Engineering and High-Performance Plastics for the Healthcare Industry
Röchling • Competence in plastics

**ISO 13485 Certified**
**Shaping the Future through Innovation!**

Roechling stands for innovation in plastics – we continuously strive to offer new and inventive products that meet the highest standards and changing requirements of the Life Science industry.

We are certified to ISO 13485: 2016, the Medical Device Standard that represents the requirements for a comprehensive management system for the design and manufacturing of medical devices. As a stock shape manufacturer of rods and sheets, our quality procedures include key elements such as resin and process validation, product inspection, lot and batch traceability (Device History Record), risk management analysis, corrective and preventive actions and customer feedback.

Our customers can be assured that our medical-grade products meet their expectations and comply with regulatory requirements. We offer full lot and batch traceability, detailed certification documents and raw material Certificate of Analysis.

We manufacture a complete line of engineering and high-performance plastics in rods and sheets for applications including surgical instruments and devices, orthopedic sizing trials, imaging and monitoring components, surgical trays and caddies, diagnostic and analytical equipment, pharmaceutical manufacturing and biotechnology equipment.

Roechling Industrial North America shares your commitment to innovation and quality. Increasingly, the physical properties of high-performance plastics such as impact resistance, dimensional stability and resistance to autoclaving pave the way for new medical device and component designs. For questions relating to material selection and specifications, rely on our expert knowledge.

As a member of Roechling, we are a leading global plastics manufacturer with a workforce of nearly 11,000 employees with 90 locations in 25 countries. Today, Roechling ranks among the top international leaders in the field of plastics processing.
Requirements for medical technology

With medical-grade (MG) materials, our product range that has been specifically developed with orthopedic, healthcare technology in mind is at your disposal.

**Biocompatibility**
With the raw materials deployed, we utilize plastics and additives that have already been in use in medical-technical applications for many years now. The raw materials are FDA-compliant, free from heavy metals, and comply with the EU directives.

**Chemical resistance**
Medical-grade materials possess good chemical resistance to various conventional disinfectants and cleansers.

**Sterilization performance**
A great number of our high-performance materials are very easily sterilizable by means of hot steam, ethylene oxide, plasma and gamma rays.

**Traceability**
Roechling offers complete lot and batch traceability for medical-grade materials, from semi-finished products right up to the raw materials.

**FDA compatibility**
The majority of our high-performance materials meet FDA requirements.

Many colors of our MG products have also been tested in regards to these guidelines.1

**Hydrolysis resistance**
The excellent hydrolysis resistance of our MG products is indispensable for the ability to sterilize with hot steam.

**Physiological harmlessness**

**ISO 10993**
Biological evaluation of Medical Devices per ISO 10993

ISO 10993 includes a series of tests for evaluating the biocompatibility of a medical device prior to a clinical trial. Compliance to ISO 10993-5 (tests for in vitro cytotoxicity) is typically the most requested for applications requiring high-performance plastics.

Many of our products have been tested and approved by an independent lab to ISO 10993. Our products are often produced with raw materials that are certified for approval.

**USP Class VI**
Biological tests per USP Class VI

USP tests are used to determine the biological reactivity of plastic materials. Most of our products are USP VI (systemic and intracutaneous toxicity) compliant.

**Food contact guidelines**
Compliance with FDA Regulation CFR21

Based on the data from the raw material supplier, we can provide a certification for direct contact with food for a large number of our materials. It is derived from the FDA regulation "Code of Federal Regulations" 21 CFR, Part 177.

**Biocompatibility**
Within the scope of selecting the suitable material for a medical-technical application, not only the technical requirements have to be considered, but often it is necessary to ensure the material is compatible with the human organism.

The biological assessment of a product is invariably requisite, if there is direct contact of the material or product with the patient.

The extent of such tests depends particularly on the experience already gained for this material in the particular application concerned, and the precise intended use of the medical product (in particular the nature and duration of the physical contact).

