

MEETING  
STATE OF CALIFORNIA  
ENVIRONMENTAL PROTECTION AGENCY  
OFFICE OF ENVIRONMENTAL HEALTH HAZARD ASSESSMENT  
ENVIRONMENTAL CONTAMINANT BIOMONITORING PROGRAM  
SCIENTIFIC GUIDANCE PANEL

CALIFORNIA DEPARTMENT OF PUBLIC HEALTH  
AUDITORIUM  
850 MARINA BAY PARKWAY  
RICHMOND, CALIFORNIA

THURSDAY, JULY 20, 2017  
10:01 A.M.

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A P P E A R A N C E S

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Megan R. Schwarzman, M.D., M.P.H., Chairperson

Carl Cranor, Ph.D., M.S.L.

Marion Kavanaugh-Lynch, M.D., M.P.H.

Ulrike Luderer, M.D., Ph.D.

Penelope (Jenny) Quintana, Ph.D., M.P.H.

CALIFORNIA ENVIRONMENTAL PROTECTION AGENCY:

Gina Solomon, M.D., M.P.H., Deputy Secretary for Science and Health

OFFICE OF ENVIRONMENTAL HEALTH HAZARD ASSESSMENT:

Lauren Zeise, Ph.D., Director

Allan Hirsch, Chief Deputy Director

Amy Dunn, M.P.H., Research Scientist III, Safer Alternatives Assessment and Biomonitoring Section

Carl DeNigris, Staff Counsel

Alexander Hoepker,

Shoba Iyer, Ph.D., Safer Alternatives Assessment and Biomonitoring Section

DEPARTMENT OF PUBLIC HEALTH:

Robin Christensen, M.S., Biomonitoring California Grant Coordinator, Sequoia Foundation

Jianwen She, Ph.D., Chief, Biochemistry Section, Environmental Health Laboratory

A P P E A R A N C E S C O N T I N U E D

DEPARTMENT OF PUBLIC HEALTH:

Nerissa Wu, M.P.H., Ph.D., Chief, Chemical Exposure  
Investigations Unit, Environmental Health Investigations  
Branch

DEPARTMENT OF TOXIC SUBSTANCES CONTROL:

Myrto Petreas, Ph.D., M.P.H., Chief, Environmental  
Chemistry Branch

GUEST SPEAKERS:

Axel Adams, M.S., University of California, San Francisco

Roy Gerona, Ph.D., University of California, San Francisco

Jon Sobus, Ph.D., National Exposure Research Laboratory,  
US Environmental Protection Agency

ALSO PRESENT:

Susan JunFish, M.P.H., Parents for a Safer  
Environment (PASE)

Aolin Wang, Ph.D., University of California, San Francisco

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## P R O C E E D I N G S

1  
2 MS. KAUFFMAN: Good morning. I'm Duyen Kauffman  
3 from the Office of Environmental Health Hazard Assessment.  
4 And we are going to start the meeting. So today's meeting  
5 is available via webinar. And for the speakers we do ask  
6 that you please speak directly into the microphone and  
7 introduce yourself before speaking. This is for the  
8 benefit of the people participating via the webinar and  
9 for the transcriber.

10 The materials for the meeting were provided to  
11 SGP members and posted on the Biomonitoring California  
12 website. A small number of copies of the meeting  
13 materials are available at the table near the entrance of  
14 the auditorium. A sample SGP packet is also available for  
15 viewing at this table.

16 We will break at 12:45 p.m. for lunch and take  
17 another short break at 3:00 p.m.

18 The location of the restrooms are at the opposite  
19 end of the hallway where you enter the auditorium. And  
20 emergency exits are here at the front of the room, on  
21 either side, and also as you exit the auditorium on either  
22 side. And with that, I'd like to introduce Dr. Lauren  
23 Zeise, Director of the Office of Environmental Health  
24 Hazard Assessment.

25 DIRECTOR ZEISE: Good morning, everyone. I'd

1 like to welcome you to this meeting of the Scientific  
2 Guidance Panel for the California Environmental  
3 Contaminant Biomonitoring Program, also known as  
4 Biomonitoring California. So welcome to the Panel and to  
5 the audience in the room, and those attending on the web.  
6 And an early thank you for your participation in sharing  
7 your expertise with us.

8           So just to give a bit of a recap, SGP met on  
9 March 8th in Sacramento. And the Panel meeting was  
10 followed by a special event cele -- to celebrate the  
11 program's 10-year anniversary. Just to briefly recap the  
12 March meeting itself, the panel heard an update about our  
13 current program activities, and provided extensive input  
14 on the design of the multi-regional study. And that's now  
15 in our planning phases -- in the planning phase.

16           The Panel also heard a presentation from CDC, Dr.  
17 Breysse, the Director of the National Center for  
18 Environmental Health at CDC. And I heard about  
19 biomonitoring at a national level, and had a very good  
20 discussion with him about synergies between State and  
21 national efforts.

22           And then finally, we had three distinguished  
23 speakers kick off the discussion about the future program  
24 activities. That was Dr. Irva Hertz of UC Davis, Tom  
25 Webster of Brown University and Julia Brody of Silent

1 Spring, and then Dr. Gina Solomon, the CalEPA Deputy  
2 Secretary for Science and Health led the discussion.

3           Some of the key recommendations included that the  
4 Program should strategize on resource-efficient ways to  
5 choose studies that advance multiple program goals; expand  
6 efforts to generate data to evaluate regulatory actions  
7 and other interventions to reduce chemical-specific  
8 exposures; maintain the Program's strong efforts  
9 communicating study results to participants; and expand  
10 efforts to reach out to policymakers; and to continue to  
11 develop capacity for non-targeted analyses for identifying  
12 emerging chemicals of concern, and guiding targeted  
13 biomonitoring studies. And we're going to hear more about  
14 that topic later today.

15           So if you'd like more details, you can refer to  
16 the summary of the March meeting on our website, but we  
17 also have transcripts available on our website,  
18 Biomonitoring California website.

19           So just to conclude the recap of the March  
20 activities, it was a very nice event we had after the  
21 meeting to celebrate the Program's 10 years. It was a  
22 perfect way to celebrate the milestones, and we really  
23 appreciate your participation in that.

24           Now, looking forward to today's activities, we're  
25 going to start with some Panel business. As some of you

1 may know, Dr. Asa Bradman has resigned as the SGP Chair  
2 and Panel member. And so just before lunch, we're going  
3 to have a tribute to Dr. Bradman. He's going to come in  
4 and join us, and we're going to thank you -- thank him for  
5 his service.

6           Second, I have the pleasure of announcing the  
7 appointment to the Panel by Governor Brown of Dr. José  
8 Suárez. He's an Assistant Professor in the Department of  
9 Family Medicine in Public Health at UC San Diego. Due to  
10 a prior commitment, he's not going to be able to attend  
11 today's meeting, but he'll be formally sworn in and  
12 participate in our next meeting, which will be November  
13 9th here in Richmond.

14           So finally, it's my pleasure to introduce to you  
15 our new SGP Chair, Dr. Meg Schwarzman. Dr. Schwarzman has  
16 been a member of the Panel since 2014, and we're grateful  
17 she's agreed to be -- take on this additional  
18 responsibility. And so for those who -- of you who don't  
19 already know her, Dr. Schwarzman is an environmental  
20 health scientist at the UC Berkeley Center for  
21 Occupational and Environmental Health, and the Associate  
22 Director of Health and Environment at the Berkeley Center  
23 for Green Chemistry.

24           This past spring she designed and taught a new  
25 graduate class in environmental health policy at UC

1 Berkeley. And she recently returned from Alaska where she  
2 gave four talks, two radio interviews, and a webinar on  
3 environmental contributions to breast cancer.

4 So just this week Dr. Schwarzman advised the  
5 Department of Toxic Substances Control on the  
6 implementation of the Safer Consumer Products Program in  
7 her role as member of the Green Ribbon Science Panel.

8 Finally, Dr. Schwarzman received her medical  
9 degree from the University of Massachusetts Medical School  
10 and completed specialty training in family medicine at  
11 UCSF General Hospital, and received her MPH from UC  
12 Berkeley.

13 So please join me in welcoming Dr. Schwarzman to  
14 her new role as your Panel Chair.

15 (Applause.)

16 DIRECTOR ZEISE: So now I'll turn the meeting  
17 over for facilitation by Dr. Schwarzman.

18 PANEL MEMBER SCHWARZMAN: Thank you. And  
19 welcome, everybody. And I'm honored to follow in the  
20 footsteps of several Panel members who have preceded me as  
21 Chair. And I'll do my best to conduct the meetings as  
22 well as they have in the past.

23 So I want to start with the goals for the  
24 meeting. The first is our Program updates from Dr.  
25 Nerissa Wu, and then we'll go on to welcome our guest

1 speaker Dr. Jon Sobus of US EPA, who is here to discuss  
2 advances in non-targeted analyses.

3           We're going to -- there's a quick switch in the  
4 agenda. So after lunch, we're going to consider the class  
5 organophosphorus pesticides as potential designated  
6 chemicals. And then after the break, we'll then hear from  
7 Dr. Roy Gerona of UC San Francisco about his work  
8 developing and applying an analytical method to measure  
9 glyphosate in urine.

10           We just made that switch this morning -- I'm  
11 sorry. We'll wait to hear whether that switch is  
12 finalized or not. Okay. It's back and forth this  
13 morning.

14           Okay. So for each agenda topic, we do, as usual,  
15 provide time for both Panel questions, public comment, and  
16 Panel discussion and input. And so the way public comment  
17 works, for those who haven't been there before, if you  
18 would like to comment on an agenda item, you can fill out  
19 a comment card. And Amy -- someone usually has those.

20           Oh, they're on the table near the -- at the  
21 entrance of the auditorium. You can turn the cards into  
22 Duyen Kauffman, and if you're joining the meeting via  
23 webinar, and you want to provide comments, you can do so  
24 by email, and the email comments relevant to the topic  
25 under discussion will be read aloud during the meeting.

1           We do subject public comments to time limits, if  
2 needed, depending on how many there are, and we'll divide  
3 the time allotted equally among the individuals who are  
4 wishing to talk about that agenda item. So please keep  
5 comments focused on the agenda topics that are being  
6 presented. And we also include an open public comment  
7 session at the end of the day for more general comments.

8           So with that, I'd like to introduce Dr. Nerissa  
9 Wu, who is the Acting Chief of the Exposure Assessment  
10 Section at the California Department of Public Health,  
11 where we are. And is also the acting lead for  
12 Biomonitoring California.

13           So Dr. Wu is going to provide an update on  
14 Program activities.

15           (Thereupon an overhead presentation was  
16 presented as follows.)

17           DR. WU: Well, welcome, everybody, panelists, and  
18 guests and Dr. Schwarzman in your new role. It's great to  
19 have you up there.

20           How is that?

21           Okay. Great. Im going to be giving the customary  
22 overview of the Program updates, and including some  
23 administrative news and project updates. And then as  
24 always -- I can just speak into -- or I can hold this

25           So I am giving the customary Program update

1 today, administrative news, project updates, and then as  
2 we have in the last couple of Panel meetings talking more  
3 about the protocol we're developing for the multi-regional  
4 sampling plan, which is now going by its name of the  
5 California Regional Exposure, or CARE, Study.

6 --o0o--

7 DR. WU: But first I want to talk a little bit  
8 about staff changes. We have had a number of staff  
9 transitions in the past few months. We've lost some  
10 staff. You might not know their names. They may not have  
11 presented to the SGP, but they're all part of the  
12 biomonitoring fabric and have all contributed in their own  
13 way to the Program: Susannah McKay, Suwati Anand,  
14 ShiZhong Wang, Sissy Petropoulou, Julie Frankenfield,  
15 Astrid Zamora, Joginder Dahliwal. And I just want to say  
16 thank you to those staff who have moved on and wish them  
17 well in their new positions.

18 We do have some new staff and they're highlighted  
19 on this slide in red. Lauren Baehner, Elizabeth Hall,  
20 Juliet Kinyua, Let Zhang, Robin Christensen, who of course  
21 is not new. You're familiar with Robin. She's been here  
22 many years, as our CDC grant coordinator, but she has  
23 joined State staff. I'm very happy to say she's part of  
24 the CDPH management team for biomonitoring, which is  
25 awesome, and Suzanne Wittwer who is our new CDC grant

1 administrator, taking Robin's place.

2 --o0o--

3 DR. WU: You've seen this slide before. This is  
4 our Program budget. With the end of fiscal year 2016-17,  
5 we have come to the end of our supplemental environmental  
6 justice funding. We've also come to the end of some  
7 limited term positions, which were established as part of  
8 a budget change proposal, or BCP. These are limited term  
9 positions that ended in June 2017, which is why we see  
10 this precipitous drop in our budget between '16-'17 and  
11 the current fiscal year.

12 We do have additional limited term positions,  
13 also created by a budget change proposal, which are  
14 scheduled to expire at the end of this fiscal year, June  
15 2018. At this point, we don't seem to have a mechanism  
16 for extending those positions or making them permanent.

17 So this downward slope is projected to continue  
18 for the following fiscal year. And then in 2019 when our  
19 cooperative agreement with CDC comes to an end, we see  
20 that our budget will be less than half of what it -- of  
21 what we've had over the past few years.

22 We've talked about our budget issues in the past,  
23 and there's not a whole lot to say about it. Our staff  
24 continues to work very hard to be as effective and  
25 efficient as possible and to accomplish what we can with

1 the budget resources that we have. You'll hear throughout  
2 my description of different projects that there are  
3 choices that have to be made as a program. We just need  
4 to -- there are things that we can and can't do given the  
5 resources that we have.

6 --o0o--

7 DR. WU: Now, I want to turn to some of our  
8 project updates, starting with project BEST. And this is  
9 the Biomonitoring Exposures Study in the Central Valley in  
10 collaboration with Kaiser. We have finished analysis for  
11 the panels that were part of the original project plan.  
12 The perchlorate data is now ready to be returned to  
13 participants.

14 Those packets are actually in -- they're being  
15 created right now, and they should go out in the next week  
16 or two. Earlier this year we finished arsenic re-testing  
17 for participants who were found to have elevated arsenic  
18 levels. And those results were returned. And we reported  
19 out on that a couple sessions ago. We do have 56  
20 remaining urine samples, which were never tested for  
21 metals. And because about 13 percent of our participants  
22 were found to have inorganic arsenic levels at the level  
23 of concern, we would like to go back to those untested  
24 urine samples and test them for arsenic, identify what  
25 their exposure source might be, and get that information

1 to those participants. It seems like the right thing to  
2 do. So we will keep you updated as we proceed with that.

3 --o0o--

4 DR. WU: For the Foam Replacement and  
5 Environmental Exposure, or FREES, Study. This is the  
6 study in which we're looking at a time point before foam  
7 and furniture replacement, and then at 6, 12, and 18  
8 months after the foam or furniture is replaced from the  
9 home. We did recruit over several months, and then  
10 participants took varying amounts of time to replace their  
11 furniture or the foam in their homes. And so the time  
12 point of replacement, which spanned from October 2015 to  
13 November 2016, is quite spread out, which means their  
14 sampling at 6, 12, and 18 months, is also very spread out.

15 We have gotten all of our initial samples and  
16 returned those results to participants. We have completed  
17 the six-month sampling and are about to return about 15 of  
18 those results to participants. And we're about halfway  
19 through the 12-month sampling, and have just started the  
20 18-month sampling point for some of our participants. And  
21 we anticipate that samples will be completely collected by  
22 the spring of 2018.

23 We're looking at the change in flame retardants  
24 in blood and urine following foam replacement or furniture  
25 replacement. We know that foam -- that flame retardant

1 levels are decreasing in the environment, and that they're  
2 fluctuating over time and people as well. And so we  
3 wanted to have a control group to compare to this FREES  
4 cohort.

5 So we have recruited non-FREES participants,  
6 people who are not changing out the foam -- the furniture  
7 or foam in their homes. And their blood and urine samples  
8 were also collected in August 2016, and we're about to go  
9 back to those participants. So that will have a one year  
10 control group to compare to our FREES cohort.

11 --o0o--

12 DR. WU: In the Asian/Pacific Islander Community  
13 Exposures, or ACE, Project, this is our Chinese population  
14 in San Francisco, for which we've recruited and collected  
15 samples from 100 participants. And we're looking at lead,  
16 cadmium, arsenic and mercury, as well as the per and  
17 polyfluorinated -- fluoroalkyl substances, the PFAS  
18 compounds.

19 We are actually in the results return phase of  
20 this project. We're calling notif -- we're calling  
21 participants with notifications of elevated metal levels.  
22 We have had one participant found with an elevated blood  
23 mercury level. And of our 100 participants, 54 of them  
24 had urinary arsenic levels at the level of concern, which  
25 triggers our speciation arsenic protocol, where we

1 speciate into inorganic and organic arsenic, and take a  
2 look at the profile of the arsenic species to determine  
3 where that arsenic might be coming from.

4           Twenty-six of the participants actually had  
5 inorganic arsenic levels at the level of concern of 20  
6 micrograms per liter. And those results will be followed  
7 up with notification to the participants, and collection  
8 of some additional exposure information, so we can work  
9 with participants to really identify their exposure  
10 source.

11           We're about to complete those calls. All results  
12 will be going to all of our 100 participants the next  
13 month, August 2017. And then following that, we're having  
14 an open community meeting for the general public in  
15 September in conjunction with APA Family Services, our  
16 community partner, to talk about the significance of the  
17 findings to the community as a whole.

18                           --o0o--

19           DR. WU: Now, I want to talk about some of the  
20 activities we've been able to accomplish under our  
21 environmental justice funding for this last year. The  
22 funding enabled us to extend our ACE Project, that I've  
23 just described, into a second community. And we've worked  
24 with the Vietnamese Voluntary Foundation, or VIVO, to  
25 recruit participants from the Vietnamese community in San

1 Jose.

2           So again, it's 100 participants, looking at those  
3 four select metals and PFAS compounds. And with our  
4 community partner and our recruitment team, we were able  
5 to recruit 100 participants in about two months. And so  
6 those samples are already at the lab, and we should have  
7 some results to report out next -- by the end of this  
8 year.

9           We also have the East Bay Diesel Exposure  
10 Project, which is being led by OEHHA and UC Berkeley,  
11 that's scheduled to start up this fall. And this study it  
12 will look at toddler and parent pairs over two different  
13 seasons in different neighborhoods of the East Bay. It  
14 will analyze urine for 1-NP, the diesel marker which we've  
15 discussed here with Dr. Chris Simpson.

16           And those results are going to be combined with  
17 exposure survey data, activity trackers, environmental  
18 samples, and a really cool mapping app, which will help  
19 under -- improve our understanding of diesel exposure.  
20 And we'll be hearing more about that at our November SGP  
21 meeting, which will focus on environmental justice.

22           We've also been able to do some community  
23 outreach and engage with community partners across the  
24 state. We've collected information from 84 different  
25 organizations around the state, including at least one

1 from each of our eight California regions. And we've  
2 talked to community groups about what are your concerns,  
3 what are your priorities, and how can Biomonitoring work  
4 with you to move your agenda forward?

5 We've heard a lot about air quality, concerns  
6 about diesel exhaust in particular, pesticides, arsenic in  
7 water, gas and oil extraction. We've actually just gotten  
8 the report back from that team. And so we'll be digesting  
9 it and hopefully reporting back a little more in November.

10 We've also been able to initiate the  
11 Biomonitoring Matters Newsletter, which I think I  
12 highlighted at our last meeting. Our first issue has gone  
13 out. It's been distributed at different community events.  
14 It will be available in Spanish in the next month. And we  
15 are coming out with our second issue this fall. Both the  
16 newsletter and the community listening sessions I think  
17 are projects that help move our agenda forward, help us  
18 learn about what can be useful about biomonitoring to the  
19 community, and help bring information about the program to  
20 a wider audience, and we really hope to be able to  
21 continue these efforts despite the fact that this resource  
22 has come to an end.

23 --o0o--

24 DR. WU: And now I'd like to turn our attention  
25 to the California Regional Exposure Study, formerly known

1 as the Multi-Regional Sampling Plan. And just a brief  
2 reminder of what the CARE Study is. We are required, by  
3 our founding legislation, to conduct statewide sampling.  
4 And this was initially conceived as CalHANES with a budget  
5 of approximately \$12 million annually. The Program was  
6 never funded at that level.

7 So we've decided to approach the statewide  
8 sampling on a regional basis, dividing the state into  
9 these eight regions that you see here, and conducting the  
10 sampling on a region-by-region approach.

11 And this modular approach gives us more  
12 flexibility to adjust to our budget fluctuations, if we  
13 have a year where we can accomplish more than one region;  
14 great. But at our -- under our current budget scenario,  
15 we're looking at one region per year, collecting 300 to  
16 500 samples per region.

17 We are going to biomonitor for certain metals:  
18 mercury, arsenic, lead, and cadmium, but also molybdenum,  
19 antimony, cobalt, manganese, and uranium and for the PFAS  
20 compounds across the State. But this modular approach  
21 will also allow us to expand the parameters of the study.  
22 If there is perhaps an analytical panel of particular  
23 interest in some region, and we have the resources to do  
24 so, we can expand the study to accommodate additional  
25 panels or additional studies, intervention studies or

1 nested studies, within the CARE Study construct.

2 --o0o--

3 DR. WU: So one of the most important tasks of  
4 the study, and one of the first things we did was to try  
5 to develop a message and a study name that would  
6 adequately convey what we're trying to do, and the  
7 importance of the Biomonitoring Program, but also be  
8 accessible and understandable to the public, and be  
9 interesting enough so that it motivates them to pick up  
10 our recruitment materials and take a second look, maybe  
11 sign up for the study.

12 So we worked internally between our departments  
13 to come up with our message, what is it we're trying to  
14 say, what can Biomonitoring do for you, what it can't do.  
15 We want to be careful not to overpromise or overstate what  
16 biomonitoring can do.

17 And then we worked with communication experts to  
18 try to hone this into a message into something that would  
19 be -- that would attract attention, that would pique  
20 people's interest. And we conducted a series of focus  
21 groups, both in English and Spanish, in L.A. County to  
22 talk to people about, you know, what do you think of this  
23 image, or this name, or this motivational message? Do you  
24 like it? Do you hate it? Does it remind you of anything?  
25 What does it make you think of?

1           And our focus group participants overwhelmingly  
2 indicated to us that they would be most motivated to join  
3 a study to find out what chemicals are in their body, but  
4 also in learning ways to reduce their exposure. So we  
5 took those -- those motivational statements and rephrased  
6 it as a question, kind of to draw people into being  
7 interested in our study, but also because there is  
8 uncertainty about what a biomonitoring study will find.

9           We also wanted to link this to health. Again,  
10 not overstating what we might find in the study. And  
11 we've come up with the tag line of, "Are there chemicals  
12 in your body that could harm your health? Join the CARE  
13 Study to learn about lead and other chemicals in your  
14 body, and actions you and your family can take to help  
15 reduce your contact with these chemicals".

16           We have a longer message that's included on some  
17 of our fliers and other promotional materials.  
18 "Biomonitoring California is doing this study to measure  
19 chemicals in people across the state. This information  
20 will support efforts to reduce chemical exposure in  
21 Californians and improve public health". We do have a  
22 shorter version of this message.

23                           --o0o--

24           DR. WU: And I should point out, as I show some  
25 of our -- some of our design work, that nothing is

1 finalized. We're still tweaking it with graphic artists,  
2 but this is just a mock-up of what our materials may look  
3 like. So this message of are there chemicals in your body  
4 that could harm your health, learn more by joining the  
5 CARE study, will appear on their postcard.

6           There's very little text, because again the point  
7 is to really just get somebody's interest, get them to  
8 look at the postcard and want to find out more. We do  
9 have this short statement translated into 10 different  
10 languages. We've looked at the most common languages  
11 spoken in households across L.A. County. So we've  
12 included English, Spanish, Chinese, Tagalog, Korean,  
13 Armenian, Vietnamese, Persian, Japanese, and Russian.

14           And the point of the postcard, regardless of how  
15 somebody gets the postcard, whether it's handed to them at  
16 a community meeting or they get it in the mail, is to get  
17 people to go to the pre-screening survey, which will be  
18 available online. We'll also have a phone number on here,  
19 so if somebody doesn't want to enter data online, they can  
20 call us and we'll help them fill out that pre-screening  
21 survey.

22                           --o0o--

23           DR. WU: This is a mock-up of the website where  
24 you would go to get more information about the study.  
25 There will be frequently asked questions and answers. And

1 you see on the right, this is -- if you're interested in  
2 joining the CARE Study, the pre-screening survey is  
3 available in those ten languages that I've just described.  
4 You can click on that and go to a pre-screening survey,  
5 which will collect basic demographic information, a little  
6 bit more about the participants interested in joining up.

7 --o0o--

8 DR. WU: We will still have our Biomonitoring  
9 California website, with which you're familiar, so that if  
10 somebody is looking for biomonitoring information and  
11 learns about the CARE Study, they can click over easily to  
12 our project page. And conversely, if a participant is  
13 reading about the study and wants to learn more about  
14 biomonitoring, they can easily go to the Biomonitoring  
15 webpage and learn more about the Program as a whole.

16 --o0o--

17 DR. WU: So from this pre-screening pool, this  
18 pool of people who have filled out that pre-screening  
19 survey, we'll be selecting 500 participants, matching L.A.  
20 demographics as best we can, particularly in terms of  
21 race. L.A. County is already divided up into eight  
22 service planning areas by the L.A. County Department of  
23 Public Health. And we'll try to include participants from  
24 across all of these SPAs.

25 As people are invited to enroll, they'll receive

1 a packet of information that will include a letter of  
2 introduction, the informed consent, the exposure survey,  
3 and a calendar, which will show them where and when we'll  
4 be collecting samples, so that they can make an  
5 appointment.

6 We are going to be working with a Salesforce  
7 platform, which is a very sophisticated cloud participant,  
8 or data tracking, software. So we'll be able to run  
9 reports in real time and look at metrics like what  
10 percentage of people have responded to the postcard, and  
11 what are the demographics in the prescreening pool, and  
12 where do we need to be doing extra recruitment?

13 And so we'll be using that to select our  
14 participants and figure out what the demographics are as  
15 people go through the study of how -- are people dropping  
16 out at a different points, and do we need to add more  
17 people?

18 --o0o--

19 DR. WU: We have three parallel pathways of  
20 participation. We've got our internet -- internet  
21 participants. And we hope that most people will be  
22 participating via the internet. Internet participants  
23 will be getting their packet via email. And they'll have  
24 a log in, a secure log in, to the Salesforce platform,  
25 where they can complete their informed consent, complete

1 the survey, and make an appointment for sample collection  
2 all online.

3 We also have paper participants, so people who  
4 fill out the pre-screening and say, "I don't want to do  
5 this online. I'd like a paper version of the materials."  
6 We'll send them the informed consent and the completed  
7 survey to them in paper through the mail. They will fill  
8 it out on paper and mail it back to us. And then we'll  
9 make a phone call to make that appointment with them.

10 For people who indicate to us that they'd like to  
11 complete the study in a language other than English or  
12 Spanish, either by filling out the pre-screen in a  
13 different language or by indicating on the English or  
14 Spanish pre-screening that they would like language  
15 support, we will send them the packet, but then call with  
16 the help of an interpreter to make an appointment. Then  
17 we will complete the informed consent and the survey and  
18 complete sample collection all at that one appointment  
19 with the assistance of the interpreter. Again, this is  
20 all going to be tracked through Salesforce, so that we can  
21 keep good tabs on our data.

22 --o0o--

23 DR. WU: Salesforce will also enable us to send  
24 out automated reminders, so don't forget to fill out your  
25 informed consent, don't forget to make your appointment,

1 you have an appointment in the next couple days. We're  
2 hoping to minimize our losses of participants by having  
3 this reminder protocol. And for non-internet  
4 participants, the system will ping the staff and we'll be  
5 able to make those reminders manually.

6 --o0o--

7 DR. WU: Once participants get to the point of  
8 booking their appointment, this is either on the internet  
9 or on the phone, they'll make an appointment for sample  
10 collection. We will have sample collection events across  
11 L.A. County to make it as convenient as possible for our  
12 participants. Each appointment includes a brief survey,  
13 so activities and diet over the last three to five days, a  
14 quick blood draw, and a urine sample collection.

15 Field staff are going to be in L.A. County for 12  
16 to 15 weeks, during which time they'll be holding three to  
17 four events per week at these -- at these centralized  
18 sites. And then for participants who cannot make it to a  
19 site, either they're not convenient, the participant isn't  
20 mobile, the field staff will be reaching out to them to  
21 make home visits or to arrange a more convenient time for  
22 sample collection.

