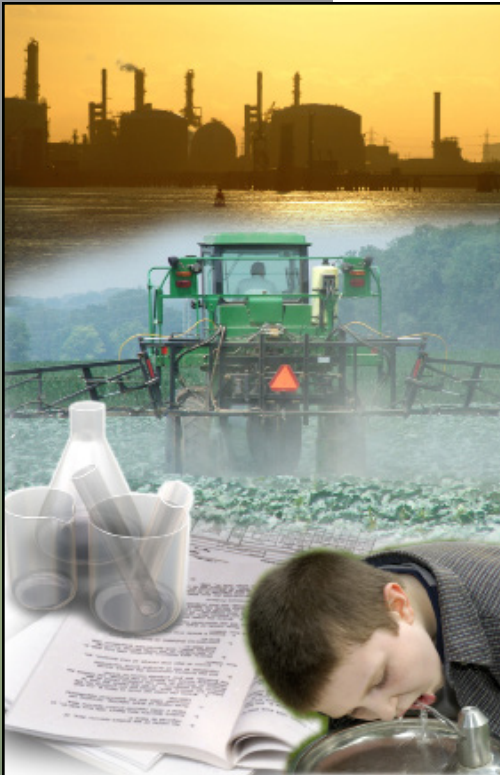




# ***TOXICOLOGY CONSULTING SERVICES***

*Regulatory & Scientific Specialists*



*Support & Planning for REACH, PPPD, BPD, GHS, EPA Registration Review Rule, HPV*

## **Preparation of Toxicology Dossiers**

Preparation of high quality active ingredient and product dossiers to meet international requirements. Formats: OECD, US EPA, EU, WHO/JMPR, IUCLID.

## **Risk Assessment**

Human Health Risk Assessments for the consumer, worker, operator and bystander: hazard identification, dose-response, human exposure modeling, risk characterization and risk mitigation.

## **Management of GLP Product Safety Studies**

Design, placement and scientific monitoring of product development programs to meet international testing requirements.

## **Chemical Candidate Selection**

Co-ordination of toxicology screening for candidate selection. Preliminary risk assessments. Regulatory acceptability of inerts.

## **Data Package Completeness**

Data gap analysis and quality checks of toxicology packages and advice on regulatory acceptance.

## **Product Defense**

Product stewardship, preparation of toxicological position papers and liaison with regulatory authorities.

## **Classification and Labeling**

Expert support on classification and labeling (including CMR) defense issues. Safety Data Sheet preparation.

## **Literature Review**

Evaluation of public domain scientific data in support of toxicological dossiers.

## **Offices in the USA and the UK**

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**Best Supporting Role**