The chief basic regulations for biological testing and assessment of materials are ISO 10993 and testing as per United States Pharmacopeia Class VI (USP Class VI for short). Despite the fact that the clearly more comprehensive ISO 10993 was originally meant to supersede testing as per USP Class VI, USP testing is very frequently referred to today to assess plastics.

There are various test results for biocompatibility according to ISO 10993 and USP Class VI available for the medical-grade materials from Roechling.

**Ultrasonic Testing**
We perform ultrasonic testing on all medical-grade products to ensure that there are no cracks, voids, or impurities.
Material selection

High performance plastics are generally defined by their ability to maintain their physical properties under thermal, chemical or electrical stress, while operating at elevated temperatures above 300° F. These materials typically feature high strength and stiffness as well as outstanding chemical resistance and electrical properties.

Engineering plastics are among the most common and useful thermoplastics and typically exhibit good mechanical properties. These materials generally have one or two main attributes that best fit the needs of the application and should be taken into consideration when deciding on a specific plastic.

Amorphous thermoplastics are mostly transparent or translucent due to their polymer structure. Their mechanical properties remain almost unchanged over a wide temperature range, frequently right up to their continuous operating temperature. They are susceptible to stress cracking and this should be taken into account when machining them. Amorphous plastics include PEI, PSU, PC and PPO.

Partially crystalline thermoplastics are a result of the molecular arrangement, usually being opaque. The mechanical properties (strength, toughness and hardness) of this group depend to a great extent on the degree of crystallinity. They feature great resistance to the formation of stress cracks and very good chemical resistance. Partially crystalline plastics include acetal, PP and PEEK.

**Sustason® PPSU MG (Radel® R5500)**

Sustason® PPSU MG offers incredible toughness in applications that receive repeated sterilization. With a high heat deflection temperature of 420° F it can absorb tremendous impact without cracking or breaking.

**PRODUCT FEATURES:**
- Excellent thermal stability
- High impact resistance
- Resistance to repeated autoclaving
- Resistance to hydrolysis

**CERTIFICATIONS:**
- ISO 10993-5 Certified
- USP Class VI ASTM D6394
- FDA compliant

<table>
<thead>
<tr>
<th>Products</th>
<th>Sizes</th>
<th>Colors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sheets</td>
<td>3/8” – 2” x 24” x 48”</td>
<td>[ ]</td>
</tr>
<tr>
<td></td>
<td>1/4” – 6” diameter</td>
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</tbody>
</table>

Colors available: Natural (bone), Black, Blue, Grey and Green.

**SustaPEI MG (ULTEM™ HU1000)**

SustaPEI MG (ULTEM™) is an amorphous transparent polyetherimide plastic that offers outstanding high heat resistance (up to 356° F), high strength and a broad chemical resistance.

**PRODUCT FEATURES:**
- Strength and modulus at elevated temperatures
- Inherent flame resistance
- Gamma radiation resistance
- Excellent resistance to steam sterilization

**CERTIFICATIONS:**
- ISO 10993 compliant, USP Class VI
- ASTM D5205
- Meets FDA 210 CFR 177.1595

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<tbody>
<tr>
<td>Sheets</td>
<td>3/8” – 4” x 24” x 48”</td>
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</tr>
<tr>
<td></td>
<td>3/8” – 6” diameter</td>
<td>[ ]</td>
</tr>
</tbody>
</table>

Colors available: Natural (amber), Black, Blue, Brown, Yellow and Green.

**SustaPEEK MG**

SustaPEEK MG is increasingly replacing metals and other lower grade plastics in the healthcare industry due to its superior physical properties, including chemical resistance and high temperature stability.

