GUIDELINES



EANM guideline on quality risk management for radiopharmaceuticals

Nic Gillings¹ · Olaug Hjelstuen² · Martin Behe³ · Clemens Decristoforo⁴ · Philip H. Elsinga⁵ · Valentina Ferrari⁶ · Oliver C. Kiss⁷ · Petra Kolenc⁸ · Jacek Koziorowski⁹ · Peter Laverman¹⁰ · Thomas L. Mindt¹¹ · Meltem Ocak¹² · Marianne Patt¹³ · Sergio Todde¹⁴ · Almut Walte¹⁵

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Abstract

This document is intended as a supplement to the EANM "Guidelines on current Good Radiopharmacy Practice (cGRPP)" issued by the Radiopharmacy Committee of the EANM (Gillings et al. in EJNMMI Radiopharm Chem. 6:8, 2021). The aim of the EANM Radiopharmacy Committee is to provide a document that describes how to manage risks associated with small-scale "in-house" preparation of radiopharmaceuticals, not intended for commercial purposes or distribution.

Keywords Risk assessment · Radiopharmaceuticals · Quality assurance

Preamble

The European Association of Nuclear Medicine (EANM) is a professional non-profit medical association that facilitates communication worldwide among individuals pursuing

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Oliver C. Kiss o.kiss@hzdr.de

> Nic Gillings ngil0001@regionh.dk

Olaug Hjelstuen olaug.hjelstuen@ge.com

Martin Behe martin.behe@psi.ch

Clemens Decristoforo clemens.decristoforo@i-med.ac.at

Philip H. Elsinga p.h.elsinga@umcg.nl

Valentina Ferrari ferrarivalentina@yahoo.it

Petra Kolenc petra.kolenc@kclj.si

Jacek Koziorowski jkoziorowski@yahoo.se

Peter Laverman peter.laverman@radboudumc.nl clinical and research excellence in nuclear medicine. The EANM was founded in 1985. This guideline has been written by members of the EANM Radiopharmacy Committee and is intended to assist professionals in the risk assessments regarding the small-scale "in-house" preparation of radiopharmaceuticals.

Thomas L. Mindt thomas.mindt@lbiad.lbg.ac.at

Meltem Ocak melocak@gmail.com

Marianne Patt marianne.patt@uniklinik-leipzig.de

Sergio Todde sergio.todde@unimib.it

Almut Walte walte.almut@mh-hannover.de

- ¹ Department of Clinical Physiology and Nuclear Medicine, Copenhagen University Hospital Rigshospitalet, Copenhagen, Denmark
- ² GE Healthcare, Pharmaceutical Diagnostics, Oslo, Norway
- ³ Center for Radiopharmaceutical Sciences, Paul Scherrer Institute, Villigen, Switzerland
- ⁴ Department of Nuclear Medicine, Medical University Innsbruck, Innsbruck, Austria
- ⁵ Department of Nuclear Medicine and Molecular Imaging, University Medical Center Groningen, Groningen, The Netherlands

Background

GMP Part I [1] and Part II [2] state that, to ensure that medicinal products are suitable for their intended use, comply with regulatory requirements and maintain product quality, safety and efficacy, the quality system must incorporate quality risk management (QRM). This highlights the importance of integrating a robust risk assessment process within the quality system.

GMP Part III [3] contains an entire section on QRM, risk management methods and tools and potential application for QRM. Its purpose is to provide guidance on more effective and consistent risk-based decisions and on some of the most frequently used risk assessment tools. The section also clarifies that it is not always appropriate to use recognised tools only, and the use of informal risk management processes can be considered acceptable.

Relevant references to the need to apply risk management and risk assessment are also included in various EU-GMP Annexes, such as Annexes 1, 3, 11, 13 and 15 [4–8]. The same principles are further stressed in other documents which, although not legally binding, nonetheless may provide useful guidance in focusing on specific aspects and related risks for radiopharmaceuticals. An example of such a document is the European Pharmacopoeia general chapter 5.19 on "Extemporaneous preparation of radiopharmaceuticals" [9], which is specifically dedicated to the preparation of small-scale, on-site preparation of radiopharmaceuticals and where risk assessment is frequently mentioned as a useful and necessary tool to determine

