

**Harvard Medical School  
Curriculum Vitae**

**Date Prepared:** January 1, 2019  
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**Education**

1996	A.B. ( <i>summa cum laude</i> )	History and Science	Harvard University, Cambridge, MA
2002	M.D.	Medicine	University of Pennsylvania School of Medicine, Philadelphia, PA
2002	J.D. ( <i>magna cum laude</i> )	Law	University of Pennsylvania Law School, Philadelphia, PA
2007	M.P.H.	Clinical Effectiveness	Harvard School of Public Health (HSPH), Boston, MA

**Postdoctoral Training**

6/02-6/03	Intern	Internal Medicine	Brigham and Women's Hospital (BWH), Boston, MA
7/03-6/05	Resident	Internal Medicine	BWH
7/05-6/07	Fellow	General Medicine and Health Care Policy Research	BWH / Harvard Medical School (HMS) / HSPH, Boston, MA

**Faculty Academic Appointments**

7/07-6/10	Instructor	Medicine	HMS
7/08-	Research Associate	Health Policy and Management	HSPH
7/10-6/14	Assistant Professor	Medicine	HMS
7/14-	Associate Professor	Medicine	HMS
7/14-7/15, 7/16-7/20	Irving S. Ribicoff Visiting Associate Professor of Law	Law	Yale Law School
8/14-	Faculty Member	Center for Bioethics	HMS

### Appointments at Hospitals/Affiliated Institutions

7/05-6/07	Associate Physician	General Internal Medicine	BWH
7/05-11/13	Associate Physician	Medicine	Harvard Vanguard Medical Associates
7/05-7/17	Staff Physician	Medicine	Dana-Farber Cancer Institute, Boston, MA
1/06-7/12	Courtesy staff	Medicine	Faulkner Hospital, Jamaica Plain, MA
7/07-	Associate Physician (research)	Pharmacoepidemiology and Pharmacoeconomics	BWH
8/10-7/15	Research Associate	Law, public health, and ethics	Edmond J. Safra Center for Ethics at Harvard University
7/12-	Staff Physician	Medicine	Faulkner Hospital
5/13-	Faculty Supervisor	Petrie-Flom Center for Health Law Policy, Biotechnology and Bioethics	Harvard Law School
9/16-	Distinguished Visitor	Solomon Center for Health Law and Policy	Yale Law School

### Other Professional Positions

2006-2007	Expert witness	Testimony in <i>IMS v. Ayotte</i> on behalf of state of New Hampshire	Concord, NH
2008-2009	Expert witness	Testimony on drug promotion	State of Texas
2008-2009	Expert witness	Testimony in <i>IMS v. Sorrell</i> on behalf of state of Vermont	Montpelier, VT
2008-2011	Consultant	Alosa Foundation	Boston, MA
2010-2011	Consultant	Robert Wood Johnson Foundation Public Health Law Research program	Temple University, Philadelphia, PA
2011, 2013-2016	Expert witness	Testimony on expert witness ethics review proceedings on behalf of American Academy of Orthopedic Surgeons	Chicago, IL
2016	Ethics review	Medical Quality Assurance Commission, Department of Health, State of Washington	Olympia, WA
2018	Outside expert	Northern District of Ohio, Judge Dan Aaron Polster, Multidistrict Litigation 2804: National Prescription Opiate Litigation	Cleveland, OH
2018	Consultant	Review of Pew Charitable Trusts' drug pricing portfolio	Washington, D.C.

### Major Administrative Leadership Positions

#### Local

2003-2005	Course director, Medico-Legal and Health Policy Curriculum for Internal Medicine Residents	BWH
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2010-2011	Admissions chair, Law and Public Health Concentration	HSPH
2012-	Site director, HMS Fellowship in General Medicine and Primary Care	Division of Pharmacoepidemiology and Pharmacoeconomics, BWH
2013-	Director, Program On Regulation, Therapeutics, And Law (PORTAL)	Division of Pharmacoepidemiology and Pharmacoeconomics, BWH
2016-	Leader, Health Policy and Bioethics Consortium monthly lecture series	HMS
<b>National</b>		
2009-2017	Chair, Council of Recent Graduates	University of Pennsylvania School of Medicine
2011	Co-organizer, national conference on conflicts of interest in medicine	American Society of Law, Medicine and Ethics, University of Pittsburgh Law School
2013	Co-organizer, national conference on blinding in biomedical research and the law	Safra Center for Ethics at Harvard University, Harvard Law School Petrie-Flom Center
2014	Co-organizer, national conference on essential evidence for new drugs and medical devices	Harvard Medical School/Brigham and Women's Hospital, American Association for the Advancement of Science (AAAS), National Center for Health Research (NCHR)
<b>International</b>		
2015-2017	Governance Board	Innovative Medicines Initiative DRIVE-AB consortium
<b>Committee Service</b>		
<b>Local</b>		
2003-2004	Resident work hours committee, Department of Medicine	BWH Member
2004-2006	Hospital work committee, Division of Pharmacoepidemiology and Pharmacoeconomics	BWH Member
2004-	Faculty committee, Division of Pharmacoepidemiology and Pharmacoeconomics	BWH Member
2009-2013	Research Ethics Working Group, Harvard Clinical and Translational Science Center	HMS Member
2011-2013	Admissions committee, Law and Public Health Concentration	HSPH Member
2011-2012	Harvard Interfaculty Working Group on Government Management of Pharmaceutical Products	Harvard University Member
2012, 2015, 2019	Honors thesis program expert reader	HMS Member
2013-	Regulatory Science Advisory Board	HMS Deputy Director
2013-2014	Clinical trial data sharing working groups	Multi-Regional Clinical Trial Center, Harvard Global Health Institute

2016	Precision Trials Challenge	Member Harvard Business School Judge
<b>Regional</b>		
2011-2012	Master's thesis overseer, Julia Kay Preis	Harvard-MIT Division of Health Sciences and Technology (HST) Biomedical Enterprise Program
2012-2013	S.J.D. thesis committee, Jonathan J. Darrow	Harvard Law School
<b>National</b>		
2007, 2012	Alumni reunion committee	University of Pennsylvania School of Med Member
2007-2008	Expert Advisory Committee	ClinicalTrials.gov Member
2008-	Medical Alumni Advisory Council	University of Pennsylvania School of Med Member
2008-	Penn Law Alumni Society of Boston	University of Pennsylvania Law School Member
2010	Task Force on Generic Immunosuppressants in Hematopoietic Cell Transplantation	American Society for Blood and Bone Marrow Transplantation Member
2011-2013	Patents for Humanity	United States Patent and Trademark Office Development Consultant and Judge
2013	Tenure review committee, Joanna K. Sax	California Western School of Law
2014	Chatham House working group on antibiotic delinkage	Observer
2015-	American Society of Law, Medicine and Ethics	Board of Directors
2015-	Food and Drug Administration (FDA) Peripheral and Central Nervous System Advisory Committee	Voting Member
2016-	Drugs and Biologics Committee, Food and Drug Law Institute	Member
2016-2017	Committee on Pain Management and Regulatory Strategies to Address Prescription Opioid Abuse, National Academies of Sciences, Engineering and Medicine	Member
<b>Professional Societies</b>		
1999-2006	American College of Legal Medicine 2003-2006: Student Awards Committee	Member
2003-	New York State Bar Association	Member
2004-2007, 2011-2013	Society of General Internal Medicine	Member
2004-2010	International Society for Pharmacoepidemiology	Member

2009-	AcademyHealth 2011-2013: Quality and Value Interest Group Advisory Committee 2012-2013: Annual Research Meeting Planning Committee 2015-2017: Alice B. Hersh Award selection committee	Member
2011-2012, 2015-	American Society of Law, Medicine & Ethics	Member

### Grant Review Activities

2010, 2013	Grant proposal reviewer ( <i>ad hoc</i> )	Robert Wood Johnson Foundation Public Health Law Research Program
2011	Grant proposal reviewer ( <i>ad hoc</i> )	Robert Wood Johnson Foundation Investigator Award in Health Policy Research
2013	Grant proposal reviewer ( <i>ad hoc</i> )	Alzheimer's Association
2015	Grant proposal reviewer ( <i>ad hoc</i> )	Harvard Clinical and Translational Science Center
2017-2018	Grant proposal reviewer ( <i>ad hoc</i> )	Greenwall Foundation Making a Difference

### Editorial Activities

#### Ad hoc peer reviewer

American Heart Journal  
American Journal of Bioethics  
American Journal of Respiratory and Critical Care Medicine  
American Journal of Tropical Medicine & Hygiene  
Annals of Internal Medicine  
BioMed Central (BMC) Medical Ethics  
BMC Medical Research Methodology  
British Medical Journal (BMJ)  
BMJ Quality & Safety  
Canadian Medical Association Journal Open  
Circulation  
Clinical Pharmacology and Therapeutics  
Current Medical Research and Opinion  
Drug Discovery Today  
Drug Testing and Analysis  
Expert Review of Molecular Diagnostics  
Expert Review of Pharmacoeconomics & Outcomes Research  
Family Practice Essentials  
Genome Biology  
Health Affairs  
Health Policy  
Journal of the American Medical Association (JAMA)  
JAMA Cardiology  
JAMA Internal Medicine  
JAMA Oncology

Journal of General Internal Medicine  
 Journal of Health Politics, Policy, and Law  
 Journal of Law and Biosciences  
 Journal of Law, Medicine, and Ethics  
 Kennedy Institute of Ethics Journal  
 Medical Letter  
 Milbank Quarterly  
 Nature  
 New England Journal of Medicine (NEJM)  
 Pharmacoeconomics  
 Pharmacoepidemiology & Drug Safety  
 Pharmacy & Therapeutics  
 Public Library of Science (PLOS) Biology  
 PLoS Medicine  
 PLoS One  
 Science  
 Science Translational Medicine  
 Social Science & Medicine  
 Yale Journal on Regulation

**Other Editorial Roles**

1999-2000	Associate Editor	<u>University of Pennsylvania Law Review</u>
2000-2002	Senior Editor	<u>University of Pennsylvania Law Review</u>
2008	Faculty articles reviewer	<u>Harvard Law Review</u>
2009	Executive Board, review of antibiotic incentive policy	London School of Economics
2012	Guest co-editor, <u>Journal of Law, Medicine and Ethics</u> , Volume 40, Issue 3 (title: "Conflict of Interest in the Practice of Medicine")	American Society of Law, Medicine and Ethics
2012-	Academic Editor, <u>PLOS Medicine</u>	Public Library of Science
2012-	Editorial Board, <u>Expert Opinion on Orphan Drugs</u>	Taylor & Francis Online
2012-	Advisory Board, Perspectives section, <u>New England Journal of Medicine</u>	Massachusetts Medical Society
2013-2015	Editorial Board, working paper series	Edmond J. Safra Center for Ethics at Harvard University
2013-2016	Health Policy Brief external editor, <u>Health Affairs</u>	Project Hope
2014	Co-editor, <u>Journal of General Internal Medicine</u> , Volume 29, Suppl 3 (title: "Research Methods for Evaluating Patient Health Outcomes in Rare Diseases")	Society for General Internal Medicine, Agency for Healthcare Research and Quality
2014-	Editorial Board, <u>Clinical Pharmacology and Therapeutics</u>	American Society for Clinical Pharmacology and Therapeutics
2014, 2017	Faculty reviewer	<u>Yale Journal of Health Policy Law and Ethics</u>
2017	Prescription Drug Pricing Health Policy	Project Hope

	Brief series external editor, <u>Health Affairs</u>	
2017	External editor, “Promoting Value, Affordability, and Innovation in Cancer Drug Treatment”	President’s Cancer Panel, National Institutes of Health
2017-	Editor-in-Chief, <u>Journal of Law, Medicine, and Ethics</u>	American Society of Law, Medicine and Ethics
2018	Co-editor, <u>Journal of Law, Medicine and Ethics</u> , Volume 45, Suppl 2 (title: “Transparency at the US Food and Drug Administration”)	American Society of Law, Medicine and Ethics
2018	Invited contributor, annual editorial board meeting	<u>JAMA</u>

### Honors and Prizes

1992	Detur Book Prize	Harvard College	Academic excellence
1992	National Scholar	Harvard College	Academic excellence
1995	Harvard / Ford Foundation Samuel H. Abramson Memorial Fellowship	Harvard College	Thesis research proposal
1996	Phi Beta Kappa honor society	Harvard College	Academic excellence
1996-2002	Ben Franklin Fellow	University of Pennsylvania School of Medicine	Academic excellence
1998	History of Medicine Prize	University of Pennsylvania School of Medicine	History of science writing competition
1998-2002	James Wilson Scholar	University of Pennsylvania Law School	Academic excellence
2000	William Osler Medal	American Association of the History of Medicine	History of science writing competition
2001	Alpha Omega Alpha honor society	University of Pennsylvania School of Medicine	Academic excellence
2002	Order of the Coif honor society	University of Pennsylvania Law School	Academic excellence
2002	Burton Award	The Burton Foundation	National excellence in legal writing
2002	Schwartz Award	American College of Legal Medicine	Health law writing competition
2002	First Place	Epstein, Becker, and Green Health Law Writing Competition	Health law writing competition
2005	Karen Kaufman Memorial Book Award	BWH	Excellence in delivery of primary care
2008	Young Alumnus of the Year	University of Pennsylvania School of Medicine	Career excellence, dedication to school
2009	Top 10% of peer reviewers	Annals of Internal Medicine	Excellence in contributions to editorial decisions
2010	Alice S. Hersh New Investigator Award	AcademyHealth	Exceptional promise for future contributions to

2010, 2013, 2015	Top peer reviewer	Annals of Internal Medicine	health policy research Excellence in contributions to editorial decisions
2011	Top peer reviewer	Pharmacoepidemiology and Drug Safety	Excellence in contributions to editorial decisions
2013	30th Anniversary Award	Center for Excellence in Education's Research Science Institute	Excellence and achievement in science, technology, engineering, math and business
2013	Second place prize	Eighth Annual Massachusetts Medical Society Research Poster Symposium (health policy/medical education category)	Senior author of research poster
2014	Chair's Research Award	BWH Department of Medicine	Skill in obtaining grant funding
2015-16	Young Mentor Award	HMS	Excellence in developing quality mentoring relationships that lead to professional development and career advancement in basic/clinical medicine
2016	Research Leadership Award	BWH	Awarded to investigators who have demonstrated outstanding research leadership of new or existing programs
2017	Power List 100: Masters of the Bench	The Medicine Maker	National trade publication list of top individuals "involved in bettering the pharma industry"
2017	Leonard M. Rosen Memorial Research Award	Children's Cause Cancer Advocacy	Outstanding contribution to childhood cancer policy and advocacy
2018	Thought Leader	<u>NEJM</u> Catalyst	Demonstrating credentials, expertise, and knowledge related to the health care marketplace

## **Report of Funded and Unfunded Projects**

### **Funding Information**

#### **Past**

1999	Health care delivery systems for terminal cancer patients National Cancer Policy Board, Washington, DC / Research fellowship Co-investigator Review of current state of end-of-life care for cancer patients, including trials, physician education and patient knowledge about care options.
2000-2001	Adapting the 25 <sup>th</sup> Amendment to provide for presidential health oversight



- Philadelphia College of Physicians and Surgeons, Philadelphia, PA / Research project  
Co-principal investigator  
Organization of expert working panel to develop recommendations for health of President of the United States and role of 25th Amendment in ensuring proper oversight. Studied history of presidential health.
- 2003-2005    Developing a health policy curriculum for medical residents  
Brigham and Women’s Hospital Support for Excellence in Educational Development / Educational project  
Principal investigator  
Organization of curriculum of guest lectures to expose internal medicine residents to pressing national health policy issues; empirical analysis of reaction to curriculum.
- 2004-2005    Investigation of health policy issues in the U.S. Senate Health, Education, Labor, and Pensions Committee  
Martin P. Solomon Medical Education Scholarship / Educational project  
Principal investigator  
Full-time externship with office of Sen. Christopher Dodd (D-CT) to contribute to considerations of current health-related legislation and development of national health information technology infrastructure development bill.
- 2007-2008    Research in drug and health law policy  
Agency for Healthcare Research & Quality (AHRQ) Post-Doctoral Fellowship in Health Services Research / Mentored training grant  
Principal investigator  
Using empirical research techniques, investigated US intellectual property policies and studied how management of intellectual property rights influences worldwide access to essential medications.
- 2007-2008    Educational outreach to improve prescribing practices  
Attorney General Prescriber and Consumer Education Grant Program / Educational project  
Co-Investigator (PI: Jerry Avorn, M.D.)  
Development of an innovative series of curricula, interactive web-based programs, and educational outreach activities to equip prescribers and prescribers-in-training with the cognitive and attitudinal tools they need to make optimal drug-use decisions.
- 2007-2010    Design of a national educational curriculum, “Generics are powerful medicines”  
Cy pres award distribution from court settlement / Educational project  
Program director  
Organization of consumer education materials and website describing the safety and efficacy of generic drugs, including developing partnerships with local public health outreach organizations through a national request for proposals.
- 2008-2009    Assessment of strategies for development of novel antimicrobial products  
Resources for the Future / Commissioned study  
Co-principal investigator (Co-PI: Kevin Outterson, J.D., LL.M.)  
Descriptive analysis of current proposals to encourage antibiotic drug development, and discussion of a novel alternative, the Antibiotic Conservation and Effectiveness program,

which would combine incentives for development with reimbursement for rational drug use.

- 2009 Using market exclusivity incentives to promote pharmaceutical innovation  
Robert Wood Johnson Foundation Public Health Law Research / Commissioned study  
Principal investigator  
Study of the effect on medical innovation of statutes that provide additional intellectual property rights or related incentives to pharmaceutical developers in the US.
- 2009-2010 Patterns of use of newly approved orphan drugs for rare diseases  
Harvard Clinical and Translational Science Center / Individual investigator initiated grant  
Principal investigator  
Analysis of effectiveness of Orphan Drug Act as means of incentivizing drug development to generate treatments for rare diseases, and expansion of use of those drugs after approval.
- 2009-2014 Off-label prescribing: Comparative evidence, regulation, and utilization  
Agency for Healthcare Research & Quality K-08 Award/Training grant (5K08HS18465-04)  
Principal investigator  
Investigation of off-label prescribing and time series analysis of how legal, regulatory, and market forces affect these uses.
- 2010 Current trends in orphan drug development  
Institute of Medicine Committee on Rare Disease and Orphan Product Development /  
Commissioned study  
Principal investigator  
Study of the characteristics of the drug development and FDA review process for a selection of orphan drugs.
- 2010-2012 Varying disclosure policy for biomedical journal articles: a randomized controlled trial for remedies for financial disclosure of science  
Edmund J. Safra Center for Ethics at Harvard University / Investigator initiated grant  
Co-principal investigator (\$60,582) [with Christopher Robertson, J.D., Ph.D.]  
Randomized controlled study to test solutions to presentations of conflicts of interest in the medical literature.
- 2010-2014 Researching ways to overcome obstacles to creation of breakthrough new drugs  
Robert Wood Johnson Foundation Investigator Award in Health Policy Research /  
Individual investigator initiated grant (67487)  
Principal investigator  
Investigation of how basic, translational, and product-development research combine to create breakthrough new drugs and role of patents in facilitating or impeding this process.
- 2011 Medical device regulation in the US and EU  
Center for Devices and Radiological Health, Food and Drug Administration /  
Commissioned study (HHSF223201111374P)  
Principal investigator  
Comparative analysis of device approval and post-market surveillance and systematic review of studies of device regulatory outcomes in the US and EU.

- 2012-2013 Post-market surveillance of medical devices in the US and EU  
Pew Charitable Trust / Individual investigator initiated grant  
Principal investigator  
Cross-national comparison of systems of post-market surveillance for medical devices.
- 2012-2014 Research methods for evaluating patient health outcomes in rare diseases: symposium and journal supplement Agency for Healthcare Quality and Research/DEcIDE-2 Request for Task Order HHSA290201000006I - TO4  
Principal investigator  
Organization of expert advisory group, literature review and stakeholder focus group addressing the application of research methods to studying outcomes for patients with rare diseases, and experiences with newly approved orphan drugs
- 2012-2014 Developing and testing a decision support tool for primary medication adherence  
Patient-Centered Outcomes Research Institute (PCORI)/PI-12-001  
Contributing investigator (PI: Jennifer Polinski, Ph.D.)  
Leading conduct and analysis of patient and provider focus groups intended to inform development of tool to promote patient adherence to antihypertensive medications
- 2013-2015 Assessing clinical equivalence for generic drugs approved using innovative methods  
Food and Drug Administration (1U01FD004856-01)  
Principal Investigator  
Study of 6 generic drugs approved using non-traditional methods for determining bioequivalence, including surveys of patients and physicians, a secondary data analysis of their use, and a systematic review of published studies of the drugs.
- 2013-2016 New methods for evaluation of impact of FDA Drug Safety Communications  
Food and Drug Administration (HHSF22301001T)  
Principal Investigator  
Combined methodological approach to understanding the impact of information disseminated by FDA about prescription drug safety using qualitative analyses of traditional and social media, surveys of patients, interview of patients and physicians, and pharmacoepidemiologic analyses of drug prescribing and patient outcome trends.
- 2013-2016 Access to drugs and devices that have limited supporting data: ethical implications for patients and physicians  
Greenwall Foundation Faculty Scholar Program  
Principal Investigator  
Using orphan drugs for rare diseases and early access programs as empirical studies to build normative ethical conclusions relevant to patients, physicians, manufacturers, and payers when regulators approve experimental drugs and devices on limited premarket data
- 2013-2017 Does variation in the physical characteristics of generic drugs affect patients' experiences: A survey of pharmacists and patients  
Food and Drug Administration (HHSF223201310232C)  
Principal Investigator  
National surveys of patients and pharmacists to determine their experiences with generic medications that change shape or color during routine refills, and the association of these

episodes with nonadherence and confusion.

- 2014-2016 Studying the impact on public health of variations among states in laws regulating substitution of generic for brand-name drugs  
Robert Wood Johnson Foundation Public Health Law Research Program  
Co-investigator (Principal Investigator: Ameet Sarpatwari, J.D., Ph.D.)  
Mapping of state drug product selection laws affecting generic substitution and observational and direct national survey studies assessing the implications of these laws on access to generic drugs
- 2016 Use of patents and FDA regulatory exclusivities to set and extend brand-name drug market exclusivity: a review of the evidence  
Commonwealth Fund  
Principal Investigator  
Description of the state of the law relating to pharmaceutical market exclusivities and a review of the evidence relating to the strategies used to delay entry of generic drugs.
- 2016-2017 A Study of Pharmaceutical Pay for Outcomes Contracts in the US and their Implications for Pharmaceutical Spending  
Commonwealth Fund  
Co-Principal Investigator (with Elizabeth Seeley, Ph.D.)  
Qualitative interview-based analysis of payors, policymakers, and pharmaceutical manufacturers involved in pay-for-outcomes contracts of high-priced drugs.
- 2016-2017 Reviewing the Legal, Political and Public Health Parameters of Increasing Transparency at the Food and Drug Administration  
Laura and John Arnold Foundation  
Co-investigator (Principal Investigator: Joshua Sharfstein, M.D.)  
Review of the current status of the transparency of FDA decision-making and the potential for enhancing the public availability of key regulatory information.
- 2016-2017 Impact of Drug Innovation Incentive Strategies on Drug Development and Costs  
Laura and John Arnold Foundation  
Principal Investigator  
To examine the outcomes of programs intended to incentivize drug innovation, to identify the most successful aspects of these programs, and to determine how efficiently these programs facilitate the introduction of important new products by grading the innovativeness, efficacy, and safety of the products whose approval they have facilitated
- Current**
- 2014-2019 Examining the Impact of FDA Regulatory Policies on Therapeutic Approval  
Harvard Program in Therapeutic Science  
Principal Investigator (\$589,674)  
Conduct of research in the field of “regulatory science” evaluating the impact of FDA-imposed Risk Evaluation and Mitigation Strategies and evaluating how the FDA applies its existing rules to novel technologies.
- 2016-2018 Development of Educational Boot Camp in Methods Used in Empirical Bioethics Research

- Greenwall Foundation  
 Consultant (PI: Eric Campbell, Ph.D.)  
 To develop a recurring, year-long educational program for Greenwall fellows to introduce them to qualitative and quantitative data collection and analysis, along with pre- and post-testing, and then expand the educational program more broadly to the bioethics community
- 2017-2018 The US Government's Contribution to Transformative Drug Development  
 Open Society Foundation  
 Co-Principal Investigator (Co-PI: Ameet Sarpatwari, Ph.D., J.D. (\$125,000)  
 To study the amount of support that the US government has provided for the discovery and development of specific highly innovative and clinically important pharmaceutical products.
- 2017-2018 The Impact of Intra-Class Competition on Drug Prices  
 Anthem Public Policy Institute  
 Co-Investigator (PI: Ameet Sarpatwari, Ph.D., J.D.)  
 To assess the impact of new drug market entry on the prices of older drugs and investigate the conditions needed for prices to fall.
- 2017-2019 An International Comparison of Regulatory Risk Communication on Medicines  
 National Health and Medical Research Council (NHMRC)  
 Co-Investigator (Principal Investigator: Barbara Mintzes, Ph.D.)  
 To understand of how regulatory warnings are related to medication safety impact health care delivery, and identify a set of 'best practices' contributing to effectiveness, by comparing medication safety advisories in Australia, Canada, the US, and Europe
- 2017-2020 Creation of the PORTAL Biomarker Research Consortium  
 Laura and John Arnold Foundation  
 Principal Investigator (\$1,840,085)  
 To systematically review and meta-analyze the validity of biomarkers used in drug development and treatment in cardiovascular medicine, cancer, Alzheimer's disease, and tuberculosis, as well as to develop additional studies and reviews of biomarker and surrogate measure policy.
- 2017-2020 Prescription Drug Innovation, Availability, and Affordability: The Impact of Drug Innovation Incentive Strategies on Drug Development and Costs  
 Laura and John Arnold Foundation  
 Principal Investigator (\$2,971,681)  
 To document the impact of policy levers on innovation, access, and affordability of prescription drugs, identify how they work well, how they work sub-optimally, and what specific policy options could be implemented to improve them, characterize and critically assess key trends at each stage of the drug product life-cycle that impact expense and innovation, and develop and assess specific possible alternatives to existing policies.
- 2018-2023 Incentivizing the Development of Effective and Safe Antibiotics  
 Collaborative Research Programme in Biomedical Innovation Law at the University of Copenhagen (supported by grant NNF17SA027784 from the Novo Nordisk Foundation)  
 Subcontract Principal Investigator (\$348,456)

To study effects of intellectual property laws and regulatory policies on pharmaceutical development, drug approval processes, and the costs, availability, and use of prescription drugs, with a particular focus on antibiotic drug development.

