



HelicsWin.Net 2.3 User manual

Point prevalence survey of healthcare-associated infections and antimicrobial use in European acute care hospitals

Surveillance of healthcare-associated infections in European intensive care units

Surveillance of *Clostridium difficile* infections in European hospitals

Edition notice

This document applies to HelicsWin.Net (HWN) version 2.3.

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Document's intended audience and purpose

The document is intended for hospital staff that use HelicsWin.Net to collect or administer data collected using ECDC's HAI-Net surveillance of healthcare-associated infections methods (HAI-Net PPS and ICU).

This document describes the HelicsWin.Net v2.3 user interface, and provides users with step-by-step instructions in its use, and conceptual information about data storage.

About HelicsWin.Net

HelicsWin.Net is a software application developed for the manual entry of data of the Healthcare-Associated Infections surveillance Network (HAI-Net).

The current version includes three surveillance modules:

- HAI-Net PPS: point prevalence survey (PPS) of healthcare-associated infections and antimicrobial use in European acute care hospitals (Protocol version 5.2), and
- HAI-Net ICU: surveillance of healthcare-associated infections in European intensive care units (ICUs) (Protocol version 2.0), and
- HAI-Net CDI: surveillance of *Clostridium difficile* infections (CDI) in the EU (pilot protocol version 2.2).

HelicsWin.Net enables local users, typically in a hospital, to collect surveillance data at the hospital and ward levels; these data are stored internally by HelicsWin.Net, but the data can be exported to other applications in a variety of formats, including formats compatible with Microsoft Access (.mdb) and Microsoft Excel (.csv).

For nominated representatives of the EU Member States only, a TESSy compatible format is provided to facilitate the importation of data to The European Surveillance System (TESSy) database.

Background

The name *HelicsWin.Net* originates from the Microsoft Access application *HelicsWin* developed by the former European HELICS (Hospitals in Europe for Infection Control through Surveillance) network for the surveillance of HAI. HelicsWin was originally developed as part of an ECDC contract for further Hospital Software Support for the European Surveillance of HAI, which included *HelicsWin for the surveillance protocols Surveillance of healthcare-associated infections in Intensive Care Units (ICU) and Surveillance of surgical site infections (SSI).*

HelicsWin.Net was originally developed by the ICT department of the Scientific Institute of Public Health, Brussels, Belgium under contract ECD.2218 and its amendment ECD.2764 until September 2011. In September 2011, development of HelicsWin.Net was transferred to ECDC.

Technology

HelicWin.Net is a standalone application developed in Microsoft .Net framework. Data are stored in an .mdb file that is in Microsoft Access format. This file is stored on the computer on which HelicsWin.Net is installed.

Users should also note that:

- HelicsWin.Net is supplied free of charge and can be freely distributed to participating hospitals.
- The text displayed on the data entry forms can be translated to meet language requirements of the participating hospitals.
- HelicsWin.Net can be installed and run from a server, but simultaneous users are not supported, i.e. only one user can run the software at any one time.
- The current version does not require Microsoft Access to be installed on the computer on which it runs.

 Microsoft .Net Framework version 3.5 SP1 or later (available free of charge from Microsoft) is required.

HelicsWin.Net versions

HelicsWin.Net has been published in incremental versions from 1.0 to the current version 2.3.

What's new in HelicsWin.Net version 2.3

Version 2.3 is a major release with the following features:

- ✓ Addition of the HAI-Net CDI pilot module.
- ✓ The HAI-Net PPS module has been updated to match the current ECDC PPS 2016-2017 protocol

What's new in HelicsWin.Net version 2.0

Version 2.0 is a major release with the following features:

- ✓ Addition of the HAI-Net ICU surveillance module for the surveillance of healthcare-associated infections in European intensive care units. The current version contains the variables for the pilot study of the new HAI-Net ICU surveillance protocol. It does not yet contain the features **TESSy export** and **Reports**, which will be added after the pilot study.
- ✓ The **General** tab in the PPS Hospital Surveys form has been split into three tabs (Form H1, H2 and H3) to allow for more detailed and accurate surveillance data to be recorded.
- ✓ The **General** tab in the Ward PPS Data form has been split into two tabs (Form W and Form W (2)) to allow for more detailed and accurate surveillance data to be recorded.
- ✓ Functionality of version 1.3.8 preserved.

What's new in HelicsWin.Net version 1.3.8

Version 1.3.8 was a minor release with the following features:

- ✓ Major functionality of version 1.3.0 preserved.
- ✓ Maintains compatibility with existing v1.3.0 data; new data is automatically converted to the new format on import.
- ✓ New reporting facilities including both predefined and user-defined report templates.
- ✓ Reports can be printed directly or exported as CSV files.
- ✓ Improved data merge facilities. Now you can merge HelicsWin.Net data from different computers, either from the same ward or from different wards in the same hospital. You can then create reports from the merged data.
- ✓ Enhanced data entry and print options.

What's new in HelicsWin.Net version 1.3.0

The key features introduced in v1.3.0 were:

- ✓ Keyboard shortcuts
- ✓ Data quality checks during data entry
- ✓ Separate data quality check function before export
- ✓ Printable search lists for all levels
- ✓ Antimicrobial brand lookup tool
- ✓ Translation possible for all forms, messages and different kinds of controls Integration of PPS validation protocol variables
- ✓ TESSy export for Standard and Light protocol
- ✓ Improved speed

✓ Integrity of the database is ensured; if a user changes the ward ID or survey date, the corresponding data in the underlying records (for example. patient data) is automatically updated.

Related documents

For information about how to complete the forms in this application, please refer to the following documents:

HAI-net CDI:

- European Centre for Disease Prevention and Control. European surveillance of Clostridium difficile infections, Surveillance protocol version 2.2 (pilot study).
- Forms 1215-TED-HAI-Net-CDI-pilot-forms.xlsx: Master copies of data collection forms for the CDI protocol v2.2 (in editable format for translation purposes).

HAI-Net ICU:

- European Centre for Disease Prevention and Control. Surveillance of healthcare-associated infections and prevention indicators in European intensive care units HAI-Net ICU protocol version 2.0 (pilot study).
- Forms 0515-TED-HAI-Net-ICUv2-pilot-forms.xls: Master copies of data collection forms for the ICU protocol v2.0 (in editable format for translation purposes).

HAI-Net PPS:

- European Centre for Disease Prevention and Control. Point prevalence survey of healthcareassociated infections and antimicrobial use in European acute-care hospitals – protocol version 5.2. Stockholm: ECDC; 2015.
- Forms 0512-TED-PPS-HAI-antimicrobial-use-forms.PPT: Master copies of data collection forms for PPS protocol v5.2 (in editable format for translation purposes).

These documents are available from within the software, under Settings > About > *Documentation* or in the Documentation subfolder of the installation folder of the HelicsWin.Net software (by default C:\HWN2).

HelpDesk

For any questions, please refer to your National HAI surveillance (and/or PPS) coordinating centre that may refer questions to mailto:HAI-Net@ecdc.europa.eu.

Feedback

ECDC welcomes any feedback from users to help us to improve future versions of this software and documentation. Please send your comments to the HelpDesk.

Installation

This section describes the installation requirements and procedures for HelicsWin.Net.

Installation Requirements

The following requirements must be met on the PC on which you are installing the HelicsWin.Net software:

- Windows XP or later Personal computer running Windows XP or later.
- Administrator rights You must have administrator rights on your PC.

Note: If you do not have administration rights on your computer, contact your system administration for further help.

• **Microsoft .NET Framework 3.5 SP1** or later (.NET 3.5) must be installed on the PC on which HelicsWin.Net runs. The installation program checks for the presence of this software component and cannot complete without it.

If you were able to run the previous version of HelicsWin.Net 1.3 on the same machine, you will be able to install the new version.

If your PC does not already have .NET 3.5, it is installed automatically during HelicsWin.Net installation, providing the PC is connected to the internet and downloads are allowed.

If your PC does not have internet access...

Check whether.NET 3.5 or later is already installed (see below).

Alternatively, if your PC is not connected to the internet, and you don't have .NET, you can download the .NET software to another PC (that does have internet access) and copy the .NET installation files to portable memory device, such as a USB stick, transfer the device to your PC, and install .NET from there. You can download this software from this URL:

http://www.microsoft.com/en-us/download/details.aspx?id=22

Runtime Requirements

Access permissions

You do not need administrator rights to your PC to run the application once it is installed, but you must have full access (read/write/execute permission) to the installation folder, which by default is C:\HWN2.

If you install HelicsWin.Net in a different folder, you will need full access to that folder.

Microsoft Access

Although the application database HelicsWinNet.mdb is in Microsoft Access format, you do not need Microsoft Access installed on your computer to run HelicsWin.Net.

You can open the database files in Microsoft Access, but you could also use other compatible software such as Microsoft Excel.

.NET

To check whether .NET 3.5 SP1 is installed on your PC

In **Windows 7**, do the following:

- Click Start > Control panel
 The control panel opens.
- 2. Click Programs.
- 3. Click **Programs and Features** and wait until the installed programs list is populated.
- **4.** Scroll down the list to the Microsoft entries.

If **Microsoft .NET Framework** is in the list, it is installed. If the version is **3.5 SP1** or higher, you have the correct version, and you can install HelicsWin.Net.

If the correct version of .NET is not installed, the included setup.exe file installs it for you, although you must have administration rights on your computer (see below).

Alternatively, to install .NET Framework 3.5 SP manually, go to this link and follow the on-screen instructions:

http://www.microsoft.com/en-us/download/details.aspx?id=22

In **Windows XP**, do the following:

Click Start > Settings > Control panel
 The control panel opens.

- 2. Click Add or remove programs and wait until the installed programs list is populated.
- **3.** See step 4 above.

Installing HelicsWin.Net

The application comes as two files, one of which is an .msi file that contains the full application. To install HelicsWin.Net 2.3, you must have administration rights to your PC.

HelicWin.Net.msi	Installation files Administration rights required
setup.exe	Installer program

To install HelicsWin.Net 2.3:

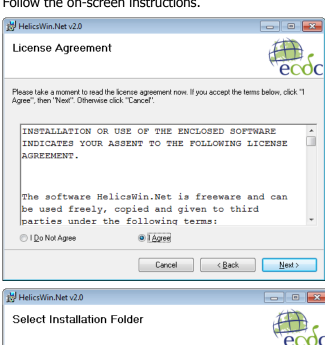
 Download the HelicsWin.Net zip file to your PC from the ECDC website at: http://ecdc.europa.eu/en/healthtopics/Healthcare-associated_infections/HELICSWin-Net/ Pages/HELICSWinNet-download-page-HWN.aspx

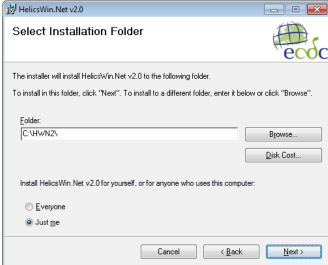
- 2. Open the zip file and extract the installation files to a folder to which you have full access.
- **3.** Make sure that after extracting the files, both installation files are in the same folder.
- 4. Click setup.exe.

The installer opens.



5. Follow the on-screen instructions.





6. By default, the installer creates a shortcut to HelicsWin.Net on your desktop. Simply click the shortcut icon to open the HWN application. The icon also appears in your taskbar when HelicsWin.Net is running.

The application files are installed, by default, in the folder C:\HWN2. Table 1 describes the purpose of each file.

Table 1: Installed files

File or folder	Contents
DatabaseBackups	Database backup file, for example, files saved by a merge operation. This folder is not visible immediately after installation, but is created automatically by the application when needed.
Documentation	User guide; HAI-Net PPS and ICU main protocols and forms.
Log	Log files: level of detail depends on user's specification.
Res	Documents enabling view of antimicrobial resistance markers and codes and ECDC's Privacy Policy from within the software.
HelicsWin.Net.Common.dll HelicsWin.Net.Components.Log.dll HelicsWin.Net.Helpers.dll	Application extension files – needed for HelicsWin.Net to work.
HelicsWin.Net.exe	Executable for HelicsWin.Net.
HelicsWin.Net.exe.config	Configuration file
license.rtf license_3rd_party_notices.rtf	License files
HelicsWinNetCDI.mdb HelicsWinNetICU.mdb HelicsWinNetPPS.mdb	Database file: HelicsWinNetXXX.mdb: is initially the empty database (Microsoft Access format) for this application, where XXX represents the module name (CDI, ICU, or PPS). These files are the most important in the application, because they contain all the survey data entered to date. Recommendation: make regular backups of these files.
Reference.mdb	Database file containing values and labels (Microsoft Access format).
Translation.mdb	Database file containing translation data (Microsoft Access format).

Warning:	Manually changing the structure, or the data, in any of the .mdb databases — for example, within MS Access — may cause the programme to stop working. If you do decide to make manual changes to the database files, always make a backup copy of the database first.
Important:	The software must be installed in a path (folder) where the user has write rights/access (for example, C:\HWN2), otherwise save errors may occur.
	Any existing data will be overwritten when copying new .mdb files! If applicable, make a backup of your existing data first (HelicsWinNet.mdb for the PPS database, Traduction.mdb for translations).

Network installation

You can also install HelicsWin.Net on a network drive in the hospital and then users can run the application from there. Users must have write access to the installation folders.

Network installation can be used when data needs to be entered from different wards; this may be preferred to having to work with multiple local ward copies (and databases); however, with the merge facility, it is relatively easy to merge these data into a single database for the hospital as a whole (see *Merging data*).

It is not possible to enter data from two or more computers simultaneously into the same database.

Getting started

To launch the software, open installation folder (default is HWN2) and run the file *HelicsWin.Net.exe*.

1. From your desktop, click the HWN2 icon on your desktop.



or

1. Select **Start** > **My Computer** > **System** (C:) > **HWN2**> **HelicsWin.Net.exe**.

The splash screen appears:



2. Click **Continue** to go to the first (login) form.



- 3. Select the surveillance module (HAI-Net CDI, ICU, or PPS) from the **Module** drop-down list.
- **4.** Select your country and language. You can add a language by translating the software.

The first time anyone uses this installation, the default password is *helics*. You will be prompted to change this the next time you log in. You can change your password in **Settings** later on.

Caution If you change your password, make sure you can remember it because there is no easy way to access your database files without that password.

The default language on installation is English. If you change to another language while logging in, HelicsWin.net will be remember your change for subsequent logins.

5. Click Login.

If you have not already defined any hospitals, you must define one now.

6. Click the **Add item** icon **.**

The fields in the upper part of the form open for editing.

- **7.** Enter the hospital code—as provided by your National HAI Surveillance/PPS Coordinating Centre—and the hospital name.
- **8.** Click the **Save** icon **I** or press **Ctrl+S** to save the data **as a record in the database**.
- **9.** Click **Select this hospital** to make changes to or view/enter data on this hospital.

Working with HelicsWin.Net forms

In HelicsWin.Net you enter your data in a series of forms. These forms have standard features, such as labelled text boxes, drop-down lists, and option buttons, to help you become familiar with the user interface so that you can enter your data quickly and accurately.

General guidelines

Caution:

Each time you log in, the software opens the first record in the database. Therefore, if you start entering data without creating a new empty record, the first record will be overwritten when data are saved!

You use the toolbar's functions to perform basic tasks:

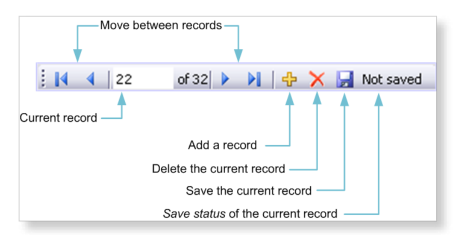


Table 2 describes these functions.

Table 2: Toolbar functions

Icon	Function	Description
22 of 33	Current record	Indicates which record is currently open for editing.
[4 4 ▶ M	Move between records	Use the Backwards ◆ and Forwards ▶ buttons on the toolbar to move one record at a time; alternatively, press Ctrl + left and right arrow
		or use Back I and Forward ► to go the start or end record respectively.
÷	Add a record	On all data entry forms, first click the yellow Add item + icon to activate the relevant input fields.
		Click to add a new record.
		Make sure the current record is saved first, before creating a new record.
		Alternatively, press Ctrl+N to create a new record.

Icon	Function	Description
Delete the current record	Most deletions are preceded by a warning. Note that if you delete a record (e.g. a ward), with dependent records (e.g. patient records), all the dependent records will be deleted.	
	However, if you deleted a record by mistake, exit the programme without saving and re-start HWN to undo the deletion. Click the Save icon to confirm the permanent deletion of the record.	
Not saved	Save status of the current record	The Not saved indicator text appears whenever you have changed data in a data record but not saved it. This indicator disappears as soon as you save the record, and reappears if you make any changes.
	Save the current record	Alternatively, press Ctrl+S to save the current record.

Table 3: Keyboard shortcuts

Keyboard shortcut	Action
Ctrl + Left arrow	Previous record
Ctrl + Right arrow	Next record
Ctrl + N	New record
Ctrl + S	Save record
Ctrl + D	Delete record

When a record is saved, the programme performs some validation checks on the data before actually saving. See *Checking data quality*.

Understanding reported errors

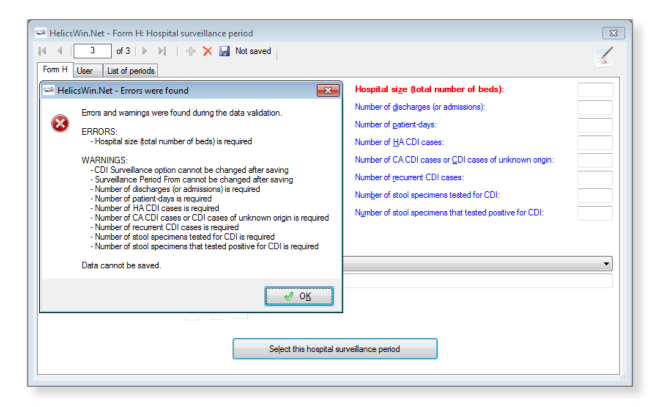
The application checks the data you supply as soon as you attempt to save it. The purpose of this check is to help you get your data right. If data quality checks and data entry validation identify errors, the application displays a message before saving.

There are three types of field validation in HWN:

- Type 1: **mandatory fields** are left blank or have been assigned invalid values: an **error message** is displayed, the relevant field labels are shown in red and you cannot save the data.
- Type 2: required fields are left blank: a warning message is displayed, the relevant field labels are shown in blue, but saving is possible. This will typically occur when required data are not available until the end of the surveillance period, e.g. the number of discharges in the surveillance period.
- Type 3: **optional fields** that can be empty: no warning or error message is displayed.

Figure 1 illustrates possible errors and warnings that could arise when you save a record in the CDI hospital data form. Note that in this case, there is one error and the data cannot be saved before the mandatory data is entered.

Figure 1: Example of errors and warnings



Note: Not all validation rules are implemented at data entry; some more complex rules are implemented only in the *data quality check*.

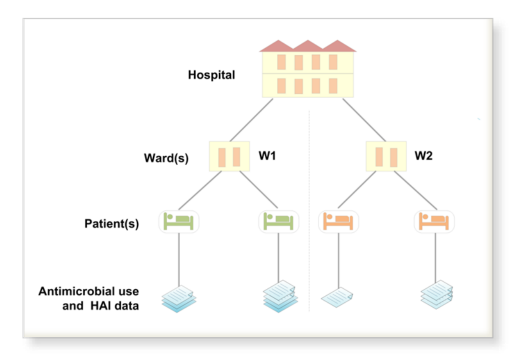
General features of the user interface

- General outlook of data forms: directly underneath the toolbar, the selected hospital code is displayed.
- You have to enter most categorical variables through drop-down lists (starting with blank line), so that you can enter only a predefined value from the list.
- Tabbing: keyboard tabbing first follows the order of input fields, after which focus will be put on toolbar.
- You can access fields with keyboard shortcuts made up of the Alt key and the underlined letter
 on the field label. For example, if you press Alt+P in the Hospital form, the focus moves to the
 PPS Protocol field.
 - Once the focus is on the field, enter the first letter of the required value, for example, **S** for Standard protocol, alternatively, you can select the value from the drop-down list.
- The default buttons in the warning messages—for example, **Yes** and **No** buttons—are displayed in the language of the installed Windows operating system.

Data hierarchy

Data in HWN is stored hierarchically, with the hospital at the top and patient data at the bottom. The definition of data at any level depends on the earlier creation of data for the level immediately above.

Figure 2: Data hierarchy



One consequence of this data hierarchy is that if you delete a definition at one level you automatically delete the data that depend on that definition, at the lower levels. For example, if you delete a ward definition (W2), all the data belonging to patients on that ward are also deleted (Figure 4). Furthermore, if you delete the whole hospital record, you would delete all the related ward and patient data as well.

HWN warns you before it deletes anything, but you need to be vigilant to ensure that you do not lose your work or that of others.

Figure 3: Data hierarchy – warning message

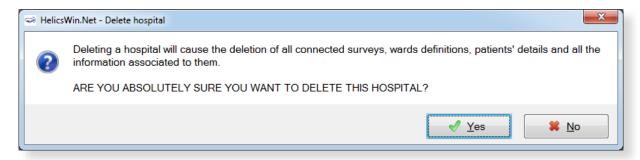
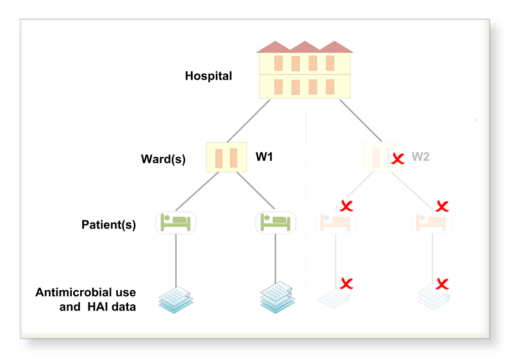


Figure 4: Data hierarchy – the effect of deleting a ward definition



Using the main menu

Use the main menu to open the main data entry forms for the different data levels. These forms are arranged in hierarchical order, based on hospital-, ward-, and patient-level data.

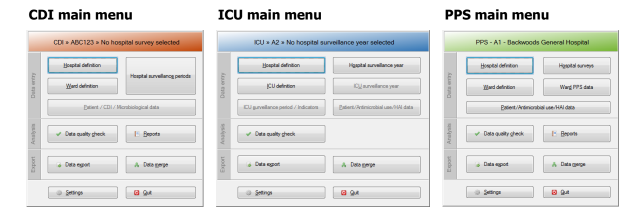


Table 4: Summary of main menu functions

Button	Actions(s)	
Hospital definition	Enter the name and code of a new hospital in the system. You can enter the data for different hospitals (sites) in the same database, e.g. hospitals within hospital trusts or organizations. This also enables national/regional coordinating centres to enter different hospitals data centrally, into a single database. If more than one hospital has been entered, you select individual hospitals here, by double clicking on its name in the Hospitals list.	
Ward definition ICU definition	Define all ward/ICU ID codes (abbreviated names). Optionally and in addition, full ward names can be entered.	
Hospital surveys	Enter data from the hospital questionnaire. These data include the hospital's survey	
Hospital surveillance periods / year	dates (or surveillance year), hospital size and type, and, for the PPS, a number of infection control structure and process indicators.	
Ward PPS data	PPS: Enter the date the PPS was performed and the ward specialty for each ward.	
ICU surveillance year	ICU: Enter data that remain constant for the entire ICU surveillance year, e.g. ICU size and specialty.	
Patient/CDI /Microbiological data	CDI: For the MINIMAL option: Enter aggregated nominator and denominator data. For the LIGHT option: Enter aggregated nominator and denominator data, and casebased data. For the ENHANCED option: Enter aggregated nominator and denominator data, casebased data, and <i>C. difficile</i> isolate data.	
ICU surveillance period/indicators	ICU: Enter the start date and end date of the surveillance period. For the LIGHT option: Enter the denominator data (required). In ICU, optionally enter denominator data in STANDARD surveillance. In ICU, this level also contains the structure and process indicators for HAI prevention.	

Button	Actions(s)	
Patient/Antimicrobial use /HAI data	PPS: For the STANDARD option: Enter each patient's demographic data and risk factors. For the LIGHT option: Enter demographic data for each patient with at least one HAI (or antimicrobial, in the case of PPS). For both options: Access the healthcare-associated infection form and/or antimicrobial use form from this patient form, through separate buttons.	
Data quality check	Analyse the data and report any missing or impossible values, and missing records. These checks go beyond the automatic validation checks that take place when individual records are saved; for example, cross-checks between different data levels.	
Reports	Create pre-formatted reports containing patient- or ward-based data related to HAIs and/or antimicrobial usage. This feature enables you to use off-the shelf reports or reports that you customise yourself. Currently only available for the PPS module.	
Data export	Export data as (i) raw data (as stored in the HelicsWinNet.mdb access format database), with or without user or validation variables, or as (ii) CSV (comma separated text) files in ECDC's TESSy CSV format (always without user variables). Export in TESSy CSV format is currently only available for the PPS module.	
Data merge	Merge HelicsWin.Net databases; this is useful if you want to consolidate the data from different hospitals, or from different wards within one hospital, into a single database. This application can (i) analyse the data in the other database to ensure that they are consistent with the data being merged; and (ii) if necessary, add user defined prefixes to user IDs to prevent overlapping patient IDs from different hospitals or wards.	
Settings	From this form you can:	
	Change your password;	
	 Specify how HelicsWin.Net sorts lists that contain data values (for example, sorting antimicrobials alphabetically by ATC5 code or alphabetically by antimicrobial agent) 	
	Translate labels on HelicsWin.Net forms	
	This feature enables you to translate labels on data fields and controls, such as tabs and buttons. You can access the text definitions for all the labels on the forms and validation messages and translate them in any language. You can also rename user field labels so that a local or national extra data collection modules can be implemented. The system also checks problems in the translation files, such as an accidental deletion of labels, and automatically fixes inconsistencies;	
	 Reset window sizes to their original values; 	
	 Define the level of detail of error logging (for debugging purposes). 	
Quit	Shut down the programme.	

Defining hospitals

If you are working in a single hospital, create the definition for that hospital. Once a hospital definition has been created, you can then add its ward and survey data.

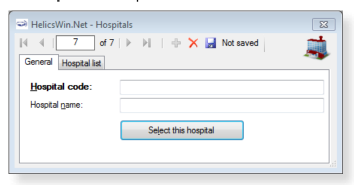
If necessary, you can add additional hospitals to the same HelicsWin.Net application. In doing this you are able to store the data for the additional hospitals in the same HelicsWin.Net database. If more than one hospital has been defined, you can only enter information for one hospital at a time.

To create a hospital definition:

1. Click Hospital definition in the main menu.



The **Hospitals** form opens.

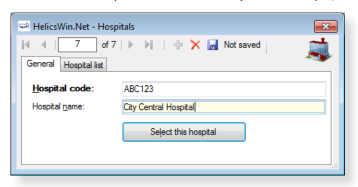


If no hospitals have been defined yet, the fields are blank.

2. Click the **Add item** icon .

The **Hospital code** and **Hospital name** fields clear.

3. Enter the new hospital code and name (in this example, 'ABC123' and 'City Central Hospital').



4. Click the **Save** icon III or press **Ctrl+S**.

The hospital record is added to the internal database. From now on, you can access this record from this form through the Forwards > and Backwards < buttons on the toolbar.

5. Click Select this hospital.

The *main menu* re-opens.

