FROM BEGINNING TO END: AN EXAMINATION
OF AGENCIES EARLY PUBLIC ENGAGEMENT AND
RETROSPECTIVE REVIEW

HEARING
BEFORE THE
SUBCOMMITTEE ON
REGULATORY AFFAIRS AND FEDERAL
MANAGEMENT
OF THE
COMMITTEE ON
HOMELAND SECURITY AND
GOVERNMENTAL AFFAIRS
UNITED STATES SENATE
ONE HUNDRED SIXTEENTH CONGRESS
FIRST SESSION
MAY 7, 2019

Available via http://www.govinfo.gov

Printed for the use of the Committee on Homeland Security
and Governmental Affairs
OPENING STATEMENT OF SENATOR LANKFORD

Senator LANKFORD. Good morning. Welcome to today's hearing entitled From Beginning to End: An Examination of Agencies Early Public Engagement and Retrospective Review. I would like to first welcome Senator Sinema to the dais. I look forward to working with you in the Senate, and am grateful, specifically, for your work that you have done on these topics in the House, and coming over here and working on this in the Senate. I know the most coveted ranking position, is to be able to work on this Subcommittee, working on regulatory issues and Federal management. And what is wonderful is that you worked on these issues a lot in the House, and I am grateful that you have come over to be able to work on this here.

Thank you, as well, to our witnesses who bring a lot of expertise into this, and we are very grateful for the time that you have spent on it. Former Office of Information and Regulatory Affairs (OIRA) administrators are a very valuable resource to us in this Committee because you have walked through this process and you bring some unique insight.

I also jokingly say all the time I can ask former administrators questions that they can actually answer, that current administrators say, “I will have to get back to you with my legislative staff,” and note that. So we are grateful to be able to get the insight.

Today we are focusing on how agencies conduct public outreach at the beginning of a rulemaking, and how real success or failures are measured years later. While these could be considered narrow issues they are by no means small. Advance Notice of Proposed

---

1The prepared statement of Senator Lankford appears in the Appendix on page 25.
Rulemaking (ANPRM) are important tools that only a few agencies are statutorily required to actually utilize.

Administrative Procedures Act (APA) sets out a process where agencies propose a rule, listen to comments from the public. They have the opportunity to make changes and then issue a final regulation. While the system looks good on paper, in practice agencies typically conduct a significant amount of work before formally engaging the public. They will consider various regulatory schemes and conduct economic analysis. Ideally, agencies would engage the public early, but there is no Administrative Procedures Act requirement to do so.

Turning to the other end of the rulemaking process, retrospective review is a process to ensure rules achieve their intended goal in the least burdensome way. Over time, changing circumstances, improved technologies may render some regulations ineffectual or unnecessary. An agency’s job is not done after the final rule is published. As initial estimates of both costs and benefits prove inaccurate, agencies should revisit a rule to ensure the desired effects are actually achieved.

Every President since Jimmy Carter has urged agencies to utilized retrospective review to examine existing regulations. While these directives were issued with good intentions, they gave agencies a significant amount of discretion in selecting which and how many rules to review.

The focus of this hearing is two bills that Senator Sinema and I will introduce shortly, that codify the best practices for both procedures. The Early Participation in Rulemaking Act direct agencies to issue advance notices for rules costing more than $100 million annually. The agency must outline what problem the rule intends to solve and listen to the public’s input on the subject. The idea behind this bill is to require agencies to listen to the public before they craft the regulation.

Washington does not have all the answers. Taking time to work with stakeholders, particularly our small businesses, is vital in crafting effective regulations. Less burdensome is not less effective. Business owners want to be good citizens, follow along, and have a safe and clean workspace.

Setting Manageable Analysis Requirements in Text (SMART) Act, is a retrospective review bill that looks ahead. It requires agencies to set metrics for how a rule will be measured for success in the future. It is hard to imagine the measure of success of anything unless it is defined. This bill instructs regulators to define what success is for a given rule and then requires them to grade that rule within 10 years.

This Subcommittee has been working on both of these issues for a while. Both bills have bipartisan support in the past two Congresses. I look forward to working with my colleagues to push both of them across the finish line.

With that I would like to recognize Senator Sinema for her opening statement.
OPENING STATEMENT OF SENATOR SINEMA

Senator Sinema. Well, thank you, Chairman Lankford, and thank you to our witnesses for joining us today to help fix our regulatory process.

I am happy to join Chairman Lankford as the Ranking Member of the Regulatory Affairs and Federal Management Subcommittee. As a Congresswoman I promoted policies that expanded business opportunities and fueled innovation in Arizona and beyond, and my goal for this Subcommittee is to continue that work and to find and push for targeted common-sense reforms to regulations that help hard-working Arizonans build better lives.

With Chairman Lankford’s leadership we are off to a strong start. We have already succeeded in moving two common-sense regulatory transparency bills through committee markup, and that is only the beginning.

The Providing Accountability Through Transparency Act and the GOOD Act will both help make government more accessible to Arizona businesses, and I am excited to accomplish even more.

Today we continue the work of advocating for a modernized rule-making process. Advance notice of proposed rulemaking allows agencies to engage with businesses, nonprofits, academics, and other everyday people, so an idea that may later become a rule is heading in the right direction.

Retrospective review requires agencies to look back at a rule, to make sure it does what was intended and that the benefits outweigh the costs. Doing a retrospective review can be time-consuming and complex, but if an agency prepares in advance for the review it can be even more beneficial and efficient.

Advance notice of proposed rulemaking and planning for retrospective review have been encouraged for decades but never enacted into law, and that should change. And soon I will be introducing the SMART Act to incorporate planning for retrospective review into major rules.

Planning for the future is common sense, which is why planning for retrospective review has been promoted by the Government Accountability Office (GAO), the Administrative Conference of the United States (ACUS), and the American Bar Association (ABA). By requiring agencies to plan for review, the reviews will be more thorough and accurate, and less expensive and time-consuming. Our legislation will improve regulations, remove unnecessary burdens, and increase transparency and accessibility for Arizona businesses, communities, and others, and I look forward to hearing from our witnesses.

Thank you, Mr. Chair.

Senator Lankford. Let me proceed to testimony from our witnesses. The Hon. Susan Dudley is the Director of the Regulatory Studies Center and Distinguished Professor of Practice at the Trachtenberg School of Public Policy and Public Administration at George Washington (GW) University. Before joining the faculty at GW, she served as the Administrator of the Office of Information and Regulatory Affairs from 2007 to 2009.

The Hon. Sally Katzen is a Professor of Practice and Distinguished Scholar in Residence at New York University School of Law. She served in multiple roles during the Clinton Administra-
tion, including Administrator of the Office of Information and Regulatory Affairs, Deputy Director for Management of the Office of Management and Budget (OMB).

Thank you both for bringing your experience and your insight, for working with our staff leading up to this hearing, and continuing to contribute to your Nation. Thank you for that continued engagement.

As you both know, because you have both been here before, it is the custom of this Subcommittee to swear in all witnesses before they testify. So if you would please stand and raise your right hand.

Do you swear that your testimony given before this Subcommittee will be the truth, the whole truth, and nothing but the truth, so help you, God?

Ms. DUDLEY. I do.

Ms. KATZEN. I do.

Senator LANKFORD. Thank you. You may be seated. Let the record reflect both answered in the affirmative.

We are using a timing device today, and you will have 6 minutes for your opening statement. We do have a mercy rule here that if you go beyond that you are fine, because we have this hearing to get your testimony, and so if you go a little bit over we are going to be OK on both of those.

But we will be glad to be able to receive your testimony now. Ms. Dudley, you are first.


Ms. DUDLEY. Thank you. Do I get bonus points if I go under?

Senator LANKFORD. Yes, clearly.

Ms. DUDLEY. Thank you very much, Chairman Lankford, Ranking Member Sinema, Senator Carper, and Senator Scott for inviting me to talk about one of my favorite subjects. I appreciate your interest in improving how the U.S. Government develops and evaluates regulatory policy. You are continuing a long bipartisan tradition of efforts to make regulation well informed, transparent, and accountable to the American people.

Agencies have long been required to seek public comment on proposed regulatory notices, yet these opportunities for public engagement often come after agencies have made key policy decisions. Proposed rules are legal documents, written to defend a selected approach against possible litigation. So this motivates agencies to circle the wagons, narrowing the menu of alternatives and the evidence they consider before the public has an opportunity to engage.

The draft Early Participation in Regulations Act would require agencies to issue for public comment advanced notices of proposed rulemakings for major rules. This could free them to share their early thinking on whether a problem requires a regulatory solution, what objectives could be achieved, and what different options are available.

1 The prepared statement of Ms. Dudley appears in the Appendix on page 28.
These ANPRMs and any differences between them and subsequent APA rulemaking steps are wisely exempt from judicial review in your draft bill. One virtue of the ANPRM is that it provides an opportunity for agencies to share their preliminary thinking about a problem and get input on potential solutions at a stage when they are truly open to feedback, analysis, and evidence. If agencies had reason to fear that this early notice could later be used against them in court that would discourage objective queries and underline those benefits.

I do not think the ANPRM requirement would significantly slow agencies’ rulemaking. For one thing, 90 days is not a long time considering that agencies often take years studying a problem and evaluating options before they issue that first proposal.

But probably more importantly, to the extent the ANPRM invites comments on preliminary deliberations that would otherwise have taken place behind closed doors, it may make the overall regulatory process more efficient. So rather than tacking 90 days on at the end of a rulemaking—or at the beginning, that makes it 90 days longer, it may provide valuable input that ends up streamlining the subsequent notice and comment process. In many cases, early engagement could lead to more efficient analysis at the proposal stage and fewer surprises during public comment.

That said, there are cases where an ANPRM would not serve the public interest and bill provides for exceptions for those.

Retrospective review is also very important. Program evaluation has a long tradition in the private sector and in activities financed through the fiscal budget, but it has received little attention in the regulatory arena, even though every President since Carter has asked agencies to evaluate existing rules. President Obama added new emphasis on evaluation, yet most regulations continued to be issued without a plan for review.

As ex ante analysis, regulatory impact analyses (RIAs) are necessarily hypotheses of the effects that regulations will have if implemented. Better regulatory evaluation would allow us to test those hypotheses against actual outcomes. That feedback would not only help with decisions on current policies but it would improve future RIAs and future regulations.

Your draft bill would require agencies to include in major regulations a framework for how they will measure effectiveness, benefits and costs, and a plan for gathering the information necessary for ex post review. It would also require agencies within 10 years, to assess a rule’s benefits and costs, evaluate how well it accomplishes its objectives, and determine whether it could be modified to achieve better outcomes.

The draft bill focuses not just on reducing regulatory burdens but on improving outcomes by subjecting rules to rigorous evaluation and feedback. It could create an evaluation mindset where agencies learn from reviewing regulatory actions and apply those lessons to improve future rules.

These two draft bills offer relatively modest yet potentially powerful changes to the rulemaking process. By engaging public input earlier in the process and providing for retrospective review of regulations to evaluate whether they are achieving their objectives, they can help ensure that regulations are based on the best avail-
able evidence and that they are working as intended. These bills
could make regulatory decisions more transparent and accountable,
leading to improved outcomes for the American people.
Thank you.
Senator LANKFORD. Thank you. Ms. Katzen.


Ms. KATZEN. Thank you, Chairman Lankford, Ranking Member Sinema, Senator Scott, and Senator Carper. Thank you for inviting me to testify today.

I have been generally skeptical, if not highly critical, of the many attempts over the last several decades to rewrite the APA and update the process that produces the regulations that translate general statutory directives into concrete requirements that the public must comply with. Many of those attempts were highly partisan and would have converted the regulatory roadmap into an obstacle course, with the effect, if not the intent, of greatly delaying or shutting down the regulatory process rather than contributing to good decisionmaking.

I understand that the effort of this Subcommittee is different. I understand that you are looking for surgical fixes to improve discrete problems with the goal of enlisting support by both Democrats and Republicans, and specifically you are looking at the beginning and the end of the process with suggestions that could be achieved by legislation, by Executive Order (EO), by OMB guidance, by agency practices.

But regardless of the vehicle, it is important to be clear about what the problem is and how to best solve that problem without introducing unintended consequences.

So at the beginning, the first official step in a rulemaking is the issuance of a notice of proposed rulemaking (NPRM), lawyers in practice and the Academy generally agree that by the time the agency issues the NPRM, the staff involved have invested so much time and energy in developing the proposal and supporting data as they are required to do—and analyzing the likely effects of the proposal—as they are required to do—and justifying the proposal—as they are often called upon to do by agency decisionmakers and during OIRA review, that they are virtually locked into their proposal and are less receptive to new ideas or even significant modifications of their proposal.

While all of the up-front work is desirable, it often has the unintended consequence of restricting the options going forward. As you mentioned, both Republican and Democratic Presidents have tried to counter this tendency by encouraging agencies to consult with the public even before they have made the decisions reflected in the NPRM. Those admonitions have produced some additional outreach to the public, but the effect has been inconsistent and less productive than had been expected.

1The prepared statement of Ms. Katzen appears in the Appendix on page 36.
Now several agencies do use an advance notice of proposed rule-making to solicit ideas at the outset, but it is not a universal panacea. It is especially useful when the agency is unsure what direction to take, what data to consider, how prescriptive to be. It is also useful when it is done before or at the outset of agency deliberations.

It is less useful when the authorizing statute is itself prescriptive or there is genuine consensus about what is needed to respond to the identified problem. In short, it can be helpful at times, but at other times it may just add an unproductive but a time-consuming step to the already extended process.

