

Influenza (Flu) (/flu/index.htm)



Weekly U.S. Influenza Surveillance Report

Updated May 3, 2024



A Weekly Influenza Surveillance Report Prepared by the Influenza Division

Key Updates for Week 17, ending April 27, 2024

Seasonal influenza activity continues to decline in most areas of the country.

Viruses

Clinical Lab

3.9% (Trend ♣)
positive for influenza
this week

(/flu/weekly/index.htm#ClinicalLaboratories)

Public Health Lab

Influenza A(H1N1)pdm09, A(H3N2), and B viruses were all co-circulating this week.

(/flu/weekly/index.htm#PublicHealthLaboratori es)

Virus Characterization

Genetic and antigenic characterization and antiviral susceptibility are summarized in this report.

(/flu/weekly/index.htm#VirusCharacterization)

Illness

Outpatient Respiratory Illness

2.2% (Trend →)

of visits to a health care provider this week were for respiratory illness *(below baseline).*

(/flu/weekly/index.htm#ILINet)

Outpatient Respiratory Illness: Activity Map

0 moderate jurisdictions and **0** high or very high jurisdictions

(/flu/weekly/index.htm#ORIAM)

FluSurv-NET

79.8 per 100,000

cumulative hospitalization rate.

(/flu/weekly/index.htm#FluSurvNet)

NCHS Mortality

0.2% (Trend **♣**)

of deaths attributed to influenza this week.

(/flu/weekly/index.htm#NCHSMortality)

NHSN Hospitalizations

2,302 (Trend **♣**)

patients admitted to hospitals with influenza this week.

(/flu/weekly/index.htm#NHSN)

Pediatric Deaths

10

influenza-associated deaths were reported this week for a total of 158 deaths this season.

(/flu/weekly/index.htm#PedMortality)

All data are preliminary and may change as more reports are received.

Directional arrows indicate changes between the current week and the previous week. Additional information (/flu/weekly/index.htm#Trends) on the arrows can be found at the bottom of this page.

A description of the CDC influenza surveillance system, including methodology and detailed descriptions of each data component is available on the surveillance methods (/flu/weekly/overview.htm) page.

Additional information on the current and previous influenza seasons for each surveillance component are available on FluView Interactive (/flu/weekly/fluviewinteractive.htm).

Key Points

- Seasonal influenza activity continues to decline nationally and in most areas of the country.
- Nationally, percent positivity for both influenza A and B decreased compared to last week.
- Nationally, outpatient respiratory illness remained stable and is below baseline for the fourth week in a row. All 10 HHS regions are below their region-specific baselines.
- Nationally, the number of weekly flu hospital admissions has been decreasing since January.
- During Week 17, of the 157 viruses reported by public health laboratories, 101 (64.3%) were influenza A and 56 (35.7%) were influenza B. Of the 69 influenza A viruses subtyped during Week 17, 23 (33.3%) were influenza A(H1N1)pdm09 and 46 (66.7%) were A(H3N2).
- Ten influenza-associated pediatric deaths occurring during the 2023-2024 season were reported to CDC during Week 17, bringing the season total to 158 pediatric deaths.
- CDC estimates that there have been at least 34 million illnesses, 380,000 hospitalizations, and 24,000 deaths from flu so far this season.
- CDC recommends that everyone 6 months and older get an annual flu vaccine as long as flu activity continues.²
- There also are prescription flu antiviral drugs that can treat flu illness; those should be started as early as possible and are especially important for higher risk patients.³
- Flu viruses are among several viruses contributing to respiratory disease activity. CDC is providing updated, integrated information (https://www.cdc.gov/respiratory-viruses/index.html) about COVID-19, flu, and RSV activity on a weekly basis.

