General Notes	
The level of measure for NWSS DCIPHER SARS-CoV-2 reporting:	Each SARS-CoV-2 PCR target ('pcr_target') that is measured should be each be reported on a separate row. Other fields that correspond to t "wide" with respect to most measurements made on a given sample,
Data types:	Fields are read in as strings; data type casting happens after file read i
Submission Requirements	
Required fields:	These fields must be included in the data file; if they are omitted, file file even if all row values are [empty].
Non-required but non-empty fields:	Some fields are "not required" but also must have non-empty values, cause them to be flagged for quality control issues once uploaded.
Value Sets	
Missing values:	Missing values should be left blank, denoted as [empty] in the Value S
Requesting additional values for categorical fields:	For categorical fields, restrict values to those listed in the Value Sets.
Case sensitivity:	Fields are not case sensitive.
Data Type Definitions	
string	ASCII-encoded characters; do not use line breaks within a value
float	Floating-point (i.e., decimal) number
integer	Integer number
ZIP code	5-digit US ZIP code (#####)
date	yyyy-mm-dd
time	hh:mm (24-hr format in the local time zone)
list	Comma-separated strings
category	Categorical variable with a defined value set of strings
NPDES permit number	National Pollutant Discharge Elimination System (NPDES) permit num
EPA registry ID number	EPA Registry ID or Facility Registry Service (FRS) identifier (<########
time zone	Time zone, represented as a UTC time offset (UTC-[hh]:[mm], e.g., UT

Field Name Groups

jurisdiction id

Reporter Collection Site WWTP Collection Method Processing Method SARSCoV2 Quantification Method Sample SARSCoV2 Quantification Results

Terminology

PCR:

Throughout this data dictionary, the term "PCR" (polymerase chain re transcriptase PCR)-based method for RNA targets, such as SARS-CoV-z quantitative (real time) PCR or digital PCR.

String 20 characters or less, containing only numbers, English alphabe

Field Name Color Key	
	Data that may be generated by the reporting jurisdiction public health
	Data likely generated by testing laboratories
	Data likely generated by wastewater utilities
	New fields or changes to existing fields

specified on a separate row in the CSV file. For example, if N1 and N2 are both measured for the same san his sample, including 'sample_id', should be repeated across those rows. In general, this format can be tho but "long" with respect to SARS-CoV-2 PCR targets.

n.

upload will be prohibited. For required fields that allow [empty] values, the field column header must be in

, conditional on the completion of another field. Omission of these fields will not interfere with upload of th

iets. Do not use other strings, such as "NA" or "[empty]", to denote missing values." To request that values be added to the vocabulary, please contact NWSS staff.

ber (<2-letter abbreviation><#######>). Search engine to locate this identifier is available here: https://ech ####>). Search engine to locate this identifier is available here: https://echo.epa.gov/facilities/facility-search C-06:00); offsets can be found at https://www.timeanddate.com/time/zone/usa tic characters, underscores, and hyphens (white space is not allowed); not case sensitive

action) is used as a shorthand for "PCR-based quantification methods", even when the method is an RT-PC 2. In addition, the term PCR does not refer to end-point PCR methods, but rather quantitative PCR methods

n agency, possibly in collaboration with either wastewater utilities or testing laboratories

nple, they should ught of as being

cluded in the data

ne data file, but will

10.epa.gov/facilities 1?mediaSelected=cwa

R (reverse s, such as

E	ield Name	<u>Data Type</u>
R	eporter eporting_jurisdiction	category
C	ollection Site	
S	ite_id	<#####-###-##-##
С	ounty_names	list (comma-separated integers)
z	ipcode	ZIP code (#####)
p	opulation_served	integer
S	ewage_travel_time	float
S	ample_location	category
S	ample_location_specify	string
ir	nstitution_type	category

WWTP		
epaid	NPDES permit number (<2-letter	
epa_registry_id	EPA Registry ID (<###########>)	
wwtp_name	string	
wwtp_jurisdiction	category	
capacity mgd	float	
capacity_inga	licat	
inductrial input	flast	
industriai_input	noat	
stormwater_input	category	
influent_equilibrated	category	

Collection Method

sample_type	category
composite_freq	float
sample_matrix	category
collection_storage_time	float
collection_storage_temp	float
pretreatment	category
pretreatment_specify	string
Processing Method solids_separation	category
concentration_method	category
extraction_method	category

pre_conc_storage_time

float

pre_conc_storage_temp	float
pre_ext_storage_time	float
pre_ext_storage_temp	float
tot_conc_vol	float
ext_blank	category
rec_eff_target_name	category
rec_eff_spike_matrix	category
rec_eff_spike_conc	float
pasteurized	category
SARSCoV2 Quantification Method	
pcr_target pcr_gene_target	category category
pcr_gene_target_ref	string
pcr_type	category

lod_ref	string
hum_frac_target_mic	category
hum_frac_target_mic_ref	string
hum_frac_target_chem	category
hum_frac_target_chem_ref	string
other_norm_name	category
other_norm_ref	string
quant_stan_type	category
stan_ref	string
inhibition_method	string
num_no_target_control	category
Sample	
sample_collect_date	date ([yyyy]-[mm]-[dd])
sample collect time	time. 24-hr ([hh]:[mm])

sample_collect_time	time, 24-hr ([hh]:[mm])
time_zone	time zone (UTC-[hh]:[mm])

flow_rate	float
ph	float
conductivity	float
tss	float
collection_water_temp	float
equiv_sewage_amt	float
sample_id	jurisdiction id (a string 20 characters or less, containing only numbers, English alphabetic characters, underscores, and hyphens; white space is not allowed; not case sensitive)
lab_id	jurisdiction id (a string 20 characters or less, containing only numbers, English alphabetic characters, underscores, and hyphens; white space is not allowed; not case sensitive)

qc_ignore	category
dashboard_ignore	category
analysis_ignore	category
SARSCoV2 Quantification Results test_result_date	date ([yyyy]-[mm]-[dd])

category

pcr_target_units

pcr_target_avg_conc	float
pcr_target_std_error	float
pcr_target_cl_95_lo	float
pcr_target_cl_95_up	float
pcr_target_below_lod	categ
lod_sewage	float

oat oat

ategory

ntc_amplify	category
rec_eff_percent	float
inhibition_detect inhibition_adjust	category category
hum_frac_mic_conc	float
hum_frac_mic_unit	category
hum_frac_chem_conc	float
hum_frac_chem_unit	category
other_norm_conc	float
other_norm_unit	category
quality_flag	category

major_lab_method	integer
major_lab_method_desc	string
	Ū

Value Set

The CDC Epidemiology and Laboratory Capacity (ELC) jurisdiction, most frequently a state, reporting these data (2-letter abbreviation)	[See Value Sets: vs_reporting_jurisdiction]
Nationally unique sampling site identifier assigned following NWSS format. <5 digit county FIPS code for WWTP address>-<3 digit facility code>-<2 digit interceptor and sewershed code>-<2 digit subsewershed code>-<2 digit site code>	
5-digit numeric FIPS codes of all counties and county equivalents served by this sampling site (i.e., served by this wastewater treatment plant or, if 'sample_location' is "upstream", then by this upstream location). Note that most jurisdictions are covered by counties, but some are covered by county equivalents, such as independent cities, parishes, or census areas.	[5-digit integers]
Zip code in which this sampling site is physically located (one 5-digit zip code)	[5-digit integers]
Estimated number of persons served by this sampling site (i.e., served by this wastewater treatment plant or, if 'sample_location' is "upstream", then by this upstream location)	[greater than or equal to 0]
What is the approximate sewage travel time, on average, from sewage source to this sampling site (i.e., this wastewater treatment plant or, if 'sample_location' is "upstream", then this upstream location)? This should be specified as a duration in hours, not a time of day.	[greater than or equal to 0]; [empty]
Sample collection location in the wastewater system, whether at a wastewater treatment plant (or other community level treatment infrastructure such as community-scale septic) or upstream in the wastewater system	[See Value Sets: vs sample location]
If 'sample_location' is "upstream", specify the collection location in the wastewater system; an arbitrary name may be used if you do not wish to disclose the real name.	[string, length less than or equal to 40 characters]; [empty]
If this sample represents wastewater from a single institution, facility, or building, specify the institution type; otherwise, specify "not institution specific"	[See Value Sets: vs institution type]

