

US EPA ARCHIVE DOCUMENT

UNITED STATES ENVIRONMENTAL PROTECTION AGENCY

DATE: December 28, 1977

SUBJECT: Lonza Formulation 70-12-Addition of Dermal LD<sub>50</sub> Data to  
EPA Reg. No. 6836-33, Caswell#613A,392H,331A,846,509,430,16E  
Shaughnessy No.069165,069166,039107,047501,069149,001501,169101

FROM: Toxicology Branch  
Registration Division

TO: J. Tavano  
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

A. Acute Dermal LD<sub>50</sub> Study (Leberco Laboratories, Assay#731375, 2/7/77, submitted by Lonza, Inc., 3/22/77, Acc.#232265).

1. Procedure

a). Twelve female rabbits, unspecified strain, 2.2-3.1 Kg, <sup>were</sup> 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.

2. <u>Results:</u>	Dose (ml/kg)	Deaths
	1	2
	2	3
	5	3

LD<sub>50</sub> > 1 ml/kg.

3. Conclusions

- a). Classification: Supplementary
- i). No data of toxic signs, necropsies, or statistics were submitted.
- ii). Doses used appear to be in the upper portion of the LD<sub>50</sub> range. The study should be extended to include doses below 1 ml/kg to permit a more precise estimate of the LD<sub>50</sub>, taking submitted results into consideration.
- iii). No males were used, and no test sites were indicated to have been abraded.

Larry Anderson  
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U.L.G.  
1/14/77