inhalation)</td>
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<td>11</td>
<td>nasal spray</td>
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<tr>
<td>15</td>
<td>mm</td>
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<td>other</td>
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#### Codes for Frequency

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<td>QD 1 time a day</td>
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<tr>
<td>2</td>
<td>BID 2 times a day</td>
</tr>
<tr>
<td>3</td>
<td>TID 3 times a day</td>
</tr>
<tr>
<td>4</td>
<td>QID 4 times a day</td>
</tr>
<tr>
<td>5</td>
<td>q4h every 4 hours</td>
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<tr>
<td>6</td>
<td>q5h every 5 hours</td>
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<tr>
<td>7</td>
<td>q6h every 6 hours</td>
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<tr>
<td>8</td>
<td>q8h every 8 hours</td>
</tr>
<tr>
<td>9</td>
<td>q12h every 12 hours</td>
</tr>
<tr>
<td>10</td>
<td>q24h every 24 hours</td>
</tr>
<tr>
<td>11</td>
<td>hs every night at bedtime</td>
</tr>
<tr>
<td>12</td>
<td>PRN as required</td>
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<tr>
<td>13</td>
<td>qod every other day</td>
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<td>qw once a week</td>
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<td>biw 2 times per week</td>
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<td>tiw 3 times per week</td>
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<td>5 times per week</td>
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<td>every 5 days</td>
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<tr>
<td>19</td>
<td>once a month</td>
</tr>
<tr>
<td>20</td>
<td>taper dose</td>
</tr>
<tr>
<td>21</td>
<td>other</td>
</tr>
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</table>
1. Has the subject had anything other than water to drink or eat in the past hour?
   - If YES, STOP HERE. Subject is ineligible to continue with ebc collection. If possible, wait until the full hour has passed, then proceed with collection.

2. Was ebc collection attempted at this visit?
   - If NO, complete Question #2a and STOP.
   - If YES, proceed to Question #3

   2a. Check the primary reason ebc collection was not attempted.
   - 1 Subject Refusal
   - 2 Equipment Unavailable
   - 3 Clinic Oversight
   - 4 Other ____________________

3. Time ebc collection started (based on 24-hour clock).

4. Time ebc collection stopped (based on 24-hour clock).

   - If collection time exceeds ten minutes, please provide an explanation below.

5. Did the subject experience any of the following during the collection process...
   - 5a. Sneezing?
   - 5b. Coughing?
   - 5c. Burping?
6. Were six eppendorf tubes aliquoted at this visit?

   If YES, proceed to Question #7.

6a. Which of the following explain why six tubes were not collected?

   Equipment Malfunction
   If YES, explain ________________________________

   Low Yield
   If YES, explain ________________________________

   Subject could not tolerate procedure
   If YES, explain ________________________________

   Other
   If YES, explain ________________________________

6b. Record the number of tubes aliquoted. __ tubes (1130)

   ➔ If '0', STOP HERE.

7. Was nitrogen gas layered on the tubes before closing them?

   If YES, explain ________________________________

8. Were the tubes stored immediately at -70° Celsius or colder?

   If NO, at what temperature were the tubes stored? _______ ° C (1170)

9. Attach one barcode label/dot pair from the subject’s visit-specific strip here. Write the barcode number from the label in the spaces provided.

   ________________________________ (1180)

Comments:___________________________________________________________________________
____________________________________________________________________________________
____________________________________________________________________________________
____________________________________________________________________________________
**EXHALED NITRIC OXIDE**

(Technician completed)

*Complete this form only if the subject is eligible according to the appropriate Pulmonary Procedures Checklist (PULMONARYCHK) form.*

1. Was the exhaled nitric oxide (ENO) procedure performed?
   
   ➔ **If NO,** complete Question #1a and **STOP.**
   
   ➔ **If YES,** proceed to Question #1b.

1a. What was the reason the ENO procedure could not be performed?

   ➔ **Equipment failure, please specify**

   ➔ **Equipment not calibrated**

   ➔ **Subject refusal**

   ➔ **Clinic oversight**

   ➔ **Other**

   ➔ **1** Yes ➔ **0** No (1000)

1b. Was the exhaled nitric oxide (ENO) procedure performed on the NIOX Mino?

   ➔ **If YES,** do not complete Question #11 on the next page since only 1 acceptable maneuver was obtained.

   ➔ **1** Yes ➔ **0** No (1015)
For each maneuver, record the time and $FE_{NO}$ value. If the maneuver was not accepted by the NIOX machine, record the time and select the 'Maneuver Not Acceptable' check box.

For a procedure done on the ACRN NIOX Machine, when TWO reproducible measurements are achieved, select the 'Reproducible Measurements' check box for both maneuvers. The two measurements are considered reproducible when they are within 5% of their mean or 1.25 ppb of their mean.

<table>
<thead>
<tr>
<th>Maneuver #</th>
<th>Time (based on 24-hour clock)</th>
<th>Measured $FE_{NO}$ (ppb)</th>
<th>Maneuver Not Acceptable</th>
<th>Clinic Use Only Reproducible Measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. Maneuver #1</td>
<td>1020</td>
<td>1030</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Maneuver #2</td>
<td>1060</td>
<td>1070</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Maneuver #3</td>
<td>1100</td>
<td>1110</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Maneuver #4</td>
<td>1140</td>
<td>1150</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Maneuver #5</td>
<td>1180</td>
<td>1190</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Maneuver #6</td>
<td>1220</td>
<td>1230</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Maneuver #7</td>
<td>1260</td>
<td>1270</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Maneuver #8</td>
<td>1300</td>
<td>1310</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Maneuver #9</td>
<td>1340</td>
<td>1350</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

11. Did the subject achieve two reproducible outcomes?  
   If NO, explain ____________________________________________  
   □ 1 Yes  □ 0 No (1380)
Complete this form only if the subject is eligible according to the appropriate Mannitol Challenge Testing Checklist form.

Asthma
Clinical
Research
Network

MANNITOL
CHALLENGE
TESTING

(Technician completed)

**Clinic Use Only** (Technician completed)
Use the FEV₁ value from the appropriate spirometry testing form as the baseline reference.

A. Baseline FEV₁ prior to mannitol challenge  ____ . ____ ____ L
   (obtained on the ACRN KoKo machine)
B. Mannitol Reversal Reference Value (Question A x 0.90 =  ____ . ____ ____ ____ L)
C. Target FEV₁ Value (Question 1000 x 0.8549 =  ____ . ____ ____ ____ L)

1. Reference (0 mg) FEV₁  ____ . ____ ____ ____ L (1000)

2. Did the subject drop ≥ 14.51% at the 0 mg dose?
   - If YES, proceed to Question #4 and record 0 for Question #4a.

   □ 1 Yes  □ 0 No (1010)

3. Mannitol Dispensation Coordinator Cough Evaluation
   (At each dose, indicate the severity of the subject’s cough)

<table>
<thead>
<tr>
<th>Dose</th>
<th>FEV₁</th>
<th>No Cough</th>
<th>Moderate</th>
<th>Extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td>3a. 5 mg</td>
<td>____ . ____ ____ L (1020)</td>
<td>____ (1030)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3b. 10 mg</td>
<td>____ . ____ ____ L (1040)</td>
<td>____ (1050)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3c. 20 mg</td>
<td>____ . ____ ____ L (1060)</td>
<td>____ (1070)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3d. 40 mg</td>
<td>____ . ____ ____ L (1080)</td>
<td>____ (1090)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3e. 80 mg</td>
<td>____ . ____ ____ L (1100)</td>
<td>____ (1110)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3f. 160 mg</td>
<td>____ . ____ ____ L (1120)</td>
<td>____ (1130)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3g. 160 mg (second)</td>
<td>____ . ____ ____ L (1140)</td>
<td>____ (1150)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3h. 160 mg (third)</td>
<td>____ . ____ ____ L (1160)</td>
<td>____ (1170)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
4. Did the subject achieve a PD$_{15}$?
   ➔ If NO, proceed to Question #5.

4a. PD$_{15}$ (mg)  

5. Time mannitol challenge ended (based on 24-hour clock)  

6. Subject’s FEV$_1$ after standard reversal from mannitol challenge  
   
   **Standard reversal = 2 puffs albuterol**  

   6a. FEV$_1$  

   6b. Time of FEV$_1$ in Question #6a (based on 24-hour clock)  

   6c. Was the FEV$_1$ from Question #6a ≥ the mannitol reversal reference value (B) in the gray box above?  
   ➔ If YES, STOP HERE and continue with remaining visit procedures.  
   ➔ If NO, proceed to the Additional Treatment for Mannitol Challenge Testing (MANNITOL_ADD_TRT) form.
**ADDITIONAL TREATMENT POST MANNITOL CHALLENGE TESTING**

(Technician completed)

*Complete this form only if the subject did not reverse to 90% of baseline FEV₁ after the first post-challenge treatment of albuterol.*

1. Was an additional treatment used in the first hour?
   - **If NO, skip to Question #3.**
   - **If YES, proceed to Question #3.**

   1a. Additional albuterol by MDI
       - **If NO, skip to Question #1b.**
       - Number of additional puffs of albuterol administered
         - ☐ 1 two
         - ☐ 2 four
         - ☐ 3 > four

   1b. Nebulized Beta-agonist
       - ☐ 1 Yes
       - ☐ 0 No

   1c. Subcutaneous epinephrine
       - ☐ 1 Yes
       - ☐ 0 No

   1d. Implementation of clinic emergency protocol or algorithm
       - ☐ 1 Yes
       - ☐ 0 No

   1e. Other (specify)
       - ☐ 1 Yes
       - ☐ 0 No

2. Subject’s FEV₁ after additional treatment within first hour.
   - 2a. FEV₁
       - ___. ___ ___ ___ L
   - 2b. Time of FEV₁ in Question #2a (based on 24-hour clock)
       - ___ ___ ___ (1090)
   - 2c. Was the FEV₁ from Question #2a ≥ the mannitol reversal reference value?
       - **If YES, STOP HERE and continue with remaining visit procedures.**
       - **If NO, proceed to Question #3.**
3. Was additional treatment used after one hour?  
   ➔ If NO, skip to Question #4.

3a. Additional albuterol by MDI  
   ➔ If NO, skip to Question #3b.
   Number of additional puffs of albuterol administered

   1  Yes  0  No (1110)

3b. Nebulized Beta-agonist  

   1  Yes  0  No (1120)

3c. Subcutaneous epinephrine  

   1  Yes  0  No (1140)

3d. Implementation of clinic emergency protocol or algorithm  

   1  Yes  0  No (1150)

3e. Treatment in the emergency room  

   1  Yes  0  No (1160)

3f. Overnight hospitalization
   ➔ If YES, please complete the Serious Adverse Event (SERIOUS) form.

3g. Other (specify) ____________________________  

   1  Yes  0  No (1190)

4. Subject’s final FEV1 after mannitol challenge.

4a. FEV1

   ___ . ___ ___ ___ L (1200)

4b. Time of FEV1 from Question #4a (based on 24-hour clock)

   ___ ___ ___ (1220)

4c. Was the FEV1 from Question #4a ≥ the mannitol reversal reference value?  
   ➔ If NO, complete the source documentation box below.

Physician Source Documentation

Physician’s signature: ____________________________ (1240)

Date: ___ / ___ / ___ ___ ___ (1250)

Time: ___ ___ ___ (based on 24-hour clock) (1260)
Complete this form only if the subject is eligible according to the Methacholine Challenge Testing Checklist (METHACHK) form.

Clinic Use Only (Technician completed)

Use the FEV1 value from the appropriate spirometry testing form as the baseline reference.

<table>
<thead>
<tr>
<th>Question</th>
<th>Description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>Baseline FEV1 prior to methacholine challenge</td>
<td>___ . ___ ___ L</td>
</tr>
<tr>
<td>B.</td>
<td>Methacholine Reversal Reference Value (Question A (\times 0.90) = ___ . ___ ___ L)</td>
<td></td>
</tr>
<tr>
<td>C.</td>
<td>Diluent FEV1 Reference Value (Question 1000 (\times 0.8049) = ___ . ___ ___ L)</td>
<td></td>
</tr>
</tbody>
</table>

1. Post Diluent FEV1 | ___ . ___ ___ L (1000)
2. Did the subject drop ≥ 20% at the diluent stage? | □ 1 Yes □ 0 No (1010)

   ➤ If YES, proceed to Question #5 and record 0 for Question #5a.
3. Last concentration of methacholine administered | ___ . ___ ___ ___ mg/ml (1020)
4. FEV1 after last concentration of methacholine administered | ___ . ___ ___ L (1030)
5. Did the subject achieve a PC20? | □ 1 Yes □ 0 No (1040)

   ➤ If NO, proceed to Question #6.

   5a. PC20 | ___ . ___ ___ ___ mg/ml (1050)
6. Time methacholine challenge ended (based on 24-hour clock) | ___ ___ ___ (1060)
7. Subject’s FEV1 after standard reversal from methacholine challenge
   - If subject is continuing with sputum induction, standard reversal = 4 puffs albuterol.
   - If subject is not continuing with sputum induction, standard reversal = 2 puffs albuterol.

   7a. FEV1 | ___ . ___ ___ L (1070)
7b. Time of FEV1 in Question #7a (based on 24-hour clock) | ___ ___ ___ (1090)
7c. Was the FEV1 from Question #7a ≥ the methacholine reversal reference value (B) in the gray box above? | □ 1 Yes □ 0 No (1100)

   ➤ If YES, STOP HERE and continue with remaining visit procedures.

   ➤ If NO, proceed to the Additional Treatment for Methacholine Challenge Testing (METHA_ADD_TRT) form.
**ADDITIONAL TREATMENT**
**POST METHACHOLINE CHALLENGE TESTING**

(Technician completed)

Complete this form only if the subject did not reverse to 90% of baseline FEV₁ after the first post-challenge treatment of albuterol.

1. Was an additional treatment used in the first hour?  
   - **Yes**  
   - **No** (1000)

   
   ➔ If NO, skip to Question #3.

1a. Additional albuterol by MDI  
   - **Yes**  
   - **No** (1010)

   Number of additional puffs of albuterol administered
   - **two**  
   - **four**  
   - **> four** (1020)

1b. Nebulized Beta-agonist  
   - **Yes**  
   - **No** (1030)

1c. Subcutaneous epinephrine  
   - **Yes**  
   - **No** (1040)

1d. Implementation of clinic emergency protocol or algorithm  
   - **Yes**  
   - **No** (1050)

1e. Other (specify) ________________________________  
   - **Yes**  
   - **No** (1060)

2. Subject’s FEV₁ after additional treatment within first hour.

2a. FEV₁  
   ___ . ___ ___ L (1070)

2b. FEV₁ (% predicted)  
   ___ ___ ___ % predicted (1080)

2c. Time of FEV₁ in Question #2a (based on 24-hour clock)  
   ___ ___ ___ (1090)

2d. Was the FEV₁ from Question #2a ≥ the methacholine reversal reference value (B) in the gray box on the Methacholine Challenge Testing (METHA) form?  
   - **Yes**  
   - **No** (1100)

   ➔ If YES, STOP HERE and continue with remaining visit procedures.

   ➔ If NO, proceed to Question #3.
3. Was additional treatment used after one hour?
   - If NO, skip to Question #4.

   a. Additional albuterol by MDI
      - If NO, skip to Question #3b.