NPDES permit number for the wastewater treatment plant specified in 'wwtp_name'. If 'upstream location' use the NPDES permit number for the wastewater treatment plant in which the water flows. If facility does not have an NPDES permit number, enter -1.	[NPDES permit number]; -1 if not permitted
EPA registry ID for the wastewater treatment plant specified in 'wwtp_name'. If 'upstream location' use the registry ID number for the wastewater treatment plant in which the water flows. If the water does not flow to a treatment plant, enter -1.	[EPA Registry ID]; -1 if not registered with EPA
The name of the Wastewater Treatment Plant (WWTP) to which this wastewater flows. If this wastewater does not flow to a WWTP, specify an identifiable name for the septic or other treatment system to which this wastewater flows. An arbitrary name may be used if you do not wish to disclose the real name.	[string, length less than or equal to 40 characters]
State, DC, US territory, or Freely Associated State jurisdiction name (2-letter abbreviation) in which the wastewater treatment plant provided in 'wwtp_name' is located	[See Value Sets: vs_wwtp_jurisdiction]
Wastewater treatment plant design capacity. This should be the capacity for which the plant is permitted. If 'upstream location', use the design capacity for the wastewater treatment plant to which the water flows.	[greater than or equal to 0]
Approximate average percentage of wastewater from industrial sources that is received by the wastewater treatment plant specified in 'wwtp_name'	[0-100]; [empty]
Does the wastewater treatment plant specified in 'wwtp_name' treat water from a combined sewer system (i.e., a sewer system that collects both sewage and stormwater)?	[See Value Sets: vs_yne]
Is influent to the wastewater treatment plant specified in 'wwtp_name' ever stored prior to treatment to equilibrate or modulate the influent flow rate?	[See Value Sets: vs_yne]

Type of sample collected, whether grab or composite. [See Value Sets: vs sample type] If composite, also provide the duration of sampling and type of composite, as listed in the Value Set (e.g., "24-hr flow-weighted composite"). A grab sample is defined as an individual sample collected without compositing or adding other samples, regardless of whether the sample matrix is liquid wastewater or sludge.

Frequency of sub-sample collection (for composite samples only): for flow-weighted, the number of subsamples collected per million gallons of flow; for timeweighted, the number of sub-samples per hour. Flowweighted example: a value of 5 would indicate 5 subsamples per million gallons, or 1 sub-sample per 200,000 gallons

Wastewater matrix from which the sample was collected

Duration of time from sample collection start time to time sample reached the lab.

Temperature at which the sample was stored after collection and prior to reaching the lab

Was the sample treated with any chemicals prior to reaching the lab? These could include chemicals, such as stabilizers, added to the sample or chemicals, such as chlorine, added to the wastewater treatment train upstream of the sample collection point. Do <u>not</u> be specify pasteurization here; it should be specified in the 'pasteurized' field.

If 'pretreatment' is "yes", then specify the chemicals used

[See Value Sets: vs_sample_matrix]

[greater than or equal to 0]; [empty] [float]; [empty]

[greater than or equal to 0];

[empty]

[See Value Sets: vs_yn]

[string]; [empty]

Process used to separate solid and liquid phases of the sample, either prior to or in the absence of the concentration method specified in 'concentration_method'

Method used to concentrate the sample prior to analysis of the concentrate

Method used for nucleic acid extraction from the sample

The approximate average duration of time between when samples reach the lab and when they are concentrated (if concentrated) [See Value Sets: vs concentration method]

[See Value Sets: vs extraction method]

[greater than or equal to 0]; [empty]

The storage temperature of samples after reaching the lab and prior to concentration (if concentrated)	[float]; [empty]
The approximate average duration of time between when samples are concentrated (if concentrated) and when they are extracted	[greater than or equal to 0]; [empty]
The storage temperature of samples after concentration (if concentrated) and prior to extraction	[float]; [empty]
Total volume of sample concentrated (if concentrated); this total volume is not necessarily assayed and is not necessarily equal to the value specified in 'equiv_sewage_amt'	[greater than or equal to 0]; [empty]
Are extraction blanks included in the extraction process?	[See Value Sets: vs yne]
Name of the recovery efficiency control target that is spiked in	<u>[See Value Sets:</u> <u>vs_rec_eff_target_name]</u>
Matrix into which the recovery efficiency control target is spiked	<u>[See Value Sets:</u> <u>vs_rec_eff_spike_matrix]</u>
Spike concentration, on average, of the recovery control on a per sample volume basis	[float]; [empty]
Was the sample pasteurized?	[See Value Sets: vs_yn]
The target of the DCD muchtification	
The Larget of the PCK quantification	[See Value Sets: vs pcr target]
The PCR gene used to quantify PCR target	<u>[See value Sets: vs_pcr_gene_target]</u>
A publication, website, or brief description of the PCR gene target used	[string]

The type of PCR used to quantify the PCR target [See Value Sets: vs pcr type]

A publication, website, or brief description of the method used to calculate the limit of detection	[string]
Name of microbial target used to estimate human fecal content	[See Value Sets: vs hum frac target mic]
A publication, website, or brief description of the microbial target specified in 'hum_frac_target_mic'	[string]; [empty]
Name of chemical compound used to estimate human fecal content	[See Value Sets: vs hum frac target chem]
A publication, website, or brief description of the chemical compound specified in 'hum_frac_target_chem'	[string]; [empty]
Name of a target or compound not specified in 'hum_frac_target_mic' or 'hum_frac_target_chem' used to estimate human fecal content	[See Value Sets: vs_other_norm_name]
A publication, website, or brief description of the target or compound specified in 'other_norm_name'	[string]; [empty]
The type of nucleic acid used as a standard for the PCR target quantification	<u>[See Value Sets:</u> <u>vs_quant_stan_type]</u>
A publication, website, or brief description of the quantitative standard material used	[string]
A publication, website, or brief description of the method used to evaluate molecular inhibition	[string]; none (if inhibition not tested)
Number of no-template controls (NTC) per instrument run	[See Value Sets: vs num no target control]
The date of sample collection; for composite samples, specify the date on which sample collection began	[date not after tomorrow's date]
The local time of sample collection; for composite samples, specify the time at which sample collection began	[time]
Current local time zone corresponding to the time specified in 'sample_collect_time', represented as a UTC time offset (e.g., UTC-06:00)	[time zone]

Wastewater volumetric flow rate at the sample collection location over the 24-hr period during which the sample was collected. If only an instantaneous flow measurement is available, it may be reported in units of million gallons per day.

[greater than or equal to 0]; [empty]

pH of wastewater sample (if sludge, pH of influent at time of collection)

Specific conductivity of wastewater sample (if sludge, conductivity of influent at time of collection)

Total suspended solids of raw (or, if unavailable, postgrit removal) wastewater

Sample temperature at time of collection

Equivalent unconcentrated volume of wastewater or mass of sludge in PCR reaction

An ID assigned to a wastewater sample. It must be unique for this NWSS reporting jurisdiction. Wastewater samples that are split and measured by different labs should have the same sample ID but different lab IDs. Wastewater samples for which multiple PCR targets are measured should also have the same sample ID. Note: do not include PII in this field.

An ID assigned to a testing lab. It must be unique [lab id] across labs used for this NWSS reporting jurisdiction's testing. If the same lab is used across multiple NWSS reporting jurisdictions, each NWSS reporting jurisdiction may assign that lab a different lab ID. Note: including PII in this field is discouraged.

[float]; [empty]

[greater than or equal to 0]; [empty]

[sample id]

Should the DCIPHER QC report ignore this sample? The [See Value Sets: vs yne] purpose of this field is to allow for removal of samples from the QC report so that samples with unresolvable QC issues do not clutter the report. This field only affects whether samples are shown in the QC report, not how they are otherwise processed or analyzed. If the value is "yes", then this sample will be excluded from the QC report. If the value is "no" or [empty], then this sample will be included in the QC report for up to 90 days. Samples will only be included in the QC report for up to 90 days, regardless of the value specified in 'qc_ignore'.

Should the DCIPHER dashboard ignore this sample? If [See Value Sets: vs vne] the value is "yes", then this sample will be excluded from the dashboard. If the value is "no" or [empty], then this sample will be included in the dashboard. In addition to excluding the sample from the DCIPHER dashboard, this field will also prevent the sample from public data release.

Should DCIPHER data analysis ignore this sample? If the value is "yes", then this sample will be excluded from DCIPHER data analysis, which will also cause it to be excluded from the dashboard. If the value is "no" or [empty], then this sample will be included in DCIPHER data analysis.

[See Value Sets: vs yne]

The date on which this PCR target measurement was made	[date greater than sample collect date and less than DCIPHER upload date]
Units of PCR target sample concentration	[See Value Sets: vs mic chem units]

Concentration of the PCR target back-calculated to unconcentrated sample basis; enter "0" if no amplification occurred, using the definition of amplification described in 'ntc_amplify'; otherwise, enter the estimated concentration; do not adjust for matrix recovery efficiency. Only the technical or biological replicate concentrations for the same pcr_gene_target should be averaged. For example, if there are three replicates of a given PCR target, average those concentrations and report the average value only. Do not include the three individual concentrations. Different pcr_gene_target concentrations **should not** be averaged for this field (e.g., do not average concentrations for N1 and N2).

