

SECURE THE DRUG COMPONENTS USING NAÏVE BAYES AND SVM

##### A PROJECT REPORT

###### ***Submitted by***

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**ABSTRACT**

In this paper, we propose a framework to secure drug components in the cloud. Specifically, we design for multiple drug formula providers’ to use the cloud securely. In our approach, the analyzer trains the drug formulas using Support Vector Machine (SVM) and naïve Bayes. To perform integer and fraction computations in the cloud server, we designed secure computation protocols. We securely train the SVM to privately refresh the selected SVM parameters using the two protocols which are SVM parameter selection protocol and sequential minimal optimization protocol. We train NB based on bayes theorem with an assumption of independence among predictors. To determine whether a drug compound is active or inactive in a cloud, the trained SVM and NB  classifier is used.Lastly, we prove that the proposed framework achieves the goal that facilitate drug manufacturers to securely outsource their formulas without privacy leakage to unauthorized parties in the cloud for storage and  for SVM and NB training.

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**LIST OF ABBREVIATIONS**

JDK Java Development Toolkit

DEX Dalvik Executables

TCP Transmission Control Protocol

IP Internet Protocol

**HTTP** Hyper Text Transfer Protocol

**ADT** Android Development Tool

**INTRODUCTION**

**CHAPTER 1**

**INTRODUCTION**

* 1. **OVERVIEW OF THE PROJECT**

Drug discovery is the process through which potential new therapeutic entities are identified, using a combination of computational, experimental, translational, and clinical models. Drug discovery is generally the process of identifying one or more active ingredients from traditional remedies, and includes the identification of screening hits, medicinal chemistry and optimization of these hits to increase the affinity, selectivity (to reduce the potential of side effects), bioavailability, and metabolic half-life. Drug discovery and development is a long-term, competitive, expensive and complicated process. Bringing the drugs from the bench to the market, that is, from screening and identification of the drug to its introduction to the market, takes several years of efforts. The process of discovering and developing a new drug involves an intricate interaction between investors, industry, academia, patent laws, regulatory authorities, marketing and the necessity to balance confidentiality with communication. The complete process of presenting a drug to the patients involves four stages- drug discovery, drug development, regulatory review and approval, and marketing.

* 1. **PROBLEM DEFINITION OF THE PROJECT**

In the drug discovery process, the first step is to identify an appropriate 'drug gable' target, which can be a biomolecule or a protein receptor. After the target has been identified, the next step involves target validation and the confirmation. This is followed by testing of the target against different small molecule compounds to identify lead compounds. The lead compounds can be identified by screening a library of compounds through various methods, such as high-throughput screening, de novo synthesis and isolation from the natural products. The drug development phase involves stringent testing and optimization of the selected compounds to identify the 'drug candidate’ which might be most effective in terms of safety, toxicity, dosage, and efficacy. For this purpose, the selected lead compounds are tested in cells (in vitro) and in animals (in vivo) to study their pharmacodynamics and pharmacokinetic properties. The successful lead candidate must be non-toxic and should be absorbed into the bloodstream, can be distributed to the proper action site in the body, and can be metabolized efficient and effective as well as successfully excreted from the body. This part of the development process is referred to as the ‘preclinical phase’. The outcome of the ‘clinical trial’ decides whether the drug candidate is safer and effective enough in treating the disease. At this point, new drug applications with all the essential evidence, including quality, preclinical and clinical data collected during development of the drug candidate, are submitted to the relevant regulatory authorities. At this point, there may be a chance where the drug information gets disclosed. Due to the significant investments and high commercial values involved in drug discovery, privacy is an important factor .To secure the drug components, a data mining tools is used.

Of the data mining tools, Support Vector Machine (SVM) has a relatively high decision rate and has been widely used in recent times to predict ligand-based chemical compounds in drug discovery. In approaches using SVMs,known drug formulas datasets is used to train the SVM classifier, and the new drug compound visual scanning is done by trained SVM.As privacy is prime, how can we minimize the risk of unauthorized disclosure during the SVM training phase? In this context, when a researcher sends some chemical compounds to the cloud for SVM classification, it is important to ensure that the potential new drug compounds will not be leaked to a third party, such as a competing pharmaceutical corporation. Furthermore, to train the SVM, multiple pharmaceutical corporations may collaborate in order to increase the SVM decision rate. At the same time, these corporations do not wish to reveal their datasets. How to achieve secure SVM training and decision under multiple data sources without compromising, the privacy of each individual party remains a research and operational challenge. Thus, in this paper, we propose secure drug components using SVM and Naïve Bayes for Securing Drug discovery in the cloud environment. Unlike existing drug discovery frameworks, our framework seeks to achieve the following

• Secure Outsourced Data Storage: The drug formula owner can securely outsource the data (e.g. drug formula) to the cloud for storage without leaking the data to unauthorized third parties.

• Secure Multi-Source SVM Training: The POD allows an authorized model provider to use other drug formula owners’ encrypted data to train the SVM on the fly. The model provider can decrypt and obtain the trained model without knowing (contents of) the training dataset.

• Secure SVM Drug Decision: An authorized tester can securely upload his/her drug chemical compounds to the cloud and determine whether the compound is active or not in a privacy-preserving way.

Commercialization is the last phase of drug development process. Once the drug has been approved, it is marketed or commercialized. The drug manufacturers need to submit marketing authorization applications in every country in which they want to sell the drug. As the drug is typically targeted to a very large number of patients, the manufacturer is expected to monitor this stage cautiously and submit reports to the FDA. The reports include evidence for medicine-related problems, e.g., treatment failure, adverse reaction, counterfeit/poor quality medicines, drug interactions, or incorrect use. These reports are significant in terms of generating proof of efficacy that will inspire public confidence and trust.

**LITERATURE**

**SURVEY**

**CHAPTER 2**

**LITERATURE SURVEY**

**DEEP LEARNING IN DRUG DISCOVERY AND MEDICINE; SCRATCHING THE SURFACE**

The practice of medicine is ever evolving. Diagnosing disease, which is often the first step in a cure, has seen a sea change from the discerning hands of the neighborhood physician to the use of sophisticated machines to use of information gleaned from biomarkers obtained by the most minimally invasive of means. The last 100 or so years have borne witness to the enormous success story of allopathy, a practice that found favor over earlier practices of medical purgatory and homeopathy. Nevertheless, failures of this approach coupled with the omics and bioinformatics revolution spurred precision medicine, a platform wherein the molecular profile of an individual patient drives the selection of therapy. Indeed, precision medicine-based therapies that first found their place in oncology are rapidly finding uses in autoimmune, renal and other diseases. More recently a new renaissance that is shaping everyday life is making its way into healthcare. Drug discovery and medicine that started with Ayurveda in India are now benefiting from an altogether different artificial intelligence (AI)—one which is automating the invention of new chemical entities and the mining of large databases in health-privacy-protected vaults. Indeed, disciplines as diverse as language, neurophysiology, chemistry, toxicology, biostatistics, medicine and computing have come together to harness algorithms based on transfer learning and recurrent neural networks to design novel drug candidates, a priori inform on their safety, metabolism and clearance, and engineer their delivery but only on demand, all the while cataloging and comparing omics signatures across traditionally classified diseases to enable basket treatment strategies. This review highlights inroads made and being made in directed-drug design and molecular therapy.[1]

**A NOVEL NEUTROSOPHIC WEIGHTED EXTREME LEARING MACHINE FOR IMBALANCED DATA SET**

Extreme learning machine (ELM) is known as a kind of single-hidden layer feedforward network (SLFN), and has obtained considerable attention within the machine learning community and achieved various real-world applications. It has advantages such as good generalization performance, fast learning speed, and low computational cost. However, the ELM might have problems in the classification of imbalanced data sets. In this paper, we present a novel weighted ELM scheme based on neutrosophic set theory, denoted as neutrosophic weighted extreme learning machine (NWELM), in which neutrosophic c-means (NCM) clustering algorithm is used for the approximation of the output weights of the ELM. We also investigate and compare NWELM with several weighted algorithms. The proposed method demonstrates advantages to compare with the previous studies on benchmarks.[2]

**DRUG DESIGN AND DISCOVERY: PRINCIPLES AND APPLICATIONS**

Drug discovery is the process through which potential new therapeutic entities are identified, using a combination of computational, experimental, translational, and clinical models Despite advances in biotechnology and understanding of biological systems, drug discovery is still a lengthy, costly, difficult, and inefficient process with a high attrition rate of new therapeutic discovery. Drug design is the inventive process of finding new medications based on the knowledge of a biological target. In the most basic sense, drug design involves the design of molecules that are complementary in shape and charge to the molecular target with which they interact and bind. Drug design frequently but not necessarily relies on computer modeling techniques and bioinformatics approaches in the big data era. In addition to small molecules, biopharmaceuticals and especially therapeutic antibodies are an increasingly important class of drugs and computational methods for improving the affinity, selectivity, and stability of this protein-based therapeutics have also gained great advances [3]. Drug development and discovery includes preclinical research on cell-based and animal models and clinical trials on humans, and finally move forward to the step of obtaining regulatory approval in order to market the drug. Modern drug discovery involves the identification of screening hits, medicinal chemistry and optimization of those hits to increase the affinity, selectivity (to reduce the potential of side effects), efficacy/potency, metabolic stability (to increase the half-life), and oral bioavailability. Once a compound that fulfills all of these requirements has been identified, it will begin the process of drug development prior to clinical trials.[3]

**PRIVACY-PRESERVING CLINICAL DECISION SUPPORT SYSTEM USING GAUSSIAN KERNEL-BASED CLASSIFICATION**

A clinical decision support system forms a critical capability to link health observations with health knowledge to influence choices by clinicians for improved healthcare. Recent trends towards remote outsourcing can be exploited to provide efficient and accurate clinical decision support in healthcare. In this scenario, clinicians can use the health knowledge located in remote servers via the Internet to diagnose their patients. However, the fact that these servers are third party and therefore potentially not fully trusted raises possible privacy concerns. In this paper, we propose a novel privacy-preserving protocol for a clinical decision support system where the patients’ data always remain in encrypted form during the diagnosis process. Hence the server involved in the diagnosis process is not able to learn any extra knowledge about the patient data and results. Our experimental results on popular medical data sets from UCI database demonstrate that the accuracy of the proposed protocol is up to 97:21% and the privacy of patient data is not compromised.[4]

**PRIVACY-PRESERVING MULTI-CLASS SUPPORT VECTOR MACHINE FOR OUTSOURCING THE DATA CLASSIFICATION IN CLOUD**

Emerging cloud computing infrastructure replaces traditional outsourcing techniques and provides flexible services to clients at different locations via Internet. This leads to the requirement for data classification to be performed by potentially untrusted servers in the cloud. Within this context, classifier built by the server can be utilized by clients in order to classify their own data samples over the cloud. In this paper, we study a privacy-preserving (PP) data classification technique where the server is unable to learn any knowledge about clients’ input data samples while the server side classifier is also kept secret from the clients during the classification process. More specifically, to the best of our knowledge, we propose the first known client-server data classification protocol using support vector machine. The proposed protocol performs PP classification for both two-class and multi-class problems. The protocol exploits properties of Pailler homomorphic encryption and secure two-party computation. At the core of our protocol lies an efficient, novel protocol for securely obtaining the sign of Pailler encrypted numbers.[5]

**DRUG DISCOVERY AND DEVELOPMENT: AN INSIGHT INTO PHARMACOVIGILANCE**

Drug discovery and development is a long-term, competitive, expensive and complicated process. Bringing the drugs from the bench to the market, that is, from screening and identification of the drug to its introduction to the market, takes several years of efforts [1]. The process of discovering and developing a new drug involves an intricate interaction between investors, industry, academia, patent laws, regulatory authorities, marketing and the necessity to balance confidentiality with communication [2]. The complete process of presenting a drug to the patients involves four stages- drug discovery, drug development, regulatory review and approval, and marketing.[6]

**DRUG DISCOVERY AND DEVELOPMENT: ROLE OF BASIC BIOLOGICAL RESEARCH**

This article provides a brief overview of the processes of drug discovery and development. Our aim is to help scientists whose research may be relevant to drug discovery and/or development to frame their research report in a way that appropriately places their findings within the drug discovery and development process and thereby support effective translation of preclinical research to humans. One overall theme of our article is that the process is sufficiently long, complex, and expensive so that many biological targets must be considered for every new medicine eventually approved for clinical use and new research tools may be needed to investigate each new target. Studies that contribute to solving any of the many scientific and operational issues involved in the development process can improve the efficiency of the process. An awareness of these issues allows the early implementation of measures to increase the opportunity for success. As editors of the journal, we encourage submission of research reports that provide data relevant to the issues presented.[7]

**MACHINE LEARNING CLASSIFICATION OVER ENCRYPTED DATA**

Machine learning classification is used for numerous tasks nowadays, such as medical or genomics predictions, spam detection, face recognition, and financial predictions. Due to privacy concerns, in some of these applications, it is important that the data and the classifier remain confidential. In this work, we construct three major classification protocols that satisfy this privacy constraint: hyperplane decision, Naïve Bayes, and decision trees. We also enable these protocols to be combined with AdaBoost. At the basis of these constructions is a new library of building blocks, which enables constructing a wide range of privacy-preserving classifiers; we demonstrate how this library can be used to construct other classifiers than the three mentioned above, such as a multiplexer and a face detection classifier. We implemented and evaluated our library and our classifiers. Our protocols are efficient, taking milliseconds to a few seconds to perform a classification when running on real medical datasets.[8]

**SAFE DRUG ADMINISTRATION: RIGHT OF EVERY PATIENT**

Medication errors are well known problem in hospitals. Registered nurses are accountable for their actions and omissions when administering any medication. To assess and compare the knowledge of nursing personnel regarding injection administration before and after demonstration of safe injection administration techniques. Experimental research approach was used with one group pretest- posttest design. Data was collected from 163 nursing personnel working in different ward of the hospital were which was selected by using total Enumeration sampling technique. The mean with standard deviation post-implementation knowledge score (8.0+1.5) was higher than the mean with standard deviation of pre-implementation knowledge scoring (4.2+1.5) scores. “t” value” calculated between the pre-implementation practice sores and post implementation practice scores with a df (162) was found to be statistically significant at 0.00\*\*(p>0.05). Thus, the study showed that along with theoretical education, special attention must be paid to the practical aspect of education. Demonstration and other such educational programs are very effective methods in updating the knowledge of the health care professionals.[9]

**NEEDLE FREE DRUG DELIVERY DEVICES MARKET SIZE, INDUSTRY OUTLOOK AND OPPORTUNITY ANALYSIS REPORT**

According to the global Needle-Free Drug Delivery Devices Market report published by Value Market Research, the market is expected to touch USD XX.X MN by 2025, with a CAGR of X.X% growing from valued USD XX.X MN (by revenue) in 2018. This is a tailored made research service providing informative data and various critical aspects of the market such as market outlook, market share, growth, and trends. Further, the report also offers evidence-based information that helps to transform clients business and achieve their business goals. Moreover, the report also highlights the key strategy of top players. Additionally, this report covers a wide spectrum of services such as the latest technology trend, market opportunity analysis, and competitive landscape.[10]

