

U2 Knee MDT™

Single-Use Modular Disposable Trial



U2 Knee MDT (Modular Disposable Trial) –

Enables surgeons to complete all trial evaluation surgical technique steps with pre-sterilized, single-use components.

MDT includes femoral trial sets, tibial trial sets and patellar trial sets designated by implant size.

When using the U2 Knee System's AiO Block and MDT Implant Trials together, the number of required instrument trays can be reduced from 6 to 1.5.

U2 Knee MDT and U2 Knee AiO was awarded the Bronze Winner at the 21st Annual Medical Design Excellence Awards (MDEA).

20  19
BRONZE WINNER
MEDICAL DESIGN
EXCELLENCE AWARDS



The U2 Knee AiO Block and MDT

Reduced Instrument Cleaning / Sterilization Costs and Faster Room Turnover

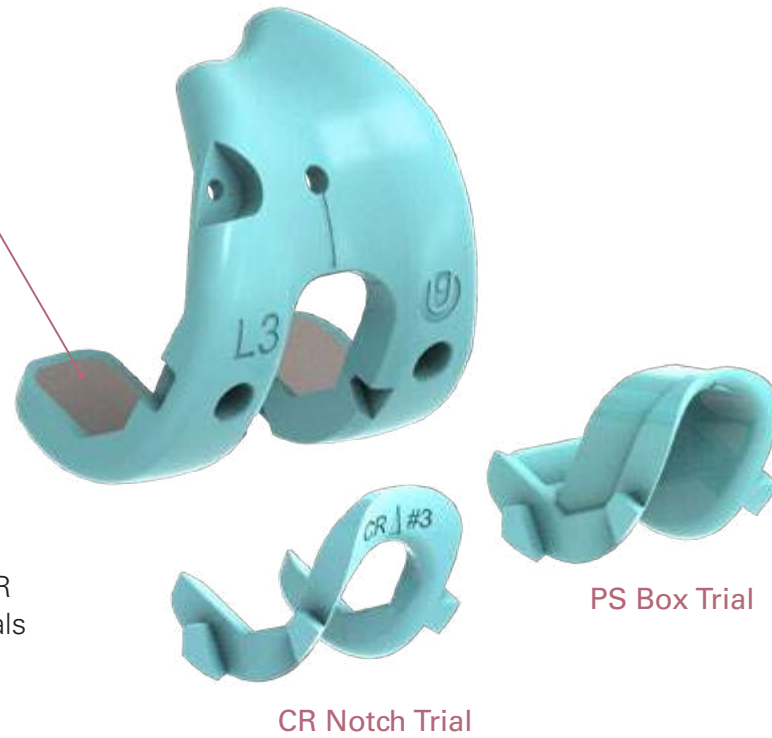


Femoral Trial Set



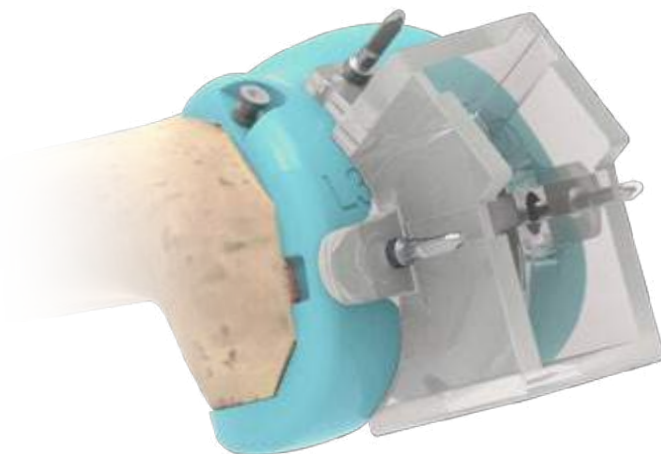
Metal-backed reinforcement

Surgeons can easily convert PS and CR versions by simply switching notch trials



CR Notch Trial

PS Box Trial



PS Notch Cutting Guide

A PS notch cutting guide is available for PS box preparation



Tibial Trial Set



PS Post Trial

Enables transformation of CR insert trial into PS version

9 mm Insert Trial

2 mm Spacers

3 mm Spacer

Thickness
9, 11, 12, 13, 14, 15,
16, 18 mm

Baseplate Trial





Patellar Trial Set



All trial patella sizes are included in each package.



Order Information

Catalog Number	Description
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Femoral Trial Set



Left	Right	
2104 - 2110	2104 - 2210	# 1
2104 - 2115	2104 - 2215	# 1.5
2104 - 2120	2104 - 2220	# 2
2104 - 2125	2104 - 2225	# 2.5
2104 - 2130	2104 - 2230	# 3
2104 - 2135	2104 - 2235	# 3.5
2104 - 2140	2104 - 2240	# 4
2104 - 2145	2104 - 2245	# 4.5
2104 - 2150	2104 - 2250	# 5
2104 - 2155	2104 - 2255	# 5.5
2104 - 2160	2104 - 2260	# 6
2104 - 2165	2104 - 2265	# 6.5
2104 - 2170	2104 - 2270	# 7

Tibial Trial Set



2204 - 4000	# 0
2204 - 4010	# 1
2204 - 4020	# 2
2204 - 4030	# 3
2204 - 4040	# 4
2204 - 4050	# 5
2204 - 4060	# 6
2204 - 4070	# 7

Patellar Trial Set



2404 - 2000	Onset (7 sizes)
2404 - 2010	Inset (4 sizes)

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