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AMENDMENT 1
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Pen systems —

Part 2:

**Plunger stoppers for pen-injectors
for medical use**

AMENDMENT 1

Systèmes de stylos-injecteurs —

Partie 2: Bouchons-pistons pour stylos-injecteurs à usage médical

AMENDEMENT 1



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The committee responsible for this document is ISO/TC 76, *Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use*.

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