

Screenshots of Feedback Survey for Community Counts Data Viz Tool

View the developer webpage for the feedback survey here:

<https://wwwdev.cdc.gov/ncbddd/hemophilia/communitycounts/data-viz-feedback.html>

The picture below is a screenshot of the survey link on the CC Data Viz webpage that users will click on to take the feedback survey.

Welcome to the Community Counts Data Visualization Tool



The Community Counts data visualization tool displays de-identified data on patients with bleeding disorders who are enrolled in Community Counts in a new interactive, visual format. This tool can be used by clinicians, patients, and policymakers to learn more about the burden of bleeding disorders in the United States.

Data Visualization Tool



Access the Community Counts data visualization tool.

Having trouble accessing the tool? Try this.

[Submit feedback here](#)

[Access the tool](#)

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Community Counts Data Visualization Tool Feedback

We welcome your feedback!

1. Thank you for using the Community Counts Data Visualization Tool. What is your primary role? (optional)

- ☐ Healthcare provider
- ☐ Member of the media
- ☐ Scientist or researcher
- ☐ Educator
- ☐ General public
- ☐ Bleeding disorder patient
- ☐ Patient advocacy organization/chapter
- ☐ Pharmaceutical company representative
- ☐ Other - please specify below

Other Role: (optional)

2. How do you rate the tutorials? If you have not used or seen the tutorials, please select this option below. (optional)

- ☐ I have not used the tutorials
- ☐ Excellent
- ☐ Good
- ☐ Average
- ☐ Poor
- ☐ Very poor

If "Poor" or "Very Poor," please describe how the tutorials can improve: (optional)

3. Did you experience any technical issues with the Community Counts Data Visualization Tool? For example, a graph did not load or text not displaying properly. (optional)

- ☐ No
- ☐ Yes

If yes, please describe the issues: (optional)

4. How frequently do you access the Community Counts Data Visualization Tool? (optional)

- ☐ This is my first time
- ☐ Daily
- ☐ Weekly
- ☐ Monthly
- ☐ Infrequently

5. How likely are you to revisit the Community Counts Data Visualization Tool? (optional)

- ☐ Not likely at all
- ☐ Somewhat unlikely
- ☐ Not sure
- ☐ Somewhat likely
- ☐ Extremely likely

6. Please briefly explain how you are using this data. This helps us learn how and why people are using the Community Counts Data Visualization Tool. (optional)

"I'm interested in using this data for research" or "I have a bleeding disorder and I want to learn more about it."

7. What would enhance your experience with the Community Counts Data Visualization Tool? (optional)

8. If you have any additional comments, please write them below. (optional)

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