

DPCC Data Standard Reference for Virus Reagent v1.2

	Project_Identifier	Contributing_Institution	Reagent_Name	Sample_Identifier
Input Type	Text Field	Text Field	Text Field	Text Field
Definition	A unique Project Identifier generated by the DPCC by combining the Center-generated Project Code and a random 4-digit number	The institution code for the group that CREATED THE REAGENT. Must be one of the CEIRS institution codes assigned by NIAID.	The WHO strain name of the virus submitted as a reagent	Identifier assigned to the virus used to create the reagent
Format	Project_Code_XXXX Maximum length: 21 characters	Center three-letter code followed by three digits. Maximum length: 6 characters	Antigenic Type/Host of Origin/Geographical Origin/Strain Number/Year of Isolation (Subtype) rg-Antigenic Type/Host of Origin/Geographical Origin/Strain Number/Year of Isolation [rg details] (Subtype) Maximum length: 150 characters	Center-specific Allowed characters include alphanumeric, hyphen, and underscore: a-z, A-Z, 0-9, -, _ Maximum length: 50 characters
Value List	None	None	Text U	Text NA
Curation	The entry must be a Project Identifier value registered with the DPCC.	The entry must be an Institution Code value registered with the DPCC. Center 3-letter codes are case-sensitive and must be entered in all-caps.	WHO strain naming convention: Fields must be ordered as follows and separated with the '/' character: 1. The antigenic type (e.g., A, B, C, D) 2. The host of origin (e.g., swine, equine, chicken. For human-origin viruses, no host of origin designation is given.) 3. Geographical origin (e.g., Denver, Taiwan) 4. Strain number (e.g., 15, 7) 5. Year of isolation (e.g., 2009, 1934) 6. For influenza A viruses, the hemagglutinin and neuraminidase antigen description in parentheses (e.g., (H1N1), (H3N2)) If the strain is a reverse genetic virus, prefix Strain_Name with rg-. Provide reverse genetic details in brackets between year of isolation and subtype.	It should match with the Sample_Identifier provided in related data submissions.
Examples	SJCProj02_4001	SJC101	A/Hong Kong/1/1968 (H3N2), A/chicken/Fujian/4/2002 (H3N6), A/chicken/Fujian/4/2002 (HxNx), A/chicken/Fujian/4/2002 (mixed), A/swine/Iowa/233-56/2011 (H3N2v), A/duck/Alberta/35/1976 (H1N1), B/Hong Kong/432/2014, C/Texas/19876/2011, rg-A/Puerto Rico/8/1934 (H1N1), or rg-A/Egypt/N03072/2010 [PR8 internal R 6+2] (H5N1)	21145-001a
Notes			Reference: http://www.cdc.gov/flu/about/viruses/types.htm (HxNx) can be used in cases where a partial subtype has been determined (e.g., H5Nx). If there are mixed subtypes contained within a sample use A/chicken/Fujian/4/2002 (mixed) for Strain_Name or list multiple comma-separated Strain_Names: A/mallard/Alaska/2/2007 (H3N8), A/mallard/Alaska/2/2007 (H3N9). For other virus types, enter the common Strain_Name.	Enter NA if the reagent is not related to a surveillance submission.
Dependent Fields				
Validation	Project_Identifier should be a valid project identifier.	Must be an Institution Code value registered with the DPCC	Validate field length	Validate field length
Message Code	Error_9_PROJECT_NOT_FOUND	Error_1_INVALID_VALUE	Error_70_INVALID_FIELD_LENGTH	Error_70_INVALID_FIELD_LENGTH
Validation	Project_Identifier exists but user does not have permission to access or edit the project.		Host of origin element of strain name must be alphanumeric, underscore, dash, period, or single quote: a-z, A-Z, 0-9, -, ., ' ,	
Message Code	Error_4_DENIED_USER_ACCESS		Error_137_INVALID_HOST_SPECIES_STRAIN_TEXT	
Validation			Geographical origin element of strain name must be alphanumeric, underscore, dash, period, or single quote: a-z, A-Z, 0-9, -, ., ' ,	
Message Code			Error_138_INVALID_LOCATION_STRAIN_TEXT	
Validation			Strain number element of strain name must be alphanumeric, underscore, dash, or period: a-z, A-Z, 0-9, -, . ,	
Message Code			Error_139_INVALID_STRAIN_NUMBER_STRAIN_TEXT	
Validation			Year of isolation element of strain name must be 4-digit year.	
Message Code			Error_110_INVALID_STRAIN_YEAR	
Validation			If strain is influenza A, subtype must be present as text between parentheses.	
