An Efficient Approach for Dengue Mitigation

A Computational Framework

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# INTRODUCTION

Presence of dengue in Sri Lanka links back to the early 1960s and yet dengue has become a major public health issue at present with a high morbidity and mortality. In 2009, dengue infections increased at an alarming rate across Sri Lanka. By the end of the year 2009, 35,095 people were infected 346 fatalities reported. The number of infections has never been lower than 28,000 since 2009. In 2013, 47,246 infections were reported. Number of fatalities were 83 in 2013. During the first 10 months of the year 2017, 158854 suspected dengue cases have been reported to the Epidemiology Unit of Sri Lanka from all over the island keeping the mortality rate at an alarming level which was about 300 deaths. There is an urgent need for a comprehensive mitigation plan to manage the impact of the epidemic.

Sri Lanka has not been laxing on tackling mosquito breeding grounds. The Presidential Task Force initiated every approach from Dengue Prevention, to fines for those who neglect breeding grounds, to declaring national dengue eradication programs. Unexpectedly, the combined result of these projects is that the rate of infection is exactly what it was five years ago, or in areas where slight reductions are reported still alarmingly high. The situation is more intensive and alarming in the Western Province, home to over 25 percent of the country’s population of over 20 million people, and to 60 percent of all reported dengue cases since 2009. Western Province (43%) has the highest number of dengue cases reported. Colombo district is the most affected area.

All the possible strategies have been tried and failed, experts are now suggesting getting help from the national Meteorological Bureau for the fight against the virus. In the work presented in [1] stated that there is a clear connection between changing climate patterns and the spread of dengue. The authors further presented that mosquito breeding grounds increased following heavy rains, pointing out that the two annual peaks in infections were recorded soon after the two annual monsoons. This work also found that warming weather patterns increased the distribution of the dengue-carrying mosquito. Researchers pointed out that detailed weather forecasts could help health authorities to better allocate resources and strategically implement prevention campaigns.

With all the information on hand mentioned above, I propose a well formulated computational approach to predict and provide guidance in mitigating dengue epidemic in Sri Lanka and Thailand. In my proposed model, I take into consideration all the environmental factors as well as geological factors that may positively affect dengue vector population and the dengue cases. Contributions of my research proposal are as given below.

## 

## 1.1 Contribution

1. **To provide a comprehensive insight into dengue epidemic and spread of the vector population based on stimulating factors.**

The foremost task to be completed before building a framework to identify and mitigate dengue epidemic is to analyze the epidemic thoroughly. There may be various factors that stimulating the spread of the dengue epidemic. These factors may contain directly related factors and hidden factors that are playing a major role in spread of the epidemic. Several major contributors of dengue epidemic have been identified by several research works conducted. Among them, temperature, rainfall, and land use appeared top in the list. All the research work conducted so far take into consideration only the global factors. And also, treat every part of the country homogenously. In reality, there is a great variation in climate, land elevation, population, temperature, and rainfall parameters among different regions in two countries. In this study, we consider global parameters and treat each region with different strategy to profoundly represent the state of the particular region throughout the country.

1. **To predict the upcoming dengue epidemic and its severity using a support vector regression and Micro Ensemble Architecture.**

There is a handful of work conducted to find the major stimulating factor of dengue epidemic. Despite the work that has been done, there is still a lack of dengue mitigation strategy implemented based on the research findings available in Sri Lanka and Thailand. The government of Sri Lanka is deploying numerous projects to mitigate the dengue epidemic. There is still no sign of dengue in under controlled so far. The main reason behind this failure is not to identify the severity of the epidemic and its epicenter. It is very questionable to use global parameters to predict the epidemic with the fact that each region is different from every other region in terms of population density, rainfall, temperature variation, land use, etc. For the identification of upcoming dengue epidemic, we propose a micro ensemble architecture in which each district is modelled with a small-scale ensemble. The result is obtained by applying a strategy on all the results obtained from each output of the ensemble.

1. **To allocate limited resources efficiently to effectively mitigate the dengue epidemic**

Resources are very limited especially in countries like Sri Lanka. It is very important to utilize the available resources effectively. This poses a challenge of how to allocate limited resources among very high demanding facilities. In severe situations such as the one happened in year 2017 where all the hospitals ran out of spaces, posed a significant threat to patients due to lack of resources. All the resources were lacking including number of beds in each hospital. Resource allocation plan is a major component in response planning. Hence, I proposed a resource allocation strategy based on modified genetic algorithm (GA) optimization. The proposed resource allocation scheme can generate a resource allocation plan in a lesser amount of amount of time that is near optimum. The proposed GA modification performed higher than the standard GA.

