

INSTRUMENTATION AND CONTROL SYSTEMS AND SOFTWARE
IMPORTANT TO SAFETY FOR RESEARCH REACTORS

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INSTRUMENTATION AND CONTROL SYSTEMS AND SOFTWARE IMPORTANT TO SAFETY FOR RESEARCH REACTORS

SPECIFIC SAFETY GUIDE

INTERNATIONAL ATOMIC ENERGY AGENCY

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FOREWORD

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1. INTRODUCTION

BACKGROUND

1.1. Requirements for the safety of research reactors, with particular emphasis on their design and operation, are established in IAEA Safety Standards Series No. SSR-3, Safety of Research Reactors [1].

1.2. This Safety Guide provides recommendations on [the](#) design and operation of instrumentation and control systems for research reactors.

1.3. This Safety Guide was developed in parallel with seven other Safety Guides on the safety of research reactors, as follows:

- (a) IAEA Safety Standards Series No. [SSG-80](#), Commissioning of Research Reactors [2];
- (b) IAEA Safety Standards Series No. [SSG-81](#), Maintenance, Periodic Testing and Inspection of Research Reactors [3];
- (c) IAEA Safety Standards Series No. [SSG-82](#), Core Management and Fuel Handling for Research Reactors [4];
- (d) IAEA Safety Standards Series No. [SSG-83](#), Operational Limits and Conditions and Operating Procedures for Research Reactors [5];
- (e) IAEA Safety Standards Series No. [SSG-84](#), The Operating Organization and the Recruitment, Training and Qualification of Personnel for Research Reactors [6];
- (f) IAEA Safety Standards Series No. [SSG-85](#), Radiation Protection and Radioactive Waste Management in the Design and Operation of Research Reactors [7];
- (g) IAEA Safety Standards Series No. SSG-10 (Rev. 1), Ageing Management for Research Reactors [8];

1.4. Additional recommendations on the safety of research reactors are provided in IAEA Safety Standards Series Nos SSG-20 (Rev. 1), Safety Assessment [for](#) Research Reactors and Preparation of the Safety Analysis Report [9], and SSG-24 (Rev. 1), Safety in the Utilization and Modification of Research Reactors [10].

1.5. The terms used in this Safety Guide are to be understood as defined and explained in the IAEA [Nuclear Safety and Security](#) Glossary [11].

1.6. This Safety Guide supersedes IAEA Safety Standards Series No. SSG-37, Instrumentation and Control Systems and Software Important to Safety for Research Reactors¹.

¹ INTERNATIONAL ATOMIC ENERGY AGENCY, Instrumentation and Control Systems and Software Important to Safety for Research Reactors, IAEA Safety Standards Series No. SSG-37, IAEA, Vienna (2015).

OBJECTIVE

1.7. The objective of this Safety Guide is to provide recommendations on instrumentation and control systems and software important to safety for research reactors, including instrumentation and control system architecture and associated components, ~~from sensors to actuators, human-machine interfaces and auxiliary equipment,~~ to meet the relevant requirements of SSR-3 [1], in particular Requirements 49, 51 and 52.

1.8. The recommendations provided in this Safety Guide are aimed at operating organizations of research reactors, regulatory bodies and other organizations involved in a research reactor project, including suppliers of instrumentation and control systems.

SCOPE

1.9. This Safety Guide is primarily intended for use for heterogeneous, thermal spectrum research reactors ~~having that have~~ a power rating of up to several tens of megawatts. For research reactors of higher power, specialized reactors (e.g. fast spectrum reactors) and reactors ~~having that have~~ specialized facilities (e.g. hot or cold neutron sources, high pressure and high temperature loops), additional guidance may be needed. For such research reactors, the recommendations provided in IAEA Safety Standards Series No. SSG-39, Design of Instrumentation and Control Systems for Nuclear Power Plants [12], might be more suitable. Homogeneous reactors and accelerator driven systems are ~~out of~~ outside the scope of this publication.

1.10. Some research reactors, critical assemblies and subcritical assemblies with a low hazard potential might need a less comprehensive approach. While all recommendations in this Safety Guide are to be considered, some might not be applicable to such research reactors, critical assemblies and subcritical assemblies (see Requirement 12 and paras 2.15–2.17 ~~and Requirement 12~~ of SSR-3 [1], ~~and as well as~~ IAEA Safety Standards Series No. SSG-22, (Rev. 1), Use of a Graded Approach in the Application of the Safety Requirements for Research Reactors [13]).

1.11. In this Safety Guide, subcritical assemblies will be mentioned separately only if a specific recommendation is not relevant for, or is applicable only to, subcritical assemblies.

1.12. The recommendations apply to both the design and the configuration management of instrumentation and control systems and software for new research reactors and the modernization of the instrumentation and control systems and software of existing research reactors.

1.13. This Safety Guide also provides recommendations on human factors engineering and human-machine interfaces for instrumentation and control systems and software important to safety. ~~This~~ The Safety Guide also provides guidance ~~related~~ relating to the consideration of the safety and security interface in the design of instrumentation and control systems for research reactors.

STRUCTURE

1.14. Section 2 provides recommendations on the safety classification of instrumentation and control functions, systems, and components. Section 3 provides recommendations on how instrumentation and control systems are to be arranged into a hierarchy. Sections 4 and 5 provide recommendations on meeting design requirements for instrumentation and control systems. Recommendations on the operational aspects of instrumentation and control systems are provided in Section 6. Section 7 expands on the recommendations provided in Section 4 in the area of human-machine interfaces. Section 8 provides recommendations on the design aspects and other aspects of computer based systems and software. Section 9 provides recommendations on configuration management for instrumentation and control systems. Section 10 provides recommendations on the modification and modernization of instrumentation and control systems. The Annex identifies instrumentation and control systems that can be used in a research reactor.

2. SAFETY CLASSIFICATION OF INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

2.1. Instrumentation and control functions, systems and components may be classified into two categories: those that are important to safety and those that are not important to safety (see Fig. 1 and the Annex). Instrumentation and control systems not important to safety are those that are used to support the operation of the facility, while having but that have no impact on the safety of the reactor.

2.2. Functions, systems and components important to safety are those that contribute to the following main safety functions (see Requirement 7 of SSR-3 [1]):

- (a) Control of reactivity, safely shutting down the reactor and maintaining it in a safe shutdown condition in operational states, accident conditions and post-accident conditions;
- (b) Removal of heat from the reactor and from the fuel storage, in all operational states and accident conditions;
- (c) Confinement of radioactive material, shielding against radiation and control of planned radioactive releases, as well as limitation of accidental radioactive releases.

2.3. Systems and components important to safety are further categorized into safety systems, safety features for design extension conditions (DEC), and safety related systems in accordance with the IAEA Nuclear Safety and Security Glossary [11], as follows:

- (a) Safety systems consist of: the protection system, the safety actuation systems and the safety system support features;

- (b) Safety features for design extension conditions are: items that are designed to perform a safety function for or that have a safety function in design extension conditions;
- (c) Safety related systems are: systems important to safety that are not part of a safety system, such as systems for monitoring the availability of safety systems.

Plant equipment					
Items important to safety					Items not important to safety
Safety features for DEC	Safety systems			Safety related items	
	Protection systems	Safety actuation systems	Safety system support features		
I&C for:	Initiation I&C for: Reactor trip Emergency core cooling Decay heat removal	Actuation I&C for: Reactor trip Emergency core cooling Decay heat removal	I&C for: Emergency power supply	Reactor control systems Plant control systems Control rooms I&C Radiation monitoring system	I&C for: Off-line water demineralizing plant Off-line water treatment systems
	Mobile/ supplementary emergency power generators Emergency water makeup Instrumentation for operating under extreme conditions Additional power backup and control measures	Dynamic confinement isolation Confinement heat removal I&C for command and monitoring: Safety parameter command and display consoles and panels		Confinement isolation Confinement heat removal	

~~Note: CCTV — closed circuit television; HVAC — heating, ventilation and air conditioning; I&C — instrumentation and control.~~

FIG. 1. Examples of instrumentation and control systems of a research reactor classified in accordance with their importance to safety. CCTV — closed circuit television; DEC — design extension conditions; HVAC — heating, ventilation and air-conditioning; I&C — instrumentation and control.

2.4. A graded approach to the application of safety requirements for a research reactor is required: (see Requirement 12 of SSR-3 [1]). For instrumentation and control systems important to safety, the extent to which a graded approach is applied should be clearly justified in the safety analysis report (the factors to be considered are listed in para. 2.17 of SSR-3 [1]). Recommendations on the application of the graded approach are provided in SSG-22 [13].

CLASSIFICATION OF INSTRUMENTATION AND CONTROL SYSTEMS

2.5. Paragraph 6.29 of SSR-3 [1] states (footnote omitted):

“The method for classifying the safety significance of items important to safety shall be based primarily on deterministic methods complemented, where appropriate, by probabilistic methods (if available), with due account taken of factors such as:

- (a) The safety function(s) to be performed by the item;
- (b) The consequences of failure to perform a safety function;
- (c) The frequency with which the item will be called upon to perform a safety function;
- (d) The time following a postulated initiating event at which, or the period for which, the item will be called upon to perform a safety function.”

2.6. An instrumentation and control system wherein which a failure on demand, or a spurious actuation, might cause an initiating event, or make the consequences of a postulated initiating event worse, should be classified in a high safety class. A similar approach should be applied to the instrumentation and control aspects of reactivity control systems whose failure might lead to accident conditions.

2.7. The instrumentation and control functions for all facility states of the research reactor should be identified. For a research reactor, the instrumentation and control functions necessary to mitigate the consequences of design extension conditions could be assigned to a lower safety class than the functions necessary for the control of anticipated operational occurrences and design basis accidents to reach a safe state.

DESIGN, CONSTRUCTION, COMMISSIONING, OPERATION AND MAINTENANCE OF INSTRUMENTATION AND CONTROL SYSTEMS

2.8. All instrumentation and control systems and equipment should be designed, constructed, commissioned, operated and maintained in such a manner that their specification, verification and validation, and as well as their quality and reliability, are commensurate with their safety classification. The specifications should consider sufficient functional margins for their safety system design. These margins should be verified at both component level and system level by testing and analysis.

2.9. Paragraph 6.30 of SSR-3 [1] states:

“The design shall be such as to ensure that any interference between items important to safety will be prevented, and in particular that any failure of items important to safety in a system in a lower safety class will not propagate to a system in a higher safety class.”

All instrumentation and control systems performing that perform functions important to safety should have appropriately designed interfaces with systems and equipment of different safety classes (by using through appropriate physical or logical barriers) in order to meet the above requirement. Equipment that prevents such a propagation of failure should be treated as being in the higher class.

2.10. The safety class of an instrumentation and control system should be the same as the highest safety class of the systems or equipment ~~that~~ it controls.

3. OVERALL INSTRUMENTATION AND CONTROL SYSTEM ARCHITECTURE FOR A RESEARCH REACTOR

3.1. The instrumentation and control system architecture should support all the instrumentation and control functions necessary to ensure the safety of the research reactor. Requirement 49 of SSR-3 [1] states:

“Instrumentation shall be provided for a research reactor facility for monitoring the values of all the main variables that can affect the performance of the main safety functions and the main process variables that are necessary for its safe and reliable operation, for determining the status of the facility under accident conditions and for making decisions for accident management. Appropriate and reliable control systems shall be provided at the facility to maintain and limit the relevant process variables within the specified operating ranges.”

The instrumentation and control systems for ensuring the safety of the research reactor in normal operation should cover startup, operation at power, shutting down, refuelling and maintenance.

3.2. Paragraph 6.168 of SSR-3 [1] states that “The reactor shall be provided with appropriate controls, both manual and automatic as appropriate, to maintain parameters within specified operating ranges.” The instrumentation and control systems should be able to automatically initiate reactor shutdown, emergency core cooling, residual heat removal and the confinement of radioactive material, although manual operation ~~action~~ may be permitted as described in para. 5.14. The instrumentation and control system architecture should provide sufficient capabilities to cover all anticipated operational occurrences, accident conditions, and post-accident conditions.

3.3. The instrumentation and control system architecture (see paras 3.15–3.18) should provide a high level definition of the instrumentation and control systems, the assignment of instrumentation and control functions to these systems, and the communications (interfaces) between instrumentation and control systems and the reactor operators and users. The architecture of highly integrated systems should be carefully considered to ensure proper implementation of the defence in depth concept (see paras 3.4–3.6). The architectural design should include the rational allocation of functions, ensuring that these functions are only ~~assigned to~~ the systems wherein which they are needed. The specific instrumentation and control systems that can be included within the overall system architecture of a particular research reactor depend on the type of reactor, its purpose and its operating modes. Different instrumentation and control systems are described in the Annex.

~~THE APPLICATION~~APPLICATION OF ~~THE~~ DEFENCE IN DEPTH ~~CONCEPT~~ TO INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

3.4. Paragraph 2.11 of SSR-3 [1] states:

“Application of the concept of defence in depth throughout design and operation provides protection against anticipated operational occurrences and accidents, including those resulting from equipment failure or inappropriate human actions within the installation and events induced by external hazards.”

3.5. Requirement 10 of SSR-3 [1] states that **“The design of a research reactor shall apply the concept of defence in depth. The levels of defence in depth shall be independent as far as is practicable.”**

3.6. The objectives of the instrumentation and control system architecture should include the following:

- (a) To apply the defence in depth concept. For instrumentation and control, defence in depth includes implementing successive instrumentation and control functions designed to limit the consequences of a postulated initiating event despite the failure of the instrumentation and control systems designed to respond first.
- (b) To not compromise the overall application of the defence in depth concept to the design of the research reactor.

INDEPENDENCE OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

3.7. The principle of independence is intended to prevent the propagation of failures from the item affected by the failure to other redundant items, or from one system to another system.

3.8. The instrumentation and control system architecture should not compromise the independence of the different levels of defence in depth.

3.9. Safety systems should be independent of systems of a lower safety class to prevent faults ~~from~~ propagating from ~~thesethe~~ systems of ~~athe~~ lower safety class and to ensure that the safety systems can perform their safety functions as necessary.

3.10. The design of safety system support features should not compromise the independence between redundant components of safety systems or between safety systems and systems of a lower safety class.

CONSIDERATION OF COMMON CAUSE FAILURE IN INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

3.11. A common cause failure is defined as the failure of two or more structures, systems and components due to a single event or cause [11]. Common cause failure might happen, for example, because of the following:

- (a) Human errors in operation or maintenance;
- (b) A design deficiency;
- (c) A manufacturing deficiency;
- (d) Inadequate specification;
- (e) Inadequate qualification for or protection against internal or external hazards, ~~a~~-human induced ~~eventevents~~, high voltages, data errors, data communication errors, or failure propagation between systems or components.

3.12. Requirement 26 of SSR-3 [1] states:

“The design of equipment for a research reactor facility shall take due account of the potential for common cause failures of items important to safety, to determine how the concepts of diversity, redundancy, physical separation and functional independence have to be applied to achieve the necessary reliability.”

Electrical isolation is used to prevent electrical failures in one system from affecting connected systems or redundant elements within a system and should be considered as a means to achieve the necessary reliability.

3.13. Latent failures and common failure modes that could potentially result in a common failure of redundant items should be identified. Justification should be provided ~~for~~ before declaring that a source of common cause failure between systems or individual components is not considered credible and need

not be considered further. This justification ~~can~~ could be based, for example, on the assigned level of defence in depth of the instrumentation and control function, the dependability of the components or the technology applied.

3.14. An analysis should be conducted of the consequences of each postulated initiating event within the scope of the safety analysis in combination with common cause failures that will prevent a protection system from performing the necessary safety functions.