23 --o0o--

24 DR. WU: So looking at our proposed timeline, we  
25 have submitted our IRB per protocol, and we are scheduled

1 to appear before our IRB on August 5th. We will be  
2 conducting outreach to community partners and refining the  
3 materials and tools that we have, and we hope to be in the  
4 field starting our recruitment in January 2018.

5 This 12 to 15 weeks will go by. We'll collect  
6 our samples. And we want to be returning those results by  
7 January, as in '19. We also want to be in the field for  
8 region 2 starting our recruitment in January 2019, which  
9 means that we pretty soon need to be reaching out to our  
10 community partners and developing those relationships in  
11 region 2. We're very busy. We have a lot to do over the  
12 next six months.

13 --o0o--

14 DR. WU: Finally, I just want to say that the  
15 comments from the Panel have been really helpful. You'll  
16 notice that we've incorporated a number of the comments.  
17 Dr. McKone is not here today, but he had some great  
18 suggestions about external reviewers for our surveys, and  
19 questionnaire materials, which we've done and that was  
20 useful. So we do continue to tweak the protocol, and I'm  
21 happy to hear your suggestions and continue honing our  
22 protocol.

23 CHAIRPERSON SCHWARZMAN: Thank you.

24 Okay. Is that better?

25 Thank you so much for that presentation. We have

1 a few minutes for just clarifying questions from the Panel  
2 before we go on to public comment, and then a full Panel  
3 discussion period about this update.

4 So any clarifying questions for Dr. Wu?

5 PANEL MEMBER QUINTANA: Is this on?

6 Okay. Thank you for that presentation. I just  
7 had a quick question. I'm sure this is being thought of,  
8 but if you could just comment on the extent to which the  
9 website for the participants is mobile friendly, since  
10 that will be a major mechanism by which many of the  
11 participants will access the materials.

12 DR. WU: That's a good question. I don't know  
13 the answer to that. We have just started our -- so there  
14 are two parts to that. One is our website, where you go  
15 for the general information. And our website at CDPH is  
16 kind of new, so I'm not sure how it translates to a mobile  
17 device or, you know, whether it has to be -- whether that  
18 has to be developed separately.

19 For Salesforce, that is also a new relationship.  
20 We're just working with them on our tools. So for  
21 participants who sign up, we would like those materials to  
22 be mobile accessible. It would obviously be very small.  
23 The questionnaires will be difficult to go through. You'd  
24 have to do a lot of scrolling, but we are exploring that  
25 as an option.

1 CHAIRPERSON SCHWARZMAN: Did you have a question?

2 PANEL MEMBER LUDERER: Yeah. Is this on?

3 It sounds like it is.

4 Okay. Thanks for that presentation. It's so  
5 exciting that this is starting. I'm really looking  
6 forward to seeing the results soon.

7 I just had a question. I know that you have a  
8 limited number of analytes that are going to be analyzed  
9 initially, and whether there are plans to, you know,  
10 collect additional specimens that will be archived and,  
11 you know, just tell us a little bit more about that?

12 DR. WU: Sure. We will have plenty of sample.  
13 We are taking blood, serum, and urine. And I can't  
14 remember the volumes off the top of my head of what we're  
15 collecting. But there will be plenty left over to archive  
16 for subsequent analysis, if that -- you know, partly if we  
17 want to add on analytical panels, but also, in the future,  
18 if we wanted to go back and do some longitudinal moni --  
19 longitudinal monitoring over a region, we would have the  
20 availability of samples to do that.

21 CHAIRPERSON SCHWARZMAN: Other questions from the  
22 Panel?

23 Yes, Dr. Cranor?

24 PANEL MEMBER CRANOR: This is an easy question.  
25 Are you pretty happy with the postcards that you send out

1 and the response people will give to them? I see you've  
2 consulted with postcard specialists, as well as  
3 communication specialists. I ask that, because I'm aware  
4 of other organizations that design terrible cards for  
5 trying to urge people to do things. And I just wondered  
6 what you feel about these?

7 DR. WU: Well, this is not a final design. We  
8 are still working with the graphic artist. So I think  
9 they will improve. This was sort of our mock-up of let's  
10 get the information we want onto the postcards.

11 Postcards I don't expect to be hugely successful.  
12 And we want to try it as a randomized sampling technique.  
13 But postcards, in general, I think don't have a huge  
14 response. I think they will have a much better response  
15 when we're giving them out, when there's sort of a  
16 community organization or a face associated with them. So  
17 the postcards have dual use.

18 We also have a flier with a little more  
19 information on it, which I think will be effective. We've  
20 seen bad postcards and good postcards. We've spent a fair  
21 amount of time trying to figure out what do we like or not  
22 like about postcards, what would make me pick this up and  
23 toss it in the garbage or pick it up and take a second  
24 look. And that's why we did the focus group.

25 So having -- having pictures that are appealing

1 to people, having a language that somebody recognizes and  
2 says, oh, I'm going to take another look at this. We've  
3 done a lot of work to figure out what those things are  
4 that will be pique somebody's interest, and we're  
5 continuing to tweak that design.

6 PANEL MEMBER CRANOR: Thank you.

7 CHAIRPERSON SCHWARZMAN: Dr. Quintana, did you  
8 have a question?

9 PANEL MEMBER QUINTANA: I might discuss this  
10 further during the other session. But in terms of the  
11 postcards recruiting technique, I think that if you, of  
12 course, only rely on that, you will miss a lot of the  
13 population, especially lower income or not as competent  
14 English speaking.

15 So I'm wondering how -- if you've -- how much  
16 emphasis you plan to having community groups for everyone,  
17 kind of signs up right there, you know, kind of groups --  
18 group information sessions through environmental justice  
19 groups, for example, that could really augment that  
20 sampling.

21 DR. WU: I would anticipate that the majority of  
22 our pre-screening people will come from community groups.  
23 As I said, I don't think the postcards will be -- I think  
24 we're hoping for a one percent response rate maybe. We're  
25 aiming pretty low.

1           Our community partnerships, which we are in  
2 development -- which are in development right now are  
3 mostly with environmental justice groups. And we're  
4 hoping to really draw from the population through our  
5 community meetings, being present at meetings, having our  
6 partners do a lot of recruiting for us, but also being at  
7 these events like you mentioned, and having somebody  
8 helping somebody with a laptop right there to fill out the  
9 pre-screening. So I think we are counting on the  
10 community outreach that we're doing to do most -- to  
11 contribute most to our recruitment.

12           PANEL MEMBER QUINTANA: Just a really quick  
13 follow up question.

14           CHAIRPERSON SCHWARZMAN: Yeah.

15           PANEL MEMBER QUINTANA: Some of the issues I've  
16 come across in community groups are this system that you  
17 have requires them to kind of sign up to be part of a pool  
18 and then maybe or maybe not get picked. And so I think  
19 that messaging has to be very careful, so that people  
20 aren't upset when they sign up and they aren't picked, you  
21 know, because they get all excited about this, what a  
22 great idea, and then my friend got picked and I didn't.  
23 We have that even for our small-scale studies. How come  
24 they get to participate and we don't, even if we -- they  
25 just hear about their friend participating and they want

1 to sign up, but we don't have funding for additional  
2 participants, for example. And so just some thought maybe  
3 to that messaging to make sure it's not upsetting people,  
4 I guess.

5 DR. WU: We have talked about this quite a bit.  
6 There are communities who have been very enthusiastic  
7 about partnering with us on the recruitment phase. And  
8 their constituents have been like, great, we get to be  
9 biomonitored. And I think we do run the risk of  
10 over-promising.

11 So we are trying to craft language, which is  
12 clear, that this is a regional study. That we're really  
13 trying to represent across a county. It's very difficult.  
14 We want people to be excited enough to sign up, but not  
15 count on this as a community. This is not a community  
16 study. So your point is well taken, and it's something  
17 we're thinking a lot about.

18 CHAIRPERSON SCHWARZMAN: If other Panelists don't  
19 have questions, I have one question.

20 Anyone else?

21 Okay. I think -- you hinted at this a little  
22 bit, but I'm wondering if you could say a little more  
23 about your goals for recruitment in terms of  
24 representation of the community. I mean you hinted that  
25 you wanted to be representational. And it's obviously

1 part of -- a big part of how you're targeting your  
2 outreach and recruitment, and the multiple languages that  
3 you're presenting the materials in.

4 But can you say a little bit more about the goal  
5 for representation? And are you over-sampling or are you  
6 just trying to get a representative population for the  
7 county or the region?

8 DR. WU: We looked at -- when we consider the  
9 parameters that we would like to cover in our  
10 representation, of course, race, socioeconomic status,  
11 geography across the region, age, sex, community, we  
12 realize with 500 samples we are not going to be able to do  
13 that. And we are also very much subject to who signs up  
14 through the pre-screening.

15 So we are -- we're putting a lot of emphasis on  
16 racial representation, just trying to match L.A. County  
17 demographics. And I think to whatever extent we can,  
18 our -- for example, our postcard distribution will be  
19 based on some socioeconomic status indicators, so that  
20 we're trying to get the materials out to a wide range of  
21 platform, a wide range of demographics.

22 But I think the reality is that we won't be able  
23 to really stratify across many of those parameters. We  
24 just don't have enough samples in the county.

25 CHAIRPERSON SCHWARZMAN: And then within the --

1 you said prioritizing racial representation, and then  
2 within that, you'll do a bit about age and gender or --

3 DR. WU: I think --

4 CHAIRPERSON SCHWARZMAN: -- how do this -- how do  
5 the categories follow kind of?

6 DR. WU: You know, we're still working that out  
7 how we set our goals, and then how we turn that into an  
8 algorithm for selection of the participants. It would be  
9 great if we could get some distribution across age and  
10 sex. But again, we -- we're sort of balancing the need to  
11 be representative with the need to just get 500  
12 participants. And we're still working that out.

13 CHAIRPERSON SCHWARZMAN: Okay. Great.

14 One other question on that is are you thinking  
15 about using occupation and geographic location also as  
16 ways of sort of broadening the diversity of the people  
17 that you're sampling or not?

18 DR. WU: Geographic location certainly. By  
19 recruiting in each of the service planning areas, we're  
20 hoping to get interested participants from across the  
21 county. And we would want to -- we will want to select to  
22 make sure that we're getting people from different parts  
23 of the county. I think occupation is harder, because we  
24 are collecting occupational information, but it would be  
25 harder to stratify based on that.

1           CHAIRPERSON SCHWARZMAN: Right. So, but there's  
2 a difference -- I mean, I guess there is a question of  
3 stratifying -- your ability to stratify is related to your  
4 desire to recruit maybe. So I was going to say that, you  
5 know, the ability to stratify isn't quite as -- it's a --  
6 it's a -- it's a longer-term or more ambitious goal than  
7 recruitment. But maybe I am thinking now that it is right  
8 to think about that first, because if you have  
9 representation, but it's not sufficient to stratify on,  
10 it's not that meaningful to have that representation.

11           Yeah. Okay. That's interesting. Thank you.

12           Any other Panel questions before we move on to  
13 public comment?

14           Okay. Duyen, do we have public comment from  
15 within the room?

16           MS. KAUFFMAN: No.

17           CHAIRPERSON SCHWARZMAN: Is there any public  
18 comment from online or from the webinar?

19           MS. DUNN: No.

20           CHAIRPERSON SCHWARZMAN: Okay. Then in that  
21 case, we can move on to the Panel discussion. And we're a  
22 little bit ahead of time, so we have 15 minutes.

23           Is this on?

24           So anyway, we have plenty of time for Panel  
25 discussion.

1 Dr. Quintana.

2 PANEL MEMBER QUINTANA: Hi. I have another  
3 question, I guess, about the study design. You said that  
4 you've been going around the state collecting information  
5 for community groups and environmental justice groups  
6 about their priorities. And I'm wondering to the extent  
7 to which those priorities have fed into what's being  
8 measured in this study.

9 And I'm thinking specifically, you know, I think  
10 the metals are a great thing to measure, because partly  
11 there's known health effects and you can interpret results  
12 to the community quite well. And you've seen, you know,  
13 elevated arsenic in preliminary data from different  
14 population groups, mercury. But for example if the  
15 community groups have been saying diesel exposure is a  
16 huge concern, have you thought about, you know, why PFASs  
17 versus 1-nitropyrene, for example.

18 I know the resources are -- and this question is  
19 being asked in the context of very limited resources, and  
20 I understand that.

21 DR. WU: We actually are considering adding 1-NP  
22 to the L.A. County. It's obviously a huge issue in L.A.  
23 County, or at least to a subset for our participants.  
24 Some of it is a logistics issue collecting enough urine  
25 samples, and then trying to figure out how to select

1 participants for that particular analysis. But it is  
2 something -- diesel is actually the analyte we're  
3 considering adding at this point.

4 PFAS, I mean, the recent work has shown that  
5 there is a geographic contribution, because of your  
6 drinking water source. And we're very interested to see  
7 how PFASs vary across the state, and whether we can  
8 continue the work that DTSC has done to look at the  
9 connection between drinking water source and PFASs. So  
10 you're right, it has not risen to the top of our community  
11 discussions, but I think it's an important one for us to  
12 look at for public health issues.

13 PANEL MEMBER QUINTANA: So, yeah, I was not  
14 questioning the interest of that. I think it is extremely  
15 interesting. I guess my question was more your vision of  
16 how the community concerns kind of fed into the study.

17 Thank you.

18 DR. WU: Well these two things were developed  
19 somewhat concurrently, so we had selected metals, as you  
20 said, for public health reasons, and also because there is  
21 so much interest. We were also looking for analytes that  
22 would be sort of unifying to the State, things that are  
23 interesting across the different regions. But I think as  
24 we have heard more from the EJ communities, it will --  
25 it's more of what we can add on to those statewide

1 analytes. And there will be regions where -- who are very  
2 interested in pesticides. And that's something we want to  
3 drop into those regions, as opposed to diesel, which are  
4 L.A. County and other parts of Southern California.

5 Go ahead.

6 PANEL MEMBER QUINTANA: Just one last follow-up  
7 comment. I'm sorry, Panel. I guess the point of the  
8 study is to be statewide, it seems if you're going to  
9 measure 1-nitropyrene in L.A., you'd also want to measure  
10 it in Kern, if nothing else to show the huge difference  
11 that might exist or not, you know, tractors kick out  
12 diesel too, if you're a farmer, but -- so I think I'm not  
13 sure you should be -- have regional specific additions,  
14 because you do need a comparison. NHANES, one of the  
15 major wonderful uses of that is to be a comparison.

16 You know, so I just want to throw that out there  
17 that I would encourage if you do it one place, to do it  
18 all places.

19 DR. WU: That's a really good point about having  
20 a comparison group, but I wouldn't want to limit us to  
21 only doing statewide just because that will probably keep  
22 us from adding any additional panels just because of the  
23 resources issue. I mean, I wouldn't want -- I mean,  
24 there's so much uncertainty about our budget, so if we had  
25 money this year and the ability to add something on in

1 this region, I feel like we should grab that opportunity.

2 CHAIRPERSON SCHWARZMAN: Are you done with that  
3 or should --

4 PANEL MEMBER QUINTANA: I'll stop now.

5 (Laughter.)

6 CHAIRPERSON SCHWARZMAN: Okay.

7 Dr. Cranor.

8 PANEL MEMBER CRANOR: This is a difficult  
9 question I realize, but hang on. California is not a poor  
10 state. The budget for this program has been going down.  
11 There are obviously other needs to protect the public  
12 health, so I suppose I have two questions.

13 To what extent -- I mean, let me -- one more  
14 background piece of information. I did have some  
15 conversations with people at the last meeting where there  
16 were lots of public folks. And I thought maybe they  
17 should speak to the Legislature. And I don't know,  
18 probably not much came of that.

19 But what are the prospects for better funding for  
20 the program, and what would you do with it? I guess those  
21 are the two -- the two questions.

22 In Calif -- I've talked to school teachers and  
23 they say, well, there's no money. Well, that's -- that's  
24 false. There may be no tax money, but California is not a  
25 poor state. So there could be funding there if it were

1 needed, and what would you do with it?

2 I said it was hard.

3 (Laughter.)

4 DR. WU: I think this is a much longer discussion  
5 than we have time for. Everyone here works in public  
6 health and environmental health. And you all know the  
7 context within which we're all working. It's a difficult  
8 time. There's a lot of uncertainty about what future  
9 budgets may hold.

10 I think part of our challenge is to help define  
11 biomonitoring and environmental health as a wider  
12 discipline as important parts of public health, because  
13 the things that I think the public recognizes -- public  
14 and legislators recognize are more traditional public  
15 health and more traditional medical models of health.

16 And so my hope is that by starting our statewide  
17 program, we'll develop data and people will be like, well,  
18 when is our region going to get done? Why don't we have  
19 the statewide sampling plan that can be done in a one- to  
20 two-year cycle. And it's creating the awareness that this  
21 really is a critical part of public health. This is a  
22 critical part of why people get sick that will create that  
23 demand. You know, who knows if that's going to happen.

24 But in terms of what we would do, you've seen the  
25 study here. There are a number of things we have had to

1 reduce in the study, in terms of numbers of participants,  
2 number of analytical panels, our ability to sample the  
3 state in a one- to two-year cycle. These are all -- these  
4 are all compromises. And I think the study is still  
5 valuable and will produce very interesting and valuable  
6 data, but it is not what we originally envisioned as a  
7 statewide sampling plan.

8           If we had the resources, we would cover the  
9 regions more quickly. We would have the ability to do  
10 1-NP, and the phenols, and pesticides, and all of the  
11 analytical panels that -- which we have determined here  
12 have been -- are so important to public health.

13           What we're doing, it's interesting and valuable,  
14 but it -- it is not really statewide sampling as initially  
15 envisioned.

16           PANEL MEMBER CRANOR: Thank you.

17           CHAIRPERSON SCHWARZMAN: Panel questions?

18           Yeah.

19           PANEL MEMBER LUDERER: I have kind of a more  
20 specific question regarding kind of going back to your ACE  
21 Study, and -- you know, I under -- you found the -- quite  
22 a large number of high inorganic arsenic levels. And I'm  
23 wondering if you could talk a little bit about how kind of  
24 those findings have informed, you know, the kinds of  
25 questions you're going to be asking people about exposure

1 in the CARE Study, you know, given that you've, you know,  
2 found these levels, and also if you have any information  
3 about what you found the source might be. You might not  
4 have that yet.

5 DR. WU: I think from our conversations with the  
6 participants, it's mostly a rice diet that's contributing  
7 to the arsenic levels. It's an Asian population. We eat  
8 lots of rice and fish.

9 So that has informed how we ask the questions in  
10 the CARE Study, but it's a very different population. So  
11 whereas our ACE questionnaire was many questions about  
12 species of fish, and how you eat your rice, and really on  
13 a micro-level focus on your rice diet and rice noodles and  
14 all this other rice product.

15 The CARE Study, partly because it's a different  
16 population, but also because we're trying to do these  
17 surveys online instead of with an interviewer, we've had  
18 to shorten that. And so there is -- there are fewer  
19 questions focused on specific analytes.

20 We do have the ability to do follow up though.  
21 If somebody is in the high-exposure group, we will do  
22 our -- have our usual follow-up protocol and then we'll be  
23 able to get into some more detail about what your  
24 potential exposures might be.

25 CHAIRPERSON SCHWARZMAN: Other Panel questions?

1           If not, I have one about the FREES Study. And I  
2 know you've presented this information to us before, but I  
3 was hoping you could just say again something about which  
4 flame retardants you're looking at. And obviously, in the  
5 older samples, the pre-samples, we would be looking at  
6 some of the chlorinated and brominated flame retardants  
7 primarily. But I'm wondering if you're also looking at  
8 some of the replacement flame retardants, so that we can  
9 see if those were lower before and higher after, even if  
10 they're not related to the foam replacement, but if like  
11 they were lower before, and then they predominate later  
12 unrelated to the furniture replacement.

13           DR. WU: So we are looking at PBDEs in blood and  
14 serum. And then we'll be looking -- there are three  
15 metabolites of organophosphate flame retardants that we're  
16 looking at in the urine samples. When people replace  
17 their foam or furniture, they were purchasing things that  
18 were labeled. Because of the new labeling law, they were  
19 able to purchase furniture, which should not have had any  
20 flame retardants in it. So our hope is that we would see  
21 a decrease in both.

22           CHAIRPERSON SCHWARZMAN: So -- and there's not  
23 actual testing of the foam, right, because the TB117-2013  
24 label means it doesn't -- like it means that different  
25 flammability standard, so it presumably doesn't have to

1 have the added chemicals, but it doesn't like exactly  
2 guarantee that there aren't compounds in it.

3 DR. WU: We were able to get foam samples -- the  
4 old foam samples from all of our participants. And  
5 actually, DTSC -- Myrto's lab is analyzing those foams.  
6 We don't have all foam from all of the new foam samples.  
7 Not everyone wanted us to cut into their new sofas.

8 (Laughter.)

9 DR. WU: So -- but we do have a number of the  
10 samples, and we'll be able to report out on that.

11 CHAIRPERSON SCHWARZMAN: That's great. Thank  
12 you.

13 Other questions?

14 Yeah, Dr. Quintana.

15 PANEL MEMBER QUINTANA: Do we have time?

16 CHAIRPERSON SCHWARZMAN: Yes, we do. We have  
17 plenty of time.

18 PANEL MEMBER QUINTANA: Hi. This is going back  
19 to the CARE Study. I'm curious about your -- I know  
20 you've submitted the materials to the IRB already, so I'm  
21 not saying you should go back and change things. But I'm  
22 just curious the language that you put in terms of  
23 permitting further analyses on these samples, and  
24 especially non-targeted analysis on the samples.

25 And as we discussed in previous meetings, there

1 is an ethical issue, because you can pick up drugs of  
2 abuse, for example, when you run a non-targeted screen.  
3 Even if you don't want to see them, they're in the  
4 database for that sample, and they could be picked out by  
5 somebody. So I'm just curious of your comments about  
6 that.

7 DR. WU: Well, it's never too late to edit  
8 materials, even if an IRB reviews them. And they will  
9 probably have comments for us as well, but we do -- as we  
10 do in all of our informed consents, we have an option for  
11 people to donate their samples for storage after our  
12 initial samples are completed.

13 So they can archive. They can volunteer to  
14 archive them for additional sampling or analyses, or they  
15 can opt out of that, and then we destroy their samples  
16 right after we're completed with our initial analyses.

17 We do -- we describe them as additional  
18 environmental chemicals. And we state that we will not  
19 measure for any pharmaceutical, or recreational, or  
20 illegal drug. I can't remember exactly what the wording  
21 is, but we're clear to specify that we are not going to be  
22 looking for particular classes of drugs. We're only  
23 looking for environmental contaminants.

24 CHAIRPERSON SCHWARZMAN: Did that answer --

25 DR. WU: I realize that doesn't preclude that,

1 like if it's in the data that comes out with non-targeted  
2 screening. I realize that is still an issue. We would  
3 not be finalizing that data. So even if the lab has it,  
4 it would not be reported to us as a -- to EHIP as a  
5 program. And so that information would not exist, for  
6 example, in the results return packets.

7 PANEL MEMBER QUINTANA: No, I agree. It wouldn't  
8 exist -- be reported. But it's just something to think  
9 about in terms of keeping data secure -- and, you know, it  
10 is -- it is an ethical issue. We came across it with  
11 house dust. We're finding all kinds of interesting stuff  
12 in house dust that we didn't expect. That's why it really  
13 came up to me, you know, so...

14 CHAIRPERSON SCHWARZMAN: Maybe, if I can add on  
15 to that question. Maybe the sort of bottom line is do you  
16 still have the opportunity to do non-targeted screening,  
17 if budget and interest kind of permits and suggests that,  
18 based on your IRB approval? Do you feel like that's  
19 consistent?

20 DR. WU: Yeah, I think we could do that. And I  
21 think we'll hear more about the non-targeted screening in  
22 our labs. Somebody may ask for that in the afternoon  
23 session.

24 But, yeah, I think our IRB does allow for these  
25 additional analyses, and we're not very specific in that,

1 except to say that it is these environmental contaminants.  
2 We haven't gotten into the details of -- and I'm -- I'm  
3 not the right person to answer the question about at what  
4 point the lab could say, okay, this data we're not going  
5 to continue to identify or we're not going to report it  
6 out. Maybe actually Jianwen or June-Soo could report out  
7 and give you a little more information on that.

8 CHAIRPERSON SCHWARZMAN: Other Panel questions?  
9 Oh, yes. Please.

10 PANEL MEMBER KAVANAUGH-LYNCH: Hi. Mel  
11 Kavanaugh-Lynch. I'm chewing on this idea that we'd like  
12 to analyze more things and sample more people, and not  
13 turn potential participants away. And I'm thinking about  
14 ways to leverage the public's interest in both other  
15 analytes and in bigger samples and particular populations.

16 Some of the things that are coming to my mind is  
17 offering the opportunity for crowdsourcing for saying to  
18 groups that say we want -- why don't you -- why aren't you  
19 sampling more of our people, or why aren't you sampling  
20 more of the things we're interested in? Saying we would  
21 be happy to do that, if you can help us find the funding  
22 for that, and even have participants pay for adding to  
23 their samples. On a -- and moving that into advocacy.

24 So in community meetings, where the community is  
25 bringing up wanting more people or wanting more analytes,

1 encouraging them and even giving them sample letters to  
2 send to local, regional, and State bodies that have the  
3 opportunity to add funding, to advocate for more funding  
4 to the Biomonitoring Program. So kind of creating --  
5 leveraging that -- those requests into actual advocacy.

6 DR. WU: I think it is important that we get  
7 creative about how we fund the Program. There is not  
8 really a State mechanism for us to function in that way,  
9 to bring in -- to like charge money for samples basically.  
10 And we're certainly not able to lobby or be part of a  
11 lobbying effort to encourage people to send letters to  
12 their legislators. That's something a community partner  
13 or an advocate would be able to do. And for those of you  
14 listening, that would be awesome.

15 (Laughter.)

16 DR. WU: But as the State, there are restrictions  
17 on what we can do.

18 CHAIRPERSON SCHWARZMAN: Other Panel questions or  
19 comments?

20 Anything else?

21 So you looked like you were thinking about  
22 something.

23 PANEL MEMBER KAVANAUGH-LYNCH: No.

24 CHAIRPERSON SCHWARZMAN: Okay. In that case, we  
25 are going to go on to our next presentation, which is by

1 Dr. Jon Sobus.

2 I'm pleased to welcome him here from US EPA.  
3 He'll talk about advancing and integrating non-targeted  
4 analysis research at US EPA. Dr. Sobus is a physical  
5 scientist at US EPA's Office of Research and Development,  
6 which is in the National Exposure Research Library --  
7 Laboratory located in Research Triangle Park, North  
8 Carolina. He currently serves as task lead for the Rapid  
9 Exposure and Dosimetry Project within the Chemical Safety  
10 for Sustainability National Research Program. And we  
11 further serves as principal investigator and team lead for  
12 the Non-Targeted Analysis Research Project.

13 (Thereupon an overhead presentation was  
14 Presented as follows.)

15 CHAIRPERSON SCHWARZMAN: Thank you for being  
16 here.

17 DR. SOBUS: Thank you so much.

18 Does that work?

19 Okay. Is that better?

20 Thank you so much for the invitation to be here,  
21 for that nice introduction. I was actually here three  
22 years ago talking about a completely unrelated topic, but  
23 it's great to be back. I'm very excited to hear some  
24 really encouraging discussions from this first session. I  
25 think it's a really excellent transition into what I'll be

1 talking about today.

2 I had the opportunity yesterday to actually speak  
3 with some of the staff scientists about common interests,  
4 common work. So that will be echoed in the presentation  
5 today. This is a fairly technical talk. I hope that's  
6 okay, but I really wanted to get into the specifics of  
7 what we're doing in ORD, and to clearly communicate that  
8 the purpose of building some of these tools, and  
9 databases, and dashboards is to make those resources  
10 available to the public, including State programs.

11 So keep in mind, everything I present today are  
12 tools that we're building to make available to you all.  
13 So again, I'm from US EPA Office of Research and  
14 Development.

15 Today, I'll be talking about tools that we're  
16 building to advance and integrate non-targeted analysis  
17 research.

18 --o0o--

19 DR. SOBUS: So we are building our portfolio in  
20 non-targeted analysis for two key reasons. One, we need  
21 to have a better understanding of the drivers of disease.  
22 We now believe that the majority of chronic disease risk  
23 is caused by differences in environments, rather than  
24 differences in genetics. But we don't have a great handle  
25 on what are the environmental components that drive



1                   --o0o--

2           DR. SOBUS: My understanding of much of what is  
3 discussed amongst the Scientific Guidance Panel and much  
4 of Biomonitoring California is reflecting targeted  
5 analysis. So this is looking at the known knowns. These  
6 are the compounds that everyone is very familiar with, and  
7 compounds for which targeted analytical chemistry methods  
8 have been developed.

