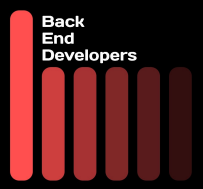
Hazard Analysis Mechatronics Engineering



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Table 1: Revision History

Date	Developer(s)	Change
2022-10-19	Back End Developers	Initial documentation for Hazard Analysis
2023-03-15	Jessica Bae	Minor improvements and proof reading for revision 1
2023-03-17	Jessica Bae	Removed and modified requirements to align with VnV
2023-04-03	Jonathan Hai	Incorporated TA comments
2023-04-04	Jessica Bae	Added logo and style to the document

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1 Introduction

In today's society, technology and engineering solutions are simply expected to work. The multitude of complex engineering designs and systems created to meet the needs of the world are expected to be infallible in the eyes of the public. When failures do occur, society considers them it shocking. When these failures happen in engineering systems critical to an aspect of safety, health, or some other critical role, people can die. In many cases, these failures are predictable and preventable in the design phase. The causes of such failures are known as hazards.

1.1 Purpose of Hazard Analysis

It is therefore necessary for engineers to perform extensive and thorough assessments of the systems they design in the aim to eliminate as many failures as reasonably possible. This is process is called hazard analysis.

More formally, hazard analysis is a step in the process to assess risk within an engineering system. Its aim is to identify and assess the potential conditions which may cause failure. These hazards can exist and cause failures alone, or in combination with other hazards or conditions. Once completed, a hazard analysis should provide a comprehensive assessment of the hazards within a system according to the system's components and boundaries, the assumptions critical in performing judgments regarding hazards and the scenarios they may occur in, and the requirements necessary to ensure that these hazards will be mitigated within the realm of reasonable possibility.

2 Scope

The purpose of this document is to perform this hazard analysis on the system to be designed by the Back End Developers. This document will first provide a description of the system boundaries and components of the system on an abstract level (both the hardware and software components), and then will list and justify the assumptions made in order to perform this hazard analysis. These assumptions will be kept to a minimum, in the hope to reduce the number of potentially overlooked hazards. The document will then describe the Failure Mode and Effect Analysis (FMEA) done by the team of the Back End Developers, and then detail the specific safety and security requirements that have been discovered in the process of performing the hazard analysis. The document will end with a roadmap describing the steps which will be taken in order to implement the novel discovered requirements, the timeline in which they will be implemented, and what considerations must be made regarding said requirements.

3 System Boundaries and Components

The system consists of several components that make up the entire system.

3.1 Battery/Power Management system

This component facilitates stepping down/up the source voltage from the battery to the necessary values required by different parts of the device. Moreover, it consists of a charge protection circuit for battery protection (Over-voltage and Discharge). Finally present is a battery level indicator that will generate an alert should the battery life fall below a certain threshold.

3.2 Sensor Array system

This component represents all the various sensors that will be used to collect the state information about the user. Also consists of various filters to facilitate smooth and accurate data collection.

3.3 Prompt generation system

This component handles all prompt generation, from the detection of when a prompt occurs, to its specific creation and finally its display on the screen.

3.4 Display System

This system manages all functionality of the device's display, such as prompt display, showing basic user feedback such as date, time, temperature, etc.

3.5 Data Storage system

This system handles all logging and storage of data collected by both the sensor array and the prompt generator. Data is stored along with an indication of which system it came from and all prompts will be stored with the data and time of entry.

3.6 Device Manager

This system handles all connection and communication between the device and the host software.

3.7 Error Handler - Hardware

This component constantly checks the states of every system present and ensures that if any of them fail or return an error, an alert is generated. Moreover the system will also try to fix the problem wherever possible.

3.8 Error Handler - Software

This component monitors the state of the host software and will attempt to solve any errors that arise and will alert the user should the attempts fail.

3.9 Host Software

This system is the primary interface for the researchers to analyze the data collected by the device. It consists of several features that allows the researcher to set different thresholds for activity tracking, calibrate all the sensors, update and create new records for participants, and finally interact with the data stored on the device.

3.10 Configuration

This system allows researchers to calibrate the sensors on the device, set and modify the different thresholds for activity tracking and create new prompts.

3.11 Records

This system stores all information about users present in the study. Moreover it provides functionality for researchers to create new records for new users.