      Number of additional puffs of albuterol administered
      - 1 two
      - 2 four
      - 3 > four

   b. Nebulized Beta-agonist
   c. Subcutaneous epinephrine
   d. Implementation of clinic emergency protocol or algorithm
   e. Treatment in the emergency room
   f. Overnight hospitalization
      - If YES, please complete the Serious Adverse Event (SERIOUS) form.

   g. Other (specify)

4. Subject’s final FEV₁ after methacholine challenge.
   a. FEV₁
      - ______ ______ L (1200)
   b. FEV₁ (% predicted)
      - ______ ______ % predicted (1210)
   c. Time of FEV₁ from Question #4a (based on 24-hour clock)
      - ______ ______ (1220)
   d. Was the FEV₁ from Question #4a ≥ the methacholine reversal reference value (B) in the gray box on the Methacholine Challenge Testing (METHA) form?
      - If NO, complete the source documentation box below.

   - If YES, please complete the Serious Adverse Event (SERIOUS) form.

   Physician Source Documentation
   - Physician’s signature: ________________________________ (1240)
   - Date: ______ / ______ / ________ (1250)
   - Time: ______ ______ (based on 24-hour clock) (1260)
Complete this form only if the subject is eligible according to the appropriate Pulmonary Procedure Checklist form and successfully completed baseline spirometry session(s).

1. Has the subject had any severe acute illness in the past 4 weeks?
   - Yes ☐  No ☐ (1000)

   If YES, has the subject received permission from the supervising physician to proceed with the methacholine challenge testing?
   - Yes ☐  No ☐ (1010)
   
   Physician’s Signature: ____________________________ (1015)

2. Does the subject have a baseline (pre-diluent) FEV₁ less than 55% of predicted?
   - Yes ☐  No ☐ (1020)

   Use the FEV₁ value from the appropriate spirometry testing form as the baseline reference.

3. Does the subject have a history of urinary retention?
   - Yes ☐  No ☐ (1030)

   If NO, proceed to Question #4.

   3a. If YES, is the subject randomized?

   If NO, proceed to Question #4 and complete the appropriate Termination of Study Participation form.

   3b. Was written medical clearance obtained from the study physician?

   If YES, obtain physician’s signature:

   ____________________________ (1055)

4. Is there any other reason the subject should not proceed with the methacholine challenge testing?
   - Yes ☐  No ☐ (1060)

   If YES, explain ____________________________

5. Is the subject eligible to proceed with the diluent (solution #0) spirometry testing for the methacholine challenge?
   - Yes ☐  No ☐ (1070)

   If any of the shaded boxes are completed, the subject is NOT eligible for the methacholine challenge.

   ➔ If YES, proceed to the Methacholine Challenge Testing (METHA) form.
### BASALT/TALC ALLOCATION CHECKLIST

(Clinic Coordinator completed)

<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Since Visit 1, has the subject experienced a significant asthma exacerbation as defined in the protocol?</td>
<td></td>
<td></td>
<td>(1000)</td>
</tr>
<tr>
<td>2. Since Visit 1, has the subject received treatment with any excluded medications (P16_EXCLDRUG)?</td>
<td></td>
<td></td>
<td>(1010)</td>
</tr>
<tr>
<td>3. Using the history stored in the DOSER™, did the subject take at least 75% of the required puffs from his or her QVAR inhaler during the interval between Visits 2 and 3?</td>
<td></td>
<td></td>
<td>(1020)</td>
</tr>
<tr>
<td>➔ Use Question #3 from P16_COMPLY to answer this question</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Using the history stored in the DOSER™, did the subject take 4 puffs per day (correct daily dose) on at least 75% of the days during the interval between Visits 2 and 3?</td>
<td></td>
<td></td>
<td>(1030)</td>
</tr>
<tr>
<td>➔ Use Question #6 from P16_COMPLY to answer this question</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. Did the subject record both AM and PM peak flow measurements and symptoms on his or her Diary Cards (P16_DIARY) on at least 75% of the days during the interval between Visits 2 and 3?</td>
<td></td>
<td></td>
<td>(1040)</td>
</tr>
<tr>
<td>6. Did the subject measure his or her AM and PM peak flow on schedule and accurately transcribe the measurements on his or her Diary Cards (P16_DIARY) on at least 75% of the days during the interval between Visits 2 and 3?</td>
<td></td>
<td></td>
<td>(1050)</td>
</tr>
<tr>
<td>7. Is the subject’s prebronchodilator FEV₁ &gt; 40% of predicted?</td>
<td></td>
<td></td>
<td>(1060)</td>
</tr>
<tr>
<td>8. Does the subject wish to withdraw consent from the study?</td>
<td></td>
<td></td>
<td>(1070)</td>
</tr>
<tr>
<td>9. Is there any new information that makes the subject ineligible according to the eligibility criteria? If YES, describe: ____________________________</td>
<td></td>
<td></td>
<td>(1080)</td>
</tr>
<tr>
<td>10. Is there any other reason why this subject should not be allocated to the BASALT or TALC study? If YES, describe: ____________________________</td>
<td></td>
<td></td>
<td>(1090)</td>
</tr>
</tbody>
</table>
ALLOCATION CHECKLIST

11. Is the subject eligible for allocation?
   If any of the shaded boxes is completed, the subject is ineligible.
   → If YES, continue with rest of form.
   → If NO, complete the BASALT/TALC RUNIN Termination of Study Participation (P16_TERM) form.

Yes ☐  No ☐ (1100)

12. Is the subject’s prebronchodilator FEV₁ > 70% of predicted?
   → If NO, skip to Question # 14.

Yes ☐  No ☐ (1110)

13. Did the subject answer 0 or 1 for each of the three questions on the ACRN Asthma Evaluation Questionnaire (AEQ) at this visit?
   → If YES, skip to Question # 15. Subject should be allocated to the BASALT study.

Yes ☐  No ☐ (1120)

14. Does the subject have any medical contraindications for tiotropium use (i.e., narrow angle glaucoma, prostatic hypertrophy, bladder-neck obstruction, renal insufficiency)?
   → If YES, subject is ineligible to continue in the BASALT/TALC studies. Complete a BASALT/TALC RUNIN Termination of Study Participation (P16_TERM) form.
   → If NO, subject should be allocated to the TALC study.

Yes ☐  No ☐ (1130)

15. Indicate the study into which the subject is enrolling.

BASALT ☐  TALC ☐

17  18 (1140)

16. Record the date on which the subject originally signed the informed consent document for the study to which he or she has been allocated.

Month / Day / Year (1150)

After study allocation, complete the following procedures:

→ Record study to which the subject was allocated (P16_LOG).
→ Enroll subject in appropriate protocol.
Check the following compliance criteria at Visits 2 and 3.

DOSE™ Compliance for QVAR MDI

1. Total number of scheduled puffs since the last visit
   \[ \text{Value obtained from Question #1 on P16\_COMPLY\_WKS} \]
   \[ \text{___ ___ ___ puffs (1000)} \]

2. Total number of puffs in DOSE™ history
   \[ \text{Value obtained from Question #2 on P16\_COMPLY\_WKS} \]
   \[ \text{___ ___ ___ puffs (1010)} \]

3. Percent compliance = \[ \frac{\text{Question #2}}{\text{Question #1}} \times 100 \]
   \[ \text{___ ___ ___ \% (1020)} \]
   \[ \text{If the subject took less than 75\% of the scheduled QVAR puffs, re-emphasize the importance of maintaining the daily dosing schedule.} \]

4. Total number of full days since the last visit
   \[ \text{Value obtained from Question #4 on P16\_COMPLY\_WKS} \]
   \[ \text{___ ___ (1030)} \]

5. Total number of compliant days
   \[ \text{Value obtained from Question #5 on P16\_COMPLY\_WKS} \]
   \[ \text{___ ___ (1040)} \]

6. Percent compliance = \[ \frac{\text{Question #5}}{\text{Question #4}} \times 100 \]
   \[ \text{___ ___ ___ \% (1050)} \]
   \[ \text{If the subject took the correct daily dose less than 75\% of the days, re-emphasize the importance of maintaining the daily dosing schedule.} \]
**To the subject:** If your peak flow is below ___ liters/minute, use your RESCUE albuterol inhaler as instructed in the handout “If Your Asthma Gets Worse.” Contact study personnel if your peak flow does not increase to this value after 1 hour of RESCUE use, or if you are experiencing extreme symptoms. If you have taken at least 10 puffs/24 hours for the past 48 hours from your RESCUE inhaler, contact study personnel.

Please use black ink to complete.

<table>
<thead>
<tr>
<th>Date</th>
<th>Day 1</th>
<th>Day 2</th>
<th>Day 3</th>
<th>Day 4</th>
<th>Day 5</th>
<th>Day 6</th>
<th>Day 7</th>
</tr>
</thead>
<tbody>
<tr>
<td>(dd/mm)</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
</tr>
</tbody>
</table>

**MORNING EVALUATION** (Between 5 AM and 10 AM)

1. Number of times you woke up last night due to asthma (1010)
2. Time of AM Peak Flow (within 15 minutes of awakening) (1020)
3. AM Peak Flow (liters/min)** (1030) / (1035)
4. Total number of puff(s) from QVAR Inhaler (AM) (1040)

**Symptoms**++ during the night

5. Shortness of Breath (1050)
6. Chest Tightness (1060)
7. Wheezing (1070)
8. Cough (1080)
9. Phlegm/Mucus (1090)

**NIGHT-TIME EVALUATION** (Between 8 PM and 1 AM)

10. Time of PM Peak Flow (between 8 PM and 1 AM) (1100)
11. PM Peak Flow (liters/min)** (1110) / (1115)
12. Total number of puff(s) from QVAR Inhaler (PM) (1120)

**Symptoms**++ since you woke

13. Shortness of Breath (1130)
14. Chest Tightness (1140)
15. Wheezing (1150)
16. Cough (1160)
17. Phlegm/Mucus (1170)

**24 HOUR EVALUATION**

18. Total number of puffs from albuterol (RESCUE) inhaler during past 24 hours. *(Do not record preventive use.)* (1180)
19. Total number of times you dosed from your albuterol (RESCUE) inhaler during past 24 hours. *(Do not record preventive use.)* (1190)

**Symptom Severity Rating Scale**

0 = Absent  No symptom
1 = Mild  Symptom was minimally troublesome, i.e. not sufficient to interfere with normal daily activity or sleep.
2 = Moderate  Symptom was sufficiently troublesome to interfere with normal daily activity or sleep.
3 = Severe  Symptom was so severe as to prevent normal activity and/or sleep.

++ Record the best of three attempts. Circle the value if you have taken any medication from your RESCUE albuterol inhaler in the last 2 hours.
(Clinic Coordinator completed)

The subject should rest quietly for at least five minutes before the electrocardiogram.

1. Time electrocardiogram started *(based on 24-hour clock)*   _ ___ ___ ___ (1000)

2. Ventricular heart rate   _ ___ ___ beats/min (1010)

3. Cardiac cycle measurements
   3a. P - R Interval   _ ___ ___ milliseconds (1020)
   3b. QRS Duration   _ ___ ___ milliseconds (1030)
   3c. Q - T Interval   _ ___ ___ milliseconds (1040)

Physician Source Documentation

Physician's signature: __________________________ (1050)
Date: ___ / ___ / ___ ___ (1060)
(Subject Interview completed)

1. **Did the subject sign either (or both) of the BASALT/TALC Informed Consent(s)?**
   - [ ] Yes [ ] No (1000)
   
   If YES, record the date the form(s) was (were) signed.
   
   ➔ **Consent should be reviewed and signed on the day Visit 1 is performed.**

2. **Are you planning to move away from this clinical center in the upcoming year such that your ability to complete the study will be jeopardized?**
   - [ ] Yes [ ] No (1020)

3. **Have you had a respiratory tract infection in the past 4 weeks?**
   - [ ] Yes [ ] No (1030)

4. **Have you experienced a significant asthma attack in the past 4 weeks?**
   - [ ] Yes [ ] No (1040)

5. **Do you work the night shift or have an altered day/night cycle for other reasons?**
   - [ ] Yes [ ] No (1050)

6. **Is the subject eligible to proceed?**
   - [ ] Yes [ ] No (1060)
   
   If any of the shaded boxes are completed, the subject is ineligible.
   
   ➔ If YES, proceed with remaining Visit 1 procedures.

**Subject Source Documentation**

Subject Initials: [ ] [ ] [ ] (1070)

Date: [ ] [ ] / [ ] [ ] / [ ] [ ] [ ] (1080)
1. Is the subject 18 years of age or older?
   - Yes ☐ No ☐

2. Does the subject have current evidence of any of the conditions listed on the Exclusionary Medical Conditions (P16_EXCLMED) reference card?
   If YES, describe ____________________________
   - Yes ☐ No ☐

2a. Does the subject have unstable or severe coronary artery disease or a history of myocardial infarction within 6 months of Visit 1?
   - Yes ☐ No ☐

3. Has the subject taken any medications listed on the Exclusionary Drugs (P16_EXCLDRUG) reference card within the specified time periods?
   If YES, describe ____________________________
   - Yes ☐ No ☐

4. Is the subject currently taking prescription or over-the-counter medication(s) other than those listed on the Allowed Medications (P16_MEDALLOW) reference card?
   If YES, describe ____________________________
   - Yes ☐ No ☐

5. Based on input from the subject and the study physician, will the subject need to use intranasal steroids at any time during the study?
   - Yes ☐ No ☐

5a. If YES, is the subject willing to use a single intranasal steroid at a stable dose continuously for the duration of the study?
   - Yes ☐ No ☐

6. Is the subject regularly using inhaled corticosteroids?
   - Yes ☐ No ☐

   ➤ If NO, skip to Question #7 and complete rest of form.

   ➤ If YES, answer Questions #6a and 6b, then skip to Question #8.

6a. Has the subject been on a stable dose of inhaled corticosteroids for at least 2 weeks?
   - Yes ☐ No ☐

6b. Has the subject been using greater than the equivalent of 1000 μg inhaled fluticasone daily?
   - Yes ☐ No ☐

   ➤ Refer to the ICS Equivalency (P16_ICS_EQUIV) reference card.
7. Has the subject used or received a prescription for an asthma controller (inhaled corticosteroids, leukotriene modifier, and/or long-acting beta-agonist) during the past year?
   - Yes □ 0 No □

7a. If **NO**, does the subject report experiencing asthma symptoms more than twice a week?
   - Yes □ 0 No □

8. Is the subject currently receiving hyposensitization therapy (e.g., allergy shots) other than an established maintenance regimen implemented continuously for a minimum of 3 **months**?
   - Yes □ 0 No □

9. Has the subject experienced a life-threatening asthma exacerbation requiring treatment with intubation and mechanical ventilation in the past **5 years**?
   - Yes □ 0 No □

10. Has the subject smoked cigarettes, a pipe, cigar, marijuana, or any other substance in the past year?
    - Yes □ 0 No □

11. Record smoking history in pack-years. (Enter 00.0 if subject never smoked.)
    - __ __ __ (1150)

   Is Question #11 ≥ 10?
    - Yes □ 0 No □

12. Is the subject potentially able to bear children?
    - If subject is male, check N/A and go to Question #13.
    - Yes □ 0 No □ □ N/A (1170)

   12a. If **YES**, is the subject using one of the approved methods indicated on the Birth Control (BIRCTRL) reference card?
    - Yes □ 0 No □

   12b. If **YES**, is the subject currently pregnant or lactating?
    - Yes □ 0 No □

13. Is the subject eligible to proceed?
   - Yes □ 0 No □

   **If any of the shaded boxes are completed, the subject is ineligible.**

   **If YES, proceed with remaining Visit 1 procedures.**

---

Subject Source Documentation
Subject Initials: __ __ __ (1210)
Date: ___ / ___ / ________ (1220)
Section 1

1. Is the subject’s prebronchodilator FEV₁ > 40% of predicted?

   ➔ If NO, STOP here. Subject is ineligible for the study.