U.S. Virologic Surveillance

(https://www.cdc.gov/flu/weekly/overview.htm#LabSurveillance)

Nationally, the percentage of respiratory specimens testing positive for influenza in clinical laboratories decreased (change of >0.5 percentage points) compared to the previous week. Regions 1, 2, 3, 5, 6, 7, and 8 decreased in percent positivity while regions 4, 9, and 10 remained stable during Week 17 compared to Week 16. The regions with the highest percent positivity were regions 7 (5.9%), 1 (5.0%), and 5 (4.4%). Nationally, influenza A(H1N1)pdm09, A(H3N2), and B/Victoria viruses are all co-circulating. However, the distribution of circulating viruses varies by region. For regional and state level data and age group distribution, please visit FluView Interactive

(https://gis.cdc.gov/grasp/fluview/fluportaldashboard.html). Viruses known to be associated with recent receipt of live attenuated influenza vaccine (LAIV) or found upon further testing to be a vaccine virus are not included, as they are not circulating influenza viruses.

Clinical Laboratories

The results of tests performed by clinical laboratories nationwide are summarized below. Data from clinical laboratories (the percentage of specimens tested that are positive for influenza virus) are used to monitor whether influenza activity is increasing or decreasing.

	Week 17	Data Cumulative since October 1, 2023 (Week 40)
No. of specimens tested	58,844	3,084,016
No. of positive specimens (%)	2,263 (3.8%)	339,304 (11.0%)
Positive specimens by type		
Influenza A	1,323 (58.5%)	234,716 (69.2%)
Influenza B	940 (41.5%)	104,577 (30.8%)

Public Health Laboratories

The results of tests performed by public health laboratories nationwide are summarized below. Data from public health laboratories are used to monitor the proportion of circulating influenza viruses that belong to each influenza subtype/lineage.

	Week 17	Data Cumulative since October 1, 2023 (Week 40)
No. of specimens tested	915	104,214
No. of positive specimens	157	34,456
Positive specimens by type/subtype		
Influenza A	101 (64.3%)	26,333 (76.4%)
Subtyping Performed	69 (68.3%)	21,818 (82.9%)
(H1N1)pdm09	23 (33.3%)	15,115 (69.3%)
H3N2	46 (66.7%)	6,703 (30.7%)
H3N2v	0 (0.0%)	0 (0.0%)
Subtyping not performed	32 (31.7%)	4,515 (17.1%)
Influenza B	56 (35.7%)	8,123 (23.6%)
Lineage testing performed	29 (51.8%)	6,716 (82.7%)
Yamagata lineage	0 (0.0%)	0 (0.0%)
Victoria lineage	29 (100.0%)	6,716 (100.0%)
Lineage not performed	27 (48.2%)	1,407 (17.3%)

Additional virologic surveillance information for surrent and past seasons:
Additional virologic surveillance information for current and past seasons: Surveillance Methods (/flu/weekly/overview.htm#LabSurveillance) FluView Interactive: National, Regional, and State Data http://gis.cdc.gov/grasp/fluview/fluportaldashboard.html) or Age Data (https://gis.cdc.gov/grasp/fluview/flu_by_age_virus.html)
nfluenza Virus Characterization
/flu/weekly/overview.htm#VirusCharacterization)

CDC performs genetic (https://www.cdc.gov/flu/about/professionals/genetic-characterization.htm) and antigenic (https://www.cdc.gov/flu/about/professionals/antigenic.htm) characterization of U.S. viruses submitted from state and local public health laboratories according to the Right Size Roadmap submission guidance. These data are used to compare how similar the currently circulating influenza viruses are to the reference viruses representing viruses contained in the current influenza vaccines. The data are also used to monitor evolutionary changes that continually occur in influenza viruses circulating in humans. CDC also tests susceptibility of circulating influenza viruses to antiviral medications including the neuraminidase inhibitors (oseltamivir, zanamivir, and peramivir) and the PA endonuclease inhibitor baloxavir.

CDC has genetically characterized 4,012 influenza viruses collected since October 1, 2023.