Standard error (SE) of the PCR target in wastewater sample, or best estimate that is consistently available. If sample replicates are always performed, use SE of sample replicates; else, if processing replicates are always performed, use SE of processing replicates; else, if qPCR is performed, use SE of PCR replicates; else, if digital PCR is performed, use error from multiple replicates if available, and Poisson error if not

[greater than or equal to 0]; -1 (if cannot be calculated, such as when no amplification observed)

[any float other than 0];

0 (if no amplification observed)

Lower bound of 95% confidence interval of the PCR target in wastewater sample, or best estimate that is consistently available. Follow the same hierarchy as described for standard error. (Note: 'cl' stands for confidence limit)

Upper bound of 95% confidence interval of the PCR target in wastewater sample, or best estimate that is consistently available. Follow the same hierarchy as described for standard error. (Note: 'cl' stands for confidence limit)

Was the concentration of the PCR target below the limit of detection?

[any float other than -1]; -1 (if cannot be calculated, such as when no amplification observed)

[any float other than -1]; -1 (if cannot be calculated, such as when no amplification observed)

[See Value Sets: vs_yne]

PCR target limit of detection back-calculated to unconcentrated sample basis

[float]

For qPCR, did any no-template controls on this instrument run have a Ct value less than 40? For ddPCR, did any no-template controls on this instrument run have 3 or more positive droplets?	[See Value Sets: vs yn]
Percent of spiked recovery control, specified in 'rec_eff_target_name', that was recovered	[greater than or equal to 0.0]; -1 (if not tested)
Was molecular inhibition detected? Was inhibition incorporated into the PCR target concentration calculation?	[See Value Sets: vs ynn] [See Value Sets: vs yne]
Concentration of microbial target specified in 'hum_frac_target_mic'; follow the same guidelines outlined for 'pcr_target_avg_conc'	[float]; [empty]
Concentration units of microbial target specified in 'hum_frac_target_mic'	[See Value Sets: vs mic chem units e]
Concentration of chemical target specified in 'hum_frac_target_chem'	[float]; [empty]
Concentration units of chemical target specified in 'hum_frac_target_chem'	[See Value Sets: vs mic chem units e]
Concentration of target spcified in 'other_norm_name'	[float]; [empty]
Concentration units of target spcified in 'other_norm_name'	[See Value Sets: vs_mic_chem_units_e]
Does this observation have quality control issues?	[See Value Sets: vs_yne]

A number used to distinguish major lab methods at the [greater than or equal to 0]; reporting jurisdiction level. Differences in lab methods [empty] may limit the comparability of PCR target concentrations. Reporting jurisdictions should use this field to indicate that lab methods are sufficiently different that caution should be used when comparing PCR target concentrations across them. These differences may result from any lab processing or quantification steps, including different PCR gene targets. Note that 'major_lab_method' may be the same for different laboratories if the reporting jurisdiction believes concentrations measured by those laboratories are comparable; conversely, 'major lab method' may be different for the same laboratory if changes made to the laboratory protocol rendered concentrations not comparable.

Brief description of the basis for assigning a [string]; 'major_lab_method', e.g., "this lab uses a lab method distinct from other labs in this reporting jurisdiction", "this lab changed lab methods - new concentration method" or "lab method is same as other labs in this reporting jurisdiction but performance differs"

Units	Submission Requirement	Dependent Fields
[none]	Required	None
	Required	None
[none]	Required	None
[none]	Required	None
[none]	Required	None
Hours	Not required	None
[none]	Required	If 'sample_location' is "upstream", specify in 'sample_location_specify'
[none]	Required	If sample_location is "upstream", then this must have a non-empty value
[none]	Required	None

[none]	Required	None
[none]	Not required	None
[none]	Required	None
[none]	Required	None
Million gallons per day (MGD)	Required	None
percent	Not required	None
[none]	Not required	None
[none]	Not required	None

Required

None

If flow-weighted composite: number per million gallons; if time- weighted or manual composite: number per hour	Not required	None
[none]	Required	None
Hours	Not required	None
Celsius	Not required	None
[none]	Required	if 'pretreatment' is "yes", then specify in 'pretreatment_specify'
[none]	Not required	if 'pretreatment' is "yes", then 'pretreatment_specify' must have a non-empty field
[none]	Networkingd	None
נווטוופן	NOT required	NOTE
[none]	Required	None
[none]	Required	None
Hours	Not required	None

Celsius	Not required	None
Hours	Not required	None
Celsius	Not required	None
mL	Not required	None
[none]	Not required	None
[none]	Required	If 'rec_eff_percent' is equal to a value other than "-1", then this must have a non- empty value
[none]	Required	If 'rec_eff_target_name' has a non-empty value, then this must have a non- empty value
log10 copies/mL	Required	If 'rec_eff_target_name' has a non-empty value, then this must have a non- empty value
[none]	Required	None
[none] [none]	Required Required	None The value selected for pcr_gene_target should align with the value selected for pcr_target
[none]	Required	None
[none]	Required	If 'pcr_target' is not "sars- cov-2", "delta", "omicron", "hMPXV", "hMPXV Clade I", or "hMPXV Clade II" then this must be a type of digital PCR, i.e. "ddpcr", "qiagen dpcr", "fluidigm dpcr", "life technologies dpcr", "raindance dpcr", or

"dpcr"

[none]	Required	None
[none]	Not required	If 'hum_frac_mic_conc' has a non-empty value, then this must have a non- empty value
[none]	Not required	If 'hum_frac_mic_conc' has a non-empty value, then this must have a non- empty value
[none]	Not required	If 'hum_frac_chem_conc' has a non-empty value, then this must have a non- empty value
[none]	Not required	If 'hum_frac_chem_conc' has a non-empty value, then this must have a non- empty value
[none]	Not required	If 'other_norm_conc' has a non-empty value, then this must have a non-empty value
[none]	Not required	If 'other_norm_conc' has a non-empty value, then this must have a non-empty value
[none]	Required	None
[none]	Required	None
[none]	Required	None

Million gallons per day (MGD)	Required.	If 'sars_cov2_units' is on a per volume wastewater basis (e.g., copies/l wastewater), then this must have a non-empty value (i.e., measurements of wastewater solids are permitted empty values for 'flow_rate'). If 'hum_frac_mic_conc' has a non-empty value and it is not possible to measure flow rate at all, then this may have an empty value.

pH units	Not required	None
microsiemens/cm	Not required	None
mg/L	Not required	None
Celsius	Not required	None
mL wastewater or g sludge	Not required	None
[none]	Required	None

[none]

Required

None

one]	Not required	None
one]	Not required	None
one]	Not required	None
one]	Not required	None

[none]	Required	None
[none]	Required	The units should relate to the PCR target indicated in pcr_target

[units specif	ied in
'pcr_target_	_units']

Required

The concentration should relate to the PCR target indicated in pcr_target

[units specified in 'pcr_target_units']	Not required	The standard error should relate to the PCR target indicated in pcr_target
[units specified in 'pcr_target_units']	Not required	The lower bound of the 95% confidence interval should relate to the PCR target indicated in pcr_target
[units specified in 'pcr_target_units']	Not required	The upper bound of the 95% confidence interval should relate to the PCR target indicated in pcr_target
[none]	Not required	Whether the concentration was below the LOD should be based on the LOD of the PCR target indicated in lod_sewage
[units specified in 'pcr_target_units']	Required	The LOD should relate to the PCR target indicated in pcr_target

[none]	Required	None
percent	Required	None
[none] [none]	Required Required	None If 'inhibition_detect' is "yes", then this must have a non-empty value
[units specified in 'hum_frac_mic_unit']	Not required	If 'hum_frac_mic_unit' has a non-empty value, then this must have a non- empty value
[none]	Not required	If 'hum_frac_mic_conc' has a non-empty value, then this must have a non- empty value
[units specified in 'hum_frac_chem_unit']	Not required	If 'hum_frac_chem_unit' has a non-empty value, then this must have a non- empty value
[none]	Not required	If 'hum_frac_chem_conc' has a non-empty value, then this must have a non- empty value
[units specified in 'other_norm_conc']	Not required	If 'other_norm_unit' has a non-empty value, then this must have a non-empty value
[none]	Not required	If 'other_norm_conc' has a non-empty value, then this must have a non-empty value
[none]	Not required	None

Required

None

[none]

