UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

December 28, 1977 DATE:

Lonza Formulation 70-12-Addition of Dermal LD₅₀ Data to SUBJECT:

EPA Reg. No. 6836-33, Caswell#613A,392H,331A,846,509,430,16E Shaughnessy No.069165,069166,039107,047501,069149,001501,169101

Toxicology Branch FROM:

Registration Division

J. Tavano TO:

Product Manager #31

Thru: Mary Quaife, Ph.D

Acting Branch Chief

Recommendation: The study is considered Supplementary Data because of reasons stated in the review.

*No RPAR criteria have been exceeded.

Review

Acute Dermal LD₅₀ Study (Leberco Laboratories, Assay#731375, 2/7/77, Α. submitted by Lonza, Inc., 3/22/77, Acc.#232265).

1. Procedure

Twelve female rabbits, unspecified strain, 2,2-3.1 Kg, divided into 3 groups of 4 animals each which received dermal applications of 1 ml, 2 ml, or 5 ml/kg of test material under occlusive dressing. Backs were shaved prior to applications. No test sites were abraded. At 24 hours post-treatment dressing was removed, and animals were washed and towel-dried. Animals were observed 14 days, post-treatment.

Dose (m1/ka) Deaths Results:

 $LD_{50} > 1 \text{ ml/kg}.$

- 3. Conclusions
- a). Classification: Supplementary
- No data of toxic signs, necropsies, or statistics were submitted. Doses used appear to be in the upper portion of the LD range. The study should be extended to include doses below 1 ml/kg to permit a

more precise estimate of the LD_{50} , taking submitted results into consideration.

No males were used, and no test sites were indicated to have been abraded.

Larry Anderson Uly

EPA FORM 1320-6 (REV. 3-76)