ARCHITECTURAL DESIGN OF THE INSTRUMENTATION AND CONTROL SYSTEM FOR A RESEARCH REACTOR

3.15. The instrumentation and control system architecture should include the following provisions:

- (a) It should include all the instrumentation and control functions necessary to ensure the safe operation of the research reactor and to manage anticipated operational occurrences and accident conditions.
- (b) It should include the systems necessary to support the defence in depth strategy for the research reactor.
- (c) It should ensure that the actions needed for the functioning of safety systems have priority over opposite actions needed for the functioning of systems with a lower safety class.
- (d) It should ensure a suitable arrangement of systems and components so that they can be adequately tested and maintained at regular intervals, and ~~be~~ capable of self-checking as appropriate, in accordance with Requirement 51 of SSR-3 [1].
- (e) It should divide the overall instrumentation and control system into individual systems as necessary to achieve the following objectives:
 - (i) To fulfil design basis requirements for independence between functions at different levels of the defence in depth concept;
 - (ii) To adequately separate systems and functions of different safety classes;
 - (iii) To establish the redundancy necessary to fulfil design basis reliability requirements;
 - (iv) To support the compliance of safety systems or groups of safety systems with the single failure criterion and the fail-safe concept.
- (f) It should define the human-machine interface as well as the interfaces between the individual instrumentation and control systems.
- (g) It should consider the future utilization and modification of the research reactor, and potential changes to the configuration of the instrumentation and control systems, to enable configuration management throughout the lifetime of the reactor.

- (h) It should include the necessary information and operator controls in the main control room and the supplementary control room (if applicable) and other areas where information is needed for operation or for managing management of an accident²;
- (i) It should include the automatic controls necessary to maintain and limit the process variables important to safety within the specified normal operational ranges.

3.16. The inputs to the design of the instrumentation and control system architecture should refer to the documented design basis for the research reactor, which should provide the following information:

- (a) The application of the defence in depth concept to the research reactor;
- (b) The safety functions to be fulfilled to address postulated initiating events;
- (c) The safety classification and the functional and performance requirements of items important to safety;
- (d) The assignment of functions to manual means and to automatic means, and the role of automation and the actions of reactor operators in the management of anticipated operational occurrences and accident conditions;
- (e) The information to be provided to operating personnel²;
- (f) The criteria for prioritizing automatically initiated and manually initiated actions;
- (g) Regulatory requirements, including those for the authorization of instrumentation and control systems;
- (h) Operational features (e.g. the design of the instrumentation and control system with regard to the human-machine interface) for systems important to safety.

3.17. The use of diversity, redundancy and independence (i.e. physical separation² and electrical and functional isolation) in the architecture of the instrumentation and control systems should be consistent with the safety classification of each instrumentation and control system² and with the application of the defence in depth concept, both for the overall facility and for the instrumentation and control systems. With regard to redundancy, other factors — such as reliability³ and the availability of instrumentation and control systems — should be considered.

3.18. The instrumentation and control system is required to have a fail-safe design (see Requirement 28 of SSR-3 [1]) such that any malfunction within the system caused solely by variations in conditions within the ranges detailed in the design basis would not result in an unsafe condition or failure.

² The operating personnel comprise the reactor manager, the reactor supervisor, the shift supervisors, the reactor operators, the maintenance personnel and the radiation protection personnel.

³ Reliability is the probability that a system or component or an item will meet its minimum performance requirements when called upon to do so, for a specified period of time and under stated operating conditions [11].

4. DESIGN GUIDELINES FOR INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

4.1. Instrumentation and control systems are required to conform to their design bases (see paras 4.3 and 4.4 of this Safety Guide), established in accordance with Requirement 17 of SSR-3 [1]. The origin of, and the objective for, each element of the design basis should be specified and documented to facilitate verification and traceability and to demonstrate that all relevant design requirements have been met.

4.2. The design of the instrumentation and control systems should be as simple as possible while still ensuring that their safety functions are fulfilled. Simplicity of design results in fewer components, simpler interfaces, easier verification and validation, and easier maintenance of the hardware and software. The design requirements for instrumentation and control systems should be carefully analysed to ensure simplicity of design.

DESIGN BASIS FOR INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

4.3. The design basis for each instrumentation and control system important to safety for a research reactor should specify the following:

- (a) The facility states (operational states and accident conditions) in which the system is required to function;
- (b) The various configurations of the research reactor ~~and, as well as~~ experimental configurations, that the instrumentation and control system needs to accommodate;
- (c) Functional requirements for the system in each facility state and operating mode, including extended shutdown;
- (d) Performance requirements for fulfilment of safety functions, including the guaranteed response time, latency, precision and instrument error;
- (e) The facility conditions during which manual control is allowed for each manual protective action;
- (f) The postulated initiating events to which the system is required to respond;
- (g) The variables, (or combination of parameters) to be monitored, the control actions required, and the identification of actions to be performed automatically, manually or both;
- (h) The necessary ranges, rates of change, and accuracy of input and output signals of the system;
- (i) Constraints on the values of process variables in all postulated conditions;
- (j) Criteria for periodic testing, self-diagnostics and maintenance;

- (k) System reliability levels, which may be specified using deterministic criteria, probabilistic criteria or both;
- (l) Requirements for system availability;
- (m) The range of transient and steady state environmental conditions under which the system is required to perform a safety function;
- (n) The range of environmental conditions, including those hazards arising from natural phenomena, under which the system is required to perform a safety function;
- (o) Any conditions with the potential to degrade the functional performance of systems important to safety and the provisions necessary to retain their capability to perform a safety function;
- (p) Operational constraints, such as the need to interface with other systems.

4.4. The design bases for reactor protection systems should specify the following:

- (a) The settings for the actuation of safety systems, which should be derived from the safety analysis;
- (b) The variables to be displayed so that the reactor operators can confirm the operation of the reactor protection system or ~~to enable them to~~can initiate manual actions;
- (c) The conditions (including duration) under which a bypass of instrumentation and control safety functions is to be permitted to allow for changes in operating modes, testing or maintenance.

DESIGN FOR RELIABILITY OF INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

4.5. Several measures should be used, if necessary in combination, to achieve and maintain the required reliability (see Requirement 24 of SSR-3 [1]) of instrumentation and control systems.

Application of redundancy and the single failure criterion to instrumentation and control systems for a research reactor

4.6. A single failure is a failure that results in the loss of capability of a component to perform its intended safety ~~function(s)~~functions and any consequential ~~failure(s)~~failures that ~~results~~result from this loss of capability [11]. The single failure could occur when the safety function needs to be performed or at any time prior to that.

4.7. The single failure criterion is required to be applied to each safety group in the design of a research reactor (see Requirement 25 of SSR-3 [1]) and involves a deterministic approach to determine the necessary degree of redundancy for items important to safety. For safety systems, the single failure criterion should be applied so that the system is capable of performing its intended safety function in the presence of any single failure. A single failure in the system should be considered together with ~~(i)~~ (a) other failures as a consequence of postulated initiating events; and ~~(ii)~~ (b) any credible undetected fault in the system.

4.8. Redundancy is an important design principle for enhancing the safety and reliability of systems important to safety. The concept of redundancy should be applied through the adequate provision of alternative (identical or diverse) structures, systems or components (SSCs) so that any single structure, system or component can perform the required function regardless of the state of operation or failure of any other-.

4.9. Paragraph 6.79 of SSR-3 [1] states that “The degree of redundancy adopted shall reflect the potential for undetected failures that could degrade reliability.” For instrumentation and control systems important to safety, redundancy should be applied (at the system level or component level, or at both) to the extent necessary to conform to the design basis for reliability and availability. For instrumentation and control systems that are safety systems, redundancy should also be applied to the extent necessary to comply with the single failure criterion when systems or components are removed from service for planned surveillance or testing.

4.10. Where compliance with the single failure criterion is not sufficient to meet reliability requirements, additional design features should be provided, or modifications to the design should be made.

Common cause failure in instrumentation and control systems for a research reactor

4.11. Paragraph 6.80 of SSR-3 [1] states that “The principle of diversity shall be adopted wherever practicable, after consideration of its possible disadvantages from complications in operating, maintaining and testing the diverse equipment.” The design of instrumentation and control systems important to safety should minimize the possibility of common cause failures by applying the criteria of independence and diversity. Safety systems should be designed in such a manner that common cause failures are prevented or mitigated.

Physical separation and independence of instrumentation and control systems for a research reactor

4.12. Requirement 27 of SSR-3 [1] states:

“Interference between safety systems or between redundant elements of a system for a research reactor facility shall be prevented by means such as physical separation, electrical isolation, functional independence and independence of communication (data transfer), as appropriate.”

4.13. Independence is a design feature that reduces the risk of common cause failures and failure propagation. As far as practicable, redundant safety systems should be physically separated and electrically isolated from each other and from systems of a lower safety class. The concept of independence should be applied to the entire safety system, for example between redundant trains within the same system and across diverse systems fulfilling the same function, such as first and second shutdown systems. Safety Different safety functions ~~which are required to be independent~~ should be

performed by different modules, components or systems to avoid the ~~effects on each other of the~~ failure of any one of these items affecting the performance of the others.

4.14. Physical separation should be considered as a means of avoiding common cause failures resulting from fire, flooding ~~and/or~~ other external events or accident conditions. Physical separation also reduces the likelihood of inadvertent human errors.

4.15. The extent to which independence might be lost after a postulated initiating event should be considered in the design of specific parts of the research reactor, such as confinement penetrations, cable spreading rooms, equipment rooms and control rooms.

4.16. Electrical connections and data connections between redundant divisions within a safety system should be designed so that no credible failure in one redundant division would prevent the other redundant ~~division(s)~~divisions from meeting design requirements for performance and reliability.

4.17. Electrical connections and data connections between safety systems and systems of a lower safety class should be designed so that no credible failure in the system of a lower safety class would prevent the safety systems from meeting design requirements for performance and reliability.

4.18. Electrical isolation should be used to control or prevent adverse interactions between equipment and components caused by factors such as electromagnetic interference, electrostatic pickup, short circuits, open circuits, grounding and the application of the maximum credible voltage (AC or DC). Examples of provisions for electrical isolation are electronic isolating devices, optical isolating devices (including optical fibres), relays, shielding of cables or components, separation and distance, or combinations of these.

4.19. When isolation devices are used between safety systems and systems of a lower safety class, the isolation devices should be part of the system having a higher safety class.

4.20. When it is not feasible to provide adequate physical separation or electrical isolation between safety systems and systems of a lower safety class, the following recommendations should be implemented:

- (a) The system of a lower safety class should be identified as part of the safety system with which it is associated.
- (b) The system of a lower safety class should be independent of other systems that are also in the lower safety class.
- (c) The system of a lower safety class should be analysed or tested to demonstrate that it does not unacceptably degrade the safety system with which it is associated.

4.21. If data communication channels are used in safety systems between items required to be independent, their design should also apply the concept of independence (functional independence, electrical isolation and physical separation, as appropriate). This includes independence from the effects of communication errors.

Diversity of instrumentation and control systems for a research reactor

4.22. Diversity is the presence of two or more redundant systems or components to perform an identified function, where the different systems or components have different attributes so as to reduce the possibility of common cause failure, including common mode failure. Diversity thus increases the likelihood that appropriate safety actions will be performed when necessary. This includes functional diversity, [and equipment diversity \(see para. 4.25\)](#). [Examples include](#) the provision of types of equipment that use different physical methods to provide physical diversity, different working principles, different hardware and/or software designs, different design teams using different development methods, and different manufacturers using different designs.

4.23. Diversity in instrumentation and control systems should be applied through monitoring and processing parameters using different methods or technologies, different logic or algorithms, and/or different means of actuation in order to provide more than one way to detect and respond to a specific event. It should be ensured that the necessary diversity of instrumentation and control systems is achieved in the design and preserved throughout the lifetime of the research reactor.

4.24. [WhereWhen](#) independence is claimed between two systems (e.g. a research reactor with a primary reactor protection system and a second diverse reactor protection system; see para. 5.11) through multiplying their failure probabilities within the probabilistic safety assessment, then the full instrumentation and control chain — from the sensors, signal conditioning devices, and signal processors and calculators to the actuator drivers — should be used when demonstrating diversity.

4.25. Diversity applied to instrumentation and control systems should include one or both of the following means:

- (a) Functional diversity: This could be achieved by systems providing different physical functions or physical means, resulting in the same safety effects.
- (b) Equipment diversity: This could be achieved by sensors and systems using different technologies or [being](#) designed and produced by different manufacturers.

4.26. In assessing equipment diversity with claimed technological diversity, attention should be paid to the components of equipment to ensure that diversity actually exists. For example, different manufacturers might use the same processor or the same operating system, thereby potentially incorporating common failure modes. Claims for technological diversity that are based only on a difference in manufacturers' names are insufficient without [thesesuch](#) considerations.

Failure modes of instrumentation and control systems for a research reactor

4.27. The possible failure modes of instrumentation and control systems important to safety should be identified during the design process. Systematic failure modes should be eliminated. Non-systematic failure modes should be properly documented using methods of failure mode analysis and cause-and-

effect analysis. The more probable non-systematic failure modes should neither place the system in an unsafe state nor cause spurious actuation of safety systems.

4.28. The identification and analysis of failure modes of instrumentation and control systems important to safety should consider human factors and the human-machine interface.

4.29. Failures of components of instrumentation and control systems should be self-indicating or should be detectable by means of periodic testing or by an alarm or other indication.

4.30. The design of instrumentation and control systems important to safety should include provisions for detecting all postulated (identified) failure modes in the system by self-checking, preferably involving a combination of failure alarms, and testing the credibility of readings, as appropriate. This is usually in addition to periodic testing to demonstrate system performance.

Fail-safe design of instrumentation and control systems for a research reactor

4.31. Fail-safe design is required to be considered and incorporated, as appropriate, in the design of instrumentation and control systems— (see Requirement 28 in SSR-3 [1+]). The design should ensure that in the event of a failure of a system, the system enters a safe state, with no necessity for any action to be initiated by any system or by reactor operators.

DESIGN CONSIDERATIONS FOR AGEING OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

4.32. The service life of electrical and electronic systems and components might be considerably less than the lifetime of the facility. Ageing effects that impair the ability of a qualified safety system component to function under severe service conditions could occur well before there is any detectable effect on the component's functional capabilities under normal conditions. Degradation mechanisms that could affect instrumentation and control system components and the means for detecting the resulting ageing effects should be identified during design. Ageing is commonly due to heat and to radiation exposure; however, the possibility that other phenomena (e.g. mechanical vibration, chemical degradation) might be relevant to a specific component should be considered.

4.33. Obsolescence is required to be considered in the design of instrumentation and control systems; (see Requirement 37 and para. 6.112 of SSR-3 [1+]). Factors to be considered include planning for and managing reductions in service life, diminishing manufacturing sources and material shortages. Special attention should be given to the obsolescence of computer-based equipment.

4.34. Potentially significant ageing effects (e.g. thermal ageing, radiation ageing) should be addressed in the design to ensure that the necessary functionality, in operational states and accident conditions, is maintained up to the end of the service life of the system or component. Further conservatism should be applied, where appropriate, to allow for unanticipated degradation mechanisms.

4.35. Examples of means to address the impacts of ageing include the following:

- (a) Replacement of a component before the end of its qualified service life;
- (b) Adjustment of functional characteristics (e.g. recalibration) to take into account the effects of ageing;
- (c) Changes to maintenance procedures or environmental conditions that have the effect of slowing the ageing process;
- (d) Monitoring of the condition of equipment, including self-checking, for ageing characteristics.