	Genetic Characterization				
Virus Subtype or Lineage	Total No. of Subtype/Lineage Tested	HA Clade	Number (% of subtype/lineage tested)	HA Subclade	Number (% of subtype/lineage tested)
A/H1	1,582				
		6B.1A.5a	1,582 (100%)	2a	366 (23.1%)
				2a.1	1,216 (76.9%)
A/H3	1,293				
		3C.2a1b.2a	1,293 (100%)	2a.1b	1 (0.1%)
				2a.3a	1 (0.1%)
				2a.3a.1	1,290 (99.7%)
				2b	1 (0.1%)
B/Victoria	1,137				
		V1A	1,137 (100%)	3a.2	1,137 (100%)
B/Yamagata	0				
		Y3	0	Y3	0 (0%)

CDC antigenically characterizes (https://www.cdc.gov/flu/about/professionals/antigenic.htm) influenza viruses by hemagglutination inhibition (HI) (H1N1pdm09, H3N2, B/Victoria, and B/Yamagata viruses) or neutralization-based HINT (https://pubmed.ncbi.nlm.nih.gov/30804469/) (H3N2 viruses) using antisera that ferrets make after being infected with reference viruses representing the 2023-2024 Northern Hemisphere recommended cell or recombinant-based vaccine viruses. Antigenic differences between viruses are determined by comparing how well the antibodies made against the vaccine reference viruses recognize the circulating viruses that have been grown in cell culture. Ferret antisera are useful because antibodies raised against a particular virus can often recognize small changes in the surface proteins of other viruses. In HI assays, viruses with similar antigenic properties have antibody titer differences of less than or equal to 4-fold when compared to the reference (vaccine) virus. In HINT, viruses with similar antigenic properties have antibody neutralization titer differences of less than or equal to 8-fold. Viruses selected for antigenic characterization are a subset representing the genetic changes in the surface proteins seen in genetically characterized viruses.

Influenza A Viruses

• A (H1N1)pdm09: 325 A(H1N1)pdm09 viruses were antigenically characterized by HI, and all were well-recognized (reacting at titers that were within 4-fold of the homologous virus titer) by ferret antisera to cell-grown

A/Wisconsin/67/2022-like reference viruses representing the A(H1N1)pdm09 component for the cell- and recombinant-based influenza vaccines.

• **A (H3N2):** 306 A(H3N2) viruses were antigenically characterized by HI or HINT, and 302 (98.7%) were well-recognized (reacting at titers that were within 4-fold of the homologous virus titer in HI or reacting at titers that were less than or equal to 8-fold of the homologous virus in HINT) by ferret antisera to cell-grown A/Darwin/6/2021-like reference viruses representing the A(H3N2) component for the cell- and recombinant-based influenza vaccines.

Influenza B Viruses

- **B/Victoria:** 235 influenza B/Victoria-lineage virus were antigenically characterized by HI, and all were well-recognized (reacting at titers that were within 4-fold of the homologous virus titer) by ferret antisera to cell-grown B/Austria/1359417/2021-like reference viruses representing the B/Victoria component for the cell- and recombinant-based influenza vaccines.
- **B/Yamagata:** No influenza B/Yamagata-lineage viruses were available for antigenic characterization.

Assessment of Virus Susceptibility to Antiviral Medications

CDC assesses susceptibility of influenza viruses to the antiviral medications including the neuraminidase inhibitors (oseltamivir, zanamivir, and peramivir) and the PA endonuclease inhibitor baloxavir using next generation sequence analysis supplemented by laboratory assays. Information about antiviral susceptibility test methods can be found at U.S. Influenza Surveillance: Purpose and (https://www.cdc.gov/flu/weekly/overview.htm) Methods | CDC (https://www.cdc.gov/flu/weekly/overview.htm).

