
Zunajtelesni pretočni sistemi za čiščenje krvi - 2. del: Zunajtelesni krvni in tekočinski obtok za hemodializatorje, hemodiafiltre, hemofiltre in hemokoncentratorje (ISO/DIS 8637-2:2025)

Extracorporeal systems for blood purification - Part 2: Extracorporeal blood and fluid circuits for haemodialysers, haemodiafilters, haemofilters and haemoconcentrators (ISO/DIS 8637-2:2025)

Extrakorporale Systeme zur Blutreinigung - Teil 2: Extrakorporaler Blut- und Flüssigkeitskreislauf bei Hämodialysatoren, Hämodiafiltern, Hämofiltern und Hämokonzentratoren (ISO/DIS 8637-2:2025)

Systèmes extracorporels pour la purification du sang - Partie 2: Circuits sanguins extracorporels et liquidiens pour les hémodialyseurs, les hémodiafiltres, les hémo-filtres et les hémoco concentrateurs (ISO/DIS 8637-2:2025)

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DRAFT International Standard

ISO/DIS 8637-2

Extracorporeal systems for blood purification —

Part 2:

Extracorporeal blood and fluid circuits for haemodialysers, haemodiafilters, haemofilters and haemoconcentrators

Systèmes extracorporels pour la purification du sang —

Partie 2: Circuits sanguins extracorporels et liquidiens pour les hémodialyseurs, les hémodiafiltres, les hémofiltres et les hémoconcentrateurs

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