3.12 Data View

This is where all data stored on the device can be viewed/ analyzed by the researchers in a graphical manner. The system also contains some functionality for data sorting/parsing and statistical analysis.

3.13 System Boundary Diagram

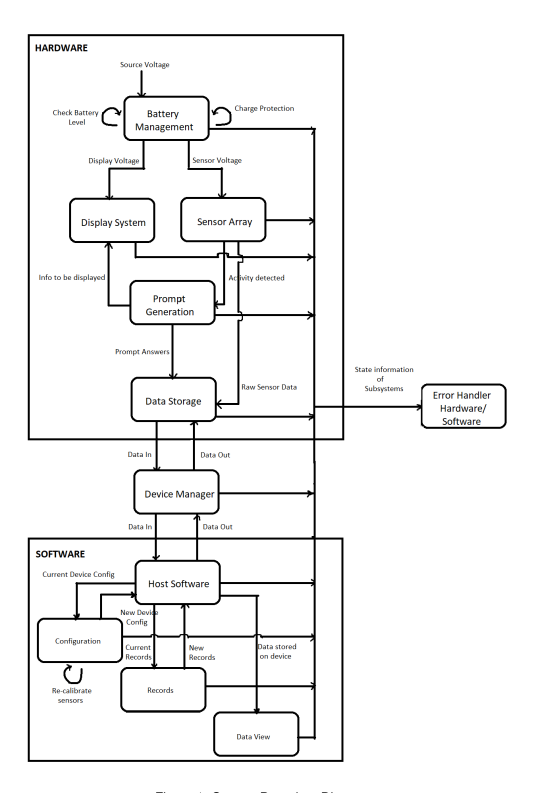


Figure 1: System Boundary Diagram

4 Critical Assumptions

- No wires will come loose during use.
- Batteries are plugged in correctly (the positive and negative ends are aligned as intended).
- All data are stored in the correct memory location.
- All subsystems work as intended.
- All off-the-shelf components work was intended.

5 Failure Mode and Effect Analysis

Table 2: Display FMEA Req: SR1, SR2

		: Display FME.		eq: SR1, SR2	
Design Component	Failure Modes	Causes of Failure	Effects of Failure	Detection	Recommended Action
Display System	Display not working	Improper circuit connection. Battery Level too low. Code malfunction. Physical damage to display.	Users cannot answer prompts Users cannot view any information on the display.	Display system returns an error code. Display Led is OFF	 Users cannot answer prompts. Users cannot view any information on the display. Check wiring and run a diagnostic on the display system. Replace faulty hardware.
	Incorrect information displayed.	 Display Driver faulty. Improper interaction between Display system and Prompt generation. 	Users face un- expected outputs causing improper use of device.	Display system returns an error code.	 Let Error Handler try to solve issue. Perform a manual overview of code. Perform a system reboot.

Table 3: Prompt Generation FMEA Req: SR1, SR2, SR3

Design	Failure	Causes of	Effects of	Detection	Recommended Ac-
•				Detection	
Component	Modes	Failure	Failure	Donata	tion
Prompt Generation System	Prompt not generated	 Prompt generation code faulty System stuck in an idle state where no activity is detected. 	 Display system will not produce an output. Users will be unable to provide feedback regarding activity 	Prompt Generation system returns an error code.	Let Error Handle try to solve the issue
Prompt (Incorrect Prompt Generated	 Prompt generation code faulty Improper interaction between Prompt generation and Sensor Array 	Prompt generated produces unex- pected outputs causing improper use of device	Prompt Generation system returns an error code. Test prompt produces unex- pected outputs	 Let Error Handler try to solve issue. Check System Array State Perform a system reboot.

Table 4: Sensor Array FMEA Req: HR1, SR1, SR2

Design Component	Failure Modes	Causes of Failure	Effects of Failure	Detection	Recommended Action
Sensor Array	Heart rate not detected Skin voltage not detected Muscle Movement not detected No motion detected	 Device not worn properly Sensor breaks down, due to passing thresholds limits Device surface is dusty or contaminated 	EMA may not be triggered	Software check to see if any sensor data is being collected	 Ensure the device has been wrapped around properly for more accurate detection Reboot the system Let error handler try to solve the issue

Table 5: Battery Management FMEA Req: HR2, SR1

Design	Failure Modes	Causes of	Effects of	Detection	Recommended Ac-
Component		Failure	Failure		tion
Battery management system	Device not starting	 Device is faulty Battery has died Device may have overheated Water may have damaged the battery 	Device is not turned on, result- ing in no monitoring	Battery dead in- dicator appears when turn- ing on the device	Contact the Back End Developers for maintenance.