4.36. Further recommendations on ageing management and obsolescence management are provided in SSG-10 (Rev. 1) [8].

CONSIDERATION OF THE SAFETY AND SECURITY INTERFACE IN THE DESIGN OF INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

4.37. The purpose of nuclear security (including physical and computer security) applied to instrumentation and control systems of research reactors is to prevent, detect and, when detected, eliminate or reduce the vulnerabilities that could be exploited from either outside or inside the site area of the protected facility, or those associated with material, equipment, software and data (see also Section 8). The instrumentation and control systems for a research reactor fulfil functions of both safety and nuclear security. The architectural and functional vulnerabilities of ~~these~~the instrumentation and control systems and ~~their~~the consequences of those vulnerabilities for the safety and nuclear security of the research reactor should be assessed.

4.38. Security provisions need to be considered in the instrumentation and control system from the beginning of the design of the system. One of the primary security considerations is the potential for the failure or manipulation of an instrumentation and control system due to an external or internal malicious act. The design of the instrumentation and control systems for a research reactor needs to consider and include measures to prevent malicious acts or exploitations of the system.

4.39. Many design concepts and components in the overall architecture of the instrumentation and control systems contribute to enhancing both safety and nuclear security; however, to meet Requirement 90 of SSR-3 [1], an assessment of the system architecture should be performed to ensure that safety measures and security measures do not compromise one another. ~~Where~~If potential conflicts are identified, compensatory measures should be considered during the design, so as not to weaken the safety or nuclear security of the systems.

4.40. Neither the operation nor the failure of any computer security feature should adversely affect the ability of an instrumentation and control system to perform its safety function. Similarly, the performance of a safety function should not affect the nuclear security of the research reactor.

4.41. If computer security measures are included in the human–machine interface, they should be designed to ensure that they do not adversely affect the ability of operating personnel to maintain the safety of the research reactor.

4.42. Where practicable, security measures that do not also provide a benefit for safety should be implemented in devices that are separate from instrumentation and control systems.

4.43. The programmes and procedures used by the operating organization to ensure safety in the design of instrumentation and control systems should not create adverse effects on the security system.

4.44. Operating organizations and designers should consider safety and nuclear security ~~and, as well as~~ computer safety and security, in all phases of the ~~I&C~~instrumentation and control system ~~life cycle~~, ~~namely life cycle~~: specification of design requirements; conceptual, preliminary and detailed design; and the procurement, fabrication, integration, installation, commissioning, operation and maintenance, and decommissioning of the instrumentation and control systems.

4.45. Nuclear security recommendations for nuclear facilities are provided in Ref. [14], and guidance on computer security is provided in Refs [15–17]. National requirements for the security of information technology also need to be considered.

EQUIPMENT QUALIFICATION OF INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

4.46. Instrumentation and control systems and components important to safety are required to be qualified for their intended functions: ~~(see Requirement 29 in of SSR-3 [17]).~~ The qualification should provide a degree of confidence commensurate with the safety class of the system or component. Components should meet all design basis requirements when subjected to the range of service conditions specified in the design basis. The basis for qualification should be documented. Recommendations on equipment qualification are provided in IAEA Safety Standards Series No. SSG-69, Equipment Qualification for Nuclear Installations [18].

4.47. The qualification programme ~~(s)~~ should address all topics affecting the suitability of the system or component to fulfil the intended safety functions, including the following:

- (a) The suitability and correctness of systems and components to perform the intended safety ~~function(s);~~functions;
- (b) Environmental qualification (including for radiation resistance, if applicable);
- (c) Seismic qualification;
- (d) Qualification for electromagnetic compatibility of systems and components.

4.48. ~~4.x~~—Qualification should be based ~~upon~~on a combination of methods, including the following:

- (a) The use of engineering and manufacturing processes in compliance with established codes and standards;
- (b) A demonstration of reliability;
- (c) ~~Using~~The use of operating experience from similar applications;
- (d) Testing of the equipment;
- (e) Analysis to extrapolate test results or operating experience under pertinent conditions;
- (f) Ageing analysis, as applicable.

4.49. Traceability should be established between all installed structures, systems and components important to safety and the applicable evidence of qualification. This includes traceability not only to the component itself, but ~~traceability~~ between the tested configuration and the installed configuration.

4.50. The equipment qualification programme should demonstrate that the as built instrumentation and control systems and installed components correctly implement the qualified design.

Suitability and correctness of instrumentation and control systems for a research reactor

4.51. The design of instrumentation and control systems and components important to safety of a research reactor should meet all functional, performance and reliability requirements contained in the design basis and in the equipment specifications. Examples of functional requirements include those for the functionality of the application and support systems, for equipment operability and the human-machine interface, and for the input and output range. Examples of performance requirements include those for accuracy and response time. Examples of reliability requirements include those for fail-safe behaviour, conformance with the single failure criterion, independence, failure detection, maintainability and service life.

Consideration of internal and external hazards in the design of instrumentation and control systems for a research reactor

4.52. Instrumentation and control systems and components should be protected against, ~~and~~or should be designed and qualified to withstand, internal and external hazards and their credible combinations, ~~(including seismic hazards)~~ that are included in the design basis and in the safety analysis.

Environmental qualification of instrumentation and control systems for a research reactor

4.53. The qualification programme is required to include the environmental conditions for which ~~the~~ equipment is qualified ~~– (see para. 6.82 of SSR-3 [1]).~~ Environmental conditions include temperature, pressure, humidity, chemicals and radiation, and degradation mechanisms that might affect the proper functioning of components under those conditions. Instrumentation and control systems important to safety are required to be designed to withstand the effects of, and to operate under, the environmental

conditions associated with normal operation, anticipated operational occurrences and design basis accidents; (see para. 6.83 of SSR-3 [17]).

Qualification of instrumentation and control systems for a research reactor for electromagnetic compatibility

4.54. The reliable operation of electrical and electronic systems and components depends on their electromagnetic compatibility with components located nearby or with which they are connected. Electromagnetic interference could be caused by sources internal or external to the facility, ~~for example,~~ Examples include fault current clearance by the operation of switchgear ~~or,~~ circuit breakers or fuses; electromagnetic fields caused by radio transmitters; and natural sources, such as lightning strikes and geomagnetically induced currents. Instrumentation and control systems and components, including associated cables, should be designed, installed and tested to withstand the conditions of their electromagnetic environment.

4.55. The types of electromagnetic interference to be considered in the design of instrumentation and control systems and components include the following:

- (a) Emission and conduction of electromagnetic disturbances via cables;
- (b) Electrostatic discharge.

4.56. The qualification of instrumentation and control systems and components for electromagnetic compatibility depends ~~upon~~ a combination of design measures for systems and components to minimize the coupling of electromagnetic noise to electrical components. Testing should be conducted to demonstrate that systems and components can withstand the expected levels of electromagnetic radiation and to demonstrate that their own electromagnetic emissions are within tolerable levels. Instrumentation and control systems and components that are already qualified should be accompanied by the corresponding qualification certificate.

4.57. The electromagnetic emission characteristics of wireless systems and devices used at the research reactor, as well as those of devices used for repair, maintenance and measurements, should be taken into consideration. Such wireless systems and devices could include; mobile telephones, radio transmitters and receivers, and wireless data communications networks. Testing for electromagnetic emissions should be applied to systems and components both important to safety and not important to safety.

4.58. Any electrical or electronic equipment at the facility can contribute to the electromagnetic environment. The contribution of electromagnetic emissions from all equipment — not only equipment important to safety — should be evaluated, ~~as well as it should~~ the impact of such emissions on the performance of instrumentation and control systems important to safety.

4.59. Equipment and systems, including associated cables, should be designed ~~and,~~ installed and qualified to appropriately limit the propagation (both by radiation and by conduction) of electromagnetic interference to equipment at the research reactor. Special consideration should be given to areas where

equipment converges (e.g. containment penetrations, motor control centres, switchgear areas, cable spreading rooms, the control room). National and international industry standards for electromagnetic emissions should be referenced.

TESTING AND TESTABILITY OF INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

4.60. Arrangements for testing of instrumentation and control systems at a research reactor include: interfaces with test equipment, installed test equipment, and built-in test facilities and procedures. The design of instrumentation and control systems important to safety is required to include provisions that enable periodic testing (see Requirements 31 and 51 of SSR-3 [1]). Ideally, testing should be possible during reactor operation, or, if justified, during shutdown only. Many research reactors are operated on relatively short operating cycles and; therefore, provisions for testing during operation might not be necessary for such research reactors. Recommendations on periodic testing of research reactors are provided in DS509BSSG-81 [3].

Test provisions for instrumentation and control systems for a research reactor

4.61. The provisions for the testing of instrumentation and control systems and components important to safety should include the following:

- (a) The test provisions should have appropriate test interfaces and status indications. Test interfaces should include the capability to introduce simulated process conditions or electrical signals.
- (b) The test provisions should operate in such a manner that any faults in the equipment are readily detectable.
- (c) The systems should have features to prevent unauthorized access.
- (d) The systems should be located so that test equipment and the components to be tested are readily accessible.
- (e) The systems should be located so that neither the testing nor access to the testing location exposes operating personnel to hazardous environmental conditions. WhereIf the equipment to be tested is located in a hazardous areasarea, provisions for testing from outside the hazardous area should be considered in the design.
- (f) The systems should have communications facilities as necessary to support the tests.

4.62. It should be ensured in the design that systems cannot be unknowingly left in a test configuration. Inoperability or bypassing of safety system components or channels should be indicated in the control room. For frequently bypassed items, such indications should be auto-announcing.

4.63. Self-checking features of instrumentation and control systems important to safety are required to be used where practicable (see para. 6.183 of SSR-3 [1]). In meeting this requirement, it is necessary

to balance the provision of self-checking features against the need for simplicity in design (see para. 4.2).

4.64. Built-in test facilities should themselves be capable of being checked at regular intervals to ensure continued correct operation.

PreservingPreservation of control functions for instrumentation during testing

4.65. The testing of instrumentation and control systems at a research reactor should neither compromise the performance of a safety function, nor ~~should it~~ introduce the potential for common cause failures. Safety aspects should be considered prior to the testing of systems important to safety during operation.

4.66. Test facilities that are permanently connected to safety systems should be considered part of the safety systems. Installed test facilities should be tested independently on a regular basis against another calibrated source.

Considerations for the testing of instrumentation and control systems at a research reactor

4.67. Considerations for the testing of instrumentation and control systems at a research reactor should include the following:

- (a) Locating and installing sensors such that their testing and calibration can preferably be performed at their location, including ~~at those for~~ equipment for draining, drying, decontamination, isolation and ventilation, where applicable;
- (b) Locating test devices and test equipment in areas ~~with~~ that have sufficient space and that are convenient to the equipment to be tested;
- (c) ~~Provisions to ensure~~ Ensuring the safety of operating personnel during the test, including through measures to ~~deenergize~~ energize equipment and prevent its inadvertent use;
- (d) ~~The~~ Ensuring convenience ~~of~~ in the indications of component status and the test connections.

4.68. Communications equipment is necessary to support the testing of instrumentation and control systems at a research reactor. The design of instrumentation and control systems important to safety should include provisions to automatically alert (e.g. by the use of alarms) operating personnel that channels or components are in test mode.

4.69. When channels of safety systems are tested, the performance of the safety function should not be compromised. In particular, the single failure criterion should still be fulfilled; ~~for example,~~ channels of safety systems being tested should be placed in trip condition during the testing, when relevant, considering the ~~implemented~~ protection logic implemented in the system.

4.70. Any impacts that the tests on instrumentation and control systems might have on assumptions made in the safety analysis should be considered.

4.71. Administrative controls should be considered prior to performing on-line tests on safety systems ~~should be considered~~.

Test programme for instrumentation and control systems at a research reactor

4.72. The design of instrumentation and control systems should include the specification of a test and calibration programme. The scope and frequency of testing and calibration should be consistent with functional requirements and availability requirements (see also para. 7.72 of SSR-3 [1]). In determining the frequency of testing, the necessary accuracy and the stability of the instruments chosen should be taken into account. Stable instruments with low drift may need to be tested less frequently.

4.73. The test programme should include the following:

- (a) A description of the programme objectives;
- (b) An identification of the systems and channels to be tested;
- (c) The master test schedule;
- (d) The reasons and justification for the tests to be conducted and the test intervals;
- (e) A description of the test documentation and reports;
- (f) The arrangements for periodic review of the effectiveness of the programme;
- (g) A specification of the individual test procedures to be used in the conduct of tests.

4.74. The tests defined in the test programme should ensure by means of clear procedures that during the tests and after their completion the following are demonstrated:

- (a) The overall functional capabilities of the systems are not degraded;
- (b) The instrumentation and control systems continue to meet their design basis requirements for performance and reliability;
- (c) -The instrumentation and control systems are correctly returned to operation.

4.75. The test programme should arrange tests into a sequence such that the overall condition of the system or component under test can be assessed without, as far as practicable, further testing of other components or systems.

4.76. The test programme should define processes for periodic tests and calibration of instrumentation and control systems that have the following objectives:

- (a) To specify checks of all functions, from the sensors to the actuators, that are capable of being performed in situ and with a minimum of effort;

- (b) To confirm that functional requirements and performance requirements⁴ for the design basis are met by documenting the results of a test showing compliance with tolerance requirements;
- (c) To test all inputs and output functions, such as alarms, indicators, control actions and the operation of actuation devices;
- (d) To provide post-maintenance testing to ensure that systems are returned correctly to operation;
- (e) To ensure the safety of the research reactor during the ~~conduct of the~~ test;
- (f) To minimize the possibility of spurious initiation of any safety action and minimize any other adverse effect of the tests on the safety and availability of the research reactor.

4.77. The conduct of the test programme should not cause any deterioration of any system or component.

4.78. ~~Where~~If temporary connections of equipment are necessary for periodic testing or calibration, the reactor operator should be alerted by alarms and/or warning lights of the presence of the temporary connection, and the use of such equipment should be subject to appropriate administrative controls.

4.79. The temporary modification of computer codes in instrumentation and control systems for testing purposes should not be allowed.

4.80. The time interval for which equipment is removed from service should be minimized, and each sensor should be individually tested to the extent practicable.

4.81. Tests of safety system channels should preferably be single on-line tests. When a single on-line test is not practicable, the test programme may combine overlapping tests to achieve the test objectives. For tests of safety system channels, documented justification for the use of overlapping tests should be provided.

4.82. Tests of a safety system should independently confirm the functional requirements and the performance requirements of (a) each channel of sensing devices, and ~~of~~(b) command, execution and support functions.

4.83. Tests of a safety system should include as much of the function under test as practicable (including sensors and actuators), with due consideration of the wear on actuators when tested excessively.

4.84. Wherever possible, tests of a safety system should be accomplished under actual or simulated service conditions, including the sequence of operations. Precautions should be taken in testing safety systems that are especially sensitive.

⁴ Requirements for testing of the response time need to be strictly based on the assumptions made in the safety analysis report and need to be limited to parameters that involve special consideration for testing of the response time because their timely response is important to the safety of the facility.

4.85. After a failed test, the reasons for the failure, its root causes and the actions taken afterwards should be evaluated and documented before the results of a repeated test can be used to demonstrate the operability of the system or the component involved.

4.86. Corrective actions may include, for example, maintenance or repair of components, or changes to test procedures. If corrective actions are determined to be unnecessary, the reasons should be documented.

MAINTAINABILITY OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

4.87. Provisions for the maintenance of instrumentation and control systems are required to be considered in the design:– (see Requirement 31 of SSR-3 [17]). The design of instrumentation and control systems should include maintenance plans for all systems and components.

4.88. Instrumentation and control systems and components are required to be designed to avoid undue exposure of maintenance personnel:– (see para. 6.88 of SSR-3 [17]). The operating organization is also required to ensure that non-radiation-related risks are as low as reasonably achievable:– (see Requirement 80 of SSR-3 [17]). The design should also facilitate preventive maintenance, troubleshooting and timely repair.

