Pathway-based Approaches to Safety Assessment: development and use

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Outline

• The need for a new approach to toxicology
• The Adverse Outcome Pathway concept
• Precedents and projects
• Thoughts on how to build a pathway
• OECD guidance
• Examples
  • Skin sensitization
  • ER mediated reproductive impairment
• Requirements for different uses
• What’s needed for the future
The argument for a new approach

Pharmaceuticals:
- 92% of drug candidates fail in clinical studies
- “The average drug developed by a major pharmaceutical company costs at least $4 billion, and it can be as much as $11 billion” (Forbes 2012)
- Need to assess novel chemistries (i.e. nanomaterials)

Industrial chemicals:
- Growing concern over lack of data (> 10K chemicals worldwide)
- Large-scale regulatory programs: REACH (EU, China, S.Korea)

Pesticides:
- Registration requires the use of approximately 10,000 animals, millions of USD, and many years (decades)
- Need to identify “greener” chemistries

Cosmetics:
- European Cosmetics Directives ban on animal testing
- Consumer concern over safety and animal testing worldwide
The opportunity for a new approach

- Capitalize on advances in chemistry, biology, and engineering (since ~1970)
- Fully utilize all existing knowledge
- Increase relevance to humans
- Increase assessment capacity (“throughput”)
- Increase efficiency (benefit/cost)
- Increase predictivity

Decrease uncertainty in hazard and risk assessment
The Adverse Outcome Pathway concept

A conceptual construct that portrays knowledge of linkages between a direct molecular initiating event and an adverse outcome at a level of biological organization relevant to risk assessment.

Precedents for pathway-based toxicology

1. Dose-response modeling
   • Using pharmacokinetic and mechanistic information

2. IPCS/WHO mode of action frameworks
   • Human relevance of rodent cancer findings
   • Extrapolated to non-cancer endpoints

3. Mode of action pathways in drug development
   • Drug and target-specific

   “envisions a new toxicity-testing system that evaluates biologically significant perturbations in key toxicity pathways by using new methods in computational biology and a comprehensive array of in vitro tests based on human biology”
Pathway projects and workshops

1. OECD Test Guidelines Programme
   - 2010 Workshop on using mechanistic information in forming chemical categories
   - Extended Advisory Group on Molecular Screening and Toxicogenomics,
     - VMG-non-animal under the EDTA-AG

2. JRC-SEURAT 2012 workshop: Describing mode-of-action in liver toxicity using adverse outcome pathways
   - Fibrosis and steatosis as prototypes
   - Way to organize and integrate SEURAT data

3. CAAT 2012 Workshop: Concept and Tools for Pathways of Toxicity
   - Combination of toxicity pathway and ‘omics approaches
   - Estrogen signaling as prototype

4. The Hamner Institutes: “Tier 1 and Done”
   - Estrogen signaling pathway as prototype
   - Including dose-response extrapolation modeling

5. HTPC 2013 workshop: Building Shared Experience to Advance Practical Application of Pathway-Based Toxicity: Liver Toxicity Mode-of-Action
Building a Pathway

1. What basic elements are needed for pathway development?

2. Where do you start?
   - How do you determine which MIEs, pathways to focus on?
   - E.g. do you start with one MIE and develop pathways for each AO?

3. How and where to limit the pathway
   - Must every pathway begin at the MIE and end with an AO
     o at the individual or population level?
   - When is it appropriate to include branching?

4. How to evaluate and assessed the completeness and confidence of a pathway?

5. How to identify the level of completeness and which uses would be appropriate?
OECD AOP project

Guidance
• Template for building
• Criteria for evaluating
• Glossary of terms

OECD Series on Testing and Assessment No. 184. 2013. Guidance Document on Developing and Assessing Adverse Outcome Pathways (available online)
OECD AOP development template

1. Three basic elements:
   a. MIE $\leftrightarrow$ intermediate events $\leftrightarrow$ Adverse Outcome

2. Begin from any of these elements
   1. MIE: molecular description of how the chemical interacts with the initial biomolecule
   2. AO: specific and well-defined outcome, associated with OECD TG endpoint

3. An AO results from a finite number of MIEs, and conversely an MIE results in a finite number of AOs, but an AOP is limited to a single MIE $\rightarrow$ a single AO

4. Information from different levels of biological organization are integrated into a single description
OECD AOP reporting

Data summation:
  • Assays that are fit for purpose, repeatable, reproducible, and directly or indirectly linked to AO
  • WoE supports the evidence used

AOP assessment:
  • Reliability and robustness
  • Strength of qualitative and quantitative understanding (Bradford-Hill criteria):
    • strength of association
    • consistency of the evidence
    • specificity of the relationship
    • consistent temporal relationships
    • dose-response relationships
    • biological plausibility
    • coherence of the evidence
    • and consideration of alternative explanations
Other practical considerations

1. Annotation of pathway should include
   a. Well defined terminology
   b. Diagram(s)
   c. Language for representing multi-dimensions including temporal
   d. Explanation of how the each step was deduced

2. Quality assessment of input data
   a. Evidence Based Toxicology?
   b. Klimisch score?

3. Quality of and confidence in causal linkages
   a. Bradford-Hill criteria
   b. Human relevance

4. Consideration of Scope
   a. species, developmental stage, sex, chemical space limitations

5. Temporal hierarchy
   a. E.g. gene expression changes that precede cellular changes)

6. Quantitative linkages
   a. Threshold and scale
OECD AOP sensitization project

OECD 2012. The Adverse Outcome Pathway for Skin Sensitisation Initiated by Covalent Binding to Proteins.
OECD sensitization project

