



Global Influenza Hospital-based Surveillance Network (GIHSN)

Core Protocol

Rationale

To establish the *[specify country/city]* branch of the global influenza hospital-based surveillance network. The Global Influenza Hospital Surveillance Network (GIHSN) is a platform able to generate strong epidemiological and medical evidence on **influenza severity** and to support vaccine strain selection through **timely sharing of clinical and laboratory data**. The GIHSN is a network of not-for profit institutions coordinating local hospitals in several countries following the same core protocol¹.

The GIHSN is a unique hospital active surveillance network using a standard protocol complementary to WHO GISRS, offering:

- The largest yearly case series of patients hospitalized with influenza worldwide from all age groups for both NH and SH allowing to better understand flu severity and related risk factors;
- A strong opportunity to inform WHO vaccine strains selection by linking clinical data with viral genome sequencing information;
- An alert system in case of pandemic/strain mutation, contributing to improve countries response and international collaboration.

Note: Main parts requiring country/site adaptations are specified in *blue*

Study objectives

1. Support international capacities developed through the Global Influenza Surveillance and Response System (GISRS) of laboratories to increase the availability of clinical information linked with genetic sequencing of influenza strains to expand the support of the biannual vaccine strain selection process of

¹ This core protocol has been adapted from the initial version developed by Joan Puig-Barberà (Centre for Public Health Research, Valencia, Spain).



the WHO's formal recommendation for the composition of human influenza vaccines.

2. Link clinical and virological data in Influenza positive patients with an emphasis on severe cases and on vaccine failure.
3. To quantify the distribution of the different influenza strains (A/H1N1, A/H3N2, B/Yamagata, B/Victoria) among these severe cases

Design: Prospective epidemiological **active** surveillance study



Study setting and population

The study will take place in *[specify number]* hospitals. *[Describe further the hospitals: names, catchment area, specialty, size]*. The study period will be organized to cover the main influenza season ie. *[complete with planned start and end date for the study – can be informed by virologic surveillance data]*

This study will focus on *[select population category among the following options: (i) all ages, (ii) elderly (60+), (iii) adults (18+), (iv) children (<18) (v) high risk groups (to be further defined)]*

Eligibility criteria

Enrolment will be based on:

- Patients with an acute process
- Patients whose indication for admission was any of a predefined set of conditions, described as possibly associated with a recent influenza infection (see table 1).
- In this case, *[a study nurse, doctor...]* will identify by hospital admission registries, chart review or available records, **all** eligible patients hospitalized in the previous 72 hours and has stayed in hospital for at least 1 night (therefore a patient admitted before midnight of the previous day).

Table 1. Admission diagnoses possibly associated with an influenza infection.
International Classification of Diseases Code version 9 and 10.

For Patients less than 5 years old	ICD 9 Codes	ICD 10 Codes
Acute upper or lower respiratory disease	382.9; 460 to 466	J00-J06, J20-J22
Dyspnea, breathing anomaly, shortness of breath, tachypnea (polypnea)	786.0; 786.00; 786.05-786.07; 786.09; 786.9	R06.0, R06, R06.9, R06.3, R06.00, R06.09, R06.83, R06.02, R06.82, R06.2, R06.89
Acute asthma or exacerbation	493.92	J45.901
Pneumonia and influenza	480 to 488	J09-J18
Acute respiratory failure	518.82	J96
Acute heart failure	428-429.0	I50-I50.9; I51.4
Myalgia	729.1	M79.1



Altered consciousness, convulsions, febrile convulsions	780.01-780.02; 780.09; 780.31- 780.32	R40.20, R40.4, R40.0, R40.1, R56.00, R56.01
Fever or fever unknown origin or non specified	780.6-780.60	R50, R50.9
Cough	786.2	R05
Gastrointestinal manifestations	009.0; 009.3	A09.0; A09.9
Sepsis, Systemic inflammatory response syndrome, not otherwise specified	995.90-995.94	R65.10, R65.11, R65.20, A41.9
Nausea and vomiting	078.82; 787.0; 787.01-787.03	R11; R11.0; R11.10 - R11.12; R11.2