9           In my experience, these compounds, the known  
10 knowns, represent less than one percent of what's  
11 typically in any given sample, environmental or  
12 biological. So by extension, we're looking at far less  
13 than one percent of the exposome.

14           So based on that, I would make the argument that  
15 we can't solve 21st century public health problems if  
16 we're kind of blinded to 99 plus percent of exposure. So  
17 we are very much looking at the tip of iceberg, and I  
18 think we need to dig a little bit deeper.

19                   --o0o--

20           DR. SOBUS: So moving on to the known unknowns.  
21 These are things that we know exist, but for which there  
22 aren't currently target methods, and there isn't currently  
23 exposure data. To give you some example, there are large  
24 public inventories of chemicals, millions of chemicals,  
25 that are known to exist that are manufactured.

1           We can develop what are called suspect screening  
2 analysis methods. So this is kind of a subsection of  
3 non-targeted analysis research. So a lot of what I'll  
4 talk about today focuses on these high throughput suspect  
5 screening analysis methods that try and take data that we  
6 generate on high-resolution mass spectrometry platforms  
7 and kind of cross-reference that to these large databases  
8 of known chemicals to try and find hits. So are any of  
9 these chemicals on this list present in any of these  
10 samples that we're examining?

11           Base on my experience, that gets us about five to  
12 ten percent of what's in a given sample, or five to ten  
13 percent of the exposome. So we made a lot of headway in  
14 developing efficient methods to do suspect screening  
15 analysis.

16                           --o0o--

17           DR. SOBUS: The final chunk is the unknown  
18 unknowns. So this is about 90 to 95 percent of what's in  
19 a given sample. This is often your biological  
20 metabolites, your environmental transformation products,  
21 things that we don't yet know exist.

22           To get at these compounds, since they're not in a  
23 database, you have to use true non-targeted analysis  
24 methods. These methods are very time and resource  
25 intensive. We definitely need ways to speed up that

1 process. But certainly, we need to go down this path to  
2 really get at the bulk of the exposome. So I just want to  
3 make a final point that at a high level we use the term  
4 non-targeted analysis to really encompass both the suspect  
5 screening and the true non-targeted.

6 Most of what I'll talk about today is technically  
7 suspect screening analysis, but I'm just going to use the  
8 term generally "NTA".

9 --o0o--

10 DR. SOBUS: So there are a variety of tools that  
11 are required when doing non-targeted analysis research.  
12 We can start on the left-hand side with analytical  
13 instrumentation. By and large, most groups that are doing  
14 NTA use mass spectrometry, and most use high-resolution  
15 Mass spectrometry. So this is what's used to generate  
16 information about unknown chemicals in any given sample.

17 Again, when we're doing suspect screening, we're  
18 going to use large chemical databases like PubChem or  
19 ChemSpider. Some of these databases have millions of  
20 compounds listed in them. But to kind of have that  
21 linkage between what's generated on a mass spectrometer  
22 and what's in these chemical databases, we need workflows  
23 to process that data to make some sense of that data.  
24 There are actually quite a few computational workflows  
25 that exist. Some are proprietary, some are open.



1 So for those in the room, or on the line, that are looking  
2 to do non-targeted analysis, we can do searches based on  
3 accurate mass, plus or minus some amount of error. You  
4 can do searches based on formula. And as you'll see in an  
5 upcoming slide, we can actually do batch searches on  
6 thousands of formula at once. And it will return all the  
7 structures that are consistent with those formulas. So  
8 it's a very powerful tool.

9           On the bottom we show a landing page for a given  
10 chemical. You can see it draws the structure. We have  
11 information on intrinsic properties, structural  
12 identifiers. And very importantly -- I'm not sure how  
13 many people are knowledgeable about our ExpoCast and  
14 ToxCast programs, but the dashboard also hosts exposure  
15 predictions for thousands of chemicals, as well as  
16 bioactivity data for thousands of chemicals from the  
17 ToxCast program and from Tox21. So all of these resources  
18 are available in the dashboard, which is why we're using  
19 this as a primary tool in our workflow.

20           For anyone that has interest in learning more  
21 about or using the dashboard, please feel free to contact  
22 Tony Williams, who's the project lead. He's a phenomenal  
23 guy and has built a really phenomenal product.

24   --o0o--

25           DR. SOBUS: So I want to step back and talk about

1 what we actually do when we do non-targeted analysis.

2           This field is borne out of metabolomics. And a  
3 lot of the workflow in metabolomics is consistent with  
4 NTA, but there are some inconsistencies, so I want to kind  
5 of go through just a very simple kind of five-step  
6 procedure that outlines how we go about doing NTA.

7           We can start with the image here, which is a  
8 total ion chromatogram that was analyzed on a mass  
9 spectrometer. This was an extract of one house dust  
10 sample. It was run in negative ionization mode. And  
11 there's about 300, what we call, molecular features shown  
12 here.

13           The molecular features are these peaks, which  
14 basically represent some unidentified chemical. They are  
15 defined at this time by an accurate mass, retention time  
16 and a mass spectrum. The goal is to look at these 300,  
17 and if we could, to try and figure out what they all are.  
18 That's very difficult to do, so we try and first  
19 prioritize them. We say which are of greatest interest to  
20 us?

21           By default, a lot of people look at the biggest  
22 ones first, which is a pretty good idea, but it's not  
23 always the best idea. So we've come up with a number of  
24 different procedures and algorithms for actually  
25 prioritizing these molecular features to figure out which



1 DR. SOBUS: So I showed just one example of one  
2 sample that was analyzed in one mode, and it had 300  
3 features. Many of the samples that we look at have  
4 hundred, if not thousands, of features. And we look at  
5 tens, or hundreds, or thousands of samples across multiple  
6 modes. So there are thousands and thousands and thousands  
7 of data points to deal with. And this becomes very  
8 cumbersome in terms of data processing and data storage.  
9 This is one big challenge of NTA.

10 Another big challenge is that, again as I  
11 mentioned in the previous slide, there needs to be some  
12 prioritization. And historically, out of the metabolomics  
13 field, biases have been introduced to look at only high  
14 intensity compounds, and lots of experiments are done to  
15 compare groups.

16 So biological samples collected from a diseased  
17 population versus a control population, or upstream versus  
18 downstream of a water treatment plant. So these types of  
19 comparisons -- statistical comparisons are done to kind of  
20 identify features that are enriched. So this is very  
21 valuable to do. But again, you're biasing only to a very  
22 small group of compounds, and we need to step back and  
23 look at bit more holistically.

24 One of the biggest problems that the field faces  
25 at their -- there is no standardized methods. Every lab

1 has their own way of doing this, which makes it extremely  
2 difficult to compare results from one lab to the next.  
3 That is potentially the biggest challenge that we face  
4 right now.

5           We need to come up with ways to overcome the lack  
6 of standardization. And then finally, most of the  
7 chemists in the room know very well from the targeted  
8 experiments that there needs to be some confirmation at  
9 the end of the day before some action can be taken. This  
10 is time-consuming. We're talking about thousands and  
11 thousands of chemicals, and we don't have standards for  
12 all these chemicals. How do we overcome that challenge?

13           --o0o--

14           DR. SOBUS: So I'm going to talk about two  
15 different things we're doing to address these four primary  
16 challenges.

17           One, I'm going to go through as quickly as I can  
18 some of the workflows and tools that we're building at the  
19 agency, again with the intention of making available to  
20 the public. And then I want to finish by covering a  
21 collaborative trial that we've been leading now for about  
22 a year and a half that involves about 30 different  
23 institutions, including international partners.

24           --o0o--

25           DR. SOBUS: So this is a broad workflow that is a

1 little bit complicated so I'm going to step through it bit  
2 by bit hopefully without going into too much detail.

3 --o0o--

4 DR. SOBUS: But as we begin this process for  
5 doing non-targeted analysis we always start with the raw  
6 samples. And I just want to make a point that these  
7 methods can be used basically for anything. All the  
8 images we show here are things that we are currently  
9 evaluating with our NTA methods. So a very, very  
10 versatile approach.

11 --o0o--

12 DR. SOBUS: We take the raw samples, as we always  
13 do in analytical experiments, we do extractions, different  
14 clean-up steps, and we generate these total ion  
15 chromatograms, which represent, what we call, the raw  
16 features. These are just kind of unfiltered chemicals  
17 that are in these samples.

18 We need to do a bit of processing to kind of get  
19 down to again the things that are of interest, the things  
20 that are real, and the things that we need to take a  
21 closer look at.

22 --o0o--

23 DR. SOBUS: So we have spent a good bit of time  
24 combining some vendor software applications, as well as  
25 writing our own programs that we intend to make open to

1 the public to do processing of these raw features. So  
2 we're kind of getting rid of the stuff that isn't real,  
3 get rid of the stuff that isn't reproducible, and kind of  
4 hone in on the things that we really need to take a closer  
5 look at. So the output of these programs that we've  
6 written are basically data matrices that say here's a  
7 feature with an accurate mass, and a retention time.  
8 Here's where it shows up in all the different samples at  
9 all the different intensities.

10 --o0o--

11 DR. SOBUS: So we can take that processed feature  
12 file, and then restrict it to things where we've been able  
13 to assign a formula to those features with some level of  
14 confidence. So now we've gone from potentially 50,000  
15 things down to 20,000 things, maybe down to 5,000 things.  
16 So we are losing a little bit of the information. But  
17 again, we're focusing on the things that we have the most  
18 confidence in at this point.

19 So here we have a list of formulas This is just  
20 a snapshot. Typically, these lists of formulas are  
21 hundreds to thousands of formulas long. The next step is  
22 we use our Chemistry Dashboard. We have the batch search  
23 function that I mentioned before, and we will actually  
24 search these thousands of formulas and try and pull back  
25 all the candidate structures that are consistent with each

1 formula.

2 --o0o--

3 DR. SOBUS: Some of these formulas can have 10,  
4 20, 30, 100 different structures. So this becomes the  
5 critical challenge of NTA, which chemical is right?

6 Assuming you got the formula right, which  
7 chemical is right? This is extremely difficult to do, and  
8 this is just one example where we pulled back nine  
9 structures for a given formula.

10 --o0o--

11 DR. SOBUS: This gets complicated, so I won't  
12 really get into it, but this is where we spend a lot of  
13 our time. To date, we use something that's called a data  
14 source ranking procedure to nominate the most likely  
15 candidate structure.

16 Data sources is basically a popularity contest.  
17 So when you build a database with chemical structures,  
18 pulling information from different sources, from different  
19 lists, different inventories, the chemical is on every  
20 single inventory that you pull from. Those are all data  
21 sources. It means it's a very popular chemical, many  
22 people know about, probably high-production volume, and  
23 thus it's most likely to be the correct compound.

24 So we initially do a data source ranking, and we  
25 actually spit out on the dashboard the top ranked



1 candidate structure, or perhaps a few candidate structures  
2 for each of the formulas that we've identified in the  
3 samples. Which ones do we care about?

4           We're still talking about thousands and thousands  
5 of features. Which ones do we care about trying to do  
6 additional experiments, confirmatory experiments on. This  
7 goes back a couple years where we had the idea of using  
8 ExpoCast data, and using ToxCast data about these  
9 chemicals that we're tentatively identifying to figure out  
10 which ones we care about.

11           So we took some information from our experiments,  
12 so the detection frequency of these compounds, the  
13 abundance of these compounds, how big is the signal in the  
14 sample, and we also pulled exposure information and  
15 bioactivity information about these compounds from the  
16 dashboard. And we build these nice little ToxPi's, which  
17 is basically a graphical with a numerical representation  
18 of how important the compounds are from a health risk  
19 standpoint.

20           So if they have high bioactivity and high  
21 exposure, and if they're in every sample at a really high  
22 concentration, that's a really important compound. That's  
23 something we really want to take a look at as soon as  
24 possible. If it has no evidence of bioactivity, limited  
25 exposure, if it's only in five percent of the samples at a



1 possible, we want to use, what I would call, high  
2 throughput semi-quantitative methods to try and predict  
3 concentrations for these compounds. Whether we know what  
4 the structure is or not, we can predict the concentration.  
5 So this is a heat map that I made just using what I would  
6 call a global calibration curve.

7           So I took many, many, many chemicals and  
8 basically took the average linear trend across all those  
9 chemicals and then applied it across several thousand  
10 compounds. So this is just a heat map with some  
11 hierarchical clustering of samples -- or of chemicals  
12 measured in dust samples. You can see on this very simple  
13 example, we're still looking at six orders of magnitude  
14 range in estimated concentration.

15           The precision is terrible, but you're at least  
16 getting some number with which you can then do additional  
17 work.

18                           --o0o--

19           DR. SOBUS: So that is the basis for our  
20 workflow. Again, everything that we're building as part  
21 of that workflow, the goal is to make available to the  
22 public as open code, or as some function available in the  
23 dashboard. So hopefully, we can continue to work with the  
24 State scientists, other state scientists, local labs, et  
25 cetera in trying to implement these tools on a wide scale.

1           So I want to shift gears and talk about  
2 applications very quickly. I'll get moving.

3           There are three applications that I will cover  
4 very briefly. Exposure surveillance. So can we use these  
5 tools just to look for what chemicals are in different  
6 foods, products, et cetera. Again, chemical  
7 prioritization. Can we not only look at what's there, but  
8 figure out what's most important. And very importantly,  
9 look at what chemicals co-occur, where we might have these  
10 mixture effects from complicated exposures. And then I'll  
11 touch on exposure forensic. So can we identify chemical  
12 signatures of specific exposure sources?

13                           --o0o--

14           DR. SOBUS: So I'm very briefly going to cover  
15 applications of the workflow in three different instances.  
16 These are the results from a consumer product analysis  
17 pilot study that recently completed. We've submitted the  
18 article for publication. We worked with an analytical  
19 contractor using GC/GC low resolution time of flight mass  
20 spectrometry to evaluate 20 different product categories.

21           We have 12 different formulation categories, such  
22 as lipstick, tooth paste, sunscreen; seven different  
23 article categories, like carpets, cotton clothing, fabric  
24 upholstery; and then one food category.

25           Our contractor went out, they picked five random

1 products from each of these categories. They did Soxhlet  
2 extraction and then again they analyzed these extractions  
3 using NTA.

4 Ten to hundreds of compounds were tentatively  
5 identified or confirmed in all of these products. If you  
6 look on the table on the left, you can see that the number  
7 of tentatively identified compounds was typically about an  
8 order of magnitude higher than those explicitly listed on  
9 ingredient list. So there's a lot more there than you may  
10 have thought.

11 When we took the tentative NTA hits and hit it  
12 against a consumer product database that we built  
13 internally, we found only about 20 percent of those  
14 chemicals show up on that inventory.

15 The take-home point here is there's a lot more  
16 stuff in products than we currently know about. And just  
17 very quickly on the right-hand side again, we can estimate  
18 concentrations for some of these chemicals and products,  
19 but there's a lot of work to be done considering things  
20 like extraction efficiency to improve the accuracy and  
21 precision of those predictions.

22 --o0o--

23 DR. SOBUS: The second pilot study that I'll talk  
24 about was our seminal piece of work. This was published I  
25 think in 2015 or 2016. It was actually featured in a

1 National Academies Report.

2           This was a house dust pilot study where we looked  
3 at samples from 56 homes. We tentatively identified  
4 thousands and thousands of structures, and we used the  
5 ToxPi approach that I mentioned before to prioritize all  
6 of these candidate structures. We took, as a proof of  
7 concept, the top 100 candidates. We acquired standards  
8 for those, and we tried to confirm as many as we could.

9           We wound up confirming about 35 of the top 100  
10 compounds. And through a lit search, we found 45 percent  
11 of those had never been reported. So as a proof of  
12 concept, many thousands of things in dust, many of the  
13 things that we find have never been looked for before.

14           And again, we're in the process now through some  
15 hierarchical clustering and some supervised and  
16 unsupervised techniques of trying to dig down into these  
17 samples using some meta-data about the homes, the age of  
18 the home, heating source, smoking, pesticide use, pets, et  
19 cetera, to try and see if we can have chemical  
20 fingerprints of some of these exposure sources. And if  
21 we're successful in that, that certainly applies to things  
22 like surface water, sediment samples, et cetera.

23                           --o0o--

24           DR. SOBUS: The last example that I'll talk about  
25 is a drinking water pilot study, where we actually

1 implemented Brita filters. So these are filters that  
2 attach to a tap. We collected nine Brita samples over a  
3 period of about a month in the North Carolina triangle  
4 area, representing four municipalities and two private  
5 wells.

6           Again, we found thousands of features. We did  
7 our prioritization using the ToxPi approach here. Rather  
8 than showing the actual pi's themselves, we actually made  
9 a stacked bar plot.

10                   --o0o--

11           DR. SOBUS: But the same principles apply, we  
12 looked at the top 100 priority compounds. We only had 16  
13 standards for the top 100 compounds, 15 of them were  
14 correct. So we can be very successful. But again, you're  
15 limited to the standards that you have when doing  
16 confirmation experiments.

17           Many of the compounds that we were able to  
18 confirm aren't on any current monitoring lists. So as a  
19 proof of concept, again, this is a very good approach for  
20 identifying emerging contaminants, in this case, in  
21 drinking water samples.

22           I'll point out that when we matched these  
23 compounds to some of our databases in-house, we found that  
24 the origin of these chemicals are from consumer product  
25 uses, as well as some industrial processes.

1                   --o0o--

2                   DR. SOBUS: So I want to transition briefly, as  
3 briefly as I can, to kind of these lingering science  
4 questions. So we've come a long way, but there's much,  
5 much more to be done.

6                   We know that methods are extremely variable from  
7 lab to lab. So what are the consequences of that  
8 variability. Are some of these methods better than others  
9 overall? Are some better than others in only specific  
10 instances for specific chemical classes, and how does  
11 sample complexity affect our ability to do NTA? We really  
12 don't have any handle on this right now.

13                   How many methods would we really need to apply to  
14 comprehensively characterize any given sample? We have no  
15 idea. And then I made a point before that we don't have a  
16 lot of good experimental MS/MS spectra to help in  
17 candidate identification. And we certainly don't have  
18 standards for many of these compounds.

19                   So how do -- how do we overcome that hurdle? And  
20 to go back to a point that I think was made earlier. Is  
21 there a way we can actually crowdsource exposure data, if  
22 we can make resources available to public laboratories.

23                   --o0o--

24                   DR. SOBUS: So to address these lingering science  
25 questions, about a year and a half, two years ago, we

1 established, what we call, ENTACT. So this EPA's  
2 Non-Targeted Analysis Collaborative Trial.

3 --o0o--

4 DR. SOBUS: This is a visualization of kind of  
5 what it's about, who's involved, and what we expect it to  
6 produce. For those of you that are familiar with the  
7 ToxCast program, at EPA, this includes about 4,600  
8 chemicals right now that are undergoing in vitro screening  
9 to get some type of bioactivity response data. We are  
10 taking advantage of the availability of these chemicals,  
11 and using it for this trial.

12 In addition to those chemicals, we have reference  
13 house dust from NIST, referenced human serum from NIST.  
14 We've actually had, through a contract, reference silicone  
15 wrist bands that were made for use in this study.

16 I have probably about 15 to 20 different  
17 institutions represented here. I think we're closer to  
18 about 30 now. We have many people that continue to keep  
19 joining to participate. And again, the goal of this is to  
20 kind of let everyone use their method to see what works  
21 best. What are the best tools in specific instances?

22 From all the data that's generated, can we have  
23 experimental data, these MS/MS reference libraries, and  
24 can that benefit the public? And then how can we support  
25 modeling for future applications where we're just never

1 going to have data.

2 --o0o--

3 DR. SOBUS: So part one is completely based on  
4 these synthetic mixtures. We have taken 1,200 of the  
5 4,600 ToxCast chemicals. These are the highest quality  
6 chemicals that we have in terms of their stability,  
7 purity, et cetera. And we've included them in 10  
8 different synthetic mixtures. The mixtures have anywhere  
9 from 100 to 400 individual chemicals, and they're actually  
10 shown here, and they're very lovely looking due to dyes, I  
11 imagine.

12 So we've given all these mixtures out to the  
13 collaborators. Everyone is currently doing a blinded  
14 analysis. They have no idea what's in them. They're  
15 applying their methods, and they're reporting to us what  
16 they think is in the mixture.

17 We then tell them here's what's actually in the  
18 mixture, and they go back and do an unblinded evaluation  
19 and say what did we get right, what did we get wrong, what  
20 did we miss completely? All that data comes back to me  
21 and I pull all my hair out.

22 (Laughter.)

23 DR. SOBUS: A subset of these groups have  
24 actually gotten multi-well plates that include all of the  
25 individual compounds that are in the mixtures. And this





1 community. Tony Williams and his team have done a  
2 phenomenal job of building these massive rich chemistry  
3 databases, and we're sharing that internationally. We,  
4 internally, are developing these tools to allow rapid  
5 identification, characterization, and semi-quantitation,  
6 which I think is critically important.

7 We are working very hard integrating with lots of  
8 international partners that are very skilled in doing,  
9 particularly sediment and drinking water, which is really,  
10 really wonderful. And then finally, we're trying to apply  
11 this everywhere we can for discovery, surveillance,  
12 prioritization, and some of the things I didn't talk  
13 about, effect-directed analysis and biomonitoring.

14 So I think this is a very powerful approach that  
15 we're taking. We're very excited about having partners in  
16 the States, in academia and in local labs, and we're here  
17 to help anyway we can. And if there's any questions I can  
18 answer now or down the road, please don't hesitate to  
19 contact me.

20 --o0o--

21 DR. SOBUS: As I close, I just want to thank our  
22 research team in RTP. We have a terrific group of  
23 chemists, cheminformaticists, and modelers. So I just want  
24 to acknowledge all of their great efforts and open it up  
25 to questions.

1 CHAIRPERSON SCHWARZMAN: Thank you so much for  
2 that presentation. It's the clearest explanation of NTA  
3 I've ever heard.

4 (Laughter.)

5 DR. SOBUS: Thank you.

6 CHAIRPERSON SCHWARZMAN: So I want to start with  
7 the opportunity for Panel members to ask clarifying  
8 questions, and then we'll have a chance for public  
9 comment.

10 It was so clear, we don't have any questions.

11 Oh, Dr. Cranor.

12 PANEL MEMBER CRANOR: A really naive question.  
13 So there are -- presumably, there are 84,000 substances  
14 registered for use in commerce. You claim there are tens  
15 and tens of thousands of them out there. What are they?  
16 Are they re-combined things? Are they rogue things? What  
17 are they?

18 DR. SOBUS: I think, A, there's million of them  
19 out there --

20 PANEL MEMBER CRANOR: Millions. Sorry.

21 DR. SOBUS: -- in my estimation. So tens of  
22 thousands registered for use. What's -- I think anyone  
23 that you talk to that's been doing this work for a period  
24 of time will tell you again that regardless of how big our  
25 screening database is -- ours is 750,000 substances, we're

1 only seeing ten percent or less of what's in a sample. So  
2 what is that 90 percent?

3 PANEL MEMBER CRANOR: Yeah.

4 DR. SOBUS: Most people, A, are calling it  
5 exposure dark matter, which I think is pretty neat.

6 (Laughter.)

7 DR. SOBUS: So I've started to use that term too.  
8 But most people think that these are degradation products,  
9 transformation products, biological metabolites. So for  
10 the last two years --

11 PANEL MEMBER CRANOR: But if they're in -- if  
12 they're in commercial products, are they degraded or have  
13 they been modified to be in those products?

14 DR. SOBUS: That's a complicated question that I  
15 don't want to dig too deep in, but I'm -- I'm thinking  
16 more of when we have the inability to characterize a large  
17 percentage of the sample contents, I'm thinking more  
18 biologically and more environmentally and less based on  
19 the products.

20 PANEL MEMBER CRANOR: Yeah.

21 DR. SOBUS: I'm trying to remember back that --  
22 the data that we got from consumer products came from  
23 contractors, so I don't remember what the break out in  
24 terms of percentages are. I would expect that there would  
25 be fewer degradation products, but certainly some, and

1 certainly some interaction products as well.

2           So the percentage may be a little bit higher in  
3 products, but then you have issues -- issues related to  
4 packaging, and leaching, and migration, et cetera. So if  
5 we're ever going to get at that, I guess -- I guess the  
6 initial question is are those things important? And I  
7 guess the answer to that is you don't know until you have  
8 some idea of what they are.

9           So I'd certainly like to have an idea of what  
10 they are. So, you know, our thinking behind that, and I  
11 think some groups are starting to implement approaches to  
12 do this, are to figure out ways to use modeling systems to  
13 predict what likely degradation transformation metabolism  
14 products are.

15           And if you can generate them within reason  
16 without it exploding, you can then add them to a screening  
17 library and start to look for evidence of them being  
18 there. So if you take, you know, a list of 100 compounds  
19 and many of them you believe to be in a given sample, and  
20 many of them you believe under the conditions that they've  
21 experienced might lead to transformation products, you  
22 know, use some modeling tools to predict what those  
23 products might be, add them to the screening list, and  
24 then see if they're there.

25           So I think that's a first step, and that's

1 something I've been trying to work towards for the last  
2 two years. Finding the appropriate models to do that has  
3 proven to be a challenge for a number of reasons. Again,  
4 we always want to use something that's open, so that if we  
5 can find that it works particularly well, we can then make  
6 it available to folks.

7           So finding appropriate tools that actually  
8 predict the correct compounds, and do it in a manner that  
9 it's consistent with our workflow is something that we're  
10 working towards. Hopefully, we'll have some success with  
11 that in the next year or so, but I think that's the path  
12 to doing it.

13           And even taking it a step further, I've seen some  
14 great presentations from some colleagues that are  
15 participating in our trial. If you can predict what those  
16 compounds might be, you then have a structure -- a  
17 theoretical structure of that predicted compound. Groups  
18 are beginning to predict the theoretical spectra of the  
19 predicted compounds from the known chemical products.

20           So computationally, you can really take this all  
21 the way down the line. So I think a lot of groups right  
22 now are in the proof of concept stage to say can we even  
23 do it, can we make the prediction? And then once the  
24 prediction is made, how valuable is that prediction?

25           So we very much have that in mind to do, to

1 explore, because it's the gaping hole in the work that we  
2 do right now. It's the 90 to 95 percent.

3 PANEL MEMBER CRANOR: That suggests one follow-up  
4 question, if I might. Do you have an example of something  
5 that you either have found or suspect that is pretty toxic  
6 that we didn't know about that wasn't registered, or it's  
7 one of these degradation products that turns out to be  
8 surprisingly toxic? Let's put it -- let me -- an extreme.

9 DR. SOBUS: Short answer, yes.

10 PANEL MEMBER CRANOR: Pardon?

11 DR. SOBUS: The short answer is yes, we do see  
12 that thing quite often. I really don't want to get into  
13 naming specific compounds. I have -- I will say generally  
14 I have colleagues at EPA RTP -- again, where I talked  
15 mostly about suspect screening, so is anything on the list  
16 that we know about present? I have colleagues that focus  
17 more on the true NTA.

18 So basically, if we match against a list,  
19 anything that doesn't match, that's what they want to look  
20 at. And it's a -- it's a laborious project, because you  
21 have to kind of take a very large number of compounds and  
22 kind of go through it manually and say what seems to be  
23 important?

24 In doing that, I know my colleague Mark Strynar,  
25 in particular, has found a number of perfluorinated

1 compounds that aren't registered, that are potentially  
2 by-products of manufacturing processes. There is a decent  
3 number of these things at, what appears to be, decent  
4 concentrations in some water systems. So that would be  
5 one very good example. And I think there's been some  
6 media coverage of that of late, so there's a lot more  
7 follow-up work being done right now with several states  
8 and specifically with our lab as well.

9           So that's one great example of not necessarily  
10 knowing what you're looking for, but looking for the  
11 things that couldn't be matched that appear to be in high  
12 concentration, getting some idea that it could have a  
13 fluorinated signature, and then doing some additional  
14 experiments to kind of fragment that molecule and  
15 reconstruct it and figure out what the structure might be.

16           So that's a great chunk of work that's come out  
17 of our group in RTP over the last really five years or so.  
18 That's probably the best example I have.

19           PANEL MEMBER CRANOR: Thank you.

20           CHAIRPERSON SCHWARZMAN: Other questions from the  
21 Panel?

22           PANEL MEMBER LUDERER: Yeah. Thank you for that  
23 great presentation. And this dashboard just sort of makes  
24 me want to go out and try to use it.

