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Once for Wales Concerns Management System

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## Datix Cymru

# INCIDENTS MODULE

## Full Reporter / Investigator / Handler Guide

A guide to the reporting, investigating, and handling of the Incidents Module.

### Functionality User Guide

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*The screenshots and examples used in this guide are drawn from test records within the All-Wales Demonstration system and do not include identifiable information.*

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## Introduction

This document provides you with a guide for how to use the Incident Reporting and Management Functionality. Incident records allow you to capture and manage all details relating to an incident record and any relevant external reporting. You can also capture and manage actions and action plans that have been assigned.

You can review a full notification and message history for the record, information about and the response, also save attachments or reports with additional information.

The information entered into the system is also used for reporting and analysis purposes and so it is vital that the data is accurate and useful.

For additional information on features such as reporting and other functionality common across the Capture module, there will be a separate guide developed.






Appendix One contains more detailed information on the focus reviews required for incidents relating to patient falls, medication, equipment, sharps injuries, manual handling, and pressure damage. There is also a focused review additional information section that requires completion following a focused review.

Appendix Two provides details on the Once for Wales Code Set and further information regarding the Dictionary of Medication and Devices.

## Logged in Users

Before you add a new incident record to the system you must do a full search using the contacts module to establish if the person affected or anyone else involved is already a contact in the system. If an incident record is found you will need to click on the 'generate from' key to create a new record, this will also link the records together. More information on how to generate from can be found under 'How to add a new incident record'.

## Aspects of the Form

-  This icon indicates that a field is mandatory, and you are required to complete it before saving or submitting the form.
-  This icon indicates that the field you are completing is a dropdown list. Clicking this icon will allow you to select the relevant option(s).
-  This icon indicates a date field. Clicking the icon will allow you to select a date from a calendar, or you can simply type the date in using the dd/mm/yyyy format.
-  Any field that shows this icon next to it indicates that there is additional information available to help you complete it correctly. Click the icon to view the additional guidance.
-  In a multi-select field, where you can choose more than one option from a dropdown, clicking this icon will remove the currently selected value(s)



Save button

## How to Access the System

You can record an incident record by clicking on the URL link.

If you are sent an email notifying you of the incident, the email will contain the URL link that will take you directly to the incident record once you login.

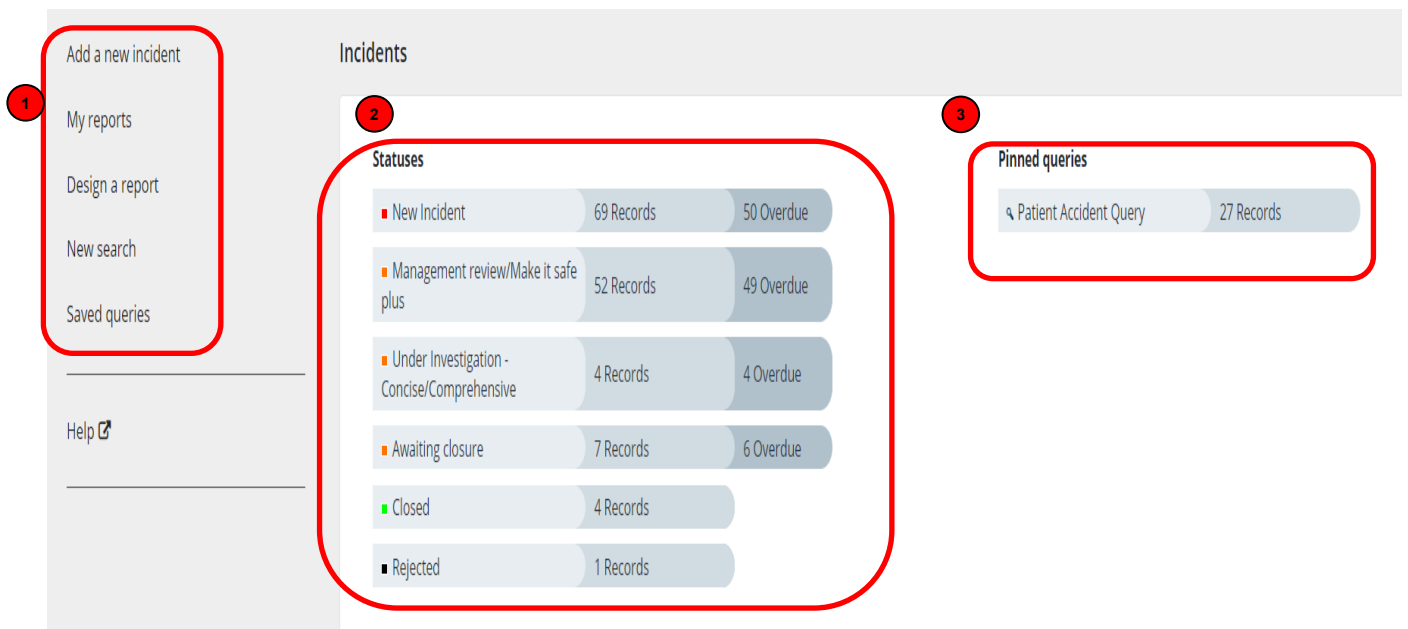
If you are an authorised user and need to login to the system, you can do this by using your Active Directory account. This is the same login name and password you use to login to your PC.

If you encounter any issues, then please contact your Local System Lead at the Health Board / Trust.

## Landing Page

When you login to the Incident Reporting and Management functionality you will be presented with the landing page below, the status area contains all the tools necessary to add a new record, manage a record and produce statistical and listing reports.

1. The left-hand navigation panel contains all the tools necessary to add a new incident, manage a record and produce statistical and listing reports.
2. The status area records where the incident record sits within the workflow.
3. Pinned queries allow instant access to the most frequently used searches.



The below table provides the information that is displayed on the landing page:

	<b>Field name</b>	<b>Explanation</b>
Left -hand navigation panel	Add new Incident	If you click on this, you can add a new Incident record
	My reports	If you click on this, you can run saved custom reports
	Design a report	If you click on this, you can create a custom report
	New Search	If you click on this, you can search for records
	Saved queries	If you click on this, you can use saved queries for re-use
Statuses	New incident	This is a holding area when a new incident is reported.
	Management Review/Make it safe plus	These records are awaiting a review and make it safe actions by a manager within 2 working days.
	Under investigation	This status indicates that the incident has undergone a management review and initial make it safe actions and has been identified for further investigation within 25 working days.
	Awaiting Closure	This status indicates that all appropriate actions have been taken and the incident is awaiting final review before closure within 30 working days.
	Closed	All actions are complete, and record has been closed.
	Rejected	This status is used for any rejected incidents, these can be duplicate incidents or those not identified as incidents.
Pinned Queries	Pinned Queries	Pinned queries allow instant access to the most frequently used searches. If you click on any of the pinned queries it will open the records within the query

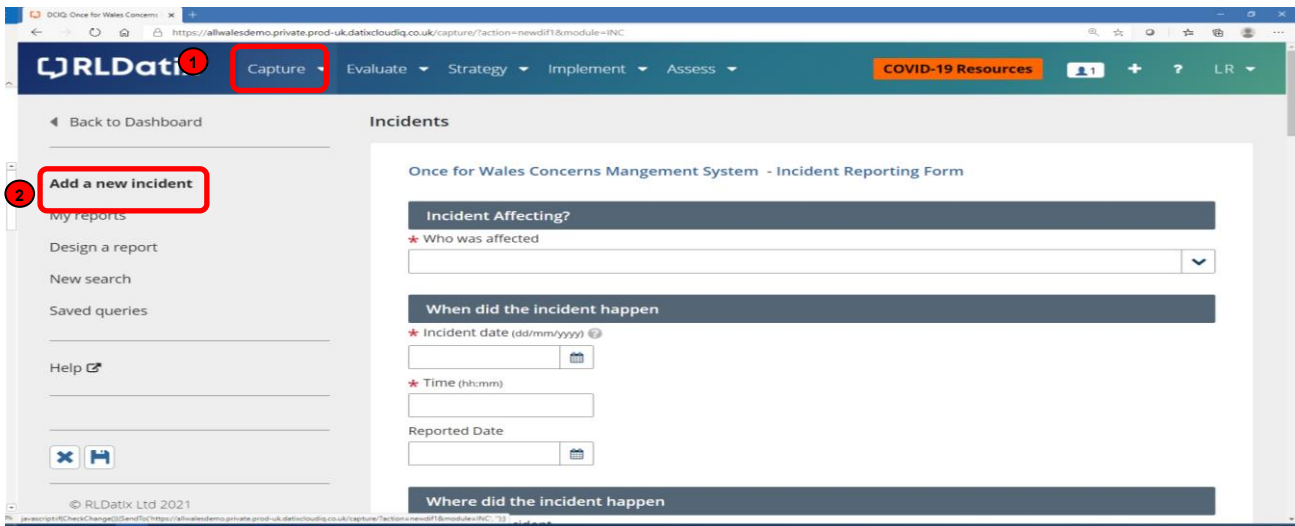
If you click on any of the status areas, it will take you to a listing page. The records displayed can be sorted by clicking on any column heading. Select the record you wish to view by clicking on any of the numbers/words.

## How to Add a New Incident Record

If you logged out of the system you can click on the link to the reporting form.