**2.11 COMPARISON TABLE**

**Table 2.1 Comparison Table for Literature Survey**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **TITLE** | **AUTHOR** | **YEAR** | **INFERENCE** | **DRAWBACKS** |
| Deep Learning in Drug Discovery and Medicine; Scratching the Surface | Dibyendu Dana , Satishkumar V. Gadhiya , Luce G. St. Surin , David Li , Farha Naaz ,Quaisar Ali | British journal of pharmacology, vol. 162, no. 6, pp. 1239–1249, 2018 | Drug discovery and medicine that started with Ayurveda in India are now benefiting from an altogether different artificial intelligence (AI)—one which is automating the invention of new chemical entities and the mining of large databases in health-privacy-protected vaults. Indeed, disciplines as diverse as language, chemistry, toxicology, biostatistics, medicine and computing have come together to harness algorithms based on transfer learning and recurrent neural networks to design novel drug candidates but only on demand. | 1)The molecular profile of an individual patient drives the selection  of therapy.  2)Post-marketing surveillance data show that its benefit is not experienced by every patient. |
| A Novel Neutrosophic Weighted Extreme Learning Machine for Imbalanced Data Set | Yaman Akbulut 1 ID , Abdulkadir ¸ Sengür 1,\* ID , Yanhui Guo 2 and Florentin Smarandache | Journal of chemical information and computer sciences, vol. 44, no. 5, pp. 1630–1638, 2017 | It has advantages such as good generalization performance, fast learning speed, and low computational cost.  Neutrosophic weighted extreme learning machine (NWELM), in which neutrosophic c-means (NCM) clustering algorithm is used for the approximation of the output weights of the ELM. | 1)Problems in the classification of imbalanced data sets.  2)ELM suffers from the presence of irrelevant variables in the large and high dimensional real data set |
| Drug discovery and development: Role of basic biological research | Richard C. Mohsa, Nigel H. Greig | IEEE Journal of Biomedical and Health Informatics, vol. 20, pp. 655 – 668, 2017 | Better medicines that are iterative improvements on current medications are valuable as they may offer benefits over existing medications in terms of potency, safety, tolerability, or convenience.  The drug discovery and development process and thereby support effective translation of preclinical research to humans | The process is sufficiently long, complex, and expensive so that many biological targets must be considered for every new medicine eventually approved for clinical use and new research tools may be needed to investigate each new target. |
| Needle Free Drug Delivery Devices Market Size, Industry Outlook and Opportunity Analysis Report 2018-2025 | Aniket Sharma | 8th ACM SIGSAC symposium on Information, computer and communications security. ACM, 2013, pp. 541–546.2019 | The global Needle-Free Drug Delivery Devices Market report is a tailored made research service providing informative data and various critical aspects of the market such as market outlook, market share, growth, and trends.Moreover, the report also highlights the key strategy of top players. Additionally, this report covers a wide spectrum of services such as the latest technology trend, market opportunity analysis, and competitive landscape | This Needle free drug discovery is yet to be developed and to carry on this discovery we need advanced expensive technology. |
| Drug Design and Discovery: Principles  and Applications | Shu-Feng Zhou , Wei-Zhu Zhong | Drug discovery today, vol. 17, no. 19, pp. 1088–1102, 2017 | Drug discovery is the process through which potential new therapeutic entities are identified, using a combination of computational, experimental, translational, and clinical models.  Modern drug discovery involves the identification of screening hits, medicinal chemistry and optimization of those hits to increase the affinity, selectivity (to reduce the potential of side effects), efficacy/potency, metabolic stability (to increase the half-life), and oral bioavailability | Despite advances in biotechnology and understanding of biological systems, drug discovery is still a lengthy, costly, difficult, and inefficient process with a high attrition rate of new therapeutic discovery. |
| Privacy-Preserving Clinical Decision Support System Using Gaussian Kernel- Based Classification | Yogachandran Rahulamathavan, Suresh Veluru, Raphael | Biomedical and Health Informatics, IEEE Journal of, vol. 18, no. 1, pp. 56–66, 2015 | A clinical decision support system forms a critical capability to link health observations with health knowledge to influence choices by clinicians for improved healthcare.  Remote outsourcing can be exploited to provide efficient and accurate clinical decision support in healthcare. | Servers are third party and therefore potentially not fully trusted raises possible privacy concerns. |
| Machine Learning Classification over Encrypted Data | Raphael Bost, Raluca Ada Popa, Stephen Tu | 22nd Annual Network and Distributed System Security Symposium, NDSS 2015, San Diego, California, USA, February 8-11, 2015. | Machine learning classification is used for numerous tasks nowadays, such as medical or genomics predictions,spam detection, face recognition, and financial predictions. Due  to privacy concerns, in some of these applications, it is important  that the data and the classifier remain confidential.Three major classification protocols that satisfy this privacy constraint: hyperplane decision, Naïve Bayes, and decision trees.  The basis of these constructions is a new library of building blocks, which enables constructing a wide range of privacy-preserving classifiers. | In the most basic sense, drug design involves the design of molecules that are complementary in shape and charge to the molecular target with which they interact and bind.  No security for drug components while train the data set. |
| Privacy-Preserving Multi-Class Support Vector Machine for Outsourcing the Data Classification in Cloud | Kanapathippillai Cumanan, Muttukrishnan Rajarajan | IEEE on Dependable and Secure Computing, vol. 11, no. 5, pp. 467–479, 2014 | Classifier built by the server can be utilized by clients in order to classify their own data samples over the cloud.  A privacy-preserving (PP) data classification technique where the server is unable to learn any knowledge about clients’ input data samples while the server side classifier is also kept secret from the clients during the classification process | Emerging cloud computing infrastructure replaces traditional outsourcing techniques and provides flexible services to clients at different locations via Internet. This leads to the requirement for data classification to be performed by potentially untrusted servers in the cloud.Releasing the data samples owned by the clients to the cloud raises privacy concerns since the data processed in the cloud are often outsourced to untrusted-third-party-servers. |
| Machine learning methods in chemoinformatics | J. B. Mitchell | Wiley Interdisciplinary Reviews: Computational Molecular Science, . 4, no. 5, pp. 468–481, 2014. | Machine learning algorithms are generally developed in computer science or adjacent disciplines and find their way into chemical modeling by a process of diffusion. This discussion is concentrate on methods for supervised learning, predicting the unknown property values of a test set of instances, usually molecules, based on the known values for a training set. Particularly relevant approaches include Artificial Neural Networks, Random Forest, Support Vector Machine, k‐Nearest Neighbors and naïve Bayes classifiers. | Although numerous articles cited herein have compared performances of the various machine‐learning algorithms used in chemoinformatics, there is no single best method for all problems. The relative abilities of methods will depend on the size and distribution in chemical space of the dataset, the linearity etc. |
| Computer-aided drug design platform using pymol | M. A. Lill and M. L. Danielson | Journal of computer-aided molecular design, vol. 25, no. 1, pp. 13–19, 2011. | Over the decades, many powerful standalone tools for computer-aided drug discovery have been developed in academia providing insight into protein-ligand interactions. As programs are developed by various research groups, a consistent user-friendly graphical working environment combining computational techniques such as docking, scoring, molecular dynamics simulations, and free energy calculations is needed. | Despite advances in biotechnology and understanding of biological systems, drug discovery is still a lengthy, costly, difficult, and inefficient process with a high attrition rate of new therapeutic discovery. |

**SYSTEM**

**ANALYSIS**

**CHAPTER 3**

**SYSTEM ANALYSIS**

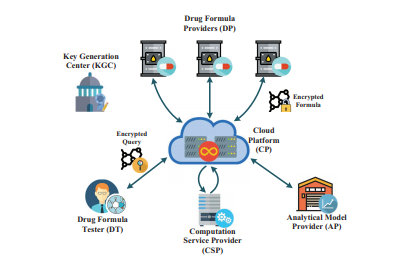
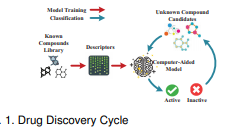
Systems design is the process of defining the architecture, components, modules, interfaces, and data for a system to satisfy specified requirements. Systems design could see it as the application of systems theory to product development.

**3.1 EXISTING SYSTEM**

In this existing system, securing drug components is designed to allow the cloud to securely use multiple drug formula providers’ drug formulas to train Support Vector Machine (SVM) provided by the analytical model provider. In this approach, it is designed to secure computation protocols to allow the cloud server to perform commonly used integer and fraction computations. To securely train the SVM, it is designed as a secure SVM parameter selection protocol to select two SVM parameters and construct a secure sequential minimal optimization protocol to privately refresh both selected SVM parameters. The trained SVM classifier can be used to determine whether a drug chemical compound is active or not in a privacy-preserving way. The existing datasets of known drug formulas to train the SVM classifier, and the trained SVM classifier can be used for new drug compound visual scanning. Due to the significant investments and high commercial values involved in drug discovery, privacy is an important factor. When a researcher sends some chemical compounds to the cloud for SVM classification, it is important to ensure that the potential new drug compounds will not be leaked to a third-party, such as a competing pharmaceutical corporation.

**3.2 PROPOSED SYSTEM**

We propose secure drug discovery components in the cloud environment. Unlike existing drug discovery frameworks, our POD seeks to achieve it efficiently. We are not using three real time datasets to check the efficiency of potential new drug component. Instead of using existing datasets, we are using another one data-mining algorithm Naïve Bayes (NB). It is a classification technique based on Bayes’ Theorem with an assumption of independence among predictors. NB classifier assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature also fast to predict class of test data set and performs well in multi class prediction. These two algorithms such as SVM and NB are used to train the uploaded drug dataset (CSV file).Thus we will get trained data and accuracy for that uploaded dataset. The trained data and accuracy will be sent to the owner from python server. Drug tester will check that new drug component. To check the new drug tester has to send request for accessing the drug component. As privacy is an important factor due to the significant investments and high commercial values involved in drug discovery, even drug tester does not know the contents of that file; they will get the trained data only. Once the Testing is completed, the tester sends result to the admin.If that particular drug component is still retained in the cloud then it is assumed that component is still active and passed the testing successfully. If not, then the drug component is removed from the cloud. Finally, admin will approve the drug component.

**Figure 3.1 Drug discovery cycle**

**3.3 REQUIREMENT ANALYSIS AND SPECIFICATION**

Requirement Specification is a complete description of the behavior of a system to be developed. It includes a set of use cases that describe all the interactions the users will have with the software. This chapter clearly depicts the software languages used in the system design and the significance of it.

The requirements specification is a technical specification of requirements for the software products. It is the first step in the requirements analysis process it lists the requirements of a particular software system including functional, performance and security requirements. The requirements also provide usage scenarios from a user, an operational and an administrative perspective.

The purpose of software requirements specification is to provide a detailed overview of the software project, its parameters and goals. This describes the project target audience and its user interface, hardware and software requirements. It defines how the client, team and audience see the project and its functionality.

**JAVA**

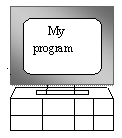
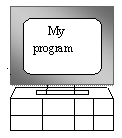
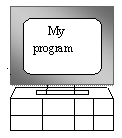
Java has been around since 1991, developed by a small team of Sun Microsystems developers in a project originally called the Green project. The intent of the project was to develop a platform-independent software technology that would be used in the consumer electronics industry. The language that the team created was originally called Oak. Web surfing has become an enormously popular practice among millions of computer users. Until Java, however, the content of information on the Internet has been a bland series of HTML documents. Web users are hungry for applications that are interactive, that users can execute no matter what hardware or software platform they are using, and that travel across heterogeneous networks and do not spread viruses to their computers. Java can create such applications. Java byte codes as the machine code instructions for the **Java Virtual Machine (JVM).** Every Java interpreter, whether it is a Java development tool or a Web browser that can run Java applets, is an implementation of JVM. That JVM can also be implemented in hardware. Java byte codes help make “write once, run anywhere” possible. Using compiler, compile the Java program into byte codes on any platform that has a Java compiler. The byte codes can then be run on any implementation of the JVM.

Complier

Interpreter

Interpreter

Interpreter

**  **

**PC-Compatible Sun Ultra Solaris Power macintosh**

**Figure 3.2 Java working process**

**APACHE TOMCAT SERVER**

Apache Tomcat (formerly under the Apache Jakarta Project; Tomcat is now a top-level project) is a web container developed at the Apache Software Foundation. Tomcat implements the servlet and the JavaServer Pages (JSP) specifications from Sun Microsystems, providing an environment for Java code to run in cooperation with a web server. It adds tools for configuration and management but can also be configured by editing configuration files that are normally XML-formatted. Because Tomcat includes its own HTTP server internally, it is also considered a standalone web server.

**DATA MINING**

In general terms, **“Mining”** is the process of extraction of some valuable material from the earth e.g. coal mining, diamond mining etc. In the context of computer science, **“**Data Mining**”** refers to the extraction of useful information from a bulk of data or [data warehouses](https://www.geeksforgeeks.org/data-warehousing/), the information gathered from Data Mining helps to predict hidden patterns, future trends and behaviors and allowing businesses to take decisions.

Technically, data mining is the computational process of analyzing data from different perspective, dimensions, angles and categorizing/summarizing it into meaningful information.

Data Mining can be applied to any type of data e.g. Data Warehouses, Transactional Databases, Relational Databases, Multimedia Databases, Spatial Databases, Time-series Databases, World Wide Web.

**SVM (Support Vector Machine)**

Support Vector Machine or SVM is one of the most popular Supervised Learning algorithms, which is used for Classification as well as Regression problems. However, primarily, it is used for Classification problems in Machine Learning.

The goal of the SVM algorithm is to create the best line or decision boundary that can segregate n-dimensional space into classes so that we can easily put the new data point in the correct category in the future. This best decision boundary is called a hyperplane.

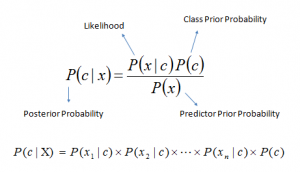
SVM chooses the extreme points/vectors that help in creating the hyperplane. These extreme cases are called as support vectors, and hence algorithm is termed as Support Vector Machine.

**Naive Bayes algorithm**

It is a classification technique based on Bayes’ Theorem with an assumption of independence among predictors. In simple terms, a Naive Bayes classifier assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature. For example, a fruit may be considered an apple if it is red, round, and about 3 inches in diameter. Even if these features depend on each other or upon the existence of the other features, all of these properties independently contribute to the probability that this fruit is an apple and that is why it is known as ‘Naive’.

Naive Bayes model is easy to build and particularly useful for very large data sets. Along with simplicity, Naive Bayes is known to outperform even highly sophisticated classification methods.

Bayes theorem provides a way of calculating posterior probability P (c|x) from P(c), P(x) and P(x|c). Look at the equation below:



Above,

* P(c|x) is the posterior probability of class (c, target) given predictor (x, attributes).
* P(c) is the prior probability of class.
* P(x|c) is the likelihood which is the probability of predictor given class.
* P(x) is the prior probability of predictor.

**3.4 HARDWARE REQUIREMENTS**

* Hard Disk : 80GB and Above
* RAM : 4GB and Above
* Processor : P IV and Above

**3.5 SOFTWARE REQUIREMENTS**

* Windows 7 and above(64 bit)
* JDK 1.8
* Python 3.6.3
* Tomcat 9.0.26
* MySQL

**3.6 TECHNOLOGIES USED**

* JAVA
* Spring Framework

**SYSTEM**

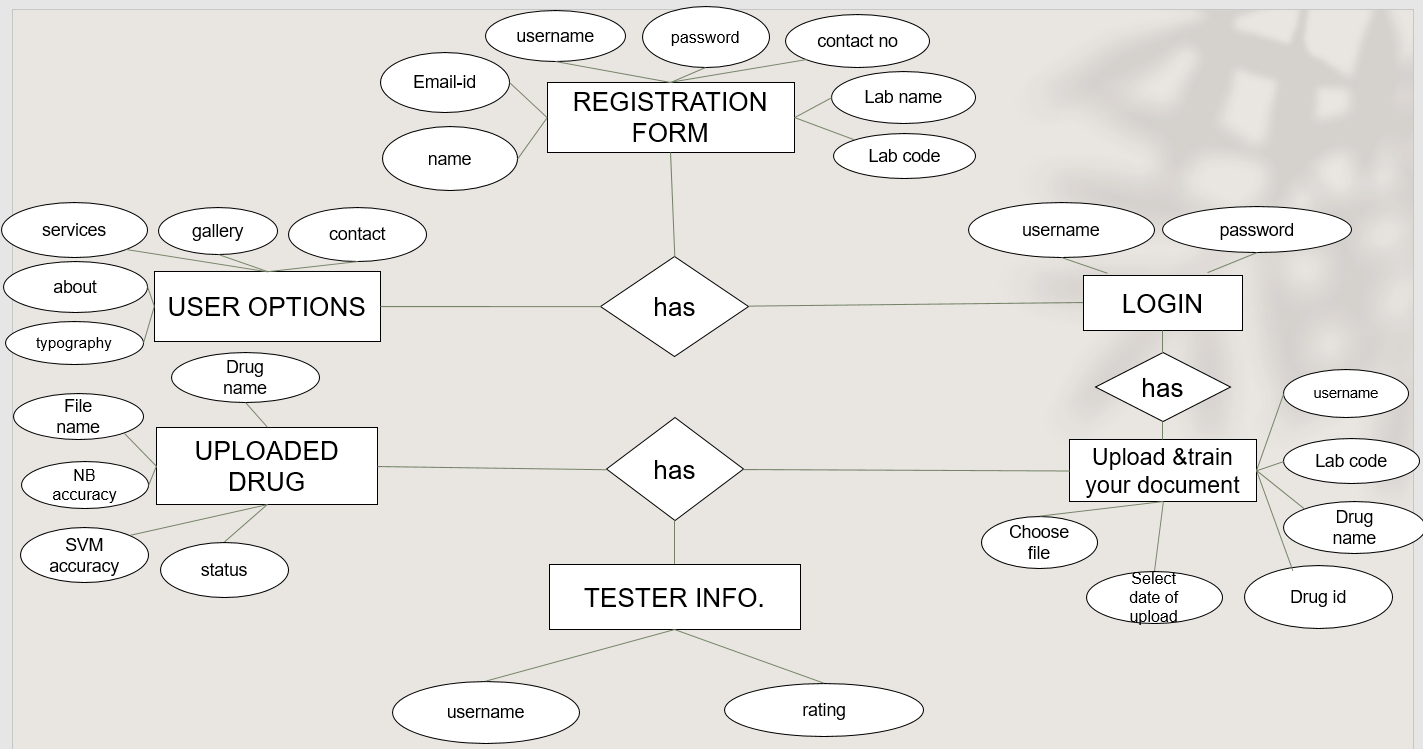
**DESIGN**

**CHAPTER 4**

**SYSTEM DESIGN**

**4.1 E-R DIAGRAM**

An entity relationship model, also called an entity-relationship (ER) diagram, is a graphical representation of entities and their relationships to each other, typically used in computing about the organization of data within databases or information systems.