Viruses collected in the U.S. since October 1, 2023, were tested for antiviral susceptibility as follows:

Antiviral Medication			Total Viruses	A/H1	A/H3	B/Victoria
Neuraminidase Inhibitors	Oseltamivir	Viruses Tested	4,011	1,585	1,293	1,133
		Reduced Inhibition	1 (0.02%)	1 (0.06%)	0 (0.00%)	0 (0.00%)
		Highly Reduced Inhibition	2 (0.05%)	2 (0.1%)	0 (0.00%)	0 (0.00%)
	Peramivir	Viruses Tested	4,011	1,585	1,293	1,133
Zanami		Reduced Inhibition	0 (0.00%)	0 (0.00%)	0 (0.00%)	0 (0.00%)
		Highly Reduced Inhibition	2 (0.05%)	2 (0.1%)	0 (0.00%)	0 (0.00%)
	Zanamivir	Viruses Tested	4,011	1,585	1,293	1,133
		Reduced Inhibition	0 (0.00%)	0 (0.00%)	0 (0.00%)	0 (0.00%)
		Highly Reduced Inhibition	0 (0.00%)	0 (0.00%)	0 (0.00%)	0 (0.00%)
PA Cap- Dependent	Baloxavir	Viruses Tested	3,887	1,534	1,254	1,099
Endonuclease Inhibitor		Decreased Susceptibility	1 (0.03%)	0 (0.0%)	1 (0.1%)	0 (0.0%)

Two A(H1N1)pdm09 viruses had NA-H275Y amino acid substitution conferring highly reduced inhibition by oseltamivir and peramivir. One (H1N1)pdm09 virus had NA-I223V and NA-S247N amino acid substitutions and showed reduced inhibition by oseltamivir. One A(H3N2) virus had PA-I38T amino acid substitution and showed reduced susceptibility to baloxavir.

High levels of resistance to the adamantanes (amantadine and rimantadine) persist among influenza A(H1N1)pdm09 and influenza A(H3N2) viruses (the adamantanes are not effective against influenza B viruses). Therefore, use of these antivirals for treatment and prevention of influenza A virus infection is not recommended and data from adamantane resistance testing are not presented.

Outpatient Respiratory Illness Surveillance (https://www.cdc.gov/flu/weekly/overview.htm#ILINet)

The U.S. Outpatient Influenza-like Illness Surveillance Network (ILINet) monitors outpatient visits for respiratory illness referred to as influenza-like illness [ILI (fever plus cough or sore throat)], not laboratory-confirmed influenza and will therefore capture respiratory illness visits due to infection with pathogens that can present with similar symptoms, including influenza viruses, SARS-CoV-2, and RSV. It is important to evaluate syndromic surveillance data, including that from ILINet, in the context of other sources of surveillance data to obtain a more complete and accurate picture of influenza, SARS-CoV-2, and other respiratory virus activity. CDC is providing integrated information about COVID-19, influenza, and RSV activity on a website (https://www.cdc.gov/respiratory-viruses/index.html) that is updated weekly. Information about other respiratory virus activity can be found on CDC's National Respiratory and Enteric Virus Surveillance System (NREVSS) website (https://www.cdc.gov/surveillance/nrevss/index.html).

Outpatient Respiratory Illness Visits

Nationally, during Week 17, 2.2% of patient visits reported through ILINet were due to respiratory illness that included fever plus a cough or sore throat, also referred to as ILI. This has remained stable (change of \leq 0.1 percentage points) since Week 16 and is below the national baseline. The percentage of visits for ILI decreased in regions 1, 2, 7, and 8 and remained stable in regions 3, 4, 6, 9, and 10 in Week 17 compared to Week 16. All 10 regions are below their respective baselines. Multiple respiratory viruses are co-circulating, and the relative contribution of influenza virus infection to ILI varies by location.

Outpatient Respiratory Illness Visits by Age Group

About 70% of ILINet participants provide both the number of patient visits for respiratory illness and the total number of patient visits for the week broken out by age group. Data from this subset of providers are used to calculate the percentages of patient visits for respiratory illness by age group.

The percentage of visits for respiratory illness reported in ILINet remained stable in all age groups in Week 17 compared to Week 16.

Outpatient Respiratory Illness Activity Map

Data collected in ILINet are used to produce a measure of ILI activity* (/flu/weekly/overview.htm#ILINet) by state/jurisdiction and Core Based Statistical Areas (CBSA).