For this reason it is important that any requirement for an ANPRM be limited to economically significant—or you call them major regulations, that are required to use notice and comment, and second—that any such provisions not impose on the agency multiple requirements for explanations, analysis, data, etc. The purpose would be to engage those affected by the rule so they can contribute to its development and formulation before the agency settles on a particular course, not to lock the agency into a particular mindset before the process begins. The more the agency has to incorporate in an ANPRM, the more the agency will become invested in a particular outcome. This is the opposite of what it should do.

With respect to retrospective review, for almost 40 years there have been concerns that there are too many rules, and so many of the rules in the books are obsolete, burdensome, unworkable. Notwithstanding the efforts of every President, from Reagan through Obama, we search and we search, and we do not find, and we do not eliminate many rules from the existing stock of regulations.

Now one reason for this may be that since 1980, new regulations are not issued unless their benefits justify their costs. To eliminate such regulations would likely mean that the costs of rescinding the regulations would be greater than the benefits, which is counter-intuitive. Other reasons for the limited success of look-back efforts are that agencies usually have not collected data along the way that would inform their retrospective reviews, and very importantly, any respective analysis requires resources, and for at least the last several decades regulatory agency budgets have generally been decreasing or straight-lined. Without the resources they cannot do the work.

Nonetheless, there is growing support for the step that you are talking about here, namely encouraging agencies to plan for retrospective review when they are in the process of developing a final rule. This idea was endorsed by many of the organizations that you mentioned, and a report prepared by the Institute for Policy Integrity, which reflected the unanimous recommendation of almost all former OIRA administrators.

Requiring agencies to provide a plan for later retrospective review of a newly issued rule would, in most instances, be salutary. If nothing else, it would require the agency personnel to focus on describing precisely what they want to accomplish and how to evaluate whether or not the rule is successful, at a time when the rule and its alternatives is foremost in their minds.
It is very important, however, to provide flexibility for the eventual implementation of the retrospective review. The agency can and should commit to a framework in the proposed and final rule, identifying the data and the metrics it anticipates using for that purpose, but this should not be cast in concrete. We learn a lot with time, including how to better analyze and measure what is going on.

This is amply demonstrated by the increased sophistication of cost benefit analysis itself over the last decade or two, and it is also demonstrated by the general preference for performance standards, which specify the desired results, rather than design standards, which lock in a particular way of getting there.

In addition, while periodic review is useful, there will likely be some, and maybe many situations where repeated retrospective reviews would yield greatly diminishing returns. After a decade or so, if rules survive a retrospective review intact, they are likely to have established their worth, and it would be wasteful to continue retrospective review after retrospective review. Some escape hatch should be provided.

I thank the Committee for its efforts and for its courtesy in allowing me to exceed my time limit. Because you are not writing on a blank slate, even this limited surgical approach will likely face an uphill battle. It might well, however, have a better chance to succeed than many that have been tried before. And for that I congratulate you both.

Senator LANKFORD. Thank you. We will take that congratulations when we are done. We are in the starting block at this point, though.

Thank you both for your testimony. I am going to defer my questions to the very end and recognize Senator Sinema.

Senator SINEMA. Thank you, Chairman. Thank you both, again, for being here today.

Starting with Executive Order 12866, which directs agencies to make sure that regulations maximize benefits while limiting costs, Administrations have discussed the merits of retrospective review and have provided guidance to agencies related to the review of existing regulations, but, of course, these directives do not have the force of law.

Over the years, as I mentioned in my opening statement, the GAO, the Administrative Conference of the United States, and the American Bar Association have all issued reports or made official comments regarding the need for post-enforcement analysis of the true costs and the benefits of regulations.

So the question for both of you is considering the long-term support for retrospective review from the Clinton and all subsequent Administrations, why do we believe that this continues to be a challenge for agencies in our Federal Government?

Ms. DUDLEY. I would say there are two reasons, lack of incentives, and then lack of data and analytical tools. I think your draft bill addresses both of those, and when I say lack of incentives for agencies—it is always more exciting to look at the next thing, the next problem to solve rather than to look back at what you have done. But it is not just the agencies because regulated parties also,
once they have complied, they are not so keen on revisiting it and
maybe making their competitor upstart not have to comply.

So by requiring this in your bill, I think you would address the
incentives but perhaps more importantly you would address the
lack of data and analytical tools. As you have said, this bill would
require agencies up front to say here are the data we need, here
is how we will collect it, and here is the outcome that we are going
to look to measure it against.

Ms. KATZEN. I agree with her response, as I often agree with
things that she say. I would just simply add the lack of resources.
It takes time. It takes money. And while you can frame it in terms
of the new guys want to do their thing, it is also fair to say that
relooking and relooking and relooking may not be as productive.
But it certainly drains resources away from what might be the cur-
rent need to address a pressing problem of health, safety, environ-
ment, open competition, whatever.

And so you are dealing with a situation where you have asked
an agency to do this and that and the next thing, and you may not
have always given them enough resources to do that.

Senator SINEMA. Based on current OMB guidance, agencies are
expected to understand the impacts of proposed regulations while
they are still under consideration, but OMB's own documents re-
lated to the realized costs and benefits of regulation allow for a
very wide range in estimates, creating ambiguity when agencies
are estimating future impacts.

So our legislation, the SMART Act, requires the solicitation of
data from regulated entities to ensure that the regulation meets its
objectives. How will the data that is collected to build retrospective
review into rules help create efficiencies at agencies when it comes
to creating new regulations or new major rules in the future?

Ms. KATZEN. I think there are two aspects. One, as Susan men-
tioned in her opening statement, as you do retrospective analyses
you learn whether the methodology you have been employing in
doing your ex ante analysis is actually flawed in any way or could
be improved in some way. So that helps the agency in thinking it
through.

It also alerts the regulated entities that they had a hand in this.
They have a joint responsibility. They are going to have to come
up with the data that show what the costs truly are. It is an urban
myth that so many of the rules that we establish, at the time we
say it is going to cost kazillions of dollars, and it turns out not to
cost quite that amount of money. We will now be able to gauge that
on a real-time basis with the kind of information that you are call-
ing for.

But one of the critical dimensions of whether a rule achieves its
objectives is whether the rule is complied with, and that is up to
the regulated entities, in large part. There is not a cop on every
corner. Most businesses do want to comply. They want to know
what the rules are and they will follow them. But it is not uni-
versal, and unless you have compliance data, you are not grappling
with one of the key conditions.

Ms. DUDLEY. I agree with Sally. I think it is not just the compli-
ance costs that you will be gathering through that process, and
that is one of the things that I really like about this draft bill, in
particular, is that it focuses on the cost, the benefits, and the outcomes. So it is a lot of information, and if agencies do not say what is expected, as Senator Lankford said in his opening statement, up front, how will we be able to measure against that?

Oh, the other thing, I think, that is valuable in your rules is the definition of major. It includes more than just the $100 million threshold. It talks about effects on competition, health, safety, etc. Those are all things that agencies should be measuring in this planning for retrospective review, and I think that is all valuable.

Senator Sinema. Thank you. I often hear from Arizona businesses that they are not opposed to regulations but they feel that the opportunities to comment during the rulemaking process are often for show rather than for substance. This is partly due, of course, to the expectation that a draft regulation, as provided in a notice to propose rulemaking, already has a selected course of action, and it usually includes documentation and data to support that coming decision.

So my question for you is how will the Early Participation in Regulations Act improve agency interactions with businesses, communities, and other regulated entities to help shape the process?

Ms. Katz. Well, it sends a message that early communication is important, that information that comes at a very early stage will be received by the agency and understood by the agency. And it does that, actually, in a number of different ways, but probably the most important is if it is done early. If you wait until you have crafted the NPRM, and then send out an ANPRM, it is useless. It is sort of like doing a cost benefit analysis after you have already decided on the approach you want to use. You should do it at the outset, when you are considering all the alternatives.

So here I would urge that you indicate—not in that statute, because it is just too much concrete—but generally suggest their agencies think about doing an ANPRM when they first get started, when they first send off notification to the Unified Agenda and say, we are about to have a rulemaking. We have not settled on a course yet. And if that message goes out, we hope it will be received and taken advantage of.

The other thing, for all regulated entities and regulatory beneficiaries, is it is possible to talk to the agencies throughout the entire process. There are ex parte rules at some agencies, but not at most. And it is sometimes difficult to move someone off the mark, but at the same time you should not be shy about having your voice heard and your points made. That is an important part of the interaction that agencies are comfortable with and should be used.

Ms. Dudley. I think that is true. I think it is important that it is not just businesses that will have an opportunity to get involved but others with information that may be relevant. And, in fact, I think that is what is really significant about the ANPRM idea. Stakeholders that have connections are involved at early stages in the rulemaking, working behind the scenes with agencies. What this does is it opens it up so that others that may have interesting, creative, new insights can get involved.

I also, as I mentioned in my opening remarks, the fact that it is not judicially reviewable, agencies really will feel much more com-
portable getting a wide range of thoughts and really thinking broadly about how to solve the problem at hand.

Senator SINEMA. Thank you. Thank you, Mr. Chair.

Senator LANKFORD. Senator Carper.

OPENING STATEMENT OF SENATOR CARPER

Senator CARPER. Thanks, Mr. Chairman, I want to commend you and the Ranking Member for holding this hearing. This is an issue of real importance. You can tell by the number of people who have packed this hearing room and the number of television cameras that are just waiting outside to interview everyone who has been a part of this hearing.

You have picked a couple of witnesses here. You could not have picked better witnesses. The balance of having Susan Dudley here and Sally Katzen shows a lot of wisdom, so thank you for that.

I would like to say everything I do I know I can do better. I think the same is true of all of us, and I think it is probably true of most of our laws and rules, including the way we create rules and regulations. So thank you for coming and joining us here today.

The last thing I am going to ask you is where do you think that you agree on some points that are most important, and where do you think maybe we do not agree on some things that are most important, in terms of what we are discussing today?

But the last several years this Subcommittee has focused on how we can work to improve the regulatory process. I have served on this Committee for 18 years. I love this Committee, and that was before it was—Homeland Security, the department was created. So it is a great Committee and this is an important part of what we do.

But the regulatory process is not perfect. No one is pretending that it is. And the Administrative Procedures Act and subsequent Executive Orders provide us with a robust roadmap for ensuring that agencies are promulgating rules fairly and we hope efficiently. The process also ensures that costs to industry are thoroughly considered and that the public has opportunities to be heard in the process.

However, under the Trump administration we have seen an unprecedented number of regulatory rollbacks, and there has also been unprecedented effort to skip steps in the normal process for considering whether the costs of regulations outweigh the benefits that protect consumers, that protect workers, protect our health, and environment.

I am especially concerned with the Office of Information and Regulatory Affairs' lack of oversight of agency rulemaking during this Administration and with the office's unwillingness to cooperate with oversight watchdogs who have legitimate requests for information.

For example, just last week, the Environmental Protection Agency Inspection General (EPA IG) sent a rare notification to Congress saying that the White House Office of Information and Regulatory Affairs was refusing to cooperate with an audit that Senator Udall and I had requested about EPA's proposal to exclude high-polluting glider trucks from emission rules. Glider trucks are not trucks that glide especially well. They are old diesel trucks that pollute a lot,
and they are cloaked in the shell of a bright, shiny, new truck. And the idea is that they are going to be able to continue to pollute. One of the greatest sources of pollution in our environment, for air pollution, is diesel engines, especially those that produce black carbon, which is far more dangerous to us and our environment than regular carbon dioxide.

But, Ms. Katzen and Ms. Dudley, do either of you recall a time when OIRA refused to cooperate with an agency Inspector General or GAO? Do you recall a time?

Ms. DUDLEY. Certainly during my tenure there was never a request from an agency IG of OIRA, and I think it may be unprecedented.

Senator CARPER. Alright.

Ms. DUDLEY. So that means I do not remember——

Senator CARPER. Yes. OK.

Ms. DUDLEY [continuing]. Not refusing, but I also do not remember it ever having occurred, such a request.

Senator CARPER. Thank you. Ms. Katzen?

Ms. KATZEN. In my recollection it did not occur where OIRA did not comply with a legitimate request from GAO, from the Congress, or from an Inspector General. I think there may have been one but I am a tad fuzzy on that.

Senator CARPER. OK.

Ms. KATZEN. To say “no” to a legitimate oversight body I think is unprecedented, to use Susan’s word.

Senator CARPER. OK. That is fine. You can hold it at that. Thank you.

OIRA seems to be failing to ensure that the EPA is conducting the cost benefit and other analyses that the law requires. I am going to just briefly describe a couple of examples, if I could.

First, during the interagency review of their proposal to exclude high-polluting glider trucks from emission rules, that I just mentioned, it was pointed out that EPA should have done more analysis because the rule was classified as, quote, “significantly, economically significant.”

Instead of requiring the analysis to be done, on the day before the rule was signed, OIRA allowed the rule to be reclassified so that analysis would no longer be required.

Second, when the EPA proposed its repeal of the clean water rule, it did not do a new cost benefit analysis to justify the rollback. Instead, EPA just deleted the benefits column of the Obama cost benefits table and OIRA allowed the proposed repeal rule to be published.

Finally, in the EPA’s proposal to remove the legal underpinnings of rules to reduce emissions of mercury and other air toxics from power plants, OIRA allowed the agency to use the agency’s old projected cost of compliance, that were three times higher than what the industry actually spent to comply with the rules and ignored the full benefits of that rule.

