A flexible approach to
Generic Product Development

QPS IS A GLOBAL CRO WITH DIVERSE CAPABILITIES THAT CAN OFFER END TO END SERVICES TO GENERIC COMPANIES, enabling them to bring products to market in a faster and more compliant manner. With operations in India, USA, Europe and Taiwan, QPS is ideally positioned to address Sponsor’s global generic product development requirements of quality, compliance, time to market.

TIME IS OF THE ESSENCE IN DRUG DEVELOPMENT. CONTACT THE QPS BUSINESS DEVELOPMENT TEAM TODAY!

CALL +1 512 350 2827   EMAIL infobd@qps.com
QPS has the experience and expertise to work with Sponsors in the global development of a diverse range of products from conventional generics to more complex products including modified release formulations, novel drug delivery systems for NCE -1, ANDA submissions.

Our comprehensive services include:

- Study Design
- Medical Writing
- Clinical Study Conduct
- Bioanalysis
- Data Management & CDISC
- PK & Statistical Analysis
- Final Report and Dossier Preparation

Our services for generic drug development include:

- BA/BE in healthy subject populations
- BA/BE in patient populations
- Clinical Endpoint Studies
- 505(b)2 NDA submissions
Clinical

Our Clinical Phase I units offer over 700 beds globally and can support a wide variety of clinical study designs:

<table>
<thead>
<tr>
<th>Clinical Unit</th>
<th># of Beds</th>
<th>Study Types</th>
</tr>
</thead>
<tbody>
<tr>
<td>QPS Missouri</td>
<td>240</td>
<td>Healthy Human BA/BE</td>
</tr>
<tr>
<td>QPS India</td>
<td>138</td>
<td>Healthy Human BA/BE, Patient Pharmacokinetic Studies, Clinical Endpoint Studies</td>
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<tr>
<td>QPS Taiwan</td>
<td>40</td>
<td>Patient Pharmacokinetic Studies and Clinical Endpoint Studies</td>
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<td>QPS Netherlands</td>
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<td>QPS Miami</td>
<td>95</td>
<td>Healthy Human BA/BE, Patient Pharmacokinetic Studies, Clinical Endpoint Studies</td>
</tr>
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Bioanalysis

We offer state of the art GLP compliant bioanalysis at multiple global locations including India, USA, the Netherlands and Taiwan. With over 46 LC-MS/MS systems installed globally and 800+ validated assays spanning all therapeutic areas including challenging and ‘hard to develop’ methods, we are positioned as the ideal partner for generic product development.

Credentials

All our global sites have an excellent regulatory track-record with multiple inspections by various agencies including US FDA, EMA, UK MHRA, WHO and CEB, resulting in numerous product approvals for our Sponsors.

QPS is your ideal CRO partner for generic product development and can also assist in understanding the regulatory requirements of different countries for generic drug approval. QPS can partner with your product development groups in taking you from study design to providing a final dossier for regulatory submission.

AGILITY. FLEXIBILITY. SPEED.
QPS is a Global CRO with locations around the world to serve the evolving needs of the Pharmaceutical and Biotech industries.