**PRODUCT FEATURES:**
- Continuous use temperature of 480°F
- Outstanding dimensional stability
- High chemical resistance
- Excellent resistance to sterilization

**CERTIFICATIONS:**
- ISO 10993-5 compliant, USP Class VI
- ASTM D6262
- FDA compliant

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</tr>
<tr>
<td></td>
<td>1/4” – 6” diameter</td>
<td>[ ]</td>
</tr>
</tbody>
</table>

Colors available: Natural (tan), Black, Blue, Brown, Yellow and Green.

**Sustason® PSU MG (Polysulfone)**

Sustason® PSU MG is an amorphous semi-transparent thermoplastic with an amber tint. Due to its inherent resistance to hot water and steam, it is regularly used in medical applications where repeated sterilization is required.

**PRODUCT FEATURES:**
- Continuous use temperature of 300°F
- Long-term resistance to steam sterilization
- Resistant to hydrolysis
- Strength and dimensional stability

**CERTIFICATIONS:**
- ISO 10993 compliant, USP Class VI
- ASTM D6394
- FDA compliant

<table>
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<th>Colors</th>
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<tbody>
<tr>
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<td>[ ]</td>
</tr>
<tr>
<td></td>
<td>3/8” – 6” diameter</td>
<td>[ ]</td>
</tr>
</tbody>
</table>

Colors available: Natural (amber) and White
Sustarin® C MG is easy to machine to close tolerances and is very dimensionally stable. The material has excellent wear properties and low coefficient of friction.

**PRODUCT FEATURES:**
- Excellent dimensional stability
- Easy to machine to close tolerances
- Porosity free
- Available in multiple colors

**CERTIFICATIONS:**
- ISO 10993-5 compliant, USP Class VI
- ASTM D6100
- FDA 21 CFR 177.2470 compliant

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**Sustarin® PC MG (Polycarbonate)**

Sustarin® PC is ideally suited for applications that demand high impact strength. It is an amorphous, transparent product that exhibits good electrical and mechanical properties along with excellent dimensional stability.

**PRODUCT FEATURES:**
- Continuous use temperature of 250° F
- Easy to machine to close tolerances
- High impact strength
- Good electrical insulation

**CERTIFICATIONS:**
- ASTM D6098 PC 0111
- ASTM D6098 PC 0111
- Natural meets FDA 21 CFR 177.1582
- USP Class VI compliant

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**Polystone® P MG (Polypropylene)**

The superior physical properties of Polystone® P MG is a compression molded polypropylene that is manufactured by a unique heat stabilization process. This product is easily machined and is specifically designed for surgical trays and caddies.

**PRODUCT FEATURES:**
- Excellent dimensional stability
- Resistant to steam autoclaving
- Laser markable
- Low moisture absorption

**CERTIFICATIONS:**
- ISO 10993-5 certified
- FDA compliant

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**General notes**

All the information contained in this product range has been researched to the best of our knowledge. Nonetheless, errors cannot be completely precluded. For this reason, the information contained in the present product range does not involve any kind of obligation or warranty. Accordingly, we therefore do not undertake any responsibility nor any resultant or any other liability, arising in any manner from utilisation of this information. No responsibility is undertaken either for the completeness of the products, processes, properties, etc. covered. Data concerning weights are purely computed values, ensuing from the density and the mean value of the tolerance dimensions.

Our medical grade semi-finished products are not intended for the manufacture of implants, permanent oral implants, or dental contact of more than twenty-four (24) hours with body fluids and/or tissues/bones. The processors and marketers of medical devices are advised that any intended application must be carefully examined. In the EU, medical products may only be placed on the market if they have been CE-marked in the procedure provided for them and also fulfill all the requirements of the respective relevant legislation. The sole responsibility for the evaluation of the end product for the intended use and compliance with the applicable requirements lies solely with the manufacturer/marketer of the respective product/end product.

**Application of Roehling materials not intended for implants**

The materials described in this product range are not suitable for application as medical implants. Furthermore, they should not be put to use in medical technical fields, necessitating direct, long-term contact of the material with the patient.

