

Survey 1: Registration Survey

We would like to invite you to participate in the Extension for Community Healthcare Outcomes (ECHO) Project Biosafety Community of Practice sessions facilitated by the Division of Laboratory Systems, Centers for Disease Control and Prevention.

Survey questions will take no more than 1 minute to complete. Your responses will be anonymous, and no unique identifying information will be sought or kept. The feedback we receive will be summarized in aggregate only and used for program improvement. If you encounter issues with this evaluation, please email dlbiosafety@cdc.gov

CDC estimates the average public reporting burden for this collection of information as 1 minute per response, including the time for reviewing instructions, searching existing data/information sources, gathering and maintaining the data/information needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC/ATSDR Information Collection Review Office, 1600 Clifton Road NE, SD-74, Atlanta, Georgia 30333; ATTN: PRA (0920-1050).

Please click "next" to begin the survey.

1. What type of facility do you work in?

Public Health Laboratory

Hospital (Clinical) Laboratory

Commercial/Private Laboratory

Point-of-Care Testing Site

Other

2. What is your current role?

Laboratory Director

Quality Manager

Biosafety Officer/Professional

Laboratory Technician

Other

3. How many years have you been performing in this role?

0-2

3-5

>5

4. How many personnel work within your facility?

Less than 25

25-50

More than 50

5. Do you have a dedicated personnel for biosafety?

Biosafety

Quality Management

None of the above

6. Which ECHO Biosafety session(s) are you interested in participating? Select all that apply*

Safely Implementing New Diagnostics in Platforms Commonly Used in Clinical Laboratories (August 2022)

Decontamination of Laboratory Equipment (September 2022)

PPE Use (who, when, how) (October 2022)

Risk Assessment in Clinical Laboratories (November 2022)

Laboratory Acquired Infections (December 2022)

Diagnostics in Remote Areas (January 2023)

Safety Challenges with Specimen Collection, Processing, and Storage (February 2023)

Waste Management (March 2023)

Public Health Laboratory Professional Burnout and Effect on Safety (April 2023)

Quality in Biosafety (May 2023)

Laboratory Professional Vaccine Compliance and Effect on Safety (June 2023)

7. Which session(s) are you interested in receiving PACE credit for?

	Yes	No
Safely Implementing New Diagnostics in Platforms Commonly Used in Clinical Laboratories (August 2022)	☐	☐
Decontamination of Laboratory Equipment (September 2022)	☐	☐
PPE Use (who, when, how) (October 2022)	☐	☐
Risk Assessment in Clinical Laboratories (November 2022)	☐	☐
Laboratory Acquired Infections (December 2022)	☐	☐
Diagnostics in Remote Areas (January 2023)	☐	☐
Safety Challenges with Specimen Collection, Processing, and Storage (February 2023)	☐	☐
Waste Management (March 2023)	☐	☐
Public Health Laboratory Professional Burnout and Effect on Safety (April 2023)	☐	☐
Quality in Biosafety (May 2023)	☐	☐
Laboratory Professional Vaccine Compliance and Effect on Safety (June 2023)	☐	☐

Thank you!

When you click "Done," your answers will be recorded, and you will exit this questionnaire.

Survey 2: 2022 ECHO Post-Session Survey Questions

Thank you for taking this voluntary survey to help us understand to what extent the ECHO Biosafety Community of Practice sessions facilitated by the Division of Laboratory Systems, Centers for Disease Control and Prevention have been serving the needs of the participants. The feedback you provide will also inform future session development.

Survey questions will take no more than 2 minutes to complete. Your responses will be anonymous, and no unique identifying information will be sought or kept. The feedback we receive will be summarized in aggregate only and used for program improvement. If you encounter issues with this evaluation, please email dlsbiosafety@cdc.gov

CDC estimates the average public reporting burden for this collection of information as 2 minutes per response, including the time for reviewing instructions, searching existing data/information sources, gathering and maintaining the data/information needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC/ATSDR Information Collection Review Office, 1600 Clifton Road NE, SD-74, Atlanta, Georgia 30333; ATTN: PRA (0920-1050).

Please click "next" to begin the survey.

1. To what extent do you agree with the following statements:

	Agree	Neither/Undecided	Disagree
Content was relevant to my current work	☐	☐	☐
Session addressed a gap in my understanding of the topic	☐	☐	☐
Session significantly improved my knowledge level	☐	☐	☐
Speaker was knowledgeable	☐	☐	☐
Speaker was organized and communicated clearly	☐	☐	☐
Session materials supported content	☐	☐	☐
Session learning objectives were clear	☐	☐	☐
Session had the right balance of lecture and interactivity	☐	☐	☐
Session length was appropriate	☐	☐	☐
Session pace was appropriate	☐	☐	☐
The case study presented was useful	☐	☐	☐
The session was well planned and organized	☐	☐	☐