****

**Figure 4.1 E-R Diagram for Drug Secure System**

**4.2 DATA DICTIONARY**

A data dictionary, or metadata repository, as defined as a centralized repository of information about data such as meaning, relationships to other data, origin, usage, and format. It is also defined as a collection of tables with metadata. The term can have one of several closely related meanings pertaining to databases. A document describing a database or collection of databases. An integral component of a DBMS that is required to determine its structure. A piece of middleware that extends or supplants the native data dictionary of a DBMS.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **FIELD NAME** | **DATATYPE** | **LENGTH** | **PRIMARY KEY** | **CONSTRAINS** |
| Username | Varchar | 255 | Yes | Primary key |
| Password | Varchar | 255 | No | Not null |
| Name | Varchar | 255 | No | Not null |
| Email | Varchar | 255 | Yes | Primary key |
| Phone | Varchar | 255 | Yes | Primary Key |
| Status | Varchar | 255 | No | Not null |
| Labname | Varchar | 255 | No | Not null |
| Labcode | Varchar | 255 | Yes | Primary Key |
| Drugname | Varchar | 255 | No | Not null |
| Drugid | Varchar | 255 | Yes | Primary Key |
| Date | Date | 255 | No | Not null |

**Table 4.1 Fields used in forms**

**4.3 TABLE NORMALIZATION**

**Normalization** is a database design technique that reduces data redundancy and eliminates undesirable characteristics like Insertion, Update and Deletion Anomalies. Normalization rules divides larger tables into smaller tables and links them using relationships. The purpose of Normalization in SQL is to eliminate redundant (repetitive) data and ensure data is stored logically. The inventor of the relational model Edgar Codd proposed the theory of normalization of data with the introduction of the First Normal Form, and he continued to extend theory with Second and Third Normal Form. Later he joined Raymond F. Boyce to develop the theory of Boyce-Codd Normal Form.

**4.3.1 FIRST NORMAL FORM**

First normal form (1NF) is a property of a relation in a relational database. A relation is in first normal form if and only if the domain of each attribute contains only atomic (indivisible) values, and the value of each attribute contains only a single value from that domain .The first definition of the term, in a 1971 conference paper by Edgar Codd, defined a relation to be in first normal form when none of its domains have any sets as elements .These two definitions do not conflict each other. First normal form is an essential property of a relation in a relational database. Database normalization is the process of representing a database in terms of relations in standard normal forms, where first normal is a minimal requirement.

First normal form enforces these criteria:

1)Every column/attribute must be unique in each table

2)Create a separate table for each set of related data[definition needed]

3)All entries must be single-valued and atomic



**Table 4.2 First normalized table**

**4.3.2 SECOND NORMAL FORM**

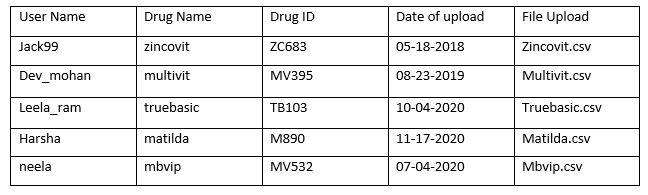
Second normal form (2NF) is a normal form used in database normalization. 2NF was originally defined by E. F. Codd in 1971.A relation is in the second normal form if it fulfills the following two requirements :

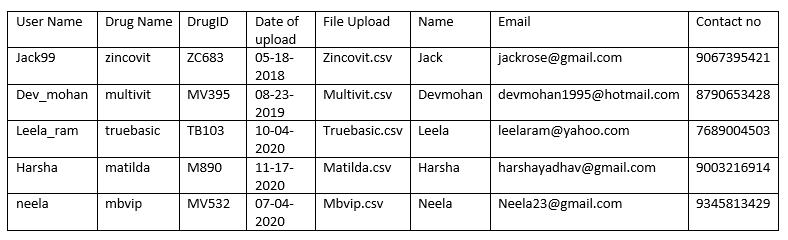
1)It is in first normal form.

2)It does not have any non-prime attribute that is functionally dependent on any proper subset of any candidate key of the relation. A non-prime attribute of a relation is an attribute that is not a part of any candidate key of the relation.

Put simply, a relation is in 2NF if it is in 1NF and every non-prime attribute of the relation is dependent on the whole of every candidate key. Note that it does not put any restriction on the non-prime to non-prime attribute dependency. That is addressed in third normal form.







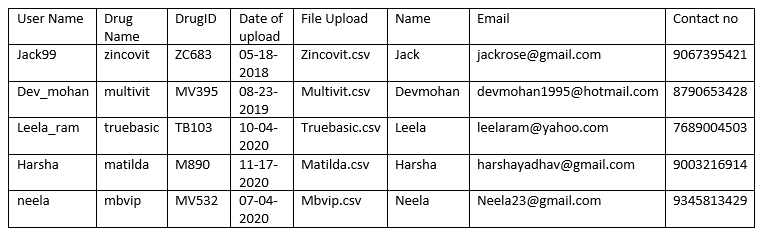
**Table 4.3 Second normalized table**

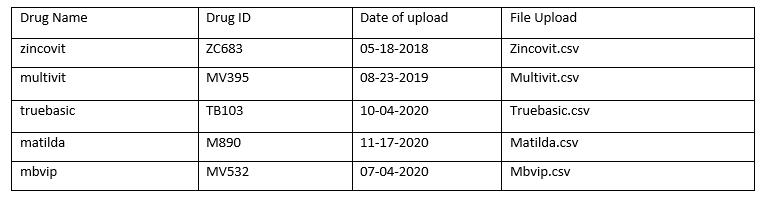
**4.3.3 THIRD NORMAL FORM**

Third normal form (3NF) is a database schema design approach for relational databases, which uses normalizing principles to reduce the duplication of data, avoid data anomalies, ensure referential integrity, and simplify data management. Edgar F. Codd, an English computer scientist who invented the relational model for database management, defined it in 1971. A relation is in third normal form, if there is no transitive dependency for non-prime attributes as well as it is in second normal form.

A relation is in 3NF if at least one of the following condition holds in every non-trivial function dependency X –> Y:

1. X is a super key.
2. Y is a prime attribute (each element of Y is part of some candidate key).







**Table 4.4 Third normalized table**

**4.4 DATA FLOW DIAGRAM**

A data flow diagram (DFD) is a graphical representation of the "flow" of data through an information system, modelling its process aspects. A DFD is often used as a preliminary step to create an overview of the system without going into detail, which can later be elaborated.

**4.4.1 CONTEXT LEVEL DIAGRAM**

Drug Owner

Registration

Login

Drug Tester

Registration

Login

**Figure 4.2 Context Level Diagram for registration**

**4.4.2 LEVEL 1 DFD**

Drug Owner

Registration

Login

Drug Tester

Registration

Login

Upload Drug Details

Train the drug document using svm and nb

**Figure 4.3 Level 1 DFD for module description**

**4.4.3 LEVEL 2 DFD**

Drug Owner

Registration

Login

Drug Tester

Registration

Login

Upload Drug Details

Train the drug document using svm and nb

Send request to Test document

Accept or Delete Request



**Figure 4.4 Level 2 DFD for module description**

**4.4.4 LEVEL 3 DFD**

Drug Owner

Registration

Login

Drug Tester

Registration

Login

Upload Drug Details

Train the drug document using svm and nb

Send request to Test document

Accept or Delete Request

If Accepted Test the components

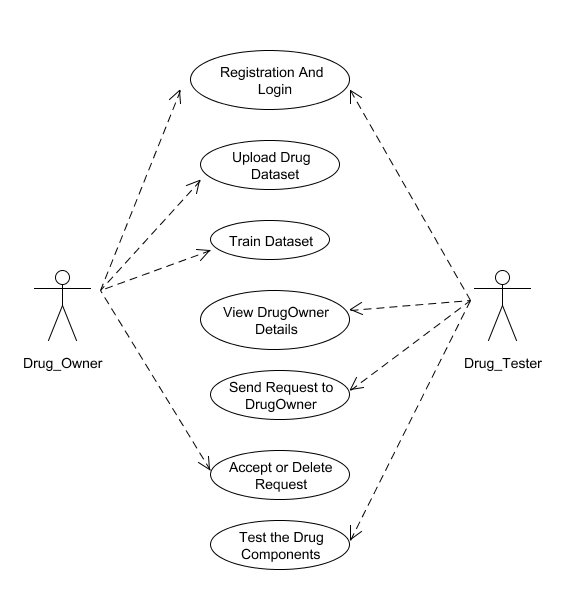
If Declined End the whole process

**Figure 4.5 Level 3 DFD for overall description**

**4.5 UML DIAGRAM**

**4.5.1 USE CASE DIAGRAM**

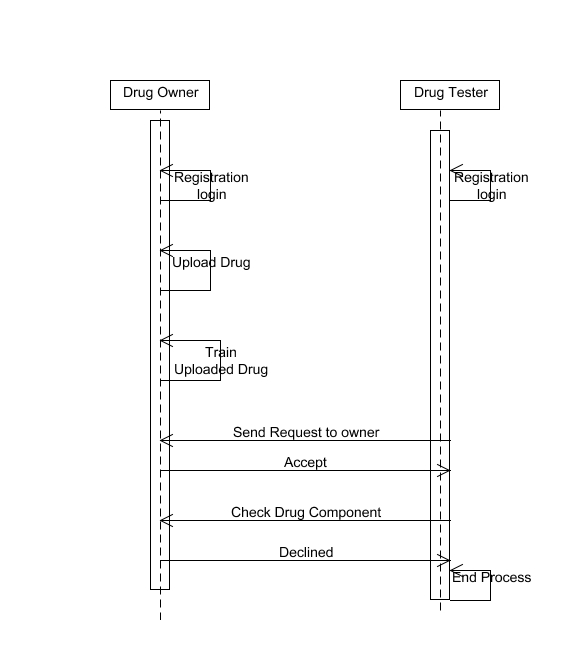
A use case diagram at its simplest is a representation of a user's interaction with the system and depicting the specifications of a use case. A use case diagram can portray the different types of users of a system and the various ways that they interact with the system. This type of diagram is typically used in conjunction with the textual use case and will often be accompanied by other types of diagrams as well.

****

**Figure 4.6 Use Case diagram for securing drug components**

**4.5.2 SEQUENCE DIAGRAM**

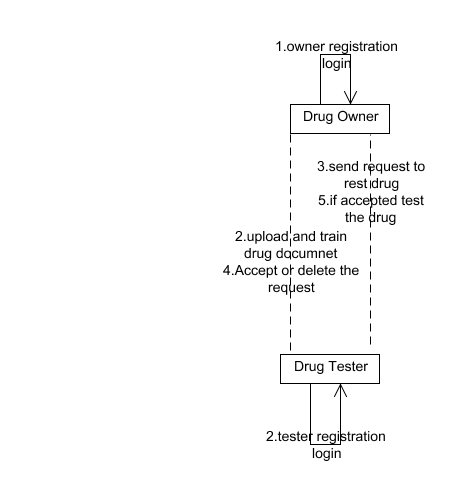
A sequence diagram is a kind of interaction diagram that shows how processes operate with one another and in what order. It is a construct of a Message Sequence Chart. A sequence diagram shows object interactions arranged in time sequence. It depicts the objects and classes involved in the scenario and the sequence of messages exchanged between the objects needed to carry out the functionality of the scenario. Sequence diagrams are typically associated with use case realizations in the Logical View of the system under development.

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**Figure 4.7 Sequence diagram for securing drug components**

**4.5.3 COLLABORATION DIAGRAM**

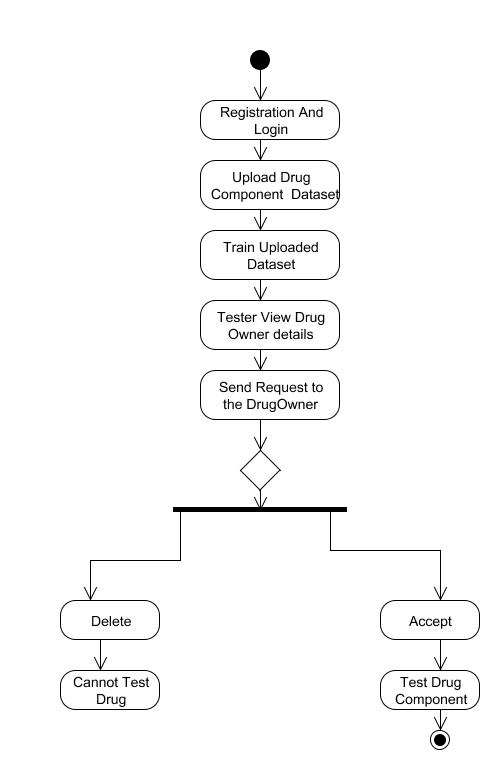
Collaboration diagrams offer a better view of a scenario than a Sequence diagram when the modeler is trying to understand all of the effects on a given object and are therefore good for procedural design. Objects are connected by links, each link representing an instance of an association between the respective classes involved. The link shows messages sent between the objects, and the type of message passed.

****

**Figure 4.8 Collaboration diagram for securing drug components**

**4.5.4 ACTIVITY DIAGRAM**

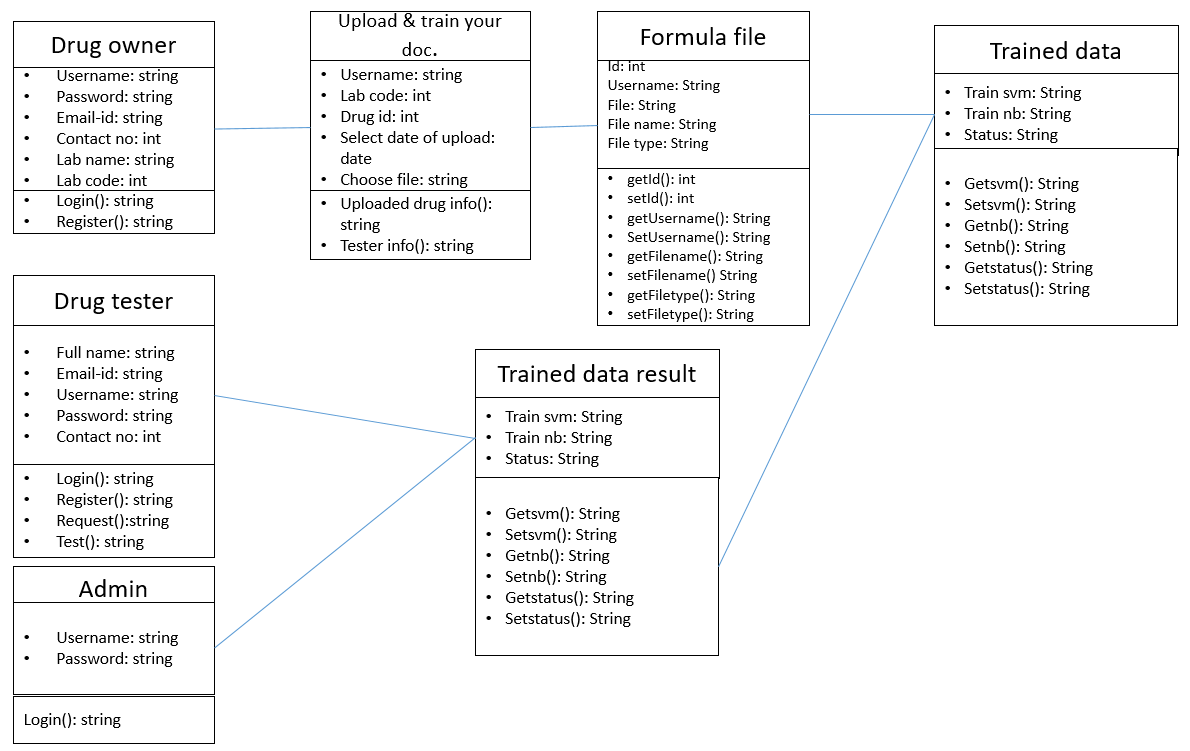
Activity diagrams are graphical representations of workflows of stepwise activities and actions with support for choice, iteration and concurrency. In the Unified Modeling Language, activity diagrams can be used to describe the business and operational step-by-step workflows of components in a system. An activity diagram shows the overall flow of control.

****

**Figure 4.9 Activity diagram for securing drug components**

**4.5.5 CLASS DIAGRAM**

Class Diagram provides an overview of the target system by describing the objects and classes inside the system and the relationships between them. It provides a wide variety of usages; from modeling the domain-specific data structure to detailed design of the target system. With the share model facilities, you can reuse your class model in the interaction diagram for modeling the detailed design of the dynamic behavior. The Form Diagram allows you to generate diagram automatically with user-defined scope.



**Figure 4.10 Class Diagram for securing drug components**

**SYSTEM**

**ARCHITECTURE**

**CHAPTER 5**

**SYSTEM ARCHITECTURE**

**5.1 ARCHITECTURE OVERVIEW**

LIST OF MODULES

* Drug Owner & Tester Registration
* Drug Component Uploading
* Train dataset
* Drug Testing

This system secure drug components for Secure Drug discovery in the cloud environment. Unlike drug discovery frameworks, the secure drug discovery seeks to achieve it efficiently. We are not using three real time datasets to check the efficiency of potential new drug component. Instead of using existing datasets, we are using another one data-mining algorithm Naïve Bayes (NB). These two algorithms are used to train the uploaded drug dataset (CSV file). In final, we will get trained data and accuracy for that uploaded dataset. Drug tester will check that new drug component. Drug tester does not know the contents of that file; they will get the trained data only. Then they let us know the file was active or not. Finally, admin will approve the drug component.