And, if I could, I would just like to ask both of you, and I do not expect you to be experts on any or all of the examples that I have just cited, but when each of you were privileged to run OIRA, would you have agreed to authorize the release of proposed rules that failed to perform a credible cost benefit analysis?
Ms. Katzen.

Ms. KATZEN. I would have fought hard to live up to what I thought the office stood for, which is good analysis, good data, and the kind of approach to rulemaking that produces good decisions. Having said that, I want to just add two qualifications. One, I am not there now, and if you are not there, it is sometimes very difficult to know what pressures are being exerted, who is saying what, who is doing what. In terms of your specific examples, however, I could give you three or four more.

I am an OIRA supporter. I am an OIRA booster. It was with great difficulty that I actually made an address earlier this year to the American Bar Association Ad Law Section criticizing OIRA, and saying that I thought it was not doing enough to ensure good analysis and being faithful to the cost benefit regime that it has guardianship of.

There are many disturbing stories. By the same token, when I was in the government I read stories that were not quite right. And so it is not fair, necessarily, to say that something is smelly in Denmark. It may well be, and I am concerned, as well.

Senator CARPER. OK. Thank you. Thank you.

Ms. Dudley, this is not a trick question. I am not trying to put you on the spot. I think you know me. Both of you know me pretty well. But when you ran OIRA would you have agreed to authorize the release of proposed rules that failed to perform a credible cost benefit analysis?

Ms. DUDLEY. I liked Sally’s comment that I would have fought hard, so I agree. I would have fought hard. It is important to understand that OIRA does wear several hats. One of them is interagency coordination. Another one is the analysis, the requirements of Executive Order 12866. And then the third is that they are part of the White House, and so sometimes those are in conflict.

So in my experience there—I remember times when we held our nose and said, alright, but before you get to the final stage you must do an analysis and put teed up for questions the things that the agency thought they needed that would allow them to do the analysis.

Senator CARPER. Good. Thank you for those responses. Mr. Chairman, thank you for being generous with the time and giving us a chance to actually have a conversation.

Senator LANKFORD. That is great. Thank you.

Thank you again. I am going to go ahead and say my questions now as well. I appreciate very much you coming through this process. Help us understand the inside of the machine. How long does it take to develop a rule and the economic impact, when you are talking about something that may be $100 million of economic impact? How much time is spent on the research side of that for the staff? And I know it is going to depend from rule to rule and agency to agency, but give me your ballpark guess.

Ms. KATZEN. Well, you are right that it will vary, depending upon the seriousness of the issue, depending upon whether it is a new issue or something that is being revisited, whether there is already expertise, or whether it has to be newly learned.

It can take a year or 2 years to do the background research. It can also be done in 4 or 5 months. You get a lot of help from the
outside sometimes, and a lot of obstruction from the outside sometimes. The whole process can take, as Susan said, years, occasionally.

Senator LANKFORD. Susan.

Ms. DUDLEY. It is hard to know exactly how long it takes, from the time an agency starts to think about the rule and the proposal. I can think of some that have been 10 years. Others can be shorter. There was a study that cite in my testimony that said, on average, 5.2 years is how long it takes before the proposal is issued.

Senator LANKFORD. So certainly 90 days is not too much time to be able to block up to get additional input. My thought on this is how to help agencies, OIRA, the future, to be able to think about, when is the moment to be able to put this out here? If there is going to be an advance notice of proposed rulemaking they are going to think about it for a while, maybe a year even. They are doing some of their own research and then they decide we may be headed toward a rulemaking here. This is a lot of information and it is all directed toward the same spot.

Where is the appropriate moment for them to go drop out an advance notice of proposed rulemaking to make sure that the insight in the information is coming at the right time, rather than, again, they form everything they need to do over 2 years, or 5 years of time, and 90 days before they plan to do it anyway they just drop this out there because it is formality?

Ms. KATZEN. Well, I mentioned in my testimony that I think it would be useful to issue an ANPRM as soon as the agency sends its first submission to the Unified Agenda. This is the document that is supposed to be available—well, it is available to the public. It is supposed to be a menu of everything that the agencies are thinking about doing. When they get to the point of saying, “We are thinking about doing a rulemaking on this,” that would be an appropriate time an ANPRM for for some rules. Again, I do not want to see too much of this cast in concrete——

Senator LANKFORD. Right.

Ms. KATZEN [continuing]. And prescriptive. But I think once they know they are heading down that path, putting out an ANPRM would be helpful.

Ms. DUDLEY. I think that is right. I think the first note is in the unified agenda, soon thereafter. That notice means we are thinking about this within the next 6 months. That would be a good time to do it.

I also would hesitate to see the bill be too strict about that, because different agencies have different practices. But I think the way that you have put OIRA in charge of that, or OIRA as the coordinator for the ANPRM bill, that would be something that agencies could work with OIRA on.

Senator LANKFORD. And again, the challenge that we have is we do not want to be prescriptive, because we cannot look in the future 10 years and we cannot see every rule and every entity. But the goal of this is to be able to get more dialogue early, not for the agencies to finish all their work and then go through the formalities.

We are not trying to make it longer, or, as you mentioned, creating an obstacle course here at the end. We are trying to get dia-
logue early on, and if the dialogue is too late in the process it does not matter. We are still in the same spot. But if it is too early, we do not even know if it is a $100 million rule at that point. We really do not see it.

So there has to be some level of information even to note, hey, I think this may be headed toward a major rule. They at least have to have enough information there to know what questions to ask, the size and complexity of it. This is a health and safety issue that is coming. We see some of the things that are going to be around it so let’s get more information out there. Does that make sense?

I guess I am asking a clarifying question. Are we hitting the right balance that we are not just creating an obstacle but we are getting this early enough? Should we be more prescriptive on when to be able to put this, or do we leave it at the discretion of OIRA, at a future OIRA, to say you did not do this ANPRM early enough and now we have a battle in the same process?

Ms. Katzen. I would not be prescriptive. When I was in the government, in the Administration, I took committee reports seriously, and I think to make clear your interest in having this done earlier rather than later in the committee report would be a very useful way of sending the message. But I would not cast it into legislative language.

Senator Lankford. OK. Fair enough.

Ms. Dudley. I agree.

Senator Lankford. OK. So let me ask you an even sillier question. Advance notice of proposed rulemaking just gets all of us excited here, as a term. Some agencies have just said request for information, to make it more generic and blunt. There have been a lot of different terms they have used, to basically accomplish some of the same things.

Is this the right term? The advance notice of proposed rulemaking has been used before. We have it in statutory language. Is that the right term to be able to even use on this? If we want to take off our D.C. hat and to say if someone outside of D.C. gets an advance notice of proposed rulemaking they have no idea what it is. If someone gets a request for information everyone knows what it is. Is one better than the other?

Ms. Dudley. Even though it is a mouthful I think advance notice of proposed rulemaking is the best—is a good way to say it. Agencies can always change something that they might have called a notice of inquiry before and just change the name.

One thing that I think is valuable about it, that you would not want to lose, you would not want to make it so loose that agencies could just reach out to specific stakeholders.

Senator Lankford. Right.

Ms. Dudley. You do want this to be something that——

Senator Lankford. It needs to be broad.

Ms. Dudley [continuing]. Is public, yes.


Ms. Katzen. And again I would refer to the committee report to clarify that an advance notice of proposed rulemaking, by any other name, would still accomplish the same objective. And you could call it a request for information. You could call it a request for pro-
posals. And again, that would send a message that I think would be helpful.

Senator LANKFORD. As long as it is public and as long as it is broad, because one of the challenges that we have is small business, large business, individuals, outside groups, environmental groups, think tanks, whatever it may be, they all need to have access to be able to submit ideas to it because they are all going to have different ideas and thoughts.

OK. So let me ask this. Are we overcomplicating this by adding another step with the advance notice of proposed rulemaking? Is there a way to be able to fix the notice of proposed rulemaking and to try to get more input from the public earlier in that process without having to do a formal process? I am willing to try to say let us go back and do it, because right now that process is so late in the process, or that product is so late in the process that a lot of decisions have already been made, and it is just tough to be able to engage with people.

As I think both of you mentioned in your testimony, once folks have done their own research, bounced their own ideas, have gone through the obligatory, working with all their bosses, explaining why this is the best idea, people get more and more entrenched. So what I am trying to figure out, is there a way to fix that process rather than add a new one?

Ms. KATZEN. I think we had decades of court cases that have spoken to what needs to be in the NPRM itself, and how much the final rule can deviate from the NPRM. I am referring here to the "logical outgrowth doctrine" that says that if the agency changes its mind between the NPRM and the final rule, in a dramatic fashion, that may well have deprived the commenting public of an opportunity to comment on the final product——

Senator LANKFORD. Right.

Ms. KATZEN [continuing]. And, therefore, it has to go back to the agency. And with 20, 30, or 40 years of cases having been built up, I would reluctantly say add a new step at the beginning rather than try to change the NPRM at this point in the face of those cases.

Senator LANKFORD. OK.

Ms. DUDLEY. I agree. I mean, I think it is the judicial reviewability of the NPRM that really forces agencies to have chosen what they want and defended that in a way that they are discarding other options.

Senator LANKFORD. OK. Let me switch to the end, if I can, here. OIRA has the unique opportunity to be able to see multiple agencies simultaneous, and when we talk about retrospective review some of the issue is other agencies have now formed regulatory processes, or there is something new in another entity that maybe this regulation is not as pertinent anymore because this entity has created something that is also doing something similar, or whatever it may be.

How can we help OIRA continue to be able to see all agencies, and when a retrospective review happens it is not just isolated to a particular agency but it still has that broad review? Is there anything that we need to build in?
Ms. DUDLEY. More staff. I think that part of the reason that OIRA is so transactional is because their staff must triage always to be able to be reviewing things. So to be able to step back and look across, I would say more staff.

Senator LANKFORD. OK.

Ms. KATZEN. In her testimony, Susan mentioned the function of OIRA as convening an interagency process, and I think that is the beginning of what you would like to lead to. If you look at Executive Order 12866, one of the definitions of “significant” is where there is something that an agency is proposing to do that is inconsistent with what another agency is proposing, or has already done, so that we do not have the Department of Labor (DOL) working on transporting hazmats, and the Department of Transportation (DOT) working on transporting hazmats, and they are at across purposes.

I thought that that would be a good vehicle for bringing together different agencies that were focused on a similar rule, and I see that in this context of retrospective review, where there might be similar rules at different agencies. That works sometimes, but by the time they have gone through the process of developing their rule, and how to implement it, how to enforce it, they are less willing to trade it for somebody else’s rule.

I think additional staff would help, and I think, again, oversight by the committee and language in the committee report would be helpful to clarify that this is one Administration, one Executive Branch, and that the various fingers should be working together rather than at across purposes.

Senator LANKFORD. Right.

There is some confusion on repealing a rule and going through that process, only that some people outside of government think if you walk in and do a review, then you decide you do not like it, you just take it out and the next day it is gone. Can you walk us through a little bit of the process that happens to actually go through a rule repeal?

Ms. KATZEN. OK. This is under the State Farm decision of the Supreme Court in the 1980s, having to do with airbags. And what it says is that if you are going to repeal or modify a regulation, you have to use the same processes that you used to create that regulation. Therefore, if you wanted to repeal or modify something that is on the books, you would need a notice of proposed rulemaking which sets forth the objective to modify an existing rule. It would need to be accompanied by data that documents the need for, the appropriateness of, making that modification.

And then you would get comments. So you have a notice of proposed rulemaking, and possibly now an advance notice of proposed rulemaking, and then the notice of proposed rulemaking, and then the comment period, and then the final rule, which would repeal the earlier rule.

Under a different Supreme Court case called Fox News, the Supreme Court has said that when you are changing your mind you have to (1) acknowledge that you are changing positions, and (2) if the previous rule was based on science, engineering, technical information, or economic information, you must show how that has changed, in what way is it different, why it is that you no longer
want to rely on the previous rule. Because there is new data? Because there is a new approach? Or whatever reason, and that explanation has to be part of the reason and basis, as it is called, of the new final rule.

So that is a whole new rulemaking proceeding, which can take a year, 2 years, and depending upon the basis for it, the assemblage of a lot of information and analysis. I hope that was clear.

Senator LANKFORD. That is exceptionally helpful, actually.

Susan, do you want to add anything to that?

Ms. DUDLEY. No. That was very clear.

Senator LANKFORD. Is it the right process to go through, you think?

Ms. DUDLEY. Yes.

Senator LANKFORD. I would assume that you would say that, because there is a great need to be careful. When you change a regulatory scheme of things that affects a lot of people, and you want to have as much impact as you can. Taking a rule out is the same as putting a rule in, and that creates uncertainty in the environment, and you have to be able to make sure everyone has input through that process.

Ms. KATZEN. I agree completely.

Senator LANKFORD. OK. So the question about retrospective review really goes back to the question of how do you determine whether things are working, and the metrics. I jokingly used the term, if there is a high school basketball team that is really bad at free throws, and the coach decides every time you miss a free throw you are going to take a lap, and that is his way to fix it, but a year later they are still not any better at free throws. They may be better cardiovascular but their free throws are still bad—we did one action to try to fix a problem but it did not work. There has to be some metrics at some point that have to decide, are we better at free throws on this?

I look at that in the same way with regulations, to say at some point we have a goal to say this is what we are trying to do, whether it clean the air or safer work environment, more consistency, better science. Whatever it may be there is a standard for what we are trying to accomplish. I think it is reasonable to say 10 years from now we should go back and look at it, and say did it actually accomplish that? The hard part is developing that metric.