**Sterilization and multiple use of medical products**

For classification of the sterilization resistance of our materials, various criteria were referred to, such as change to the mechanical properties, change in weight or loss in transparency (amorphous materials). For these reasons, this assessment only represents recommendations and not definite commitment for the suitability of a material for a specific reprocessing procedure. Should the medical product be reused, it is incumbent upon the manufacturer of the product to determine the suitability and the number of possible reprocessing cycles for a process.
Material properties and applications

<table>
<thead>
<tr>
<th>Material</th>
<th>Trade Name</th>
<th>Common Name</th>
<th>Specific Gravity</th>
<th>Tensile Strength</th>
<th>Tensile Modulus</th>
<th>Tensile Elongation Break</th>
<th>Flexural Strength</th>
<th>Flexural Modulus</th>
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</thead>
<tbody>
<tr>
<td>PPSU</td>
<td>Sustason® PPSU MG</td>
<td>Radel® R5500</td>
<td>1.29</td>
<td>11,000 PSI</td>
<td>390,000 PSI</td>
<td>30</td>
<td>15,500 PSI</td>
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<tr>
<td>POM C</td>
<td>Sustarin® C MG</td>
<td>Hostaform® MT2U06</td>
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<tr>
<td>PEEK</td>
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<tr>
<td>PEI</td>
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<td>16,700</td>
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<td>80</td>
<td>20,000</td>
<td>500,000</td>
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<tr>
<td>PSU</td>
<td>Sustason® PSU MG</td>
<td>Polysulfone</td>
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<td>360,000</td>
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<tr>
<td>PC</td>
<td>Sustanat® PC MG</td>
<td>Polycarbonate</td>
<td>1.20</td>
<td>10,000</td>
<td>320,000</td>
<td>75</td>
<td>13,000</td>
<td>340,000</td>
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<tr>
<td>PP</td>
<td>Polystone® P MG</td>
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<td>0.91</td>
<td>4,700</td>
<td>232,000</td>
<td>–</td>
<td>–</td>
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<table>
<thead>
<tr>
<th>Material</th>
<th>Trade Name</th>
<th>Common Name</th>
<th>Rockwell Hardness</th>
<th>Izod Impact Notched</th>
<th>Heat Deflection Temp @ 66 psi</th>
<th>Heat Deflection Temp @ 264 psi</th>
<th>Continuous Use</th>
<th>Volume Resistivity</th>
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<tbody>
<tr>
<td>PPSU</td>
<td>Sustason® PPSU MG</td>
<td>Radel® R5500</td>
<td>R120</td>
<td>13.0</td>
<td>417</td>
<td>420</td>
<td>320</td>
<td>10^16</td>
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<tr>
<td>POM C</td>
<td>Sustarin® C MG</td>
<td>Hostaform® MT2U06</td>
<td>R120</td>
<td>1.2</td>
<td>320</td>
<td>225</td>
<td>180</td>
<td>10^16</td>
</tr>
<tr>
<td>PEEK</td>
<td>SustaPEEK MG</td>
<td>PEEK</td>
<td>R126</td>
<td>1.2</td>
<td>360</td>
<td>320</td>
<td>480</td>
<td>10^16</td>
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<tr>
<td>PEI</td>
<td>SustaPEI MG</td>
<td>ULTEM™ HUI000</td>
<td>R123</td>
<td>0.6</td>
<td>405</td>
<td>395</td>
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<td>10^16</td>
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<td>PSU</td>
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<td>Polysulfone</td>
<td>R125</td>
<td>1.3</td>
<td>359</td>
<td>345</td>
<td>300</td>
<td>10^16</td>
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<td>PC</td>
<td>Sustanat® PC MG</td>
<td>Polycarbonate</td>
<td>R126</td>
<td>10.0</td>
<td>295</td>
<td>280</td>
<td>250</td>
<td>&gt;10^13</td>
</tr>
<tr>
<td>PP</td>
<td>Polystone® P MG</td>
<td>Heat-stabilized Polypropylene</td>
<td>–</td>
<td>1.9</td>
<td>221</td>
<td>149</td>
<td>180</td>
<td>–</td>
</tr>
</tbody>
</table>

Statistics based on resin data, not stock shape. Individual testing for stock shapes can be provided upon request.