25           (Laughter.)

1 DR. SOBUS: Oh, and there's a mobile app.

2 (Laughter.)

3 PANEL MEMBER LUDERER: But, you know, so -- so I  
4 have some more -- some questions. I mean, you were  
5 talking about all the many different platforms that there  
6 are, and everyone has their own way of doing these things.  
7 And so the -- you know, the study that you're doing where  
8 you're basically sending out the same mixtures to all  
9 these different people, I mean, I think that's going to go  
10 a long way towards start -- trying to start to resolve  
11 that and come up, as you said, with best practices.

12 My question is right now in the dashboard, is  
13 there a way, for example, for -- you know, if a user can,  
14 for example, I don't know, enter -- enter data about what  
15 was the platform they used, or how the samples were  
16 prepared to, you know, help narrow down what the unknowns  
17 are, or is that something that would be --

18 DR. SOBUS: No, that's the goal.

19 PANEL MEMBER LUDERER: -- in the future, but  
20 that's not there?

21 DR. SOBUS: That's the goal. So I guess the  
22 process that we're taking -- to let me step back and say  
23 the dashboard was meant to serve a different client.  
24 Tony -- so Tony was the original developer of ChemSpider,  
25 and we were Lucky enough to get him, I think, two years

1 ago now. And he's been developing this dashboard for us,  
2 which is phenomenal.

3           So I'm very lucky to work with him and his team.  
4 And, you know, I kind of bring in the NTA users group  
5 internally, and say these are some things that we would  
6 love to see the dashboard. So we worked very closely to  
7 try and prioritize which of those things need to happen  
8 first.

9           So it's been about a year and a half now where  
10 we've been trying to integrate these batch searches based  
11 on formula. Another big thing is we want to do batch  
12 searches based on mass. As we're doing that, as we're  
13 building the functionality in the dashboard, in the  
14 background, we have a modeling team that are building the  
15 retention time prediction models, that are building the  
16 media occurrence prediction models.

17           So to the extent that it serves the public, we  
18 try and host those predictions on the dashboard as is,  
19 because we think it's particularly valuable. But even  
20 when we build -- let's say we build a retention time  
21 model, that model, at least initially, is benefiting only  
22 our group because it was built on the conditions of the  
23 analyses that we ran.

24           So kind of step two of that is how do you then  
25 figure out how to adapt that model to support all users?

1 And if you can build that model to support all users, then  
2 you need to have added functionality to allow people to  
3 upload the information that they uses. So I would say  
4 that we're -- that's down the road a little bit, and I'm  
5 probably speaking on Tony's behalf here. I really hope  
6 that we get there. I think he has the same vision for  
7 that, but a lot of things will happen between now and  
8 then. So we're -- we're very much in that proof of  
9 concept stage to say if we build this model, is it  
10 helpful? And if it's helpful, can we make it more  
11 versatile? And if it's more versatile, how do we build  
12 the tools in the dashboard to allow people to enter in  
13 their information. So that's where we're going. It's  
14 just a little bit down the road.

15 CHAIRPERSON SCHWARZMAN: Yeah. Dr. Cranor.

16 PANEL MEMBER CRANOR: One more question your  
17 comments just raised. Is there a way to shortcut some of  
18 this process? You mentioned in your answer to me that a  
19 number of these compounds were fluoride -- had fluorides  
20 in them.

21 Now, presumably lots of the fluoride compounds  
22 are very stable, and so is there a way to do a detection  
23 method for fluorides so you can pick them up or detection  
24 method for other chemical compounds that we know are  
25 toxic? I mean, there's a -- I'm not -- so I'm not a

1 chemist -- neither chemist nor toxicologist, but I've read  
2 in toxicological books or talked to colleagues who have  
3 pictures of the molecules that have toxicological  
4 cancerous endpoints on them.

5 DR. SOBUS: Sure, sure.

6 PANEL MEMBER CRANOR: Are there ways to short  
7 circuit the process and pick out some of those things  
8 early?

9 DR. SOBUS: Yeah, so I have two answers to that.

10 PANEL MEMBER CRANOR: So a speculative question.

11 DR. SOBUS: And again, I'll address this from the  
12 suspect screening, things that hit against the database  
13 versus things that doesn't.

14 You need to do prioritizations in both. Okay.  
15 So I presented on that ToxPi approach. Okay. So these  
16 are for things that hit against the database. We've  
17 written code to automatically process this stuff. So you  
18 hit the database, you pull everything back. In pulling it  
19 back, you pull back the exposure data, you pull back the  
20 toxic data. It goes into a calculation in the program and  
21 spits out those things that are most interesting from a  
22 risk standpoint, taking into account both exposure and  
23 hazard. Okay. So that's -- that's done programmatically.  
24 That's easy.

25 Currently, I want to say we've got bioactivity

1 predictions for fewer than 10,000 compounds. So 750,000  
2 substances in the database. So it stands to reason to say  
3 that the Majority of things in the database don't  
4 currently have exposure and hazard data. So this is where  
5 you have to come up with computational approaches. So we  
6 are working with people within the National Center for  
7 Computational Toxicology - so those are Tony Williams  
8 colleagues and mine - that have built chemical read-across  
9 approaches and QSAR approaches for predicting bioactivity  
10 using models that are based on existing ToxCast data, as  
11 well as in vivo results.

12 So that is kind of something that we are actively  
13 working on now to use QSARs for bioactivity prediction, so  
14 at least we have some estimate of whether or not chemicals  
15 could be bioactive for things that hit against the  
16 database.

17 You know, we're still in proof of concept there  
18 as well. So where the data exists, programmatically we  
19 can pull it back and calculate it very quickly. Where the  
20 data doesn't exist, we're going to have to introduce the  
21 QSARs to make the predictions. So it will happen over the  
22 course of the next year. And then again, it will be done  
23 programmatically over that.

24 In terms of the true non-targeted work, things  
25 that don't hit the database, there aren't that many great

1 approaches. Again, I'll mention two things that are  
2 commonly done, and then the third thing is something that  
3 we do. Number one is look for the biggest things. A lot  
4 of groups out there -- we -- we begin in what's called an  
5 MS-only analysis. So we're looking basically at parent  
6 ions. We try and collect as much information about those  
7 compounds as possible, and then we do our MS/MS  
8 experimentation where we generate fragment ions.

9           Many groups just kind of bypass the first step,  
10 and do an all ions, where they're basically fragmenting  
11 everything, and looking at all the fragments of all the  
12 compounds, which is an overwhelming amount of data, where  
13 they do what's called data-dependent acquisition, where  
14 they say whatever comes in as a parent ion, if it's above  
15 some intensity threshold, we're going to automatically do  
16 the fragmentation and then try and so the annotation  
17 downstream. So there, you're prioritizing based only on  
18 intensity. So that's approach one.

19           Approach two is, again, if you have groups of  
20 compounds, if you want to say -- and this is what's done  
21 in metabolomics, what are the molecules that are  
22 up-regulated in a diseased population versus a  
23 non-diseased population. Again, you're going to take  
24 everything that's here, everything that's here and make a  
25 statistical comparison, and what's left might be 50, 100,

1 200 things that are significantly different.

2           So now you're prioritized to look at those  
3 compounds. On the environmental end, you specifically  
4 mentioned fluorinated compounds. In terms of identifying  
5 halogenated compounds, which are always of great interest,  
6 we use something called a mass defect filter, which -- I'm  
7 not a physicist, I'm not a chemist, but my understanding  
8 is for compounds below a specific molecular weight,  
9 typically 600 or 700 daltons, if you -- if you round to an  
10 integer mass and then subtract from that the accurate  
11 mass, when you have a negative number, that's going to  
12 represent a negative mass defect.

13           And as it turns out, for halogenated compounds,  
14 you very often have this negative mass defect. So we've  
15 introduced into our programs a very quick calculation to  
16 calculate whether or not you have a positive mass defect,  
17 or a negative mass defect.

18           So a lot of my colleagues that studied the  
19 perfluorinated compounds implement the calculation and  
20 immediately look at the negative mass defect compounds and  
21 kind of sort it by intensity and say, here is the highest  
22 intensity negative mass defect compounds. These are  
23 halogenated compounds that are in the sample. Let's start  
24 there.

25           PANEL MEMBER CRANOR: Thank you.

1           CHAIRPERSON SCHWARZMAN: I think I'm going to  
2 save my question for the discussion, because we actually  
3 have a half hour of discussion allotted. And I want to go  
4 to public comment, and then we can get back to the  
5 discussion, because these are all really interesting  
6 questions and I think it's very applicable.

7           We have two comments, one of which is a question  
8 that -- to just be read.

9           So one is a question, which is what is  
10 bioactivity a parameter of the ToxPi key, and how is  
11 bioactivity measured. And the question is specifically in  
12 dust. And I think this is reference to slide 17  
13 describing the workflow. So I think that's just a  
14 reference to bioactivity in the ToxCast data. Maybe you  
15 can just elaborate on that.

16          DR. SOBUS: Sure. I'll do the best I can at  
17 remembering this. So the ToxCast -- so there's a Tox21  
18 effort that is a multi-federal institutional effort that  
19 has looked at a very, very large number of chemicals  
20 across a decent number of bioassays. The ToxCast program  
21 is specific to EPA. It's looked at a fewer number of  
22 chemicals. Again, I think it's about 4,000 right now  
23 across hundreds of bioassays.

24          So each one of these bioassays that I think  
25 were -- many of them were originally developed for pharma

1 applications, basically look at the bioactivity of a given  
2 tested chemical in a concentration response format.

3           So you basically are testing a specific assay at  
4 increasing concentrations to try and determine a value  
5 called an AC-50. So that's an activation concentration,  
6 50 percent. That gives you some information again kind of  
7 on this concentration response format. I'm not exactly  
8 sure how it's done, but it's determined from that  
9 concentration response information, whether it's a hit or  
10 not.

11           For our application, we are basically taking  
12 assay hits, so -- and different chemicals have been tested  
13 across different numbers of assays. So we basically take  
14 a percentage of active assays across all tested assays.  
15 So if something has been tested across five bioassays, and  
16 it was a hit in each assay, it's active 100 percent of the  
17 time. And that's going to be basically the number one  
18 bioactive compound according to our score.

19           So the dashboard, when you enter in your formulas  
20 as a batch search, there is a little button that let's you  
21 export it is a CSV file or in several other formats. You  
22 click on a variety of different data options that you can  
23 actually export. And one of those is on number of tested  
24 assays, and one of those is on number of active assays.

25           So you export that data, and then our programs

1 automatically calculate percentage of active assays. And  
2 that gets rolled into our ToxPi calculation, which is  
3 basically bioactivity, exposure, detection frequency, and  
4 abundance all get normalized to a value of 0 to 1, and  
5 then you sum across all four things. So the most  
6 interesting compound would have a value of 4. It would be  
7 the number one compound bioactivity, number one exposure,  
8 et cetera, et cetera.

9           That's basically how it's done. I have very nice  
10 graphics that shows that in other presentations. So I'm  
11 sorry I don't have that with me today, but hopefully  
12 that's clearly enough.

13           CHAIRPERSON SCHWARZMAN: Also, for the public  
14 member who asked the question, but maybe if I could add  
15 just a quick follow-up question. It sounds like that's  
16 basically prioritizing, in terms of bioactivity, the  
17 higher number of assays on which it's positive, not degree  
18 of positivity. So -- or like lower AC. You know, lower  
19 AC would be more --

20           DR. SOBUS: That is correct.

21           CHAIRPERSON SCHWARZMAN: -- potent. And so if  
22 something is very potent at fewer say receptors, or in  
23 fewer assays, it wouldn't necessarily rise to the top as  
24 much as something that is active in many different assays  
25 and potentially different receptors.

1 DR. SOBUS: That is absolutely 100 percent  
2 correct. And I'll make the pitch out there for anyone who  
3 has interest in optimizing, you know, how we go about  
4 pulling, utilizing that type of information, every single  
5 component of this needs optimization. And at this point,  
6 it's kind of let's show that it works, lets try and apply  
7 it as broadly as we can to figure out where we are, and  
8 then we will work on the optimization of each area. That  
9 is absolutely an area that needs optimization.

10 CHAIRPERSON SCHWARZMAN: So the other written  
11 public comment that I should read is a question actually.  
12 Does current NTA at US EPA focus on parent compounds or  
13 metabolites as well?

14 DR. SOBUS: So we have our core group in RTP.  
15 There are other groups in Athens and Cincinnati and Las  
16 Vegas, and I believe other places that do -- that take  
17 similar approaches. Some of these are a bit more focused  
18 to metabolites. A lot of our ecological studies that may  
19 focus on fish species, looking at fish tissues and stuff  
20 like that, they're going to be a little bit more focused  
21 on the metabolite side of things.

22 Specifically, the work that we do, because I'm in  
23 the Chemical Safety for Sustainability Program, is very  
24 much focused on the parent compounds. Again, the goal is  
25 to try to bridge those two things to the best of our

1 ability, which is going to involve some different modeling  
2 platforms to predict what the likely metabolites  
3 transformation products are going to be. So we're working  
4 towards it, but the work is that I've talked about today  
5 is more or less focused on the parent compounds for the  
6 time being.

7 CHAIRPERSON SCHWARZMAN: Okay. For the moment,  
8 maybe I'll have you surrender the microphone to our two  
9 public comment readers and then have you come back up for  
10 our discussion.

11 Oh, great. Never mind. Hang on to the mic.

12 DR. SOBUS: Good. I didn't want to surrender it  
13 yet.

14 (Laughter.)

15 CHAIRPERSON SCHWARZMAN: Sorry about that.

16 Okay. So we have -- Dr. Alex Hoepker from OEHHA  
17 and Aolin Wang from UCSF.

18 DR. WANG: Thank you so much for a very exciting  
19 and interesting talk. I just wonder, so before you  
20 mentioned that you cross reference a very huge database in  
21 the dashboard. Have you considered whether it would be  
22 helpful to cross-reference curated like library because  
23 some of the -- under some of the platforms like LC/Q-TOF?  
24 There are certain compounds that cannot be detected. So  
25 in order to like reduce false positive and also maybe to

1 put -- to pick out candidates that are of interest to  
2 maybe scientists or of concern say the compounds that are  
3 used in baby products, particularly, would that be anymore  
4 helpful or...

5 DR. SOBUS: It's a great question. And I'll say  
6 this is a question that's debated a lot. A lot of our  
7 European colleagues are -- are pretty adamant that when  
8 you talk about suspect screening, they have to  
9 legitimately be suspects. You have to have some reason to  
10 believe that they might be there. You can't just take  
11 some massive list of things and say, is any of this there?

12 I disagree. I think if you have the means to  
13 screen against a large list, you should. I'm not seeing  
14 any negative consequence of doing that. You make a very  
15 good point that you could have some false assignments in  
16 doing that. We're evaluating to what extent that happens.  
17 But as a get around, Tony and his team have been working  
18 on basically anyone that contacts him that says, I have a  
19 listing of compounds that I have developed as a screening  
20 library.

21 He is accepting that information. He's  
22 developing linkages to our identifiers within the  
23 dashboard and then he's basically tagging that group of  
24 compounds as a specific list.

25 So I haven't spoken to him. We've been missing

1 each other of late. But I think potential future  
2 functionality would be, rather than using everything to  
3 screen, to be able to select specific lists that you want  
4 to hit against as a subset of everything. So I think that  
5 functionality will be there, if people choose to use it.

6 Again, I'm of the mindset that I haven't really  
7 seen a lot of evidence that we're going to get a lot of  
8 mis-assignments or false positives by using a larger list.  
9 Again this is something I've gone back and forth on a lot  
10 of my colleagues about.

11 I think potentially, an important practical issue  
12 is, depending upon which piece of software you're using  
13 and which vendor instrument you're using, some physically  
14 can't handle a list of 750,000 substances. And this is  
15 something that we're dealing with right now. And this  
16 is -- it's really -- it's what's cool about the ENTACT  
17 Study is you're engaging everyone. And everyone is going  
18 to defend their approach for doing it, right?

19 But you have these discussions, and at the end of  
20 those discussions, you hopefully come to some consensus.  
21 Now, let me say with that, we're going to have a workshop  
22 in May of 2018 for all the participants and anyone else  
23 that wants to be there to talk about all of this,  
24 successes, hardships, path forward.

25 So we wound up in ENTACT, we had given the

1 listing of 750,000 compounds to everyone. I had four  
2 people come back and say, I just can't do it. We  
3 physically -- our software will not allow us to do it. So  
4 we made a concession and gave them an abbreviated list of  
5 about 5,000 compounds.

6 Again, I had some people yell at me for doing  
7 that, because we're making it too easy. And I had other  
8 people praise me for saying thank you, now I can actually  
9 do this.

10 So it's yet to be determined what the best  
11 approach is, if there is a best approach. My mindset is I  
12 want to look for everything I can possibly look for and  
13 try and be very careful that I'm not getting a lot of  
14 these false assignments.

15 Good question.

16 CHAIRPERSON SCHWARZMAN: And our final public  
17 comment is from Alexander Hoepker from OEHHA.

18 DR. HOEPKER: Thank you very much for that  
19 presentation. That was really insightful.

20 I wanted to ask about there's really three steps  
21 issued that are cited, right, there's extraction, there's  
22 separation, and then there's detection with mass  
23 spectrometry, and sometimes the extraction and separation  
24 parts are not addressed as much.

25 DR. SOBUS: A little overlooked.

1 DR. HOEPKER: Yeah. And so I was wondering if  
2 EPA is currently developing orthogonal methods that are  
3 trying to fill some of these gaps in extracting and  
4 separating these chemicals. That was one sort of really  
5 big question.

6 But the other one -- the other question I had was  
7 GC seems to be overrepresented as a method of separation,  
8 I think, primarily because of the resolution and the large  
9 spectral databases. LC always seems to fall short. And I  
10 know we have a lot of capacity in LC of course at EPA.  
11 But I'm wondering for GC, because it is so over-emphasized  
12 right now, if there are thermal degradation issues, and if  
13 there are guidelines, red flags for certain chemical  
14 groups that exist that researchers need to be cautious of.  
15 I can think of peroxy acids, other organic acids that are  
16 prone to degradation. But I was wondering if there was  
17 any guidelines for other classes of compounds?

18 DR. SOBUS: There's a lot of good questions.

19 I don't know about any guidelines for thermal  
20 degradation. I would say to the extent that anyone doing  
21 this work has concerns and has an idea of something that  
22 we could look into, I think part of our job is to work  
23 with folks to try and look at what data we have in hand to  
24 see if we can do some experiments to perhaps model that  
25 type of activity.

1           It's interesting that you say that the GC is  
2 perhaps better represented than LC. My perspective is a  
3 little bit of the opposite, I think. We actually -- we're  
4 a fairly small group. We have an older straight TOF, and  
5 we have a newer QTOF and we're getting an Orbitrap.

6           We're trying to build our capacity. We  
7 currently -- really, we have -- we have a GC triple quad,  
8 and we're looking at getting a GC high resolution  
9 instrument, but that's potentially down the road a bit.  
10 So the bulk of the work that we're doing to date is LC  
11 based.

12           So -- and I think a lot of our collaborators  
13 particularly in Europe are primarily focused on LC. The  
14 nice thing is in ENTACT, we have a pretty good split. I'd  
15 say maybe 20 to 40 percent of the methods that will be  
16 employed are going to be GC based and the rest LC based.  
17 But we are working -- again, I think there's going to be  
18 three or four vendors that are applying GC approaches to  
19 characterizing these mixtures.

20           So again, there's going to be a lot of really  
21 interesting data that comes back from the analyses of  
22 these 4,600 samples. And again, someone like you that has  
23 a research hypothesis about thermal degradation, you know,  
24 to the extent that we can use the data that comes out of  
25 that trial to evaluate that hypothesis, that's something

1 we want to do.

2           And we can even make those -- well, we have made  
3 those samples available to folks here for investigations.  
4 So that potentially is a good application. Help me with  
5 your first question again.

6           DR. HOEPKER: In some ways, you're addressing  
7 that already. I'm presuming there's going to be overlap  
8 between LC and GC, so there's ways of validating --

9           DR. SOBUS: Yes.

10          DR. HOEPKER: -- across methods too.

11          DR. SOBUS: Yes. There's -- and I will say,  
12 we're neck deep into our analyses of the SRM's, so that's  
13 the serum and the dust sample. And you've got -- we went  
14 and looked in the literature and tried to pull every  
15 compound that's been identified and quantified with  
16 targeted methods in the SRMs, and basically seeing how  
17 many of them can we see with this method, how many of them  
18 can we see with this method, how many can we see with both.

19           So that speaks to the kind of -- the reach of a  
20 given method for a particular chemical space, which gets  
21 to one of the questions I asked of we don't know how many  
22 different methods would be needed to kind of fully  
23 characterize a sample, if that's even necessary.

24           Did I miss anything as part of that first  
25 question? I feel like I did.

1 DR. HOEPKER: The extraction part.

2 DR. SOBUS: The extraction part. Good.

3 (Laughter.)

4 DR. SOBUS: No, that's a great question. And we  
5 went back and forth on this. The reality is you are one  
6 hundred percent correct. And I think more attention needs  
7 to be paid to that. Most people right now just try and  
8 use something that is going to give them, you know, kind  
9 of the biggest swath of information.

10 The biggest number of chemicals without biasing  
11 against any particular group. You're never going to get  
12 it all. Ideally, you would be doing multiple extraction  
13 procedures, multiple columns for chromatography, you know.

14 But it's all about bandwidth, right? If you do  
15 multiple extractions, multiple clean-ups, multiple columns  
16 for chromatography, multiple ionization sources, multiple  
17 platforms, LC and GC. You know, it gets multiple  
18 databases, big, small. You know, it just explodes.

19 So you've kind of got to do what you can to prove  
20 the methods using something that's a little bit more  
21 tangible. And then once you get a handle on that and have  
22 some type of benchmark for doing it, then I think you can  
23 extend. Like we want to get in to using HILIC columns,  
24 rather than C18 and C8 or in addition to. But it's just  
25 you've only got so many people, you've only got so much

1 time. You can only do so much in a period of time, but  
2 that's absolutely something that I think has been not  
3 considered enough and something that needs to be  
4 considered more in future experiments.

5 Great questions.

6 CHAIRPERSON SCHWARZMAN: So we have time now for  
7 general Panel discussion, as well as audience discussion.

8 So, Dr. Cranor.

9 PANEL MEMBER CRANOR: Yes a follow-up question.  
10 You've given a very exciting and interesting  
11 presentation -- closer. Okay. At the same time, we know  
12 that there are a number of substances out there that are  
13 fairly toxic that we haven't done much with. Of the  
14 things that you've discovered, how do they compare in  
15 terms of toxicity, or do you even have any idea? This  
16 really goes to the question of the social benefits of the  
17 elaborate programs you're putting together, which are very  
18 exciting, I think, but in terms of the public health  
19 benefit.

20 DR. SOBUS: Um-hmm. So I guess I would  
21 potentially restate that as you asking me of all of the  
22 things we find, are the things that are currently looked  
23 for amongst the very top of that list? So if you find  
24 5,000 things, and there's 300 things in NHANES, are those  
25 300 things at the list of the -- top of the list of the

1 5,000 that we're finding?

2 PANEL MEMBER CRANOR: Yeah, and how they compare  
3 with the things we already know about?

4 DR. SOBUS: Right. So I'd say the things that we  
5 know to look for, that we've already determined we care  
6 about due tend to be at the top of the list, but they're  
7 amongst many other things that are also at the top of the  
8 list, and in many cases at much higher concentrations.

9 And --

10 PANEL MEMBER CRANOR: The things you found are at  
11 higher concentrations?

12 DR. SOBUS: The things that we're finding.

13 PANEL MEMBER CRANOR: Wow.

14 DR. SOBUS: And, you know, I'm not a chemist, but  
15 it's interesting when we -- when I work with the chemists,  
16 and, you know, my job is kind of to facilitate to write  
17 some of these programs, to kind of make everything happen  
18 quickly, when this stuff comes back, and I share a results  
19 file with a chemist and they look at what's at the top of  
20 the list, they're often blown away, because they -- you  
21 know, they may recognize a compound, but never thought  
22 that it might be present, and certainly not present at  
23 that concentration.

24 So it is really, really eye opening. And a lot  
25 of stuff makes sense, you know. I want to -- I always

1 show this example of piperine. Piperine is what's in  
2 black pepper. It's basically in every home that we look  
3 at. That makes sense. You know, basically everyone uses  
4 pepper.

5 (Laughter.)

6 DR. SOBUS: Nobody ever really thinks about, you  
7 know, if you do this, basically anywhere in your home,  
8 you've got a bunch of piperine on your finger.

9 (Laughter.)

10 DR. SOBUS: So there's -- you know, there's a  
11 thousand chemicals just like that that are in every home.  
12 And when you sit back and when you look at the list of  
13 things that are on your finger, you're like, wow, that  
14 makes sense. So then you have to step back and say, what  
15 are the potential biological consequences of me being  
16 around that all the time.

17 So there are some really cool things that you  
18 find, but I think a lot of it makes a lot of sense. You  
19 know, the consumer product example that we gave I think is  
20 really cool of, you know, basically 80 percent of the  
21 things that came back aren't previously associated with  
22 consumer products. So what the heck are they and how are  
23 they getting there? You know, are -- you know for  
24 formulations are they coming from packaging?

25 So there's a lot of investigative work to be done

1 I think when you find some of these things. And that's  
2 why you want to be pretty careful about confirmation. You  
3 don't want to go, you know, chasing some compound that you  
4 think is there that you could be wrong on.

5 But I think there's -- there's a ton of work to  
6 be done as you generate these data sets to really dig in  
7 and say what does this mean? And that's what I would --  
8 even if people don't have the capacity to make the numbers  
9 themselves, that's why we're trying to build the tools so  
10 that we can do the analyses or provide the tools to folks,  
11 so they can do the analyses, but also make the data  
12 available, so people can begin to mine it and say what  
13 does this really mean? Because there's just a ton of it  
14 there, and no one person or even small group of people can  
15 fully understand what's going on in any reasonable amount  
16 of time. So we need to kind of take it on as a big team  
17 effort, I think.

18 PANEL MEMBER CRANOR: Thank you.

19 CHAIRPERSON SCHWARZMAN: I'm going to ask -- take  
20 the Chair's prerogative and ask a question I didn't get to  
21 ask before. So one is just sort of a follow up on that,  
22 but then it kind of connects to my other main question,  
23 which -- so is it fair to say do you think that we don't  
24 have a very clear idea about toxicity for many of the  
25 compounds that you're detecting precisely, because

1 they're -- they're the known unknowns, in a sense. Like,  
2 you find them, you can identify them, but they weren't the  
3 things that we thought to look for, because we have an  
4 understanding about how they're used and whether they're  
5 particularly toxic.

6           So would you say that you're generating sort of a  
7 list of commonly found compounds for which we don't have  
8 much toxicity data.

9           DR. SOBUS: You nailed it. And I think I view  
10 that as a big part of my job. Okay. So the ToxCast  
11 program, you know, you can only procure and evaluate so  
12 many chemicals, right? And they're up to, like I said,  
13 close to 5,000. So that's a huge undertaking. It  
14 involves tons of resources and a lot of time.

15           So given that there are 4,600 ToxCast chemicals,  
16 and 750,000 substances in DSSTox, what's the next list of  
17 things you want to screen in ToxCast?

18           So again, I view that as part of my job is to say  
19 here's the things that we routinely find. Let me work  
20 with some QSAR modelers and say does this have any  
21 potential to be bioactive? So that is a very important  
22 approach for kind of nominating things that should go in  
23 the next cycle of the program.

24           I didn't mention, but I think some of the  
25 criticisms of existing in vitro high throughput screening

1 programs is the lack of capability for testing metabolic  
2 capacity. We're constantly testing the parent compounds,  
3 and the limited testing on mixtures.

4           So this is something that again we really hope to  
5 contribute to these high throughput screening programs is  
6 to say here are our things that we're finding over and  
7 over again. We believe this to be the structure. This is  
8 not a parent chemical. This is not registered for use,  
9 but it's something that's out there everywhere at very  
10 high concentrations.

11           Perhaps, we should test the activity of that  
12 compound in ToxCast. At the same time, you know, if you  
13 look at every possible combination of every possible  
14 chemical, you have an untestable number of mixtures,  
15 right? So how do we use our empirical data to say here's  
16 a cluster of compounds that always co-occurs and it always  
17 co-occurs when you have this exposure source. If this is  
18 relevant say to children's health, maybe this is a mixture  
19 of seven things that we really need to test the activity  
20 of to see if it's additive or if there's a synergy,  
21 something like that. So that's something through our work  
22 we hope to contribute back to the ToxCast program.