You can now enter this hospital's ward and survey information – see:

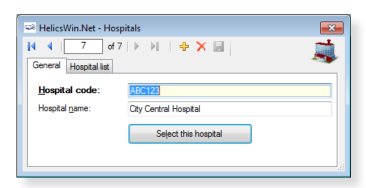
- Defining wards and ICUs and
- Creating a PPS point prevalence survey or
- Creating an ICU survey or
- Creating a CDI survey

Selecting a defined hospital

If you have more than one hospital defined, you can enter information for one hospital at a time. This includes (i) adding or editing survey data and (ii) adding or editing ward definitions.

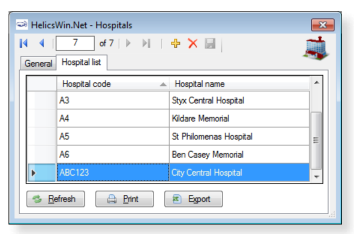
To select a hospital:

- **1.** Click **Hospital definition** in the *main menu* (as in step 1 above).
 - **2.**The **Hospitals** form opens.



2. Click the **Hospitals list** tab.

A list of all defined hospitals opens.

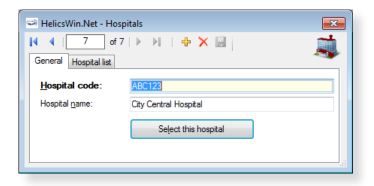


3. Click Refresh.

This ensures that hospitals defined since the last refresh are included in this list.

4. To select the hospital, double-click on its code.

This re-opens the **General** tab, and shows the hospital you chose.



5. Click **Select this hospital** to select it.

Hospital types

You do not enter details about the type of hospital while defining a hospital. The type is defined later when you create a survey.

Table 5: Hospital types

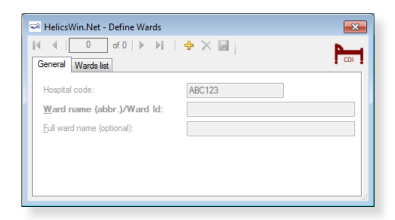
Primary	Often referred to as a 'district hospital' or 'first-level referral' hospital.
	Few specialities (mainly internal medicine, obstetrics-gynaecology, paediatrics, general surgery or only general practice).
	Limited laboratory services are available for general, but not for specialised pathological analysis.
	Often corresponds to a general hospital without teaching function.
Secondary	Often referred to as a 'provincial hospital'.
	Hospital is highly differentiated by function with five to ten clinical specialities, such as haematology, oncology, nephrology, ICU.
	Takes some referrals from other (primary) hospitals.
	Often corresponds to a general hospital with teaching function.
Tertiary	Often referred to as a 'central', 'regional' or 'tertiary-level' hospital.
	Highly specialised staff and technical equipment (ICU, haematology, transplantation, cardio-thoracic surgery, neurosurgery); specialised imaging units.
	Clinical services are highly differentiated by function.
	Specialised imaging units.
	Provides regional services and regularly takes referrals from other (primary and secondary) hospitals.
	Often a university hospital or associated with a university.
Specialised	Single clinical specialty, possibly with sub-specialties.
	Highly specialised staff and technical equipment.
	Specialisation (e.g. paediatric hospital, infectious diseases hospital) should be specified in a form when this is possible.

Defining wards and ICUs

You can define wards (units) to be included in the Point Prevalence Survey or different intensive care units (ICUs) in the ICU module. For each ward/unit, you have to enter an abbreviated name (the Unit ID or code) that will be used in all levels of the database and, optionally, a full ward name. The procedure to define wards in the PPS module described below is similar to the procedure to define ICUs in the ICU module.

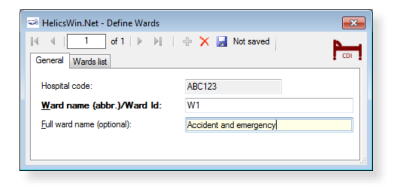
To create a ward or ICU definition:

Click Ward definition (PPS and CDI) or ICU definition (ICU) in the main menu.
 The Wards or Define ICUs form opens showing the current hospital code.



If this is not the correct hospital, click the **Close** icon to cancel. To select the correct hospital, see *Selecting a defined hospital*.

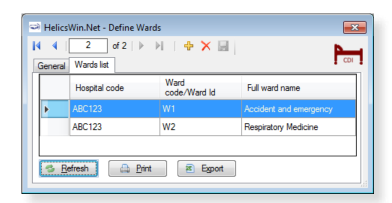
- Click the Add item icon [♣] or press Ctrl+N.
 The form opens for editing.
- **3.** In the **Ward name** or **ICU name** field, enter the ward/unit name (up to 20 characters) or ID.



You cannot enter the same ID twice. The ward/unit list can be used for different hospital surveys within the same hospital.

- **4.** Optionally, specify the full (expanded) name in the **Full ward name** or **Full ICU name** field.
- 5. Click the **Save** icon or press **Ctrl+S**.

6. Click the **Wards list** or **ICU list** tab to view all the wards/units for which a record has been created.



Note: The ward specialty is defined later (because it may change from one survey to another).

You can now enter this hospital's survey information – see:

- Creating a CDI survey or
- Creating an ICU survey or
- Creating a PPS point prevalence survey

McCabe score

The examples in Table 6, and in particular those of the second (ultimately fatal) category, are not meant to be exhaustive, but rather to serve as guidance when using the protocols.

Table 6: McCabe score examples

McCabe score categories	Examples
Non-fatal (> five years)	Diabetes Carcinoma/haematological malignancy with > 80% five-year survival Inflammatory disorders Chronic GI, GU conditions Obstetrics Infections (including HIV, HCV, HBV – unless in above categories) All other diseases
Ultimately fatal (one year to four years)	Chronic leukaemias, myelomas, lymphomas, metastatic carcinoma, end-stage kidney disease (without transplant). Motor neuron disease, multiple sclerosis non-responsive to treatment. Alzheimer's/dementia. Diabetes requiring amputation or post amputation.
Rapidly fatal (< one year)	End-stage haematological malignancies (unsuitable for transplant, or relapsed), heart failure (EF < 25%) and end-stage liver disease (unsuitable for transplant with recalcitrant ascites, encephalopathy or varices). Multiple organ failure on intensive care unit – APACHE II score > 30, SAPS II score > 70. Pulmonary disease with cor pulmonale.

Creating a CDI survey

There are three *Clostridium difficile* surveillance options:

- Minimal collects:
 - Aggregated numerator and denominator data
- Light collects:
 - Aggregated numerator and denominator data
 - Case-based numerator data
- Enhanced
 - Aggregated numerator and denominator data
 - Case-based numerator data
 - Microbiological data

Please refer to the CDI Protocol for detailed descriptions of these options.

Please bear the following in mind when planning your CDI survey:

- Continuous surveillance for 12 months is recommended.
- The recommended minimum surveillance period is three consecutive months, preferably from 1 October to 31 December, or from 1 January to 31 March.
- Starting on the first day of the month is recommended (the pilot study demonstrated that completing Form H is much easier by starting surveillance on the first day of a month).
- You cannot change the start date of the survey once it has been defined. When creating your CDI survey, it is therefore important to make sure its start date is earlier than any ward survey date.
- You can have multiple surveys, but two surveys cannot start on the same date.
- You can later change from Minimal to Light, or from Light to Enhanced, but not vice versa.

CDI survey data collection is described in the following sections:

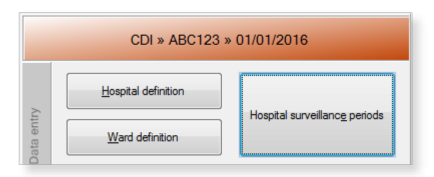
- Entering CDI hospital-based data (Form H)
- Entering CDI patient, CDI and microbiological data (Form C)
- Entering CDI microbiological data (Form M)

Form H - entering CDI hospital-based data

Follow the procedure in this section to create a new **CDI** survey for each defined hospital, entering information from Form H. Before you can create your survey, you must *define your hospital and wards*.

To create a CDI hospital survey:

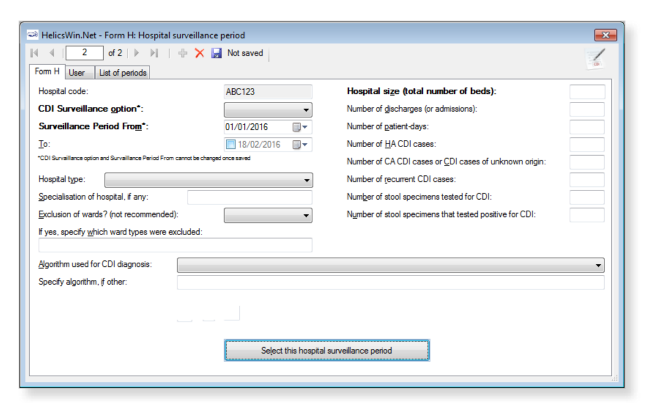
1. Click Hospital surveillance periods in the CDI main menu.



The **Hospital survey** form opens for the hospital you have selected.

- **2.** If not already open, click the **Form H** tab.
- 3. In the Form H form, click the Add item icon ♣ to open the form for editing.

 The data entered here is hospital data applicable to the Minimal, Light and Enhanced surveillance options.



4. Specify the values for the fields.

Descriptions of the variables in Form H are given in Table 7. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

5. Click the **Save** icon or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

6. Click **Select this hospital surveillance period** to return to the main menu and proceed to *enter Form C data*.

You can now see the surveillance period in the **List of periods** tab (click **Refresh**).

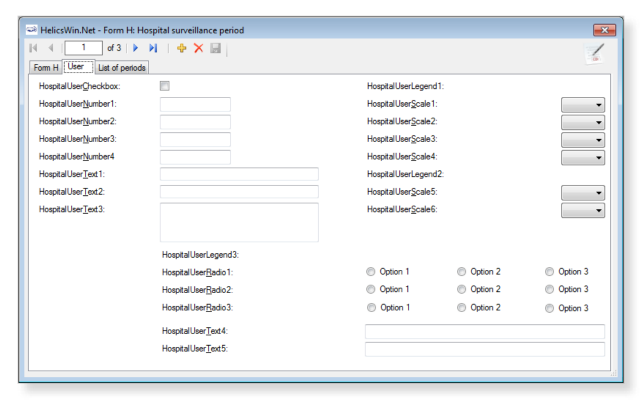
Table 7: CDI Form H variables

Variable	Description
Hospital code *	The hospital identifier/code assigned by the national/regional CDI surveillance coordinator. Hospital codes should be unique within each surveillance network, and kept constant between the ECDC Antimicrobial Resistance and Healthcare-Associated Infection (ARHAI) surveillance protocols and from one year to the next. Inserted automatically based on the hospital selected during Hospital definition.
CDI Surveillance option *	Specify the CDI surveillance option to be used. You can select either Minimal , Light or Enhanced . Note that you can later change from Minimal to Light, or from Light to Enhanced, but not vice versa. Refer to the CDI Protocol for detailed descriptions of the three options.
Surveillance period from *	The start date of the survey. You can have multiple surveys, but two surveys cannot start on the same date. Warning: You cannot change the Surveillance period from date after you have saved it. If you save without having selected the correct date, today's date is applied by default. Before saving, make sure that the start date of the hospital CDI survey is earlier than any ward survey date. If you need to change the start date, you will have to
То	contact your national or regional CDI co-ordinator. The end date of the survey. Can be entered at a later time and can also later be changed.
Hospital type *	Designate the hospital type – PRIM: primary, SEC: secondary, TERT: tertiary, SPEC: specialised, missing=UNK. See <i>Hospital types</i> .
Specialisation of hospital	Free text. Indicate the hospital specialty for a specialised hospital (e.g. paediatric, infectious diseases, etc.); please use specialty codes if possible.
Exclusion of wards	All wards should be included for the surveillance of CDI, exclusion of wards is strongly discouraged. If, despite this recommendation, certain wards were excluded, it is crucial that the aggregated denominator data are provided for the included wards only.
If yes, specify wards	Free text field. Enter the name or ID of any ward excluded from the survey. Max. 100 characters.
Hospital size (total number of beds) *	Number of hospital beds for the current surveillance period. If, despite the recommendation to the contrary, certain wards were excluded, it is crucial that the aggregated denominator data are provided for the included wards only.
Number of discharges (or admissions) *	Number of hospital discharges in the current surveillance period. Use number of admissions if discharges are not available.

Variable	Description
Number of patient-days *	Number of hospital patient-days in the current surveillance period.
Number of HA CDI cases *	Number of healthcare-associated CDI cases within the surveillance period (i.e. with onset on day three or later, following admission to a healthcare facility on day one, OR in the community within four weeks of discharge from any healthcare facility). Exclude recurrent cases.
Number of CA CDI cases or CDI cases of unknown origin *	Number of community-associated CDI cases and cases of unknown origin within the surveillance period i.e. onset outside of healthcare facilities, AND without discharge from a healthcare facility within the previous 12 weeks, OR onset on the day of admission to a healthcare facility or on the following day AND not resident in a healthcare facility within the previous 12 weeks, OR a CDI case discharged from a healthcare facility 4–12 weeks before the onset. Exclude recurrent cases.
Number of recurrent CDI cases *	Number of CDI episodes with onset within two and eight weeks of a previous episode (including both healthcare-associated and community-associated recurrent cases).
Number of stool specimens tested *	Number of stool specimens tested for CDI in the surveillance period. Each specimen should only be counted once, even if more than one test was performed on that specimen.
Number of stool specimens that tested positive for CDI *	Number of stools tested for CDI with a positive test result in the surveillance period. Each specimen should only be counted once.
Algorithm used for CDI diagnosis	The laboratory test(s) applied on faeces samples to recognise the presence of toxin-producing <i>C. difficile</i> , either as a solitary test or as a combination of screening and confirmatory tests. If multiple algorithms are applied (i.e. depending on work hours or patient categories), please indicate the most frequently applied algorithm(s), that is/are used for more than 80% of the samples tested for <i>C. difficile</i> .
	Toxin A/B EIA Enzyme immunoassays, including enzyme-linked immunosorbent assays (ELISA), that test for both toxins A and B in stool samples or cultures.
	GDH EIA Enzyme immunoassays, including enzyme-linked immunosorbent assays (ELISA), that test for both Glutamate dehydrogenase in stool samples or cultures.
	NAAT Nucleic acid amplification tests.
	Cytotoxicity assay Demonstration that stool sample supernatant kills a cell monolayer in the absence of a <i>C. difficile</i> toxin-neutralising antibody.
	Toxigenic culture Demonstration that a <i>C. difficile</i> culture is able to produce toxins in vitro, e.g. by cytotoxicity assays, Toxin A/B EIA or NAAT from colonies.
	Toxin detection Detection of toxins, in stool samples or cultures, e.g. by toxin A/B EIA or cell cytotoxicity assays.
Specify algorithm, if other	Free text field. If none of the listed algorithms matches your algorithm, indicate the algorithm which matches most closely. Max. 255 characters.

Surveillance period User tab

The **Surveillance period User** tab contains fields that can be renamed to enable collection of information during the survey that is outside the scope of the CDI protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.

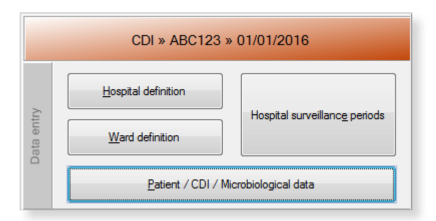


Form C - entering CDI patient and infection data

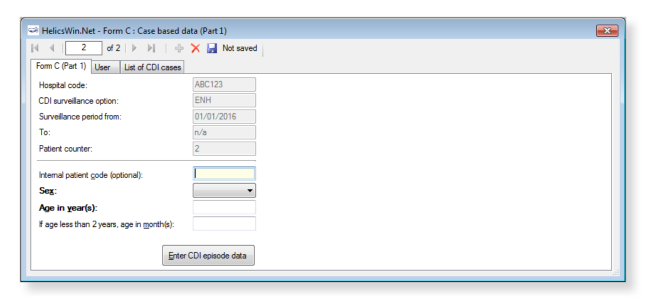
Form C is applicable only to the **Light** and **Enhanced** surveillance options. The **Minimal** surveillance option does not record case-based data

To enter CDI patient and infection data:

1. Click Patient / CDI / Microbiological data in the CDI main menu.



A form equivalent to the upper section of Form C opens.



- 2. Click the **Add item** icon $\stackrel{1}{+}$ or press **Ctrl+N** to open the form for editing.
- **3.** Enter the patient's age and gender and, optionally, an internal patient code.

The fields are described in Table 8. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

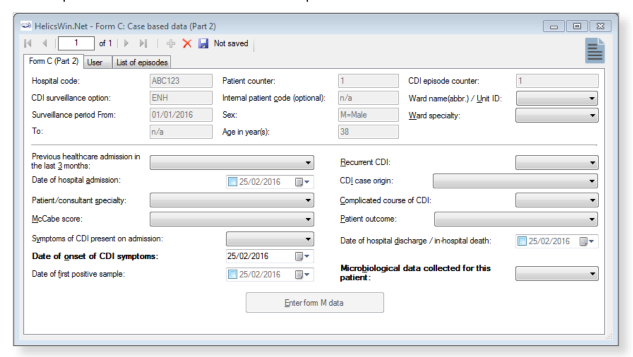
Important: please note that to avoid data privacy infringement, internal patient codes must be removed from any data submitted to ECDC/TESSy.

Table 8: CDI Form C variables, part 1

Variable	Description
Hospital code	The hospital identifier/code for the hospital selected for the survey. Inserted by the system.
CDI Surveillance option	Minimal, Light or Enhanced. From Form H.
Surveillance period from	The start date for the survey. From Form H.
То	The end date for the selected survey. From Form H. The field is blank if the end date has not yet been defined for this survey.
Patient counter	A system allocated sequential patient number for anonymised identification. In enhanced surveillance, this number should permit linkage of patient data with microbiological typing/susceptibility data and patient data from enhanced surveillance.
Internal patient code	For hospital internal use only. Optional. Must be removed from any data submitted to ECDC/TESSy.
Sex *	Gender of the patient: M (male), F (female).
Age in year(s) *	Patient's age in years; if missing=unknown (UNK). Enter respectively 0 or 1 if the patient is less than two years old.
If age less than 2 years	Patient's age in months. Mandatory if Age in years is less than 2 (note that you must <i>also</i> enter the baby's Age in years).

4. Click Enter CDI episode data.

A form equivalent to the lower section of Form C opens.



Note that the **Enter form M data** button is only displayed for the **Enhanced** surveillance option. See also *Form M - entering CDI microbiological data*.

- **5.** Enter the patient's CDI data, see Table 9. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- **6.** Enter Yes, No, or Unknown for **Microbiological data collected for this patient**. If you have entered:
 - No or Unknown then click the **Save** icon III or press **Ctrl+S** to save and exit.
 - Yes and you are working in the Light option then click the Save icon or press
 Ctrl+S to save and exit.
 - Yes and you are working in the Enhanced option click Enter form M Data and proceed to Entering CDI microbiological data.



If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

You can now see the Form C data in the **List of episodes** tab (click **Refresh**).

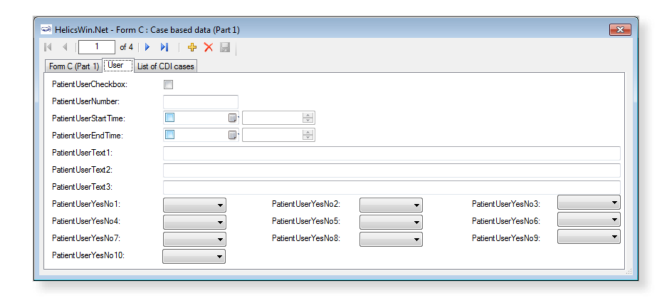
Table 9: CDI Form C variables, part 2

Variable	Description
Hospital code	The hospital identifier/code for the hospital selected for the survey. Inserted by the system.
CDI surveillance option	Minimal, Light or Enhanced. From Form H.
Surveillance period from	The start date previously selected for the survey. From Form H.
То	The end date for the survey. From Form H. The field is blank if the end date has not yet been defined for this survey.
Patient counter	System assigned sequential patient number for anonymised identification. In Enhanced surveillance the counter links patient data with microbiological typing/susceptibility data.
Internal patient code	Optional. From Form C, part 1. Must be removed from any data submitted to ECDC/TESSy.
Sex	From Form C, part 1.
Age in year(s)	From Form C, part 1.
CDI episode counter	System assigned sequential number for episode identification.
Ward name/unit ID	Abbreviated name of hospital ward; it should be used consistently and should remain the same in different surveillance periods/years.
Ward specialty	Enter the code for the main ward specialty. See Protocol for the ward specialty code list.
Previous healthcare admission	Previous admission in a healthcare facility in the last three months relative to the onset of CDI: Yes/No/Unknown. If yes: admission in a hospital or another healthcare facility (long-term care, outpatient department, etc.). Collect from electronic records and/or patient notes, and/or by asking the patient.
Date of hospital admission *	Date patient was admitted to the hospital for the current hospitalisation (dd/mm/yyyy).
Patient/ consultant specialty	Enter the code for the specialty of the physician in charge of the patient; this may differ from the ward/unit specialty. See Protocol for the ward specialty code list.
McCabe score	Classification of the severity of underlying medical conditions. Disregard the influence of an active CDI, i.e. estimate the score the patient had before the infection. Some examples of diseases and their different McCabe score categories are shown in Table 6. These examples, in particular those of the second (ultimately fatal) category, are not meant to be exhaustive but rather to serve as a guidance tool for the current protocol.
Symptoms of CDI present on admission	Patient had CDI symptoms when admitted for this episode. Yes/No/Unknown.
Date of onset of CDI symptoms *	Mandatory if symptom onset was during current hospitalisation, but not recorded if signs/symptoms were present on admission. Record the date of the first signs or symptoms of the infection (dd/mm/yyyy). If unknown, record the date treatment was started for this infection or the date the first diagnostic sample was taken. If no treatment or sample, please estimate. If no date is entered, today's date is selected by default. This date can later be changed.
Date of first positive sample	Date on which the first positive diagnostic stool sample was taken from the patient referred to on this form.

Variable	Description
Recurrent CDI *	Choose yes if the patient had an episode of CDI (return of diarrhoeal stools with a positive laboratory test after the end of treatment) more than two weeks and less than eight weeks following the onset of a previous episode.
CDI case origin *	 Choose one (for detailed definitions, see Definitions section): Healthcare-associated CDI: A case with onset of symptoms on day three or later, following admission to a healthcare facility on day one, OR in the community within four weeks of discharge from any healthcare facility. This may apply to the current hospital or a previous stay in another healthcare facility, e.g. in another hospital, a long-term care facility or other healthcare facilities (e.g. outpatient departments etc.). Community-associated CDI: A case with [onset outside of healthcare facilities, AND without discharge from a healthcare facility within the previous 12 weeks] OR [onset on the day of admission to a healthcare facility or on the following day AND not resident in a healthcare facility within the previous 12 weeks]. Unknown association: A case discharged from a healthcare facility 4–12 weeks before symptom onset.
Complicated course of CDI *	 Yes / No / Unknown. CDI leading to any of the following: Admission to a healthcare facility for treatment of community-associated CDI. Admission to an intensive care unit for treatment of CDI or its complications (e.g. for shock requiring vasopressor therapy). Surgery (colectomy) for toxic megacolon, perforation or refractory colitis. Death within 30 days after diagnosis if CDI is either a primary or contributing cause.
Patient outcome *	 Status of the patient at hospital discharge or at end of follow-up in the hospital Discharged alive: patient was discharged alive; OR patient was still in the hospital and alive at end of follow-up during this hospital stay. Death, CDI definitely contributed to death: use this category if a causal link between CDI and death can be demonstrated.
Date of discharge/ in- hospital death	Date the patient was discharged from the hospital; OR date of end of follow-up if the patient was still hospitalised and alive; OR date of death if patient died during the current hospitalisation.
Microbiological data collected for this patient *	Yes/No/UNK. Indicate whether Form M has been completed.

Case-based data User tab

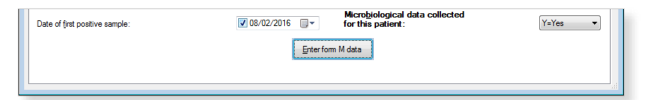
The **Case-based data User** tab contains fields that can be renamed to enable collection of information during the survey that is outside the scope of the CDI protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.



Form M - entering CDI microbiological data

Only the **Enhanced** surveillance option supports recording microbiological data – the **Minimal** and **Light** options do not support this.

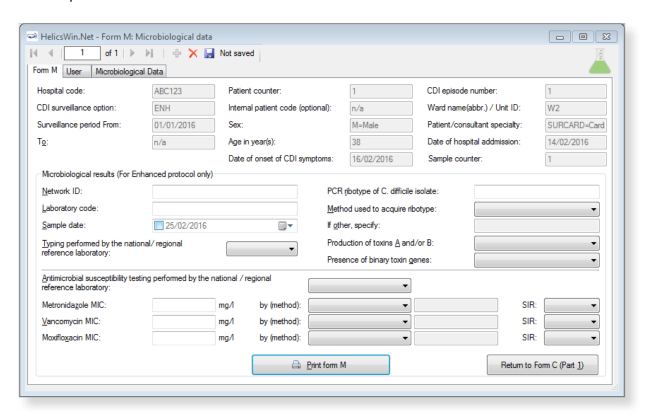
If you are working in the **Enhanced** option, the **Enter form M data** button is displayed at the bottom of Form C, part 2.



To enter microbiological data:

1. Click Enter form M data in Form C.

Form M opens.



- **2.** Enter the microbiological data. The fields are described in Table 10.

 Descriptions of the variables in Form H are given in Table 10. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- 3. Click the **Save** icon III or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

You can now see the Form M data listed in the Microbiological data tab (click Refresh).

Table 10: CDI Form M variables

Variable	Description
Hospital code	The hospital identifier/code for the hospital selected for the survey.
CDI surveillance option	Minimal, Light or Enhanced. From Form H.
Surveillance period from	The start date for the survey. From Form H.
То	The end date for the survey. From Form H. The field is blank if the end date has not yet been defined for this survey.

Variable	Description
Patient counter	System assigned sequential patient number for anonymised identification. In Enhanced surveillance the counter links patient data with microbiological typing/susceptibility data.
Internal patient code	From Form C. Must be removed from any data submitted to ECDC/TESSy.
Sex	From Form C.
Age in year(s)	From Form C.
Date of onset of CDI symptoms	From Form C.
CDI episode number	System assigned sequential number for identification.
Ward name/unit ID	From Form C.
Patient/consultant specialty	From Form C.
Date of hospital admission	From Form C.
Sample counter	System assigned sequential number for identification. Assigned when Form M is created.
Network ID	Unique identifier for each surveillance network within a Member State, selected and generated by the Member State, e.g. for UK, EN, NI, SC or WA; for France, different CClin networks. This field is combined with the hospital identifier to create a unique hospital code, because different networks within one Member State may use the same hospital code. Can be omitted if the hospital identifiers are unique within the reporting Member State.
Laboratory code *	Local laboratory identifier/code assigned by national/regional CDI surveillance coordinating centre. For the primary laboratory responsible for microbiological confirmation of the CDI (not the code of the national/reference laboratory). It is recommended to use the same laboratory codes as in EARS-Net. Required for data export.
Sample date	Date on which the first positive diagnostic stool sample was taken from the patient referred to in this form, if available. Otherwise, the date the stool sample was taken resulting in the results referred to in this form.
Typing performed by a national/ regional reference laboratory	Typing of <i>C. difficile</i> isolates performed by a laboratory that provides diagnostic, analytical and advisory services to other laboratories, nationally or sub-nationally.
PCR ribotype of <i>C.</i> difficile isolate	<i>C. difficile</i> PCR ribotype as determined by conventional gel-electrophoresis or capillary-based PCR ribotyping.
Method used to acquire ribotype	Method used to acquire PCR ribotype information, e.g. capillary-based PCR ribotyping; conventional gel-electrophoresis.
If other	Other method used to acquire PCR ribotype (please specify, e.g. whole genome sequencing).
Production of toxins A and/or B	Production of toxins A and/or B as determined by PCR of tcdA and tcdB or by EIA for TcdA and TcdB.
Production of binary toxin genes	Production of binary toxin (CDT) as determined by PCR of cdtA and cdtB
Antimicrobial susceptibility testing performed by reference laboratory	Testing of <i>C. difficile</i> isolates for their susceptibility to antimicrobial agents performed by a laboratory that provides diagnostic, analytical and advisory services to other laboratories, nationally or sub-nationally. Yes/No/Test not performed/Unknown.