**Report of Local Teaching and Training**

**Teaching of Students in Courses at HMS/HSDM/DMS**

2002-2005	Core Medicine Clerkship I Third- and fourth-year medical students	HMS 9 hrs per day for 12 wks per year
2002-2005	Core Medicine Clerkship II Third- and fourth-year medical students	HMS 9 hrs per day for 12 wks per year
2005-2009	Core Medicine Clerkship I Third- and fourth-year medical students	HMS 13 hrs per wk for 4 wks per year
2005-2009	Core Medicine Clerkship II Medical students	HMS 13 hrs per wk for 4 wks per year
2009	Health Care Policy Second-year medical students	HMS 6 hrs per lecture for 1 guest lecture
2009-2014	Health Care Policy First-year medical students	HMS 3 hrs per lecture for annual guest lecture
2015	HMS Health Policy Student Interest Group 50 first-year medical students	HMS 3 hrs per lecture for 1 guest lecture
2016	HMS BCMP 311qc: Unmet Medical Needs and Translational Solutions 25 medical and PhD students	6 hrs per lecture for 1 guest lecture
2017	Essentials of Professions: Health care policy First-year medical students	HMS 3 hrs per lecture for 1 guest lecture
2018	Essentials of the Professions II: Everything you need to know about prescription drug policy in 60 minutes 25 medical and PhD students	3 hrs per lecture for 1 guest lecture

**Other Harvard University Courses**

2005	Public Health Law Masters students	HSPH 8 hrs per wk for 1 semester
2006	Law and Public Health Masters students	HSPH 5.5 hrs per lecture for 2 lectures
2007-2009	Public Health Law Law students	Harvard Law School 5.5 hrs for annual guest lecture
2008-2014	Advanced Pharmacoepidemiology Masters students	HSPH 4 hrs for annual guest lecture
2012-2013	GHHP 91r Seminar Undergraduate student independent study	Harvard Faculty of Arts and Sciences 25 hrs per semester for 2 semesters
2013	Law and Public Health (HPM 213) Masters students	HSPH 6 hrs for 1 guest lecture
2014	EPI 502 Antibiotic Epidemiology Masters students	HSPH 4 hrs for 1 guest lecture
2016-2018	HPM 213 Public Health Law Masters students	HSPH 4 hrs for 1 guest lecture

2016	Navigating the American Pharmaceutical Sector Executive education students	Executive and Continuing Professional Education, Harvard T.H. Chan School of Public Health 4 hrs for 1 guest lecture
2016, 2017, 2018	Bioethics 706.0 Health Law, Policy, and Bioethics (Co-taught with H.F. Lynch, J.D., M.B.E. [2016-17] and Brendan Abel, J.D. [2018]) Masters students	HMS Center for Bioethics 4 credit spring semester-long seminar
2016-2017, 2017-2018	Bioethics 742: Policy & Ethics Consortium Masters students	HMS Center for Bioethics 2 credit year-long tutorial

### **Courses Taught While Appointed as Visiting Faculty at Yale**

2015	Law 21767 FDA Law Law students	Yale Law School 2 credit semester-long seminar
2016	Law 20616 FDA Law Law students	Yale Law School 2 credit semester-long seminar
2017-2018	Law 20616/HPM 595 FDA Law and Policy Law and School of Public Health students	Yale Law School, cross-registered with Yale School of Public Health 2 credit semester-long seminar

### **Formal Teaching of Residents, Clinical Fellows and Research Fellows (post-docs)**

2004	Primary care in the White House 30-50 residents	BWH and Faulkner Hospital Guest lecture, 5 hrs
2005	The health care of our political leaders 30-50 residents	BWH and Faulkner Hospital Guest lecture, 3 hrs
2004-2009	Medico-legal issues for medicine residents 30-50 residents	BWH and Faulkner Hospital Annual guest lecture, 5 hrs
2005-2008	Ambulatory care rotation Residents	Massachusetts General Hospital, Boston 4 hrs per wk for 3 wks per year
2011-	Partners Center of Expertise in Health Policy and Management: Health Policy Certificate Course 30-50 residents	HMS-affiliated teaching hospitals Annual guest lecture, 3 hrs
2015	What do we know about diabetes drugs? 60 residents	BWH Guest lecture, 2 hrs
2017-2018	Understanding Biomarker Science: From Molecules to Images 120 graduate students	Harvard Catalyst Guest lecture, 2 hrs

### **Clinical Supervisory and Training Responsibilities**

2005-2009	General Medical Service Attending / Brigham and Women's Hospital	5 hrs per day for 4 wks per year
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### **Laboratory and Other Research Supervisory and Training Responsibilities**

2008-	Supervision of college students, medical students, medical and neurology interns/residents, post-doctoral fellows,	Varied levels of mentorship, from daily to weekly, lasting from a few months to several years.
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visiting scholars, and junior faculty members on intersections between law and medicine, pharmaceutical and medical device law and policy, legal research methodology, qualitative data collection, manuscript preparation, career development. Brigham and Women's Hospital

2013- Initiated Program On Regulation, Therapeutics, And Law (PORTAL) to bring together post-doctoral fellows trained in law and medicine, along with students with law, public health, and/or public policy interest, to study questions related to regulatory and drug development and delivery. Brigham and Women's Hospital

Close mentorship on daily basis, weekly lab meetings, lasting from a few months to several years.

### **Mentored Trainees and Faculty**

2005-2009 Rahul Rajkumar, M.D., J.D. / Senior vice president/Chief Medical Officer at CareFirst BlueCross BlueShield, Baltimore, MD  
Career stage: medical resident (BWH). Oversight of research program in intellectual property issues affecting availability of drugs in resource-poor settings, leading to 3 publications.

2006-2012 Dave A. Chokshi, M.D., M.Sc. / Assistant Professor of Medicine, New York University Langone Medical Center, New York, NY  
Career stage: medical student (University of Pennsylvania) and resident (BWH). Oversight of research program in access to and study of drugs and vaccines, leading to 2 publications. Dave served as 2012-2013 White House Fellow.

2008-2013 Alex Misono, M.D., M.B.A. / Resident, Massachusetts General Hospital, Boston, MA  
Career stage: medical student (HMS). Research on generic and brand-name drug policy, including evidence of relative efficacy of generic and brand-name drugs and study of effect of generic/brand color changes on medication adherence, leading to 3 publications.

2009-2010 Devan D. Bartels, M.D., M.P.H. / Resident, Massachusetts General Hospital, Boston, MA  
Career stage: medical student (HMS). Oversight of research project in effect of legal, social, and medical market events on off-label use of Neurontin, leading to 1 publication.

2010-2011 Kirsten E. Austad, M.D. / Attending physician, BWH, Boston, MA  
Career stage: medical student (HMS). Oversight of Safra Center-funded fellowship on medical school education and changes in attitudes about the pharmaceutical industry, leading to 8 publications.

2010-2012 Julia Kay Preis, S.M., M.B.A. / Consultant, The Frankel Group, Boston, MA  
Career stage: masters student (HMS). Oversight of honors master thesis on innovation in influenza vaccine development.

2010- Daniel B. Kramer, M.D., M.P.H. / Assistant Professor of Medicine, Division of Cardiovascular Medicine, Beth Israel-Deaconess Medical Center, Boston, MA  
Career stage: Junior faculty. Supervision of series of projects relating to medical device regulation and ethics, leading to 14 publications.

2011-2012 Adam Licurse, M.D. / Assistant Medical Director, Brigham and Women's Physician's Organization, BWH, Boston, MA



- Career stage: medical resident (BWH). Oversight of research on conflicts of interest and physician disclosure of industry relationships, leading to 1 publication.
- 2011-2014, 2016- Jonathan J. Darrow, J.D., M.B.A., S.J.D. / Instructor in Medicine, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: S.J.D. student (Harvard Law School) and post-doctoral fellow and junior faculty (BWH). Supervision of thesis and post-doctoral work on history of drug efficacy study and regulation, leading to S.J.D. thesis and 11 publications.
- 2011-2014 Shuai Xu, M.D., M.Sc. / Instructor in Dermatology, Northwestern Feinberg School of Medicine, Chicago, IL  
Career stage: medical student (HMS). Oversight of HMS/HSDM Scholars in Medicine-funded research and honors thesis on medical device innovation, leading to 5 publications, a *cum laude* medical school thesis, and 2012 Soma Weiss Research day finalist.
- 2011-2016 Bo Wang, M.D., Pharm.D. / Internal medicine resident, Stanford, Palo Alto, CA  
Career stage: medical student (HMS). Oversight of course of research related to drug policy issues, leading to 17 publications. Bo won the 2015 Robert Wood Johnson Foundation Public Health Law Research Program Young Investigator Award.
- 2012 Kyle D. Checchi, M.Sc., M.D. / Resident, San Diego, CA  
Career stage: medical student (HMS). Oversight of HMS/HSDM Scholars in Medicine-funded research internship on use of pill bottle-related medical device innovation to improve medication adherence, leading to 1 publication.
- 2012-2013 Colin Schwartz / Senior Associate for Policy and Advocacy, American Association of People with Disabilities, Washington, D.C.  
Career stage: masters student (Harvard Kennedy School). Oversight of research on development of transformative HIV drugs (zidovudine and protease inhibitors)
- 2012-2015 Yongtian T. Tan / M.D./M.B.A. student, HMS and Harvard Business School, Boston, MA  
Career stage: medical student (HMS). Oversight of research on medical device innovation in resource-poor settings and comparison of medical device regulation in China and US, leading to 5 publications.
- 2012-2015 Evan S. Caplan, M.D., M.B.A. / Consultant, McKinsey & Co.  
Career stage: medical student (HMS). Investigation of sources of innovation leading to development of vascular endothelial growth factor inhibitors for use in ophthalmologic disease, leading to 1 publication.
- 2012- Thomas J. Hwang / Research Assistant, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: undergraduate (Harvard) and research associate (BWH). Oversight of coursework and thesis research on Food and Drug Administration rulemaking, regulation, and biopharmaceutical innovation, leading to 14 publications.
- 2013 Nathan Shiu, J.D., M.P.H. / Lawyer at FDA  
Career stage: law student (University of California-Los Angeles). Oversight of summer research fellowship on adjudication of truth and scientific certainty in the federal courts, leading to 2 publications.
- 2013-2015 James S. Yeh, M.D. / Instructor in Medicine, Massachusetts General Hospital, Boston, MA  
Career stage: post-doctoral fellow (BWH). Oversight of post-residency general medicine fellowship in health services research, leading to 6 publications.
- 2013-2016 Carolyn Treasure, M.D. / Resident, BWH, Boston, MA  
Career stage: medical student (HMS) and resident (BWH). Oversight of HMS/HSDM Scholars in Medicine-funded research internship on university patenting and government march-in rights, leading to 4 publications.

- 2013- Ameet Sarpatwari, Ph.D., J.D. / Instructor in Medicine, Division of Pharmacoeconomics and Pharmacoepidemiology, Boston, MA  
Career stage: post-doctoral fellow and junior faculty (BWH). Oversight of post-doctoral research program on law and public health topics, leading to 7 publications.
- 2013- Ben Rome, M.D. / Resident, BWH, Boston, MA  
Career stage: medical student (HMS) and resident (BWH). Oversight of HMS/HSDM Scholars in Medicine-funded research internship on US high-risk medical device regulation, leading to 3 publications.
- 2014 Prashant Rajan / Orthopedic surgery resident, Cleveland Clinic, Cleveland, OH  
Career stage: medical student (HMS). Oversight of project on current and future prospects for FDA postmarket regulation of medical devices, and the FDA regulation of medical device approval, leading to 2 publications.
- 2014-2016 Laura E. Bothwell, Ph.D. / Assistant Prof, Worcester State University, Worcester, MA  
Career stage: post-doctoral fellow (BWH). Oversight of project on adaptive design clinical trials, leading to 2 peer-reviewed publications.
- 2014- Jing Luo, M.D. / Instructor in Medicine, Division of Pharmacoeconomics and Pharmacoepidemiology, Boston, MA  
Career stage: post-doctoral fellow and junior faculty (BWH). Oversight of post-residency general medicine fellowship in health services research, leading to 10 publications.
- 2015 Audrey D. Zhang / Student, New York University School of Medicine, New York, NY  
Career stage: undergraduate (Harvard). Oversight of projects on use of biomarkers in FDA decision-making about investigational drugs, and tracing their conceptual evolution as shaped by academia, industry, and regulatory agencies.
- 2015 Vincent C. Capati, Pharm.D., M.S. / Associate, Wiley Rein LLP, Washington, D.C.  
Career stage: law student (University of New Hampshire) Oversight of project examining interaction of antitrust law and pharmaceutical manufacturer marketing behavior, leading to 1 publication.
- 2015-2016 Nicole L. Levidow, J.D., M.P.H. / Compliance administrator, Massachusetts Institute of Technology Office of Sponsored Programs, Cambridge, MA  
Career stage: post-doctoral fellow (BWH). Oversight of project examining characteristics of clinical trials used to evaluate drugs moving through the Accelerated Approval pathway at FDA, leading to 2 publications.
- 2015-2017 Dalia M. Deak, M.P.H. / Law student, Harvard Law School, Cambridge, MA  
Career stage: masters student (HSPH) and law student (Harvard Law School). Oversight of projects examining, drug rediscovery and repurposing, the state of antibiotic development, the ethics of FDA approval pathways, and the history of biotechnology innovation, leading to 2 publications.
- 2015-2017 Mallika L. Mundkur, M.D., M.P.H. / Medical Officer, FDA, White Oak, MD  
Career stage: post-doctoral fellow (BWH). Oversight of projects on trends in high-risk medication use, including antibiotics and opioids, leading to 1 peer-reviewed publication.
- 2015- Spencer Phillips Hey, Ph.D. / Research Scientist, Division of Pharmacoeconomics and Pharmacoepidemiology, Boston, MA  
Career stage: post-doctoral fellow (BWH) and staff (HMS). Oversight of projects at intersection of ethics and regulation involving personalized medicine and biomarker, leading to 8 peer-reviewed publications.
- 2016-2017 Sana Mostaghim, Dr.P.H. / Vaccines Business Unit, Takeda, Cambridge MA  
Career stage: doctoral student (HSPH). Oversight of projects on regulatory approval pathways and prescription drug safety, leading to 2 publications.

- 2016-2018 Chana A. Sacks, M.D., M.P.H. / Instructor in Medicine, Massachusetts General Hospital  
Career stage: post-doctoral fellow (BWH). Oversight of projects on drug prices and off-label use of drugs for rare diseases, leading to 3 peer-reviewed publications.
- 2016- Kerstin N. Vokinger, M.D., J.D., Ph.D., LL.M. / Instructor in Medicine, University of Zurich, Switzerland  
Career stage: post-doctoral fellow (BWH). Oversight of projects on differences between U.S. and European drug regulation, market exclusivity and second-generation brand-name drugs, leading to 2 peer-reviewed publications.
- 2016- Michael S. Sinha, M.D., J.D., M.P.H. / Research Fellow, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: post-doctoral fellow (BWH). Oversight of projects on market exclusivity extensions applied to drugs studied in pediatric trials, use of social media in communicating about drug safety, leading to 6 peer-reviewed publications.
- 2016- Emily Jung / Research Assistant, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: undergraduate (Harvard). Oversight of projects on racial, ethnic, and gender diversity in pivotal clinical trials used for FDA drug approval, leading to 1 peer-reviewed publication.
- 2016- Nina Jain, M.D., M.B.A., M.Sc. / Resident, BWH, Boston, MA  
Career stage: medical student (HMS) and resident (BWH). Oversight of projects on incentives for drug innovation, leading to 2 peer-reviewed publications.
- 2016-2018 Michael Fralick, M.D., M.P.H. / Clinician Scientist Training Program, Department of Medicine, University of Toronto, Canada  
Career stage: post-doctoral fellow (BWH). Oversight of projects on drug safety monitoring and evaluation of drug clinical trials, leading to 12 peer-reviewed publications.
- 2017-2018 Reed F. Beall, M.A., Ph.D. / Assistant Professor, University of Calgary, Alberta, Canada  
Career stage: post-doctoral fellow (BWH). Oversight of projects on impact of patents and market exclusivity on availability of essential medical products, leading to 5 peer-reviewed publications.
- 2017- Chintan Dave, Pharm.D., Ph.D. / Research Fellow, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: post-doctoral fellow (BWH). Oversight of projects on prescription drug pricing, generic drug availability, drug shortages, and pharmacoepidemiology, leading to 2 publications.
- 2017- Elvira D'Andrea, M.D., M.P.H. / Research Fellow, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: post-doctoral fellow (BWH). Oversight of projects on biomarkers and their use in drug development, leading to 1 peer-reviewed publication.
- 2017- Huseyin Naci, Ph.D., M.H.S. / Assistant Professor, London School of Economics, UK.  
Career stage: Harkness fellow (BWH). Oversight of projects on FDA expedited approval pathways and insurance coverage of high-priced drugs, leading to 1 peer-reviewed publication.
- 2018- Bishal Gyawali, M.D., Ph.D. / Research Fellow, Division of Pharmacoepidemiology and Pharmacoeconomics, Boston, MA  
Career stage: post-doctoral fellow (BWH). Oversight of projects on biomarkers and their use in oncology drug development, leading to 3 peer-reviewed publications.
- 2018- William B. Feldman, M.D. / Fellow, Division of Pulmonary and Critical Care, BWH, Boston, MA

Career stage: fellow (BWH). Oversight of projects on ‘exceptions from informed consent’ clinical trials and evidence-based use and cost of pulmonary disease medications, leading to 1 peer-reviewed publication.

### **Local Invited Presentations**

Those presentations below sponsored by outside entities are so noted and the sponsor(s) is (are) identified.

- 2004 Two medico-legal cases / Medicine Grand Rounds (with James T. Hilliard)  
Department of Medicine, BWH
- 2004 Patents, academic research, and drug discovery / Research Rounds  
Division of Pharmacoepidemiology and Pharmacoeconomics, Department of Medicine, BWH
- 2006 Characteristics of physicians who frequently act as expert witnesses in neurological birth injury litigation / Research Rounds  
Department of Medicine, BWH
- 2007 Patent extensions and public health: an empirical analysis / Research Rounds  
Department of Health Care Policy and Management, HSPH
- 2007 Patents and public health: balancing innovation and access / Research Rounds  
Center for Outcomes and Policy Research, Dana-Farber Cancer Institute
- 2008 Balancing drug development and public health / Invited Lecture  
Department of Medicine, Massachusetts General Hospital
- 2008 The insiders: a decade of health care whistleblowers and Department of Justice investigations of health care fraud / Research Rounds  
Department of Medicine, BWH
- 2008 Industry sponsorship in medicine and medical research / Grand Rounds  
Department of Geriatric Medicine, Hebrew Rehabilitation Center, Jamaica Plain, MA
- 2008 Patents and public health: balancing access and incentives for innovation / Plenary Talk  
Harvard Interfaculty Initiative for Medicines and Society conference, Harvard University
- 2009 Patents and cancer drug development / Research Rounds  
Center for Outcomes and Policy Research, Dana-Farber Cancer Institute
- 2009 Patents, innovation, and public health / Invited Lecture  
Department of Medicine, Massachusetts General Hospital
- 2009 Intellectual property issues limiting access to essential medicines / Panel  
Journal of Law and Technology annual symposium, Harvard Law School
- 2009 Health metrics evaluation workshop / Panel  
Petrie-Flom Center for Health Policy, Biotechnology, and Bioethics, Harvard Law School
- 2010 Intellectual property and health care delivery / Invited Speaker  
Harvard Law School Conference on Intellectual Property Law, Cambridge, MA
- 2010 Market exclusivity incentives for drug development: perils and promise / Invited Lecture  
Department of Medicine, Massachusetts General Hospital
- 2011 Legal ecology of resistance / Invited Speaker  
Antimicrobial resistance: biology, population dynamics and policy options, HSPH Center for Communicable Disease Dynamics annual symposium, Boston, MA
- 2011 Patents and public health: what are the limits / Invited Lecture  
Department of Biostatistics, HSPH
- 2011 The Orphan Drug Act and transformative drug development in oncology / Research rounds  
Center for Outcomes and Policy Research, Dana-Farber Cancer Institute
- 2011 Medical malpractice as a health policy issue / Invited Lecture  
Department of Medicine, Massachusetts General Hospital

- 2011 Legislative incentives for pharmaceutical innovation / Invited Lecture  
Department of Medicine, Massachusetts General Hospital
- 2011 Making drug approval and surveillance less scary / Invited Lecture  
Harvard Interfaculty Initiative on Drug Development, Harvard University
- 2012 Legislative incentives for pharmaceutical innovation / Invited Lecture  
Health Policy Certificate Program, Partners Graduate Medical Education
- 2012 Influence of conflict of interest disclosure on physicians' interpretation of clinical  
research: a randomized controlled trial / Research Rounds  
Center for Outcomes and Policy Research, Dana-Farber Cancer Institute
- 2013 Association for Molecular Pathology v. Myriad Genetics, the Supreme Court, and the  
ongoing fight over breast cancer patents / Research Rounds  
Center for Outcomes and Policy Research, Dana-Farber Cancer Institute
- 2013 Health law year in p/review: gene patents / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics, Cambridge, MA
- 2013 Legal and ethical issues in therapeutic development and regulation / Invited Speaker  
Harvard Program in Therapeutic Science, Boston, MA
- 2013 Bayh-Dole march-in rights and the public's access to medical products based on federally-  
funded research / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics Health Law Policy and Bioethics Workshop, Cambridge, MA
- 2014 Second Annual Health law year in p/review: breakthrough drugs / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics, Cambridge, MA
- 2014 Patents without patents / Moderator  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics, Cambridge, MA
- 2014 Overview of current issues facing biosimilar regulation / Featured Speaker  
Mini-Course to Visiting Members of Chinese FDA, Boston, MA (sponsored by Charles  
Institute of Management)
- 2014 Accelerated FDA approval of new drugs and devices: what are the medical, legal, and  
ethical risks? / Grand Rounds  
Beth Israel Deaconess Medical Center Department of General Medicine and Primary Care,  
Boston MA
- 2014 Are stem cells patentable? / Invited lecture  
Harvard Department of Stem Cell and Regenerative Biology-Laboratory of Systems  
Pharmacology Research Day, Cambridge, MA
- 2014 Studies in regulatory science / Invited lecture  
Therapeutic Science Advisory Council Meeting, HMS, Boston MA
- 2014 Hepatitis C drugs: what price progress? / Medicine Grand Rounds (with Paul E. Sax)  
Department of Medicine, BWH
- 2015 Updating the HMS conflicts of interest policy / Invited speaker  
HMS Standing Committee on Conflicts of Interest and Commitment, Boston MA
- 2015 Brain hacking to boost your A-game: the ethics of cognitive enhancement in gaming and  
competition / Invited Speaker  
HMS Center for Bioethics neuroethics seminar series, Boston MA
- 2015 FDA in the 21st Century / Invited panelist  
Harvard Law School, Cambridge MA

- 2015 Regulatory science and the 21st Century Cures Act / Invited lecture  
Therapeutic Science Advisory Council Meeting, HMS, Boston MA
- 2015 Specimen science: background and foundations / Invited panel moderator  
Harvard Law School, Cambridge MA
- 2015 Ethical issues in expanded access to investigational drugs / Invited discussant  
HMS Center for Bioethics, Boston MA
- 2015 Institutional corruption and public health: the case of FDA expedited review and  
development programs/Invited speaker  
Edmond J. Safra Center for Ethics at Harvard University, Cambridge, MA
- 2016 Health law year in p/review: 21st Century Cures Act / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics, Cambridge, MA
- 2016 High-cost drugs: origins, impacts, prospects for reform / Cardiovascular Grand Rounds  
Division of Cardiovascular Medicine, BWH
- 2016 Should cost matter in the care of patients with advanced cancer? / Featured discussant  
Harvard Center for Bioethics Clinical Ethics Consortium, HMS
- 2016 Regulatory environment around cancer drug development / Featured speaker  
HMS External Education: Cancer Care in 2025, Boston MA
- 2016 Current Legal and Ethical Issues Affecting Prescription Drugs / Featured speaker  
HMS Media Fellowship on Bioethics, Boston MA
- 2016 Fostering innovation in early stage bio-pharma / Featured speaker  
Harvard Business School Health Care Initiative and Harvard Kennedy School Healthcare  
Policy Program, Cambridge MA
- 2016 FDA regulation, innovation, and the 21st Century Cures Act / Featured speaker  
Pharmaceutical Policy Research Seminar, Department of Population Medicine, HMS and  
the Harvard Pilgrim Health Care Institute, Boston MA
- 2016 Patient involvement with the FDA / Discussant and Moderator  
Health Policy and Bioethics Consortium, HMS, Boston MA
- 2016 Regulatory science and precision medicine: the tale of eteplirsen / Invited lecture  
Regulatory Science Advisory Council Meeting, HMS, Boston MA
- 2016 What is the proper role of patient advocacy in FDA approval decisions? / Grand Rounds  
Henry Hardy Lecture in Bioethics and Public Policy, Beth Israel Deaconess Medical  
Center, Boston MA
- 2017 Prescription drug policy: The past, present and future / Invited Lecture  
Harvard Graduate School of Arts and Sciences Science Policy Group, Cambridge, MA
- 2017 Looking forward: the next generation of biosimilars / Moderator  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics, Cambridge, MA
- 2017 The future of the FDA / Medicine Grand Rounds  
Department of Medicine, Brigham and Women's Faulkner Hospital, Boston, MA
- 2017 Global health challenge: 2017 and beyond / Panelist  
Harvard Kennedy School Global Development Conference, Cambridge, MA
- 2017 Prescription drug prices: controversies and potential solutions / Grand Rounds  
Department of Medicine, BWH, Boston MA
- 2017 The Cost of Medications: Current Realities and the Future of Pharmaceutical Pricing  
Regulations in the United States / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and  
Bioethics, Cambridge, MA

- 2018 Prescription Drug Prices and “Value” / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and Bioethics, Cambridge, MA
- 2018 Patients’ Role in FDA Drug Approval Decisions / Ethics Grand Rounds  
Dana-Farber Cancer Institute, Boston, MA

**Report of Regional, National and International Invited Teaching and Presentations**

**Invited Presentations and Courses**

Those presentations below sponsored by outside entities are so noted and the sponsor(s) is (are) identified.