For Patients 5 years old or older	ICD 9 Codes	ICD 10 Codes
Acute upper or lower respiratory disease	382.9; 460-466	J00-J06, J20-J22, H66.90
Acute myocardial infarction or acute coronary syndrome	410-411 and 413- 414	I20-I25.9
Acute asthma or exacerbation	493.92	J45.901
Acute Heart failure	428-429.0	I50-I50.9; I51.4
Pneumonia and influenza	480-488	J09-J18
Bronchitis and exacerbations of Chronic Pulmonary Obstructive disease	490, 491.21 and 491.22,	J40; J44.0; J44.1
Acute respiratory failure	518.82	J96
Myalgia	729.1	M79.1
Acute metabolic failure (diabetic coma, renal dysfunction, acid-base disturbances, alterations to the water balance)	250.1- 250.3; 584- 586; 276-277	E11.9, E10.9, E11.65, E10.65, E10.11, E11.01, E10.641, E11.641, E10.69, E11.00, E10.10, E11.69, N17.0, N17.1, N17.2, N17.8, N17.9, N18.1, N18.2, N18.3, N18.4, N18.5, N18.6M N18.9, N19, E87.0, E87.1, E87.2, E87.3, E87.4, E87.5, E87.6, E87.70, E87.71, E87.79,



		E86.0, E86.1
Altered consciousness, convulsions, febrile convulsions, syncope and collapse	780.01-780.02; 780.09; 780.2; 780.31-780.32	R40.20, R40.4, R40.0, R40.1, R55, R56.00, R56.01
Dyspnea/respiratory abnormality	786.0	R06.0, R06-R06.9
Respiratory abnormality	786.00	R06.9
Shortness of breath	786.05	R06.02
Respiratory abnormality not otherwise specified	786.09	R06.3, R06.00, R06.09, R06.83
Respiratory symptoms/chest symptoms	786.9	R06.89
Fever or fever unknown origin or non-specified	780.6-780.60	R50, R50.9
Cough	786.2	R05
Sepsis, Systemic inflammatory response syndrome	995.90-995.94	R65.10, R65.11, R65.20, A41.9

Inclusion criteria

Patients 5 years old and more will be included in the study if they refer to seven days or less antecedent of a community onset influenza like-illness (see definition in table 2).

Table 2. Modified European Centre for Diseases Control definition of influenza like-illness (ILI)

Combination of:

- at least one of the following four systemic symptoms (ICD-9-CM code): Fever or feverishness (780.6), headache (784.0), myalgia, (729.1) or malaise (780.79);
- at least one of the following three respiratory symptoms (ICD-9-CM code): b) Cough (786.2), sore throat (787.2) or shortness of breath (786.05), Nasal Congestion (478.19)

Patients less than 5 years will be included if indications for admission (table 1), occurred within seven days or less between the beginning of symptoms and admission to hospital.



Swabbing procedures

A nasopharyngeal swab for all patients and in addition a pharyngeal swab for patient 14 years of age or older and a nasal sample for children (less than 14 years old) will be obtained from each patient in case they comply with inclusion criteria and give consent.

Sample management and laboratory procedures

All samples will be kept at -20°C until sent to reference laboratory. Multiplex real-time RT-PCR will be performed on the samples to detect the presence of:

- influenza A (H1N1n and H3N2), influenza B (B/Yamagata, B/Victoria)

Notice: the GIHSN goals are related to influenza epidemiology [*If testing for other respiratory viruses is performed, the following can be considered coronavirus, metapneumovirus, bocavirus, respiratory syncytial viruses, adenovirus, parainfluenza viruses, rhinovirus. Analysis for respiratory viruses other than influenza can be carried out after the study ends if samples are stored appropriately*].

Sequencing must be done of selected positive specimens according to agreed schedule (see table 3). If the site has no capacities to generate genetic sequence data (GSD), the site may ship its specimens to the GIHSN sequencing platform at the National Influenza Center in Lyon, France, under the Terms of Reference for sharing materials in GISRS. Shipments are organized by OpenHealth.