23           CHAIRPERSON SCHWARZMAN: So that's really  
24 helpful, and it kind of connects to my earlier question,  
25 which is you mentioned that you didn't talk a lot about

1 the application of these methods in biomonitoring studies.  
2 And it's really exciting to hear about, you know,  
3 environmental applications, and consumer product, and  
4 household dust and all of that sort of thing. It's lots  
5 of interesting stuff.

6 But I wonder if you could elaborate a little bit  
7 about some of the challenges and opportunities in applying  
8 NTA in biomonitoring samples. And it occurs to me just  
9 from a not very expert perspective in this that there's a  
10 lot of -- but as a physician, I have some sense of the  
11 number of biological compounds that are in our bodies and  
12 what is -- is there -- is it a ton of work to distinguish  
13 among or between sort of big classes of chemicals to which  
14 we're exposed from the environment, and our own endogenous  
15 chemistry?

16 DR. SOBUS: Great question. So I actually had  
17 dinner with I Steve Rappaport last night, and we talked  
18 about that for about two hours. The -- you know, the  
19 definition of the exposome, as representing the totality  
20 of exposures is -- is conceived, construed differently by  
21 groups of people. And it's done in such a way that it's  
22 kind of fit for purposes for their application.

23 So whereas, you know, my mandate -- agency's  
24 mandate, my activities to support that mandate are to  
25 evaluate a lot of the environmental component of the

1 exposome. Certainly, there's that endogenous component  
2 that's critically important, and often the focus of many  
3 exposomics, and certainly metabolomic studies.

4           So I think -- I haven't heard any naysayers yet,  
5 but my thinking is a lot of the effort to date done in  
6 biological samples, particularly serum, has focused on the  
7 higher concentration compounds, more often the endogenous  
8 molecules, because they're easier to detect, because  
9 they're at much higher concentrations.

10           Steve has pointed out through publications,  
11 through presentations that it is harder to get at the  
12 environmental chemicals because they are at lower levels,  
13 and you have some signal suppression. You've got all of  
14 this other biological stuff that's really highly  
15 concentrated, and then the environmental stuff is lower  
16 level.

17           So that's why we're looking at serum right now.  
18 And I will say it is more of a challenge to characterize  
19 the environmental chemicals in biological samples,  
20 particularly serum, because of some of those signal  
21 suppression issues. We do tend to see lots more stuff and  
22 see stuff more successfully in the environmental samples.  
23 But there's no reason that we can't do xenobiotic, a  
24 component of biological sample at the same time that we do  
25 the endogenous component.

1           So, you know, the path forward -- and we're  
2 working with some partners in the field like Scripps that  
3 manage XCMS and METLIN. You know, we've incorporated into  
4 METLIN, the DSSTox compounds. So now folks that do  
5 metabolomics and that look routinely at the endogenous  
6 compounds for some type of enhancement that could be  
7 mapped to a biological pathway, and to an apical endpoint,  
8 you know, we're providing them with the ability to also  
9 look for the environmental chemicals.

10           So I think in the near future, we're going to be  
11 in a paradigm where you either have individual groups that  
12 look at both sides of it, and then can draw relationships  
13 between the xenobiotic, exogenous, as well as the  
14 endogenous. But even if you have, you know, distinct  
15 communities, they're at least aware of what's going on on  
16 the other side, and then folks that can bridge the data  
17 across, that's where we need to be.

18           But I don't think we're too off -- too far off  
19 from doing it. So again, you can -- you can do what you  
20 can do. And from my standpoint, you know, there needs to  
21 be more work done on the environmental end, just because  
22 it's lagging behind the metabolomics community, but I  
23 think we're really close to having kind of a bridging of  
24 those efforts. And I really don't see a whole lot  
25 standing in the way of it.

1 CHAIRPERSON SCHWARZMAN: Other Panel comments.  
2 Dr. Quintana.

3 PANEL MEMBER QUINTANA: Do we have time?

4 CHAIRPERSON SCHWARZMAN: Yes, we have time.

5 PANEL MEMBER QUINTANA: Hi. When you were  
6 talking, I was thinking about the old days. I think I'm  
7 old enough to talk about the old days in, for example,  
8 genotoxicity testing. In a way, we used to do the  
9 opposite of what you're doing. So instead of looking for  
10 everything in a sample, we might take say urine from a  
11 group of nurses that handle cancer drugs, and compare it  
12 to urine of nurses who didn't, and see whether the urine  
13 was more mutagenic, for example.

14 DR. SOBUS: Right.

15 PANEL MEMBER QUINTANA: So not even looking for  
16 what was in there.

17 DR. SOBUS: Right.

18 PANEL MEMBER QUINTANA: Or the same thing for  
19 house dust. Is an extract of house dust more mutagenic  
20 than this house dust? I'm just wondering, if you take,  
21 for example, your house dust samples, 56 samples, and you  
22 tested the extracts say in ToxCast or something, and you  
23 looked at the 10 most mutagenic -- just taking that as an  
24 endpoint. There's many other endpoints, reproductive,  
25 whatever. Like the 10 most mutagenic compared to the 10

1 least mutagenic, could you subtract out a bunch of  
2 compounds and see what was different or how much have you  
3 explored that kind of approach to kind of use a downstream  
4 gatekeeper on environmental samples, like house dust, like  
5 urine, like other things.

6 DR. SOBUS: Sure. Sure. It's a great idea.  
7 It's something that I'll say we're actively pursuing.  
8 We're not far enough along on any particular study where  
9 we have results to report, but this is something we're  
10 working towards.

11 I had one other slide on NTA applications, and  
12 the bottom two bullets were grayed out. The first of  
13 those two bullets is called the effect-directed analysis.  
14 So that's exactly what you were describing, where you  
15 effectively take an extract of a sample of interest,  
16 introduce it to some type of test system, look for a  
17 response in that test system, and then you fractionate at  
18 different stages the mixture itself. So making it into  
19 two fractions and testing the two fractions. And then if  
20 one of the two fractions is bioactive break that active  
21 fraction into two, and you just successively break it into  
22 smaller fractions, so you figure out what are the primary  
23 components of that mixture that are driving the  
24 bioactivity.

25 So this is something that's done a fair bit. We

1 have a lot of European colleagues that are absolutely  
2 experts in this field. It's something we want to work  
3 towards. I believe there are actually platforms now where  
4 this can be done in an automated fashion. I think  
5 that's -- we talked about this when we had a workshop. I  
6 think our workshop was two or three years ago now in RTP.  
7 And we had a whole discussion about the specific issue.  
8 And again, the field was kind of split between some people  
9 being adamant that this is the correct approach, and other  
10 people saying it does work very well, but it's extremely  
11 time-consuming.

12 I think there was a comment that said, one  
13 effect-directed analysis, one Ph D. This is --

14 (Laughter.)

15 DR. SOBUS: It takes that long to -- to do a  
16 successful experiment. Now, people really pushed back on  
17 that and said that, you know, the tools have been  
18 developed in an automated way, so that this can be done  
19 fairly quickly. So I think you're absolutely on point. I  
20 think now that we have the means -- again, you can't  
21 really do effect-directed analysis without having the  
22 underlying non-targeted analysis tools. You can identify  
23 the active fraction, but if you can't figure out what the  
24 chemicals are in that fraction, you know, have you really  
25 succeeded.

1           So I think now that we have the underlying tools  
2 to do the NTA, we can start to introduce it for different  
3 effect-directed analysis applications.

4           PANEL MEMBER QUINTANA: Just a very quick follow  
5 up that I guess the care to which the samples were  
6 collected, if you're using real-world samples, that would  
7 be to me proof of something we should be worried about.  
8 If it's in real people's homes, you know, like you said,  
9 or in urine for kids, I'm just wondering if you've thought  
10 about using some of the National Children's Study samples  
11 that were collected with such care from Irvine and other  
12 places, because they do have the air that -- the water,  
13 the -- all the components that might feed into that  
14 exposure. So I'm just curious if you've explored that.

15           DR. SOBUS: We absolutely have. Not necessarily  
16 National Children's Study, but certainly other well-known  
17 programs where samples have been collected and archived  
18 with care. We've had some -- some conversations with  
19 folks across the pond. So EPA and NIEHS share a campus.  
20 So we've been having some conversations lately about  
21 opportunities for some existing studies and for some  
22 upcoming studies, where we can start to kind of go down  
23 that path.

24           I think one of the important things about me  
25 being here is, you know, we're not CDC. We have a fairly

1 small group of people. We have really two instruments  
2 right now that we're using. We're not in a position to be  
3 looking at thousands of samples right now. Our role is  
4 really to help build the methods to do proof of concept,  
5 to build the computational tools, and to make that  
6 available to the public.

7           And again, you know, what we've found is be it  
8 through, you know, data coming back from contractors or  
9 from collaborators, the real challenge to the community  
10 right now is folks saying, yeah, I can do that. We can do  
11 NTA. And then you have no expectation of what's going to  
12 come back.

13           And then sometimes, money is spent, time goes by,  
14 data comes back, and you say this wasn't what I was  
15 expecting, because there's no way of really evaluating  
16 when you say you can do NTA, what is it that you can  
17 really do. And that's why there's -- it's critical to  
18 have again some standards to which we hold laboratories to  
19 say if you can do it, you need to be able to at least do  
20 this.

21           And at the same time, there really needs to be a  
22 network of laboratories, so that when we have big studies  
23 with thousands of samples, maybe millions of samples, we  
24 have the capacity to take that on, because this isn't  
25 something -- you know, it's neat when we make these

1 presentations and people get excited and, oh, you can do  
2 this work. And, you know, our initial inclination is,  
3 yeah, let's do it. And then my managers always look at me  
4 and they're just like what are you talking about? You  
5 can't that do that. You don't have the capacity to do  
6 that.

7 (Laughter.)

8 DR. SOBUS: So you -- you need to pick and  
9 choose, but I think the path forward for that is building  
10 a network of practitioners so that you can spread samples  
11 around, and then adhere to some agreed-upon standards for  
12 performing the work. And I think that's how -- the only  
13 way we're going to handle doing rigorous analyses on some  
14 of these precious samples for which great care has been  
15 taken to collect and store.

16 CHAIRPERSON SCHWARZMAN: Dr. Cranor, and then Dr.  
17 Luderer.

18 PANEL MEMBER CRANOR: One more hard question.

19 (Laughter.)

20 PANEL MEMBER CRANOR: I think many of us thinking  
21 about toxic substances grew up on carcinogens, and  
22 mutagens, and such things as that. On the -- of current  
23 concern, are xenohormones, you might say. And they  
24 typically operate at much lower con -- much lower  
25 concentrations, and they can disturb things. Can your

1 system accommodate those, incorporate them, and identify  
2 them? I mean, this is a really speculative question,  
3 but --

4 DR. SOBUS: When you -- so when you say identify  
5 them, is this under the assumption that they've been  
6 identified as being bioactive for endocrine effects, or  
7 something like that, at low concentrations, are you asking  
8 if we can identify the effects or are you asking --

9 PANEL MEMBER CRANOR: Can we identify other  
10 ones --

11 DR. SOBUS: -- can we identify the chemicals,  
12 given that --

13 PANEL MEMBER CRANOR: -- that will have those  
14 effects?

15 DR. SOBUS: Broadly I think, yeah. I think to  
16 the extent that -- I mean, you say the chemicals are very  
17 low level. Obviously, we're not going to have the  
18 sensitivity with a broad non-targeted method on a QTOF  
19 that you would on a dedicated triple quad where you're  
20 focusing on 10 compounds. You're just not going to get  
21 it, but you're not too far off.

22 You know, we're -- in these SRM samples, we're  
23 picking up on pretty much everything that people have seen  
24 with targeted, trace level, triple quad analyses. So I  
25 think, you know, the vendors are really pushing the

1 sensitivity on these instruments to where we can see them.

2           Whether or not we would anticipate if they would  
3 have effects at low concentration, like similar compounds,  
4 truthfully, we have a staff of computational modelers  
5 whose job it is to figure that out. So that's why I kind  
6 of turned to them to say, what do you have available to  
7 you to predict bioactivity, so that if I find this  
8 compound, I can come to you and say is this something I  
9 need to be worried about? Do you anticipate this thing  
10 could be bioactive in any way? Does it resemble anything  
11 else you've looked at before that was bioactive with a low  
12 concentration? So that is the workflow.

13           PANEL MEMBER CRANOR: Thank you.

14           CHAIRPERSON SCHWARZMAN: Dr. Luderer.

15           PANEL MEMBER LUDERER: Yeah, I wanted to just  
16 kind of get back to this -- the question that we were  
17 talking about with the metabolomics versus the, you know,  
18 exposome, and that, you know, a lot of -- there's maybe  
19 more work that already that has been done on the  
20 metabolomics side. And so one question I had in terms  
21 of -- I mean, obviously, if you're biomonitoring, it is  
22 really important to, I think, integrate those two things,  
23 and whether in the dashboard, in your 750,000 compounds,  
24 does it include any endogenous compounds or is it only,  
25 you know, xenobiotics --

1 DR. SOBUS: I'm pretty sure it has some, but I  
2 could -- I could not even take a guess at how many.

3 The reality is Tony and his team have so many  
4 connections in the field, and it's kind of -- and I  
5 believe -- I forget who the person was at CalEPA. He was  
6 just in contact with someone last week about an OEHHA  
7 chemical list, that was integrated in the dashboard over  
8 night. So it's -- his job is kind of networking and  
9 identifying people with rich databases that he can map to  
10 through the dashboard.

11 So when we started this work, we were looking at  
12 a list of, I want to say, like 7,000 compounds. And then  
13 version 2 was 32,000 compounds, and then it was 300,000  
14 compounds, and now it's 750,000 compounds. This has  
15 happened over the course of two years.

16 So -- and again, this isn't just random pull  
17 stuff in and it's there. They've built automated programs  
18 and protocols for doing curation of that chemical  
19 information. And with that, they put flags on it. So if  
20 you go out to a massive public database with millions of  
21 compounds, there are often inconsistencies in the drawn  
22 structure, in the CAS number, which may no longer be  
23 registered in synonyms, in SMILES notation, and InChIKey.

24 So the programs effectively look across all of  
25 those components to make sure that they are consistent.

1 And if they're not consistent, they flag them. And once  
2 they're flagged, there's at least some attempt for at  
3 least a subset of the compounds to fix the errors. So  
4 those data quality flags exist in the database.

5           So to the extent that Tony and the team can kind  
6 of map out or link in and at least run the automated  
7 programs to kind of check all of these different pieces of  
8 information, that's how stuff gets pulled in. So how much  
9 of that that's got pulled in over the last two years that  
10 represents endogenous, I really don't know. I would say,  
11 by no means, is it anywhere near the majority, but some of  
12 it's probably in there.

13           Again, I think the most explicit action to kind  
14 of bridge those two things was Tony sharing the DSSTox  
15 content with those that manage METLIN which is a  
16 repository for mostly endogenous compounds.

17           CHAIRPERSON SCHWARZMAN: This may be an overly  
18 simplistic question, but I'm wondering how -- like for the  
19 purposes of this Program, of the Biomonitoring Program,  
20 something that sounds interesting to me to think about as  
21 a Panel would be to start from -- get our hands on a list  
22 of compounds that is like -- this one that I was referring  
23 to kind of the known unknowns of it's not the ones that  
24 are looked at in NHANES.

25           DR. SOBUS: Sure.

1           CHAIRPERSON SCHWARZMAN:  It's like what is the  
2 kind of next set of compounds that we don't know a lot  
3 about that's showing up in environmental samples, or  
4 consumer product samples, dust samples, or biological  
5 samples that we should be thinking about is this something  
6 we should be looking at in a biomonitoring study in  
7 California

8           DR. SOBUS:  That's a fantastic question.  And I  
9 mean it's a very logical and important question, and  
10 perhaps surprisingly something that we haven't really  
11 talked about doing.  I think everyone is so kind of  
12 focused and nested in their individual application that no  
13 one has really stepped back to say, okay, broaden it to  
14 what should we be looking for on a wide scale.  I mean, we  
15 certainly from -- you know, you could do that a number of  
16 ways.  You could say, do you want to focus that on a  
17 medium or do you want just want to -- to just broadly say  
18 give me a list of a hundred chemicals that you think are  
19 the next biggest deal?

20           That's something that I think any one group could  
21 kind of nominate their list.  But what potentially would  
22 be more powerful is to look to the community, the people  
23 that are doing this, and say what is it that you think for  
24 any reason is the most important thing?  And then, you  
25 know, it's kind of -- it's kind of a voting and a tally.

1 And we could put together that list.

2 And I think the network of participants would be  
3 a fantastic group to kind of query and say what would you  
4 nominate? From that, we could absolutely put together a  
5 really interesting list.

6 CHAIRPERSON SCHWARZMAN: That's my wish list.

7 (Laughter.)

8 DR. SOBUS: That's a -- that's a fantastic  
9 question.

10 CHAIRPERSON SCHWARZMAN: That's what I would like  
11 to suggest we do, because it's like using non-targeted  
12 analysis just direct targeted analysis to inform targeted  
13 analysis.

14 DR. SOBUS: That's absolutely the idea.

15 CHAIRPERSON SCHWARZMAN: And that's what, I  
16 think, would be a really exciting kind of next step.

17 DR. SHE: Jianwen She. I like to also help  
18 answer this question that the California Biomonitoring  
19 Program would use low targeted approach to select the  
20 chemical groups for us to work on.

21 So Dr. Sobus presented a very excellent talks.  
22 For the Biomonitoring Program, we need to look at what's  
23 practical for us, what the EPA can do, what can we do? So  
24 I think for us generally, we will focus on application of  
25 the tools that are already built in.

1           And incidentally, like some of the -- for me,  
2 some of the unknown unknown problems that's like artificial  
3 intelligence. We can build off a goal, narrow down on the  
4 board game, or we can broaden the search, like an  
5 exhaustive search that you can find all of the unknowns or  
6 you can do your ..... search, so you focus on specific  
7 areas. As artificial, that's how human solves unknown  
8 problems.

9           So for -- so I use this energy to compare mass  
10 spectrometry based non-targeted search for us. Instead of  
11 the broadest first search, we need a depths-first search.  
12 What I mean is depth-first search. Depth first search  
13 first, we focus on class of chemicals. That many class of  
14 chemicals you can pick up. For mass spectrometry, like  
15 Dr. Sobus said, all halogenated chemical compound that  
16 have specific mass spectrometer features, mass  
17 deficiencies, isotope profiles, mix and match easier to  
18 search.

19           But as poly-halogenated compound the ECL is  
20 working on, they may consider this group of chemicals who  
21 find a chemical maybe unique, new there. Like, we found  
22 poly-halogenated dioxins that's not registered. Aside far  
23 dioxins not registered, mainly by-product from the  
24 chemical or industrial process can be a group for us to  
25 look at.

1           For us, we -- in the past, we proposed using BP-3  
2 analogs. So we still working on this group of chemicals.  
3 So that's my general comment.

4           And anything regarding specific progress, like  
5 Dr. Yu-Chen Chang can comment, if you are here. Do you  
6 want to make any comment on this part of the work?

7           DR. CHANG: So I'm Yu-Chen. In response to one  
8 of the SGP meeting that California EPA presented, there  
9 was a concern for UV filters. So at EHLB, our primary  
10 focus is to do untargeted screening primarily focused on  
11 the class with a structurally-related compound, which is  
12 benzophenone class.

13           So right now, since we have a very limited man  
14 power, so there are very limited compound we can screen.  
15 And database is right now limited, and we have a whole  
16 list of structural library available screen for this class  
17 of compounds. And we are in the process to expand our  
18 class to environmental -- to phthalates, which also listed  
19 as designated chemical for the list as a whole class,  
20 which we don't have the capability to screen for the whole  
21 class.

22           So right now, that's what our focus is on, the  
23 UVA filter, benzophenone structurally-related class.

24           Thank you.

25           CHAIRPERSON SCHWARZMAN: Any further comment from

1 the Panel or from the audience that we should include  
2 before we move on?

3 I would just -- oh, there's one more.

4 Martha.

5 DR. SANDY: Hi. I'm Martha Sandy from OEHHA.

6 I just wanted to follow up on what -- one of the  
7 points that Alex had made about the extraction method for  
8 non-targeted analysis. And you have also talked about,  
9 well, what is the chemical space we're covering?

10 I would like to suggest, if possible, you could  
11 come up with a couple different ways of extraction to try  
12 to cover as much of the chemical space as you can and just  
13 remind folks of an example, where looking for food  
14 mutagens for years and years and years, researchers did  
15 extractions and looked at one type of extract, and they  
16 found a lot of things but they missed acrylamide.

17 (Laughter.)

18 DR. SOBUS: It's a point well taken. And, you  
19 know, I don't know the best approach, if, you know, there  
20 can be composite extracts. But we were -- we're very  
21 conscious of this. And again, we could have bit off a bit  
22 more and shared the raw samples with participants in our  
23 trial and asked them to kind of apply their own extraction  
24 procedures.

25 If it turns out that we should have done that, I

1 will take all the blame for missing an opportunity. And  
2 perhaps we should have, but there's so much variability  
3 downstream on processing that we thought if we had too  
4 many undefined variables - this is a classic statistics  
5 problem, right - that if you've got 10 variables and 30  
6 methods, it's going to be very difficult to pinpoint which  
7 of those variables is more largely responsible for  
8 affecting the outcome than something else.

9           There's got to be a reasonable relationship  
10 between the number of variables and the number of -- in  
11 this case, of laboratories. So I think -- we collectively  
12 made the decision, but I can take the blame for it if it  
13 was a bad decision, to kind of just make one extraction  
14 and to send that out.

15           I think in follow-up efforts that could be done  
16 individually by any laboratory, but certainly by a  
17 consortium as well, that is something we need to look into  
18 more, because obviously there's going to be some issues  
19 with being too narrow in the wet chemistry procedures.

20           CHAIRPERSON SCHWARZMAN: All right. Thank you  
21 very much. I'm very intrigued by this notion of both how  
22 we can expand our non-targeted analysis studies, and also  
23 how we might inform some subsequent decisions about more  
24 targeted screening based on this future collaborate on  
25 getting this wish list -- my wish list of up and coming

1 compounds.

2           So I want to trans -- thank you so much for your  
3 presentation --

4           DR. SOBUS: Thank you all very much.

5           (Applause.)

6           CHAIRPERSON SCHWARZMAN: -- and I want to  
7 transition now to turn the mic back over to Dr. Lauren  
8 Zeise.

9           So this is the part where we're going to honor  
10 Dr. Asa Bradman for his service to the Committee.

11           DIRECTOR ZEISE: Yes. We wanted to take a few  
12 moments to thank Asa. Asa, would you like to stand up?

13           Great. Okay.

14           (Thereupon an overhead presentation was  
15 presented as follows.)

16           DIRECTOR ZEISE: So on behalf of the  
17 Biomonitoring California program, I'd like to thank you  
18 for your outstanding service as a member of the Scientific  
19 Guidance Panel.

20           Asa was on the original Panel appointed in 2007  
21 by Governor Schwarzenegger. And he assumed the Chair  
22 position in 2015. So in this program's 10th year -- it's  
23 Asa's 5th -- 10th year as serving on the Panel. And we  
24 really do have deep appreciation for all of your  
25 outstanding guidance, hearing your wide-ranging knowledge

1 of assessing exposures to chemical hazards, long  
2 experience garnered as a researcher at UC Berkeley, and a  
3 lot of groundbreaking studies, particularly in children,  
4 including the CHAMACOS Study.

5           So we really did benefit so much from -- the  
6 Program did benefit so much from your unique insights and  
7 advice as an SGP member, and then your enthusiastic  
8 member -- your enthusiastic leadership as Chair.

9           So as I think we all know, you've long been an  
10 advocate of biomonitoring of children. And again, your  
11 many thoughtful contributions during SGP meetings have  
12 helped us shape key aspects of the Program's work over the  
13 past 10 years, and put special attention on exposures  
14 affecting disadvantaged groups and children.

15           So I know I speak for all Program staff in saying  
16 we'll miss working with Dr. Bradman as Panel members and  
17 SGP Chair. And we wish him the very best as he takes on  
18 fresh challenges, including some collaborations with State  
19 researchers. And we look forward from hearing about your  
20 future contributions.

21           Thank you, Asa.

22           (Applause.)

23           DR. BRADMAN: I just wanted to thank you, Lauren,  
24 and everyone on the Panel, and the Program, and the  
25 community that's been so supportive of this Program. And

1 again, you know, I just want to emphasize how important  
2 this effort really is. And I have a feeling -- I'm here  
3 today. I have a feeling I'll be back at future meetings,  
4 or at least participating online, because these issues are  
5 so important to me, and also we'll have ongoing  
6 collaboration.

7 Thank you.

8 (Applause.)

9 CHAIRPERSON SCHWARZMAN: So that concludes the  
10 morning portion of the meeting. And I think we're ending  
11 about 15 minutes before the Program says so, but I have  
12 permission from the powers that be to do so with hope that  
13 we can actually move the afternoon session a little bit  
14 earlier, just because some Panel members have to leave a  
15 little bit early and we want to get to the discussion that  
16 has to do with a Panel vote in time to make sure we have  
17 all the Panel members here.

18 So we have an hour for lunch, which means we'll  
19 resume at 1:30. And that's -- just to highlight for  
20 people, that's a little bit shorter lunch than we've had  
21 in the past. So it's recommended that folks eat in the  
22 cafeteria that's right here and be back to resume the  
23 meeting right at 1:30 for our afternoon session.

24 And before we break for lunch, I just want to  
25 invite Fran Kammerer to give us our -- oh, it's Carl

1 today. Sorry. Changed staff -- to remind us about the  
2 Bagley-Keene.

3 STAFF COUNSEL DeNIGRIS: Carl DeNigris, staff  
4 counsel for OEHHA. Just a reminder that the Panel is  
5 subject to Bagley-Keene, and refrain from discussing any  
6 matters that are before the Panel outside of this forum.

7 Thanks. Have a good lunch.

8 CHAIRPERSON SCHWARZMAN: Okay. So we'll conclude  
9 the meeting for this morning and resume at 1:30.

10 Thank you.

11 (Off record: 12:32 p.m.)

12 (Thereupon a lunch break was taken.)

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1                   A F T E R N O O N   S E S S I O N .

2                   (On record: 1:34 p.m.)

3                   CHAIRPERSON SCHWARZMAN: Okay. We're going to  
4 start. I'm going to bring the afternoon session to being.  
5 Thank you for coming back from lunch.

6                   Before we start, I just wanted to mention  
7 apparently we're out of the packets of materials at the  
8 table when you walk in. But just so that everybody knows,  
9 all of the materials are posted online. So if you're  
10 wanting access to anything and it's not available in  
11 physical form here, you can get it on the Internet at the  
12 Biomonitoring California website.

13                   So I'm -- I'd like to welcome Dr. Roy Gerona. He  
14 is here from UCSF. He runs the Clinical Toxicology and  
15 Environmental Biomonitoring Lab at UCSF, and has  
16 collaborated with the Program on Reproductive Health and  
17 the Environment at UCSF on suspect screening for  
18 endocrine-disrupting chemicals in biological samples. And  
19 most recently, Dr. Gerona's lab has been developing  
20 biomonitoring methods for pesticides including glyphosate,  
21 glufosinate and other organophosphate metabolites. And  
22 that's bio -- in biological samples obviously.

23                   So he'll be presenting today his work on  
24 glyphosate. And thank you so much for coming to be with  
25 us.

1 (Thereupon an overhead presentation was  
2 presented as follows.)

3 DR. GERONA: Okay. Am I on?

4 All right. I'm on.

5 Okay. First of all, I'd like to thank Sara and  
6 the Advisory Panel for actually inviting us here to  
7 present the work that we're doing on glyphosate. I will  
8 not be the only one presenting this afternoon. There has  
9 been changes in the schedule of the graduate student who  
10 has done a lot of work in the method development for this  
11 work. So I'm going to be introducing him first, let him  
12 talk, and then I will follow it up with the applications  
13 that we've been working on.

14 Axel Adams graduated from the University of  
15 Wisconsin - Madison in 2014, which is where I came from as  
16 well. I did grad school there. And immediately after  
17 being admitted to the Joint Medical Program between UC  
18 Berkeley and UCSF, approached me and has in mind that what  
19 we wanted to do for his Master's thesis is really to  
20 develop methods for biomonitoring.