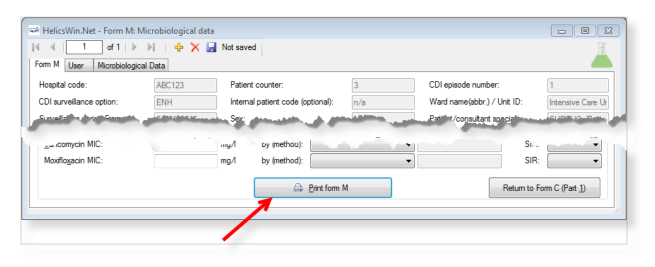
Variable	Description
Antimicrobial susceptibility testing	 For the relevant antimicrobial: Enter the MIC (minimum inhibitory concentration) in mg/l. Select the method used for the determination of the MIC from the dropdown list. If Other is selected, then the free-text field to the right of the list is enabled – enter a description of the method used. Max. 255 characters. Select the interpretation S, I or R, i.e. susceptible, intermediate or resistant.

Printing Form M

25%

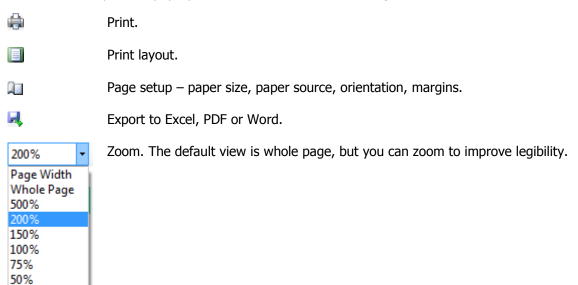
You might require to print a patient's microbiological data in order to send it to a laboratory, for example.

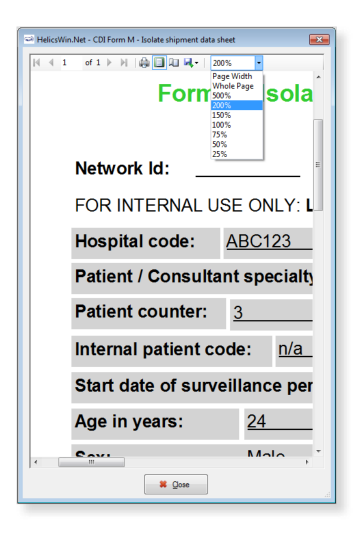
To do this, click $\operatorname{\textbf{Print}}$ form $\operatorname{\textbf{M}}$ at the bottom of the form.



The patient's microbiological data is entered into the hardcopy template and displayed in a pop-up window.

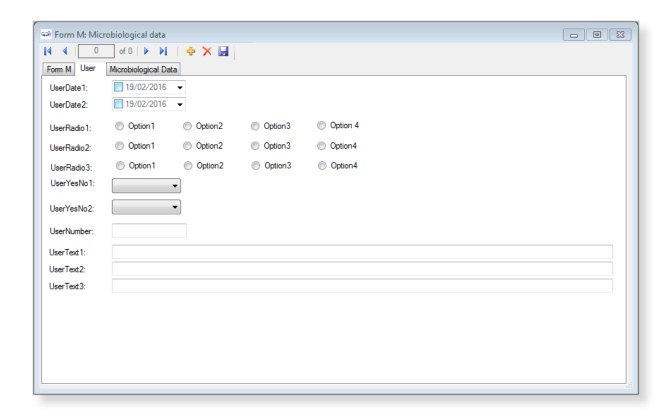
Controls at the top of the pop-up allow for basic document management:





Microbiological data User tab

The **Microbiological data User** tab contains fields that can be renamed to enable collection of information during the survey that is outside the scope of the CDI protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.



CDI validation tabs

There are no CDI Validation tabs or CDI validation protocol for HWN2.2.

Creating an ICU survey

There are two Intensive Care Unit (ICU) survey options:

- **Standard**: Patient-based option allows advanced risk adjustment of healthcare-associated infection rates for inter-hospital comparisons. You enter all patient data for all patients, including those without HAI and/or antimicrobial use. Risk factors are collected for each patient (infected or not).
- **Light**: Unit-based option is less labour-intensive solution, produces partially the same indicators as Standard option for follow-up of trends, as well as the same descriptive results about infections and antimicrobial resistance, but with less possibility for risk-adjusted comparisons. You only enter patient data for patients with any antimicrobial use and/or a healthcare-associated infection. Consequently, in the Light protocol, each patient record must include details of at least one antimicrobial use or HAI. Denominator data are aggregated at the unit (ICU) level.

Case definitions, included patients, and infection data (including antimicrobial resistance data and mortality review data) are identical in both options.

Further details are provided in the HAI-Net ICU protocol (see Related documents).

PPS data collection is described in the following sections:

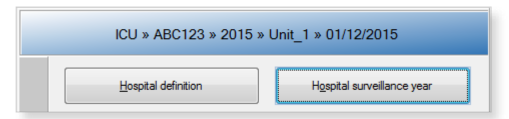
- Entering an ICU hospital surveillance year
- Entering ICU surveillance year and surveillance period data
- Entering ICU patient, HAI and antimicrobial use data

Entering an ICU hospital surveillance year

Follow the procedure in this section to create a new **ICU** survey for each defined hospital, entering information from Form H. Before you can create your survey, you must *define your hospital and wards*.

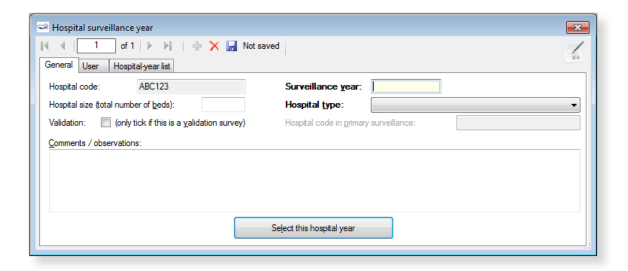
To create a hospital surveillance year:

1. Click **Hospital surveillance year** in the ICU *main menu*.



The **Hospital survey** form opens (for the hospital you have selected).

- 2. If not already open, click the **General (1)** tab.
- 3. In the General (1) form, click the Add item icon $\frac{1}{2}$ to open the form for editing.



The **Light** or **Standard** protocol option is selected later at the ICU surveillance period level.

- **4.** Specify values for the fields. Descriptions of the variables in this form are given in Table 11. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- **5.** Click **Select this hospital year** to be able to enter (an) ICU surveillance period(s).
- **6.** Click the **Save** icon **I** or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

Table 11: Hospital surveillance year ICU General tab variables

Variable	Description
Hospital code	The hospital identifier/code assigned by national/regional PPS coordinating centre; unique code per surveillance/PPS network. Hospital codes should be unique within each surveillance network, and kept constant between the ECDC Antimicrobial Resistance and Healthcare-Associated Infection (ARHAI) surveillance protocols and from one year to the next. Inserted automatically based on the hospital selected during Hospital definition.
Hospital size (total number of beds) *	Total number of beds in the hospital. If, for reasons of confidentiality, the exact number of hospital beds cannot be given, please enter the number of beds rounded up to the nearest 50.
Validation	If the current survey or surveillance year is a validation study performed by a validation team, the Validation checkbox should be checked to indicate that this is not a primary data collection but a repeated data collection for validation purposes of the primary PPS data.
	Checking the Validation checkbox will change the messages on data entry for the entire application, especially for the variables displayed on the Validation tabs at different levels (see <i>Validation tab</i> below).
	Validation cannot be changed after saving.
Surveillance year *	The year in which the survey will be performed.
	Warning: The surveillance year cannot be changed after saving. Before saving, make sure that the start date of the hospital ICU survey is within the year you are specifying. If you need to change the surveillance year, you will have to contact your national or regional PPS co-ordinator.

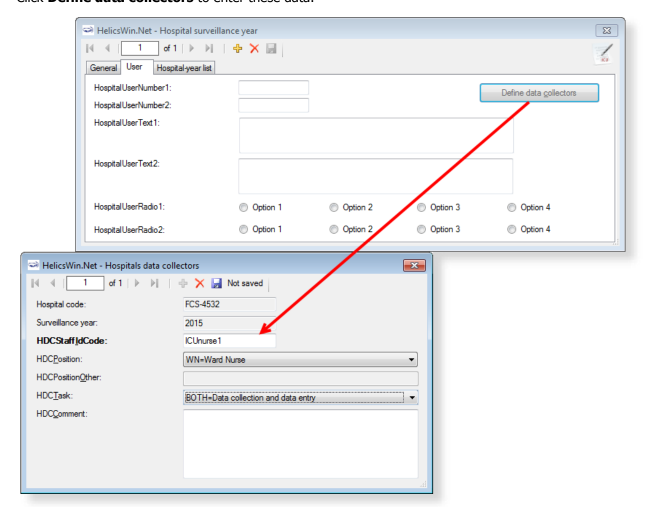
Variable	Description
Hospital type *	Designate the hospital type – PRIM: primary, SEC: secondary, TERT: tertiary, SPEC: specialised, missing=UNK. See <i>Hospital types</i> .
Hospital code in primary surveillance	
Comments/ observations	Free text field. Enter any information relevant to the survey.

User tab

The **User** tab contains fields useful for users at national, regional or hospital level. They can be renamed and personalised to enable collection of information during the PPS that is outside of the PPS protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.

Additionally, the **User** tab of the **Hospital surveys** form also enables you to record the details of all hospital data collectors (HDC): their ID Code, Position (function) and survey-related task(s). This is optional.

Click **Define data collectors** to enter these data.



Data collectors for ICU surveillance defined here can be used later in the **Pilot: Feasibility** tab of the ICU surveillance period / Indicators screen, see below.

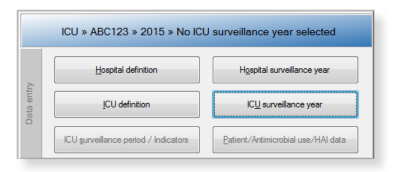
Entering ICU surveillance year and surveillance period data

How you enter these data depends on the protocol being used:

- In the (unit-based) **Light protocol**, you only enter patient data for patients with any antimicrobial use and/or a healthcare-associated infection. Therefore, in the Light protocol, each patient record must include details of at least one antimicrobial use or HAI.
- In the (patient-based) Standard protocol, you enter all patient data for all patients, including
 those without HAI and/or antimicrobial use. Additional risk factors (not applicable to the Light
 protocol) appear once you select the ward.

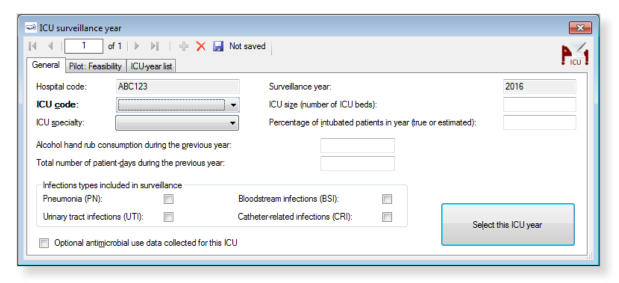
Surveillance year data

To enter surveillance year data, click **ICU surveillance year** in the *main menu*.



After entering the surveillance year data, save and click **Select this ICU year** in the **General** tab to be able to enter surveillance period data. Specifying surveillance year data, General tab

The **General** tab contains the ICU ward data that remain constant for all surveillance periods in the surveillance year. This tab also contains one of the structure and process indicators, i.e. the consumption of alcohol-based hand rub in the ICU in the previous year. Table 12 describes the variables in this form. Further details are provided in the HAI-Net ICU protocol (see *Related documents*).



To define the ICU surveillance year:

- **1.** Specify values for the fields in the **ICU surveillance year General** tab. Descriptions of the variables in this form are given in Table 12. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- 2. Click the **Save** icon or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

3. Click **Select this ICU year** to be able to enter an ICU surveillance period.

Table 12: ICU surveillance year General tab variables

Variable	Description
Hospital code	Inserted automatically based on the hospital selected during Hospital definition.
Surveillance year	The year previously selected for the survey. Inserted by system.
ICU code *	Unique identifier for each intensive care unit within a hospital, should remain identical in different surveillance periods/years.
ICU size *	Number of beds in the ICU.
ICU speciality *	If 80% of the patients belong to a particular category, the ICU falls within that category, otherwise the specialty is 'Mixed'. MIX = Mixed; MED = Medical; SURG = Surgical; CORO = Coronary; BURN = Burns; NEUR = Neurosurgical; PED = Paediatric; NEON = Neonatal; O = Other; UNK = Unknown.
Percentage of intubated patients in the ICU *	Percentage of intubated patients over the past year in the ICU. Measured or estimated average percentage (not: proportion) of patients with an invasive respiratory device over the last year in the current ICU. Number from 0.00 to 100.00. This variable is used as a proxy for severity of ICU case-mix and should also be collected if pneumonia is not included in the surveillance.
Alcohol hand rub consumption during the previous year *	Total number of litres of alcohol-based hand rub delivered to the intensive care unit (usually by the hospital pharmacy) during the previous year.
Total number of patient-days during the previous year *	Total number of patient-days during the year prior to the current surveillance year (patient-days for all patients, not only for patients staying more than 2 days in the ICU). Short interruptions are not taken into account. Partial days count as one patient day. This variable is the denominator of the indicator 'alcohol-based hand rub consumption in the ICU per 1000 patient-days'.
Infections types included in the surveillance *	Check the relevant checkbox(s) to indicate which of the four types of infection are included in the current ICU surveillance year. Included types should remain constant between different surveillance periods within the same surveillance year.
Optional antimicrobial use data collected	Check the checkbox to indicate that antimicrobial use data was collected at the patient level (blank = not collected).

Specifying surveillance year data, Pilot: Feasibility tab

The **Pilot: Feasibility** tab contains the surveillance-year part of the feasibility questionnaire (Form F) of the HAI-Net ICU pilot study. Table 13 describes the variables in this form. Further details are provided in the HAI-Net ICU protocol (see *Related documents*). Since this is a pre-configured User variables screen, please remember to include User variables when you export the data.

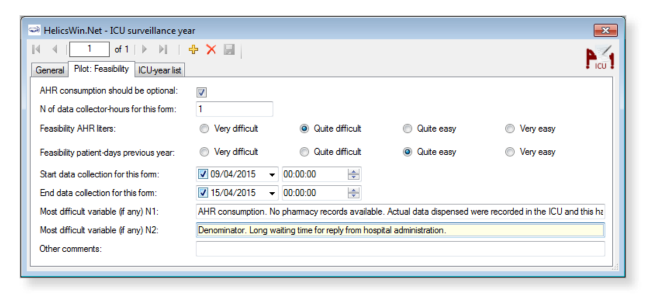
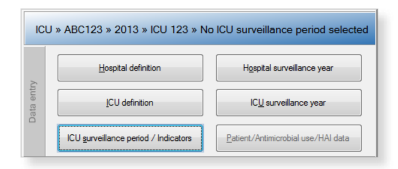


Table 13: ICU surveillance year Pilot: Feasibility tab variables

Variable	Description
AHR consumption should be optional	Should the AHR consumption indicator be kept optional in the ICU protocol? YES/NO (NO=required for all participating ICUs, leave checkbox unchecked).
Number of data collector-hours for this form	(Estimated) number of person-hours of surveillance staff (data collectors) actually spent by one or more of the data collectors, for ICU surveillance-year data only. In practice, this will be the time spent on the AHR indicator alone, e.g. time to write and send the requests to the pharmacist (for the numerator) and possibly hospital administration (for the denominator) plus the time actually spent by the pharmacist to extract the amount of litres delivered to the ICU during the previous year plus the time needed to extract the denominator data – do NOT include the waiting time needed for the reply (see dates below). Round to the nearest hour (e.g. round 45 minutes to 1 hour).
Feasibility AHR liters	Level of difficulty of collecting data the total number of litres of alcohol-based hand rub delivered to the ICU during the last year (numerator of the AHR consumption indicator). Click the appropriate choice.
Feasibility of the total number of patient-days during previous year	Level of difficulty of collecting data on the total number of ICU patient-days during the last year (denominator of the AHR consumption indicator). Click the appropriate choice.
Start/end data collection for this form	Start date and end date for the data collection of the ICU characteristics (type, size) and alcohol-based hand rub consumption during the previous year. Because ICU type and size are in principle readily available, these dates allow the calculation of the number of days (including waiting time) to receive the numerator and denominator data for the indicator AHR consumption.
Most difficult variable (if any) 1/2	Free text. Indicate the two most difficult data items of the ICU surveillance-year data and the reasons why or encountered problems (up to 255 characters). These fields may be left empty.
Other comments	Comments regarding variables or HelicsWin.Net software. Collected once per year or on the software interface. Free text, max 255 characters. This field may be left empty.

Surveillance period data

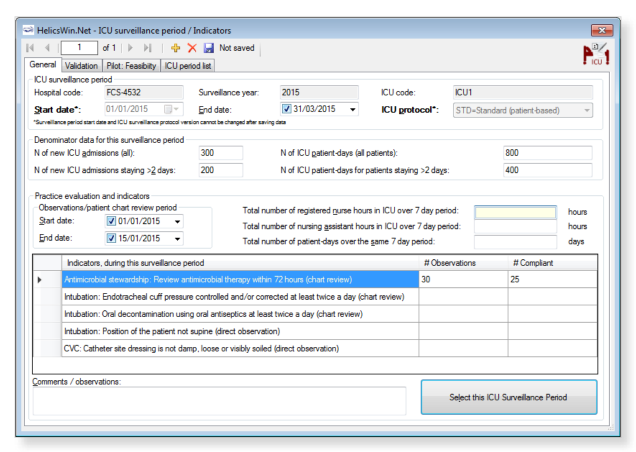
To enter surveillance period data, click **ICU surveillance period/Indicators** in the *main menu*.



After entering the surveillance period data, save and click **Select this ICU surveillance period** in the **General** tab to continue.

Specifying surveillance period data, General tab

The **General** tab contains the surveillance protocol option (Light or Standard), the denominator data (optional in Standard surveillance, required in Light surveillance) and the other structure and process indicators: nurse-to-patient ratio and five indicators collected during an evaluation of practices (or audit) in the ICU. Table 14 describes the variables in this form. Further details are provided in the HAI-Net ICU protocol (see *Related documents*).



To define the ICU surveillance period:

- Specify values for the fields in the ICU surveillance period General tab. Descriptions of the variables in this form are given in Table 14. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- 2. Click the **Save** icon or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

3. Click **Select this ICU surveillance period** to be able to enter Patient/ Antimicrobial use/ HAI data.

Table 14: ICU surveillance period General tab variables

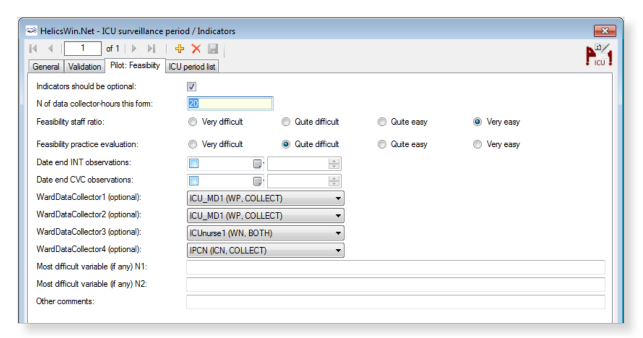
Variable	Description
Hospital code	Inserted automatically based on the hospital selected during Hospital definition.
Surveillance year	The year previously selected for the survey. Inserted by system.
ICU code	Unique identifier for each intensive care unit within a hospital, should remain identical in different surveillance periods/years. Inserted by system.
Start date *	Start date of the ICU surveillance period. Cannot be changed after saving.
End date *	End date of the ICU surveillance period. The recommended surveillance period is minimum 3 months, maximum 1 year. Can be entered later.
ICU protocol *	Select either Standard or Light . Cannot be changed after saving. Standard : Patient-based option allows advanced risk adjustment of healthcareassociated infection rates for inter-hospital comparisons. Risk factors are collected for each patient (infected or not). Light : Unit-based option is less labour-intensive solution, produces partially the same indicators as Standard option for follow-up of trends, as well as the same descriptive results about infections and antimicrobial resistance, but with less possibility for risk-adjusted comparisons. Denominator data are aggregated at the unit (ICU) level. Case definitions, included patients, and infection data (including antimicrobial resistance data and mortality review data) are identical in both versions. Refer to the HAI-Net ICU protocol (see <i>Related documents</i>) for further details.
Number of new ICU admissions (all) *	Total number of new admissions in the intensive care unit during the period. Used for burden estimates of HAIs in ICUs, assessing the ICU workload for patients staying 1 or 2 days in the ICU and comparing some indicators with ICU surveillance systems that include all ICU patients. Optional, but strongly recommended.
Number of ICU patient-days (all patients) *	Total number of patient-days in the intensive care unit during the period. Used for burden estimates of HAIs in ICUs, assessing the ICU workload for patients staying 1 or 2 days in the ICU, comparing some indicators with ICU surveillance systems that include all ICU patients and cross-checking the plausibility of the denominator of the alcohol hand rub consumption and nurse-to-patient ratio indicators. Optional, but strongly recommended.
Number of new ICU admissions staying > 2 days *	Required for light surveillance. Number of new admissions of patients staying more than 2 days in the intensive care unit during the period. Main denominator for the indicator 'cumulative incidence of HAIs'. In standard surveillance, this variable is optional and allows verifying the exhaustiveness of the entered patient-based data.
Number of ICU patient-days for patients staying > 2 days *	Required for light surveillance. Number of patient-days for patients staying more than 2 days in the intensive care unit during the period. Main denominator for the indicator 'incidence density of HAIs'. In standard surveillance, this variable is optional and allows verifying the sum of patient-days reported on patient level.

Variable	Description
Observations/ patient chart review period	Start date and end date of the period during which HAI prevention and antimicrobial stewardship practices are evaluated.
Total number of registered nurse hours in ICU over 7 day period	Total number of hours of real presence of registered nurses during a period of 7 days, including hours of presence during the night (presence of 1 full-time nurse 24/7=168 hours). Only include registered nurses involved in bedside patient care. Students are not included. A 'registered nurse' is a nurse who has graduated from a college's nursing program or from a school of nursing and has passed a national licensing exam to obtain a nursing license. Also include 'agency nurses', 'bank nurses', 'interim nurses' or other registered nurses who are not permanently employed for that position in the hospital.
Total number of nursing assistant hours in ICU over 7 day period	Total number of hours of real presence of nursing assistants during a period of 7 days, including hours of presence during the night (presence of 1 full-time nursing assistant 24/7=168 hours). Only include nursing assistants involved in bedside patient care. Students are not included. A 'nursing assistant' is also referred to as 'nurses' aide', 'healthcare assistant', 'nursing auxiliary', 'auxiliary nurse', 'patient care assistant' or similar terms. Also include nursing assistants who are not permanently employed for that position in the hospital.
Total number of patient-days over the same 7 day period	Total number of patient-days (all patients) over the same 7 days used for the number of (registered/assistant) nurse hours. Short interruptions are not taken into account. Partial days count as one patient day.
Antimicrobial stewardship: Review antimicrobial therapy within 72 hours (chart review)	Verify, for 30 (minimum 20) consecutive patients with antimicrobial therapy whether the therapy was evaluated within 72 hours after the start of the antimicrobial and has been documented in the patient file. Only consider first empiric or documented antimicrobial therapies that were started in the current ICU. Only systemic antimicrobial therapy (IV, IM, SC, oral) started since more than 72 hours are eligible for evaluation. Number of observations (denominator) = Total number of audited antimicrobial therapies that were started more than 3 days ago; Number compliant (numerator) = Number of antimicrobial therapies that were started more than 3 days ago and were reassessed within 72 hours after start of the antimicrobial.
Intubation: Endotracheal cuff pressure controlled and/or corrected at least twice a day (chart review)	Numerator=Number of intubation days (days of patients with intubation) during which the endotracheal cuff pressure was verified and maintained between 20 and 30 cm H2O (and documented in the patient file) at least twice per day; Denominator=Total number of observed intubation days. Source: medical or nurse patient file, prospective review of 30 patient-days with intubation. One patient with intubation is included only once a day, but the same patient can be included for several consecutive days.
Intubation: Oral decontamination using oral antiseptics at least twice a day (chart review)	Numerator=Number of intubation days (days of patients with intubation) during which oral decontamination with oral antiseptics has been performed (and documented in the patient file) at least twice per day; Denominator=Total number of observed intubation days. Source: medical or nurse patient file, prospective review of 30 patient-days with intubation. One patient with intubation is included only once a day, but the same patient can be included for several consecutive days.
Intubation: Position of the patient not supine (direct observation)	Numerator=Number of days of patients with intubation during which the patient's position was not supine (= was either prone or recumbent); Denominator=Total number of observed intubation days. Source: Direct observation of the position of the patient with intubation (in bed), up to 30 patient observations. One patient with intubation is included only once a day, but the same patient can be included for several consecutive days. Observations should as much as possible be perform at the same time during the day (e.g. at 16:00 in the afternoon). Patients in strict supine (dorsal decubitus) position for specific indications (e.g. certain trauma patients) should be excluded.

Variable	Description
CVC: Catheter site dressing is not damp, loose or visibly soiled (direct observation)	Numerator = Number of days of patients with a central vascular catheter during which the dressing of the CVC was not loose, damp or visibly soiled; Denominator = Total number of observed CVC days. Source: Direct observation of 30 patients with at least one CVC in place, up to 30 patient observations. One patient with one or several CVCs is included only once a day, but the same patient can be included for several consecutive days. Observations should as much as possible be perform at the same time during the day (e.g. at 16:00 in the afternoon). For patients with several CVCs in place, all CVC dressings need to be ok (not loose, damp nor visibly soiled).
Comments/ observations	Free text field.

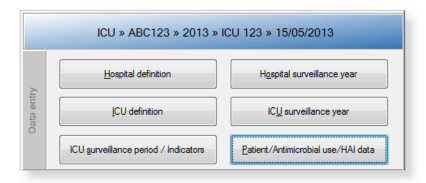
Specifying surveillance period data, Pilot: Feasibility tab

The **Pilot: Feasibility** tab contains the surveillance-year part of the feasibility questionnaire (Form F) of the HAI-Net ICU pilot study. Full descriptions of all the variables in this form are provided in the HAI-Net ICU protocol (see *Related documents*). Data collectors should first be defined in the **User** tab of the hospital-year screen, by clicking the **Define data collectors** button (see *User tab*). Please remember to include User variables when you export the data.