Message Code			Error_112_INVALID_STRAIN_SUBTYPE	
Validation			Brackets cannot be present if strain does not have rg- prefix.	
Message Code			Error_119_INVALID_STRAIN_BRACKETS	
Message Code				

	Passage_History	Parent_Sample_Identifier	Parent_Strain_Name
Input Type	Text Field	Text Field	Text Field
Definition	Description of the passage history of the virus strain	A series of identifiers referencing the parent virus or viruses used to create the reagent	The name of the parent strains used to create the reverse genetic or reassortant virus strain
Format	Text Maximum length: 50 characters	Maximum of eight comma-separated Sample_Identifier. Allowed characters include alphanumeric, hyphen, and underscore: a-z, A-Z, 0-9, -, _ Maximum length: 800 characters	Eight comma-separated Strain_Names Antigenic Type/Host of Origin/Geographical Origin/Strain Number/Year of Isolation (Subtype) Maximum length: 800 characters
Value List	Text X	Text U NA	Text U NA
Curation	To indicate a change in passage history at one institution, combine passage host and number: E5C2 To indicate a change in passage history for samples between users or institutions, separate passage host and number with a slash '/': E5/C2	Each Sample_Identifier provided must match with a Sample_Identifier provided in related data submissions. For virus reagents derived from a surveillance sample, provide the Sample_Identifier of the surveillance record. For reverse genetic or reassortant virus reagents, provide the Sample_Identifier associated with each gene segment in the following comma-separated order: PB2, PB1, PA, HA, NP, NA, MP, NS	The entry must list a value for each of the eight gene segments. Strain_Names for each gene segment must be listed in the following comma-separated order: PB2, PB1, PA, HA, NP, NA, MP, NS Strain_Names must follow WHO strain naming convention.
Examples	E3, E2C2, EX, or E5/C2	13250,13250,U,21145,13250,21145,13250,13250, or NA	A/Puerto Rico/8/1934 (H1N1),A/Puerto Rico/8/1934 (H1N1),U,A/Puerto Rico/8/1934 (H1N1),A/Puerto Rico/8/1934 (H1N1),A/Vietnam/1203/2004 (H5N1),A/Puerto Rico/8/1934 (H1N1),A/Puerto Rico/8/1934 (H1N1), or NA
Notes	Indicate lab host and number of passages: E = Embryonated chicken eggs C = MDCK cells S = MDCK-SIAT cells M = Monkey kidney cells For other hosts, write out common name and include passage number e.g., Mouse2, E(duck)1, 293T1. Use X if the host or passage number is unknown, e.g., EX, CXE2. Enter X if passage history information is unknown. Enter NA if virus was not passaged. Additional information relating to viral passaging can be entered in the Comments field.	If the reagent is a reverse genetic or reassortant virus, eight comma-separated entries must be present. If an individual Sample_Identifier is unknown, enter U for the specific gene segment. Enter NA once if the strain is not a reverse genetic or reassortant virus.	The Strain_Name for each segment of the reverse genetic or reassortant virus must be named. If an individual Strain_Name is unknown, enter U for the specific gene segment. Enter NA once if the strain is not a reverse genetic or reassortant virus.
Dependent Fields		Parent_Strain_Name	Parent_Sample_Identifier
Validation	Validate field length	Validate field length	Validate field length
Message Code	Error_70_INVALID_FIELD_LENGTH	Error_70_INVALID_FIELD_LENGTH	Error_70_INVALID_FIELD_LENGTH
Validation		Fields Parent_Sample_Identifier and Parent_Strain_Name must have same number of values	Fields Parent_Sample_Identifier and Parent_Strain_Name must have same number of values
Message Code		Error_63_INVALID_NUMBER_ENTRIES	Error_63_INVALID_NUMBER_ENTRIES
Validation			Count number of segments. It should be 8
Message Code			Error_1_INVALID_VALUE
Validation			If Parent_Sample_Identifier is NA then Parent_Sample_Strain should be NA
Message Code			Error_1_INVALID_VALUE
Validation			Host of origin element of strain name must be alphanumeric, underscore, dash, period, or single quote: a-z, A-Z, 0-9, -, ., ', _
Message Code			Error_137_INVALID_HOST_SPECIES_STRAIN_TEXT
Validation			Geographical origin element of strain name must be alphanumeric, underscore, dash, period, or single quote: a-z, A-Z, 0-9, -, ., ', _
Message Code			Error_138_INVALID_LOCATION_STRAIN_TEXT
Validation			Strain number element of strain name must be alphanumeric, underscore, dash, or period: a-z, A-Z, 0-9, -, ., _
Message Code			Error_139_INVALID_STRAIN_NUMBER_STRAIN_TEXT
Validation			Year of isolation element of strain name must be 4-digit year.