21 So the work that he will be presenting here is  
22 his Master's thesis. I am lucky to have him in the lab.  
23 His is a very good analytical chemist. And what he'll  
24 present -- or what you'll -- what you'll hear from him  
25 this afternoon are the challenges part, because often the

1 paradigm here in America is you give the challenges to  
2 your graduate students --

3 (Laughter.)

4 DR. GERONA: -- and then you seek out the  
5 opportunities later once the challenges have been  
6 resolved. So she will be -- he will be presenting the  
7 challenges part in developing the method for glyphosate,  
8 and I will be following it up with opportunities where we  
9 think we would be able to contribute in epidemiological  
10 studies concerning glyphosate.

11 Axel.

12 MR. ADAMS: Yeah. Thank you so much for having  
13 me here today.

14 --o0o--

15 MR. ADAMS: So to begin with the analyte in  
16 question, N-(Phosphonomethyl)glycine, I'll be referring to  
17 it as PMG, it's IUPAC name, also generically known by the  
18 trade name glyphosate.

19 It's a broad spectrum herbicide. It was  
20 originally discovered by a Swiss chemist, and was  
21 initially patented as a chelating agent. And it actually  
22 works as a chelating agent to induce its herbicidal action  
23 by chelating a manganese co-factor for one of the enzymes  
24 involved in aromatic amino acid production in plants.

25 It's the most widely used pesticide in the world

1 with 113.4 million kilograms used in the United States in  
2 2014 alone. That represents about a 30,000 percent  
3 increase from its original usage in 1974. And it's the  
4 most widely used domestic pesticide. It's also -- which  
5 means that in terms of like home usage, applying on lawns,  
6 applying in home gardens, things of that nature.

7           So it's used in a couple different ways. The  
8 primary way, and probably the way most people are  
9 familiar, is in conjugation with PMG-tolerant crops. So  
10 things like Roundup Ready soybeans, Roundup Ready corn  
11 that are developed to have enzymes that are resistant to  
12 the herbicidal action of PMG.

13           But it's also widely used with non-PMG tolerant  
14 crops. For instance, it's used as a preceding application  
15 to kill down weeds in fields. It's also used as a  
16 desiccation agent. So they'll apply it to, for instance,  
17 grain to hasten drying prior to harvest. And it's also  
18 used as a spot application in different agricultural  
19 settings like orchards, and also municipally, so like  
20 along railroad tracks, and parks, things like that.

21                   --o0o--

22           MR. ADAMS: So lately, the safety status of this  
23 pesticide has been contested. It was classified as not  
24 likely to be carcinogenic to humans by the US EPA in 2015.  
25 In 2014, the International Agency for Research on Cancer

1 found an association between PMG exposure and  
2 non-Hodgkin's lymphoma in a meta-analysis, that looked at  
3 a number of different studies. It was slightly  
4 problematic because some of the studies were rather old at  
5 that point, and did not really assess exposure at moderate  
6 levels. And then there were issues with kind of the  
7 distribution of these settings and where they took place.  
8 So it may not have been representative of exposure in the  
9 general sense.

10 But nevertheless, in 2015, IARC performed its  
11 first review of PMG and classified it as Group 2A probable  
12 carcinogen. And there have been a number of different in  
13 vitro, in vivo studies kind of looking at different ways  
14 in which it might be carcinogenic. And there are a number  
15 of different proposed mechanisms, which is kind of beyond  
16 the scope of this talk, as we are analytical chemists.

17 --o0o--

18 MR. ADAMS: Most methods for measuring PMG have  
19 been indirect methods. So what that means is you have the  
20 analyte in question, you conjugate a different moiety to  
21 it, so you're able to measure it better, and then you're  
22 able to do, for instance, gas chromatography-mass  
23 spectrometry. And there have been very few direct  
24 methods.

25 Direct methods are advantageous because they

1 require less processing, which makes it easier to do  
2 large -- a large number of -- or very high throughput  
3 analysis, which is important for biomonitoring. They also  
4 generate fewer kind of toxic waste products. And green  
5 chemistry is kind of the goal in this day and age.

6           Of the direct methods, even fewer are applied in  
7 human matrices. So 2008, there's Wang et al. that looked  
8 at serum. It had a limit of quantitation of 5 nanograms  
9 per ml with ion-pair chromatography. You had Yoshioka et  
10 al. who used Obelisc N column, which is also the column we  
11 used. They had a limit of detection of 20 nanograms per  
12 ml, limit of quantitation of 90 nanogram per ml.

13           And then you had Jensen et al. 2016 that were  
14 looking at urine and also milk matrix. And their LOD and  
15 LOQ were reported based on the individual transitions. So  
16 I'll talk about this a little bit more. That's kind of  
17 problematic, and isn't very common in the literature. And  
18 there are several other problems with that paper that kind  
19 of call into question whether or not it's a good method.

20                           --oOo--

21           MR. ADAMS: So PMG itself is a very difficult  
22 analyte. It's kind of like -- being able to measure it  
23 effectively is sort of like the holy grail of small  
24 molecule mass spectrometry. And the reasons for that is  
25 that it's very low molecular weight. It has basic and

1 acidic features. It doesn't have an analytically useful  
2 group like a fluorophore or chromophore. It has several  
3 different pKa's, and it's also a chelating agent, as I  
4 mentioned previously, for both bivalent and trivalent  
5 cations.

6 But fortunately, it's a very stable molecule. So  
7 the half-life in water is about 33 days. In soil, it's  
8 half-life has been reported up to about 250 days. And  
9 there's not very much data in half-life in other, you  
10 know, human or biological matrices.

11 Also, it is very stable in terms of in the soil.  
12 So it's a little bit unique among pesticides as it adsorbs  
13 very strongly to soil particles. And they can stay, you  
14 know, for great periods of time in soil. They can also be  
15 degraded on the soil absorption surface, so AMPA, which is  
16 the most commonly reported breakdown product, but also to  
17 sarcosine and formaldehyde that are subsequently broken  
18 down just to CO2 and water.

19 --o0o--

20 MR. ADAMS: So what we did is we used liquid  
21 chromatography tandem-mass spectrometry, a very high  
22 fidelity method for analyzing small molecules. And the  
23 basic principle is it's a series of a different separation  
24 techniques. You separate first based on the overarching  
25 chemical features of the molecule, so its polarity. And

1 then you separate based on the mass-to-charge ratio of the  
2 parent ion. You fragment that and then you separate it,  
3 based on the mass-to-charge ratio of the daughter ions.  
4 So it's kind of like a -- like a Russian nesting doll of  
5 separation science.

6 --o0o--

7 MR. ADAMS: So when we were developing the  
8 method, first we looked at tap water, because that would  
9 be a very simple method to do. We were able to do that  
10 with external calibration, kind of a standard method. We  
11 just acidified the water, add our C-13 internal standard,  
12 and run a calibration curve, kind of very bread and butter  
13 analytical toxicology.

14 --o0o--

15 MR. ADAMS: Add then what we found is in terms of  
16 precision accuracy studies, it's very straightforward, as  
17 you would imagine, not a complex matrix.

18 --o0o--

19 MR. ADAMS: Now, the issue was in terms of  
20 transferring it to urine and breast milk, those are much  
21 more complex matrices than tap water, as you'd imagine.  
22 And there were several different problems with both  
23 matrix. So usually what we want to do is we want to, if  
24 we're doing external calibration, run a calibration curve  
25 and a matrix blank. So in terms of urine, we would get

1 drug-free urine or urine that's free of the analyte.

2 But it had been reported in the literature  
3 actually by Jensen et al, that drug-free urine that was  
4 commercially available had trace -- a trace signal of PMG  
5 in it. So that's problematic, because then where are you  
6 getting your drug-free urine?

7 We also reproduced that and found that in the  
8 drug-free urine we were obtaining, we could also detect a  
9 signal. You can use synthetic urine which is not quite as  
10 good as human urine, because you're not capturing all the  
11 different aspects of the metabolome.

12 Similarly with milk, there's no commercially  
13 available like breast milk or even bovine milk reference  
14 standard. So really what that leaves you is internal  
15 standard addition. So that's internal calibration.

16 So basically what you're doing is you're spiking  
17 the analyte at different levels into the individual sample  
18 that you have. So if you have it for Mr. A, you're going  
19 to have four aliquots of Mr. A's urine spiked at different  
20 concentrations. It's really a robust method. It's a  
21 little bit annoying because you have to do more aliquots  
22 and submit it for analysis, but it is very robust.

23 The other problems with PMG is it elutes very  
24 near the solvent front. So in the liquid chromatography,  
25 you basically have a dumping of a bunch of different



1 addition. There's also translational matrix effects,  
2 which you can't deal with standard addition. So we were  
3 able to demonstrate that it was a very valid way of  
4 dealing with matrix effects to use standard addition.

5 And we were able to achieve a limit of detection  
6 of 0.1 nanograms of PMG per ml, so 0.1 parts per billion.

7 --o0o--

8 MR. ADAMS: Moving on to milk, which is a very,  
9 very difficult matrix. It's far more difficult -- a far  
10 more difficult matrix than, for instance, blood. And the  
11 reason for that is it's a complex mixture. So in milk you  
12 have a bunch of different fractions. You have a solution  
13 of globular whey proteins, you have colloidal casein  
14 suspension, and you have lipid emulsion. The casein  
15 suspension is very interesting, because in the casein  
16 micelle, which is a protein micelle, you have a lot of  
17 different cations.

18 And as you recall, PMG is a chelating agent. So  
19 there's -- it's thought that it could be interacting with  
20 these casein micelle. So you have to drop out the casein  
21 micelles, drop out the whey proteins, deal with the  
22 lipids, because the lipids are a very problematic source  
23 of matrix effect, in analytical chemistry, and then  
24 finally, you know, get as much of your analyte as you can  
25 in the aqueous solution.

1           And there is a number of different ways you can  
2 deal with each one of these different sources of obstacles  
3 separately. But what we did is we diluted it to deal  
4 with -- basically to separate out the mixture, acidifying  
5 it, basically dropping out the casein micelles, going past  
6 isoelectric points so they drop out solution. You can  
7 then centrifuge it to bring those to the top of the  
8 solution, because they're a very low density protein.

9           You can then denature the globular proteins with  
10 dichloromethane, centrifuge it, drop that to the bottom,  
11 and then dilute the aqueous solution again. The DCM also  
12 acts as a liquid-liquid extraction, so you're able to  
13 remove a lot of the phospholipids, and then just have the  
14 PMG in your aqueous fraction.

15                               --o0o--

16           MR. ADAMS: So what we were able to find is that  
17 when we looked at precision and accuracy studies in milk,  
18 we were able to get good precision, good accuracy, very  
19 good linear range across a pretty wide range -- dynamic  
20 range of 1 to 40 parts per billion. We were also getting  
21 relatively good recoveries, as low as 1 part per billion,  
22 and a limit of detection of slightly under 0.2 parts per  
23 billion for milk and a limit of quantitation of 1 part per  
24 billion for milk.

25           There are two statistics that are important for

1 standard addition, the cap on Q statistics, which are  
2 calculated based on the level that you're expecting to see  
3 in your sample. And there haven't been good studies on  
4 actual levels that are observable in milk, so these are  
5 uncalculated. We can kind of estimate what they would be.

6 And the reason you would do that is you would  
7 adjust the fortifications for the standard addition in  
8 order to ensure that you have like the maximally valid  
9 internal calibration curve.

10 --o0o--

11 MR. ADAMS: But in summary, it looks like it --  
12 this way of dealing with the complexities of the milk  
13 matrix, in conjunction with the complexities of PMG as an  
14 analyte, is a relatively reliable way of doing it -- doing  
15 so.

16 So problems with our method, there are several.  
17 One is that the Obelisc N column, which is a HILIC column  
18 that we used, and I mentioned previously is also used by  
19 Yoshioka et al. for dealing with blood as a matrix,  
20 degrades relatively rapidly. And it's an expensive  
21 column. We've been able to do about 1,000 injections in  
22 milk before it degraded.

23 So it is kind of an expensive method in that  
24 regard. And that was also reported by Yoshioka that they  
25 had problems with degradation just from how dirty the

1 matrix is. And HILIC columns tend to be a little bit more  
2 fragile. The standard addition also necessitates 4  
3 aliquots per individual sample. So if you're analyzing  
4 Mr. A's urine, you're going to have 4 aliquots of his  
5 urine that you have to run.

6           So if this is -- our method is 6 minutes, so you  
7 have 4 by 6 minutes of time on the machine, and as you  
8 multiply that out for large and larger data sets, that's a  
9 substantial increase in time, and also time in preparation  
10 for the lab technician who's preparing the samples.

11           However, it's slightly counterbalanced, because  
12 you don't have the external calibration curve to create.  
13 And we figured that, you know, the increase in validity of  
14 the method counteracted the extra time it took. Also,  
15 with the current method, AMPA was not detectable due to  
16 the short retention time. I mentioned previously that  
17 it's eluting very close to the elution front. So you're  
18 basically having a dumping of all these different analytes  
19 at the same time, and AMPA's was in there.

20           And as result, you have a higher background noise  
21 that makes it -- your limit of detection and your limit of  
22 quantitation rise higher. Also, whether or not AMPA is a  
23 good analyte to look at is up for debate. It had  
24 initially been thought that AMPA was the degradation  
25 product of PMG. However, there's more and more evidence

1 suggesting that in, you know, for instance, soils or  
2 environments that are naive to PMG, it's more commonly  
3 broken down by soil bacteria sarcosine and formaldehyde,  
4 and subsequently to CO2 and water.

5 In which case, you don't have a very good stable  
6 degradation product to analyze. And so it may be in terms  
7 if you're actually looking at accessing exposure, it may  
8 just be worthwhile to focus on PMG itself, as opposed to  
9 this other degradation product, which may not represent  
10 the major pathway of degradation.

11 --o0o--

12 DR. GERONA: Thanks for that clear presentation.

13 I think that it's the most exciting part of the  
14 presentation that we have this afternoon.

15 (Laughter.)

16 DR. GERONA: So now that the challenge has been,  
17 you know, answered, or responded to, our task in the lab  
18 is to find ways by which we can apply this method. And  
19 then fortunately, there are a lot of studies. One thing  
20 to use the method.

21 I would preface this -- this set of slides by  
22 saying that we are analytical chemists. And so what we  
23 intend to do is to present you the data and have other  
24 experts start the dialogue on how that data should be  
25 interpreted. We will not comment on the implications of a

1 lot of this data, because we think that it is outside our  
2 area of expertise. So we're presenting you the data. So  
3 let's start the dialogue and let's plan on what to do with  
4 the data.

5 Obviously, some of this data needs to be  
6 verified. That's why we're also looking for partner  
7 laboratories that can verify the results that we will be  
8 presenting to you this afternoon.

9 So immediately, or even before the method  
10 development started, we already had partners who are  
11 actually require -- who require that the data be  
12 developed, so that we can provide precise and accurate  
13 data on glyphosate in biological samples.

14 We have five cohorts that we're currently working  
15 on. Some of them have been completed. Some are ongoing,  
16 and some are about to start. So for the first cohort, the  
17 Detox Project, it actually started as the Feed the World  
18 Study. And this is a study that is sponsored by the  
19 Organic Consumers Association.

20 These are crowdsourced urine and tap water  
21 samples collected across the United States. At the end of  
22 the -- at the end of the project, we have about 252 urine  
23 samples collected. This study has been completed. And so  
24 we'll present to you the data that we've obtained from  
25 this work.

1           Now, one of the lessons that we learned from the  
2 Feed the World or the Detox Project is that the midwest  
3 is, as expected I guess, has -- basically, the cohort from  
4 the midwest from The Detox Project has the highest levels  
5 of glyphosate that we're detecting. And so we were very  
6 interested on finding specific cohorts in the midwest to  
7 see what would be the levels of glyphosate that we will be  
8 measuring.

9           So there are two cohorts that we were able to  
10 find. The group of Paul Winchester in Indiana has  
11 provided us with urine samples from pregnant women. The  
12 total number of samples is about 283. We will be  
13 presenting the results of the first 83 samples this  
14 afternoon. We have run all 283. The remaining 200 needs  
15 to be analyzed.

16           The third cohort is very interesting. This  
17 cohort was not really meant for studying glyphosate. This  
18 was actually a cohort from Gail McCarver at the Medical  
19 College of Wisconsin. And this was a cohort where Gail  
20 was looking at BPA levels the first year of life.

21           So Gail has time points from day 0 to day 365  
22 urine samples from these infants, where we previously  
23 measured BPA. But when the -- when the project was over,  
24 we have a lot of left-over urine samples. And so when  
25 this opportunity came in, we thought this is an excellent

1 set of samples to look at whether -- especially the day 0  
2 urine, day 0, day 2 urine.

3           And so we collected and gathered leftover urine  
4 samples. We were able to actually collect 36 samples --  
5 my timer is not flashing yet, so that's good -- 36  
6 infants. And we have complete samples for day 0, day 2 as  
7 the first time point; the first week, so that's about day  
8 7, day 10; days 180 to 185, that's six months; and then  
9 day 360 to 365, that's about one year.

10           It's ongoing. We have done our initial studies,  
11 and I would just say that we have some interesting  
12 results, in the sense that we're finding glyphosate even  
13 in some day 0 urine samples. So we know that this is  
14 controversial. It's going to be controversial. I guess  
15 if we were to report it. And that is the reason why it's  
16 very important that our study is verified by an  
17 independent lab, that the blind samples can be forwarded  
18 to the independent lab to see whether they can reproduce  
19 the results that we're getting in the laboratory.

20           I'm not going to be discussing more of cohort 3,  
21 except saying that, you know, that's our initial finding  
22 that we're getting.

23           Cohort 4 -- and this is again -- you know, we're  
24 very lucky to be able to collaborate with the MARBLES  
25 Study. Cheryl Walker from UC Davis will be able to

1 provide us with 60 breast milk samples, 30 from children  
2 with -- that were diagnosed with autism, and 30 that are  
3 age-matched control. So we're getting the breast milk of  
4 obviously the mom.

5 We're also going to be getting the urine samples  
6 collected when these children are being conceived, first  
7 trimester, second trimester, and third trimester urine  
8 samples from this specific cohort. We haven't received  
9 the samples. We're very excited to receive them and start  
10 the analysis.

11 And ultimately, our group wanted to actually  
12 demonstrate that -- or, not demonstrate, but ask the  
13 question would the mom's exposure, is that actually being  
14 delivered to the baby in the first year of life. If the  
15 only source of food of the baby is mom's breast milk, are  
16 we going to be able to find glyphosate in the infant's  
17 urine.

18 So it's a study where we're collecting matched  
19 serum, urine, breast milk, infant urine. And we're  
20 recruiting for this particular study.

21 --o0o--

22 DR. GERONA: So let me -- let me present the  
23 first cohort. So as I've said, the Detox Project is a  
24 crowdsourced project. What we did here is the Organic  
25 Consumers Association partnered with us. We were the lab

1 testing the urine samples for the individuals who wanted  
2 to actually have their glyphosate level in their urine  
3 tested by a laboratory.

4           So in this particular study, we were limited, of  
5 course, with data that we can get from individuals. So  
6 unfortunately, there's no questionnaire that is associated  
7 with this study. We were given limited demographic  
8 information like the gender, the age, where the samples  
9 came from, and some times we're given tap water as well.

10           So most of the samples came from the United  
11 States.

12           Is this also a pointer?

13           Oh. Okay. So we were able -- so the collection  
14 or the analysis of sample -- Five minutes. I can go over  
15 by another five, right, so I have 10 minutes?

16           (Laughter.)

17           DR. GERONA: So there are 252 urine samples. We  
18 actually collected samples between April of 2015 to  
19 February of 2016. As you can see, the number of samples,  
20 a bit lopsided with the west coast. There's about 124  
21 samples. It's very interesting that when we did this  
22 work, what we were expecting is that most of these people  
23 who are sending us samples are very much aware of the  
24 debate that is going on on glyphosate.

25           So we were expecting that these people are



1           So one of the things that you worry about,  
2 especially when you report back to the lay public is if  
3 you tell them, oh, your glyphosate is 3.1 ppb, how will  
4 that person interpret it, right? It's always the  
5 interpretation of that data that is an issue.

6           So when we give them the report backs, we give  
7 them some reference points on how they could possibly  
8 interpret the data. This is not the only -- the only  
9 information that we provide. There are -- they get a  
10 sheet where they have the averages for all the different  
11 parts of the United States, the averages for the months  
12 that we were collecting, but they also get this table  
13 where we have their values in green, the running average  
14 at the time that we actually gave the results to them.  
15 And these all other dots is in that batch where is  
16 there -- where is their value with reference to the other  
17 people that we measured for that particular batch.

18           We also compare what we're getting to other  
19 studies that have already been reported in the  
20 literature --

21           Seven minutes now.

22           -- a European study, which is by Hoppe et al.,  
23 and then the averages of the farmer study, where  
24 they -- these are farmers from Minnesota and North  
25 Carolina, where the method, I think, is HPLC, where they



1 see, this is a heat map of the agricultural use of  
2 glyphosate in the United States. There is a particular  
3 interest in the midwest, because that is where most of the  
4 agricultural use is happening. So we were thinking, okay,  
5 so it would be very interesting to see the exposure levels  
6 of people in the midwest.

7           And that's what we did here. So we have, in this  
8 particular table, we're comparing it with the Detox  
9 Project, where we have level -- the statistics for the  
10 whole U.S., the whole midwest, and what we saw in Indiana.  
11 You have 83 samples from Indiana. The detection frequency  
12 is about similar to the detection frequency of -- for the  
13 midwest, which is slightly higher than the average for the  
14 whole United States. And the geometric mean is slightly  
15 higher also for Indiana.

16                           --o0o--

17           DR. GERONA: I have three slides, so you will let  
18 me finish, okay?

19                           (Laughter.)

20           DR. GERONA: So this study we were able to  
21 contextualize the results a little bit more, primarily  
22 because when they collected the urine samples, they were  
23 already looking at potential association between pesticide  
24 use -- and not necessarily just glyphosate -- of pesticide  
25 use and potential sources of exposure. So they have a



1 DR. GERONA: And so the other thing that we did  
2 is we divided the levels into terciles. And as you can  
3 see here, what's interesting is when we did that  
4 particular division of the data, we found out that -- we  
5 looked at full-term births only -- that there is a trend  
6 in this particular data. I won't comment in the  
7 statistical significance yet, because we're still adding  
8 on data on this particular cohort.

9 But you can already see a trend that the ones  
10 with low values have longer pregnancy length than the ones  
11 with the higher values. We actually did a different type  
12 of analysis -- or they did a different type of analysis  
13 which I will show in the next slide.

14 For the birth weight percentile, how did we get  
15 the birth weight percentile? So we looked at published  
16 curves of birth weight at specific gestation age, and  
17 looked at the birth weight percentile of each of the  
18 individual babies that are part of this particular study.

19 And again, so what you're seeing here is that the  
20 ones -- the ones that have high levels of -- the high  
21 tercile has the lowest birth weight percentile average.

22 --o0o--

23 DR. GERONA: And then finally the last slide that  
24 I'm presenting, so I told you that, you know, we did the  
25 opposite. We looked at -- instead of dividing the values,



1 DR. GERONA: And so what are the future  
2 directions? We would want to revalidate the milk method  
3 with actual milk samples from UC Davis. And Axel is still  
4 not off the hook. He will still come back to the lab and  
5 do this.

6 (Laughter.)

7 DR. GERONA: We will finish the current  
8 collaborative epidemiological studies that we're doing.  
9 We want to assist another lab with methods run transfer.  
10 This is key because we think we think -- we really wanted  
11 another lab to independently validate the results that  
12 we're getting.

13 There have been several groups that have already  
14 approached us to do work. Aside from those five, there  
15 are three more groups that we're probably working with in  
16 terms of urine, and breast milk, and tap water levels.  
17 And we would incorporate this method in a multi-analyte  
18 method for polar pesticides. We're also -- these are some  
19 of the analytes that we are working on as well. These are  
20 only some. There are several we have about, I think, 25  
21 now, organophosphate metabolites, glufosinate, and 2,4-D.

22 --o0o--

23 DR. GERONA: And with that, I would like to thank  
24 all our funders and collaborators. This was Axel when  
25 he's much younger.

1 (Laughter.)

2 DR. GERONA: Some people who have done a lot of  
3 work on this also. It's not -- Matt Friesen is my RA who  
4 started the lab with me. So he initially was developing  
5 the method for tap water. And Ann Gordon is the QA  
6 manager of the lab. Anita Wen is our expert in method  
7 development and validation, and acts as a consultant in  
8 the lab.

9 These are the collaborators that we have in those  
10 cohorts, which I think has already been mentioned. And  
11 special thanks to Asa, because we have been consulting him  
12 also in this study.

13 And with that, we will -- Axel will take all the  
14 questions.

15 (Laughter.)

16 (Applause.)

17 CHAIRPERSON SCHWARZMAN: Thank you very much.  
18 We'll start with clarifying questions from the Panel.

19 PANEL MEMBER LUDERER: I have one.

20 Can you hear me now?

21 Okay. I thought -- and I may have just  
22 misunderstood this, but when you were talking about the  
23 milk analyses -- by the way this is really exciting work  
24 and thank you very much for this -- the great  
25 presentation.

1           I thought you said something about that PMG is  
2 interacting with the casein and the micelles. So I --  
3 what -- and maybe I didn't understand what you meant by  
4 interacting. But then if you're centrifuging to remove  
5 the micelles, aren't you going to be removing some of the  
6 PMG in the sample?

7           MR. ADAMS: Yeah. So that's a very good point.  
8 And so the reason -- the way that we deal with that is by  
9 adding the heavy isotope of the PMG itself. So basically  
10 you add that at the beginning, and then you're able to do  
11 recovery studies. And we were finding that we weren't  
12 losing that much during the analysis of that heavy  
13 isotope.

14           The other good thing about standard addition is  
15 it kind of takes into account the losses that you're going  
16 to have within the sample itself. So you might have --  
17 you know, depending on the external matrix that you're  
18 using, you may have different casein concentrations, and  
19 then you'd expect a difference. So standard addition deals  
20 with that in and of itself.

21           The other thing is that the casein interaction is  
22 more of theoretical one. It hasn't been well  
23 demonstrated, and well -- because there isn't very much  
24 data on analytical chemistry in milk as a matrix. So what  
25 we did is we looked at analyses for other different

1 smaller organic acids in milk.

2           And the thought is that -- so casein is a linear  
3 protein -- low density and linear protein that organizes  
4 itself in micelles in milk. And if you can bring it  
5 down -- bring down the pH of the solution past the  
6 isoelectric point, you basically disrupt the associations  
7 between the proteins. And since it's such a low density,  
8 it actually floats to the surface.

9           And so presumably you'd have freeing of the  
10 glyphosate within the aqueous layer, and then you're able  
11 to dip your pipette tip, sample the aqueous, and then do  
12 your subsequent liquid-liquid. But the internal standard  
13 is how we deal with losses.

14           PANEL MEMBER LUDERER: That was added before you  
15 did all the process?

16           MR. ADAMS: Yeah, exactly. That's how we  
17 calculated recovery.

18           DR. GERONA: It may -- I may just add to this.  
19 You know, when -- we started developing the method in  
20 2014, because we actually have collaborators already in  
21 2014. When we look at all the methods that are published  
22 in the literature, they're all using external calibration.

23           And so we tried our darndest to actually repeat  
24 or reproduce some of the -- some of the work that they  
25 were doing. But with tap water we're able to do that.

1 With urine and breast milk, we were not able to do that.  
2 And our findings consistently is that every urine sample  
3 is unique.

4 And so you would get very different matrix  
5 effects from one urine sample to the other. That's when  
6 we basically made the decision and say, okay, what other  
7 alternative methods can we use? If external calibration  
8 is not working, then what other -- what other armaments  
9 does analytical chemists have to approach this problem?

10 And that's when we shifted to standard addition.  
11 That's the reason why, you know, every time we were asked  
12 about the method -- and we have been requested by several  
13 different laboratories to basically just tell them what  
14 our method was. And we say -- we're telling them that,  
15 you know, we want our work published, because it's -- in  
16 the end, it's really a simple method that we're able to  
17 work on, at least for urine, not for milk. Milk, as you  
18 see, is quite complicated in that.

19 MR. ADAMS: And something, too, I failed to  
20 mention is another reason why internal calibration is kind  
21 of perfect for milk as a matrix is during the lactation  
22 course for an individual, the constitution of the milk  
23 varies. So like early on, you have the cholesterol. It's  
24 a more heavy lipid milk. And then it goes and you get  
25 increasing protein, and then you get increasing levels of

1 lactose, all of which build in background signal.