So my question for you is, the developing the metric portion of it, is there any other definition that we need to put into this to give to OIRA or to the agencies to say when you develop a metric make sure you are thinking about this. Do you think what we have put into place, between Senator Sinema and I, is broad enough but is also clear enough that it is going to lay the groundwork for them to be able to do metrics, when you are looking a decade in advance?

Ms. DUDLEY. I do think it is so important to be able to define clearly what outcome you expect to get from the regulation, and a surprisingly limited number of regulations do that in a clear way.

I thought that what you had in the draft made sense. It covers the things that people talk about—the outcomes, which is general enough to cover a lot of things; benefits costs.

I would love to take the opportunity to look at it more and see if I do have some specific ideas I am in a public policy school where
a big focus is program evaluation. So some of the experts on that are at GW. I would love to talk with them, and say, “OK, does this get you what you would need 5 or 10 years down the road, to be able to do that evaluation?”

Senator LANKFORD. That would be great. We would be glad to be able to have that input.

Ms. DUDLEY. OK.

Senator LANKFORD. Should I put out a formal request for information to be able to get to that?

Ms. DUDLEY. An advance notice of a formal—yes.

Senator LANKFORD. OK. Thank you.

Ms. Katzen.

Ms. KATZEN. I agree with Susan on that, and my only caveat would be not to be too prescriptive. Ten years is actually a long time, given how we learn and how we develop. And we learn so much that to try to be specific might be greatly counterproductive here.

Senator LANKFORD. The hard part is trying to balance. You want to be specific enough that you actually know if you get there. I will use my free throw example. If the free throw percentage does not go up then that method did not work, so let’s go back and review it because we are trying to get to the end method. But general enough that the unknowns that are still out there can still be determined.

I guess what I am trying to say is if you make the metrics so vague, anything can hit it. It is not really a metric then. It is improved energy in the country. OK, what does that mean? So there has to be something specific enough.

Ms. KATZEN. I thought the proposal to include the objectives and the metrics at the NPRM stage was beneficial, because then those who are affected by the regulation, whether they be the regulated entities or the regulatory beneficiaries, will be able to comment on that during the comment period and lend their insight and their expertise to the agency, so that there would be a finite review of those two pieces when it is fresh in their minds, when they are still thinking about what they are trying to accomplish. I think that should be sufficient.

Senator LANKFORD. OK. Great.

Senator Sinema, you have a question as well?

Senator SINEMA. Yes. Thank you, Mr. Chairman.

For Ms. Katzen, you discussed escape hatches for repeated reviews of rules. In the SMART Act we have included a provision that allows the agency to create a list of circumstances in consultation with the OIRA administrator, which would require the performance of a subsequent review.

Do you think that provision adequately guards against unreasonable review requirements?

Ms. KATZEN. I believe so. I think the agency should be the responsible official who says we have looked at this, we are not going to make any changes, and further review would not be very helpful. I mean, I always use the example of airbags or seatbelts. Do we want to look at them in 10 years, then 10 years more, then 10 years more? Well, assuming we do not have V2V, assuming we do not have autonomous vehicles—but even with autonomous vehicles...
I want my seatbelt and airbag in the car. Why ask the agency to go through that? I think the agency, Secretary of the Department or the agency official, is the person to be able to say enough is enough.

Senator Sinema. Regulatory impact analysis documents, which accompany regulations, provide wide ranges of estimated costs and benefits. The uncertain nature of forecast-based analysis hides the true impacts and benefits for rules. So through the data collection requirements of the SMART Act, can we expect that, over time, agencies will become better positioned to accurately forecast both costs and benefits?

Ms. Dudley. I would hope so. I think one of the most important potential outcomes of retrospective review is that it is going to make us better at predicting things in future regulations. So both thinking about the data and analysis and what models we use to make those predictions? They are so uncertain and we never go back and check. So I think that is a key benefit.


Senator Sinema. A final couple of questions. In written testimony you discussed a number of rules that would be applicable under the Early Participation in Regulations Act, and I believe you said 70— is that correct, 70?—but that many rules would not benefit from the bill's requirements.

Could you discuss these rules that would be major but would not benefit from an ANPRM?

Ms. Dudley. Yes, the 70 includes all the independent regulatory agencies. If you just look at the Executive Branch agencies it is closer to 40 or 50 that would fall under the rule.

One set of rules are hunting bag limits, and there probably a half a dozen of those every year. OIRA has long let the Department of Interior (DOI) use the same regulatory impact analyses that they probably prepared 15 or 20 years ago, as a regulatory analysis. That would not be the kind of thing that advance notice.

I would expect, although I am not positive, that a lot of the Medicare and Medicaid rules—the regulations that are determining what fees different doctors or services should receive—again, that is not the kind of thing that advance notice might be valuable for. So I can imagine that early on OIRA and the agencies would streamline that and identify some exemptions.

Ms. Katz. I agree with that, and in my testimony I mentioned that there are occasions when the congressional delegation, which is the basis for any rulemaking, it itself highly prescriptive. I am thinking of probably the unregulated example of positive train control, which did not give the agency any discretion to do anything other than what was specified in the statute. That being the case, an advance notice of proposed rulemaking would not make a whole lot of sense. The agency’s hands are tied by its authorizing statute.

Senator Sinema. Thank you. Thank you, Mr. Chair.

Senator Lankford. Senator Carper.

Senator Carper. Thank you both again for your testimony today. As I am sure you know, under the Paperwork Reduction Act (PRA), the Office of Management and Budget must review and approve Federal collections of information before they are, I guess, con-
ducted. After reviewing the agency request, the OMB may approve or disapprove that request, or can go ahead and define conditions that need to be met for approval.

The OMB is required to ensure that any information collection maximizes practical utility and public benefits and protects integrity, objectivity, and impartiality of collected statistical information.

Last May, I led a letter with 34 of our colleagues, both in the House and the Senate, requesting information from the Commerce Department and OIRA as to how they planned to ensure that the Paperwork Reduction Act requirements were met with respect to the addition of a question on citizenship on the 2020 Census. That was last May, a year ago. We have not received a response to that letter.

Let me ask, first, if I could, Ms. Katzen, as you know, concerns have been raised about adding a question on citizenship, due to the potential negative consequences, including a lower self-response rate, which would lead to a less accurate and more costly Census. I would just ask would you please weigh in for us and explain OIRA's role in reviewing information collections, and what OIRA should be reviewing with respect to the citizenship question and the 2020 Census? Could you take a shot at that?

Ms. KATZEN. I will try. Thank you.

Senator CARPER. Thank you.

Ms. KATZEN. Under the Paperwork Reduction Act, any question being posed to the public, identical questions to 10 or more people, has to receive, in effect, a comment period by the agency, and then if the agency decides to go ahead with it, the agency will then send it to the OMB, which has a second comment period. And as you correctly stated, one of the conditions for approval by OMB is to maximize practical utility.

There is also a requirement to minimize burden. It is a modified cost benefit analysis. Are we going to get something from this paperwork requirement that will be useful and well worth whatever offsetting cost there is?

Now with the Decennial Census, for 2020, there has been a lot of publicity, and I have actually read the three district court cases, one of which is 267 pages. So I have some knowledge about the specifics of that particular paperwork requirement.

I think the most significant aspect is that the primary purpose of the Decennial Census, which is embodied in the Constitution—it is the only paperwork requirement embodied in the Constitution—is for the enumeration every 10 years. That is the primary purpose.

The addition of a question relating to citizenship is a question which, at least pretextually, has been justified by assisting the Department of Justice (DOJ) in better enforcing voting rights cases. That is a secondary purpose. If the secondary purpose is going to have an adverse effect on the primary purpose, one would have a very hard time justifying it under the Paperwork Reduction Act, because every past Census Bureau director, and vast numbers of statisticians, including the National Academy of Science, has said that this will decrease the response rate significantly for the enumeration purposes, and that the data are not needed in the Decen-
nial Census because there are alternative places where this data reside, in the ACS study and other kinds of statistical compilations. With that, it is hard to see how the Decennial Census with the citizenship question would pass muster under the Paperwork Reduction Act. Now there are all sorts of politics, policies, whatever, but I was just trying to focus——

Senator CARPER. OK. Good.

Ms. KATZEN [continuing]. On the PRA.

Senator CARPER. Thank you. Mr. Chairman, you have been generous with your time. I would like to ask, if I could, for the witnesses to each give us like maybe one change, amendment that you would suggest to the legislation that Mr. Chairman has designed, just one change. And as I said earlier, I never introduced a perfect bill, or probably a perfect amendment, and as good as these two legislators are, these two Senators are, there is probably room for improvement. Can you think of one thing that you would suggest that we amend as we take it to markup?

Ms. Dudley.

Ms. DUDLEY. Perhaps that the 10-year window for review, it provides for one review at 10 years, and then nothing thereafter. And that may be appropriate for some types of regulations but not for others.

So I think Sally and I disagree on this. I might like to see a continued requirement for that retrospective analysis that would continue to measure and continue to observe whether we are achieving the desired objective.

Senator CARPER. All right. Thank you. Just one idea, Sally, if you would.

Ms. KATZEN. Susan is correct that I disagree with her suggestion. I was looking at what was supposed to go in the ANPRM, and I came across something which I question its utility, and that was in the ANPRM to list the legal authority under which a major rule may be proposed. And then the following language: “including whether a rulemaking is required by statute, and, if so, whether by a specific date, whether the agency has discretion to commence a rulemaking.”

I can envision circumstances where the discretion is reduced with greater incidence of death or harm or safety of some sort. I had not really focused on that particular language, but I am not sure that, if I were giving you one thing that think I might want to change from the latest draft—which looks very good, indeed—it might be to stop that little a iii, whatever it is, little iii sooner.

Senator CARPER. Alright. Thank you.

For some people, you remember the saying—this is about as exciting—talking about an experience in their life—they said it was about as exciting as watching wet paint dry. I suspect for some people a hearing on this subject is just like that. I think it is a terrifically exciting hearing, and I applaud our Chair and Ranking Member for bringing us together, and look forward to working with you to—what does it say in the Constitution?—“We the People of the United States of America, in order to form a more perfect union,” to maybe work with you to see if we can form an even more perfect bill. Thank you.
Senator LANKFORD. Sure. By the way, my oldest daughter graduated and is out and has a real life, and my wife and I repainted her sunshine-yellow bedroom into a color where guests could actually sleep. I put the initial primer coat on, and put the regular coat on, and watched the yellow disappear, and as I watched the paint dry I thought, I am standing here watching paint dry, and it is exciting. [Laughter.]

Because I feel like we are getting a room back. So watching paint dry is not always bad.

Senator CARPER. Well, hopefully they will come back and sleep in these rooms sometime. That sounds like a lot of fun.

Senator SINEMA. I think I would like the yellow.

Senator LANKFORD. The sunshine yellow?

Senator SINEMA. Yes.

Senator LANKFORD. It was a happy color.

Senator SINEMA. Yes, I would like that.

Senator LANKFORD. She loved it, all the way through high school. She loved it.

Senator LANKFORD. Let me say thank you to both of you. I would ask you both two quick questions, and one you can think about and send back your ideas, just as a message to say this Committee is always interested in your input. So as you have good ideas we are always listening. So consider this your advance notice of proposed legislating, that we are interested in the ideas.

One is, Ms. Katzen, several times you have mentioned surgical changes to the APA. That is what we are trying to do, not a massive wholesale shift but surgical changes. If you have thoughts on other surgical changes where we need to legislate on—we focused today on beginning and end, but if there are other areas we are interested, and we are trying to be able to work through to try to find ways to be able to help fix the process long-term. So that is my homework assignment to you. If you think of anything away from here, contact us or our team, and we would be glad to be able to hear that.

The other one is, based on where the text is right now—I got your comments from Senator Carper—do you support these bills where they are right now, and continue to move forward in the process?

Ms. DUDLEY. Absolutely.

Senator LANKFORD. Yes.

Ms. KATZEN. I would indeed.

Senator LANKFORD. Yes. Thank you. Thanks for all your input. You have both been exceptionally professional with our team in trying to be able to provide some additional input and thought. You both have very busy lives on your own. You do not work for us, but you do work for the American people still—that is pretty obvious—of your continuing engagement in policy areas. So thanks for continuing to be able to give your time, to be able to help the Nation in the future.

Let me make a quick closing statement and we will shut us down. Before we adjourn I do want to announce that on May the 22 this Subcommittee intends to hold a joint hearing with the Senate Small Business and Entrepreneurship Committee to examine the disproportionate impact regulations have on small businesses.
Both of you have dealt with that quite a bit, actually, in your time at OIRA. All members of the Homeland Security and Governmental Affairs Committee (HSGAC) are invited to attend this hearing.

That concludes today’s hearing, though. I do want to thank again our witnesses for their testimony. The hearing record will remain open for 15 days until the close of business on May the 22, for the submission of statements and questions for the record.

This hearing is adjourned.

[Whereupon, at 11:20 a.m., the Subcommittee was adjourned.]
Appendix

Opening Statement
Hearing before the Regulatory Affairs and
Federal Management Subcommittee
Tuesday, May 7th at 10:00 am

“From Beginning to End: An Examination of Agencies’ Early Public Engagement and Retrospective Review.”

- Good Morning, welcome to today’s hearing titled “From beginning to end: an examination of agencies early public engagement and retrospective review.”

- I’d like to welcome Senator Sinema to the dais and I look forward to working with you in the Senate and specifically in this Subcommittee to improve our regulatory process and support our federal workforce.

