

This form allows you to notify us of a significant event as required by Regulation 17(1)(e). This form must be used when notifying HIQA of significant events involving **Dental, Radiology** or **Nuclear Medicine** (therapeutic and diagnostic). Significant events should be notified to HIQA within three working days of the discovery of the significant event.

Undertaking and medical radiological installation details			For official use
Undertaking name			
Undertaking address (include Eircode)			
Undertaking email address			
Undertaking contact number			
Medical radiological installation name where incident occurred			
Address incident occurred (include Eircode)			
Designated manager name			
Designated manager email address			
Designated manager contact number			

Section 1. Significant event details				For official use
Exact location incident occurred (area or department or room or unit)				
Date incident occurred				
Time incident occurred (HH:MM)				
Date incident discovered				
Multiple service users affected	Yes	No	Number[†]	
Service user details	Gender		Age	
	Male	Female		

* Please complete this form with HIQA's statutory notification guidance. You can download the guidance at www.hiqa.ie.

† If multiple service users are affected, please contact HIQA at radiationprotection@hiqa.ie for further advice

Please provide brief details of the incident - no personally identifiable information (PII) should be submitted in line with General Data Protection Regulations (GDPR)				
Have appropriate actions been taken to mitigate against immediate recurrence of this incident?	Yes		No	
Please provide brief details of the initial actions taken to mitigate against immediate recurrence of this incident				

Significant event category		
Administered activity variation of 20% from intended dose during use of therapeutic nuclear medicine		
Administration of a Reference Point Air Kerma ($K_{a,r}$) of 15 Gray (Gy) or greater as a result of a single interventional radiological procedure (including interventional cardiology) or a cumulative $K_{a,r}$ dose of 15 Gy arising from a series of interventional radiological procedures carried out over a six month period		
Dose to a breastfed child greater than 1 millisievert (mSv)		
Dose given to comforters and carers greater than 3 millisievert (mSv) for adults under 60 years of age and 15 millisievert (mSv) for those over 60 years of age		
Inadvertent dose to a foetus greater than 1 milligray (mGy)		
Incorrect anatomy greater than 1 millisievert (mSv)		
Incorrect procedure greater than 1 millisievert (mSv)		
Incorrect radiopharmaceutical		
No dose intended/incorrect service user exposed to greater than 1 millisievert (mSv)		
Overexposure of a child of more than twice the exposure intended that leads to a dose that is greater than 3 millisievert (mSv) or 15 times the dose intended		
Overexposure of an adult of more than twice the exposure intended that leads to a dose that is greater than 10 millisievert (mSv) or 20 times the dose intended		
Therapeutic dose given instead of diagnostic dose, for example, in the use of radioiodine		
Tissue reactions (deterministic effects) as a result of interventional radiology/cardiology		
Any other radiation exposure incident considered to have serious service user safety implications, for example, multiple non-notifiable incidents of a similar nature		

Type of procedure or treatment involved in the incident				
Computed Tomography (CT)		Interventional radiology		
Dental		Mammography		
Dual-energy X-ray absorptiometry (DXA)		Nuclear medicine		
Fluoroscopy		Positron Emission Tomography/CT		
Interventional cardiology		Radiology - general		

Section 2. Dental/Radiology/Nuclear Medicine incident details				For official use
Radiology, diagnostic nuclear medicine and interventional procedures Please provide an initial estimated dose	1 to 5 mSv			
	Over 5 to 10 mSv			
	Over 10 to 15 mSv			
	Over 15 to 20 mSv			
	Greater than 20 mSv			
	Other, please specify:			
Therapeutic nuclear medicine procedures Please provide an initial estimated radiation dose variation	Greater than 20% total dose, please specify:			
Type of incident	Hardware/software - Ancillary equipment			
	Hardware/software - Medical radiological equipment			
	Inappropriate or incorrect justification			
	Incorrect protocol selection			
	Optimisation error (practical aspects)			
	Quality assurance error			
	Referral error - wrong patient			
	Referral error - wrong procedure			
	Scheduling error			
	Service user movement			
	Service user related circumstance			
	Wrong anatomical site			
	Wrong service user			
	Wrong service user setup			
	Wrong side (laterality)			
Other, please specify:				

Section 3. Open Disclosure				For official use
Was the incident that occurred considered to be a clinically significant unintended or accidental exposure?		Yes	No	
Did you inform the following individuals of this incident?	Service user/service user representative	Yes	No	
	Referrer	Yes	No	
	Practitioner	Yes	No	

Section 4. Notification of stakeholders				For official use
Please indicate, where applicable , if the following stakeholders have been notified of the incident:				
Medical Physics Expert	<input type="checkbox"/>	Radiation Therapy Services Manager	<input type="checkbox"/>	
Practitioner	<input type="checkbox"/>	Radiography Services Manager	<input type="checkbox"/>	
Risk Manager	<input type="checkbox"/>	Referrer	<input type="checkbox"/>	
Radiation Safety Officer	<input type="checkbox"/>	Radiation Safety Committee or equivalent	<input type="checkbox"/>	
Undertaking	<input type="checkbox"/>			
Other regulatory agencies where necessary, please list if applicable:				

Section 5. Follow-up documentation
The results of the investigation into the significant event and corrective measures to avoid such events must be submitted within the next 120 calendar days.

Section 6. Declaration		For official use
By submitting, I declare that the information I have provided in this form is true to the best of my knowledge and belief. The undertaking is aware that I am making this submission on its behalf.		
Name (print)		
Job Title		
Contact number		
Signed		
	Type your name in the signature field	
Date		

- **Email** form to: radiationprotection@hqa.ie
- **Telephone:** 01 8286750.