**Regional**

- 2008 Pressing issues in health care and pharmaceutical policy / Invited Lecture  
Massachusetts Attorney General Health Care Division offices, Boston, MA
- 2009 Access to human papillomavirus vaccines: human rights and global health / Plenary talk  
American Journal of Law and Medicine annual symposium, Boston University School of Law, Boston, MA
- 2009 Clinical and policy rationales for legislation banning the commercial sale of physician-identified prescription data / Invited Lecture  
Massachusetts state legislature Joint Committee on Health Care Financing, Boston, MA
- 2011 Public health goals and commercial speech in off-label drug promotion / Plenary talk  
American Journal of Law and Medicine annual symposium, Boston University School of Law, Boston, MA
- 2012 The past, present and future of pay-for-delay settlements between brand-name and generic manufacturers / Invited Speaker  
Boston Intellectual Property Law Association, Antitrust Division, Boston, MA
- 2012 Incentivizing research in rare diseases / Invited Plenary Speaker  
Pharmaceutical Research and Manufacturers of America Annual Meeting, Boston, MA
- 2012 Health policy visiting scholar / Invited Speaker  
Yale College, Yale School of Management, and Robert Wood Johnson Clinical Scholars Program, New Haven, CT
- 2013 Implementing conflicts of interest policies at academic medical centers / Invited Speaker  
New England Medical School and Academic Medical Center Roundtable, Community Catalyst, Boston, MA
- 2013 Public health implications of the Supreme Court’s decision in *Federal Trade Commission v. Actavis* / Invited Speaker  
Boston Intellectual Property Law Association, Antitrust Division, Boston, MA
- 2013 Opening up translational research / Featured Speaker  
Universities Allied for Essential Medicines joint MIT-Harvard conference, Cambridge, MA
- 2013 Overview of current issues facing biosimilar regulation in the US / Featured Speaker  
Days of Molecular Medicine Global Foundation, Boston, MA [sponsored by Sectoral Asset Management]
- 2013 Antibiotics: Issues in the Development and Evidence-Based Use / Guest Course Lecture  
Massachusetts Institute of Technology Introductory Biology 7.015, Cambridge, MA
- 2013 Prescription Drugs: Intersections with Patents and Public Health / Guest Course Lecture  
Boston University School of Public Health Epidemiology 748 Masters Seminar, Boston, MA
- 2014 Patents and public health / Guest Course Lecture  
Northeastern University School of Law 7606: Health Law, Boston, MA

- 2015 Is there a myth of data exclusivity?/Invited speaker  
2nd Annual BioIP conference, Boston University School of Law, Boston, MA
- 2016 The Future of Drug Promotion and Public Health / Invited Speaker  
Northeastern University School of Law Conference on the Future of Public Health Law,  
Boston, MA
- 2016 Government Interventions to Address High Drug Prices / Invited Speaker  
American Society of Law, Medicine and Ethics' Health Law Professors' Conference,  
Boston, MA
- 2016 Developing Legal and Policy Responses to Drug-Resistant Bacteria / Panelist  
Yale Global Health Justice Partnership Forum, New Haven, CT
- 2016 The Legal Causes of – and Solutions to – High Drug Prices / Panelist  
Yale Global Health Justice Partnership Forum, New Haven, CT
- 2017 Myths and realities of FDA drug regulation / Featured speaker  
Pharmaceuticals Certificate Program, Global Health Department at Boston University  
School of Public Health
- 2017 Physicians and Their Role in Reducing Drug Costs / Featured speaker  
Massachusetts Medical Society Ethics Forum, Boston, MA
- 2017 Managing High Prescription Drug Prices / Featured speaker  
Institute for Healthcare Improvement Leadership Conference, Boston, MA

### **National**

- 2000 End-of-life care report: information for patients and families / Invited Lecture  
National Cancer Policy Board, Woods Hole, MA
- 2001 Gleevec (STI-571), a new treatment for chronic myelogenous leukemia: the science of  
drug discovery and FDA approval / Grand Rounds  
M.D./Ph.D. program, University of Pennsylvania School of Medicine
- 2004 Deoxyribonucleic Acid (DNA) in civil litigation / Invited Lecture  
American College of Legal Medicine annual meeting, Las Vegas, NV
- 2005 Financial impact of current drug patent policy on Medicaid drug spending / Invited Lecture  
Society of General Internal Medicine annual meeting, New Orleans, LA
- 2006 Update on DNA in civil litigation / Invited Lecture  
American College of Legal Medicine annual meeting, Las Vegas, NV
- 2006 The price of innovation: the effect of patents on medical practice / Plenary Lecture  
American Association of Pharmaceutical Scientists annual meeting, San Antonio, TX
- 2007 Presenting truthful information to physicians / Invited Lecture  
National State Attorney General Program at Columbia Law School, New York, NY
- 2008 Local prescribing practices and access to drugs in resource-poor settings / Plenary Talk  
American Journal of Law and Medicine symposium, Boston University School of Law
- 2008 Free speech and pharmaceutical promotion to physicians / Invited Lecture  
American University Washington College of Law Conference, Washington, DC
- 2008 Pharmaceutical policy issues and points of interest for Attorneys General / Invited Lecture  
National Teleconference of Attorneys General
- 2008 Should FDA drug and device regulation bar liability claims? / Congressional Testimony  
House of Representatives Committee on Oversight and Government Reform (Rep.  
Waxman, Chairman), Washington, DC
- 2008 Global Health Frontiers Workshop / Panel  
Center for Global Development, Warrenton, VA
- 2008 Pharmaceutical development: innovation vs. public health / Invited Lecture



- Leonard Davis Institute, University of Pennsylvania
- 2008 The priority review vouchers: questions and concerns / Invited Lecture  
Knowledge Ecology International meeting on incentivizing drug development for neglected diseases, Washington, D.C.
- 2008 The risks and benefits of follow-on biologics legislation for Medicare / Panel  
Medicare Payment Advisory Commission, Washington, DC
- 2010 Constitutional health law: pharmaceutical regulation and commercial speech / Panel  
Association of American Law Schools Annual Meeting, New Orleans, LA
- 2010 Using market exclusivity to incentivize drug development / Invited Speaker  
University of Pennsylvania Law School Center for Technology, Innovation, and Competition, Philadelphia, PA
- 2010 Implementation of and innovation within the Orphan Drug Act / Invited Speaker  
Committee Accelerating Rare Disease Research and Orphan Product Development, Institute of Medicine, Washington, D.C.
- 2010 Legal issues in drug development and drug use / Invited Speaker  
Robert Wood Johnson Clinical Scholars Policy Speaker Series, Philadelphia, PA
- 2010 Methodological issues in comparative effectiveness research / Invited Speaker  
Health Affairs Comparative Effectiveness Research consortium, Washington, D.C.
- 2010 Sources of transformative innovation in drug development / Invited Plenary Speaker  
Robert Wood Johnson Investigator Award in Health Policy Research Meeting, Itsaca, IL
- 2011 Insiders' perspectives on off-label drug promotion / Invited Speaker  
Food and Drug Administration Drug Safety Oversight Board, White Springs, MD
- 2011 Transformative drug and device development / Invited Plenary Speaker  
Robert Wood Johnson Investigator Award in Health Policy Research Meeting, Princeton, NJ
- 2011 Institutional challenges at the FDA / Invited Plenary Speaker  
FDA at Crossroads National Meeting, Union of Concerned Scientists and GW School of Public Health, Washington, D.C.
- 2012 Asymmetry in the ability to communicate CER findings / Invited Speaker  
National Pharmaceutical Council, Washington, DC
- 2012 Reauthorization of the Medical Device User Fees Amendments: what it means for jobs, innovation and patients / Congressional Testimony  
House of Representatives Committee on Energy and Commerce Subcommittee on Health (Rep. Pitts, Chairman), Washington, DC
- 2012 Restrictions on promoting comparative effectiveness research (CER) / Invited Speaker  
Health Affairs kick-off symposium on promotion of CER, Washington, D.C.
- 2012 The roles of academia, industry, and patents in transformative drug development in oncology / Invited Plenary Speaker  
Robert Wood Johnson Investigator Award in Health Policy Research Annual Meeting, Princeton, NJ
- 2012 Patents and market exclusivity: a lever for incentivizing drug development? / Keynote  
18<sup>th</sup> Annual Thomas Langfitt Symposium on Health Care Policy, College of Physicians of Philadelphia and the University of Pennsylvania, Philadelphia, PA
- 2013 Research on COI: results from two national surveys / Invited Keynote Speaker  
FOCI Academe Meeting, Association of American Medical Colleges, Baltimore, MD
- 2013 The Food and Drug Administration in the 21st century / Invited Speaker  
Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and Bioethics, Cambridge, MA [national attendees]
- 2013 Issues and case studies in clinical trial data sharing: lessons and solutions / Invited Panelist

- 2013 Multi-Regional Clinical Trial Center, Harvard Global Health Institute [national attendees]  
Patient-centered outcomes research in rare diseases / Keynote Speaker
- 2013 14th Annual North American Lysosomal Storage Disease Registries Meeting, Chicago, IL  
Effect of drug detailing restrictions on prescribing of antidepressants and antipsychotics in children / Invited Lecture  
AcademyHealth annual meeting, Baltimore, MD
- 2013 High Priority Research Topics in Regulatory Science Related to Generic Drugs / Featured Speaker [with William Shrank]  
FDA Office of Generic Drugs Generic Drug User Fee Act (GDUFA) Regulatory Science Initiatives Public Meeting, Silver Spring, MD
- 2013 FDA Safety and Innovation Act (FDASIA) and the breakthrough drug designation: the risks of approving drugs on the basis of limited supporting data / Featured Speaker  
Briefings for Senate and House of Representative Congressional Staff, Washington, D.C.
- 2013 The practices and perils of “non-traditional” drug promotion / Invited Panelist  
Food and Drug Law Institute Advertising and Promotion for the Pharmaceutical, Medical Device, Biological, and Veterinary Medicine Industries, Washington, D.C.
- 2013 Prospects for regulation of off-label drug promotion in an era of expanding commercial speech protection / Featured Speaker  
University of North Carolina School of Law Annual Symposium, Chapel Hill, NC
- 2013 Are biomarkers patentable? / Keynote Speaker  
Global Biomarkers Consortium 2nd Annual Conference, Boston, MA [national attendees]
- 2013 Approval of new drugs on the basis of extremely limited data / Invited Speaker  
Center for Excellence in Education’s 30th Anniversary Celebration, Cambridge, MA [national attendees]
- 2013 Ethical implications of approval of drugs on the basis of limited data / Invited Speaker  
Greenwall Foundation Scholar Annual Meeting, New York City, NY
- 2013 Alternative or additional incentives for drug development / Invited Speaker  
Duke Law School Center for Innovation Policy Annual Meeting, Washington, D.C.
- 2014 Lessons for Follow-On Biologics from Generic Small Molecules / Speaker and Panelist  
Federal Trade Commission Follow-On Biologics Workshop, Washington, D.C.
- 2014 Specialty pharmaceuticals / Round table discussant  
Health Affairs Planning Meeting, Bethesda, MD
- 2014 Is sunshine the best disinfectant? Promise and perils of the Sunshine Act / Invited speaker  
American College of Physicians Internal Medicine 2014 annual meeting, Orlando, FL
- 2014 Ethical approaches to expanded access of investigational drugs / Round table discussant  
Engelberg Center for Health Care Reform, Brookings Institution, Washington, D.C.
- 2014 Tackling generic drug safety / Featured Speaker  
FDA Office of Generic Drugs Generic Drug User Fee Act Regulatory Science Initiatives Public Meeting, Silver Spring, MD
- 2014 Using ‘big data’ to change policy: physician financial relationships and prescribing practices / Invited panelist  
AcademyHealth Annual Research Meeting, San Diego, CA
- 2014 Generating evidence for use of new drugs and devices: what are the issues? / Keynote  
PORTAL/AAAS/NCHR conference on evidence development and FDA policy, Washington, D.C.
- 2014 21st Century Cures: Modernizing Clinical Trials / Congressional Testimony  
House of Representatives Committee on Energy and Commerce Subcommittee on Health (Rep. Pitts, Chairman), Washington, DC

- 2014 Lessons from the development of the most transformative drugs of the past 25 years / Invited speaker  
Robert Wood Johnson Foundation Investigator Award in Health Policy Research Annual Meeting, Indianapolis, IN
- 2014 FDA regulation of specialty drugs/ Invited Speaker  
Health Affairs kick-off symposium on specialty drugs, Washington, D.C.
- 2014 Health policy implications of FDA approval of new drugs and devices/ Grand Rounds  
Department of Health Services, Policy & Practice, Brown University School of Public Health, Providence, RI
- 2014 Preparing for biosimilars in the U.S.: what are the controversies?/ Invited Speaker  
Academy of Managed Care Pharmacy 2014 annual meeting, Boston, MA
- 2014 Regulation of off-label drug promotion and the First Amendment/ Invited Speaker  
Public Health in the Shadow of the First Amendment symposium at Yale Law School, New Haven, CT
- 2014 Regulation of new technologies: vaccines for non-communicable diseases/ Invited Speaker  
Emerging Issues and New Frontiers in FDA Regulation, Food and Drug Law Institute/Petrie-Flom Center Symposium, Washington, D.C.
- 2014 Subcommittee Hearing Investigating Generic Drug Prices / Congressional Testimony  
Senate Committee on Health, Education, Labor and Pensions Subcommittee on Primary Health and Aging (Sen. Sanders, Chairman), Washington, DC
- 2014 Ethical and clinical implications of expedited regulatory development and approval of new drugs and medical devices / Invited speaker  
Arthur & Ilene Dalinka Penn Grand Rounds Series, Hospital of the University of Pennsylvania Department of Medicine, Philadelphia, PA
- 2015 Adjusting regulatory standards to promote development of new CNS drugs  
Financial Incentives to Support Unmet Medical Needs for Nervous System Disorders: A Workshop, Institute of Medicine, Washington, D.C.
- 2015 Roles of academia, repurposing and orphan drugs in transformative drug development / Invited Speaker  
Health Affairs kick-off symposium on innovation, Washington, D.C.
- 2015 Expanded access to investigational drugs and other health policy topics / Invited Speaker  
National Physician's Alliance FDA task force, Boston, MA [national attendees]
- 2015 Managing uncertainty and reproductive rights with new technology / Invited speaker  
Institute of Medicine Workshop: Ethical and Social Policy Considerations of Novel Techniques for Prevention of Maternal Transmission of Mitochondrial DNA Diseases, Washington, D.C.
- 2015 Prospects for use of march-in rights to affect pricing of drugs emerging from government-sponsored research/Invited speaker  
Yale Health Law and Policy Society Guest Lecture Series, New Haven, CT
- 2015 Lessons from the most transformative drugs of the past 25 years / Invited speaker  
Michael M. Davis Lecture Series, Center for Health Administration Studies, University of Chicago School of Social Service Administration, Chicago, IL
- 2015 Does controversy during generic drug approval affect outcomes? Results from observational data, a systematic review, and surveys of patients and physicians/Invited speaker [with Joshua Gagne]  
FDA Office of Generic Drugs (OGD)/Office of Research & Standards, Rockville, MD
- 2015 Studying the post-market safety and rational use of generic drugs / Featured Speaker  
FDA Office of Generic Drugs Generic Drug User Fee Act (GDUFA) Regulatory Science

- Initiatives Public Meeting, Silver Spring, MD
- 2015 Assessing PDUFA 2012: breakthrough therapy and other expedited review and approval designations / Invited Speaker  
FDA Center for Drug Evaluation and Research PDUFA Reauthorization Public Meeting, Silver Spring, MD
- 2015 Role of Public Funding in the Development of Transformative Drugs / Invited Speaker  
Middle Class Prosperity Project Forum, U.S. Senate, Washington, D.C.
- 2016 Law and humanities: Blinding images in the law and other disciplines / Panel  
Association of American Law Schools Annual Meeting, New York, NY
- 2016 Innovation, Safety, and Value: The 21st Century Cures Bill / Invited Speaker  
Center for Drug Safety and Effectiveness, Johns Hopkins Bloomberg School of Public Health, Baltimore, MD
- 2016 Prescription Drug Prices: Origins and Options for Reform / Plenary speaker  
American Heart Association Quality of Care and Outcomes Research Annual Meeting, Phoenix, AZ
- 2016 Hospital administration and prescription drug prices / Plenary speaker  
American Hospital Association Annual Meeting, Washington, D.C.
- 2016 Balancing speed vs. evidence in cancer drug development / Grand Rounds speaker  
Memorial Sloan Kettering Cancer Center Survivorship, Outcomes, and Risk Seminar Series, New York, NY
- 2016 Pharma, Science, and Innovation: What Does the Future Hold for the Health Care Industry and for Patients? / Speaker and moderator (with Peggy Hamburg and Ken Frazier)  
Yale Law School Solomon Health Law and Corporate Law Centers' Craig Wasserman '86/Wachtell, Lipton, Rosen & Katz Alumni Breakfast, New York, NY
- 2016 High Drug Prices: Sources and Solutions / Invited Speaker  
American Medical Association Board of Delegates, Chicago, IL
- 2016 Regulatory Review Times and Adverse Event Reports in Cardiovascular Devices / Speaker  
American Society of Health Economics Biannual Meeting, Philadelphia PA
- 2016 Transforming Data to Inform Value: Balancing Innovation with Access / Panelist  
American Heart Association Corporate Forum Policy Dialogue, Washington, DC
- 2016 High Drug Prices and State-Based Solutions / Speaker  
Council of State Governments Medicaid Leadership Policy Academy, Washington, D.C.
- 2016 High-Cost Drugs: Ensuring Access without Hampering Innovation / Speaker  
Yale Law School, New Haven, CT
- 2016 Strategies for Ensuring Patient Access to Affordable Drug Therapies / Speaker  
National Academies of Science, Engineering and Medicine, Washington, D.C.
- 2016 Limiting Off-Label Promotion is Needed to Protect Patients / Speaker  
Part 15 Public Hearing: Manufacturer Communications Regarding Unapproved Uses of Approved or Cleared Medical Products, Food and Drug Administration, Silver Spring, MD
- 2016 Emerging Opportunities to Streamline Cancer Drug Development / Panelist  
President's Cancer Panel, Arlington, VA
- 2017 Expedited FDA approval and stem cell therapies / Keynote speaker  
International Society for Stem Cell Research Nucleus Forum, Berkeley, CA
- 2017 March-In Rights: Experiences and Prospects for Reducing Drug Prices / Speaker  
Knowledge Ecology International, Washington, D.C.
- 2017 Prescription Drug Pricing / Featured Speaker  
American Medical Association National Advocacy Conference, Washington, D.C.
- 2017 Right to Try and Expanded Access to Investigational Drugs / Featured speaker

- Pew Prescription Project: Framing the Debate on Right to Try, Washington, D.C.
- 2017 Ensuring Availability of Innovation and Prescription Drugs to Patients / Featured speaker  
America's Health Insurance Plans National Health Policy Conference, Washington, D.C.
- 2017 An Overview of the 21st Century Cures Act / Featured speaker  
National Comprehensive Cancer Network Institutional Review Board Directors Forum,  
Orlando, FL
- 2017 The Future of Prescription Drug Prices / Keynote Speaker  
Distinguished Lecture Series, Florida Hospital, Orlando, Florida
- 2017 Regenerative Medicine and the 21st Century Cures Act / Featured speaker  
National Academies of Science, Engineering, and Medicine Forum on Regenerative  
Medicine, Washington, D.C.
- 2017 Physicians' Knowledge and Perceptions about FDA Approval Standards / Invited Speaker  
Committee for Advanced Scientific Education Seminar, FDA, Silver Spring, MD
- 2017 Can Importation Address High Generic Drug Prices? / Featured Speaker [with Thomas J.  
Bollyky]  
Brookings Institution "Reining in Prescription Drug Prices", Washington, D.C.
- 2017 What is the Price of a Drug? / Invited panelist  
Financial Times US Healthcare & Life Sciences Summit, New York City, NY
- 2017 Prescriptions Drug Prices and Policy Reform Options / Keynote Speaker  
340B Coalition Summer Conference, Washington D.C.
- 2017 Generic drug competition: understanding demand, price, and supply / Invited Speaker  
Federal Trade Commission Workshop, Washington, D.C.
- 2017 An Interview with Rep. Henry Waxman / Interviewer  
Next Steps in Health Reform Conference, Washington College of Law at American  
University, Washington, D.C.
- 2018 FDA's Breakthrough Therapy Designation: Origins, (Early) Outcomes / Guest speaker  
Stanford Law School Law and Biosciences Workshop, Palo Alto, CA
- 2018 Prescription Drug Prices: Problems and Potential Solutions / 2018 Stuart Rome Lecture  
University of Maryland Francis Carey King School of Law, Baltimore, MD
- 2018 Prescriptions for Lowering Drug Prices / 2018 Rodman Lecture  
St. Jude Children's Research Hospital Grand Rounds, Memphis, TN
- 2018 Promoting Competition in the Prescription Drug Market / Invited speaker  
House of Representatives Antitrust Caucus Briefing, Washington, D.C.
- 2018 The Breakthrough Therapy Pathway: Policy Goals and Outcomes / Invited speaker  
The Commonwealth Fund Harkness Fellow Orientation Meeting, New York City, NY
- 2018 Ethical role of patients in FDA approval decisions / Invited speaker  
Stanford Center for Biomedical Ethics, Palo Alto, CA

### **International**

- 2005 Economic impact of patent extension on Medicaid drug expenditures / Invited Lecture  
International Society for Pharmacoepidemiology 21st annual meeting, Nashville, TN  
[international attendees]
- 2007 The patentability of pharmacoepidemiology methods / Invited Lecture  
International Society for Pharmacoepidemiology 23rd annual meeting, Quebec City,  
Canada
- 2007 Balancing drug innovation and cost-effective medical treatment in the US / Invited Lecture  
European Science Foundation semiannual meeting, Kiel, Germany
- 2009 Roundtable on delinking research and development incentives from prices: designing

- innovation inducement prizes for tuberculosis diagnostics and new drugs for tuberculosis and Chagas disease / Invited Panelist  
Knowledge Ecology International, Geneva, Switzerland
- 2010 The prevalence and cost of unapproved and non-evidence-based uses of selected orphan drugs / Invited Lecture  
International Society for Pharmacoepidemiology 26th annual meeting, Brighton, England
- 2013 Five models of incentives for drug innovation: successes, collateral effects, and lessons / Invited Lecture  
Médecins Sans Frontières, New York City, NY [international attendees]
- 2013 Intersection of market exclusivity and access to medicines / Roundtable Participant  
University of Melbourne-Vanderbilt International Roundtable Meeting, Honolulu, HI
- 2015 Eye of the beholder: legal views on drugs risks and causation / Plenary lecture  
International Society for Pharmacoepidemiology 31st annual meeting, Boston, MA [international attendees]
- 2015 Regulatory and legal issues for follow-on biologic drugs / Course faculty speaker  
International Society for Pharmacoepidemiology 31st annual meeting, Boston, MA [international attendees]
- 2015 Rethinking the economics of pharmaceutical innovation / Roundtable participant  
Open Society Foundations, New York, NY [international attendees]
- 2017 Drug regulation in the US: past, present, and future / Keynote speaker  
London School of Economics International Health Policy Conference, London, England
- 2018 Generic Drug Price Changes: Should the US be Looking to Canada? / Guest speaker  
York University, Toronto, Canada
- 2018 FDA's Breakthrough Therapy Designation: Origins, (Early) Outcomes / Keynote speaker  
University of Toronto Faculty of Law Health Law, Ethics & Policy Seminar, Canada
- 2018 Antibiotics and Innovation / Invited speaker  
Innovation Gaps and Life Sciences Frontiers, University of Copenhagen, Denmark

## **Report of Clinical Activities and Innovations**

### **Current Licensure and Certification**

- 2002 United States Patent and Trademark Office (Patent attorney license)
- 2004 National Board of Medical Examiners (Physician license)
- 2004 New York State Bar (Attorney license)
- 2005 American Board of Internal Medicine (Diplomate)
- 2005 Massachusetts Board of Registration in Medicine (License)

### **Practice Activities**

- |           |                     |  |  |
|-----------|---------------------|--|--|
| 2005-2009 | Attending physician | Internal Medicine Inpatient Ward, BWH                    | 15 hours per week / 4 weeks per year           |
| 2005-2011 | Attending physician | Hospitalist Service, Harvard Vanguard Medical Associates | 20 hours per month / 12 months per year        |
| 2005-     | Ambulatory Care     | Phyllis Jen Center for Primary Care, BWH                 | 1 half-day session per week / 4 hours per week |
| 2011-2013 | Attending physician | Hospitalist Service, BWH                                 | 20 hours per month / 12 months per year        |

## **Report of Education of Patients and Service to the Community Activities**

No activities or materials below were sponsored by outside entities.

2000-2001      Pennsylvania Health Law Project / Volunteer

### **Educational Material for Patients and the Lay Community**

#### **Monographs, articles and presentations in other media**

1. **Kesselheim A** and Outterson K. Super bugs call for super changes in drug-sale rules. [Op-Ed] *Boston Globe*, 15 Nov 2010, at A11.
2. **Kesselheim AS**. Does pharmaceutical industry marketing to medical students affect their prescribing choices as physicians? [Invited commentary] *Robert Wood Johnson Foundation Human Capital Blog*. 28 Jun 2011. Available at: <http://blog.rwjf.org/humancapital/?p=887>.
3. **Kesselheim AS**, Shiu N. *FTC v. Actavis*: the Supreme Court issues a reversal on reverse payments. [Invited commentary] *Health Affairs Blog*. 21 Jun 2013. Available at: <http://healthaffairs.org/blog/2013/06/21/ftc-v-actavis-the-supreme-court-issues-a-reversal-on-reverse-payments/#more-32326>
4. Lipsitch M, **Kesselheim AS**, Bell B, Levy S. Battling drug-resistant superbugs: can we win? [Webcast] *The Forum at the Harvard T.H. Chan School of Public Health*. February 5, 2014. Available on-line at: <http://theforum.sph.harvard.edu/events/battling-drug-resistant-superbugs/>.
5. **Kesselheim AS**, Tan YT. Accelerating Medicines Partnership: a new public-private collaboration for drug discovery. [Invited commentary] *Health Affairs Blog*. 8 Apr 2014. Available at: <http://healthaffairs.org/blog/2014/04/08/accelerating-medicines-partnership-a-new-public-private-collaboration-for-drug-discovery/>.
6. Darrow JJ, **Kesselheim AS**. New drug and device approval: what is sufficient evidence? [Invited commentary] *Health Affairs Blog*. 1 July 2014. Available at: <http://healthaffairs.org/blog/2014/07/01/new-drug-and-device-approval-what-is-sufficient-evidence/>
7. Maggs LR, **Kesselheim AS**. The role of Black Box Warnings in safe prescribing practices. [Invited commentary] *Health Affairs Blog*. 20 Aug 2014. Available at: <http://healthaffairs.org/blog/2014/08/20/the-role-of-black-box-warnings-in-safe-prescribing-practices/>
8. Maggs LR, **Kesselheim AS**. The short-term and long-term outlook of drug coupons. [Invited commentary] *Health Affairs Blog*. 12 Nov 2014. Available at: <http://healthaffairs.org/blog/2014/11/12/the-short-term-and-long-term-outlook-of-drug-coupons/>
9. Thacker PD, **Kesselheim AS**, Campbell EG. Will a new website empower patients to ask their physicians about financial relationships with industry? *JAMA Forum*. 17 Dec 2014. Available at: <http://newsatjama.jama.com/2014/12/17/jama-forum-will-a-new-website-empower-patients-to-ask-their-physicians-about-financial-relationships-with-industry/>
10. Sarpatwari A, **Kesselheim AS**. Ensuring timely approval of generic drugs. [Invited commentary] *Health Affairs Blog*. 24 March 2015. Available at: <http://healthaffairs.org/blog/2015/03/24/ensuring-timely-approval-of-generic-drugs/>
11. **Kesselheim AS**, Sarpatwari A. To spur innovation, make corporate cheaters pay. [Invited commentary] *Health Affairs Blog*. 30 April 2015. Available at: <http://healthaffairs.org/blog/2015/04/30/to-spur-medical-innovation-make-corporate-cheaters-pay/>
12. Greene J, **Kesselheim AS**. Selfie-medication: regulation of drug promotion in the Instagram era. *The Atlantic* 10 September 2015. Available at: <http://www.theatlantic.com/health/archive/2015/09/fda-drug-promotion-social-media/404563/>
13. Carrier MA, **Kesselheim AS**. The Daraprim price hike and a role for antitrust. [Invited

- commentary] *Health Affairs Blog*. October 21, 2015. Available at: <http://healthaffairs.org/blog/2015/10/21/the-daraprim-price-hike-and-a-role-for-antitrust/>.
14. Terry NP, Pasquale F, **Kesselheim AS**. Episode 26: EHR gag clauses, ACOs, the state of drug safety & price regulation & Kim Kardashian. [Podcast] *This Week in Health Law Podcast*. 2015 Sept 17. Available on-line at: <http://twihl.podbean.com/e/26-guest-aaron-kesselheim-ehr-gag-clauses-acos-the-state-of-drug-safety-price-regulation-kim-kardashian/>
  15. Pearson S, **Kesselheim AS**, Rosenthal M, Schnipper L. Drug pricing: public health implications. [Webcast] *The Forum at the Harvard T.H. Chan School of Public Health*. 2015 Oct 23. Available on-line at: <https://theforum.sph.harvard.edu/events/drug-pricing/>.
  16. Sax PE, Gallant JA, **Kesselheim AS**. Episode 8: Daraprim price hike. [Podcast] *Open Forum Infectious Diseases Podcast*. 2015 Nov 20. Available on-line at: [http://www.oxfordjournals.org/our\\_journals/ofid/podcasts.html](http://www.oxfordjournals.org/our_journals/ofid/podcasts.html)
  17. **Kesselheim AS**, Leape L, Gutierrez A, Arnaout R. Medical tests: inaccuracies, risks, and the public's health. [Webcast] *The Forum at the Harvard T.H. Chan School of Public Health*. 2015 Dec 11. Available on-line at: <https://theforum.sph.harvard.edu/events/medical-tests/>.
  18. **Kesselheim AS**. Why are we years away from a Zika vaccine? [Webcast] *Health Affairs*. 2016 Feb 11. Available on-line at: <http://healthaffairs.org/blog/2016/02/11/why-are-we-years-away-from-a-zika-vaccine/>.
  19. Engelberg AB, Avorn J, **Kesselheim AS**. Addressing generic drug unaffordability and shortages by globalizing the market for old drugs. [Invited commentary] *Health Affairs Blog*. 2016 Feb 23. Available on-line at: <http://healthaffairs.org/blog/2016/02/23/addressing-generic-drug-unaffordability-and-shortages-by-globalizing-the-market-for-old-drugs/>
  20. Sperling R, **Kesselheim A**, Tenaerts P, Goldstein J. Drug trials: challenges for Alzheimer's and other urgent needs. [Webcast] *The Forum at the Harvard T.H. Chan School of Public Health*. 2016 April 18. Available on-line at: <https://theforum.sph.harvard.edu/events/drug-trials/>.
  21. Rizvi Z, Kapczynski A, **Kesselheim A**. A simple way for the government to curb inflated drug prices. [Op-Ed] *Washington Post*. 13 May 2016. Available at: [https://www.washingtonpost.com/opinions/a-simple-way-for-the-government-to-curb-inflated-drug-prices/2016/05/12/ed89c9b4-16fc-11e6-aa55-670cabef46e0\\_story.html](https://www.washingtonpost.com/opinions/a-simple-way-for-the-government-to-curb-inflated-drug-prices/2016/05/12/ed89c9b4-16fc-11e6-aa55-670cabef46e0_story.html).
  22. Luo J, **Kesselheim AS**. Setting prescription drug prices: a comparison of strategies in the US, UK, Canada, Australia, and Germany. *Harvard Health Policy Review* 2016;15(2):4-9.
  23. **Kesselheim A**, Hey SP, Deak D, Lo B. Ethical tensions in expedited regulatory approval of new prescription drugs. [Invited commentary] *Health Affairs Blog*. 2016 June 23. Available on-line at: <http://healthaffairs.org/blog/2016/06/23/four-ways-to-address-the-ethical-tensions-around-expedited-approval-of-new-prescription-drugs/>
  24. Goldman AS, **Kesselheim AS**, Davis MH, Sachs RE, Singhroy D, Basey M, Maybarduk P. NIH patent policy. [Conference call] *Knowledge Ecology International IP Health Policy Update*. 2016 June 29. Available on-line at: <http://keionline.org/node/2608>.
  25. Pollack HA, Rector B, **Kesselheim AS**, Conti R. Drug Pricing: Value, Affordability, and Advocacy. [Webinar] *Doctors for America Expert Policy Webinar*. June 29, 2016.
  26. Court E, **Kesselheim AS**. Drugs that could cut billions from health costs. [Podcast] *Wall Street Journal Money, Markets, and More*. 2016 July 21. Available on-line at: <http://www.wsj.com/podcasts/drugs-that-could-cut-billions-from-health-costs/CA501FCE-5C22-41B2-A18B-294271722CDA.html>
  27. **Kesselheim AS**. The newest antibiotics on the block. [Podcast] *NEJM Journal Watch*. 15 Jul 2016. Available on-line at: [www.audiogest.org/NEJMJWinterviews](http://www.audiogest.org/NEJMJWinterviews).
  28. Hey SP, **Kesselheim AS**. Imprecise research threatens precision medicine. [Invited commentary] *STAT: First Opinion*. 2016 Aug 11. Available at:



- <https://www.statnews.com/2016/08/11/precision-medicine-research/>.
29. **Kesselheim AS**. Featured expert: NEJM Group Open Forum “Drug pricing: de-mystifying the power, politics, and practice behind today’s pharmaceutical economy.” 2016 Oct 12-22. [Web discussion] Available at: <https://medstro.com/groups/nejm-group-open-forum/discussions/300>
  30. **Kesselheim AS**. Juno trial deaths underscore need for greater transparency by FDA. [Invited commentary] *STAT: First Opinion*. 2016 Nov 24. Available at: <https://www.statnews.com/2016/11/24/deaths-juno-trial-transparency-fda/>
  31. Darrow J, **Kesselheim A**, Laskey-Su J. The future of precision medicine: great promise, significant challenges. [Invited commentary] *Health Affairs Blog*. 2017 Feb 28. Available at: <http://healthaffairs.org/blog/2017/02/28/the-future-of-precision-medicine-great-promise-significant-challenges/>
  32. Klitzman RL, **Kesselheim AS**, Holcombe K, Dehoney E. Implications of the 21st Century Cures Act: a webinar. *Columbia University Bioethics Program*. 2017 April 20. Available on-line at: <http://sps.columbia.edu/bioethics/events/04-20-2017-webinar-implications-of-the-21st-century-cures-act>.
  33. Hwang TJ, **Kesselheim AS**. Taxing drug price spikes: assessing the impact. [Invited commentary] *Health Affairs Blog*. 2017 May 12. Available on-line at: <http://healthaffairs.org/blog/2017/05/12/taxing-drug-price-spikes-assessing-the-potential-impact/>.
  34. Sarpatwari A, **Kesselheim AS**. Get generics to market faster. [Op-Ed] *Bloomberg View*. 2017 June 20. Available on-line at: <https://www.bloomberg.com/view/articles/2017-06-20/get-generic-drugs-to-market-faster>.
  35. **Kesselheim AS**, Lacey P. Fast-tracked drugs can save lives, but are they safe? [Televised roundtable] *Greater Boston with Jim Braude*. WGBH News. 2017 July 18. Available on-line at: <http://news.wgbh.org/2017/07/18/local-news/fast-tracked-drugs-can-save-lives-are-they-safe>
  36. Sinha MS, **Kesselheim AS**. The future of American science and medicine. [Blog post] *The Spoke*. 2017 July 25. Available on-line at: <http://www.wellesley.edu/albright/about/blog/3516-future-american-science-and-medicine>
  37. Terry NP, Pasquale F, **Kesselheim AS**, Sarpatwari A. Episode 107: Prescription Drug Price Metrics. [Podcast] *This Week in Health Law Podcast*. 2017 August 4. Available on-line at: <https://www.podbean.com/media/share/pb-vjp26-6db941>
  38. **Kesselheim AS**. The complexities behind high prescription prices: a conversation with Dr Aaron Kesselheim. [Podcast] *Medscape Cardiology: The Bob Harrington Show*. 2017 Aug 14. Available on-line at: <http://www.medscape.com/viewarticle/883281>.
  39. Redberg RF, **Kesselheim AS**, Califf RM. Are they safe? Drugs and devices receiving accelerated approval by the FDA. [Podcast] *JAMA Podcast*. 2017 Aug 15. Available on-line at: <http://jamanetwork.com/learning/audio-player/14650592>
  40. Darrow J, **Kesselheim A**. Nearly one-third of new drugs are no better than older drugs, and some are worse. [Invited commentary] *Health Affairs Blog*. 2017 Oct 6. Available on-line at: <http://healthaffairs.org/blog/2017/10/06/nearly-one-third-of-new-drugs-are-no-better-than-older-drugs-and-some-are-worse/>.
  41. Sherman M, Chandra A, **Kesselheim AS**. Navigating Payment Reform for Providers, Payors, and Pharma. [Web series] *NEJM Catalyst*. 2017 Nov 2. Available on-line at: <https://catalyst.nejm.org/events/navigating-payment-reform-providers-payers-pharma/>.
  42. Kapczynski A, **Kesselheim AS**. Three things Trump can do to bring drug price ‘way down.’ [Op-Ed] *Washington Post*. 2017 Nov 21. Available on-line at: [https://www.washingtonpost.com/opinions/what-trump-should-do-if-he-actually-wants-to-cut-drug-prices/2017/11/21/f7522422-be4f-11e7-8444-a0d4f04b89eb\\_story.html?utm\\_term=.3ef061b32123](https://www.washingtonpost.com/opinions/what-trump-should-do-if-he-actually-wants-to-cut-drug-prices/2017/11/21/f7522422-be4f-11e7-8444-a0d4f04b89eb_story.html?utm_term=.3ef061b32123).

43. Bollyky TJ, **Kesselheim AS**, Sharfstein JM. What Trump should actually do about the high cost of drugs. [Op-Ed] *New York Times*. 2018 May 14. Available on-line at: <https://www.nytimes.com/2018/05/14/opinion/trump-costs-drugs-pricing.html>.
44. **Kesselheim AS**, Mitchell D, Thomas K. Runaway train: America's drug price problem. [Webcast] *The Center for Health Journalism at the USC Annenberg School of Journalism*. 2018 May 17. Available on-line at: <https://www.centerforhealthjournalism.org/content/runaway-train-americas-drug-price-problem>.
45. Dafny L, Frank R, **Kesselheim AS**, Pearson S. U.S. drug prices: why are they so high? [Webcast] *The Forum at the Harvard T.H. Chan School of Public Health*. September 26, 2018. Available on-line at: <https://theforum.sph.harvard.edu/events/u-s-drug-prices/>.
46. Cohen M, Gupta R, Bollyky TJ, Ross JS, **Kesselheim AS**. Policy options for increasing generic drug competition through importation. [Invited commentary] *Health Affairs Blog*. 2019 January 7. Available on-line at: <https://www.healthaffairs.org/doi/10.1377/hblog20190103.333047/full/>.

### Patient educational materials

1. **Kesselheim AS**, Avorn J. True or false: common myths about generic drugs. [Patient monograph] September 2008. Available at: <http://www.mmcpo.org/resource/d/13754/commonmythsaboutgenericdrugs.pdf>.
2. **Kesselheim AS**, Avorn J. What are generic drugs? How can they help me? [Patient monograph] September 2008. Available at: <http://www.mmcpo.org/resource/d/60045/whataregenericdrugs.pdf>.
3. **Kesselheim AS**, Avorn J. Generics are powerful medicines. [Patient monograph] September 2008. Available at: <http://www.mmcpo.org/resource/d/68289/Genericsarepowerfulmeds.pdf>.
4. **Kesselheim AS**, Avorn J. Frequently asked questions about generic drugs. [Patient monograph] September 2008. Available at: <http://www.mmcpo.org/resource/d/48358/faqgenericdrugs.pdf>.
5. **Kesselheim AS**, Avorn J. What brand-name drug companies don't want you to know: how they keep generics off the market. [Patient monograph] September 2008. Available at: [http://www.usaindiana.org/document/generics/Dont\\_want.pdf](http://www.usaindiana.org/document/generics/Dont_want.pdf)<http://www.genericsarepowerful.org/learn?d=0002>

There are numerous articles, interviews, and information products in the national and global popular media (on-line, in print, broadcast news, etc.) related to my work or for which I have served as a contributor, including the following selected samples:

1. Press M. Interview with Aaron Kesselheim, M.D., J.D., M.P.H: patent attorney, general internist and health services researcher from Harvard Medical School. *RWJF Clinical Scholars Health Policy Broadcast*. Broadcast May 17, 2010. Available at: <http://rwjcsp.unc.edu/resources/podcast/archive.html>
2. Rooney E. Antibiotics reform. *The Emily Rooney Show*. Broadcast November 30, 2010. Available at: <http://www.wgbh.org/programs/The-Emily-Rooney-Show-Podcast-1162/episodes/Airport-Security-Boston-Accent-Reduction-Antibiotics-Reform-22067>.
3. Kolata G. Pills morph as patients try to cope. *New York Times*. July 11, 2011. Available on-line at: [http://www.nytimes.com/2011/07/12/health/12pills.html?\\_r=1&](http://www.nytimes.com/2011/07/12/health/12pills.html?_r=1&)
4. Song S. Lipitor vs. Crestor: cholesterol drugs on a par. *Time Magazine*. November 16, 2011. Available on-line at: <http://healthland.time.com/2011/11/16/lipitor-vs-crestor-cholesterol-drugs-are-on-a-par/>
5. Understanding how 'the system' can be made to work better for patients. *Health Affairs*. December 2011.
6. Investigator examines path to more affordable and effective drugs. *Robert Wood Johnson Foundation*. January 30, 2012. Available at: <http://www.rwjf.org/content/rwjf/en/about->

- rwjf/newsroom/newsroom-content/2011/12/breaking-new-ground-in-research/investigator-examines-path-to-more-affordable-and-effective-drug.html?cid=XEM\_205596
7. National Pharmaceutical Council. *CER & academic detailing: Harvard's Dr. Kesselheim explains*. YouTube. Posted February 16, 2012. Available on-line at <http://www.youtube.com/watch?v=e0Xs4dH5F8U>.
  8. AJMCTv. *Dr. Aaron Kesselheim discusses comparative effectiveness research*. YouTube. Posted April 3, 2012. Available on-line at <http://www.youtube.com/watch?v=vTdIZ0d934w>.
  9. Out of the mire?: The justice department may spoil the drugmaker's fresh start. *The Economist*. April 28, 2012. Available on-line at: <http://www.economist.com/node/21553512>
  10. Krumholz H. A suggestion to restore faith in pharma studies. *Pharmalot*. September 20, 2012. Available at: [www.pharmalot.com/2012/09/the-op-ed-a-suggestion-to-restore-faith-in-pharma-studies/](http://www.pharmalot.com/2012/09/the-op-ed-a-suggestion-to-restore-faith-in-pharma-studies/)
  11. Conaboy C. Study: physicians give less credence to studies funded by pharmaceutical industry. *Boston Globe*. September 20, 2012. Available on-line at: <http://www.boston.com/whitecoatnotes/2012/09/20/study-physicians-give-less-credence-studies-funded-pharmaceutical-industry/xd8MAnN5SqizUiBO6kTTOJ/story.html>
  12. Rehman J. Can the source of funding for medical research affect the results? *Scientific American*. September 23, 2012. Available at: <http://blogs.scientificamerican.com/guest-blog/2012/09/23/can-the-source-of-funding-for-medical-research-affect-the-results/>
  13. Chen P. Are doctors too wary of drug companies? *New York Times*. October 18, 2012. Available on-line at: <http://well.blogs.nytimes.com/2012/10/18/are-doctors-too-wary-of-drug-companies/>
  14. Lyons C. What's on your doctor's mind? *Boston Magazine*. December 2012. Available on-line at: <http://www.bostonmagazine.com/articles/2012/11/boston-best-doctors-how-doctors-think-top-docs/>
  15. Tinker B. Don't judge that generic pill by its color. *CNN.com*. December 31, 2012. Available on-line at: <http://thechart.blogs.cnn.com/2012/12/31/hfr-123112-4pet-dont-judge-your-generic-pill-by-its-color/>
  16. Bakalar N. The confusion of pill coloring. *New York Times*. December 31, 2012. Available on-line at: <http://well.blogs.nytimes.com/2012/12/31/the-confusion-of-pill-coloring/>
  17. Conaboy C. Study authors: On medical school conflict of interest policies, more enforcement needed. *Boston Globe*. February 28, 2013. Available on-line at: <http://www.boston.com/whitecoatnotes/2013/02/28/study-authors-medical-school-conflict-interest-policies-more-enforcement-needed/q06z6QUFQdyfUq36H8pahK/story.html>
  18. Britt R. Will drug makers see other nations challenge drug patents? *Wall Street Journal MarketWatch*. April 1, 2013. Available on-line at: <http://blogs.marketwatch.com/health-exchange/2013/04/01/will-drug-makers-see-other-nations-challenge-drug-patents/>.
  19. Conaboy C. 1 in 4 Mass. physicians received industry gifts, payments. *Boston Globe*. May 1, 2013. Available on-line at: <http://www.boston.com/lifestyle/health/blogs/white-coat-notes/2013/05/01/mass-physicians-received-industry-gifts-payments/RbxWq4YEEvJMwk0Xt6N0QL/blog.html>
  20. Rabin RC. Doctors' lucrative industry ties. *New York Times*. May 13, 2013. Available on-line at: <http://well.blogs.nytimes.com/2013/05/13/doctors-lucrative-industry-ties/>
  21. Chen P. For med students, love from the drug rep. *New York Times*. October 4, 2013. Available on-line at: [http://well.blogs.nytimes.com/2013/10/03/for-med-students-love-from-the-drug-rep/?\\_r=0](http://well.blogs.nytimes.com/2013/10/03/for-med-students-love-from-the-drug-rep/?_r=0)
  22. Schwarz A. The selling of attention deficit disorder. *New York Times*. December 15, 2013. Available on-line at: [http://www.nytimes.com/2013/12/15/health/the-selling-of-attention-deficit-disorder.html?\\_r=1&](http://www.nytimes.com/2013/12/15/health/the-selling-of-attention-deficit-disorder.html?_r=1&)

23. New York Times Video. *How drug companies sell A.D.H.D.* New York Times website. Posted December 15, 2013. Available on-line at [http://www.nytimes.com/2013/12/15/health/the-selling-of-attention-deficit-disorder.html?\\_r=1&#videoModal](http://www.nytimes.com/2013/12/15/health/the-selling-of-attention-deficit-disorder.html?_r=1&#videoModal).
24. Rabin RC. The device maker's shortcut. *New York Times*. April 1, 2014, at D4. Available on-line at: [http://well.blogs.nytimes.com/2014/03/31/the-device-makers-shortcut/?\\_php=true&\\_type=blogs&hpw&rref=health&\\_r=0](http://well.blogs.nytimes.com/2014/03/31/the-device-makers-shortcut/?_php=true&_type=blogs&hpw&rref=health&_r=0)
25. Silverman E. Can pharma sales reps influence prescribing for unapproved uses? *Wall Street Journal*. June 9, 2014. Available on-line at: <http://blogs.wsj.com/pharmalot/2014/06/09/can-pharma-sales-reps-influence-prescribing-for-unapproved-uses>
26. Rosenthal E. Rapid price increases for some generic drugs catch users by surprise. *New York Times*. July 9, 2014 at A16.
27. Dennis B. FDA has free speech, safety issues to weigh in review of 'off-label' marketing rules. *Washington Post*. July 9, 2014. Available on-line at: [http://m.washingtonpost.com/national/health-science/2014/07/09/3708dd6a-fbc4-11e3-8176-f2c941cf35f1\\_story.html](http://m.washingtonpost.com/national/health-science/2014/07/09/3708dd6a-fbc4-11e3-8176-f2c941cf35f1_story.html)
28. Thomas K. Financial ties between doctors and health care firms detailed. *New York Times*. October 2, 2014 at B1.
29. LaPook J [video]. Why some generic drug prices are skyrocketing. *CBS Evening News*. November 12, 2014. Available on-line at: <http://www.cbsnews.com/news/generic-drug-prices-skyrocketing/>
30. Godman H. Best tips to stay on your medication and stay healthy. *Harvard Health Letter*. 2015;40(6):1,7.
31. Perrone M. Obama nominates Dr. Robert Califf, cardiologist and FDA deputy, to be next FDA commissioner. *Associated Press/US News & World Report*. September 15, 2015. Available on-line at: <http://www.usnews.com/news/business/articles/2015/09/15/obama-to-nominate-fdas-no-2-official-to-lead-agency>.
32. Tavernise S. F.D.A. nominee Califf's ties to drug makers worry some. *New York Times*. September 19, 2015. Available on-line at: <http://www.nytimes.com/2015/09/20/health/fda-nominee-califfs-ties-to-drug-industry-raise-questions.html>
33. Flatow I [radio segment]. Why generic doesn't mean cheap. *PRI's The World Science Friday*. September 25, 2015. Available on-line at: <http://www.sciencefriday.com/#path/segment/09/25/2015/why-generic-doesn-t-mean-cheap.html>
34. Loftus P. Drug firms buy pricey vouchers to speed products to market. *Wall St Journal*. October 20, 2015.
35. Pollack A. Martin Shkreli's arrest gives drug makers cover. *NY Times*. December 18, 2015. Available on-line at: <http://www.nytimes.com/2015/12/18/business/martin-shkreli-arrest-gives-drug-makers-cover.html>.
36. Span P. A prescription for confusion: when to take all those pills. *NY Times*. December 22, 2015. Available on-line at: [http://www.nytimes.com/2015/12/22/health/a-prescription-for-confusion-when-to-take-all-those-pills.html?\\_r=0](http://www.nytimes.com/2015/12/22/health/a-prescription-for-confusion-when-to-take-all-those-pills.html?_r=0).
37. Sweetland Edwards H. Why can't drug costs be reined in? *Time*. May 30, 2016. Available on-line at: <http://time.com/4341416/why-cant-drug-costs-be-reined-in/>.
38. Timmerman L and Tirrell M [podcast]. The death of a child and a golden ticket for drug makers. *STAT Signal Podcast*. January 18, 2017. Available on-line at: <https://www.statnews.com/2017/01/18/cancer-children-priority-review-vouchers/>.
39. Dankosky J [radio segment]. Does faster drug approval lead to better medicine? *PRI's The World Science Friday*. November 18, 2017. Available on-line at: <https://www.sciencefriday.com/segments/does-faster-drug-approval-lead-to-better-medicine/>

40. Burton TM. Tax overhaul looks set to cut credits for drugs targeting rare diseases. *Wall St. Journal*. December 8, 2017. Available on-line at: <https://www.wsj.com/articles/tax-overhaul-looks-set-to-cut-credits-for-drugs-targeting-rare-diseases-1512745739>.
41. Karlin-Smith S. How Trump's HHS nominee's drug company 'gamed' a patent. *Politico*. January 9, 2018. Available on-line at: <https://www.politico.com/story/2018/01/08/trump-azar-patent-drug-company-268942>.
42. Green J and Cortez M. Trump is getting a physical. What about your CEO? *Bloomberg News*. January 12, 2018. Available on-line at: <https://www.bloomberg.com/news/articles/2018-01-12/the-ceo-president-is-getting-a-physical-what-about-your-ceo>.
43. Burton TM. Trump moves to cut costs for prescription drugs. *Wall St. Journal*. February 9, 2018. Available on-line at: <https://www.wsj.com/articles/trump-moves-to-cut-costs-for-prescription-drugs-1518187386>.
44. Thomas K. Patients eagerly awaited a generic drug. Then they saw the price. *New York Times*. February 24, 2018. Available on-line at: <https://www.nytimes.com/2018/02/23/health/valeant-drug-price-syprine.html>.
45. Ellis Nutt A. One of America's most popular drugs — first aimed at schizophrenia — reveals the issues of 'off-label' use. *Washington Post*. March 30, 2018. Available on-line at: [https://www.washingtonpost.com/national/health-science/one-of-americas-most-popular-drugs--first-aimed-at-schizophrenia--reveals-the-issues-of-off-label-use/2018/03/28/78a538ca-1e27-11e8-b2d9-08e748f892c0\\_story.html?utm\\_term=.32ba5b2078d6](https://www.washingtonpost.com/national/health-science/one-of-americas-most-popular-drugs--first-aimed-at-schizophrenia--reveals-the-issues-of-off-label-use/2018/03/28/78a538ca-1e27-11e8-b2d9-08e748f892c0_story.html?utm_term=.32ba5b2078d6).
46. Wapner J. Medication just keeps getting more expensive—and big pharma won't explain why. *Newsweek* April 24, 2018. Available on-line at: <http://www.newsweek.com/medication-keeps-getting-more-expensive-and-big-pharma-wont-explain-why-897925>.
47. Johnson CY. The truth about 'breakthrough' drugs. *Washington Post*. July 17, 2018. Available on-line at: [https://www.washingtonpost.com/news/to-your-health/wp/2018/07/17/the-truth-about-breakthrough-drugs/?noredirect=on&utm\\_term=.6e934aabbacc](https://www.washingtonpost.com/news/to-your-health/wp/2018/07/17/the-truth-about-breakthrough-drugs/?noredirect=on&utm_term=.6e934aabbacc).
48. Frakt A. Something happened to US drug costs in the 1990s. *NY Times*. November 12, 2018. Available on-line at: <https://www.nytimes.com/2018/11/12/upshot/why-prescription-drug-spending-higher-in-the-us.html>.

## **Report of Scholarship**

\*\* *Indicates mentee as co-author*

### **Peer reviewed publications in print or other media**

#### Research investigations

1. **Kesselheim AS**. Deception and presidential disability: a historical analysis. *Trans & Stud Coll Phys Phila* 2001;23:87-98.
2. **Kesselheim AS**. What's the appeal? Trying to control managed care medical necessity decisionmaking through a system of external appeals. *Univ of Penn Law Rev* 2001;149:873-920.
3. **Kesselheim AS**. Privacy versus the public's right to know: presidential health and the White House physician. *J Legal Med* 2002;23:523-545.
4. Rudnick MR, **Kesselheim A**, Goldfarb S. Contrast-induced nephropathy: how it develops, how to prevent it. *Cleveland Clinic J Med* 2006;73:75-80, 83-87.
5. **Kesselheim AS** and Studdert DM. Characteristics of physicians who frequently act as expert witnesses in neurological birth injury litigation. *Obstet & Gyn* 2006;108:273-279.
  - Stokes RL 3rd. Characteristics of physicians who frequently act as expert witnesses in neurologic birth injury litigation. [Letter to the editor] *Obstetrics & Gynecology* 2006;108(6):1552-1553.

- **Kesselheim AS**, Studdert DM. Characteristics of physicians who frequently act as expert witnesses in neurological birth injury litigation. [Author reply] *Obstetrics & Gynecology* 2006;108:1552-3.
6. **Kesselheim AS**, Fischer MA, Avorn J. Extensions of intellectual property rights and delayed adoption of generic drugs: effects on Medicaid spending. *Health Affairs* 2006;25:1637-1647.
    - Copeland KR, Boccuzzi SJ. Brand-name versus generic. [Letter to the Editor] *Health Affairs* 2007;26(4):1198-1199.
    - **Kesselheim AS**, Fischer MA, Avorn J. Brand name versus generic: the authors respond. [Author reply] *Health Affairs* 2007;26:1199.
  7. **Kesselheim AS** and Avorn J. The role of litigation in defining drug risks. *JAMA* 2007;297:308-311.
  8. Shrank WH, Agnew-Blais J, Choudhry NK, Wolf MS, **Kesselheim AS**, Avorn J, Shekelle P. The variability and quality of medication container labels. *Archives of Internal Medicine* 2007;167:1760-1765.
  9. Outterson K and **Kesselheim AS**. Putting patients first: a market-based licensing proposal for HPV vaccines and other patented medical products in developing countries. *Health Affairs* 2008;27:130-139.
  10. **Kesselheim AS** and Studdert DM. Whistleblower-initiated enforcement actions against health care fraud and abuse in the United States, 1996-2005. *Annals of Internal Medicine* 2008;149:342-349.
  11. **Kesselheim AS**, Misono AS\*\*, Lee JL, Stedman MR, Brookhart MA, Choudhry NK, Shrank WH. The clinical equivalence of generic and brand-name drugs used in cardiovascular disease: a systematic review and meta-analysis. *JAMA* 2009;300:2514-2526.
    - Carter BL. Equivalence of generic and brand-name drugs for cardiovascular disease. [Letter to the Editor] *JAMA* 2009;301(16):1654.
    - Reiffel JA. Equivalence of generic and brand-name drugs for cardiovascular disease. [Letter to the Editor] *JAMA*. 2009;301(16):1655.
    - Zema MJ. Equivalence of generic and brand-name drugs for cardiovascular disease. [Letter to the Editor] *JAMA*. 2009;301(16):1654-5.
    - **Kesselheim AS**, Misono AS, Shrank WH. The clinical equivalence of generic and brand-name drugs used in cardiovascular disease: a systematic review and meta-analysis. [Author reply] *JAMA* 2009;301:1655-1656.
    - Manns B. ACP Journal Club. Brand-name drugs are not more effective than generic versions for treating cardiovascular disease. [Review] *Annals of Internal Medicine* 2009;150(8):JC4-6.
    - Manns B. Review: brand-name drugs are not more effective than generic versions for treating cardiovascular disease. [Review] *Evidence Based Medicine* 2009;14(3):81.
  12. **Kesselheim AS** and Studdert DM. Professional oversight of physician expert witnesses: an analysis of complaints to the Professional Conduct Committee of the American Association of Neurological Surgeons, 1992-2006. *Annals of Surgery* 2009;249:168-172.
  13. **Kesselheim AS** and Avorn J. Using patent data to assess the value of pharmaceutical innovation. *Journal of Law, Medicine, and Ethics* 2009;37(2):176-183.
  14. **Kesselheim AS**, Stedman MR, Bubrick EJ, Gagne JJ, Misono AS\*\*, Lee JL, Brookhart MA, Avorn J, Shrank WH. Seizure outcomes following use of generic vs. brand-name antiepileptic drugs: a systematic review and meta-analysis. *Drugs* 2010;70(5):605-621.
  15. **Kesselheim AS**, Studdert DM, Mello MM. Whistle-blowers' experiences in fraud litigation against pharmaceutical companies. *New England Journal of Medicine* 2010;362:1832-1839.
  16. Kramer DB\*\*, **Kesselheim AS**, Brock DW, Maisel WH. The ethical and legal views of physicians