Table 6: Error Handler FMEA Req: SR1,SR2, DSR2

Design	Failure Modes	Causes of	Effects of	Detection	Recommended Ac-
Component		Failure	Failure		tion
Error Handler	Security Compromised	Fail-open security check Error data improperly protected Malicious cyber attack	Participant data will be made vulnerable to exploita- tion	Error handler returns strange or incomplete results	Device enters data- lockdown mode, preventing data from being ac- cessed until security issue is resolved.
	Errors are strange or incomplete	Stack overflow Memory leak Error previously unac- counted for	Persons respon- sible for responding to errors will be unable to diagnose and ad- dress the underlying issue	Error comes in an unex- pected form, or returns no value	Use different channels to handle device logic and error handling Ensure that strange errors either return as Optional or Empty List (i.e. any value but null)

Table 7: Data Storage FMEA Req: SR1, DSR1

Design	Failure Modes	Causes of	Effects of	Detection	Recommended Ac-
Component		Failure	Failure		tion
Data Storage	Data stored at wrong memory location	Incorrect software commands Memory space doesn't exist (invalid memory selected) Insufficient memory space Physical damage to hardware memory chip	Lost and unsaved data	Set up er- ror handler to check if each data point is suc- cessfully stored at the correct memory location each time	Replace faulty hard- ware or set up cor- rect memory path
	Data Stored with incorrect	Wrong data type used	Analysis program	Failed data analysis	Convert data to cor- rect type
	type	for storing data	can't inter- pret data		

Table 8: Device Manager FMEA Req: SR1,SR2

Design	Failure Modes	Causes of	Effects of	Detection	Recommended Ac-
Component	Tandre Modes	Failure	Failure	Detection	tion
Device manager	Unable to establish connection	Loose wires Incorrect communication protocol Incorrect parameters for serial packets (size, format, etc.)	Data can't be trans- ferred between hard- ware and software Lost data	Check list of connected devices on device manager Visual inspection of wiring and circuitry Attach an error detection LED on the device	Make sure all necessary connections are made Reboot device Restart host software

Table 9: Record System FMEA Req: SR1,SR2, DSR2,DSR3

	le 9: Record S			: SR1,SR2, D	
Design Component	Failure Modes	Causes of Failure	Effects of Failure	Detection	Recommended Action
	Records not cre- ated and deletion of records	Code that writes to database is faulty. Connection between software systems are incorrect.	Present users lose data, new users cannot have data stored. Time loss.	Records system returns an error code using error checking. Database trigger for stagnant data.	Create redundant data sets and storage units, or cloud storage of data.
Records	Corrupted records	Code is faulty. Database structure is faulty. Wrong datatype used for storing data.	Data stored for various users is unusable.	Detect if data records are re- turning unusable values, e.g. NaN, Inf, 0, garbage values. View values stored in database to check for unex- pected outputs.	 Prevent users from accessing database. Ensure database is robust prior to deployment through testing.

Table 10: Data View System FMEA Reg: SR1.SR2

Doolan		a View System		Req: SR	_ ·
Design Component	Failure Modes	Causes of Failure	Effects of Failure	Detection	Recommended Action
Data View	Data not visible	Data collection failure. Poorly aggregated data. Incorrect software implementation. Statistical analysis of data is faulty.	Information provided is not useful to researchers or participants.	Error handling to detect if the software is not performing as it should be. Check error codes of device manager.	Visual inspection to make sure device is working. Ensure database is robust prior to deployment through testing
	Graphical data representation incorrect.	Improper communication with device manager. Device malfunction, no correct data.	Incorrect observa- tions could be made based on the data.	Error handling to detect if the software is not performing as it should be. Check error codes of device manager	 Visual inspection to make sure device is working. Use error handler to solve issue.