2. Does the subject have valid source documentation for a methacholine challenge (ACRN systems and procedures only) within the past 6 months?

   ➔ If NO, skip to Question #3.

   ➔ If YES, record values below:

   PC₂₀: ___ ___ ___ ___ ___ ___ ___ mg/ml (1002)

   Source Documentation Date: ___ / ___ / ___ ___ (1003)

   Technician ID: ___ ___ ___ ___ ___ ___ ___ (1004)

   Supervisor ID: ___ ___ ___ ___ ___ ___ ___ (1005)

2a. Was the subject using ICS regularly at the time the challenge was performed?

   ➔ If YES, complete Question #2b and skip to Question #2d.

   ➔ If NO, complete Question #2c and continue with rest of form.

2b. Does the subject have source documentation of a methacholine PC₂₀ ≤ 16 mg/ml?

   ➔ If YES, complete Question #2b and skip to Question #2d.

   ➔ If NO, complete Question #2c and continue with rest of form.

2c. Does the subject have source documentation of a methacholine PC₂₀ ≤ 8 mg/ml?

   ➔ If YES, complete Question #2c and continue with rest of form.

2d. Is the subject eligible to proceed?

   ➔ If YES, continue with remaining visit procedures and complete Section 4.

   ➔ If NO, complete Question #3 on the next page and proceed accordingly.
3. Is the subject’s prebronchodilator FEV$_1$ $\geq$ 55% of predicted and he/she qualifies for methacholine challenge?
   - If **YES**, complete Section 2.
   - If **NO**, complete Section 3.

<table>
<thead>
<tr>
<th>Subject ID:</th>
<th>Visit Number:</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

**ELIGIBILITY CHECKLIST 3**

### Section 2

4. Is the subject regularly using ICS at this time?
   - If **YES**, complete Question #5 and skip to Question #7.
   - If **NO**, complete Question #6 and continue with rest of form.

<table>
<thead>
<tr>
<th>1</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

5. Does the subject have a methacholine PC$_{20}$ $\leq$ 16 mg/ml?
   - Yes
   - No

6. Does the subject have a methacholine PC$_{20}$ $\leq$ 8 mg/ml?
   - Yes
   - No

7. Is the subject eligible to proceed?
   - If either shaded box in Section 2 is completed, the subject is ineligible at this point.
   - If **YES**, continue with remaining visit procedures and complete Section 4.
   - If **NO**, the subject may return at a later date for a continuation visit to perform albuterol reversibility testing to qualify. Complete Question #8 and proceed accordingly.

<table>
<thead>
<tr>
<th>1</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

8. Will the subject complete reversibility testing?
   - If **NO**, STOP here. Subject is ineligible for the study.
   - If **YES**, continue with visit procedures on the P16_VISITA checklist and complete Section 3.

<table>
<thead>
<tr>
<th>1</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>

### Section 3

9. Did the subject’s FEV$_1$ improve $\geq$ 12% in response to four puffs of albuterol?
   - If **YES**, continue with remaining visit procedures and complete Section 4 on the next page.
   - If **NO**, STOP here. Subject is ineligible for the study.

   - Yes
   - No

<table>
<thead>
<tr>
<th>1</th>
<th>0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>No</td>
</tr>
</tbody>
</table>
### Section 4

10. Is the subject able to use the AM1® device correctly, as evidenced by achieving a satisfactory rating on the AM1® Performance Checklist (PERF_AM1)?

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>0</td>
</tr>
</tbody>
</table>

11. Is the subject able to use a metered dose inhaler (MDI) properly, as evidenced by achieving a score of 6 on two consecutive, separate inhalations using the MDI Inhalation Technique Checklists (SCORE, TECH_MDI)?

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>0</td>
</tr>
</tbody>
</table>

12. Is the subject eligible to proceed?

If either shaded box in Section 4 is completed, the subject is ineligible.

- If YES, continue with remaining visit procedures.
- If NO, STOP here. Subject is ineligible for the study.
1. Eosinophils (absolute count)  ___ ___ ___ ___ /mm$^3$ (1000)
(Subject Interview completed)

**COLD HISTORY**

1. On average, how many respiratory tract infections/colds do you experience per year? __ __ (1000)

2. How severe are your colds usually?

- □ 1 Extremely severe
- □ 2 Very severe
- □ 3 Severe
- □ 4 Moderate
- □ 5 Mild
- □ 6 Extremely mild (1010)

3. When you get colds, how often do they make your asthma worse?
   - □ 1 Always
   - □ 2 Usually
   - □ 3 Sometimes
   - □ 4 Rarely
   - □ 5 Never (1020)

   ➔ If NEVER, skip to Question # 5.

4. When colds make your asthma worse, how severe does your asthma usually get?

- □ 1 Extremely severe
- □ 2 Very severe
- □ 3 Severe
- □ 4 Moderate
- □ 5 Mild
- □ 6 Extremely mild (1030)

**PRIOR ASTHMA TREATMENT**

*I will read a list of medications. Indicate if you have ever used each medication. If you have, please indicate, to the best of your knowledge, the date last taken.*

If Yes, indicate date medication was last taken

- month / day / year

5. Non-long-acting Inhaled Beta-Agonists (Bronkaid Mist, Duo-Medihaler, Medihaler-Epi, Primatene Mist, Alupent, Brethaire, Brethine, Bronkometer, Maxair, Metaprel, Proventil, Tornalate, Ventolin, Xopenex and others)
   - □ 1 Yes □ 0 No □ 8 Unknown __ / __ / ______ (1040) (1050) (1060) (1070)

5a. If YES, indicate average daily puffs in the past month. (Enter ‘00’ if none used.) ___ ___ puffs (1080)

6. Long-acting Inhaled Beta-Agonists (Serevent, Foradil, Advair Diskus Symbicort Turbuhaler)
   - □ 1 Yes □ 0 No □ 8 Unknown __ / __ / ______ (1090) (1100) (1110) (1120)
<table>
<thead>
<tr>
<th></th>
<th>Medical History</th>
</tr>
</thead>
<tbody>
<tr>
<td>7</td>
<td>Asthma medication via a Nebulizer Machine</td>
</tr>
<tr>
<td>8</td>
<td>Oral Beta-Agonists</td>
</tr>
<tr>
<td></td>
<td>(Alupent, Brethine, Bricanyl, Metaprel, Proventil, Ventolin, Repetabs, Volmax and others)</td>
</tr>
<tr>
<td>9</td>
<td>Short-acting Oral Theophylline</td>
</tr>
<tr>
<td></td>
<td>(Aminophylline, Slo-Phyllin and others)</td>
</tr>
<tr>
<td>10</td>
<td>Sustained release Oral Theophylline</td>
</tr>
<tr>
<td></td>
<td>(Slo-bid, Theo-Dur, Uniphyl and others)</td>
</tr>
<tr>
<td>11</td>
<td>Inhaled Anticholinergic</td>
</tr>
<tr>
<td></td>
<td>(Atrovent, Combivent, Spiriva)</td>
</tr>
<tr>
<td>12</td>
<td>Anti-allergic Inhaled Medications</td>
</tr>
<tr>
<td></td>
<td>(Intal, Tilade and others)</td>
</tr>
<tr>
<td>13</td>
<td>Anti-allergic Nasal Medications</td>
</tr>
<tr>
<td></td>
<td>(Nasalcom, Astel and others)</td>
</tr>
<tr>
<td>14</td>
<td>Anti-allergic Oral Medications</td>
</tr>
<tr>
<td></td>
<td>(Allegra, Claritin, Zyrtec, Chlor-Trimeton and others)</td>
</tr>
<tr>
<td>15</td>
<td>Leukotriene Antagonist / 5LO Inhibitors</td>
</tr>
<tr>
<td></td>
<td>(Accolate, Zyflo, Singulair)</td>
</tr>
<tr>
<td>16</td>
<td>IgE Blocker</td>
</tr>
<tr>
<td></td>
<td>(Xolair)</td>
</tr>
<tr>
<td>17</td>
<td>Topical Steroids - Prescription</td>
</tr>
<tr>
<td></td>
<td>(Synalar, Lidex, Dermacin, Fluocinonide and others)</td>
</tr>
<tr>
<td>18</td>
<td>Topical Steroids - OTC</td>
</tr>
<tr>
<td></td>
<td>(Hydrocortisone - multiple strengths and products)</td>
</tr>
<tr>
<td>19</td>
<td>Nasal Steroids</td>
</tr>
<tr>
<td></td>
<td>(Beconase, Vancenase, Flonase, Nasacort, Nasalide, Nasarel, Rhinocort, Nasonex and others)</td>
</tr>
<tr>
<td>20</td>
<td>Oral Steroids</td>
</tr>
<tr>
<td></td>
<td>(Prednisone, Medrol and others)</td>
</tr>
</tbody>
</table>
MEDICAL HISTORY

Subject ID:  6 - ____________
Visit Number:  1

21. Inhaled Steroids

☐ 1 Yes  ☐ 0 No  ☐ 8 Unknown

☐ 1 beclomethasone MDI (1 puff = 42 μg)  
(e.g., Beclovent, Vanceril)
☐ 2 beclomethasone MDI (1 puff = 84 μg)  
(e.g., Vanceril-DS)
☐ 3 beclomethasone HFA (1 puff = 40 μg)  
(e.g., QVAR)
☐ 4 beclomethasone HFA (1 puff = 80 μg)  
(e.g., QVAR)
☐ 5 budesonide DPI (1 puff = 80 μg)  
(e.g., Symbicort Turbuhaler)
☐ 6 budesonide DPI (1 puff = 160 μg)  
(e.g., Symbicort Turbuhaler)
☐ 7 budesonide DPI (1 puff = 200 μg)  
(e.g., Pulmicort Turbuhaler)
☐ 8 budesonide DPI (1 puff = 320 μg)  
(e.g., Symbicort Turbuhaler)
☐ 9 flunisolide MDI (1 puff = 250 μg)  
(e.g., Aerobid, Aerobid - M)
☐ 10 fluticasone MDI (1 puff = 44 μg)  
(e.g., Flovent)
☐ 11 fluticasone MDI (1 puff = 110 μg)  
(e.g., Flovent)
☐ 12 fluticasone MDI (1 puff = 220 μg)  
(e.g., Flovent)
☐ 13 fluticasone DPI (1 puff = 50 μg)  
(e.g., Flovent Rotadisk)
☐ 14 fluticasone DPI (1 puff = 100 μg)  
(e.g., Advair Diskus)
☐ 15 fluticasone DPI (1 puff = 250 μg)  
(e.g., Advair Diskus)
☐ 16 fluticasone DPI (1 puff = 500 μg)  
(e.g., Advair Diskus)
☐ 17 mometasone DPI (1 puff = 220 μg)  
(e.g., Asmanex Twishtaler)
☐ 18 triamcinolone acetonide MDI (1 puff = 100 μg)  
(e.g., Azmacort)
☐ 19 other __________________________

If Yes, indicate date medication was last taken month / day / year

(1690) (1700) (1710) (1720)

If NO or unknown, skip to Question #22.
If YES, complete Questions #21a - 21c.

21a. Indicate most recent type of inhaled steroid taken
21b. Indicate number of daily puffs used  ___ ___ puffs (1740)

21c. Indicate how long you used the inhaled steroid (duration of use)

- 1 - less than 1 month
- 2 - 1 - 6 months
- 3 - greater than 6 months (1750)

PRIOR CHOLESTEROL TREATMENT WITH STATIN DRUGS
Indicate if you have ever used statin medications. If you have, please indicate, to the best of your knowledge, the most recent drug taken, date last taken, and the total daily dose.

22. Statin medications

☑ 1 Yes ☐ 0 No ☐ 8 Unknown

☐ _ _ / _ _ / _ _ _ _ (2030) ☐ _ _ / _ _ / _ _ _ _ (2040) ☐ _ _ / _ _ / _ _ _ _ (2050) ☐ _ _ / _ _ / _ _ _ _ (2060)

If Yes, indicate date medication was last taken month / day / year

- 1 atorvastatin (e.g., Lipitor)
- 2 cerivastatin (e.g., Baycol)
- 3 fluvastatin (e.g., Lescol)
- 4 lovastatin (e.g., Advicor, Mevacor)
- 5 pravastatin (e.g., Pravachol)
- 6 simvastatin (e.g., Vytorin, Zocor)
- 7 rosuvastatin (e.g., Crestor) (2070)

22b. Indicate the total daily dose used (If dosage is unknown enter ‘999’)

☐ _ _ _ _ . ___ mg (2080)

PRIOR DISEASES, ILLNESSES AND SURGERIES
Have you had any diseases, illnesses, conditions or surgeries related to the following areas?

23. Skin

☑ 1 Yes ☐ 0 No (1760)

24. Blood, Lymph, or Immune Systems

☑ 1 Yes ☐ 0 No (1770)

25. Eyes

☑ 1 Yes ☐ 0 No (1780)

26. Ears, Nose, or Throat

☑ 1 Yes ☐ 0 No (1790)

26a. Have you ever had nasal polyps?

☐ 1 Yes ☐ 0 No ☐ 9 Don’t know (1800)

26ai. If YES, have you ever had any nasal polyps removed?

☐ 1 Yes ☐ 0 No (1810)

26b. Do you have chronic or recurrent sinusitis (treated with antibiotics)?

☐ 1 Yes ☐ 0 No ☐ 9 Don’t know (1820)
<table>
<thead>
<tr>
<th>Subject ID: 16 - - -</th>
<th>Visit Number: 1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MEDICAL HISTORY</strong></td>
<td></td>
</tr>
<tr>
<td>27. Breasts</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>28. Endocrine Systems</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>29. Lung - other than asthma</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>29a. Have you ever had pneumonia?</td>
<td>1 Yes 0 No 9 Don’t know</td>
</tr>
<tr>
<td>29b. Have you ever had bronchitis?</td>
<td>1 Yes 0 No 9 Don’t know</td>
</tr>
<tr>
<td>30. Heart and Blood Vessels</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>31. Liver or Pancreas</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>32. Kidneys or Urinary Tract System</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>33. Reproductive System</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>34. Stomach or Intestines</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>34a. Do you have gastroesophageal reflux disease (GERD)?</td>
<td>1 Yes 0 No 9 Don’t know</td>
</tr>
<tr>
<td>35. Muscles or Bones</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>36. Nervous System</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>37. Psychiatric</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>38. Other</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>39. Drug Allergies</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td>40. Food Allergies</td>
<td>1 Yes 0 No</td>
</tr>
<tr>
<td><strong>SUBJECT’S WEIGHT</strong></td>
<td></td>
</tr>
<tr>
<td>(Clinic Coordinator completed)</td>
<td></td>
</tr>
<tr>
<td>41. Weight (without shoes or heavy clothing)</td>
<td>___ ___ . ___ kg</td>
</tr>
</tbody>
</table>

Subject Source Documentation

Subject’s Initials: ___ ___ (2010)

Date: ___ / ___ / _____ (2020)
Please reference the Drug Classifications list for a complete list of examples for the questions below. If any medications other than study QVAR or rescue albuterol were used, record the medication(s) on the Concomitant Medications for Asthma/Allergies and Adverse Events (CMED) form.