- regarding deactivation of cardiac implantable electrical devices: a quantitative assessment. *Heart Rhythm* 2010;7(11):1537-1542.
17. **Kesselheim AS**, November MT, Lifford KL, McElrath TF, Puopolo AL, Orav EJ, Studdert DM. Risk factors for neurological impairment among infants following nonreassuring fetal heart rate patterns during labor. *Journal of Evaluation in Clinical Practice* 2010;16(3):476-483.
  18. Shrank WH, Choudhry NK, Agnew-Blais J, Federman AD, Liberman JN, Liu J, **Kesselheim AS**, Brookhart MA, Fischer MA. State generic substitution laws can lower drug outlays under Medicaid. *Health Affairs* 2010;29(7):1383-1390.
  19. Kramer DB\*\*, **Kesselheim AS**, Brock DW, Maisel WH. Ethical and legal views regarding deactivation of cardiac implantable electrical devices in patients with hypertrophic cardiomyopathy. *American Journal of Cardiology* 2011;107(7):1071-1075.
  20. **Kesselheim AS**, Mello MM, Studdert DM. Strategies and practices in off-label marketing of pharmaceuticals: a retrospective analysis of whistleblower complaints. *PLoS Medicine* 2011;8(4):e1000431.
  21. Shrank WH, Liberman JN, Fischer MA, Avorn J, Kilabuk E, Chang A, **Kesselheim AS**, Brennan TA, Choudhry NK. The consequences of requesting “dispense as written.” *American Journal of Medicine* 2011;124(4):309-317.
  22. Austad KE\*\*, Avorn J, **Kesselheim AS**. Medical students’ exposure to and attitudes about the pharmaceutical industry: a systematic review. *PLoS Medicine* 2011;8(5):e1001037.
  23. **Kesselheim AS**, Myers JA, Avorn J. Characteristics of clinical trials to support approval of orphan vs nonorphan drugs for cancer. *JAMA* 2011;305:2320-2326.
    - Saltonstall PL. Clinical trials of orphan drugs for cancer. [Letter to the editor] *JAMA* 2011;306(14):1545.
    - **Kesselheim AS**, Avorn J. Approval of orphan drugs for cancer. [Author reply] *JAMA* 2011;306:1545-1546.
  24. Avorn J and **Kesselheim AS**. The NIH’s new drug-development initiative: socialization of risk and privatization of gain? *Nature Medicine* 2011;17(10):20.
  25. **Kesselheim AS**, Darby DL\*\*, Studdert DM, Glynn RJ, Levin RL, Avorn J. False Claims Act prosecution did not deter off-label drug use in the case of Neurontin. *Health Affairs* 2011;30(12):2318-2327.
    - Greenberg PE, Pike C, Sisitsky T. Confounding factors in off-label drug use. [Letter to the editor] *Health Affairs* 2012;31(2):460.
    - **Kesselheim AS**, Darby DL, Avorn J. Off-label use of Neurontin. [Author reply] *Health Affairs* 2012;31(2):460.
  26. **Kesselheim AS**. An empirical review of major legislation affecting drug development: past experiences, effects, and unintended consequences. *Milbank Quarterly* 2011;89(3):450-502.
  27. **Kesselheim AS**, Lee JL, Avorn J, Servi A, Shrank WH, Choudhry NK. Conflict of interest in oncology publications. *Cancer* 2012;118(1):188-195.
  28. **Kesselheim AS**, Myers JA, Solomon DH, Winkelmayr WC, Levin R, Avorn J. The prevalence and cost of unapproved uses of top-selling orphan drugs. *PLoS One* 2012;7(2):e31894.
  29. Kramer DB\*\*, Xu S, **Kesselheim AS**. How does medical device regulation perform in the United States and the European Union? A systematic review. *PLoS Medicine* 2012;9(7):e1001276.
  30. **Kesselheim AS**, Wang B, Studdert DM, Avorn J. Conflict of interest reporting by authors involved in promotion of off-label drug use: an analysis of journal disclosures. *PLoS Medicine* 2012;9(8):e1001280.
  31. Greene JA, Choudhry NK, **Kesselheim AS**, Brennan TA, Shrank WH. Changes in direct-to-consumer pharmaceutical advertising following Rx to OTC shift. *JAMA* 2012;308(10):973-975.
  32. **Kesselheim AS**, Robertson CT, Myers JA, Rose SL, Gillet V, Ross KM, Glynn RJ, Joffe S, Avorn

- J. A randomized study of how physicians interpret research funding disclosures. *New England Journal of Medicine* 2012;367:1119-1127.
- Drazen JM. Believe the data. [Editorial] *New England Journal of Medicine* 2012;367(12):1152-1153.
  - Feuerstein JD, Leffler DA, Cheifetz AS. How physicians interpret research funding disclosures. [Letter to the Editor] *New England Journal of Medicine* 2012;367(24):2358-2359.
  - Fonseca R. How physicians interpret research funding disclosures. [Letter to the Editor] *New England Journal of Medicine* 2012;367(24):2358.
  - **Kesselheim AS**, Avorn J. How physicians interpret research funding disclosures. [Author reply] *New England Journal of Medicine* 2012;367:2360.
  - Reider B. Do We Care? Should We? [Comment] *American Journal of Sports Medicine* 2016;44(4):835-837.
33. Amin T and **Kesselheim AS**. Secondary patenting of branded pharmaceuticals: a case study of how patents on two HIV drugs could be extended for decades. *Health Affairs* 2012;31:2286-2294.
34. Robertson CT, Rose SL, **Kesselheim AS**. Effect of financial relationships on the behaviors of health care professionals: a review of the evidence. *Journal of Law, Medicine & Ethics* 2012;40(3):452-466.
35. Xu S\*\*, Avorn J, **Kesselheim AS**. Origins of medical innovation: the case of coronary artery stents. *Circulation: Cardiovascular Quality and Outcomes* 2012;5(6):743-749.
- Tomaselli GF. Origins of medical innovation: the case of coronary artery stents. [Editorial] *Circulation: Cardiovascular Quality and Outcomes* 2012;5(6):741-742.
36. **Kesselheim AS**, Misono AS\*\*, Shrank WH, Greene JA, Doherty M, Avorn J, Choudhry NK. Variations in pill appearance of antiepileptic drugs and the risk of non-adherence. *JAMA Internal Medicine* 2013;173(3):202-208.
- Yu LX, Geba GP. Generic pills from the patient perspective: dressed for success? [Editorial] *JAMA Internal Medicine* 2013;173(3):208-209.
  - Covinsky KE. Debating effect sizes. [Editorial] *JAMA Internal Medicine* 2013;173(3):209.
  - Appel K. Einfluss von Tablettenform und -farbe. [Commentary in German] *Fortschritte der Neurologie-Psychiatrie* 2013;81(6):301.
37. **Kesselheim AS**, Robertson CT, Siri K, Batra P, Franklin JM. Distributions of industry payments to Massachusetts physicians. *New England Journal of Medicine* 2013;368(22):2049-2052.
38. **Kesselheim AS**, Avorn J. The most transformative drugs of the past 25 years: a survey of physicians. *Nature Reviews: Drug Discovery* 2013;12(6):425-431.
39. Austad KE\*\*, Avorn J, Franklin JM, Kowal MK, Campbell EG, **Kesselheim AS**. Changing interactions between physician trainees and the pharmaceutical industry: a national survey. *Journal of General Internal Medicine* 2013;28(8):1064-1071.
- Ramachandran R, Hams M, Silver-Isenstadt J. Physician trainees' interactions with the pharmaceutical industry. [Letter to the Editor] *Journal of General Internal Medicine* 2013;28(10):1266.
  - Austad KE\*\*, Avorn J, Franklin JM, **Kesselheim AS**. Physician trainees' interaction with the pharmaceutical industry. [Author reply] *Journal of General Internal Medicine* 2013;28:1267.\*\*
40. **Kesselheim AS**, Franklin JM, Avorn J, Duke JD. Speaking the same language?: International variations in the safety information accompanying top-selling prescription drugs. *BMJ Quality & Safety* 2013;22:727-734.
41. **Kesselheim AS**, Wang B\*\*, Avorn J. Defining 'innovativeness' in drug development: a



- systematic review. *Clinical Pharmacology and Therapeutics* 2013;94(3):336-48.
42. Wang B\*\*, Avorn J, **Kesselheim AS**. Clinical and regulatory features of drugs not initially approved by the FDA. *Clinical Pharmacology and Therapeutics* 2013;94(6):670-677.
  43. Ross JS, **Kesselheim AS**. Prescription-drug coupons—no such thing as a free lunch. *New England Journal of Medicine* 2013;369:1188-1189.
  44. Outterson K, Powers JH, Seoane-Vazquez E, Rodriguez-Monguio R, **Kesselheim AS**. Approval and withdrawal of new antibiotics and other anti-infectives in the US, 1980-2009. *Journal of Law, Medicine, and Ethics* 2013;41(3):688-696.
  45. Gagne JJ, Polinski JM, **Kesselheim AS**, Choudhry NK, Hutchins D, Matlin OS, Tong A, Shrank WH. Patterns and predictors of generic narrow therapeutic index drug use among older adults. *Journal of the American Geriatrics Society* 2013;61(9):1586-1591.
  46. Hwang TJ\*\*, Carpenter D, **Kesselheim AS**. Assessment of US pathway for approving medical devices for rare conditions. *BMJ* 2014;348:g217.
  47. Rome BN\*\*, Kramer DB\*\*, **Kesselheim AS**. FDA approval of cardiac implantable electrical devices via original and supplement pre-market approval pathways, 1979-2012. *JAMA* 2014;311(4):385-391.
    - Goodman SN, Redberg RF. Opening the FDA black box. [Editorial] *JAMA* 2014;311(4):361-363.
    - McCarthy M. Quality of evidence behind FDA approvals varies widely. [Review] *BMJ* 2014;348:g1075.
  48. Hwang TJ\*\*, Avorn J, Carpenter DP, **Kesselheim AS**. Quantifying the Food and Drug Administration's rulemaking delays highlights need for transparency. *Health Affairs* 2014;33(2):309-315.
  49. **Kesselheim AS**, Xu S\*\*, Avorn J. Clinicians' contributions to the development of coronary artery stents: a qualitative study of transformative device innovation. *PLoS One* 2014;9(2):e88664.
  50. Hwang TJ\*\*, **Kesselheim AS**, Bourgeois FT. Postmarketing trials and pediatric device approvals. *Pediatrics* 2014;133(5):e1197-202.
  51. Gellad WF, Choi PC, Mizah M, Good CB, **Kesselheim AS**. Assessing the chiral switch: approval and use of single-enantiomer drugs, 2001-2011. *American Journal of Managed Care* 2014;20(3):e90-e97.
  52. Hwang TJ\*\*, Carpenter D, **Kesselheim AS**. Target small firms for antibiotic innovation. *Science* 2014;344(6187):967-969.
  53. Larkin I, Ang D, Avorn J, **Kesselheim AS**. Restrictions on pharmaceutical detailing reduced off-label prescribing of antidepressants and antipsychotics in children. *Health Affairs* 2014;33(6):1014-1023.
  54. Austad KE\*\*, Avorn J, Franklin JM, Campbell EG, **Kesselheim AS**. Association of marketing interactions with medical trainees' knowledge about evidence-based prescribing: results from a national survey. *JAMA Internal Medicine* 2014;174(8):1283-1290.
    - Ross JS. Restricting interactions with industry to promote evidence-based prescribing. [Editorial] *JAMA Internal Medicine* 2014;174(8):1290.
  55. Polinski JM, **Kesselheim AS**, Frolkis JP, Wescott P, Allen-Coleman C, Fischer MA. A matter of trust: patient barriers to primary medication adherence. *Health Education Research* 2014;29(5):755-763.
  56. Darrow JJ\*\*, **Kesselheim AS**. Drug development and FDA approval, 1938-2013. *New England Journal of Medicine* 2014;370(26):e39.
  57. **Kesselheim AS**, Bykov K, Tong A, Doherty M, Avorn J, Choudhry NK. Burden of changes in pill appearance for patients receiving generic cardiovascular medications after myocardial infarction: cohort and nested case-control studies. *Annals of Internal Medicine* 2014;161(2):96-103.

- Uhl K, Peters JR. Burden of changes in generic pill appearance. [Letter to the Editor] *Annals of Internal Medicine* 2014;161(11):839.
  - Pauker S. Burden of changes in generic pill appearance. [Letter to the Editor] *Annals of Internal Medicine* 2014;161(11):839.
  - **Kesselheim AS**, Choudhry NK, Avorn J. Burden of changes in generic pill appearance. [Author reply] *Annals of Internal Medicine* 2014;161(11):840.
58. Gagne JJ, Choudhry NK, **Kesselheim AS**, Polinski JM, Hutchins D, Matlin OS, Brennan TA, Avorn J, Shrank WH. Comparative effectiveness of generic and brand-name statins on patient outcomes. *Annals of Internal Medicine* 2014;161:400-407.
  59. Checchi KD\*\*, Huybrechts KF, Avorn J, **Kesselheim AS**. Electronic medication packaging devices and medication adherence: A systematic review. *JAMA* 2014;312(12):1237-1247.
  60. **Kesselheim AS**, Tan YT, Darrow JJ, Avorn J. Existing FDA pathways have potential to ensure early access to, and appropriate use of, specialty drugs. *Health Affairs* 2014;33(10):1770-1778.
  61. Yeh JS\*\*, Austad KE, Franklin JM, Chimonas S, Campbell EG, Avorn J, **Kesselheim AS**. Association of medical students' reports of interactions with the pharmaceutical and medical device industries and medical school policies and characteristics: a cross-sectional study. *PLoS Medicine* 2014;11(10):e1001743.
  62. Gagne JJ, Thompson L, O'Keefe K, **Kesselheim AS**. Innovative research methods for studying treatments for rare diseases: methodological review. *BMJ* 2014;349:g6802.
  63. Xu S\*\* and **Kesselheim AS**. Medical innovation then and now: perspectives of innovators responsible for transformative drugs. *Journal of Law, Medicine & Ethics* 2014;42(4):564-575.
  64. **Kesselheim AS**, Huybrechts KF, Choudhry NK, Fulchino LA, Isaman DL, Kowal MK, Brennan TA. Prescription drug insurance coverage and patient health outcomes: A systematic review. *American Journal of Public Health* 2015;105(2):e17-30.
  65. Rajan PV\*\*, Kramer DB\*\*, **Kesselheim AS**. Medical device postapproval safety monitoring: where does the United States stand? *Circulation Cardiovascular Quality & Outcomes* 2015;8:124-131.
  66. **Kesselheim AS**, McGraw S, Thompson L, O'Keefe K, Gagne JJ. Development and use of new therapeutics for rare diseases: views from patients, caregivers, and advocates. *The Patient: Patient-Centered Outcomes Research* 2015;8(1):75-84.
  67. Polinski JM, **Kesselheim AS**, Seeger JD, Connolly JG, Choudhry NK, Shrank WH. A cross-national comparison of 17 countries' insulin glargine drug labels. *Pharmacoepidemiology and Drug Safety* 2015;24(2):159-165.
  68. **Kesselheim AS**, Tan YT, Avorn J. The roles of academia, rare diseases, and repurposing in the development of the most transformative drugs. *Health Affairs* 2015;34:286-294.
  69. Wang B\*\*, Liu J, **Kesselheim AS**. Variations in time of market exclusivity among top-selling prescription drugs in the United States. *JAMA Internal Medicine* 2015;175(4):635-637.
    - Downing NS. Market exclusivity for top-selling pharmaceuticals: too long, too short, or just right? [Editorial] *JAMA Internal Medicine* 2015;175(4):637-638.
  70. Sarpatwari A\*\*, Franklin JM, Avorn J, Seeger JD, Landon JE, **Kesselheim AS**. Are Risk Evaluation and Mitigation Strategies associated with less off-label use of medications? The case of ITP. *Clinical Pharmacology and Therapeutics* 2015;97(2):186-93.
  71. **Kesselheim AS**, Connolly JC, Rogers J, Avorn J. Despite mandatory disclaimers on dietary supplements, many consumers remain unaware or overlook the information. *Health Affairs* 2015;34(3):438-446.
  72. Wang B\*\*, Choudhry NK, Gagne JJ, Landon J, **Kesselheim AS**. Availability and utilization of cardiovascular fixed-dose combination drugs in the United States. *American Heart Journal* 2015;169(3):379-386.

73. **Kesselheim AS**, Polinski JP, Fulchino LA, Isaman DL, Gagne JJ. Modified regulatory pathways to approve generic drugs in the US and a systematic review of their outcomes. *Drugs* 2015;75(6):633-650.
74. **Kesselheim AS**, Campbell EG, Schneeweiss S, Rausch P, Lappin BM, Zhou EH, Seeger JD, Brownstein JS, Woloshin S, Schwartz LM, Toomey T, Dal Pan G, Avorn J. Methodological approaches to evaluate the impact of FDA Drug Safety Communications. *Drug Safety* 2015;38(6):565-575.
75. Darrow JJ\*\*, **Kesselheim AS**. A new wave of vaccines for non-communicable diseases: what are the regulatory challenges? *Food and Drug Law Journal* 2015;70(2):243-258.
76. **Kesselheim AS**, Franklin JM, Kim SC, Seeger JD, Solomon DH. Reductions in use of colchicine after FDA enforcement of market exclusivity in a commercially insured population. *J Gen Intern Med* 2015;30(11):1633-1638.
77. Luo J\*\*, Avorn J, **Kesselheim AS**. Trends in Medicaid reimbursements for insulin from 1991 through 2014. *JAMA Internal Medicine* 2015;175(10):1681-1686.
- Bryant J. Inaccurate Reporting of Insulin Reimbursement. [Letter to the Editor] *JAMA Internal Medicine* 2016;176(3):408.
  - Alatorre CI, Schultz EH. Inaccurate Reporting of Insulin Reimbursement. [Letter to the Editor] *JAMA Internal Medicine* 2016;176(3):407-408.
  - Luo J, Avorn J, **Kesselheim AS**. Inaccurate reporting of insulin reimbursement. [Author reply] *JAMA Internal Medicine* 2016;176(3):408-409.\*\*
78. Gagne JJ, **Kesselheim AS**, Choudhry NK, Polinski JM, Hutchins D, Matlin OS, Brennan TA, Avorn J, Shrank WH. Comparative effectiveness of generic versus brand-name antiepileptic medications. *Epilepsy & Behavior* 2015;52(Pt A):14-18.
79. **Kesselheim AS**, Wang B\*\*, Franklin JM, Darrow JJ\*\*. Trends in utilization of FDA expedited drug development and approval programs, 1987-2014: cohort study. *BMJ* 2015;351:h4633.
- Frakt A. JAMA Forum: The risks and benefits of expedited drug reviews. May 23, 2018. Available from: <https://newsatjama.jama.com/2018/05/23/jama-forum-the-risks-and-benefits-of-expedited-drug-reviews/>.
80. Wang B\*\*, **Kesselheim AS**. Characteristics of efficacy evidence supporting approval of supplemental indications for prescription drugs in the United States, 2005-2014: systematic review. *BMJ* 2015;351:h4679.
81. Treasure CL\*\*, Avorn J, **Kesselheim AS**. Do march-in rights ensure access to medical products arising from federally funded research? A qualitative study. *Milbank Quarterly* 2015;93(4):761-787.
82. Luo J\*\*, **Kesselheim AS**. The evolution of insulin patents and market exclusivities in the USA, 2004-2014. *The Lancet Diabetes & Endocrinology* 2015;3(11):835-837.
- Kaplan W, Laing R, Ewen M, Beran D. Insulin patents and market exclusivities: unresolved issues. [Letter to the Editor] *The Lancet Diabetes & Endocrinology* 2016;4(2):98.
  - Luo J, **Kesselheim AS**. Insulin patents and market exclusivities: unresolved issues. [Author reply] *The Lancet Diabetes & Endocrinology* 2016;4(2):98-99.\*\*
83. **Kesselheim AS**, Hwang TJ\*\*, Franklin JM. Two decades of new drug development for central nervous system disorders. *Nature Reviews: Drug Discovery* 2015;14(12):815-816.
84. Yeh JS\*\*, Austad KE, Franklin JM, Chimonas S, Campbell EG, Avorn J, **Kesselheim AS**. Medical schools' industry interaction policies are not associated with trainees' self-reported behavior as residents: results of a national survey. *Journal of Graduate Medical Education* 2015;7(4):595-602.

85. Kim SC, Choi NK, Lee J, Kwon KE, Eddings W, Sung YK, Song HJ, **Kesselheim AS**, Solomon DH. Uptake of the first biosimilar infliximab since its approval in South Korea. *Arthritis and Rheumatology* 2016;68(5):1076-9.
86. Hwang TJ\*\*, **Kesselheim AS**. Vaccine pipeline has grown during the past two decades with more early-stage trials from small and medium-size companies. *Health Affairs* 2016;35(2):219-26.
87. **Kesselheim AS**, Gagne JJ, Franklin JM, Eddings W, Fulchino LA, Avorn J, Campbell EG. Variations in patients' perceptions and use of generic drugs: results of a national survey. *Journal of General Internal Medicine* 2016;31(6):609-14.
- Kaplan CM. Capsule Commentary on Kesselheim et al., Variations in Patients' Perceptions and Use of Generic Drugs: Results of a National Survey. [Review] *Journal of General Internal Medicine* 2016;31(6):647.
88. **Kesselheim AS**, Woloshin S, Eddings W, Franklin JM, Ross KM, Schwartz LM. Physicians' knowledge about FDA approval standards and perceptions of the Breakthrough Therapy designation. *JAMA* 2016;315(14):1516-1518.
89. Wang B\*\*, Franklin JM, Eddings W, Landon J, **Kesselheim AS**. Did FDA decisionmaking affect anti-psychotic drug prescribing in children?: A time-trend analysis. *PLoS One* 2016;11(3):e0152195.
90. Gagne JJ, Polinski JM, Jiang W, Dutcher SK, Xie J, Lii J, Fulchino LA, **Kesselheim AS**. Switch-backs associated with generic drugs approved using product-specific determinations of therapeutic equivalence. *Pharmacoepidemiology and Drug Safety* 2016;25(8):944-952.
91. **Kesselheim AS**, Gagne JJ, Eddings W, Franklin JM, Ross KM, Fulchino LA, Campbell EG. Prevalence and predictors of generic drug skepticism among physicians: results of a national survey. *JAMA Internal Medicine* 2016;176(6):845-847.
- Chee M, Ngooi S, Arora VM. Prescription Trends—Brand-name Drugs vs Generic. [Letter to the Editor] *JAMA Internal Medicine* 2016;176(10):1573-1574.
  - **Kesselheim AS**, Gagne JJ. Prescription trends—brand-name drugs vs. generic. [Author reply] *JAMA Internal Medicine* 2016;176(10):1574-1575.
92. Yeh JS\*\*, Franklin JM, Avorn J, Landon J, **Kesselheim AS**. Association of industry payments to physicians with the prescribing of brand-name statins in Massachusetts. *JAMA Internal Medicine*. 2016;176(6):763-768.
- Fagan TC. Payments to physicians, prescribing rates, and more appropriate conclusions. [Letter to the Editor] *JAMA Intern Med*. 2016;176(10):1576.
  - Feuerstein JD. Payments to Physicians, Prescribing Rates, and more appropriate conclusions. [Letter to the Editor] *JAMA Intern Med*. 2016;176(10):1576-1577.
  - Yeh JS\*\*, Franklin JM, **Kesselheim AS**. Payments to physicians, prescribing rates, and more appropriate conclusions. [Author reply] *JAMA Internal Medicine* 2016;176(10):1577.
93. Caplan ES\*\*, **Kesselheim AS**. Anti-VEGF therapy in ophthalmology—a qualitative analysis of transformative drug development. *Drug Discovery Today* 2016;21(6):1019-1026.
94. Deak DM\*\*, Powers JP, Outterson MK, **Kesselheim AS**. Progress in the fight against multidrug-resistant bacteria?: a review of U.S. Food and Drug Administration-approved antibiotics 2010-2015. *Annals of Internal Medicine* 2016;165(5):363-372.
- Weinrauch LA, D'Elia JA. The fight against multidrug-resistant bacteria. [Letter to the Editor] *Annals of Internal Medicine* 2017;166(1):77-78.
  - Boucher HW, Cosgrove SE, Cox E, Talbot GH. The fight against multi-drug resistant bacteria. [Letter to the Editor] *Annals of Internal Medicine* 2017;166(1):78-79.
  - Deak D\*\*, Powers JH, Outterson K, **Kesselheim AS**. The fight against multidrug-resistant bacteria. [Author reply] *Annals of Internal Medicine* 2017;166(1):79.

95. Hwang TJ\*\*, Sokolov E, Franklin JM, **Kesselheim AS**. Comparison of rates of safety issues and trial outcomes reporting for medical devices approved in EU and US: cohort study. *BMJ* 2016;353:i3323.\*\*
96. Sarpatwari A\*\*, **Kesselheim AS**. Navigating the dermatologic drug cost curve. *JAMA* 2016;315(24):2724-2725.
97. Luo J\*\*, Seeger JD, Donneyong M, Gagne JJ, Avorn J, **Kesselheim AS**. Effect of generic competition on atorvastatin prescribing and patients' out-of-pocket spending. *JAMA Internal Medicine* 2016;176(9):1317-1323.
98. Gupta R, **Kesselheim AS**, Downing N, Greene J, Ross JS. Generic drug approvals since the 1984 Hatch-Waxman Act. *JAMA Internal Medicine* 2016;176(9):1391-1393.
  - Toufanian M, Peters JR, Uhl K. Prioritization of generic drug review. [Letter to the Editor] *JAMA Internal Medicine* 2017;177(1):140-141.
  - Gupta R, **Kesselheim AS**, Ross JS. Prioritization of generic drug review. [Author reply] *JAMA Internal Medicine* 2017;177(1):141-142.
99. Hwang TJ\*\*, Lauffenburger JC, Franklin JM, **Kesselheim AS**. Temporal trends and factors associated with cardiovascular drug development, 1990 to 2012. *Journal of the American College of Cardiology: Basic to Translational Science* 2016;1(5):301-308.
  - Fiuzat M, Stockbridge N, Califf RM. Resourcing drug development commensurate with its public health importance. [Editorial] *JACC: Basic to Translational Science* 2016;1(5):309-312.
100. **Kesselheim AS**, Avorn J, Sarpatwari A\*\*. The high cost of prescription drugs in the United States: origins and prospects for reform. *JAMA* 2016;316(8):858-871.
  - Roy V, Hawksbee L, King L. Factors influencing prescription drug costs in the United States. [Letter to the Editor] *JAMA* 2016;316(22):2431.
  - Arbiser JL. Factors influencing prescription drug costs in the United States. [Letter to the Editor] *JAMA* 2016;316(22):2430-2431.
  - Sarpatwari A\*\*, Avorn J, **Kesselheim AS**. Factors influencing prescription drug costs in the United States. [Author reply] *JAMA* 2016;316(22):2431-2432.
  - #41 most-discussed journal article of 2016. Altmetric. Available online at: <https://www.altmetric.com/details/10782162>.
101. **Kesselheim AS**, Bykov K, Gagne JJ, Wang S, Choudhry NK. Switching generic antiepileptic drug manufacturer not linked to seizures: a case-crossover study. *Neurology* 2016;87(17):1796-1801.
  - Krauss GL, Privitera M. More data on the safety of generic substitution: yes, the blue tablet is OK? [Editorial] *Neurology*. 2016;87(17):1754-1755.
102. Hwang TJ\*\*, Carpenter D, Lauffenburger JC, Wang B\*\*, Franklin JM, **Kesselheim AS**. Failure of investigational therapeutics in late-stage clinical development and publication of trial results. *JAMA Internal Medicine* 2016;176(12):1826-1833.
103. **Kesselheim AS**, Eddings W, Raj T, Campbell EG, Franklin JM, Ross KM, Fulchino LA, Avorn J, Gagne JJ. Physicians' trust in the FDA's use of product-specific pathways for generic drug approval. *PLoS One* 2016;11(10):e0163339.
104. Hwang TJ\*\*, **Kesselheim AS**. Public referendum on drug prices in the US: will it bring relief? *BMJ* 2016;355:i5657.
105. Luo J\*\*, Gagne JJ, Landon J, Avorn J, **Kesselheim AS**. Comparative effectiveness and safety of thalidomide and lenalidomide in patients with multiple myeloma in the United States of America: a population-based cohort study. *European Journal of Cancer* 2016;70:22-33.
106. **Kesselheim AS**, Treasure CL\*\*, Joffe S. Biomarker-defined subsets of common diseases: policy and economic implications of Orphan Drug Act coverage. *PLoS Medicine* 2017;14(1):e1002190.

