

Toxicology Facilities at WuXi AppTec – Suzhou (China)

WuXi AppTec is a global leader in providing discovery, testing and manufacturing services for the pharmaceutical, biotechnology and medical device industries. Research-driven and customer-focused, with operations in China and the U.S., WuXi AppTec offers a broad and integrated portfolio of services designed to assist our customers with cost-effective and efficient outsourcing solutions.

WuXi AppTec has the vision to offer its customers fully integrated preclinical safety evaluation services with world-class capabilities and unparalleled capacity.

- Cost-effective operation
- Best-in-class preclinical CRO
- NHP center



**WX
LISTED
NYSE**

323,450 ft² – One of the largest AAALAC-accredited preclinical safety evaluation center in Asia

➤ This state-of-the-art facility, located in Suzhou, about 60 miles from Shanghai, offers:

- General Toxicology Services
- Genotoxicity Services
- Safety Pharmacology Services
- Tissue-cross Reactivity (TXR) Services
- DMPK Services

➤ Features include:

- Integrated services from API synthesis through preclinical safety evaluation
- 90 non-rodent and 38 rodent study rooms
- NHP center with unparalleled NHP resources
- Dedicated client facilities
- Board-certified toxicologists and pathologists

For more information on WuXi AppTec's services please contact:

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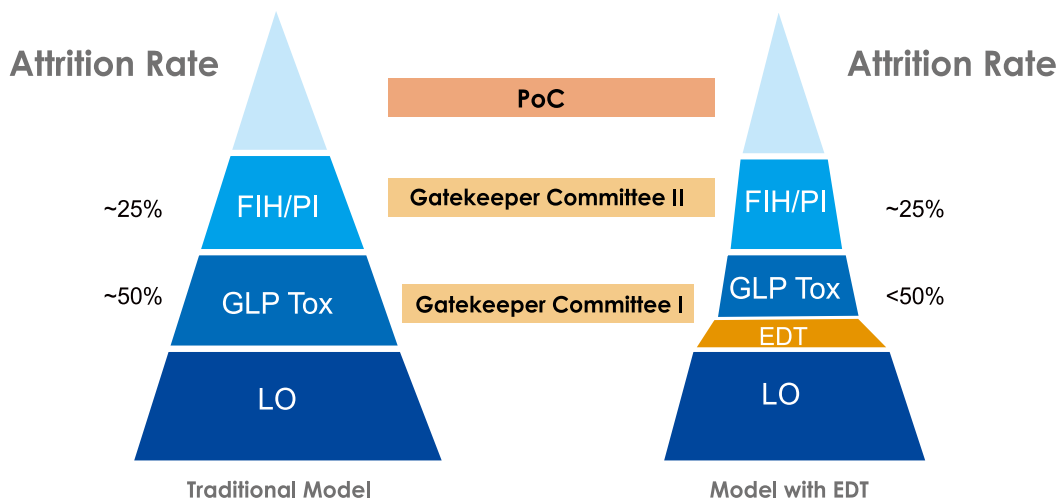
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WuXi AppTec's Suzhou Toxicology Facility

Growth Strategy

- Currently conducting non-GLP safety evaluation studies such as “early diagnosis tests” (EDT), exploratory toxicity, and dose range-finding studies in rodents and non-rodents
- GLP genotoxicity study services
- GLP General Toxicology study services and Safety Pharmacology including CV Telemetry
- Full preclinical safety evaluation services including DART in 2013



Highlights of Our Current Services

- “Early diagnosis tests” (EDT) with small quantity of API to evaluate three major liabilities in 2 ½ to 3 months from start of synthesis to report:
 - Genotoxicity *in vitro*
 - QT prolongation *in vitro* and *in vivo*
 - Target organs and dose-limiting toxicities in rodents and non-rodents
- Non-GLP exploratory/range-finding toxicity studies in rodents and non-rodents
- NHP center with >1500 consistent and high-quality NHPs available per year

Our NHP Center Offers

- Long-term, ready-source, purpose-bred NHP colonies
 - Consistent, high quality NHPs
 - 50-day quarantine period
 - Free from Herpes-B virus, TB, and diarrhea
 - >1500 available per year
- 68 NHP rooms



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LT557-11.30.09