# MOTIVATION

## 2.1 Global burden of dengue

World Health Organization reported that dengue incidence has grown exponentially around the world in recent years. Actual numbers of dengue cases are underreported as the level of education and awareness of public on the dengue epidemics is low. A survey conducted estimates 390 million dengue infections per year [53]. Another study conducted estimates that 3900 million people living in 128 countries are at risk [54]. The global risk map of dengue epidemic is shown in Figure 2.1.



Figure 2.1 Dengue risk map for year 2009

## 2.2 Status and trends of Dengue Disease

2.2.1 Dengue Status of Sri Lanka

It is clearly shown in the global risk map of dengue epidemic that Sri Lanka is in the high-risk area. This is not a coincidence and it is clearly reflected in the reports produced by various institutions in Sri Lanka. In the last quarter of the year 2015, 14776 suspected dengue cases have been reported to the Epidemiology Unit. In the Western Province, that accounted for 47.11%. The recent development of dengue epidemic in Sri Lanka is alarming. During the first 10 months of the year 2017, 158854 suspected dengue cases have been reported. The mortality rate was at an alarming level which was about 300 deaths. 37988 dengue cases have been reported January through September in the current year 2018. Distribution of cases by weeks for the year 2017 is given in the Figure 2.2.



Figure 2.2 Dengue annual case rate reported weekly in year 2017

A committee has been appointed by the government of Sri Lanka to thoroughly study the dengue epidemic and provide recommendations towards the better control of dengue epidemic. The committee comprised of professionals from several important fields including medicine, healthcare, environmental and higher research institutions. The committee was first appointed in 2001.The final report was produced in year 2005 and handed over to the government of Sri Lanka. This report proposed several action plans which included very important recommendations. Those listed below,

1. To reduce morbidity and mortality due to DF/DHF.
2. To forecast and prevent dengue epidemics.
3. To strengthen liaison with civil society groups, NGO, media and other relevant stakeholders for social mobilization in dengue control.
4. To identify and mobilize resources to carry out research on dengue.
5. To develop and sustain an effective dengue prevention and control program in Sri Lanka.

Among them there is a great attention paid to establish a forecasting model for the dengue epidemic based on the influencing factors. This item in the list of action plan has not been addressed up to now. The effect of the delay of action plan is clearly shown in the dengue trend as shown in Figure 2.3 and Figure 2.4. And also, it is clearly depicted in the Figure 2.3 the strong relation that exists between rainfall and reported dengue cases. The higher peaks occur after every monsoon season.

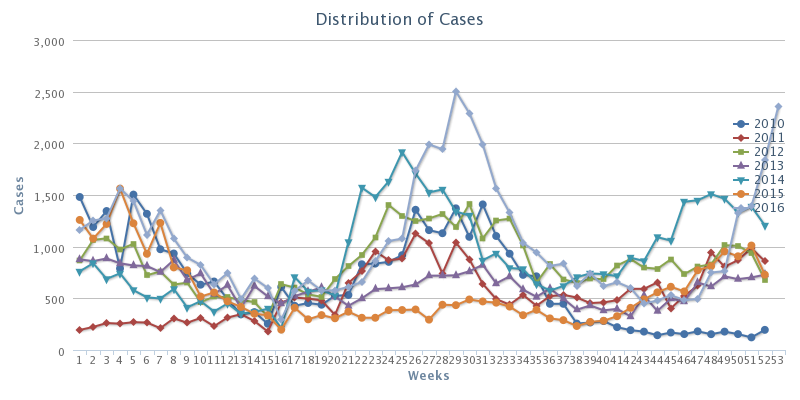


Figure 2.3 Dengue trend for years 2010-2016

Figure 2.4 Dengue case trends from 2000 to 2018

It is very crucial to design and develop a dengue epidemic forecasting model as the dengue epidemic is getting worse and fatality of the epidemic is getting increased every year since 2005.

