

## Medical selector guide



TRITAN copolyesters	Suggested application	Specific gravity ASTM D792	Tensile yield ASTM D638		Tensile break ASTM D638		Flexural modulus ASTM D790 Mpa (105 psi)	Izod impact, notched J/m (ft.lbf/in.) @ 23°C (73°F) ASTM D256	Deflection temperature °C (°F) ASTM D648		Optical properties ASTM D1003		Chemical resistance
			Elongation %	Stress Mpa (psi)	Elongation %	Stress Mpa (psi)			@ 1.82 Mpa (264 psi)	@ 0.455 Mpa (66 psi)	Transmittance %	Haze %	
MX710	Y sites	1.18	6	43 (6200)	210	53 (7700)	1550 (2.3)	980 (18.4)	85 (185)	99 (210)	91	<1	Best
MX711 (mold release)	IV components												
MX731 (mold release)	Medical devices	1.18	7	43 (6200)	210	52 (7500)	1570 (2.3)	860 (16.1)	81 (178)	94 (201)	91	<1	Better
MX810	Device housings	1.17	7	44 (6400)	140	53 (7700)	1580 (2.3)	650 (12.2)	92 (198)	109 (228)	92	<1	Good
MX811 (mold release)													



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*It is the responsibility of the medical device manufacturer ("Manufacturer") to determine the suitability of all component parts and raw materials, including any Eastman product, used in its final product to ensure safety and compliance with requirements of the United States Food and Drug Administration (FDA) or other international regulatory agencies.*

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*For manufacturers of medical devices, biological evaluation of medical devices is performed to determine the potential toxicity resulting from contact of the component materials of the device with the body. The ranges of tests under FDA-Modified ISO-10993, Part 1, "Biological Evaluation of Medical Devices" include cytotoxicity, sensitization, irritation or intracutaneous reactivity, systemic toxicity (acute), subchronic toxicity (subacute), implantation, and hemocompatibility. For Eastman products offered for the medical market, limited testing information is available on request. The Manufacturer of the medical device is responsible for the biological evaluation of the finished medical device.*

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