**Early Evidence of the SARS-CoV-2 B.1.1.529 (Omicron) Variant in Community Wastewater — United States, November–December 2021**

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The United States designated the B.1.1.529 (Omicron) variant of SARS-CoV-2 (the virus that causes COVID-19) a variant of concern on November 30, 2021, and the first U.S. Omicron COVID-19 case was reported on December 1.† By December 18, Omicron was estimated to account for 37.9% of U.S. COVID-19 cases. Early warning systems, such as sewage (wastewater) surveillance, can help track the spread of SARS-CoV-2 variants across communities.‡

The National Wastewater Surveillance System (NWSS) comprises 43 health departments funded by CDC to provide data presence of and trends in SARS-CoV-2 infections that are independent of clinical testing. In addition to total SARS-CoV-2 testing, some health departments track SARS-CoV-2 variants by detecting variant-associated mutations in wastewater. Health departments in four states (California, Colorado, New York, and Texas) were the first wastewater surveillance programs to detect evidence of Omicron in community wastewater. This report describes the initial detections in wastewater during November 21–December 16, 2021, and the interpretative framework for these types of data. This activity was reviewed by CDC and was conducted consistent with applicable federal law and CDC policy.§

**California**

The California Department of Public Health and academic partners use mutation-specific reverse transcription–polymerase chain reaction (RT-PCR) and sequencing to track variants in wastewater collected daily from 10 sewersheds.¶¶¶¶ Omicron-associated mutations delHV69–70 (also seen with Alpha variant [B.1.1.7 and Q lineages])†† and del143–145 were detected in samples collected November 25 and November 30, 2021, from two Northern California communities (Table). Results from these samples were available on December 2; at that time, two clinical COVID-19 cases attributed to Omicron had been identified in California, but none from these communities. By December 17, del143–145 mutations were detected at all 10 sampled sewersheds in California communities.

**Colorado**

The Colorado Department of Public Health and Environment conducts biweekly SARS-CoV-2 wastewater testing at 21 sewersheds, using sequencing to track variants. Thirteen Omicron-associated mutations were detected in a sample collected on December 2, 2021. At that time, only one travel-associated Omicron case had been reported in Colorado. No Omicron-associated mutations were detected in the samples collected on December 6; however, by December 16, Omicron-associated mutations were detected at 19 of 21 sewersheds.

**New York City**

The New York City Department of Environmental Protection tracks variants in wastewater by sequencing weekly samples collected from 14 sewersheds (3). Twelve Omicron-associated mutations were detected in a sample collected on November 21. By December 4, the date the wastewater data were reported, one Omicron case had been identified in a resident of the sewershed. Samples collected on November 28 from this same sewershed and from another sewershed contained Omicron-associated mutations, as reported to the health department on December 17.

**Houston, Texas**

The Houston Health Department conducts weekly wastewater testing at 39 sewersheds in the city and uses sequencing to track SARS-CoV-2 measurements in untreated sewage can provide information on changes in total SARS-CoV-2 infection in the community contributing to that wastewater treatment plant. That area is known as the sewershed.

### Notes from the Field

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### TABLE. Detection of mutations associated with the SARS-CoV-2 B.1.1.529 (Omicron) variant in wastewater — California, Colorado, New York City, and Houston, Texas, November 21–December 16, 2021