2.2.2 Dengue Status of Thailand

Thailand reported 6,565 total dengue cases from 75 of the 76 provinces from January 2018 to April 2018. There were nine dengue fatalities. The categorization of the cases based on the type of the case. Out of 6,565 total cases, there were 3,878 dengue fever cases, 2,610 dengue hemorrhagic fever and 2 reported dead and 77 dengue shock syndrome cases and 7 reported dead. Phuket has been the top in terms of the total morbidity, followed by Samutsakorn and Pangnga. Figure 2.5 shows the dengue trend in Thailand from year 2000 to 2011. Figure 2.6 shows dengue cases by month from year 2000 to 2012. Data displayed on Figure 2.5and Figure 2.6 are obtained from the survey study presented in [68].

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Object name is pntd.0003241.g002.jpg

Figure 2.5 Number of reported dengue disease cases and dengue disease incidence, Thailand, 2000–2011

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Object name is pntd.0003241.g006.jpg

Figure 2.6 Number of reported cases due to dengue disease, by month, Thailand, 2000–2012

## 2.3 Questions to be addressed

1. What are the factors that positively affect the spread of dengue epidemic?
2. Are there any local factors that are more important when modeling epidemic and missed when considering only global factors?
3. What is the effectiveness of Support Vector Regression (SVR) and micro ensemble architecture in prediction when considering global and local factors with both vector and human population considered? What parameter settings give the best fit?
4. Can we improve the result by feeding the SVR and ensemble with a combination of geographical, socio economic, and weather data?

# Background

## 3.1 The geography of Sri Lanka and Thailand

3.1.1 Sri Lanka

Sri Lanka is an island in the Indian Ocean and located in Southern Asia. It has 64,740 km² of land and 870 km² of water. Sri Lanka's climate is tropical. There are two main rainfall seasons which are the northeast monsoon (from December to March), and the southwest monsoon (from June to October). Majority of Sri Lanka’s land is flat and at sea level. The highest point is Pidurutalagala which is 2,524.13 m high.

Provinces are the highest-level administrative division in Sri Lanka. Districts are the 2nd level administrative divisions and are included in province. There are 25 districts and they are organized into 9 provinces. Administrative regions are shown in Figure 3.1.

|  |  |
| --- | --- |
|  |  |
| (a) | (b) |

Figure 3.1 (a) Provinces (b) Districts of Sri Lanka

3.1.2 Thailand

Thailand is located in Southeast Asia. With a total area of 513,000 km2 (198,000 sq mi), Thailand is the world's 50th-largest country. Its population ranked the 20th in the world, with 69 million individuals. Thailand is divided into 76 provinces as given in the Figure 3.3. The figure also shows the dengue incidence for the month May of year 2014, which are gathered into five groups of provinces by location. Bangkok (Krung Thep Maha Nakhon) and Pattaya are considered as two special districts. Bangkok is considered both a district and a province. The country of Thailand is given in the Figure 3.2.



Figure 3.2 Location of Thailand

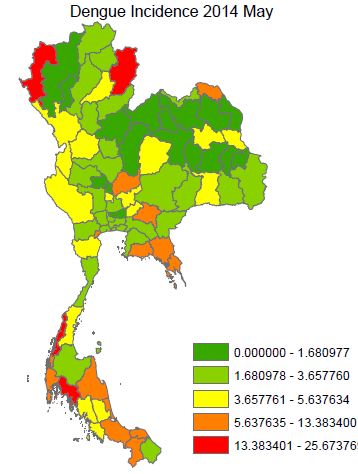


Figure 3.3 Provinces of Thailand and dengue incidence in May 2014

## 3.2 Climate of Sri Lanka and Thailand

### 3.2.1 Rainfall

The main sources of rainfall in Sri Lanka are Monsoonal, Convectional and expressional rain. The mean annual rainfall varies between 900mm to 5000mm (Figure 3.4).



Figure 3.4 **Annual rainfalls in Sri Lanka (Courtesy: Department of Meteorology Sri Lanka)**

There are three seasons of climate in Thailand. Those are Southwest monsoon season or rainy season (May - October), Northeast monsoon season or Winter season (October - February) and Pre-monsoon season or Summer (February – May).

