

Updates on Federal Biosafety and Biosecurity Policy

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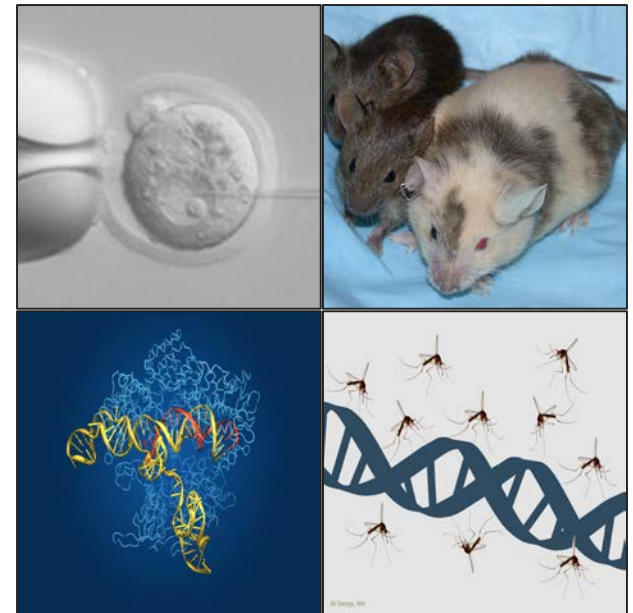
Emerging Biotechnologies: Opportunities and Challenges

An exciting and challenging time for science and policy

- Advances have dramatically altered the biotechnology landscape and hold great promise for innovation and discovery
- Some applications of new technologies are testing policies may raise questions about safety, security, and ethics

Areas of active policy discussion include:

- CRISPR/cas9 gene editing techniques
- Genome sequencing/synthesis
- Synthetic biology
- Gene drives
- Chimera research
- Human germline/mitochondrial modification
- Gain-of-function studies



Lab Incidents Prompt Renewed Safety and Security Efforts

The Washington Post

Health & Science

FDA found more than smallpox vials in storage room



The administrative building of the National Institutes of Health in Bethesda, Md. Hundreds of other potentially dangerous pathogens were in forgotten boxes on NIH campus. (J. Scott Applewhite/Associated Press)

By Brady Dennis and Lena H. Sun July 16, 2014

REUTERS

More: Reuters Health Reuters US

White House Orders U.S. Labs To Take Inventory Of Infectious Agents

BY JULIE STEENHUYSEN, REUTERS

AUG. 28, 2014, 7:20 PM | 26

GAO

United States Government Accountability Office

Testimony

Before the Subcommittee on Oversight and Investigations, Committee on Energy and Commerce, House of Representatives

For Release on Delivery Expected 10:00 a.m. ET Wednesday, July 16, 2014

HIGH-CONTAINMENT LABORATORIES

Recent Incidents of Biosafety Lapses

GAO

United States Government Accountability Office

Report to Congressional Requesters

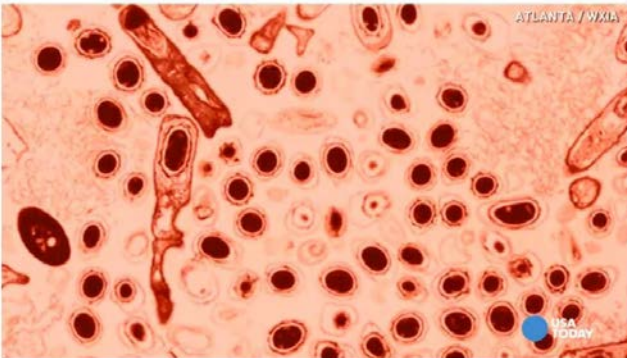
March 2016

HIGH-CONTAINMENT LABORATORIES

Comprehensive and Up-to-Date Policies and Stronger Oversight Mechanisms Needed to Improve Safety

Anthrax scare is latest safety lapse at CDC labs

ATLANTA / WXIA



The Center for Disease Control and Prevention in Atlanta says 75 workers may have been unintentionally exposed to anthrax. The CDC stressed that the public is not in any danger.

Michael Winter and Alison Young, USA TODAY 3:31 p.m. EDT August 6, 2014

Initiatives to Strengthen Biosafety and Biosecurity

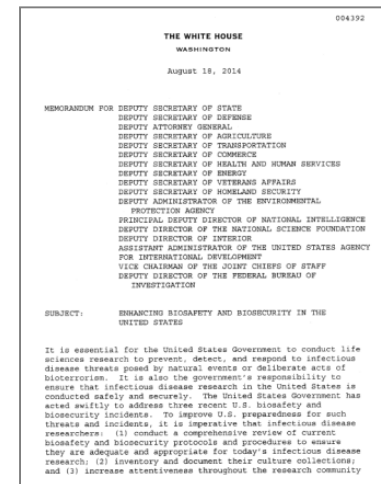
The U.S. Government has undertaken a series of complementary, short- and long-term policy activities aimed at reinforcing commitments to biosafety, biosecurity, and responsible conduct of research.

