

WORK INSTRUCTION

WI 01 – How to Enter a Safety Report onto Q-Pulse

Version	3
Effective Date	28/09/2022
Review Date	28/09/2025
Author & Position	Nicola Jones, Quality Assurance Coordinator
Signature	N.Jones
Date	11/10/2022
Approver & Position	Samantha Smart, Research Governance and Quality Manager
Signature	Shot
Date	11/10/2022

Controlled document

This document has been created following the Royal Devon University Healthcare NHS Foundation Trust Policy for the Development, Ratification & Management of Procedural Documents. It should not be altered in any way without the express permission of the author or their representative.

It is the responsibility of all users of this Work Instruction to ensure that the correct version is being used. If you are reading this in a paper format please go <u>on-line</u> to confirm you have the latest version.

DISCLAIMER

This generic R&D Work Instruction (WI) must be followed unless a study specific SOP/WI exists.

Once printed this is an uncontrolled document



Full History			
Version	Date	Author	Reason
V1.0	22 November 2012		
V1.1	14 November 2014	Lisa Treeby & Rhianne Lewis, R&D Facilitators and Joanne Lowe, Quality Assurance Coordinator	Minor clarifications to the process. Transferred content to new template.
V1.2	03 November 2015	Lisella Wilkinson, QA Coordinator	Minor additions to process to accommodate metrics required and report designing.
V2.0		Nikki Sawyer, Quality Assurance Coordinator and Elizabeth Watson R&D Facilitator	Minor additions to the process and new information on updating records. Transferred content to new template.
V3	13 July 2022	Quality Assurance Coordinator	Updated into current WI template and to reflect change in saving process.

Associated Trust Policies/ Procedural documents:	<u>S22 Safety Reporting</u>
Key Words:	R&D Work Instruction AE/AR/SAE/SAR/SUSAR Safety Reporting Q Pulse
In consultation with: Quality Assurance Group (June 2019) Directorate Governance Group Quality Assurance Group (July 2022)	



Contents

1	INTRODUCTION	4
2.	PURPOSE	4
3.	SCOPE	4
4.	DEFINITIONS & ABBREVIATIONS	4
5.	DUTIES AND RESPONSIBILITIES OF STAFF	4
6.	PROCEDURES	5
6.1	Guidance on Entering a Safety Report onto Q-Pulse	5
6.2	Updating Records on Q-Pulse	6
6.3	Useful Information when entering Safety reports on Q Pulse	6
7.	DISSEMINATION AND TRAINING	7
8.	MONITORING COMPLIANCE AND EFFECTIVENESS OF THIS WI	7
9.	ARCHIVING ARRANGEMENTS	7
10.	REFERENCES	7



1 INTRODUCTION

All safety reports which are reported to Research & Development (R&D) should be recorded on R&D's quality management system, Q-Pulse, and reviewed and followed-up appropriately. Safety reporting systems are subject to audit by the Sponsor/Host institution or inspection by the Medicines and Healthcare Products Regulatory Agency.

2. PURPOSE

The purpose of this Work Instruction (WI) is to describe the procedure for recording safety report forms on Q-Pulse.

3. SCOPE

This WI applies to the Quality Assurance (QA) Coordinator and, in their absence, the R&D Facilitators. It also applies to the Assistant R&D Manager in terms of oversight of the procedure.

4. **DEFINITIONS & ABBREVIATIONS**

CI	Chief Investigator
GOG	R&D Governance Oversight Group
PI	Principal Investigator
MHRA	Medicines and healthcare Products Regulatory Agency
SOP	Standard Operating Procedure
R&D	Research and Development
Sponsor	An individual, company, institution or organisation which takes responsibility for the initiation, management and financing of a clinical trial. Sponsorship activities may be delegated to the Investigator, CTU and/ or other organisations as appropriate
WI	Work Instruction
DTX	Datix Incident Report Number
CTIMP	Clinical Trial of an Investigational Medicinal Product
SAE	Serious Adverse Event

5. DUTIES AND RESPONSIBILITIES OF STAFF

QA Coordinator

It is the responsibility of the QA Coordinator to enter all safety reports received by R&D onto Q-Pulse in a timely manner. In the absence of the QA Coordinator, the R&D Facilitators will assume this responsibility.

Assistant R&D Manager

It is the responsibility of the Assistant R&D Manager to oversee this procedure and to review all safety reports which are received and entered onto Q-Pulse. In the absence of the Assistant R&D Manager, the Research Management and Governance Manger will assume this responsibility.



6. **PROCEDURES**

6.1 Guidance on Entering a Safety Report onto Q-Pulse

NB First check if Causality has been ticked; if not, you can't put the safety report on Q-Pulse.