Message Code			Error_110_INVALID_STRAIN_YEAR
Validation			If strain is influenza A, subtype must be present as text between parentheses.
Message Code			Error_112_INVALID_STRAIN_SUBTYPE

	Mutations	BEI Number	Contact Name
Input Type	Text Field	Text Field	Text Field
Definition	A description of the mutation(s) or non-coding change(s) made in each segment of the reverse genetic or reassortant virus	The identification number assigned to the reagent by BEI	The name of the point-of-contact for reagent requests or the name of the BEI depositor
Format	An HGVS-formatted list describing sequence changes Maximum length: 200 characters	BEI Number Maximum length: 20 characters	FirstName LastName FirstName MI. LastName Maximum length: 50 characters
Value List	Text NA	Text NA	None
Curation	The entry must use the HGVS reference for annotation guidelines to describe mutations, insertions, or deletions in any gene segment order. http://www.hgvs.org/mutnomen/quickref.html	The entry must be a valid BEI number.	None
Examples	HA:p.[N220K; Q222L],NS:p.[H155del],PB2:p.[E627K]	NR-12277	Jane Smith, or Davey M. Jones
Notes	<u>Format for amino acid mutation: protein_name.p.[mutation_information]</u> For a single mutation, list the original amino acid, its position, and the new amino acid in brackets: PB2:p.[E627K] For a multiple mutations within a single protein, separate individual mutations by a semicolon: HA:p.[N220K;Q222L] For a deletion within a single protein, list the amino acid, its position, and 'del': NS:p.[H155del] For a stop codon, use an asterisk: NA:p.[K471_*472insHHHHHH] <u>Format for non-coding mutation: gene_segment_name.c.[mutation_information]</u> For a single mutation: enter the position, the original nucleotide, >, and the new nucleotide: PB2:c.[273C>T] For multiple mutations within a single gene, separate individual mutations by a semicolon: PB1:c.[273C>T;274A>G;275G>T] For a deletion: enter the position, del, and the nucleotide deleted: NA:c.[73delT] To list both amino acid and non-coding changes, comma-separate the individual mutation information in any order: HA:p.[N220K;Q222L],HA:c.[273C>T;274A>G;275G>T],PB1:c.[273C>T;274A>G;275G>T] Enter NA for non-reverse genetic or reassortant viruses, or viruses with no known mutations.	Enter NA if the reagent was not deposited with BEI.	
Dependent Fields			
Validation	Validate field length	Validate field length	Validate field length
Message Code	Error_70_INVALID_FIELD_LENGTH	Error_70_INVALID_FIELD_LENGTH	Error_70_INVALID_FIELD_LENGTH
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	Contact_Email	Availability	Publication_PMID	Comments
Input Type	Text Field	Text Field	Text Field	Text Field
Definition	Email address for the point-of-contact	Whether the reagent is available to share with other investigators on the reagents page of the CEIRS Public Website	The PubMed Unique Identifier (PMID) for the publication in which the reagent was published.	Text describing anything else of interest related to the submission
Format	Text@domain.com Maximum length: 50 characters	Text Maximum length: 1 character	PMID Maximum length: 50 characters	Text Maximum length: 2000 characters
Value List	None	Y N	Text NA	Text NA
Curation	The entry must be a valid email address.	The entry must be one and only one member of the Value List.	The entry must be a valid PMID number: 7 or 8-digit number with no leading zeros. http://www.nlm.nih.gov/bsd/mms/medlineelements.html#pmid	None
Examples	jsmith@email.edu	Y	19465683, or NA	NA
Notes		Y = Yes N = No	If the reagent was referenced in multiple publications, comma-separate the individual PMIDs. Enter NA if the reagent was not referenced in a publication.	If there are no comments, enter NA.
Dependent Fields				
Validation	Validate field length	Field value should be one of valid values as in list.	Validate field length	Validate field length
Message Code	Error_70_INVALID_FIELD_LENGTH	Error_1_INVALID_VALUE	Error_70_INVALID_FIELD_LENGTH	Error_70_INVALID_FIELD_LENGTH
Validation	Must be a valid email address.		Must be a valid PMID.	
Message Code	Error_114_INVALID_EMAIL		Error_96_INVALID_PMID	
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