Software Requirements Specification Software Engineering

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Table 1: Revision History

Date	Version	Notes
2023-10-06	1.0	Initial revision of SRS

This document describes the requirements for the web application REACH. The template for the Software Requirements Specification (SRS) is a subset of the Volere template (Robertson and Robertson, 2012). If you make further modifications to the template, you should explicitly state what modifications were made.

1 Project Drivers

1.1 The Purpose of the Project

The purpose of this project is to create a web application, REACH, which will help people find and apply to clinical trials that they are eligible for. Currently there is no intuitive path between researchers and patients which patients can use to apply to a clinical trial. Existing databases of these clinical trials are often hard to navigate, especially for those who do not have experience with medical terminology. The completed application will allow users to find clinical trials for which they may be eligible, and give users a method of contacting the research coordinator for that trial.

1.2 The Stakeholders

1.2.1 The Clients

The clients for this project are our project supervisors: Dr. Terrence Ho, and Dr. Cieran Scallan.

1.2.2 The Customers

The customers for this project are people who are looking to take part in a clinical trial.

1.2.3 Other Stakeholders

Other stakeholders for this project are researchers who will be contacted by patients using this application.

1.3 Mandated Constraints

The constraints for this project are:

1.3.1 Implementation Constraints

The project must be accessible from the web with the need for installation of any applications other than a web browser.

1.3.2 Schedule Constraints

The project must be completed within the 8 month period of the school year.

1.3.3 Budget Constraints

The project must not use more than \$750.

1.4 Naming Conventions and Terminology

- Platform The web system/application.
- Heavy traffic When many users are accessing the platform at one time.
- NIST National Institute of Standards and Technology
- Patient An individual with a given medical condition(s) who is seeking to enroll in medical research studies.
- Clinician A medical care provider enrolling a patient in research studies on their behalf.
- Organizer Person or organization seeking patients for their research study.

1.5 Relevant Facts and Assumptions

It is assumed that users of the application will vary in age, ability, and how much experience they have in using web applications.

2 Functional Requirements

2.1 The Scope of the Work and the Product

2.1.1 The Context of the Work

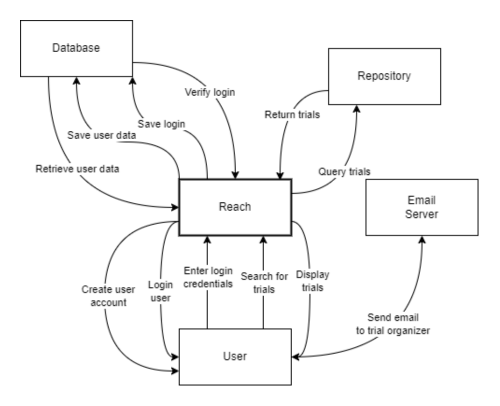


Figure 1: Work Context Diagram

2.1.2 Work Partitioning

Event Number	Event Name	Input	Output
1	Search	Keyboard and mouse	Trial list
2	Login	Keyboard and mouse	Logged-in view
3	Create account	Keyboard mouse	User account
4	Opening FAQ	Mouse	FAQ page
5	Updating user	Keyboard and mouse	Updated user
	data		data entry
6	Sending email	Keyboard and mouse	Sent email

Table 2: Work Partitioning Events

Event Number	Summary
1	The user can use their mouse to navigate to the search fields,
	use their keyboard to enter data, and click submit.
2	The user, through mouse and keyboard input, can enter their
	login details in the login screen.
3	The user can enter their new login details through mouse and
	keyboard in the account creation dialogue.
4	The user can click on the FAQ option to arrive at the FAQ
	page.
5	The user can use their mouse and keyboard to navigate to
	their profile and update their user data.
6	The user can use their mouse and keyboard to send an email
	to the trial coordinator.

Table 3: Work Partitioning Summaries

2.1.3 Individual Product Use Cases

1. Patient/Clinician Account Creation. A patient or clinician creates an account on the app. The user first selects to create an account. They provide a username and password, along with their email. Upon entering valid inputs, the user is informed that their account is created, and they are logged in. Otherwise, the user should be notified of the invalid credentials.