If you are logged into the system, please:

1. Click Capture > Incidents in the top application menu.
2. Click Add new Incident in the left-hand navigation menu.



Enter the information for the incident record ensuring all mandatory fields are completed.

The fields that you are required to complete to add a new Incident record are described in the table below, this includes the section names, field names, descriptors and what classification of incident information will display what fields within the reporting form:

Section	Field	Description
<b>Incident Affecting?</b>	Who was affected	Select the relevant field from the drop-down menu.
<b>People Affected</b>	ID	Auto populated.
	Contact Role	Auto populated.
	Contact Type	If the contact type is anything other than Service User/Relative/Public, a search of the contact module can be undertaken by entering the forename and surname of the person affected.  If no record is found, the following fields relating to the person involved in the incident are available to be completed manually: Sub type Title Forenames Date of Birth Date of Death Email Postcode.

	Field	Description
<b>People Affected</b>	NHS/ID Number	If the contact type is a Patient/Service User, the search function will trigger. You can select to search via Nadex, NHS number.
	ID Number	Enter the appropriate number and search.
	Subtype	Select from drop-down menu.
	Title	Enter title
	Forenames	Enter forename(s)
	Surname	Enter surname
	Date of birth	Enter date of birth.
	Date of death	Enter date of death, if applicable.
	Email	Enter email
	Address Line 1	Enter first line of address.
	Address Line 2	Enter second line of address.
	Address Line 3	Enter third line of address.
	Postcode	Enter postcode
	Primary Contact Number	Enter primary contact number.
	Secondary Contact Number	Enter secondary contact number.
	Was the person injured in the incident?	If yes, this will trigger an Injury Details section.
<b>Injury Details</b>	Injury	Select type of injury form drop-down menu.
	Body part	Select which body part the injury is location on in the drop-down menu.
	Treatment	Select the relevant treatment received for the injury.
	Add another	The Add Another field allows you to capture details of other persons affected.
<b>When did the incident happen</b>	Incident date (dd/mm/yyyy)	The date the incident occurred.
	Time (hh:mm)	The time the incident occurred.
	Reported Date	This record auto populated at time of reporting.
<b>Responsible Service</b>	Incident Service	The Service under which the incident occurred. <b>Not visible on Primary Care Reporting Forms.</b>
<b>Where did the Incident happen?</b>	Location of incident	The Location where the incident took place.
	Exact Location	Please enter the exact location the incident took place if not available in Location field.
<b>Incident Severity</b>	Reporter's Initial Harm Assessment	Enter the level of harm from a drop-down menu options
	Does this Incident need external reporting?	Select yes if it needs to be reported externally
<b>Incident Details/What Happened</b>	Description	Enter details of the incident, enter only the facts not personal opinions and do not enter any personal identifiable information. i.e., names.
	Brief Description of actions taken	Enter details of actions taken at the time of the incident.
	Vehicle registration number	Enter details as appropriate e.g., vehicle involved.
	Booking/CAS number if applicable (WAST)	Enter details as appropriate. This is a WAST reference number.
	Laboratory specimen number	Enter details as appropriate e.g., specimen involved.

	<b>Field</b>	<b>Description</b>
<b>Incident Type</b> (Entries in this section may trigger additional questions for completion by the reporter)	Classification	Select the Classification of Incident that has occurred from the options in the drop-down menu
	Category	Enter the Category of incident that has occurred from the drop-down menu.
	Sub Category	Enter the Sub Category of incident that has occurred from the drop-down menu.
	Fire Additional Options	Please select all appropriate options if this field has been made visible.
	Fire alarm activation: Additional Options	Please select all appropriate options if this field has been made visible.
	Was a ligature used?	Please select Yes or No as appropriate if this field has been made visible.
	Method violence and aggression was received by	Please select appropriate option if this field has been made visible.
	Was absconder detained under the Mental Health Act?	Please select Yes or No as appropriate if this field has been made visible.
	Has a Perpetrator been identified?	Select 'Yes' to make the Perpetrator form visible or 'No' if you haven't got specific details and text field will be visible to capture any known details
	Please provide as much information that is known about the perpetrator	Please provide as much detail as possible in this text field to assist the case manager.
<b>Restrictive Practice</b> (Triggered for Physical Restraint event – Care planned and Physical restraint event – Not care planned incidents).	What model of physical intervention (training) used?	Please select all appropriate options from list.
	What technique was used?	Please select appropriate option from list.
	Who took the lead in the intervention	Please select appropriate staff from options from list.
	What other staff were involved in intervention?	Please select appropriate staff from options from list.
	Time physical intervention commenced	Please end the time in the physical intervention commenced.
	Time physical intervention concluded	Please end the time in the physical intervention concluded.
	Legal status of patient / service user?	Please select appropriate option from list.
	Why was intervention used?	Please select appropriate option from list.
	Was medication used?	Please select Yes or No from drop down
	Who administered the medication?	Please enter the information of the person who administered the medication in this field.
	Was any other person injured during the intervention?	Please select appropriate option from list.
	What was the outcome for the patient / service user	Please select appropriate option from list.
	Was there a post incident welfare debrief with staff?	Please select Yes or No from drop down
<b>Additional Information</b>	Was any equipment involved in the incident?	Select yes if there is any equipment involved in the incident. If yes is selected and equipment section will be visible to complete.
	Did medication have a direct impact in this incident?	Select yes if there is any medication involved in the incident. A controlled drug field will be visible if yes is selected.
	Was there any controlled drug involved in the incident?	Select yes if there is any controlled drug involved in the incident.
	Is this Incident related to EPMA (Electronic Prescribing and Medicines Administration)?	Please select Yes or No as appropriate

	Does the incident have Information Governance considerations?	Select yes if there are any information governance considerations related to the incident, If yes is selected an Information Governance section will be visible.
	Does this incident have any safeguarding elements?	Select yes if any safeguarding elements relate to the incident. If yes, is selected a further text field will be visible to capture the process to be followed.
	Are there any Additional Factors relating to this Incident?	Please select all relevant options from the drop down menu.
	Date of Industrial Action	If visible please enter date of Industrial Action as appropriate
	Further Information pertinent to Industrial Action	If visible please enter any relevant information to the Industrial Action
	Is this incident about nursing care?	Select yes if the incident is related to nursing care.
	IPC antigens	Please select the IPC antigen from the list available
	Were temporary staff involved in the incident?	Please select Yes or No as appropriate
	Was any other contact involved in the incident	If there were other contacts involved complete the other contact fields that will be triggered by answering yes to this question.

<b>Information Governance</b>	Has personal data been disclosed outside of the organisation?	Select yes if the information has been disclosed outside of the organisation.  If yes is selected, please select the type of personal data that is included. This is a multi-pick option. If other is selected a further field will populate to capture the information.
	What type of personal data is included?	Select all that apply from drop down menu
	Has there been (or is likely to be) any impact to the individuals as a result of the incident?	This is a multi-pick option, please select all relevant options.
	Other adverse effects please specify	Enter other adverse effects.
<b>Communication</b>	Is this incident highly confidential (not for circulation)?	Select yes if this incident is not for wide circulation. This field should also be used in "Freedom to Speak Up Safely" cases.
	Incident Manager	Reporters should select their Line Manager or Departmental Manager.
	Who have you informed of the incident?	Select who you have informed about the incident from the drop-down list.
	Please select which 'Other NHS Body' has been informed	Select from drop down which other NHS body has been informed.
	Other NHS Body, please provide more detail	If 'other' is selected, please provide further details
<b>Medications</b>	Please add medication below, if you are unable to find name in medication section	Enter name of medication.
<b>Medications Administered</b>	Search for drug administered or omitted	

	Brand name of drug administered	Auto populated by search.
	Manufacturer of drug administered	Auto populated by search.
	Class of drug administered	Auto populated by search.
	Type of drug administered, prescribed, dispensed or omitted	Auto populated by search.
	Route administered, prescribed, dispensed or omitted	Select from drop-down menu.
	Dose administered, prescribed, dispensed or omitted	Select from drop-down menu.
	Form administered, prescribed, dispensed or omitted	Select from drop-down menu.
	Drug administered or omitted BNF classification (1-15)	Select from drop-down menu.
	Drug administered or omitted batch number	Enter batch number.
	Drug administered or omitted ethics committee name and reference	Enter name and reference.
	Is the drug administered or omitted under clinical trial?	Select from drop-down menu.
	Is the drug administered or omitted a manufactured special?	Select from drop-down menu.
	Is the drug administered or omitted a parallel import?	Select from drop-down menu.
	Stage at which error occurred	Select from drop-down menu.
	Type of error	Select from drop-down menu.
	Notes	Notes on type of error.
	Other Important Factors	Provide any other important factors.
<b>Medications Intended / Suspected</b>	Search for intended / suspected drug	
	Brand name of drug intended / suspected	Auto populated by search.
	Manufacturer of drug intended / suspected	Auto populated by search.
	Class of drug intended / suspected	Auto populated by search.
	Type of drug intended / suspected	Auto populated by search.
	Route intended / suspected	Select from drop-down menu.
	Dose intended / suspected	Select from drop-down menu.
	Form intended / suspected	Select from drop-down menu.
	Drug intended BNF classification (1-15)	Select from drop-down menu.
	Intended drug batch number	Enter batch number.
	Drug intended ethics committee name and reference	Enter name and reference
	Is the drug intended under clinical trial?	Select from drop-down menu.
	Is the drug intended a manufactured special?	Select from drop-down menu.
	Is the drug intended a parallel import?	Select from drop-down menu.
<b>Equipment</b>	Type of product	Select from drop-down menu.
	Manufacturer name	Enter name of manufacturer
	Serial Number	Enter serial number.
	Equipment ID	Enter equipment ID number
	Description and location of the defect/problem	Provide details of defect/problem.

