

Exhibit 209

CDC Director Admits Agency Gave False Information on
COVID-19 Vaccine Safety Monitoring

<https://yournews.com/2022/09/13/2413033/cdc-director-admits-agency-gave-false-information-on-covid-19-vaccine/>

CDC Director Admits Agency Gave False Information on COVID-19 Vaccine Safety Monitoring

Sep 13, 2022



By Zachary Stieber

The director of the [Centers for Disease Control and Prevention](#) (CDC) has acknowledged publicly for the first time that the agency gave false information about its COVID-19 vaccine safety monitoring.

Dr. [Rochelle Walensky](#), the agency's director, said in a letter [made public on Sept. 12](#) that the CDC did not analyze certain types of adverse event reports at all in 2021, despite the agency previously saying it started in February 2021.

"CDC performed PRR analysis between March 25, 2022, through July 31, 2022," Walensky said. "CDC also recently addressed a previous statement made to the Epoch Times to clarify PRR were not run between February 26, 2021, to September 30, 2021."

event reporting system, which it helps manage.

But the agency [said in June](#) that it did not perform PRRs. It also said that performing them was “outside th[e] agency’s purview.”

Confronted with the contradiction, Dr. John Su, a CDC official, [told The Epoch Times](#) in July that the agency started performing PRRs in February 2021 and “continues to do so to date.”

But just weeks later, the CDC said Su was wrong.

“CDC performed PRRs from March 25, 2022 through July 31, 2022,” a spokeswoman [told The Epoch Times](#) in August.

Walensky’s new letter, dated Sept. 2 and sent on Sept. 6 to Sen. Ron Johnson (R-Wis.), shows that Walensky is aware that her agency gave false information.



U.S. DEPARTMENT OF HEALTH AND HUMAN SERVICES

Public Health Service

Centers for Disease Control and Prevention (CDC)
Atlanta GA 30329-4027

September 2, 2022

The Honorable Ron Johnson
United States Senate
Washington, DC 20510

Dear Senator Johnson:

Thank you for your letters dated June 23 and July 25, 2022, regarding the Centers for Disease Control and Prevention’s (CDC) tracking of reports of coronavirus disease 2019 (COVID-19) vaccine adverse events.

The Vaccine Adverse Event Reporting System (VAERS) Standard Operating Procedures (SOP) for COVID-19 is a CDC planning document developed with internal and external partners, including federal entities.¹ Within the VAERS SOP disclaimer it states, the VAERS SOP was designed to be a dynamic resource that is used, revised, and implemented based on the current science of the COVID-19 pandemic and has since been updated from the version referenced in Freedom of Information Act (FOIA) Request #22-01479 and mentioned in your letters.²

The weekly data tables that were produced during the time period of February 26, 2021, to September 30, 2021, were provided to the FOIA requester and are included as an addendum to this response. The reported incident counts reflect preliminary information, details of which might not have been confirmed by a medical provider interview or medical record review.³ Revised descriptions of the weekly tables and the information they provide are also found in the updated VAERS SOP.

Regarding your question about the use of proportional reporting ratio (PRR) analysis, CDC and the Food and Drug Administration (FDA) chose to rely on Empirical Bayesian (EB) data mining—a more robust technique used to analyze disproportionate reporting—rather than PRR calculations to mitigate potential false signals. CDC performed PRR analysis between March 25, 2022, through July 31, 2022, to corroborate the results of EB data mining. Notably, results from PRR analysis were generally consistent with EB data mining, revealing no additional unexpected safety signals. CDC also recently addressed a previous statement made to the *Epoch Times* to clarify PRRs were not run between February 26, 2021, to September 30, 2021. Given the strength of the EB data mining method, CDC and FDA plan to continue relying upon EB data mining moving forward.

¹ www.cdc.gov/vaccinesafety/pdf/VAERS-v2-SOP.pdf

² www.cdc.gov/vaccinesafety/pdf/VAERS-COVID19-SOP-02-02-2022-508.pdf

³ <https://vaers.hhs.gov/data.html>

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CDC consistently performs extensive data collection and analysis to detect potential adverse

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