- ⁶ MSD Animal Health, Milton Keynes, UK
- ⁷ Department of Radiopharmaceuticals Production, Institute for Radiopharmaceutical Cancer Research, Helmholtz-Zentrum Dresden-Rossendorf, Dresden, Germany
- ⁸ Department of Nuclear Medicine and Faculty of Pharmacy, University Medical Centre Ljubljana and University of Ljubljana, Ljubljana, Slovenia
- ⁹ Independent Consultant, Linköping, Sweden
- ¹⁰ Department of Medical Imaging, Nuclear Medicine, Radboud University Medical Center Nijmegen, Nijmegen, The Netherlands
- ¹¹ Ludwig Boltzmann Institute Applied Diagnostics, Vienna, Austria
- ¹² Department of Pharmaceutical Technology, Istanbul University, Istanbul, Turkey
- ¹³ Department for Nuclear Medicine, University Hospital Leipzig, Leipzig, Germany
- ¹⁴ Tecnomed Foundation, University of Milano-Bicocca, Monza, Italy
- ¹⁵ Department of Nuclear Medicine, Hannover Medical School, Hanover, Germany

the general level of risk and to make decisions on specific potential issues arising during the preparation of this class of radiopharmaceuticals.

Thus, it is clear why risk assessment principles have gained continuous and increasing attention over the last decades, and they are now considered an important aspect in the framework of the preparation of medicinal products. Thus, QRM principles should be used to assess and control risks associated with the preparation of any pharmaceuticals, including radiopharmaceuticals.

Risk assessment in the preparation of radiopharmaceuticals

Radiopharmaceuticals are a special class of medicinal product, and, due to their inherent nature, it is commonly accepted that general rules and regulations applying to classic, non-radioactive pharmaceuticals need to be adapted in case of radiopharmaceutical preparations. Risk assessment should be viewed as a powerful tool that may help to take decisions and evaluate whether the above adaptations may be considered acceptable and ensure that they do not have an adverse impact on the patient.

There are specific characteristics of radiopharmaceutical preparations that might, in principle, decrease the risk compared with classical pharmaceutical preparations, such as:

- Radiopharmaceuticals are generally used within a few hours of their preparation (microbiological growth in case of contamination is negligible).
- Very small quantities of starting materials are normally used (e.g. in the mg scale). Masses associated with radiopharmaceuticals are often very low (micro-dosing concept), and thus toxicity concerns are often minimal.
- Toxicity is further reduced by the small number of times (often one time only) a radiopharmaceutical is typically administered during the whole life of a patient.
- Shelf lives of radiopharmaceuticals are often short or very short (from less than 1 h up to a few days), and risks related to long-term storage are negligible.
- Finally, "in-house" prepared radiopharmaceuticals are typically used internally, and there are no risks associated with a complex distribution chain.

Quality risk management

QRM should enable the identification of risks and mitigate them with appropriate and robust controls, to ensure that product quality, safety and efficacy are maintained during the product life cycle.

The process of managing, especially the evaluation of the risks associated with the preparation of any pharmaceutical, including radiopharmaceuticals, should be based on scientific knowledge and is ultimately aimed at improving patient protection. A thorough knowledge of the products and critical processes are the basis for a solid and effective QRM program.

QRM should be an integral element of the quality management system, and continuous efforts should be made to effectively manage the risks and optimise the efficiency of the preparation of radiopharmaceuticals. An approved procedure to describe the approach to risk management should be available. The structure and details of risk management procedures depend on many factors, such as site organisation, complexity and the variety of radiopharmaceutical preparation. The level of efforts placed in risk management should be commensurate with the level of risk. For smallscale "in-house" preparation of radiopharmaceuticals, a simplified QRM process may often be adequate. A schematic

[10]

representation of a typical risk management process is given in Fig. 1.

1) Initiation of a risk assessment

To initiate and plan an assessment, the most important things to do are the following:

- Clearly define the aim.
- Select people with a thorough knowledge of the topic to be involved in the assessment process. It is useful to identify a leader.
- Define timelines, if applicable. _
- Define the risk criteria and the acceptance levels (quantitative, qualitative, ranks of severity), which are needed for subsequent risk evaluation.

Defining the aim is very important. Thus, the basic question to be answered when initiating an assessment is

Fig. 1 The key steps in QRM Initiate as described in ICH guideline Quality Risk Management Process Q9 on quality risk management Risk Assessment Risk Identification **Risk Analysis Risk Evaluation** unacceptable Risk Management too Risk Communication **Risk Control Risk Reduction Risk Acceptance** Output / Result of the Quality Risk Management Process **Risk Review Review Events**

"What is the precise purpose of this assessment?" instead of "Which outcome should be made?"

Following the above steps will result in an efficient process, minimising gaps in the assessment. The level of effort, resources, formality and documentation of the assessment should be commensurate with the level of risk, in line with ICH Q9 [10].

The initiation of an assessment may be triggered by several events, such as:

Deviations:

- A deviation (either planned or unplanned) from an established procedure
- An out of specification (OOS) result
- An adverse trend
- An unusual event or abnormal result, which does not necessarily represent a deviation but needs to be assessed and investigated.