4.89. Design measures to facilitate maintenance, troubleshooting and repair of instrumentation and control systems include the following:

- (a) Avoiding locating equipment in areas of extreme temperature, extreme humidity, ~~and/~~ or high radiation levels;
- (b) Taking into account human factors in performing maintenance activities;
- (c) Leaving sufficient space around the equipment to ensure that the maintenance personnel can perform their tasks using the necessary tools;
- (d) ~~The provision of~~ Providing test panels, instrument isolation ~~and draining~~ and test connections.

4.90. If components have to be located in inaccessible areas, other design measures should be considered, for example ~~as follows~~ the following:

- (a) The installation of spare redundant devices in cold or hot standby;
- (b) The provision of facilities for remote replacement, repair and return to service.

SAFETY ASSESSMENT OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

4.91. Safety analysis is required to support the design of a new instrumentation and control system or the modification of an existing system:– (see Requirement 41 of SSR-3 [17]). The following activities should be performed to confirm that instrumentation and control systems fulfil their design basis:

- (a) Confirmation that all known and predictable failure modes are either self-revealing or detectable by planned testing, and that the system is fail-safe, as appropriate.
- (b) Verification that the overall instrumentation and control system supports the application of the defence in depth concept at the research reactor.
- (c) Verification that the vulnerabilities to common cause failures of instrumentation and control systems important to safety ~~to common cause failures~~ are known and have been adequately addressed. Vulnerabilities to common cause failures may be dealt with by eliminating the vulnerabilities, by providing diverse means of fulfilling the safety functions that are subject to the common cause failures, or by justifying acceptance of the vulnerability.
- (d) Verification that design basis reliability requirements are met. This verification may be based on deterministic criteria and quantitative reliability analysis in which design features such as redundancy and testability, failure modes, mean time between failures, and rigour of qualification are considered. For complicated systems, a combination of qualitative analysis, quantitative analysis and testing is usually needed to verify compliance with design basis reliability requirements.
- (e) Verification that the design of instrumentation and control systems includes adequate provisions for testing.
- (f) ~~In determining~~ Determining system availability, and ensuring that test facilities that are part of the safety system ~~should be~~ regarded as permanently installed test equipment.
- (g) Confirmation of functional requirements for various operational modes of instrumentation and control systems. ~~This includes~~ Such confirmation would include analysis of correct system behaviour during commissioning, during first startup when the facility is not operating under normal conditions (e.g. following trips due to low flux with a fresh core)), and during normal operation, including following power interruptions and restart (or reboot) after the execution of tests.
- (h) Verification that the effects of failures of automatic control systems will not exceed the acceptance criteria established for anticipated operational occurrences.

4.92. The methodology for any safety analysis that is conducted should be thoroughly specified and should be documented, together with the inputs for the analysis, its results and the details of the analysis itself. Traceability analysis should be used to confirm implementation and validation requirements. Each assumption made for an analysis should be justified, and this justification should be documented.

4.93. Further recommendations on safety assessment for research reactors are provided in SSG-20 (Rev. 1) [9].

SAFETY SYSTEM SETTINGS FOR INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

4.94. The operational limits and conditions established in the design of a research reactor should include safety system settings for instrumentation and control systems. Recommendations on operational limits and conditions for research reactors are provided in [DS509DSSG-83](#) [5].

4.95. The following are usually considered in the determination of settings for instrumentation and control systems that are safety systems:

- (a) Safety limits: Limits on certain operational parameters within which operation of the research reactor has been shown to be safe.
- (b) Analytical limits (of safety system settings): Limits on a measured or calculated variable established by the safety analysis to ensure that a safety limit is not exceeded.
- (c) Allowable values: The limiting values of safety system settings, beyond which appropriate action needs to be taken. The allowable value for a particular safety system setting is the value at which it is acceptable for a trip to occur when periodically testing the corresponding channel. If the point at which a protective action would be initiated is found to be beyond the allowable value, corrective action should be taken.

Figure 2 illustrates the relationship between these terms and the types of measurement uncertainty that are normally considered in establishing safety system settings.

IDENTIFICATION AND VERIFICATION OF INSTRUMENTATION AND CONTROL SYSTEMS IMPORTANT TO SAFETY AT A RESEARCH REACTOR

4.96. The safety classification of instrumentation and control systems is required to be based primarily on deterministic safety analysis complemented by probabilistic safety assessment, where appropriate: (see para. 6.29 of SSR-3 [1-1]). This should be supported by engineering judgement provided by experts, including knowledgeable personnel from design organizations and operating organizations of a research reactor.

4.97. A consistent and coherent method of naming and identifying all instrumentation and control components should be determined and followed throughout the design, construction, installation, commissioning and operation stages in the lifetime of the research reactor as well as for the labelling of controls, displays and indications. Clear identification of components should be used to reduce the likelihood of inadvertently performing installation, modification, maintenance, tests, repair or calibration of an incorrect component. Components or modules mounted in equipment or an assembly that is clearly identified might not themselves need identification.

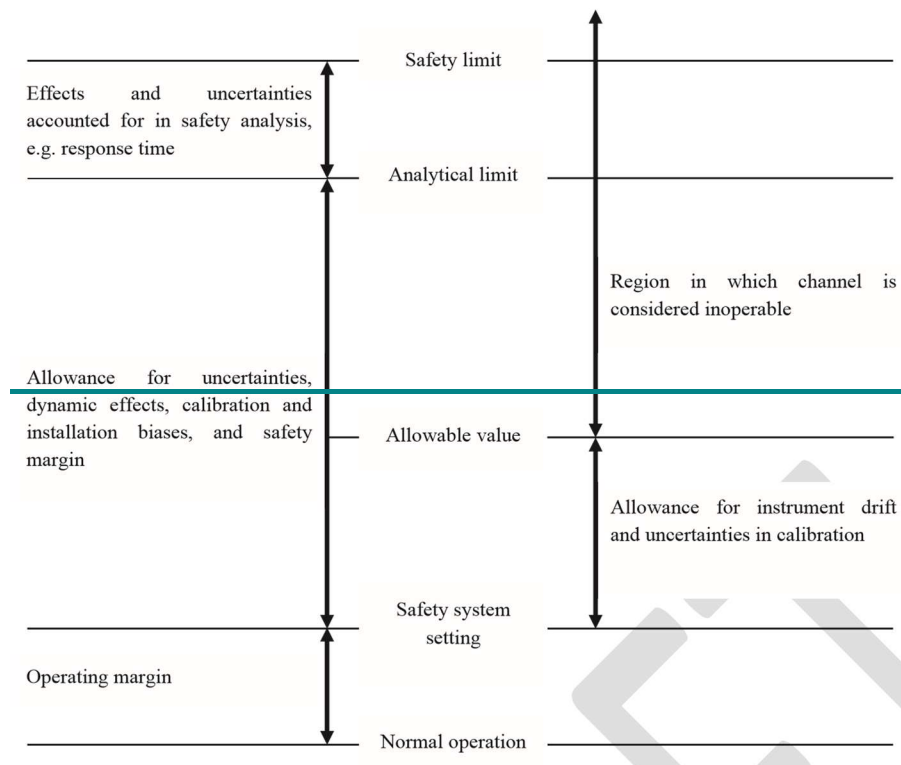


FIG. 2. Safety system setting terminology and uncertainties to be considered in determining safety system settings. *(to be revised to show “Region in which channel is may be considered inoperable”*

DESIGN OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR FOR DESIGN EXTENSION CONDITIONS

4.98. Paragraph 6.65 of SSR-3 [1] states:

“The design extension conditions shall be used to define the design specifications for safety features and for the design of all other items important to safety that are necessary for preventing such conditions from arising, or, if they do arise, for controlling them and mitigating their consequences.”

4.99. The instrumentation and control systems and equipment provided as additional safety features for design extension conditions should meet the design requirements for reliability, testability, maintainability, inspectability, ageing management and (to the extent practicable) equipment qualification. Such systems are considered to be items important to safety.

5. SYSTEM SPECIFIC DESIGN GUIDELINES FOR INSTRUMENTATION AND CONTROL SYSTEMS FOR A RESEARCH REACTOR

SENSING DEVICES FOR INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

5.1. Measurements of variables for a research reactor should be consistent with the requirements of the design basis. These measurements include both detection of the present value of a variable within a range, and detection of a discrete state such as is detected by limit switches or on/off switches (e.g. temperature, pressure, flow or level limit switches, switches for availability of the main supply or for operation of the control system, interlock on/off switches).

5.2. The measurements of variables may be made directly ~~or indirectly, such as~~ (e.g. calculation of the value by performing multiple measurements,) or indirectly by measuring other data ~~having that have~~ a known relationship to the desired variable. To the extent practicable, the reactor conditions should be monitored by direct measurements rather than being inferred from indirect measurements.

5.3. The sensor for each monitored variable and its range should be selected on the basis of the accuracy, response time and range needed to monitor the variable in normal operation and in accident conditions.

5.4. The vulnerability of sensing devices to common cause failure should be identified (e.g. saturation of radiation monitors), as they have the potential to deny reactor operators the information and parameters necessary to control and mitigate accident conditions.

5.5. If more than one sensor is necessary to cover the entire range of values for the reactor parameter being monitored, a reasonable amount of overlap from one sensor to another should be provided. Examples include source range, intermediate range and power range of neutron flux monitors.

5.6. If the monitored variables have a spatial dependence (i.e. if the measured value of a parameter depends ~~upon~~ the location of the sensor), the minimum number and locations of sensors, such as flow measurement elements, should be identified in the design and justified. The final locations also need to be tested to verify the design assumptions and to determine whether associated set points, limiting conditions and allowable values should be reassessed.

REACTOR PROTECTION SYSTEM

5.7. Requirement 50 of SSR-3 [1] states that: **“A protection system shall be provided for a research reactor to initiate automatic actions to actuate the safety systems necessary for achieving and maintaining a safe state.”** The design of this system is required to include provisions for achieving and maintaining a safe state even if the primary reactor protection system is subjected to a credible common

cause failure ~~(e.g., for example, hardware failure, failure due to human factors)~~ (see para. 6.177 of SSR-3 [17]).

5.8. Where applicable, the reactor protection system should comply with the recommendations provided in Section 4 of this Safety Guide for the design of instrumentation and control systems.

5.9. The reactor protection system should, at a minimum, include a function to initiate shutdown of the reactor. For subcritical assemblies, the shutdown may be achieved by withdrawing the neutron source. The reactor protection system may also provide other safety functions, such as initiation of emergency core cooling and confinement (the features of the reactor protection system thereby acting as extended engineered safety features of the instrumentation and control system).

5.10. The reactor protection system is required to be capable of automatic response, without the need for manual action within a short period of time, for the full range of postulated initiating events (see paras 6.173 and 6.174 of SSR-3 [17]).

5.11. As part of defence in depth and to cope with a potential common mode failure of the primary reactor protection system, the need for a second reactor protection system with all or some of the functions of the primary reactor protection system should be evaluated and assessed. Where two reactor protection systems are provided, these two systems should be independent and diverse from ~~one another~~ each other.

5.12. Actions initiated by the reactor protection system ~~is~~ should be latched so that once an action is initiated it will continue until its completion even if the initiating state is terminated (see para. 6.174 of SSR-3 [17]). Latching of actions initiated by the protection system is normally implemented at the level of actuation signals to reactor equipment, and such actions should not reduce the reliability of the system below an acceptable level.

5.13. In some cases, manual actions to initiate a protective action may be sufficient, provided that the diagnosis is simple, and the action is clearly defined, for example as follows:

- (a) The reactor operator has sufficient and clearly presented information to make valid judgements on the need to initiate the necessary safety actions.
- (b) The reactor operator is allowed sufficient time to evaluate the status of the research reactor and to complete the necessary actions.
- (c) The reactor operator is provided with sufficient means of control of the reactor to perform the necessary actions.

5.14. In addition to the automatic actions, means should be provided to manually initiate a reactor trip and any other safety actions of the reactor protection system. Such manual actions should act ~~as~~ ~~close~~ closely as practicable to the final actuation devices (e.g. reactor trip breakers) rather than being an input to the reactor protection system logic.

5.15. System functions that inhibit the tripping of the reactor protection system, including the means for activating and deactivating these functions, should be part of the reactor protection system. Sometimes, it is necessary to inhibit the actions of the reactor protection system to enable changes in reactor conditions. For example, the trips that limit reactor power during startup have to be inhibited at some point to enable power increase. Another example would be the necessity to inhibit certain actions in the case of pulsed operation of a research reactor. In this Safety Guide, such inhibit functions of the reactor protection system are called operational interlocks and are classified as components and/or functions of safety systems. The reactor protection system should prevent the enabling of an operational interlock when the applicable enabling conditions are not met. If actions are inhibited and conditions change so that this is no longer permissible, the reactor protection system should automatically disable the operational interlock, or initiate appropriate protective actions. (See also para. 6.175 of SSR-3 [1].)

5.16. Where temporary connections of equipment are needed for periodic testing or calibration of the reactor protective system, the operator should be alerted by alarms and/or warning lights of the presence of the temporary connection, and use of such equipment should be subject to appropriate interlocks or administrative controls to ensure that it is removed after use.

5.17. The design should ensure that safety system settings can be established with a margin between the initiation point and the safety limits where the action initiated by the reactor protection system will be able to control the process before the safety limit is reached. In addition, the following should be taken into account in selecting such margins:

- (a) Inaccuracy of instrumentation;
- (b) Uncertainty in calibration;
- (c) Instrument drift;
- (d) Instrument and system response time.

5.18. If a computer based system is intended to be used in a reactor protection system, the life cycle of the system is required to be systematically documented (see para. 6.180(b) of SSR-3 [1]), and independent verification and validation processes should be applied.

5.19. Diverse means of ensuring fulfilment of the protective actions should be considered. The diversity may be provided in the following ways:

- (a) Within the reactor protection system itself, or by a separate and independent system, provided that the design bases are met;
- (b) By a diverse system using distinct technology, which may be hard wired or computer based as long as adequate diversity can be justified.

5.20. Paragraph 6.180(c) of SSR-3 [1] states:

“In order to confirm the reliability of the computer based systems, a systematic, fully documented and reviewed assessment shall be undertaken by expert personnel who are independent of the designers and the suppliers.”

5.21. For computer based reactor protection systems and components, the design is required to include computer security features:– (see para. 6.180(d) of SSR-3 [1]–~~See also~~ and paras 4.37–4.45 of this Safety Guide:–).

OTHER INSTRUMENTATION AND CONTROL SYSTEMS IMPORTANT TO SAFETY OF A RESEARCH REACTOR

5.22. The reactor operator should be provided with sufficient instrumentation for monitoring the operation of the reactor systems during normal operation (including shutdown, refuelling and maintenance) and accident conditions, including the recording of all parameters important to safety.

5.23. The need for suitable startup neutron sources and dedicated startup instrumentation should be taken into account in the design.

5.24. The operating organization should establish a set of design requirements and limitations on the normal operation of the instrumentation and control systems necessary for the safe operation of the research reactor. These design requirements and limitations should address the following:

- (a) The information necessary to establish the safety limits and safety system settings;
- (b) Control system constraints and procedural constraints on process variables and other important parameters;
- (c) Maintenance, testing and inspection of the research reactor to ensure that systems, structures and components function as intended;
- (d) Clearly defined operating configurations, including operational restrictions in the event of safety system outages;
- (e) Considerations for research related tasks.

These design requirements and limitations are the bases for establishing the operational limits and conditions under which the research reactor is authorized to operate.

DESIGN OF RESEARCH REACTOR CONTROL ROOMS

5.25. In the main control room of a research reactor, measures are required to be implemented to protect the occupants against hazardous conditions:– (see para. 6.185 of SSR-3 [1]). Similar provisions should be made for a supplementary control room (see Requirement 54 of SSR-3 [1]) and other areas where operating personnel are expected to monitor and control the research reactor systems. In all these locations, the aim should be to ensure satisfactory conditions in the working environment.