<table>
<thead>
<tr>
<th>Question</th>
<th>1</th>
<th>Yes</th>
<th>0</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Have you consumed caffeine in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: Pepsi, Coke, Coffee, Mountain Dew, Tea, Rootbeer, Red Bull</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Have you used medications with caffeine in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: Anacin, Darvon compound, Esgic, Excedrin, Fiorinal, Fioricet, No Doz, Norgesic, Vivarin</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. Have you used any weight loss medications in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: bitter orange, Xenadrine EFX, Thermorexin</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. Have you consumed any food containing alcohol or beverages containing alcohol in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>5. Have you used any oral antihistamines in the past 48 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: Allegra, Chlor-Trimeton, Claritin, Tylenol PM</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6. Have you used any nasal antihistamines in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: Astelin, Livostin, Patanase</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7. Have you used any oral decongestants or cold remedies in the past 48 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: pseudoephedrine (Sudafed), Tylenol Allergy</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8. Have you used any nasal decongestants in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: oxymetazoline (Afrin)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9. Have you used any cough medicines, anti-tussives, or expectorants in the past 48 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: guaifenesin, dextromethorphan, Duratuss, Benylin, Triaminic expectorant, Dayquil Anti-Cough</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10. Have you used a rescue intermediate-acting inhaled beta-agonist in the past 6 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Example: albuterol (Ventolin or Proventil), study RESCUE</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11. Have you used any nasal steroids in the past 48 hours?</td>
<td>1</td>
<td>Yes</td>
<td>0</td>
<td>No</td>
</tr>
<tr>
<td>Examples: Flonase, Rhinocort, Nasonex</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
12. Have you used any smokeless tobacco products today?  
   Examples: chewing tobacco, snuff
   □ 1 Yes  □ 0 No (1095)

13. At this time, is your asthma worse because of recent exposure to triggers?  
   Examples: cold air, smoke, allergens, recent exercise, a recent respiratory tract infection, or other pulmonary infection
   □ 1 Yes  □ 0 No (1100)

14. Is there any other reason you should not proceed with spirometry testing?  
   If YES, explain ________________________________

15. Is the subject eligible to proceed with the spirometry testing?  
   If any of the shaded boxes are filled in, the subject is ineligible for spirometry and exhaled nitric oxide testing.
   ➤ If YES, proceed to Question #16 or the next form/procedure listed on the visit procedure checklist.
   □ 1 Yes  □ 0 No (1120)

---

Complete for all subjects at Visit 1.
If subject is less than 21 years old, complete Question #16 at each visit.

16. Height (without shoes)  _____ _____ _____ cm (1130)
This questionnaire is to be completed at Visit 12 by the ACRN study coordinator who was primarily responsible for the subject’s BASALT visits during the preceding 36 weeks. If a randomized subject terminates prior to Visit 12, this form should be completed at the time of the termination visit.

1. **Blinded Scheduled Inhalers (1000)/(1010)**
   
   Subjects in the BASALT study were randomized to receive three inhalers where one contained active drug and two contained placebo. You were blinded to the actual treatment assignments. Please check the box that most closely represents your feelings about the treatment the subject received during the randomized treatment period (Visit 4 through Visit 12).

   (Choose one statement that represents your feelings and choose one set of inhalers for the statement.)

   - 1 I am certain the inhaler that contained active drug is:
     - 1 Inhaler A
     - 2 Inhaler B
     - 3 Inhaler C

   - 2 I think the inhaler that contained active drug is:
     - 1 Inhaler A
     - 2 Inhaler B
     - 3 Inhaler C

   - 3 I have no idea which inhaler contained active drug however my best guess would be:
     - 1 Inhaler A
     - 2 Inhaler B
     - 3 Inhaler C

2. Please comment with respect to any other observations you made that helped you make your choice in Question #1.

   ________________________________________________________________

   ________________________________________________________________

   ________________________________________________________________

Clinic Coordinator Source Documentation

Coordinator’s Initials: _______ __________ (1020)

Date: _______ / _______ / _______ _______ (1030)
**BASALT**  
**CHANGE IN**  
**MEDICATIONS**  

(Clinic Coordinator completed)  

**Complete this form if the subject has experienced an adverse event that resulted in altering the dose of any of the subject’s study medications.**

1. Related Adverse Event Number  
   __ __ (1000)

2. Inhaler A  
   - □ 1 Discontinued  
   - □ 2 Reduced  
   - □ 3 Increased  
   - □ 4 Unchanged  
   - □ 5 Not Applicable (1010)

   ➔ If **Unchanged or Not Applicable**, proceed to Question #3.

2a. Date change began  
   __ __ / __ __ / __ __ ___ (1020)

2b. Date change ended  
   __ __ / __ __ / __ __ ___ (1030)

2c. Ongoing at current visit  
   □ 1 (1040)

3. Inhaler B  
   - □ 1 Discontinued  
   - □ 2 Reduced  
   - □ 3 Increased  
   - □ 4 Unchanged  
   - □ 5 Not Applicable (1050)

   ➔ If **Unchanged or Not Applicable**, proceed to Question #4.

3a. Date change began  
   __ __ / __ __ / __ __ ___ (1060)

3b. Date change ended  
   __ __ / __ __ / __ __ ___ (1070)

3c. Ongoing at current visit  
   □ 1 (1080)

4. Inhaler C  
   - □ 1 Discontinued  
   - □ 2 Reduced  
   - □ 3 Increased  
   - □ 4 Unchanged  
   - □ 5 Not Applicable (1090)

   ➔ If **Unchanged**, stop here.

4a. Date change began  
   __ __ / __ __ / __ __ ___ (1100)

4b. Date change ended  
   __ __ / __ __ / __ __ ___ (1110)

4c. Ongoing at current visit  
   □ 1 (1120)
BASALT COMPLIANCE CHECKLIST

(Clinic Coordinator completed)

Check the following compliance criteria at all scheduled Visits 4 - 12.

At Visit 4 Only, compliance is calculated for 14 full days prior to the visit (not including Visit 3 or prior).

1. **DOSER™ Compliance for Inhaler A**

   *If the interval between visits exceeds 30 days, complete Questions #1a - #1f using data for the 30 days prior to the visit.*

   1a. Total number of scheduled puffs since the last visit  ____ ____ ____ puffs (1000)

   ➔ Value obtained from Question #1 on P17_COMPLY_WKS_A

   1b. Total number of puffs in yellow DOSER™ history  ____ ____ ____ puffs (1010)

   ➔ Value obtained from Question #2 on P17_COMPLY_WKS_A

   1c. Percent compliance = \( \frac{\text{Question #1b}}{\text{Question #1a}} \times 100 \)  ____ ____ ____ \% (1020)

   ➔ *If the subject took less than 75% of the scheduled Inhaler A puffs, re-emphasize the importance of maintaining the daily dosing schedule.*

   1d. Total number of full days since the last visit  ____ ____ days (1030)

   ➔ Value obtained from Question #4 on P17_COMPLY_WKS_A

   1e. Total number of compliant days  ____ ____ days (1040)

   ➔ Value obtained from Question #5 on P17_COMPLY_WKS_A

   1f. Percent compliance = \( \frac{\text{Question #1e}}{\text{Question #1d}} \times 100 \)  ____ ____ ____ \% (1050)

   ➔ *If the subject took the correct daily dose less than 75% of the days, re-emphasize the importance of maintaining the daily dosing schedule.*
2. **DOSERTM Compliance for Inhaler B**

*If the interval between visits exceeds 30 days, complete Questions #2a - #2f using data for the 30 days prior to the visit.*

2a. Total number of scheduled puffs since the last visit

   

   ➔ Value obtained from Question #1 on P17_COMPLY_WKS_B

2b. Total number of puffs in blue DOSERTM history

   

   ➔ Value obtained from Question #2 on P17_COMPLY_WKS_B

2c. Percent compliance = \( \frac{\text{Question } \#2b}{\text{Question } \#2a} \times 100 \)

   

   ➔ *If the subject took less than 75% of the scheduled Inhaler B puffs, re-emphasize the importance of maintaining the daily dosing schedule.*

2d. Total number of full days since the last visit

   

   ➔ Value obtained from Question #4 on P17_COMPLY_WKS_B

2e. Total number of compliant days

   

   ➔ Value obtained from Question #5 on P17_COMPLY_WKS_B

2f. Percent compliance = \( \frac{\text{Question } \#2e}{\text{Question } \#2d} \times 100 \)

   

   ➔ *If the subject took the correct daily dose less than 75% of the days, re-emphasize the importance of maintaining the daily dosing schedule.*

3. **DOSERTM Compliance for Inhaler C**

*If the interval between visits exceeds 30 days, complete Questions #3a - #3f using data for the 30 days prior to the visit.*

3a. Total number of albuterol puffs in DOSERTM history

   

   ➔ Value obtained from Question #1 on P17_COMPLY_WKS_C

3b. Total number of puffs in red DOSERTM history

   

   ➔ Value obtained from Question #2 on P17_COMPLY_WKS_C
3d. Total number of full days since the last visit ___ ___ days (1140)
   ➔ Value obtained from Question #4 on P17_COMPLY_WKS_C

3e. Total number of compliant days ___ ___ days (1150)
   ➔ Value obtained from Question #5 on P17_COMPLY_WKS_C

3f. Percent compliance = $\frac{\text{Question } #3e}{\text{Question } #3d} \times 100$
   ___ ___ . ___ % (1160)
   ➔ If the subject took the correct daily dose less than 75% of the days, re-emphasize the importance of maintaining the daily dosing schedule.

4. Diary Card Compliance

   If the interval between visits exceeds 30 days, complete Questions #4a - #4e using data for all full days prior to the visit.

4a. Total number of full days since the last visit ___ ___ days (1170)

4b. Total number of days where all 3 measurements (AM PEFR, PM PEFR, and complete symptom score) have been recorded ___ ___ days (1180)

4c. Percent compliance = $\frac{\text{Question } #4b}{\text{Question } #4a} \times 100$
   ___ ___ . ___ % (1190)

4d. Total number of days where the subject measured both AM PEFR and PM PEFR on schedule and transcribed the measurements accurately on his/her diary cards ___ ___ days (1200)

4e. Percent compliance on schedule = $\frac{\text{Question } #4d}{\text{Question } #4a} \times 100$
   ___ ___ . ___ % (1210)
   ➔ If the percent compliance for either Question #4c or #4e is less than 75%, re-emphasize the importance of accurately and regularly completing diary cards.
**To the subject:** If your peak flow is below ________ liters/minute (65% of baseline), use your RESCUE albuterol inhaler as instructed in the handout “If Your Asthma Gets Worse.” Contact study personnel if your peak flow does not increase to ________ liters/minute (80% of baseline), after 1 hour of RESCUE use, or if you are experiencing extreme symptoms. If you have taken at least ________ puffs/24 hours for the past 48 hours from your RESCUE inhaler, contact study personnel.

Please use black ink to complete.

<table>
<thead>
<tr>
<th>Date (ddate)</th>
<th>Day 1:</th>
<th>Day 2:</th>
<th>Day 3:</th>
<th>Day 4:</th>
<th>Day 5:</th>
<th>Day 6:</th>
<th>Day 7:</th>
</tr>
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<tr>
<td>month</td>
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<td>day</td>
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</tr>
</tbody>
</table>

**MORNING EVALUATION (Between 5 AM and 10 AM)**

1. Number of times you woke up last night due to asthma (1010)

2. Time of AM Peak Flow (within 15 minutes of awakening) (1020)

3. AM Peak Flow (liters/min)** (1030)

4. Total number of puff(s) from yellow Inhaler A (AM) (1040)

5. Total number of puff(s) from blue Inhaler B (AM) (1045)

**Symptoms**

6. Shortness of Breath (AM) (1050)

7. Chest Tightness (1060)

8. Wheezing (1070)

9. Cough (1080)

10. Phlegm/Mucus (1090)

**NIGHT-TIME EVALUATION (Between 8 PM and 1 AM)**

11. Time of PM Peak Flow (between 8 PM and 1 AM) (1100)

12. PM Peak Flow (liters/min)** (1110)

13. Total number of puff(s) from yellow Inhaler A (PM) (1120)

14. Total number of puff(s) from blue Inhaler B (PM) (1125)

**Symptoms**

15. Shortness of Breath (PM) (1130)

16. Chest Tightness (1140)

17. Wheezing (1150)

18. Cough (1160)

19. Phlegm/Mucus (1170)

**24 HOUR EVALUATION**

20. Total number of puffs from red Inhaler C during the past 24 hours (1180)

21. Total number of puffs from albuterol (RESCUE) inhaler NOT including exercise preventive use, during the past 24 hours (1190)

22. Total number of puffs from albuterol (RESCUE) inhaler taken for exercise preventive use, during the past 24 hours (1195)

**++ Symptom Severity Rating Scale**

0 = Absent
No symptom

1 = Mild
Symptom was minimally troublesome, i.e. not sufficient to interfere with normal daily activity or sleep.

2 = Moderate
Symptom was sufficiently troublesome to interfere with normal daily activity or sleep.

3 = Severe
Symptom was so severe as to prevent normal activity and/or sleep.

**++ Record the best of three attempts. Circle the value if you have taken any medication from your RESCUE albuterol inhaler in the last 2 hours.**
1. This visit is a
   (Check one box only)
   
   ◯ Regular study visit
   ◯ Treatment Failure (TF) visit
   (Visits 90–99)  

(Clinic Coordinator completed)

(Completed at regular study and Treatment Failure visits)

2. Inhaler A adjustment
   
2a. Has inhaler A been adjusted within the last 4 weeks due to treatment failure (If Visit 4, answer No)?
   
   ➔ If YES, inhaler A should NOT be adjusted. Skip to Question #2e.
   
2b. Since the last adjustment of inhaler A, has the subject experienced a treatment failure for which an adjustment has not been considered/made (If Visit 4, answer No)?
   
   ➔ If YES, and the TF occurred within 4 weeks of an adjustment of inhaler A due to a previous TF, inhaler A should NOT be adjusted. Otherwise, increase inhaler A by one step. Skip to Question #2e.
   
2c. At this visit, is the subject’s FEV₁ < 85% of baseline? (*At Visit 4, baseline is the FEV₁ value obtained at Visit 3. After Visit 4, baseline is the FEV₁ value obtained at Visit 4.)
   
   ➔ If YES, increase inhaler A by one step. Skip to Question #2e.
   
2d. At this visit, did the subject answer the Asthma Evaluation Questionnaire (AEQ) as follows:
   
   i) 0 on each of the three questions
   
   ➔ If YES, decrease inhaler A by one step. Skip to Question #2e.
   
   ii) 1 on at least one question and 0’s or 1’s on other two questions
   
   ➔ If YES, inhaler A should NOT be adjusted. Skip to Question #2e.
   
   iii) 2 or 3 on at least one question
   
   ➔ If YES, increase inhaler A by one step.