- 2. Patient/Clinician Login. A patient or clinician logs into their account. When the user attempts to login to their account, they are required to input their credentials once the system verifies that their details are valid it will log them in to the system. If invalid credentials are entered, the user is informed of the incorrect credentials and is prompted to try again.
- 3. Profile Creation. The user can input a set of information related to themselves and their medical condition(s), and save it for later use when searching for clinical studies or trials.
- 4. Select Profile. The user can select a profile that they have previously created, which will become the basis for a study search.
- 5. Search for Studies. Using a previously created profile that is chosen by the user, studies are retrieved from the repository, filtered, and displayed to the user.
- 6. Send Email to Organizer. Once a user selects a study, they can use the provided template and contact information to send an email to the organizer of the study.
- 7. Profile Modification. A patient modifies their profile to change certain attributes (weight, height, birthday, medical conditions, address). Upon saving the changes, the server updates their information in the database, and the change is reflected in the frontend.
- 8. Bookmark Studies. Upon searching for clinical studies/trials, a user can bookmark (i.e., save) the study/trial, and should be able to view all the studies/trials they have saved previously.

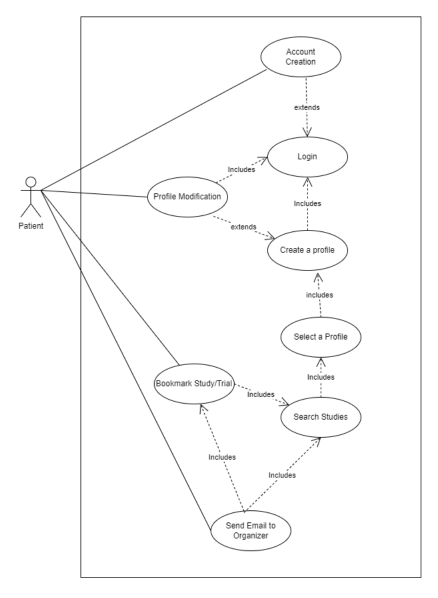


Figure 2: Patient Use Case Diagram

2.2 Functional Requirements

FR-1: The system shall enable the user to create an account before signing in to the application. Account credentials must include an email and a password satisfying the NIST password guidelines.

Rationale: Many application features will be reliant on a user account/email.

Additionally, to ensure the safety/security of sensitive information, it is important to enforce strict password guidelines.

Fit criterion: The system shall log the user in to their newly created account upon successful creation.

FR-2: The system shall ensure the same email cannot be used across multiple accounts.

Rationale: The email needs to be a unique identifier of an account.

Fit criterion: The system shall inform a user if the email they entered is already in use under another account.

FR-3: The system shall enable a user to sign in to an existing account.

Rationale: To use all of the application features a user must be signed in to their account.

Fit criterion: The system shall log the user into the application upon validation of the credentials they entered. If invalid credentials are entered, the user should be informed of this error.

FR-4: The system shall enable a user to sign in to the application as a guest. **Rationale:** Some features of the application do not require a full account, so the user should have the option to sign in as a guest if they only require a subset of the application's features.

Fit criterion: The system shall log the user into the application as a guest upon entering an identifier for the current session.

FR-5: The system shall enable a user to reset their password before signing in to the application.

Rationale: It is likely some users will forget their password, and will need to reset it to be able to log back into their account.

Fit criterion: The system shall send the user an email (to the email tied to the account they are trying to log into), allowing them to create a new password. The user should be logged into their account upon entering a new, valid password.

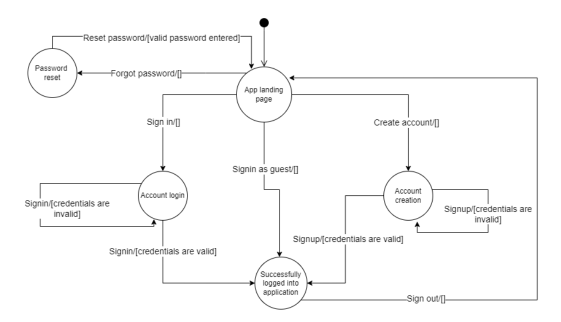
FR-6: The system shall enable a user to sign out of the application.

Rationale: Users should be able to sign out for both security reasons, and practical reasons.

Fit criterion: The system shall return the user to the main landing page upon signing out of their account.

State machine for user login/signup:

- Precondition User is on the initial page of the app
- Postcondition User is logged in to application
- Intermediary States:
 - User is in the process of signing in
 - User is in the process of creating an account
 - User is in the process of resetting their password



FR-7: The system shall prompt a user to enter information about themselves and their medical history upon creation of an account (i.e., first time

signing on), or when a user signs on as a guest.