5.26. Ergonomic principles and human factors are required to be considered in the design of control rooms: (see para. 6.104 of SSR-3 [1]). The results of task analysis should also be considered in the design of control rooms.

5.27. To meet Requirement 39 of SSR-3 [1], the design of control rooms should include adequate provisions for preventing unauthorized access and use.

5.28. The main control room is required to be designed and constructed to resist internal and external hazards that might challenge its continued operation, in particular fires: (see paras 6.185 and 6.186 of SSR-3 [1]). At least one control room (main, or supplementary) should be designed and constructed to resist design basis earthquakes.

Main control room at a research reactor

5.29. The principal location for safety actions and safety related control actions is the main control room. Requirement 53 of SSR-3 [1] states:

“A control room shall be provided at a research reactor facility, from which the reactor facility can be safely operated in all its operational states, either automatically or manually, and from which measures can be taken to maintain the research reactor in a safe state or to bring it back into a safe state after anticipated operational occurrences and accident conditions.”

Supplementary control room at a research reactor

5.30. A remote capability for reactor shutdown should be provided if the safety analysis identifies events that could inhibit the reactor operator's ability to shut down the reactor and to maintain it in a safe condition from the main control room. A supplementary control room or an emergency control console should be provided if reactor operators need to perform safety actions and the safety analysis identifies events where during which the main control room could be unavailable or operations from the main control room could be inhibited. Events that could inhibit the reactor operator's ability to shut down the reactor from the control room include, for example, a fire in the control room or a fire in a location that affects connections between the control room and devices elsewhere in the research reactor.

5.31. The instrumentation and control systems of the supplementary control room are required to be appropriately independent from the main control room (see Requirement 54 of SSR-3 [1]) to avoid common cause failures diminishing the operability of the systems of the supplementary control room. For example, the design of control system networking should ensure that there is an extremely low likelihood of being unable to use the system from either of the two control rooms. Another example is the separation of electrical power supplies for the control rooms.

5.32. A suitable provision outside the main control room for transferring priority control of the reactor to a new location and for isolating the equipment in the main control room if it is abandoned should be considered.

CONTROL SYSTEMS FOR IRRADIATION FACILITIES AND EXPERIMENTAL DEVICES AT A RESEARCH REACTOR

5.33. In many research reactors, there are special control consoles for irradiation facilities and experimental devices, which may be located in the main control room ~~and~~ or in other rooms. The operator of experimental devices should have communications links with the reactor operator to share information on experiments and on the reactor status and to make each other aware of expected actions (e.g. situations that necessitate the shutdown of the reactor).

5.34. The control consoles for irradiation facilities and experimental devices should be devoted exclusively to ~~the~~ those irradiation facilities and experimental devices if the safety analysis identifies events that show an independent instrumentation and control system is necessary for irradiation facilities and experimental devices to ensure a functional separation from other activities at the research reactor.

5.35. Parameters important to the operation of the reactor should be covered by the alarm system. Alarms of experimental devices, with no reactor safety implications, should be presented with ~~a~~ functional separation from the reactor alarms.

VOICE COMMUNICATION SYSTEM AT A RESEARCH REACTOR

5.36. Communication systems are required to be provided between the main control room and the supplementary control room (if applicable) and the emergency centre ~~+~~ (see paras 6.185 and 6.91 of SSR-3 [1]). Secure means of communication should also be provided with other locations within the facility, including with the operators of experimental devices and associated facilities, and with off-site response organizations.

5.37. Both the main control room and the supplementary control room should have at least two diverse communications links with the following:

- (a) On-site locations where communications are needed in anticipated operational occurrences and accident conditions;
- (b) Off-site emergency response organizations;
- (c) Other facilities that might be affected by the operation of the research reactor.

These communications links should be routed so that they will not all be affected by loss of the primary communications links, whatever the cause of this loss (including external events), and they should be capable of operating independently of both the research reactor power supply and the off-site power supply.

5.38. Paragraph 6.189 of SSR-3 [1] requires the provision of relevant information and means of communication from the control ~~room(s)~~ rooms to the relevant emergency response facilities. The emergency response facilities are also required to be provided with environmental monitoring

information from sources outside the research reactor. The information and communication links are required to be designed for operation during accident conditions: (see para. 7.93 of SSR-3 [1]).

FIRE DETECTION SYSTEMS AND FIRE EXTINGUISHING SYSTEMS AT A RESEARCH REACTOR

5.39. Requirement 61 of SSR-3 [1] states:

“Fire protection systems for a research reactor facility, including fire detection systems and fire extinguishing systems ... shall be provided throughout the research reactor facility, with due account taken of the results of the fire hazard analysis.”

The nature of the fire alarm system, its layout, the necessary response time and the characteristics of its detectors should be determined on the basis of the fire hazard analysis. The fire detection system should provide a warning by means of audible and visual fire alarms in the control room of the specific location of the fire.

5.40. Local audible and visual fire alarms, as appropriate, should also be provided in areas of the facility that are usually occupied. Fire alarms should be distinctive to avoid confusion with any other alarms at the facility.

5.41. The fire detection and alarm system is required to be operational at all times (see para. 6.207 of SSR-3 [1]) and should be provided with ~~non-interruptible~~uninterruptible emergency power supplies, including fire resistant cables where necessary.

5.42. Fire detectors should be located so that the flow of air due to ventilation or pressure differences that are necessary for contamination control will not cause smoke or heat energy to flow away from the detectors and thus unduly delay actuation of the detector alarm.

5.43. If the environment does not allow detectors to be placed in the area to be protected (e.g. because of increased radiation levels or high temperatures), alternative methods should be considered, such as the sampling of the gaseous atmosphere by remote detectors with automatic operation.

5.44. When items such as fire pumps, water spray systems, ventilation equipment, fire dampers and the corresponding power supplies are controlled or used by fire detection systems, and ~~where~~when spurious operation would be detrimental to the research reactor and site personnel, operation should be controlled by two diverse means of fire detection operating in series. The design should allow the operation of the system to be stopped if the actuation is confirmed to be spurious. The potential effects on the research reactor of the spurious operation of the system should also be considered; for example, gas suppression systems may be a good alternative to water sprinkler systems for rooms containing power systems and instrumentation and control systems.

5.45. The following measures should be implemented for the wiring of fire detection systems, alarm systems and actuation systems:

- (a) The wiring should be protected from the effects of fire by a suitable choice of cable type, by proper routing or by other means.
- (b) The wiring should be protected from mechanical damage.
- (c) The wiring should be continuously monitored for integrity and functionality.

5.46. The fire detection system and fire extinguishing systems should be periodically tested.

5.47. National requirements for fire protection should also be used as inputs to the design of fire detection systems and fire extinguishing systems.

POWER SUPPLIES OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

5.48. The power supplies for instrumentation and control systems (see Requirement 56 of SSR-3 [1]) should be classified and qualified, and should include provisions for reliability, isolation, testability, maintainability and indication of removal from service that are consistent with the design basis of the instrumentation and control systems ~~that~~ they serve. In addition, failure modes for these power supplies should be considered.

5.49. Instrumentation and control systems that need to be available for use at all times, in operational states and/or in accident conditions, are required to be connected to uninterruptible power supplies: (see para. 6.191 of SSR-3 [1]). Such power supplies should provide the instrumentation and control systems with power within the tolerances specified by the design basis for the systems. These tolerances should be specified for the instrumentation and control systems to ensure that the systems can withstand failures in the normal power supply as well as a facility blackout caused by an external event.

5.50. Power supplies can provide a transmission path for electromagnetic interference that might originate outside the instrumentation and control systems or might arise from other instrumentation and control systems that are connected directly or indirectly to the same power supply. Such origins of interference include electrical fault clearance associated with other equipment on the same supply. These interferences should be analysed and should be avoided to the extent possible.

6. OPERATION OF INSTRUMENTATION AND CONTROL SYSTEMS DURING THE OPERATION OF A RESEARCH REACTOR

INSTRUMENTATION AND CONTROL SYSTEMS AND OPERATIONAL LIMITS AND CONDITIONS AT A RESEARCH REACTOR

6.1. Requirement 71 of SSR-3 [1] states that “**The operating organization for a research reactor facility shall ensure that the research reactor is operated in accordance with the operational limits and conditions.**”

6.2. The instrumentation and control systems of the research reactor help to ensure that the reactor's operating parameters are kept within the operational limits and conditions. Recommendations on operational limits and conditions for research reactors are provided in [DS509DSSG-83](#) [5].

Safety limits for a research reactor

6.3. The instrumentation and control systems at a research reactor should function to prevent safety limits from being exceeded in operational states, in design basis accidents and in design extension conditions.

Safety system settings for a research reactor

6.4. Each parameter for which an analytical limit (see para. 4.9495) is required to actuate a safety system, and any other important safety related parameters, should be monitored by an instrumentation and control system. Where appropriate, the system should provide a signal that can be [utilized](#) to automatically prevent that parameter from exceeding the set limit.

Limiting conditions for safe operation for a research reactor

6.5. Acceptable margins between normal operating values and the safety system settings should be applied in the instrumentation and control systems to ensure safe operation of the reactor, while avoiding the frequent actuation of safety systems. Acceptable margins should allow for instrument accuracy, system response time, expected drift and allowable margin of error in measured signals and for all expected variations in normal operation.

CONTROL OF ACCESS TO INSTRUMENTATION AND CONTROL SYSTEMS IMPORTANT TO SAFETY AT A RESEARCH REACTOR

6.6. Requirement 39 of SSR-3 [1] states that **“Unauthorized access to, or interference with, items important to safety at a research reactor facility, including computer hardware and software, shall be prevented.”** All reasonable precautions should be taken to prevent persons from performing unauthorized actions that could jeopardize safety when accessing instrumentation and control systems or [when](#) performing tasks on instrumentation and control systems.

6.7. Access to instrumentation and control systems classified as important to safety should be controlled. Access control methods should include physical restrictions or barriers, special embedded devices, and restrictions on access to functions important to safety by means of hardware or software access keys, access alarms and administrative controls.

6.8. Wherever safety system settings are configured, only authorized personnel should be allowed to change these settings, and these settings should be checked for their integrity. Access to the safety system settings and calibration adjustments should be restricted by physical and administrative means.

6.9. The protection of computer based components of instrumentation and control systems needs to be addressed in appropriate security procedures that take into account national requirements. Guidance on computer security is provided in Refs [15–17].

6.10. Secure storage arrangements and procedural controls should be used to ensure that only authorized software versions are uploaded into instrumentation and control systems and equipment at the research reactor. Records of software versions should be maintained within the management system for the research reactor. The correct performance of the computer based system should be demonstrated before it is returned to service.

6.11. Electronic access to the software and data of computer based systems important to safety via network connections from outside the reactor should be prohibited.

6.12. Access control methods should be used to ensure that users can only access data and commands for which they have been authorized.

6.13. Suitable measures should be implemented to prevent unauthorized access to software, the use or corruption of software or data, the introduction of malicious code, unauthorized connections to external networks or other computer based attacks.

MAINTENANCE, PERIODIC TESTING, AND INSPECTION OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

6.14. Maintenance, periodic testing, and inspection are required to be conducted on instrumentation and control systems important to safety to ensure that all their components function in accordance with the design requirements and intent, and in compliance with the operating limits and conditions: (see para. 7.68 of SSR-3 [1]). The frequency of such activities is required to be consistent with the reliability requirements for such systems or components: (see para. 7.72 of SSR-3 [1]). Further recommendations are provided in DS509BSSG-81 [3].

6.15. The instrumentation and control systems at a research reactor should facilitate periodic testing and, where applicable, include on-line testing functions to reduce the number of insertions and extractions of connectors, and thereby improve the system reliability.

PROVISIONS FOR REMOVAL FROM SERVICE OF INSTRUMENTATION AND CONTROL SYSTEMS FOR TESTING OR MAINTENANCE

6.16. Removal from service of any single safety system, component or channel should not result in loss of the required minimum redundancy unless the acceptably reliable operation of the system can be adequately demonstrated.

6.17. If use of equipment for testing or maintenance can impair an instrumentation and control function, the interfaces should be subject to hardware interlocking to ensure that interaction with the test or maintenance system is not possible without deliberate manual intervention.

6.18. For safety systems, design features should ensure that during periodic tests of part of a safety system, those parts remaining in service can continue to perform the required safety ~~function(s).~~functions. For example, tripping the redundancy during the testing of a ‘two out of three’ logic leaves the system in a ‘one out of two’ logic arrangement. Administrative controls on the availability of safety systems should keep ~~them~~those systems in operation within the design basis.

INSTRUMENTATION AND CONTROL SYSTEMS DURING EXTENDED SHUTDOWN OF A RESEARCH REACTOR

6.19. The operating organization should assess and define the minimum instrumentation and control systems important to safety that need to be kept operational during an extended shutdown.

7. HUMAN FACTORS ENGINEERING AND HUMAN–MACHINE INTERFACES AT A RESEARCH REACTOR

7.1. An effective process for human factors engineering is required to be embedded into the overall design process for every aspect of the design ~~– (see Requirement 35 of SSR-3 [17]).~~ Appropriate design standards and guidelines should be identified and should be used throughout the design process. Guidance on human factors engineering aspects is provided in Ref. [2019].

7.2. Verification and validation activities ~~related~~relating to human factors engineering should be performed throughout the design process for instrumentation and control systems for a research reactor to confirm that the design adequately accommodates all necessary actions by operating personnel and all relevant administrative arrangements of the operating organization. This verification and validation could include task analysis and consideration of factors such as timing ~~and~~, human cognition and perception, operator overload, and available indications for the operator response.

7.3. If a part of an instrumentation and control system is modified, careful consideration should be given to the design of the modernized part of the system and to its compatibility with ~~and~~ human interaction with the existing systems.

CRITERIA FOR HUMAN FACTORS ENGINEERING AND DESIGN OF HUMAN–MACHINE INTERFACES AT A RESEARCH REACTOR

7.4. The design of human–machine interfaces should take into account operating experience to retain useful features and avoid problems with human factors engineering. Such design considerations should be taken into account in the system architecture for new projects as well as for modification projects. The design of human–machine interfaces should emphasize the importance of human factors and of machine features and take both into account.

7.5. Instrumentation and control functions necessary to achieve the safety objectives for the research reactor should be identified, and the human resources and system resources necessary to fulfil these functions should be allocated in accordance with a specified method and should be taken into account in the system architecture at the design stage.

7.6. All human-machine interfaces should be designed in accordance with ergonomic criteria. The operating organization should determine which information is most effectively displayed using conventional displays (e.g. panel instruments, alarm annunciators) and which information is most effectively displayed using video screens. In the design criteria for information displays and controls, the different roles and authorized access levels of operating personnel — such as reactor operators, maintenance personnel, systems managers and personnel with responsibilities in an emergency — should be taken into account.

7.7. Design requirements for human-machine interfaces should be specified on the basis of all the tasks to be supported by the interfaces, including during normal operation and during anticipated operational occurrences and accident conditions. ~~This~~ The tasks considered should include ~~tasks~~ those undertaken by reactor operators, maintenance staff, experimenters and personnel with responsibilities in an emergency.

7.8. In accordance with Requirement 49 of SSR-3 [1], the instrumentation and control systems are required to provide the information necessary to detect changes in the status of systems and to make decisions for accident management. The human-machine interfaces are required to include the instrumentation and control systems necessary to assess the general state of the research reactor in any condition, to confirm that automatic safety actions are implemented as intended and to determine both the need for and the time for manual safety actions:— (see para. 6.105 of SSR-3 [1]).

7.9. During operation of the research reactor, the reactor operator should be provided with suitable warnings or alarms when the facility is approaching a state in which operational interlocks should be enabled or should be disabled.

7.10. The reactor operator should be provided with sufficient indicators and recording instrumentation to be able to monitor relevant reactor parameters ~~in~~ during, and following, anticipated operational occurrences and accident conditions.