Changes:

- Substantial changes related to a process, product, method, equipment or material
- Introduction of a new radiopharmaceutical product, method, equipment and material
- Commissioning of a new radiopharmaceutical preparation laboratory
- Development of a new environmental monitoring program

Standard processes:

- Requirements for storage of retention samples
- Vial labelling prior to radiopharmaceutical preparation
- Batch release before completion of quality control testing
- Approval of suppliers
- Requirements for chemical precursors for small-scale radiopharmaceutical preparation
- (Re)validation of facilities, equipment and processes

2) Risk assessment

Risk assessment is the "core" part of the risk management process and includes risk identification, analysis and evaluation.

The basic questions to be answered during the stages of the assessment process are the following:

- What might go wrong (identification of hazards)?
- What will happen, if it really goes wrong (impact on product quality, patient safety and efficacy)?
- How serious are the consequences (severity)?
- What is the probability of it going wrong (occurrence)?
- How easy is it to detect (detectability)?

Risk identification

The first step is to identify and list potential hazards related to the activity to be assessed. When they have been identified, they must be accurately described to ensure that their exact nature is defined. All available information should be used systematically. If the aim (defined during initiation of the risk assessment) is not clear, the wrong risk could be assessed, and ineffective corrective and preventative actions (CAPA) could be introduced. Risk (or hazard) identification provides the basis for the next stages of the assessment process. This stage addresses the "What might go wrong" question, including identifying the possible impact.

Risk analysis

The risk analysis consists of understanding and estimating the individual parts of the risk associated with each identified hazard. It is the process of ranking and linking the occurrence of the hazard and the severity of potential consequences, either qualitative or quantitative. Risk analysis addresses the questions "What is the probability (occurrence) of it going wrong", and "What are the consequences, if it goes wrong, especially for the patient (severity)". The ability to detect the consequences early enough (detectability) is also a factor of concern at this stage. For a robust output it is useful to differentiate clearly between these three factors: occurrence, severity and detectability.

Risk evaluation

At the risk evaluation stage, the significance of each identified risk is assessed by combining the outcome of the individual parts (occurrence, severity, detectability) based on the criteria defined during the risk initiation. Every combination of the individual parts (occurrence, severity, detectability) attached with a qualitative ("high", "medium" or "low") or quantitative (numeric value) description of risk should be considered. Risk evaluation not only leads to a qualitative or quantitative output of the risk assessment process, but also forms the basis for the following stage of risk control. The robustness of the data collected during the previous stages of the process will determine the quality of the output.

Risk assessment tools

Numerous risk assessment tools are extensively described in ICH Q9 [10] and are beyond the scope of this guideline. A system based on qualitative or simple quantitative methods is generally considered to be adequate for small-scale radiopharmaceutical facilities. The choice of risk assessment tool should be made during initiation of the risk assessment. In this section, qualitative risk assessment, risk ranking and FMEA methods [11] are described.

Qualitative risk assessment

This is the simplest risk assessment method, whereby risks are classified as high, medium or low. The classification must be clearly defined before starting the risk analysis to ensure consistent results.

Risk ranking

This is the simplest quantitative assessment method, and it is applicable to assessment of simple processes. The result is a numeric output (risk priority number = RPN). The rating depends upon three factors:

- i) Severity of the event while having no installed control measures (from no impact to severe impact)
- ii) Occurrence (from highly unlikely to very probable)
- iii) Detectability of the event (are controls and procedures in place to detect the error in good time)

By multiplying the above three factors, a quantitative description of the impact rating is obtained that helps to determine the level of investigation required and whether additional control measures need to be adopted.

A typical example is as follows:

	Severity (S)	Description	
Low	1	Expected to have little negative impact	
Medium	2	Expected to have a medium negative impact	
High	3	Expected to have a high negative impact	

	Occurrence (O)	Description
Low	1	Failure expected to happen less than once per year
Medium	2	Failure expected to happen once per year
High	3	Failure expected to happen more than once per year

	Detectability (D)	Description
High	1	All failures are expected to be detected early enough
Medium	2	Some, but not all failures are expected to be detected early enough
Low	3	None of the failures are expected to be detected or are expected to be detected too late

By multiplying $S \times O \times D$, the RPN is calculated. Suitable action levels are defined based on the RPN, for example:

RPN > 12	High, unacceptable, action needed.	
12 > RPN > 5	Medium, tolerable, further investigations to	
	determine possible actions.	
RPN < 5	Low, acceptable, no action needed.	

FMEA

Failure mode and effects analysis (FMEA) is a more detailed assessment method which is used for complex processes [11]. It evaluates potential failure modes of a process and their likely effects on the product. The first step when conducting a FMEA is to break down a complex process into simpler steps; thus, this assessment tool relies on thorough product and process understanding. Subsequently, the potential failure modes (hazards and their impact) are identified and ranked, assigning an RPN to each potential risk based on previously defined scores for the severity, occurrence and detectability (see paragraph above). Once risks are identified and scored, risk reduction activities can be identified which change the numeric values of occurrence or detectability, resulting in a new RPN.