Entering ICU patient, antimicrobial use and HAI data

To enter patient data for antimicrobial use and HAI, click **Patient/Antimicrobial use/HAI data** in the *main menu*.



General tab

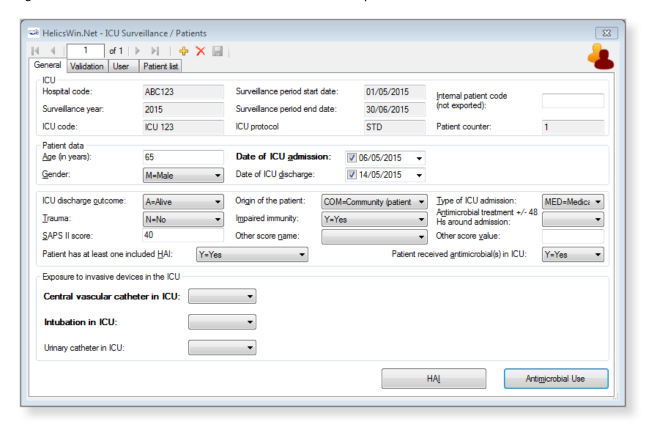
When you select **Patient/Antimicrobial use/HAI data** in the *main menu*, the **Patients** forms open at the **General** tab.

As in the PPS, the appearance of the **General** tab depends on whether Standard or Light surveillance has been chosen.

Unlike PPS, antimicrobial use data in the ICU module are optional and can only be entered in Standard surveillance.

Table 15 below describes the variables in the Standard and Light forms. Further details are provided in the HAI-Net ICU protocol (see *Related documents*).

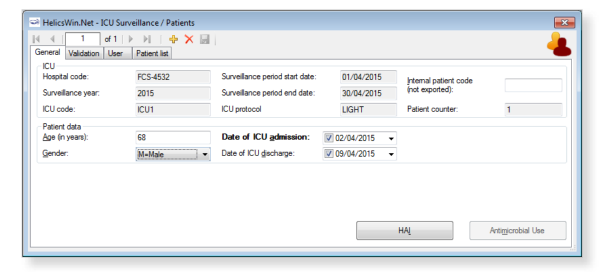
Figure 5: ICU Surveillance/Patients General tab - Standard protocol



In Standard surveillance, the **General** tab shows all the risk factors that should be entered for each patient staying more than two days in the ICU (with or without an infection).

In Standard surveillance the **HAI** button is deactivated when the variable **Patient has at least one included HAI** is set to **No**.

Figure 6: ICU Surveillance/Patients General tab - Light protocol



In Light surveillance, the **General** tab only shows a few demographic variables (see Table 15). These should only be entered for patients with an HAI. Consequently, a patient record in Light surveillance should always have at least one HAI record, and the **HAI** button is therefore activated by default each time a patient is created.

In Light surveillance, the **Antimicrobial use** button is deactivated.

To define the ICU patient, antimicrobial use and HAI data:

- Specify values for the fields in the ICU surveillance/patients General tab. Descriptions of the variables in this form are given in Table 15. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- 2. Click the **Save** icon III or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

- 3. Click the **HAI** or **Antimicrobial data** button to proceed to the respective dialog-box:
 - HAI data
 - Antimicrobial use data

Table 15: ICU patients General tab variables (Standard and Light protocols)

Variable	Description
Hospital code	Inserted automatically based on the hospital selected during Hospital definition.
Surveillance year	The year previously selected for the survey. Inserted by system.
Surveillance period start date	Start date of the ICU surveillance period. Inserted by system. Cannot be changed after saving.
Surveillance period end date	End date of the ICU surveillance period. Inserted by system. Can be blank if not yet entered.
ICU code	Identifier for the selected intensive care unit. Inserted by system.
Internal patient code	For hospital internal use only. Optional. Not exported. Must be removed from any data submitted to ECDC/TESSy.
Patient counter	A system allocated sequential patient number for anonymised identification.
Age *	Age of the patient on the date of admission to the ICU (in years).
Gender *	Gender of the patient. M = Male; F = Female; O = Other; UNK = Unknown.
Date of ICU admission *	Date of admission in the ICU. Mandatory – data cannot be saved without this date.
Date of ICU discharge	Date the patient was discharged from the ICU or date of in-ICU death or date of last follow-up in the ICU.
ICU discharge outcome	Patient status at discharge from the ICU or at end of follow-up in the ICU. A = Alive; D = Dead in ICU; UNK = Unknown.
Origin of the patient	Origin of the patient at the time he/she was admitted at the ICU: HOSP = Ward in this/other hospital. OICU = Other ICU; COM = Community (patient came from his home, via emergency or not); LTC = Long-term care/nursing home; O = Other; UNK = Unknown.
Type of ICU admission	Type of admission as defined in SAPS II score: (medical: no surgery within one week of admission to ICU; scheduled surgical: surgery was scheduled at least 24 hours in advance +/- 7 days ICU admission; unscheduled surgical: patients added to the operating room schedule within 24 hours of the operation. MED = Medical; SSUR = Scheduled surgical; USUR = Unscheduled surgical; UNK = Unknown.
Trauma patient	Intensive care unit admission resulted from blunt or penetrating traumatic injury to the patient, with or without surgical intervention. $Y = Yes; N = NO; UNK = Unknown$.

Variable	Description
Impaired immunity	Impaired immunity as defined in APACHE II score: impaired immunity due to treatment (chemotherapy, radiotherapy, immune suppression, corticosteroids long duration or high doses recently), due to disease (leukaemia, lymphoma, AIDS), or < 500 PMN/mm3. Y = Yes; N = NO; UNK = Unknown.
Antibiotic treatment +/- 48 hours around admission	Specify 'yes' if any antibiotic therapy in the 48 hours preceding ICU admission and/or during the first two days of ICU stay (=antibiotic therapy for an infectious event around ICU admission, excl. antifungal and antiviral treatment) has been given; not: antimicrobial prophylaxis, SDD, local treatment. Y = Yes; N = NO; UNK = Unknown.
SAPS II score	Simplified Acute Physiology Score II on admission (first 24h of ICU stay). Severity of illness score developed to predict mortality. Integer number from 0 to 163.
Other score name	Add alternative severity of illness score.
Other score value	Corresponding value for alternative severity of illness score. Possible scores [and possible values]: APACHE II [0-71], APACHE III [0-299], APACHE IV [0-286], MPM II [0-100], MPM III [0-100], McCabe score [0=non-fatal (survival >= 5 years); 1=ultimately fatal (survival < 5 years), 2=rapidly fatal (survival<1 year); 9=unknown], SAPS 3 [0-217]
Central vascular catheter in ICU *	Patient had a central vascular catheter during the current ICU stay. Y = Yes; N = NO; UNK = Unknown. Mandatory. If yes, enter dates for the corresponding exposure period.
Intubation in ICU *	Patient was intubated (invasive respiratory device) during the current ICU stay. Y = Yes; N = NO; UNK = Unknown. Mandatory. If yes, enter dates for the corresponding exposure period.
Urinary catheter in ICU	Patient had indwelling urinary catheter during the current ICU stay. Y = Yes; N = NO; UNK = Unknown. Required if UTI is included in surveillance. If yes, enter dates for the corresponding exposure period.
Patient received antimicrobial(s) in ICU	Patient received any antimicrobials during ICU stay. If yes, fill corresponding antimicrobial use data. Y = Yes; N = NO; UNK = Unknown.
Patient has at least one included HAI *	Patient has at least one healthcare-associated infection (with onset on day three or later, see definition) included in the current surveillance-year. If yes, fill out an HAI form for each infection. Y = Yes; N = NO; UNK = Unknown.

Antimicrobial use data

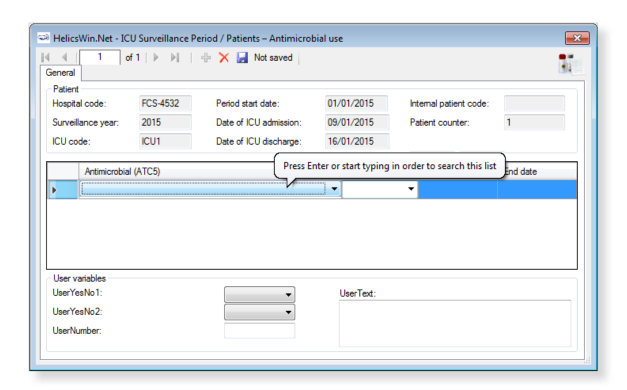
Antimicrobial use data are optional in the ICU module and can only be entered in Standard surveillance. Data entry requires that:

 The variable Patient received antimicrobial(s) in ICU is set to Yes the ICU General tab.

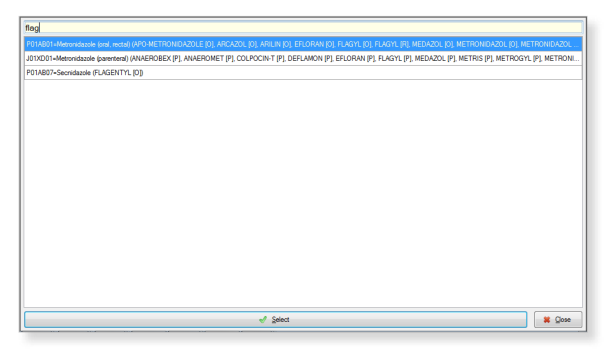
AND

 The option for antimicrobial use data has been activated in the ICU surveillance year form.

To enter antimicrobial use data, click the **Antimicrobial use** button in the ICU **General** tab to open the **Antimicrobial use** form. The fields in the form are described in Table 16 below.



Example of searching for an ATC5 code using a brand name:



To enter antimicrobial use data:

- 1. Click the **Antimicrobial use** button in the ICU **Surveillance/patients General** tab to open the form.
- **2.** Specify values for the fields in the the **Antimicrobial use General** tab. Descriptions of the variables in this form are given in Table 16 below. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

- **3.** Search for antimicrobials in the dropdown list by pressing enter when the cursor is placed over the **Antimicrobial (ATC5)** field. You can search on the ATC5 code, antimicrobial generic name, or any of the brand names included in the database.
- **4.** Click the **Save** icon **I** or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

Table 16: Patient's antimicrobial use

Variable	Description
Header section	See Table 15.
Antimicrobial (ATC5) *	Antimicrobial coded as ATC5 code. Include ATC2 classes J01 antibacterials, J02 antifungals and ATC4 A07AA, P01AB, D01BA and ATC5 J04AB02. See ATC5 list in the HAI-Net ICU protocol Annex 4 (see <i>Related documents</i>).
Indication	Indication for use of this antimicrobial episode. If the indication changes (e.g. from empiric treatment to documented treatment), enter a new line, even if the antimicrobial has not changed. If the same antimicrobial (ATC5 code) is used for different indications, enter a line for each indication. P=Prophylaxis; E=Empiric treatment (not based on microbiological results); M=Documented treatment (based on microbiological results with or without antimicrobial susceptibility results); S=Selective digestive decontamination; O=Other; UNK=Unknown.
Start date	Start date within the ICU of this antimicrobial agent/indication (days before ICU admission should not be reported). For antimicrobials present on admission, enter date of ICU admission.
End date	End date within the ICU of this antimicrobial agent/indication (days after ICU discharge should not be reported). For antimicrobials continued after discharge, enter date of ICU discharge.
User variables	As defined in ICU Surveillance/Patients Users tab.

HAI data

HAI, microorganism and antimicrobial resistance data are entered through the **Patients - Healthcare-associated infections (HAI)** form. The fields in the form are described in Table 17.

To enter HAI, microorganism and antimicrobial resistance data:

- 1. Click the **HAI** button in the ICU **Surveillance/patients General** tab to open the **HAI** form.
- 2. Specify values for the fields in the **ICU surveillance/patients HAI General** tab. Descriptions of the variables in this form are given in Table 17 below. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.
- **3.** Enter the **Case definition code** and **Date of infection onset** data (mandatory).
- **4.** Select the relevant device status from the drop-down list.
- **5.** Click the **Save** icon **J** or press **Ctrl+S**.

If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

Once the HAI record is saved, the microorganism sub-form is activated.

- **6.** Click on the drop-down list in the **Microorganism** field or enter the first letters of a microorganism to select a microorganism.
 - Once the organism is selected, the corresponding AMR markers are automatically displayed.
- 7. Enter the susceptibility results in the corresponding SIR1, SIR2, etc. fields.
- **8.** Select the relationship of the HAI to the patient outcome from the drop-down list at the bottom of the form.
- **9.** Click the **Save** icon I or press **Ctrl+S**.

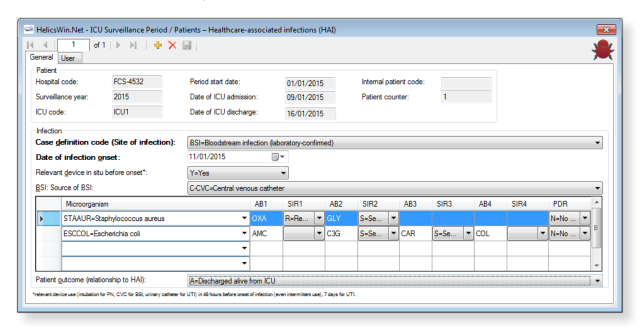


Table 17: ICU patient HAIs General tab variables

Variable	Description
Header section	See Table 15.
Case definition code (Site of Infection) *	Site of infection according the case definition (including subcategory), taking into account signs and symptoms of the entire infection episode (not just day one of the HAI). See Chapter 3 for case definitions. BSI = Bloodstream infection; PN = Pneumonia (unknown subcategory); PN1 = Pneumonia (protected sample + quantitative culture); PN2 = Pneumonia (non-protected sample (ETA) + quantitative culture); PN3 = Pneumonia (alternative microbiological criteria); PN4 = Pneumonia (sputum bacteriology or non-quantitative ETA); PN5 = Pneumonia (no microbiology); UTI = Symptomatic urinary tract infection (unknown subcategory); UTI-A = Symptomatic urinary tract infection (microbiologically confirmed); UTI-B = Symptomatic urinary tract infection (not microbiologically confirmed); CRI1-CVC = CVC-related infection (local); CRI2-CVC = CVC-related infection (generalised no positive haemoculture); CRI3-CVC = CVC-related infection (generalised with positive haemoculture). If (optional) catheter-related infections (CRIs) are included in the surveillance, report a CVC-related BSI corresponding to the case definition of CRI3-CVC as CRI3-CVC (do not report twice).
Date of infection onset *	Date of onset of symptoms or, if unknown, date treatment was started or date first diagnostic examination was done.

Variable	Description
Relevant device in situ before onset *	Relevant invasive device was present (even intermittently) in the 48 hours preceding the infection (7 days for UTIs): intubation for pneumonia, central vascular catheter for bloodstream infection, urinary catheter for urinary tract infections. Necessary to distinguish device-associated infections. Y = Yes; N = No; UNK = Unknown.
BSI: Source of BSI	Required if the case definition code is <i>BSI</i> . Source/origin of the bloodstream infection. C = The same microorganism was cultured from the catheter or symptoms improve within 48 hours after removal of the catheter. Exception: Report microbiologically confirmed CVC-related BSI as CRI3-CVC if optional CRIs are included in the surveillance. C = Catheter, catheter type unknown; C-CVC = Central venous catheter; C-PVC = Peripheral venous catheter; C-ART = Arterial catheter; S = Secondary to another site, primary site unknown; S-PUL = Pulmonary infection; S-UTI = Urinary tract infection; S-SSI = Surgical site infection; S-DIG = Digestive tract infection; S-SST = Skin/Soft Tissue infection; S-OTH = Other infection or procedure; UO= None of the above, BSI of unknown origin; UNK=Unknown/Missing.
Microorganism *	Isolate result. Microorganism (MO) six letter code or negative code including reason why the isolate result is not availableNA = Results not available; _NOEXA = Examination not done; _NONID = Microorganism not identified; _STERI = Sterile examination. See Code list in the HAI-Net ICU protocol Annexes 2 and 3 (see Related documents). It is recommended to use the extended microorganism list, even though minimal list codes are also allowed. Minimum one code per HAI is Required, the recommended maximum per HAI is 3 microorganisms.
АВ	Antibiotic code. Antimicrobial drug for which susceptibility was tested, depends on the microorganism. In HWN, recommended antimicrobial codes are automatically generated.
SIR *	Final interpretation result of all different susceptibility tests performed. If antibiotic code is $_NOTEST$, $SIR=NA$. $S=Susceptible$; $I=Intermediate$; $R=Resistant$; $UNK=Unknown$; $NA=Not$ applicable.
Patient outcome (relationship to HAI)	 Relationship of HAI to ICU outcome in patients with HAI: Discharged alive: patient was discharged alive; OR patient was still in the hospital and alive at end of follow-up during this hospital stay. Death, HAI definitely contributed to death: use this category if a causal link between CDI and death can be demonstrated. Death, HAI possibly contributed to death: use this category if no causal link between CDI and this case's death can be demonstrated, but it is still plausible that CDI was at least a contributory factor. Death, unrelated to HAI: use this category if the cause of death can be demonstrated not to be related to CDI. Death, relationship to HAI unknown: use this category if no evidence of contributory factors to the cause of death is available. Unknown: unknown patient outcome.

ICU validation tabs

The **Validation** tabs in the **ICU** forms are not currently activated for HWN2.2.

Creating a PPS point prevalence survey

There are two Point Prevalence Survey (PPS) options:

- **Standard**: Patient-based option allows advanced risk adjustment of healthcare-associated infection rates for inter-hospital comparisons. You enter all patient data for all patients, including those without HAI and/or antimicrobial use. Risk factors are collected for each patient (infected or not).
- **Light:** Unit-based option is less labour-intensive solution, produces partially the same indicators as Standard option for follow-up of trends, as well as the same descriptive results about infections and antimicrobial resistance, but with less possibility for risk-adjusted comparisons. You only enter patient data for patients with any antimicrobial use and/or a healthcare-associated infection. Consequently, in the Light protocol, each patient record must include details of at least one antimicrobial use or HAI.

Both options use the hospital data forms $\mathbf{H1} ext{-}\mathbf{H3}$, the ward data form \mathbf{W} , and the national data form \mathbf{N} .

However, the Standard option uses patient data form ${\bf A}$, whereas the Light option uses numerator data form ${\bf B}$.

Further details are provided in the HAI-Net PPS protocol (see Related documents).

PPS data collection is described in the following sections:

- Entering PPS hospital data
- Entering ward PPS data
- Entering PPS patient, antimicrobial use and HAI data

Entering PPS hospital data

Follow the procedure in this section to create a new **PPS** survey for each defined hospital, entering information from Form H. Before you can create your survey, you must *define your hospital and wards*.

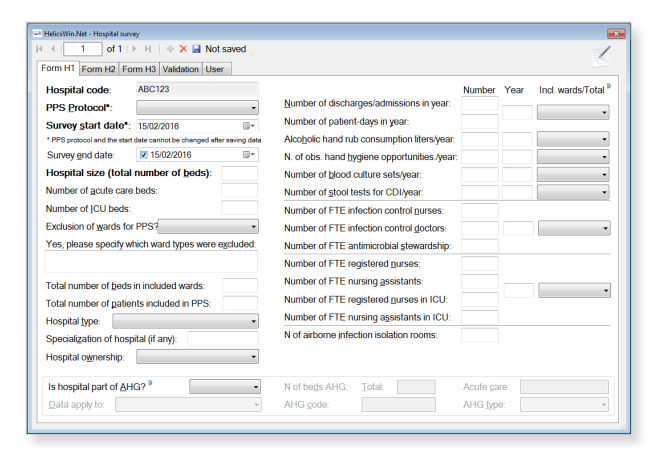
To create a hospital Point Prevalence Survey:

1. Click Hospital surveys in the PPS main menu.



The **Hospital survey** form opens (for the hospital you have selected).

- **2.** If not already open, click the **Form H1** tab.
- 3. In the Form H1 tab, click the Add item icon $\frac{1}{2}$ to open the form for editing.



4. Enter the values for the fields in the **Form H1** tab.

Note that the **Incl. wards / Total** options (for included wards only OR for hospital total) are required fields.



- **5.** Click the **Save** icon **J** or press **Ctrl+S**.
- **6.** Enter the values for the fields in the **Form H2** and **Form H3** tabs, and save.

Descriptions of the variables for Hospital survey tabs Form H1, 2, and 3 are given in Table 18. If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

Table 18: PPS Hospital survey tabs (Form H1, 2 and 3) variables

Variable	Description
Form H1 tab	
Hospital code *	The hospital identifier/code assigned by national/regional PPS coordinating centre; unique code per surveillance/PPS network. Hospital codes should be unique within each surveillance network, and kept constant between the ECDC Antimicrobial Resistance and Healthcare-Associated Infection (ARHAI) surveillance protocols and from one year to the next. Inserted automatically based on the hospital selected during Hospital definition.
PPS protocol *	Select either Standard , Light or Validation . Note that you cannot add Light protocol
	data records to a PPS data collection defined to use the Standard protocol, or vice versa. If the current survey or surveillance year is a validation study performed by a validation team, Validation should be selected to indicate that this is not a primary data collection but a repeated data collection for validation purposes of the primary PPS data. Selecting Validation will change the messages on data entry for the entire application,
	especially for the variables displayed on the Validation tabs at different levels (see Validation tab below).
	Warning: You cannot change the PPS protocol after you have saved it.
Survey start date	Start and end dates for the PPS in the entire hospital.
*	Warning: You cannot change the Survey start date after you have saved it. If you save without having selected the correct Survey start date , today's date is applied by default.
	Before saving, make sure that the start date of the hospital PPS is earlier than any ward survey date. If you need to change the survey start date, you will have to contact your national or regional PPS co-ordinator.
Survey end date	The date the last data were collected in the last ward. You can enter the Survey end date at a later time.
Hospital size *	Total number of beds in the hospital. If, for reasons of confidentiality, the exact number of hospital beds cannot be given, please enter the number of beds rounded up to the nearest 50.
Number of acute care beds	Number of acute care beds in the hospital (according to national definition).
Number of ICU beds	Number of intensive care unit beds in the hospital. No ICU=0.
Exclusion of wards for PPS?	Were any wards excluded for the PPS in your hospital? Yes/No.
Specify excluded wards	Specify which wards where excluded, if any. Free text; please use specialty codes if possible.
Total number of beds in included wards *	Sum of the number of beds in wards that were included in the PPS.
Total number of patients included in PPS *	Sum of the number of patients included in the PPS.
Hospital type	Hospital type – PRIM: primary, SEC: secondary, TERT: tertiary, SPEC: specialised, missing=UNK. See <i>Hospital types</i> .
Specialisation of hospital	Free text. Indicate the hospital specialty for a specialised hospital (e.g. paediatric, infectious diseases, etc.); please use specialty codes if possible.

Variable	Description
Hospital ownership	Hospital ownership as defined by WHO/Europe [7], Eurostat [8] and OECD [9]: PUB: Public, PRIVNFP: Private, not-for-profit, PRIVFP: Private, for-profit, OTHUNK: Other or unknown. See protocol for definitions.
Is hospital part of AHG?	The hospital is part of an administrative group of hospitals (AHG, also referred to as hospital "trust", "merger", "fusion", "board", "region" etc.). Yes/No.
Data apply to	If the hospital is part of an AHG, data apply to a single hospital (hospital with a single address, or a hospital site belonging to a trust) (S) OR a group of hospitals (including entities referred to as 'trusts' or 'chain') (T).
N of beds AHG	Total number of beds of the administrative hospital group.
AHG code	
Acute care	Total number of acute care beds of the administrative hospital group.
AHG type	If hospital is part of AHG, what is the hospital type of the administrative group: PRIM: primary, SEC: secondary, TERT: tertiary, SPEC: specialised (see above for definition of hospital type).
Incl. wards / Total *	Included wards only OR total for hospital. Select INC or TOT in the drop-down lists.
Number of discharges/admissions *	Number of hospital discharges in a given year (data from previous year if available, specify year in second column), use number of admissions if discharges are not available. Provide the number for the included wards only (if not available, provide number for entire hospital; specify 'included wards only OR total for hospital' in last column).
Number of patient-days *	Number of hospital patient-days in a given year (data from previous year if available, specify year in second column). Provide data for the same year and wards (included wards only OR total for hospital) as for the number of discharges/admissions.
Alcoholic hand rub consumption *	Total number of litres of alcoholic hand rub used in a given year (data from previous year if available, specify year in second column); provide the number for the included wards only (if available, otherwise provide number for the entire hospital; specify 'included wards only OR total for hospital' in last column).
Number of observed hand hygiene opportunities	Number of observed hand hygiene opportunities performed in the previous year if available (or the most recently available year). Report the total number of observed opportunities for hand hygiene, not only the compliant observations.
Number of blood cultures per year	Number of inpatient blood culture sets tested by the microbiological laboratory for the current hospital in a period of one year. Provide data for previous year or the most recently available data (specify year data in a separate variable). If the number of blood culture sets is not directly available, estimate by the [total number of blood culture bottles processed] divided by the [total number of bottles per blood culture request]. Count all blood culture tests per patient, not the number of patients for whom >=1 test was performed. Count the number of blood culture sets actually processed by the laboratory, not the number sent to the laboratory for analysis.
Number of stool tests for CDI per year	Number of inpatient stool tests performed for <i>Clostridium difficile</i> infections (CDI) per year. Provide data for previous year or the most recently available data (specify year data in a separate variable). Count all stool specimens per patient, not the number of patients for whom >=1 test was performed. Count the number of stool specimens actually processed by the laboratory, not the number sent to the laboratory for analysis.
Number of FTE infection control nurses	Number of full-time equivalent (FTE) infection control nurses in the hospital; infection control nurse=nurse with specialised training in infection control/hospital hygiene and usually responsible for infection control/hospital hygiene tasks such as training of hospital employees in infection control, elaboration and implementation of infection control procedures, management (implementation, follow-up, evaluation) of an infection control work plan and projects, audits and evaluation of performance, procedures for disinfection of medical devices etc. (see TRICE project report). Specify year of data collection (current year if available) and whether the number of FTE infection control nurses is provided for the entire hospital or only for the included wards.