<b>Contact (triggered from question in Additional Information)</b>	Contact Type	Select the contact type from a drop-down list.
<b>Other Contact</b>	Role (Persons)	<p>If the contact type is anything other than 'Patient/Service user', a search of the contact module can be undertaken by entering the forename and surname of the person affected.</p> <p>If no record is found, the following fields relating to the person providing feedback are available to be completed manually:  Email:  Address Line 1  Address Line 2  Address Line 3  Postcode  Primary Contact number  Secondary Contact number  Language (preferred language for contact)  Was the person injured in the incident? If yes, this will trigger an Injury detail section.</p>
<b>Injury Details</b>	Injury	Select type of injury form drop-down menu
	Body part	Select which body part the injury is located on in the drop-down menu
	Treatment	Select the relevant treatment received for the injury
	Add another	The Add Another field allows you to capture details of another injury.
<b>Openness and Transparency (Triggered for incidents graded Moderate and above).</b>	Was the patient/appropriate person informed that an incident occurred?	Yes/No drop down
	When was the Patient/Service User /appropriate person informed? (dd/mm/yyyy)	Enter date the discussion took place
	Were the members of staff involved in the incident involved in informing the Patient/Service User/appropriate person?	Yes/No drop down
	Please provide details of staff members who informed the Patient/Service User/appropriate person.	Enter details of staff who were involved
	Please provide details of the Patient/Service User/appropriate person who was informed.	Enter details of the Patient/Service User/appropriate person who was informed
	Was an apology provided to the Patient/Service User/appropriate person?	Yes/No drop down
	Was a truthful account of the facts known at the time shared with the Patient/Service User/appropriate person?	Yes/No drop down

	Was the Patient/Service User/appropriate person advised about next investigative steps to be undertaken?	Yes/No drop down
	<b>Field</b>	<b>Description</b>
<b>Documents</b>	Are there any documents to be attached to this record	Select yes if you have any documents to attach on submission, this will trigger fields at the bottom of the form. You will also be able to attach documents after submission.
<b>Documents (triggered from question in Additional Information)</b>	Link as	Select from drop down
<b>Details of Person reporting the Incident</b>	Reporter	If you are logged in your details will automatically populate. If you are not logged in, you will need to add your details in this field and then submit the Incident record
<b>Additional Reporter Details</b>	Reporters Location	Please enter the Location of the Reporter at the time the Incident took place.
	Reporters Service	Please enter the Service the Reporter was under at the time the Incident took place.

Note: For the Details of person affected section the Reporter section and any other contact section, you can clear the information that may be automatically populated by clicking Clear section, within the section. When a new Incident Record is recorded an email notification can be sent to the relevant Handler/Investigator.

## People Involved

It is important that anyone that is involved in the Incident is linked as a contact. To add any other contacts to a record, click as appropriate one of the following links:

- Create a new Person Affected link
- Create a new Other contact link

Complete all mandatory fields and click Check for Matching Contacts to ensure that the contact does not already exist. If a record exists, click Choose and complete any incident specific information as appropriate.

If a record does not exist, click Cancel and complete the Contact Details and any Link Details. Click Create New Link > Save.

## Contact Matching

The system not only integrates with the Master Patient/Service user Index but also contains a large contact database in the background which provides you with incredibly useful information. This will enable you to search on how many times an individual has been involved in an incident, complaint etc. If you need to manually enter a contact then the person responsible for managing the record will need to perform an “approval” process on each contact submitted to ensure that duplicate contact records are not created, the new contact will initially be marked as ‘unapproved’ and they will need to approve or reject them.

It is extremely important that you always search for contacts before creating any new contact records. If new contacts are created the person responsible for handling/managing the record must ensure they follow the correct process when they approve the contact. **It is extremely important that we do not create duplicate contacts in Datix Cymru wherever possible.**

In the contact type field, you must select the Type of Contact that you are searching for.

### **For Patient/Service User**

When the Contact Type of Patient/Service User is selected, you will need to select NHS number in the ID number Type field, in the ID number field enter the Patient/Service Users NHS number and select search, the system is integrated with the Master Patient Index and will pull the demographic details of the Patient/Service User into the system.

If you do not have the NHS number and the Patient/Service User’s contact information is already in the database, you can also search using the Hospital number.

If you do not have the Patient/Service User’s NHS number, you will need to look this information up in your Patient/Service User Information system (e.g. Myrddin).

Once you confirm this is the correct Patient/Service User, select choose, if it is not please click clear section. A further search will need to be carried out or manually add the Patient/Service User’s details in.

**Type** Clear section

Contact Role  
Person Affected

Contact Type  
Patient

Search

**NHS/ID Number**  
When searching for patients details please search by NHS number

ID Number Type  
NHS number

ID Number  
12345678

Search

Add more contact numbers

**Person Affected**  
When searching for staff please search using forename, surname and email  
When searching for any other contacts please search using forename & surname only

\* Subtype

Choose	ID	Surname	Forenames	Type	Subtype
Choose	1	TEST	TEST	Patient	Service User

Basic demographic information will be returned at this point, however additional Patient/Service User's demographic information will be available in the contacts database. If any data is missing from visible fields and you have this information, then please add it to the record e.g., e-mail address. The data that has been populated will be greyed out and read only on the form.

**Person Affected**  
When searching for staff please search using forename, surname and email  
When searching for any other contacts please search using forename & surname only

\* Subtype  
Service User

Title  
Mrs

\* Forenames  
TEST

\* Surname  
TEST

Date of birth (dd/mm/yyyy)  
01/01/1950

Date of death (dd/mm/yyyy)  
Fill this in for a patient who has died  
01/01/2019

Postcode  
CF33 999

\* Primary Contact Number  
01656 755555

\* Secondary Contact Number  
09874562

\* Was the person injured in the incident?  
No

Add another

### For Employee/Member of Staff

For best search results when searching for staff then please add in the forename, surname and email address. If you do not select all of these fields prior to searching you may be presented with a large number of contacts on your list to choose. If the employee/member of staff is not presented to you then you can manually input the staff details, however, this will then need to be checked and approved by the person responsible for managing the record.

### Any Other Contacts

When searching for any other contacts please search using forename & surname only. If the contact is not presented to you then you can manually input the staff details, however, this will then need to be checked and approved by the person responsible for managing the record.

\*Please note the search button is currently in the Type section of the form – we are working

to relocate this field to more suitable place on the form.

### Approval of Contacts

The person responsible for a managing the record will need to check all contacts are approved:

- Click 'People Involved' to confirm their contact details. This brings up a list of staff/people involved
- If the contact shows as unapproved (red) in the approval status – click on the name to confirm the contact details

People Involved											
Approval status	ID	Contact Type	Role (Persons)	ID Number Type	ID Numbers	Subtype	Surname	Forenames	Tel 1	Body part (primary)	Injury (primary)
Unapproved	395	Named Consultant	Person Affected				McAvoy	James			

Create a new Person Affected link

Other Contacts											
Approval status	ID	Contact Type	ID Number Type	ID Numbers	Subtype	Surname	Forenames	Date of birth	Age (Persons)	Language	Tel 1
Approved	394	Employee/Member of Staff			Additional Clinical Services	Crompton	Rachel				

Create a new Other Contact link

Check the contact details are correct and add any additional information you may have - At the bottom of form Click on the 'Check for matching contacts' button to confirm if there is already an approved contact for the contact.

**Contact Details**

\* Subtype

\* Title

Gender

Date of birth (dd/mm/yyyy)

Date of death (dd/mm/yyyy)

Tel 1

Tel 2

Disabilities

Nationality

Ethnicity

Religion

Marital status

Current approval status  
Unapproved

\* Approval status  
Unapproved

If you find the correct contact, please select choose. This contact record will then replace the contact record of the unapproved contact.

If no matching contacts are found the user needs to amend the 'Approval status' field. This is done by clicking on the drop-down arrow at the side of the field – the user should then select 'Approved' from the drop-down list.

The 'Approval status' field will then be updated. The user then needs to click the 'save' button – this will link the record to the user.

The contacts that have been searched for correctly and are available in the system will automatically be approved.

### To Generate a Record (left hand side panel)

1. Click Capture in the top application menu.
2. Select the module you want to generate a record from.
3. Find and select the appropriate record.
4. Click Generate from the left-hand navigation menu.
5. Select Incident in the Module to generate record in drop-down field.
6. Select the desired workflow status in the Approval status drop-down field.
7. Select the appropriate checkboxes as needed:
  - Copy links to contacts?
  - Copy documents?
  - Copy respondents?
  - Link generated record to master record.
8. Click Generate. The new feedback record is generated and linked to the existing record.

