

Committee of Experts on the Transport of Dangerous Goods and on the Globally Harmonized System of Classification and Labelling of Chemicals

Sub-Committee of Experts on the Transport of Dangerous Goods

18 June 2010

Thirty-seventh session

Geneva, 21–30 June 2010

Item 5 of the provisional agenda

**Miscellaneous proposals of amendments to the Model Regulations
on the Transport of Dangerous Goods**

Used Health Care Products

Transmitted by the expert from Switzerland

Introduction

1. Hospitals, physicians, pharmacies and patients use diagnostic devices to determine numerous clinically relevant parameters in various biological materials. These devices have to be transported not only when new but also in a used condition, e.g. for major repair and maintenance work, for reasons of quality verification and also for final disposal at the end of their useful lives. These devices range from small pocket-size blood-sugar test devices to complex multi-modular electronic pieces of equipment of several cubic meters of volume. If properly prepared for transportation, these instruments present no infectious risk during the transport although the presence of infectious agents (e.g. hepatitis B or hepatitis C virus) in the innermost parts of these complex devices cannot be totally excluded. Transport of such devices as non-dangerous goods has therefore so far occurred in a gray zone of the transport regulations.

2. The United States have overcome this issue by a special derogation for so called “used health care products” (reference: CFR 49, Chapter 173.134), a term that also includes diagnostic devices. In analogy to these regulations that have proven valuable in everyday practice, we submit the following proposals.

3. For land transport in Europe the RID and the ADR have also a derogation to their rules in 1.1.3.1.b).

4. No harmonized solution exists for the time being. The following proposals intend to bring an harmonized rule for such necessary carriages.

Proposals

5. Add the following definition to 2.6.3.1:

“Used health care product means a medical, diagnostic, or research device or piece of equipment, or a personal care product used by consumers, medical professionals, or pharmaceutical providers that does not meet the definition of a diagnostic specimen, biological product, or regulated medical waste. It can be contaminated with potentially infectious body fluids or materials, and is not decontaminated or disinfected to remove or mitigate the infectious hazard prior to transportation.”.

6. Add the following paragraph to 2.6.3.2.3 Exemptions:

2.6.3.2.3.x

“Used health care products are exempted from these Regulations if they have been drained of free liquid and have been decontaminated or disinfected to remove or mitigate the infectious hazard prior to transportation. Small diagnostic devices for single-patient use (e.g. devices for monitoring the blood sugar) need not to be treated by a disinfectant if they are completely free of liquid and show no visible contamination on their outsides.”.