7.11. Audible and visual alarm systems should be used to provide an early indication of changes in the operating conditions of the reactor if these changes in the operating conditions could affect ~~its~~ safety.

7.12. Careful attention should be paid in the design of human-machine interfaces to ensure that reactor operators would not be overwhelmed by large amounts of data that could be difficult to assimilate owing to the inherent limitations ~~on~~ of human perception, cognition and memory. This is particularly important for the design of alarms.

7.13. In the design of the instrumentation and control system, due account should be taken of the time periods necessary for reactor operators to perform their expected tasks.

7.14. The instrumentation and control system should protect against human errors by implementing range limits, interlocks or trips to protect the research reactor from unsafe operation.

7.15. Where a safety function is performed automatically, the instrumentation and control system should provide reactor operators with the necessary information to monitor the performance of the function. The information should be provided at a rate and to a level of detail that reactor operators can monitor effectively.

7.16. The instrumentation and control system should alert the reactor operator of a failure of an automatic control system.

7.17. The presentation of information should be harmonized to facilitate understanding of the status of the research reactor and of the activities necessary to ensure safety.

7.18. The operation and appearance of the human-machine interfaces in different locations within the research reactor should be consistent, should reflect a high degree of standardization and should be consistent with operating procedures and the training provided to operating personnel.

7.19. The human-machine interfaces should include the capability to display recorded information where such information will help reactor operators identify patterns and trends, ~~to~~ understand the past or present state of the system, or ~~to~~ predict its future progression. The design of display screens should ensure that a frozen display is easily identified.

HUMAN FACTORS ENGINEERING FOR RESEARCH REACTOR CONTROL ROOMS

7.20. Ergonomic principles are required be taken into account in the design of the research reactor control room:—(see para. 6.104 of SSR-3 [17]). Human factors considerations such as workload, possibility of human error, operator response time and minimization of the physical and mental effort of reactor operators should be taken into account ~~in order~~ to facilitate the execution of the necessary operating procedures in order to ensure safety in all operational states and accident conditions.

7.21. Acceptable working environments in the control room — in terms of radiation levels, lighting, temperature, humidity, noise, dust and vibration — should be ensured for normal operation, anticipated operational occurrences and accident conditions. The design of the main control room is required to take into account conditions resulting from internal hazards (e.g. fire ~~or~~, smoke, toxic substances in the atmosphere) and external hazards (e.g. earthquakes, flooding, extreme meteorological conditions, hazards due to human error~~;-~~)(see paras 6.185 and 6.186 of SSR-3 [17]). Similar provisions should be made for the supplementary control room, if applicable.

7.22. The layout of instrumentation and the means of presenting information to operating personnel, with both an adequate overall summary of the status and performance of the research reactor and, where

necessary, detailed information on the status and performance of particular systems or equipment, should be considered in the design.

7.23. The information displayed in the main control room should allow reactor operators to achieve the following:

- (a) To take specific manually controlled actions for which no automatic control is provided;
- (b) To confirm the availability of safety functions and the performance of automatic safety actions;
- (c) To determine the potential for the breach of a fission product barrier or to detect such a breach;
- (d) To confirm the performance of safety systems, auxiliary supporting features and other systems necessary for ~~the mitigation of~~mitigating accident conditions or for achieving and maintaining a safe state;
- (e) To determine the magnitude of any radioactive releases and to continually assess such releases.

7.24. The parameters displayed in the supplementary control room may differ from those displayed in the main control room if the supplementary control room does not need to be used to respond to the same range of anticipated operational occurrences and accident conditions as the main control room. For the supplementary control room, sufficient instrumentation and control equipment is required to be available so that, in the event of a loss of ability to perform essential safety functions from the main control room, the research reactor can be placed and maintained in a safe state, residual heat can be removed, confinement functions can be performed and the essential facility parameters can be monitored ~~in the event of a loss of ability to perform essential safety functions from the main control room~~: (see para. 6.188 of SSR-3 [1]). The instrumentation and control equipment in the supplementary control room should be physically and electrically separate from the equipment in the main control room.

8. COMPUTER BASED SYSTEMS AND SOFTWARE AT A RESEARCH REACTOR

8.1. Requirement 52 of SSR-3 [1] states:

“If a system important to safety at a research reactor is dependent upon computer based equipment, appropriate standards and practices for the development and testing of computer hardware and software shall be established and implemented throughout the lifetime of the system, and in particular throughout the software development cycle.”

8.2. Computer based systems are of increasing importance to safety in research reactors as their use is increasing in both new and older facilities. Such systems are used both in safety related applications, such as process control systems and monitoring systems, and in safety applications, such as the reactor protection system.

8.3. Paragraph 6.184~~(e) and (d)~~ of SSR-3 [1] states:

“For computer based equipment in safety systems and systems important to safety...:

.....

(c) An assessment of the equipment shall be undertaken by experts who are independent of the design team and the supplier team to provide assurance of its high reliability.

(d) When the necessary high reliability of the equipment cannot be demonstrated with a high level of confidence, diverse means of ensuring fulfilment of the safety functions shall be provided”.

The reliability of computer based systems should be evaluated in accordance with a systematic, fully documented and reviewed engineering process. This process should include the evaluation of new software and pre-existing software. Relevant operating experience may be used to evaluate pre-existing software.

8.4. Design faults, including software faults, are systematic in nature and not random, and therefore potential common cause failures are required to be considered (see para. 6.184(e) of SSR-3 [1]). In particular, safety systems employing redundant subsystems using identical copies of the hardware and software should be systematically considered.

8.5. Depending on the complexity of experimental devices in the research reactor, consideration should be given to having separate computer based instrumentation and control systems for the reactor and for experiments. In such cases, each system could be provided with its own set of design requirements and objectives.

8.6. Obsolescence management should be taken into account in the design and operation of computer based systems to plan for and manage reductions in service life, diminishing manufacturing sources and material shortages.

DESIGN CONSIDERATIONS FOR COMPUTER BASED SYSTEMS AND SOFTWARE AT A RESEARCH REACTOR

8.7. For computer based safety systems, complexity should be avoided, both in the functionality of the system and in its implementation, by the use of a structured design that follows a formal software development life cycle. Concepts such as ‘top-down’ decomposition, levels of abstraction and modular structure are important for coping with complexity. The logic behind the system modularization and the definition of interfaces should be made as simple as possible. A ‘top-down’ design process (i.e. breaking the system down to gain insight into its subsystems) for the system and its associated software should be used to facilitate the assessment of whether design objectives are being achieved.

8.8. With regard to safety systems at a research reactor, the functional requirements fulfilled by a computer system should all be essential to the fulfilment of safety functions. Functions not essential to safety should be isolated to avoid any impact on safety functions.

8.9. When a computer system incorporates two or more components in different safety classes, the computer system should meet the requirements of the higher safety class.

8.10. The use of diverse functions and system components at different levels of the design of computer based systems should be considered.

8.11. System fail-safe features, self-supervision and fault tolerant mechanisms should be incorporated into the software, but only to the extent that the additional complexity is justified by design basis functional requirements and performance requirements. Fault detection and self-supervision features should not adversely affect the ability of a computer system to perform its safety function or cause spurious actions.

8.12. Paragraph 6.184(f) of SSR-3 [1] states:

“For computer based equipment in safety systems and systems important to safety—
: ...Protection shall be provided against accidental disruption of, or deliberate interference with, system operation (computer based systems and communication and network systems important to safety, including the reactor protection system, are to be adequately protected against cyber-attacks, up to and including the design basis threat.”...).”

It should be demonstrated that measures have been taken to protect a computer based system throughout its entire life cycle against physical and computer attacks, unauthorized access, fraud, computer viruses and other malicious threats. Safety systems should not be connected to external networks. The connections for external storage devices should be locked to prevent unauthorized use.

8.13. A computer based system at a research reactor should be designed for maintainability to facilitate the detection, ~~locating~~location and diagnosis of potential or actual failures so that the system can be repaired or replaced efficiently. Software that has a modular structure may be easier to repair, review and analyse, since the design may be easier to understand. Maintainability of software also includes the concept of making changes to the functionality. The design of a computer based system should allow, as far as practicable, ~~that~~for changes ~~are to be~~ confined to a small part of the software.

8.14. Computer based systems that perform safety functions should have deterministic behaviour with regard to functions and timing, meaning that any given input sequence that is within the specification of the system will always produce the same outputs within the same response time.

8.15. Sample rates and processing speed should be consistent with requirements for accuracy and timing.

8.16. Data communications channels important to safety should satisfy the recommendations for independence in para. 4.21. The design should ensure that errors and failures of transmission equipment and data communications equipment are detected ~~and~~, that suitable alarms are provided for operating personnel and that records are made for the analysis of performance.

8.17. The communications technology for a research reactor should be chosen and configured to ensure that it is capable of a timely response under all possible conditions of data loading.

8.18. Appropriate consideration should be given to the use of redundancy in data communications.

8.19. The topology and network interface of the data communications network should be designed and implemented, from both an electrical and a communication protocol perspective, to avoid common cause failures of independent systems or subsystems.

8.20. Data flow from lower classified safety systems to higher classified safety systems should be avoided unless a decoupling device is installed, and a decoupling protocol is used.

8.21. The selection of ~~pre-developed~~predeveloped items to be included in computer based systems at a research reactor should follow a defined and documented process to guarantee their suitability.

8.22. Software tools can be used to support the life cycle of instrumentation and control systems. These tools should be verified and assessed in accordance with the reliability requirements, with the type of tool, and with the potential for the software tool to introduce errors or to fail in detecting existing errors.

PROJECT PLANNING FOR COMPUTER BASED SYSTEMS AND SOFTWARE AT A RESEARCH REACTOR

8.23. The project development process for computer based systems and software at a research reactor should be carefully planned, and clear evidence should be provided that the process has been followed to facilitate the independent assessment of systems important to safety. The development plan should identify and define the process that will be used on the particular project. Other aspects of the project that should be considered and planned are quality management, verification and validation, configuration management, installation, and commissioning.

8.24. All phases of the development process for computer based systems and software at a research reactor should be identified. The design activity of preceding phases should provide the inputs for the next phase. Verification should be performed across each phase of the development process.

8.25. The methods to be used in the development process should be identified. The selection of methods should be in accordance with a quality management programme in which processes and procedures are established. The quality management programme should be prepared and implemented before the project begins. A software quality management plan should be available at the start of the project.

Verification and validation of computer based systems and software at a research reactor

8.26. Paragraph 6.184(g) of SSR-3 [1] states that “For computer based equipment in safety systems and systems important to safety: ...Appropriate verification and validation and testing of the software systems shall be performed.” Verification and validation activities should be performed to demonstrate that the computer system achieves its overall safety objectives and functional requirements. Specific techniques and validation procedures should be included in the verification and validation plan.

8.27. Verification and validation planning should include the listing and collection of applicable codes and standards, procedures, and conventions that guide the verification process.

8.28. The teams performing verification and validation should be independent of the development team. Independence is usually ensured by having a different line management for the verification and validation teams and for the development teams. Personnel performing verification and validation should not have been involved in the design of the same project. A different organization could be used to complete the verification and validation activities.

8.29. The verification and validation plan should include a mechanism for recording all instances of non-compliance found during the analysis and ensuring that they are properly resolved by means of an approved change control process.

Configuration management of computer based systems and software at a research reactor

8.30. All items ~~related~~relating to software development, such as compilers, development tools, configuration files and operating systems, should be subject to configuration management. All identifiable items, such as documents, components of the software or data structures, should be given a unique identification, including the version number. These items should include both developed items and existing items that are being reused or reapplied.

8.31. A procedure for change control should be established. This procedure should provide for the keeping of records of any problems identified during the development process, or during the operation of the research reactor, that necessitated changes to computer based systems or software. ~~It~~The procedure should also ~~provide for documenting~~include documentation of the following: problems and their analysis; items that were affected; changes that were made to solve the problem; and which versions (~~such as e.g.~~ version numbers of software ~~or~~ components of software) and which baseline database of systems and components of the instrumentation and control systems were produced as a result. The change control procedure should specify responsibilities for approving changes.

Installation and commissioning of computer based systems and software at a research reactor

8.32. The installation and commissioning plan for computer based systems and software at a research reactor should cover the following:

- (a) The sequence of steps for proper integration of the system into the research reactor and the corresponding facility states necessary for the safe introduction of the new or changed system;
- (b) The required interactions with the regulatory body, including approvals, hold points and reports, that are necessary before the system can be put into operation;
- (c) The commissioning test cases and sequence and the corresponding facility states necessary to confirm proper functioning of the system in the environmental conditions at the research reactor;

- (d) A description of the records and reports that will be necessary to describe the results of commissioning.

SPECIFICATION OF COMPUTER BASED SYSTEM REQUIREMENTS AT A RESEARCH REACTOR

8.33. The specifications for a computer system for a research reactor should include, at a minimum, the necessary functional properties and non-functional properties of the computer system. Safety analyses (e.g. facility safety analyses, transient analyses, and accident analyses, based on postulated initiating events and safety criteria) should be used in specifying the functional requirements. Other requirements not directly associated with safety, such as requirements for availability or security, should be included at this stage of the design.

8.34. A safety analysis should be made for safety systems and safety related systems to determine the functional requirements for these systems.

8.35. The specification of non-functional requirements should include the following:

- (a) The relevant dependability attributes, such as reliability, availability and safety performance of the system;
- (b) The security requirements for the computer based systems, including security procedures;
- (c) Performance requirements (e.g. the response time for performing safety functions);
- (d) Environmental qualification requirements, such as for temperature and radiation;
- (e) Whether and where physical separation is needed (e.g. between safety functions and control functions);
- (f) A confirmation that requirements not directly associated with safety (e.g. requirements for availability or security) will not adversely affect the ability to perform a safety function.

An accurate and clear description of these requirements should be formulated before starting the next stage of the project, and this description should be subject to independent review.

SOFTWARE REQUIREMENTS FOR A RESEARCH REACTOR

8.36. The specification of software requirements should define the function(s)functions of each individual software item and how it will interact with other items of the system. Software requirements for a research reactor should be complete, unambiguous, consistent, clear and understandable to their target audience (e.g. domain experts, safety engineers, software designers), as well as verifiable and traceable. The origin of every software requirement should be sufficiently documented to facilitate verification, validation, and traceability to higher-level documents and to demonstrate that all software requirements have been addressed.

8.37. The software requirements should include a description of the allocation of system requirements to the software, with a consideration of potential failure conditions, functional requirements in each mode of operation, performance criteria, timing and constraints, detection of failures, logging of events, self-supervision, monitoring of safety, and interfaces with nuclear security.

SOFTWARE DESIGN FOR A RESEARCH REACTOR

8.38. In the software of systems important to safety at a research reactor, unnecessary complexity should be avoided at all levels of design. The simpler the design is, the easier it is to achieve and demonstrate all other attributes. ~~It~~[Having a simpler design](#) also gives greater confidence that the software is fully understood.

8.39. To facilitate tracing of compliance ~~of~~[with](#) software requirements through the design, each design element, ~~—~~ such as a software module, a procedure, a subroutine or a file, ~~—~~ should have a unique identifier.

8.40. The design of software for a research reactor should contain no contradictions and no ambiguities. The description of the interfaces between modules should be complete. In addition to internal interfaces between modules of the software, the design should explicitly specify the external interfaces of the software, such as hardware interfaces and libraries. The design and its description should demonstrate that each software design requirement has been met and should enable verification that the implementation is correct with respect to the detailed design.

8.41. The documentation of software design should provide technical information on the software architecture and on the detailed design of all software modules, and of their interactions, to demonstrate that they collectively behave as specified in all possible cases. This ~~includes~~[information should include](#) non-functional aspects such as timings and resource use. Relevant constraints on implementation should also be specified.