RLDatix | Capture Evaluate Strategy Implement Assess | COVID-19 Resources | 11 + ? CB

Claims

Generate from selected records

Generate options

5 \* Module to generate record in:  
Incidents

6 \* Approval status  
New Incident

7  Copy links to contacts?  
 Copy documents?  
 Copy causal factors?  
 Link generated record to master record?

8 Cancel Generate

## Incident Details

The information submitted from the reporting form will automatically pull through to the incident details, it is important to undertake data validation to ensure this information is correct. Additional questions are also available in incident details following the incident submission.

	Field	Description
<b>Name and reference</b>	ID Number	This is auto populated from reporting the incident
	Date Reported (dd/mm/yyyy)	This is auto populated from reporting the incident
	DWEB Reference Number (If applicable)	Enter a Datix Web reference number if applicable
	Vehicle Registration Number	Enter a vehicle registration where a vehicle was involved in the incident
	Booking/CAS number if applicable (WAST)	Enter details as appropriate. This is a WAST reference number.
	Laboratory specimen number	Enter details as appropriate e.g., specimen involved.
	Name of Person Affected	This is auto populated when a person affected is entered as a contact in the reporting form.
	Incident Manager	Please select the Line Manager or the person responsible for the management of the incident.
	<b>Incident Severity</b>	Reporters Initial harm assessment
<b>Details</b>	Incident date (dd/mm/yyyy)	The date the incident occurred.
	Time (hh:mm)	The time the incident occurred.
	Description	Enter details of the incident, enter only the facts not personal opinions and do not enter any personal identifiable information. i.e., names.
	Brief Description of actions taken	Enter details of actions taken at the time of the incident.
<b>Responsible Service</b>	Incident Service	The Service under which the incident occurred. <b>Not visible on Primary Care Reporting Forms.</b>
<b>Incident Type</b> (Entries in this section may trigger additional questions for completion by the reporter)	Who was affected?	Select who was affected by the incident from the options in the drop-down menu
	Classification	Select the Classification of Incident that has occurred from the options in the drop-down menu
	Category	Enter the Category of incident that has occurred from the drop-down menu.
	Sub Category	Enter the Sub Category of incident that has occurred from the drop-down menu.
<b>Additional Information</b>	Are there any Additional Factors relating to this Incident?	Please select all relevant options as appropriate
	Date of Industrial Action	If visible, please enter date of the industrial action
	Further information pertinent to Industrial Action	If visible, please type in any information pertinent to the industrial action
	Does the incident have	Select yes if there are any information governance

	Information Governance considerations?	considerations related to the incident. If yes is selected an Information Governance Team Initial Triage section will be visible.
	Does this incident have any safeguarding elements?	Select yes if there are any safeguarding elements related to the incident. If yes, is selected a further text field will be visible to capture the process to be followed.
	Was any equipment involved?	Select yes if there are any equipment involved in the incident.
	Did medication have a direct impact on this incident?	Select yes if there is any medication involved in the incident. A controlled drug field will be visible if yes is selected.
	Was there any controlled drug involved?	Select yes if there is any controlled drug involved in the incident.
	Is this incident related to EPMA (Electronic Prescribing and Medicines Administration)	Please select Yes or No as appropriate
	Were temporary staff involved in the incident?	Select the appropriate radio button.
<b>Information Governance Team Initial Triage</b>	IG reviewed by	Please select user who has reviewed the incident for IG
	Date IG reviewed	Please enter date IG reviewed the incident
	Confirmed IG issue	Please select Yes or No.  If 'Yes' is selected, the Information Governance Management Review section and Information Governance Team Review section become visible.
<b>Responsible Service</b>	Incident Service	Select the appropriate service if applicable (reporting form will complete the service field)
<b>Where did the Incident happen?</b>	Location of Incident	Select the appropriate location if applicable (reporting form will complete the location field)
<b>Communication</b>	Is this incident highly confidential (not for circulation)?	Please select Yes or No from the drop-down menu
	Who else needs access to this Incident?	Please select users from the drop-down menu who will need access to this incident.
	Who have you informed of the incident?	Please select who have you informed of the incident if applicable (reporting form will populate information)
<b>Openness and Transparency (Triggered for incidents graded Moderate and above).</b>	Was the patient/appropriate person informed that an incident occurred?	Yes/No drop down
	When was the Patient/Service User /appropriate person informed? (dd/mm/yyyy)	Enter date the discussion took place
	Were the members of staff involved in the incident involved in informing the Patient/Service User/appropriate person?	Yes/No drop down
	Please provide details of staff members who informed the Patient/Service	Enter details of staff who were involved

	User/appropriate person.	
	Please provide details of the Patient/Service User/appropriate person who was informed.	Enter details of the Patient/Service User/appropriate person who was informed
	Was an apology provided to the Patient/Service User/appropriate person?	Yes/No drop down
	Was a truthful account of the facts known at the time shared with the Patient/Service User/appropriate person?	Yes/No drop down
	Was the Patient/Service User/appropriate person advised about next investigative steps to be undertaken?	Yes/No drop down

## Incident Investigation/Managing an Incident

When an incident has been reported onto the system, it initially sits within the New Incident Status.

Reviewers must go into the new incident check the information is correct and complete any missing fields.

Before you make any changes, save the record in the Management Review status, the fields for the Management Review will now be visible for completion. You will be unable to save any changes you make if the record is left as a New Incident. If you do not enter the name of the Incident Manager, the system will input your name. The approval status should stay as 'Management Review/Make it safe plus until the Management Review has been completed.

**Current Approval Status**

Current approval status

Management review/Make it safe plus

**Management Review**

★ Investigator(s)  
Please add all the staff that will be reviewing or investigating this incident in this field

**Management Review**

★ Date Management Review started (dd/mm/yyyy)

★ Managers Interim Harm Assessment  
The All Wales Grading Framework is part of the PTR Regs. For a copy of the framework please click here

None

★ Following the initial review, has the grading changed

You are able and encouraged to add progress notes at this stage as you can add these without having to complete all mandatory fields.

It is the responsibility of the allocated manager to review the incident record and complete the management review section.

This involves checking the information included in the incident record, to ensure that it is accurate, complete, relevant, reliable and timely.

Section	Field	Description
<b>Management Review</b>	Investigator(s)	Enter the reviewer/investigator.
<b>Management Review</b>	Date management review started (dd/mm/yyyy)	Enter the date that the management review was started.
	Manager's Interim Assessment	Select from drop-down menu
	Following the initial review, has the grading changed	Yes/No drop down.
	Please explain why the grading has changed since the record was submitted	Enter reason for grading change.
	Was the healthcare provided a factor or may it have been a factor in a Patient/Service user suffering harm?	Please choose Yes or No from the drop-down options. Only visible for Moderate, Severe or Catastrophic graded incidents from Managers Interim Harm Assessment. If 'Yes' is selected in this field the 'Duty of Candour' section is visible for completion.
	Date NHS Body first became aware that DoC was triggered	Select date.
	Is this incident connected to the nursing care?	Yes/No drop down. If 'Yes' is selected the Nurse Staffing Act' panel is visible
	What were the findings of the management review?	Enter findings of the review.
	Are further immediate actions required at this time?	Enter all further actions required to prevent recurrence.
	Is there any early learning identified for sharing internally and externally?	
	Staff Absence	Visible if person affected is type 'Staff/Contractor. If there was a staff absence, further date fields are visible to capture the start and end date of absence.
	Does this require a focused review?	Selected the appropriate radio button (Focused Review appears in the Side Panel). If yes, please ensure you complete the focused review e.g., Inpatient Falls, Pressure Ulcer, Sharps, Manual Handling.
	Management review completed by	Select the appropriate user from the list
	Date Management review completed	Enter date the management review was completed
	Further investigations following management review?	Select close or proportionate investigation required.
Are you ready to complete the Investigation?	Yes/No drop-down menu. Select yes to trigger Proportionate Investigation Following the Management Review section.	

If following Management Review, it is felt the actions undertaken are sufficient to close the record, select 'Management Actions Sufficient – Close' from the option available. The Yorkshire Contributory Classification Framework\* and Conclusion section will appear and **must** be completed.

Further investigations following management review?

Management Actions Sufficient-Close



### Yorkshire Contributory Factors Framework

#### Domain 1: Situational Factors

★ Team Factors: Was there any failure or team function?

For example; Conflicting Team Goals, Poor Delegation, Lack of respect for colleagues, Absence of feedback

- Yes  
 No  
 Maybe

★ Individual Staff Factors: Were there any reasons this incident was more likely to occur with the particular staff involved?

For example; fatigue, stress, rushed, distraction, inexperience

- Yes  
 No  
 Maybe

★ Task characteristics: Did the task features make the incident more likely?

For example; unfamiliar task, monotonous task, difficult task

- Yes

Complete all required fields. You will also be required to complete the below sections:

Section	Field	Description
Conclusion	Is this incident related to the five harms of Covid 19?	Select from drop-down menu.
	Conclusion.	Enter conclusion following review/investigation.
	Post Investigation Harm Assessment	Enter the harm of the Incident <b>following the</b> investigation. This should be the severity of harm caused by the organisation. Help text is provided in a link on the field. If Moderate, Severe, Catastrophic / Death is selected you will need to answer a further Duty of Candour question if not already completed.
	Result	Enter the outcome of the incident following investigation.
	Recommendations	Enter recommendations.
	Lessons learned.	Enter any lessons for learning.
	Feedback to incident reporter.	Enter narrative which will be sent to the reporter.
	Date review/investigation completed.	Enter date review completed dd/mm/yyyy.
	Is this death confirmed as a suicide by the coroner?	Please select Yes or No as applicable. Only visible when Catastrophic / Death is selected in Post Investigation Harm Assessment.