Applications for medical technology

Today, finished products machined from our plastics are used in a host of medical devices and instruments. For applications requiring proven biocompatibility we offer a variety of medical-grade materials as well as a comprehensive offering of standard plastics for applications that do not come into direct contact with patients.

**Surgical instruments and supplies**
- Handles and grips for instrumentation
- Sizing trials for knee and hip replacement
- Endoscopic housings and eyepieces
- Sterilization trays and caddies

**Diagnostic**
- Parts for X-ray and MRI devices
- Components for supports and biopsy units

**Therapeutic systems**
- Blocks and housings for dialysis machines
- Pistons and valves for anesthetic equipment
- Supports and adaptors for respiratory units

**Dental**
- Grips and handles for dental instruments
- Components for treatment and therapy units

**Pharmaceutical and biotechnology**
- Components for sample changers
- Valve housings and nozzles for fluid distribution
- Spectrometer parts for chromatography systems
- Wear parts for pill and tablet production
Biocompatibility

Biocompatible implies that a medical device is safe for human use. The ISO 10993 standard plays an important role in the assessment of the biocompatibility of a medical device through a series of tests depending on the intended use and the time that it is exposed to the human body. Most importantly, the ISO 10993-5 test method assesses the in vitro cytotoxicity of medical devices and is designed to determine the biological response of mammalian cells in vitro using appropriate biological parameters.

Sterilization and disinfection

An essential aspect of selecting a suitable plastic for a medical technology application is also the requirement for repeated sterilization and disinfection of the product. The cleaning process typically occurs in autoclaving devices at elevated temperatures (greater than 250°F) with steam or suitable disinfectants. The resistance of the polymer should be checked in each case.

Selection of the tests as per ISO 10993-5

<table>
<thead>
<tr>
<th>Nature of the physical contact</th>
<th>Biological risks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin</td>
<td>Cytotoxicity</td>
</tr>
<tr>
<td></td>
<td>Irritation or inducible reactivity</td>
</tr>
<tr>
<td></td>
<td>Acute Toxicity</td>
</tr>
<tr>
<td></td>
<td>Genotoxicity</td>
</tr>
<tr>
<td></td>
<td>Hemocompatibility</td>
</tr>
<tr>
<td></td>
<td>Genotoxicity</td>
</tr>
<tr>
<td></td>
<td>Reproductive/Developmental Toxicity</td>
</tr>
<tr>
<td>Medical products with contact to body surfaces</td>
<td>Duration of contact</td>
</tr>
<tr>
<td>Skin</td>
<td>A x x x x x x x</td>
</tr>
<tr>
<td>Mucous membrane</td>
<td>B x x x x x x x</td>
</tr>
<tr>
<td>Injured surface</td>
<td>C x x x x x x x</td>
</tr>
<tr>
<td>Blood system indirectly</td>
<td>A x x x x x x x</td>
</tr>
<tr>
<td>Tissue / bone / dentin</td>
<td>B x x x x x x x</td>
</tr>
<tr>
<td>Circulating blood</td>
<td>C x x x x x x x</td>
</tr>
<tr>
<td>Implantable medical products</td>
<td>A x x x x x x x</td>
</tr>
<tr>
<td>Tissue / bone</td>
<td>B x x x x x x x</td>
</tr>
<tr>
<td>Blood</td>
<td>C x x x x x x x</td>
</tr>
</tbody>
</table>

Test to be included to ISO 10993-5
A = short-term (= 24 h)  B = protracted (> 24 h to 30 d)  C = continuous (> 30 d)

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