Sensitisation ITS
- Aeby et al. (2010). Toxicol In Vitro 24, 1465 – 1473
- Bauch et al. (2011). Toxicol In Vitro 25, 1162–1168
- McKim et al. Cutan Ocul Toxicol Apr 12. [Epub ahead of print]
US EPA ER-mediated reproductive impairment

P. Schmieder, McKim conference 2008.

IVITP Spring Meeting 2013 | May 15 - 16 | Southampton, UK
OECD AOPs in development

Extended Advisory Group on Molecular Screening and Toxicogenomics

• Mitochondrial toxicity – OECD
• Cell proliferation/differentiation – OECD
• Fish reproductive toxicity – US EPA
• Thyroid hormone pathways – US EPA
• PPARα – OEDC, Hamner
• Cancer epigenetics – S.Korea
• Germ cell mutagenicity – Canada
• Neurotoxicity and inflammation – Switzerland
• Liver Steatosis and Fibrosis – JRC
• AhR – BIAC
• Aquatic toxicity: UK and Japan
• Mutagenic MOA: US
• PPARα/CAR: US
• Embryonic vascular development: US
Uses of AOPs

Near-term use:
- Inform chemical categories and structure activity relationships
- Hazard identification
- Prioritization of chemicals for further assessment
- Increase certainty of interpretation of both existing and new information
- Develop integrated testing strategies that maximize useful information gained from minimal testing

Longer-term use:
- Identify key events for which non-animal tests can be developed, thereby facilitating mechanism-based, non-animal chemical assessment
- Create predictive toxicological assessments with low uncertainty and high human relevance
- Eventually without the use of animals
Use $\propto$ strength/type of information

Chemical categories
Hazard identification
Prioritization
Integrated strategy design

Molecular initiating event $\rightarrow$ Intermediate event(s) $\rightarrow$ Adverse outcome

Risk assessment
ID key events that link pathways
Predictive system for toxicology
Use $\propto$ strength/type of information

Chemical categories

Molecular initiating event → Intermediate event(s) → Adverse outcome
Use $\propto$ strength/type of information

Hazard identification
Prioritization

Molecular initiating event
Assay 1
Intermediate event(s)
Assay 2
Adverse outcome
Use $\propto$ strength/type of information

Integrated strategy design

Molecular initiating event

Intermediate event(s)

Adverse outcome

Assay 1

Assay 2

Assay 3

Assay n
Use $\propto$ strength/type of information

Risk assessment

Molecular initiating event $\rightarrow f(MIE)$ $\rightarrow f(I\_1)$ $\rightarrow f(I\_2)$ $\rightarrow f(I\_\ldots)$ $\rightarrow$ Adverse outcome

Assay 1 $\rightarrow$ Assay 2 $\rightarrow$ Assay 3 $\rightarrow$ Assay $\ldots$ $\rightarrow$ Assay $n$
Use $\propto$ strength/type of information

ID key events that link pathways

Pathway A

- Molecular initiating event
- Assay 1
- Assay 2
- Assay 3
- Assay n

Pathway B

- Molecular initiating event
- Assay 1
- Assay 2
- Assay 3
- Assay n

IE 1 $\rightarrow$ IE 2 $\rightarrow$ IE ... $\rightarrow$ Adverse outcome

IE 1 $\rightarrow$ IE ... $\rightarrow$ Adverse outcome
Use $\propto$ strength/type of information

Predictive system for toxicology

Pathway B
- Molecular initiating event
- $f(MIE)$
- $f(MIE)$
- $f(MIE)$

Pathway A
- Molecular initiating event
- $f(MIEa)$
- $f(MIEa)$
- $f(MIEa)$

Pathway n
- Molecular initiating event
- $f(MIEn)$
- $f(MIEn)$
- $f(MIEn)$

IE 1
- $f(IE 1)$
- $f(IE 1a)$
- $f(IE 1b)$

IE 2
- $f(IE 2)$
- $f(IE 2a)$
- $f(IE 2b)$

IE...
- $f(IE...)$
- $f(IE...)$
- $f(IE...)$

Adverse outcome
- $f(IE...)$
- $f(IE...)$
- $f(IE...)$

ETC...

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What’s needed for the future

• **Build a series of prototype pathways**
  - OECD / EPA / FDA / industry / academia
  - OECD Guidance

• **Improve predictive tools**
  - NIH National Center for Advancing Translational Sciences
  - EPA’s Computational Toxicology Research
  - OECD QSAR tool box
  - The Hamner Institutes

• **Develop assessment systems for complex endpoints**
  - Reconstructed tissues and organ systems

• **Integrate absorption, metabolism and distribution information**
  - QSAR
  - Liver cells, tissues, extracts, reconstructed tissues

• **Integrated databases and “knowledge bases”**
  - **ACToR and MetaPath**: EPA – all available chemical toxicity data on over 500,000 environmental chemicals searchable by chemical name and structure
  - **Kegg pathway database**: collection of manually drawn pathway maps representing current knowledge on the molecular interaction and reaction networks
  - **OpenTox / AOP Wiki (JRC/EPA/OECD)** - open knowledge aggregation and collaboration tools that provide a means of describing adverse outcome pathways in an encyclopedic manner.
Implementing the Science: Working together to accelerate technical and scientific advances in pathway-based approaches to chemical safety assessment.

Presentations · Workshops · Papers · Sponsorship

Articulating the Vision: Effective communication facilitates consensus building among stakeholders and is essential to building confidence in chemical and drug safety.

Website · Articles · Video

Lobbying/Funding: Advocate policy changes and cultivate funding opportunities that support pathway-based approaches in the U.S. and internationally.

Bill language · Appropriations
Thank You

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