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AMENDMENT 1  
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**Infusion equipment for medical use —**  
**Part 15:**  
**Light-protective infusion sets for**  
**single use**

**AMENDMENT 1**

*Matériel de perfusion à usage médical —*

*Partie 15: Perfuseurs photoprotecteurs à usage unique*

*AMENDEMENT 1*



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This document was prepared by Technical Committee ISO/TC 76, *Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use*, in collaboration with the European Committee for Standardization (CEN) Technical Committee CEN/TC 205, *Non-active medical devices*, in accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).

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