Variable	Description
Number of FTE infection control doctors	Number of full-time equivalent (FTE) infection control doctors (or pharmacists, hospital epidemiologists, etc.) in the hospital with specialised training in infection control/hospital hygiene and usually responsible for infection control/hospital hygiene tasks such as identification and investigation of outbreaks, analysis and feedback of infection control data, elaboration of an infection control work plan and projects, design and management of surveillance systems, elaboration of infection control procedures etc. (see TRICE project report). Please ensure that the reported number was collected for the same year and wards (included wards only OR total for hospital) as the number of FTE infection control nurses.
Number of FTE antimicrobial stewardship	Number of full-time equivalent (FTE) antimicrobial stewardship consultants in hospital. FTE antimicrobial stewardship = dedicated time of an ('external') consultant employed by the hospital and specifically paid for antimicrobial stewardship tasks, NOT the time spent by treating physicians on antimicrobial stewardship activities (e.g. post-prescription review) as part of their daily practice. Deduct FTE from FTE infection control doctor if same person: in case antimicrobial stewardship tasks are an integral part of the job description/daily activities of the infection control doctor (or equivalent), the estimated FTE (proportion of his/her time) spent on antimicrobial stewardship activities should be deduced from the FTE infection control doctors and be reported separately.
Number of FTE registered nurses	Number of full-time equivalent (FTE) registered (graduated, qualified) nurses in hospital. A 'registered nurse' is a nurse who has graduated from a college's nursing program or from a school of nursing and has passed a national licensing exam to obtain a nursing license. Also include 'agency nurses', 'bank nurses', 'interim nurses' or other registered nurses who are not permanently employed for that position in the hospital. Students are not included. Provide current situation if possible, or the situation for the earliest available year (specify year) and specify if the number of FTEs is given for the entire hospital or for the included wards only.
Number of FTE nursing assistants	Number of full-time equivalent (FTE) nursing assistants in hospital. A 'nursing assistant' is also referred to as 'nurses' aide', 'healthcare assistant', 'nursing auxiliary', 'auxiliary nurse', 'patient care assistant' or similar terms. Also include nursing assistants who are not permanently employed for that position in the hospital. Students are not included. Provide current situation if possible, or the situation for the earliest available year (specify year) and specify if the number of FTEs is given for the entire hospital or for the included wards only.
Number of FTE registered nurses in ICU	Number of full-time equivalent registered (graduated, qualified) nurses in intensive care unit(s). See also above.
Number of FTE nursing assistants in ICU	Number of full-time equivalent nursing assistants in intensive care unit(s). See also above.
Number of airborne infection isolation rooms	Number of airborne infection isolation rooms in the hospital. An airborne infection isolation room is defined as a hospital room provided with negative pressure and an anteroom.
Form H2 tab	
Annual IPC plan	Is there an annual infection prevention and control (IPC) plan and if so, was it approved by the hospital Chief Executive Officer (CEO, Managing Director) or by a senior executive officer? Yes/No/Unknown.
Annual IPC report	Is there an annual infection prevention and control (IPC) report and if so, was it approved by the hospital Chief Executive Officer (CEO, Managing Director) or by a senior executive officer? Yes/No/Unknown.
Does your hospital have a multimodal strategy for the prevention of following infections?	Indicate for each of the main HAI types and for antimicrobial stewardship which components of a multimodal strategy are available at the hospital-wide level and specifically in intensive care (presence in at least one adult, paediatric or neonatal ICU). Each cell of the table is a Yes/No/Unknown variable (28 variables for ICU + 35 variables hospital-wide/other non-ICU wards): mark Y=Yes, N=No or U=Unknown or not assessed in each cell. A multimodal strategy is defined as an intervention aiming at improving practice and offering education and training at multiple levels (e.g. written information, leaflets, posters, bedside teaching, workshops, focus groups, knowledge tests, competency assessments, surveillance and feedback, audits, checklists, care bundles).

Variable	Description
	The strategy must be rooted in written guidelines. Simple information sessions (e.g. for new staff), updating guidelines, or target setting alone (even if communicated to staff but without combining it with education and training) are not multimodal strategies.
Participation in surveillance networks	Indicate (Yes/No) if your hospital participates in a national or regional surveillance network for each of following surveillance modules: surveillance of surgical site infections (SSI), surveillance of HAIs in intensive care (ICU), surveillance of C. difficile infections (CDI), surveillance of antimicrobial resistance according to the EARS-Net protocol (surveillance of antimicrobial resistance in invasive isolates of S. pneumonia, S. aureus, Enterococcus spp., E. coli, K. pneumoniae, P. aeruginosa and/or A. baumannii), surveillance of antimicrobial consumption in the hospital (surveillance at 5th ATC level in defined daily dose (DDD) per 1 000 patient-days) and other HAI or AMR surveillance modules (national/regional protocols for which a European/ECDC protocol does not exist). Local surveillance without transmission of data to a national or regional surveillance coordination centre for comparative analysis and feedback is not sufficient.
Microbiology/ diagnostic performance	Indicate whether clinicians can request routine microbiological tests and receive back results during weekends. Report yes/no/unknown separately for Saturdays and Sundays for clinical tests and screening tests respectively.
Comments	Free text field for additional information.
Form H3 tab	
Number of beds with AHR dispensers at point of care	Number of beds in the hospital with alcohol hand rub (AHR) dispensers available at the point of care as recommended by the 2009 WHO Guidelines on Hand Hygiene in Health Care. AHR dispensers at the entrance of the patient room only are NOT considered as 'available at the point of care'. The 'point of care' is the place where three elements come together: the patient, the HCW, and care or treatment involving contact with the patient or his/her surroundings (within the patient zone). The concept embraces the need to perform hand hygiene at recommended moments exactly where care delivery takes place. This requires that a hand hygiene product (e.g. alcohol-based hand rub, if available) be easily accessible and as close as possible — within arm's reach of where patient care or treatment is taking place. Point-of-care products should be accessible without having to leave the patient zone. Dispensers available at the point of care that happen to be empty on the PPS day should be included.
Number of beds assessed for the presence of AHR dispensers	The denominator of the previous variable, i.e. the total number of beds for which the presence of alcohol hand rub dispensers at the point of care was checked. If all wards were assessed, then this number is in principle the same as the total number of hospital beds.
Total number of patient rooms	Total number of rooms in included wards or total for hospital. Data from current year if available, specify year in second column; provide the number for the included wards only (if available, otherwise provide number for the entire hospital; specify 'included wards only OR total for hospital' in last column).
Number of single patient rooms	Total number of single-bed rooms in included wards OR total for hospital. Please ensure that the number of single patient rooms was collected for the same year and wards (included wards only OR total for hospital) as the total number of patient rooms. Rooms with more than one bed that are designated for use as single occupancy and isolation rooms (e.g. for infection control purposes) should be included.
Number of single patient rooms with individual toilet and shower	Total number of single-bed rooms with individual toilet and shower in included wards OR total for hospital. Please ensure that the number was collected for the same year and wards (included wards only OR total for hospital) as the total number of patient rooms. Rooms with more than one bed that are designated for use as single occupancy and isolation rooms (e.g. for infection control purposes) should be included. Rooms which have toilet and shower in a communal area should not be counted. An individual toilet alone or a commode (toilet chair) is not sufficient to qualify for this indicator.
Number of beds occupied at 00:01 on the day of the PPS	Number of hospital beds occupied at midnight on the day of the PPS. Since the PPS usually takes several days for an entire hospital, this variable can be measured on a day in the middle of the PPS data collection period, but not during the weekends.

Variable	Description
Number of beds assessed for occupancy at 00:01 on the day of PPS	Number of hospital beds that were checked for occupancy at midnight on the day of the PPS. Denominator of the previous variable. If occupancy was checked for all beds, this variable normally equals the total number of beds in the hospital.
Do healthcare workers carry AHR dispensers?	Percentage of healthcare workers in hospital that carry alcohol hand rub dispensers. In your hospital, do healthcare workers (HCW) carry AHR dispensers (e.g. in their pockets)? (If yes, please estimate percentage). Zero=0% Q0; >0-25% Q1; >25-50% Q2; >50-75% Q3; 75-100% Q4; Yes, percentage unknown YES; Unknown.
Is there a formal procedure to review antimicrobial within 72 hours?	Is there a formal procedure for a post-prescription review? I.e. to review the appropriateness of an antimicrobial within 72 hours (three calendar days) from the initial order in the hospital? A formal post-prescription review procedure should be documented and adopted by the hospital management and should be performed by a person or team other than the treating physician. The procedure should at least address the prescription of broad-spectrum or reserve antimicrobials. Choose one answer. YESALL = Yes, in all wards; YESSEL = Yes, in selected wards only (usually, but not necessarily, including ICU); YESICU = Yes, in ICU only; NO = No; UNK=Unknown.

Further hospital data can be recorded in the **Form H2** and **H3** tabs. The fields are described in Table 18. If any mandatory or required data field is empty when you save, e.g. because you do not yet have the data, an error and/or warning message is shown and the field label is highlighted as a reminder - in red for a mandatory field (you cannot save) or blue for a required field. See *Understanding reported errors*.

Figure 7: Hospital survey Form H2 tab

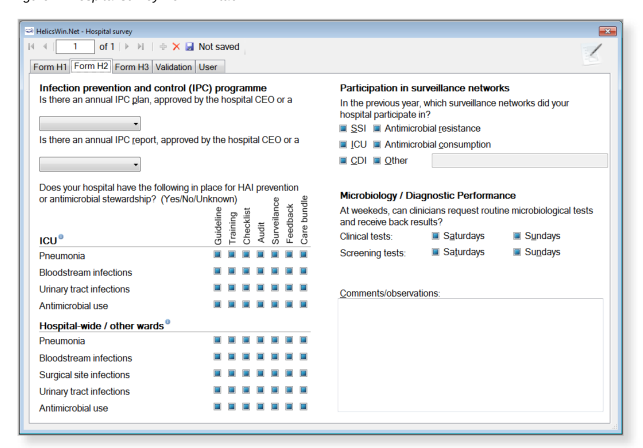
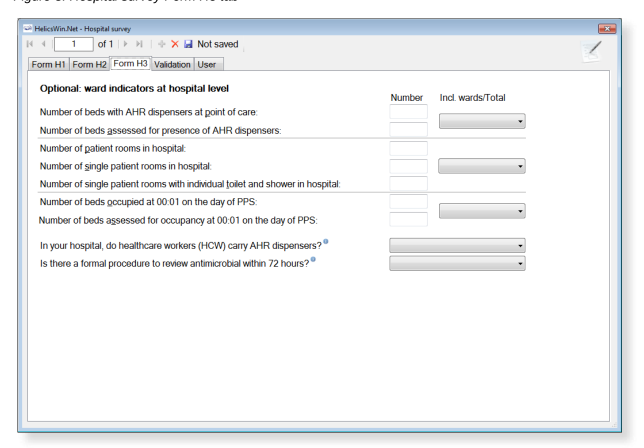
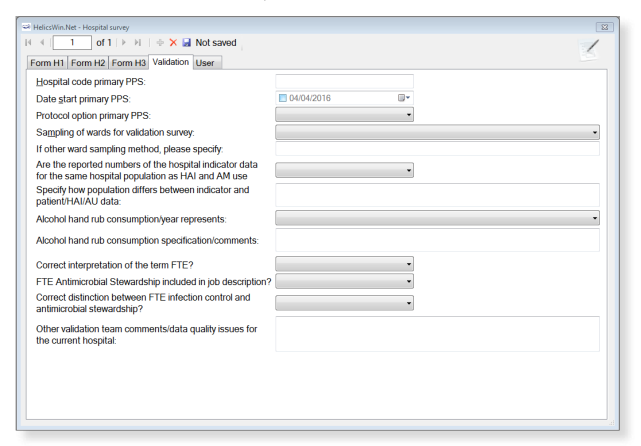


Figure 8: Hospital survey Form H3 tab



Validation tab

Validation teams use the **Validation** tab to enter data if a validation survey is performed. For the PPS module, use of this tab is described in a separate document, the **PPS Validation Protocol**.



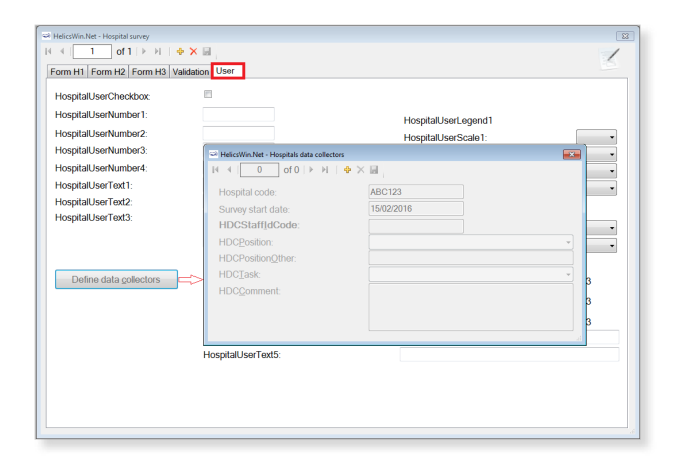
User tab

The **User** tab contains fields useful for users at national, regional or hospital level. They can be renamed and personalised to enable collection of information during the PPS that is outside of the PPS protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.

Additionally, the **User** tab of the **Hospital surveys** form also enables you to record the details of all hospital data collectors (HDC): their ID Code, Position (function) and survey-related task(s). This is optional.

You can also later assign the ID Code for the data collectors to the wards (see the **User** tab in the **Ward PPS data** form - Figure 9 on page 72).

Click **Define data collectors** to enter these data.



Entering ward PPS data

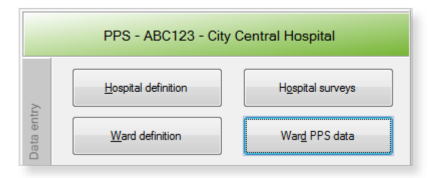
Follow the procedure in this section to create a new **PPS** survey for each defined ward, entering information from Form W. Before you can enter ward data, you must have *defined your surveys and wards*.

For each ward (unit) included in the PPS, you need to choose the ward and the date when the PPS was performed in that ward. In principle, a ward should be surveyed on a single day. However, you can enter more than one survey date for the same ward. The ward survey date and specialty are mandatory in both the **Light** and **Standard** protocols.

The ward PPS data are entered in the **Ward PPS data Form W and W (2)** tabs, the fields in which are described in Table 19.

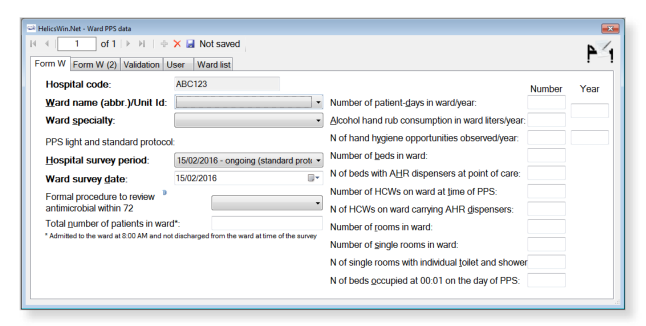
To specify the PPS date, specialty and denominator data:

1. Click Ward PPS data in the main menu.



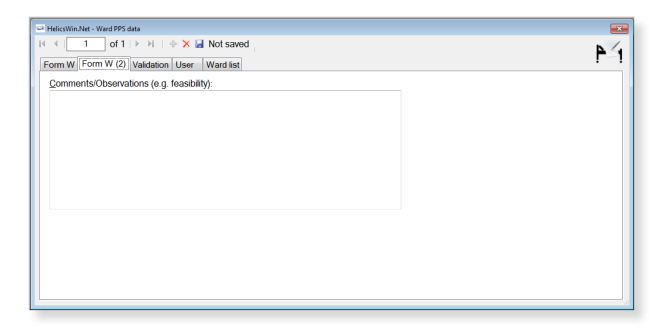
The **Ward PPS data** form opens showing the current hospital code.

2. Click the **Add item** icon $\frac{1}{2}$ or press **Ctrl+N** to open the form for editing.



You cannot change the hospital code from this form.

- **3.** For each ward (unit) included in the PPS, choose the **Ward specialty** from the drop-down list and enter the date the PPS was performed in that ward (i.e. the **Ward survey date**).
 - A ward survey should be completed within a single day, but if that is not possible, you can enter the other date(s) for the same ward on this form.
- **4.** Fill in the supplementary data for the ward, as relevant. In the **Form W (2)** you can add comments and observations for the ward, e.g. on the feasibility of the PPS.



- **5.** Click the the **Save** icon or press **Ctrl+S**.
- **6.** To start entering patient data in the Standard protocol, exit this form and click **Patient/Antimicrobial Use/HAI** data in the *main menu*.

The fields in the **Ward PPS Form W** tab are described in Table 19. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

Table 19: Ward PPS data Form W and W (2) tab variables

Variable	Description
Hospital code *	Inserted automatically based on the hospital selected during Hospital definition.
Ward name /unit ID *	Unique identifier for each unit (abbreviated ward name) within a hospital; should be used consistently on all forms (link with patient and HAI/AU data) and should remain unchanged in different PPS periods/years.
Ward specialty *	Main ward specialty (≥ 80% of patients require this specialty). If less than 80%, report 'mixed ward' (MIX). PED=Paediatric, NEO=Neonatal, ICU=Intensive Care, MED=Medicine, SUR=Surgery, GO=Gynaecology/Obstetrics, GER=Geriatrics, PSY=Psychiatry, RHB=Rehabilitation, LTC=Long-term care, OTH=Other, MIX=Mixed. Allows combination with patient specialty to refine specialties, e.g. paediatrics: ward specialty PED + patient specialty: PEDICU = paediatric ICU, NEOICU = neonatal ICU, SURCARD = paediatric cardiac surgery). A ward with healthy new-borns either must be allocated to GO (GOBAB) when it is located in obstetrics, or to PED (PEDBAB) if it is located in paediatrics. Note: how to code paediatric patients? Use the ward code PED for paediatric wards. If the ward specialty code is PED, then patients should be coded as per consultant/patient specialty MEDGEN, MEDSUR etc. The consultant/patient specialty PEDGEN should be used only for paediatric patients on adult wards.
Hospital survey period *	List of PPS surveys that the ward can participate in.
Ward survey date *	Date on which the data were collected in the ward. Data from a single ward should be collected on one day; date dd/mm/yyyy.

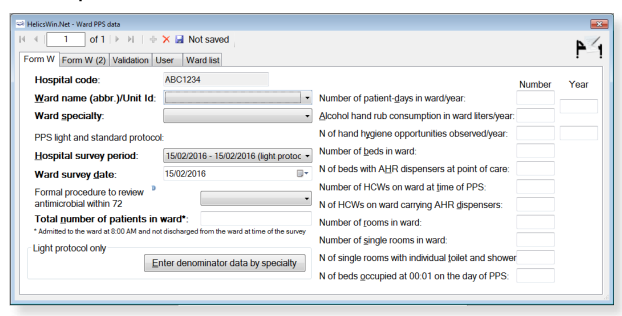
Variable	Description
Formal procedure to review antimicrobial within 72 hours?	Is there a formal procedure for a post-prescription review? I.e. to review the appropriateness of an antimicrobial within 72 hours (three calendar days) from the initial order in the hospital? A formal post-prescription review procedure should be documented and adopted by the hospital management and should be performed by a person or team other than the treating physician. The procedure should at least address the prescription of broad-spectrum or reserve antimicrobials. Yes/No/Unknown.
Total number of patients in ward	Total number of patients admitted to the ward before or at 8 a.m. that were not discharged from the ward at the time of the survey. Mandatory for light protocol, optional for standard protocol.
Number of patient-days in ward/year	Number of patient-days in one year for current ward (data from previous year if available, specify year in second column).
Alcohol hand rub consumption in ward liters/year	Number of liters of alcohol hand rub delivered to the ward in one year.
Number of hand hygiene opportunities observed in ward/year	Number of hand hygiene opportunities observed in the current ward in one year. Provide data for previous year if available or the most recent data available (specify year in second column). Report the total number of observed opportunities for hand hygiene, not only the compliant observations.
Number of beds in ward	Total number of beds in ward on the PPS day. Include 'corridor beds' and neonatal beds.
Number of beds in ward with AHR dispensers at the point of care	Number of beds in the ward with alcohol hand rub (AHR) dispensers available at the point of care as recommended by the 2009 WHO Guidelines on Hand Hygiene in Health Care. AHR dispensers at the entrance of the patient room only are NOT considered as 'available at the point of care'. The 'point of care' is the place where three elements come together: the patient, the HCW, and care or treatment involving contact with the patient or his/her surroundings (within the patient zone). The concept embraces the need to perform hand hygiene at recommended moments exactly where care delivery takes place. This requires that a hand hygiene product (e.g. alcohol-based hand rub, if available) be easily accessible and as close as possible – within arm's reach of where patient care or treatment is taking place. Point-of-care products should be accessible without having to leave the patient zone.
Number of HCWs on ward at time of PPS	Number of healthcare workers (HCWs) on ward at the time of PPS.
Number of HCWs on ward carrying AHR dispensers	Number of HCWs on ward carrying AHR dispensers (e.g. in their pocket).
Number of rooms in ward	Total number of rooms in the ward on the PPS day.
Number of single rooms in ward	Total number of single-bed rooms in the ward on the PPS day. Rooms with more than one bed that are designated for use as single occupancy and isolation rooms (e.g. for infection control purposes) should be included.
Number of single rooms with individual toilet and shower	Total number of single-bed rooms with individual toilet and shower in the ward. Rooms which have toilet and shower in a communal area should not be counted. An individual toilet alone or a commode (toilet chair) is not sufficient to qualify for this indicator.
Number of beds occupied at 00:01 on the day of PPS	Number of ward beds occupied at midnight on the day of the PPS (can also be measured at midnight after the PPS took place).
Comments/obser vations	(Form W (2) tab) Free text field to report e.g. feasibility issues, data quality problems or specific epidemiological information for the current ward.

Light protocol

The following instructions apply to the Light protocol only. In the Light protocol, the form has an additional mandatory field, **Total number of patients in ward**, and a button to enter the denominator data by consultant/patient specialty. The sum of the consultant/patient specialty denominators should be equal to the field **Total number of patients in ward**. This is verified during the *data quality check*, not during data entry.

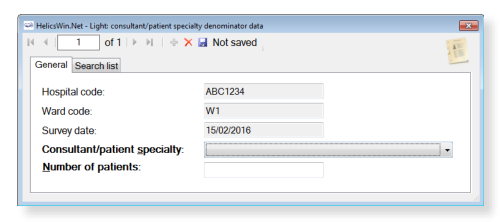
The ward PPS data are entered in the **Ward PPS data Form W** tab, the fields in which are described in Table 19. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

1. Enter the total number of included patients in ward, i.e. the denominator data in the **Total number of patients in ward** field.



2. Click **Enter denominator data by specialty** to enter denominator information separately for consultant(s)/patient speciality(s) within that same ward.

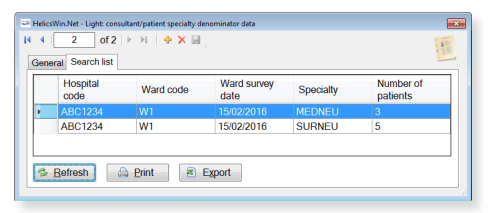
The Light: consultant/patient speciality denominator data form opens.



Warning: In the Light protocol, denominator data by ward AND by consultant/patient specialty are mandatory. If the detailed denominator data by consultant/patient

specialty are not known, enter at least one record with the total number of patients (using the unit specialty instead). Also make sure that the list of consultant/patient specialties entered in this form includes all the consultant/patient specialties that are used in the patient/AM/HAI form afterwards (this is verified by the *data quality check* before data export as well).

- 3. Click the **Add item** icon $\stackrel{1}{\leftarrow}$ or press **Ctrl+N** to open the form for editing.
- 4. For each ward (unit) included in the PPS, choose the Consultant/patient specialty from the drop-down list and enter the Number of patients of the selected specialty in that ward.
 Repeat for each different specialty within that ward.
- **5.** Finally, you can see the total number of patients entered for that ward in the **Search list** tab. Click **Refresh** to ensure that all recent data is included in this list. Remember that the sum of the consultant/patient specialty denominators should be equal to the field **Total number of patients in ward**.



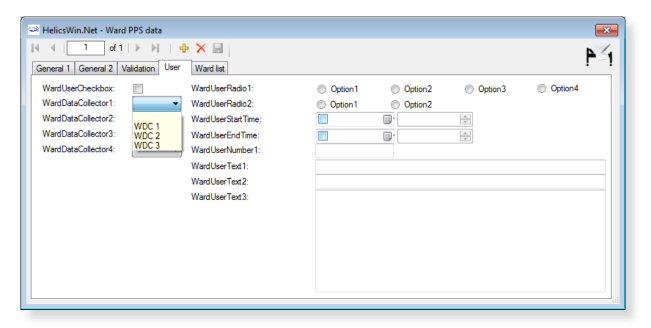
User tab

The **User** tab contains fields useful for users at national, regional or hospital level. They can be renamed and personalised to enable collection of information during the PPS that is outside of the PPS protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.

Optionally, choose to associate one or more data collectors with the ward.

ID codes of Ward data collectors entered in the **Hospital surveys** form will be present in drop down lists on the **User** tab. Optionally, choose one of these from each dropdown list, to associate them with the ward chosen in previous steps.

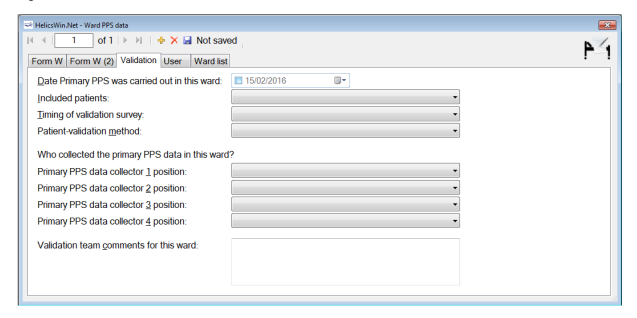
Figure 9: Ward PPS data - User tab



Validation tab

Validation teams use the **Validation** tab to enter data if a validation survey is performed. For the PPS module, use of this tab is described in a separate document, the **PPS Validation Protocol**.

Figure 10: Ward PPS data - Validation tab



Entering PPS patient, antimicrobial use and HAI data

The way in which you enter these data depends on the protocol being used:

• In the (unit-based) **Light protocol**, you only enter patient data for patients with any antimicrobial use and/or a healthcare-associated infection. Therefore, in the Light protocol, each patient record must include details of at least one antimicrobial use or HAI.

• In the (patient-based) **Standard protocol**, you enter all patient data for all patients, including those without HAI and/or antimicrobial use. Additional risk factors (not applicable to the Light protocol) appear once you select the ward.

Note:

- Antimicrobial use data and HAI data can only be accessed from the Patients form.
- Only records which have HAI/AU specific data entered will be included in prevalence results during final analysis, e.g. in reports generated by ECDC after upload to the TESSy database.
- The Antimicrobial use button is activated when the field Patient receives antimicrobial(s): is set to Yes.

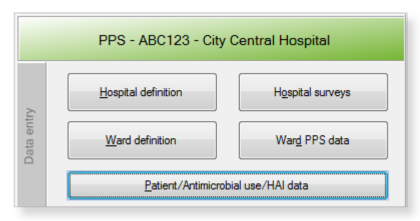


The HAI button is activated when the field Patient has at least one HAI: is set to Yes.



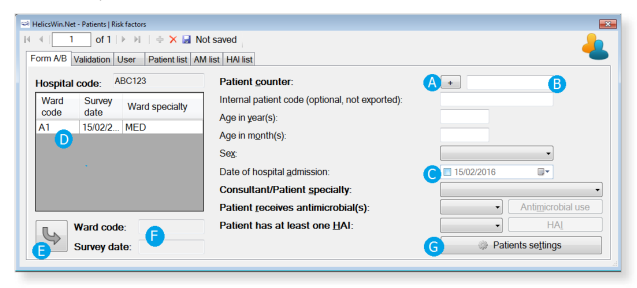
 No warning message is given if either of these fields are set to Yes and the corresponding AU/HAI data are not entered in the AU/HAI forms. These errors are only reported in the data quality check.

To enter patient data for antimicrobial use and HAI, click **Patient/Antimicrobial use/HAI data** in the *main menu*.



Form A/B tab

When you select Patient/Antimicrobial use/HAI data in the main menu, the Patients | Risk factors form opens at the Form A/B tab.