Rationale: The system must know information about the user to match them to potential clinical trials.

Fit criterion: After a user signs on for the first time, they should be asked to enter their date of birth, height, weight, gender, and current address. Additionally, they should be asked to enter any medications they take, and any medical conditions that they have.

FR-8: The system shall store the information entered by a user with an account in the database.

Rationale: The system shall only need to ask users with an account for their information once.

Fit criterion: The system shall be able to retrieve the information tied to a users account after they have signed out of the application.

FR-9: The system shall enable a user to update the personal information tied to their account.

Rationale: Personal characteristics and circumstances are always changing, and a user should be able to make these changes in the application.

Fit criterion: The updates made by the user shall be reflected in both the UI (for the user to see) and in the database.

FR-10: The system shall create an email template for the user to send their interest in participating in a trial to the research coordinator.

Rationale: Users may not know what they need to include in the email, so the template should simplify this process.

Fit criterion: The system shall output an email template. The email template shall include all relevant information that the user has provided and details about the trial they wish to participate in.

FR-11: The system shall notify the user when new medical trials matching their specifications are available, if requested by the user.

Rationale: New medical trials matching the users specifications which the user may want to participate in may be started after the user searches ini-

tially, this feature means the user would not have to search every couple of days for new trials.

Fit criterion: The system shall notify the user about new trials with parameters that they specified.

FR-12: The system shall include frequently asked questions and information about how medical trials work.

Rationale: Users may not be aware of how medical trials work exactly, and this may cause confusion for the user when searching for a medical trial.

Fit criterion: The system shall contain and display information about medical trials, and answers to questions the users may commonly have.

FR-13: The system shall allow the user to search for medical studies based on their health conditions.

Rationale: Searching for studies is the primary purpose of the application. Fit criterion: The system shall query and filter medical trials from a repository of medical studies.

FR-14: The system shall display to the user the results of their trial search, complete with information about the search, the distance to their location, and the option to contact the organizer of the clinical trial.

Rationale: Users must be able to see the trials for which they searched, and be given information pertaining to each result.

Fit criterion: The system shall display a listing of trials based on a user search.

FR-15: The system shall allow a signed-in user to automatically enter their conditions and other information into the search parameters.

Rationale: This will save users time when performing multiple searches, and fulfils the purpose of having a login system.

Fit criterion: The system shall automatically populate search fields for a logged-in user with their saved information.

FR-16: The system shall allow a signed-in user to save (or, bookmark) trials.

Rationale: This will allow users to save trials that they are interested in for later use (for example, if the trial is not recruiting yet, or if the user still needs to decide whether or not they want to partake in the study).

Fit criterion: Users should be able to know which trials are saved and which trials are not saved after saving a trial.

FR-17: The system shall allow a signed-in user to delete trials that they have previously saved.

Rationale: A user may be finished with a trial, may no longer be interested in the trial, or could have saved the trial by accident - all valid reasons for a user to want to delete the trial.

Fit criterion: Users should be able to know which trials are saved and which trials are not saved after deleting a trial.

FR-18: The system shall allow a signed-in user to access the trials that they have previously saved.

Rationale: A user will likely be saving trials for future use - so the user must have an easy way to access these saved trials when they return to the application at a later date.

Fit criterion: Users can obtain a list of all the trials that they have previously saved, and can perform any action on these trials that they would from the searching page (for example, deleting the trial).

FR-19: The system shall allow a signed-in user to filter the list of saved trials based on the different sets of search criteria that they have created on their account.

Rationale: If a user has many different sets of information that they use to conduct searches, it will be much easier to view the saved trials on a "set by set" basis, as opposed to viewing all the trials at once.

Fit criterion: A user can select which set of criteria they would like to use to filter the list of saved trials by. Upon selecting the criteria, only the trials that were saved under that "profile" shall be displayed to the user.

FR-20: The system shall display the location of a selected trial on a map for the user to see, when there is a list of trials displayed for the user to see.

Rationale: User's will not be familiar with every trial location, so giving them a map to see the location will allow them to gain a better understanding of where it is.

Fit criterion: Upon selecting a trial, the map should update with the new location, corresponding to the newly selected trial.