FMEA is a very useful tool to summarise possible failures, their causes, possible risk reducing activities and their impact on processes/products. It may potentially be applied to any equipment/facility and be used to analyse an operational step and its effect on product or process, to identify the parts of the process that are most in need of change. To perform a successful and useful FMEA, all the steps of the process must be identified with all the possible failure modes. Once the failure modes have been identified, all controls to avoid failure occurrence need to be listed, and the risk and likelihood of the failure evaluated. Examples of risk assessments using FMEA are given below in example 5 and in the literature [12].

3) Risk control

The outcome of risk control is a conscious decision regarding risk acceptance or reduction. The aim is to reduce all risks to an acceptable level, if possible and appropriate. The rationale for accepting the risk must be stated on a case-bycase basis:

- a) Which risks can be accepted because of their low level?
- b) Which risks can and need to be reduced or eliminated?
- c) Which risks cannot be reduced but are acceptable or require the cancellation of the assessed project?

In the first case (a), the risk is deemed acceptable, because its level is so low that specific actions are not necessary. In the second case (b), the impact on product quality, efficacy or patient safety is so severe (unacceptable), that control measures must be implemented to reduce or eliminate the risk. These risk reducing activities can either decrease the occurrence (by validation and training activities) or increase the detectability (by additional quality controls or checks). It is not possible to change the severity for any given risk. The decision to reduce the risk may entail implementation of corrective and preventative actions, redesign of processes and writing or reviewing of documents.

In the third case (c), the risk is estimated unacceptable, but it is not possible to reduce it, or reduction is not considered feasible, so that the proposed control measures are rejected.

4) Risk communication

The extent of risk communication is related to the size and complexity of the production site and organisation but is of utmost importance. Risk communication should occur throughout the whole risk assessment process and should therefore be embedded in the risk management process. The importance of risk communication and who should be informed at which stage of the process should be clearly described in the associated procedure. While initiating a risk assessment process, it can be helpful to share the information for risk identification and evaluation with all relevant personnel in the organisation. At the very least, the completed risk assessment document should be adequately communicated to all relevant personnel.

5) Risk review

Once an assessment has been completed and corrective/preventative actions implemented when deemed appropriate, a review of the entire assessment process must be performed to assess its effectiveness. The risk review process should be embedded in the risk management process and should be clearly described in the risk management procedure. The outcome of the assessment should be reviewed:

- · Regularly to consider new knowledge and experience
- After changes to assess their impact

Practical examples

Examples of risk assessments associated with radiopharmaceuticals are presented below, covering several different situations and factors where a risk assessment may be appropriate.

Risk assessment required for a standard process

Example 1: Qualitative output—batch release before completion of quality control testing for PET radiopharmaceuticals

Due to the short half-life of their radionuclides, PET radiopharmaceuticals may be released before the results of the test for sterility, radionuclidic purity test and environmental monitoring are available. This can only be accepted if an assessment is carried out to make a rationale decision about the associated risks. This simple assessment is carried out qualitatively in the following example:

Introduction

Annex 3 "Manufacture of Radiopharmaceuticals" outlines the requirements for manufacture of radiopharmaceuticals. Adherence to this annex is associated with the following premise (as stated in the annex): Due to short shelf-life of their radionuclides, some radiopharmaceuticals may be released before completion of all quality control tests. In this case, the exact and detailed description of the whole release procedure including the responsibilities of the involved personnel and the continuous assessment of the effectiveness of the quality assurance system is essential

Aim

The aim is to decide if and under which conditions it can be accepted to release the product before completion of all QC tests (here: test for sterility and radionuclidic purity)

Risk identification

- The following hazards can occur, when the product is released before completion of the test for sterility and radionuclidic purity test:
 - Sepsis in the case of microbial contamination

• Extra radiation dose due to radionuclidic impurities

Risk analysis

The risk of releasing radiopharmaceuticals before completion of all quality control tests is high as this could lead to a negative impact on the patient. This risk is mitigated by the fact that only small amounts of the product are administered to patients, all processes are validated, an adequate quality management system is in place and personnel involved in production, quality control and release of radiopharmaceuticals are appropriately trained in specific radiopharmaceutical aspects of the quality management system. All manufacturing steps take place in self-contained facilities dedicated to radiopharmaceuticals, accessible only by authorised personnel. Measures are established and implemented to prevent cross-contamination. Preventative maintenance, calibration and qualification programmes ensure that all facilities and equipment used in the manufacture of radiopharmaceutical are suitable and qualified. The facilities are routinely monitored so that the appropriate level of environmental cleanliness is maintained. The starting materials, packaging materials and critical process aids are purchased from approved suppliers. All documents related to the manufacture of radiopharmaceuticals are prepared, reviewed, approved and distributed according to written procedures. A written procedure detailing the assessment of production and analytical data is followed before the batch is released