- **Biosafety and Biosecurity Stewardship**
- **Research Involving Select Agents and Toxins**
 - Federal Experts Security Advisory Panel (FESAP)
 - Fast-Track Action Committee on Select Agent Regulations (FTAC-SAR)
- **Dual Use Research of Concern (DURC)**
 - National Science Advisory Board for Biosecurity (NSABB)

Initiatives to Strengthen Biosafety and Biosecurity

Holdren-Monaco memo (Aug. 2014) called on Federal agencies to:

1. Conduct a comprehensive review of current biosafety and biosecurity protocols to ensure adequacy and appropriateness for today's infectious disease research
2. Inventory and document culture collections
3. Increase attentiveness throughout research community to ensure the safety of laboratory workers and the American public



National Biosafety Month

National Biosafety Month 2014

- Review policies, procedures/practice, and training
- Conduct inventories to ensure proper storage, containment, and documentation of infectious agents and toxins

National Biosafety Month 2015

- Enhance a top-down/bottom-up culture of safety and responsibility
- Promote training and transparency regarding institutional oversight

National Biosafety Month 2016

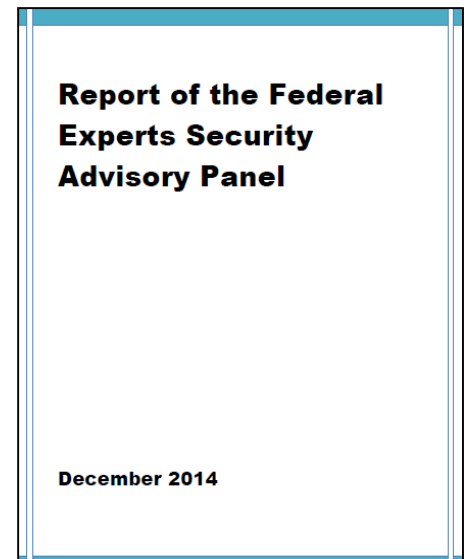
- Evaluate biosafety programs and related governance structures
- Collaborate to promote biosafety and share best practices
- Commit to providing appropriate resources to all elements of biosafety programs

Federal Experts Security Advisory Panel (FESAP)

FESAP examined policy issues relevant to the security of select agents and toxins to optimize biosafety, biosecurity, oversight, and inventories

FESAP Report (Dec. 2014)

- Concludes that, in general, the USG has a robust set of rules, regulations, and practices to inform safe, secure and responsible research with infectious agents and toxins
- Outlines recommendations to promote oversight and compliance for biosafety, biocontainment, and biosecurity; called for more applied biosafety research



Fast Track Action Committee on Select Agent Regulations (FTAC-SAR)

FTAC-SAR engaged with a wide range of stakeholders on the impact of the Select Agent Regulations on science, technology, and national security

FTAC-SAR Report (Oct. 2015)

- Outlines findings and recommendations based on stakeholder feedback
- Calls for strengthening inventory control, material accountability, outreach and education, sharing of best practices

**Fast Track Action Committee Report:
Recommendations on the Select Agent
Regulations Based on Broad
Stakeholder Engagement**

October 2015

**National Science and Technology Council
Committee on Homeland and National Security
Subcommittee on Biological Defense Research and
Development
Fast Track Action Committee on the Select Agents
Regulations**

Implementation of FESAP & FTAC-SAR Recommendations

USG is implementing FESAP and FTAC-SAR recommendations:

- Developing and optimizing biosafety and biosecurity policies and practices
- Strengthening oversight mechanisms
- Promoting outreach and education
- Enhancing the culture of responsibility

**IMPLEMENTATION OF RECOMMENDATIONS OF THE
FEDERAL EXPERTS SECURITY ADVISORY PANEL (FESAP)
AND THE FAST TRACK ACTION COMMITTEE ON SELECT
AGENT REGULATIONS (FTAC-SAR)**

October 2015

Gain-of-Function Research

NSABB led the deliberative process; developed recommendations for the evaluation and oversight of gain-of-function research involving pathogens with pandemic potential.