Required

None
Field

concentration_method	vs_concentration_method
extraction_method	vs_extraction_method
hum_frac_target_chem	vs_hum_frac_target_chem
hum_frac_target_mic	vs_hum_frac_target_mic
institution_type	vs_institution_type
sars_cov2_units	vs_mic_chem_units
hum_frac_chem_unit	vs_mic_chem_units_e
other_norm_units	vs_mic_chem_units_e
hum_frac_mic_unit	vs_mic_chem_units_e
num_no_target_control	vs_num_no_target_control
other_norm_name	vs_other_norm_name
pcr_target	vs_pcr_target
pcr_gene_target	vs_pcr_gene_target
pcr_type	vs_pcr_type
quant_stan_type	vs_quant_stan_type
rec_eff_spike_matrix	vs_rec_eff_spike_matrix
rec_eff_target_name	vs_rec_eff_target_name
reporting_jurisdiction	vs_reporting_jurisdiction
sample_location	vs_sample_location
sample_matrix	vs_sample_matrix
sample_type	vs_sample_type
solids_separation	vs_solids_separation
wwtp_jurisdiction	vs_wwtp_jurisdiction
sars_cov2_below_lod	vs_yn
ntc_amplify	vs_yn
stormwater_input	vs_yne
influent_equilibrated	vs_yne
pretreatment	vs_yne
ext_blank	vs_yne
pasteurized	vs_yne
inhibition_adjust	vs_yne
quality_flag	vs_yne
inhibition_detect	vs_ynn

vs_concentration_method

Value Set membrane filtration with addition of mgcl2 membrane filtration with sample acidification membrane filtration with acidification and mgcl2 membrane filtration with no amendment membrane filtration with addition of mgcl2, membrane recombined with separated solids membrane filtration with sample acidification, membrane recombined with separated solids membrane filtration with acidification and mgcl2, membrane recombined with separated solids membrane filtration with no amendment, membrane recombined with separated solids peg precipitation aloh3 precipitation ultracentrifugation skimmed milk flocculation beef extract flocculation promega wastewater large volume tna capture kit centricon ultrafiltration amicon ultrafiltration hollow fiber dead end ultrafiltration innovaprep ultrafiltration no liquid concentration, liquid recombined with separated solids ceres nanotrap zymo environ water rna kit/ zymo environ water rna kit (cat. r2042) membrane filtration with addition of mgcl3 water concentrating buffer (R2042-1) none

vs_extraction_method

Value Set giagen allprep powerviral dna/rna kit qiagen allprep powerfecal dna/rna kit qiagen allprep dna/rna kit qiagen rneasy powermicrobiome kit qiagen powerwater kit qiagen rneasy kit qiagen ez1 virus mini kit v2.0 promega ht tna kit promega automated tha kit promega manual tna kit promega wastewater large volume tha capture kit nuclisens automated magnetic bead extraction kit nuclisens manual magnetic bead extraction kit phenol chloroform chemagic viral dna/rna 300 kit trizol, zymo mag beads w/ zymo clean and concentrator 4s method (https://www.protocols.io/view/v-4-direct-wastewater-rna-capture-and-purification-bpdfmi3n) qiagen qiaamp buffers with epoch columns zymo quick-rna fungal/bacterial miniprep #r2014 thermo magmax microbiome ultra nucleic acid isolation kit zymo environ water rna kit/ zymo environ water rna kit (cat. r2042) luminultra wastewater extraction kit giaamp viral rna kit trizol and RNA purification kit trizol, garnet bead beating, alcohol precipitation zymo quick-rna viral 96 kit #r1041 zymo quick-rna viral kit #r1035 qiagen qiaamp dsp viral rna mini kit monarch total RNA miniprep kit (new england biolabs) + onestep PCR inhibitor removal kit (zymo) exclusions based sample preparation (ESP) sciencell viral rna isolation kit thermo magmax viral/pathogen nucleic acid isolation kit

	vs_hum_frac_target_chem		
Description	Value Set	Description	
	caffeine		
	creatinine		
	sucralose		
	ibuprofen		
	[empty]		

vs_hum_frac_target_mic	vs_institution_type
Value Set Descrip	otion Value Set
pepper mild mottle virus	not institution specific
crassphage	correctional
hf183	long term care - nursing home
f+ rna coliphage	long term care - assisted living
f+ dna coliphage	other long term care
PMMoV (GT-Digital)	short stay acute care hospital
[empty]	long term acute care hospital
	child day care
	k12
	higher ed dorm
	higher ed other
	social services shelter
	other residential building
	ship
	airplane
	other worksite

This sample does not represent wastewater from a single institution, facility, or building

A prison, penitentiary, penal facility, jail, detention unit, or other facility in which persons are incarcerated by government office A residential healthcare facility that provides 24-medical care. These are also called skilled nursing facilities. Generally licensed A residential facility that provides assistance with daily care but generally does not provide skilled nursing care. May be licensed Other residential facilities that provide daily and/or medical care, but are not defined as nursing home/skilled nursing facilities.

Acute care hospitals that provide care for patients with average length of stay longer than 25 days. LTACH patients are often t

A school serving students in the kindergarten to 12th grade range

Solely higher education dormitory buildings

Higher education buildings or facilities that do not include dorms or that include dorms and other buildings

Other type of social services shelter

Individual residential buildings or institutions not captured in other categories

Any commercial facility not captured in other commercial facility categories

vs_mic_chem_units Value Set

copies/L wastewater log10 copies/L wastewater copies/g wet sludge log10 copies/g wet sludge copies/g dry sludge log10 copies/g dry sludge micrograms/L wastewater log10 micrograms/L wastewater micrograms/g wet sludge log10 micrograms/g wet sludge micrograms/g dry sludge

Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven

Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven vs_mic_chem_units_e Value Set

copies/L wastewater log10 copies/L wastewater copies/g wet sludge log10 copies/g wet sludge copies/g dry sludge log10 copies/g dry sludge micrograms/L wastewater log10 micrograms/L wastewa micrograms/g wet sludge log10 micrograms/g wet sluc micrograms/g dry sludge log10 micrograms/g dry slud [empty]

Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven

ater

Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Wet sludge mass is based on the mass of sludge without drying the solids in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven Dry sludge mass is based on the mass of solids after drying in a drying oven

vs_num_no_target_control		vs_other_norm_name	
Value Set	Description	Value Set	Description
0		pepper mild mottle vi	rus
1		crassphage	
2		hf183	
3		f+ rna coliphage	
more than 3		f+ dna coliphage	
		caffeine	
		creatinine	
		sucralose	
		ibuprofen	
		rnase p	
		[empty]	

vs_pcr_target	
Value Set	Description
sars-cov-2	
delta	
omicron	
hMPXV	
hMPXV Clade I	
hMPXV Clade II	
caur	Candida auris. Source: KEGG
NoV GI	Norovirus GI. Source: https://www.ncbi.nlm.nih.gov/pm
NoV GII	Norovirus GII. Source: https://www.ncbi.nlm.nih.gov/pm
FLUAV	Influenza A. Source: Bacterial and Viral Bioinformatics R
FLUBV	Influenza B. Source: Bacterial and Viral Bioinformatics R
NVO	Non-Variolla Orthopoxvirus. Source: CDC https://ww
Cjejuni	Campylobacter jejuni. Source: KEGG
HAdV-F	Adenovirus Group F41. Source: PubChem Taxonomy
Col	Colistin AMR
Gly	Vancomycin AMR
Bla	Beta-Lactamase AMR
Integrase	Integrase AMR
Tet	Tetracycline AMR
RSV	Respiratory Syncytial Virus
Fluor	Fluoroquinolone AMR
Macro	Macrolide AMR
Shigella	Shigella spp.
STEC	Shiga-toxin producing or intimin-producing E. coli or othe
cycloc	Cyclospora cayetanensis
Other Bacteria	Other bacteria not listed
Other Virus	Other virus not listed
Other Parasite	Other parasite not listed
Other Fungus or V	/«Other fungus or yeast not listed
Other Eukaryote	Other eukaryote not listed
Other Archaea	Other archaea not listed

vs_pcr_gene_target Value Set n1 n2 n3 e_sarbeco n_sarbeco rdrp_sarsr niid_2019-ncov_n rdrp gene / ncov_ip2 rdrp gene / ncov_ip4 taqpath n taqpath s orf1b orf1ab n1 and n2 combined n s orf1a ddcov_n ddcov_e ip2 and ip4 combined CDC N1 (GT-Digital) CDC N2 (GT-Digital) N2 (PREvalence) E (PREvalence) a1306s p2046l p2287s t3646a v2930l g662s p100l a1918v t19r e156g del 157/158 l452r p681r d950n i82t d63g r203m g215c d377y wt214 del156-157

k856r s2083i del2084/2084 a2710t p3395h del3674/3676 i3758v i1566v a67v del69/70 a67v/del69/del70 del143/145 n211i del212/212 g339d s371l s373p s375f s477n e484a q493r g496s q498r y505h t547k n679k d796y n856k q954h n969k l981f t9i d3g q19e a63t ins214epe del142-144 del31-33 l24s lppa24s p13L E9L-NVAR G2R_G G2R_WA C3L E9L-OPX3 B6R

gtmol_hMPXV MCR-1.1 VanA-A blaNDM blaKPC blaOXA-48 blaVIM blaCTX-M-1 blaCMY blaTEM Class I Integrase blaSHV TetW blaIMP caur NoV GI NoV GII InfA1 InfA2 InfA1 and InfA2 combi InfB Cjejuni . HAdV-F RSVL1 ipaH 18S rRNA mph(A) stx1 stx2 eae RSV fluor Other