**Drug Owner**

**Drug Tester**

Registration & Login

Registration& Login



Upload Drug Dataset

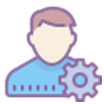
Store it into the cloud Server



Train dataset using dataming algorithms

Store it into the cloud server

Send Request to the owner if request has accepted tester can test the drug component



Test the drug component



Sent the results to admin

Login



View Results

**Admin**

**Figure 5.1 System Architecture for secure drug component system**

**5.2 MODULE DESIGN SPECIFICATION**

**5.2.1 DRUG OWNER AND TESTER REGISTRATION**

Drug Owner will register in the service provider platform. MySQL database is used to store the drugs Meta details. This drug owner registration process will involves few entities such as 1.Name(drug owner’s)2.Email(drug owner’s)3.Contact number(drug owner’s)4.lab name(where the drugs has been discovered)5.lab code(unique number of that lab) for this process.

The drug tester will also be registered in the drug tester registration platform. MySQL database is used to store the drugs Meta details. This drug tester registration process will involves few entities such as 1.Name(drug tester’s)2.Email(drug tester’s)3.Contact number(drug tester’s) and 4.TesterID( unique number of that tester) for this process.

**5.2.2 DRUG COMPONENT UPLOADING**

Once the Registration is completed, the drug owner should upload the drug component. For uploading the drug component, the owner must provide the Drug Name, Drug Id and Date of Uploading. Now we have to choose the file that contains the drug data sets and drug components. That data set contains the formula and we have to mention the type of class (Class A, Class B). While uploading the file we will read the content and store into the database and store that .csv file in cloud.

**5.2.3 TRAIN DATASET**

As the drug owner has successfully uploaded the drug components, now we will train the uploaded data using python. For this part, we will use two algorithms, SVM and Naïve Bayes. The trained data and accuracy will be sent to the owner from python server.

**5.2.4 DRUG TESTING**

As the drug dataset has been trained and uploaded in the cloud, the drug tester now send request to the owner for testing. Only after accepting the request, the test can test the uploaded drug components. Once the Testing is completed, the drug tester sends result to the admin.If that particular drug component is retained in the cloud then it is assumed that component is still active and passed the testing successfully. If not, then the drug component is removed from the cloud.

**5.3 PROGRAM DESIGN LANGUAGE**

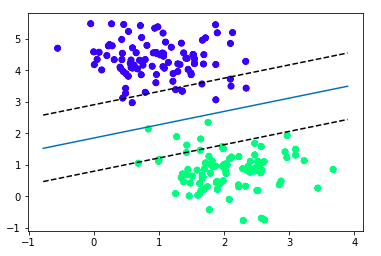
**5.3.1 SVM (Support Vector Machine)**

Support Vector Machine or SVM is one of the most popular Supervised Learning algorithms, which is used for Classification as well as Regression problems. However, primarily, it is used for Classification problems in Machine Learning.

The goal of the SVM algorithm is to create the best line or decision boundary that can segregate n-dimensional space into classes so that we can easily put the new data point in the correct category in the future. This best decision boundary is called a hyperplane.

SVM chooses the extreme points/vectors that help in creating the hyperplane. These extreme cases are called as support vectors, and hence algorithm is termed as Support Vector Machine.

class SVM:def fit(self, X, y):  
 n\_samples, n\_features = X.shape# P = X^T X  
 K = np.zeros((n\_samples, n\_samples))  
 for i in range(n\_samples):  
 for j in range(n\_samples):  
 K[i,j] = np.dot(X[i], X[j])P = cvxopt.matrix(np.outer(y, y) \* K)# q = -1 (1xN)  
 q = cvxopt.matrix(np.ones(n\_samples) \* -1)# A = y^T   
 A = cvxopt.matrix(y, (1, n\_samples))# b = 0   
 b = cvxopt.matrix(0.0)# -1 (NxN)  
 G = cvxopt.matrix(np.diag(np.ones(n\_samples) \* -1))# 0 (1xN)  
 h = cvxopt.matrix(np.zeros(n\_samples))solution = cvxopt.solvers.qp(P, q, G, h, A, b)# Lagrange multipliers  
 a = np.ravel(solution['x'])# Lagrange have non zero lagrange multipliers  
 sv = a > 1e-5  
 ind = np.arange(len(a))[sv]  
 self.a = a[sv]  
 self.sv = X[sv]  
 self.sv\_y = y[sv]# Intercept  
 self.b = 0  
 for n in range(len(self.a)):  
 self.b += self.sv\_y[n]  
 self.b -= np.sum(self.a \* self.sv\_y \* K[ind[n], sv])  
 self.b /= len(self.a)# Weights  
 self.w = np.zeros(n\_features)  
 for n in range(len(self.a)):  
 self.w += self.a[n] \* self.sv\_y[n] \* self.sv[n]  
   
 def project(self, X):  
 return np.dot(X, self.w) + self.b  
   
   
 def predict(self, X):  
 return np.sign(self.project(X))

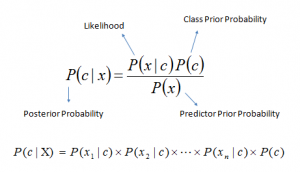


**Figure 5.2 2D Diagram of SVM algorithm**

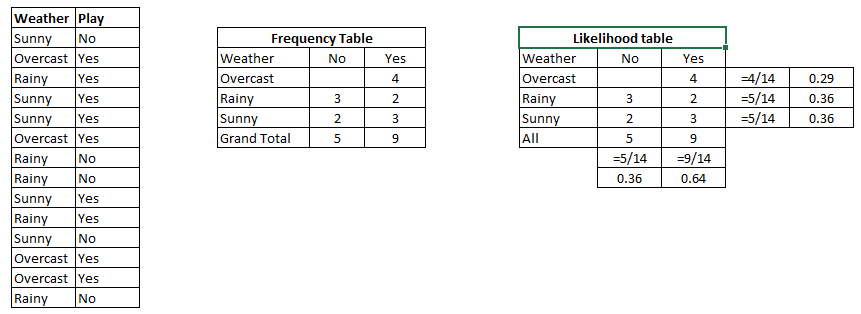
**5.3.2 Naïve Bayes**

It is a classification technique based on Bayes’ Theorem with an assumption of independence among predictors. In simple terms, a Naive Bayes classifier assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature. For example, a fruit may be considered to be an apple if it is red, round, and about 3 inches in diameter. Even if these features depend on each other or upon the existence of the other features, all of these properties independently contribute to the probability that this fruit is an apple and that is why it is known as ‘Naive’.

Bayes theorem:



Let’s understand it using an example. Below we have a training data set of weather and corresponding target variable ‘Play’ (suggesting possibilities of playing). Now, we need to classify whether players will play or not based on weather condition.



**Problem:**Players will play if weather is sunny. Is this statement is correct?

We can solve it using above discussed method of posterior probability.

P(Yes | Sunny) = P( Sunny | Yes) \* P(Yes) / P (Sunny)

Here we have P (Sunny |Yes) = 3/9 = 0.33, P(Sunny) = 5/14 = 0.36, P( Yes)= 9/14 = 0.64

Now, P (Yes | Sunny) = 0.33 \* 0.64 / 0.36 = 0.60, which has higher probability.

Naive Bayes uses a similar method to predict the probability of different class based on various attributes. This algorithm is mostly used in text classification and with problems having multiple classes.

**SYSTEM IMPLEMENTATION**

**CHAPTER 6**

**SYSTEM IMPLEMENTATION**

**6.1 CLIENT-SIDE CODING**

import org.springframework.stereotype.Controller;

import org.springframework.validation.BindingResult;

import org.springframework.web.bind.annotation.PathVariable;

import org.springframework.web.bind.annotation.RequestMapping;

import org.springframework.web.bind.annotation.RequestMethod;

import org.springframework.web.servlet.ModelAndView;

import com.drug.dao.OwnerDao;

import com.drug.model.OwnerModel;

import com.drug.model.testerModel;

@Controller

public class TesterController {

@Autowired

private OwnerDao dao;

@RequestMapping(value = "/testerInfo/testerView/{testername}/{username}", method = RequestMethod.GET)

public ModelAndView tester(@PathVariable("testername") String testername,

@PathVariable("username") String username) {

ModelAndView model = new ModelAndView();

List<testerModel> tester = dao.getTester(testername);

List<testerModel> list = dao.getTester();

List<OwnerModel> owner = dao.findbyid(username);

if (tester != null) {

model.addObject("msg", tester);

model.addObject("owner", owner);

model.setViewName("testerslist");

} else {

model.addObject("msg1", list);

model.setViewName("testerPage");

}

return model;

}

@RequestMapping(value = "/testerInfo/testerView/{testername}/accept/{username}/{testername}",method = RequestMethod.GET)

public ModelAndView acceptRequest(@PathVariable("username") String username,

@PathVariable("testername") String testername) {

ModelAndView model = new ModelAndView();

String status="Accepted";

int count=dao.updateOwner(username, status);

int counter=dao.updateTester(testername, status);

if(count>0) {

model.setViewName("Success");

}

return model;

}

@RequestMapping(value = "/testerInfo/testerView/{testername}/delete/{username}/{testername}",method = RequestMethod.GET)

public ModelAndView deleteRequest(@PathVariable("username") String username,

@PathVariable("testername") String testername) {

ModelAndView model = new ModelAndView();

String status="No action Taken";

String status1="No action";

int count=dao.updateOwner(username, status);

int counter=dao.updateTester(testername, status1);

if(count>0) {

model.setViewName("Awesome");

}

return model;

}

}

package com.drug.controller;

import java.io.BufferedReader;

import java.io.InputStreamReader;

import java.net.HttpURLConnection;

import java.util.ArrayList;

import java.util.List;

import org.apache.http.HttpResponse;

import org.apache.http.NameValuePair;

import org.apache.http.client.HttpClient;

import org.apache.http.client.entity.UrlEncodedFormEntity;

import org.apache.http.client.methods.HttpPost;

import org.apache.http.impl.client.DefaultHttpClient;

import org.apache.http.message.BasicNameValuePair;

@SuppressWarnings("deprecation")

public class PythonCall {

@SuppressWarnings("deprecation")

public List<String> executeMultiPartRequest(String urlString, String path) throws Exception {

String output = null;

List<String> respons = new ArrayList<String>();

HttpClient client = new DefaultHttpClient();

HttpPost postRequest = new HttpPost(urlString);

ArrayList<NameValuePair> postParameters;

try {

System.out.println("============path===========" + path);

// Send request

postParameters = new ArrayList<NameValuePair>();

postParameters.add(new BasicNameValuePair("path", path));

// postParameters.add(new BasicNameValuePair("courseid", courseid));

// postParameters.add(new BasicNameValuePair("activityid", actid));

postRequest.setEntity(new UrlEncodedFormEntity(postParameters, "UTF-8"));

HttpResponse response = client.execute(postRequest);

System.out.println("-Response-----" + response.getStatusLine().getStatusCode());

// Verify response if any

if (response != null) {

BufferedReader br = new BufferedReader(new InputStreamReader((response.getEntity().getContent()))); // Getting

System.out.println("Output from Server .... \n");

while ((output = br.readLine()) != null) {

respons.add(output);

System.out.println(output); // Instead of this, you could append all your response to a StringBuffer

// and use `toString()` to get the entire JSON response as a String.

// This string json response can be parsed using any json library. Eg. GSON from

// Google.

br.close();

return respons;

}

} else {

return null;

}

} catch (Exception ex) {

return null;

}

return respons;

}

}

**6.2 SERVER-SIDE CODING**

package com.drugTest.controller;

import java.io.InputStream;

import java.util.List;

import org.springframework.beans.factory.annotation.Autowired;

import org.springframework.stereotype.Controller;

import org.springframework.web.bind.annotation.PathVariable;

import org.springframework.web.bind.annotation.RequestMapping;

import org.springframework.web.bind.annotation.RequestMethod;

import org.springframework.web.servlet.ModelAndView;

import com.drugTest.dao.testerDao;

import com.drugTest.model.FormulaFile;

import com.drugTest.model.OwnerModel;

import com.drugTest.model.Requestmodel;

import com.drugTest.model.Traineddata;

@Controller

public class DrugOwnerController {

@Autowired

private testerDao dao;

@RequestMapping(value = "/ownerDetails/{username}/{username1}/{status}", method = RequestMethod.GET)

public ModelAndView owner(@PathVariable("username") String username, @PathVariable("username1") String username1,

@PathVariable("status") String status) {

ModelAndView model = new ModelAndView();

List<OwnerModel> owner = dao.getOwnerList(username);

List<OwnerModel> list = dao.getOwnerList();

if (owner != null) {

model.addObject("msg", owner);

model.addObject("username", username1);

model.addObject("status", status);

model.setViewName("ownerDetails");

} else {

model.addObject("msg1", list);

model.setViewName("ownerPage");

}

return model;

}

@RequestMapping(value = "/fileDetails/{username}/{status}", method = RequestMethod.GET)

public ModelAndView file(@PathVariable("username") String username, @PathVariable("status") String status) {

ModelAndView model = new ModelAndView();

OwnerModel owner = new OwnerModel();

owner.setUsername(username);

owner.setTestername(status);

String result = dao.testerstatus(owner);

System.out.println("----result----" + result);

List<FormulaFile> file = dao.getFileList(username);

List<OwnerModel> list = dao.getOwnerList();

if (file != null) {

model.addObject("msg", file);

model.addObject("result", result);

model.setViewName("fileDetails");

} else {

model.addObject("msg1", list);

model.setViewName("ownerPage");

}

return model;

}

@RequestMapping(value = "ownerDetails/{username}/{username1}/request/{username1}/{username}", method = RequestMethod.GET)

public ModelAndView request(@PathVariable("username1") String testername,

@PathVariable("username") String username) {

ModelAndView model = new ModelAndView();

String status = "Request pending";

Requestmodel request = new Requestmodel();

request.setUsername(username);

request.setTestername(testername);

request.setStatus(status);

// OwnerModel owner=new OwnerModel();

// owner.setUsername(username);

// owner.setStatus(status);

// owner.setTestername(testername);

int result = dao.uodateStatus(username, status, testername);

int count = dao.updateStatus(testername, status);

System.out.println("------------------" + count);

int counter = dao.saveRequest(request);

if (count > 0) {

model.addObject("msg", "success");

model.setViewName("Success");

} else {

model.addObject("msg", "failed");

model.setViewName("ownerPage");

}

return model;

}

@RequestMapping(value = "fileDetails/{username}/testFile/{username}/{drugname}", method = RequestMethod.GET)

public ModelAndView testing(ModelAndView model, @PathVariable("drugname") String drugname,

@PathVariable("username") String username) {

String status="checked";

String statuss="Active";

List<Traineddata> dataList=dao.getDataList(username);

List<FormulaFile> fileList=dao.getFileList(drugname);

for (FormulaFile formulaFile : fileList) {

for (Traineddata traineddata : dataList) {

String dnb=formulaFile.getTrain\_nb();

String tnb=traineddata.getTrain\_NB();

String dvm=formulaFile.getTrain\_svm();

String tvm=traineddata.getTrain\_SVM();

if(dataList.isEmpty()) {

model.addObject("msg","Cannot Find the Data");

model.setViewName("fileDetails");

}else if (dnb!=null || dvm!=null || tnb!=null || tvm!=null) {

int result=dao.formulaStatus(drugname, status);

int count=dao.dataStatus(drugname, statuss);

System.out.println("------------------------");

model.addObject("msg","success");

model.setViewName("checked");

}

}

}

return model;

}

}

package com.gts.controller;

import java.util.List;

import org.springframework.beans.factory.annotation.Autowired;

import org.springframework.stereotype.Controller;

import org.springframework.web.bind.annotation.PathVariable;

import org.springframework.web.bind.annotation.RequestMapping;

import org.springframework.web.bind.annotation.RequestMethod;

import org.springframework.web.bind.annotation.RequestParam;

import org.springframework.web.servlet.ModelAndView;

import com.gts.dao.AdminDao;

import com.gts.model.Traineddata;

@Controller

public class AdminController {

@Autowired

private AdminDao dao;

@RequestMapping(value = "/adminlogin",method = RequestMethod.POST)

public ModelAndView admin(ModelAndView model, @RequestParam("username") String username,

@RequestParam("password") String password) {

System.out.println("====="+username+"====="+password);

if(username.equalsIgnoreCase("admin")&&password.equalsIgnoreCase("admin")) {

List<Traineddata> list=dao.getAlldata();

model.addObject("msg", list);

model.setViewName("dashboard");

}else {

model.setViewName("index");;

}

return model;

}

@RequestMapping(value = "/accept/{drugname}",method = RequestMethod.GET)

public ModelAndView OK(ModelAndView model,@PathVariable("drugname")String drugname) {

String status="Accepted";

int result=dao.accept(drugname, status);

int ok=dao.ok(drugname, status);

model.setViewName("dashboard");

return model;

}

@RequestMapping(value = "/decline/{drugname}",method = RequestMethod.GET)

public ModelAndView NO(ModelAndView model,@PathVariable("drugname")String drugname) {

String status="Decliened";

int result=dao.decline(drugname, status);

int notok=dao.notok(drugname, status);

model.setViewName("dashboard");

return model;