- Herder M. What is the purpose of the Orphan Drug Act? *PLoS Medicine* 2017;14(1):e1002191.
107. Stern AD, Kramer DB\*\*, Ouellet M, **Kesselheim AS**. Review times and adverse events for cardiovascular devices. *Nature Biomedical Engineering* 2017;1:0013.
  108. **Kesselheim AS**, Rome BN\*\*, Sarpatwari A\*\*, Avorn J. Six-month market exclusivity extensions to promote research offer substantial returns for many drug makers. *Health Affairs* 2017;36(2):362-370.
    - Bronstein M, Kakkis E, Fajgenbaum D, Chambers C. For rare disease patients, a pathway to hundreds of new therapies. [Editorial] *Health Affairs Blog*. March 21, 2017. Available online at: <https://www.healthaffairs.org/doi/10.1377/hblog20170321.059289/full/>
  109. Rathi VK, Wang B, Ross JS, Downing NS, **Kesselheim AS**, Gray ST. Clinical evidence supporting US Food and Drug Administration premarket approval of high-risk otolaryngologic devices, 2000-2014. *Otolaryngology Head & Neck Surgery* 2017;156(2):285-288.
  110. Rathi VK, Wang B, Ross JS, Downing NS, **Kesselheim AS**, Gray ST. Clinical evidence supporting US Food and Drug Administration approval of otolaryngologic prescription drug indications, 2005-2014. *Otolaryngology Head & Neck Surgery* 2017;156(4):683-692.
  111. Gagne JJ, Polinski JM, Jiang W, Dutcher SK, Xie J, Lii J, Fulchino LA, **Kesselheim AS**. Outcome associated with generic drugs approved using product-specific determinations of therapeutic equivalence. *Drugs* 2017;77(4):427-433.
  112. **Kesselheim AS**, McGraw SA, Dejene SZ, Rausch P, Dal Pan GJ, Lappin BM, Zhou EH, Avorn J, Campbell EG. Patient and physician perceptions of drug safety information for sleep aids: a qualitative study. *Drug Safety* 2017;40(6):531-542.
  113. Woloshin S, Schwartz LM, Dejene SZ, Rausch P, Dal Pan GJ, **Kesselheim AS**. Media coverage of FDA Drug Safety Communications about zolpidem: a quantitative and qualitative analysis. *Journal of Health Communication* 2017;22(5):365-372.
  114. Luo J\*\*, **Kesselheim AS**, Avorn J. Medicaid expenditures and estimated rebates for epinephrine autoinjectors, 2012 to 2016. *JAMA Internal Medicine*. 2017;177(5):734-736.
  115. **Kesselheim AS**, Gagne JJ, Franklin JM, Eddings W, Fulchino LA, Campbell EG. Do patients trust the FDA?: A survey assessing how patients view the generic drug approval process. *Pharmacoepidemiology and Drug Safety* 2017;26(6):694-701.
  116. Wang B\*\*, Studdert DM, Sarpatwari A, Franklin JM, Landon J, **Kesselheim AS**. The effect of federal and state off-label marketing investigations on drug prescribing: the case of olanzapine. *PLoS One* 2017;12(4):e0175313.
  117. Hwang TJ\*\*, **Kesselheim AS**. Effect of US Food and Drug Administration's cardiovascular safety guidance on diabetes drug development. *Clinical Pharmacology and Therapeutics* 2017;102(2):290-296.
  118. Davies BJ, Hwang TJ\*\*, **Kesselheim AS**. Ensuring access to injectable generic drugs—the case of intravesical BCG for bladder cancer. *New England Journal of Medicine* 2017;376(15):1401-1403.
  119. Sacks CA\*\*, Avorn J, **Kesselheim AS**. The failure of solanezumab—how the FDA saved taxpayers billions. *New England Journal of Medicine* 2017;376(18):1706-1708.
  120. Bollyky TJ, **Kesselheim AS**. Can drug importation address high generic drug prices? Brookings Institution White Paper. May 2 2017. Available at: <https://www.brookings.edu/research/can-drug-importation-address-high-generic-drug-prices/>.
    - Khan J. Elizabeth Warren on drugs. *National Review* January 4, 2019. Available from: <https://www.nationalreview.com/2019/01/elizabeth-warren-generic-drugs-proposal-not-serious/>

121. Dave C\*\*, **Kesselheim AS**, Fox E, Hartzema AG. High generic drug prices and market competition levels: a retrospective cohort study. *Annals of Internal Medicine* 2017;167(3):145-151.
122. Fralick M\*\*, Avorn J, **Kesselheim AS**. The price of crossing the border for medications. *New England Journal of Medicine* 2017;377(4):311-313.
  - Ashley DD. The price of crossing the border for medications. [Letter to the Editor] *New England Journal of Medicine* 2017;377(17):1699.
  - Fralick M\*\*, Avorn J, **Kesselheim AS**. The price of crossing the border for medications. [Author reply] *New England Journal of Medicine* 2017;377(17):1699-1700.
123. Jain N\*\*, Hwang TJ\*\*, Franklin JM, **Kesselheim AS**. Association of the priority review voucher with neglected tropical disease drug and vaccine development. *JAMA* 2017;318(4):388-389.
124. Naci H\*\*, Smalley KR, **Kesselheim AS**. Characteristics of preapproval and postapproval studies for drugs granted accelerated approval by the US Food and Drug Administration. *JAMA* 2017;318(7):626-636.
  - Califf RM. Balancing the need for access with the imperative for empirical evidence of benefit and risk. [Editorial] *JAMA* 2017;318(7):614-616.
  - Tanimoto T, Kosugi K, Tsuda K. [Letter to the Editor] Evidence required for drugs granted accelerated approval. *JAMA* 2017;318(24):2492-2493.
  - Naci H\*\*, **Kesselheim AS**. Evidence required for drugs granted accelerated approval [Author Reply] *JAMA* 2017;318(24):2493-2494.
125. Mostaghim S\*\*, Gagne JJ, **Kesselheim AS**. Safety-related label changes for new drugs after approval in the US through expedited regulatory pathways: retrospective cohort study. *BMJ* 2017;358:j3837.
126. **Kesselheim AS**, Donneyong M, Dal Pan GJ, Zhou EH, Avorn J, Schneeweiss S, Seeger JD. Changes in prescribing and healthcare resource utilization after FDA Drug Safety Communications involving zolpidem-containing medications. *Pharmacoepidemiology and Drug Safety* 2017;26(6):712-721.
127. Alpern JD, Zhang L, Stauffer WM, **Kesselheim AS**. Trends in pricing and generic competition within the oral antibiotic drug market in the United States. *Clinical Infectious Diseases* 2017;65(11):1848-1852.
128. Hwang TJ\*\*, Darrow JJ\*\*, **Kesselheim AS**. The FDA's expedited pathways and clinical development times for novel therapeutics, 2012-2016. *JAMA* 2017;318(21):2137-2138.
129. Hey SP\*\*, Franklin JM, Avorn J, **Kesselheim AS**. Success, failure, and transparency in biomarker-based drug development: a case study of Cholesteryl Ester Transfer Protein inhibitors. *Circulation Cardiovasc Quality and Outcomes*. 2017;10(6):e003121.
130. Darrow JJ\*\*, Avorn J, **Kesselheim AS**. Speed, safety, and industry funding — from PDUFA I to PDUFA VI. *New England Journal of Medicine* 2017;377(23):2278-2286.
131. Dave CV\*\*, Hartzema A, **Kesselheim AS**. Prices of generic drugs associated with numbers of manufacturers. *New England Journal of Medicine* 2017;377(26):2597-2598.
132. Hwang TJ\*\*, Sachs RE, **Kesselheim AS**. Public participation in drafting of the 21st Century Cures Act. *Journal of Law, Medicine and Ethics* 2017;45(2):212-220.
133. Sinha MS\*\*, Freifeld CC, Brownstein JS, Donneyong MM, Rausch P, Lappin BM, Zhou EH, Dal Pan GJ, Pawar AM, Hwang TJ\*\*, Avorn J, **Kesselheim AS**. Social media impact of the Food and Drug Administration's drug safety communication messaging about zolpidem: mixed-methods analysis. *JMIR: Public Health Surveillance* 2018;4(1):e1.
134. Fralick M, **Kesselheim AS**, Avorn J, Schneeweiss S. Use of health care databases to support supplemental indications of approved medications. *JAMA Internal Medicine* 2018;178(1):55-63.

- Califf RM. Comparison of observational data and the ONTARGET results for telmisartan treatment of hypertension: bull’s-eye or painting the target around the arrow? [Editorial] *JAMA Internal Medicine* 2018;178(1):63-65.
135. Beall RF\*\*, **Kesselheim AS**. Tertiary patenting on drug-device combination products in the United States. *Nature Biotechnology* 2018;36(2):142-145.
    - Brennan Z. Increase in combo product patents delay competition further, researchers find. *Regulatory Focus*. April 3, 2018. Available from: <https://www.raps.org/news-and-articles/news-articles/2018/4/increase-in-combo-product-patents-delay-competitio>
  136. Bothwell LE\*\*, Avorn J, Khan N, **Kesselheim AS**. Adaptive design clinical trials: a review of the literature and *ClinicalTrials.gov*. *BMJ Open* 2018;8(2):e018320.
  137. Fralick M\*\*, Avorn J, Franklin JM, Abdurrob A, **Kesselheim AS**. Application and impact of run-in studies. *Journal of General Internal Medicine* 2018;33(5):759-763.
  138. Fralick M\*\*, Avorn J, Franklin JM, Bartsch E, Abdurrob A, **Kesselheim AS**. Application and impact of run-in studies for the evaluation of statin efficacy and safety. *Journal of General Internal Medicine* 2018;33(6):792-794.
  139. Gupta R, Bollyky T, Cohen M, Ross JS, **Kesselheim AS**. Availability of US off-patent drugs at risk for acute price increases or shortages from manufacturers approved by non-US regulators. *BMJ* 2018;360:k831.
    - Gupta R, **Kesselheim AS**. Importing drugs—a solution to high priced generics in the US. [Blog] *The BMJ Opinion*. 2018 Mar 19. Available from: <http://blogs.bmj.com/bmj/2018/03/19/ravi-gupta-and-aaron-kesselheim-importing-drugs-a-solution-to-high-priced-generics-in-the-us/>
  140. Desai RJ, Sarpatwari A\*\*, Dejene S, Khan NF, Lii J, Rogers JR, Dutcher SK, Raofi S, Bohn J, Connolly J, Fischer MA, **Kesselheim AS**, Gagne JJ. Differences in rates of switchbacks after switching from branded to authorized generic and branded to generic drug products: Cohort study. *BMJ*. 2018;361:k1180.
  141. Jung E\*\*, Zettler PJ, **Kesselheim AS**. Prevalence of publicly available expanded access policies. *Clinical Pharmacology and Therapeutics* 2018;104(5):1016-1021.
  142. Darrow JJ\*\*, Avorn J, **Kesselheim AS**. The FDA’s “Breakthrough Drug” designations: four years of experience. *New England Journal of Medicine* 2018;378(15):1444-1453.
    - Corrigan-Curay J, McKee AE, Stein P. Breakthrough-therapy designation—an FDA perspective. *New England Journal of Medicine* 2018;378(15):1457-1458.
    - Allen J. The breakthrough therapy designation for promising cancer drugs is good for patients. *STAT: First Opinion*. April 27, 2018. Available online at: <https://www.statnews.com/2018/04/27/breakthrough-therapy-designation-helps-cancer-patients/>
  143. Hwang TJ\*\*, Franklin JM, Chen CT, Lauffenburger JC, Gyawali B\*\*, **Kesselheim AS**, Darrow JJ\*\*. Efficacy, safety, and regulatory approval of Food and Drug Administration-designated breakthrough and nonbreakthrough cancer medicines. *Journal of Clinical Oncology* 2018;36(18):1805-1812
    - Kuderer NM, Lyman GH. Evolving landscape of US Food and Drug Administration Drug approval in the era of precision oncology: finding the right balance between access and safety. *Journal of Clinical Oncology* 2018 May 9;JCO2018785592.
    - Rawson NSB. New cancer breakthrough therapies at the United States Food and Drug Administration. *Journal of Hospital Management Health Policy* 2018;2:43.
  144. Sarpatwari A\*\*, Beall R\*\*, Abdurrob A, He M, **Kesselheim AS**. Evaluating the impact of the Orphan Drug Act’s seven-year market exclusivity period. *Health Affairs* 2018;37(5):732-737.



145. Sarpatwari A\*\*, Lee M, Gagne JJ, Lu Z, Dutcher SK, Jiang W, Campbell EG, **Kesselheim AS**. Generic versions of narrow therapeutic index drugs: a national survey of pharmacists' substitution beliefs and practices. *Clinical Pharmacology and Therapeutics* 2018;103(6):1093-1099.
146. Pregelj L, Hwang TJ\*\*, Siegel E, Barnard R, Hine D, Darrow JJ\*\*, **Kesselheim AS**. Precision medicines have faster approvals based on fewer and smaller trials than other medicines. *Health Affairs* 2018;37(5):724-731.
147. Mundkur ML\*\*, Franklin J, Huybrechts KF, Fischer MA, **Kesselheim AS**, Linder JA, Landon J, Patorno E. Changes in outpatient use of antibiotics by adults in the United States, 2006-2015. *Drug Safety* 2018 Jul 9. doi: 10.1007/s40264-018-0697-4.
148. Beall RF\*\*, Darrow JJ\*\*, **Kesselheim AS**. Patent term restoration for top-selling drugs in the USA. *Drug Discovery Today* 2018 Jul 25. pii: S1359-6446(18)30141-7.
149. Gyawali B\*\*, Hey SP\*\*, **Kesselheim AS**. A comparison of response patterns for progression-free survival and overall survival following treatment for cancer with PD-1 inhibitors: a meta-analysis of correlation and differences in effect sizes. *JAMA Network Open* 2018;1(2):e180416.
150. Rogers JR, Sarpatwari A\*\*, Desai RJ, Bohn JM, Khan NF, **Kesselheim AS**, Fischer MA, Gagne JJ, Connolly JG. Effect of lawyer-submitted reports on signals of disproportional reporting in the Food and Drug Administration's Adverse Event Reporting System. *Drug Safety* 2018 Aug 1.
151. Sinha MS\*\*, Jain N\*\*, Hwang T\*\*, **Kesselheim AS**. Expansion of the Priority Review Voucher program under the 21st Century Cures Act: implications for innovation and public health. *American Journal of Law and Medicine* 2018;44:329-341.
152. Sacks CA\*\*, Lee CC, **Kesselheim AS**, Avorn J. Medicare spending on brand-name combination medications vs their generic constituents. *JAMA* 2018;320(7):650-656.
153. Hernandez I, Sampathkumar S, Good CB, **Kesselheim AS**, Shrank WH. Changes in drug pricing after drug shortages in the United States. *Annals of Internal Medicine* 2018 Sep 18. doi: 10.7326/M18-1137.
154. Sinha MS\*\*, Najafzadeh M, Rajasingh EK, Love J, **Kesselheim AS**. Labeling changes and costs for clinical trials performed under the US Food and Drug Administration pediatric exclusivity extension, 2007 to 2012. *JAMA Internal Medicine* 2018;178(11):1458-1466.
  - Ross JS. Clinical trials—we get what we pay for. *JAMA Internal Medicine* 2018;178(11):1457.
155. Feldman WB\*\*, Hey SP\*\*, **Kesselheim AS**. A systematic review of the Food and Drug Administration's 'exception from informed consent' pathway. *Health Affairs* 2018;37(10):1605-1614.
156. **Kesselheim AS**, Woloshin S, Lu Z, Tessema FA, Ross KM, Schwartz LM. Internal medicine physicians' financial relationships with industry: an updated national estimate. *Journal of General Internal Medicine* 2018 Oct 5 doi: 10.1007/s11606-018-4688-z.
157. Sinha MS\*\*, **Kesselheim AS**. The Tax Cuts and Jobs Act of 2017 and the pharmaceutical industry. *Journal of Law, Medicine, and Ethics* 2018;46(3):806-808.
158. Hwang TJ\*\*, Orenstein L, **Kesselheim AS**, Bourgeois FT. Completion rate and reporting of mandatory pediatric postmarketing studies under the US Pediatric Research Equity Act. *JAMA Pediatrics* 2018 Nov 19. [Epub ahead of print]
  - Cohen-Wolkowicz M, Benjamin DK Jr. Development of therapeutics for children—a tricky balancing act. *JAMA Pediatrics* 2018 Nov 19. [Epub ahead of print]
159. Dave CV\*\*, Pawar A, Fox ER, Brill G, **Kesselheim AS**. Predictors of drug shortages and association with generic drug prices: a retrospective cohort study. *Value in Health* 2018;21(11):1286-1290.

160. Gyawali B\*\*, **Kesselheim AS**, D'Andrea E\*\*. Does *Helicobacter pylori* eradication therapy to prevent gastric cancer increase all-cause mortality? *International Journal of Cancer*. 2019;144(2):411-412.
161. Hwang TJ\*\*, Jain N, Lauffenburger JC, Vokinger KN\*\*, **Kesselheim AS**. Analysis of proposed Medicare Part B to Part D shift with associated changes in total spending and patient cost-sharing for prescription drugs. *JAMA Intern Med*. 2019 Jan 14. doi: 10.1001/jamainternmed.2018.6417.
162. **Kesselheim AS**, Woloshin S, Lu Z, Tessema FA, Ross KM, Schwartz LM. Physicians' perspectives on FDA approval standards and off-label drug marketing. *JAMA Internal Medicine* 2019 Jan 22. doi: 10.1001/jamainternmed.2018.8121.
163. Hwang TJ\*\*, **Kesselheim AS**, Gyawali B\*\*. Affordability and price increases of new cancer drugs in clinical guidelines, 2007-2016. *JNCI Cancer Spectrum* 2019 [in press]
164. Darrow JJ\*\*, Beall RF\*\*, **Kesselheim AS**. The generic drug industry embraces a faster, cheaper pathway for challenging patents. *Applied Health Economics and Health Policy* 2019 [in press].
165. Bourgeois FT, Graham DA, **Kesselheim AS**, Randolph, AG. Cost implications of escalating intravenous acetaminophen use in children. *JAMA Pediatrics* 2019 [in press]
166. Sarpatwari A\*\*, Gagne JJ, Lu Z, Campbell EG, Carman WJ, Enger CL, Dutcher SK, Jiang W, **Kesselheim AS**. A survey of patients' perceptions of pill appearance and responses to changes in appearance for four chronic disease medications. *Journal of General Internal Medicine* 2019 Jan 10. doi: 10.1007/s11606-018-4791-1. [Epub ahead of print]
167. Luo J, Khan NF, Manetti T, Rose J, Kaloghlian A, Gadhe B, Jain SH, Gagne JJ, **Kesselheim AS**. Association between implementation of a health plan program for switching from analog to human insulin with glycemic control among Medicare beneficiaries with type 2 diabetes. *JAMA* 2019 [in press]
168. Dave C\*\*, Brill G, **Kesselheim AS**. Changes in price for generic drugs in the United States, 2008-2016. *Journal of General Internal Medicine* 2019 [in press]

#### Other peer-reviewed publications

1. Kissick WL and **Kesselheim AS**. Presidential disability: the panel of physicians. *Trans & Stud Coll Phys Phila* 2001;23:113-119.
2. **Kesselheim AS**. Perspectives: ode to a short white coat. *Pharos* 2003;66:35.
3. **Kesselheim AS** and Avorn J. University-based science and biotechnology research: defining the boundaries of intellectual property. *JAMA* 2005;293:850-854.
  - Sobolski GK. Biotechnology products and university-based science. [Letter to the Editor] *JAMA* 2005;293(23):2862.
  - Korn D, Heinig SJ. Biotechnology products and university-based science. [Letter to the Editor] *JAMA* 2005;293(23):2862-3.
  - Lempert P. Biotechnology products and university-based science. [Letter to the Editor] *JAMA* 2005;293(23):2861-2
  - **Kesselheim AS**, Avorn J. Biotechnology products and university-based science. [Author reply] *JAMA* 2005;293:2863.
4. **Kesselheim AS** and Brennan TA. The swinging pendulum: the Supreme Court reverses course on ERISA and managed care. *Yale Journal of Health Policy, Law, and Ethics* 2005;5:451-463.
5. **Kesselheim AS** and Brennan TA. Overbilling vs. downcoding — the battle between physicians and insurers. *New England Journal of Medicine* 2005;352:855-857.
6. **Kesselheim AS** and Avorn J. Biomedical patents and the public's health: is there a role for eminent domain? *JAMA* 2006;295:434-437.
7. **Kesselheim AS**, Ferris TG, Studdert DM. Will physician-level measures of clinical performance be used in medical malpractice litigation? *JAMA* 2006;295:1831-1834.

- Luft HS. Clinical performance measures and medical malpractice. [Letter to the editor] JAMA 2006;296(13):1589.
  - **Kesselheim AS**, Studdert DM. Clinical performance measures and medical malpractice. [Author reply] JAMA 2006;296:1589-90.
8. **Kesselheim AS**, Fischer MA, Avorn J. The rise and fall of Natrecor for congestive heart failure: implications for drug policy. Health Affairs 2006;25:1095-1102.
  9. **Kesselheim AS** and Mello MM. Medical process patents – monopolizing the delivery of health care. New England Journal of Medicine 2006;355:2036-2041.
  10. Reese PP, Caplan AL, **Kesselheim AS**, Bloom RD. Creating a medical, ethical, and legal framework for complex living kidney donors. Clinical Journal of the American Society of Nephrology 2006;1:1148-1153.
  11. **Kesselheim AS** and Mello MM. Confidentiality laws and secrecy in medical research: improving access to drug safety data. Health Affairs 2007;26:483-491.
  12. **Kesselheim AS**. Intellectual property policy in the pharmaceutical sciences: the effect of inappropriate patents and market exclusivity extensions on the health care system. AAPS Journal 2007;9:Article 33.
  13. **Kesselheim AS** and Studdert DM. Role of medical professional organizations in regulating physician expert witness testimony. JAMA 2007;298:2907-2909.
    - Larriviere D, Williams MA, Sagsveen M. Regulating physician expert witness testimony. [Letter to the editor] JAMA 2008;299(14):1667.
    - Hartz AJ, Green M. Regulating physician expert witness testimony. [Letter to the editor] JAMA 2008;299(14):1667-1668.
    - **Kesselheim AS**, Studdert DM. Role of medical professional organizations in regulating physician expert witness testimony. [Author reply] JAMA 2008;299:1667-1668.
  14. Dudzinski DM and **Kesselheim AS**. Scientific and legal viability of follow-on protein drugs. New England Journal of Medicine 2008;358:843-849.
  15. Chokshi DA\*\* and **Kesselheim AS**. Rethinking global access to vaccines. BMJ 2008;336:750-753.
  16. **Kesselheim AS** and Choudhry NK. The international pharmaceutical market as a source of low-cost prescription drugs for U.S. patients. Annals of Internal Medicine 2008;148:614-619.
  17. **Kesselheim AS** and Avorn J. Pharmaceutical promotion to physicians and First Amendment rights. New England Journal of Medicine 2008;358:1727-1732.
    - Troy DE, Gottlieb S. Pharmaceutical promotion and First Amendment rights. [Letter to the Editor] New England Journal of Medicine 2008;359(5):536.
    - **Kesselheim AS**, Avorn J. Pharmaceutical promotion and First Amendment rights. [Author reply] New England Journal of Medicine 2008;359:536-537.
  18. **Kesselheim AS**. Think globally, prescribe locally: how rational pharmaceutical policy in the U.S. can improve global access to essential medicines. American Journal of Law and Medicine 2008;34:125-139.
  19. **Kesselheim AS**. Encouraging drug development for neglected diseases — the trouble with FDA review vouchers. New England Journal of Medicine 2008;359:1981-1983.
    - Moe J, Grabowski H, Ridley D. FDA review vouchers. [Letter to the Editor] New England Journal of Medicine 2009;360(8):837.
    - **Kesselheim AS**. FDA review vouchers. [Author reply] New England Journal of Medicine 2009;360:837-838.
  20. **Kesselheim AS** and Avorn J. A national essay contest on the intersection between pharmaceutical marketing and medical student education. Academic Medicine 2009;84:228-235.

21. **Kesselheim AS**. Priority review vouchers: an inefficient and dangerous way of promoting neglected disease drug development. *Clinical Pharmacology and Therapeutics* 2009;85:373-375.
22. **Kesselheim AS** and Studdert DM. The Supreme Court, preemption, and malpractice liability. *New England Journal of Medicine* 2009;360:559-561.
  - Hauser RG. Preemption and malpractice liability. [Letter to the Editor] *New England Journal of Medicine* 2009;360(21):2257-2258.
  - **Kesselheim AS**, Studdert DM. Preemption and malpractice liability. [Author reply] *New England Journal of Medicine* 2009;360:2257-2258.
23. Outterson MK and **Kesselheim AS**. Evaluating Medicare Part D drug price reform options. *Health Affairs* 2009;28(5):w832-w841.
24. Amin T, Rajkumar R\*\*, Radhakrishan P, **Kesselheim AS**. Expert participation as a strategy for improving the quality of pharmaceutical patents. *Health Affairs* 2009;28(6):w948-w956.
25. Engelberg AB, **Kesselheim AS**, Avorn J. Balancing innovation, access, and profits — market exclusivity for biologics. *New England Journal of Medicine* 2009;361:1917-1919.
  - Pollock A, Zagari M. Market exclusivity for biologics. [Letter to the Editor] *New England Journal of Medicine* 2010;362(7):661.
  - Wheadon DE. Market exclusivity for biologics. [Letter to the Editor] *New England Journal of Medicine* 2010;362(7):661.
  - Engelberg AB, **Kesselheim AS**. Balancing innovation, access, and profits. [Author reply] *New England Journal of Medicine* 2010;362:661-662.
26. **Kesselheim AS**. The Supreme Court, process patents, and medical innovation. *New England Journal of Medicine* 2009;361:2303-2306.
27. **Kesselheim AS** and Maisel WH. Conflict of interest in health care delivery: protecting patients' interests. *American Journal of Therapeutics* 2010;17(4):440-443.
28. **Kesselheim AS** and Mello MM. Gene patenting — is the pendulum swinging back? *New England Journal of Medicine* 2010;362(20):1855-1858.
29. **Kesselheim AS** and Solomon DH. Incentives for drug development: the curious case of colchicine. *New England Journal of Medicine* 2010;362(22):2045-2047.
  - Woodcock J, Okada S. Incentives for drug development—the curious case of colchicine. [Letter to the Editor] *New England Journal of Medicine* 2010;363(15):1484.
  - **Kesselheim AS**, Solomon DH. Incentives for drug development—the curious case of colchicine. [Author reply] *New England Journal of Medicine* 2010;363(15):1484-1485.
30. **Kesselheim AS**. Permitting products-liability litigation for FDA-approved drugs and devices promotes patient safety. *Clinical Pharmacology and Therapeutics* 2010;87(6):645-647.
31. **Kesselheim AS** and Outterson MK. Fighting antibiotic resistance: marrying new financial incentives to meeting public health goals. *Health Affairs* 2010;29(9):1689-1696.
32. Chokshi DA\*\*, Avorn J, **Kesselheim AS**. Designing comparative effectiveness research on prescription drugs: lessons from the clinical trial literature. *Health Affairs* 2010;29(10):1842-1848.
33. **Kesselheim AS**. Using market exclusivity incentives to promote pharmaceutical innovation. *New England Journal of Medicine* 2010;363:1855-1862.
34. Greene JA and **Kesselheim AS**. Pharmaceutical marketing and the new social media. *New England Journal of Medicine* 2010;363:2087-2089.
35. Polinski JM and **Kesselheim AS**. Where cost, medical necessity, and morality meet: should US government insurance programs pay for erectile dysfunction drugs? *Clinical Pharmacology and Therapeutics* 2011;89:17-19.
36. Cutler C, **Kesselheim A**, Gabardi S, Andersson BS, Carpenter P, Khoury HJ, Litzow M, Rowley SD, Lanum S, Leather H, Shih YT, Gale RP, Wingard JR, Appelbaum FR, Anasetti C. Generic immunosuppressants in hematopoietic cell transplantation. *Biol Blood Marrow Transpl*

- 2011;17(3):285-290.
37. **Kesselheim AS** and Outterson MK. Improving antibiotic markets for long-term sustainability. *Yale Journal of Health Policy Law & Ethics* 2011;11:101-167.
  38. **Kesselheim AS** and Austad KE. Residents: workers or students in the eyes of the law? *New England Journal of Medicine* 2011;364:697-699.
  39. **Kesselheim AS**. Covert pharmaceutical promotion in free medical publications. *CMAJ: Canadian Medical Association Journal* 2011;183(5):534-535.
  40. Avorn J and **Kesselheim A**. A hemorrhage of off-label use. *Annals of Internal Medicine* 2011;154:566-567.
    - Karkouti K, Levy JH. A hemorrhage of off-label use. [Letter to the Editor] *Annals of Internal Medicine* 2011;155(5):339.
    - Avorn J and **Kesselheim AS**. The off-label use of recombinant activated Factor VII. [Author reply] *Annals of Internal Medicine* 2011;155(5):339-340.
  41. **Kesselheim A**. Safety, supply, and suits: litigation and the vaccine industry. *New England Journal of Medicine* 2011;364:1485-1487.
  42. Greene JA and **Kesselheim AS**. Why do the same drugs look different? Pills, trade dress, and public health. *New England Journal of Medicine* 2011;365:83-89.
  43. Carpenter D, **Kesselheim AS**, Joffe S. Reputation and precedent in the bevacizumab decision. *New England Journal of Medicine* 2011;365(2):e3.
  44. **Kesselheim AS**. The backlash against bioequivalence and the interchangeability of brand-name and generic drugs. *CMAJ: Canadian Medical Association Journal* 2011;183(12):1350-1351.
  45. **Kesselheim AS**. Off-label drug use and promotion: balancing public health goals and commercial speech. *American Journal of Law and Medicine* 2011;37:225-257.
  46. Austad KE\*\* and **Kesselheim AS**. Conflict of interest disclosure in early medical education: should medical students stay in the dark? *JAMA* 2011;306:991-992.
  47. **Kesselheim AS** and Rajkumar R\*\*. Who owns federally funded research? The Supreme Court and the Bayh-Dole Act. *New England Journal of Medicine* 2011;365(13):1167-1169.
  48. **Kesselheim AS**, Murtagh L, Mello MM. “Pay-for-delay” settlements of disputes over pharmaceutical patents. *New England Journal of Medicine* 2011;365(15):1439-1445.
  49. **Kesselheim AS**, Cresswell K, Phansalkar S, Bates DW, Sheikh A. Clinical decision support systems could be modified to reduce “alert fatigue” while still minimizing the risk of litigation. *Health Affairs* 2011;30(12):2310-2317.
  50. Cole LW, Kesselheim JC, **Kesselheim AS**. Ethical issues in new drug prescribing. *Journal of Bioethical Inquiry* 2012;9(1):77-83.
  51. Kramer DB\*\*, Xu S\*\*, **Kesselheim AS**. Medical device regulation in the United States and European Union. *New England Journal of Medicine* 2012;366(9):848-855.
  52. **Kesselheim AS**, Karlawish J. Biomarkers unbound — the Supreme Court’s ruling on diagnostic-test patents. *New England Journal of Medicine* 2012;366:2338-2340.
  53. **Kesselheim AS**. Ethical considerations in orphan drug approval and use. *Clinical Pharmacology and Therapeutics* 2012;92:153-155.
  54. Truog RD, **Kesselheim AS**, Joffe S. Paying patients for their tissue: the legacy of Henrietta Lacks. *Science* 2012;337(6090):37-38.
    - Hayflick L. Paying for tissue: the case of WI-38. [Letter to the Editor] *Science* 2012;337(6100):1292.
    - Kominers SD, Becker GS. Paying for tissue: net benefits. [Letter to the Editor] *Science* 2012;337(6100):1292-1293.
    - Truog RD, **Kesselheim AS**, Joffe S. Paying for tissue: net benefits – response. [Author reply] *Science* 2012;337(6100):1293.