2           And so if you're going to be using, you know, a  
3 commercially available drug-free human milk, where is that  
4 milk from? And also, there are variations in the  
5 composition of milk throughout the day. There are  
6 variations based on what the mother is eating. And so  
7 internal calibration is a way of dealing with all those  
8 different factors at the same time.

9           CHAIRPERSON SCHWARZMAN: Other Panel questions?

10           Okay. Then we have some time for public comment.  
11 Is there any public comment?

12           There's one in the room. Did you want to --

13           DR. GERONA: So you believed our presentation?

14           (Laughter.)

15           DR. SHE: Thank you very much for the excellent  
16 presentation regarding delineating a very polar compound  
17 in the very complex mixture like urine.

18           So as you're very well aware, the limitation of  
19 the standard addition method, which is throughput. But as  
20 far as you already know, the standard addition method  
21 solves the rotational matrix effect perfectly, which was  
22 traditionally used for the metals analysis, because they  
23 do not require longer analysis.

24           So basically, I think you find a very smart  
25 approach to solve the problem. And my only comment is how

1 you resolve the throughput problem for the wide, large  
2 epidemiological research, because basically, every single  
3 sample you do a calibration on it. That's the first  
4 comment.

5           The second one is we did try using it because for  
6 the OPFR. And I discuss with Myrto, we use the standard  
7 addition, but we haven't summarized data, because OPFR  
8 tend to be very polar.

9           And another approach I think you are already  
10 searched. You referred as indirect analysis, which is  
11 more kind of -- I guess, you mean is derivatized, which  
12 after derivatized analytes, the polarity has changed. And  
13 then now you can extract the analyte and separate it from  
14 matrix, so -- which is indirect, but when the industry  
15 start with the PMG, that's EPA request that provide method  
16 on the derivatized method that did that. Our laboratory  
17 used derivatized approach, for example, to analyze the  
18 DAPs. That's the CDC's method.

19           And also, we use derivatized method to analyze  
20 the PAH. And so it seem to be okay after you derivatize.  
21 Now, suppose you eliminate the matrix effect. One is you  
22 compensate matrix effect by vertical approach. So you  
23 can -- can you give us some -- a little bit more details  
24 in review of this method?

25           And also, we -- I'd like to emphasize, we may be

1 a candidate to get your method transferred, because we  
2 work on different disease, but if that's Nerissa and other  
3 program leaders' decision, but we are possibly a good  
4 candidate, because we work on the urine, and on the  
5 similar chemicals.

6 DR. GERONA: You know, thanks for the -- for your  
7 offer, because we're actually -- so to answer your first  
8 question about throughput. That is the reason why we need  
9 partners, and that is the reason why we're looking for  
10 labs where we can assess whether the method that they will  
11 be developing its -- if it's standard addition, we want to  
12 assess whether there's concordance in the data between our  
13 lab and your lab.

14 The high throughput problem with standard  
15 addition, I mean, that's one of the reasons why we --  
16 originally, the Feed the World Project really wanted us to  
17 continue the work, but they were predicting thousands of  
18 samples every month. And I told them with -- with the  
19 approach that we have, it will not just be compatible with  
20 the number of -- that number of samples.

21 For one, this is just one part of several studies  
22 that we're doing in the lab. We have probably a dozen  
23 projects going on at the same time. And so you're right,  
24 you know, one of the disadvantages of the, you know,  
25 standard addition method is instead of analyzing one, for

1 every sample we analyze four, and then we -- obviously, we  
2 run that in replicates also. I would let Axel comment on  
3 some of the -- some of the methodologies that were also in  
4 the literature, and I'll also supplement it if needs to  
5 be.

6 MR. ADAMS: Yeah. So thank you. In terms of the  
7 throughput, it's really not that bad for the urine  
8 preparation, because that's basically dilute and shoot  
9 just by four.

10 With milk, it's a pretty arduous method, because  
11 it's -- you know, it's -- as you precipitate out the  
12 casein and phospholipids, you have to make sure not to  
13 disrupt it back into the solution when you sample the  
14 aqueous layer from below. And it's a very -- it's kind of  
15 a technical method from start to finish, and it takes a  
16 lot of time.

17 That we don't have a good answer for. It's --  
18 you know, the method is at it stands. And it's a little  
19 bit slower. But that being said, I think it's a pretty  
20 robust approach for a lot of different analytes in milk.  
21 And there are not very many methods published for  
22 different analytes in milk, because it's such a difficult  
23 matrix.

24 With regards to conjugation derivatization  
25 methods, when we approached this, we were initially

1 thinking of throughput and we wanted a direct method for  
2 throughput reasons. And so from the very beginning from  
3 method development, we were reviewing direct methods and  
4 trying to figure out a direct method that didn't require a  
5 derivatization step and the additional time and cost  
6 involved with that.

7           And so there are -- there are a lot of  
8 derivatization methods out there that are very good. But  
9 from our end, it was more kind of like you had to pick a  
10 problem with throughput on one end or the other. And so  
11 we opted for something that was maybe a little bit more  
12 tedious in terms of time, but didn't require those other  
13 resources. And so it was something of an arbitrary  
14 decision, I guess.

15           DR. GERONA: Yeah. And in addition to that, you  
16 know, I know that throughput is always -- with  
17 epidemiological studies, it's always something that drives  
18 the study that, you know, you need to be able to analyze  
19 this number of samples in this given amount of time. But  
20 for us, when we were starting it, the question that we  
21 were asking is has somebody got it correctly already?

22           And so to us what is more important is if we're  
23 going to be developing the method, we don't care how slow  
24 it is. We would rather give you the correct answer in the  
25 accurate and precise method, then -- and then worry about

1 the high throughput later.

2           There are now -- so now that we think that we  
3 actually have a method that is reliable, we can start  
4 thinking and we can start the dialogue with different  
5 laboratories to see whether are there ways by which we can  
6 actually shorten the method or maybe do it faster, if  
7 there is really a need to do that based on the  
8 interactions that we have with different groups?

9           CHAIRPERSON SCHWARZMAN: Moving on from  
10 clarifying questions, we have time now for discussion, and  
11 if you wouldn't mind staying up. It often turns into  
12 discussion with the presenters.

13           So I'll open it up to discussion from the Panel  
14 and also welcome discussion from the people in the room.

15           CHAIRPERSON SCHWARZMAN: Dr. Quintana.

16           PANEL MEMBER QUINTANA: Hi. Thank you for the  
17 presentation. And I know that your samples, especially  
18 the first ones you show, were not randomly selected or  
19 anything. But you were speaking about the study, and I  
20 was expecting more variability across the regions than I  
21 saw. And I'm wondering if you were surprised by that in  
22 general? Even the data you showed later, kind of a  
23 surprising amount in lots of people and quite similar.

24           DR. GERONA: Yeah. So -- and if I can comment  
25 about that. So there are a lot of surprises in that Detox

1 Project results. I mean, I've already pointed out one,  
2 right? Why are we detecting glyphosate in 86.1 percent of  
3 this population?

4 If the people that we're already -- we're already  
5 screening for are people who are already aware, and most  
6 likely are eating organic food, who are very much aware of  
7 their day-to-day exposure, we were thinking that we will  
8 get so much less.

9 But I guess if you look -- so the FDA started --  
10 actually, I might have it here. So these are -- these are  
11 some of the other interesting data. If we were given one  
12 hour presentation --

13 (Laughter.)

14 DR. GERONA: -- we would actually be able to  
15 present all this data.

16 So if you look at -- so there has been effort in  
17 actually looking at what could products contain -- contain  
18 glyphosate, and how much levels are you actually getting  
19 from each one of these?

20 And so -- and you -- can

21 --o0o--

22 DR. GERONA: You know, I'm presenting this to  
23 you -- so some of the product -- common products --  
24 Cheerios is 1,125.3 ppb. Ritz crackers. These brownies  
25 from Kashi, who doesn't eat that, right?

1 (Laughter.)

2 --o0o--

3 DR. GERONA: Some honey has 41 ppb glyphosate.

4 --o0o--

5 DR. GERONA: Quaker Oats, Stacy's Organic -- this  
6 is organic, Stacy's Organic Simply Naked Pita Chips has  
7 one. California -- California wines have it.

8 --o0o--

9 DR. GERONA: And all of this. I mean, who  
10 doesn't eat Oreos, Lay's? I eat Lay's all the time, and  
11 so --

12 (Laughter.)

13 --o0o--

14 DR. GERONA: So as you can see, we present you  
15 with all this data on the sources and the levels that  
16 we're measuring. I think it's easy to understand why  
17 we're getting what we're getting. That pretty much  
18 everything you eat -- almost everything you eat might have  
19 some -- the worst thing is this, there are food that we --  
20 you know, you would think -- so, for example, protein  
21 bars, for example, right? It's like before I go to the  
22 gym, I'll eat my protein bar. That's healthy. Well, what  
23 kind of protein is a protein bar?

24 Soy. And how much of the soy in the United  
25 States is GMO soy?

1 MR. ADAMS: Ninety-six percent.

2 DR. GERONA: This is why he's my graduate  
3 student.

4 (Laughter.)

5 DR. GERONA: And so I think that in itself  
6 answers the question. We looked at tap water. Tap water,  
7 as -- I don't know if I mentioned this -- we analyzed 110  
8 tap water samples from The Detox Project, only two  
9 actually have glyphosate, one from Simi Valley and one  
10 from Quincy, Washington. The levels is 0.02 ppb. And we  
11 were asking ourselves why is it that tap water won't have  
12 it?

13 Well, if you think about the treatment of water  
14 for your tap water, there is a step of phosphate --  
15 phosphate precipitation. And glyphosate has phosphate.  
16 So we do think that, you know, well it's good that our tap  
17 water does not have glyphosate, but the sad -- the  
18 corollary to that is even if your tap water doesn't have  
19 it, you still have it. Where else is this coming from,  
20 right? I think it's clear that the answer is probably the  
21 food that we eat.

22 MR. ADAMS: Yeah. The other thing I'd like to  
23 add is that in addition to these reference values, the UN  
24 FAO basically set an ADI for PMG of 60 milligrams per  
25 person per day. And they estimated the average intake for

1 this pesticide for 17 model diets. And those range from  
2 around 88 to 530 micrograms per person per day.

3 So that's a -- you know, we're reporting in parts  
4 per billion or nanograms per ml. So really it kind of  
5 makes sense, the levels we see based on, you know,  
6 residues that are already in different food products and  
7 even, you know, estimates of dietary intake.

8 DR. GERONA: Can you mention something about why  
9 -- where we're seeing it in organic food?

10 MR. ADAMS: Oh, yeah. So there was a study done  
11 by the USGS, in I believe 2014, looking at levels in about  
12 4,000 hydrologic samples. And what they found is that  
13 they had the highest detection frequency as you might  
14 expect in drainage ponds, things like that where there's  
15 high levels of agricultural runoff.

16 But something that was quite interesting was that  
17 they were detecting it relatively -- at relatively high  
18 levels in terms of detection frequency. About 71 percent  
19 of precipitation samples had detectable levels of PMG.  
20 And the reason for that is pesticide drift.

21 And so even if, you know, say there is an organic  
22 production field where they're producing organic soy for  
23 Kashi Brownies or whatever, you're still going to be  
24 exposed to pesticide drift. There's also -- there's also  
25 an issue where, as I mentioned previously this is a very

1 stable pesticide in terms of soil adsorption. It's rather  
2 unique in that aspect. The phosphate forms associations  
3 with different aspects of the basic crystal matrix of the  
4 soil and can stay there for very long periods of time. So  
5 even if it was a conventional field in the past, there may  
6 still be residues that are being incorporated.

7           There was also a study in Norway, in I believe  
8 2015 or 2016, that looked at levels in different soybeans.  
9 There was -- there was organic soybeans, conventional, and  
10 Roundup Ready. And they were still detecting residues in  
11 organic soy. So there's still, you know, exposure from  
12 pesticide drift, runoff, and also just residues in soil.

13           DR. GERONA: So -- and that is probably the  
14 reason why, you know, in these 83 urine samples, we  
15 actually also surveyed organic food consumption of these  
16 individuals. And so even though this obviously relied on  
17 whether participants are telling the truth or not, you  
18 have the always/frequently/rarely/never, there was no  
19 significant difference in terms of the levels that we're  
20 finding versus those that are frequently organic --  
21 consuming organic food and those that are not consuming  
22 organic food.

23           CHAIRPERSON SCHWARZMAN: May I ask just a quick  
24 follow-up question on that? You mentioned this study of  
25 the different soys. And there was glyphosate residues in

1 the organic soy, but was there -- were there differences  
2 in the Roundup Ready versus the --

3 MR. ADAMS: Yeah, that was the conclusion that  
4 they had as well, that in Roundup Ready, there are higher  
5 residues than conventional soy. So as I mentioned  
6 previously, you're not going to be using PMG as an  
7 herbicide in non-Roundup Ready soy, because you'll kill  
8 the soy, but it's still used as a desiccation agent, and a  
9 field pre-treatment agent. So there are still residues  
10 that are being -- there are still going to be residues on  
11 the crop itself.

12 CHAIRPERSON SCHWARZMAN: But there was a  
13 difference observed?

14 MR. ADAMS: Yes. Yep.

15 CHAIRPERSON SCHWARZMAN: Other comments or  
16 questions?

17 DR. PARK: DTSC, June-Soo Park. It was a great  
18 presentation. Thank you.

19 This might be a very naive question too. First,  
20 in the -- it's quite an acidic compound. I don't know  
21 what was the motivation you developed, the method to  
22 analyze breast milk? The urine sound good matrix for this  
23 kind of compound, Mr. Adams, as an alternative?

24 But you struggled and spend a lot of time and  
25 effort to develop this method due to the complication of

1 matrix I believe. So that was the first question. But  
2 I -- my main question was the -- also you developed the  
3 method. You know, that you have -- you sound like you  
4 have a very robust one.

5 I wonder, you know, the non-persistent chemicals  
6 like glyphosate, I don't know if this -- have you tried  
7 some -- tried to measure some depuration effect or anybody  
8 done it? Like you kind of have a different period of, you  
9 know, the breast feeding and any decreasing over the time.  
10 So that was kind of main -- my major question.

11 DR. GERONA: So let me address the motivation  
12 first, why we developed a method for breast milk. When we  
13 were approached, there are two questions that our  
14 collaborators are asking. At the time that they  
15 approached us, there has been a lot of claims that because  
16 glyphosate is polar, that you will not find it in a matrix  
17 that is quite hydrophobic. And as you can -- as you know,  
18 breast milk has a lot of nonpolar compounds.

19 The second concern is when you talk about  
20 compounds that could potentially be an endocrine disruptor  
21 or toxic, the concern always is which is the most  
22 vulnerable part of the population, right? The vulnerable  
23 part of the population, of course, are infants. And then  
24 we were asking also the question, well, but if the  
25 infant -- so if the source is food, infants are not

1 necessarily eating all this -- eating all this solid food,  
2 so the main food for the baby is either formula or breast  
3 milk.

4           And so what we wanted to -- the reason why we  
5 started developing the method actually is to ask that  
6 question, whether the exposure of the mom, which we were  
7 predicting to be a high detection frequency. This  
8 translates to an exposure of the infant.

9           And so as you can see in the slide that I  
10 presented, that particular study is still ongoing. And  
11 ultimately, that's really the study that we wanted to  
12 publish -- to actually say that in these matched samples  
13 from different biological matrices between the mother and  
14 the infant that we may or may not be seeing glyphosate.

15           MR. ADAMS: Yeah. The other aspect of that is  
16 that there was a unpublished study that was performed by  
17 an advocacy group, Moms Across America, that used  
18 immunoassay to detect PMG in breast milk. And they found,  
19 I think it was, 7 out of 11 samples. Immunoassay is not  
20 going to work very well for an analyte like this, because  
21 it's small. It's N-(Phosphonomethyl)glycine, so it's very  
22 similar to amino acids. It is a phosphate group.  
23 Everything in your body has a phosphate group.

24           And so we wanted a mass spec method that could  
25 basically double check those kind of studies. The other

1 thing is it is very polar, and you wouldn't expect that to  
2 necessarily partition into milk. But there were a couple  
3 kind of biological reasons that we were wondering whether  
4 or not it was going into milk.

5           And one was whether or not milk was acting as an  
6 ion trap. So the idea that this compound might be going  
7 across into a new environment that has a different pH,  
8 changing its polarity by its protonation status and not  
9 diffusing back. So there are a couple different systems  
10 that that acts on, like the uterus can act like that, and  
11 you can have crop passage of drugs that accumulate in the  
12 amniotic fluid, because it's acting as an ion trap.

13           The other thing is that so there are all type of  
14 amino acid transporters, LAT1 and LAT2. And those have  
15 been shown to transport PMG across epithelial barriers.  
16 And those are present in mammary epithelium. So we know  
17 that there's a transporter by which it could possibly get  
18 into milk. We think that there may be this acid trap idea  
19 or acid trap mechanism happening as well. So we would --  
20 just wanted to develop a method that could be able to  
21 answer that question, if anyone wanted to ask it.

22           Further more, dairy in Wisconsin -- we're both  
23 from Wisconsin, so, you know, dairy --

24           (Laughter.)

25           MR. ADAMS: -- dairy is big, and we wanted to

1 develop a method that could be also applied in the dairy  
2 industry for quality assurance for, you know, milk  
3 samples.

4 DR. GERONA: We also have happy cows in  
5 Wisconsin, not only in California.

6 (Laughter.)

7 CHAIRPERSON SCHWARZMAN: Do you have a follow-up  
8 question?

9 DR. PARK: Yeah, just one more thing. I'm from  
10 Wisconsin too.

11 (Laughter.)

12 DR. GERONA: Oh, so maybe we should collaborate.

13 (Laughter.)

14 CHAIRPERSON SCHWARZMAN: Other questions or  
15 discussion points, comments?

16 Okay. One more.

17 PANEL MEMBER LUDERER: I do have one more quick  
18 one. In your comparison in the Detox Project graph, I  
19 noticed that the European study levels were really low.

20 DR. GERONA: 0.17.

21 PANEL MEMBER LUDERER: Yeah, and is that using  
22 the same method?

23 DR. GERONA: No.

24 PANEL MEMBER LUDERER: No. That was my question.

25 DR. GERONA: So -- so -- yeah, yeah, yeah.

1           PANEL MEMBER LUDERER: How comparable are the  
2 different studies?

3           DR. GERONA: So that's actually -- so when you  
4 compare -- especially when you compare between studies.  
5 The first thing that an analytical chemist could ask is  
6 what's the method, right? And what's the LOQ of the  
7 method?

8           Because I think what gets lost in the translation  
9 of a lot or interpretation of a lot of the studies, I mean  
10 there have been plenty of studies that have looked at  
11 biological samples. And the conclusion of the study is  
12 glyphosate is not detectable in serum. That glyphosate is  
13 not detectable in urine. Glyphosate is not detectable  
14 breast milk.

15           Well, the general public might be fooled to  
16 actually accept that. But if you're talking to an  
17 analytical chemist, the first question that an analytical  
18 chemist would ask is what's your limit of quantification?

19           Because a lot of the studies that are being  
20 published where the conclusion is that if you look at the  
21 limit of quantification, it's 20 nanogram per ml or 50  
22 nanogram per ml. Of course, you know, you will not detect  
23 glyphosate, because the levels, as we looked at in urine,  
24 we -- the levels that we're seeing are below that.

25           Now, so the European study used -- so the report

1 was, and we cannot verify it. The report was GC-MS/MS  
2 with an LOQ of 0.15 nanogram per ml. You are  
3 asking about -- you're asking about the difference, right?  
4 There -- so if you actually contextualize that result, and  
5 look at the values based on which country, because there  
6 are several countries, so that's -- I think, if I'm not  
7 mistaken, that's about 183 individuals from 18 different  
8 countries, right?

9           There is a correlation on how lax the regulation  
10 is for using GMO products containing glyphosate, and the  
11 levels that you're seeing. The more lax the country is,  
12 the higher is the level.

13           Well, we are very strict here in the United  
14 States, right? And so that's -- that's sarcasm.

15           (Laughter.)

16           DR. GERONA: And that's probably the reason why,  
17 you know, you're seeing a higher level here in the United  
18 States. That's potentially the reason why, you know,  
19 you're asking about what is the difference, right?

20           The levels that -- the geometric mean is about  
21 0.70 nanogram per ml, but I've also seen other higher  
22 levels depending on the country that sent us the sample.

23           MR. ADAMS: And if I can add on, so with the IARC  
24 ruling that ruled it is a probable carcinogen. That  
25 meta-analysis looking at PMG exposure and non-Hodgkin's

1 lymphoma -- I re-did the meta-analysis a couple years ago.  
2 And the issue with that meta-analysis is that there are  
3 relatively few studies but the distribution of the studies  
4 is -- you know, there's heavy weighting of studies in  
5 Sweden, and France, than there is Canada, and then there  
6 are two -- two or three studies from the United States.

7           The United States studies are pre-1995. The  
8 importance of that is that's the emergence of PMG-tolerant  
9 crops. And so it doesn't really capture modern exposure  
10 levels in the United States, and its heavily weighted  
11 towards countries that have more stringent regulation with  
12 regard to pesticide application.

13           The other problem with that meta-analysis and  
14 that collection of studies is most of it's questionnaire  
15 based. Well, we know that farmers are pretty good at  
16 reporting questionnaires and exposures, but there's really  
17 no -- I guess no double checking in terms of actual levels  
18 of the analyte in their system.

19           And so, yeah, I guess that was another main --  
20 major impetus for us was to study this very difficult  
21 analyte, and see what we can learn from developing methods  
22 about it for application in the other small molecules we  
23 look at. And then also develop, you know, methods and  
24 tools for people to kind of tease apart some of these  
25 issues in the epidemiology regarding this particular

1 compound.

2 CHAIRPERSON SCHWARZMAN: Great. Well, thank you  
3 both very much for your interesting work and for coming  
4 here to talk with us about it.

5 (Applause.)

6 CHAIRPERSON SCHWARZMAN: Okay. So I think we're  
7 actually -- we have the go ahead to continue without a  
8 break, which we were -- the reason that we're trying to  
9 move this discussion up is just to make sure that we have  
10 a chance to get a vote among the Panel members on our next  
11 topic before some have to leave early.

12 So we're going to move right into the next agenda  
13 item, which is the potential designated chemicals:  
14 organophosphorus pesticides. And I want to introduce Dr.  
15 Shoba Iyer who is a staff toxicologist in the Safer  
16 Alternatives Assessment and Biomonitoring Section at  
17 OEHHA. And she will be present a brief summary of  
18 information on the class organophosphorus pesticides as  
19 potential designated chemicals.

20 (Thereupon an overhead presentation was  
21 presented as follows.)

22 DR. IYER: Thank you. Is this on?

23 Testing.

24 Sounds okay?

25 All right. So the purpose of this agenda item is

1 for the Panel to consider the class of organophosphorus  
2 pesticides as potential designated chemicals.

3 --o0o--

4 DR. IYER: As a brief reminder, designated  
5 chemicals can be considered for biomonitoring by the  
6 Program. Chemicals are designated based on inclusion in  
7 CDC's National Reports on Human Exposure to Environmental  
8 Chemicals program and on recommendations by the Scientific  
9 Guidance Panel, or SGP, for Biomonitoring California.

10 --o0o--

11 DR. IYER: Some organophosphate insecticides are  
12 already on the list of designated chemicals based on their  
13 inclusion in CDC's National Reports on Human Exposure to  
14 Environmental Chemicals program. Some examples of already  
15 designated organophosphate insecticides that are still in  
16 use in California are acephate, chlorpyrifos, diazinon,  
17 dimethoate, malathion, and naled.

18 --o0o--

19 DR. IYER: The class we are discussing today is  
20 organophosphorus pesticides for which we are using a  
21 structure-based definition, that is phosphorus-containing  
22 organic compounds used as pesticides. The class of  
23 organophosphorus pesticides does include all of the  
24 organophosphate insecticides already on the designated  
25 list, as well as other subclasses, such as

1 organophosphinates and organophosphonates.

2           The class also includes any organophosphorus  
3 pesticide currently in use or that may be introduced in  
4 the future. Shifts in pesticides being used are likely to  
5 occur. For example, the emergence of herbicide-resistant  
6 weeds is a factor that can affect pesticide selection.

7   --o0o--

8           DR. IYER: With regard to past SGP actions on  
9 this class of pesticides, in March 2009, the Panel  
10 recommended that all already designated organophosphate  
11 insecticides be added to the list of priority chemicals.  
12 And last July, the SGP reviewed our preliminary screening  
13 of three pesticide classes. The Panel recommended that  
14 OEHHA prepare a potential designated chemical document on  
15 organophosphorus pesticides. We followed this  
16 recommendation, and the document we prepared was posted  
17 two weeks ago on the SGP meeting page.

18   --o0o--

19           DR. IYER: Here is a list of the criteria for  
20 recommending designated chemicals, which also applies for  
21 classes of designated chemicals. The criteria are:  
22 Exposure or potential exposure, known or suspected health  
23 effects, the need to assess the efficacy of public health  
24 actions, availability of a biomonitoring analytical  
25 method, availability of adequate biospecimen samples, and

1 incremental analytical cost.

2           And please note that these criteria are not  
3 joined by the term "and".

4   --o0o--

5           DR. IYER: We selected the seven organophosphorus  
6 pesticides listed on this slide to highlight in our  
7 designated chemical document. These are all in current  
8 use in California. In my talk today, I'll be giving an  
9 overview of information on these seven pesticides relevant  
10 to the criteria for designated chemicals.

11   --o0o--

12           DR. IYER: On this slide and the next slide, I  
13 have the chemical structures of the seven highlighted  
14 organophosphorus pesticides. Here are the structures of  
15 bensulide, ethephon, ethoprop, and fosetyl-aluminum.

16   --o0o--

17           DR. IYER: And here are the structures of  
18 glufosinate-ammonium, glyphosate and tribufos. And I want  
19 to show you just for reference the structure of an  
20 organophosphate pesticide, tetrachlorvinphos, which is  
21 already a designated chemical. The organophosphate  
22 component of the structure is outlined in the blue box.

23   --o0o--

24           DR. IYER: This table shows you the ranks of the  
25 highlighted pesticides that were in the top 100 used

1 agriculturally in California in 2015. These ranks were  
2 obtained from the California Department of Pesticide  
3 Regulation, or DPR, and they are based on the number of  
4 pesticide pounds applied for agricultural use. Ethoprop  
5 and tribufos did not rank in the top 100. And among  
6 organophosphorus pesticides only, glyphosate is the most  
7 highly used agriculturally in the State.

8 --o0o--

9 DR. IYER: On this slide, time trends for six of  
10 the highlighted organophosphorus pesticides are displayed.  
11 I'll show glyphosate on the next slide. The graph here on  
12 the left plots agricultural use based on data from DPR.  
13 The time trend for glufosinate-ammonium here is  
14 particularly interesting. There were supply chain issues  
15 that resulted in significantly decreased agricultural use  
16 of this herbicide a few years ago with use increasing  
17 again in 2015.

18 And the graph on the right shows you the time  
19 trend of pesticide pounds sold for these same highlighted  
20 pesticides. The information in the graph on the right is  
21 compiled by DPR based on self-reported data from pesticide  
22 registrants, pest control dealers, and pesticide brokers.

23 Note that these data include pounds sold for any  
24 use in California, including agricultural, institutional  
25 and home use. So comparing the time trends for

1 agricultural pesticide use with those for pesticides sold  
2 can provide a rough idea of the potential for  
3 non-agricultural use. Of note here, are the apparent  
4 increases in pounds of the glufosinate-ammonium and  
5 fosetyl-aluminum sold in 2015.

6 --o0o--

7 DR. IYER: On this slide, I'm showing you the  
8 same type of time trend information for glyphosate. The  
9 isopropylamine and potassium salts of glyphosate are the  
10 primary forms used. The trends for each of these salts  
11 are shown in red and blue in the graphs, respectively.  
12 And the black triangles in each of these graphs show the  
13 sum of all forms of glyphosate used or sold, including the  
14 two primary salts as well as other salts.

15 The trend in the sales graph on the right  
16 indicates the magnitude of glyphosate use in  
17 non-agricultural applications in California.

18 --o0o--

19 DR. IYER: We searched for information on retail  
20 availability of products containing the highlighted  
21 organophosphorus pesticides to determine the potential for  
22 home use. We located herbicide products containing  
23 glufosinate-ammonium and glyphosate for gardens and lawns;  
24 a turf herbicide with bensulide; a plant growth regulator  
25 containing ethephon, and a fosetyl-aluminum fungicide for

1 plants and lawns. Some of these products are widely  
2 available and home use could represent a potentially  
3 significant exposure pathway.