	Number of Jurisdictions		Number	of CBSAs
Activity Level	Week 17	Week 16	Week 17	Week 16
	(Week ending	(Week ending	(Week ending	(Week ending
	Apr. 27, 2024)	Apr. 20, 2024)	Apr. 27, 2024)	Apr. 20, 2024)

Very High	0	0	0	0
High	0	0	3	1
Moderate	0	1	4	11
Low	5	8	49	57
Minimal	49	45	645	639
Insufficient Data	1	1	228	221

*Data collected in ILINet may disproportionally represent certain populations within a jurisdiction or CBSA, and therefore, may not accurately depict the full picture of influenza activity for the entire jurisdiction or CBSA. Differences in the data presented here by CDC and independently by some health departments likely represent differing levels of data completeness with data presented by the health department likely being the more complete.
Additional information about medically attended visits for ILI for current and past seasons: Surveillance Methods (/flu/weekly/overview.htm#ILINet) FluView Interactive: National, Regional, and State Data (http://gis.cdc.gov/grasp/fluview/fluportaldashboard.html) or ILI Activity Map (https://gis.cdc.gov/grasp/fluview/main.html)
Hospitalization Surveillance (http://www.cdc.gov/flu/weekly/overview.htm#HospitalizationSurv)
(http://www.cdc.gov/flu/weekly/overview.htm#HospitalizationSurv)

FluSurv-NET

The Influenza Hospitalization Surveillance Network (FluSurv-NET) conducts population-based surveillance for laboratory-confirmed influenza-related hospitalizations in select counties in 14 states and represents approximately 9% of the U.S. population. FluSurv-NET hospitalization data are preliminary. As data are received each week, prior case counts and rates are updated accordingly.

A total of 24,385 laboratory-confirmed influenza-associated hospitalizations were reported by FluSurv-NET sites between October 1, 2023, and April 27, 2024. The weekly hospitalization rate observed in Week 17 was 0.5 per 100,000 population. The weekly hospitalization rate observed during Week 52 is tied with the 2014-2015 season for the second highest peak weekly rate observed during all seasons going back to 2010-2011, and only lower than the 2017-2018 season. The overall cumulative hospitalization rate was 79.8 per 100,000 population. This cumulative hospitalization rate is the second highest cumulative hospitalization rate when compared against previous end-of-season rates for Week 17, and it is the second highest cumulative in-season hospitalization rate observed in Week 17, following the 2017-2018 season (106.0). Cumulative in-season hospitalization rates observed in Week 17, from 2010-2011 through 2022-2023 (excluding 2017-2018) ranged from 0.8 to 69.0.

When examining rates by age, the highest cumulative hospitalization rate per 100,000 population was among adults aged 65 years and older (218.2), followed by adults aged 50-64 years (94.7) and children aged 0-4 years (82.0). When examining age-adjusted rates by race and ethnicity, the highest rate of hospitalization per 100,000 population was among non-Hispanic Black persons (145.5), followed by non-Hispanic American Indian or Alaska Native persons (109.7), Hispanic persons (75.4), non-Hispanic White persons (61.6), and non-Hispanic Asian/Pacific Islander persons (42.9).

Among 24,385 hospitalizations, 20,666 (84.7%) were associated with influenza A virus, 3,546 (14.5%) with influenza B virus, 46 (0.2%) with influenza A virus and influenza B virus co-infection, and 127 (0.5%) with influenza virus for which the type was not determined. Among those with influenza A subtype information, 3,487 (69.9%) were A(H1N1) pdm09 and 1,501 (30.1%) were A(H3N2).

Among 3,450 hospitalized adults with information on underlying medical conditions, 95.6% had at least one reported underlying medical condition, the most commonly reported were hypertension, cardiovascular disease, obesity, and metabolic disease. Among 1,509 hospitalized women of childbearing age (15-49 years) with information on pregnancy status, 22.7% were pregnant. Among 1,015 hospitalized children with information on underlying medical conditions, 68.6% had at least one reported underlying medical condition; the most commonly reported was asthma, followed by obesity and neurologic disease.