Risk evaluation and control

With the adopted risk reducing activities, the risks are considered to be tolerable. All risks are accepted, on the condition that the described risk reducing activities are conducted Example 2: Quantitative output—requirements for chemical precursors for small-scale radiopharmaceutical preparation

When using a chemical precursor for a radiopharmaceutical preparation, there is no need for risk assessment if the chemical precursor complies with a monograph in the European Pharmacopoeia (in this case, the risk assessment was done by people with acknowledged expertise). When there is no individual monograph in the European Pharmacopoeia, then the general monograph "Chemical precursors for radiopharmaceutical preparations" (2902) is applicable. Points which do not comply with the monograph should be risk assessed on a case-by-case basis. This simple assessment is carried out quantitatively in the following example:

Introduction:

The precursor used for the preparation of PET radiopharmaceutical XXX is fully compliant with Ph. Eur. monograph 2902 except for the requirements for microbial contamination and bacterial endotoxins

Aim

The aim is to decide if and under which conditions the chemical precursor for a radiopharmaceutical preparation can be used, although there is no data on microbiological and endotoxin contamination

Risk identification:

- The product may not be sterile and contain excessive levels of bacterial endotoxins
- Risk analysis:

Severity: (3)

The risk of using a product that does not meet the sterility and endotoxin requirements is high as this could be detrimental to the patients health

Detectability: (1)

Controls and procedures are in place which will detect any microbial or endotoxin contamination of the precursor. A bioburden test of the radiopharmaceutical product (without terminal sterile filtration) is performed for every new batch of precursor as part of the incoming goods approval process. Endotoxin testing is performed on the radiopharmaceutical product before release

Occurrence: (1)

This failure is expected to happen less than once a year. The precursor is provided with a certificate of analysis which certifies the chemical purity which is specified at > 97%. YYY is a well trusted supplier and has been audited regularly and found to have a well-established quality management system. Each batch of product is sterile filtered, and each filter is tested for integrity after use. Each batch of product is tested for endotoxins prior to release

$RPN = 3 \times 1x1 = 3$ Low, acceptable

Risk evaluation and control

The risk is considered acceptable with the installed reducing activities including at least a bioburden test of the product with every new batch of precursor and an endotoxin test of the radiopharmaceutical product before release

Risk assessment triggered by deviations

In case of a deviation, a risk assessment may be needed to assess the impact that the event can have on the process/ product and ultimately on patients, to identify the root cause and to identify corrective/preventative actions when necessary. The outcome of the assessment may be to accept the deviation if it is considered an isolated event and its impact is of no concern.

In the case of minor, but repeated deviations, the outcome of the assessment may still be to accept the deviation, but corrective/preventative actions may be considered necessary to prevent reoccurrence. For instance, if the reactor of a radiosynthesis module is not performing as expected and validated and it takes longer to heat up, it is possible that it will reach the point where it fails to reach the required temperature and the radiosynthesis fails to produce a radiopharmaceutical that meets the required specifications. Finally, if the deviation is considered to have a medium/major impact, the product may be rejected, and a thorough investigation may be needed to understand the root cause and resolve the problem.

A deviation procedure must be available within the quality management system, which describes how to identify and evaluate deviations, how to document them (i.e. a deviation form), the approval flow and the responsibilities of personnel involved in the investigation. The classification of the deviations (i.e. minor/medium/major, planned/unplanned) must also be well described. The deviation procedure should also describe how to perform a root cause analysis and how to identify corrective and preventative actions when necessary. The most used root cause analysis tool is the "5 whys". When using the 5 whys tool, the aim is to identify the most probable root cause(s) by starting with a defined problem statement and asking "why did this happen" until you cannot ask the question anymore, as you have reached the final reason that caused the event. The number "5" is not restrictive; it can take more questions or even less questions to identify the most probable root cause.

In the following two examples, a deviation form is used to record the deviation, the assessment, the root cause analysis (using the 5 whys tool) and any corrective actions, according to an approved deviations procedure.