\* Acknowledgement to the Yorkshire and Humber Improvement Academy.

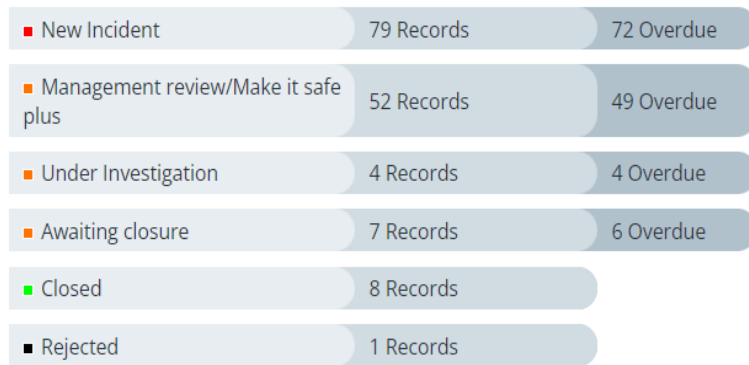
If following Management Review/Make it Safe Plus, it is identified a proportionate investigation is required, the form will ask the user to select one of proportionate investigation options.

\* Further investigations following management review?

- Management Actions Sufficient-Close
- Proportionate Investigation Required

When you are ready to complete a Proportionate Investigation, you can access this by clicking on the status area below and selecting the appropriate record.

**Statuses**



You will then need to click on the Incident Investigation Side Panel.

- Incident Details
- People Involved
- Incident Investigation**
  - Corporate Review
  - External Reporting
  - Progress notes
  - Actions

Section	Field	Description
<b>Proportionate Investigation Following the Management Review</b>	Date review started	Enter the date that the investigation was started
	Narrative of Patient/Person Affected Journey/Timeline/Key Issues	Enter details of the Patient/Service User journey
	Incident consequences	Enter incident consequences
	Narrative of Findings	Enter findings
	Responsible Monitoring Committee	Enter responsible monitoring committee
	Just Culture Tool considered	Select from drop-down box
	Following the Proportional Investigation are any further investigations required?	Select from drop down box
	What is the chance of this incident re-occurring	Drop down of matrix Consequence Likelihood Grade
	Responsible Directorate/Unit	Enter responsible directorate
	Further Duty of Candour Discussion Undertaken	Select Yes or No from drop down box

If the incident is ready to close following the Proportionate Investigation, the Yorkshire Contributory Classification Framework and Conclusion will appear for completion, as noted above.

## Duty of Candour

The Duty of Candour panel is triggered when Moderate, Severe or Catastrophic/Death is selected in the field 'Manager's Interim Harm Assessment' and 'Yes' is selected in the field 'Was the healthcare provided a factor or may it have been a factor in a Patient/Service user suffering harm?'

<b>Management Review</b>	Managers Interim Harm Assessment	Enter the potential harm of the incident from the drop-down menu
	Following the initial review, has the grading changed	Please select Yes or No from the drop-down menu
	Please explain why the grading has changed since the record was submitted.	Please enter a reason why the grading has changed since the incident was submitted
	Was the healthcare provided a factor or may have been a factor in a Patient/Service User	Please select Yes or No as appropriate. If Yes is chosen the following narrative box is visible
	Please explain why the grading has changed since the record was submitted	Complete narrative box with full explanation to cover the reason for change.

You can access the Duty of Candour panel by clicking onto this on the left-hand side.



It is the responsibility of the allocated manager to review the incident record and complete the Duty of Candour section. This involves checking the information included in the incident record, to ensure that it is accurate, complete, relevant, reliable, and timely.

Section	Field	Description
<b>Duty of Candour (DoC)</b>	Who is the Duty of Candour Point of Contact for this case?	Select the Duty of Candour Point of contact from the list of staff available
	Brief description of the circumstances in which the duty came into effect	Free text box – Please type the brief description of the circumstances in which the duty came into effect
	Has the Duty of Candour Point of Contact for this case made initial 'in person' contact with the Service User or the person acting on their behalf?	Please select from the drop-down options available  No / Unable to make in person notification / Unable to obtain contact details. All 3 codes trigger text box 'Please provide further details on attempts made to contact the service user or the person acting on their behalf'.
	Please provide further details on attempts made to contact the service user or person acting on their behalf or the reasons why a decision was made not to make contact	Enter the details of further attempts made to contact the service user or person acting on their behalf.
<b>Duty of Candour - Initial Notification</b>	Date of 'in person' initial notification (dd/mm/yyyy)	Enter the date of 'in person' initial notification
	Was the 'in person notification' made within 30 working days of the Date the NHS Body first became aware that DoC was triggered	Please select Yes or No as appropriate
	Please explain why the 'in person notification' was made after 30 working days of the Date the NHS Body first became aware that DoC was triggered	Text field to be completed if visible giving reason why 'in person' notification was made after 30 working days of the Date the NHS Body first became aware that DoC was triggered

	Method of initial notification	Please select the method of how the initial notification was conveyed to the Service user or the person acting on their behalf.  If 'Other Method' is selected, the below question appears.
	Other method (further details)	Free text box – Please describe how the initial notification was conveyed if this option has been selected
	Following the 'in person' initial notification, has written correspondence been sent to the Service User or person acting on their behalf	Please select Yes/No/Patient did not want to receive written correspondence' from the drop down  If 'Yes' is selected, the question 'Was the written correspondence sent to the service user or the person acting on their behalf in Welsh?' appears  If 'No' or "Patient did not want to receive written correspondence" is selected, the question 'Please explain why written correspondence has not been sent' appears  If 'Yes' is selected, Duty of Candour - Investigation Response section also appears.  If 'Yes' is selected 'Date written notification sent' is visible
	Was the written correspondence sent to the service user or the person acting on their behalf in Welsh?	Please select 'Yes' or 'No' from the drop down
	Date written notification sent (dd/mm/yyyy)	Enter Date written notification sent
	Please explain why written correspondence has not been sent	Please enter further details explaining why written correspondence has not been sent
	Was the 'written notification' sent within 5 working days following the in-person contact being made	Please select Yes or No as appropriate
	Please explain why the 'written notification' was sent after 5 working days following the in-person contact being made	Text field to be completed if visible giving reason why the 'written notification' was sent after 5 working days following the in-person contact being made
Duty of Candour - Investigation response	Has the investigation response been sent to the Service User or person acting on their behalf?	Please select 'Yes' or 'No' from the drop down  If 'Yes' is selected, a date field will be visible to capture the date and another field 'Type of Response' will also be visible  If, 'No' is selected, a text field is visible to capture explanation of why final response has not been sent.
	Final response done date (dd/mm/yyyy)	Enter Date Final response sent
	Please explain, why no final response has been sent	Please enter an explanation of why a final response has not been sent
	Type of response	Please select the type of response sent to the service user of the person acting on their behalf.

Time Chain Welsh Duty of Candour	The time chain is not a mandatory section; however, this could be used to support calculation of the essential criteria regarding the in-person contact being made within 30 working days of the organisation becoming aware that the Duty was triggered, and 5 working days to complete the written notification. This will only be triggered if the question: Was Healthcare provided a factor or may it have been a factor in a patient/service user suffering harm and answered Yes.
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## Radiology Section

This section is currently being reviewed by All Wales Imaging forum and this section will be updated when fully reviewed.

## Information Governance Management Review Section

The Information Governance section is triggered when the reporter selects **YES** to the question “Does this Incident have Information Governance considerations?”. The Information Governance Management Review Section is then visible in the left hand panel

The screenshot shows a web application interface with a top navigation bar (Capture, Evaluate, Strategy, Implement, Assess) and a right-hand button for COVID-19 Resources. A left-hand navigation menu is visible, with 'Information Governance Management Review' highlighted in a red box and a red circle with the number '1' next to it. The main content area shows a form for 'Additional Information' with several questions and dropdown menus, including 'Is there any factor relating to Emergency Planning for this incident?', 'Does this Incident have Information Governance considerations?', 'Does this incident have any safeguarding elements?', and 'Was any equipment involved in the incident?'.