The five divisions of Thailand are Northern, Northeastern, Central, Eastern and Southern Parts. The rainfall for each part is given in Table 3.1

Table 3.1 Seasonal Rainfall (mm) in Thailand

|  |  |  |  |
| --- | --- | --- | --- |
| Region | Winter | Summer | Rainy |
| North | 100.4 | 187.3 | 943.2 |
| Northeast | 76.3 | 224.4 | 1,103.80 |
| Central | 127.3 | 205.4 | 942.5 |
| East | 178.4 | 277.3 | 1,433.20 |
| South |  |  |  |
| - East Coast | 827.9 | 229 | 680 |
| - West Coast | 464.6 | 411.3 | 1,841.30 |

### 3.2.2 Temperature

Altitude is the main cause of regional differences observed in air temperature over Sri Lanka. The mean monthly temperatures slightly differ time to time based on the seasonal changes due to movement of the sun. The mean annual temperature in Sri Lanka rapidly decreasing moving towards highlands from low lands. At the altitude of 100 m to 150 m, the mean annual temperature is between 26.5 0C to 28.5 0C. The temperature falls rapidly as the altitude increases as it moves towards the highlands. The town Nuwara Eliya is at 1800 m sea level and its mean annual temperature is 15.9 0C. The coldest month is January, and April and August are the warmest.



(c)

(b)

(a)

Figure 3.5 Average temperatures from 1961 to 2015 for (a) April (b) August and (c) January **(Courtesy: Department of Meteorology Sri Lanka)**

The temperature of Thailand for each part is given in Table 3.2.

Table 3.2 Temperature (Celsius Degree - °C ) in Thailand

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Temperature | Region | Winter | Summer | Rainy |
| Mean | North | 23.4 | 28.1 | 27.3 |
|  | Northeast | 24.2 | 28.6 | 27.6 |
|  | Central | 26.2 | 29.7 | 28.2 |
|  | East | 26.7 | 29.1 | 28.3 |
|  | South |  |  |  |
|  | - East Coast | 26.3 | 28.2 | 27.8 |
|  | - West Coast | 27 | 28.4 | 27.5 |

### 3.2.3 Climate Seasons

The Climate experienced for 12 months period in Sri Lanka can be characterized in to 4 climate seasons as follows.

*First Inter-monsoon Season (March - April)*



Figure 3.6 Distribution of rainfall in First Inter-monsoon Season**(Courtesy: Department of Meteorology Sri Lanka)**

*Southwest -monsoon Season (May - September)*



Figure 3.7 rainfall distributions for Southwest -monsoon Season**(Courtesy: Department of Meteorology Sri Lanka)**

*Second Inter-monsoon Season (October-November)*



Figure 3.8 rainfall distributions for Second Inter-monsoon Season **(Courtesy: Department of Meteorology Sri Lanka)**

*Northeast -monsoon Season (December - February)*



Figure 3.9 rainfall distributions for Northeast -monsoon Season **(Courtesy: Department of Meteorology Sri Lanka)**

## 3.3 The dengue epidemic of Sri Lanka

Dengue viruses (DENV) are mosquito-borne flaviviruses that have plagued humans for centuries. Urbanization and human population growth in the tropical regions of the world have produced favorable conditions for DENV transmission. Moreover, changes due to urbanization and human population growth have led to the current global dengue pandemic, characterized by a dramatic increase in DENV infections and an expanding geographic distribution of both DENV and the mosquito vectors, *Aedes aegypti* and*Aedes albopictus*, which transmit DENV among humans.

In Sri Lanka, 48 Aedes species belonging to 11 subgenera have been reported to date. The subgenera are *Aedimorphus, Cancraedes, Christophersiomyia, Diceromyia, Finlaya, Mucidus, Neomelaniconion, Paraedes, Rhinoskusea, Stegomyia, and Verrallina.* The established DENV vectors *A. aegypti* and *A. albopictus* belong to the subgenus *Stegomyia*. However, nothing is known about the role of the remaining 46 *Aedes* species in DENV carriage and transmission, an area that might shed some light on how DENV survive the intra-epidemic periods.

Sri Lanka has been affected by dengue fever (DF)/dengue hemorrhagic fever (DHF) epidemics for over two decades. DENV infections have been endemic in Sri Lanka since the mid-1960s. DF was serologically confirmed in the island in 1962. The presence of DF in all of the major towns situated below 1200 m elevation was confirmed in 1966 and in 1976–1978.

In Sri Lanka, DF control efforts have been targeted at the disease and vector, including laboratory surveillance for DENV infections in patients and vectors, vector control, social mobilization, clinical management of DF/DHF patients, and public awareness through the media. A multidisciplinary task force on DF/DHF has been established to moderate the DF/DHF control activities. Furthermore, there are provincial and district-level DF/DHF control activities in place. Training clinicians on clinical management has been carried out continually to bring the DF/DHF mortality to zero, or to a minimum level. It is hoped that with the implementation of collective control programs in collaboration with other governmental and non-governmental organizations, with maximum cooperation from the community, the morbidity and mortality of DF/DHF will be reduced soon.