NSABB Report (May 2016)

- Central finding: Studies anticipated to enhance pathogens with pandemic potential have potential public health benefits but also entail significant potential risks
- Recommended additional, multidisciplinary Department-level evaluation prior to funding decision, and appropriate ongoing oversight if funded





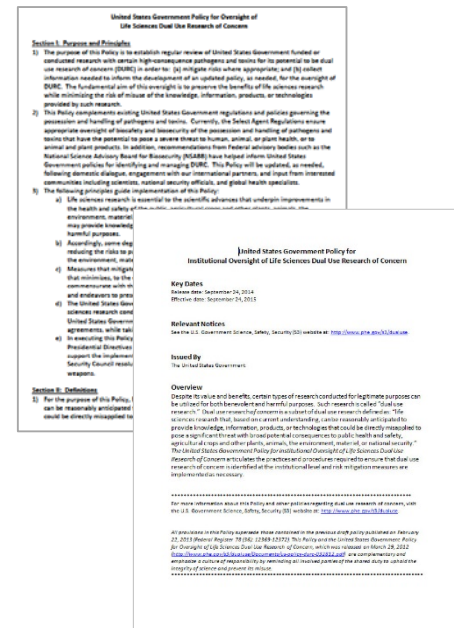
Looking Ahead: New Tasks for the NSABB

USG DURC Policies

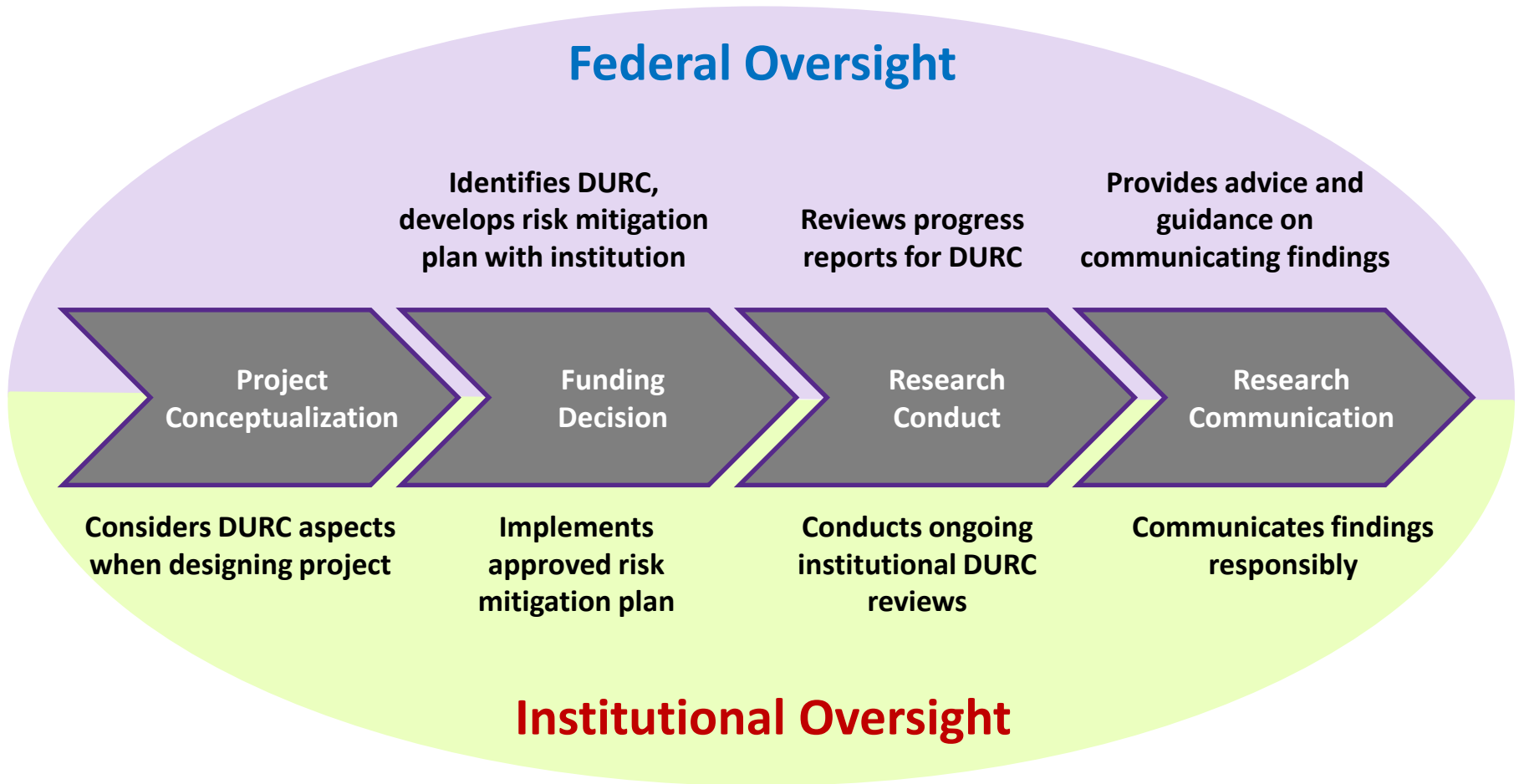
USG has released two policies for the oversight of dual use research of concern:

- **USG Policy for Oversight of Life Sciences DURC (March 2012)** – Addresses roles and responsibilities of Federal funding agencies; requires identification and oversight of DURC in research portfolios

- **USG Policy for Institutional Oversight of Life Sciences DURC (issued 2014; effective Sept. 2015)** – Addresses roles and responsibilities of USG-funded research institutions; requires agencies to establish a system for the identification and oversight of DURC



USG DURC Policies: Collaborative Oversight Throughout the Research Life Cycle



Evaluating the Institutional DURC Policy

Questions to Address

- What are the strengths and limitations of the institutional DURC policy?
- What challenges are institutions facing as they implement the policy?
- Are there best practices for identifying DURC, managing risks, training researchers, etc?
- What are the effects of the policy on scientific research and publishing?

NSABB New Task: DURC Policy Stakeholder Engagement

NSABB Task

To host a series of regional meetings to gather feedback from stakeholders implementing the U.S. Government Policy for Institutional Oversight of Life Sciences Dual Use Research of Concern

Anticipated deliverables

- A series of regional stakeholder meetings, video archived
- Meeting summaries describing stakeholder feedback

NSABB Working Group

- Chair: Joseph McDade, Ph.D.

NSABB New Task: DURC Policy Stakeholder Engagement

Objective

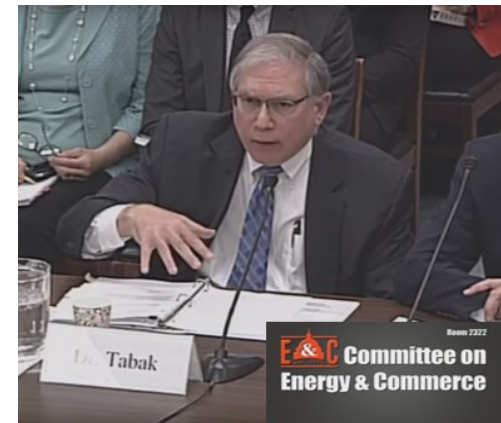
To obtain feedback from those involved in the implementation of the institutional DURC policy to inform USG policy review and development efforts.

Topic areas include:

- Experiences implementing the policy at the institutional level
- Challenges (and any steps taken to address them)
- Best practices and novel strategies for managing DURC
- Effective strategies for training investigators and laboratory staff
- Impact of the policy on researchers and institutions

NIH Conducting an External Review of 2014 Smallpox Incident

- **July 2014:** 327 sealed vials of pathogens and biological materials were discovered on a cold room on the NIH main campus; 6 vials contained variola virus
- Samples were promptly secured, transported to CDC; smallpox samples were tested and destroyed, witnessed by officials from WHO
- Incident has resulted in a number of reports/investigations and generated considerable interest from members of Congress
- NIH is conducting a review of this incident



Blue Ribbon Panel Reviewing 2014 Smallpox Incident

- NIH Director has appointed a Blue Ribbon Panel to review the incident and to report its findings to the NSABB
- **Blue Ribbon Panel is to:**
 - Review the incident and the immediate response
 - Consider whether systemic issues may have contributed to the presence of variola virus and to the delay in its discovery
 - Determine whether NIH policies have been appropriately strengthened, if necessary, in response to the incident
- **NSABB is to:**
 - Provide additional subject matter expertise and input to the Blue Ribbon Panel and a forum for public discussion of the issue

Resources

❖ NSABB

- Submit public comments to NSABB: nsabb@od.nih.gov
- Website: <http://osp.od.nih.gov/office-biotechnology-activities/biosecurity/nsabb>



❖ Science, Safety, Security (S3)

- <http://www.phe.gov/s3/Pages/default.aspx>



❖ NIH Office of Science Policy

- Website: <http://osp.od.nih.gov/>
- Blog: <http://osp.od.nih.gov/under-the-poliscope>
- Twitter: <https://twitter.com/cwolinetznih>
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Bringing Science Policy Into Focus