2e. i) Dosing step for inhaler A prior to this visit
   
   ii) Dosing step for inhaler A after adjustment, if any, at this visit
(Completed at regular study visits only)

3. Inhaler B adjustment

3a. Record the average ENO value

   ➔ This value is the average of two reproducible values obtained at this visit.

3ai. Is Question #3a < 22.0 ppb?

   ➔ If YES, decrease inhaler B by one step.
   Skip to Question #3b.

3a(ii). Is Question #3a between 22.0 - 35.0 ppb (inclusive)?

   ➔ If YES, inhaler B should NOT be adjusted.
   Skip to Question #3b.

3a(iii). Is Question #3a > 35.0 ppb?

   ➔ If YES, increase inhaler B by one step.

3b. i) Dosing step for inhaler B prior to this visit

   __________________________ (1130)

ii) Dosing step for inhaler B after adjustment, if any, at this visit

   __________________________ (1140)
The dose adjustment was corrected...

1. (Check one box only)
   - On the phone
   - At the visit

2. The dose adjustment was corrected for ...
   - Inhaler A only
   - Inhaler B only
   - Inhalers A and B

3. Inhaler A
   a. Dosing step for inhaler A prior to the correction
      - Value obtained from Q1080 on the previous P17_DOSE_ADJUST form
   b. Dosing step for inhaler A after the correction
   c. Effective start date of the dose adjustment
      - month / day / year
   d. Were any additional inhaler A's dispensed?
      - Yes
      - No

4. If NO, and inhaler B needs to be corrected, skip to Question #4.
   If NO, and inhaler B does not need to be corrected, skip to Source Documentation.

5. Number of inhaler(s) dispensed

6. Box number from which the inhaler(s) has(have) been taken

7. Number of puffs written on the label
   - AM
   - PM

8. If adjusting inhaler A only, skip to Source Documentation.

Affix the box # label(s) for inhaler A below:
4. **Inhaler B**
   
4a. Dosing step for inhaler B prior to the correction
   
   \[\text{Value obtained from Q1140 on the previous P17_DOSE_ADJUST form}\]
   
4b. Dosing step for inhaler B after the correction
   
4c. Effective start date of the dose adjustment
   
   [Day] / [Month] / [Year] (1120)

4d. Were any additional inhaler B’s dispensed?

   ➔ **If NO, skip to Source Documentation.**

4e. Number of inhaler(s) dispensed
   
   [Number] (1140)

4f. Box number from which the inhaler(s) has(have) been taken
   
   [Box Number] (1150)

4g. Number of puffs written on the label

   (Must be same as on the Daily Activities handout for Inhaler B)
   
   [Number] AM (1160)

   [Number] PM (1170)

Affix the box # label(s) for inhaler B below:


Coordinator’s
Signature: ___________________ (1180)

Date: ___ / ___ / ___ ___ (1190)
(Clinic Coordinator completed)

1. Since Visit 3, has the subject experienced a treatment failure as defined in the protocol?
   
   ➔ If NO, skip to Question #2.

1a. Has the subject received oral, parenteral, inhaled corticosteroids, or another new asthma medication?
   
   ➔ If NO, and budesonide will be prescribed according to the protocol, Visit 4 should be rescheduled 4 weeks after the last dose.

1ai. If NO, and budesonide will not be prescribed, has at least 4 weeks passed since the treatment failure?
   
   ➔ If YES, skip to Question #2.
   
   ➔ If NO, Visit 4 should be rescheduled 4 weeks after the date of the event.

1aii. If YES, record the date of the subject’s last dose from the Concomitant Medications for Asthma/Allergy and Adverse Events (CMED) form. month / day / year

1aiii. Was the last dose taken at least 4 weeks prior to today’s date?
   
   ➔ If NO, Visit 4 should be rescheduled 4 weeks after the last dose.

2. Using the history stored in the yellow DOSER™ from Inhaler A, did the subject take at least 75% of the required puffs from his or her inhaler A during the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?
   
   ➔ Use Question #1c from P17_COMPLY to answer this question

3. Using the history stored in the yellow DOSER™ for Inhaler A, did the subject take 4 puffs per day (correct daily dose) on at least 75% of the days during the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?

   ➔ Use Question #1f from P17_COMPLY to answer this question

4. Using the history stored in the blue DOSER™ from Inhaler B, did the subject take at least 75% of the required puffs from his or her inhaler B during the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?

   ➔ Use Question #2c from P17_COMPLY to answer this question
5. Using the history stored in the blue DOSER™ for Inhaler B, did the subject take 4 puffs per day (correct daily dose) on at least 75% of the days during the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?

   Use Question #2f from P17_COMPLY to answer this question

6. Using the history stored in the red DOSER™ from Inhaler C and the DOSER™ from albuterol, did the subject take the same number of puffs from red inhaler C as from the albuterol inhaler on at least 75% of the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?

   Use Question #3f from P17_COMPLY to answer this question

7. Did the subject record both AM and PM peak flow measurements and symptoms on his or her Diary Cards (P17_DIARY) on at least 75% of the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?

   Use Question #4c from P17_COMPLY to answer this question

8. Did the subject measure his or her AM and PM peak flow on schedule and accurately transcribe the measurements on his or her Diary Cards (P17_DIARY) on at least 75% of the days during the 14 days prior to Visit 4 (only use the full days between Visit 3 and Visit 4, exclude visit dates)?

   Use Question #4e from P17_COMPLY to answer this question

9. Is the subject able to use a metered dose inhaler (MDI) properly, as evidenced by achieving a score of 6 on two consecutive, separate inhalations using the MDI Inhalation Technique Checklist (SCORE, TECH_MDI)?

10. Does the subject wish to withdraw consent from the study?

11. Is there any new information that makes the subject ineligible according to the eligibility criteria?

   If YES, describe: ______________________________

12. Is there any other reason why this subject should not be included in the study?

   If YES, describe: ______________________________

13. FEV₁ (% predicted) category at Visit 4:

   Refer to Question #2c on the SPIRO form
14. Is the subject eligible?

- **If any of the shaded boxes are completed, the subject is ineligible. If the subject is not eligible due to the compliance and this is the first compliance assessment at V4, reschedule Visit 4 in 2 weeks. Otherwise, complete the BASALT Termination of Study Participation (P17_TERM) form.**

- **If the subject is eligible and will participate in BASALT, randomize the subject.**

15. Inhalers Box Numbers *(record on P16_LOG):*

<table>
<thead>
<tr>
<th>Inhalers</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Step</td>
<td>1-4</td>
<td>5</td>
<td>1-4</td>
</tr>
<tr>
<td>Box #</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(1180)</td>
<td>(1190)</td>
<td>(1200)</td>
</tr>
</tbody>
</table>
This form should only be completed if the subject has been hospitalized during the BASALT Study. Obtain hospital discharge summary or abstract to complete this form. Complete Serious Adverse Event Reporting (SERIOUS) form.

DO NOT ENTER. FOR REFERENCE PURPOSES ONLY.

Hospital Name: ________________________________
Hospital Address: ________________________________

1. Admission date
   ___ ___ / ___ ___ / ___ ___ ___ (1000)

2. Discharge date
   ___ ___ / ___ ___ / ___ ___ ___ (1010)

3. Number of days in ICU/CCU/Stepdown Unit
   ___ ___ . ___ days(s) (1020)

4. Number of days in regular care unit
   ___ ___ . ___ days(s) (1030)

5. Was intubation or ventilation assistance required?
   □ 1 Yes  □ 0 No (1040)

6. Was the hospitalization due to...
   (Check one box only)
   ➔ If asthma related, complete BASALT Significant Exacerbation (P17_SIGEX) and BASALT Treatment Failure (P17_TF) forms.
   □ 1 Asthma
   □ 2 Other _________________ (1050)

7. What was the subject’s status at discharge?
   □ 1 Alive
   □ 2 Deceased (1060)
I am going to ask you some questions based on several events which may have occurred since your last study contact that took place on:

___ / ___ / ___

1. Since your last study contact, were you admitted to a hospital for an overnight stay of at least one night? ☐  Yes ☐ No (1000)
   ➔ If YES, how many times were you admitted due to...

1a. asthma-related events
   ➔ Complete the BASALT Hospitalization Summary Report (P17_HOSPITAL), BASALT Significant Exacerbation (P17_SIGEX), BASALT Treatment Failure (P17_TF) and Serious Adverse Event (SERIOUS) forms for each event.

1b. other (describe) ──────────────────────── ☐ ☐ time(s) (1020)
   ➔ Complete the BASALT Hospitalization Summary Report (P17_HOSPITAL) and Serious Adverse Event (SERIOUS) forms for each event.

2. Since your last study contact, did you go to an emergency room? ☐  Yes ☐ No (1030)
   ➔ If YES, how many times due to...

2a. asthma-related events
   ➔ Complete the BASALT Significant Exacerbation (P17_SIGEX) and BASALT Treatment Failure (P17_TF) forms for each event.

2b. other (describe) ──────────────────────── ☐ ☐ time(s) (1050)

3. Since your last study contact, did you go to an urgent care visit to a physician? ☐  Yes ☐ No (1060)
   ➔ If YES, how many times were the visits due to...

3a. asthma-related events
   ➔ Complete the BASALT Significant Exacerbation (P17_SIGEX) and BASALT Treatment Failure (P17_TF) forms for each event.

3b. other (describe) ──────────────────────── ☐ ☐ time(s) (1080)

DO NOT ENTER. FOR REFERENCE PURPOSES ONLY.
4. Since your last study contact, did you have a regular clinic/office visit to a physician (does not apply to study visits)?
   - If YES, how many times were the visits due to...
   4a. asthma-related events
   4b. other (describe)

5. Since your last study contact, did you miss at least a half-day of work, house work, or school because of your health (does not apply to time off for study visits)?
   - If YES, complete BASALT School/Work Absenteeism (P17_SWA) form.

6. Since your last study contact, were you prescribed any new medication?
   - If YES, record the new medication on the Concomitant Medications for Asthma/Allergy and Adverse Events (CMED) form.

7. Since your last study contact, did you purchase any over-the-counter (OTC) medication?
   - If YES, record the new medication on the Concomitant Medications for Asthma/Allergy and Adverse Events (CMED) form.
Complete this form only if the subject is eligible according to the appropriate Pulmonary Procedure Checklist form and successfully completed baseline spirometry session(s).

1. **(Complete at Visit 4A Only)** Did the subject consent to perform a mannitol challenge?
   - Yes [ ] No [□] 1000
   - If **YES**, record the date the consent form was signed.
   - ___ ___ / ___ ___ / ___ ___ ___ 1010

2. **(Complete at Visit 4A Only)** Has the subject performed a methacholine challenge at Visit 4?
   - Yes [ ] No [□] 1020
   - If **YES**, proceed to Question #2a.

2a. Did the subject require > 4 puffs of albuterol for reversal after Visit 4 methacholine challenge testing?
   - Yes [ ] No [□] 1030

3. **(Complete at Visits 8 and 10 Only)** Has the subject performed a mannitol challenge at Visit 4A?
   - Yes [ ] No [□] 1040

4. Has the subject had any severe acute illness in the past 4 weeks?
   - Yes [ ] No [□] 1050
   - If **YES**, has the subject received permission from the supervising physician to proceed with the mannitol challenge testing?
   - Yes [ ] No [□] 1060

   Physician’s Signature: ______________________ 1070

5. Does the subject have a baseline FEV₁ < 70% of predicted?
   - Yes [ ] No [□] 1080

6. Is there any other reason the subject should not proceed with the mannitol challenge?
   - Yes [ ] No [□] 1090
   - If **YES**, explain ______________________
7. Record the date and time (based on a 24 hour clock) the subject last took a dose from each inhaler:
   ➔ If the subject did not take a dose from an inhaler in the past 24 hours, check the box "Did not take" and do not complete the date and time fields.

7a. Inhaler A  
   Did not take  
   Date: ___/___/_____  
   Time: _______ (1100)

7b. Inhaler B  
   Did not take  
   Date: ___/___/_____  
   Time: _______ (1130)

7c. Inhaler C  
   Did not take  
   Date: ___/___/_____  
   Time: _______ (1160)

7d. Open label ICS  
   Did not take  
   Date: ___/___/_____  
   Time: _______ (1190)

8. Is the subject eligible to proceed with the reference (0 mg) spirometry testing for the mannitol challenge?  
   If any of the shaded boxes are completed, the subject is NOT eligible for the mannitol challenge.  
   ➔ If YES, proceed to the Mannitol Challenge Testing (MANNITOL) form.
1. Inhaler A
   1a. Number of inhaler(s) dispensed
      ➔ *If 0, skip to Question #2.*
      __ (1000)

   1b. Box number from which the inhaler(s) has(have) been taken
      __ (1010)

   1c. Number of puffs written on the label
      AM (1020)
      PM (1025)
      *(Must be same as on the Daily Activities handout for Inhalar A)*

      Affix the box # label(s) for inhaler A below:

      |   |   |   |   |
      ---|---|---|---|

2. Inhaler B
   2a. Number of inhaler(s) dispensed
      ➔ *If 0, skip to Question #3.*
      __ (1030)

   2b. Box number from which the inhaler(s) has(have) been taken
      __ (1040)

   2c. Number of puffs written on the label
      AM (1050)
      PM (1055)
      *(Must be same as on the Daily Activities handout for Inhaler B)*

      Affix the box # label(s) for inhaler B below:

      |   |   |   |   |
      ---|---|---|---|
3. **Inhaler C**
   
   3a. Number of inhaler(s) dispensed
   
   3b. Box number from which the inhaler(s) has (have) been taken

Affix the box # label(s) for inhaler C below:

---

Coordinator’s Signature: _____________________ (1080)
Date: __ __ / __ __ / __ __ __ __ (1090)

By signing in the source documentation box you are:

1) confirming that the same number of inhalers A, B, and C were dispensed as indicated on this form.
2) confirming that the box number from which each set of inhalers were dispensed is correctly written on this form.
3) confirming that the box number from which each set of inhalers were dispensed corresponds to the box numbers for each set as indicated on the subject’s randomization report.
4) confirming that the dosing instructions for inhalers A and B are written on the labels as indicated on this form.
5) confirming that the dosing instruction label has been attached to inhaler C.
(Subject Interview completed)

Please reference the Drug Classifications list for a complete list of examples for the questions below. If any medications other than study inhalers or rescue albuterol were used, record the medication(s) on the Concomitant Medications for Asthma/Allergies and Adverse Events (CMED) form.