Description sars-cov-2 gene target sars-cov-2 gene target; Detection of N1 and N2 targets was performed on a single fluorescence channel, such that distinct me sars-cov-2 gene target delta gene target

omicron gene target omicron gene target

hMPXV gene target. orthopoxvirus DNA polymerase. CDC Assay all hMPXV and non-variola Orthopoxvirus. https://doi.org/10. hMPXV gene target. G2R_G primers and probe: detects all MPXV strains . https://doi.org/10.1016/j.jviromet.2010.07.012 hMPXV Clade II gene target. hMPXV G2R_WA primers and probe: detects Clade II viruses.CDC https://doi:10.1016/j.jviromet.2010.07 hMPXV Clade I gene target. hMPXV C3L primers and probe: detects Clade I viruses . CDC https://doi:10.1016/j.jviromet.2010.07 hMPXV gene target. CDC https://doi.org/10.4269/ajtmh.2010.09-0716

hMPXV gene target. Envelope protien gene CDC Assay; selective for MPXV. CDC https://doi.org/10.1016/j.jcv.2006.03.012

hMPXV gene target. hMPXV gene target proprietary GT Molecular Colistin AMR gene target. Source: ARG-ANNOT Vancomycin AMR gene target. Source: ARG-ANNOT New-Delhi Metallo- (NDM) Beta-Lactamase AMR gene target. Source: ARG-ANNOT Klebsiella pneumoniae Carbapenam (KPC) Beta-Lactamase AMR gene target. Source: ARG-ANNOT OXA-type Beta-Lactamase AMR gene target. Source: ARG-ANNOT Verone Integron-Encoded Metallo- (VIM) Beta-Lactamase AMR gene target. Source: ARG-ANNOT CTX-M Beta-Lactamase AMR gene target. Source: ARG-ANNOT Cephamycin Beta-Lactamase AMR gene target. Source: ARG-ANNOT Beta-Lactamase AMR gene target. Source: ARG-ANNOT Integrase AMR gene target. Beta-Lactamase AMR gene target. Source: ARG-ANNOT Tetracycline AMR gene target. Source: ARG-ANNOT Impenemase (IMP) Beta-Lactamase AMR gene target. Source: ARG-ANNOT Candida auris gene target. Source: https://www.cdc.gov/fungal/candida-auris/pdf/Real-time-PCR-based-Id-C-auris-508.pdf Norovirus GI ORF1-ORF2 junction gene target. Source: https://wwwnc.cdc.gov/eid/article/17/8/10-1837-t2#r27 Norovirus GII ORF1-ORF2 junction gene target. Source: https://wwwnc.cdc.gov/eid/article/17/8/10-1837-t2#r27 Influenza A 1 gene target. Source: https://www.cdc.gov/coronavirus/2019-ncov/lab/multiplex.html?CDC_AA_refVal=https%3. Influenza A 2 gene target. Source: https://www.cdc.gov/coronavirus/2019-ncov/lab/multiplex.html?CDC_AA_refVal=https%3. Influenza A gene target; Detection of InfA1 and InfA2 targets was performed on a single fluorescence channel, such that distir Influenza B gene target. Source: https://www.cdc.gov/coronavirus/2019-ncov/lab/multiplex.html?CDC_AA_refVal=https%3A Campylobacter jejuni gene target. Source: https://www.sciencedirect.com/science/article/pii/S0882401017303728?via%3Dih Adenovirus Group F41 Fiber gene target. Source: https://www.sciencedirect.com/science/article/pii/S0022347608006744?via RSV-A and RSV-B gene tartget. Primer sets are reactive to both viruses. Two separate probes for RSV-A and RSV-B. https://cdr Shigella gene target Eukaryotic gene target (e.g., Cyclospora) Macrolide resistance target STEC gene target STEC gene target STEC gene target **RSV** gene target Fluoroquinolone gene target Please contact NWSS staff to submit data using the "Other" category, or to request that values be added to the vocabulary.

vs_pcr_type Value Set qpcr ddpcr qiagen dpcr fluidigm dpcr life technologies dpcr raindance dpcr dpcr 9-ncov%2Flab%2Fmultiplex-primer-probes 9-ncov%2Flab%2Fmultiplex-primer-probes (ist. An example is the GT-Digital Influenza •ncov%2Flab%2Fmultiplex-primer-probes.l

/rsv-surveillance/who-rsv-surveillance-stra

Real-time PCR, also called 'quantitative' PCR Used to refer specifically to BioRad digital droplet emulsification technology

Generic digital PCR

s.html s.html and SARS-CoV-2 Wastewater Surveillance Multiplex Assay Kits. html

```
ategy-phase-26mar2021.-final.pdf?sfvrsn=d8b1c36a_9;%20https://journals.plos.org/plosone/article
```

vs_quant_stan_type		vs_rec_eff_spike_matrix	
Value Set	Description	Value Set	Description
dna raw samp		raw sample	
rna		raw sample post pasteurizat	ion
		clarified sample	
		sample concentrate	
		lysis buffer	
		dewatered solids	
		[empty]	

e?id=10.1371/journal.pone.0015098

Value Set	Description
bcov vaccine	
bcov culture	
brsv vaccine	
brsv culture	
murine coronavirus	
oc43	
phi6	
puro	
ms2 coliphage	
hep g armored rna	
heat inactivated sars-cov-2 virus	
murine hepatitis virus	
MHV (PREvalence)	
BCoV (GT-Digital)	
[empty]	
parapoxvirus	

vs_reporting_jurisdiction Value Set Description AL Alabama AK Alaska AS American Samoa ΑZ Arizona AR Arkansas CA California CI Chicago, IL CO Colorado MP Commonwealth of Northern Mariana Islands СТ Connecticut DE Delaware DC **District of Columbia** FΜ Federated States of Micronesia FL Florida GA Georgia GU Guam HI Hawaii Houston, TX HO ID Idaho IL Illinois Indiana IN IA Iowa KS Kansas КΥ Kentucky LC Los Angeles County, CA LA Louisiana ME Maine MD Maryland MA Massachusetts MI Michigan MN Minnesota MS Mississippi MO Missouri MT Montana Nebraska NE Nevada NV NH **New Hampshire** NJ New Jersey NM New Mexico NY New York New York City, NY NZ NC North Carolina ND North Dakota OH Ohio Oklahoma OK

OR	Oregon
PA	Pennsylvania
PH	Philadelphia, PA
PR	Puerto Rico
MH	Republic of the Marshall Islands
PW	Republic of Palau
RI	Rhode Island
SC	South Carolina
SD	South Dakota
TN	Tennessee
ТХ	Texas
VI	U.S. Virgin Islands
UT	Utah
VT	Vermont
VA	Virginia
WA	Washington
WV	West Virginia
WI	Wisconsin
WY	Wyoming
AP	Acoma Pueblo
CN	Catawba Nation
CVBPI	Coyote Valley Band of Pomo Indians
EBCI	Eastern Band of Cherokee Indians
GTBI	Grand Traverse Band of Ottawa and Chippewa Indians
MAT	Mescalero Apache Tribe
NPT	Nez Perce Tribe
OMT	Otoe-Missouria Tribe
SBT	The Shoshone Bannock Tribes
TRIT	Tule River Indian Tribe
ZP	Zuni Pueblo
PBCI	Poarch Band of Creek Indians

vs_sample_location Value Set wwtp upstream

A sampling location at a wastewater treatment plant or other community-scale treatment infrastructure specified in 'wwtp_n A sampling location other than "wwtp"

vs_sample_matrix Value Set

raw wastewater post grit removal primary sludge primary effluent secondary sludge secondary effluent septage holding tank