}

}

package com.gts.model;

public class FormulaFile {

private int id;

private String username;

private String labcode;

private String drugname;

private String drugid;

private String date;

private String filename;

private String file;

private String filetype;

private String train\_svm;

private String train\_nb;

private String status;

public String getStatus() {

return status;

}

public void setStatus(String status) {

this.status = status;

}

public String getTrain\_svm() {

return train\_svm;

}

public void setTrain\_svm(String train\_svm) {

this.train\_svm = train\_svm;

}

public String getTrain\_nb() {

return train\_nb;

}

public void setTrain\_nb(String train\_nb) {

this.train\_nb = train\_nb;

}

public FormulaFile() {

}

public String getDrugid() {

return drugid;

}

public void setDrugid(String drugid) {

this.drugid = drugid;

}

public int getId() {

return id;

}

public void setId(int id) {

this.id = id;

}

public String getUsername() {

return username;

}

public void setUsername(String username) {

this.username = username;

}

public String getLabcode() {

return labcode;

}

public void setLabcode(String labcode) {

this.labcode = labcode;

}

public String getDrugname() {

return drugname;

}

public void setDrugname(String drugname) {

this.drugname = drugname;

}

public String getDate() {

return date;

}

public void setDate(String date) {

this.date = date;

}

public String getFilename() {

return filename;

}

public void setFilename(String filename) {

this.filename = filename;

}

public String getFile() {

return file;

}

public void setFile(String file) {

this.file = file;

}

public String getFiletype() {

return filetype;

}

public void setFiletype(String filetype) {

this.filetype = filetype;

}

}

package com.gts.model;

public class Traineddata {

private int id;

private String username;

private String drugname;

private String filename;

private String accuracy\_nb;

private String accuracy\_svm;

private String Train\_SVM;

private String Train\_NB;

private String status;

public String getStatus() {

return status;

}

public void setStatus(String status) {

this.status = status;

}

public int getId() {

return id;

}

public void setId(int id) {

this.id = id;

}

public String getUsername() {

return username;

}

public void setUsername(String username) {

this.username = username;

}

public String getDrugname() {

return drugname;

}

public void setDrugname(String drugname) {

this.drugname = drugname;

}

public String getFilename() {

return filename;

}

public void setFilename(String filename) {

this.filename = filename;

}

public String getAccuracy\_nb() {

return accuracy\_nb;

}

public void setAccuracy\_nb(String accuracy\_nb) {

this.accuracy\_nb = accuracy\_nb;

}

public String getAccuracy\_svm() {

return accuracy\_svm;

}

public void setAccuracy\_svm(String accuracy\_svm) {

this.accuracy\_svm = accuracy\_svm;

}

public String getTrain\_SVM() {

return Train\_SVM;

}

public void setTrain\_SVM(String train\_SVM) {

Train\_SVM = train\_SVM;

}

public String getTrain\_NB() {

return Train\_NB;

}

public void setTrain\_NB(String train\_NB) {

Train\_NB = train\_NB;

}

}

package com.drugTest.model;

public class Requestmodel {

private String testername;

private String username;

private String status;

public Requestmodel() {

}

public String getTestername() {

return testername;

}

public void setTestername(String testername) {

this.testername = testername;

}

public String getUsername() {

return username;

}

public void setUsername(String username) {

this.username = username;

}

public String getStatus() {

return status;

}

public void setStatus(String status) {

this.status = status;

}

}

**SYSTEM**

**TESTING**

**CHAPTER 7**

**SYSTEM TESTING**

**7.1 UNIT TESTING**

Unit Testingis a level of the software testing process where individual units/components of a software/system are tested. The purpose is to validate that each unit of the software performs as designed. The goal of unit testing is to isolate each part of the program and show that the individual parts are correct. In our application, the modules are taken separately and tested. In the module of registration each input is denoted with specify constraint and the violation of this constraint will display error message. Thus, it verify the unit testing. Likewise, the other modules are tested separately to test their functionality. The developer usually performs unit testing.

**7.2 INTEGRATION TESTING**

Integration testing is a systematic technique for construction the program structure while at the same time conducting tests to uncover errors associated with interfacing. i.e., integration testing is the complete testing of the set of modules, which makes up the product. The objective is to take untested modules and build a program structure tester should identify critical modules. Critical modules should be tested as early as possible. One approach is to wait until all the units have passed testing, and then combine them and then tested. This approach is evolved from unstructured testing of small programs. Another strategy is to construct the product in increments of tested units. A small set of modules are integrated together and tested, to which another module is added and tested in combination. And so on. The advantages of this approach are that, interface dispenses can be easily found and corrected.

The major error that was faced during the project is linking error. When all the modules are combined, the link is not set properly with all support files. Then we checked out for interconnection and the links. Errors are localized to the new module and its intercommunications. The product development can be staged, and modules integrated in as they complete unit testing. Testing is completed when the last module is integrated and tested.

**7.3 TEST CASES & REPORTS**

Testing is a process of executing a program with the intent of finding an error. A good test case is one that has a high probability of finding an as-yet –undiscovered error. A successful test is one that uncovers an as-yet- undiscovered error. System testing is the stage of implementation, which is aimed at ensuring that the system works accurately and efficiently as expected before live operation commences. It verifies that the whole set of programs hang together. System testing requires a test consists of several key activities and steps for run program, string, system and is important in adopting a successful new system. This is the last chance to detect and correct errors before the system is installed for user acceptance testing. The software testing process commences once the program is created and the documentation and related data structures are designed. Software testing is essential for correcting errors. Otherwise the program or the project is not said to be complete. Software testing is the critical element of software quality assurance and represents the ultimate the review of specification design and coding. Testing is the process of executing the program with the intent of finding the error.

**TEST CASE DESIGN**

A test case is a detailed procedure that fully tests a feature or an aspect of a feature. Whereas the test plan describes what to test, a test case describes how to perform a particular test. You need to develop a test case for each test listed in the test plan. The below table shows the possible Inputs and their corresponding Expected and Obtained output for the given modules with their status.

**Table 7.1: Test Case Design**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Action** | **Input** | **Expected Output** | **Actual Output** | **Test Result** |
| Enter the username,password for registration. | Username:XXX  Password:  \*\*\* | XXX  \*\*\* | XXX  \*\*\* | passed |
| Compare username and password with registered field | Username:XXX  Password:  \*\*\* | It Redirects to drug component uploading page | It redirects todrug component uploading page | passed |
| Compare username and password with registered field | Username:XXy  Password:  \*\*\* | Invalid username and password. | Invalid username and password. | passed |
| Drug Owner should provide name, mail id ,contact no,Lab Name and Lab Id. | Name:Jayalakshmi  Mail id: [jay@gmail.com](mailto:jay@gmail.com)  Contact no:9876543210  Lab Name:ABC  Lab ID:1234 | These details will be stored in drug Owner database. | These details will be stored in drug owner database | passed |
| Drug Owner should provide name, mail id ,contact no,Lab Name and Lab Id.  ACTION | Name:Jayalakshmi  Mail id: [jay@gmail.com](mailto:jay@gmail.com)  Contact no:9876543210  Lab Name:ABC  Lab ID:1234  INPUT | Please match the Requested format for  Mail Id.  EXPECTED OUTPUT | Please match the requested format for  Mail Id.  ACTUAL OUTPUT | Passed  TEST RESULT |
| Drug Owner should provide name, mail id ,contact no,Lab Name and Lab Id. | Name:Jayalakshmi  Mail id: [jay@gmail.com](mailto:jay@gmail.com)  Contact no:9876543210  Lab Name:ABC  Lab ID:1234 | Please match the requested format for  Contact number. | Valid Us Please match the requested format for  Contact number.  er | passed |
| Drug Tester should provide name, mail id ,contact no and Tester Id. | Name:Saiveena  Mail id: saiveena@gmail.com  Contact no:9876543210  Tester ID:1234 | These details will be stored in drug Owner database | These details will be stored in drug Owner database | passed |
| Drug Tester should provide name, mail id ,contact no and Tester Id. | Name:Saiveena  Mail id: saiveena@gmail.com  Contact no:9876543210  Tester ID:1234 | Please match the requested format for  Mail Id. | Please match the requested format for  Mail Id. | passed |
| Drug Tester should provide name, mail id ,contact no and Tester Id. | Name:Saiveena  Mail id: saiveena@gmail.com  Contact no:9876543210  Tester ID:1234 | Please match the requested format for  Contact number. | Please match the requested format for  Contact number. | passed |
| Drug Owner should provide drug name, drug id, date of upload and drug component file. | Drug Name:XYZ  Drug ID:12345  Date of Upload:  MM-DD-YYYY  File:drug1.csv | These details will be stored in drug Owner database. | These details will be stored in drug owner database. | passed |
| Drug Owner should provide drug name, drug id ,date of upload and drug component file. | Drug Name:Zincovit  Drug ID:1234  Date of Upload:  06-22-2020  File:drug1.csv | Please match the requested format for Drug Id. | Please match the requested format for Drug Id. | passed |
| Drug Owner should provide drug name, drug id , date of upload and drug component file. | Drug Name:Zincovit  Drug ID:1234  Date of Upload:  22-06-2020  File:drug1.csv | Please match the requested format for Date of upload. | Please match the requested format for Date of upload. | passed |
| Drug Owner should provide drug name, drug id , date of upload and drug component file. | Drug Name:Zincovit  Drug ID:1234  Date of Upload:  22-06-2020  File:drug1.csv | Please match the requested format for  File upload. | Please match the requested format for file upload. | passed |

**CONCLUSION**

**CHAPTER 8**

**CONCLUSION**

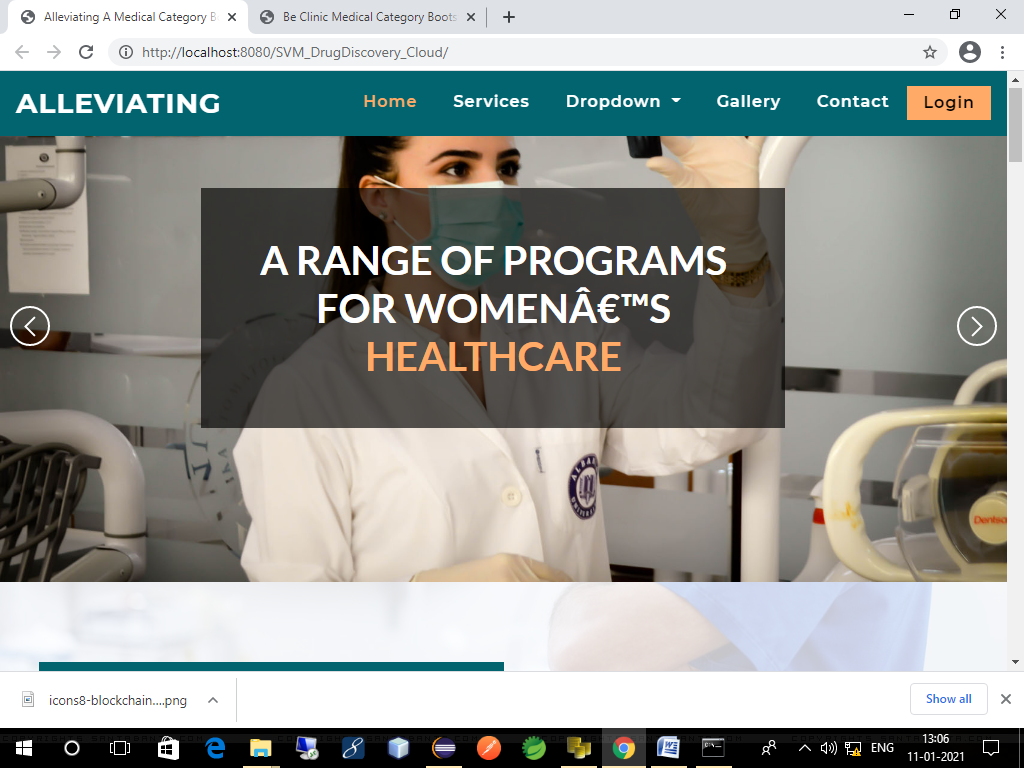
**8.1 CONCLUSION AND FUTURE ENHANCEMENTS**

This project focuses on securing the drug components in the cloud. Privacy is a major factor in drug discovery as it involves significant investment and high commercial values. Drug discovery is a long term expensive process. Bringing the drugs from the bench to the market involves a lot of threat when stored in the cloud. Thus, we proposed securing the drug components for drug discovery in the cloud. Securing the drug components is designed to facilitate drug manufacturers to securely outsource their formulas to the cloud for storage and SVM and NB training. This two algorithms such as SVM and NB are used to train the uploaded drug dataset (CSV file). As a result, we receive trained data and accuracy for that uploaded dataset. The trained model could be used for authorized client’s compound classification in a privacy-preserving way. Specifically, we designed a secure domain transformation protocol and several basic secure computation components for secure outsourced computation across different parties. We also built two key secure components (i.e. secure parameter selection and secure sequential minimal optimization) to achieve privacy-preserving SVM and NB training in drug discovery. We will be extending our approach to support more sophisticated data mining method in order to support very large dataset in drug discovery.

