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FOR IMMEDIATE RELEASE

Contact:

Diana Caldwell

Pearl Pathways

Business: (317) 899-9341

Cell: (317) 490-0511

[contact@pearlpathways.com](mailto:%20contact@pearlpathways.com)

[www.pearlpathways.com](http://www.pearlpathways.com/)

**Pearl Pathways Hires Three New Associates**

*The diverse backgrounds of our new teammates serve growing demands of Pearl Pathways' clients.*

INDIANAPOLIS, INDIANA –February 25, 2013- Pearl Pathways announces the recent hiring of Daphine Glowner, Gretchen Parker, PhD, RAC and Sarah Witwer, JD, RAC.

Glowner joined Pearl Pathways in January 2013 as an Associate and is an experienced publications specialist, editor, and project manager in the life sciences industry. Her previous roles in clinical research, publications, and regulatory have allowed her to gain expertise in database management and bibliographic coordination.

Parker received her PhD in biochemistry and molecular biology with an emphasis on endocrinology, and she is a seasoned regulatory and scientific medical writing professional. She joined the company in February 2013. In addition to her role as a Regulatory Compliance Advisor, Parker serves as a co-chair board member of [Pearl IRB](http://www.pearlirb.com/).

Witwer also joined Pearl Pathways in February 2013 as a Regulatory Compliance Advisor. She has 30 years’ experience in drug and device development and proven proficiency in authoring and negotiating submission packages, as well as managing Chemistry Manufacturing and Controls (CMC) manufacturing issues.

Glowner’s editorial and project management skills combined with Parker’s scientific technical experience and Witwer’s regulatory prowess add to the strong team at Pearl Pathways. Diana Caldwell, President and CEO shares “Last month, we were proud to announce our partnership with the state of Indiana when we received conditional tax credits and training grants based on our job creation plans. These three additions so early in the year are a great start. Our clients are fortunate to have the opportunity to work with such knowledgeable talent.”

**About Pearl Pathways**

***Pearl Pathways*** is a comprehensive life science product development services company. Our experienced team is obsessed with expediting life science product development regulatory pathways. We have three business units to serve biopharma and medical device clients:

***Pearl IRB*** is a full service commercial Independent Review Board that provides human research IRB reviews, IRB exemptions and waivers, and also offers support for research protocol/ICF medical writing, site assessments, and monitoring services.

***Pearl ReGXP*** is a regulatory and quality compliance consulting practice that provides regulatory filing guidance, conducts global health authority negotiations, develops/improves quality systems, and delivers GMP/GLP/GCP auditing services.

***Pearl IDEAS*** provides strategic product development assistance, third party vendor selection and management strategies, due diligence services, and sales and marketing services for drug, biologic and device companies.

To learn more, please visit [www.pearlpathways.com](http://www.pearlpathways.com/), call 317.899.9341, or email [contact@pearlpathways.com](mailto:%20contact@pearlpathways.com). Pearl Pathways is located in Indianapolis, Indiana, and is a WBENC certified woman owned business. For media inquiries, contact Diana Caldwell at [contact@pearlpathways.com](mailto:contact@pearlpathways.com)