- Wastewater without any form of treatment applied to it
- Wastewater after removal of large solids at a treatment plant but prior to a primary clarifier
- Sludge from the primary clarifier
- Effluent from the primary clarifier
- Sludge from the secondary clarifier
- Efluent from the secondary clarifier
- Wastewater sampled from within a septic tank
- Wastewater sampled from a holding tank, such as from an airplane or ship

vs_sample_type Value Set

grab

48-hr flow-weighted composite 47-hr flow-weighted composite 46-hr flow-weighted composite 45-hr flow-weighted composite 44-hr flow-weighted composite 43-hr flow-weighted composite 42-hr flow-weighted composite 41-hr flow-weighted composite 40-hr flow-weighted composite 39-hr flow-weighted composite 38-hr flow-weighted composite 37-hr flow-weighted composite 36-hr flow-weighted composite 35-hr flow-weighted composite 34-hr flow-weighted composite 33-hr flow-weighted composite 32-hr flow-weighted composite 31-hr flow-weighted composite 30-hr flow-weighted composite 29-hr flow-weighted composite 28-hr flow-weighted composite 27-hr flow-weighted composite 26-hr flow-weighted composite 25-hr flow-weighted composite 24-hr flow-weighted composite 23-hr flow-weighted composite 22-hr flow-weighted composite 21-hr flow-weighted composite 20-hr flow-weighted composite 19-hr flow-weighted composite 18-hr flow-weighted composite 17-hr flow-weighted composite 16-hr flow-weighted composite 15-hr flow-weighted composite 14-hr flow-weighted composite 13-hr flow-weighted composite 12-hr flow-weighted composite 11-hr flow-weighted composite 10-hr flow-weighted composite 9-hr flow-weighted composite 8-hr flow-weighted composite 7-hr flow-weighted composite 6-hr flow-weighted composite 5-hr flow-weighted composite

4-hr flow-weighted composite 3-hr flow-weighted composite 2-hr flow-weighted composite 1-hr flow-weighted composite 72-hr time-weighted composite 48-hr time-weighted composite 47-hr time-weighted composite 46-hr time-weighted composite 45-hr time-weighted composite 44-hr time-weighted composite 43-hr time-weighted composite 42-hr time-weighted composite 41-hr time-weighted composite 40-hr time-weighted composite 39-hr time-weighted composite 38-hr time-weighted composite 37-hr time-weighted composite 36-hr time-weighted composite 35-hr time-weighted composite 34-hr time-weighted composite 33-hr time-weighted composite 32-hr time-weighted composite 31-hr time-weighted composite 30-hr time-weighted composite 29-hr time-weighted composite 28-hr time-weighted composite 27-hr time-weighted composite 26-hr time-weighted composite 25-hr time-weighted composite 24-hr time-weighted composite 23-hr time-weighted composite 22-hr time-weighted composite 21-hr time-weighted composite 20-hr time-weighted composite 19-hr time-weighted composite 18-hr time-weighted composite 17-hr time-weighted composite 16-hr time-weighted composite 15-hr time-weighted composite 14-hr time-weighted composite 13-hr time-weighted composite 12-hr time-weighted composite 11-hr time-weighted composite 10-hr time-weighted composite 9-hr time-weighted composite 8-hr time-weighted composite 7-hr time-weighted composite

6-hr time-weighted composite 5-hr time-weighted composite 4-hr time-weighted composite 3-hr time-weighted composite 2-hr time-weighted composite 1-hr time-weighted composite 48-hr manual composite 46-hr manual composite 45-hr manual composite

44-hr manual composite
43-hr manual composite
42-hr manual composite
41-hr manual composite
40-hr manual composite
39-hr manual composite

38-hr manual composite 37-hr manual composite 36-hr manual composite 35-hr manual composite 34-hr manual composite 33-hr manual composite 32-hr manual composite 31-hr manual composite 30-hr manual composite 29-hr manual composite 28-hr manual composite 27-hr manual composite 26-hr manual composite 25-hr manual composite 24-hr manual composite 23-hr manual composite 22-hr manual composite 21-hr manual composite 20-hr manual composite 19-hr manual composite 18-hr manual composite 17-hr manual composite 16-hr manual composite 15-hr manual composite 14-hr manual composite 13-hr manual composite 12-hr manual composite 11-hr manual composite 10-hr manual composite 9-hr manual composite 8-hr manual composite 7-hr manual composite 6-hr manual composite 5-hr manual composite 4-hr manual composite 3-hr manual composite 2-hr manual composite 1-hr manual composite 72-hr passive sample 48-hr passive sample 47-hr passive sample 46-hr passive sample 45-hr passive sample 44-hr passive sample 43-hr passive sample 42-hr passive sample 41-hr passive sample 40-hr passive sample 39-hr passive sample 38-hr passive sample 37-hr passive sample 36-hr passive sample 35-hr passive sample 34-hr passive sample 33-hr passive sample 32-hr passive sample 31-hr passive sample 30-hr passive sample 29-hr passive sample 28-hr passive sample 27-hr passive sample 26-hr passive sample 25-hr passive sample 24-hr passive sample 23-hr passive sample 22-hr passive sample 21-hr passive sample 20-hr passive sample 19-hr passive sample 18-hr passive sample 17-hr passive sample 16-hr passive sample 15-hr passive sample 14-hr passive sample 13-hr passive sample 12-hr passive sample
11-hr passive sample
10-hr passive sample
9-hr passive sample
8-hr passive sample
7-hr passive sample
6-hr passive sample
5-hr passive sample
4-hr passive sample
3-hr passive sample
2-hr passive sample
1-hr passive sample
120-hr passive sample

Description

An individual sample collected without compositing or adding other samples

E.g., Moore swabs E.g., Moore swabs

E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs E.g., Moore swabs vs_solids_separation Value Set filtration centrifugation gravity settling none [empty] Description

Filtration to remove solids from a wastewater sample prior to further concentration or analysis of the liquid filtrate Centrifugation prior to or in the absence of a concentration step; centrifugation as part of a flocculation or precipitation proce Solids separated from liquid by allowing solids to settle by gravity

	vs_wwtp_jurisdiction		
	Value Set	Description	
	AL	Alabama	
oncentration proce	e AK	Alaska	
	AS	American Samoa	
	AZ	Arizona	
	AR	Arkansas	
	CA	California	
	СО	Colorado	
	MP	Commonwealth of Northern Mariana Islands	
	СТ	Connecticut	
	DE	Delaware	
	DC	District of Columbia	
	FM	Federated States of Micronesia	
	FL	Florida	
	GA	Georgia	
	GU	Guam	
	HI	Наwаіі	
	ID	Idaho	
	IL	Illinois	
	IN	Indiana	
	IA	lowa	
	KS	Kansas	
	КҮ	Kentucky	
	LA	Louisiana	
	ME	Maine	
	MD	Maryland	
	MA	Massachusetts	
	MI	Michigan	
	MN	Minnesota	
	MS	Mississippi	
	MO	Missouri	
	MT	Montana	
	NE	Nebraska	
	NV	Nevada	
	NH	New Hampshire	
	NJ	New Jersey	
	NM	New Mexico	
	NY	New York	
	NC	North Carolina	
	ND	North Dakota	
	OH	Ohio	
	ОК	Oklahoma	
	OR	Oregon	
	PA	Pennsylvania	
	PR	Puerto Rico	
	MH	Republic of the Marshall Islands	

PW	Republic of Palau
RI	Rhode Island
SC	South Carolina
SD	South Dakota
TN	Tennessee
ТХ	Texas
VI	U.S. Virgin Islands
UT	Utah
VT	Vermont
VA	Virginia
WA	Washington
WV	West Virginia
WI	Wisconsin
WY	Wyoming

vs_yn Value Set Description	vs_yne Value Set Description	vs_ynn Value Set Description
yes	yes	yes
no	no	no
	[empty]	not tested

Use of semantic versioning began with v2.0.0. A description follows; see https://semv Major changes (indicated by X in version X.Y.Z): not backwards compatible. In the con Minor changes (indicated by Y in version X.Y.Z): backwards compatible functionality a Patches (indicated by Z in version X.Y.Z): backwards compatible bug fixes. In the conte

Date of change 20200923 20201020	New file name NWSS_DCIPHERForm_DataDictionary_Sent-to-states_9.23.20 NWSS_DCIPHERForm_DataDictionary_20201020
20201021	NWSS_DCIPHERForm_DataDictionary_20201021
20201103	NWSS_DCIPHERForm_DataDictionary_20201103
20201105	NWSS DCIPHERForm DataDictionary 20201104

- 20201201 NWSS_DCIPHERForm_DataDictionary_20201201
- 20201204 NWSS_DCIPHERForm_DataDictionary_20201204
- 20201215 NWSS_DCIPHERForm_DataDictionary_20201215
- 20201218 NWSS_DCIPHERForm_DataDictionary_20201218
- 20210111 NWSS_DCIPHERForm_DataDictionary_20210111
- 202101222 NWSS_DCIPHERForm_DataDictionary_20210122

202101225 NWSS_DCIPHERForm_DataDictionary_20210125

20210127 NWSS_DCIPHERForm_DataDictionary_20210127

202102110 NWSS_DCIPHERForm_DataDictionary_20210210

20210319 NWSS_DCIPHER_Data_Dictionary_v2.0.0_20210323

20210416 NWSS_DCIPHER_Data_Dictionary_v2.0.1_20210416

20210512 NWSS_DCIPHER_Data_Dictionary_v2.0.2_20210512

20210621 NWSS_DCIPHER_Data_Dictionary_v2.0.3_20210621

20210818 NWSS_DCIPHER_Data_Dictionary_v2.1.0_20210818

20211028 NWSS_DCIPHER_Data_Dictionary_v2.1.1_20211028

2021202 NWSS_DCIPHER_Data_Dictionary_v3.0.0_2021202

20211208 NWSS_DCIPHER_Data_Dictionary_v3.1.0_20211208

20220524 NWSS_DCIPHER_Data_Dictionary_v3.1.1_20220524

20220906 NWSS Data Dictionary_v3_1_3_20220906

20221018 NWSS Data Dictionary_v3_1_4_20221018

20221122 NWSS Data Dictionary_v3_1_5_20221122

20230322 NWSS Data Dictionary_v4.0.0_20230322

20230606 NWSS Data Dictionary_v5.0.0_20230606

'er.org/ for further details.