FR-21: The system shall enable a user to switch between multiple pre-saved sets of information which will act as the 'autofill' for a trial search.

Rationale: It is possible that a user may have multiple sets of information that they want to conduct a search for (i.e., on behalf of family members, or perhaps for multiple medical conditions), so they must be able to select which set of information will be used for the search.

Fit criterion: Upon selecting a set of information, any search must be done with the information tied to that set, until a new set is selected.

3 Non-functional Requirements

3.1 Look and Feel Requirements

NFR-1: The system's interface must be formatted neatly and not be cluttered.

Rationale: The user should find the appearance of the system's interface attractive to further their interest in using the system.

Fit criterion: Repeated or grouped sections will have consistent measurements and styling. Additionally, at least 25% of every page should be empty (i.e., just whitespace).

Traceability: FR-1, FR-3, FR-4, FR-5, FR-6, FR-7, FR-9, FR-12, FR-13, FR-14, FR-15

NFR-2: The font and colors of the system's interface must be simple and

allow users to easily read the text or interpret the images.

Rationale: If the interface is not simple, it may cause the users to strain their eyes or distract them from the main content displayed on the interface.

Fit criterion: The interface will utilize solid colors with no patterns. The font used for the interface will be Roboto.

Traceability: FR-1, FR-3, FR-4, FR-5, FR-6, FR-7, FR-9, FR-12, FR-13, FR-14, FR-15

3.2 Usability and Humanity Requirements

NFR-3: The application interface shall be intuitive to use to a point where users do not require a tutorial or help from another person.

Rationale: The interface will be used by users with a wide range of technical ability, so making the interface as easy to learn as possible will make the application maximally accessible.

Fit criterion: 95% of users tested should be able to accomplish the main function of the application without need for outside intervention.

Traceability: FR-1, FR-3, FR-4, FR-5, FR-6, FR-7, FR-9, FR-12, FR-13, FR-14, FR-15

NFR-4: The application interface shall only include the minimum necessary elements for the system to function effectively.

Rationale: The simplicity of the system is important to ensure that the system is easy for all users to interact with.

Fit criterion: Testers should not be able to identify any element of the user interface that does not serve any immediate and apparent use.

Traceability: FR-1, FR-3, FR-4, FR-5, FR-6, FR-7, FR-9, FR-12, FR-13, FR-14, FR-15

NFR-5: Users are able to read the UI text without issue.

Rationale: The average user cannot make full use of the features if they find it difficult to read the instructions, options, or descriptions.

Fit criterion: The UI text size will be 12 pt.

Traceability: FR-1, FR-3, FR-4, FR-5, FR-6, FR-7, FR-9, FR-12, FR-13,

3.3 Performance Requirements

NFR-6: The system shall load and display trials to the user in a timely manner.

Rationale: In addition to obvious usability reasons, there is a potential for users (especially clinicians) to rapidly change the criteria for eligibility, meaning the time spent waiting for trials to load can grow very fast if it is slow.

Fit criterion: The system shall not take longer than 5 seconds to load and display a set of trials, during any given search made by a user.

Traceability: R-13, R-14, R-15

NFR-7: The system shall remain performant during times of heavy traffic. **Rationale:** The system will experience varying amounts of traffic, and it should be prepared to handle all cases (heavy traffic being the one to likely cause issues).

Fit criterion: Any system API call shall take less than 1 second to respond for up to 1000 concurrent users, and less than 2 seconds to respond for anything greater (assuming the system will not see > 10000 users).

Traceability: Traces to pretty much all FRs that are mentioned in the other performance requirements

NFR-8: The systems UI elements shall acknowledge all forms of user input in a timely manner.

Rationale: Timely response from the system is necessary for a good user experience. Additionally, things like keyboard shortcuts should be just as performant as using a mouse (and vice-versa).

Fit criterion: The system shall take less than 150ms to acknowledge user input, and make it clear to the user that it has registered the request.

Traceability: FR-1, FR-3, FR-4. FR-5, FR-6, FR-7, FR-9, FR-13, FR-14, R-15

NFR-9: The system shall efficiently store/retrieve user data from the database.

Rationale: For the system to remain performant, it must handle data efficiently, especially as the total number of users increase.

Fit criterion: The system shall take no longer than 500ms when querying the database for user info (includes query execution time + any overhead of executing the query).