- Thank you to our witnesses who bring a unique insight into the regulatory process. Former OIRA Administrators have been an incredibly valuable resource to this subcommittee and we thank you for your willingness to impart your knowledge and share your unique perspectives.

- Today we are focusing on how agencies conduct public outreach at the beginning of a rulemaking and how a rule’s success or failure is measured years later.

- While these could be considered narrow issues, they are by no means small.

- Advanced Notices of Proposed Rulemaking, or ANPRMs, are important tools that only a few agencies are statutorily required to utilize.
• The Administrative Procedure Act sets out a process where agencies propose a rule, listen to comments from the public, have the opportunity to make changes, and then issue a final regulation.

• While this system looks good on paper, in practice, agencies conduct a significant amount of work before formally engaging the public – they will consider various regulatory schemes and conduct economic analysis. Ideally, agencies would engage the public early, but there is no APA requirement to do so.

• Turning to the other end of the rulemaking process, retrospective review is a process to ensure rules achieve their intended goal in the least burdensome way. Over time, changing circumstances and improved technology may render some regulations ineffectual or unnecessary.

• An agency’s job is not done after a final rule is published. As initial estimates of both costs and benefits can prove inaccurate, agencies should revisit a rule to ensure the desired effects are achieved.

• Every President since Jimmy Carter has urged agencies to utilize retrospective review to examine existing regulations. While these directives were issued with good intentions, they gave agencies a significant amount of discretion in selecting which, and how many rules to review.

• The focus of this hearing is two bills Senator Sinema and I will introduce shortly that codify the best practices for both procedures.

• The Early Participation in Rulemaking Act directs agencies to issue advanced notices for rules costing more than $100 million annually.

• The agency must outline what problem the rule intends to solve and listen to the public’s input on the subject.
• The idea behind this bill is to require agencies to listen to the public before they craft a regulation – Washington does not have all the answers, taking time to work with stakeholders, particularly our small businesses, is vital in crafting effective regulations.

• Less burdensome does not mean less effective. Business owners want to be good citizens, follow the law, and have safe and clean workplaces.

• The Setting Manageable Analysis Requirements in Text Act (or SMART Act) is a retrospective review bill that looks ahead – it requires agencies to set metrics for how a rule will be measured for success in the future.

• It is hard to measure the success of anything unless it is defined. This bill instructs regulators to define what “success” is for a given rule and then requires them to grade the rule within 10 years.

• This Subcommittee has been working on both of these issues for a while and both bills have had bi-partisan support for the past two Congresses. I look forward to working with my colleagues to push them both across the finish line. With that, I recognize Senator Sinema for her opening statement.
Prepared Statement of Susan E. Dudley

Director, GW Regulatory Studies Center
Distinguished Professor of Practice,
Trachtenberg School of Public Policy and Public Administration

Hearing on

From Beginning to End: An Examination of Agencies’ Early Public Engagement and Retrospective Review

Homeland Security and Governmental Affairs Committee
Regulatory Affairs and Financial Management Subcommittee
United States Senate

May 7, 2019
From Beginning to End: An Examination of Agencies’ Early Public Engagement and Retrospective Review

Prepared Statement of Susan E. Dudley

Thank you, Chairman Lankford, Ranking Member Sinema, and Members of the Subcommittee for inviting me to share my thoughts on early public engagement and retrospective review of regulations. I am Director of the George Washington University Regulatory Studies Center, and Distinguished Professor of Practice in the Trachtenberg School of Public Policy and Public Administration. From April 2007 to January 2009, I oversaw federal executive branch regulations as Administrator of the Office of Information and Regulatory Affairs (OIRA) in the Office of Management and Budget (OMB). I have studied regulations and their effects for over three decades, from perspectives in government (as both a career civil servant and political appointee), the academy, and private consulting.

I appreciate the Subcommittee’s interest in improving how the U.S. government develops and evaluates regulatory policy. Your efforts continue a long bipartisan tradition in the United States of efforts to make regulation well-informed, transparent, and accountable to the American people. By 1) engaging public input earlier in the regulatory development process and 2) providing for retrospective review of regulations to evaluate whether they are achieving their objectives, the bills you have proposed can help ensure that regulations are based on the best evidence available and that they are working as intended for the American people.

My testimony reviews the problems necessitating the practices required by your legislation and addresses and examines each bill’s requirements and impacts. It concludes with some cross-cutting comments and observations.

Engaging Public Input Early in Rulemaking

The Problem

The Administrative Procedure Act of 1946 requires agencies to publish a “general notice of proposed rule making ... in the Federal Register,” and “give interested persons an opportunity to participate in the rule making through submission of written data, views, or arguments with or
without opportunity for oral presentation." 2 In addition, every president since Jimmy Carter 3 has required agencies to examine expected regulatory impacts before issuing proposed and final regulations; Executive Order (E.O.) 12866 has guided this analysis for more than 25 years. 4 Despite these longstanding requirements, agencies often develop regulatory impact analyses (RIAs) after they have made key policy decisions; many analyses appear to be designed to justify regulatory actions, rather than inform them. 5 Agencies view their RIAs and preambles to proposed rules as legal documents, prepared in anticipation of litigation. 6 The need to defend their selected regulatory approach motivates agencies to “circle the wagons,” narrowing the menu of alternatives and the evidence they consider before the public has an opportunity to engage. As a result, changes in response to notice and comment “tend to be small and painful, and they are often subtractive rather than innovative or additive.” 7

The Early Participation in Regulations Act of 2019

The draft “Early Participation in Regulations Act of 2019” would require agencies to issue for public comment advance notices of proposed rulemaking (ANPRMs) for major rules. This requirement could free the agency to share its early thinking on whether a problem requires a regulatory solution and what different options are available. As such, ANPRMs could be valuable for soliciting input from knowledgeable parties on a range of possible approaches, data, models, etc., before policy decisions are framed, or positions established. 8 As the President’s Jobs Council noted in 2011, resulting regulations would benefit from critiques, feedback, and other public input provided by ANPRMs. 9

---

2 5 U.S. Code § 553(b) and (c).

The George Washington University Regulatory Studies Center
ANPRMs could encourage better, more informed regulatory analyses. Experts have suggested that “back of the envelope” analyses could encourage agencies to consider the effects of a wide range of alternatives before they narrow their focus to just a few options.\textsuperscript{10} Empirical research found that “pre-proposal notice[s] requesting comment from the public... are associated with higher quality regulatory impact analyses” supporting final regulations.\textsuperscript{11}

The bill would require an ANPRM for a major rule to identify the problem that may call for regulation, and data or information that supports that regulatory need. This is an essential first step for developing effective regulation. E.O. 12866 calls on each agency to “identify the problem that it intends to address (including, where applicable, the failures of private markets or public institutions that warrant new agency action) as well as assess the significance of that problem.”\textsuperscript{12} Yet, in my experience, agencies too often proceed with developing a regulatory solution without first clearly articulating the nature and significance of the problem to be solved.\textsuperscript{13} The bill’s related requirement to identify “an achievable objective for the major rule” would also serve to focus public comment, subsequent analysis, and evaluation.\textsuperscript{14}

ANPRMs subject to the bill would also present a general description of alternatives the agency has identified for consideration. Laying out a range of preliminary alternatives early in the regulatory development process could elicit invaluable input from the public, not only on the merits of those alternatives (including data, analysis, experience) but suggestions of other options for agency consideration.

Together, these ANPRM elements should not be unduly demanding or burdensome; they would merely make the factors influencing the agency’s thinking more transparent to the public at a stage when public input could be very valuable.

Over the past few decades, agencies have issued an average of 13 significant ANPRMs per year, representing less than 5% of their significant regulatory actions.\textsuperscript{15} As noted below, given the bill’s definition of “major rule,” as many as 70 regulatory actions a year\textsuperscript{16} could begin with an

\textsuperscript{10} Carrigan and Shapiro (2017).
\textsuperscript{12} E.O. 12866 Sec. 1(b)(1).
\textsuperscript{13} Dudley et al. (2017).
\textsuperscript{16} Major rules include regulations issued by independent regulatory agencies, which are not captured in the above counts. Executive branch agencies issue an average of 40 regulations per year that would likely meet the bill’s definition of major. See the data maintained by OIRA and the General Services Administration at www.RegInfo.gov.
ANPRM. However, the number would likely be less than that since the bill provides for exceptions.

The bill appropriately gives OIRA authority for determining whether a rule is major under the section, and whether an exemption should apply. OIRA is well-positioned to make those determinations given its authorities under the Congressional Review Act (CRA) and presidential executive orders.

The bill also wisely precludes judicial review of any differences between an agency's ANPRM and subsequent NPRM. One virtue of the ANPRM is that it provides an opportunity for agencies to share their preliminary thinking about a problem and get input on potential solutions at a stage when they are truly open to feedback, analysis, and evidence. If agencies had reason to fear this early notice could later be used against them in court, that would discourage objective queries and undermine these benefits.

The requirement to issue and accept comment on an ANPRM before proceeding to a proposal should not unduly slow agencies' regulations for several reasons. First, 90 days is not a long time considering that agencies often take years studying a problem and evaluating regulatory options before they issue a proposed rule. One study found that "the average interval between the formal initiation of research on a policy issue...and the publication of a proposed rule...was 5.3 years." 17

Perhaps more important, to the extent the ANPRM serves to open up for public engagement preliminary deliberations that would otherwise have taken place behind closed doors, it may make the overall rulemaking process more efficient. Rather than tacking 90 days onto the rulemaking schedule, it may provide valuable input that ends up streamlining the subsequent notice-and-comment process. In many cases, early engagement could lead to more efficient analysis at the NPRM stage and fewer surprises during public comment. While there will certainly be cases where an ANPRM would not serve the public interest, the bill provides for those exceptions.

Creating an Evaluation Mindset

The Problem

Ex post evaluation has a long tradition in other areas (particularly in programs financed through the fiscal budget),18 but it has received little attention in the regulatory arena, despite government

17 West (2004).
RIAs are an important part of the regulatory process, but as ex ante analyses, they are necessarily hypotheses of the effects regulatory actions will have if implemented. Better regulatory evaluation would allow agencies and others to test those hypotheses against actual outcomes. Retrospective review would not only inform decisions related to the benefits and costs of existing policy but would provide feedback that would improve future RIAs and future policies.

Setting Manageable Analysis Requirements in Text (SMART)

While no one questions the importance of evaluation, a lack of methods and data make retrospective review of regulations challenging. The draft SMART bill addresses that problem at the outset of rulemaking by requiring agencies to include in major regulations a framework for how they will measure effectiveness, benefits, and costs, and to incorporate plans for gathering the information necessary for ex post evaluation. It would also require agencies, within 10 years of a rule’s effective date, to assess its benefits and costs, evaluate how well it accomplishes its objectives, and determine whether it could be modified to achieve better outcomes.

This would fill an important gap in current regulatory practice. The GW Regulatory Studies Center reviewed all major rules proposed in 2014 and found that, despite President Obama’s requirements to do so, none of them included a plan for retrospective review, and not one was written and designed to facilitate review of its impacts. While we have not conducted a similarly comprehensive review since that year, case-by-case analysis suggests that most regulations continue to be issued without any plan for review.

The bill would ensure not only that existing major regulations are being evaluated, but that new major rules are designed to facilitate such evaluation in the future. It focuses not just on reducing regulatory burdens, but on improving regulatory outcomes by subjecting regulatory programs to rigorous evaluation and feedback. Institutionalizing a requirement to evaluate whether the predicted effects of the regulation were realized would provide a powerful incentive to improve the ex-ante RIA tools used to predict the impacts of regulatory alternatives. The bill would

19 Joseph Aldy, Learning from Experience: An Assessment of the Retrospective Reviews of Agency Rules and the Evidence for Improving the Design and Implementation of Regulatory Policy, ADMIN. COMP. U.S. (Nov. 18, 2014). Aldy writes that federal regulatory agencies have a mixed record on exp post review, despite their “long track record of prospective analysis of proposed regulations that can address these questions.”
22 Dudley (2017).
create an evaluation mindset and a feedback mechanism where agencies learn from evaluating regulatory outcomes and apply those lessons to improve future rules.

Other Observations

These two draft bills offer relatively modest, yet potentially powerful, changes to current rulemaking practices. If enacted, they could make regulatory decisions more transparent and accountable, leading to improved regulatory outcomes for the American people. My testimony concludes with observations on features common to both bills.

The definition of “major rule” in both bills appropriately captures what are likely to be the most significant regulatory actions, while not unduly burdening agencies with additional procedures for all their rules. It maintains the annual $100 million annual impact trigger embodied in the CRA and E.O. 12866 (Sec. 4(f)(i)), but it is not purely a monetary test. It recognizes that rules that are likely to significantly affect consumer prices, competition, productivity, innovation, the environment, public health, or safety deserve greater public engagement and ex post evaluation.

According to records kept by the Government Accountability Office (GAO), agencies issue approximately 3,000 regulations each year. Of those, GAO classifies around 900 as “substantive or significant.” An average of 70 of those meet the CRA definition of major, which closely resembles the bills’ definition of major rule. This likely overstates the number of regulations that would be subject to these bills’ requirements since many of them are routine (such as annual hunting and fishing limits) or affect annually recurring monetary transfers from taxpayers to program recipients (for example, Medicaid and Medicare payment rules). For these, the requirement to issue ANPRMs and develop a retrospective review framework could either be streamlined or they might qualify for an exemption.