Itext of NWSS DCIPHER data submission, these are changes that we dded. In the context of NWSS DCIPHER data submission, these ma ext of NWSS DCIPHER data submission, these may be clarifications

Version			
0.1.0			
0.2.0			
0.3.0			
0.3.1			
0.4.0			

\mathbf{n}	E	\mathbf{n}
U	. ר.	U
-		~

1.0.0

1.0.1

1.0.2

1.0.3

1.0.6

1.0.7

2.1.0

2.1.1

3.0.0

3.1.0

3.1.2

3.1.3

3.1.4

3.1.5

ould cause submission (see note on Submission Requirements) of a y be additions of non-required fields, name changes to non-required of field descriptions, additions to value sets, or reorganization of the

Major changes

NA None

Replaced Pascal Case variable names with snake case variable names to align with bulk upload template variables

None

None

Changed "Required Field" value for the 'state' variable in the "Wastewater Treatment Plant" table from blank to "Required"

None

None

None

None

None

None

None

 New required fields (3): 'institution_type', 'sample_id', 'lab_id';
 Name changes to required fields (2): 'state' changed to 'wwtp_jurisdiction', 'reporting_state' changed to 'reporting_jurisdiction';

None

None

1. Addition of a new required field 'pcr_gene_target' which should contain the information previously captured in 'pcr_target'. 2. Renaming of fields: 'sars_cov2_units' renamed to 'pcr_target_units' 'sars_cov2_avg_conc' renamed to 'pcr_target_avg_conc' 'sars_cov2_std_error' renamed to 'pcr_target_std_error' 'sars_cov2_cl_95_lo' renamed to 'pcr_target_cl_95_lo' 'sars_cov2_cl_95_up' renamed to 'pcr_target_cl_95_up' 'sars_cov2_below_lod' renamed to 'pcr_target_below_lod'

1. Renaming of fields: 'pcr_target_ref' renamed to 'pcr_gene_target_ref'

none

none

none

none

1. Changing existing fields to required (5): 'epaid', 'pretreatment', 'pasteurized', 'major_lab_method', 'major_lab_method_desc'

1. Add new required field: 'site_id'

data file, if unaltered, to fail upload. These may include addition of r 1 fields, or changes to value sets, which will not prohibit upload but v = metadata.

Minor changes

NA 1. Changed to allow for sampling location not associated with treatment plant; 2. EPAID no longer required

None

None

A number of changes were made to reflect the shift from formbased data submission, which had built-in validation prior to submission, to CSV file submission to DCIPHER: 1. Recharacterized "Pre-populated value set" values as "Allowed values";

2. Added variable type column

Changed 'composite_freq' from "Required" (if composite) to "Not required"

None

None

None

None

None

None

1. Important description change relevant for upstream sites for 3 fields: 'county_names', 'other_jurisdiction', and

'sewage_travel_time' (these previously pertained to the treatment plant but have been changed to pertain to the sampling site, which is more relevant);

2. Removal and addition of values in value sets (>2 fields): 'sample_location' ("primary influent" and "primary sludge" changed to "wwtp"), 'rec_eff_target_name' ("bovine coronavirus" changed to "bcov vaccine"), other fields where similar values with the same meaning were submitted have been consolidated into one value, and will be indicated in QC checks;

3. Addition of non-required fields (3): 'pasteurized', 'time_zone', 'solids_separation'

4. Name changes to non-required fields (4):

'hum_frac_target_phys' changed to 'other_norm_name',

'hum_frac_target_phys_ref' changed to 'other_norm_ref',

'hum_frac_phys_conc' changed to 'other_norm_conc',

'hum_frac_phys_unit' changed to 'other_norm_unit'

None

1. Addition of non-required fields:

'qc_ignore' has been added to allow samples to be intentionally excluded from the DCIPHER QC report; 'analysis_ignore' has been added to allow samples to be intentionally excluded from DCIPHER data analysis; 'dashboard_ignore' has been added to allow samples to be intentionally excluded from the DCIPHER dashboard; 'major_lab_method' has been added to allow reporting jurisdictions to identify major lab methods in their data, which may be useful for determining groups of SARS-CoV-2 results that can be reasonably compared;

'major_lab_method_desc' has been added to allow a brief description of the rationale for assigning a 'major_lab_method'.

1. Repurposing of existing fields:

'pcr_target' will be repurposed from the field where PCR gene target information is submitted to now capturing what the overall PCR target was. For example, 'n1' was previously submitted to indicate the target gene for total SARS-CoV-2 was n1. Now that information will be captured in 'pcr_gene_target' and 'pcr_target' will be 'sars-cov-2'. For variants, you would put the variant target gene in 'pcr_gene_target' and 'pcr_target' would be the variant name (currently 'delta' or 'omicron').

None

none

none

none

none
1. Changes to value sets: 'pcr_target' now allows 'caur', 'NoV GI', 'NoVGII', 'FLUAV', 'FLUBV', 'NVO', 'Cjejuni', 'HAdV-F', 'Col', 'Gly', 'Bla', 'Integrase', 'Tet'. 'pcr_gene_target' now allows 'MCR-1.1', 'VanA-A', 'blaNDM', 'blaKPC', 'blaOXA-48', 'blaVIM', 'blaCTX-M-1', 'blaCMY', 'blaTEM', 'Class I Integrase', 'blaSHV', 'TetW', 'blaIMP', 'caur', 'NoV GI', 'NoV GII', 'InfA1', 'InfA2', 'InfB', 'Cjejuni', 'HAdv-F'. 'sample_type' now allows '72-hr time-weighted composite'. 'pasteurized' no longer allows [empty]. 'pretreatment' no longer allows [empty]. 'extraction_method' now allows 'exclusions based sample preparation (ESP)'

 Changes to dependencies of existing field: 'pcr_type' must be a type of dpcr if 'pcr_target' is <u>not</u> 'sars-covs-2', 'delta', 'omicron', '
Added 'epa_registry_id' field

1. Changes to value sets: 'extraction method' now allows 'sciencell viral rna isolation kit', 'thermo magmax viral/pathogen nucleic acid isolation kit'.

equired fields or name changes to required fields. will be flagged in on-platform QC checks.

Patches

NA

Description of treatment plant changed to be 'identifiable name'

Added red to data source color scheme to denote DCIPHER generated variables.

Clarified that state variable response for WWTP should be 2-letter abbreviation.

A number of changes were made to reflect the shift from formbased data submission, which had built-in validation prior to submission, to CSV file submission:

1. Changed "Forced column" to "Required Field";

2. Changed possible responses for "Required Field" to "Not required" and "Required";

3. Changed "DCIPHER Form Format", "DCIPHER Form Prepopulated Value Set", and "DCIPHER Form Data Validation" column names to denote Internal-Use only:

4. Made all values lower case (not a major change because all values are cast to lower during file parsing);

Changed 'pcr_target_ref' description to "A publication, website, or description of the PCR gene target used"

Changed "Allowed values" from "[any]" to "0-100" for 'industrial_input' to clarify percentage units

Changed "Tooltip / descriptive text" for standard error and confidence interval fields so that '-1' is only entered when these values cannot be calculated

Changed "Tooltip / descriptive text" for 'ntc_amplify' field to be more specific: "For qPCR, did any no-template controls on this instrument run have a Ct value less than 40? For ddPCR, did any no-template controls on this instrument run have 3 or more positive droplets?" This definition aligns with the definition of positivity given on the NWSS webpage for Data Reporting and Analytics, and aligns with guidance provided by a ddPCR instrument manufacturer. Added clarification to "Tooltip / descriptive text" for 'population_served' field to consider population estimates for upstream sampling locations: "if the sampling location is upstream of a treatment plant (i.e., sample_location = "upstream"), please use the estimated population served by the corresponding service area"

Added additional text and example to "Tooltip / descriptive text" for 'composite_freq' to clarify definition

1. Removed "Internal Use" columns;

2. Updated "Allowed values" column to make 'suggested' values more explicit (e.g., for fields that do not have a finite value set but have values that are commonly used) and to indicate where '-1' should be used to indicate unavailable data for required numeric fields;

