Toxikon is a preclinical contract research organization (CRO). We contract and partner with biotech, pharmaceutical and medical device industries to deliver exceptional product development services from concept to final product.

No matter what your development needs, you can rely on Toxikon to be a flexible partner with technical research and development expertise to meet your project goals for timely market launch.

We not only provide you with excellent service, we also work closely and collaborate with you so that we understand your needs to develop a comprehensive testing program well aligned to meet your development objectives. As a leading preclinical CRO, Toxikon has the experience and is a proven life science leader with a staff of scientific distinction who can provide in vivo, in vitro, analytical, and a host of other services central to the development of products designed to improve the quality of life.

With Toxikon you gain a strategic and responsive scientific partner, who not only brings reliability, trust, and flexibility to your projects, but also the assurance to align with your corporate goals, achieve product safety and regulatory compliance, and maintain the highest levels of environmental and manufacturing quality control.

Whether you are a pharmaceutical, medical device or biotech company, collaborating with Toxikon provides you access to a team of experts experienced in a comprehensive range of compound structures, therapeutic targets as well as medical devices and combination products.

**A Resourceful CRO**

- Toxikon’s Massachusetts campus, located in the heart of Bay State’s life science research hub, has more than 125,000 square feet of facility space dedicated to the development of compounds, biologics, and medical devices;
- Toxikon Europe operates a 14,000-square-foot facility in Leuven, Belgium, to meet the needs of European pharmaceutical, biotech and medical device industries;
- An experienced international team;
- A highly educated and trained workforce;
- Broad technical expertise with recognized scientific excellence;
- World-class facilities in the United States and Europe;
- Client/Sponsor-centric;
- Collaborative and research-oriented culture.
Services at a Glance
Toxikon is an ISO/IEC 17025-accredited Contract Research Organization (CRO) registered with the FDA for drug, biologics, and medical device testing. With approximately 200 employees, Toxikon is committed to staying at the forefront of life science technology development. Toxikon's safety services include toxicology (acute, subchronic, and chronic toxicity, reproductive toxicity, genetic toxicology, carcinogenicity), pharmacokinetics, toxicokinetics, bioavailability, ADME, chemical characterization, impurities analysis and synthesis, bioanalysis, and microbiology. Toxikon’s extensive certifications and licenses allow it to meet international regulatory requirements for data acceptability and harmonization.

To meet all applicable international GxP requirements, Toxikon’s facilities are fully registered with FDA and USDA, and accredited with the Association for the Assessment and Accreditation of Laboratory and Animal Care International (AAALAC), and the International Organization for Standardization (ISO). Toxikon also has assurance with the National Institutes of Health so federally funded work can be performed at our facilities in compliance with NIH’s Public Health Service Policy on Human Care and Use of Laboratory Animals (PHS Policy).

Please visit www.toxikon.com to learn more about our preclinical development capabilities.