Example 3: Risk assessment within a deviation form due to a minor deviation

DEVIATION FORM	Refer to Deviation Procedure SOP xxx
Deviation number: DEV 01/YYYY	Raised by:
Date of occurrence: DD/MM/YYYY	
Planned Unplanned Adverse trend	Other (Tick as appropriate)
·	
Product:	Batch number:
Equipment/material/process involved:	_1
Deviation description	
The standard transfer line used to dispense PET	radiopharmaceuticals is not available as the
supplier has stopped manufacturing it. The ma	nufacturing procedure states the specific transfer
line used. An alternative transfer line which cou	uld be suitable and is available has been sourced
from a different but non-approved supplier. Th	e use of an alternative transfer line from a non-
approved supplier is proposed.	
The aim of this risk assessment is to assess the	risk of using a transfer line which is different from
the standard one and not provided by an appro	oved supplier. The CoAs of the current and
proposed transfer line have been compared an	d the items are equivalent (same dimensions and
material).	
Assessment	
Risk Identification:	
The proposed transfer line may interact with the pr	oduct.
Risk Analysis:	
SEVERITY: (1)	
Using an alternative line that is not suitable for	or the process could create numerous risks as the
material from the new supplier could contain p	plasticisers which may interfere with the synthesis
causing a low yield. Based on the CoA the cha	aracteristics of the new transfer line appear to be
	r lines, therefore the proposed transfer line is
	roposed line is not the standard one, there is no
foreseeable impact from not using it.	
DETECTABILITY: (1)	
	are subjected to incoming materials checks (CoA
check, visual inspection of material).	
OCCURRENCE: (1)	
This failure is expected to happen less than one	ce a year.
RPN= 1x1x1= 1 Low, Acceptable	
Risk evaluation	
The risk is considered acceptable.	
Root cause analysis	
Problem statement: use of an alternative trans	fer line.
WHY: because the standard transfer line is obs	olete.
WHY: because the approved supplier has stopp	ed manufacturing.
WHY: this is outside of our control.	
Most probable root cause: the supplier stopped	d manufacturing.
Corrective actions taken	
The supplier of the proposed transfer line shou	ld be assessed before adding to the list of
approved suppliers. The procedure should ther	
processes should be controlled through a chan	
Conclusion	
The proposed transfer line has been ordered a	nd will be implemented without any further
	ia will be implemented without any further
actions.	
Deviation number noted in batch record? Y	N N/A C (Tick as appropriate)
APPROVED BY QA:	APPROVED BY DEPARTMENT:

APPROVED BY QA: APPROVED BY DEPARTMENT: DATE: DATE:

Example 4: Risk assessment within a deviation form due to a major deviation

DEVIATION FORM	Refer to Deviation Procedure SOP xxx			
Deviation number: DEV 01/YYYY	Raised by:			
Date of occurrence: DD/MM/YYYY				
Planned Unplanned Adverse trend	□ Other □ (Tick as appropriate)			
Product:	Batch number:			
Equipment/material/process involved:				
Deviation description				
After synthesis of F-18 tracer XXX using a non-ca	ssette-based system, the final radioactive			
product was only partially transferred to the pro	duct vial, and it could not be completely			
recovered. The radioactivity recovered was not s				
product was not delivered/released. It was imme				
was used for the process, therefore the different	t material could have interacted with the final			
product and reduced the yield.				
Assessment				
Risk Identification:				
The production of F-18 tracer XXX failed due to t	rapping of the product on the sterilising filter			
material.				
Risk Analysis:				
SEVERITY: (2)				
The non-conformity affects the quality or effica	acy of the product as the final radioactivity was			
lower than required. The non-conformity affecte	ed the patient investigation as the dose could not			
be delivered.				
DETECTABILITY: (3)				
There is no opportune detectability as there is	no check in place to ensure use of the correct			
filter during production.				
OCCURRENCE: (3)				
This failure is expected to happen more than one	ce a year.			
RPN= 2x3x3= 12 High, Not Acceptable				
Risk evaluation				
The risk is not considered acceptable. Root cause analysis				
Problem statement: The production of F-18 trace	er XXX failed.			
WHY: because the wrong sterilising filter was use	ed.			
WHY: this was caused by operator error.				
WHY: no check to ensure use of the correct filter				
	Most probable root cause: No check is in place to ensure the correct filter is used.			
Corrective actions taken				
CAPA to be raised to:				
1) Add specific details of the filter to the production worksheet.				
 Amend the SOP to include which filter should be used. Train operators in the new procedure. 				
3) Train operators in the new procedure. Conclusion				
	patient to quoid reasoning and this is still at the			
A CAPA has been raised to implement corrective actions to avoid reoccurrence of this incident. A filter check has been added to the production worksheet, the SOP has been amended to include				
miter check has been added to the production worksneet, the SOP has been amended to include which filter needs to be used and operators have been trained in the procedure.				
Deviation number noted in batch record? Y \square N \square N/A \square (Tick as appropriate)				
APPROVED BY QA:	APPROVED BY DEPARTMENT:			