In these figures, cumulative and weekly rates for all seasons prior to the 2023-2024 season reflect end-of-season rates. For the 2023-2024 season, rates for recent hospitals admissions are subject to reporting delays. As hospitalization data are reviewed each week, prior case counts and rates are updated accordingly.
Additional FluSurv-NET hospitalization surveillance information for current and past seasons and additional age groups:
Surveillance Methods (https://www.cdc.gov/flu/weekly/overview.htm#FluSurvNet) FluView Interactive: Rates by Age, Sex, and
Race/Ethnicity (http://gis.cdc.gov/GRASP/Fluview/FluHospRates.html) or Data on Patient Characteristics
(http://gis.cdc.gov/grasp/fluview/FluHospChars.html) RESP-NET Interactive (https://www.cdc.gov/surveillance/resp-net/dashboard.html)

National Healthcare Safety Network (NHSN) Hospitalization Surveillance

Hospitals report to NHSN the weekly number of patients admitted with laboratory-confirmed influenza. During Week 17, 2,302 patients with laboratory-confirmed influenza were admitted to a hospital. The number of patients admitted to a hospital with laboratory-confirmed influenza for Week 17 decreased (change of >5%) compared to Week 16 nationally and in regions 1, 2, 3, 4, 5, 6, 7, and 8. The number of hospitalizations reported in regions 9 and 10 increased slightly compared to last week.

Effective May 1, 2024, hospitals are no longer required to report hospital admissions, hospital capacity, or hospital occupancy data to HHS through NHSN. After May 3, 2024, the NHSN flu hospitalization data will not be included in FluView/FluView Interactive for the remainder of the 2023-2024 season.

Additional NHSN Hospitalization Surveillance information:

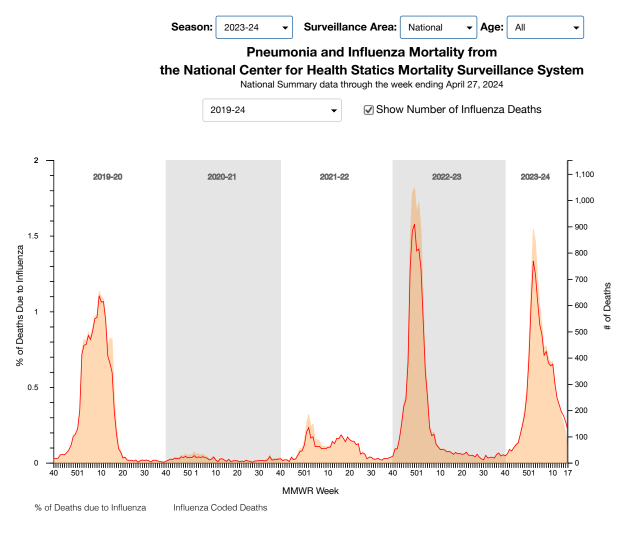
Surveillance Methods (https://www.cdc.gov/flu/weekly/overview.htm#NHSN) | Additional Data (https://data.cdc.gov/Public-Health-Surveillance/Respiratory-Virus-Response-RVR-United-States-Hospi/9t9r-e5a3) | FluView Interactive (http://gis.cdc.gov/grasp/fluview/FluView12.html)

Mortality Surveillance

(https://www.cdc.gov/flu/weekly/overview.htm#MortalitySurveillance)

National Center for Health Statistics (NCHS) Mortality Surveillance

Based on NCHS mortality surveillance data available on May 2, 2024, 0.2% of the deaths that occurred during the week ending April 27, 2024 (Week 17), were due to influenza. This percentage decreased (≥ 0.1 percentage point change) compared to Week 16. The data presented are preliminary and may change as more data are received and processed.