4 And ethoprop and tribufos are restricted  
5 pesticides and are generally not available for purchase or  
6 use by the general public.

7 --o0o--

8 DR. IYER: This slide summarizes detections of  
9 the highlighted organophosphorus pesticides in  
10 environmental samples collected in California. DPR's  
11 report on 2015 data collected as part of the Air  
12 Monitoring Network noted one detection of bensulide that  
13 year. The detection was in a Salinas air sample, but was  
14 below the limit of quantitation of 9.3 nanograms per  
15 meters cubed. The air samples collected that year were  
16 also tested for tribufos, but that pesticide was not  
17 detected.

18 We located two detections of glyphosate from tap  
19 water testing in the state from 2004 to 2009: one in the  
20 Imperial Irrigation District in 2005 at 16.5 micrograms  
21 per liter, and another in Bakersfield in 2006 at 32  
22 micrograms per liter.

23 In reviewing groundwater data from the State  
24 Water Resources Control Board, we identified two  
25 detections of glyphosate in the last 10 years: one in Los

1 Angeles County in 2009 at 1.3 micrograms per liter, and  
2 another in Santa Barbara County in 2013 at 20 micrograms  
3 per liter.

4 We did not locate any California studies of the  
5 highlighted pesticides in dust. For more details on these  
6 studies in the slides, as well as for details from some  
7 non-California studies, you can refer to the document we  
8 posted online.

9 --o0o--

10 DR. IYER: Potential toxicity concerns for the  
11 highlighted organophosphorus pesticides include  
12 neurotoxicity, carcinogenicity, developmental effects,  
13 endocrine effects, and respiratory effects. These effects  
14 are consistent with those associated with organophosphate  
15 pesticide exposure. We covered known or suspected health  
16 effects of the highlighted pesticides in more detail in  
17 our document, so you can find additional relevant details  
18 there.

19 --o0o--

20 DR. IYER: This table summarizes the  
21 organophosphorus pesticides listed under Proposition 65  
22 and classifications by the International Agency for  
23 Research on Cancer, or IARC. The highlighted pesticides  
24 are in the top rows of this table, and the pesticides  
25 already on the designated list are in these bottom rows.

1 All but one of the pesticides in this table have  
2 been identified as carcinogens. And oxydemeton-methyl is  
3 listed as a reproductive toxicant under Proposition 65.

4 --o0o--

5 DR. IYER: The next few slides will cover  
6 information we located on the potential to biomonitor the  
7 highlighted organophosphorus pesticides.

8 --o0o--

9 DR. IYER: I'll start by showing you the  
10 structures of metabolites and breakdown products for  
11 selected pesticides. The major human metabolite and  
12 breakdown product of glufosinate-ammonium is  
13 3-methylphosphinopropionic acid, or 3-MPPA.

14 For glyphosate, aminomethylphosphonic acid, or  
15 AMPA, is the primary metabolite and also a breakdown  
16 product.

17 --o0o--

18 DR. IYER: The major human and rat metabolite of  
19 ethoprop is O-ethyl-S-propyl phosphorothioate. And  
20 bensulide oxon is a rat metabolite, and also a breakdown  
21 product of bensulide.

22 --o0o--

23 DR. IYER: This table summarizes serum and urine  
24 detections of the highlighted organophosphorus pesticides  
25 in past biomonitoring studies for the parent compounds and

1 some metabolites. The serum detections of bensulide,  
2 ethoprop, and tribufos were reported in a doctoral thesis  
3 we located. Ethoprop's metabolite, O-ethyl-S-propyl  
4 phosphorothioate, was detected in a different study in  
5 urine. Glufosinate-ammonium and its metabolite 3-MPPA  
6 have been detected in serum. Glyphosate has been detected  
7 in serum and in urine, and its metabolite AMPA has been  
8 detected in urine.

9 One study looked for AMPA in serum, but did not  
10 detect it. And, of course, Dr. Gerona and Axel Adams just  
11 presented their interesting findings from their recent  
12 work on glyphosate.

13 Back here.

14 The designated chemical document provides details  
15 on other studies summarized here, including detection  
16 frequencies and ranges of levels measured. We did not  
17 locate any California biomonitoring publications.

18 --o0o--

19 DR. IYER: In considering the potential for  
20 bioaccumulation, we obtained log octanol-water partition  
21 coefficients, or log KOWs, based on experimental data that  
22 were tabulated by US EPA. OEHHA has identified a log KOW  
23 of greater than or equal to 4 as indicating potential  
24 concern for bioaccumulation. Bensulide and tribufos are  
25 the only highlighted pesticides that have log KOWs greater

1 than 4.

2 With regard to potential for bioaccumulation in  
3 fish, US EPA concluded that this was not a concern for  
4 bensulide. US EPA determined that tribufos could  
5 accumulate in exposed aquatic or terrestrial organisms,  
6 but also noted that metabolism and rapid elimination are  
7 expected to mitigate the bioaccumulation concern.

8 --o0o--

9 DR. IYER: In evaluating environmental  
10 persistence, we focused mostly on soil half-lives. Each  
11 bullet in this column of soil half-lives for the  
12 highlighted pesticides shows results from one study. We  
13 also looked up some persistence information from US EPA's  
14 ecological risk assessments.

15 US EPA's assessment of bensulide predicted it to  
16 be extremely persistent in terrestrial ecosystems. US EPA  
17 concluded, based on studies across various media, that  
18 ethoprop is moderately to strongly persistent in the  
19 environment, but they also noted that there is significant  
20 uncertainty in the estimates of ethoprop's environmental  
21 half-lives due to observed variability in persistence  
22 measured at different sites, and the potential impact of  
23 previous use history at the sites.

24 Glufosinate-ammonium degrades moderately rapidly  
25 in soil to 3-MPPA. US EPA concluded that glufosinate

1 residues may persist in aquatic environments. And they  
2 also reported that glufosinate is expected to remain  
3 primarily in the water column rather than in sediment with  
4 concentrations decreasing over time via dilution and  
5 metabolic degradation.

6 US EPA reported that the persistence of  
7 glyphosate in soil appears to correlate with climate. A  
8 publication we located found that glyphosate persists  
9 longer in soil under cool and dry conditions, and that  
10 same study also found that AMPA persisted in soil longer  
11 than glyphosate did, even under warm and moist conditions.

12 And US EPA noted that tribufos appears to be more  
13 persistent than is typical for most chemicals in this  
14 class, which is illustrated by the very long soil  
15 half-life of 745 days reported in one study.

16 --o0o--

17 DR. IYER: Moving along to analytical  
18 considerations; the California Department of Public  
19 Health's Environmental Health Laboratory currently  
20 measures two specific urinary metabolites of the  
21 organophosphate pesticides chlorpyrifos and diazinon, and  
22 they also measure four non-specific dialkyl phosphate  
23 metabolites.

24 Additional method development would be required  
25 for Biomonitoring California to measure any of the

1 highlighted organophosphorus pesticides. But as you heard  
2 earlier, there are methods available for a number of  
3 these.

4 --o0o--

5 DR. IYER: One of the criteria for recommending  
6 designated chemicals that I showed you in an earlier slide  
7 is assessing the efficacy of public health actions to  
8 reduce exposure. For the class of organophosphorus  
9 pesticides, adding them to the designated chemicals list  
10 would allow the Program to select any member of the class  
11 to be included in a future study, to have the flexibility  
12 to choose analytes appropriate to the particular scenario  
13 of interest, and to track the levels of exposure and how  
14 they change over time and by region. The results of  
15 biomonitoring studies can inform ongoing State efforts to  
16 reduce pesticide exposures of concern.

17 --o0o--

18 DR. IYER: The slide shows the Panel's options  
19 for this agenda item. The Panel could recommend adding  
20 organophosphorus pesticides as a class to the list of  
21 designated chemicals; the Panel could defer, pending more  
22 information; or the Panel could recommend against adding  
23 organophosphorus pesticides as a class to the list of  
24 designated chemicals.

25 This concludes my talk, and I'm happy to take any

1 Panel questions.

2 CHAIRPERSON SCHWARZMAN: Yeah. We have a  
3 question.

4 Dr. Cranor.

5 PANEL MEMBER CRANOR: Let's see if I can turn  
6 this -- oh, is this on?

7 You mentioned that you have in here that some of  
8 the organophosphates linger in the soil much longer. I  
9 didn't see that apparently they can also be stored in  
10 human fat for a period of time, is that correct?

11 DR. IYER: I didn't look into that aspect of the  
12 pesticides. We were looking primarily at potential for  
13 biomonitoring, so the biomonitoring studies are what I  
14 focused on, aside from soil.

15 PANEL MEMBER CRANOR: I guess, was -- I'd looked  
16 at a toxicology book in looking at risk, and if they're  
17 stored and there is other exposures, they can interact,  
18 they can -- they're add -- they can be additive. And so  
19 it does seem important to know what's going on out there,  
20 which the Biomonitoring Program would do, given these --  
21 the variety of facts about it.

22 DR. IYER: Yeah. My best guess would be that  
23 some of the information we located on log KOW might give  
24 you the best prediction of what might stick around in the  
25 fat. So bensulide and tribufos were the two that we've

1 found had log KOWs greater than the cutoff of concern.

2 PANEL MEMBER CRANOR: I just saw a couple of them  
3 had long -- long half-lives.

4 DR. IYER: Yeah. Yeah. The others, aside from  
5 bensulide and tribufos, had very low log KOWs.

6 CHAIRPERSON SCHWARZMAN: Do you have a question?

7 PANEL MEMBER LUDERER: Well, yeah, I guess --  
8 just kind of related to that too, I mean, looking at the  
9 data that we were just presented in the last presentation,  
10 I think it was from the agricultural study where they had  
11 the day of the application of the glyphosate, and then  
12 subsequent days. So that one doesn't, you know, seem to  
13 go down relatively quickly, at least in the applicators.  
14 But yeah, I agree that the log KOWs that are elevated for  
15 those do lead to the one to be concerned about  
16 accumulation in humans, as well as other organisms.

17 I just -- I guess -- are we supposed to start  
18 talking about whether we want to designate yet or are we  
19 still on questions?

20 CHAIRPERSON SCHWARZMAN: No, this is clarifying  
21 questions at this time.

22 (Laughter.)

23 CHAIRPERSON SCHWARZMAN: Any other clarifying  
24 questions?

25 We have time now for public comment, and then

1 we'll have our conversation.

2           So are there any public comments from the room?

3           There will be one.

4           Is there anything on the web that we should be  
5 aware of?

6           MS. DUNN: There are none.

7           CHAIRPERSON SCHWARZMAN: Nothing. So we'll have  
8 one public comment.

9           Dr. Bradman.

10          DR. BRADMAN: Hi. I just want to comment a  
11 little bit on this issue, and also perhaps related to  
12 Roy's presentation too. I think one thing we know with  
13 pesticides that when they're used, we're exposed to them.  
14 I mean, in our studies, we found materials that had been  
15 used in the field that historically never -- no one would  
16 even think they would get into people's homes or get into  
17 people's bodies outside the fields, but they do. Whether  
18 those are of concern or not is, of course, another issue,  
19 when we look at risk and health effects.

20          But I think understanding exposures to these  
21 materials is really important. And like consumer  
22 products, like air pollutants, these are things that we're  
23 all exposed to. So I would encourage real -- the  
24 opportunity to really carefully look, particularly at the  
25 cholinesterase-inhibiting compounds, and there's several

1 on here that are included. And then other heavy-use  
2 compounds. I think it's really important that they be  
3 looked at and at least the beginning stages of  
4 understanding of what the exposures are.

5 CHAIRPERSON SCHWARZMAN: Do we have any other  
6 public comments?

7 One more.

8 MS. JUNFISH: Hi. My name is Susan JunFish. And  
9 I'd like to express support for also monitoring  
10 organophosphates, particularly the ones that are used by  
11 the mosquito vector control districts throughout the  
12 State, such as malathion is listed on that list. And I  
13 believe another very commonly used organophosphate. I  
14 think it's of serious concern what's happening when there  
15 is evidence from a very important study that was done in I  
16 think it was 2006 at Harvard School of Public Health  
17 showing an efficacy of ULV spraying for West Nile Virus.  
18 They showed that reproductive rates in communities that  
19 did ULV spraying versus communities that did not spray,  
20 there was difference in the reproductive rate -- the  
21 number of eggs being laid by West Nile Virus-carrying  
22 mosquito species.

23 DR. IYER: I'll just add a --

24 CHAIRPERSON SCHWARZMAN: Yeah, I was going to say  
25 about the presence of malathion on the list.

1           Go ahead.

2           DR. IYER: Yeah. Yeah, just to clarify,  
3 malathion was included on the slide as an example of a  
4 chemical that is on our designated list, and it also falls  
5 under the broader class of organophosphorus pesticides.  
6 So just to clarify.

7           CHAIRPERSON SCHWARZMAN: It is already currently  
8 on the designated chemical list?

9           DR. IYER: That is correct.

10          CHAIRPERSON SCHWARZMAN: Okay. Any other public  
11 comment before we move on to general discussion?

12          Okay. So, why don't you go first, Dr. Luderer.

13          PANEL MEMBER LUDERER: I mean, I was just looking  
14 at the criteria list. So, you know, from what we've --  
15 we've heard in the document that was given to us, as well  
16 as the earlier presentation, there's -- certainly the  
17 exposure to the public criteria I think is met by these  
18 compounds, given the wide detection of the ones that have  
19 been measured in biomonitoring specimens and the use in  
20 California in agriculture, and other uses, that there  
21 is -- I think we -- it meets the known or suspected health  
22 effects. You know, we've heard about several -- many  
23 of -- of the ones that we talked about today, how many of  
24 them are considered to be carcinogens, as well as, you  
25 know, other neurological endocrine disruption effects that

1 have been reported for many of these compounds.

2 I think there's also the need -- it meets the  
3 criteria for the need to assess the efficacy of public  
4 health actions. You know, some of these organophosphates  
5 have -- are now not permitted to be used, you know,  
6 indoors in residential settings. And so will we see a  
7 decline in levels of those -- you know, that's something  
8 that we need to be able to assess.

9 And then we've also heard that the biomonitoring  
10 analytical methods, you know, exist to be able to analyze  
11 these specimens. So I think, based on all of these  
12 things, that we certainly meet multiple criteria for  
13 designating these compounds - the organophosphorus  
14 compounds.

15 CHAIRPERSON SCHWARZMAN: Thank you for that  
16 summary of the evidence and how it meets with our  
17 criteria.

18 Before -- I guess I'll jump in with a comment  
19 that goes off of one of the things that Asa just mentioned  
20 about the document that you prepared I noticed there was  
21 the mention of the -- looking at how many cholinesterase  
22 inhibitors were applied in proximity to schools. I wonder  
23 actually if you would speak about that data for a minute,  
24 because that was a really interesting sort of subset to  
25 me, of this class.

1 Do you mind? Sorry to put you on the spot.

2 DR. IYER: No problem. I'm -- we were trying to  
3 focus on a selection of information to present in the  
4 slides. And I didn't cover that in the slides, but yes --  
5 so we took a look at the report that the California  
6 Environmental Health Tracking Program put together, where  
7 they examined the use of agricultural pesticides used near  
8 California public schools. The -- that work was done in  
9 2010.

10 And so I tried to get a feel for what -- which of  
11 our highlighted pesticides were included or were used  
12 close to schools. So of those organo -- highlighted  
13 organophosphorus pesticides, bensulide, ethephon, and  
14 glufosinate-ammonium were among the top 10 pesticides  
15 applied within a quarter mile of schools in one or more of  
16 the counties they assessed in the State. And bensulide  
17 was additionally one of the top 10 pesticides classified  
18 as a cholinesterase inhibitor used within a quarter mile  
19 of schools in the counties they looked at.

20 I'm aware that there's ongoing work that I think  
21 will give us some more information on more recent expo --  
22 potential for exposure near school sites. So that's what  
23 I'll add to that.

24 CHAIRPERSON SCHWARZMAN: Thank you.

25 I think one of the other things that was striking

1 to me, in addition to your sort of rundown of how the  
2 evidence really meets a lot of the criteria - although, of  
3 course, we aren't required to meet all of them - is the  
4 getting back to this concept of designating a class as  
5 opposed to designating individual chemicals. And I think  
6 this is a conversation that we have each time we look at a  
7 class, but it seems as relevant for this class of  
8 chemicals as for every other class that we've looked at  
9 that there is the -- there are these dramatic shifts in  
10 the market that can happen relatively quickly, based on  
11 sort of external pressures or even like that supply chain  
12 issue that you highlighted that happened where one drops  
13 out and others presumably fill the place in the  
14 short-term, and that are dictated by factors that we  
15 wouldn't predict or know about in time to like designate a  
16 chemical.

17           And particularly, I'm sort of curious about  
18 what's going to happen with the increasing public  
19 attention on glyphosate, and Roundup Ready crops, and  
20 the -- with the listing under Prop 65 what's going to  
21 happen in terms of public pressure on the manufacturer.

22           And it just, you know, glyphosate is obviously of  
23 concern because of its sheer volume of use, as we've heard  
24 about. And it's -- not persistence. That's using the  
25 wrong word, but it's presence in so many of the -- both

1 environmental monitoring and biomonitoring samples.

2           But I'm -- I'm interested not just in -- it  
3 raises the point to me not just of pointing toward  
4 glyphosate, but thinking, well, these could shift pretty  
5 quickly, and the emphasis on what's being used could shift  
6 pretty quickly. And I -- for that reason I'm really, I  
7 guess, intrigued by and supportive of the Program's  
8 continued movement toward highlighting a whole class of  
9 chemicals, and sort of enabling the Program to look at  
10 those as the market dictates.

11           Any other comments or thoughts?

12           Yeah.

13           PANEL MEMBER QUINTANA: Hi. Is this on?

14           After all that drama, I was just going to say I  
15 agree with you.

16           (Laughter.)

17           PANEL MEMBER QUINTANA: No, just to second the  
18 idea of this -- that graph that you prepared. And I want  
19 to also congratulate you on, and your -- all the staff on  
20 another really excellent and thorough report. But I think  
21 that the graph on how the changes and use over time, you  
22 know, can be very dramatic, even year to year, I think  
23 that's such an important point that supports the idea of  
24 designating a class.

25           CHAIRPERSON SCHWARZMAN: Okay. Thoughts or

1 points for discussion?

2 I'm getting the sense that the Panel is ready to  
3 make a motion. Would anybody like to make a motion?

4 PANEL MEMBER LUDERER: I move designate -- to  
5 designate the organophosphorus -- that the Panel vote to  
6 designate the organophosphorus pesticides as a class.

7 PANEL MEMBER CRANOR: I'll second.

8 CHAIRPERSON SCHWARZMAN: So individual member  
9 votes. Maybe we'll start at this end.

10 PANEL MEMBER QUINTANA: Yes.

11 PANEL MEMBER KAVANAUGH-LYNCH: Yes.

12 CHAIRPERSON SCHWARZMAN: I will vote for that  
13 also.

14 PANEL MEMBER CRANOR: Since I second, yes.

15 (Laughter.)

16 PANEL MEMBER LUDERER: Yes.

17 CHAIRPERSON SCHWARZMAN: So I think I'm supposed  
18 to say that the Panel has therefore moved to -- that the  
19 chemical class organophosphorus pesticides be included as  
20 designated chemicals as a class in the California  
21 Environmental Contaminant Biomonitoring Program. And so I  
22 think that actually concludes our -- this decision portion  
23 around this topic, and we can -- we're doing beautifully  
24 for time.

25 And we have some time designated now that's for

1 open public comment that could be about the items we've  
2 discussed today, and specifically this afternoon, or other  
3 public comments related to the Biomonitoring Program. And  
4 we have time for that now. And because I think we're a  
5 little early, I'm -- I don't -- I don't want to -- we have  
6 one. There we go.

7           And if people can continue to suggest that they'd  
8 like to make a public comment because you didn't have much  
9 heads up about this.

10           MS. BUERMEYER: Hi. Nancy Buermeyer with the  
11 Breast Cancer Prevention Partners. I didn't have a  
12 comment before you voted, but now I do, which is to say  
13 I'm really glad you voted the way you did. We, at the  
14 Breast Cancer Prevention Partners, have concern about  
15 these pesticides, because there have been some indications  
16 of connection to breast cancer, which is obviously of  
17 concern to us. And I just -- I was prompted to say  
18 something by the reference to the Health Tracking  
19 Program's report on pesticide use within a quarter mile of  
20 schools.

21           And I think that report has generated interest in  
22 changing the regulations around pesticide use and  
23 reporting through the Department of Pesticide Regulation.  
24 I actually haven't tracked that as carefully as I would  
25 like to, but that report is suggesting there's going to be

1 a change in public policy. And so it would be great if  
2 there could be a commensurate tracking of changes in  
3 exposures based on that change in policy.

4 I think the challenge, as we all know, is the  
5 funding to do that, right, because you'd want to get that  
6 baseline now, so you'd have a point of comparison, or at  
7 least be able to go back and find samples from now. And I  
8 think it just -- it raises the ongoing challenge of the  
9 ability of the Program to do all that it can and should  
10 without the resources to do that.

11 And I don't have an answer to that obviously, but  
12 it's a challenge that continues. But I -- but we also  
13 strongly support the designation of classes of chemicals,  
14 as you did here, and as you have for phthalates and some  
15 other things because of that volatility of how companies  
16 are using, changing, moving from chemical to chemical or  
17 even just tweaking a particular chemical to a different  
18 very similar chemical, and being able to try to keep up  
19 with that along the way. So the short version is thank  
20 you for voting for that.

21 CHAIRPERSON SCHWARZMAN: We have a public comment  
22 here.

23 CAL/EPA DEPUTY DIRECTOR SOLOMON: Hi. This is  
24 Gina Solomon of EPA and -- It's green. In response to the  
25 previous comment related to a pending regulatory action on

1 pesticide use around schools, the California Department of  
2 Pesticide Regulation did propose to take several actions  
3 related to pesticide use around schools and day care  
4 centers, public schools and public -- publicly licensed  
5 day care facilities in California that would ban most  
6 types of applications of all pesticides within a quarter  
7 mile, or at least any dispersive applications.

8           Then there are some exemptions that allow smaller  
9 buffer zones for less dispersive type applications. And  
10 then there's also a warning or information --  
11 informational component to the regulation.

12           The regulation underwent two public comment  
13 periods, and is expected to be finalized fairly shortly.  
14 So it would come into effect in the coming year. So if  
15 there were interest in looking at any potential changes  
16 over time, that would be something that could happen  
17 fairly soon.

18           CHAIRPERSON SCHWARZMAN: Great. Thank you.

19           Other comments?

20           DR. HOEPKER: Alex Hoepker, OEHHA. Just a  
21 clarifying question. Do the criteria of recommending  
22 designated chemicals equally apply to chemical classes? I  
23 think it does, but I just wanted to clarify that. And a  
24 second question related to that, how do we group  
25 chemicals? Obviously, this is an obvious example of

1 organophosphate as a class of pesticides, but I'm  
2 wondering if there's space to classify chemicals also by  
3 receptor endpoints relating to cancer and reproductive  
4 toxins?

5 CHAIRPERSON SCHWARZMAN: Shoba, do you want to  
6 start with that?

7 DR. IYER: Yeah. So the answer to the first part  
8 of your question, yes, the same criteria applies to both  
9 individual chemicals and chemical classes, the criteria  
10 for recommending them as a designated chemical or class.

11 And my understanding is that thus far we have  
12 defined some chemical classes as structure-based  
13 definitions or chemical function-based definitions, but at  
14 this time no toxicity endpoint-based definitions.

15 CHAIRPERSON SCHWARZMAN: Can you say anything  
16 about whether that's possible, or is there anything that's  
17 preventing the program from doing that?

18 DR. IYER: I think it gets -- it gets to be more  
19 complicated doing that. You know, known or suspected  
20 health effects is one of the multiple criteria we have for  
21 recommending a chemical or chemical class. And that  
22 information might not exist for a chemical. So this is  
23 one of the challenges around considering that, that I'm  
24 aware of.

25 CHAIRPERSON SCHWARZMAN: I know this isn't

1 exactly public comment, but I just remember something that  
2 I was going to say earlier and I forgot, but I think it's  
3 become salient. I'm thinking about, you know, on part of  
4 the -- one of the criteria for recommending a chemical as  
5 a designated chemical is the potential to monitor levels  
6 around an intervention study. And whenever I think of  
7 glyphosate, I think of the intriguing example of the City  
8 of San Francisco, and what -- the action they took under  
9 the -- their precautionary principle ordinance, in which  
10 they targeted the use of glyphosate in public areas in the  
11 city.

12           And since they weren't taking legal action on it,  
13 but they were implementing their precautionary principle  
14 ordinance about its use, what they did is gather the  
15 gardeners together who used glyphosate in the city and  
16 said where do you use it? And, you know, they came up  
17 with a whole list of areas like right of ways, and areas  
18 where there's invasive -- natural areas where there's  
19 invasive plants and airport, and median strips, and -- et  
20 cetera, et cetera, you know, public parks and things like  
21 that. And then they asked the gardeners what else could  
22 you use, like what are the alternatives?

23           And they came up with a long list of alternatives  
24 from like flaming weeds, to hand weeding, to just feeding  
25 them compost, and making the substrate strong, to putting

1 goats out. And this whole creative range of possible  
2 alternatives.

3 And they ended up -- through looking at that,  
4 they ended up, you know, identifying other alternatives  
5 for most uses. And there were a few uses for which they  
6 retained glyphosate, for example, medians of busy roads,  
7 where it's too dangerous to put workers in the median and  
8 do hand weed, and also like SFO, where they can't shut  
9 down the runways and put goats out.

10 (Laughter.)

11 CHAIRPERSON SCHWARZMAN: And -- but as a result  
12 of that intervention, they reduced glyphosate use by 90  
13 percent in the city. And that was a completely  
14 non-regulatory intervention. And it just -- I mention it  
15 here, because I think it's an intriguing model that  
16 imagine if Biomonitoring California could have gotten in  
17 on that, and looked at before and after and what we could  
18 learn by that kind of intervention.

19 So just as long as we're thinking of  
20 organophosphorus pesticides, it's a story that sticks with  
21 me, because of its creativity and its non-regulatory  
22 approach and how much we might be able to learn from  
23 looking at similar situations. So that's my addition.  
24 Does anyone else have any comments before we close the  
25 meeting?

1           And nothing from the web that we need to be aware  
2 of?

3           MS. DUNN: Nothing from the web.

4           CHAIRPERSON SCHWARZMAN: Sorry, one more in the  
5 room.

6           MS. JUNFISH: I appreciate your comment about San  
7 Francisco. I think they're a great role model for the  
8 rest of the cities in California, and the world over  
9 actually. This is not related to biomonitoring, but just  
10 as a quick addition to what you said, the County of Irvine  
11 has been testing steamers. And so hot steaming to kill  
12 weeds. And that has been a really great alternative to  
13 most emergent herbicides -- broad spectrum herbicides.

14           Susan JunFish.

15           CHAIRPERSON SCHWARZMAN: I think my favorite  
16 option in that was the flamethrower.

17           (Laughter.)

18           CHAIRPERSON SCHWARZMAN: That's what I would want  
19 to use.

20           (Laughter.)

21           CHAIRPERSON SCHWARZMAN: Okay. With that, thank  
22 you all for your contributions for today's meeting. I'm  
23 supposed to announce that the transcript of the meeting  
24 will be posted to the Biomonitoring California website  
25 when it's available. And that our next Scientific

1 Guidance Panel meeting will be on November 9th here in  
2 Richmond.

3 And with that, I think we'll adjourn the meeting.  
4 Thank you all.

5 (Thereupon the California Environmental  
6 Contaminant Biomonitoring Program, Scientific  
7 Guidance Panel meeting adjourned at 3:27 p.m.)  
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## 1 C E R T I F I C A T E O F R E P O R T E R

2 I, JAMES F. PETERS, a Certified Shorthand  
3 Reporter of the State of California, do hereby certify:

4 That I am a disinterested person herein; that the  
5 foregoing California Environmental Contamination  
6 Biomonitoring Program Scientific Guidance Panel meeting  
7 was reported in shorthand by me, James F. Peters, a  
8 Certified Shorthand Reporter of the State of California,  
9 and thereafter transcribed under my direction, by  
10 computer-assisted transcription.

11 I further certify that I am not of counsel or  
12 attorney for any of the parties to said meeting nor in any  
13 way interested in the outcome of said meeting.

14 IN WITNESS WHEREOF, I have hereunto set my hand  
15 this 28th day of July, 2017.

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22 JAMES F. PETERS, CSR  
23 Certified Shorthand Reporter  
24 License No. 10063  
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