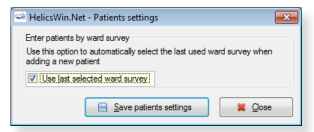


- 1. Click the **Add item** icon $\frac{1}{2}$ to activate the form. Click **(A)** to increment the counter for the next patient number, or **(B)**, enter the number manually.
 - The patient counter defines a unique record within the hospital (not within the ward). The patient counter is an anonymous patient identifier (that is, it is not the true patient number), and it can contain only numbers; other characters are not allowed.
- 2. Optionally, you can also enter a patient identifier in the **Internal patient code** field.

 This field is for local (hospital) use only. By default, data in this field is not exported as it may contain personal identifiers. However, HelicsWin.Net's export function allows you to choose to export it (see Exporting data from the database). To preserve patient confidentiality in compliance with Data Protection principles, the internal patient code data should not be included in the export file sent to the regional, national or EU level.
- **3.** Select the check box **(C)** to confirm the proposed date or select a date in the calendar pop-up. The date of hospital admission defaults to the ward survey date once the ward is selected. If this is not the correct date, select the actual admission date from the calendar.
- **4.** Select the ward code (ID) from the **Ward list (D)**, and then click the transfer button **(E)** to update the **Ward code** and **Survey date** fields **(F)**.
 - Only wards for which the survey date and ward specialty were previously entered appear in the **Ward list**.

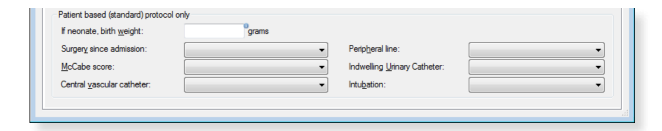
5. Optionally, to save time when entering subsequent patients, click on **Patient settings (G).** The Patients settings box opens.

Select **Use last selected ward survey**, and click **Save patient settings** to ensure that the chosen ward **(D)** is automatically entered in the **Ward code** and **Survey date** fields both when the transfer button **(E)** is clicked, and when subsequent patient records are added.



- **6.** Complete the remaining fields as necessary.

 Note that bold field labels denote mandatory fields.
- **7.** If you are using the Standard protocol, complete the additional risk factors that are displayed when the ward is selected (when the transfer button **(E)** is clicked).



8. Click the **Save** icon I or press **Ctrl+S**.

Descriptions of the variables in the **Form A/B** tab are given in Table 20. Mandatory fields are indicated by a red asterisk *. Required fields by a black asterisk *.

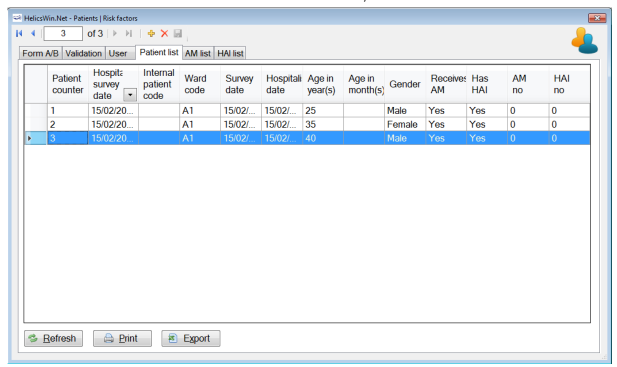
Table 20: Patients risk factors Form A/B tab

Variable	Description				
Hospital code	Inserted automatically based on the hospital selected during Hospital definition.				
Ward code	From ward PPS data.				
Survey date	From ward PPS data. Date on which data were collected in this ward. Data from a single ward should be collected on one day (dd/mm/yyyy).				
Ward specialty	From ward PPS data.				
Patient counter *	A system allocated sequential patient number for anonymised identification. Click to apply a number to the patient.				
Internal patient code	For hospital internal use only. Optional. Must be removed from any data submitted to ECDC/TESSy.				
Age in years	Patient's age in years. Enter respectively 0 or 1 if the patient is less than two years old.				
Age in months	Patient's age in months. Mandatory if Age in years is less than 2 (note that you must <i>also</i> enter the baby's Age in years).				
Sex	Select from the listed options.				

Variable	Description					
Date of hospital admission	Date patient was admitted to the hospital for the current hospitalisation (dd/mm/yyyy).					
Consultant/ patient speciality *	Specialty of physician in charge of the patient or specialty of main disease of the patient. If the consultant specialty differs from the patient specialty, give priority to the patient specialty. For paediatric patients on a PED ward, use the subspecialty (MEDGEN, MEDSUI etc.) (see ward specialty). LTC is in principle a ward specialty and should only exceptionally be used as a patient/consultant specialty.					
Patient receives antimicrobial(s) *	Patient receives at least one systemic antimicrobial agent on the date of the survey (given or planned treatment, including intermittent treatments, e.g. alternate day; or medical prophylaxis); for surgical antimicrobial prophylaxis, check whether any surgical prophylaxis was given in the 24 hours prior to 8 a.m. on the day of the survey; yes/no/unknown. If yes, collect <i>antimicrobial use data</i> .					
Patient has at least one HAI *	Patient has an active healthcare-associated infection on survey date; yes/no/unknown. If yes, collect <i>HAI data</i> .					
If neonate, birth weight	Birth weight in grams for neonates (patients who are \leq 30 days of age); the birthweight is the weight of the infant at the time of birth and should not be changed as the infant gains or loses weight.					
Surgery since admission	Patient has undergone surgery during current hospitalisation. Surgery is defined as a procedure where an incision is made (not just a needle puncture), with breach of mucosa and/or skin – not necessarily in the operating theatre. Answer categories: No surgery; yes, minimally invasive/non-NHSN surgery (examples see annex); yes, NHSN surgery – optionally specify NHSN surgery code (ICD9-CM code of the intervention is listed for the surveillance of surgical site infections in the NHSN system, see codebook); unknown.					
McCabe score	Classification of the severity of underlying medical conditions. Disregard the influence of acute infections, e.g. if the patient has an active HAI, estimate the score the patient had before the infection. Some examples of diseases and their different McCabe score categories are shown in Table 6. These examples, in particular those of the second (ultimately fatal) category, are not meant to be exhaustive but rather to serve as a guidance tool for the current protocol. Answer categories: Non-fatal disease (expected survival at least five years); ultimately fatal disease (between one and five years); rapidly fatal disease (expected death within one year); unknown.					
Central vascular catheter	Patient has central vascular catheter in place on survey date; yes/no/unknown. A central vascular catheter is defined by the CDC as an intravascular catheter that terminates at or close to the heart or in one of the great vessels which is used for infusion, withdrawal of blood, or hemodynamic monitoring. The following are considered great vessels for the purpose of reporting central-line BSI and counting central-line days in the NHSN system: Aorta, pulmonary artery, superior vena cava, inferior vena cava, brachiocephalic veins, internal jugular veins, subclavian veins, external iliac veins, common femoral veins, and in neonates, the umbilical artery/vein.					
Peripheral line	Patient has peripheral vascular catheter or arterial catheter in place; yes/no/unknown.					
Indwelling urinary catheter	Patient has indwelling urinary catheter in place at the date of the survey; yes/no/unknown.					
Intubation	Patient is under intubation with or without mechanical ventilation (endotracheal tube or tracheostomy) on survey date; yes/no/unknown.					

Patient list, AM list and HAI list tabs

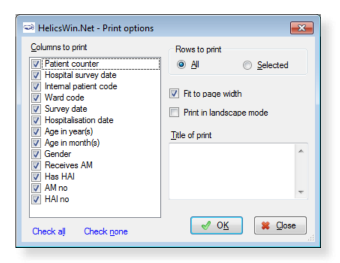
You can view, export and print listed information entered on all patients, all antimicrobial use, and all healthcare-associated infections within the tabs **Patient list**. **AM list** and **HAI list**.



- 1. Click **Refresh** to show the latest added records.
- **2.** Click any column header in the first row change the sort order of the list.
- **3.** Click **Print** to print the list.

The **Print options** form opens.

4. Select the fields and rows to be printed.

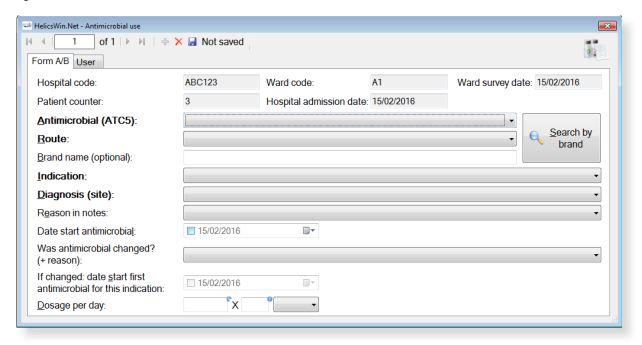


5. To add data for the next patient, click the **Add item** icon **.**

Antimicrobial use data

Antimicrobial use data are entered through the **Antimicrobial use Form A/B** tab. This form is opened by clicking the **Antimicrobial use** button in the **Patients | Risk factors Form A/B** tab.

Figure 11: The Antimicrobial use Form A/B tab



A unique antimicrobial record is defined by the fields with a bold label in the **Antimicrobial use** form. These fields create a unique combination of:

 ATC5 code + route + indication + diagnosis site (for treatment only – otherwise code NA=not applicable).

This means that it is possible to enter the same antimicrobial agent for more than one indication, for the treatment of more than one infection site, or for more than one route, in the same patient. Deduplication of the data will be done at the analysis level as necessary.

No warning message is given if the corresponding AU data are not entered in the AU form. These errors are only reported in the *data quality check*.

Typically you add patient data and antimicrobial consumption data, when you enter the survey data.

If data is incomplete, a Warning/Error message will open, listing required data items. Table 21 describes the variables in the **Antimicrobial use** form. Further details are provided in the HAI-Net PPS protocol (see *Related documents*).

To specify antimicrobial use data for a patient:

- Click Patient/Antimicrobial use/HAI data in the main menu.
 The Patients | Risk factors form opens displaying the Form A/B tab.
- **2.** Scroll through the list of patients to locate the patient you are interested in and confirm that the patient details match the patient whose data you want to update.
- **3.** Select **Yes** for the **Patient receives antimicrobial(s)** field to activate the **Antimicrobial Use** button.

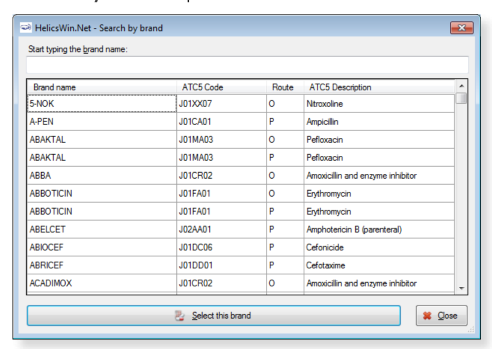


4. Click Antimicrobial use.

The **Antimicrobial use Form A/B** tab (Figure 11) opens.

5. Click Search by brand.

The **Search by brand** form opens.



- **6.** Enter the brand name in the field or click the required brand in the list.
- 7. Click Select this brand.

You return to the **Antimicrobial use** form, where the ATC5 code and route have been inserted.

- **8.** Optionally, click the *User tab* to view or edit the antimicrobial use user variables.
- **9.** Click the **Save** icon III or press **Ctrl+S**.

Descriptions of the variables in the **Antimicrobial use General** tab are given in Table 21. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

Table 21: Antimicrobial use Form A/B tab variables

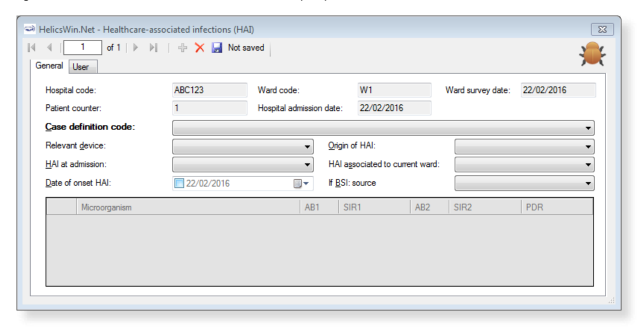
Variable	Description				
Header section	n ward PPS data (Table 19) and PPS patient risk data (Table 20).				
ANTIMICROBIAL (ATC5) *	Antimicrobial generic or brand name. Allowed are, for example, amoxicillin, but also national brand names; include ATC codes (ATC2: J01 antibacterials, J02 antifungals; ATC4: A07AA, P01AB, D01BA; ATC5: J04AB02). See codebook for included antimicrobial agents.				
Route *	Route of administration of the antimicrobial agent; P =parenteral; O =oral; R =rectal; I =inhalation.				

Variable	Description				
Brand name	Retrieved from stored brand names.				
Indication *	 Patient receives systemic antimicrobials for: Treatment intention: CI: community-acquired infection; LI: infection acquired in long-term care facility (e.g. nursing home) or chronic-care hospital; HI: acute-hospital-acquired infection. Surgical prophylaxis: SP1: single dose; SP2: one day; SP3: > 1 day: check if given in the 24 hours prior to 8 a.m. on the day of the survey – if yes, check if given on the day before yesterday or on the day of the survey in order to determine duration. MP. Medical prophylaxis. O. Other indication (e.g. erythromycin use as a prokinetic agent). UI. Unknown indication/reason (verified during PPS). UNK. Unknown/missing, information on indication was not verified during PPS. If treatment intention for infection, fill in site of infection (diagnosis). 				
Diagnosis (site) *	Diagnosis group by anatomical site. Should only be recorded if there is an intention to treat an infection; not recorded for prophylaxis or other indications (use code NA=not applicable). See codebook for code list.				
Reason in notes *	The reason for antimicrobial use was documented in the patient chart/notes. Yes/No/Unknown .				
Date start antimicrobial	Start date of the current antimicrobial. If the patient received the antimicrobial on admission, record the date of admission.				
Was antimicrobial changed? (+reason)	 admission, record the date of admission. Was the antimicrobial (or the route of administration) changed for this indication, and if so, what was the reason? If the antimicrobial was changed more than once for the current indication, report the reason of the last change. N=no change, antimicrobial was not changed. E=Escalation: antimicrobial was escalated on microbiological and/or clinical grounds, i.e. the isolated microorganism was not susceptible to the previous antimicrobial and/or lack of clinical effect of previous antimicrobial; includes switch from oral to parenteral for the same antimicrobial. D=De-escalation: antimicrobial was de-escalated on microbiological and/or clinical grounds, i.e. the isolated microorganism was susceptible to more narrow-spectrum or first-line antimicrobials than the previous antimicrobial and/or the clinical situation of the patient allows changing to more narrow-spectrum or first-line antimicrobial. S=switch IV to oral; route of administration of same antimicrobial was changed from parenteral to oral. A=adverse effects; antimicrobial was changed because of observed or expected side or adverse effects of the antimicrobial. OU=change for other or unknown reason: the antimicrobial for that indication was changed for another reason or the antimicrobial was changed but the reason why could not be determined by the surveyor. 				
If changed; date start first antimicrobial for this indication	Start date of the first antimicrobial prescribed before the current antimicrobial for the same indication if the current antimicrobial replaced a previous one. If the antimicrobial was changed more than once for the current indication, report the start date of the first (not the previous) antimicrobial.				
Dosage per day	Number and strength (in milligrams, grams, IU or MU) of doses of the antimicrobial given per day. Report as e.g. 4 x 1 g per day (three variables: number of doses, strength of one dose, unit of one dose). When one dose of an antimicrobial is given every other day, report the number of doses as 0.5 (e.g. 0.5 x 1 g/day).				

HAI data

HAI data are entered through the **Healthcare-associated infections (HAI)** form. This form is opened by clicking the **HAI** button in the **Patients | Risk factors General** tab.

Figure 12: The Healthcare-associated infections (HAI) form



In the **HAI** form, a unique record is defined by the **Case definition code**. The same HAI case definition code cannot be reported twice in the same patient record, even with a different date of onset, because this is not possible according to the ECDC-PPS protocol. In fact, even the same infection site should not be allowed, but currently the programme looks only at the case definition codes. For example:

- A PN1 and a PN4 should not be reported in the same patient record although both definitions were met.
- Related codes for the same site should not be reported: e.g.,
 - A PN1-5 should not be reported together with a NEO-PNEU (the latter has priority in neonates);
 - A BSI should not be reported together with a NEO-CNSB or NEO-LCBI, and so on.

If data is incomplete, a Warning/Error message will open, listing required data items. Table 22 describes the variables in the HAI form. Further details are provided in the HAI-Net PPS protocol (see *Related documents*).

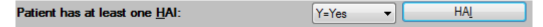
To specify healthcare-associated infection (HAI) data for a patient:

1. Click Patient/Antimicrobial use/HAI data in the main menu.

The **Patients** | **Risk factors** form opens displaying the **General** tab.

2. Scroll through the list of patients to locate the patient you are interested in and confirm that the patient details match the patient whose data you want to update.

3. Select Yes for the Patient has at least one HAI field to activate the HAI button.



4. Click HAI.

The Healthcare-associated infections (HAI) form (Figure 12) opens.

Note

- The date of onset of the HAI should be specified only if the HAI was not present at admission, that is, date of onset ≥ date of hospital admission
- The resistance markers (phenotypes) do not have labels depending on the microorganism.
- **5.** Optionally, click the *User tab* to view or edit the HAI user variables.
- **6.** Click the **Save** icon **J** or press **Ctrl+S**.

Descriptions of the variables in the **HAI General** tab are given in Table 22. Mandatory fields are indicated by a red asterisk *. Required fields are indicated by a black asterisk *.

Table 22: HAI General tab variables

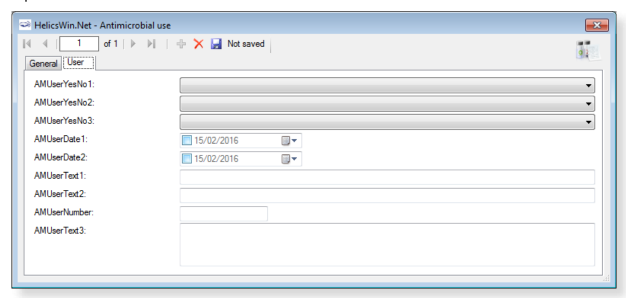
Variable	Description				
Header section	From ward PPS data (Table 19) and PPS patient risk data (Table 20).				
Case definition code *	HAI case definition codes: specify subcategory, e.g. PN4 (see codebook), CVS-VASC. A single-case definition code should only be provided once per patient (no different infection episodes).				
Relevant device *	To be specified for PN, BSI, NEO-LCBI, NEO-CNSB and UTI only. Relevant invasive device was in situ (even intermittently) for 48 hours (seven days for UTI) before onset of the infection, i.e. intubation for pneumonia, central/peripheral vascular catheter for bloodstream infections, urinary catheter for UTI; Unk=unknown. Yes/No/Unknown.				
Origin of HAI *	Infection is associated with (1) current hospital; (2) another acute-care hospital; (3) other origin or unknown. Record 'HAI present at admission' if associated with an earlier stay in surveyed hospital or transferred from another facility.				
HAI at admission	Signs and symptoms of the infection were present at admission to the hospital; if not, provide date onset of infection. Yes/No/Unknown.				
An HAI is associated with the current ward if the infection started on day 3 or after admission to the current ward (where the date of admission to the ward day 1) OR if the infection started on day 1 or 2 after a placement of an invasion device on the current ward OR if the patient was readmitted with an HAI presson admission associated to a previous stay in the same ward (within 30 days operation for surgical site infections (or 90 days for deep and organ/space SS after implant surgery), less than 28 days after discharge for C. difficile infection less than 48 hours (two calendar days) after discharge for other HAIs).					
Date of onset HAI	Date of onset of the infection (dd/mm/yyyy). Not to be recorded if signs/symptoms are present at admission, but mandatory if onset during current hospitalisation. Date of first signs or symptoms of the infection; if unknown, record date treatment was started for this infection or the date the first diagnostic sample was taken. If no treatment or sample, please estimate.				
If BSI: source	If lab-confirmed bloodstream infection, specify the origin: catheter-related (central: C-CVC, peripheral C-PVC), secondary to another infection: pulmonary (S-PUL), urinary tract (S-UTI), digestive tract (S-DIG), surgical site infection (S-SSI), skin and soft tissue infection (S-SST), other infection (S-OTH), or BSI of (confirmed)				

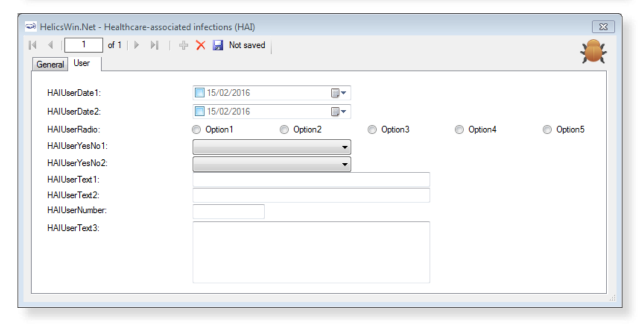
unknown origin (UO); missing data, no information available=UNK; Secondary BSI reported as separate HAI, in addition to the primary infection if it matches the case definition.

User tab

The **User** tab contains fields useful for users at national, regional or hospital level. They can be renamed and personalised to enable collection of information during the PPS that is outside of the PPS protocol. These modifications are made using the **Translation** functionality in the **Settings** form, see *Translating the text in user forms*.

Skip the **User** tab.





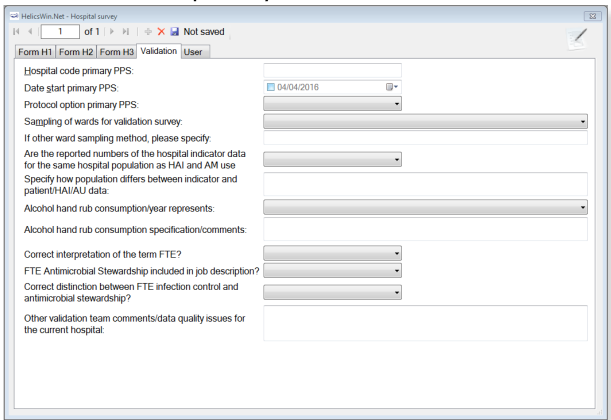
PPS validation tabs

The **Validation** tabs in the **Hospital surveys**, **Ward PPS data** and **Patient** forms are used by validation teams, only if the current survey is a validation study of the primary PPS for the hospital (validation checkbox in hospital data form checked), following the validation protocol.

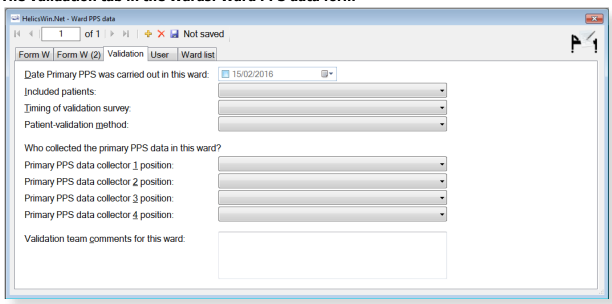
PPS validation is in principle performed by **an external validation team** trained by the national/regional PPS coordinating centre. The objective of the validation study is to assess the sensitivity, specificity and/or reproducibility of the data collected in the ECDC PPS.

See also: Further details are contained in the **Validation Protocol**.

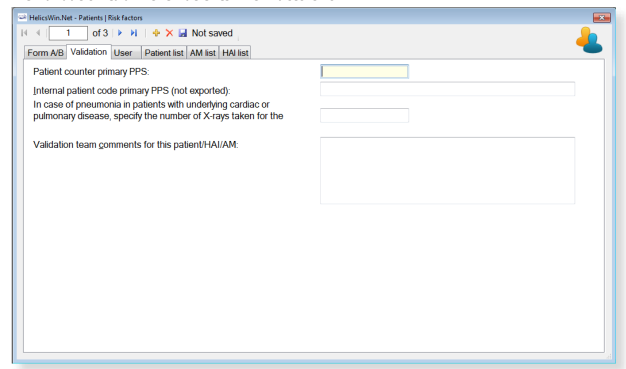
The Validation tab in the Hospital surveys form



The Validation tab in the Wards: Ward PPS data form



The Validation tab in the Patients: Risk factors form



Checking data quality

HelicsWin.Net performs a selected number of automatic validation checks during data entry. These checks do not pick up all possible problems. You are strongly recommended to perform a further level of in-depth checking *before* creating a report, exporting or merging data. The HelicsWin.Net *data quality check* function performs these in-depth checks. The checks must be started manually.

The data quality check verifies the internal consistency of all information entered into your database. This verification helps you to identify any problems, in turn greatly reducing the chances of producing inaccurate or inappropriate data outputs, and helping to ensure functional export of your data. As with all such data operations, you are advised to check your final data and ensure that the data make sense.

When you perform a data quality check, HelicsWin.Net produces messages at three severity levels: Error, Warning and Success. A data quality check output is produced, with each message including details of the issue identified.

Table 23: Data quality check messages

Severity	Code	Description
Error ERR		A fatal inconsistency has been identified in a record or across multiple records. The severity of the inconsistency is such that the data could be misreported, or there could be a failure of a merge, report or export operation. User action: You must locate and fix all such errors before proceeding with any merge,
		report or export operations.
Warnings	WARN	A significant inconsistency has been detected that indicates that data may be compromised in merge, report or export operations.
		User action: You must identify the inconsistency and then decide whether the issue identified might invalidate your data.
Success	SUCC	The data quality check found no issues with the record; the check was therefore deemed to be a success.
		User action: None.

Some types of errors detected

One common warning occurs when merging ward or patient data, when one of the two datasets has less detailed hospital data. In this situation, if you are sure that the hospital definitions relate to the same hospital, it would be safe to merge the data.

Errors detected by the data quality check (but not detected on data entry) include the following:

- The variable *has_amu* is reported as *Yes*, but there are no antimicrobial use data reported in the **Antimicrobial use** form.
- The variable *has_hai* is recorded as *Yes*, but there are no healthcare-associated data reported in the **HAI** form.
- In the Light protocol only:
 - A warning is generated when the user did not report denominator data by consultant/patient speciality for a given ward survey.
 - An error is generated if the total of the denominator data by consultant/patient speciality does not equal the value of *Total number of patients in ward* for a given ward survey.

• An error is generated if the consultant/patient specialty of a patient does not have a corresponding consultant/patient specialty record at the ward level.

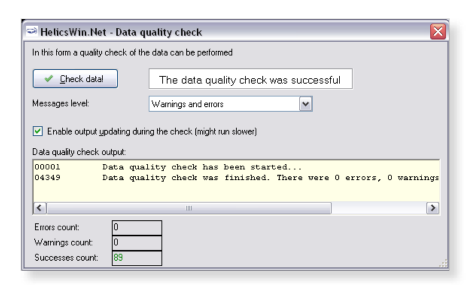
Running a data quality check

A data quality check checks the whole of the current HelicsWin.Net database.

To run a data quality check:

1. Click Data quality check in the main menu.

The **Data quality check** form opens.



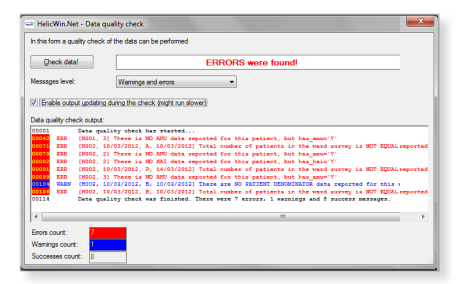
- 2. In the Messages level field, select Warnings and errors (recommended).
- 3. Deselect Enable output updating during the check (recommended).