1. Have you consumed caffeine in the past 6 hours?  
   Examples: Pepsi, Coke, Coffee, Mountain Dew, Tea, Rootbeer, Red Bull

2. Have you used medications with caffeine in the past 6 hours?  
   Examples: Anacin, Darvon compound, Esgic, Excedrin, Fiorinal, Fioricet, No Doz, Norgesic, Vivarin

3. Have you used any weight loss medications in the past 6 hours?  
   Examples: bitter orange, Xenadrine EFX, Thermorexin

4. Have you consumed any food containing alcohol or beverages containing alcohol in the past 6 hours?

5. Have you used any oral antihistamines in the past 48 hours?  
   Examples: Allegra, Chlor-Trimeton, Claritin, Tylenol PM

6. Have you used any nasal antihistamines in the past 6 hours?  
   Examples: Astelin, Livostin, Patanase

7. Have you used any oral decongestants or cold remedies in the past 48 hours?  
   Examples: pseudoephedrine (Sudafed), Tylenol Allergy

8. Have you used any nasal decongestants in the past 6 hours?  
   Examples: oxymetazoline (Afrin)

9. Have you used any cough medicines, anti-tussives, or expectorants in the past 48 hours?  
   Examples: guaifenesin, dextromethorphan, Duratuss, Benylin, Triaminic expectorant, Dayquil Anti-Cough

10. Have you used a rescue intermediate-acting inhaled beta-agonist in the past 6 hours?  
    Example: albuterol (Ventolin or Proventil), study RESCUE

11. Have you used any nasal steroids in the past 48 hours?  
    Examples: Flonase, Rhinocort, Nasonex
12. Have you used any smokeless tobacco products today?
   Examples: chewing tobacco, snuff  
   □ 1 Yes □ 0 No (1095)

13. At this time, is your asthma worse because of recent exposure to triggers?
   Examples: cold air, smoke, allergens, recent exercise, a recent respiratory tract infection, or other pulmonary infection  
   □ 1 Yes □ 0 No (1100)

14. Is there any other reason you should not proceed with spirometry testing? 
   If YES, explain ____________________________________________
   ____________________________________________  
   □ 1 Yes □ 0 No (1110)

15. Is the subject eligible to proceed with the spirometry testing? 
   If any of the shaded boxes are filled in, the subject is ineligible for spirometry and exhaled nitric oxide testing. 
   ➔ If YES, proceed to Question #16 or the next form/procedure listed on the visit procedure checklist.
   □ 1 Yes □ 0 No (1120)

If subject is less than 21 years old, complete Question #16 at each visit.

16. Height (without shoes) _______ ___ ___ cm (1130)
This form must be completed each time a subject experiences an asthma exacerbation according to the definition below.

1. Did the subject experience an increase in symptoms of cough, phlegm/mucus, chest tightness, wheezing, or shortness of breath along with any of the following conditions?

   1a. An increase in rescue inhaler use of > 8 puffs per 24 hours over baseline rescue inhaler use for a period of 48 hours?  
       □ 1 Yes □ 0 No (1000)

   1b. Use of rescue inhaler > 16 total puffs per 24 hours for a period of 48 hours?  
       □ 1 Yes □ 0 No (1010)

   1c. A fall in pre-bronchodilator PEFR to < 65% of baseline on 2 consecutive scheduled morning measurements?  
       □ 1 Yes □ 0 No (1020)

   1d. A fall in post-bronchodilator PEFR (any time of day) to < 80% of baseline despite 60 minutes of rescue beta-agonist treatment (≥ 6 puffs of albuterol in one hour)?  
       □ 1 Yes □ 0 No □ 9 N/A (1030)

       1di. If YES, record the post-bronchodilator PEFR value that qualified the subject as a treatment failure.  

       __ __ __ L/min (1040)

   1e. A fall in pre-bronchodilator FEV₁ to < 80% of baseline?  
       □ 1 Yes □ 0 No □ 9 N/A (1060)

2. Did the subject require an urgent medical care visit for asthma in an office setting or emergency room?  
   □ 1 Yes □ 0 No (1060)

3. Were systemic corticosteroids (oral, IM, or IV), given to the subject for his/her asthma as a result of rescue intervention or by the opinion of the treating physician?  
   ➔ If YES, please complete the CMED form.

4. Was the subject hospitalized for his/her asthma?  
   □ 1 Yes □ 0 No (1080)

5. Did the subject experience a significant asthma exacerbation?  
   If any of the shaded boxes are completed, the subject experienced a significant asthma exacerbation.  
   ➔ If NO, STOP HERE. DO NOT SUBMIT THIS FORM TO THE DCC.  
   ➔ If YES, complete the rest of this form. Also, complete BASALT Treatment Failure Assessment (P17_TF) form.
6. Date significant asthma exacerbation occurred

7. Did the subject seek care for the asthma exacerbation?
   If NO, skip to Question #14.

8. What type of care was sought?
   8a. Study Investigator or Clinic Coordinator?
       If YES, indicate type of contact.

   8b. Primary Care or Other Physician?
       Name of physician: ______________________
       If YES, indicate type of contact.

   8c. Emergency Room visit?
       Name of hospital: ______________________

9. Was spirometry performed at the visit?
10. Was peak flow measured at the visit?
11. Were any treatments given during visit?
   If NO, skip to Question #12.
   If YES, please complete the CMED form, if needed.
11a. Nebulizer (i.e., breathing) treatment
11b. IM steroids
11c. IV steroids
11d. IV aminophylline
11e. Other ______________________

11f. Other ______________________
12. Were any medications prescribed at discharge?
   ➔ If NO, skip to Question #13.
   ➔ If YES, please complete the CMED form, if needed.

12a. Oral steroids
   ➔
   1 Yes
   0 No (1250)

12b. Antibiotics
   ➔
   1 Yes
   0 No (1270)

13. Was the subject hospitalized?
   ➔ If YES, please complete Serious Adverse Event Reporting (SERIOUS) and BASALT Hospitalization Summary Report (P17_HOSPITAL) Forms.

14. Was the asthma exacerbation resolved solely by increasing PRN use of the rescue inhaler?
   ➔
   1 Yes
   0 No (1290)

15. Was the asthma exacerbation treated as outlined in the protocol?
   If NO, describe

16. Was budesonide prescribed for management of the significant exacerbation?
   ➔ If YES, please complete CMED form.

17. Was the significant asthma exacerbation related to the routine pulmonary function testing, including the collection of exhaled nitric oxide? (Check one box only)
   1 Definitely related
   2 Probably related
   3 Relationship undetermined
   4 Probably not related
   5 Definitely not related (1310)

18. Was the significant asthma exacerbation related to the methacholine challenge testing? (Check one box only)
   1 Definitely related
   2 Probably related
   3 Relationship undetermined
   4 Probably not related
   5 Definitely not related (1320)
<table>
<thead>
<tr>
<th>Question</th>
<th>Options</th>
</tr>
</thead>
</table>
| 19. Was the significant asthma exacerbation related to the sputum induction procedure? *(Check one box only)* | 1. Definitely related  
2. Probably related  
3. Relationship undetermined  
4. Probably not related  
5. Definitely not related *(1330)* |
| 20. Was the significant asthma exacerbation related to the collection of exhaled breath condensate? *(Check one box only)* | 1. Definitely related  
2. Probably related  
3. Relationship undetermined  
4. Probably not related  
5. Definitely not related *(1340)* |
This questionnaire is to be completed by the BASALT subject at the end of Visit 12. If a randomized subject terminates prior to Visit 12, please ask him or her to complete this form during the termination visit.

1. Scheduled Inhalers (1000)/(1010)

As a BASALT study participant you were randomized to receive three inhalers A, B, and C where one was an active (i.e. real) inhaled steroid inhaler and two were a look-alike placebo (i.e. inactive) inhalers. Please choose the statement that more closely represents you feelings about the scheduled inhalers.

(Choose one statement that represents your feelings and choose one set of inhalers for the statement.)

☐ 1 I am certain the inhaler that contained active drug is:
    ☐ 1 Inhaler A
    ☐ 2 Inhaler B
    ☐ 3 Inhaler C

☐ 2 I think the inhaler that contained active drug is:
    ☐ 1 Inhaler A
    ☐ 2 Inhaler B
    ☐ 3 Inhaler C

☐ 3 I have no idea which inhaler contained active drug however my best guess would be:
    ☐ 1 Inhaler A
    ☐ 2 Inhaler B
    ☐ 3 Inhaler C
2. Please comment with respect to the taste of the medication you received from your scheduled...

<table>
<thead>
<tr>
<th>Inhaler A (1020)</th>
<th>Inhaler B (1030)</th>
<th>Inhaler C (1040)</th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ 1 Tasted good</td>
<td>☐ 1 Tasted good</td>
<td>☐ 1 Tasted good</td>
</tr>
<tr>
<td>(Describe)</td>
<td>(Describe)</td>
<td>(Describe)</td>
</tr>
<tr>
<td>☐ 2 No noticeable taste</td>
<td>☐ 2 No noticeable taste</td>
<td>☐ 2 No noticeable taste</td>
</tr>
<tr>
<td>☐ 3 Tasted bad</td>
<td>☐ 3 Tasted bad</td>
<td>☐ 3 Tasted bad</td>
</tr>
<tr>
<td>(Describe)</td>
<td>(Describe)</td>
<td>(Describe)</td>
</tr>
</tbody>
</table>

3. Please comment with respect to the smell of the medication you received from your scheduled...

<table>
<thead>
<tr>
<th>Inhaler A (1050)</th>
<th>Inhaler B (1060)</th>
<th>Inhaler C (1070)</th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ 1 Smelled good</td>
<td>☐ 1 Smelled good</td>
<td>☐ 1 Smelled good</td>
</tr>
<tr>
<td>(Describe)</td>
<td>(Describe)</td>
<td>(Describe)</td>
</tr>
<tr>
<td>☐ 2 No noticeable smell</td>
<td>☐ 2 No noticeable smell</td>
<td>☐ 2 No noticeable smell</td>
</tr>
<tr>
<td>☐ 3 Smelled bad</td>
<td>☐ 3 Smelled bad</td>
<td>☐ 3 Smelled bad</td>
</tr>
<tr>
<td>(Describe)</td>
<td>(Describe)</td>
<td>(Describe)</td>
</tr>
</tbody>
</table>

4. Please comment with respect to the physical sensations of the medication you received from your scheduled...

<table>
<thead>
<tr>
<th>Inhaler A (1080)</th>
<th>Inhaler B (1090)</th>
<th>Inhaler C (1100)</th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ 1 Pleasant sensations</td>
<td>☐ 1 Pleasant sensations</td>
<td>☐ 1 Pleasant sensations</td>
</tr>
<tr>
<td>(Describe)</td>
<td>(Describe)</td>
<td>(Describe)</td>
</tr>
<tr>
<td>☐ 2 No noticeable sensations</td>
<td>☐ 2 No noticeable sensations</td>
<td>☐ 2 No noticeable sensations</td>
</tr>
<tr>
<td>☐ 3 Unpleasant sensations</td>
<td>☐ 3 Unpleasant sensations</td>
<td>☐ 3 Unpleasant sensations</td>
</tr>
<tr>
<td>(Describe)</td>
<td>(Describe)</td>
<td>(Describe)</td>
</tr>
</tbody>
</table>
5. Please comment with respect to any other observations you have made regarding your scheduled...

<table>
<thead>
<tr>
<th>Inhaler A (1110)</th>
<th>Inhaler B (1120)</th>
<th>Inhaler C (1130)</th>
</tr>
</thead>
<tbody>
<tr>
<td>☐ 1 I have no further comments</td>
<td>☐ 1 I have no further comments</td>
<td>☐ 1 I have no further comments</td>
</tr>
<tr>
<td>☐ 2 I observed the following:</td>
<td>☐ 2 I observed the following:</td>
<td>☐ 2 I observed the following:</td>
</tr>
<tr>
<td><em>(Describe below)</em></td>
<td><em>(Describe below)</em></td>
<td><em>(Describe below)</em></td>
</tr>
</tbody>
</table>

Subject Source Documentation
Subject’s Initials: ___ ___ (1140)
Date: ___ / ___ / ___ ___ (1150)
1. How many full or half days of school/work/housework did the subject miss because of his/her health? (do not count time missed due to study visits) (Indicate full or half days in increments of 0.5 days)

IF LESS THAN 0.5, STOP HERE. DO NOT SUBMIT THIS FORM TO THE DCC.

2. Primary activity missed. (Check one box only)
   - Work
   - School
   - Housework

3. Was the activity missed due to...
   (Check one box only)
   - Asthma
   - Other ____________________

If asthma, was it...
3a. due to worsening symptoms caused by your asthma?
   If YES, please complete Clinical and Laboratory Adverse Events (AECLIN) form.
   - Yes
   - No

3b. to see an MD or health-care provider about your asthma (does not apply to time off for study-related visits)?
   - Yes
   - No

3c. due to side effects related to asthma medication?
   If YES, please complete Clinical and Laboratory Adverse Events (AECLIN) form.
   - Yes
   - No

3d. Other ________________________________
   - Yes
   - No
(Clinic Coordinator completed)

Complete this form only for those subjects who have been allocated to BASALT study and have been terminated or deemed ineligible.

1. Has the subject completed the study through Visit 12?  
   ➔ If YES, skip to the SIGNATURES section.
   - 1 Yes  
   - 0 No  

2. Who initiated termination of the subject?  
   ➔ If subject withdrew due to impending clinical staff termination, please indicate termination by clinical staff.
   ➔ If Clinical Staff, skip to Question #4.
   - 1 Subject
   - 2 Clinical Staff

3. Indicate the primary reason the subject has withdrawn from the study.
   - 1 no longer interested in participating*
   - 2 no longer willing to follow protocol*
   - 3 difficult access to clinic (location, transportation, parking)
   - 4 unable to make visits during clinic hours
   - 5 moving out of the area
   - 6 unable to continue due to personal constraints*
   - 7 unable to continue due to medical condition unrelated to asthma*
   - 8 side effects of study medications*
   - 9 dissatisfied with asthma control*
   - 10 other  

* Additional explanation required:

__________________________________________________________________________________________

__________________________________________________________________________________________

➔ Skip to the SIGNATURES section.
4. Did clinical staff terminate the subject due to ...

4a. pregnancy? (Check N/A if the subject is male.)
    ☐ 1 Yes ☐ 0 No ☐ 9 N/A

4b. ineligibility during the adherence period (Visits 3-4) * other than compliance
    ☐ 1 Yes ☐ 0 No

4c. loss to follow-up? *
    ☐ 1 Yes ☐ 0 No

4d. an asthma-related adverse event? *
    ☐ 1 Yes ☐ 0 No

4e. a medication-related adverse event? *
    ☐ 1 Yes ☐ 0 No

4f. an adverse event not related to asthma or medications? *
    ☐ 1 Yes ☐ 0 No

4g. non-compliance with dosing of inhalers? *
    ☐ 1 Yes ☐ 0 No

4h. non-compliance with diary completion? *
    ☐ 1 Yes ☐ 0 No

4i. non-compliance with visit attendance? *
    ☐ 1 Yes ☐ 0 No

4j. non-compliance with peak flow monitoring? *
    ☐ 1 Yes ☐ 0 No

4k. subject experienced 3 treatment failures that required prednisone treatments?
    ☐ 1 Yes ☐ 0 No

4l. allocated to BASALT in error
    ☐ 1 Yes ☐ 0 No

4m. other reason? *
    ☐ 1 Yes ☐ 0 No

* If YES, additional explanation required:

________________________________________

________________________________________

4n. Indicate the letter corresponding to the primary reason the subject was terminated. ___ (1170)

SIGNATURES

* Please complete the following section regardless of the reason for termination of study participation.

I verify that all information collected on the ACRN BASALT data collection forms for this subject is correct to the best of my knowledge and was collected in accordance with the procedures outlined in the ACRN BASALT Protocol.