55. Kramer DB\*\*, **Kesselheim AS**. User fees and beyond — the FDA Safety and Innovation Act of 2012. *New England Journal of Medicine* 2012;367:1277-1279.
56. **Kesselheim AS** and Avorn J. The Food and Drug Administration has the legal basis to restrict promotion of flawed comparative effectiveness research. *Health Affairs* 2012;31:2200-2205.
57. **Kesselheim AS**, Orentlicher D. Insights from a national conference: “Conflicts of Interest in the Practice of Medicine.” *Journal of Law, Medicine & Ethics* 2012;40(3):436-440.
58. **Kesselheim AS**, Avorn J, Greene JA. Risk, responsibility, and generic drugs. *New England Journal of Medicine* 2012;367(18):1679-1681.
59. **Kesselheim AS**, Mello MM, Avorn J. FDA regulation of off-label drug promotion under attack. *JAMA* 2013;309(5):445-446.
60. **Kesselheim AS**. Drug company gifts to medical students: the hidden curriculum. *BMJ* 2013;346:f1113.
61. Kramer DB\*\* and **Kesselheim AS**. Medical device excise tax — over before it begins? [Letter to the Editor] *New England Journal of Medicine* 2013;368:1767-1769.
  - Sorensen AG. The medical device excise tax—over before it begins? [Letter to the Editor] *New England Journal of Medicine* 2013;369(10):982-983.
  - Prabhakar AM, Harvey HB, Oklu R. The medical device excise tax—over before it begins? [Letter to the Editor] *New England Journal of Medicine* 2013;369(10):983-984.
  - Ubl SJ. The medical device excise tax—over before it begins? [Letter to the Editor] *New England Journal of Medicine* 2013;369(10):984.
  - Kramer DB\*\*, **Kesselheim AS**. The medical device excise tax—before it begins? [Author reply] *New England Journal of Medicine* 2013;369(10):984-985.
62. **Kesselheim AS**. Rising health care costs and life-cycle management in the pharmaceutical market. *PLoS Medicine* 2013;10(5):e1001461.
63. Rajkumar R and **Kesselheim AS**. Balancing access and innovation: India’s Supreme Court rules on Gleevec. *JAMA* 2013;310(3):263-264.
64. **Kesselheim AS**, Green MD, Avorn J. Who is now responsible for discovering and warning about adverse effects of generic drugs? *JAMA* 2013;310(10):1023-1024.
65. **Kesselheim AS**, Cook-Deegan RM, Winickoff DE, Mello MM. Gene patenting—the Supreme Court finally speaks. *New England Journal of Medicine* 2013;369(9):869-875.
66. Kramer DB\*\*, Tan YT, Sato C, **Kesselheim AS**. Postmarket surveillance of medical devices: a comparison of strategies in the US, EU, Japan, and China. *PLoS Medicine* 2013;10(9):e1001519.
67. Wang B\*\* and **Kesselheim AS**. The role of direct-to-consumer pharmaceutical advertising in patient consumerism. *AMA Journal of Ethics* 2013;15(11):960-965.
68. Yeh JS\*\* and **Kesselheim AS**. Same song, different audience: pharmaceutical promotion targeting non-physician health care providers. *PLoS Medicine* 2013;10(11):e1001560.
69. **Kesselheim AS**, Gagne JJ. Strategies for post-market surveillance of drugs for rare diseases. *Clinical Pharmacology and Therapeutics* 2014;95(3):265-268.
70. **Kesselheim AS** and Avorn J. New and unproved medical devices. *BMJ* 2013;347:f7413.
71. Shah AK, Warsh J, **Kesselheim AS**. Ethics of intellectual property rights in an era of globalization. *Journal of Law, Medicine, and Ethics* 2013;41(4):841-851.
72. **Kesselheim AS** and Shiu N\*\*. Evolving role of biomarker patents in personalized medicine. *Clinical Pharmacology and Therapeutics* 2014;95(2):127-129.
73. Kramer DB\*\*, Tan YT\*\*, Sato C, **Kesselheim AS**. Ensuring medical device effectiveness and safety — a cross-national comparison of approaches to regulation. *Food and Drug Law Journal* 2014;69(1):1-24.
74. Treasure CL\*\*, Avorn J, **Kesselheim AS**. What is the public’s right to access medical discoveries based on federally-funded research? *JAMA* 2014;311:907-909.

75. **Kesselheim AS** and Mello MM. Prospects for regulation of off-label drug promotion in an era of expanding commercial speech protection. *University of North Carolina Law Review* 2014;92:1539-1604.
76. Darrow JJ\*\*, Avorn J, **Kesselheim AS**. New FDA breakthrough-drug category—implications for patients. *New England Journal of Medicine* 2014;370:1252-1258.
- McClellan M, Sigal E. New FDA breakthrough-drug category—implications for patients. [Letter to the Editor] *New England Journal of Medicine* 2014;371(1):87-88.
  - Murray BE. New FDA breakthrough-drug category—implications for patients. [Letter to the Editor] *New England Journal of Medicine* 2014;371(1):88.
  - Velleca M. New FDA breakthrough-drug category—implications for patients. [Letter to the Editor] *New England Journal of Medicine* 2014;371(1):88-89.
  - Ricart AD. New FDA breakthrough-drug category—implications for patients. [Letter to the Editor] *New England Journal of Medicine* 2014;371(1):89.
  - Darrow JJ, Avorn J, **Kesselheim AS**. New FDA breakthrough-drug category—implications for patients. [Author reply] *New England Journal of Medicine* 2014;371(1):89-90.\*\*
77. Sarpatwari A\*\*, Avorn J, **Kesselheim AS**. Using a drug-safety tool to prevent competition. *New England Journal of Medicine* 2014;370:1476-1478.
78. Rome BN\*\*, Kramer DB\*\*, **Kesselheim AS**. Approval of high-risk medical devices in the US: implications for clinical cardiology. *Current Cardiology Reports* 2014;16(6):489-498.
79. Hwang TJ\*\*, Avorn J, **Kesselheim AS**. Lifecycle of medical product rules issued by the US Food and Drug Administration. *Journal of Health Politics, Policy, and Law* 2014;39(4):751-780.
80. Duke J and **Kesselheim AS**. The Food and Drug Administration's role in promoting consistent labels for generic drugs. *JAMA Internal Medicine* 2014;174(8):1213-1214.
81. Cortez NG, Cohen IG, **Kesselheim AS**. Regulation of mobile health technologies. *New England Journal of Medicine* 2014;371(4):372-379.
82. **Kesselheim AS** and Gagne JJ. Introduction to a special issue on innovative approaches to studying health outcomes in rare diseases. *Journal of General Internal Medicine* 2014;29 Suppl 3:709-711.
83. Kirschner N, Sulmasy LS, **Kesselheim AS**. Health policy basics: the Physician Payment Sunshine Act and Open Payments program. *Annals of Internal Medicine* 2014;161:519-521.
84. **Kesselheim AS** and Rajan PV\*\*. Regulating incremental innovation in medical devices. *BMJ* 2014;349:g5303.
85. Wang B\*\*, Joffe S, **Kesselheim AS**. Chemotherapy parity laws: a remedy for high drug costs? *JAMA Internal Medicine* 2014;174(11):1721-1722.
86. Sarpatwari A\*\*, **Kesselheim AS**, Malin BA, Gagne JJ, Schneeweiss S. Ensuring patient privacy in data sharing for post-approval research. *New England Journal of Medicine* 2014;371(17):1644-1649.
87. Alpern JD, Stauffer WB, **Kesselheim AS**. High-cost generic drugs: implications for patients and policymakers. *New England Journal of Medicine* 2014;371(20):1859-1862.
- Uhl K, Peters JR, Flanagan K. High-cost generic drugs—implications for patients and policymakers. [Letter to the Editor] *New England Journal of Medicine* 2015;372(7):685-686.
  - **Kesselheim AS**, Alpern JD, Stauffer WM. High-cost generic drugs—implications for patients and policymakers. [Author reply] *New England Journal of Medicine* 2015;372(7):686.
88. Miller FG, Joffe S, **Kesselheim AS**. Evidence, errors, and ethics. *Perspectives in Biology and Medicine* 2014;57(3):299-307.
89. Darrow JJ\*\*, Sarpatwari A, Avorn J, **Kesselheim AS**. Practical, legal, and ethical issues in

- expanded access to investigational drugs. *New England Journal of Medicine* 2015;372(3):279-286.
- Lurie P, Chan-Tack KM, Woodcock J. [Letter to the Editor] Expanded access to investigational drugs. *New England Journal of Medicine* 2015;372(15):1473.
  - Sarpatwari AS\*\*, Darrow JJ\*\*, **Kesselheim AS**. Legal, ethical, and practical implications of expanded access to investigational drugs. [Author reply] *New England Journal of Medicine* 2015;372(15):1473-1474.
90. **Kesselheim AS** and Darrow JJ. FDA designations for therapeutics and their impact on drug development and regulatory review outcomes. *Clinical Pharmacology and Therapeutics* 2015;97(1):29-36.
  91. Hwang TJ\*\*, Carpenter D, **Kesselheim AS**. Paying for innovation: reimbursement incentives for antibiotics. *Science Translational Medicine* 2015;7(276):276fs9.
  92. Kramer DB\*\*, **Kesselheim AS**. The Watchman saga: closure at last? *New England Journal of Medicine* 2015;372(11):994-995.
  93. Sarpatwari A\*\*, Choudhry NK, Avorn J, **Kesselheim AS**. Paying physicians to prescribe generic drugs and follow-on biologics in the United States. *PLoS Medicine* 2015;12(3):e1001802.
  94. Hwang TJ\*\*, Lehmann LS, **Kesselheim AS**. Precision medicine and the FDA's draft guidance on regulating laboratory-developed tests. *Nature Biotechnology* 2015;33(5):449-451.
  95. Sarpatwari A\*\*, Avorn J, **Kesselheim AS**. Progress and hurdles in follow-on biologics. *New England Journal of Medicine* 2015;372(35):2380-2382.
  96. Avorn J and **Kesselheim AS**. The 21st Century Cures Act—will it take us back in time? *New England Journal of Medicine* 2015;372(36):2473-2475.
    - McClellan MB, Sigal EV. The 21st Century Cures Act. [Letter to the Editor] *New England Journal of Medicine* 2015;373(17):1677-1678.
    - Kakkis E, Bronstein MG. The 21st Century Cures Act. [Letter to the Editor] *New England Journal of Medicine* 2015;373(17):1678.
    - Calderwood SB, Murray BE, Chambers HF. The 21st Century Cures Act. [Letter to the Editor] *New England Journal of Medicine* 2015;373(17):1679.
    - **Kesselheim AS**, Avorn J. The 21st Century Cures Act. [Author reply] *New England Journal of Medicine* 2015;373(17):1679-1680.
  97. Hwang TJ\*\*, Powers JH, Carpenter D, **Kesselheim AS**. Accelerated pathway for rapid diagnostics and targeted antibacterial agents for serious and life-threatening diseases. *Nature Biotechnology* 2015;33(6):589-590.
  98. Sharfstein JM and **Kesselheim AS**. The safety of prescription drugs. *JAMA* 2015;314(3):233-234.
  99. **Kesselheim AS** and Gagne JJ. Product-specific regulatory pathways to approve generic drugs: the need for follow-up studies to ensure safety and effectiveness. *Drug Safety* 2015;38(10):849-853.
  100. Sarpatwari A\*\* and **Kesselheim AS**. The 21st Century Cures Act: opportunities and challenges. *Clinical Pharmacology and Therapeutics* 2015;98(6):575-577.
  101. Luo J\*\* and **Kesselheim AS**. The Trans-Pacific Partnership Agreement and implications for access to essential medicines. *JAMA* 2015;314(15):1563-1564.
  102. **Kesselheim AS** and Darrow JJ\*\*. Hatch-Waxman turns 30: do we need a re-designed approach for the modern era? *Yale J Health Policy Law Ethics* 2015;15(2):293-347.
  103. Avorn J, Sarpatwari A\*\*, **Kesselheim AS**. Forbidden and permitted statements about medications—rules in evolution. *New England Journal of Medicine* 2015;373(10):967-973.
  104. Ross JS and **Kesselheim AS**. FDA policy and cardiovascular medicine. *Circulation* 2015;132:1136-1145.
  105. **Kesselheim AS**, Maggs LR, Sarpatwari A. Experience with the priority review voucher program for drug development. *JAMA* 2015;314(16):1687-1688.



- Ridley DB, Dent J, Egerton-Warburton C. Efficacy of the priority review voucher program. [Letter to the Editor] *JAMA* 2016;315(15):1659-1660.
  - Sarpatwari A\*\*, **Kesselheim AS**. Efficacy of the priority review voucher program. [Author reply] *JAMA* 2016;315(15):1659-1661.
106. **Kesselheim AS**, Sinha MS\*\*, Joffe S. Physicians and insider trading. *JAMA Internal Medicine* 2015;175(12):1955-1959.
  107. Luo J\*\*, Sarpatwari A, **Kesselheim AS**. Regulatory solutions to the problem of high generic drug costs. *Open Forum Infectious Diseases* 2015;2(4):ofv179.
  108. **Kesselheim AS** and Hwang TJ. Breakthrough medical devices and the 21st Century Cures Act. *Annals of Internal Medicine* 2016;164(7):500-502.
  109. Podolsky SH and **Kesselheim AS**. Regulating homeopathic products — a century of dilute interest. *New England Journal of Medicine* 2016;374(3):201-203.
  110. Sarpatwari A\*\* and **Kesselheim AS**. The case for reforming drug naming: should brand-name trademark protections expire upon generic entry? *PLoS Medicine* 2016;13(2):e1001955.
  111. Wang B\*\* and **Kesselheim AS**. Promoting therapeutic innovation: what do we do about drug-device combinations? *JAMA* 2016;315(9):857-858.
  112. Rathi V, **Kesselheim AS**, Ross JS. The FDA 515 Initiative: addressing the evidence gap for high-risk cardiovascular devices? *JAMA Cardiology* 2016;1(2):117-118.
  113. Hey SP\*\* and **Kesselheim AS**. An uninformative truth: the logic of Amarin’s off-label promotion for prescription fish oil. *PLoS Medicine* 2016;13(3):e1001978.
  114. Yeh JS\*\*, Sarpatwari A, **Kesselheim AS**. Ethical and practical considerations in removing Black Box Warnings from drug labels. *Drug Safety* 2016;39(8):709-714.
  115. Capati VC\*\* and **Kesselheim AS**. Drug product life-cycle management as anticompetitive behavior: the case of memantine. *Journal of Managed Care & Specialty Pharmacy* 2016;22(4):339-344.
  116. Kapczynski A and **Kesselheim AS**. Government ‘patent use’: a legal approach to reduce drug spending. *Health Affairs* 2016;35(5):791-797.
    - Grabowski H. Government appropriation of breakthrough drug patent rights would deter biopharmaceutical R&D and innovation. [Commentary] *Health Affairs Blog*. June 20, 2016. Available on-line at: <https://www.healthaffairs.org/doi/10.1377/hblog20160620.055440/full/>
    - Kapczynski A, **Kesselheim AS**. Why government patent use to lower drug costs won’t stifle innovation. [Author response] *Health Affairs Blog*. July 28, 2016. Available at: <http://healthaffairs.org/blog/2016/07/28/why-government-patent-use-to-lower-drug-costs-wont-stifle-innovation/>
  117. Treasure CL\*\* and **Kesselheim AS**. How patent troll legislation can increase timely access to generic drugs. *JAMA Internal Medicine* 2016;176(6):729-30.
  118. Engelberg AB and **Kesselheim AS**. Use the Bayh-Dole Act to lower drug prices for government healthcare programs. *Nature Medicine* 2016;22(6):576.
  119. Sarpatwari A\*\*, Avorn J, **Kesselheim AS**. State initiatives to control medication costs — can transparency legislation help? *New England Journal of Medicine* 2016;374(24):2301-2304.
  120. Luo J\*\* and **Kesselheim AS**. Protecting pharmaceutical patents and data: How the Trans-Pacific Partnership Agreement could affect access to medicines in the US and abroad. *AMA Journal of Ethics* 2016;18:727-735.
  121. Hwang TJ\*\* and **Kesselheim AS**. Leveraging novel and existing approval pathways to combat antimicrobial resistance. *American Journal of Law & Medicine* 2016;42:429-450.
  122. Hey SP\*\* and **Kesselheim AS**. Countering imprecision in precision medicine. *Science* 2016;353(6298):448-449.

123. Hey SP\*\* and **Kesselheim AS**. The FDA, Juno Therapeutics, and the ethical imperative of transparency. *BMJ* 2016;354:i4435.
124. Sinha MS\*\* and **Kesselheim AS**. Regulatory incentives for antibiotic development: a review of recent proposals. *Bioorganic & Medicinal Chemistry*. 2016;24(24):6446-6451.
125. Kimmelman J and **Kesselheim AS**. Translational research and the U.S. federal elections. *Science Translational Medicine* 2016;8(361):361ed13.
126. **Kesselheim AS** and Avorn J. Approving a problematic muscular dystrophy drug: implications for FDA policy. *JAMA* 2016;316(22):2357-2358.
  - Nelson SF, Miceli MC. FDA approval of eteplirsen for muscular dystrophy. [Letter to the Editor] *JAMA* 2017;317(14):1480.
  - Sackner-Bernstein J. FDA approval of eteplirsen for muscular dystrophy. [Letter to the Editor] *JAMA* 2017;317(14):1480-1481.
  - **Kesselheim AS**, Avorn J. FDA Approval of eteplirsen for muscular dystrophy. [Author reply] *JAMA* 2017;317(14):1481-1482.
127. Robertson CT and **Kesselheim AS**. Regulating off-label promotion—a critical test. *New England Journal of Medicine* 2016;375(24):2313-2315.
128. Sarpatwari A\*\*, Gagne JJ, Levidow NL\*\*, **Kesselheim AS**. Active surveillance of follow-on biologics: a prescription for uptake. *Drug Safety* 2017;40(2):105-108.
129. Carrier MA, Levidow NL\*\*, **Kesselheim AS**. Using antitrust law to challenge Turing’s Daraprim price increase. *Berkeley Technology Law Journal* 2016;31(2):1379-1408.
130. Mostaghim S\*\* and **Kesselheim AS**. Suitability of expanding the priority review voucher into rare disease drug development. *Expert Opinion on Orphan Drugs* 2016;4:10:1001-1003.
131. **Kesselheim AS** and Avorn J. New “21st Century Cures” legislation—speed and ease vs. science. *JAMA* 2017;317(6):581-582.
132. Guo E, Jacobs DB, **Kesselheim AS**. Eliminating coverage discrimination through the essential health benefit’s anti-discrimination provisions. *American Journal of Public Health* 2017;107(2):253-254.
133. Luo J\*\*, **Kesselheim AS**, Greene J, Lipska KJ. Strategies to improve the affordability of insulin in the USA. *Lancet Diabetes and Endocrinology* 2017;5(3):158-159.
134. Fralick M\*\* and **Kesselheim AS**. Periodic Benefit Risk Evaluation Reports have substantial promise to guide patient care and should be made publicly available. *Pharmacoepidemiology and Drug Safety* 2017;26(5):597-599.
135. Hey SP\*\* and **Kesselheim AS**. Reprioritizing research activity for the post-antibiotic era: ethical, legal, and social considerations. *Hastings Center Report* 2017;47(2):16-20.
136. Fralick M\*\* and **Kesselheim AS**. FDA approval of desmopressin for nocturia. *JAMA* 2017;317(20):2059-2060.
  - Fein S, Herschkowitz S. Low-dose desmopressin nasal spray and FDA approval. [Letter to the Editor] *JAMA* 2017;318(11):1070-1071.
  - Fralick M\*\*, **Kesselheim AS**. Low-dose desmopressin nasal spray and FDA approval. [Author reply] *JAMA* 2017;318(11):1071-1072.
137. Kramer DB\*\* and **Kesselheim AS**. Coverage of magnetic resonance imaging for patients with cardiac devices: improving the coverage with evidence development program. *JAMA Cardiology* 2017;2(7):711-712.
138. Sharfstein JM, Miller JD, Davis AL, Ross JS, Alexander GC, **Kesselheim AS**. FDA Transparency Working Group. Blueprint for Transparency at the U.S. Food and Drug Administration: Recommendations to Advance the Development of Safe and Effective Medical Products. *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):7-23. Available at <http://jhsph.edu/BlueprintFDA>.

- Califf RM. Transparency at the U.S. Food and Drug Administration. [Commentary] *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):24-28.
  - Carpenter D. FDA transparency in an inescapably political world. [Commentary] *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):29-32.
  - Kapczynski A and Kim J. Clinical trial transparency: the FDA should and can do more. [Commentary] *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):33-38.
  - Cortez N. FDA and the marketplace of ideas for medical products. [Commentary] *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):39-41.
  - Doshi P and Jefferson T. Disclose data publicly, without restriction. [Commentary] *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):42-45.
  - Almashat S and Carome M. Withholding information on unapproved drug marketing applications: the public has a right to know. [Commentary] *Journal of Law, Medicine, and Ethics* 2017;45(4 suppl 2):46-49.
139. Sinha MS\*\* and **Kesselheim AS**. The effects of the Sunshine Act: what can and should we expect? *American Journal of Bioethics* 2017;17(6):22-24.
  140. Chen CT and **Kesselheim AS**. The journey of generic imatinib: a case study in oncology drug pricing. *Journal of Oncology Practice* 2017;13(6):352-355.
  141. Gellad WF and **Kesselheim AS**. Accelerated approval and expensive drugs — a challenging combination. *New England Journal of Medicine* 2017;376(21):2001-2004.
  142. Hwang TJ\*\* and **Kesselheim AS**. Challenges in the development of novel cardiovascular therapies. *Clinical Pharmacology and Therapeutics* 2017;102(2):194-196.
  143. Hwang TJ\*\*, **Kesselheim AS**, Sarpatwari A. Value-based pricing and state reform of prescription drug costs. *JAMA* 2017;318(7):609-610.
  144. Bonnie RJ, **Kesselheim AS**, Clark DJ. Both urgency and balance needed in addressing opioid epidemic. *JAMA* 2017;318(5):423-424.
  145. Sarpatwari A\*\*, Sinha MS, **Kesselheim AS**. The opioid epidemic: fixing a broken pharmaceutical market. *Harvard Law and Policy Review* 2017;464(11):463-484.
  146. Hey SP\*\*, Cohen IG, Adashi EY, **Kesselheim AS**. Influence, integrity, and the FDA: an ethical framework. *Science* 2017;357(6354):876-877.
  147. **Kesselheim AS**, Sinha MS\*\*, Avorn J. Determinants of market exclusivity for prescription drugs in the United States. *JAMA Internal Medicine* 2017;177(11):1658-1664.
  148. Vokinger KN\*\*, **Kesselheim AS**, Avorn J, Sarpatwari A\*\*. Strategies that delay market entry of generic drugs. *JAMA Internal Medicine* 2017;177(11):1665-1669.
  149. Seeley E, **Kesselheim AS**. Outcomes-based pharmaceutical contracts: an answer to high U.S. drug spending? *Commonwealth Fund Issue Brief*. 2017 Sep;2017:1-8. Available from: <http://www.commonwealthfund.org/publications/issue-briefs/2017/sep/outcomes-based-contracts-high-drug-spending>
  150. Bothwell LE\*\*, **Kesselheim AS**. The real-world ethics of adaptive-design clinical trials. *Hastings Center Report* 2017;47(6):27-37.
  151. Darrow JJ\*\*, Beall RF\*\*, **Kesselheim AS**. Will inter partes review speed US generic drug entry? *Nature Biotechnology* 2017;35(12):1139-1141.
  152. Sinha MS\*\*, **Kesselheim AS**, Darrow JJ. Pharmaceutical advertising in medical journals: revisiting a long-standing relationship. *Chest* 2017;153(1):9-11.
  153. Sommers BD, **Kesselheim AS**. Massachusetts' proposed Medicaid reforms — cheaper drugs and better coverage? *New England Journal of Medicine* 2018;378(2):109-111.
  154. Darrow JJ\*\*, Fuse Brown EC, **Kesselheim AS**. The Regulatory Accountability Act of 2017 — implications for FDA regulation and public health. *New England Journal of Medicine* 2018;378(5):412-414.