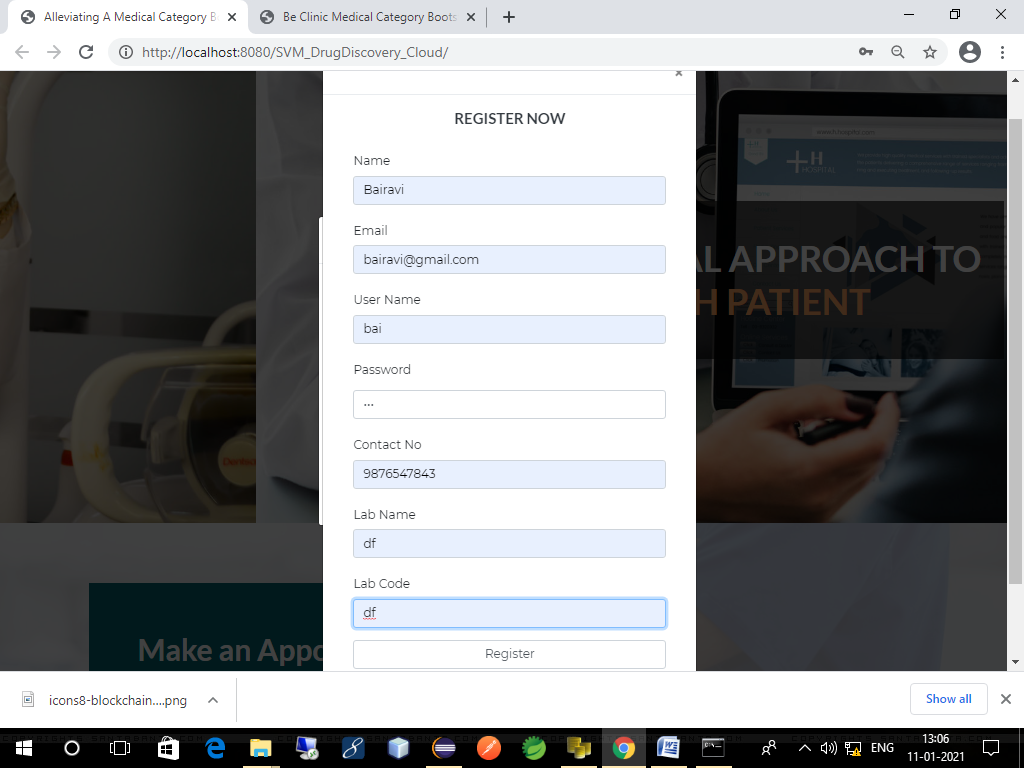
**APPENDICES**

**A.1 SAMPLE SCREENS**

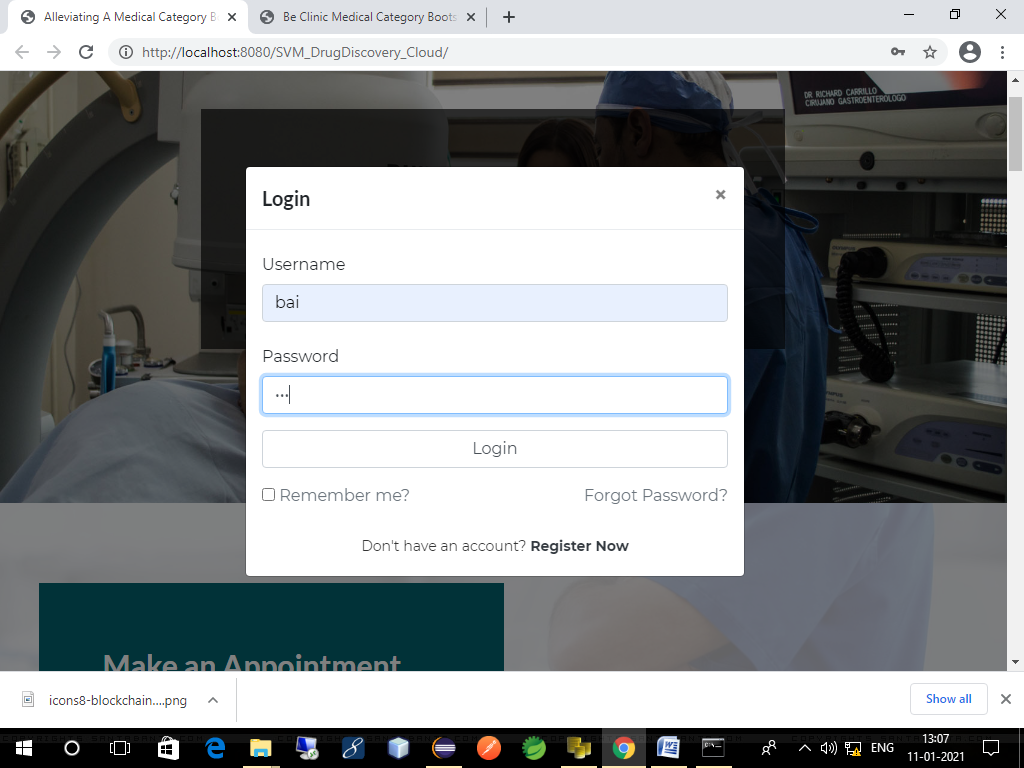
**HOME PAGE FOR DRUG OWNER**



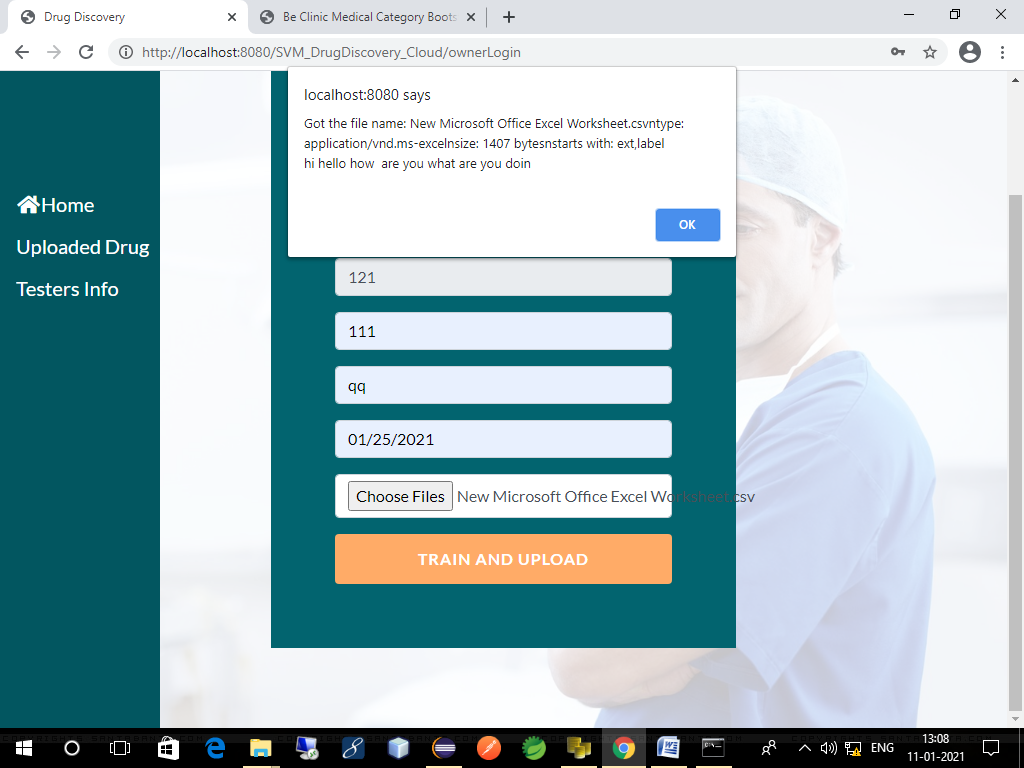
**REGISTRATION PAGE FOR DRUG OWNER**



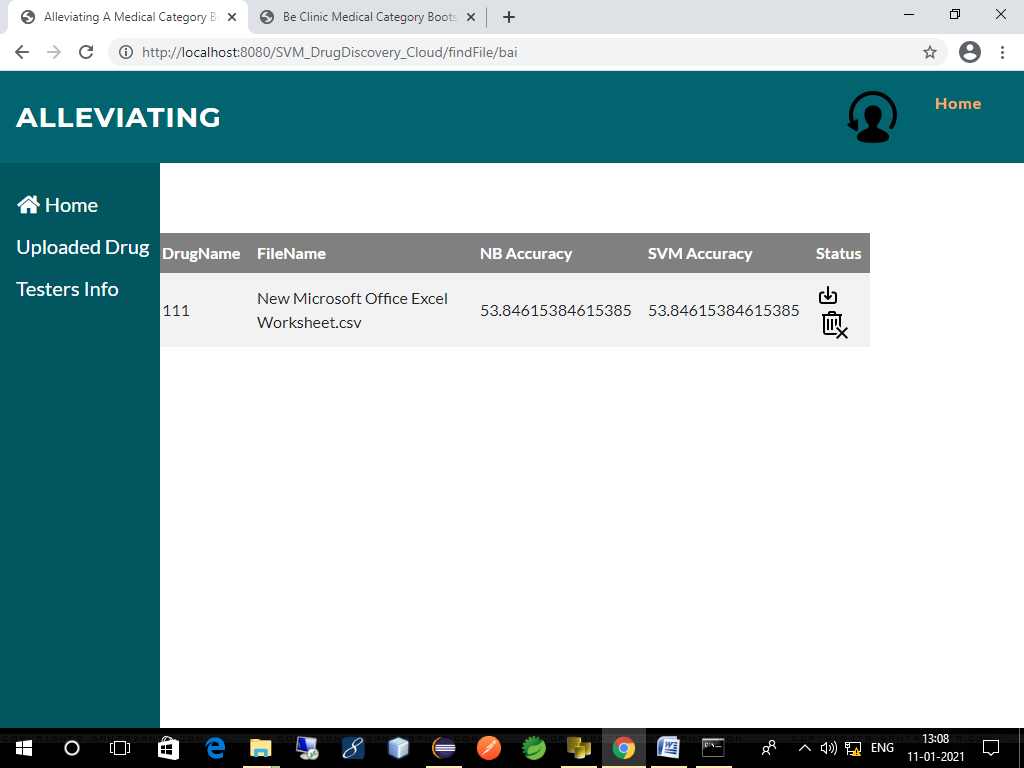
**LOGIN PAGE FOR DRUG OWNER**



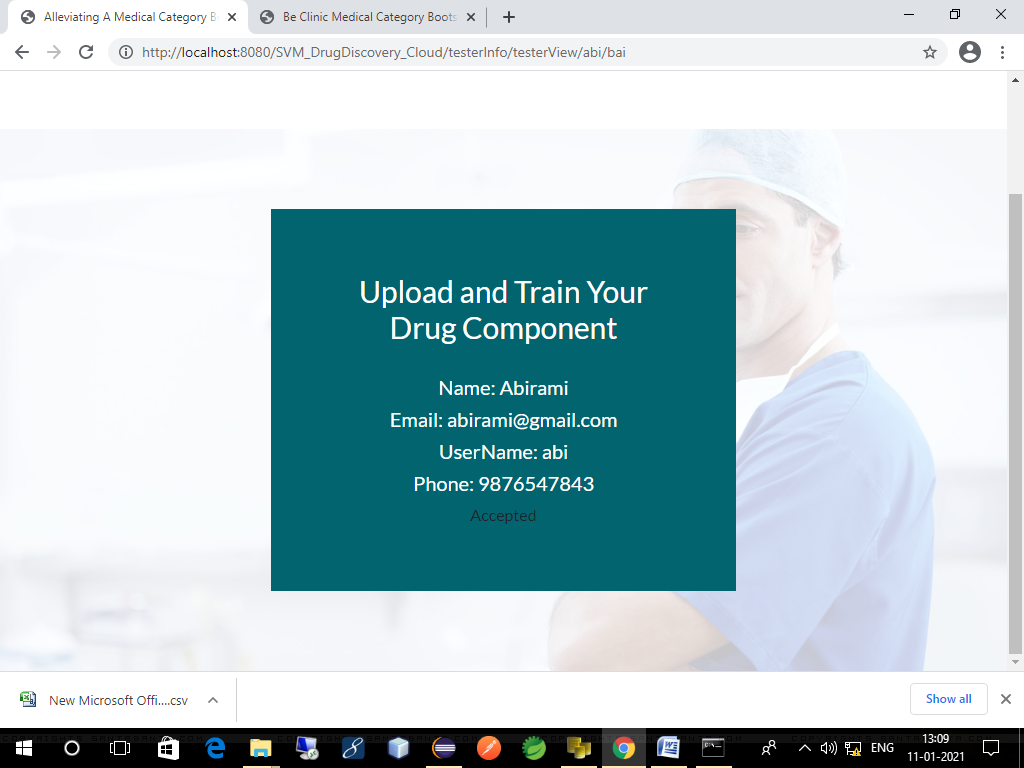
**FILE UPLOADING MODULE**



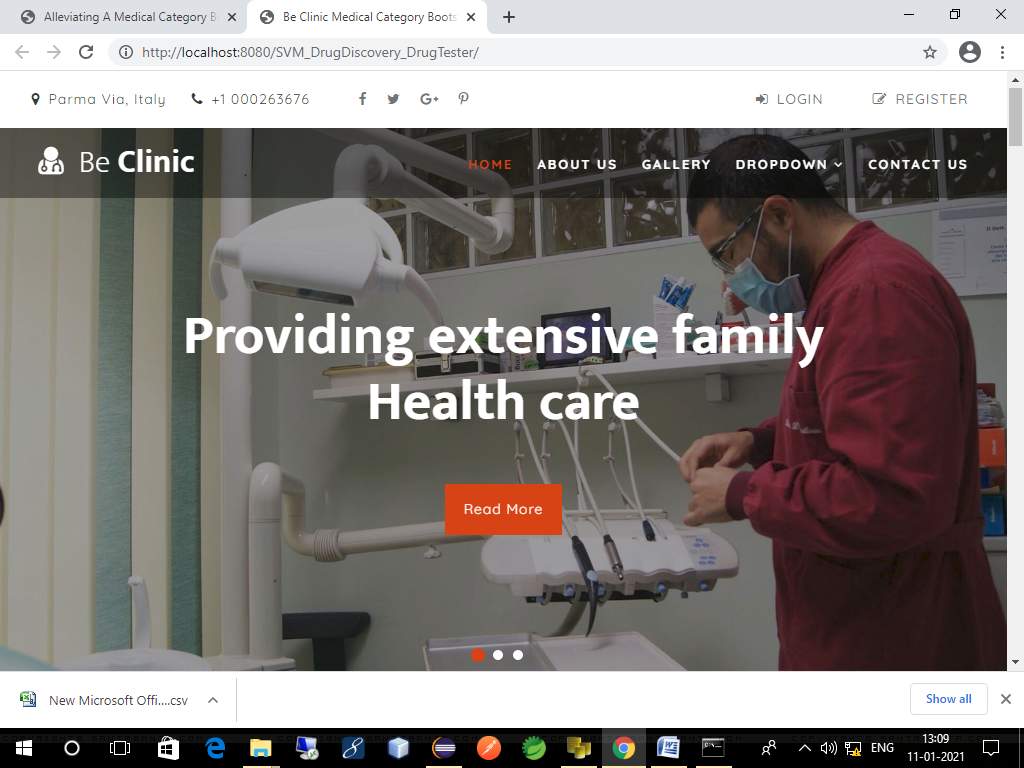
**ACCURACY DETECTION**



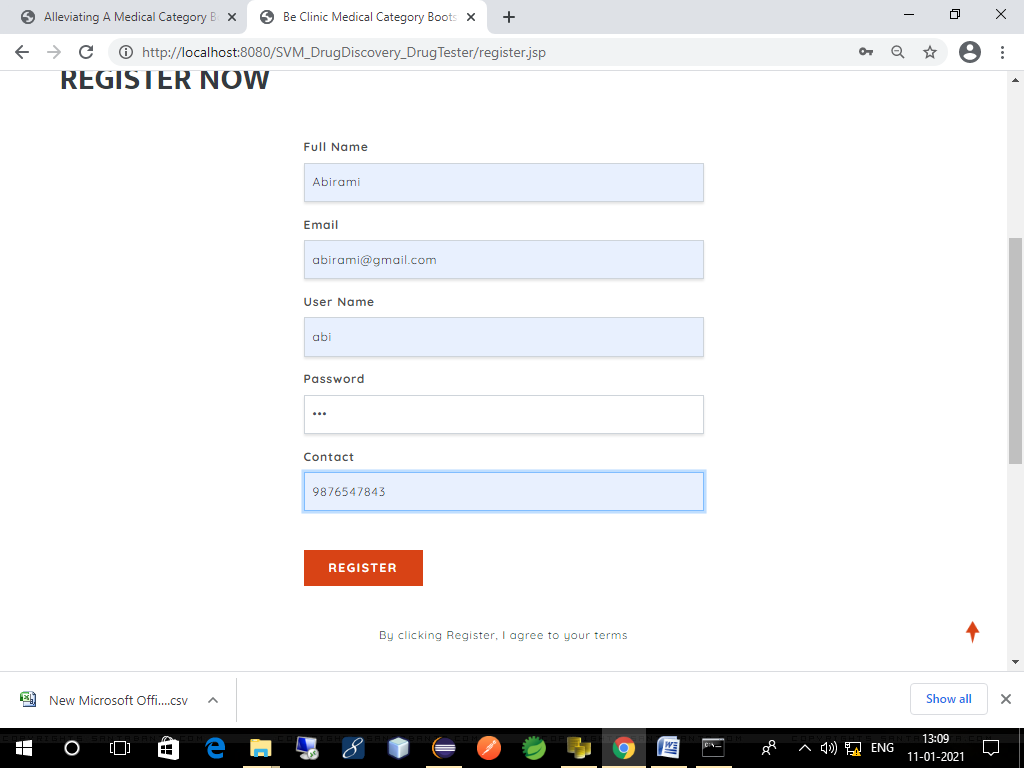
**STATUS PAGE**



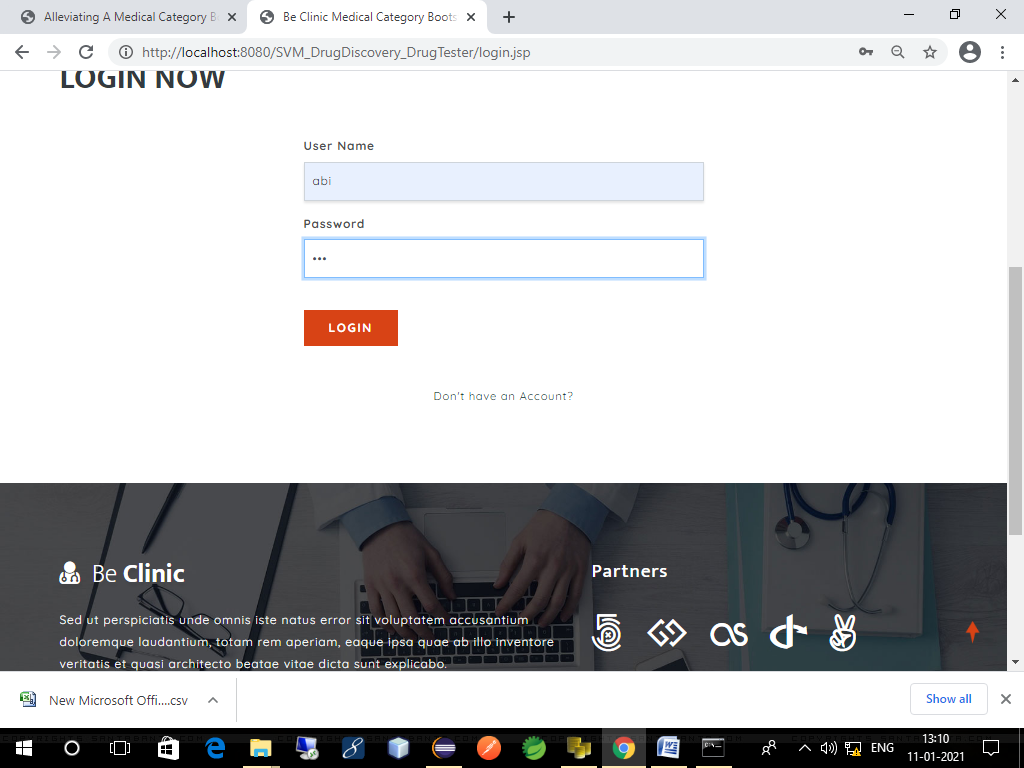
**HOME PAGE FOR DRUG TESTER**



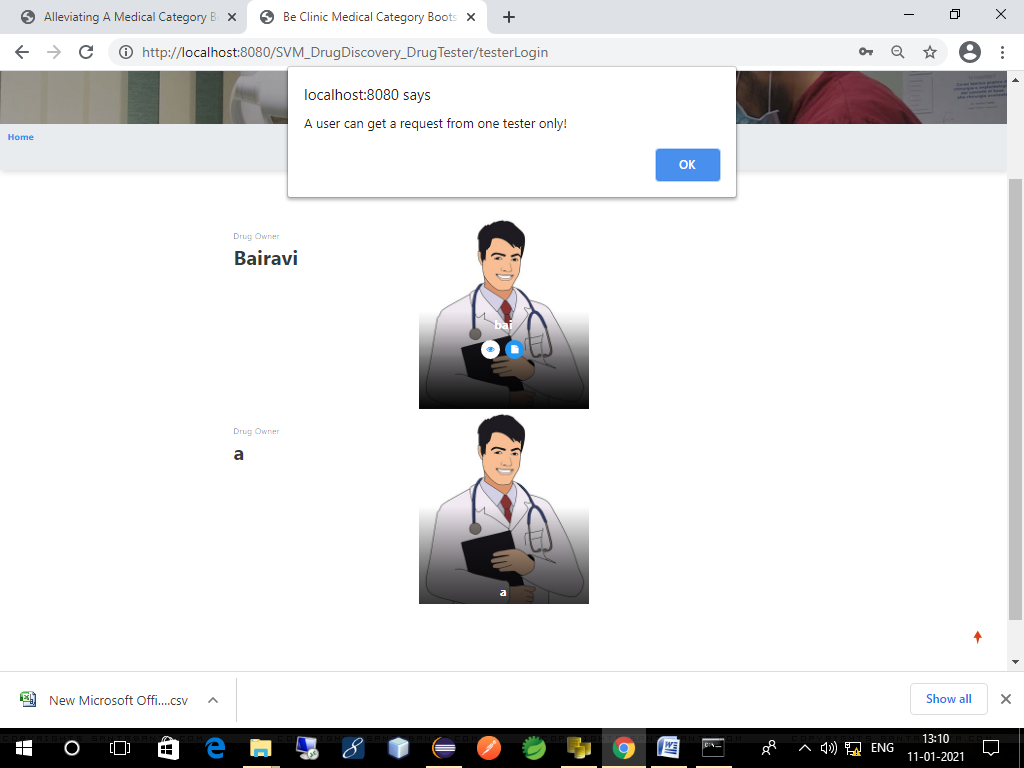
**REGISTRATION PAGE FOR DRUG TESTER**



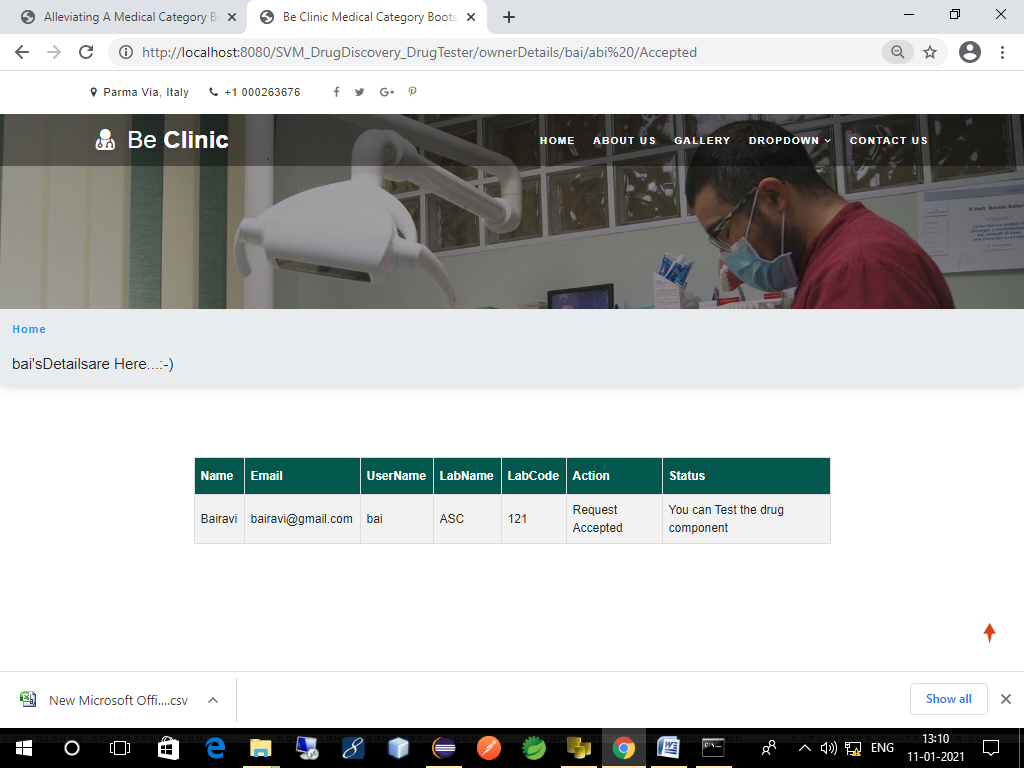
**LOGIN PAGE FOR DRUG TESTER**



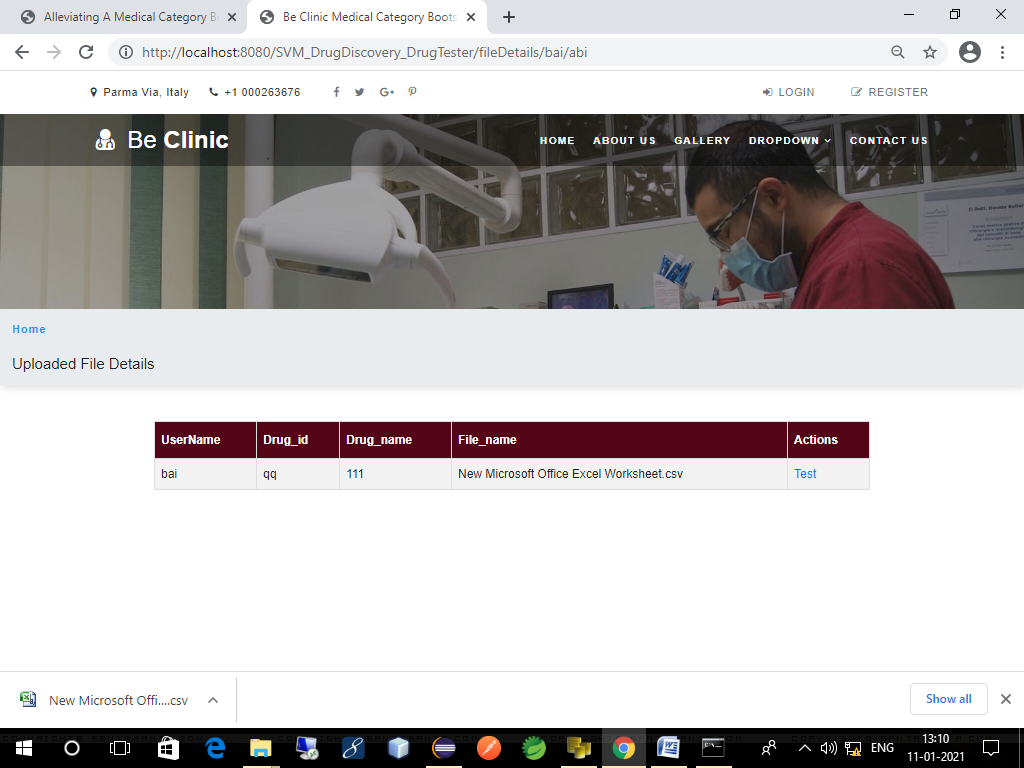
**REQUEST PAGE**



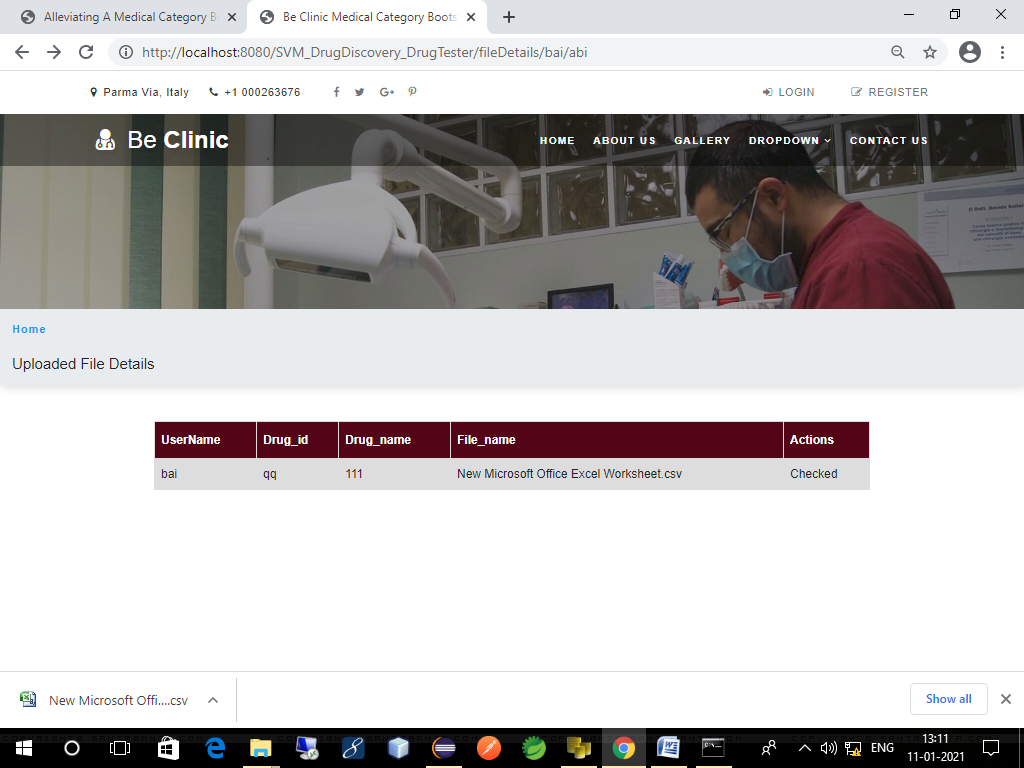
**STATUS MODULE**



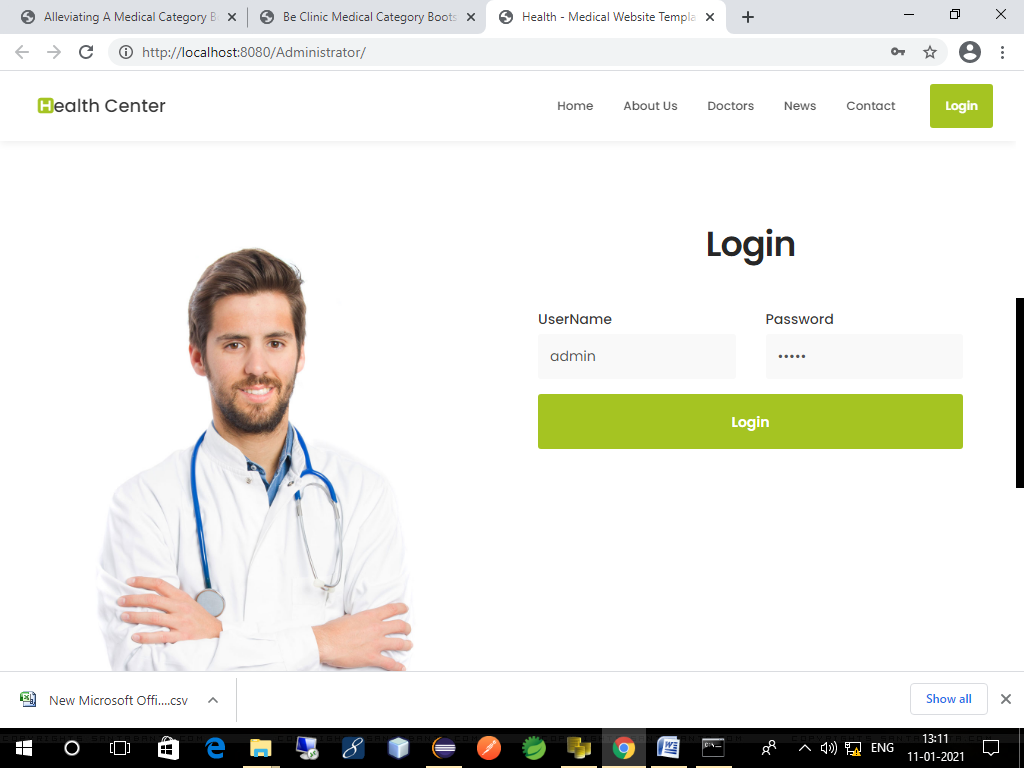
**TESTING PAGE**



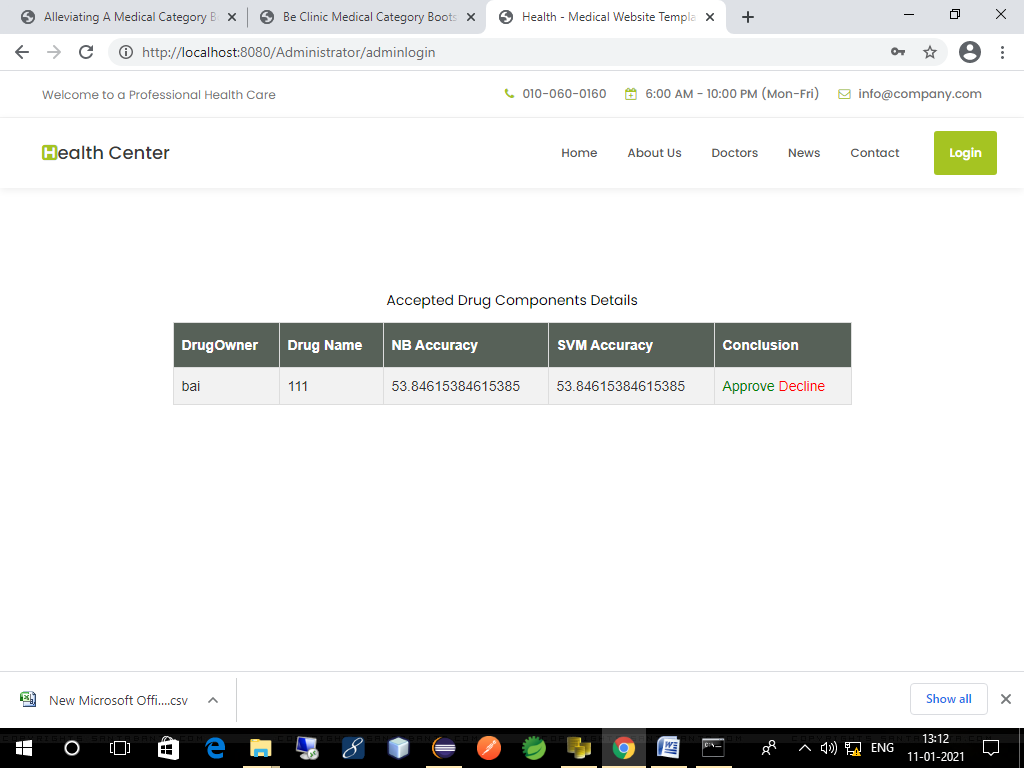
**DRUG TESTED STATUS PAGE**



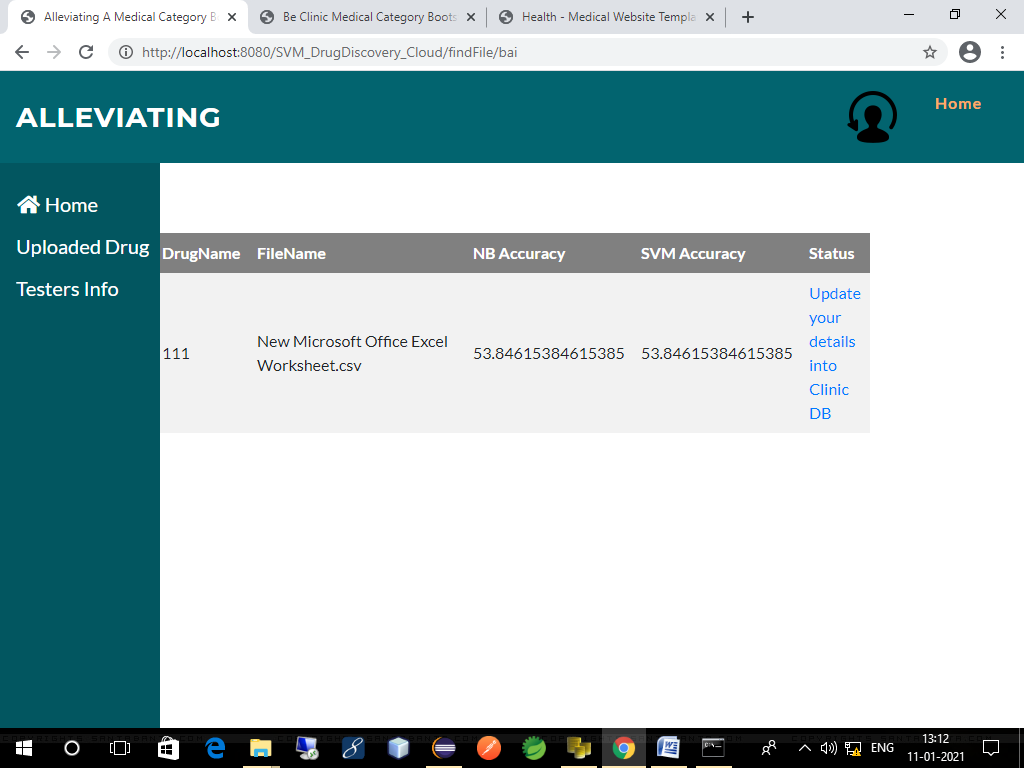
**LOGIN PAGE FOR ADMIN**



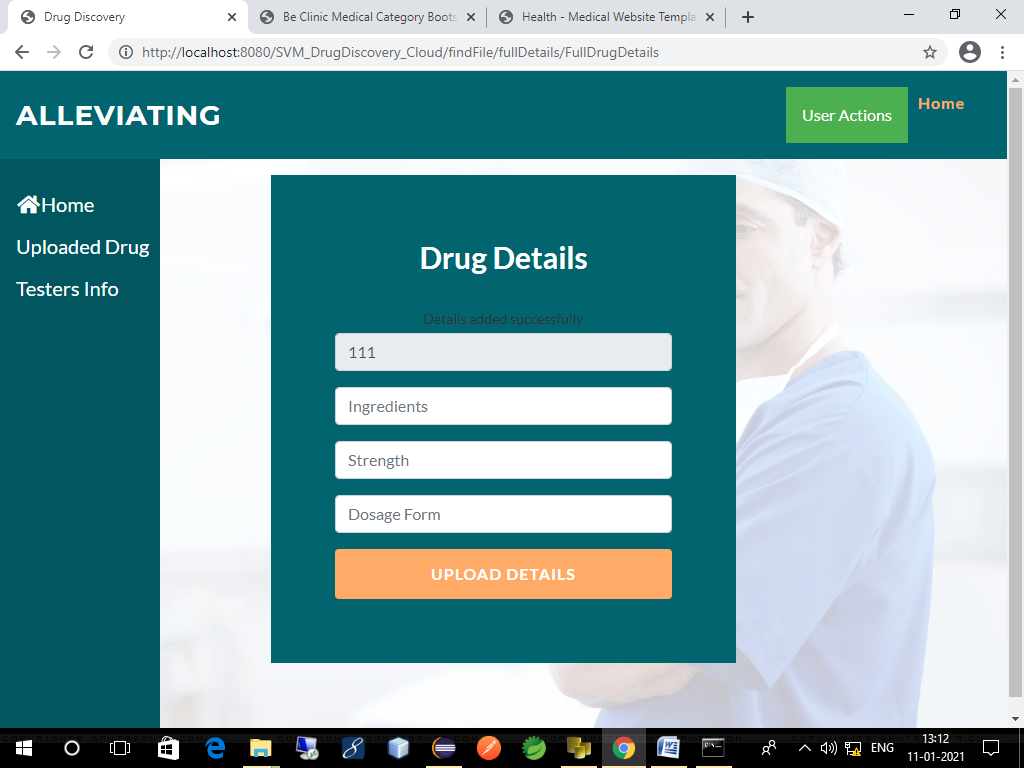
**ADMIN APPROVAL PAGE**



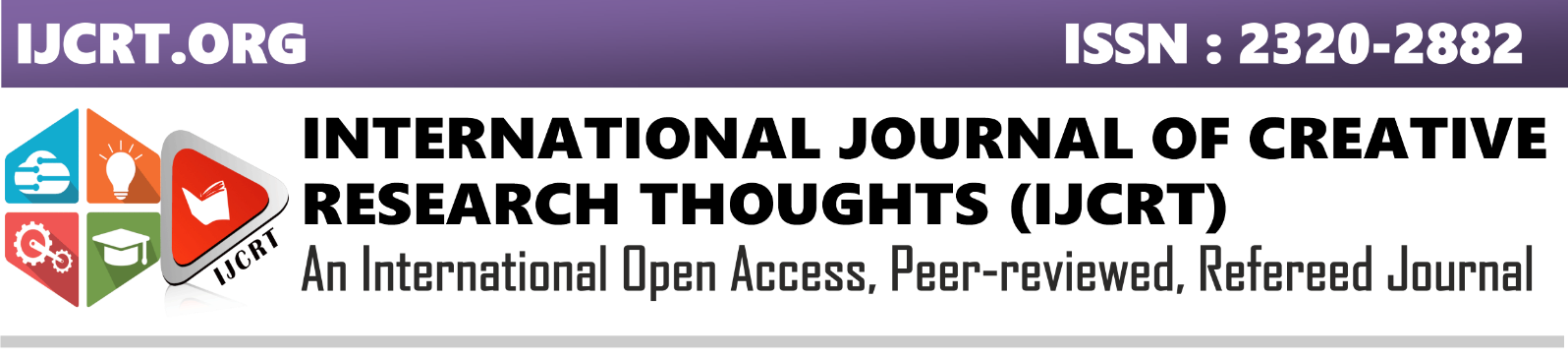
**STATUS MODULE**



**INCLUDING DRUG DETAILS PAGE**



**A.2 PUBLICATIONS**

SECURE THE DRUG COMPONENTS USING NAÏVE BAYES AND SVM

Lavanya T1 , Bhagya sree C R2 , Vinmathi M S3  , Dr Kavitha Subramani4

1,2Final year students, 3,4Assistant Professor,

1,2,3,4Department of Computer Science and Engineering, Panimalar Engineering College, Chennai, Tamil Nadu, India

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

*Abstract:*  In this paper, we propose a framework to secure drug components in the cloud. Specifically, we design for multiple drug formula providers’ to use the cloud securely. In our approach, the analyzer trains the drug formulas using Support Vector Machine (SVM) and naïve Bayes. To perform integer and fraction computations in the cloud server, we designed secure computation protocols. We securely train the SVM to privately refresh the selected SVM parameters using the two protocols, which are SVM parameter selection protocol and sequential minimal optimization protocol. We train NB based on Bayes theorem with an assumption of independence among predictors. To determine whether a drug compound is active or inactive in a cloud, the trained SVM and NB classifier is used. Lastly, we prove that the proposed framework achieves the goal that facilitate drug manufacturers to securely outsource their formulas without privacy leakage to unauthorized parties in the cloud for storage and  for SVM and NB training.

***Index Terms - Securing drug components, Naïve Bayes, SVM, training datasets***

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# Introduction

The initial phase in the drug discovery process is to find a proper 'drug gable' target, which is either a protein receptor or biomolecule. Once the target is found, the second phase is the validation and confirmation of the target. The next phase involves identifying lead compound of a drug followed by testing the target compound. The library of compounds are screened to identify lead compound in various methods. The various methods are screening high-throughput, isolating the natural products etc. After drug discovery, stringent testing and optimization techniques identify process, in the drug development phase the effectiveness of the drug. To study the properties of the lead compounds, it is tested in cells and in animals. To consider lead candidate successful, it should be non-toxic, absorbed, distributed, metabolized and excreted from the body. The result of a development phase concludes if the drug candidate is best for treatment of disease. All these phases marks an end by submitting the drug components to the specific regulatory authority. There may be a leakage of drug information in this phase. Due to the significant investments and high commercial values involved in drug discovery, privacy is an important factor .To secure the drug components, a data mining tools is used.

Of the data mining tools, Support Vector Machine (SVM) has a relatively high decision rate and has been widely used in recent times to predict ligand-based chemical compounds in drug discovery. In approaches using SVMs,known drug formulas datasets is used to train the SVM classifier, and the new drug compound visual scanning is done by  trained SVM.As privacy is prime, how can we minimize the risk of unauthorized disclosure during the SVM training phase? In this context, when a researcher sends some chemical compounds to the cloud for SVM classification, it is important to ensure that the potential new drug compounds will not be leaked to a third party, such as a competing pharmaceutical corporation. Furthermore, to train the SVM, multiple pharmaceutical corporations may collaborate in order to increase the SVM decision rate without revealing their datasets. How to achieve secure SVM training and decision under multiple data sources without compromising, the privacy of each individual party remains a research and operational challenge. Thus, in this paper, we propose secure drug components using SVM and Naïve Bayes for Securing Drug discovery in the cloud environment. Unlike existing drug discovery frameworks, our framework seeks to achieve the following

• Secure Outsourced Data Storage: The drug formula owner can securely outsource the data (e.g. drug formula) to the cloud for storage without leaking the data to unauthorized third parties.

