RISK ASSESSMENT RECORD FORM

Please refer to your local guidance and risk scoring template when completing this form

Section 1: Administrative D	Details	
Name of Assessor:	Job Title:	Date of Assessment:
Section 2: Activity/Task		
Activity /Task		
Provision of an MRI service		
Risk:		
Adverse reaction to MRI conf	trast agents	
Area affected:		
MRI		
Source of Biok/Bookgroups	<u> </u>	
Source of Risk(Background MRI contrast agents are often	-	enously to patients as part of their MRI scan. These
		chelates. A very small proportion of people will
		agents. This is usually minor, but in rare cases a severe
		phrogenic Systemic Fibrosis (NSF) has been associated
		gnetic resonance imaging (MRI) contrast agents in
		so growing evidence for accumulation of gadolinium in se agents, the clinical significance of which is unclear.
the brain following repeated a	administration of thes	e agents, the chilical significance of which is unclear.
Supporting Evidence:		
MHRA guidance on the admi	inistration of MRI con	trast agents [1].
Factors the risk contains: ((if for COSHH include re	oute of exposure, length of exposure time and exposure limits)
- patient renal function		
Potential Consequence if r	ick is realised:	
Adverse reaction, risk of NSF		
·		
Section 3: Current Control		
when to administer contrast a		tant radiologist, in conjunction with renal physicians, on
when to administer contrast a	agent, taking into acc	ount patient's GFN.
MRI safety screening form in	cludes a question sp	ecifically about known kidney problems.
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MR staff are trained in respon	nding to anaphylactic	reactions.
Section 4: Risk Rating		
-	d and risk score tables	in your local guidance to identify the scores below.
Consequence Score:		
Likelihood Score:		
Risk Score:		

Initial Risk Grading:	

Section 5: Risk Reduction Options								
Options			d Risk Score	Cost				
No further reduction needed								
Section 6: Directorate/Divisional Agreed Actions								
Actions		Lead		Target Date				
Section 7: Risk Grading								
	Consequence		Likelihood	Score	Grade			
Initial:								
Current (will be the same as initial to begin with):								
Residual:								
Section 8: Review								
Risk Owner:								
Planned Review Date:								
Reference								
[1] D. Grainger, "Safety Guidelines for Magnetic Resonance Imaging Equipment in Clinical Use,"								
Medicines and Healthcare Products Regulatory Agency, Mar 2015								