Section	Field	Description
Information Governance Management Review	Has personal data been disclosed outside of the organization?	None/Yes/No/Don't know radio buttons
	What type of personal data is included?	Select all that apply from drop down menu
	Has there been (or is there likely to be) any impact to the individuals as a result of the incident?	Please select the relevant impact. This is a multi-pick option
	Who does this incident relate to?	Please select who the incident relates to. This is a multi-pick option. If 'other' is selected please see box below
	Who does this incident relate to? Other, please specify	Please type who the incident relates to
	Where has the data been disclosed accidentally or deliberately?	Please select where the data has been disclosed to. This is a multi-pick option. If 'other' is selected please see box below
	Where has the data been disclosed accidentally or deliberately? Other, please specify	Please type where the data has been disclosed to
How many personal data records are concerned?	Please type the number of personal data records that are concerned with the incident	

	How many individuals are affected by this incident?	Please type the number of individuals that are affected by this incident
	Is the recipient aware of the breach?	Yes/No/Don't know drop down
	Is the data subject(s) aware of the breach?	Yes/No/Don't know drop down
	Is it likely that the recipient knows the data subject(s)?	Yes/No/Don't know drop down
	Has a complaint been received by the organisation?	Yes/No/Don't know drop down If yes is selected, an alert message pops up reminding the reviewer to link the complaint to the record
	Has the data been recovered?	Yes/No/Don't know drop down
	Has any inaccurate data been removed/updated/corrected to avoid further incidents?	Yes/No/Don't know drop down
	Has there been or is there likely to be press or media interest relating to the breach?	Yes/No/Don't know drop down If yes is selected, an alert message pops up reminding the reviewer to complete and Early Warning Notification for submission
	Has there been or is there likely to be social media activity relating to the breach?	Yes/No/Don't know drop down
	Have individuals directly involved with the incident completed their statutory and mandatory information governance training within the last 2 years?	Yes/No/Don't know drop down

**The information Governance Team Review section will only show to those users who have permitted access**

Section	Field	Description
Information Governance Team Initial Triage	IG Reviewed by	Please select from drop down menu
	Date IG reviewed	Please enter date IG was reviewed
	Confirmed IG issue	Yes/No option. If yes is selected, Information Governance Team Review section is visible.
Information Governance Team Review	What is the type of Information Governance Incident?	Please select what type of information governance the incident relates to
	What are the potential consequences of the incident?	Please type what the potential consequences of the incident are
	What preventative measures are in place?	Please type what preventative measures are in place
	Does this personal data breach require notification to the ICO?	None/Yes/No If yes is selected please see boxes below
	Date ICO notified	Please enter date ICO was notified of the IG breach

ICO reference number	Please type ICO reference number
Where notification exceeds the 72-hour timescale, please explain why?	Please type reasons for notifications exceeding 72-hour timescale
ICO Status	Open/Closed drop down
Recommendations/actions provided by ICO	Please type recommendations/actions provided by ICO
Record of correspondence with ICO	Please type any record of correspondence with ICO
Does this personal data breach require notification to the data subject(s)?	None/Yes/No If yes is selected please see boxes below
Date data subject notified	Please enter date that the data subject was notified
Record of correspondence with data subjects	Please type any record of correspondence with the data subjects
Summary of IG actions taken	Please type the summary of IG actions taken

## Corporate Review (left hand side panel)

The Corporate Review panel has been added for corporate teams (including Health & Safety, Patient Safety, Tissue Viability Nurses, RRails Review and other corporate specialist teams) to keep track of the corporate management of various incidents.

Corporate Review

Type of Corporate Specialist Review

Health and Safety

Other

Patient Safety

RRails Review

Specialist Palliative Care (SPC)

TVN Corporate Review

Section	Field	Description
<b>Corporate Specialist Reviews for:</b> <ul style="list-style-type: none"> <li>• <b>Health &amp; Safety</b></li> <li>• <b>Patient Safety</b></li> </ul>	Reviewer	Enter the reviewer
	Actions taken	Enter actions taken
	Who has been informed	Enter communication information
	Date Closed	Enter date the incident is closed
	Reviewer Notes	Enter any notes in relation to the management of the incident
<b>TVN Corporate Specialist Review</b>	TVN actions taken	Enter actions taken
	Date Closed	Enter date the incident is closed
<b>Other Corporate Specialist Review</b>	Other - actions taken	Enter actions taken
<b>RRAILS Review</b>	Resus Issue Confirmed	Select from drop down
	RRAILS Review: Reviewed by	Enter reviewer information
	RRAILS Review: Reviewed date	Enter review date (dd/mm/yyyy)
	Was there failure to follow the NEWS protocol	Select from drop down
<b>Specialist Palliative Care (SPC)</b>	Reviewed by (SPC)	Please select from drop-down menu
	Review Date (SPC)	Enter review date (dd/mm/yyyy)
	Does this incident relate to the Specialist Palliative Care Team?	Please select from drop-down menu

	SPC Incident relates to the following Themes	Please select from drop-down menu. This field can select multiple options.
	Feedback to Department	Please enter text
	Date closed (SPC)	Enter closed date (dd/mm/yyyy)
<b>The Datix Cymru LSL panel is read only by default</b>		
<b>Datix Cymru LSL</b>	Was a review undertaken within 72 hours?	Select from drop down
	Corrections made to records?	Select Yes or No. If yes is selected a further 3 fields are visible to capture the information changed and the member of staff making the changes
	What corrections were made to the record?	Select from drop-down menu.
	Supporting information on what corrections were made	Enter corrections made.
	Local System Lead making correction(s)	Select from drop-down menu.

## External Reporting (left hand side panel)

The External Reporting Panel has been added for corporate teams to keep track of the management of various incidents that require reporting to external bodies.

**External Reporting**

To whom are you externally reporting?

---

Counterfraud  
HIW  
HSE  
HTA  
Information Commissioner  
MHRA  
Natural Resource Wales (NRW)

Section	Field	Description
<b>External Reporting</b>	To whom are you externally reporting?	Multi pick selection: Counter fraud NHS Wales Executive HIW HSE HTA Information Commissioner MHRA MHRA Yellow Card Public Health Wales SABRE SHOT Welsh Government WHSSC Other Health Body Natural Resources Wales
	What is being externally reported?	Multi pick selection from the following: Adverse reaction Early Warning Notification Fatal Drug Poisoning HTA Breach IG Breach IRMER Breach National Reportable Incident RIDDOR Originated in other Health Body License Breach
	Is a joint review required?	Select 'Yes' or 'No' from the drop-down menu If 'Yes' is selected, the below question appears
	If yes, please specify which organisation(s)	Aneurin Bevan University Health Board Welsh Ambulance Service NHS Trust Other NHS Organisation Betsi Cadwaladr University Health Board Cardiff & Vale University Health Board Cwm Taf Morgannwg University Health Board Hywel Dda University Health Board Powys Teaching Health Board Public Health Wales Swansea Bay University Health Board Velindre University NHS Trust
	If yes, brief description of action taken in relation to joint investigation	Free text – Please type a brief description of action taken in relation to joint investigation
	Are you the lead Organisation for Duty of Candour?	Select 'Yes' or 'No' from the drop-down menu

Depending on what is being reported externally various extra questions trigger for completion.

## Case Management (left hand side panel)

The Case Management Panel is available as required for Case Managers to assist with the management of violence and aggression incidents in the NHS.

**Case Management**

★ Were security in attendance?  ▼

Is there Case Management Involvement  ▼

Case Manager  ▼

Immediate action taken by the Case Manager

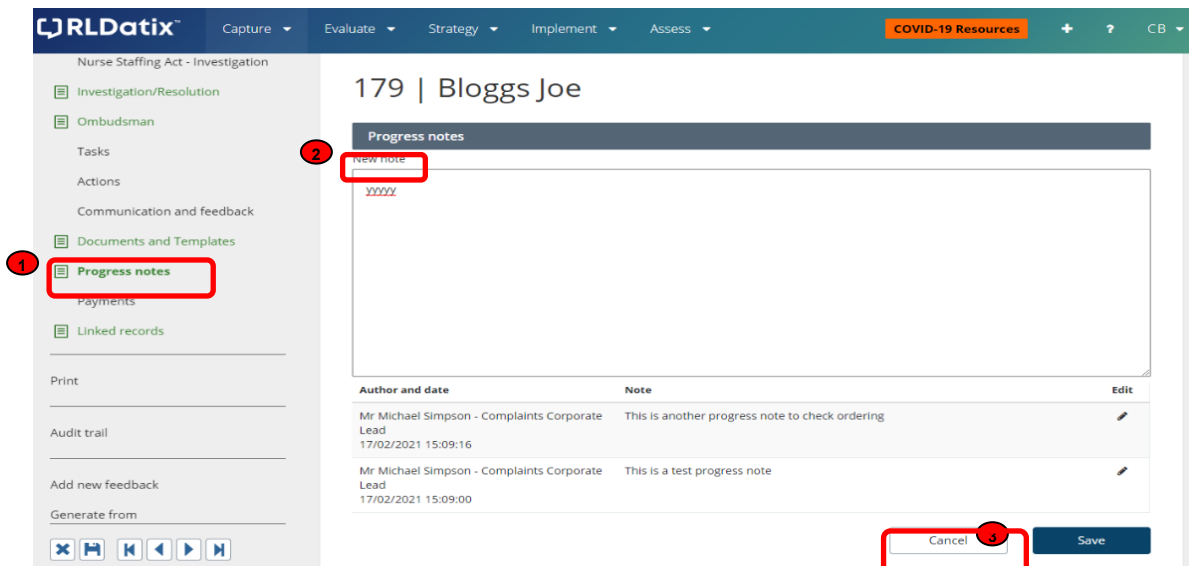
Section	Field	Description
<b>Case Management</b>	Were security in attendance	Select from drop down
	Is there Case Management Involvement	Select from drop down
	Case Manager	Select Case Manager from drop down
	Immediate action taken by the Case Manager	Enter any action taken from Case Manager
	Progress Notes (sec incident)	Enter progress notes confidential to Case Manager only
	Exact Category of Behaviour	Select from drop down
	Repeat offender	Select from drop down
	Was this a hate incident?	Select from drop down
	Type of hate crime?	Select from drop down
	If Annex G Completed, what was the outcome?	Select from drop down
	Was Restrictive Physical Intervention used during this incident?	Select from drop down
	Was NHS property damaged or stolen?	Select from drop down
	Contributory factors	Select from drop down
	Contact made with	This is a multi – coded field, please select from drop down options
	Feedback given to	Select from drop down
	Action Taken?	Enter action taken
	Were the Police in attendance?	Select from drop down
	Outcome	Select from drop down
	Status of case	Select from drop down
	Has a management plan been completed to assist in preventing recurrence?	Select from drop down

## Progress Notes (left hand side panel)

Progress notes are a way of adding notes to records that will be given a time stamp.