View Regional and State Level Data (https://gis.cdc.gov/grasp/fluview/Mortality.html) | Download Chart Data | Download PowerPoint Presentation

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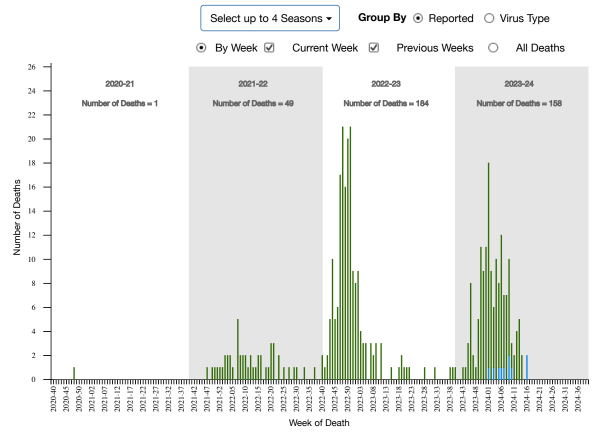
Additional pneumonia, influenza and COVID-19 mortality surveillance information for current and past seasons:

 $Surveillance\ Methods\ (https://www.cdc.gov/flu/weekly/overview.htm\#NCHSMortality)\ \mid\ FluView\ Interactive\ (https://gis.cdc.gov/grasp/fluview/mortality.html)$

Influenza-Associated Pediatric Mortality

Ten influenza-associated pediatric deaths occurring during the 2023-2024 season were reported to CDC during Week 17. The deaths occurred between weeks 1 and 10 (the weeks ending January 6, 2024, and March 9, 2024) and during Week 16 (the week ending April 20, 2024). Six deaths were associated with influenza A viruses. Four of the influenza A viruses had subtyping performed; three were A(pdm09H1N1) viruses and one was an A(H3N2) virus. Three deaths were associated with influenza B viruses with no lineage determined. Lastly, one death was associated with a co-infection of influenza A and B viruses.

Influenza-Associated Pediatric Deaths by Week of Death, 2020-21 season to 2023-24 season



View FluView Interactive (https://gis.cdc.gov/grasp/fluview/PedFluDeath.html) | Download Chart Data | Download PowerPoint Presentation

Additional pediatric mortality surveillance information for current and past seasons:

Surveillance Methods (https://www.cdc.gov/flu/weekly/overview.htm#PediatricMortality) | FluView Interactive (https://gis.cdc.gov/GRASP/Fluview/PedFluDeath.html)

Trend Indicators

Increasing: Decreasing: Stable:

Indicators Status by System

Clinical Labs: Up or down arrows indicate a change of greater than or equal to 0.5 percentage points in the percent of specimens positive for influenza compared to the previous week.

Outpatient Respiratory Illness (ILINet): Up or down arrows indicate a change of greater than 0.1 percentage points in the percent of visits due to respiratory illness (ILI) compared to the previous week.

NHSN Hospitalizations: Up or down arrows indicate change of greater than or equal to 5% of the number of patients admitted with laboratory-confirmed influenza compared to the previous week.

NCHS Mortality: Up or down arrows indicate change of greater than 0.1 percentage points of the percent of deaths due to influenza compared to the previous week.

Reference Footnotes

¹U.S. Influenza Surveillance: Purpose and Methods (2023 Oct). Centers for Disease Control and Prevention. https://www.cdc.gov/flu/weekly/overview.htm#ILINet (https://www.cdc.gov/flu/weekly/overview.htm#ILINet).

³Influenza Antiviral Medications: Summary for Clinicians (2023 Sept). Centers for Disease Control and Prevention. https://www.cdc.gov/flu/professionals/antivirals/summary-clinicians.htm (https://www.cdc.gov/flu/professionals/antivirals/summary-clinicians.htm).

Additional National and International Influenza Surveillance Information

FluView Interactive: FluView includes enhanced web-based interactive applications that can provide dynamic visuals of the influenza data collected and analyzed by CDC. These FluView Interactive applications (http://www.cdc.gov/flu/weekly/fluviewinteractive.htm) allow people to create customized, visual interpretations of influenza data, as well as make comparisons across flu seasons, regions, age groups and a variety of other demographics.

National Institute for Occupational Safety and Health: Monthly surveillance data on the prevalence of health-related workplace absenteeism among full-time workers in the United States are available from NIOSH (https://www.cdc.gov/niosh/topics/absences/default.html).

