



Safe equipment and systems

All parts of a production line that come into contact with drug substances or drug products must be designed, tested, and documented as not altering the safety or quality of the finished product.

SAXOCON provides you with everything you need to establish and document compliance with regulatory requirements for pharmaceutical manufacturing systems, including:

- The selection and characterisation of construction materials
- A risk analysis in accordance with European and US Pharmacopeia standards
- Extractables test strategies
- A review of protocols and monitoring of extractables studies
- A toxicological risk assessment of the results

Why choose us?

SAXOCON services for manufacture of pharmaceutical products give you access to:

- A multidisciplinary team of highly skilled toxicologists, material scientists, and supply chain professionals
- Extensive experience with pharmaceutical product safety assessments
- Safety evaluations based on proprietary information obtained from our worldwide network of materials suppliers
- State-of-the-art solutions according to international standards and regulatory guidelines
- Smart, cost-effective test strategies that bring your products to market in a timely manner

Delivery

SAXOCON compiles all necessary documentation in a written report for you to use in your Good Manufacturing Practice file.





