

Log APHIS Form 2008

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Source Document: CVB-SOP-0032, *Processing Serial Records*

Complete hard copy APHIS Form 2008s (Form 2008s) are received and entered by the Inspection and Compliance group. Most Form 2008s are received from the licensed manufacturers through the NCAH Portal, in which case this work instruction is not applicable. The Form 2008s are logged into LSRTIS, which is typically done between 10 a.m. and 1 p.m. CST.

The assigned Logger enters the Form 2008s into LSRTIS under the **Serial Release** module, **Log APHIS 2008** section. The following are procedures to perform this duty.

A. Click Log APHIS 2008 and enter the following within the Create APHIS 2008 screen.

The screenshot shows the 'Create APHIS 2008' form with the following fields and options:

- Establishment***: Text input field.
- Product***: Text input field.
- Serial Number***: Text input field.
- Expiration Date**: Text input field.
- Site**: Dropdown menu.
- Fill Date**: Text input field.
- Is this a subserial?**:
- Autogenous or Prescription Product?**: **First Serial** **Fifty Or Less Vials**
- Doses**: **# Containers*** (input: 0) and **Container Size*** (input: 0). Below is a blue **+ Add** button and the text **Number Of Total Doses: 0**.
- Doses Type ***: Dropdown menu (selected: Doses).
- Firm Disposition ***: Dropdown menu (selected: Eligible for Release).
- Received Date***: Text input field (value: 03/10/2017).
- VBI Number**: Text input field.
- Related Submissions**: Text input field with a blue **+** button.
- Comments**: Large text area.

1. **Establishment number** (Block 2 on the Form 2008)
2. **Product code** (Block 5)
 - No decimal point. Ensure this is a valid product code for the selected establishment or an error message will appear after you try and save.
3. **Serial number** (Block 7)
 - a. Limit of 20 characters
 - b. Alpha/numeric characters only – no spaces or hyphens

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- c. If allergenic extract, no serial number is assigned by the manufacturer. CVB identifies each serial by designating it a number for each product code listed in their submission. The serial number will be the same for all; the only difference is the product code, i.e., 13Q4 translates:
 - last 2 digits of the calendar year of the submission
 - Q for quarter year
 - 1-4 for the quarter year it covers
 - Jan-Mar = 1
 - Apr-Jun = 2
 - Jul-Sept = 3
 - Oct-Dec = 4
4. **Expiration Date** (if applicable) (Block 6)
i.e., MM/DD/YYYY
5. **Site** – drop down field of available sites for the firm
6. **Fill Date** (if applicable) (Block 4)
i.e., MM/DD/YYYY
7. **Is this a subserial?** – An optional field to indicate if a serial is a part of a subserial product
8. **Autogenous or Prescription Product**
 - a. Check if the Product is an autogenous product (1015.xx, 2051.00, 2052.xx, 4500.xx) or a prescription product (9Pxx.xx)
 - b. First Serial – check if the firm indicates the serial is a 1st serial of a harvest (no samples are needed for release)
 - c. Fifty or Less Vials – check if the inventory is less than 50 containers (if yes, no samples are needed for release)
9. **Inventory**
 - a. Doses:
 - # Containers (Block 10A)
 - Container Size (Block 10B)
 - Select +Add if there are multiple container sizes
 - b. If no inventory is shown, enter 0
 - c. **NOTE:** If the manufacturer is a permittee, there will be two separate dose field entries (Permittee Total Doses Manufactured and doses received in the United States).
 - Follow the A.5.a. directions above for container/container size(s) for doses received in the United States (Block 11)
 - Total Doses the foreign entity manufactured - amount in Block 10C

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Containers* 500 Container Size* 10
+Add Number Of Total Doses: 5000
Doses Type* Doses Firm Disposition* Eligible for Release
Received Date* 09/14/2015
Permitee Total Doses Manufactured 10000
VBI Number

d. If allergenic extracts, enter:

- Total # containers
- 1 in container size

e. Check running total shown against total inventory in Block 10C

f. If a serial is a first serial autogenous serial, the only inventory field that will show is the number of total doses

Autogenous or Prescription Product (platform?) First Serial Fifty Or Less Vials
Number of Total Doses
Doses Type* Doses Firm Disposition* Eligible for Release
Received Date* 09/14/2015

10. Dose Type

- doses, mL, or units only (see Unit of Measure work instructions **CVB-WI-0157**, *Unit of Measure Use and Conversion for Data Entry into LSRTIS*)

11. Firm Disposition (Block 12)

- Click on the LOV to select firm's disposition (See **CVB-WI-0129**, *APHIS Dispositions and Associated Information on Form 2008s*, for the entire list of Firm Dispositions)
- “Reprocess & Retest”
The new product code and serial number indicated in Block 11 of Form 2008 appears at the bottom of the screen. Select **+Add** if there is more than one new serial number.
- “Other - Rebottling”
The new product code and serial number indicated in Block 11 of Form 2008 appears at the bottom of the screen. Select **+Add** if there is more than one new serial number.
- “Other - Transfer Request”
Once you enter this disposition, enter the establishment, product code, and serial that the inventory is being transferred to as indicated in Block 11 of Form 2008.

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12. Received Date (Stamp date on the Form 2008)

- a. Default as today's date
- b. Ability to back date is available

13. VBI Number

Format (10-001). This field points directly to the investigation module within LSRTIS, use only valid investigation numbers.

14. Related Submissions (mail log #) This field will point directly to either IC or PEL Mail log numbers, if references on the Form 2008.

15. Comments

Add any applicable comments in comment field

16. Click Create complete the entry

B. Once the Form 2008s have been logged in, the Logger will distribute all Form 2008s to the assigned Biologics Compliance Assistant in their Incoming 2008s folder.

C. Form 2008s received after 2 p.m. may be held, stamped, and logged into LSRTIS the following day. Hard copy 2008s may not be entered the day of receipt, though they will be backdated to the date of actual receipt.