• Secure Multi-Source SVM Training: The POD allows an authorized model provider to use other drug formula owners’ encrypted data to train the SVM on the fly. The model provider can decrypt and obtain the trained model without knowing (contents of) the training dataset.

 • Secure SVM Drug Decision: An authorized tester can securely upload his/her drug chemical compounds to the cloud and determine whether the compound is active or not in a privacy-preserving way.

Commercialization is the last phase of drug development process. The drug is either marketed or commercialized when it is approved. The drug manufacturers need to submit marketing authorization applications in every country in which they want to sell the drug. As the drug is typically targeted to a very large number of patients, the manufacturer is expected to monitor this stage cautiously and submit reports to the FDA. The reports include evidence for medicine-related problems, e.g., treatment failure, adverse reaction, counterfeit/poor quality medicines, drug interactions, or incorrect use. These reports are significant in terms of generating proof of efficacy that will inspire public confidence and trust.

1. **EXISTING SYSTEM:**

In the existing system, it is proposed to secure drug components in the cloud. Specifically, it is designed for multiple drug formula providers’ to use the cloud securely. In this approach, the analyzer trains the drug formulas using Support Vector Machine (SVM). To perform integer and fraction computations in the cloud server, secure computation protocols is designed. The SVM is trained securely to privately refresh the selected SVM parameters using the two protocols, which are SVM parameter selection protocol and sequential minimal optimization protocol. To determine whether a drug compound is active or inactive in the cloud, the trained SVM classifier is used. The existing datasets of known drug formulas to train the SVM classifier, and the trained SVM classifier can be used for new drug compound visual scanning. Due to the significant investments and high commercial values involved in drug discovery, privacy is an important factor. When a researcher sends some chemical compounds to the cloud for SVM classification, it is important to ensure that the potential new drug compounds will not be leaked to a third party, such as a competing pharmaceutical corporation.

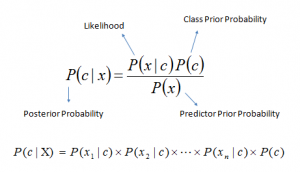
**III.PROPESED SYSTEM:**

We propose secure drug discovery components in the cloud environment. Unlike existing drug discovery frameworks, our POD seeks to achieve it efficiently. We are not using three real time datasets to check the efficiency of potential new drug component. Instead of using existing datasets, we are using another one data-mining algorithm Naïve Bayes (NB), which is based on Bayes’ Theorem .Bayes’ Theorem, is a classification technique, which considers independent predictor’s assumption. NB classifier considers both the presence of a particular feature and any other feature of a class, which are unrelated. It performs well in multi class prediction since it predicts class of test data sets quickly. These two algorithms such as SVM and NB are used to train the uploaded drug dataset (CSV file).Thus we will get trained data and accuracy for that uploaded dataset. The trained data and accuracy will be sent to the owner from python server. Drug tester will check that new drug component. To check the new drug tester has to send request for accessing the drug component. As privacy is an important factor due to the significant investments and high commercial values involved in drug discovery, even drug tester does not know the contents of that file; they will get the trained data only. Once the Testing is completed, the tester sends result to the admin.If that particular drug component is retained in the cloud then it is assumed that component is still active and passed the testing successfully. If not, then the drug component is removed from the cloud. Finally, admin will approve the drug component.

**IV.NAÏVE BAYES:**

The simplest solutions are usually the foremost powerful ones, and Naïve Bayes may well be a model of that. Despite the advances in Machine Learning within the past few years, it is well tried to not only be straightforward but also in addition fast, accurate, and reliable. It has been with success used for many functions, but it works notably well with Natural language processing (NLP) issues. Naïve Bayes may well be a probabilistic machine learning formula supported by the Bayes Theorem, utilized in an exceedingly properness of classification tasks. Bayes’ Theorem is a classification technique that considers independence among predictors’ assumption. Naive mathematician model is simple to make and notably useful for extraordinarily big info sets. Beside simplicity, Naive mathematician is known to defeat even very refined classification ways.

Naïve Bayes uses the formula to predict the accuracy



Above,

* P(c|x) is the posterior probability of class (c, target) given predictor (x, attributes).
* P(c) is the prior probability of class.
* P(x|c) is the likelihood which is the probability of predictor given class.
* P(x) is the prior probability of predictor.

**V.SUPPORT VECTOR MACHINE:**

Support Vector Machine or SVM is one in every of the foremost well-liked supervised Learning algorithms, that is used for Classification equally as Regression issues. However, it is used for clearing Classification issues in Machine Learning.