Accomplishing the important goals of these bills would require resources. As noted above with respect to timing, the ANPRM requirement need not impose additional resource costs to the extent it engages public participation in preliminary deliberations that have traditionally been closed. To support more rigorous retrospective review, Congress and OMB could reallocate resources from ex ante analysis to allow agencies to gather the information and evaluation tools necessary to validate ex ante predications. In the long run, shifting resources from ex ante analysis to ex post review would not only help with evaluation, but could improve agencies’ ex ante hypotheses of regulatory effects.27

26 The OIRA/GSA RegInfo.gov database classifies less than 300 rules as significant on average each year. Fewer than 50 of those would meet the bills’ definition of major. See GW Regulatory Studies Center “Reg Stats” for more detail.

Both bills would make OIRA responsible for overseeing compliance, which is appropriate. Executive branch oversight of regulatory actions has proven valuable, but it is not sufficient. Congress may also want to assign a congressional body, perhaps a new regulatory office in the Congressional Budget Office (CBO), responsibility for reviewing these assessments. Just as the CBO provides independent estimates of the on-budget costs of legislation and federal programs, a congressional regulatory office could provide Congress and the public feedback on legislation that enables regulation, as well as serve as an independent check on the analysis and decisions of regulatory agencies and OIRA.28


The George Washington University Regulatory Studies Center
Statement of Sally Katzen  
Professor, NYU School of Law  

before the  
Senate Homeland Security and Governmental Affairs Committee  
Subcommittee on Regulatory Affairs and Federal Management  

on  
“From Beginning to End: An Examination of Agencies Early Public Engagement and Retrospective Review”  

May 7, 2019
Chairman Lankford, Ranking Member Sinema, Members of the Subcommittee. I understand that this hearing is intended to explore specific areas of the process for federal rulemaking that would likely garner bipartisan support. To that end, I am pleased to participate and thank you for inviting me to testify today.

I have worked on regulatory issues during most of my career in private practice, government service, and teaching and writing. I served as the Administrator of the Office of Information and Regulatory Affairs (OIRA) at the Office of Management and Budget (OMB) for the first five years of the Clinton Administration, then as the Deputy Assistant to the President for Economic Policy and Deputy Director of the National Economic Council, and then as the Deputy Director for Management of OMB. Before entering government service in 1993, I was a partner at the law firm Wilmer, Cutler & Pickering (now WilmerHale), specializing in regulatory and legislative issues, and among other professional activities, I served as the Chair of the American Bar Association Section on Administrative Law and Regulatory Practice (1988-89). During my government service, I was the Vice Chair (and Acting Chair) of the Administrative Conference of the United States (ACUS). After leaving the government in January 2001, I have been teaching courses in administrative law at various law schools, and since 2011 I have been at New York University School of Law, where I am currently a Professor of Practice and Distinguished Scholar in Resident.

As this Committee well knows, the regulatory system — and the rules that it develops, promulgates and enforces — is an integral component of governance. Congress makes the law but it typically does not have the time, the expertise, or sometimes the ability to identify and resolve all the details. That responsibility is usually delegated to the agencies that are expected to issue regulations that translate general statutory directives into concrete requirements or prohibitions with which the public must comply. There are appreciably more regulations than statutes: some have depicted it as a pyramid, with the Constitution, the supreme law of the land on top, hundreds of statutes enacted by Congress on the next level, and then thousands of regulations issued by the agencies.

Apart from the delegations from Congress — which provide the primary direction and constraint on an agency’s substantive and procedural authority — the principal law that governs the development and promulgation of regulations is the Administrative Procedure Act (“APA”). It was enacted in 1946 and, with relatively few amendments — mostly having to do with the Freedom of Information Act — and with a series of federal
court cases fleshing out the general terms of the Act, it has generally withstood the test of time.

Most of the criticism and praise of the administrative state is focused on regulations produced by notice-and-comment rulemaking – the subject of Section 553 of the APA. While it is commonly referred to as “informal rulemaking,” there is nothing informal about the process. It is resource intensive and time consuming – some rulemakings take years rather than months to go from concept to a final rule, plus whatever additional time and effort goes into judicial review.

The current process has its critics, from both conservatives and progressives. That suggests that perhaps the process is just about right. Conservatives are concerned that there are not sufficient checks along the way; progressives are concerned that the number of checkpoints has created ossification. In my view, process is good, but too much process can be counterproductive. The issuing agency should think, research, consult, analyze, question and continually refine. The public, both those who would bear the costs or burden of the regulation and those that would benefit from it, should be informed of what the agency is thinking and why, and be engaged in supplying and reviewing the information the agency is relying on and in critiquing the options the agency is considering. The public has a great deal to contribute, but the process should not be so extended as to unduly delay or disrupt the work of the agencies. And, most importantly, each step in the process (and any new steps imposed) should be evaluated in terms of its contribution to good decision-making, to what would help produce the most sensible, effective and efficient way forward.

I understand that this Committee is focusing today on two specific aspects of the rulemaking process: the beginning and the end. I think that each of these pieces can be improved, whether by legislation, executive order, OMB guidance or simply agency practices. Regardless of the vehicle, however, it is important to be clear about what the problem is and how best to solve that problem without introducing unintended consequences.

**The Beginning: an Advanced Notice of Proposed Rulemaking.**

The first official step in a rulemaking proceeding is the issuance of a notice of proposed rulemaking ("NPRM"). Lawyers in practice and in the academy generally agree that by the time the agency issues the NPRM, the staff involved have invested so much time and energy in developing the proposal and supporting data (as they are required to do) and analyzing the likely effects of the proposal (as they are required to do) and justifying their proposal (as they are often called upon to do by their agency
decision-makers and OIRA review) that they are virtually locked into their proposal and are less receptive to new ideas (or even significant modifications of their proposal). While all the up-front work (and documentation) is desirable, it often has the unintended result of restricting the options going forward.

To counter this tendency, there have been various efforts to encourage the agencies to consult with the public even before they have essentially made their decisions reflected in the NPRM. This has been true of both Democratic and Republican Administrations. For example, Executive Order 12866 clearly states that “before issuing a notice of proposed rulemaking, each agency should . . . [consistent with its own rules] seek the involvement of those who are intended to benefit from and those expected to be burdened by any regulation (including, specifically, State, local, and tribal officials).” Section 6(a) emphasis added. And Executive Order 13563 expanded on this concept with an entire section devoted to “Public Participation.” See Section 2. Among other things, it states in subsection (a) that “regulations shall be based, to the extent feasible and consistent with law, on the open exchange of information and perspectives among State, local, and tribal officials, experts in relevant disciplines, affected stakeholders in the private sector, and the public as a whole.” And it specifically provides in subsection (c) that “[b]efore issuing a notice of proposed rulemaking, each agency, where feasible and appropriate, shall seek the views of those who are likely to be affected, including those who are likely to benefit from and those who are potentially subject to such rulemaking.” Section 2, emphasis added.

Several agencies use an Advance Notice of Proposed Rulemaking (ANPRM) to solicit ideas at the outset of a rulemaking proceeding. It is especially useful when the agency is unsure what direction to take, what data to consider, how prescriptive to be, and the like. It is also more useful when it is done early in the process or even at the outset of agency deliberations (for example, when the agency first sends notice of its undertaking a rulemaking for inclusion in the Unified Agenda). It is less useful when the authorizing statute is itself prescriptive or if there is general agreement about the nature of the regulation necessary to respond to the identified problem. In short, it can be helpful at times; at other times, it may just add an unproductive, but time consuming, step to the already extended process.

For this reason, it is important that any requirement for an ANPRM should be limited to economically significant (or “major”) regulations that are required to use notice and comment under Section 553. Similarly, it is important that any such provision should not impose on the agency multiple requirements for explanations, analysis, data, etc. The purpose would be to alert those affected by the regulation, so
they can contribute to its development and formulation in advance of the NPRM, not to lock the agency into a particular mind-set before the process even begins. The more the agency has to incorporate in an ANPRM, the more the agency will become invested in a particular outcome. This is the opposite of what an ANPRM should do—namely, obtain ideas and information from interested entities before the agency settles on a particular course.

I understand that your and your staffs have been working on a draft bill that reflects these considerations and takes a sensible (but appropriately limited) step towards expanding the opportunity for public participation at the pre-NPRM stage. If the product of those efforts adheres to the principles (and specific provisions) we have been discussing, I would be supportive of the effort.

The End: Retrospective Review

For almost 40 years, there have been concerns that there are too many rules, and that so many of the rules on the books are obsolete, unnecessarily burdensome, unworkable, or just plain wrong. This was one of the themes President Reagan campaigned on, and, after his election, he set on a course to deregulate. President George H.W. Bush followed the same path with his Competitiveness Council, searching the existing stock of regulations for those that could be eliminated. President Clinton ordered agencies “to submit to OIRA a program . . . under which [they] will periodically review existing significant regulations to determine whether any such regulations should be modified or eliminated . . .” E.O. 12866, Section 5. President George W. Bush launched a similar effort. President Obama also emphasized the need for retrospective review of rules in his Executive Order 13563. And President Trump’s “two-for-one” Executive Order is designed in part to accomplish the same objective—weed out unnecessary, out-of-date, ineffective rules.

Despite these efforts, it doesn’t happen. One reason may be that since 1980, new regulations are not issued unless their benefits justify their costs; to eliminate such a regulation would likely mean that the costs of rescinding the regulation would be greater than the benefits. [Note: This is so because removing a rule means that the foregone benefits of the existing rule become the costs of the new rule, and the foregone costs of the existing rule become the benefits of the new rule]. Other reasons for the limited success of these efforts are that agencies have not undertaken to collect data along the way that would inform their retrospective reviews and, importantly, any retrospective analysis requires resources and, for at least the last few decades,
regulatory agency budgets have generally been decreasing or straight-lined, with the situation compounded by continuing resolutions and sequestration.

Nonetheless, there is growing support for one step that can be taken – namely, encouraging agencies to plan for retrospective review when they are in the process of developing a final rule. This idea came from a study by Joseph Aldy for the Administrative Conference of the United States (ACUS), Learning from Experience: An Assessment of Retrospective Reviews of Agency Rules & the Evidence for Improving the Design and Implementation of Regulatory Policy (Nov. 2014). This led to a series of ACUS recommendations designed to promote “a culture of retrospective review at agencies,” which stressed the need to carefully select regulations for reevaluation and to coordinate with OIRA, other agencies and outside entities (including stakeholders) when designing and conducting retrospective reviews.” For present purposes, it is instructive that, among other things, ACUS adopted a specific recommendation for “agencies to plan for retrospective review when drafting new regulations.” A similar recommendation was part of a report to the 2016 transition teams developed by the Institute of Policy Integrity after consultation with almost all of the past OIRA Administrators. Strengthening Regulatory Review (2016).

Requiring agencies to provide, along with the NPRM and the final rule, a plan for a later retrospective review of a newly issued regulation would in most circumstances be salutary. If nothing else it would force agency personnel to focus on describing precisely what they want to accomplish and how to evaluate whether or not the rule is successful in achieving that objective, at a time when the rule (and all its alternatives) is foremost in their minds. It also would enable those affected by the rule to participate in the framing of the subsequent retrospective review while they too are keenly focused on the provisions of the proposed rule.

It is very important, however, to provide flexibility for the eventual implementation of the retrospective review. The agency can (and should) commit to a framework in the proposed and final rule, identifying the data and the metrics it anticipates using for that purpose. But this should not be cast in concrete. We learn a lot with time, including how to better analyze and measure what is going on around us. This is amply demonstrated by the increased sophistication of cost-benefit analysis itself over the last decade or two. It is also demonstrated by the general preference for performance standards, which specify the desired results, rather than design standards, which lock in a particular way to achieve those results.

In addition, while periodic review is useful, there will likely be some (or many) situations where repeated retrospective reviews will yield greatly diminishing returns.
After a decade or so, rules that survive a retrospective review intact are likely to have established their worth, and it would be wasteful to continue retrospective review after retrospective review. In this connection, Section 553(e) of the APA provides for petitions for rulemaking that can be used if, at some later point in the future, a consensus develops that a particular rule should be modified or rescinded. This would place some responsibility on the regulated entities, but that is appropriate because they are in the best position to identify (and document) rules that have outlived their usefulness.

As above, I understand that you and your staffs have been working on a draft bill that would implement these recommendations in a straightforward, targeted way. Again, if the product of those efforts reflects the principles (and specific provisions) we have been discussing, I believe such a bill would be a constructive addition to the rulemaking process.

Again, I appreciate this Committee's efforts to fine tune critical steps rather than redesign the whole rulemaking process. I emphasize this because, unfortunately, you are not writing on a blank slate. For the past three decades, there have been concerted efforts underway to revise and revamp the rulemaking process that have mostly proceeded on a highly partisan path. Not only have they not been successful in terms of being enacted into law, but they have also sown suspicion and distrust. As a result, bipartisan efforts on regulatory reform have been difficult to achieve. The limited, surgical approach that you are considering will likely face an uphill battle, but it might well have a better chance to succeed.

Thank you again for giving me an opportunity to speak to these issues. I look forward to any comments or questions you may have.
Susan Dudley Responses to
Post-Hearing Questions for the Record
“From Beginning to End: An Examination of Agencies’ Early Public Engagement and Retrospective Review”
From Senator Thomas R. Carper
For the Honorable Susan Dudley and for the Honorable Sally Katzen

**OIRA’s role in reviewing agency rulemaking and EPA’s “Strengthening Transparency in Regulatory Science”**

On May 9, 2018 I joined Senator Hassan and several other Senators in a letter to then-OIRA Administrator Neomi Rao raising concerns and seeking answers as to how OIRA reviewed an EPA proposed rulemaking entitled: “Strengthening Transparency in Regulatory Science.”