Traceability: FR-1, FR-2, FR-3, FR-4, FR-5, FR-8, FR-9, FR-15

NFR-10: The system shall be available to users 99% of the time.

Rationale: Since users will likely make infrequent visits to the platform, it is necessary that the platform will be available at those times (i.e., if the system is frequently unavailable, it increases the chances that users will never use the platform).

Fit criterion: The system shall experience no more than 15 hours of down-time throughout the year.

Traceability: N/A

3.4 Operational and Environmental Requirements

NFR-11: The system shall interface with data from ClinicalTrials.gov and/or some other medical trials repositories.

Rationale: The system needs to gather trials from a preexisting data source so that it can match patients to them.

Fit criterion: The system should have a similar number (within order of magnitude) of currently running clinical trials as Clinical Trials.gov for users to search from.

Traceability: FR-13

NFR-12: The system shall be able to be used on all modern devices which are capable of browsing the web.

Rationale: Users will likely use a large variety of devices from phones to PCs, so it is necessary to support them, (i.e. users are less likely to use the application if it doesn't work on mobile).

Fit criterion: The system shall work on smartphones, personal computers (including laptops and desktops), and tablets (such as an ipad). In all cases,

the user must be accessing the app through a web browser (including google chrome, firefox, safari, and microsoft edge), using a version which was released no earlier than 2019.

Traceability: N/A

3.5 Maintainability and Support Requirements

NFR-13: The system shall be easy to fix in the event it experiences unexpected downtime.

Rationale: While it may not happen often, an unexpected system failure is definitely possible. As the time it takes to fix the issue increases, so does the number of users negatively affected.

Fit criterion: On average, it should take less than 1 hour to restore the system in the event of a failure.

Traceability: N/A

NFR-14: The system shall be adaptable to new requirements.

Rationale: It is likely new requirements will be discovered once users begin to use the platform, and there will be a need/desire to implement these quickly and effectively.

Fit criterion: The system must satisfy all of its existing requirements after the implementation of a new requirement/feature.

Traceability: N/A

3.6 Security Requirements

NFR-15: The interface shall ensure that the (account holding) users creates a strong password according to the NIST guidelines for strong password creation.

Rationale: The NIST is a reputable organization creating strong standards for all manners of security compliance. Following their standards for password creation will ensure that the users' accounts are secure.

Fit criterion: The interface shall not allow the user to create an account with a password that does not meet the NIST guidelines. All passwords will

be 12 characters long with at least one uppercase letter, one lowercase letter, one number, and one special character. Additionally, the interface will recommend against using common passwords, or known words.

Traceability: FR-1, FR-5

NFR-16: The system shall ensure database access is controlled by a password and cannot be accessed by anyone except for the database manager.

Rationale: Although the development team should initially have access to the database for testing purposes. Only a database manager should be able to access individual account information. Role-based Access Control (RBAC) can increase security and ensure information is not dispersed unnecessarily to team members.

Fit criterion: The system will have admin roles for database access and shall ensure that there is authentication with strong credentials. The system will also restrict database access to known IPs to ensure personnel authorization.

Traceability: FR-8, FR-9

3.7 Cultural Requirements

NFR-17: The interface shall be useable by users who speak a language other than English or French.

Rationale: Restricting the interface's language to only official languages will exclude many ethnic minorities from being able to access clinical studies. Making the program multilingual will ensure that a minimal number of ethnic minorities will be excluded.

Fit criterion: Users who speak the languages in the top 5 most spoken non-official languages in Canada should be able to use the interface without issue.

Traceability: FR-1, FR-3, FR-4, FR-5, FR-6, FR-7, FR-9, FR-12, FR-13, FR-14, FR-15

3.8 Legal Requirements

NFR-18: The system shall conform to regulations regarding the collection, storage, and usage of medical information.

Rationale: Since the data stored pertains to personal medical information, we are legally required to manage the data in such a way that protects the personal information of the patients.

Fit criterion: The system should be able to pass an independent audit to ensure the data is being collected and secured in an appropriate manner.

Traceability: FR-7, FR-8, FR-9, FR-13, FR-14, FR-15.

3.9 Health and Safety Requirements

N/A

4 Project Issues

4.1 Open Issues

N/A

4.2 Off-the-Shelf Solutions

There are currently some websites with a repository of active research studies and clinical trials that users can search through, such as clinicaltrials.gov. These current solutions display the studies and allow the users to filter them based on some general information such as condition, treatment, age, etc.