Example 5: Risk assessment within a change control form: introduction of a new product synthesised using a cassette system

	Refer to Change Control SOP xxx
Change control number: CC 01/YYYY	Raised by:
	DATE:
	DATE:
Proposed change:	
Introduction of a new product in the radiop	harmaceutical production facility, synthesised using a
cassette-based system. (Add more details	including the type of product, what equipment is
needed, what area will be used for the prepa	aration.)
Reviewed and approved by QA:	
DATE:	
DATE:	
Assessment of the proposed change:	
The product is similar to products already	prepared in the facility; the equipment used is the
same as used for existing products. Therefore	ore, no equipment validation needs to be performed,
only process validation for the specific radio	opharmaceutical. Cross contamination is controlled by
documented transfer line clearance betwee	en different batches and single use cassettes (closed
system). Therefore, the risk of cross contam	ination is minimal.
Operators need to be trained. New proced	ures need to be written. FMEA should be performed
(see attached FMEA). The actions identified	in the EMEA will form the basis of the proposed plan
, ,	In the PIVEA will form the basis of the proposed plan.
Validation required? Y \square N \square N/A	
Validation required? Y DND N/A	 (Tick as appropriate)
Validation required? Y N N/A	 (Tick as appropriate)
Validation required? Y N N/A Proposed plan: • Manufacturing process to be validate	 (Tick as appropriate)
Validation required? Y N N/A Proposed plan: Manufacturing process to be validated QC process to be validated	 (Tick as appropriate) ed
Validation required? Y N N/A Proposed plan: Manufacturing process to be validated QC process to be validated New labels to be created	 (Tick as appropriate) ed
Validation required? Y N N/A Proposed plan: Manufacturing process to be validated QC process to be validated New labels to be created New release procedure to be created	 (Tick as appropriate) ed
Validation required? Y N N/A Proposed plan: Manufacturing process to be validated QC process to be validated New labels to be created New release procedure to be created Personnel to be trained	 (Tick as appropriate) ed
Validation required? Y N N/A Proposed plan: Manufacturing process to be validated QC process to be validated New labels to be created New release procedure to be created Personnel to be trained Conclusion:	 (Tick as appropriate) ed
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Risk assessment triggered by changes

When introducing a new radiopharmaceutical product, method, equipment or material or in the case of a substantial change, a risk assessment may be needed to assess the impact on the patients' safety and to assess and identify the level of validation required [13]. Commissioning of a new radiopharmaceutical preparation laboratory should also use change management, and a risk assessment should be performed to assess and identify the level of validation required.

A change control procedure should be available to ensure changes are implemented in a controlled manner. If a change is not performed using change management, it is possible that the impact on the validation status of the equipment/process has not been considered, and therefore, the equipment/process is no longer compliant. For example, if a transfer line is changed without an assessment and a material that is not compatible with the radiopharmaceutical is used, the final product could stick to the new material, and the radioactivity recovered in the final product vial could be insufficient.

The change control procedure should specify the workflow and the responsibilities involved. The proposed change should be reviewed and approved by relevant personnel (e.g. process/equipment owner and QA). The proposed change should be risk-assessed by the relevant personnel (e.g. QC or production managers), to ensure it is sufficiently detailed to make an informed decision. Consequently, the change could be either rejected (e.g., because it is not necessary, not sufficiently detailed or holds unacceptable risks) or approved. If the change is approved, a plan should be created and approved by suitable personnel including OA. At this stage, the change is ready to be implemented in a controlled way. Example 5 below includes an example of change control form. After implementation, the change and the associated risk assessment should be reviewed to ensure the change has been implemented correctly.

In the following example, a new radiopharmaceutical product is introduced in a facility. This change is introduced via change management, using a change control form. As it is a complex change with several different factors to be considered, a FMEA is used to perform the risk assessment and to identify any required actions.

If a new radiosynthesis module, new materials and new QC equipment and methods are used, the risks identified in the assessments will usually be considerable, and thus, a full validation will be required (IQ, OQ, PQ). If the new radiopharmaceutical is prepared using a fully validated radiosynthesis module already in use, fully validated QC equipment but different analytical methods, the risks identified in the assessment will usually be lower, and a lower level of validation will be required.