According to Reginfo.gov, OIRA received the EPA’s proposed rule on Thursday, April 19, 2018. Then-EPA Administrator Scott Pruitt publicly announced this proposed rule and signed it three business days later on Tuesday, April 24, 2018. Reginfo.gov initially stated that OIRA’s review was completed on Wednesday, April 25, 2018– the day after Administrator Pruitt announced and signed the rule. Later, following press inquiries, Reginfo.gov was changed to indicate that OIRA’s review was completed on Monday, April 23, 2018.

Our letter sought detailed information on the scope and substance of OIRA’s review of this proposed rulemaking. However, in a response to Senator Hassan on July 12, 2018, former Administrator Rao provided little in way of response and simply noted that the conclusion date for the OIRA review of this proposed rule of April 25, 2018 was due to a clerical error that was subsequently corrected.

With this in mind, I ask that you review and respond to the following:

1. Please comment on the general process by which OIRA would employ when reviewing a complex and far-ranging agency proposal such as this.

   **OIRA facilitates interagency review of agencies’ draft proposed and final regulations. While this often occurs after the responsible agency has submitted the draft formally to OIRA, in many cases, that formal submission is preceded by substantial interagency discussion and planning.**

2. During your tenure as Administrator, how long, on average, would OIRA take to review complex and far-ranging agency proposals akin to the above mentioned EPA proposal?

   a. Similarly, do you ever recall during your time as OIRA Administrator where OIRA signed off on an agency rulemaking decision within 3 business days of receiving a rule?

   **During my tenure, the average formal review time for proposed economically-significant EPA rules was 31 days (RegInfo.gov). The shortest review was 0 days (lead NAAQS, subject to a**
judicial deadline), and the longest was 122 days (Revisions to the Spill Prevention, Control, and Countermeasure rule).

3. If then-Administrator Pruitt did indeed sign off on the proposed rule prior to OIRA completing its review, would that violate sections 7 and 8 of Executive Order 12866?

Without knowledge of the discussions between EPA and the Executive Office of the President, it's hard to know. In my experience, decisions on roll out and timing are context-specific, which makes commenting on a hypothetical difficult.

a. What processes are generally in place to prevent agencies from proceeding with rules without OIRA approval?

Communications and repeat interactions between agencies, OIRA, and other parts of the White House generally ensure agencies do not issue rules before review has concluded.

4. Do you believe agencies should use best available evidence in the rulemaking process?


5. Similarly, do you think it is appropriate to restrict the scientific evidence that agencies can consider during rulemaking?

Agencies should rely on well-supported scientific evidence and they should be transparent regarding the models, data, studies, assumptions, and judgments they relied on to translate scientific inputs for use in regulatory analysis.

My comments submitted on the record of EPA’s proposed “Strengthening Transparency in Regulatory Science” offer more detail. They are attached and available on the GW Regulatory Studies Center website.
THE GEORGE WASHINGTON UNIVERSITY

WASHINGTON, DC

Public Interest Comment on
The Environmental Protection Agency’s Proposed Rule
Strengthening Transparency in Regulatory Science
Docket ID No. EPA-HQ-OA-2018-0259; FRL-9977-40-ORD
RIN: 2080-AA14

May 18, 2018
Susan E. Dudley, Director

The George Washington University Regulatory Studies Center

The George Washington University Regulatory Studies Center improves regulatory policy through research, education, and outreach. As part of its mission, the Center conducts careful and independent analyses to assess rulemaking proposals from the perspective of the public interest. This comment on the Environmental Protection Agency’s proposed rule to strengthen the transparency of its regulatory science does not represent the views of any particular affected party or special interest, but is designed to evaluate the effect of the proposal on the reliability of the scientific information underlying EPA’s regulatory decisions.

Introduction

In this proposal, EPA aims to strengthen the transparency of the science it considers “pivotal” to its significant regulatory actions by ensuring that “the data and models underlying the science is publicly available in a manner sufficient for validation and analysis.” It cites existing authorities and policies, but acknowledges, “EPA has not previously implemented these policies and guidance in a robust and consistent manner.”

---

1 This comment reflects the views of the author and does not represent an official position of the GW Regulatory Studies Center or the George Washington University. The Center’s policy on research integrity is http://regulatorystudies.columbian.gwu.edu/policy-research-integrity.

2 Director, the George Washington University Regulatory Studies Center and Distinguished Professor of Practice in the Trachtenberg School of Public Policy and Public Administration.
The rule would not directly regulate non-governmental entities, but instead would require "EPA to ensure that the regulatory science underlying its actions is publicly available in a manner sufficient for independent validation." The preamble says the policy is "designed to provide a mechanism to increase access to dose response data and models underlying pivotal regulatory science in a manner consistent with statutory requirements for protection of privacy and confidentiality of research participants, protection of proprietary data and confidential business information, and other compelling interests." In the long run, through this rule, EPA aims "to change agency culture and practices regarding data access so that the scientific justification for regulatory actions is truly available for validation and analysis."

The proposal defines "dose response data and models" as those used to characterize the quantitative relationship between the amount of dose or exposure to a pollutant, contaminant, or substance and the magnitude of a predicted health or environmental impact. "Pivotal regulatory science" refers to "specific scientific studies or analyses that drive the requirements and/or quantitative analysis of EPA final significant regulatory decisions." "Regulatory decisions" are limited to "significant regulatory actions" as defined in Executive Order 12866. "Regulatory science means scientific information, including assessments, models, criteria documents, and regulatory impact analyses, that provide the basis for EPA final significant regulatory decisions," and research data is used as "defined in Uniform Administrative Requirements, Cost Principles, and Audit Requirements for Federal Awards."

The rule would require EPA to:

1. Clearly identify all regulatory science it relied on in selecting a regulatory action and make those studies available to the public "to the extent practicable."
2. Consistent with laws and sensitive to privacy, confidentiality, and national and homeland security, "ensure that dose response data and models underlying pivotal regulatory science are publicly available in a manner sufficient for independent validation."
3. "Describe and document any assumptions and methods used," including the scientific basis for those assumptions, as well as analysis "showing the sensitivity of the modeled results to alternative assumptions."
4. “Give explicit consideration to high quality studies” that may challenge existing default assumptions including “parametric dose-response or concentration-response models; a robust set of potential confounding variables; nonparametric models that incorporate fewer assumptions; various threshold models across the dose or exposure range; and models that investigate factors that might account for spatial heterogeneity.”

5. “Conduct independent peer review on all pivotal regulatory science used to justify regulatory decisions,” consistent with existing peer review requirements and exemptions.

It is unusual for an agency to provide guidelines such as these through a rulemaking rather than internal guidance; however, the transparency of this rulemaking approach is consistent with the goals of the proposal, and the robustness and legitimacy of any final policy will be enhanced if it is supported by the record the agency develops as it solicits public comment. A 30-day comment period may not provide enough time for constructive input on key issues, however, and a May 12, 2018 memorandum from a working group of EPA’s Science Advisory Board (SAB) argues that this “action merits further review by the SAB.”

EPA also seeks comment on its legal authority for the rulemaking, and the appropriateness of the proposed policies to regulatory decisions developed pursuant to its various statutory mandates. This is important, but this comment will leave those issues to others and focus on the intrinsic merits of the proposal.

**Merits of the Proposed Rule**

1) **EPA will clearly identify and make publicly available the studies and science relied on for significant regulatory actions.**

When the Office of Information and Regulatory Affairs (OIRA) tallies up the total benefits and costs of all federal regulations each year, EPA’s regulations always comprise the bulk of those figures. In OIRA’s most recent draft report, for example, EPA’s estimated annual benefits of the regulations it issued between fiscal years 2007 and 2017 ranged from $196 billion to $707 billion.
billion, constituting 68% to 78% of all the regulatory benefits agencies estimated during that 10-year window. OIRA reported corresponding cost estimates for those EPA rules of $54 billion to $65 billion per year, which comprise between 57% and 69% of all the rules OIRA reviewed during that period.17

Given the significance of these estimates, documenting and making available for public review the underlying science supporting them is essential. EPA estimates that its National Ambient Air Quality Standards (NAAQS), for example, are among the most beneficial regulations issued. In setting NAAQS, EPA is statutorily proscribed from considering the costs of meeting standards, but it must evaluate available data on health and welfare impacts in presenting alternatives for the administrator to consider. EPA’s formulation and presentation of the studies and data necessarily involve judgments about which studies to consider and which to exclude, but these decisions and their rationale are often not transparent.18 Increasing transparency regarding which science it considered; how it weighted and combined individual studies; what competing theories were included, etc. would allow broader review and analysis, and improve the rigor of regulatory decisions.19

EPA’s proposal to clearly identify available studies and make them available for public review is not only important for ensuring decisions are supported by the best information, but also consistent with policies on scientific integrity espoused by previous administrations. OMB’s 2002 Information Quality Guidelines directed agencies to make publicly available “peer-reviewed studies known to the [agency] that support, are directly relevant to, or fail to support any estimate of [risk] effects and the methodology used to reconcile inconsistencies in the scientific data.”20 In 2009, President Obama issued a memorandum to agencies that encouraged “transparency in the preparation, identification, and use of scientific and technological information in policymaking,” and affirmed that “scientific and technological information ... should ordinarily be made available to the public.”21


The George Washington University Regulatory Studies Center • 4
2) EPA will make dose response data and models underlying pivotal regulatory science publicly available for independent validation.

The selection of the model used to estimate responses to different exposures to contaminants can have significant impacts on estimated regulatory benefits. In 2007, OIRA and the Office of Science and Technology Policy (OSTP) observed in a memorandum to agency heads on risk assessment that a “high degree of transparency with respect to data, assumptions, and methods will increase the credibility of the risk analysis, and will allow interested individuals, internal and external to the agency, to understand better the technical basis of the analysis.”

In 2010, OSTP directed agencies to develop policies to “facilitate the free flow of scientific and technical information, consistent with privacy and classification standards.” In a memorandum to department and agency heads, President Obama’s science advisor John Holdren stated:

Open communication among scientists and engineers, and between these experts and the public, accelerates scientific and technological advancement, strengthens the economy, educates the Nation, and enhances democracy. Consistent with the Administration’s Open Government Initiative, agencies should expand and promote access to scientific and technological information by making it available online in open formats. Where appropriate, this should include data and models underlying regulatory proposals and policy decisions.

In a 2013 memorandum, OSTP directed all Executive Departments with greater than $100 million in yearly research and development expenditures to prepare plans for improving the public’s access to the results of that research. EPA did not publish a plan to comply with the OSTP directive until November 29, 2016, years after many other agencies had begun to implement their plans.

EPA’s proposal to make the data and models underlying its pivotal regulatory science public is also consistent with developments in scholarly journals. In recent years, scientific publishing has

focused more on the sharing of data and experimental transparency. Indeed, disclosure of underlying data and computer code has become standard among the more prestigious scientific and technical journals, which allow for data sharing agreements when personally identifiable information prevents public disclosure. These disclosure policies appear to improve the reproducibility of the results of published papers. In 2013, for example, the journal Nature took steps to ensure it reported “key methodological details and encourage[d] authors to be transparent by including the raw data used in their studies.” While recognizing that experimental studies vary, the editors concluded that variation does not preclude “a full report of how a study was designed, conducted and analysed that will allow reviewers and readers to adequately interpret and build on the results.”

The journal Science has also undertaken “initiatives to increase transparency and promote reproducibility in the published research literature... Connected to that progress, and an essential element to its success, an additional focus will be on making data more open, easier to access, more discoverable, and more thoroughly documented.” EPA’s proposal states that it would consider information to be “‘publicly available in a manner sufficient for independent validation’ when it includes the information necessary for the public to understand, assess, and replicate findings.” This emphasis on replicability can encourage challenge and validation that is so important to the scientific method. It is consistent with OMB’s 2002 Information Quality Guidelines, which require that significant information disseminated to the public be “capable of being substantially reproduced, subject to an acceptable degree of imprecision.”

As the Science editors observe, “When the greatest number of creative and insightful minds can find, access, and understand the essential features that led to the collection of a data set, the data reach their highest potential.”

References:

29 Science 2 January 2015: Vol. 347 no. 6217 p. 7
31 OMB 2002, p. 8460

The George Washington University Regulatory Studies Center
3) EPA will describe and document its assumptions and methods and show how sensitive modeled results are to those and alternative assumptions.

This requirement comports with recommendations from various sources. For example, in a recent article, 19 regulatory analysis experts warned:

Analyses that do not provide information on how sensitive the primary estimate is to assumptions, data, and models, and the range of outcomes possible under reasonable alternative analytic assumptions should raise questions. Sensitivity analysis examines different “what if” scenarios to see how changes in key assumptions (or combinations of assumptions) influence estimated outcomes. Because many uncertain factors determine the impact of any regulation, one should look for a convincing justification regarding which uncertain parameters have the most consequential effects on outcomes, and a sensitivity analysis that varies these factors over a reasonable range to gauge their effects on the rule’s net benefits. 33

In 2010, OSTP directed agencies to communicate scientific and technological findings to the public “by including a clear explication of underlying assumptions; accurate contextualization of uncertainties, and a description of the probabilities associated with both optimistic and pessimistic projections, including best-case and worst-case scenarios where appropriate.” 34

This is important, because, as Dudley and Peacock explain, “scientists will never have complete information to predict outcomes with certainty, so analysts rely on what the [National Research Council] 35 calls ‘risk assessment policy’—assumptions, judgments, and rules of thumb—to guide the use of scientific information in analyses that inform policy in the face of uncertainty.” 36 The Institute of Medicine observed in 2013:

Uncertainty is inherent in the scientific information upon which health risk estimates are based. Uncertainties enter the health risk assessment process at


34 Holdren 2010.


every step and can be caused by the potential confounders in observational studies, by extrapolation from animal studies to human studies, by extrapolation from high to low dose exposures, by inter-individual variability, and by modeling the relationships between concentrations, human exposures, and human health responses and evaluating the effect of interventions or risk control options on public health risk.37

Assumptions and judgments made in each of these steps get embedded in predictions of health risk under different policy options. Intentionally or not, they can bias the ultimate advice provided to decision-makers and the public. Documenting those assumptions and estimating how sensitive predicted outcomes are to them and alternative assumptions and judgments could greatly improve the transparency and quality of EPA’s decisions.