155. Hey SP\*\*, Weijer C, Taljaard M, **Kesselheim AS**. Research ethics for emerging trial designs: does equipoise need to adapt? *BMJ* 2018;360:k226.
156. Lee TT, **Kesselheim AS**, Kapczynski A. Legal challenges to state drug pricing laws. *JAMA* 2018;319(9):865-866.
157. Powers JH, Evans SR, **Kesselheim AS**. Studying new antibiotics for multidrug resistant infections: are today's patients paying for unproved future benefits? *BMJ* 2018;360:k587.
158. Darrow JJ\*\*, **Kesselheim AS**. Promoting competition to address pharmaceutical prices. *Health Affairs Health Policy Brief* March 15, 2018. DOI:10.1377/hpb20180116.967310
  - Lott R. Three new *policy options* papers examine factors driving prescription drug prices. *Health Affairs Blog*. March 15 2018. Available from: <https://www.healthaffairs.org/doi/10.1377/hblog20180313.644056/full/>
  - Inzerro A. Health policy experts voice ways to tame rising drug costs, protect patients. *AJMC*. April 3, 2018. Available at: <http://www.ajmc.com/newsroom/health-policy-experts-voice-ways-to-tame-rising-drug-costs-protect-patients>.
159. Lee TT, **Kesselheim AS**. U.S. Food and Drug Administration's PreCertification Pilot Program for digital health software: weighing the benefits and risks. *Annals of Internal Medicine* 2018;168(10):730-732.
  - Howard NL, Ackerman W. What physicians are reading about digital health. *National Law Review*. June 4, 2018. Available at: <https://www.natlawreview.com/article/what-physicians-are-reading-about-digital-health>.
160. Sinha MS\*\*, **Kesselheim AS**. The next forum for unraveling FDA off-label marketing rules: state and federal legislatures. *PLoS Medicine* 2018;15(5):e1002564.
161. Sacks CA\*\*, **Kesselheim AS**, Fralick M\*\*. The shortage of normal saline in the wake of Hurricane Maria. *JAMA Internal Medicine* 2018;178(7):885-886.
162. Luo J\*\*, **Kesselheim AS**. Delayed generic market saturation after patent expiration—a billion-dollar problem. *JAMA Internal Medicine* 2018;178(5):721-722.
163. Sarpatwari A\*\*, Avorn J, **Kesselheim AS**. An incomplete prescription: President Trump's plan to address high drug prices. *JAMA* 2018;319(23):2373-2374.
164. Zettler PJ, Foster Riley M, **Kesselheim AS**. Implementing a public health perspective in FDA drug regulation. *Food and Drug Law Journal* 2018;73(2):221-256.
165. Gyawali B\*\*, **Kesselheim AS**. Reinforcing the social compromise of accelerated approval. *Nature Reviews Clinical Oncology* 2018;15(10):596-597.
166. Wang A, **Kesselheim AS**. Government patent use to address the rising cost of naloxone: 28 U.S.C. § 1498 and *Evzio*. *Journal of Law, Medicine, and Ethics* 2018;46(2):472-484.
167. Hey SP\*\*, **Kesselheim AS**. Defining “true and non-misleading” for pharmaceutical promotion. *Journal of Law, Medicine, and Ethics* 2018;46(2):552-554.
168. Avorn J, **Kesselheim A**, Sarpatwari A\*\*. The FDA Amendments Act of 2007—assessing its effects a decade later. *New England Journal of Medicine* 2018;379(12):1097-1099.
169. Sarpatwari A\*\*, Barenie R\*\*, Curfman G, **Kesselheim AS**. The US biosimilar market: stunted growth and possible reforms. *Clinical Pharmacology and Therapeutics* 2019;105(1):92-100.
170. Minssen T, **Kesselheim AS**, Darrow JJ\*\*. An export-only exception to pharmaceutical patents in Europe: should the United States follow suit? *Nature Biotechnology* 2019;37(1):21-22.
171. Sarpatwari A\*\*, **Kesselheim AS**. **Tepid Steps on Drug Pricing**. *JAMA Internal Medicine* 2019 Jan 22. doi: 10.1001/jamainternmed.2018.6593.
172. Gyawali B\*\*, **Kesselheim AS**. The promise of ESCAT: a new system for evaluating cancer drug-target combination. *Nature Reviews Clinical Oncology* 2019 [in press]
173. Gyawali B\*\*, **Kesselheim AS**. FDA approval of new drugs based on non-inferiority trials in oncology: a dangerous precedent? *JAMA Oncology* 2019 [in press]

#### Scholarship without named authorship

1. Stanic Benic M, Milanic R, Monnier AA, Gyssens IC, Adriaenssens N, Versporten A, Zanichelli V, Le Maréchal M, Huttner B, Tebano G, Hulscher ME, Pulcini C, Schouten J, Vlahovic-Palcevski V; DRIVE-AB WP1 group. Metrics for quantifying antibiotic use in the hospital setting: results from a systematic review and international multidisciplinary consensus procedure. *Journal of Antimicrobial Chemotherapy* 2018;73(suppl\_6):vi50-vi58.
2. Monnier AA, Schouten J, Le Maréchal M, Tebano G, Pulcini C, Stanic Benic M, Vlahovic-Palcevski V, Milanic R, Adriaenssens N, Versporten A, Huttner B, Zanichelli V, Hulscher ME, Gyssens IC; DRIVE-AB WP1 group. Quality indicators for responsible antibiotic use in the inpatient setting: a systematic review followed by an international multidisciplinary consensus procedure. *Journal of Antimicrobial Chemotherapy* 2018;73(suppl\_6):vi30-vi39.

#### **Non-peer reviewed scientific or medical publications/materials in print or other media**

##### Proceedings of meetings or other non-peer reviewed research publications

1. **Kesselheim AS**. Should FDA drug and medical device regulation bar state liability claims? Hearing before the House of Representatives Committee on Oversight and Government Reform (Rep. Waxman, Chairman). 14 May 2008. United States Congressional Record. Serial No. 110-112. Available on-line at: <http://www.gpo.gov/fdsys/pkg/CHRG-110hhr56191/pdf/CHRG-110hhr56191.pdf>.
2. Carpenter DM, **Kesselheim AS**, Avorn J, Law MT, Polsky D. State failure-to-warn litigation plays a critical role in ensuring drug safety. Amicus curiae brief to the Supreme Court in the case of *Wyeth v. Levine*. Aug 2008.
3. Coukell AG, Avorn J, **Kesselheim AS**, Psaty B, Nissen S, Turner E, Dickersin K, Ross J, Mann H, Caplan A, Maschke K. Comments to the FDA Transparency Task Force. 24 June 2009.
4. **Kesselheim AS**. The essential role of the FDA in certifying innovative device safety and effectiveness. Hearing before the House of Representatives Committee on Energy and Commerce Subcommittee on Health (Rep. Pitts, Chairman). 15 Feb 2012. United States Congressional Record. Serial No. 112-116. Available on-line at: <http://www.gpo.gov/fdsys/pkg/CHRG-112hhr76255/pdf/CHRG-112hhr76255.pdf>.
5. **Kesselheim AS**, Avorn J, Engelberg AB. Comments on draft guidance: ‘Size, Shape, and Other Physical Attributes of Generic Tablets and Capsules’ (FDA-2013-B-1434-0001). 2014 Mar 10. Available on-line at: <http://www.regulations.gov/#!documentDetail;D=FDA-2013-N-1434-0022>.
6. **Kesselheim AS**, Avorn J, Duke J. Comments on proposed rule: ‘Supplemental Applications Proposing Labeling Changes for Approved Drugs and Biologic Products’ (RIN 0910-AG94). 2014 Mar 10. Available on-line at: <http://www.regulations.gov/#!documentDetail;D=FDA-2013-N-0500-0023>
7. **Kesselheim AS**. 21st century cures: modernizing clinical trials. Hearing before the House of Representatives Committee on Energy and Commerce Subcommittee on Health (Rep. Pitts, Chairman). 9 Jul 2014. United States Congressional Record. Serial No. 113-157. Available on-line at: <https://www.gpo.gov/fdsys/pkg/CHRG-113hhr91847/html/CHRG-113hhr91847.htm>.
8. **Kesselheim AS**. Why are generic drugs suddenly skyrocketing in price? Hearing before the Senate Committee on Health, Education, Labor and Pensions Subcommittee on Primary Health and Aging (Sen. Sanders, Chairman). 20 Nov 2014. United States Congressional Record. Senate Hearing 113-859. Available on-line at: <https://www.gpo.gov/fdsys/pkg/CHRG-113shrg24459/html/CHRG-113shrg24459.htm>.
9. Alpern JD, Stauffer WB, **Kesselheim AS**. Comments on proposed rule: ‘Proposed Criteria for “First Generic” Submissions for Purposes of Abbreviated New Drug Application Review

- Prioritization Under the Generic Drug User Fee Amendments’ (RIN FDA-2014-N-1741). 2014 Dec 7. Available on-line at: <http://www.regulations.gov/#!documentDetail;D=FDA-2014-N-1741-0003>.
10. Sarpatwari A\*\*, **Kesselheim AS**. Further comments on proposed rule: ‘Supplemental Applications Proposing Labeling Changes for Approved Drugs and Biologic Products’ (Docket FDA-2013-N-0500). 2015 Apr 8. Available on-line at: <https://www.regulations.gov/document?D=FDA-2013-N-0500-0106>.
  11. Shachar C, Greenwald R, Sarpatwari A, **Kesselheim AS**, et al. All Payer Claims Databases are important for public health and are not preempted by ERISA. Amicus curiae brief to the Supreme Court in the case of Gobeille v. Liberty Mutual. 2015 Sept.
  12. Sarpatwari AS\*\*, **Kesselheim AS**, Avorn J. Comments to CMS re: follow-on biologics reimbursement policy. 2015 Sept 8. Available on-line at: <http://freepdfhosting.com/660d337b8f.pdf>.
  13. **Kesselheim AS**. S.1048: A Bill to Improve Transparency of Drug Research and Development Costs. Hearing before the Massachusetts Legislature Joint Committee on Health Care Financing (Rep. Sanchez and Sen. Welch, Co-Chairman). 2016 Apr 11.
  14. Constitutional, Administrative, Contracts, and Health Law Scholars. In support of respondents. Amicus curiae brief to the Supreme Court in the case of Expressions Hair Design v. Schneiderman. 2016 Dec. Available on-line at: <http://www.scotusblog.com/wp-content/uploads/2016/12/15-1391-amicus-respondents-Constitutional-Administrative-Contracts-and-Health-Law-Scholars.pdf>.
  15. Sharfstein JM, Anderson G, Ballreich JM, Brennan HW, Greene JA, Jackson MR, Kapczynski A, **Kesselheim AS**, Lee JJ, Sen AP, Trujillo AJ. Hepatitis C in Louisiana: recommendations on drug availability. Memo to Secretary of Health of Louisiana. May 4, 2017. Available on-line at: <http://ldh.la.gov/assets/docs/HepatitisC/ResponsememotoSecretaryGeeHCV.pdf>.
  16. Sinha MS\*\*, **Kesselheim AS**. Re: FDA-2013-N-0402-0029, Generic Drug User Fee Amendments of 2012; Regulatory Science Initiatives; Public Workshop; Request for Comments. 2017 June 5. (Docket FDA-2013-N-0402-0034). Available on-line at: <https://www.regulations.gov/document?D=FDA-2013-N-0402-0034>\*\*
  17. **Kesselheim AS**. The public health implications of reducing restrictions on off-label promotion. Hearing before the House Committee on Energy and Commerce Subcommittee on Health (Rep. Burgess, Chairman). 12 July 2017. United States Congressional Record. [In press]
  18. **Kesselheim AS**. Strategies that delay timely entry of generic drugs and potential policy solutions: the CREATES Act and beyond. Hearing before the House Committee on the Judiciary Subcommittee on Regulatory Reform, Commercial and Antitrust Law (Rep. Marino, Chairman). 27 July 2017. United States Congressional Record. [In press]
  19. Sarpatwari A\*\*, Sinha MS\*\*, **Kesselheim AS**. Administering the Hatch-Waxman Amendments: Ensuring a Balance Between Innovation and Access; Public Meeting; Request for Comments. November 10, 2017. (Docket FDA-2017-N-3615-0001). Available on-line at: <https://www.regulations.gov/document?D=FDA-2017-N-3615-0001>\*\*

#### Reviews, chapters, monographs and editorials

1. Gelband H, Greene A, **Kesselheim AS**, Kidd E. High quality end-of-life care for individuals dying from cancer: current beliefs and barriers. [White Paper] Nat’l Cancer Policy Board 2000 Jan. Available at [http://www4.nationalacademies.org/iom/iomhome.nsf/WFiles/ncpbeol/\\$file/ncpbeol.pdf](http://www4.nationalacademies.org/iom/iomhome.nsf/WFiles/ncpbeol/$file/ncpbeol.pdf)
2. **Kesselheim AS**. Ensuring quality end-of-life cancer care: an analysis of the current state of patient and family information about end-of-life care issues, its deficiencies, and recommendations for the

- future. [Textbook chapter] In: Gelband H & Foley KM, eds. Excellent end-of-life care for cancer patients: a policy agenda for action and research. Washington, D.C.: National Academy Press; 2001. Pp. 132-152.
3. **Kesselheim AS**. Patents and intellectual property in medicine. [Textbook chapter] In: Sanbar SS and Firestone MH, eds. Legal Medicine, 7th ed. Philadelphia, PA: Mosby Press; 2007. Pp. 151-158.
  4. **Kesselheim AS** and Avorn J. Patents and public health. [Book chapter] In: Galea S, ed. Macrosocial Determinants of Health. New York, NY: Springer Science; 2007. Pp. 233-246.
  5. **Kesselheim AS**. An idea takes root: hold expert witnesses accountable. OBG Management 2007;19(11):30-36.
  6. **Kesselheim AS**. Resident supervision and patient safety. CRICO/RMF Insight Summer 2008;1.
  7. Gagne JJ and **Kesselheim AS**. Experimenting with the consumer: the mass testing of risky products on the American public (Shapo, 2008). Journal of Law Medicine & Ethics 2010;38(2):432-435.
  8. **Kesselheim AS**. Innovation and the Orphan Drug Act, 1983-2009: the regulatory and clinical characteristics of approved orphan drugs. [Book chapter] In: Field MJ & Boat TF, eds. Accelerating rare diseases research and orphan product development. Washington, D.C.: National Academy Press, 2010. Pp. 291-308.
  9. Licurse A and **Kesselheim AS**. Conflicts of Interest and the Future of Medicine (Rodwin, 2011). Journal of Bioethical Inquiry 2011;8(4):383-386.
  10. **Kesselheim AS**. Pharmacoepidemiology and the law. [Textbook chapter] In: Strom B, Kimmel S, Hennessy S, eds. Pharmacoepidemiology, 5th ed. Chichester, UK: John Wiley & Sons, 2012. Pp. 117-134.
  11. **Kesselheim AS**. Institutional corruption and perceptions of methodological rigor. [Book chapter] In: Lessig L and Somos M, eds. Research in Action: Lab Dispatches, Vol. 1. Cambridge, MA: Harvard University, 2013. Pp. 19-21.
  12. Avorn J, Mo J, Reynolds RF, Dal Pan GJ, Arlett P, **Kesselheim AS**. Views from academia, industry, regulatory agencies, and the legal system. [Textbook chapter] In: Strom BL, Kimmel SE, Hennessy S, eds. Textbook of Pharmacoepidemiology, 2nd ed. Chichester, UK: John Wiley & Sons, 2013. Pp. 63-97.
  13. **Kesselheim AS**. The pharmaceutical market's adverse effects (Bad Pharma (Goldacre, 2012)). Health Affairs 2014;33:179-180.
  14. **Kesselheim AS**, Mello MM. Prospects for regulation of off-label drug promotion in an era of expanding commercial speech protection. [Textbook chapter] In: Lynch HF and Cohen IG, eds. FDA in the 21st Century: The Challenges of Regulating Drugs and New Technologies. New York, NY: Columbia University Press, 2015. Pp. 184-203.
  15. **Kesselheim AS**, Yeh JS. Interactions with Pharmaceutical and Medical Device Industry Representatives [Textbook chapter]. ACP Smart Medicine. <http://smartmedicine.acponline.org/content.aspx?gbosId=393>. Updated May 7, 2015. doi 10.7326/el057.
  16. Sarpatwari AS, Choudhry NK, Avorn J, **Kesselheim AS**. Using behavioral economics to promote physicians' prescribing of generic drugs and follow-on biologics: what are the issues? [Textbook chapter] In: Cohen IG, Lynch HF, Robertson CT, eds. Behavioral Economics, Law, and Health Policy. Baltimore, MD: Johns Hopkins University Press, 2016. Pp. 158-171.\*\*
  17. **Kesselheim AS**. Behavioral economics and the doctor-patient relationship [Textbook chapter introduction] In: Cohen IG, Lynch HF, Robertson CT, eds. Behavioral Economics, Law, and Health Policy. Baltimore, MD: Johns Hopkins University Press, 2016. Pp. 219-221.
  18. Kapczynski A, **Kesselheim AS**, Ezer T. Submission to the Department of Trade and Industry on

the intellectual property consultative framework. Oct 28 2016. Available on-line at:  
[http://media.wix.com/ugd/148599\\_721218272a1c43cbaee5ff8fda3a00cf.pdf](http://media.wix.com/ugd/148599_721218272a1c43cbaee5ff8fda3a00cf.pdf).

19. **Kesselheim AS**. Background and foundations: Introduction [Textbook chapter introduction] In: Lynch HF, Bierer BE, Cohen IG, Rivera SM, eds. Specimen Science: Ethics and Policy Implications. Cambridge, MA: MIT Press, 2017. Pp. 21-23.

#### Books/Textbooks for the medical or scientific community

1. Robertson CT, **Kesselheim AS**, eds. Blinding as a Solution to Bias: Strengthening Biomedical Science, Forensic Science, and Law. London, UK: Elsevier Academic Press, 2016.
2. National Academies of Sciences, Engineering, and Medicine Committee on Pain Management and Regulatory Strategies to Address Prescription Opioid Abuse (Bonnie RJ, Amaro H, Burnes Bolton L, Caulkins JP, Clark D, Eliav E, FitzGerald G, Green TC, Hernan M, Hoffer LD, Jarris PE, Kaltenbach K, **Kesselheim AS**, Mackenzie-Brown AM, Moron-Concepcion J, Paltiel AD, Schumacher M, Reyna V). Pain Management and the Opioid Epidemic: Balancing Societal and Individual Benefits and Risks of Prescription Opioid Use. Washington, D.C.: National Academies Press, 2017.

#### Letters to the Editor

1. Stedman MR, Elbourne DR, Curtain F, **Kesselheim AS**, Brookhart MA. Meta-analyses involving cross-over trials: methodological issues. [Letter to the Editor] International Journal of Epidemiology 2011;40(6):1732-1734.
2. Outterson K, Powers III JH, Gould IM, **Kesselheim AS**. Questions about the 10 x '20 initiative. [Letter to the Editor] Clinical Infectious Diseases 2010;51(6):751-752.
3. **Kesselheim AS**. Adalimumab pricing and market exclusivity for biologics. [Letter to the Editor] New England Journal of Medicine 2010;363(24):2374.
4. **Kesselheim AS**, Avorn J. Drug labels: a flawed source of data for studying orphan drug approvals. [Letter to the Editor] Clinical Pharmacology and Therapeutics 2012;92(6):694.
5. Luo J\*\*, **Kesselheim AS**. Underrepresentation of older adults in cancer trials [Letter to the Editor] JAMA 2014;311:965-967.
6. Luo J\*\*, **Kesselheim AS**. Transparency in drug costs may improve cost-effectiveness studies. [Letter to the Editor] JAMA 2015;314(20):2191.
7. Fralick M\*\*, **Kesselheim A**, Avorn J. Applying academic detailing and process change to promote Choosing Wisely. [Letter to the Editor] JAMA Internal Medicine 2017;177(2):282.
8. Fralick M\*\*, **Kesselheim AS**. Three design aspects for high quality post-marketing cohort studies. [Letter to the Editor] BMJ 2017;357:j1851.

#### Professional educational materials or reports, in print or other media

1. Zerden M, Avorn J, **Kesselheim AS**. Pharmaceutical marketing practices towards physicians. Written for Improvehealthcare.org; 2008.
2. Gold L, Avorn J, **Kesselheim AS**. Postmarketing review and analysis of new drugs. Written for Improvehealthcare.org; 2008.
3. Wheeler LA, Avorn J, **Kesselheim AS**. The FDA and the drug approval process. Written for Improvehealthcare.org; 2008.
4. Rivara M, Mello MM, **Kesselheim AS**. Medical malpractice: Current issues and policies. Written for Improvehealthcare.org; 2008.
5. Song Z and **Kesselheim AS**. Conflicts of interest policies and federal antikickback and fraud laws. Written for Improvehealthcare.org; 2008.

Teaching cases



All written for medical students affiliated with Improvehealthcare.org, a student-run organization based at Harvard Medical School, with 19 affiliated chapters, that uses case-based learning to teach physicians-in-training about health policy issues. At the time it was developed, it was available to all interested medical students on integrated website.

### **Clinical guidelines and reports**

1. **Kesselheim AS**, Stevenson LW, Nohria A, Fischer MA, Avorn J. Assessing patients with decompensated congestive heart failure. Brigham and Women's Hospital medication use guidelines. Feb 2005.  
Clinical algorithm  
Written for cardiologists, emergency department physicians, intensive care physicians and house officers in the Partners healthcare system.
2. Choudhry NK, Fischer MA, Hoge E, **Kesselheim AS**, Parikh S, Shrank WH. The pursuit of happiness: management of depression in the elderly. Independent Drug Information Service; 2008: available at: [www.rxfacts.org](http://www.rxfacts.org).  
Evidence-based care guidance document  
Written for physicians caring for elderly patients, for use in academic detailing programs in Pennsylvania and available via Internet website for all interested clinicians. Academic detailing program extending to other states, including New York, South Carolina, Washington D.C., and other countries including Canada and Brazil.
3. Jackowski L, Avorn J, Choudhry NK, Fischer M, **Kesselheim A**, May F, Parikh S, Rowett D, Shrank W. Preventing falls and enhancing mobility in the community dwelling elderly. Independent Drug Information Service; 2009: available at: [www.rxfacts.org](http://www.rxfacts.org).  
Evidence-based care guidance document  
Written for physicians caring for elderly patients, for use in academic detailing programs in Pennsylvania and available via Internet website for all interested clinicians. Academic detailing program extending to other states, including New York, South Carolina, Washington D.C., and other countries including Canada and Brazil.
4. Jackowski L, Avorn J, Choudhry NK, Fischer M, **Kesselheim A**, Parikh S, Shrank W. Maximizing function in the patient with impaired cognition and behavior: What the primary care physician needs to know to help patients and caregivers. Independent Drug Information Service; 2009: available at: [www.rxfacts.org](http://www.rxfacts.org).  
Evidence-based care guidance document  
Written for physicians caring for elderly patients, for use in academic detailing programs in Pennsylvania and available via Internet website for all interested clinicians. Academic detailing program extending to other states, including New York, South Carolina, Washington D.C., and other countries including Canada and Brazil.

### **Thesis**

1. **Kesselheim AS**. A method to their madness: Greek Methodism in its social context. [Honors undergraduate thesis]. On file, Department of History of Science, Cambridge, MA: Harvard University, 1996.

### **Narrative Report**

I have established a program of research within the Division of Pharmacoepidemiology and Pharmacoeconomics at BWH and as a faculty member at HMS that combines the fields of medical practice, law and regulation, pharmacoepidemiology, and health services research. My work analyzes how

prescribing and other aspects of medication use – and their resulting clinical outcomes – are shaped by drug and device policies, laws, and ethical norms. This work has four interrelated areas of focus.

The first is studying how laws and regulations affect access to and use of therapeutic interventions, as well as drug approval and promotion. This work has led to grant funding from the Laura and John Arnold Foundation to develop empirical research on drug development and the effects of patents and other forms of market exclusivity on medication access, prices, and utilization. Another component of this work studies the role of biomarkers and other surrogate measures in FDA drug approval. The FDA has implemented several policy proposals related to our work through these grants, including a) expediting the review of generic drugs when there are 3 or fewer manufacturers in the field to enhance competition and control costs; b) increasing generic drug competition by issuing guidances on generic drug interchangeability for complex products soon after their initial approval; and c) allowing greater therapeutic substitution across drugs within the same drug class when clinically appropriate.

Second, drawing on my training as a patent attorney, I have studied the effects of market exclusivity on drug innovation, development and use. I have reviewed the impact of patents and legislative incentive programs including the Orphan Drug Act to analyze their strengths and weaknesses in contributing to the discovery and approval of new drugs. Through this work, we have documented the strategies used to delay generic drug availability, and described the role that Orphan Drug Act and other incentives play in the development, evaluation, and approval of new drugs. In work funded by an Investigator Award in Health Policy Research from the Robert Wood Johnson Foundation, I examined the origins and development of the most transformative drugs and devices of the past 25 years. By mapping patents and conducting interviews with key inventors, I described the roles played by academic and private-sector researchers in moving innovation forward, and defined the contribution of patents and other incentives to this work. My studies on the contribution of government-funded research to the development of transformative drugs has been widely cited in the national debate on the proper level of public funding of science in the US.

Third, I have analyzed the clinical, ethical, and economic consequences of regulatory decisions that are based on limited pre-approval clinical studies, and considered the implications for patients, physicians, and payors of making such drugs and devices widely available. This work has examined the increasing use of expedited drug development and regulatory review pathways in the US as well as issues in post-approval followup and the risk-benefit tradeoffs for patients that these products and procedures can pose. In 2013, I was selected to join the Greenwall Faculty Scholar program in Bioethics to study the ethical considerations involved in regulatory determinations about new medications. I have continued pursuing this work through the Program On Regulation, Therapeutics, And Law (PORTAL) that I developed within the Division, which now encompasses a team of junior faculty members, post-doctoral fellows, and students focused on this area and a \$1 million annual budget.

Finally, I have conducted empirical research into other intersections of public health, law, and medication use and outcomes, including showing that disclosures about funding directly influence the interpretation of clinical trial data, often counterproductively (*New England Journal of Medicine*, 2012), and how conflict of interest disclosure policies such as state and federal open payments legislation influence physician reporting and brand-name drug prescribing.

In recognition of the impact of my research, I have been invited to speak at numerous national and international meetings, and to consult for expert bodies such as the US Patent and Trademark Office and ClinicalTrials.Gov. In 2016, I was appointed to a committee of the National Academies of Science, Engineering, and Medicine and contributed expertise on prescription drug regulation to help shape recommendations on how FDA oversight of opioid medications can best promote public health goals. I currently serve as a Deputy Director of the HMS Regulatory Sciences Advisory Group, as a member of the *New England Journal of Medicine* Perspectives Advisory Board, as a faculty affiliate of the Harvard Law School Petrie-Flom Center for Health Law Policy, Biotechnology, and Bioethics, and as a core

faculty member of the HMS Center for Bioethics. In 2015, I was invited to serve on an FDA Advisory Committee and to join the Board of Directors of the American Society of Law, Medicine, and Ethics. In 2017, I was appointed editor-in-chief of its *Journal of Law, Medicine, and Ethics*.

Clinically, I practice internal medicine in the Phyllis Jen Center for Primary Care at BWH, where I manage a panel of primary care patients with a wide range of acute and chronic primary care problems. I have cared for many of these patients since my residency, and the ways that they have benefitted from new drug treatments, as well as struggled with issues related to drug costs and side effects, has inspired my work.

My administrative and institutional leadership has included several novel contributions to the BWH and HMS communities. The PORTAL program, which is among the largest independent research centers in the US focusing on drug policy issues, has attracted numerous talented trainees and faculty and is widely known as a center for expertise on drug regulatory science and policy. As an outgrowth of my PORTAL work, I have become a Deputy Director of the HMS Regulatory Science initiative. I established a monthly Policy and Ethics Consortium series at HMS in 2016 that attracts experts in the field to wrestle with challenging current health policy topics; we routinely receive 100-150 audience members from the community at each public session.

Finally, I have been committed to teaching throughout my career. As founder and director of PORTAL, I have been directly responsible for the oversight of numerous post-doctoral fellows, who have gone on to academic and government positions, as well as HMS students interested in prescription drug policy and law. I have consistently taught in the HMS Health Policy course as well as lectured on prescription drug policy issues in annual seminars for medical residents and fellows across the Harvard teaching hospitals. In 2015-2016, I originated a class on Health Law, Policy, and Bioethics for the HMS Center for Bioethics, and in 2016-2017 I initiated a monthly health policy and bioethics seminar for the entire Harvard community that is also offered for class credit for Bioethics Masters students. In 2014-2015, I was first invited by Yale Law School to teach a class on FDA law. Receiving top student reviews, I was re-appointed as Irving S. Ribicoff Visiting Associate Professor of Law in 2016-2017, 2017-2018, and 2018-2019. Because of growing demand, we doubled the class size and opened it up to cross-registrants from Yale Medical School and Yale School of Public Health.