Users can be given rights to edit all progress notes or just their own by their administrator.

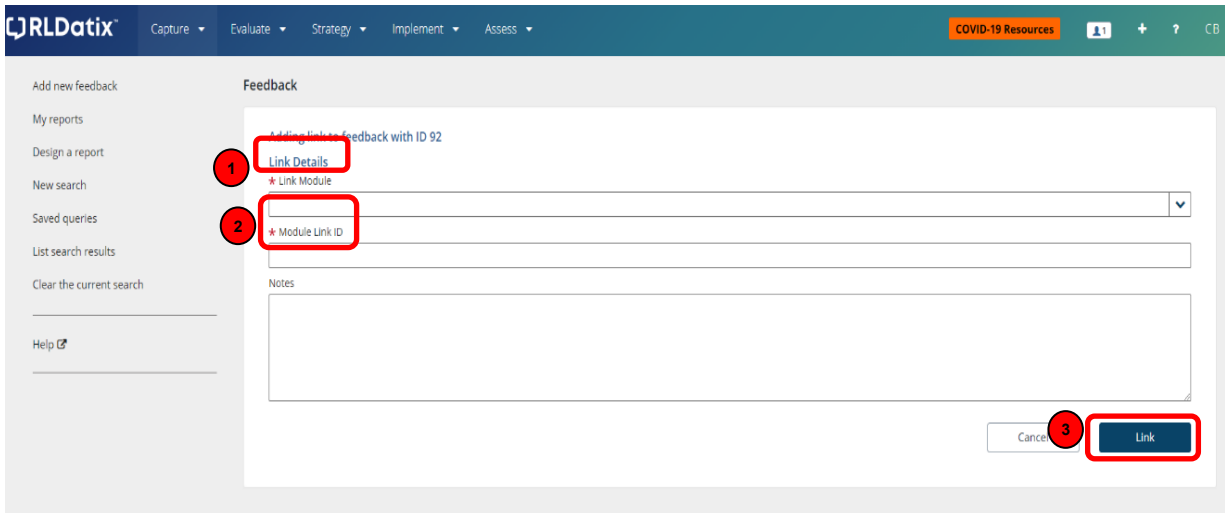
1. Click Progress notes in the left-hand navigation menu.
2. Type into the Notepad field
3. Click Save.



## Linked Records (left hand side panel)

This gives you the ability to link records together, e.g. where a feedback record is linked to an incident. Make a note of the Function (Module) and Record ID number you want to link to the record.

1. From the Link Module drop-down list, select the module.
2. In the Module Link ID enter the Record ID to link to this feedback record.
3. Click Link.
4. Click Save

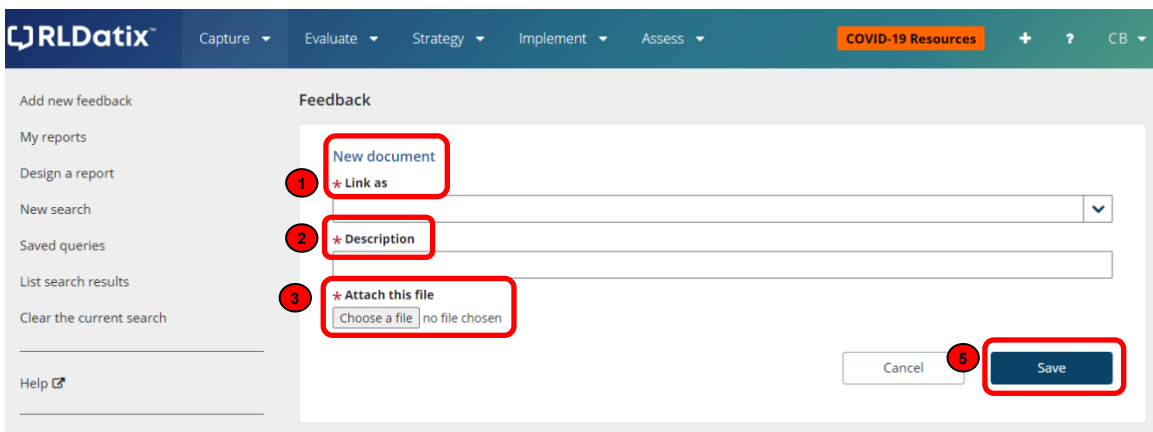


## Documents and Templates (left hand side panel)

To add a new document Click Attach a new document

1. Select a format from the Link as drop-down list.
2. Type a description of the document in the Description field.
3. Click Browse from the Attach this file field and navigate to the file on your computer.
4. Select the file and click Open.
5. Click Save

6.



To add a new template

1. Select the document template you want to merge from the Word Document drop-down list.
2. Click Merge into a template. The relevant field values from the feedback record are merged into the document template.

## Notifications, Communication and Feedback (left hand side panel)

The Notification, Communication and feedback section is used to send emails directly from Datix Cymru.

When you send an email from Datix Cymru, the system logs the email directly in the record as message history. Responses to the email will be sent to the sender's usual email account and will need to be added to the record as a document.

In addition to sending an email, you can include a document that has already been saved to the record.

To do this, please select a document (or multiple documents if necessary) from the drop-down list in the '**Communication & Feedback Document Attachment**' of the Communication and Feedback section, amend the title and body of the email message as appropriate and click send.

The screenshot shows the 'Communication and feedback' interface. It includes a 'Recipients' section with a dropdown menu set to 'All users' and an 'Additional recipients' field. The 'Message' section has a 'Subject' field with the text 'Datix Cloud IQ feedback message' and a 'Body of Message' field with a rich text editor containing the text: 'This is a feedback message from Maria Stolzenberg. Incident form reference is HDD13 The feedback is: Please go to <https://hyweldda.private.prod-uk.datixcloudiq.co.uk/capture?action=incident&recordid=13> to view the incident.' Below the message field is a 'Send message' button. At the bottom, there is a 'Message history' table with columns for Date/Time, Sender, Recipient, Body of Message, and Attachments.

If you select notifications, it will show you a list of staff that were notified at the time the Incident was reported.

## Actions (left hand side panel)

The Actions module will need to be completed where any issues/failures have been identified. All actions must be added to the record separately. Example of two actions have been identified

- The need to update a policy
- The need to deliver training

You will need to add these actions in one at a time and ensure you assign the correct staff member responsible for managing the action through to the completion against the record. You can do this by:

1. Click Create a new action.
2. Complete all the required fields.
3. In the Assigned To section, click Search.

4. In the Forename and Surname fields, enter a name.
5. Click Search.
6. Select a contact. You can add multiple contacts.
7. Click Save Action.

## Final Check list before closing down the Incident Record

- All the field have been completed accurately
- All relevant panels have been completed and reflects the outcome of the Incident
- All actions that have been identified from the Investigation have been added to the record and closed if possible
- All relevant documentation has been uploaded.
- Any Unapproved Contacts highlighted in RED will need to be approved or updated
- All staff involved will need to be added as a contact, including the completion of their role they undertook within the incident, i.e., witness, provided comments, etc.

## Basic Search and Reporting

When performing a search, you will be shown a copy of your incident form. You can select any values from a field in order to search for them. For example, if you wanted to see a specific record and you had the ID number you could enter this into the ID field and Click on Search. If it is relating to a particular type and service, you would simply select that Incident Classification and the service in the drop down and Click Search.

Below are some handy hints for searching:

Symbol	Explanation
*	Wildcard searches are particularly useful for searching within text boxes. For example, to show all complaints where the word "Needle" appears in the description, you would search for*needle*. This would locate the word where any text can come before it, or after it.
=	Searches for all records where a particular field has no value recorded in it. For example, an = sign in the "Closed Date" field would return all records where there is no recorded closed date.
==	Searches for all records where a particular field has a value recorded in it, regardless of what it is. For example, == in the "Closed Date" field would return all records where a closed date has been recorded.

For additional information on features such as reporting and other functionality common across the Capture module, a separate guide will be produced. Alternatively, contact your Local System Lead in the Health Board / Trust.

## Appendix One

### Medication Incidents

You can record incidents that involve medication in two ways. Either via incident Classification

Incident Type
* <b>Who was affected?</b>
Patient/Service User
* <b>Classification</b>
Medication, IV Fluids
* <b>Category</b>
Medication prescribing error
* <b>Sub Category</b>
Duplication of medication

or by indicating medication was involved.

Additional Information
* <b>Were there any medications involved?</b>
Yes

When medication has been indicated in the incident a medication form triggers on the reporting form (overleaf).