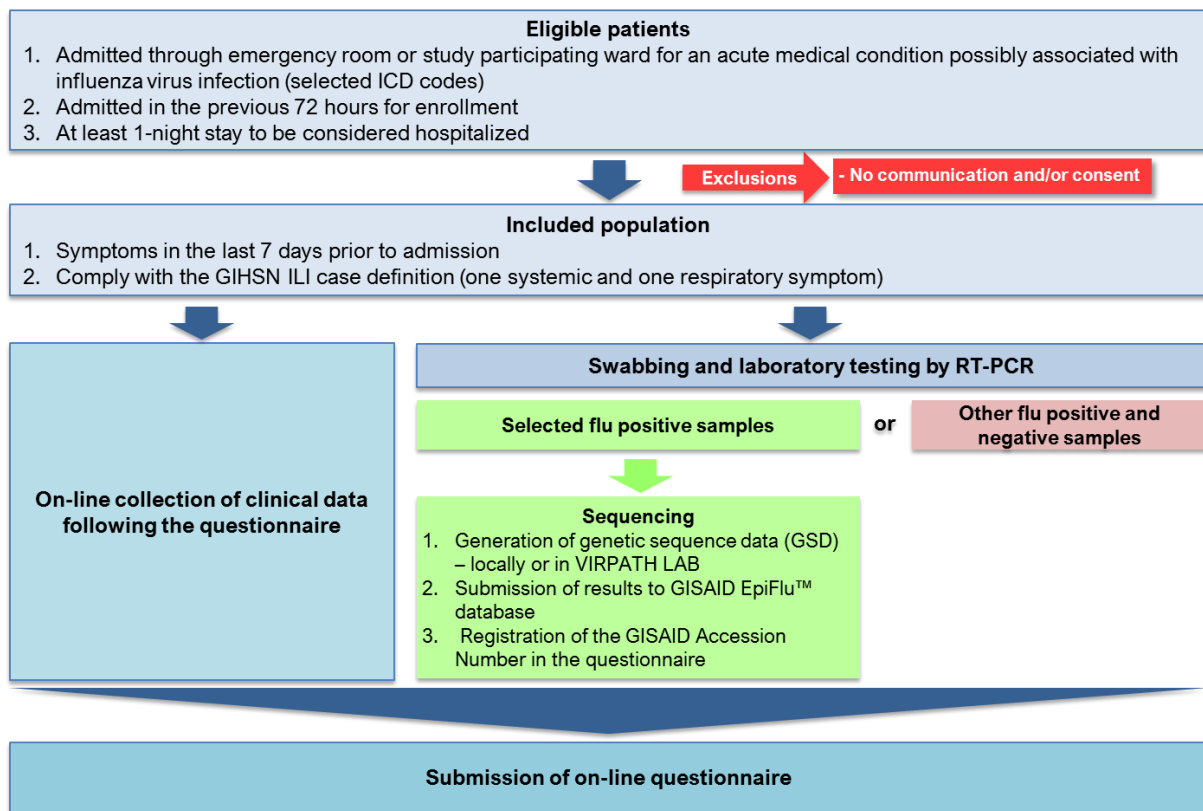
All sites must submit GSD data to the GISAID EpiFlu™ database (http://gisaid.org/EPI_ISL/123456).

Table 3. Sequencing scheme for all samples (patients of all ages):

<i>Hemisphere</i>	<i>Early season</i>	<i>ICU/deaths and vaccine failures</i>	<i>Samples per month</i>
<i>Northern</i>	<i>all samples until 15 January</i>	<i>All</i>	<i>10-30 (during season)</i>
<i>Southern</i>	<i>all samples until 15 July</i>	<i>All</i>	<i>10-30 (during season)</i>
<i>Intertropical</i>	<i>NA</i>	<i>All</i>	<i>5-15 (all year)</i>



Study process



Sample size, data collection and analysis

Sample size

The number of **sequenced** laboratory confirmed influenza cases we expect per site is around 50-100. The number of hospitals (**study setting and population**) to involve in this study should be planned to reach the agreed minimum target.

Data collection

Trained [study nurses, doctor....] collect relevant information by a combination of face-to-face interviews of patients and attending physicians, and by reviewing clinical records (refer to both questionnaires, younger than 5 years old and 5 years and older).

Influenza vaccination status is obtained by asking the patient (or representative) if he or she had received the influenza vaccine of the current season, the date of vaccination, and if the vaccine had been administered at least two weeks before the onset of symptoms. Whenever possible, this information will be validated by existing



registers, vaccination cards or through contacting the place where the vaccine was administered.

Data analysis

A descriptive analysis of the frequency of laboratory results by epidemiological week, age group, and comorbidities will be conducted. If possible, hospitalization rates by age group according to population denominators (based on the size of the hospital catchment area) and for each different virus will be also estimated.

Ethical considerations

Approval by the local Research Ethics Committee will be obtained. The confidentiality legislation and requirements in the handling in personal information will be strictly followed. Informed written consent will be required for enrolment. No intervention, apart the nasopharyngeal, nasal and pharyngeal sampling is associated with the study.

Good Epidemiological Practice procedures will be implemented in all the study process.