U.S. State and local influenza surveillance: Select a jurisdiction below to access the latest local influenza information.

Alabama (http://adph.org/influenza/)	Alaska (http://dhss.alaska.gov/dph/Epi/id/Pages/influenza/fluinfo
Colorado (https://www.mass.gov/info-details/influenza-reporting)	Connecticut (https://portal.ct.gov/DPH/Epidemiology-ar Emerging-Infections/Influenza-Surveillance-and-Statistics)
Georgia (https://dph.georgia.gov/flu-activity-georgia)	Hawaii (http://health.hawaii.gov/docd/resources/reports/influenz reports/)

lowa (https://idph.iowa.gov/influenza/reports)	Kansas (http://www.kdheks.gov/flu/surveillance.htm)
Maryland (https://phpa.health.maryland.gov/influenza/fluwatch/)	Massachusetts (https://www.mass.gov/infodetails/influenza-reporting)
Missouri (http://health.mo.gov/living/healthcondiseases/communicable/influenza/reports.php)	Montana (https://dphhs.mt.gov/publichealth/cdepi/diseases/Pan- Respiratory/index)
New Jersey (http://www.nj.gov/health/cd/topics/flu.shtml)	New Mexico (https://nmhealth.org/about/erd/ideb/isp/
Ohio (http://www.flu.ohio.gov)	Oklahoma (https://oklahoma.gov/health/health-education/acute-disease-service/disease-information/influenza-home-page.html)
South Carolina (http://www.scdhec.gov/Health/DiseasesandConditions/InfectiousDiseases/Flu/FluData/)	South Dakota (https://doh.sd.gov/health-data-reports/cdashboards/influenza-dashboard/)
Vermont (http://www.healthvermont.gov/immunizations-infectious-disease/influenza/flu-activity-and-surveillance)	Virginia (http://www.vdh.virginia.gov/epidemiology/influeflu-in-virginia/influenza-surveillance/)
Wyoming (https://health.wyo.gov/publichealth/infectious-disease-epidemiology-unit/disease/influenza/)	New York City (http://www1.nyc.gov/site/doh/providers/health-topics/flualerts.page)

World Health Organization:

Additional influenza surveillance information from participating WHO member nations is available through FluNet (https://www.who.int/tools/flunet) and the Global Epidemiology Reports. (https://www.who.int/teams/global-influenza-programme/surveillance-and-monitoring/influenza-surveillance-outputs)

WHO Collaborating Centers for Influenza:

Australia (http://www.influenzacentre.org/Surveillance_Samples_Received.html), China (http://www.chinaivdc.cn/cnic/), Japan (http://idsc.nih.go.jp/index.html), the United Kingdom (https://www.crick.ac.uk/research/worldwide-influenza-centre), and the United States (http://www.cdc.gov/flu/) (CDC in Atlanta, Georgia)

Europe:

The most up-to-date influenza information from Europe is available from WHO/Europe and the European Centre for Disease Prevention and Control (http://www.flunewseurope.org/).

Public Health Agency of Canada:

The most up-to-date influenza information from Canada is available in Canada's weekly FluWatch report (http://www.phac-aspc.gc.ca/fluwatch/).

Public Health England:

The most up-to-date influenza information from the United Kingdom is available from Public Health England (http://www.hpa.org.uk/Topics/InfectiousDiseases/InfectionsAZ/SeasonalInfluenza/).

Any links provided to non-Federal organizations are provided solely as a service to our users. These links do not constitute an endorsement of these organizations or their programs by CDC or the Federal Government, and none should be inferred. CDC is not responsible for the content of the individual organization web pages found at these links.

A description of the CDC influenza surveillance system, including methodology and detailed descriptions of each data component is available on the surveillance methods (http://www.cdc.gov/flu/weekly/overview.htm) page.

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Source: Centers for Disease Control and Prevention (https://www.cdc.gov/), National Center for Immunization and Respiratory Diseases (NCIRD) (https://www.cdc.gov/ncird/index.html)