8.42. Each software module identified in the software architecture should be described in the detailed design. Diagrams and flow charts can be used, provided ~~that~~ the meaning of the elements of the diagrams is well defined. Common techniques ~~used~~ for describing design include data flow diagrams, ~~and~~ structure diagrams or graphics.

8.43. Review should be conducted at the software design stage to avoid potential errors and to assess the software quality.

SOFTWARE IMPLEMENTATION FOR A RESEARCH REACTOR

8.44. The production of software code should be verifiable against the software design requirements. The code should be readable, adequately commented and understandable. Validated software tools can be used to facilitate the code verification process.

8.45. A formal system for requesting changes and controlling modifications to software for a research reactor should be used during the implementation phase to deal with omissions and inconsistencies. Up to date records of changes should be kept available for reviews and audits.

8.46. The code of each computer program should be kept simple and easy to understand, both in its general structure and in its details. Data structures and their naming conventions should be used uniformly throughout the whole computer based system.

VERIFICATION AND ANALYSIS OF SOFTWARE FOR A RESEARCH REACTOR

8.47. Techniques for verification and analysis should be used to provide assurance of software quality and compliance with instrumentation and control system requirements.

8.48. A software verification plan should be produced that documents the following:

- (a) The verification techniques to be used;
- (b) Details of, or references to, the procedures to be used in applying each technique, including its scope and depth;
- (c) How non-functional requirements and constraints will be demonstrated to be met;
- (d) The criteria for determining when sufficient verification has taken place, including targets for completeness with respect to the outputs of the previous phase and for structural coverage of the functional tests, and how these will be demonstrated;
- (e) The means by which results will be recorded;
- (f) The means by which non-compliance and faults will be recorded and resolved;
- (g) The team(s) performing the verification and their independence from the designers of the software (see para. 8.50);
- (h) The functionality of any software tool for verification, including expectations and limitations on how it is to be used (e.g. domain, language, process);
- (i) The rationale for each of the elements listed in items (a)–(h) above, and justification that the verification will be sufficient for software in the system of the safety class to which it is applied.

8.49. A software test plan should be developed covering all testing to be done, including unit level tests, integration tests, factory acceptance tests and installation tests.

8.50. Verification should be performed by teams, individuals or organizational groups that are independent of the designers and developers.

8.51. The software code should be reviewed to check for security vulnerabilities using automated software tools and complemented by manual review of the critical sections of the code (e.g. input/output handling, exception handling).

8.52. All outputs of the instrumentation and control system should be monitored during the verification of the software code, and any deviation from the expected results should be investigated and documented.

8.53. Any shortfall in the verification results against the verification plan (e.g. in terms of the test coverage achieved) should be documented and resolved or justified.

8.54. Any errors detected should be analysed for cause and should be corrected by means of agreed modification procedures and regression tested, as appropriate, to ensure that previously developed and tested software still performs correctly after a change. The error analysis should include an evaluation of applicability to other parts of the instrumentation and control systems.

8.55. Records of the number and types of anomalies should be maintained. These records should be reviewed to determine whether ~~or not~~ any lessons ~~could~~can be learned and whether appropriate process improvements should be made.

8.56. Techniques such as reviews, inspections or audits should be applied to the verification of all phases in the life cycle of computer based systems and software. The method by which the results of the reviews, inspections or audits are to be recorded should be stated in the verification plan, together with a justification of the method used.

8.57. Review of the documentation on software design and software implementation should be performed. Software test cases should be designed on a functional basis only (i.e. without knowing the structure of the software under test) to preserve the independence of the testing activity and to enable analysis of the structural coverage of the tests performed after their design. The test case specifications should be fully documented and reviewed.

8.58. Test plans should be designed to facilitate regression testing by ensuring that tests are repeatable and involve minimal human intervention.

8.59. Any anomalies in the performance of testing should be reviewed and, if it is determined that there is a need for a modification to the test procedure, an appropriate procedure for change control should be applied.

8.60. Any anomaly in the software test performance should be documented in a report that includes the nature of the problem, the identified corrective action, the retest arrangements and ~~the completion~~confirmation of a satisfactory retest. In addition, a cross-reference record of software corrections and software builds should be maintained for configuration management of the installed software.

THIRD PARTY ASSESSMENT OF SOFTWARE FOR A RESEARCH REACTOR

8.61. A third party assessment of software for safety systems at a research reactor should be conducted concurrently with the software development process.

8.62. The objective of such a third party assessment is to provide a view on the adequacy of the system and its software that is independent of both the ~~suppliers~~suppliers of the system and ~~or~~ software and the

operating organization. Such an assessment may be undertaken by the regulatory body or by a body acceptable to the regulatory body.

8.63. It is important that proper arrangements ~~are be~~ made with the software originator to permit third party assessment.

8.64. The assessment should involve an examination of the following:

- (a) The development process (e.g. through quality assurance audits and technical inspections, including examination of life cycle documents, such as plans, software design requirements and the full scope of test activities);
- (b) The final software (e.g. through static analysis, inspection, audit and testing), including any subsequent modifications.

COMPUTER SYSTEM INTEGRATION AT A RESEARCH REACTOR

8.65. The software version integrated into the computer system of a research reactor should be the latest version to have been verified and validated.

8.66. The computer system integration phase should encompass at least three sequenced activities: software ~~test~~testing; hardware testing and integration; and hardware–software integration. The hardware–software integration should consist of three parts: loading of all software into the hardware system; testing that the software–hardware interface requirements are satisfied; and testing that all the software can operate in the integrated software–hardware environment.

8.67. During the verification of the computer system, evidence should be generated to demonstrate that the system integration has been properly checked and verified. A traceability analysis should be performed and documented as part of this verification to demonstrate that the system integration requirements are complete with respect to the design specification for the computer system.

Integrated computer system tests at a research reactor

8.68. The integrated computer system tests should be performed before the system is transferred to the research reactor and installed. The final integrated computer system test is often combined with the factory acceptance test to form a single test activity.

8.69. In constructing test cases, special consideration should be given to the following:

- (a) Coverage of all design requirements for the hardware and software (including robustness tests and features);
- (b) Coverage of the full ~~range~~range of values for input signals;
- (c) Handling of exceptions (e.g. demonstration of acceptable behaviour when input failure occurs);
- (d) Timing related requirements (e.g. response time, input signal scanning, synchronization);
- (e) Accuracy;

- (f) All interfaces (e.g. the hardware–software interface in system integration ~~and~~, external interfaces during validation);
- (g) Stress testing and load testing;
- (h) Requirements for security (e.g. logging of user activities);
- (i) All modes of operation of the computer system, including transition between modes and recovery after failure of the power supply.

8.70. A traceability analysis should be performed to demonstrate that the validation process for testing and evaluation is complete.

Validation and commissioning tests of computer based systems and software at a research reactor

8.71. Validation and commissioning tests should be performed to verify that the computer system has been correctly connected at the research reactor and to confirm the correct functioning of the system.

8.72. The validation and commissioning tests should usually be combined with the site acceptance test, which includes verification of the operation of the equipment.

8.73. Strict configuration control of the computer system (both hardware and software) should be maintained during the commissioning programme. Any changes made should be subject to a formally documented change process.

8.74. Sufficient documentation should be produced to demonstrate the adequacy of the commissioning programme for the installed computer based system and software.

Operation, maintenance and modification of computer based systems and software at a research reactor

8.75. The following activities should be considered with regard to the operation, maintenance ~~and/or~~ modification of computer based systems and software at a research reactor:

- (a) Periodic tests, performed to verify that the systems are not degrading;
- (b) Regression testing because of modifications; performed to enhance or change the functionality or to correct errors;
- (c) Change of operating parameters;
- (d) Diagnostic activities, for example the execution of special diagnostic programmes;
- (e) Replacement of hardware components because of failures.

8.76. All software tools used in software development, testing, installation, integration, operation and maintenance should be qualified.

8.77. The life cycle of the systems should include the processes for implementing modifications. This life cycle should include the main phases of ~~the~~ development, including verification and validation.

These activities, together with an impact analysis and regression testing, will be necessary to ensure that the modifications have been correctly implemented and no new errors have been introduced.

8.78. After failure of a hardware component, corrective actions should be limited to one-by-one replacements of hardware components and to the reloading of the existing software modules. These actions should not include any modification unless analysis of the failed components reveals a need for modification.

9. CONFIGURATION MANAGEMENT OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

9.1. A full set of documents reflecting the configuration and status of instrumentation and control systems at the research reactor should be available prior to the commissioning of the facility. The documentation should be kept up to date throughout the lifetime of the research reactor.

9.2. A baseline database of systems and components of the instrumentation and control systems should be established and should include the following information:

- (a) General information (e.g. system identification, serial number, manufacturer, supplier support, location, safety class);
- (b) System summary (e.g. functions, configuration, impacts of the system on safety, current performance, loss of operational availability in the event of unavailability of the system, interfaces, documentation);
- (c) Physical characteristics (e.g. number of cabinets, detailed component inventory, operational limits);
- (d) Boundaries (e.g. environmental conditions, power supply, grounding, power supply margins necessary in the cabinets and rooms, the amount of information exchanged with other systems);
- (e) System constraints (e.g. licensing conditions, technical specifications, design constraints, operating characteristics);
- (f) Obsolescence issues (e.g. maintenance costs, replacement parts, performance degradation);
- (g) Measures for improvements (e.g. functionality, configuration, performance, maintenance);
- (h) Supporting references.

9.3. Reactor operators and maintenance personnel should collaborate in the improvement and the updating of documentation for configuration management of instrumentation and control systems. The documentation and database described in paras 9.1 and 9.2 should be considered to be sensitive

information and should be protected in accordance with national requirements for security of information.

10. MODIFICATION AND MODERNIZATION OF INSTRUMENTATION AND CONTROL SYSTEMS AT A RESEARCH REACTOR

10.1. The main reasons to modernize instrumentation and control systems at a research reactor are obsolescence of the existing systems (e.g. as a result of the unavailability of spare parts) or an increased failure rate of the systems. These developments can lead to frequent reactor shutdowns and long repair periods, resulting in increasing unavailability of the research reactor. Recommendations on ageing management for research reactors are provided in SSG-10 (Rev. 1) [8].

10.2. ~~Additional reasons~~An additional reason for modernization ~~are~~is the technological progress in instrumentation and control systems, which can provide greater reliability, improved human–system interfaces, and more extensive and faster data collection and data processing capabilities. Other factors (e.g. new regulatory requirements) may also influence the final decision for modernization of the instrumentation and control systems at a research reactor.

10.3. Before modernizing an instrumentation and control system, information on current limitations and future needs should be collected using experience from the existing system. This includes information from past failures and incidents, which should be documented by event recording systems used at the research reactor. Other weaknesses could be identified from the self-assessment of operational performance, including analysis of even small deviations from normal operation. In addition, possible future problems should be considered, and the potential impact of these problems should be assessed.

10.4. The modification and modernization of instrumentation and control systems should be performed in accordance with the recommendations provided in SSG-24 (Rev. 1) [10], on planning, organizational aspects, safety assessment, implementation and post-implementation aspects, training, and documentation of modifications to the research reactor. Vigorous independent verification and validation should be performed for every change associated with the modification and modernization of instrumentation and control systems at a research reactor.

10.5. A modification to an instrumentation and control system might include a complete replacement of the components of the system. Modifications to existing systems ~~that are~~important to safety are required to follow the same procedures for design, construction and commissioning that were applied to the original equipment—(see para. 7.101 of SSR-3 [1]). Recommendations on the design of instrumentation and control systems are provided in Section 4 of this Safety Guide.

10.6. Modification of instrumentation and control equipment is expected over the lifetime of ~~the~~^a research reactor. In each case, consideration should be given to the function of the equipment being modified⁷ and the possible impacts of changing from one technology to another. This may include changing from an analogue system to a digital system or replacing an instrumentation and control system because of a lack of spare parts (i.e. obsolescence).

10.7. When a decision is made to implement a modification to an instrumentation and control system important to safety, the possible effects on the safety of the research reactor are required to be considered and assessed⁸ ~~(see para. 7.101 of SSR-3 [1].)~~.

10.8. The operating organization should verify that every modification to an instrumentation and control system has been properly assessed, documented and reported in terms of its potential effects on safety⁷ and should ensure that the reactor is not restarted without formal approval after the completion of modifications.

10.9. The design documentation for older instrumentation and control systems might be incomplete or inaccurate. Consequently, major modifications to, or replacement of, such systems might involve some degree of 'reverse engineering' to recreate the original design bases and specifications. A full set of documentation reflecting the present configuration of the instrumentation and control systems at the research reactor should be made available.

10.10. A process of verification and updating of the existing documentation in accordance with para. 7.99(e) of SSR-3 [1] should be undertaken prior to commencing any modernization of instrumentation and control systems at a research reactor. Reactor operators and maintenance personnel should collaborate on the updating of this documentation to ensure that all modernization activities are captured in the documentation on configuration management of the systems (see Section 9). The baseline database of instrumentation and control systems described in para. 9.2 should also be updated.

10.11. Verification and updating of existing documentation should start with a high level functional description of the instrumentation and control system architecture, preferably in the form of a diagrammatic representation with an accompanying list of all instrumentation and control systems.

10.12. The operating organization should designate a designer to be responsible for design, integration, documentation and maintenance of modifications to instrumentation and control systems, as well as for training of operating personnel in the use of the new systems. Reference [¹⁹20] provides details on the responsibilities ~~that~~⁹ the designated designer should assume.

10.13. For modifications to an instrumentation and control system, the duties and ~~the~~¹⁰ responsibilities of the operating personnel and personnel with responsibilities in an emergency should be taken into consideration. The effects of the modifications on the interactions of these persons with the systems should be taken into consideration so as to achieve an effective human-machine interface. Section 7 provides recommendations on human factors engineering and the human-machine interface.

10.14. The reliability of new or modified equipment should be taken into consideration, as ~~well~~ ~~asshould~~ the effects of the modification on overall system reliability. Performance of a qualitative analysis (e.g. analysis of failure modes and their effects) may be helpful in determining which parts of the instrumentation and control system might be affected by the modifications and the implications in terms of the ability of the system to perform its safety function.

10.15. In modifying an instrumentation and control system, the effect on the implementation of the concepts of defence in depth and independence should be considered. Safety systems are required to be independent, as far as practicable, of other reactor systems ~~÷~~ (see Requirement 27 of SSR-3 [17]). In modifying an instrumentation and control system, the development of design guidelines should be considered.

10.16. The effects of complex modifications to an instrumentation and control system will be more difficult to analyse. Careful consideration should be given to the addition of any new functions and to the ability to expand the capabilities of the existing system in the future.

10.17. The qualification of modifications for the expected environmental conditions of the system should be considered. Modifications should be qualified for the relevant service conditions (including the operational environment), and the qualification programme should be based on the safety analysis of the proposed modifications.

10.18. Procedures for change control for instrumentation and control systems at a research reactor should be established, including appropriate procedures and organizational structures for the review and approval of the safety aspects of the modification.

10.19. The following should be considered in the design of upgrades and modifications to instrumentation and control systems:

- (a) Any limitations due to the physical characteristics of the research reactor that might restrict the design options for instrumentation and control systems;
- (b) The need to maintain consistency between the design of replacement equipment and that of existing instrumentation and control systems (e.g. to reduce the complexity of the overall human-machine interface and the maintenance of the system);
- (c) The commercial availability of equipment or technology and the future availability of technical support by manufacturers or third parties for the service life of the equipment.

10.20. The benefits of modifications should be weighed against potential negative consequences for safety, and this assessment should be documented as part of the justification for the changes. Compensatory actions should be implemented to anticipate and prevent negative safety consequences.

10.21. Appropriate training on modified instrumentation and control systems should be provided to relevant personnel to minimize or ~~to~~ eliminate the potential for errors.

