Promoting The Integrity of Pharmaceutical Research: What Role for Academia?

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Dominant Discourse Academia-Industry Partnerships
- Partnerships are motor for new health product (& economic) development
  - Align agendas of industry—funding agencies-academia
  - Promote collaborative research & health product development
- Changing Nature Academic Institutions
  - Co-funding arrangements
  - Commercialization Hubs (e.g. MaRs)
  - Industry-funded Research & Education, industry-funded Chairs
Drug Regulatory Context

- Focus on Initial Drug Review and Approval: based on limited data of safety and effectiveness; creates public expectation that pharmaceuticals on market are safe and effective
- Information on post-marketing drug safety & effectiveness: largely provided by industry
- Clinical practice (including Off-Label Prescription) largely informed by medical literature and CPGs, with significant role of academic research community

“Problems Current System”


1. Initial Review: too limited
2. PMS: out of FDA’s control; lack of authority to enforce PMS by companies
3. Lack of accountability for suppression or delaying unfavourable trial information
4. Structure of FDA: Potential Conflict may prevent action post marketing
5. Lack of expertise FDA drug safety panels
6. Independence; underreporting adverse drug reactions; slow withdrawals

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How Reliable is Publicly Available Information on Drug Safety and Effectiveness?

- Subtle impact of source of funding clinical trials
- Troublesome Publication Practices Involving Academic Researchers

Subtle Impact of Interactions with Industry: Bias

- Source of Funding Impacts on Outcome of Research: industry funded research more likely to be favourable to products of industry sponsors
- Meta-analyses by
  - Bekelman et al. (JAMA 2003)
  - Lexchin et al. (BMJ 2003)
  - Schott et al. (Dtsch Arztebl Int 2010):

Publication Bias

  - Published literature: 94% studies positive
  - Analysis 74 FDA registered studies:
    - only 51% positive
    - 33 negative studies:
      - 22 not published
    - 11 published conveying a positive outcome
Scientific Publications Strategy: Managing Reputation, Clinical Trial Results and Commercial Relevance, Best Practices LLP ($3,695)

"While picking and choosing favorable findings may have been acceptable a decade ago it is now considered unethical and potentially illegal."

A.G. New York v. GlaxoSmithKline: June 2004

- Elliott Spitzer (AG): "GSK has engaged in repeated and persistent fraud by misrepresentation, concealing and otherwise failing to disclose to physicians information in its control concerning the safety and effectiveness of its antidepressant medication paroxetine in treating children and adolescents."
- "GSK has allowed positive information … to be disclosed publicly, but has withheld and concealed negative information concerning the safety and effectiveness"
Consequences are serious

- D. Graham: 'VIOXX may have caused 140,000 serious injuries or deaths between 1999-2004'
- Hormone Replacement Therapy: NIH study 2002
  - 2 most common HRT: 62 million prescriptions 2000
  - Breast cancer, ovarian cancer, stroke, pulmonary embolism (8+)
  - Increased risk of heart disease (7+ per 10,000)
  - Fewer hip fractures (5<) and colon cancer (6<)
- NOTE: statistics US confirm drop of 8.6% in breast cancer rates since reduction hormone replacement prescription

Remedy: Comprehensive and Integrated Approach

1. Conflict of Interest Rules
2. Clinical Trial Registration and Results Reporting
3. Strengthen Power & Independence Drug Regulator
5. Professional & Academic Sanctions
6. Criminal Law
7. Use of Whistle-blowers: US Qui Tam procedures
8. Structural Reform: PMS system
   - Canadian Drug Safety and Effectiveness Network

2. Trial Registration & Results Disclosure
   (Krleza-Jerić K, Lemmens T et al. PAHO J. Public Health)
US FDA Amendment Act 2007

- Expands ClinicalTrials.gov Registry: obligation to register clinical trials (exc. Phase I)
- Obligatory results reporting of clinical trials of approved drugs and devices (basic results)
  - "Basic" Results: Baseline Characteristics, Key Outcomes, Statistical Analyses (and Adverse Events)
- Serious Penalties for non-compliance
  - Withdrawal of funding
  - Penalty of 10,000 per violation/per day

Limits of Registration

- Enforceability registration requirement?
  - ICMJE: enforcement related to publication
  - WHO: no enforcement other than 'moral authority'
- Penalties Necessary: Register Trials for Serious & Life-Threatening Diseases US: significant non-compliance (prior to FDA Amendment Act!): Only 46% of 127 cancer trial protocols sponsored by pharmaceutical companies were submitted to the registry (Derbis J, et al (2003): reported by Turner PloSMedicine 2004)

Limits of Current Registration

- Access to protocol and full data?
  - S. Vedula, L. Bero et al. (2009 NEJM 361): comparison internal research reports – published reports Neurontin
    - Several reports not published
    - 6/20 identified primary outcomes not in published report; 4/20 reported as secondary outcomes; new primary outcomes; underreporting of negative outcomes
  - "[these] reporting biases increase the likelihood that interventions will appear to be effective when they are not" registration should include registration of the full study protocol and amendments"
- Sponsor still controls clinical trials
New Approaches Necessary

New approaches for the conduct, oversight, and reporting of industry-sponsored trials, are necessary. A clinical trials system in which sponsors fund the trials that are conducted by independent investigators would provide additional protections.


7. Independent Drug Safety Agency

  - Center for Drug Approval
  - Center for Post-Marketing Studies
  - Center for Drug Information

Canada’s Drug Safety and Effectiveness Network (DSEN)

- Network of centres for PM pharmaceutical research
- Mandate:
  - provide strategic direction and common research agenda, in collaboration with national partners
  - organize funding
  - govern independent PM research of centres
- Funding federal government: 35 million per 5 years
Good Governance Framework
(Ferris & Lemmens, Open Med. Forthcoming)

Key Principles DSEN and Participating Research Centres
- Transparency and Openness
- Accountability
- Independence
- Commitment to Scientific Integrity
- Freedom of Action

Challenges DSEN
- Funding: limited when compared with industry
- Stability of Funding?
- Governance and Independence from Industry-Funding Agency Connections
- How Stringent Can COI Rules Be?

Conclusion
- Strengthen Regulatory Tools
- Promote Transparency
  - Financial Relations
  - Research Results
- Use of Existing Legal Tools
  - Separation of academic & regulatory interests AND RESEARCH CONDUCT from industry interests
- Enhance public interest research by reinvigorating partnership public interest science and academic sector
“The piecemeal organizational modifications and short-lived programmatic initiatives of the past and the current, seemingly fragmented and reactive initiatives . . . are not sufficient to meet the need to improve postmarket drug safety activities and protect the public health better.” Institute of Medicine The Future of Drug Safety 2007

“Knowing is not enough; we must apply. Willing is not enough; we must do.” Goethe