PRODUCT CLINICAL DATA SUMMARY
NO. 1538L
3M Woven Medical Tape
Effective: January 1996

No. 1538L 3M Woven Medical Tape has been subjected to the following safety evaluations:

**In Vitro Cytotoxicity (Agar Overlay)**
**Results:** 1.0/0.5

**Acute Primary Skin Irritation in Albino Rabbits**
**Results:** 0.7/8.0

**Modified Buehler Sensitization in Guinea Pigs**
**Results:** Weak sensitizer.

**Repeated Insult Patch Test (Draize) in Humans**
**Results:** No induced contact sensitization.

**21-day Cumulative Irritation in Humans**
**Results:** Consistent with responses characteristic of adhesive material; within historically acceptable levels for surgical tapes.
These tests are in accordance with the ISO 10993 Part-1 "Biological Evaluation of Medical Devices", as put forth by the FDA. No. 1538-L has satisfied the requirements for devices in contact with intact skin for short term application (up to 29 days). All laboratory testing was conducted in accordance with the FDA Good Laboratory Practices Regulation of 1978.

The use of the term "hypoallergenic" has come to indicate a product which is non-sensitizing to the general public. The hypoallergenic claim for this product is supported by clinical evaluation using the repeated insult patch test in humans, commonly known as the Draize test. This protocol involves repeated application of samples on 200 healthy volunteers for a 2- to 3-week induction period, followed by a 2-week rest period and a challenge application. To be termed hypoallergenic, 3M Medical Specialties products are required to show no evidence of sensitization potential under these test conditions.

It is the responsibility of our customers to determine the final suitability of our products for their application.