Gray and Cohen suggest:

Fundamentally, the EPA should replace risk values that are built on science-policy assumptions with risk estimates that acknowledge underlying uncertainties. For instance, the agency could follow the example of the Intergovernmental Panel on Climate Change and report a range of risks that correspond to different models. Users would then be able to see whether a value is sufficiently precise to support a particular course of action.38

They recognize that policymakers will face more difficult choices when faced with a range of reasonable estimates but argue “that is how it should be.”

The EPA’s definitive values are illusions: they conceal uncertainty that cannot be resolved scientifically. Bringing conflicting value judgements into the open will enable honest debate and improve public health.39

4) EPA will explicitly consider high quality studies that offer new dose-response information that may allow the agency to move away from default assumptions.

In estimating adverse effects of exposure to many pollutants (especially potential carcinogens, but also fine particles), EPA’s default dose-response function is assumed to be linear within the range of exposures under consideration. Both theory and observation suggest that thresholds

39 Gray and Cohen, p. 28.
exist below which further reductions in exposure do not yield changes in mortality response, however more accurate dose-response functions are elusive. The default linear, no threshold assumption is convenient in that it allows EPA to estimate incremental health improvements in proportion to estimated reductions in exposure; but, if it is inaccurate, it can lead to under or over estimates of risks at relevant exposure levels, and to a misallocation of resources.

EPA’s proposed commitment to consider research that can help clarify the effect of low-dose exposure to key pollutants would not only improve short term policy outcomes, but it would also provide incentives for researchers to devote attention and resources to exploring and reducing this key uncertainty. As the 19 regulatory experts observe, “if expected outcomes hinge on the value assumed for a particular uncertain variable, it might be appropriate to gather more information regarding that variable prior to making a decision, or to ask what policies would generate the information necessary to reduce that uncertainty.”

5) EPA will conduct independent peer review on all pivotal regulatory science used to justify regulatory decisions.

Peer review is a fundamental component of the scientific process. Concerns over the extent and rigor of review of important scientific analyses led OMB in 2004 to issue a memorandum establishing guidelines for the use of external peer review at all federal agencies and departments. EPA’s proposed approach is consistent with those guidelines, and the exemptions therein.

When engaging experts in peer review, EPA should consider the recommendations of recent interdisciplinary efforts regarding scientific advisory panels. Such advisors can provide a necessary and valuable source of information and peer review for agency science, but care should be taken in both the composition of the panels and the charges they are given.

An important 2012 Keystone Center report offers a series of recommendations on “the composition of committees that are empaneled to review the science behind a regulatory

---

40 See, for example Texas Commission on Environmental Quality, “PM2.5 Standards may be set Lower than Scientifically Justifiable,” noting that “extrapolations to current exposure levels” can be contrary to the basic principles of toxicology where the biological threshold (a level below which no effect is apparent) is a key concept. http://www.tceq.texas.gov/assets/public/comm_exec/pubs/pdf/020/2013/Outlook-Mar-2013-x.pdf

The George Washington University Regulatory Studies Center ◆ 9
decision.”45 Acknowledging the importance of choosing panelists that “have the knowledge, training, and experience needed to address the charge to the panel,”46 it encouraged agencies “to recognize that all potential panelists will have conscious and unconscious biases.” It recommended, “because biases exist, an agency should strive to engage a wide range of perspectives of qualified scientific experts.”47

Both the Keystone group and a group convened by the Bipartisan Policy Center in 2009 recommend that scientific peer reviewers restrict their advice to matters of science, and not be asked to recommend regulatory policies. EPA’s Clean Air Scientific Advisory Committee (CASAC), for example, is tasked with advising on policy choices, which creates incentives to present policy views as scientific recommendations.48 When drafting charge questions for individual peer reviewers and scientific advisory committees, EPA should be careful to solicit their scientific expertise without encouraging them to blur the lines between scientific expertise and policy judgment.49 As both the BPC and Keystone reports emphasized, the questions posed to experts “should be clearly articulated, and ‘explicitly differentiate, to the extent possible, between questions that involve scientific judgments and questions that involve judgments about economics, ethics, and other matters of policy.”'50

Applications of the Proposed Policies to Existing Regulatory Science

When possible, EPA should apply the new guidelines to existing regulations and the regulatory science that supports them.51 This may not always be feasible, especially for regulatory science

47 Keystone, 2012. p. 15
51 The Keystone Center, 2012: 8. (Internal citation to BPC at 5.)
52 83 FR 18772
that was developed under conditions that would limit disclosure, and EPA should develop clear
criteria for the types of research that would be eligible for exemptions to its transparency policy.

Nevertheless, EPA has an opportunity, when conducting retrospective evaluation of regulatory
impacts (as required by Executive Orders 13563 and 13771), to reexamine the studies, data,
models and assumptions that generated ex-ante estimates of regulatory benefits and costs. Such
retrospective review can provide data that either corroborates or raises questions about the
assumptions on which previous estimates were based. As such, ensuring transparency and
opportunities for independent analysis and evaluation could be particularly valuable in
retrospective review, not only for decisions regarding continuation of existing policies, but also
for improving the quality of the science used to design new policies.53

For regulatory programs, like the NAAQS, that periodically review and update standards based
on a record that has been built over decades, application of the proposed transparency and
integrity procedures to that record, when feasible, could allow a broader group of experts access
to pivotal regulatory science, including data, models, and assumptions. Such review would be
consistent with EPA’s statutory mandate for review under the Clean Air Act, and allow EPA to
make future decisions on better data.

Protecting Privacy and Confidentiality

EPA acknowledges concerns that increased transparency and public access to data may risk
exposing confidential or private information, but it points to practices at other federal agencies
and in scientific publishing that can ensure confidential or personally-identifiable information is
not disclosed. Lutter and Zorn review some of these practices and conclude that, depending on
the situation and sensitivity of the information, a range of measures is available to share data in a
way that allows access for replication and validation purposes while protecting personally
identifiable information.54

The SAB working group memo raises concerns that “for studies already completed or underway,
the participation of human subjects is undertaken according to terms approved by the cognizant
IRB” (institutional review board) which may constrain data sharing.55 Input from the SAB and
public comments may elucidate specific concerns, as well as methods EPA can use to address
them.

54 Randall Lutter and David Zorn, “On the Benefits and Costs of Public Access to Data Used to Support Federal
55 Cullen, 2018.
Conclusion

Regulations aimed at addressing public health and environmental risks depend heavily on scientific information. Regulatory impact analyses often hinge on assessments of risk that necessarily involve assumptions and judgments but often generate precise-sounding predictions that hide not only considerable uncertainty about the actual risk, but hidden policy judgements. A lack of transparency surrounding these judgments harms the credibility of scientific advice and results in poorer policy decisions.

Former EPA scientist Robert T. Lackey cautions against what he calls “normative science”:

Science should be objective and based on the best information available. Too often, however, scientific information presented to the public and decision-makers is infused with hidden policy preferences. Such science is termed normative, and it is a corruption of the practice of good science. Normative science is defined as “information that is developed, presented or interpreted based on an assumed, usually unstated, preference for a particular policy choice.”

EPA’s proposal to strengthen the transparency of its regulatory science includes reasonable steps that could improve the evidential basis for its regulatory policies, and thus improve regulatory outcomes by targeting resources to where they can achieve the largest benefits. As President Obama’s science advisor observed, “open communication among scientists and engineers, and between these experts and the public, accelerates scientific and technological advancement, strengthens the economy, educates the Nation, and enhances democracy.”

The requirements proposed here are not a radical departure from existing guidelines. For example, since 2004, OMB has directed agencies to issue information quality guidelines to, among other things, ensure the objectivity of information, including “a high degree of transparency about data and methods to facilitate the reproducibility of such information by qualified third parties.” President Obama in 2011 encouraged an “open exchange of information and perspectives among State, local, and tribal officials, experts in relevant disciplines, affected stakeholders in the private sector, and the public as a whole, ... including relevant scientific and technical findings.”

---

56 Dudley and Peacock, 2017, p. 36.
58 Holdren memo, 2010.
Greater transparency in the studies, models, assumptions, and risk assessment policy choices used in regulatory decisions could encourage more open, constructive debate on those choices. The scientific method depends on falsifiable hypotheses, data gathering, replication, dissent, and challenge, to ensure objective analysis and to minimize bias in the interpretation of results. Greater transparency is an essential step in improving scientific integrity. Clearer explanations regarding the policy rationales for choosing one set of assumptions or models over another would encourage more openness and constructive discussion about science and policy, improving the ultimate policy decision and engendering greater acceptance of that policy choice.

61 Open Data Initiative https://www.whitehouse.gov/open
Sally Katzen
Post-Hearing Questions for the Record
“From Beginning to End: An Examination of Agencies’ Early Public Engagement and Retrospective Review”
From Senator Thomas R. Carper
For the Honorable Susan Dudley and for the Honorable Sally Katzen

OIRA’s role in reviewing agency rulemaking and EPA’s “Strengthening Transparency in Regulatory Science”

On May 9, 2018 I joined Senator Hassan and several other Senators in a letter to then-OIRA Administrator Neomi Rao raising concerns and seeking answers as to how OIRA reviewed an EPA proposed rulemaking entitled: “Strengthening Transparency in Regulatory Science.”

According to Reginfo.gov, OIRA received the EPA’s proposed rule on Thursday, April 19, 2018. Then-EPA Administrator Scott Pruitt publicly announced this proposed rule and signed it three business days later on Tuesday, April 24, 2018. Reginfo.gov initially stated that OIRA’s review was completed on Wednesday, April 25, 2018 – the day after Administrator Pruitt announced and signed the rule. Later, following press inquiries, Reginfo.gov was changed to indicate that OIRA’s review was completed on Monday, April 23, 2018.

Our letter sought detailed information on the scope and substance of OIRA’s review of this proposed rulemaking. However, in a response to Senator Hassan on July 12, 2018, former Administrator Rao provided little in way of response and simply noted that the conclusion date for the OIRA review of this proposed rule of April 25, 2018 was due to a clerical error that was subsequently corrected.

With this in mind, I ask that you review and respond to the following:

1. Please comment on the general process by which OIRA would employ when reviewing a complex and far-ranging agency proposal such as this.

A: There is substantial variability in OIRA’s review of draft proposed and final rules, depending on, among other things, the significance and complexity of the issues being addressed, the potential interests and equities of other agencies, the need for expedition (e.g., statutory or judicial deadlines or imminent health or safety concerns), and the extent to which OIRA staff had been informed or involved during the agency drafting process. Typically, when a draft is received, it is reviewed by a desk officer at OIRA and is also sent to other agencies for their comments. The process can move quickly if no concerns are raised and can be quite extended if any of the reviewers has serious issues that need to be addressed or resolved. The applicable Executive Order provides for a 90-day review time, which is generally adhered to; there are many examples of drafts being reviewed in substantially less and substantially more time.
2. During your tenure as Administrator, how long, on average, would OIRA take to review complex and far-ranging agency proposals akin to the above mentioned EPA proposal?

A: During my tenure, the average time for review of complex and far-ranging proposals would typically be closer to (or even exceed) the 90-day limit, especially where there was significant interest in the proposal by other agencies with equities in the matter. Averages do not, however, reflect what might be the appropriate review time for a particular proposal.

a. Similarly, do you ever recall during your time as OIRA Administrator where OIRA signed off on an agency rulemaking decision within 3 business days of receiving a rule?

A: While data on review times may be available from OIRA, I can recall a very limited number of instances where the review time was quite abbreviated – including for example, DOT’s response to several deaths caused by airbags’ expanding with infants or short-statured women in the passenger seat. As a rule, however, we took the time necessary to conduct a meaningful review, including the inter-agency process.

3. If then-Administrator Pruitt did indeed sign off on the proposed rule prior to OIRA completing its review, would that violate sections 7 and 8 of Executive Order 12866?

A: If the facts were as alleged, that would be inconsistent with the terms of Executive Order 12866.

a. What processes are generally in place to prevent agencies from proceeding with rules without OIRA approval?

A: During my tenure as Administrator, we worked with the agencies to ensure that did not occur, and we had the support of the Director of OMB when necessary. On the one occasion that I recall where an NPRM for a significant regulation was released by the agency without having been approved by OIRA, the Secretary (or the General Counsel) of the Department immediately withdrew the proposal and submitted the draft to OIRA for review.

4. Do you believe agencies should use best available evidence in the rulemaking process?

A: Yes, unequivocally.
5. Similarly, do you think it is appropriate to restrict the scientific evidence that agencies can consider during rulemaking?

A: All available relevant and authoritative scientific evidence should be considered in a rulemaking; if certain material or studies are considered less trustworthy or probative the reasons for that judgment should be set forth in the preamble to the proposal.