Enabling this option updates the data displayed in the *Data quality check output* field every time a new line is added. If this option is not enabled, updates are only made periodically.

If the output consists of only 50 lines, for example, you can evaluate the output displayed in the *Data quality check output* field. However, if you have verbose output (Message level = all) and you have many patients, etc., thousands of lines of output might be displayed in the field. In such a case, if you enable this option (i.e. if you check this check-box), system response time can be very slow; whereas leaving this option unchecked will improve system speed.

4. Click Check data.

When the processing has finished, the resulting messages are displayed in the data quality check output text box, and the (colour-coded) breakdown of the Error, Warning, and Success counts are shown underneath.



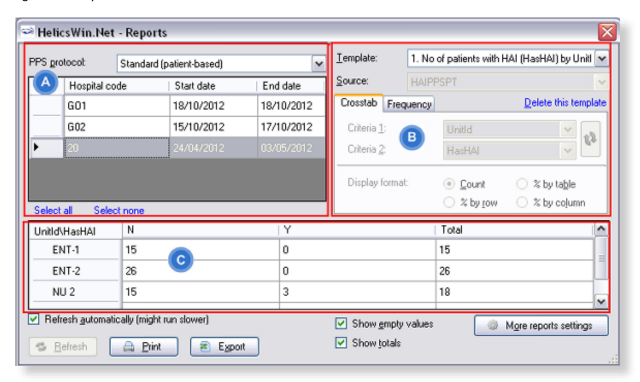
Creating reports from survey data

HelicsWin.Net provides a form for extracting data from existing surveys and collating these data into reports that may be viewed on a screen, printed, or exported as CSV files. You can create standard pre-formatted reports from templates that you select from a drop-down list. You can also customise your own specialised reports, and save these customisations for future use.

The Reports functionality is not yet available for the ICU module.

You create reports in the **Reports** form, which you access by clicking **Reports** in the *main menu*. This form has three main parts labelled **A**, **B**, and **C** in Figure 13.

Figure 13: Reports form



On this form...

- Select PPS protocol and hospital(s). Select a single or multiple hospitals. You can select multiple hospitals by pressing the CTRL key and clicking on any additional hospitals you want to report on. Reports will show data for all selected hospitals together.
- Define outputs. The dropdown menu for Templates contains four pre-prepared templates for tables, and the option Custom. Choosing Custom enables you to either (i) choose two variables to crosstabulate in a table, or (ii) choose one variable for a tabulation of frequency.

 Note: you can only edit the type of data in a table when you are creating a custom template.
- **Review the tabular results** that are displayed on the form. You can then use the buttons at the bottom of the form to print a hard copy of your report or export the report data as a CSV (comma separated variable) file for subsequent analysis in Microsoft Excel or other CSV compatible program.

Producing a report

To produce a report, you must first select the PPS protocol and hospital(s):

- 1. Click **Reports** in the *main menu*.
- 2. In the PPS protocol field, select Standard or Light from the drop-down menu.

Once you have made this selection, only Hospitals that used the Standard/Light protocol will be displayed.

3. Select a hospital you want to create a report on.

The row containing the currently selected hospital is shaded.

4. To select additional hospitals for your reports, press the **CTRL** key and then click each of the additional hospitals you want to select, in turn.

Creating and printing a report from a template

After the protocol and hospitals have been selected in the previous procedure, then:

1. Select a report template from the Templates list.

The following default options are available.

- a. Number of patients with HAI (HasHAI) by UnitID
- b. Number of patients with HAI (HAI data) by UnitSpecialty
- c. Number of patients receiving antimicrobials (ReceiveAntimicrobials) by MainUnitSpecialty
- d. Number of patients receiving antimicrobials (AM use data) by MainPatientSpecialty

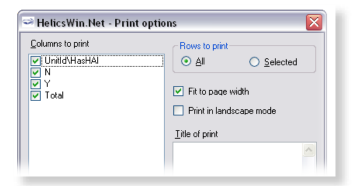
Note: MainUnit/PatientSpecialty = the first three characters of the full specialty code, e.g. 'surgery' for 'orthopaedic surgery' (SURORT)

Tip: If you cannot read the complete text on screen, expand the form by dragging the right edge to the right of the screen.

The report is data is automatically retrieved and, where necessary, calculated and displayed in the table. The **Crosstab** and **Frequency** tabs are not active for this option.

2. Click **Print** to initiate the printing the report.

HelicsWin.Net prompts you to select which columns you want to print:



3. Make your selections, and click **OK**.

4. Follow the on-screen prompts and check the print preview before deciding to print.



5. In the **Print preview** window, click the **Print** icon to complete the print request, and then close the window.

Creating custom reports and templates

Use the **Crosstab** or the **Frequency** tab – see area **B** in the **Reports** form (see Figure 13 on page 90).

As with the provided templates, custom reports are viewed in the tabular results area **C** in the **Reports** form, and optionally printed or exported as a CSV file. Once you have defined your custom report, you can either save it as a named template, or simply discard it. If you save the template, its name will appear as an option for future use in the Template drop-down list in area **B** in the **Reports** form.

The two tabs represent two different ways of viewing the selected data.

Crosstab	Create a table with two different variables on the each axis (and shown in the row and column headers). You then have the option of specifying whether the cells contain numbers or percentages; and if the latter, what the percentages are how they are displayed.
Frequency	Create a table with the values that can be taken by a single variable (for example, HasHAI has possible values of Yes (Y) and No (N)) on one axis and the frequency of occurrence on the

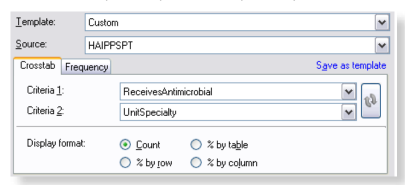
To create a custom report:

1. Click Reports in the main menu.

other.

- **2.** Select one or more hospitals and the protocol.
- **3.** In the **Template** field, select **Custom**.
- **4.** In the **Source** field select your source for your data.

The number of options depends on the protocol you have selected.



5. Select the **Crosstab** tab, and specify the variables you want to see in a table, and the format of the data within the cells in that table.

Option	Defines the vertical axis, i.e. row headings			
Criterion 1				
Criteria 2	Defines the horizontal axis, i.e. column headings			
Count	The cells contain raw* data; and the column and row totals are unscaled numerical values.			
% by table Each cell contains a percentage of the raw* to whole table.				
% by row	Each cell in a row contains a percentage of the sum of the raw* values for that row.			
% by column	Each cell in a column contains a percentage of the sum of the raw* values for that column.			

^{*}the raw value is the numerical value in the table before any calculations are made.

6. Alternatively, select the **Frequency** tab and select the variable (criterion) whose data you want to display, in this case HasHAI (from the field **Patient has active HAI**). The frequencies, in terms of the count and percentage, are shown for each value of the selected variable.

HasHAI	Frequency	Frequency (%)
N	202	91%
Y	19	9%
Total	221	100%

- **7.** Optionally, as for the default templates, click **Print** to print the table, or **Export** to export the data as a CSV file.
- **8.** Click **Save a template**, to save these table settings as a named templete.

The template is automatically added to the Template drop-down list, so that you can reuse these settings again later.



Note: You cannot save a template that is identical in function to an existing template; if you try this, you simply get an error message.

Changing the display precision settings

You can change the number of decimal places on the reports, and whether sub-precision values are displayed. On the **Reports** form, click **More report settings** to change the number of decimal places displayed, and whether sub-precision values are displayed.

Precision

Precision refers to the number of decimal places in the displayed results. The following table shows the displayed frequency results for the quotient 202 / 221 for different display precisions.

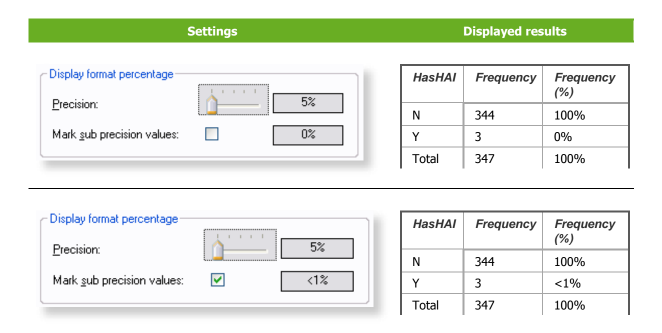
Table 24: Precision of displayed results

Decimal places	0	1	2	3	4
Displayed results	91%	91.4%	91.40%	91.403%	91.4027%

Sub-precision values

Sub-precision values are those values that would give a value of zero when expressed as percentages, when rounded down. This can be misleading if the results tables shows non-zero results.

The following tables demonstrate the effect of selecting and deselecting the **Mark sub precision values** checkbox when the precision set to the lowest level. A frequency of 3 in 347 corresponds to 0.86%. With **Mark sub precision values** deselected, this value is displayed as 0%; if **Mark sub precision values** is selected, the value is displayed as <1%, which most users will find more informative.



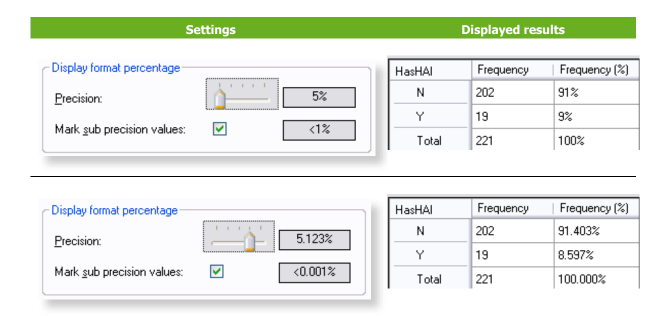
Changing precision values in reports

To change the precision, and sub-precision values in reports:

- 1. Click **Reports** in the *main menu*.
- 2. Click on the More report settings button.

A form opens with contains a slider control and checkbox, as shown below.

Sliding the slider fully to the left gives zero decimal places (e.g. 5%). Sliding it three steps to the right gives three decimal places (e.g. 5.123%).



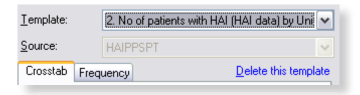
3. Tick the option **Mark sub-precision values:** to change the format of data below the minimum value.

Removing report templates

You can remove built-in and custom report templates one at a time if you find you do not need those templates again. Alternatively, you can reset the list of templates to its initial status: any missing built-in templates will be restored; all custom templates will be permanently deleted.

To remove a report template:

1. Open the **Reports** form:



2. In the Template drop-down list, select the template you want to delete.

You can select either a custom template or a built-in template.

3. Click Delete this template.

HelicsWin.Net prompts you to confirm the deletion.

Caution: The selected template will be deleted from the list.

If you delete an existing custom template, you cannot get it back without

recreating it from scratch.

4. Click OK.

The selected template is deleted, and you return to the **Reports** form.

Resetting the templates list

To return your templates to their initial status, you need to reset the templates list. This procedure

- Restores any built-in templates that have been deleted.
- Permanently deletes all custom report templates from the list.

To reset the templates list:

- 1. In the Reports form, click More reports settings.
- **2.** Click the **Templates** tab.



3. Click Reset the templates list.

HelicsWin.Net asks you to confirm your request, and then confirms that all user-defined templates are deleted.

Exporting data from the database

Before exporting data, always perform a data quality check and correct any data errors that you find. We recommend using the HelicsWin.Net *data quality check* functionality to help you achieve this (see Section 0).

You can export in the original Access database format (as an .mdb file compressed in a ZIP file) or in the TESSy CSV format.

Exporting data in Microsoft Access format

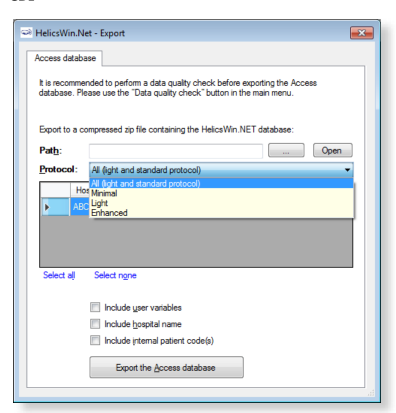
You can export data from the HelicsWin.Net database files as a zipped Access .mdb file.

Before you start, make sure that the survey start and end dates are defined in the **Hospital data** form for each hospital for which you want to export data.

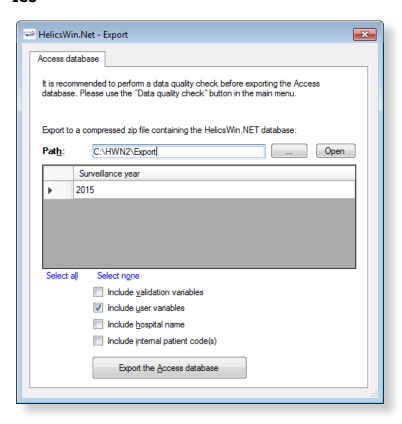
To export data in Microsoft Access format:

Click Data export in the main menu.
 The Export form opens, by default in the Access database tab.

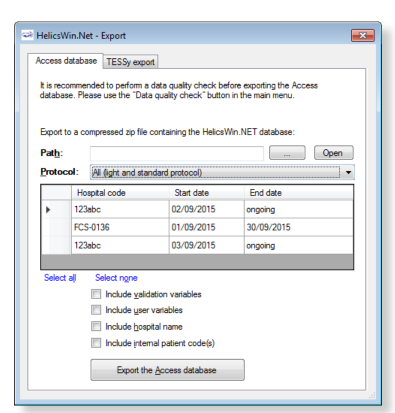
CDI



ICU



PPS



• Click the **Open** button (to the right of the **Path** field), to locate the folder to which you want to send the exported file, and then click **OK**.

For example, to use the folder C:\HWN2\Export, go to the folder C:\HWN2, and create the folder **Export** manually.

• CDI and PPS: In the data grid, select the code(s) and survey dates for the hospital-survey(s) for which you are exporting the data. You can select more than one hospital-survey by holding down the control button, and then clicking on each hospital code you wish to select or select all hospitals by clicking the **Select all**.

Make sure that the CDI / PPS start and end dates are defined in the **Hospital data** form.

- ICU: The entire selected surveillance year for the current hospital will be exported.
- Optionally, use the checkboxes to choose whether to include additional sets of variables in the exported files. These are data entered relating to:
 - a. Validation studies (not for CDI)
 - b. Users
 - c. Hospital names
 - d. Internal patient codes (for both primary and validation data).

As some of these may contain confidential information, they are not exported by default.

If a checkbox is left unchecked, the corresponding variables will be empty in the export file.

For the <u>pilot study of the HAI-Net ICU protocol v2.0</u>, user variables include the feasibility questionnaire data and need to be included in the export.

To preserve patient confidentiality in compliance with Data Protection principles, the internal patient code data must not be included in the export file sent to the regional, national or EU level, and must not be included in any data sent to ECDC/TESSy.

• Click **Export the Access database**.

HNW generates a ZIP file containing the exported database file **HelicsWinNet_export.mdb** or **HelicsWinNetICU_export_yyyymmdd_hhmmss.mdb**

The ZIP file is date and time stamped with the file name having the format: **HWN_yyyymmdd_hhmmss.zip** (**PPS**) or **HWN_ICU_yyyymmdd_hhmmss.zip**

- In the **Access database** form, click **Open** to view the contents of the folder you have exported your file to.
- ICU pilot: attach the exported zip file containing the Access database to an email and send it to vour national HAI surveillance coordinator.

Exporting data to TESSy CSV format

This feature is not yet available for the CDI or ICU modules.

You can also export data for a given hospital and survey in a TESSy CSV (comma-separated values) file. TESSy (The European Surveillance System) is ECDC's online system for upload and analysis of data for different networks for surveillance of communicable diseases, and can only be accessed by nominated national/regional contact points. The format of the exported data is defined in the CDI / ICU / PPS protocol, and in the supporting documents on the TESSy website at

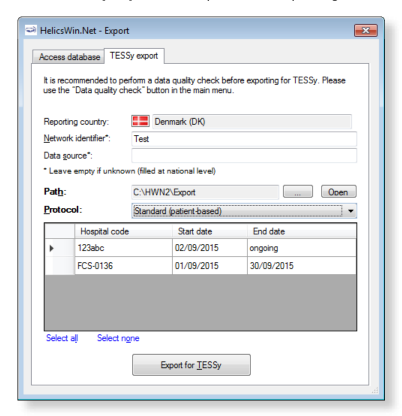
https://tessy.ecdc.europa.eu/TessyWeb.

Some variables need to be added to the TESSy format, such as the data source (usually the national surveillance institute) or the network ID (for example, if there is more than one surveillance network for the same disease in the same country). It is possible for any user to enter values for the variables *Network identifier* and *Data Source*. In general, this specific information is not known at the hospital level, and so these variables can be left empty, e.g. until entry by the nominated contact points. The freeware tool *csved* can be used to add the information to the CSV file.

To export data in TESSy format:

Click Data Export in the main menu.
 The Export form opens.

2. Click the **TESSy export** tab to open the corresponding form.



3. Enter the **Network identifier** and **Data source**, if known.

Usually, this information is not known at the hospital level, in which case these variables should be left empty and subsequently entered by the nominated contact points.

- **4.** Click the **Browse** button (... to the right of the **Path** field), to locate the folder to which you want to send the exported file. You can either use an existing folder or create a new folder by clicking **Make New Folder** to create e.g. C:\HWN2\Export. You can also open the selected path by clicking the **Open** button.
- **5.** In the **Protocol** field, select the protocol for the file to export.
- **6.** In the data grid, select the code for the hospital(s) for which you are exporting the data. You can select more than one hospital-survey by holding down the control button, and then clicking on each hospital code you wish to select or select all hospitals by clicking **Select all**.

Make sure that the start and end dates are defined for your PPS.

7. Click Export for TESSy.

The CSV export may take some time to finish. HWN generates five CSV files, with following names:

Standard protocol:

- 1.HAIPPS.csv
- 2.HAIPPSPT.csv
- 3.HAIPPSPTAM.csv
- 3.HAIPPSPTINF.csv
- 4.HAIPPSPTINFRES.csv

Light protocol:

- 1.HAIPPSLIGHT.csv
- 2.HAIPPSLIGHTDENO.csv
- 3.HAIPPSLIGHTDENOAM.csv
- 3.HAIPPSLIGHTDENOINF.csv
- 4.HAIPPSLIGHTDENOINFRES.csv

The numbers at the beginning of the file names designate the level of the data in the hierarchical database, for example, the first level 1.HAIPPS.csv file contains the hospital data.

Note:

Values for national and some hospital variables are not included in the first level export file (1.HAIPPS.csv). These values should be completed by the National/Regional PPS Co-ordinating Centre before uploading to TESSy HAIPPS: DataSource, NetworkID, HospitalLocation (NUTS1 code) and SampleHospital (which specifies whether this hospital belongs to a national sample for the PPS/survey).

Warning:

The file names do not contain the hospital code or the time stamp, so when data are exported for a second hospital, any previous TESSy export csv files in the same directory are overwritten.

To prevent this, compress the files into a zip file and include the name of the hospital in the file name, before sending the data to the national/regional PPS coordinating centre. It is also recommended that you transfer the TESSy export files together with the Access database export file.

Converting to TESSy CSV using Stata

Stata conversion programmes are available from ECDC for national PPS coordinating centres to convert both Standard- and Light-protocol Access data to TESSy CSV files. These Stata programmes ask for user input for the data that must be completed by the national coordinating centre.

Recommendation:

Transfer the TESSy export files together with the Access database export file to your National/Regional PPS Coordinating Centre.

Merging data

The data merge facility enables you to merge data in different databases that relate to the same ward or hospital. At the national or regional level, you can merge data from different hospitals so that you can collect combined statistics for all the hospitals, and run reports on the combined database.

Recommendations

Before you attempt to perform a merge on two or more databases:

- You must have a complete backup of each database file (HelicsWinNet.mdb) in its current state.
- You must run a *data quality check* on each database before you perform the merge. In addition, you must identify the causes of all errors and warnings in the results list.

Merging two database files for the same hospital

Typically you need to merge data whenever you have collected survey data on more than one computer. In the scenario described here, there are two computers, named A and B, each of which contains data collected from the same hospital (hospital code G01), but different wards. We are going to merge the data from computer B with that on computer A. The computers share a networked drive P.

The following table summarises the database names, location, and content:

Table 25: Properties used for merging Ward data

Hospital code	Computer	Wards	Original database	iginal database Location of database files	
G01	A	W01, W02	HelicsWinNet.mdb	C:\HWN2	Yes
G01	В	W11, W12	HelicsWinNet.mdb	P:\HWN2	Yes

On computer A

Backup the current database and select hospital G01 as the active hospital

- Copy the file HelicsWinNet.mdb to a backup folder (on computer A).
 The merge process modifies the original file, so if anything does go wrong, you may need this backup.
- **2.** Click Hospital definition in the *main menu*.

The **Hospitals** form opens.

3. Select the hospital with code G01, and then click **Select this hospital**.

The form closes and the code G01 is displayed at the top of the main menu.

On computer B

Export the file HelicsWinNet.mdb database to the common P drive.

Note: You do not need a backup of the database on computer B, because you will be working with a copy and the original will remain unchanged.

- 1. Create a folder HWN2 on the P drive.
- 2. Click Hospital definition in the main menu.

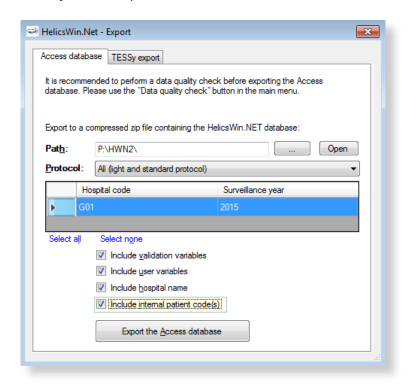
The **Hospitals** form opens.

3. Select the hospital with code G01, and then click **Select this hospital**.

The **Hospitals** form closes.

4. Click **Data Export** in the main menu.

The **Export** form opens.



5. In the **Access database** tab, click the **Open** button (situated to the right of the **Path** field), and navigate to the folder you just created on the P drive, and then click **OK**.

The text P:\HWN2 is added to the path field. You cannot edit this field directly.

Note: The following steps match those described above in the section *Exporting data from the HelicsWin.Net database.*

- **6.** In the **Protocol** field, select the protocol(s) as necessary for your survey(s).
- **7.** Select the hospital code **G01**, and check the start and end dates to confirm that you have the correct survey.
- **8.** Optionally, tick relevant checkboxes.

9. Click Export the Access database.

The database is exported as a zip file.

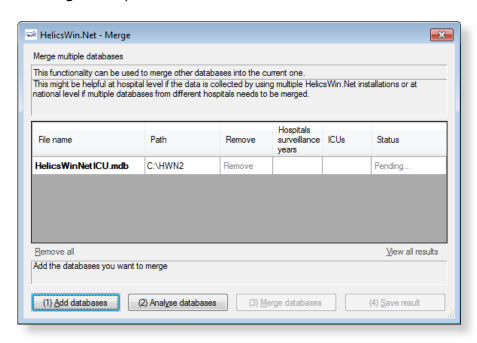
10. Note the name of the zip file and click **OK**, and then close the **Export** form. Navigate to the P folder and unzip the file (HelicsWinNet.mdb) to P:\HWN2.

On computer A

With hospital G01 selected, open the Merge form and add the database from computer B, now stored on the P drive.

1. Click **Data merge** in the *main menu*.

The **Merge** form opens.



- 2. Click (1) Add databases.
- **3.** Navigate to the folder containing the file you want to merge, in this case it is HWN2 on the P drive, and then select **HelicsWinNet**.mdb.

The P database is added to the list.

File name	Path	Remove	Hospitals surveillance years	ICUs	Status
HelicsWinNetICU.mdb	C:\HWN2	Remove			Pending
HelicsWinNetICU.mdb	P:\HWN2	Remove			Pending

Click (2) Analyse databases.

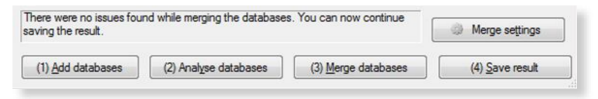
If the files are correctly formatted, the status **Analysed OK** is returned. In addition, the hospital survey dates and ward names are displayed for each database.

File name	Path	Remove	Hospitals surveillance years	ICUs	Status
HelicsWinNetICU.mdb	C:\HWN2	Remove	G01 24/0	W-01 ,W	Analyzed OK
HelicsWinNetICU.mdb	P:\HWN2	Remove	G01 26/0	W-11 ,W	Analyzed OK

4. Check that the survey dates and ward names, as displayed, represent the correct data. You can expand the column widths to read the data as necessary.

- **5.** Check that the survey dates and ward names, as displayed, represent the correct data.
- 6. Click (3) Merge databases.

If successful, the following message appears at the bottom of the **Merge** form.



7. Click (4) Save result.

A warning box opens. This warns you that saving the data will change the dataset that is active in your HelicsWin.Net to be a merged version of the datasets selected in the previous steps, i.e. from computers A and B.

- 8. Click Yes.
- 9. Optionally, restart HelicsWin.Net to ensure that the data is refreshed. This is recommended.
- **10.** Additionally, we recommend running a Data quality check from the *main menu*, following the merge, and a restart, to identify any issues.

To check patient data

You can check your new data in the **Patients** form.

- Click Hospital definition in the main menu.
 The Hospitals form opens.
- 2. Select the hospital you wish to check.
- **3.** Click Patient/Antimicrobial use/HAI data.

The **Patients | Risk Factors** form opens

4. Click the Patients list tab.

The list opens. You can sort the list by column header, if necessary.

5. To examine the list in detail, click **Export**.

HWN exports the data as a CSV file, which you can open in Excel.

Troubleshooting data merges

Warnings

I get warnings when I try to merge data from a single survey made on two different computers.

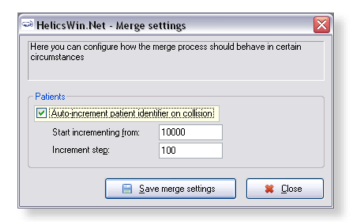
The most common reason for warnings is that your hospital definitions are not identical in the two databases. In such cases you will get a warning message, but you will still be able to merge the data. If you are sure you are not going to lose data, you can go ahead and merge. In these cases, HWN discards the second hospital definitions and uses the one in the database you are merging data into.

Duplicate patient IDs and collision detection

There are several scenarios that can lead to having duplicate patient IDs. The action you need to take depends on several factors:

Scenario 1: If you are merging two distinct databases that contain the same patient IDs, but these IDs belong to different patients:

• You need to turn on the **Auto-increment patient identifier on collision** setting in the **Merge settings** form.



1.To access the **Merge settings** form, click **Data merge** in the *main menu*, and then click **Merge settings**.

The default values usually work, but you may need to change these if you are doing multiple merges, for example.

Scenario 2: You really do have some duplicate data with the same patient records occurring more than once.

In this case, make sure you turn off (clear) the auto-increment setting.
 Any duplicates will **not** be uploaded.

Caution

It is possible to get a situation where both scenarios are true in part. If this is the case, you can only make progress by examining each record in turn.

Modifying settings

You can modify basic settings in HelicsWin.NET to suit your own personal preferences.

To change settings:

1. Click Settings in the main menu.

The **Settings** form opens; by default the content of the **General** tab is displayed.

