

**U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES
FOOD AND DRUG ADMINISTRATION**

REPORT OF TUBERCULIN TESTS OF CATTLE
*REQUIRED FOR INTRODUCTION INTO THE UNITED STATES
OF RAW MILK AND CREAM UNDER THE FEDERAL IMPORT MILK ACT*

*Form Approved; OMB No. 0910-0212
Expiration Date: 08-31-21
See Reverse for OMB Statement*

OWNER	DATE OF EXAMINATION
ADDRESS	LOCATION OF HERD

A N I M A L N O	A G E	S E X	IDENTIFICATION OF ANIMAL <i>(Accurate description or ear tag number or registration name and number)</i>	SUBCUTANEOUS								INTRADERMAL INJECTION			CONCLUSIONS ² <i>(healthy, reactor, suspect)</i>					
				TEMPERATURE BEFORE INJECTION			TIME OF INJECTION	TUBERCULIN INJECTION QTY/TYPER ¹		TEMPERATURE AFTER INJECTION						Type _____	Quantity _____	Date _____	Hour _____	
				_ M	_ M	_ M	_____ M	C.C.	TYPE	_ M	_ M	_ M	_ M	_ M		_ M	OBSERVATIONS			
															_ hrs	_ hrs	_ hrs			
1																				
2																				
3																				
4																				
5																				
6																				
7																				
8																				
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IF MORE SPACE IS NEEDED USE ADDITIONAL FORMS FDA 1994

CERTIFICATION: I HEREBY CERTIFY THAT I HAVE INSPECTED AND TESTED WITH TUBERCULIN THE _____ ANIMALS ABOVE WITH RESULTS AS STATED.

SIGNATURE OF VETERINARIAN	TITLE <i>(Official connection)</i>	MAILING ADDRESS <i>(Include ZIP Code)</i>
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1. The following symbols may be used to denote the type of tuberculin: B - Bovine; A - Avian; & H - Human.
2. All reactors & suspicious animals must be removed from herd & certificate of owner to that effect given on the other side of this form.

This section applies only to the requirements of the Paperwork Reduction Act of 1995: The public reporting burden time for this collection of information is estimated to average .25 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRAStaff@fda.hhs.gov

Do not send your completed form to the PRA Staff email address to the left.

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."

INSTRUCTIONS

Either the subcutaneous or the intradermal test may be used.

Subcutaneous Test - When the subcutaneous tuberculin test is applied, the chart on the other side of this form shall show that at least 3 temperatures were taken, 2 or 3 hours apart, before the injection of tuberculin; that at least 6 temperatures were taken 2 hours apart, beginning 8 hours after the injection; and that the test had run for a period of not less than 18 hours after injection.

Intradermal Test - When the intradermal test is applied, the chart on the other side of this form shall show that the last observation was made not earlier than the seventy-second hour after injection. At least two observations should be made on animals from infected herds.

Reactions, Subcutaneous Test - A rise of 2 degrees Fahrenheit or more above the maximum temperature observed prior to the injection of tuberculin or a temperature above 103.8°F. should be regarded as an indication of tuberculosis, provided the temperature reaction shows a characteristic curve. An elevation of temperature higher than 103.8°F. should also be regarded as an indication of tuberculosis,

even though the curve may not be regarded as typical. Animals that after injection show a rise in temperature of 2 degrees Fahrenheit, with a maximum between 103° and 103.8°F., as well as those that show a rise of less than 2 degrees Fahrenheit, with a maximum temperature of 103.8° F., are regarded as suspicious reactors.

Reactions, Intradermal Test - Animals that show at the point of injection swellings, either hard and circumscribed, or soft and infiltrated, with no distinct line of demarcation, should be classified as reactors to the intradermal tuberculin test. Such swellings may be of various sizes, from those hardly perceptible to the naked eye to those as large as the human fist or larger. The results of observations on individual animals should be reported on the test record in accordance with the prescribed code of the country in which the cattle are tested.

Identification of Animals - All animals reported on this chart must be identified by proper metal ear tags, or registration names and numbers, or by accurate descriptions.

I HEREBY CERTIFY that I am the owner of the cows described on the other side of this form; that all animals in the herd have been tested, and that the animals designated as "reactors" and "suspected" as numbers _____

_____ on this form, have been permanently removed from the herd and disposed of as follows _____

I certify further that no milk or cream from such reactors or suspected animals will be introduced into the United States.

(Signed) _____
(Name of Owner)

(Address)

(Date)