| Location                | Sample date | Test method                                                                 | Results                                                                 |
|-------------------------|-------------|-----------------------------------------------------------------------------|-------------------------------------------------------------------------|
| California              |             |                                                                             |                                                                        |
| Seawatershed A          | Nov 25, 2021| Mutation-specific RT-PCRs targeting delHV69–70 and delL143–145*             | Both mutations detected at <1,000 genomic copies/gram wastewater solids  |
| Seawatershed B          | Nov 30, 2021| Mutation-specific RT-PCRs targeting delHV69–70 and delL143–145*             | Both mutations detected at <1,000 genomic copies/gram wastewater solids  |
|                         | Dec 2, 2021 | Mutation-specific RT-PCRs targeting delHV69–70 and delL143–145*             | Both mutations detected at <1,000 genomic copies/gram wastewater solids  |
|                         |             | Partial sequencing of S-gene using ARTIC v4 73R, 74L primers               | Detected 9 bp insertion mutation in s214EPE and 3 bp N211I deletion      |
| Seawersheds (10 sites)  | Dec 17, 2021| Mutation-specific RT-PCR targeting del143–145*                             | Mutations detected at >4,500 genomic copies/gram wastewater solids       |
| Colorado                |             |                                                                             |                                                                        |
| Seawatersheds (21 sites)| Dec 2, 2021 | SARS-CoV-2-enriched tiled amplicon sequencing                               | Detected 13 of 17 Omicron-associated mutations                           |
|                         |             | One of 21 sites                                                            |                                                                         |
|                         |             | Five of 21 sites                                                            |                                                                         |
|                         | Dec 9, 2021 | SARS-CoV-2-enriched tiled amplicon sequencing                               | Detected between four and 13 of 17 Omicron-associated mutations depending on the site |
|                         | Dec 13, 2021| SARS-CoV-2-enriched tiled amplicon sequencing                               | Detected between six and 14 of 17 Omicron-associated mutations, depending on the site |
|                         | Dec 16, 2021| SARS-CoV-2-enriched tiled amplicon sequencing                               | Detected between 12 and 14 of 17 Omicron-associated mutations, depending on the site |
| New York City           |             |                                                                             |                                                                        |
| Seawatershed A          | Nov 21, 2021| Short-read sequencing of S-gene amplicon†,§                                  | Detected 12 Omicron-associated mutations including eight mutations unique to Omicron |
| Seawatershed B          | Nov 28, 2021| Short-read sequencing of S-gene amplicon†,§                                  | Detected 12 Omicron-associated mutations including eight mutations unique to Omicron |
| Houston, Texas          |             |                                                                             |                                                                        |
| Seawatersheds (39 sites)| Nov 29, 2021| SARS-CoV-2-enriched tiled amplicon sequencing using ARTIC v3 primers*       | Detected six Omicron-associated mutations                              |
|                         |             | Seven of 39 sites                                                           |                                                                         |
|                         | Dec 6, 2021 | SARS-CoV-2-enriched tiled amplicon sequencing using ARTIC v3 primers*       | Detected 14 Omicron-associated mutations                              |
|                         | Dec 13, 2021| SARS-CoV-2-enriched tiled amplicon sequencing using ARTIC v3 primers*       | Detected 18 Omicron-associated mutations                              |
|                         |             | 35 of 39 sites                                                              |                                                                         |

Abbreviation: RT-PCR = reverse transcription–polymerase chain reaction.

* [https://www.protocols.io/view/quantification-of-sars-cov-2-variant-mutations-hv6-b2mqd6u6](https://www.protocols.io/view/quantification-of-sars-cov-2-variant-mutations-hv6-b2mqd6u6)

††† [https://www.medrxiv.org/content/10.1101/2021.09.08.21263279v1](https://www.medrxiv.org/content/10.1101/2021.09.08.21263279v1)

Sequencing detected six Omicron-associated mutations in samples collected on November 29 from seven sewersheds across the city. The first clinical detection of Omicron in the city was reported on December 1. The number of Omicron-positive sites, as well as the number of Omicron-associated mutations detected, increased over the subsequent 2 weeks.

### Discussion

The wastewater surveillance programs in these four states were the first to detect evidence of Omicron in community wastewater. Variant tracking data from wastewater cannot confirm the presence of a specific variant because the methods used cannot determine whether all variant-defining mutations are present on a single genome. However, conditions that increase confidence in the results include detection of multiple variant-associated mutations; linked mutations (i.e., on the same sequence read), or unique mutations not shared by other known variants; RNA concentration data consistent with emergence (e.g., low initial concentrations, increasing over time); the reporting of clinical cases in the area; detections in consecutive samples or via multiple methods; and RNA concentration or sequence abundance data for multiple variant-associated mutations trending together. Limitations of variant tracking in wastewater include detections inconsistent with the current epidemiology, low quality sequence data, sporadic detections, detection of a single variant-associated mutation, and conflicting trends in concentration or abundance data for...
mutations associated with the same variant. Reporting times >1 week can limit the usefulness of this data.

The detection of Omicron-associated mutations in community wastewater provides strong early evidence that the Omicron variant was likely present or more widely distributed in these communities than originally indicated by clinical testing alone; Omicron-associated mutations were documented during November 2021, at least a week before the first U.S. case identified via clinical testing on December 1. Variant tracking data from wastewater can be used as a complement to clinical testing for early detection of emerging variants, which can help guide decisions about allocation of clinical and public health resources, testing strategies, and public health messaging.

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