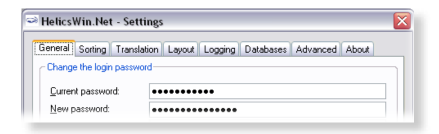


Table 26 outlines which settings you can change.

Table 26: Settings forms

Tab	Functions
General	- Reset password and restore hidden notification messages
Sorting	- Set the sort order for data display and reports
Translation	Edit translation texts that appear in the user interface.Check the translation for missing or hidden text.Automatically fix inconsistencies in the translation file.
Layout	 Restore the application to its default layout settings. The application remembers your screen layout if you resize any screen elements. Scale the application, including text size and button size, to suit your screen. A set of miscellaneous controls.
Logging	View the location of the folder where the logs are stored; open the folder.Set the level of detail collected in log files. Note: setting the level to trace may slow down the application.
Databases	Lists the current database in use.Backs up all databases to the back folder to a chosen backup folder.
Advanced	- Resets all customizations made to your application.
About	Contact information.Links to documentation, i.e. the protocol and forms, and this manual.Version number and license agreement

Changing your log-in password

You can replace your existing password with a new one from the **General** tab in the **Settings** form. However, you do need to know your existing password to be able to replace it.

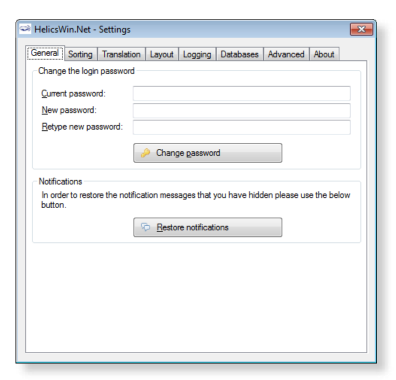
Important:

For security reasons, if you forget your password you will not be able to recover it and so you will be locked out of the application. Re-setting your password requires special intervention from your National PPS Co-ordinating Centre or equivalent.

To change your log-in password:

1. Click **Settings** in the *main menu*.

The **Settings** form opens; by default the **General** tab is displayed.

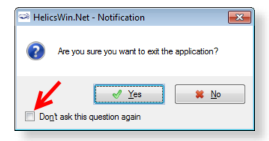


- **2.** In the **Current password** field, enter your existing password.
- **3.** In the **New password** field, enter your new password, and then enter it again in the **Retype new password field**.
- **4.** Click **Change password** to implement the change.

The **General** tab also allows you to restore notifications.

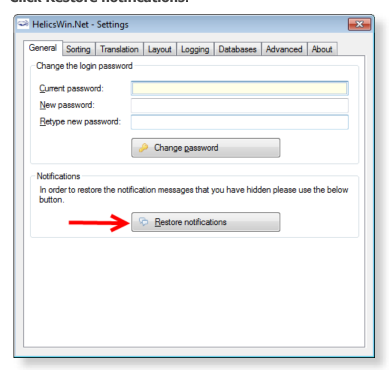
Notifications

A number of dialog boxes allow you to select not to see that particular notification again.



If you later wish to see all the notifications that you have selected not to see:

- Click Settings in the main menu.
 The Settings form opens; by default the General tab is displayed.
- 2. Click Restore notifications.

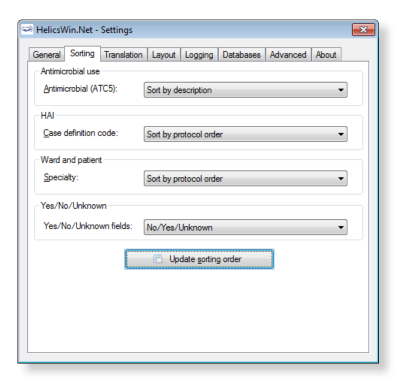


Sorting drop-down lists

To change the sorting order for options displayed in drop-down lists:

Click Settings in the main menu.
 The Settings form opens; by default the General tab is displayed.

2. Click the Sorting tab.



- **3.** For each section on this form, select your preference for the sorting order, see Table 27.
- 4. Click **Update sorting order**.

The preferences you have set are now available in the software application.

Table 27: Sorting order alternatives

List	Options
Antimicrobial use	Sort by description. Sort by code (alphabetically).
HAI – Case Definition	Sort by protocol. Sort by code (alphabetically).
Ward and patent - Speciality	Sort by protocol. Sort by code (alphabetically).
Yes/No/Unknown fields	Either 'No/Yes/Unknown' or 'No/Unknown/Yes'. Also, you will be requested to choose this if you translate the HelicsWin.Net user interface.

Translating the text in user forms

HelicsWin.Net is very flexible when it comes to translation. You can translate the texts manually from the default language, which is English. The items that can be translated include: Labels on the forms, Buttons, Form titles, and Error and warning messages.

The customizations you implement apply only to your current PC. However, it is possible to export a language translation file **Translation.mdb**. This pre-prepared file can then be run on any other PC running HelicsWin.Net. ECDC invites users and coordinating centres who have made a translation to send their zipped translation file **Translation.mdb** to <code>hainet@ecdc.europa.eu</code>. In this way, a central language database can be made available to all users.

Recommendation: To avoid duplication of work, translations are best performed at the national level. The National PPS Coordinating Centre can then distribute the database file Translation.mdb to the hospitals.

You can either translate user form labels and texts:

Directly in HelicsWin.Net – see Translating user forms in HelicsWin.Net

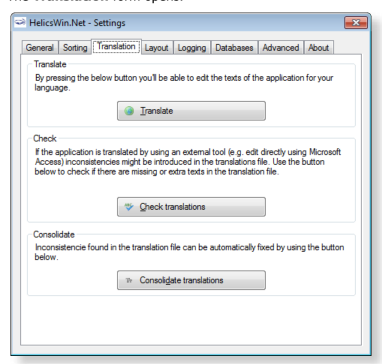
or

In either MS Access or MS Excel – see Translating user forms using a pre-prepared file.

Translating user forms in HelicsWin.Net

To translate the default labels to your language in HelicsWin.Net:

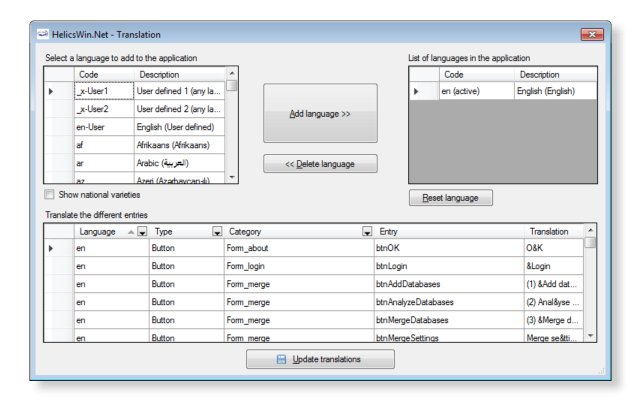
Click Settings in the main menu, and then click the Translation tab.
 The Translation form opens.



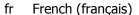
Please note that the Check translations and Consolidate translations options are only used in connection with *translating user forms using a pre-prepared file*.

2. Click Translate.

The **Translation** form opens.



3. In the Select a language... list, select the language you want to translate to, for example,



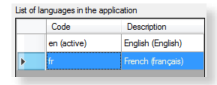


4. Click Add language.

The selected language is added to the list under **List of Languages in the application.**

5. Select the new language from this list.

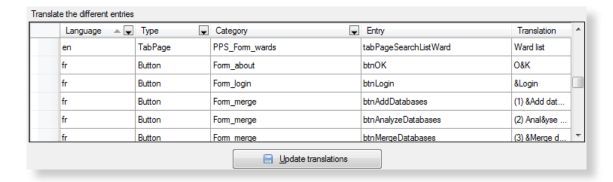
The selection indicator ▶ points to the selected language.



The labels for the foreign language are shown in the **Translate the different entries** grid at the bottom of the form.

The entries in the **Translate** column are initially simply the English labels, and you must translate them one by one.

6. You can make the data easier to manage if you apply filters to the column headers to reduce the number of items visible in the table, for example:



You can filter each column just as would do in Excel or Access, by clicking the down arrow, and making a selection. **Examples:**

To show only the names in the **Reports** form:

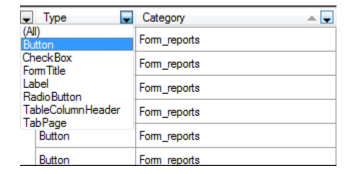
Click the down arrows to the right of **Category**, and select **Form-reports**.

Category

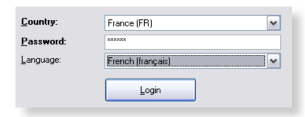
Fom_about
FORM_ABOUT
FORM_CHECK
Fom_login
FORM_LOGIN
FORM_MAIN
Fom_merge
FORM_MERGE
Fom_merge_settings
Fom_reports

To show only the button names in the **Reports** form:

Expand the **Type** column header and select **Button**.



- **7.** In the translation column, replace the English text, for example, "Cancel" with the corresponding text is your language, for example, "Annuler" in French.
- **8.** Repeat these steps for all labels in the user interface that you want to change.
- **9.** After each update, restart the application. When when you login, select the language which you translated (for example, French).



Creating and translating keyboard shortcuts

In any user form, any text character that is preceded by "%" (ampersand character) is displayed underlined; HWN automatically assigns a keyboard shortcut *Alt+underlined_character*. The characters preceded by "%" do not need to be used in the English language (e.g. D&éfinition).

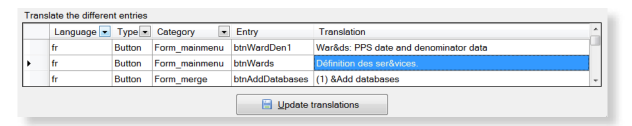
The behaviour for each control type is as follows:

Control type	Behaviour
Field	Cursor moves into the field
Button	Same as clicking the button
Tab	Same as clicking the tab

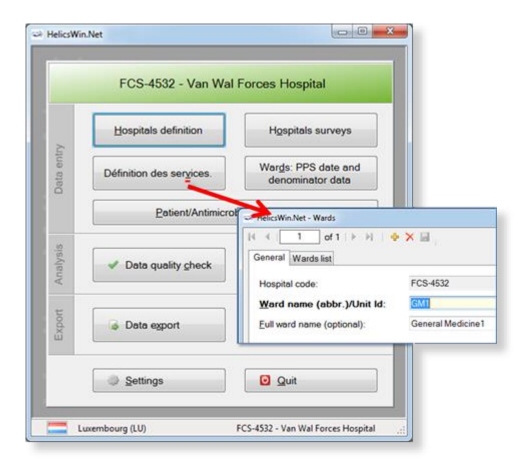
Translating keyboard shortcuts

When you switch to another language, translating text will remove the original keyboard shortcuts connected to that text. Keyboard shortcuts can be implemented in the translation tool, within the text entered in the column labelled **Translation**, by entering '&' (the ampersand symbol) before the letter for the shortcut. This is illustrated in the following example.

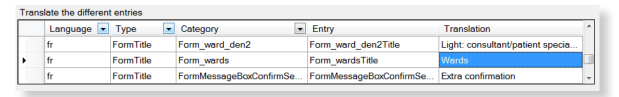
The following screenshot shows that the French translation of the button in the *main menu* for **Ward definition** has been translated to **Définition des services.** The shortcut has been set to **v** by placing '&' before the v in services, i.e. **Définition des ser&vices.**



The following screenshot shows the result of the translation. The button Ward definition has been changed to Définition des services. The shortcut V is indicated by underlining, i.e. Définition des services. Clicking v here will open the **Wards** form.

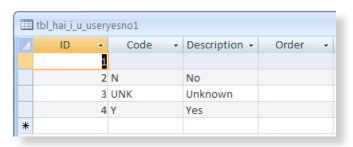


Note that the **Wards** form is still labelled HelicsWin.Net – Wards. This field is translated in the Translation form in Type *FormTitle*.



Translating drop-down list items

The labels in the drop-down lists are not stored in the file Translation.mdb, and therefore you cannot find them in the **Translation** form. These labels are stored in a separate database file, **Reference.mdb**. These can be translated there using Microsoft Access. See also *Translating user forms using a pre-prepared file*.



For example, in a French translation, you can change the value in the **Description** field from "Unknown" to "Inconnu".

Warning: Do not change the values of any code (for example, "UNK") in Reference mdb. If you

do, the application may not work properly, and you will also have problems when you $\,$

come to upload your data to TESSy (nationally nominated users only.)

Note: If you change the labels (descriptions) in the Reference.mdb file, the change is

implemented for all languages, independently of the language chosen at login.

Therefore, always make a backup copy of the original Reference.mdb before starting

to translate the value labels.

Translating user forms using a pre-prepared file

As an alternative to *translating user forms directly in HelicsWin.Net*, you can edit the translation file, **Translation mdb** in Microsoft Access.

To translate HelicsWin.Net labels in Microsoft Access:

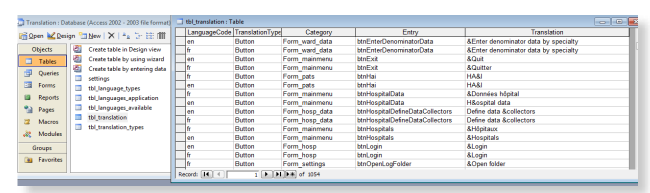
- **1.** Repeat steps 1 to 5 in the procedure *Translating user forms in HelicsWin.Net*.
- **2.** Click **Update**.
- 3. Close HelicsWin.Net.

This ensures that the labels for your language will be available in Microsoft Access. HelicsWin.Net only saves your changes in the **Transaction.mdb** file when HelicsWin.Net is closed.

4. Open the **Transaction.mdb** file in Microsoft Access.

This file is located in the HWN folder (the default is C:\HWN2).

- **5.** Open the **tbl_translation** table.
- **6.** Filter the **Language** column to show only entries for you language (for example, "fr" for French).



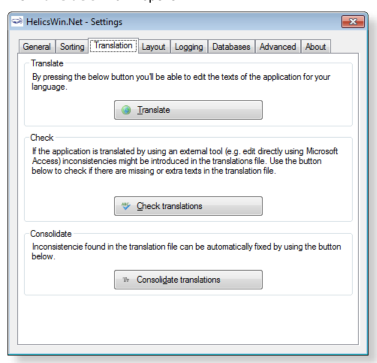
- **7.** Add your translations to **Translation** column.
- 8. Save Transaction.mdb.
- **9.** Copy your updated **Transaction.mdb** to the HWN folder and overwrite the existing Translation.mdb.
- 10. Restart HelicsWin.Net.
- **11.** Check the translation in HelicsWin.Net, see *Checking your translation*.

Checking your translation

When you have translated outside HelicsWin.Net, there is a risk of errors occurring in the **Transaction.mdb** file – for example, an accidental change to a field outside the Translation column, or the unintentional deletion or insertion of a row.

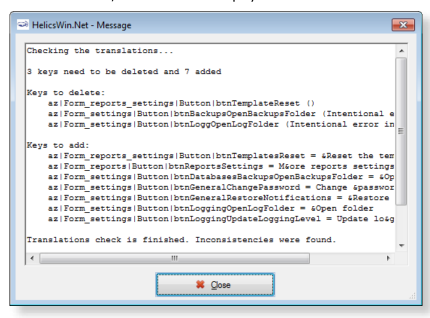
Once you have replaced the **Transaction.mdb** file in in the HWN folder (the default is C:\HWN2), you should check the translation database for errors:

- **1.** Close and restart HelicsWin.Net to reload the translation database.
- Click Settings in the main menu, and then click the Translation tab.The Translation form opens.



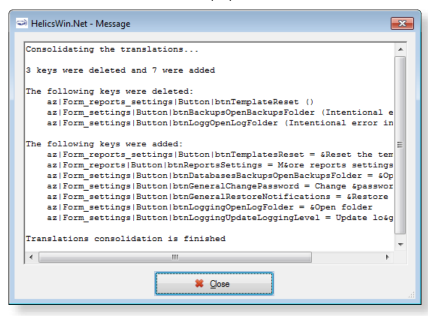
3. Click Check translations.

If there are errors, a list of these is displayed.



- 4. Click Close.
- Click Consolidate translations.

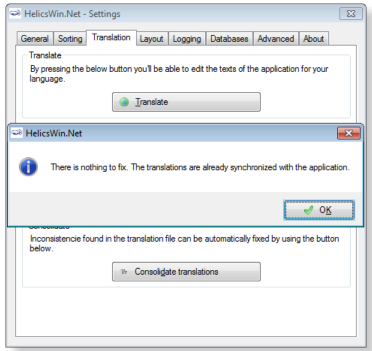
A list of the corrections made is displayed.



The corrections made do not, of course, insert a translation, but only restore the default setting for the incorrect key.

- 6. Note the errors.
- **7.** Finalise correction of the errors by manually translating the keys using the procedure described in *Translating user forms in HelicsWin.Net*).

If your translation of the **Transaction.mdb** file contains no errors, the following message is displayed when you click **Check translations**.

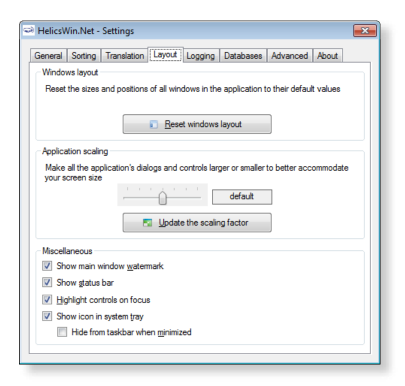


Resetting form layout options

If you change the size or the position of the windows, HWN remembers the windows settings the next time HWN is launched.

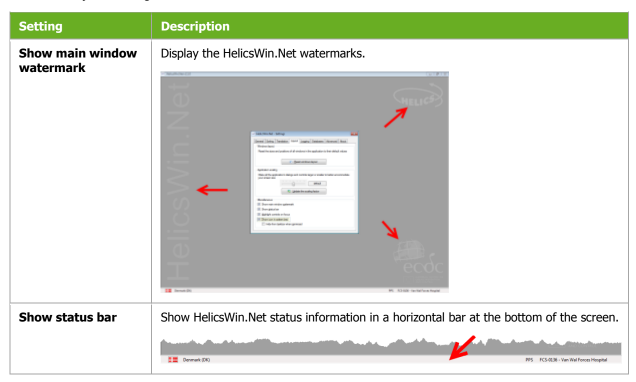
To reset the size and position of all HWN windows to their original values:

Click Settings in the main menu, and then click the Layout tab.
 The Layout form opens.



- Click Reset windows layout to revert to HelicsWin.Net's default display settings.Or
- **3.** Use the slider to increase or decrease the size of the HelicsWin.Net windows (requires you to log in again).
- **4.** Select or deselect the miscellaneous settigns options, see Table 28.

Table 28: Layout settings



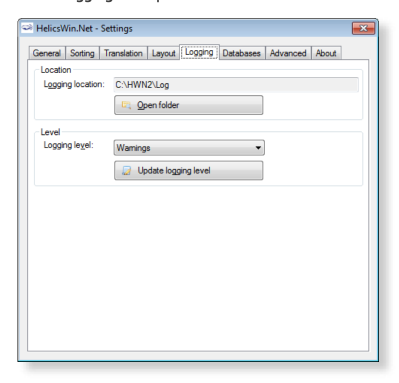
Highlight controls on focus	A control is highlighted when the cursor hovers over it, or when you tabulate to it.		
Show icon in system tray	Add the HelicsWin.Net icon in the system tray (located in the Windows taskbar at the bottom of the screen next to the clock) for quick access to HelicsWin.Net.		
	② ② ③ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □		
Hide from taskbar when minimized	When HelicsWin.Net is minimized, the HelicsWin.Net icon is not displayed on the taskbar at the bottom of the screen.		

Log file for debugging

Each session of HWN is logged so potential issues can be easily diagnosed. Using these files can save debugging time in the event of a crash of the application at user level.

The log files are saved in the **Logs** folder in the HelicsWin.Net installation folder. You can set the logging level to produce logs at different levels of detail.

- 1. Click **Settings** in the *main menu*, and then click the **Logging** tab.
 - **1.**The **Logging** form opens.



2. Click **Open folder** to access the log files.

- **3.** Adjust the logging level, if desired.
- 4. Click Update logging level.

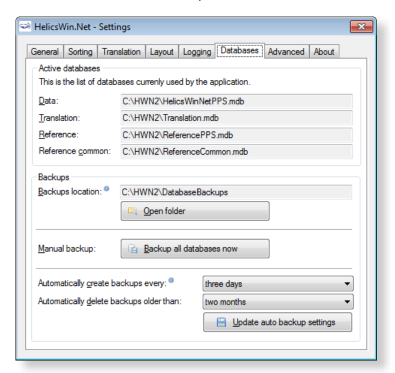
Table 29: Logging levels

Level	Logging
None	No messages are logged.
Errors	Only error messages are logged.
Warnings	Warning and error messages are logged.
Info	Information, warning and error messages are logged.
Trace	Trace, information, warning and error messages are logged.

See Table 23: Data quality check messages for a description of error and warning messages.

HelicsWin.Net databases

The **Databases** tab shows the paths to the database **.mdb** files and the backup locations.



- **1.** Click **Open folder** to select a backup location.
- 2. Click **Backup all databases now** to make a backup.

HelicsWin.net allows you to select automatic backup creation and deletion:

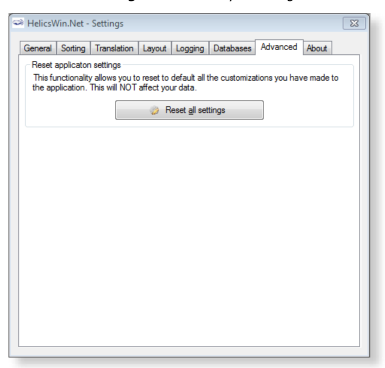
1. Select a frequency for the automatic backup creation and deletion from the respective drop-down lists.

2. Click **Update auto backup settings** to apply your choice.

Resetting default settings

The **Advance** tab enables you to remove all configurations you have made to HelicsWin.Net by restoring the default settings.

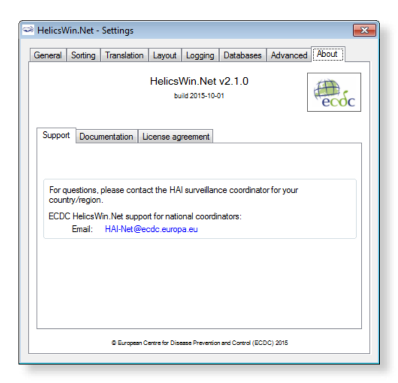
Click **Rest all settings** to remove all your configurations.



About HelicsWin.Net

The **About** tab in the **Settings** form contains three tabs:

- Support
- Documentation
- License agreement



HelicsWin.Net HelpDesk support

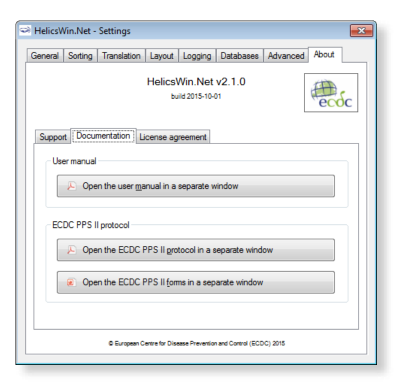
The first-line HelpDesk for HelicsWin.Net is at the national level, ensured by the national/regional PPS coordinator. These coordinators may refer questions to ECDC by posting them on the HAI-Net Extranet, or by sending an email to <code>HAINET@ecdc.europa.eu</code>.

Accessing documentation, protocols and forms

The **Documentation** tab contains links to:

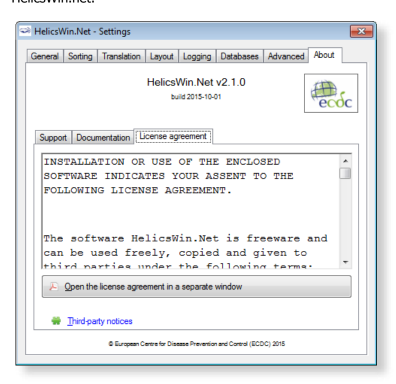
- A PDF version of the user manual.
- The PPS and ICU protocols.
- The PPS and ICU forms.

These documents can also be found in the **Documentation** folder (e.g. C:\HWN2\Documentation).



License agreement

The **License agreement** tab contains a transcript of the HelicsWin.Net license terms (which you are requested to read closely) and a link to information on the third-party products included in HelicsWin.net.



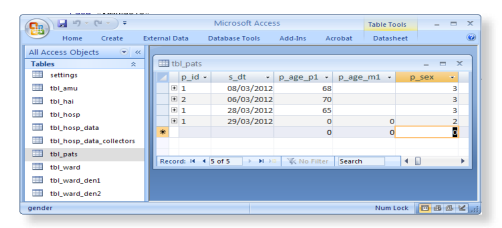
HelicsWin.Net database files

The database files used to store your data and settings are accessible to anyone who has access to the PC on which the application is installed.

Structure of the HelicsWinNet.mdb database

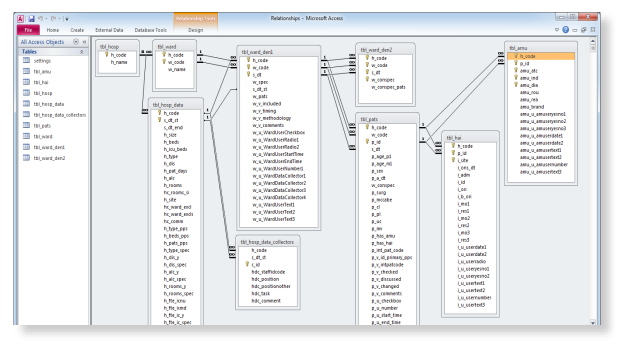
The data are stored in the internal software format in the HelicsWinNet.mdb database (Microsoft Access 2000 format) which is stored in the folder where the programme was installed. For data transfer to a national coordinating centre, this .mdb file can be sent by email, preferably after compression using a password.

If you open the file in Microsoft Access, you can see the tables and their data, as shown below:



Relationships

You can view the relationships between the tables in the HelicsWinNet.mdb database using standard Access tools.

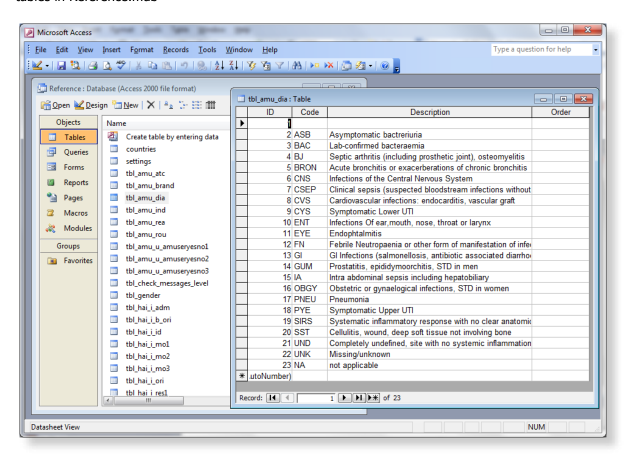


HelicsWin.Net variables and values

The internal data structure of the HWN databases differs from that of the TESSy database. The variable names and code values stored in the database are also different. The internal code values of HWN are stored in the Reference.mdb database.

For a description of the HWN variable names and the corresponding TESSy variables, see the HelicsWin.Net protocols.

The following screenshot shows the internal code values in HWN that are stored in the ID field in the tables in Reference.mdb



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