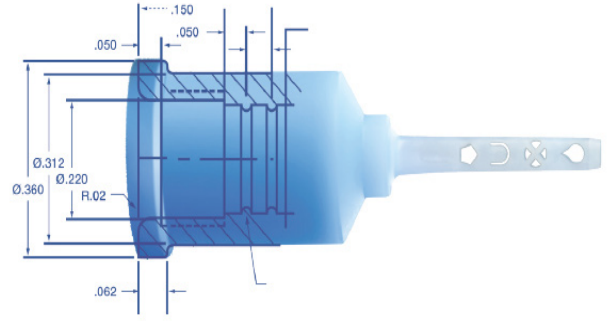


Tooling Prototyping

From initial concept to
full scale production



At Saint-Gobain Medical, we know that speed matters during early-stage medical device development. Our prototyping capabilities were designed with this in mind, allowing us to move fast without compromising precision or quality. With dedicated project engineers, deep materials expertise, and in-house tooling, we help you iterate quickly and make smarter design decisions up front.

Our team collaborates closely with you to select the right materials, optimize for manufacturability, and reduce downstream risk. Depending on the complexity of your design, we can deliver functional silicone components in as little as 5-10 business days, helping you maintain momentum through the development cycle.

Saint-Gobain is prepared to help you on your next project. When you're ready to scale, we offer full validation support and a seamless path to production—so you can launch with confidence.

PROTOTYPING CAPABILITIES

- Functional silicone components delivered in as little as 5-10 business days, depending on part complexity
- Dedicated project engineer ensures responsive communication and technical alignment
- Expert material selection support to meet performance and regulatory requirements
- In-house tooling and molding enables rapid iteration and design optimization
- Seamless transition to production with full IQ/OQ/PQ validation support

Saint-Gobain Performance Plastics Corporation's Life Science ("Saint-Gobain") products that are used as components in the manufacture of any Medical Devices (as defined by the FDA) are sold by Saint-Gobain only and exclusively to Medical Device manufacturers for use in the manufacture, assembly or distribution of their medical devices. Medical Device manufacturers, to whom Saint-Gobain sells components or for whom Saint-Gobain acts as a subcontractor for finished products, are solely responsible for determining whether their finished products are a medical device and complying with the appropriate certifications and registrations.

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