3. Added "Notes" column to provide additional guidance, such as where appropriate to leave blank;

4. Clarified "Tooltip / descriptive text" for 'sars_cov2_std_error' and 'tss';

5. Changed "Display Variable Name" for 'population_served' for clarity

1. Additions to Value Sets:

'sample_type' now allows composite sample values for any integer duration from 1 to 30 hrs;

'flow_rate' now allows empty values to accommodate a 'sample_matrix' that is not a flowing liquid, such as "primary sludge" or "holding tank";

'concentration_method' now allows values that capture when solids and liquid are separated but both are tested, e.g., "membrane filtration with addition of mgcl2, membrane recombined with separated solids";

'pcr_target' now allows these additional values: "n", "s," "orf1a"; 'rec_eff_spike_matrix' now allows [empty] values; 'hum_frac_target_chem' now allows [empty] values; 'hum_frac_target_mic' now allows [empty] values;

'other_norm_name' now allows [empty] values;

'hum_frac_mic_unit' now allows [empty] values;

'hum_frac_chem_unit' now allows [empty] values;

'other_norm_unit' now allows [empty] values;

2. Description clarification:

'other_jurisdiction' has been further clarified to indicate that this refers to locations that exist outside of county boundaries; 'sample_id' has been clarified to indicate that PII must not be included in this field;

'lab_id' has been clarified to indicate that PII is discouraged from being included in this field;

the 'jurisdiction id' data type description has been modified to explicitly state that is is not case sensitive, which was previously and is still described in the note on all data types in the Read Me tab

1. Additions to Value Sets:

'extraction_method' now allows "chemagic viral dna/rna 300 kit", "trizol, zymo mag beads w/ zymo clean and concentrator", "4s method (https://www.protocols.io/view/v-4-direct-wastewaterrna-capture-and-purification-bpdfmi3n)"; 'pcr target' now allows "ddcov n" and "ddcov e" 1. Additions to Value Sets:

'rec_eff_spike_matrix' now allows "dewatered solids";

'flow_rate' now allows empty values when SARS-CoV-2 units are provided on a per mass sludge basis;

'concentration_method' now allows "innovaprep ultrafiltration"; 'extraction_method' now allows "qiagen qiaamp buffers with epoch columns";

'hum_frac_target_mic' and 'other_norm_name' both now allow "f+ rna coliphage" and "f+ dna coliphage";

2. Description clarification:

'pretreatment' has been clarified to indicate that this field pertains to both chemicals added to the sample and chemicals added as part of the treatment train upstream of the sample collection point, but that pasteurization should be specified in the 'pasteurized' field

1. Additions to Value Sets:

'extraction_method' now allows "zymo quick-rna fungal/bacterial miniprep #r2014";

'pcr_target' now allows "ip2 and ip4 combined";

'sars_cov2_below_lod' now allows [empty] values;

'concentration_method' now allows "ceres nanotrap";

2. Corrections to Value Sets (value yet to be used):

'reporting_jurisdiction' value for Arkansas has been corrected to "AR";

'extraction_method' value "qiange allprep dna/rna kit" has been corrected to "qiagen allprep dna/rna kit";

3. Dependent Fields clarification:

Dependent Fields for 'flow_rate' has been clarified to indicate that 'flow_rate' can have an [empty] value when SARS-CoV-2 RNA is measured on a per unit wastewater solids basis, rather than a per volume wastewater basis;

4. Dependent Fields change:

All uncertainty fields now allow [empty] values, regardless of the values submitted for the other uncertainty fields. In other words, the three fields 'sars_cov2_std_error', 'sars_cov2_cl_95_lo', and 'sars_cov2_cl_95_up' are now all allowed to be [empty];

5. Requirement dropped:

'sars_cov2_std_error', 'sars_cov2_cl_95_lo', 'sars_cov2_cl_95_up', and 'sars_cov2_below_lod' are no longer required.

1. Additions to Value Sets:

'concentration_method' now allows "aloh3 precipitation";

'rec_eff_target_name' now allows "heat inactivated sars-cov-2 virus";

'extraction_method' now allows "qiagen ez1 virus mini kit v2.0" and "thermo magmax microbiome ultra nucleic acid isolation kit"; 'sample_type' now allows composite sample values for any integer duration from 1 to 48 hrs;

'vs_rec_eff_target_name' now allows "bcov culture" and "brsv culture";

'extraction_method' now allows "zymo environ water rna kit/ zymo environ water rna kit (cat. r2042)";

'concentration_method' now allows "zymo environ water rna kit/ zymo environ water rna kit (cat. r2042)";

2. Description clarification:

'capacity_mgd' has been clarified to indicate that this should be the capacity for which the plant is permitted;

3. Description change:

'county_names' has been changed to ask for FIPS codes rather than names; it also now takes values for both counties and county equivalents;

'other_jurisdiction' has been deprecated, since all jurisdictions previously specified in this field should now be specified in the 'county_names' field. Submitting this field will not interfere with data upload, but this field will no longer be used;

4. Requirement dropped:

'other_jurisdiction' is no longer required;

1. Additions to Value Sets: 'vs_other_norm_name' now allows "rnase p"; 'vs_institution_type' now allows 'other_worksite' 'vs_pcr_gene_target' now allows 'delta' and 'omicron' genes targets. See 'vs_pcr_gene_target' for the updated valueset options 'vs_pcr_target' now <u>only</u> allows 'sars-cov-2', 'delta', and 'omicron' 2. Field description modifications: Many PCR fields have been modified to reflect the new PCR

Many PCR fields have been modified to reflect the new PCR targets now able to be submitted using the dynamic 'pcr_target' fields as opposed to the total SARS-CoV-2 specific descriptions.

None

1.Additions to Value Sets:

'vs_concentration_method' now allows 'membrane filtration with addition of mgcl3'

'vs_extraction_method' now allows 'luminultra wastewater extraction kit', 'qiaamp viral rna kit', 'trizol and RNA purificaton kit', 'trizol, garnet bead beating, alcohol precipitation', 'zymo quick-rna viral 96 kit #r1041', 'zymo quick-rna viral kit #r1035', 'qiagen qiaamp dsp viral rna mini kit'

'vs_pcr_gene_target' now allows 'wt214', 'ins214epe', 'del142-144', 'del156-157', 'del31-33'

'vs_pcr_type' now allows 'qiagen dpcr', 'fluidigm dpcr', 'life technologies dpcr', 'raindance dpcr', 'dpcr'

'vs_rec_eff_target_name' now allows 'murine hepatitis virus' 'vs_reporting_jurisdiction' now allows 'AP', 'CN', 'CVBPI', 'EBCI', 'GTBI', 'MAT', 'NPT', 'OMT', 'SBT', 'TRIT', 'ZP'

'vs_sample_type' now allows '120-hr passive sample', from '48-hr passive sample' to '1-hr passive sample' in one hour increments 2. Field description modified:

'pcr_gene_target' description modified to clarify different gene target concentrations should not be averaged

1. Additons to Value Sets: 'pcr_gene_target' now allows 'CDC N1 (GT-Digital)', 'CDC N2 (GT-Digital)', 'N2 (PREvalence)', 'E (PREvalence)'. 'rec_eff_target_name' now allows: 'MHV (PREvalence)', 'BCoV (GT-Digital)'. 'hum_frac_target_mic' now allows: 'PMMoV (GT-Digital)'

1. Additions to Value Sets: 'pcr_target' now allows: 'hMPXV', 'hMPXV Clade I', and 'hMPXV Clade II'. 'rec_eff_target_name' now allows: parapoxvirus'. 'pcr_gene_target' now allows: 'E9L-NVAR', 'G2R_G', 'G2R_WA', 'C3L', 'E9L-OPX3', 'B6R', 'B2R', and 'gtmol_hMPXV'

1. Additions to Value Sets: 'pcr_gene_target' now allows 'a67v/del69/del70'. Reporting_jurisdiction now allows: 'PBCI'.

1. Addition to Value Sets:'concentration_methods' now allows: 'water concentrating buffer (R2042-1)'; 'extraction_method' now allows 'monarch total RNA miniprep kit (new england biolabs) + onestep PCR inhibitor removal kit (zymo)'; 'pcr_gene_target' now includes 'p13L'; 'sample_type' now includes '72-hr time-weighted composite'. Clarifications of field or value set descriptions (2): 'zipcode', 'capacity_mdg', 'test_result_date', 'pcr_target_avg_conc', 'pcr_target_std_error', 'pcr_target_cl_95_lo', 'pcr_target_cl_95_up'
Added website to look-up NPDES ID and EPA Registry ID to 'Read Me' and to description in 'Metadata' tabs.
Removed space from "other long term care" value for 'institution type'