



Technical Data Sheet

INNATE™ TF220 Precision Packaging Resin

Main Characteristics

- Blown and cast extrusion process. Monolayer and co-extrusion film for packaging applications.
- Critical for BOPE application.

Complies with

- U.S. FDA 21 CFR 177.1520(c)3.2a
- EU, No 10/2011

Consult the regulations for complete details.

Properties¹

Physical	Nominal Value	Unit (English)	Nominal Value	Unit (SI)	Test Method
Density	0.955	g/cm ³	0.955	g/cm ³	ASTM D792
Melt Index (190°C/2.16 kg)	1.5	g/10 min	1.5	g/10 min	ASTM D1238

1. Typical properties: these are not to be construed as specifications. Users should confirm the results by their own tests.
2. ASTM: American Society for Testing and Materials.

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