Table 11: Host Software FMEA Req: SR1,SR2

Design	ign Failure Causes of Effects of Failure Detection Recomm				
Com- po- nent	Modes	Failure	Lifects of Failure	Detection	Action
Host Software	Calculation and float- ing point error	Improper units used (metric vs Imperial) Rounding error Floating point conversion error	 Wrong information can be reported. Rounding errors can propagate through different formulae resulting in data inaccuracies. Data calculated will be completely wrong due to binary overflow. 	 Unit and integration testing for checking proper metrics and calculation. Boundary condition testing for detecting binary overflow/floating point conversion errors. 	 Avoid/fix type conversion and export raw data collected through hardware in case calculation of other metrics for tracker fails. Use error handler to solve issue.
	Memory allocation failed	Memory fragmentation or overflow due to Dynamic memory allocation.	This can cause data to be overwritten and also failure of storing data.	Host software stops running and system fails.	Reboot Soft- ware and make sure to free memory if not using it during memory alloca- tion.
	Security threats and data loss	Malware injected. SQL injection.	Software could have catastrophic failure and device could stop running. SQLi cyberattack can let attackers view or modify the database causing data leaks.	Unrecognizable patterns and processes running in the host software.	 Run a closed-loop network to avoid backend database manipulation. Limit access of tracker to Internet/IoT devices to avoid hacking and data leakage. Perform a manual overview of system and processing running.

Table 12: Configuration System FMEA Req: SR1,SR2,SR4

Table 12: Configuration System FMEA Req: SR1,SR2,SR4							
Design Com- po- nent	Failure Modes	Causes of Failure	Effects of Failure	Detection	Recommended Action		
Configuration of device	Threshold adjustment fails	 Threshold settings changed by Researcher not registered correctly. Code fails to save and keep settings for threshold. 	Wrong configuration of device can lead to false measurements and data reported.	 Boundary condition check to see if threshold values entered are reasonable. Systemwide check for checking last modified date of configuration of device. 	Report error if threshold values are incorrect and Reboot system.		
Con	Calibration fails	 System stuck in idle state and does not record reference for calibration. Not enough power for sensors/system to calibrate. Code to calibrate sensors and system fails. 	 Wrong Calibration of system will result in incorrect activity detection. Data will be false/will not be measured correctly. 	Display message to check if all calibration is completed correctly.	Reboot system and check wiring to sensors. Let error handler solve issue.		

6 Safety and Security Requirements

6.1 Hardware Requirements

- 1. HR1: Device components will have sufficient electrical tolerances for various voltage levels.
- 2. HR2: Charge protection circuity will be integrated to avoid overcharging of the battery.

6.2 Software Requirements

- 1. SR1: The Error Handler will initially try to solve all Non-Hardware issues.
- 2. SR2: Tests will be conducted for all Software before deployment to the device.
- 3. **SR3**: All users are required to complete the test prompt.
- 4. SR4: Configuration setting will fall within accepted boundaries.

6.3 Data Requirements

- 1. **DSR1**: All data to be backed up, in case of power failure.
- 2. DSR2: Access to records will be restricted to administrative users with encrypted ssh access key.
- 3. **DSR3**: The device will not modify user data unnecessarily.

7 Roadmap

Table 13: Road Map

When to Implement	Requirement	Priority
During Course	HR1	High
	HR2	Very High
	SR1	Low
	SR2	High
	SR4	High
	DSR1	Very High
	DSR3	Very High
Postponed*	SR2**	High
	SR3	Low
	DSR2	Very High

^{*}Requirement implementation postponed past course completion.

Postponed requirements are not of lower priority than the requirements to be completed during the course. However, these postponed requirements can only come into effect once the device begins operation in its intended use cases. Otherwise, they are meaningless during the development phase. In addition, low priority values do not mean that the requirement may be dropped. Instead, it means that the implementation of said requirement may be easier, or less critical to the safety of the device, safety of the user, or functionality in general.

^{**}Regarding software imported to device after course completion.

Appendix

Therac-25 case study

The Back End Developers would like to take a section of this document to describe the importance of hazard analyses (and other relevant documentation). A perfect example of this is the Therac-25.

The Therac-25 was a radiotherapy machine designed by Atomic Energy Canada Limited (AECL) in 1982. It was designed to treat cancer by sending beams of high-energy particles (either electrons or x-rays) through a patient's tumor, thereby killing it. This machine was one of the first of its kind to be computer controlled; using software to direct its functions rather than interdependent physical mechanisms. [Leveson(1999)] Considered state-of-the-art at the time, the Therac-25 promised to revolutionize radiotherapy.