4.3 New Problems

N/A

4.4 Tasks

Please refer to the *Project Scheduling* section in our Development Plan.

4.5 Migration to the New Product

N/A. REACH will be a new, standalone product.

4.6 Risks

The major risks associated with this project are as follows:

Table 4: Risk Table

Risk	Probability	Impact
Not being able to have the app deployed on the cloud.	Moderate	High
Not being able to allow for account creation and subse-	Moderate	Moderate
quent saving of information.		
Python or Typescript functions being used in the back-	Moderate	High
end and frontend get deprecated.		
Inaccurate parsing of information through API calls.	Low	High
Database is not properly secured and information is	Low	High
available to the public.		

4.7 Costs

No large-scale monetary costs should be involved in the development of this project. The only costs that may be incurred are the costs of hosting the application on a cloud platform, and the time cost for developers.

4.8 User Documentation and Training

A set of documentation pages with examples will be created in Confluence as part of project completion. This will be tailored (in part) towards the end user, and will be used to help them understand how to use the application. Additional sections will be created for developers in order to facilitate merge requests and validate results. Training will also be provided to Dr. Ho and his team to ensure they are able to use the application.

4.9 Waiting Room

FR-4, FR-5: These requirements are not critical to the core functionality of the application, and will be implemented after the core functionality is

complete.

FR-11: This requirement is a low priority, and will be implemented after higher priority requirements are completed.

4.10 Ideas for Solutions

N/A - There is no additional commentary to add here that has not been included in the requirements.

4.11 Anticipated Changes

Below are some changes that could potentially be made to future revisions of the SRS for this application, depending on several factors, such as unexpected costs/issues arising during development (or even the lack of expected costs/issues during development).

- Two different types of accounts one for patients and one for clinicians
- Clinician account specifically designed for finding clinical trials on behalf of patients
- Clinician account enables saving patient's information (i.e., clinicians can have their own personal database of patients)
- The types of information collected should be expected to change frequently, especially as development begins and users begin testing the application

References

James Robertson and Suzanne Robertson. Volere Requirements Specification Template. Atlantic Systems Guild Limited, 16 edition, 2012.

5 Appendix

Reflection:

Below are some skills that all the team members will need to acquire, two ways the team can learn the skill, and how the team actually plans on developing/learning the skill.:

- How to deploy apps to the cloud
 - Free videos/tutorial's on YouTube (or other similar platforms)
 - Documentation surrounding the cloud platform (or platform's) of interest
- How to use existing NLPs to parse/process data
 - Online classes related to Machine learning and natural language processing
 - Free videos/tutorial's on YouTube (or other similar platforms)

For both of the skills mentioned above, the team has collectively decided to use YouTube (and other free resources) to develop the skills, since there is an abundance of information available, which would likely be more than enough for what is needed for the application being developed.

Additionally, below is a skill that each individual will need to develop, two ways they can learn that skill:

- David Typescript
 - Free videos/tutorials on YouTube (or other similar platforms)
 - Websites such as "codeacademy" or "freecodecamp"
 - Typescript documentation
- Deep React
 - Free videos/tutorials on YouTube (or other similar platforms)

- Online classes on platforms like "Courseera"
- Alan Django
 - Free videos/tutorials on YouTube (or other similar platforms)
 - Online classes on platforms like "Courseera"
- Aamina Backend development in python
 - Free videos/tutorials on YouTube (or other similar platforms)
 - Online classes on platforms like "Courseera"
 - Textbooks
- Anika Backend development in Python
 - Free videos/tutorials on YouTube (or other similar platforms)
 - Code Academy, Pluralsight, Udemy
 - Textbooks

How each member is actually planning on learning the skill:

- David Will use YouTube, due to the abundance of videos/tutorials made on Typescript
- Deep Will use online classes in a more formalized setting (like "Courseera") since it is more structured than something like youtube, and React is a complex tool to learn.
- Alan Similar to React, Django is a pretty complex web platform, so will use online classes due to the structure it provides.
- Aamina Similar to Typescript, will use YouTube due to the abundance of videos made on Python/backend development in python.
- Anika Will use YouTube and Code Academy to learn the fundamentals and will supplement with Pluralsight.

5.1 Symbolic Parameters

N/A