The SVM formula's goal is to create the foremost effective line or decision boundary that is in a position to segregate n-dimensional house into categories so we will simply place the new information within the correct category among the long-standing run. This best decision boundary is termed as a hyperplane.

The hyperplane is formed with the assistance of acute points/vectors chosen by SVM. These extreme cases are noted as support vectors, thus this algorithmic program is termed as Support Vector Machine or SVM.

**VI.ALGORITHM USED:**

Instead of using existing datasets, we are using another one data-mining algorithm Naïve Bayes (NB). It is a classification technique based on Bayes’ Theorem with an assumption of independence among predictors. NB classifier assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature also fast to predict class of test data set and performs well in multi class prediction. These two algorithms such as SVM and NB are used to train the uploaded drug dataset (CSV file).Thus we will get trained data and accuracy for that uploaded dataset.

**VII.SYSTEM DESCRIPTION:**

This system secure drug components for Secure Drug discovery in the cloud environment. Unlike drug discovery frameworks, the secure drug discovery seeks to achieve it efficiently. We are not using three real time datasets to check the efficiency of potential new drug component. Instead of using existing datasets, we are using another one data-mining algorithm Naïve Bayes (NB).  These two algorithms are used to train the uploaded drug dataset (CSV file). In final, we will get trained data and accuracy for that uploaded dataset. Drug tester will check that new drug component. Drug tester does not know the contents of that file; they will get the trained data only. Then they let us know the file was active or not. Finally, admin will approve the drug component.

LIST OF MODULES:

* Drug Owner & Tester Registration
* Drug Component Uploading
* Train dataset
* Drug Testing

DRUG OWNER AND TESTER REGISTRATION:

Drug Owner will register in the service provider platform. MySQL database is used to store the drugs Meta details. This drug owner registration process will involves few entities such as 1.Name(drug owner’s)2.Email(drug owner’s)3.Contact number(drug owner’s)4.lab name(where the drugs has been discovered)5.lab code(unique number of that lab) for this process.

The drug tester will also be registered in the drug tester registration platform. MySQL database is used to store the drugs Meta details. This drug tester registration process will involves few entities such as 1.Name(drug tester’s)2.Email(drug tester’s)3.Contact number(drug tester’s) and 4.TesterID( unique number of that tester) for this process.

DRUG COMPONENT UPLOADING:

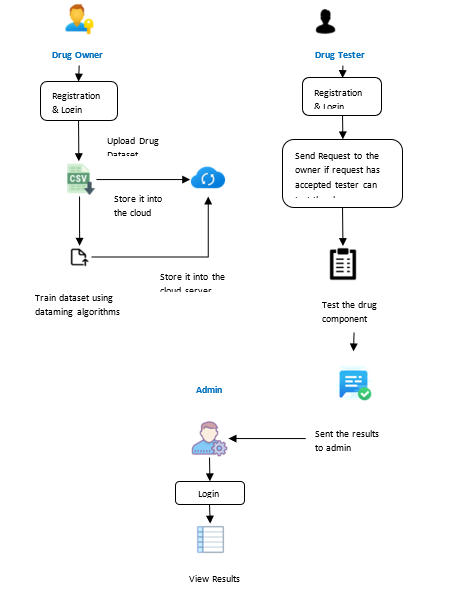
Once the Registration is completed, the drug owner should upload the drug component. For uploading the drug component, the owner must provide the Drug Name, Drug Id and Date of Uploading. Now we have to choose the file that contains the drug data sets and drug components. That data set contains the formula and we have to mention the type of class (Class A, Class B). While uploading the file we will read the content and store into the database and store that .csv file in cloud.

TRAIN DATASET:

As the drug owner has successfully uploaded the drug components, now we will train the uploaded data using python. For this part, we will use two algorithms, SVM and Naïve Bayes. The trained data and accuracy will be sent to the owner from python server.

DRUG TESTING:

As the drug dataset has been trained and uploaded in the cloud, the drug tester now send request to the owner for testing. Only after accepting the request, the test can test the uploaded drug components. Once the Testing is completed, the drug tester sends result to the admin.If that particular drug component is retained in the cloud then it is assumed that component is still active and passed the testing successfully. If not, then the drug component is removed from the cloud.



**VIII. CONCLUSION AND FUTURE ENHANCEMENT:**

This project focuses on securing the drug components in the cloud. Privacy is a major factor in drug discovery as it involves significant investment and high commercial values. Drug discovery is a long-term expensive process. Bringing the drugs from the bench to the market involves a lot of threat when stored in the cloud. Thus, we proposed securing the drug components for drug discovery in the cloud. Securing the drug components is designed to facilitate drug manufacturers to securely outsource their formulas to the cloud for storage and SVM and NB training. These two algorithms such as SVM and NB are used to train the uploaded drug dataset (CSV file). As a result, we receive trained data and accuracy for that uploaded dataset. The trained model could be used for authorized client’s compound classification in a privacy-preserving way. Specifically, we designed a secure domain transformation protocol and several basic secure computation components for secure outsourced computation across different parties. We also built two key secure components (i.e. secure parameter selection and secure sequential minimal optimization) to achieve privacy-preserving SVM and NB training in drug discovery. We will be extending our approach to support more sophisticated data mining method in order to support very large dataset in drug discovery.

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