However, there were problems. Operators of the machine would find that they encountered many error messages from the computer console, of which none were explained by AECL. These errors were non-descript, and often simple read 'MALFUNCTION' followed by a number. The same operators stated that they had grown used to these messages, and often ignored them and proceeded with treatments. Also, none really knew how the Therac-25 was programmed. The original developer had developed the software for the previous Therac-6 and Therac-20 models of radiotherapy machines, and had since left the company. AECL had directly ported the software into the Therac-25, and did not test the combination of the software and the new hardware properly.

Cancer patients treated using the Therac-25 began to report issues. They entered the machine thinking that the procedures would be painless (which is usually true for radiotherapy). However, they often felt powerful shocks and intense burning sensations around the areas that the Therac-25 would administer radiation. Later on, these same patients would fall heavily sick with symptoms identical to those with acute radiation poisoning. Some patients had to have entire limbs removed due to intense radiation damage, and none knew why. The AECL assured the operators that the machine was incapable of administering deadly doses of radiation, and attested to its safety.[Leveson and Turner(1993)] In reality, the machine was exposing patients to doses of radiation hundreds of times greater than what was prescribed.[Baase and Henry(2019)] Between 1985 and 1987, three people died of radiation exposure due to the Therac-25, and two others sustained lifelong injuries.

A later investigation revealed that when operators changed from the electron mode of the machine to the x-ray mode in a certain way, the Therac-25 would fail to move a dampening shield between the fully powered x-ray and the patient. This dampening shield served to lower intensity of the x-ray beam to safe levels. The software would correctly detect an error, but simply displayed a window with the text 'MALFUNCTION 54'. Having grown used to these messages, the operators would proceed with treatment without the shield in place. In addition, a flag variable which would instruct the machine to perform safety checks was designed to be set by incrementation rather than being set to a fixed value. This flag would often overflow from its maximum value of 255 to zero, which instructed the machine to bypass safety checks. A later review would reveal that the Therac-25 had eleven separate critical engineering and institutional failures which would have put a patient's life in danger. The combination of these factors caused the Therac-25 to give unlucky patients massive radiation overdoses. In response to this incident, the International Electrotechnical Commission (IEC) released standard IEC 62304; a standard which details development lifecycle standards specifically for software present on medical devices. It also describes the dangers and guidance regarding Software of Unknown Pedigree (SOUP).

The worst part is that none of this had to happen. Should a hazard analysis have been performed, the team at AECL who designed the Therac-25 would have known that unaccounted arithmetic overflows could bypass safety features. They would have known that the software written by the original Therac-6 and Therac-20 developer could not be maintained and troubleshot even for the most basic errors. They would have known that non-descript errors in a system like the Therac-25 could potentially be life-threatening.

They would never have assured machine operators that overdoses were impossible. Had they done so, they would have never decided that a direct port of the Therac-20 software would have been appropriate for the system they were designing. They may have thrown out the software all together and started from scratch. This would have taken much more time and many more resources than a simple port, but that team had a responsibility to ensure that the potential hazards of the Therac-25 were accounted for and mitigated. It would have been a good decision. As they did not make it, three patients died from treatments meant to save their lives.[Rose(1994)]

Events like these are a sobering reminder of the responsibilities that engineers have to the products they design, the users of said products, and to the world that their systems exist in. A hazard analysis is an essential document to ensure that catastrophic failures do not occur as a result of an engineer's work. It can act as a reminder to engineers that there is a reason to all of this procedure, and that documentation is essential because the risks are real. It is the responsibility of every engineer to ensure cases such as the Therac-25 never happen again.

Error Codes

To facilitate error handling, every system will return an error code that represents the status of that system. These error codes follow the following format, BED_ERR_ERRORTYPE For example BED_ERR_NONE represent that the system is working correctly. Some more examples of error codes are:

- BED_ERR_PARAM_ERR: Represents an error with the parameters of the function.
- BED_ERR_GENERAL: Represents a generic error that has not been cataloged yet.
- BED_ERR_INVALID_DATA_SIZE : Represents an error related to a size mismatch in what is stored within a block of memory or variable.

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