Step 1	Open Q-Pulse and in the 'CAPA' module:
	Open 'New'
	Click 'From wizard'
	 Click 'Safety report'
Step 2	You should now be on the 'Initial Details' page.
	 Details – Enter in a line the title of the study followed by the patient ID, then the summary of the event detailed on the safety report form, divided by a few spaces. R&D Number – Enter the R&D number, which should be located on
	the safety report (can also be searched for on EDGE)
	 Sponsored or hosted coloct
	 Sponsored of hosted – select Click 'Next'
Step 3	This will take you to the 'Additional Details' page.
	• Causality – select the causality from the drop-down-list, this can be
	identified from the safety report.
	NB: If it's a registry study causality will always be not related (there is
	no intervention on the protocol).
	• Source – Select 'Datix' from the drop-down-list and expand the list.
	Select which type of report it is, this can be found on the safety report.
	 Expected or Not – Select whether the event/reaction was expected or unexpected, this can be found on the safety report.
	NB : Can be skipped over if not required (for Not Related or Unlikely).
	If causality is possibly, probably, or definitely related expectedness
	should always be ticked. Contact the person who submitted the form
	if not.
	Click 'Next'
Step 4	This will take you to the 'Date incident occurred' page.
	 Incident Date – This is the 'date of onset' on the safety report.
	• Target Date – This is the 'date reported to R&D' on the safety report.
	NB: Remember to check the date as the month automatically comes
	up as the following month.
	Raised Date – Select the date you are entering the safety report onto
	Q-Pulse.
	 Investigator Aware Date – next to date of onset on the safety report.
	Click 'Next'
Step 5	The Safety Report Wizard is now complete
Sich 2	Tick 'After Finish _ Display Details'



	 Click 'Finish' The safety report details will be displayed. Enter the 'DTX' number on the top left of the safety report form before saving in the R&D study folder and send a copy to the Assistant R&D Manager for review.
--	--

6.2 Updating Records on Q-Pulse

Open records on Q-Pulse

To search for a record, click on the CA/PA module on the LaunchPad Q-Pulse screen. This brings up a Filter screen. Enter the study name, a specific search word/number or the DTX number in the keyword search box.

Select the required AE/SAE by double clicking on it. It will open in a new screen.

Within 'Details' Box add the update. Type in 'Update received' and the date. Then add the extra information from the updated safety report. Save what has been entered before exiting the screen.

Enter the DTX number of the initial report and 'follow up' or 'final report' onto the top left corner of the form before saving and forward a copy to the Assistant R&D Manager to review.

Updates to closed records on Q-Pulse

Closed report will need re-opening. Select the required AE/SAE and open in a new screen, click on 'Actions', click 'Re-Open Record'. You will be prompted to enter a reason, write 'Update received'.

Within 'Details' Box add the update. Type in 'Update received' and the date. Then add the extra information from the updated safety report. Close the report with a final comment eg 'Issue resolved'. Save what has been entered before exiting the screen.

Enter the DTX number of the initial report and 'follow up' or 'final report' onto the top left corner of the form before saving and forward a copy to the Assistant R&D Manager to review.

6.3 Useful Information when entering Safety Reports on Q Pulse

Study specific safety report forms agreed by the Sponsor can be used to notify R&D in place of the R&D template forms.

If the CI/PI hasn't signed the safety report and isn't available to sign it the sub or coinvestigator should sign instead (if delegated to do so).

If the safety report is for a CTIMP and the CI/PI has not signed, check if it is a first report. If it is a first report the CI/PI needs to sign. If it is a follow up report and if no changes have been made to causality, expectedness or severity then no CI/PI



signature is needed. The person submitting the report does however need to evidence that the CI/PI has seen and considered the updated report, this can be done by copying them into the email to R&D.

7. DISSEMINATION AND TRAINING

- 7.1 This WI and associated templates and forms will be uploaded to the <u>RDE Research</u> <u>website</u> shortly after having been released.
- 7.2 All staff whose activities are subject to this WI should ensure that they take time to read and understand the content of this WI.

8. MONITORING COMPLIANCE AND EFFECTIVENESS OF THIS WI

8.1 In order to monitor compliance with this WI, the auditable standards will be monitored as follows:

No	Minimum Requirements	Evidenced by
1.	Enter the 'DTX' number on the top left of the	Review forms saved on
	safety report form before saving in the R&D	Research_Develop\$ shared
	study folder.	drive, study folders.
2.	Reports should be sent to the Assistant R&D Manager for review.	Assistant R&D Manager sign off on Q-Pulse

- 8.2 Outcomes from audit will be presented to the R&D Governance Oversight Group (GOG) which will monitor any resulting action plans until all issues have been addressed to satisfaction.
- 8.3 Issues identified via the audit process which require escalation will be referred to GOG.

9. ARCHIVING ARRANGEMENTS

- 9.1 The original of this document will remain with the R&D Quality Assurance Coordinator. An electronic copy will be maintained on the R&D section of the Q-Pulse document management system and a pdf copy on the <u>RDE Research</u> website.
- 9.2 Archive copies must be maintained for any documents which have been superseded. Archive copies in electronic format should be retained indefinitely.

10. REFERENCES

RDE Research Website: https://rderesearch.co.uk/home/