Administered	Intended / Suspected
<input type="button" value="Clear"/>	<input type="button" value="Clear"/>
<input type="button" value="Duplicate"/>	
*Search for drug administered or omitted	*Search for intended / suspected drug
<input type="text"/>	<input type="text"/>
Brand name of drug administered	Brand name of drug intended / suspected
<input type="text"/>	<input type="text"/>
Manufacturer of drug administered	Manufacturer of drug intended / suspected
<input type="text"/>	<input type="text"/>
Class of drug administered	Class of drug intended / suspected
<input type="text"/>	<input type="text"/>
Type of drug administered, prescribed, dispensed or omitted	Type of drug intended / suspected
<input type="text"/>	<input type="text"/>
Route administered, prescribed, dispensed or omitted	Route intended / suspected
<input type="text"/>	<input type="text"/>
Dose administered, prescribed, dispensed or omitted	Dose intended / suspected
<input type="text"/>	<input type="text"/>
Form administered, prescribed, dispensed or omitted	Form intended / suspected
<input type="text"/>	<input type="text"/>
Drug administered or omitted BNF classification (1-15)	Drug intended BNF classification (1-15)
<input type="text"/>	<input type="text"/>
Drug administered or omitted batch number	Intended drug batch number
<input type="text"/>	<input type="text"/>
Drug administered or omitted ethics committee name and reference	Drug intended ethics committee name and reference
<input type="text"/>	<input type="text"/>
Is the drug administered or omitted under clinical trial?	Is the drug intended under clinical trial?
<input type="text"/>	<input type="text"/>
Is the drug administered or omitted a manufactured special?	Is the drug intended a manufactured special?
<input type="text"/>	<input type="text"/>
Is the drug administered or omitted a parallel import?	Is the drug intended a parallel import?
<input type="text"/>	<input type="text"/>
Stage at which error occurred	
<input type="text"/>	
Type of error	
<input type="text"/>	

You can search for the administered medication and the intended. You should complete all required fields.

If you are unable to find the medication you require to report, please complete the section below the medication table:

✦ Other Important Factors



### New Medication

Please add medication below, if you are unable to find name in medication section

A radio button allows the Reporter to indicate if the medication involved is a controlled drug

## Equipment Incidents

You can record incidents that involve equipment in two ways. Either via incident Classification or by indicating equipment was involved.

### Incident Type

✦ Who was affected?

Patient/Service User

✦ Classification

Equipment, Devices

✦ Category

Medical devices

✦ Sub Category

Medical device user error

### Additional Information

✦ Was any equipment involved in the incident?

When equipment has been indicated in the incident, an equipment section is visible. You should complete all required fields.

### Equipment

Type of product



Manufacturer name

Serial Number

Equipment ID

Description and location of the defect/problem

## Focused Review Incidents

Some Incidents may require a focused review as part of the proportionate investigation.

Within the Management Review section there is a question asking if a Focused Review is required (select Yes).

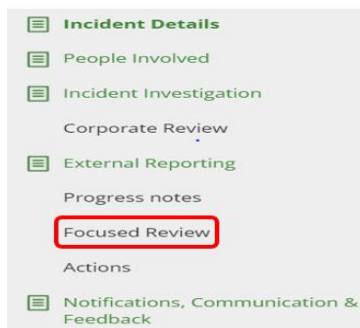
✳ Does this require a focused review?

If yes, please ensure you complete the focused review panel  
e.g Falls, Pressure Ulcer, Sharps, Manual Handling

Yes

No

This will then trigger the Focused Review option in the side panel and a selection of Focused Reviews will appear.



The question 'What type of Focused Review is this?' is now visible in this panel. Please select the type of review to be conducted.

A screenshot of a form titled 'Focused Review'. Below the title is a question: '✳ What type of Focused Review is this?'. Below the question is a dropdown menu. The dropdown menu is open, showing a list of options: 'Extravasation', 'In Patient Falls', 'Manual Handling', 'Pressure Damage', and 'Sharps'. The dropdown menu has a light blue background and a thin border.

Please select the type of review that is to be conducted and the appropriate Investigation tool will be triggered for completion. You should complete all the required fields.

## In Patient Falls Incidents

You can record incidents that involve falls under the incident Category on the reporting form. The Falls Investigation Tool is to be completed as part of the proportionate investigation. You should complete all the required fields, the section is visible following selection at Focused Review

**Falls**

Was there anything particular about the time of day that might have contributed to this fall?

What were the available staff on the ward doing at the time of the fall / where were they?

## Pressure Damage Incidents

You can record incidents that involve pressure damage under the incident Classification on the reporting form this includes device related (d) and non-device related pressure damage.

**Incident Type**

\* Who was affected?

Patient/Service User ▼

\* Incident Type

Pressure Damage, Moisture Damage ▼

\* Sub Type

Pressure ulcer category 1| ▼

- Device-related pressure ulcer category 3 (d)
- Device-related pressure ulcer category 4 (d)
- Device-related suspected deep tissue injury (d)
- Device-related unstageable pressure ulcer (d)
- Moisture-associated skin damage (MASD) (incontinence-associated)
- Moisture-associated skin damage (MASD) (not incontinence-associated)
- Pressure ulcer category 1


Please complete all required fields in the Pressure Damage tool as part of the proportionate investigation, following selection at Focused Review.

**Pressure Damage**

**Details of the pressure damage you are investigating**

\* How long has the individual been in this care environment

\* Reason why the individual is receiving care? 

## Manual Handling Incidents

You can record incidents that involve manual handling under the incident category on the reporting form.

**Incident Type**

\* **Who was affected?**

\* **Classification**

\* **Category**

\* **Sub Category**

Please complete all required fields in the Manual Handling tool as part of the proportionate investigation, following selection at Focused Review.

**Manual Handling**

\* Description of Activity (e.g. what was being done, method/technique used, how many staff involved etc.)

\* Load / Patient Factors (e.g. weight, size, shape, contents, compliance etc.)

## Sharps Incidents

You can record incidents that involve sharps under the incident category on the reporting form.

Incident Type	
* Who was affected?	Staff/Contractor
* Classification	Accident, Injury
* Category	Contact with needles or medical sharps
* Sub Category	During disposal - No safety fitting on sharp contaminated/used

Please complete all required fields in the Sharps Incident tool as part of the proportionate investigation, following selection at Focused Review.

Sharps	
1. What was the risk from the sharps injury?	
(*) Where a patient or the source of blood or body fluids is known this will be reportable to the Health and Safety Executive, contact the H&S team and Occupational Health for advise VIRUS	
<input type="checkbox"/> Clean medical sharp (no blood, body fluid or drugs/chemical contamination)	
<input type="checkbox"/> Contaminated with body fluids (patient does not have a blood bourne virus)	
<input type="checkbox"/> Contaminated with drugs/chemical	
<input type="checkbox"/> Patient has or suspected to have a blood Bourne virus (*)	
<input type="checkbox"/> Were bloods taken from the staff member and patient?	
2. What type of medical sharp was involved?	
Describe device type	

# Extravasation

You can record incidents that involve Extravasation under the incident category on the reporting form.

Incident Type	
* Who was affected?	Patient/Service User
* Classification	Treatment, Procedure
* Category	Treatment or procedure issues
* Sub Category	Extravasation (infusion injury)

Please complete all required fields in the Extravasation tool as part of the proportionate investigation, following selection at Focused Review

Extravasation	
What type of extravasation is this?	
Size of Extravasation	
Is this a late presentation of Extravasation?	
What Extravasation symptoms are present? If you have a photograph of the Extravasation site, please upload in the document panel	
Does this meet the criteria for a plastic surgeon? As a general guideline, advice should be obtained from a plastic surgeon for: Extravasation of vesicant involving an area greater than 2cm	
Name of drug causing Extravasation Vesicants are designated with (V)	
Type of IV access?	
Drug used to treat the Extravasation?	

What treatment was applied to the Extravasation?

Any other information regarding the Extravasation and its treatment?

Was the IV flushed before use?

Rate of Injection

Estimated volume of extravasation

Was a peripheral vascular/intravenous cannula care bundle in place and completed to establish if checks have been undertaken?

Was an infusion device in use at the time of the extravasation?

Was the extravasation avoidable?

## Focused Review Additional Information

Please complete all required fields in the focused review additional information as part of the proportionate investigation, following selection of the appropriate Focused Review

### Focused Review Additional Information

Name and designation of Focused Review investigator

Name and Designation of Senior Person approving the Focused Review investigation & outcome decision

Date focused review investigation completed



## Appendix Two – Additional Information

### National Once for Wales Code Set

The current version of the code set is held in the Coding Workstream shared folder in the Once For Wales Resource Library. This is accessible via your Local System Lead in the Health Board / Trust.

### Dictionary of Medications and Devices

Datix Cymru utilises a database table of medications for all users. This is known as the Medications List.

RLDatix obtain this information automatically from the Dictionary of Medications & Devices (DM&D) list provided by the NHS Business Services Authority.

Following upload of the DM&D list, the NHS Wales Central Team will be required to identify which medications need to be visible within NHS Wales. It is currently the intention of the central team to initially include all available medications and will liaise with the Chief Pharmaceutical Officer and the All-Wales Medicines Management Network to determine which medications are appropriate to be hidden.

The NHS Wales Central Team will provide a full summary of the DM&D list once it has been successful imported and highlight the additions and amendments.