________________________________________
Clinic Coordinator Signature

___ / ___ / ___ (1180)

month day year

________________________________________
ACRN Investigator Signature

___ / ___ / ___ (1200)

month day year
BASALT TREATMENT FAILURE ASSESSMENT

Subject ID: - - -
Subject Initials: -
Visit Number: -
Visit Date: / / Year

Coordinator ID: -

(Clinic Coordinator completed)

1. (Visits 90-99 Only) Complete the number of the last regular visit completed

2. Did the subject experience any of the following conditions WITHOUT increase in symptoms (e.g., cough, phlegm/mucus, chest tightness, wheezing, or shortness of breath)?
   - An increase in rescue inhaler use of > 8 puffs per 24 hours over baseline rescue inhaler use for a period of 48 hours?
   - Use of rescue inhaler > 16 total puffs per 24 hours for a period of 48 hours?
   - A fall in pre-bronchodilator AM PEFR to < 65% of baseline on two consecutive scheduled morning measurements?
   - A fall in post-bronchodilator PEFR (any time of day) to 80% of baseline despite 60 minutes of rescue beta-agonist treatment (≥ 6 puffs of albuterol in one hour)?
     - If YES, record the post-bronchodilator PEFR value that qualified the subject as a treatment failure.
   - A fall in pre-bronchodilator FEV₁ to < 80% of baseline on two consecutive spirometric determinations measured 24-72 hours apart?

3. Did the subject experience a significant asthma exacerbation?
   - If YES, please complete the BASALT Significant Asthma Exacerbation (P17_SIGEX) form.

4. Were open-label inhaled corticosteroids or another (non-systemic corticosteroid) new asthma medication (e.g., montelukast) given to the subject for his/her asthma as a result of rescue intervention or by the opinion of the treating physician?
   - If YES, please complete the CMED form.

5. Based on clinical judgement, did the physician deem this subject a treatment failure for safety reasons?

6. Was the subject dissatisfied with asthma control achieved with study regimen? (Check N/A if it is not the subject’s last visit.)

7. Is the subject a treatment failure? If any of the shaded boxes are completed, the subject is a treatment failure.
   - If YES, please complete the rest of this form. Also, complete the BASALT Healthcare Utilization Review (P17_HUR) form.
   - If NO, STOP HERE and continue with remaining visit procedures.
8. Date treatment failure conditions were met

(Physician completed)

9. From a clinical perspective, would you have considered this subject to be a 'treatment failure' if he/she were not participating in BASALT trial and, instead, you were seeing him/her in your outpatient clinic?

10. Based on the subject’s clinical status at the time he/she met one of the treatment failure criteria, when do you think the subject reached this status?

11. What was the subject’s opinion of his/her asthma at the time he/she reached treatment failure?

12. Based on your experience with this subject, are you satisfied with the BASALT treatment failure criteria?

If NO, explain _____________________________________________________________

__________________________________________________________

13. Physician Narrative Assessment

__________________________________________________________

__________________________________________________________

__________________________________________________________

(1150)

Physician Source Documentation

Physician’s signature: ________________________________ (1160)

Date: ___ / ___ / ___ ___ (1170)

Time: ___ ___ ___ (based on 24-hour clock) (1180)
Complete Questions #14 - #22 only for subjects who did not meet Treatment Failure status by experiencing a significant asthma exacerbation (i.e., Question #3 is No).

14. Did the subject seek care for treatment failure conditions?
   ➔ If NO, skip to Question #17.

15. Did the subject require an urgent medical care visit for asthma in an office setting or emergency room?
   ➔ If YES, STOP HERE. Complete the BASALT Significant Asthma Exacerbation (P17_SIGEX) form and continue according to the protocol.

16. What type of care was sought?
   16a. Study Investigator or Clinic Coordinator?

   If YES, indicate type of contact.

   1 Yes
   0 No (1190)

   1 Scheduled clinic visit
   2 Unscheduled clinic visit
   3 Scheduled phone contact
   4 Unscheduled phone contact (1220)

   16b. Primary Care or Other Physician?

   Name of physician: ________________________________

   If YES, indicate type of contact.

   1 Yes
   0 No (1230)

   1 Scheduled clinic visit
   2 Unscheduled clinic visit
   3 Scheduled phone contact
   4 Unscheduled phone contact (1240)

17. Were the subject’s treatment failure conditions treated as outlined in the protocol?

   If NO, describe ________________________________

   ________________________________

18. Was budesonide prescribed for management of the treatment failure?

   ➔ If YES, please complete the CMED form.

   1 Yes
   0 No (1250)

   1 Yes
   0 No (1255)
<table>
<thead>
<tr>
<th>Question</th>
<th>Options</th>
</tr>
</thead>
</table>
| 19. Was the treatment failure related to the routine pulmonary function testing, including the collection of exhaled nitric oxide? (Check one box only) | 1  Definitely related  
2  Probably related  
3  Relationship undetermined  
4  Probably not related  
5  Definitely not related (1260) |
| 20. Was the treatment failure related to the methacholine challenge testing? (Check one box only) | 1  Definitely related  
2  Probably related  
3  Relationship undetermined  
4  Probably not related  
5  Definitely not related (1270) |
| 21. Was the treatment failure related to the sputum induction procedure? (Check one box only) | 1  Definitely related  
2  Probably related  
3  Relationship undetermined  
4  Probably not related  
5  Definitely not related (1280) |
| 22. Was the treatment failure related to the collection of exhaled breath condensate? (Check one box only) | 1  Definitely related  
2  Probably related  
3  Relationship undetermined  
4  Probably not related  
5  Definitely not related (1290) |
Complete this form only if the subject is eligible according to the appropriate Pulmonary Procedure Checklist form and successfully completed baseline spirometry session(s).

Administer 4 puffs of albuterol and wait 15 minutes, then perform spirometry.

1. Time albuterol administered (based on 24-hour clock) ____________ (1000)

2. Time post-albuterol spirometry started (based on 24-hour clock) ____________ (1010)

The best effort reflects the trial where the sum of FEV₁ and FVC is maximized.

3. Results of best effort post-albuterol:
   3a. FVC ____________ L (1020)
   3b. FEV₁ ____________ L (1030)
   3c. FEV₁ (% predicted) ____________ % predicted (1040)
   3d. PEFR ____________ L/S (1050)
   3e. FEF₂₅⁻⁷₅ ____________ L/S (1060)

4. In your judgment, was the subject’s spirometry technique acceptable? □ 1 Yes □ 0 No (1070)
   4a. If NO, why was it unacceptable?
      Inadequate inspiratory effort □ 1 Yes □ 0 No (1080)
      Inadequate expiratory effort □ 1 Yes □ 0 No (1090)
      Inadequate duration of expiration □ 1 Yes □ 0 No (1100)
      Cough during procedures □ 1 Yes □ 0 No (1110)
      Other (specify) ___________________________ □ 1 Yes □ 0 No (1120)
Complete this form only if the subject is eligible according to the appropriate Pulmonary Procedure Checklist form and successfully completed baseline spirometry session(s).

Note: Ipratropium should NOT be administered to subjects who have a hypersensitivity/allergy to soy or peanuts.

Administer 4 puffs of ipratropium and wait 30 minutes, then perform spirometry.

1. Time ipratropium administered (based on 24-hour clock) — — — — (1000)
2. Time post-ipratropium spirometry started (based on 24-hour clock) — — — — (1010)

The best effort reflects the trial where the sum of FEV₁ and FVC is maximized.

3. Results of best effort post-ipratropium:
   3a. FVC — — — — L (1020)
   3b. FEV₁ — — — — L (1030)
   3c. FEV₁ (% predicted) — — — — % predicted (1040)
   3d. PEFR — — — — L/S (1050)
   3e. FEF₂₅₋₇₅ — — — — L/S (1060)

4. In your judgment, was the subject’s spirometry technique acceptable?
   4a. If NO, why was it unacceptable?
      □ 1 Yes □ 0 No (1070)
      Inadequate inspiratory effort
      □ 1 Yes □ 0 No (1080)
      Inadequate expiratory effort
      □ 1 Yes □ 0 No (1090)
      Inadequate duration of expiration
      □ 1 Yes □ 0 No (1100)
      Cough during procedures
      □ 1 Yes □ 0 No (1110)
      Other (specify) ______________________
      □ 1 Yes □ 0 No (1120)
Complete this form for female subjects only.

1. Is the subject unable to bear children due to any of the following reasons?
   
   1a. Post-menopausal (at least one year since last menses) [ ] Yes [ ] No (1000)
   
   1b. Hysterectomy [ ] Yes [ ] No (1010)
   
   1c. Tubal ligation [ ] Yes [ ] No (1020)
   
   ➔ *If any of the shaded boxes are filled in, a pregnancy test is not required. Proceed to the source documentation box.*

2. Pregnancy test results
   
   ➔ *If pregnancy test results are positive, the subject must be terminated from study participation. Complete the appropriate Termination of Study Participation form and follow study termination procedures.*
   
   [ ] Positive [ ] Negative (1030)

---

Subject Source Documentation
Subject’s Initials: __ __ __ (1040)
Date: __ __ / __ __ / __ __ __ __ (1050)
Search the ACRN Registry. If the subject is either incomplete or not found in the Registry, complete the Registry form and enter/update the subject’s information appropriately.

ADMINISTRATIVE

1. Did the subject sign an ACRN Protocol Informed Consent and HIPAA Authorization form? □ 1 Yes □ 0 No (1000)
   If NO, stop here. Data cannot be entered into the ACRN Registry.
   If YES, record the signature date. □ 1 Male □ 2 Female (1030)

DEMOGRAPHICS

2. Subject’s date of birth (Ask the subject his/her date of birth.) □ 1 Hispanic or Latino □ 2 Not Hispanic or Latino (1040)
3. Subject’s gender □ 1 Male □ 2 Female (1030)
4. Subject’s Race and Ethnicity
   4a. Subject’s ethnic background (Ask the subject to identify his/her ethnic background.) □ 1 Yes □ 0 No (1050)
   American Indian or Alaskan Native □ 1 Yes □ 0 No (1060)
   Asian □ 1 Yes □ 0 No (1070)
   Black or African American □ 1 Yes □ 0 No (1080)
   White □ 1 Yes □ 0 No (1090)
   Native Hawaiian or Other Pacific Islander □ 1 Yes □ 0 No (1100)
   Other (specify) _____________________________
REGISTRY

Subject’s Initials: ___ ___ ___

5. Subject’s primary racial identification
   (This identification will be used for spirometry testing. Ask the subject which category best describes him or her and check only one box.)
   □ 1 American Indian or Alaskan Native
   □ 2 Asian or Pacific Islander
   □ 3 Black, not of Hispanic Origin
   □ 4 White, not of Hispanic Origin
   □ 5 Hispanic
   □ 6 Other ________________ (110)

Registry Form Storage Instructions:
Upon printing the subject’s label sheet, print the subject’s name on the upper right hand label. Attach the Registry form label to the upper left hand corner of the form. Lastly, attach the Registry Log label to the next available row on the Registry Log and complete the required fields. The Registry form should be stored alphabetically by subject’s last name in the ACRN Registry Binder. The label sheet should then be filed directly behind the Registry form.

REGISTRY FORMS SHOULD NOT BE SENT TO THE DCC.
ABOUT YOUR SLEEP

The questions below apply to the last 6 months.

Some people work the night shift or rotating shifts. Other people have a bedtime that changes a lot. If these apply to you, then questions about the time of day refer to the time when you awaken from your longest sleep and become active. Questions about time of night refer to the time when you have your longest sleep.

Please check only one box for each question.

<table>
<thead>
<tr>
<th>Question</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Usually</th>
<th>Always</th>
</tr>
</thead>
<tbody>
<tr>
<td>I am told I snore loudly and bother others.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>I am told I stop breathing (&quot;hold my breath&quot;) in sleep.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>I awake suddenly gasping for breath, unable to breathe.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>If this never happens to you, please skip to question #4.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>If this happens to you even rarely, please answer questions #3a and 3b.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>It takes just a couple of breaths to fully recover.</td>
<td>□ Yes</td>
<td>□ No</td>
<td>(1030)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>It happens when I have chest tightness, wheezing, or cough and takes more than a couple of breaths to fully recover.</td>
<td>□ Yes</td>
<td>□ No</td>
<td>(1040)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>I sweat a great deal at night.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>I have high blood pressure (or once had it).</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>I have a problem with my nose blocking up when I am trying to sleep.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>My snoring or my breathing problem is much worse if I sleep on my back.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
<tr>
<td>My snoring or my breathing problem is much worse if I fall asleep right after drinking alcohol.</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
<td>□</td>
</tr>
</tbody>
</table>
ABOUT YOUR DAYTIME ALERTNESS

9. Do you feel that you are excessively (overly) sleepy during the day? ☐ Yes ☐ No (1100)

*How likely are you to doze off or fall asleep in the following situations, in contrast to feeling just tired? This refers to your usual way of life in recent times. Even if you have not done some of these things recently, try to work out how they would have affected you.*

Please check one box that best represents the likelihood of your dozing off in each situation.

<table>
<thead>
<tr>
<th>Situation</th>
<th>Never</th>
<th>Slight</th>
<th>Moderate</th>
<th>High</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sitting and reading</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>Watching TV</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>Sitting, inactive in a public place (for example, a theater or a meeting)</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>As a passenger in a car for an hour without a break</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>Lying down to rest in the afternoon when circumstances permit</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>Sitting and talking to someone</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>Sitting quietly after a lunch without alcohol</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>In a car, while stopped for a few minutes in traffic</td>
<td>☐ 0</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>During the past 2 months, on average, how many hours of actual sleep (including daytime naps) did you get in a 24-hour period? This may be different than the number of hours you spent in bed.</td>
<td>____ ____ hours (1200)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CURRENT WEIGHT

19. What is your current weight in pounds?

_____ _____ pounds (1205)
SERIOUS ADVERSE EVENT REPORTING FORM

(Clinic Coordinator completed)

This form and a final resolution report (including relevant documents) written by the Principal Investigator should be faxed to the DCC at (717) 531-4359 within 72 hours of notification of a serious event. Also fax the Clinical Adverse Events form (AECLIN), the appropriate Concomitant Medications for Asthma and Allergies (CMED) form, and any relevant source documents.

1. Date of Adverse Event

2. Description of Adverse Event (ICD9 Code)
   Describe: ______________________________

3. Time interval between taking the study drug (last dose before symptoms) and subsequent onset of symptoms.

4. Unit of time for above interval
   - 1 second(s)
   - 2 minute(s)
   - 3 hour(s)
   - 4 day(s)

5. Why was the event serious?
   5a. Fatal Event
   - 1 Yes
   - 0 No
   5b. Life-threatening event
   - 1 Yes
   - 0 No
   5c. Inpatient hospitalization required
   → If NO, skip to Question #5d.
   Admission date
   Discharge date
   5d. Hospitalization prolonged
   - 1 Yes
   - 0 No
   5e. Disabling or incapacitating
   - 1 Yes
   - 0 No
   5f. Overdose
   - 1 Yes
   - 0 No
   5g. Cancer
   - 1 Yes
   - 0 No
   5h. Congenital anomaly
   - 1 Yes
   - 0 No
   5i. Serious laboratory abnormality with clinical symptoms
   - 1 Yes
   - 0 No
   5j. Other (specify) ______________________________
6. What, in your opinion, caused the event?