10.22. The consequences of updating or changing a software tool may be significant and should be subject to an impact assessment (e.g. the upgrade of a compiler could invalidate the results of previous analysis or verification concerning the adequacy of the compiler).

10.23. Modifications to instrumentation and control systems at a research reactor are required to be performed by qualified personnel: (see para. 7.99(d) of SSR-3 [17]). The work should be performed under the supervision of the designer and with the authorization of the reactor manager.

10.24. Once complete, and before startup of the research reactor, the installation should be functionally tested in accordance with the recommendations provided in SSG-10 (Rev. 1) [8].

10.25. When an instrumentation and control system is modified or is part of an upgrade, the effort applied in justifying and executing the change should be commensurate with the role and function of the system in ensuring the safety of the research reactor, taking into account both the existing systems and any systems that will remain in operation after the modification or the upgrade. This also applies to software based systems.

10.26. When an instrumentation and control system is replaced, the new instrumentation and control system may, where appropriate, be run in parallel with the old system for a probationary period, until sufficient confidence has been gained in the adequacy of the new system. In this configuration, only the old instrumentation system should be able to control the reactor. During this probationary period, the response of the drivers of the new instrumentation and control system should be registered in an independent acquisition system to assess and compare their responses against the responses of the old system.

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Annex

INSTRUMENTATION AND CONTROL SYSTEMS THAT CAN BE USED IN A RESEARCH REACTOR

A-1. The instrumentation and control systems of a research reactor involve many systems ~~that, which~~ may differ depending on the type of reactor, ~~the~~its purpose and its modes of operation (see Section 2). This [annex](#) identifies instrumentation and control systems, and relevant architecture, that can be used in a research reactor. Some of these instrumentation and control systems might not be used in a particular research reactor if they are not necessary for that specific type of facility.

A-2. Instrumentation and control systems can be designed in a ‘top-down’ architecture (see Fig. A-1) ~~having that has~~ different levels of monitoring, ~~and~~ processing, ~~and of~~ sensors and actuators. The design of the different architectural levels can be used to reduce the likelihood of dependent failures at each level. In this architecture, the monitoring functions are normally located at the supervision level; the calculation, algorithms, and safety and process functions are located at the control level; and sensors and actuators are located at the field level.

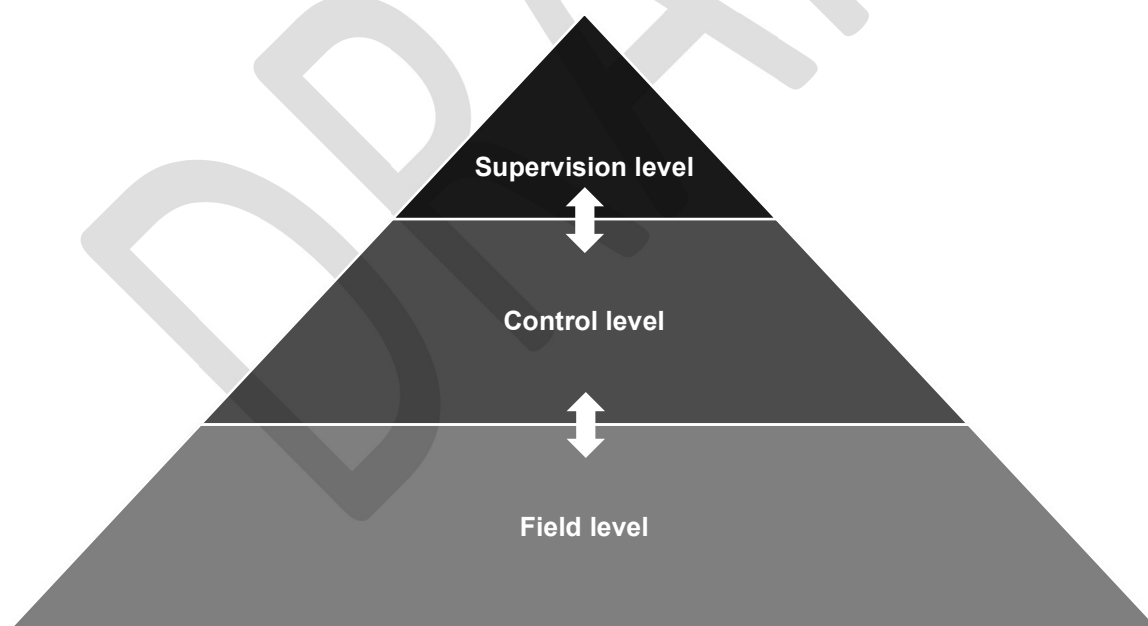


FIG. A-1. ‘Top-down’ architecture of instrumentation and control systems.

A-3. The typical instrumentation and control systems and their ~~interrelations~~interrelationships are shown in Fig. A-2.

DESCRIPTION OF THE MAIN INSTRUMENTATION AND CONTROL SYSTEMS AT A

REACTOR PROTECTION SYSTEM

A-4. The reactor protection system is a set of components designed to monitor reactor operating parameters (e.g. neutron power and period, coolant flow rate, inlet and outlet temperatures, pressure drop in the reactor core) to compare them with safety system settings and automatically initiate action of the reactor shutdown system when the parameters reach or exceed the safety system settings. Each parameter may be measured by two or more independent channels. The automatic actions are initiated on the basis that the logic arrangement for the initiation of protective actions complies with the single failure criterion. When three independent channels are available, the logic arrangement of two out of three channels may be used to prevent the initiation of protective actions by spurious signals. A reactor protection system could also be actuated manually by the reactor operator, by the experimenters or by the control and monitoring system for irradiation and experimental devices. A trip of the reactor protection system results in the shutdown of the reactor.

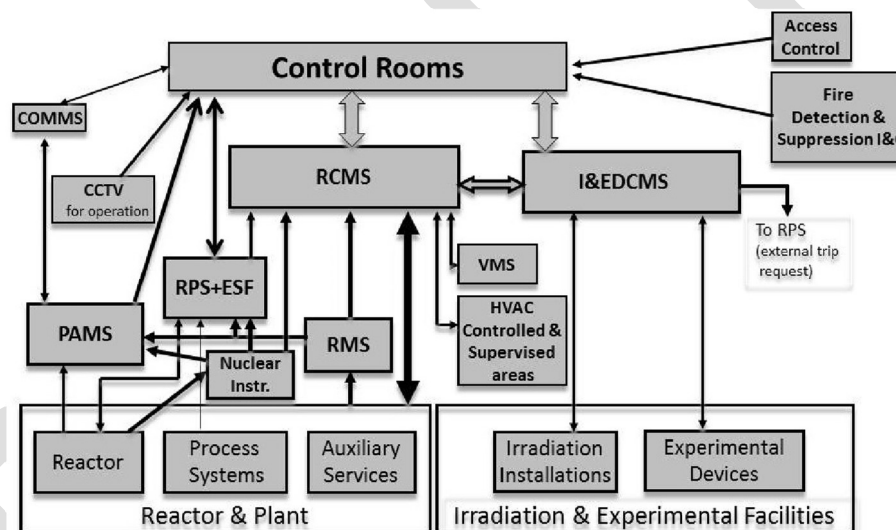


FIG. A-2. Research reactor instrumentation and control systems. Note: CCTV — closed circuit television; COMMS — communication system; ESF — instrumentation and control system for the initiation of other engineered safety features; HVAC — heating, ventilation and air-conditioning for controlled and supervised areas; I&C — instrumentation and control; I&EDCMS — control and monitoring system for irradiation facilities and experimental devices; Instr. — instrumentation; PAMS — post-accident monitoring system; RCMS — reactor control and monitoring system; RMS — radiation monitoring system; RPS — reactor protection system; VMS — vibration monitoring system.

~~FIG. A-2. Research reactor instrumentation and control systems.~~

INSTRUMENTATION AND CONTROL SYSTEM FOR THE INITIATION OF OTHER

ENGINEERED SAFETY FEATURES AT A RESEARCH REACTOR

A-5. The instrumentation and control system for the initiation of engineered safety features is a set of components designed to initiate, upon request, action of the emergency core cooling system, the decay heat removal system, the confinement isolation system and the confinement heat removal system. The system may also be actuated manually by the operator. A trip in the engineered safety features results in the initiation of one or more of the actions mentioned above. The functions of the engineered safety features may be included in the reactor protection system.

POST-ACCIDENT MONITORING SYSTEM AT A RESEARCH REACTOR

A-6. A post-accident monitoring system is an important feature of for research reactors. Its purpose is to provide the reactor operators and their backup teams with the information necessary for accident management and to ensure that this information is reliable. Under accident conditions, reactor operators may need information for the following purposes:

- (a) To perform preplanned manual control actions for which no automatic control system is provided and which are necessary to prevent or to mitigate the consequences of the accident. Such actions, specified in the safety analysis report, are described in the accident management procedures.
- (b) To determine whether important safety functions relating to reactivity control, core cooling, integrity of the reactor coolant system, the heat sink, containment integrity and surveillance for radioactivity are challenged, and whether they are being fulfilled by the reactor protection system, the engineered safety features and their essential support systems.
- (c) To relay information to, and to communicate effectively with, emergency response facilities on the site.

NUCLEAR INSTRUMENTATION AT A RESEARCH REACTOR

A-7. The nuclear instrumentation follows the value and evolution of the neutron flux of the reactor in all its operational states, since this parameter is of most relevance to ensuring the safe operation of the reactor. The nuclear instrumentation also provides the means to establish a suitable control strategy for starting up the reactor and keeping it in stable operation at different power levels.

REACTOR CONTROL AND MONITORING SYSTEM

A-8. The reactor control and monitoring system is intended for the reliable monitoring of the performance and the safe operation of the reactor. The reactor control and monitoring system provides startup and automatic adjustment of power, compensates for fuel burnup and provides interlocks for safe operation. The reactor control and monitoring system is built using fail-safe and redundant devices to receive and process signals from a large number of sensors, to actuate the corresponding control drivers and to present information on the reactor's status for the reactor operator at the main control console of the research reactor (the main human-machine interface).

A-9. The process instrumentation (e.g. detectors, sensors ~~and~~, switches), which ~~measure~~measures process parameters and the actual state (position) of actuators, and which ~~are~~is connected to the reactor control and monitoring system, ~~reside~~resides at the field level of the instrumentation and control systems.

Radiation monitoring system at a research reactor

A-10. The radiation monitoring system is designed for continuous radiation monitoring of the research reactor, as well as of surrounding areas ~~for detecting, to detect~~ the possible release of radioactive material. Such a release might arise, ~~for example~~, because of failures of technical equipment, loss of integrity of protective barriers, loss of effectiveness of water purification systems, loss of confinement isolation or failure of filters and ventilation systems.

HEATING, VENTILATION AND AIR-CONDITIONING SYSTEMS AT A RESEARCH REACTOR

A-11. Heating, ventilation and air-conditioning systems are used to ensure and maintain adequate environmental conditions for both personnel and equipment by providing temperature control and air quality control. The ventilation system also helps in maintaining the radiological conditions, for example by means of pressure gradients and the use of appropriate filters. Modern electronic equipment generates much less heat than older ~~types:equipment;~~ nevertheless, excess temperature can still degrade performance. Air-conditioning, as a means of removing excess heat from instrumentation and control systems that are safety systems, needs to meet the design requirements specified for safety system support features. In regions with a tropical climate or high levels of humidity, the proper design of ventilation systems (with physical separation, redundancy and a closed cycle) might be the only way to eliminate a source of potential common mode failure in instrumentation and control systems. In some facilities, the reactor control and monitoring system has the capability to send commands remotely to the heating, ventilation and air-conditioning systems (i.e. a command for the remote trip of the emergency ventilation system).

VIBRATION MONITORING SYSTEM AT A RESEARCH REACTOR

A-12. The vibration monitoring system provides a means of monitoring and detecting abnormal vibrations in the main rotary equipment of the reactor. The reactor control and monitoring system is used to transmit information from the vibration monitoring system to the control room.

CONTROL ROOMS FOR A RESEARCH REACTOR

A-13. Sufficient controls, indications, alarms and displays are provided in the main control room to initiate, supervise and monitor normal reactor operation and reactor shutdown to a safe state and to provide assurance that a safe state has been reached and is being maintained.

A-14. The minimum provisions for the main control room, including the human-machine interfaces, have to consider the need for the reactor operator to do the following:

- (a) ~~To operate~~Operate the reactor safely in all its operational states;
- (b) ~~To monitor~~Monitor the safe operation of the reactor;
- (c) ~~To monitor~~Monitor the appearance of alarms;
- (d) ~~To perform~~Perform and confirm a controlled shutdown;
- (e) ~~To actuate~~Actuate safety related systems;
- (f) ~~To perform~~Perform and confirm a reactor trip;
- (g) ~~To perform~~Perform and confirm the actuation of the engineered safety features;
- (h) ~~To monitor~~Monitor the status of fission product barriers;
- (i) ~~To keep~~Keep the reactor in a safe shutdown mode;
- (j) ~~To implement~~Implement emergency operating procedures.

A-15. The alarm annunciators show the status of systems. Safety systems have audible and visual alarms on the reactor operator's control console or control panel to provide a warning of any violation of the operational limits and conditions. Reactor operators can access all signals through the main control console of the reactor control and monitoring system. Control consoles and displays for the irradiation facilities and experimental devices are usually located in the main control room.

A-16. The supplementary control room, where applicable, provides a possibility for remote shutdown of the reactor in the event that this cannot be done from the main control room. Sufficient controls, indications, alarms and displays are provided in the supplementary control room to be able to initiate, supervise and monitor a reactor shutdown to a safe state and to provide assurance that a safe state has been reached and is being maintained.

CONTROL AND MONITORING SYSTEM FOR IRRADIATION FACILITIES AND EXPERIMENTAL DEVICES AT A RESEARCH REACTOR

A-17. The primary use of a research reactor is ~~for~~ the production of neutrons, for research purposes and for the neutron irradiation of materials. Irradiation facilities include the equipment that is used to place, move and arrange samples. A dedicated and tailored instrumentation and control system is designed to control and monitor these operations. Irradiation facilities and experimental devices might have an impact on the safe operation of the research reactor. The parameters of the experimental devices that affect the safety of the reactor are displayed in the main control room. Trip signals from the control and monitoring system for irradiation facilities and experimental devices to the reactor protection system can also be provided as indicated by the safety analysis.

COMMUNICATION SYSTEMS AT A RESEARCH REACTOR

A-18. Communication systems are the link between the operating personnel in the main control room and supplementary control rooms, the reactor hall and the process areas, the persons using the irradiation

facilities and experimental devices, and other internal locations (e.g. alarm stations) within the facility, and [for are the link to](#) off-site emergency response organizations. A voice announcement system is used for making announcements that can be heard by all on-site personnel, or to report an emergency or unforeseen circumstances that necessitate an immediate response.

CLOSED CIRCUIT TELEVISION SYSTEMS AT A RESEARCH REACTOR

A-19. Closed circuit television is a useful aid that allows the reactor operator to monitor and supervise relevant operational or maintenance tasks or activities that are being executed by the operating personnel and can also be used for monitoring the security status of the facility.

INSTRUMENTATION AND CONTROL SYSTEMS FOR THE DETECTION AND SUPPRESSION OF FIRES AT A RESEARCH REACTOR

A-20. The instrumentation and control system for the detection and suppression of fires is an independent system that has the capability to detect a fire at the research reactor and then initiate automatic fire suppression systems in the affected areas. Fire detection panels are located in the control rooms to provide the reactor operators with relevant information.

ACCESS CONTROL SYSTEM AT A RESEARCH REACTOR

A-21. The access control system is part of the physical protection system for a research reactor and has the capability to oversee and manage the movement of the personnel at the facility. Access control panels may be located in the control rooms and/or in the central alarm stations to provide the reactor operators with relevant information.

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