

First edition  
2003-08-01

AMENDMENT 1  
2018-01

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**Elastomeric parts for parenterals and  
for devices for pharmaceutical use —**

**Part 3:  
Determination of released-particle  
count**

**AMENDMENT 1**

*Éléments en élastomère pour administration parentérale et dispositifs  
à usage pharmaceutique —*

*Partie 3: Détermination des particules libérées*

*AMENDEMENT 1*



Reference number  
ISO 8871-3:2003/Amd.1:2018(E)

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This document was prepared by ISO/TC 76, *Transfusion, infusion and injection, and blood processing equipment for medical and pharmaceutical use*.

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