   6a. Toxicity of study drug(s) □ 1 Yes □ 0 No (1160)

   6b. Withdrawal of study drug(s) □ 1 Yes □ 0 No (1170)

   6c. Concurrent medication
       If YES, describe ____________________________ □ 1 Yes □ 0 No (1180)

   6d. Concurrent disorder
       If YES, describe ____________________________ □ 1 Yes □ 0 No (1190)

   6e. Other event
       If YES, describe ____________________________ □ 1 Yes □ 0 No (1200)

DO NOT ENTER QUESTIONS #7 - 8: FOR REPORTING PURPOSES ONLY.

7. If subject died, cause of death: ________________________________
   _____________________________________________________________

8. Was an autopsy performed? □ 1 Yes □ 0 No

   If YES, attach report or send as soon as possible.

REPORTING INVESTIGATOR:
Comments (discuss any relevant laboratory data or other assessments which help explain the event):

Name: ________________________________

Signature: ______________________________

Date: __ __/ __ __/ __ __ __ __
**SYMPTOM-FREE DAY QUESTIONNAIRE**

(Subject Interview completed)

1. In the **past 14 days**, how many days did you have wheezing, chest tightness, cough, or shortness of breath?  
   ___ ___ day(s)  

2. In the **past 14 days**, how many days did you have to slow down or stop activities because of asthma, wheezing, chest tightness, cough, or shortness of breath?  
   ___ ___ day(s)  

3. In the **past 14 days**, how many nights did you wake up because of asthma, wheezing, chest tightness, cough, or shortness of breath?  
   ___ ___ day(s)  

4. Thinking about all three asthma signs or symptoms (wheezing, chest tightness, cough, or shortness of breath; slowing down or stopping activities; nights awakened), in the **past 14 days**, how many days did you have any of these day-time or night-time symptoms?  
   ___ ___ day(s)  

5. In the **past 14 days**, how many days did you experience any day with NO day-time and night-time symptoms of asthma (including no wheezing, no cough, no chest tightness, or no shortness of breath)?  
   ___ ___ day(s)  

---

**Subject Source Documentation**

Subject’s Initials:  

Date:  

Time:  (based on a 24-hour clock)
1. Since August 2004, has the subject had an acceptable skin test for an ACRN protocol within three years of the visit date?

- If **NO**, proceed to Question #2.

   1a. Date of previous skin test
   - Month ___ / Day ___ / Year ___

   1b. Coordinator ID who performed the skin test
   - ___ ___ ___ ___ ___ ___ ___ ___

   1c. Time test sites pricked/punctured *(based on 24-hour clock)*
   - ___ ___ ___ ___ ___ ___ ___ ___

   1d. Time test sites evaluated *(based on 24-hour clock)*
   - ___ ___ ___ ___ ___ ___ ___ ___

   ➔ **STOP HERE** and attach a photocopy of pages 3 and 4 from the previous Allergy Skin Test Results (SKIN) form to this page for data entry purposes.

2. Has the subject had dermatographia **or** a significant adverse reaction to skin testing previously (e.g., anaphylaxis, angioedema, asthma, hypotension, etc.)?

- If **YES**, do not proceed with allergy skin testing.

  ➔ If **YES**, and the subject has acceptable ACRN skin testing results from a prior ACRN protocol (ACRN I or II), record Subject ID associated with the most recent acceptable test.

   - Subject ID: ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___

3. Has the subject taken any of the medications listed in the ACRN Skin Testing MOP within the exclusionary periods?

- If **YES**, the allergy skin testing procedure should be rescheduled.

   - Subject ID: ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___ ___
4. Was the subject’s most recent FEV1 below 60% predicted? ☐ 1 Yes ☐ 0 No (1072)

⇒ If NO, proceed to Question #5.

4a. Has the subject received permission from the supervising physician to proceed with the skin testing? ☐ 1 Yes ☐ 0 No (1074)

⇒ If YES, obtain physician’s signature:

______________________________ (1076)

⇒ If NO, allergy skin testing procedure should be rescheduled.

5. Is the subject eligible for allergy skin testing? ☐ 1 Yes ☐ 0 No (1080)

If any of the shaded boxes are completed, the subject is ineligible for allergy skin testing. STOP HERE.

⇒ Allergy Skin testing may be rescheduled for the next visit if the subject is ineligible due to Question #3 or Question #4a.

6. Time test sites pricked/punctured (based on 24-hour clock) ___ ___ ___ (1090)

7. Time test sites evaluated (based on 24-hour clock) ___ ___ ___ (1100)
Transfer the tracing of each measurable wheal and record the longest diameter and the diameter at the perpendicular midpoint in mm. If the wheal is not measurable, record ‘0’ for both diameters.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
<td>Largest Wheal</td>
</tr>
<tr>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
<td>Diameter: ___ ___ ___ mm</td>
</tr>
<tr>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
<td>Perpendicular Wheal Diameter: ___ ___ ___ mm</td>
</tr>
</tbody>
</table>
ALLERGY SKIN TEST RESULTS

Subject ID: __ __ __ __ __ __ __ __
Visit Number: __ __ __ __

13. Is the mean diameter for the ‘Negative Control’ < 3 mm?  
   ☐ 1 Yes ☐ 0 No (1350)
   ➔ If YES, go to Question #14.
   ➔ If NO, administer the negative control on the opposite hand and complete Question #13a and #13b.

13a. Record the measurements for the ‘Negative Control’ administered on the opposite hand:
   Largest Wheal Diameter: ___ ___ mm (1352)
   Perpendicular Wheal Diameter: ___ ___ mm (1354)

13b. Is the mean diameter calculated from the measurements in Question #13a < 3 mm?  
   ☐ 1 Yes ☐ 0 No (1360)
   ➔ If NO, go to Question #15. The subject has dermatographia and therefore, do not repeat skin testing on this subject.

14. Is the mean diameter for ‘Positive Control’ ≥ 3 mm more than the mean diameter from the ‘Negative Control’?  
   ☐ 1 Yes ☐ 0 No (1370)

15. Was this test acceptable?  
   ☐ 1 Yes ☐ 0 No (1380)
   
   If any of the gray shaded boxes are checked, this test was not acceptable.
   
   ➔ Allergy Skin testing may be rescheduled for the next visit if the subject’s test was unacceptable due to the use of exclusionary medications.
Complete this form only if the subject is eligible according to the appropriate Pulmonary Procedure Checklist form.

1. Time spirometry started *(based on 24-hour clock)*

   ___ ___ ___ ___ (1000)

*The best effort reflects the trial where the sum of FEV₁ and FVC is maximized.*

2. Results of best effort:

   2a. FVC

       ___ . ___ ___ L (1010)

   2b. FEV₁

       ___ . ___ ___ L (1020)

   2c. FEV₁ (% predicted)

       ___ ___ ___ % predicted (1030)

   2d. PEFR

       ___ . ___ ___ L/S (1040)

   2e. FEF₂₅₋₇₅

       ___ . ___ ___ L/S (1050)

3. In your judgment, was the subject’s spirometry technique acceptable?

   [ ] 1 Yes   [ ] 0 No (1060)

   3a. If NO, why was it unacceptable?

   Inadequate inspiratory effort

       [ ] 1 Yes   [ ] 0 No (1070)

   Inadequate expiratory effort

       [ ] 1 Yes   [ ] 0 No (1080)

   Inadequate duration of expiration

       [ ] 1 Yes   [ ] 0 No (1090)

   Cough during procedures

       [ ] 1 Yes   [ ] 0 No (1100)

   Other *(specify)* __________________________

       [ ] 1 Yes   [ ] 0 No (1110)
### SPUTUM INDUCTION LAB VALUES

(Technician completed)

**Processing Sample**

1. Technician ID

2. Processing Date

3. Time processing started *(based on 24-hour clock)*

4. Total Cell Count

**Differential Cell Counts**

5. Technician ID

6. Read Date

7. Squamous Cells

8. Did the subject’s sputum sample reveal ≥ 80% squamous cells?  
   □ 1 Yes  □ 0 No

   ➤ If NO, please complete Question #9 through Question #14 and send the sputum sample for overreading.

   ➤ If YES, STOP HERE and mark the samples as excluded from shipment to San Francisco in the Sample Tracking Module.

The parameters below are calculated following exclusion of squamous cells.

9. Total Cell Count

10. Epithelial Cells

11. Macrophages

12. Neutrophils

13. Eosinophils

14. Lymphocytes
SPUTUM INDUCTION
UCSF OVER-READ

(Technician completed)

1. Date of Over-Read

    ___ ___ / ___ ___ / ___ ___ ___ (1000)

2. Is the slide quality acceptable?
   ➔ If NO, please comment below. If a back-up slide is required, update the Sample Tracking Module.

   ______________________________________________________________
   ______________________________________________________________
   ______________________________________________________________

Differential Cell Counts

3. Squamous Cells

   ___ ___ . ___ % (1020)

*The parameters below are calculated following exclusion of squamous cells.*

4. Epithelial Cells

   ___ ___ . ___ % (1030)

5. Macrophages

   ___ ___ . ___ % (1040)

6. Neutrophils

   ___ ___ . ___ % (1050)

7. Eosinophils

   ___ ___ . ___ % (1060)

8. Lymphocytes

   ___ ___ . ___ % (1070)
SPUTUM INDUCTION

(Technician completed)

Complete this form only if the subject is eligible according to the Sputum Induction Checklist (SPUTUMCHK) form.

1. (If attempting sputum induction for the first time in this protocol or subject has not had an adequate sample at prior attempts, do not complete Question #1.)
   What was the duration of sputum induction the first time the subject’s sample was processed and had < 80% squamous cells for this protocol?  
   Duration of sputum induction at current visit should not exceed this.  
   _____ . ____ minutes (1000)

2. Sputum induction start time (based on 24-hour clock)  
   _____ _____ (1010)

3. Sputum induction stop time (based on 24-hour clock)  
   _____ _____ (1020)

4. Duration of sputum induction collection phase at this visit  
   _____ . ____ minutes (1030)
   4a. Was the duration ≥ 4 minutes?  
      □ 1 Yes □ 0 No (1040)

5. Volume of sputum sample at this visit  
   _____ _____ . ____ ml (1050)
   5a. Is the volume of the sample ≥ 1 ml?  
      □ 1 Yes □ 0 No (1060)

6. Is the sample adequate for laboratory analysis?  
   If either shaded box in Question #4a or #5a are completed, the sputum sample is not adequate and should not be sent for analysis of squamous cell counts.  
   ➔ If YES, the technician reading the slide should complete the Sputum Induction Lab Values (SPUTLAB) form.  
      □ 1 Yes □ 0 No (1070)
7. Subject’s FEV₁ immediately after completion of sputum induction

7a. FEV₁ ____. ____ L (1080)

7b. FEV₁ (% predicted) ____. ____ % predicted (1090)

7c. Time of FEV₁ in Question #7a (based on 24-hour clock) ___________ (1100)

7d. Percent difference in FEV₁ \( \frac{\text{Reference} - \text{Question #7a}}{\text{Reference}} \times 100 \) ____. ____ % (1110)

Reference = FEV₁ used for assessment of eligibility for SI.

7e. Did the subject’s FEV₁ drop > 10% (from post-albuterol baseline) as indicated in Question #7d?

☐ 1 Yes ☐ 0 No (1120)

➔ If NO, STOP HERE and continue with remaining visit procedures.
➔ If YES, proceed to the Additional Treatment for Sputum Induction (SPUTUM_ADD_TRT) form.
**ADDITIONAL TREATMENT POST SPUTUM INDUCTION**

Subject ID: _____ - _____ - _____
Subject Initials: _____ ______
Visit Number: ________
Visit Date: ___ / ___ / ___
Month            Day                 Year
Technician ID: ________

(Technician completed)

Complete this form only if the subject has experienced > 10% fall in FEV\textsubscript{1} from post-albuterol baseline immediately after completion of sputum induction.

> Administer 2 puffs of albuterol and wait 15 minutes, then perform spirometry.

1. Subject’s FEV\textsubscript{1} after initial 2 puffs of albuterol
   1a. FEV\textsubscript{1} _______ • _______ L (1000)
   1b. FEV\textsubscript{1} (% predicted) _______ • _______ % predicted (1010)
   1c. Time of FEV\textsubscript{1} from Question #1a *(based on 24-hour clock)* _______ • _______ (1020)
   1d. Was the FEV\textsubscript{1} from Question #1a ≥ the sputum induction reversal reference value in the gray box above?
      ➔ If YES, stop here and continue with remaining visit procedures.
      ➔ If NO, administer 2 puffs of albuterol and wait 15 minutes, then perform spirometry. Proceed to Question #2.

2. Subject’s FEV\textsubscript{1} after 2 additional puffs of albuterol
   2a. FEV\textsubscript{1} _______ • _______ L (1040)
   2b. FEV\textsubscript{1} (% predicted) _______ • _______ % predicted (1050)
   2c. Time of FEV\textsubscript{1} from Question #2a *(based on 24-hour clock)* _______ • _______ (1060)
   2d. Was the FEV\textsubscript{1} from Question #2a ≥ the sputum induction reversal reference value in the gray box above?
      ➔ If NO, complete the source documentation box below.

**Clinic Use Only**

<table>
<thead>
<tr>
<th>Sputum Induction Reversal Reference Value:</th>
<th>Reference \times 0.90 = _____ • _____ L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reference = FEV\textsubscript{1} used for assessment of eligibility for Sputum Induction.</td>
<td></td>
</tr>
</tbody>
</table>

Physician Source Documentation

Physician signature: ____________________________ (1080)
Date: ___ / ___ / ___ ______ (1090)
Time: ________ (based on 24-hour clock) (1100)
(Technician completed)

**Complete this form only if the subject successfully completed baseline spirometry session(s).**

1. **(If attempting Sputum Induction for the first time in this protocol, do not complete Question #1)**
   
   Was the subject’s sputum sample processed and had < 80% squamous cells the first time a sputum induction was attempted for this protocol?  
   
   - [ ] Yes  
   - [ ] No (1000)

2. **(Only for subjects who completed a methacholine challenge at this visit.)**
   
   Was the subject’s FEV₁ after reversal from the methacholine challenge ≥ the methacholine reversal reference value (B) in the gray box on the Methacholine Challenge Testing (METHA) form?  
   
   - [ ] Yes  
   - [ ] No (1010)

   2a. If NO, has the subject received permission from the supervising physician to proceed with sputum induction testing?  
   
   - [ ] Yes  
   - [ ] No (1020)

   **Physician’s Signature:** ____________________________ (1030)

3. Subject’s FEV₁ used for assessment of eligibility for sputum induction  
   
   ______. ______ L (1040)

4. Subject’s FEV₁ (% predicted) used for assessment of eligibility for sputum induction  
   
   ______ ______ % predicted (1050)

5. Was the subject’s FEV₁ (% predicted) from Question #4 ≥ 60% predicted?  
   
   - [ ] Yes  
   - [ ] No (1060)

6. Is there any other reason the subject should not proceed with sputum induction?  
   
   If YES, explain ____________________________

7. Is the subject eligible for sputum induction?  
   
   - [ ] Yes  
   - [ ] No (1080)

   *If any of the shaded boxes are completed, the subject is NOT eligible for sputum induction.*

   ➔ **If YES, proceed to the Sputum Induction (SPUTUM) form.**