§ 58.135 Bacterial estimate.

(a) Methods of Testing. Milk shall be tested for bacterial estimate by using one of the following methods or by any other method approved by Standard Methods for the Examination of Dairy Products.

(1) Direct Microscopic clump count;
(2) Standard plate count;
(3) Plate loop count;
(4) Pectin gel plate count;
(5) Petrifilm aerobic count;
(6) Spiral plate count;
(7) Hydrophobic grid membrane filter count;
(8) Impedance/conductance count;
(9) Reflectance calorimetry.

(b) Frequency of Testing. A laboratory examination to determine the bacterial estimate shall be made on a representative sample of each producer’s milk at least once each month at irregular intervals. Samples shall be analyzed at a laboratory in accordance with State regulations.

(c) Acceptance of milk. The following procedures shall be applied with respect to bacterial estimates:

(1) Whenever the bacterial estimate indicates the presence of more than 500,000 bacteria per ml., the producer shall be notified with a warning of the excessive bacterial estimate.

(2) Whenever two of the last four consecutive bacterial estimates exceed 500,000 per ml., the appropriate regulatory authority shall be notified and a written warning notice given to the producer. The notice shall be in effect so long as two out of the last four consecutive samples exceed 500,000 per ml.

(3) An additional sample shall be taken after a lapse of 3 days but within 21 days of the notice required in paragraph (c) (2) of this section. If this sample also exceeds 500,000 per ml., subsequent milkings shall be excluded from the market until satisfactory compliance is obtained. Shipment may be resumed when an additional sample of herd milk is tested and found satisfactory.


§ 58.136 Rejected milk.

A plant shall reject specific milk from a producer if the milk fails to meet the requirements for appearance and odor (§ 58.133(a)), if it is classified No. 4 for sediment content (§ 58.134), or if it tests positive for drug residue (§ 58.133(c)).

[58 FR 26913, May 6, 1993]

§ 58.137 Excluded milk.

A plant shall not accept milk from a producer if:

(a) The milk has been in a probational (No. 3) sediment content classification for more than 10 calendar days (§ 58.134);

(b) Three of the last five milk samples have exceeded the maximum bacterial estimate of 500,000 per ml. (§ 58.135 (c)(3));

(c) Three of the last five milk samples have exceeded the maximum somatic cell count level of 750,000 per ml. (1,000,000 per ml. for goat milk) (§ 58.133 (b)(6)); or

(d) The producer’s milk shipments to either the Grade A or the manufacturing grade milk market currently are not permitted due to a positive drug residue test (§ 58.133(c)(4)).

[58 FR 26913, May 6, 1993, as amended at 67 FR 48975, July 29, 2002]

§ 58.138 Quality testing of milk from new producers.

A quality examination and tests shall be made on the first shipment of milk from a producer shipping milk to a plant for the first time or resuming shipment to a plant after a period of non-shipment. The milk shall meet the requirements for acceptable milk, somatic cell count and drug residue level (§§ 58.133, 58.134 and 58.135). The buyer
shall also confirm that the producer’s milk is currently not excluded from the market (§58.137). Thereafter, the milk shall be tested in accordance with the provisions in §§58.133, 58.134 and 58.135.

[58 FR 26913, May 6, 1993]

§ 58.139 Record of tests.

Accurate records listing the results of quality and drug residue tests for each producer shall be kept on file at the plant. Additionally, the plant shall obtain the quality and drug residue test records (§58.148(a), (e) and (g)) for any producer transferring milk shipment from another plant. These records shall be available for examination by the inspector.

[58 FR 26913, May 6, 1993]

§ 58.140 Field service.

A representative of the plant shall arrange to promptly visit the farm of each producer whose milk tests positive for drug residue, exceeds the maximum somatic cell count level, or does not meet the requirements for acceptable milk. The purpose of the visit shall be to inspect the milking equipment and facilities and to offer assistance to improve the quality of the producer’s milk and eliminate any potential causes of drug residues. A representative of the plant should routinely visit each producer as often as necessary to assist and encourage the production of high quality milk.

[58 FR 26913, May 6, 1993]

§ 58.141 Alternate quality control program.

When a plant has in operation an acceptable quality program, at the producer level, which is approved by the Administrator as being effective in obtaining results comparable to or higher than the quality program as outlined above for milk or cream, then such a program may be accepted in lieu of the program herein prescribed.

OPERATIONS AND OPERATING PROCEDURES

§ 58.142 Product quality and stability.

The receiving, holding and processing of milk and cream and the manufac-turing, handling, packaging, storing and delivery of dairy products shall be in accordance with clean and sanitary methods, consistent with good commercial practices to promote the production of the highest quality of finished product and improve product stability. Milk should not be more than three days old when picked up from the producer and delivered to the plant, receiving station or transfer station.

§ 58.143 Raw product storage.

(a) All milk shall be held and processed under conditions and at temperatures that will avoid contamination and rapid deterioration. Drip milk from can washers and any other source shall not be used for the manufacture of dairy products. Bulk milk in storage tanks within the dairy plant shall be handled in such a manner as to minimize bacterial increase and shall be maintained at 45 °F or lower until processing begins. This does not preclude holding milk at higher temperatures for a period of time, where applicable to particular manufacturing or processing practices.

(b) The bacteriological quality of commingled milk in storage tanks shall not exceed 1,000,000/ml.


§ 58.144 Pasteurization or ultra-pasteurization.

When pasteurization or ultra-pasteurization is intended or required, or when a product is designated “pasteurized” or “ultra-pasteurized” every particle of the product shall be subjected to such temperatures and holding periods in approved systems as will assure proper pasteurization or ultra-pasteurization of the product. The heat treatment by either process shall be sufficient to insure public health safety and to assure adequate keeping quality, yet retaining the most desirable flavor and body characteristics of the finished product.

§ 58.145 Composition and wholesomeness.

All necessary precautions shall be taken to prevent contamination or