



Low Cost



Minimal Compound




Rapid Turnaround



Multiple Exposures

**devTOX<sup>qP</sup>** is a biomarker-based human *in vitro* assay for prediction of developmental toxicity that utilizes key biomarkers identified using our metabolomics platform.

## COMPARING *IN VITRO* MODELS

Assay	Model	Endpoints	Accuracy*
	Human pluripotent stem cells	Quantitative, metabolic biomarkers	89%
ZET	Non-mammalian embryos	Subjective, embryo morphology	75%
mEST	Non-human embryonic stem cells	Subjective, cardiomyocyte differentiation	74%
rWEC	Non-human embryos	Subjective, embryo morphology	73%

ZET: Zebrafish Embryotoxicity Test; mEST: Mouse Embryonic Stem Cell Test; rWEC: Rat Whole Post-Implantation Embryo Culture Assay

\* Palmer JA, Smith AM, Egnash LA, Conard KR, West PR, Burrier RE, Donley EL, Kirchner FR. Establishment and assessment of a new human embryonic stem cell-based biomarker assay for developmental toxicity screening. Birth Defects Res B Dev Reprod Toxicol. 2013; 98(4): 343-363.

## devTOX<sup>qP</sup> provides

- Accurate, sensitive, and specific results
- Data for multiple exposure levels
- Positive and negative controls
- Human test system
- Rapid turnaround

**Flexibility** Choice of human embryonic stem (hES) cells or induced pluripotent stem (iPS) cells. Assay can be **customized** for your test articles and testing requirements.

**Sensitivity** The assay is performed using human pluripotent stem cells exposed to eight different concentrations of each test compound to yield a **broad look at developmental toxicity potential**.

**UPLC/HRMS** Ultra performance liquid chromatography, coupled with high-resolution mass spectrometry, results in **selective and reproducible biomarker measurement**.

## Data Analysis & Reporting

- SOP-driven analysis
- LIMS-controlled data analysis pipeline
- Identification of critical exposure where cellular metabolism is altered
- Uniform reporting for rapid turnaround
- Custom reporting available

**Assay validation** Our assay has been internally validated with more than 80 compounds with published human/*in vitro* developmental toxicity data.

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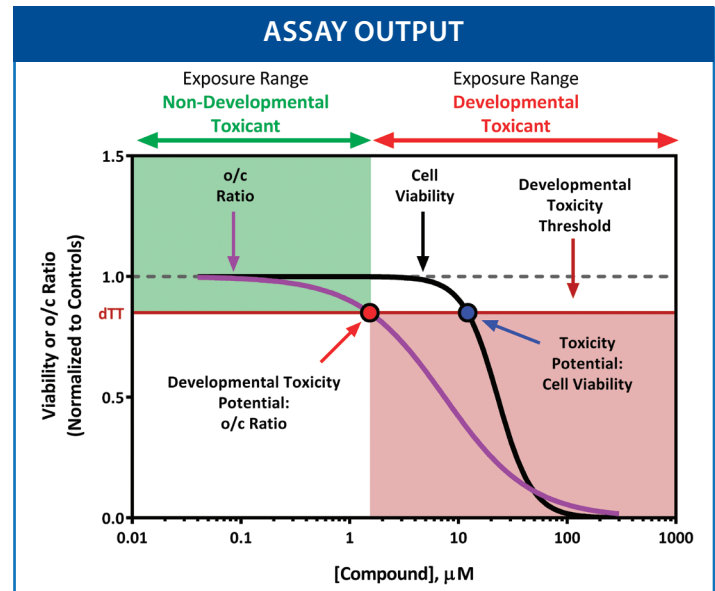
## How it works

- Human pluripotent stem cells are exposed to eight concentrations of each test article.
- Spent media is collected to determine changes in ornithine and cystine via LC-MS analysis and cell viability is measured.
- Non-linear dose-response analysis of the the biomarker ratio response.
- A test article shows developmental toxicity potential where the curve drops below the threshold.

Predictive Power				
	N	Accuracy	Sensitivity	Specificity
hESC	79	85%	83%	86%
iPSC	112	86%	84%	87%

COMPARISON TO PUBLISHED <i>IN VIVO</i> RESULTS				
Compound	devTOX <sup>OP</sup> QUICKPREDICT	<i>In vivo</i>		
		Human	Rodent	Rabbit
Diphenhydramine	NON	NON	NON	NON
Doxylamine	NON	NON	NON	NON
All- <i>trans</i> Retinoic Acid	DT	DT	DT	DT
Hydroxyurea	DT	DT	DT	DT
Methotrexate	DT	DT	DT	DT
Thalidomide	DT	DT	NON	DT
Warfarin	DT	DT	DT	NON

DT = Developmental Toxicant



## Commitment to Quality

From start to finish, Stemina has a well-defined, quality program to ensure data integrity. Compounds in our test set and controls show excellent reproducibility over time.

## Experience Counts

Our team has extensive experience in screening a wide variety of proprietary compounds including pharmaceuticals, agriculturals, tobacco products, consumer products, and cosmetic ingredients. Stemina was founded in 2006; its state-of-the-art facilities are located in the United States.

## Extending Our Global Reach

Stemina has partnered with CiToxLAB, which has facilities in Canada, France, Denmark, and Hungary, to provide worldwide service.

## EPA ToxCast™ Contractor