Process	Potential fail- ure mode	Potential failure effects	Severity	Current controls	Occurrence	Detect- ability	RPN	Actions for the proposed plan, which complete the current controls
Label produc- tion	Labels contain incorrect information	Product incorrectly labelled	Medium (2)	Current label pro- cess to be used. The labels are double checked before approval for use and before use	Very low (1)	High (1)	2	New labels needed
Manufacturing	Production fails	Radiopharmaceutical batch is not ready for the patients	High (3)	Current valida- tion policy will ensure that processes are validated	Very low (1)	High (1)	3	Validation of the new manufactur- ing process
QC	QC fails	Radiopharmaceutical batch may not be released	High (3)	Current valida- tion policy will ensure reliability of the process. The analytical method will be validated. The equipment is already validated	Very low (1)	High (1)	3	Validation of new analytical methods
Release	Product is not released	Radiopharmaceutical is not available for the patients	High (3)	The process will be performed according to approved proce- dures, by trained operators	Very low (1)	High (1)	3	New procedures needed. Training to be performed

Attached FMEA:

Appendix

Appendix 1 Abbreviations

EANM	European Association of Nuclear Medicine	
cGRPP	Current Good Radiopharmacy Practice	
QMS	Quality management system	
SOP	Standard operating procedure	
CAPA	Corrective actions and preventative actions	
FMEA	Failure mode and effects analysis	
ISO	International Organization for Standardization	
OOS	Out of specification	
OOT	Out of trend	
Ph. Eur	European Pharmacopoeia	
QC	Quality control	
QRM	Quality risk management	
RP	Radiopharmaceutical	
RPN	Risk priority number	

Abnormal result	Results that are still within specification but are unexpected, unusual	
Adverse trend	An identified trend that indicates a degradation in performance	
CAPA	Corrective actions and preventative actions. A CAPA process is a formal way to implement appropriate corrective and preventative actions with the aim of eliminating causes of non-conformity such as deviations, OOTs, deficiencies from audits and other complaints. Prior to initiation of a CAPA process, non-conformities must be investigated systemically to determine the root cause. CAPA methodology should result in product and process improvements and enhanced product and process understanding	
Change management	A systematic approach to proposing, evaluating, approving, implementing and reviewing changes that ensures that changes to methods, equipment and materials are properly documented and assessed before implementation	
Corrective actions	Actions taken to eliminate the cause of a detected non-conformity to prevent recurrence	
Detection	The process of identifying a failure mode	
Deviation	A non-conformity with respect to approved procedures or established standards or specifications. A devia tion can be planned or unplanned. A planned deviation is pre-approved and covers a specified period or number of batches	
Failure mode	Different ways that a process or sub-process can fail to yield the anticipated result	
Failure mode and effects analysis	The process of reviewing as many components, assemblies, and subsystems as possible to identify poten- tial failure modes in a system and their causes and effects	
Good radiopharmacy practice	Good radiopharmacy practice is described in the "Guidelines on current Good Radiopharmacy Practice GRPP)" by the Radiopharmacy Committee of the EANM [14]	
Occurrence	Probability that an event may happen within a given timeframe	
Preventative actions	Actions taken to eliminate the cause of a potential non-conformity to prevent its occurrence	
Product life cycle	All stages in the life of the product, from development through its clinical use	
Quality risk management (QRM)	A systematic process for the assessment, control, communication, and review of risks to the quality of the pharmaceutical product throughout the product life cycle. Refer to Fig. 1 for key steps in QRM	
Risk	Combination of the occurrence and severity of a hazard (potential source of harm) or non-conformity	
Risk assessment	Risk assessment is the initial step towards collecting information to support a risk decision. It consists of three main stages	
Risk identification	The first stage of the risk assessment process, during which hazards are identified. All available informa- tion should be used systematically	
Risk analysis	The second stage of the risk assessment process, during which the risk associated with the identified haz- ard or non-conformity is estimated	
Risk evaluation	The third stage of the risk assessment process, during which the significance of the risk is determined by comparison of the answers provided during the risk identification and risk analysis stage with given risl criteria (e.g. quantitative scale or qualitative levels such as high, medium and low)	
Risk control	This step follows the risk assessment, and the output is a decision regarding risk acceptance or reduc- tion. Which risks can be accepted because of their low level, which risks can and need to be reduced/ eliminated and which risks cannot be reduced but are accepted without reduction given an acceptable risk-benefit ratio	
Risk reduction	Actions taken to decrease risk to a lower level. This can be achieved by reducing the occurrence or enhan ing the detectability	
Risk acceptance	An active decision to accept an identified risk	
Risk communication	The act of sharing information and outcomes of a risk assessment	
Risk review	Review or monitoring of output or results of a risk assessment considering (if appropriate) new knowledg and experience	
Risk priority number (RPN)	A numeric, quantitative assignment of the level of risk associated with a process or steps in a process. Each failure mode is assigned a numeric score that quantifies the likelihood of occurrence, likelihood of detection and severity of impact. The product of these three scores is the RPN for that failure mode. RPN = severity rating × occurrence rating × detection rating	
Root cause analysis	A systematic process for identifying root causes of problems conformities or other adverse events	
Validation	The documented actions of confirming that any procedure, process, equipment, material, activity or syste leads to the expected results	

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