Case study presented was useful	☐	☐	☐
Difficulty level of case study was appropriate	☐	☐	☐
Overall, I was satisfied with the session	☐	☐	☐
I am likely to recommend this session to others	☐	☐	☐

2. Will you use what you learned in this session in your work?

Definitely not

Probably not

Possibly

Probably yes

Definitely yes

Not applicable – I did not learn anything new from this session

3. What factors will keep you from using the content of this session in your work? (Select all that apply)

I need additional training in the subject matter

I will not have the resources or opportunities to use what I learned

I will not have the time to use what I learned

My supervisor or colleagues will not support me in using what I learned

The session content is not relevant to my current work

Other

4. What specifically do you plan to use from this session?

5. How did you hear about this training? (Select all that apply)

CDC DLS

APHL workgroups or committees

Coworker/Colleague

Other, please specify:

6. What part of this session was most helpful to your learning and how could this session be improved?

Thank you!

When you click "Done," your answers will be recorded, and you will exit this questionnaire.

Survey 3: CDC Biosafety ECHO Follow-up Surveys

Thank you for taking this voluntary survey to help us understand to what extent the ECHO Biosafety Community of Practice sessions facilitated by the Division of Laboratory Systems, Centers for Disease Control and Prevention have been serving the needs of the participants. The feedback you provide will also inform future session development.

Survey questions will take no more than 2 minutes to complete. Your responses will be anonymous, and no unique identifying information will be sought or kept. The feedback we receive will be summarized in aggregate only and used for program improvement. If you encounter issues with this evaluation, please email dlsbiosafety@cdc.gov

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Please click "next" to begin the survey.

1. Which ECHO Biosafety session(s) did you attend in the past six months? Select all that apply

Safely Implementing New Diagnostics in Platforms Commonly Used in Clinical Laboratories (August 2022)

Decontamination of Laboratory Equipment (September 2022)

PPE Use (who, when, how) (October 2022)

Risk Assessment in Clinical Laboratories (November 2022)

Laboratory Acquired Infections (December 2022)

No, I haven't attended any of these ECHO Biosafety sessions

1. Which ECHO Biosafety session(s) did you attend in the past six months? Select all that apply

Diagnostics in Remote Areas (January 2023)

Safety Challenges with Specimen Collection, Processing, and Storage (February 2023)

Waste Management (March 2023)

Public Health Laboratory Professional Burnout and Effect on Safety (April 2023)

Quality in Biosafety (May 2023)

Laboratory Professional Vaccine Compliance and Effect on Safety (June 2023)

No, I haven't attended any of these ECHO Biosafety sessions

2. Select all that apply for each session attended.

After participating in this session, in my personal or facility's work practices:

	I used information from this session to identify biosafety areas for improvement	I can discuss biosafety challenges I encounter	I feel empowered to communicate acquired laboratory safety knowledge	I reviewed the current processes and procedures to determine if they are up to date.
Safely Implementing New Diagnostics in Platforms Commonly Used in Clinical Laboratories	☐	☐	☐	☐
Decontamination of Laboratory	☐	☐	☐	☐
PPE Use (who, when, how)	☐	☐	☐	☐
Risk Assessment in Clinical Laboratories	☐	☐	☐	☐
Laboratory Acquired Infections	☐	☐	☐	☐

2. Select all that apply for each session attended.

	After participating in this session, in my personal or facility's work practices:			
	I used information from this session to identify biosafety areas for improvement	I can discuss biosafety challenges I encounter	I feel empowered to communicate acquired laboratory safety knowledge	I reviewed the current processes and procedures to determine if they are up to date
Diagnostics in Remote Areas (January 2023)	☐	☐	☐	☐
Safety Challenges with Specimen Collection, Processing, and Storage (February 2023)	☐	☐	☐	☐
Waste Management (March 2023)	☐	☐	☐	☐
Public Health Laboratory Professional Burnout and Effect on Safety (April 2023)	☐	☐	☐	☐
Quality in Biosafety (May 2023)	☐	☐	☐	☐
Laboratory Professional Vaccine Compliance and Effect on Safety (June 2023)	☐	☐	☐	☐

3. What factors facilitated your decision to use information learned from the session(s) you attended? (Select all that apply)

Training in the subject matter

Availability of resources or opportunities to use what I learned

Time to use what I learned

Supervisors or colleagues supported me in using what I learned

Relevance to my current work

Other (please specify)

Not applicable

4. What factors impeded your use of the content of the session(s) you attended in your work? (Select all that apply)

I need additional training in the subject matter

Lack of resources or opportunities to use what I learned

Lack of time to use what I learned

My supervisor or colleagues will not support me in using what I learned

The course content was not relevant to my current work

Other (please specify)

Not applicable

5. Would you like to stay connected with this Biosafety Community of Practice (participants and/or speakers) in the future?

Yes

No

6. What suggestions do you have to enhance members' engagement after the Biosafety Community of Practice sessions end?

Thank you!

When you click "Done," your answers will be recorded, and you will exit this questionnaire.