## 3.4 The virus

DENV is a flavivirus which is transmitted by *Aedes* *aegypti* mosquitoes. There are four distinct DENV serotypes, DENV 1–4. Infection with a single DENV serotype leads to long-term protective immunity against that serotype. The immunity obtained from one serotype will not protect from other serotypes. All four serotypes of DENV have been co-existed in Sri Lanka for more than three decades. Despite its long-existence, their distribution has not changed in the last 30 years. Studies found the existence of two or more DENV serotype in different parts of the country. There was an epidemic of DF associated with DENV serotypes 1 and 2 from 1965 to 1968. This island-wide epidemic caused 51 DHF cases and 15 deaths.

## 3.5 REPLAN framework

RE-PLAN is a computational framework developed to create, analyze and optimize emergency response plans for public health emergencies. Specially, RE-PLAN facilitates the placement of PODs across the region of interest and establishes the geographic region that is being served by each POD. POD locations are selected to minimize the distance that the public has to travel to receive emergency services. Population distribution and geospatial data of the region are utilized for the purpose of response plan creation. Data pertaining to the infrastructure of the region, such as the road network, are utilized in analyzing the effectiveness of the resulting response plan. Specific methods have been developed as part of the RE-PLAN framework to enable creation, analysis and optimization of response plans for different scenarios [2].

A response plan developed in RE-PLAN consists of a set of PODs and their respective service areas. Each POD is a location in the region of interest defined by its geographic coordinates and attributes such as the number of service booths that the facility may accommodate. A service area of a POD is a portion of the region of interest that is serviced by the POD. Service areas consist of groups of contiguous population blocks, which are geographic entities such as represented by polygons with associated population counts. Population blocks, for instance, can be geographic entities such as census blocks or block groups used by the United States Census Bureau to represent populations. RE-PLAN facilitates response plan creation by either establishing the service areas for a set or subset of user-supplied PODs or by recommending a partitioning of the region into service areas and selecting available POD locations for each of the service areas. Methods which determine the service areas for a given set of POD locations are referred to as constrained methods. Unconstrained methods partition the region into service areas and map suitable PODs to these service areas.

The response plans created using either constrained or unconstrained methods are examined for their feasibility with respect to mandated guidelines. The feasibility of a response plan is determined subject to its performance with respect to mandated time constraints and infrastructural limitations. In general, a response plan is said to be feasible if the following conditions hold:

* All PODs that are part of the plan are capable of serving the populations of their respective service areas within a mandated time frame.
* The burden on roads or other public infrastructure that are used during the emergency is within acceptable limits.
* The infrastructure at a POD facility can accommodate the demand exerted by the population of the service area.

POD analysis tools such as the traffic analyzer or the facility analyzer, which is a part of the RE-PLAN framework, can be used to determine the feasibility of a response plan. In this proposal we have adopted a simpler form of response plan feasibility, focusing only on a POD's ability to serve the population of its service area in a mandated time frame. The main components of the RE-PLAN framework are the RE-PLAN client and the RE-PLAN spatial database. The client can be broadly structured into the following components: plan designer, logistics calculator and plan analyzer. The plan designer provides an interface for users to specify the region of interest, select POD locations and adjust POD configurations. Constrained and unconstrained methods to create and optimize response plans based on selected POD locations or service areas can be invoked through the plan designer. Once a response plan has been created, the logistics calculator incorporates data pertaining to the infrastructure of the region, such as road capacities or public transportation, into the response plan to facilitate analysis of the plan. Plan analysis tools provide methods to analyze the feasibility of response plans with respect to constraints such as time, POD capacity and infrastructural limitations of the region. The facility analyzer examines the performance of each POD facility with respect to the POD's capacity and population of its service area. RE-PLAN is written in Java and uses a PostgreSQL database with PostGIS capabilities to handle spatial data and operations. An overview of RE-PLAN's architecture has been described [2].

3.6 Genetic Algorithm

Genetic algorithm is a metaheuristic [62] belongs to the larger class of evolutionary algorithms (EA). The main idea was taken from the process of natural selection. Genetic algorithms are used to generate high-quality solutions to optimization and search problems by applying operations such as mutation, crossover and selection inspired by the natural selection process. GA can produce the near optimum solution with a sufficiently large population and careful parameter selection. The standard GA is closely resembling the natural process and hence population generation, crossover and mutations are fully random by nature. For this study, I have modified the standard GA to match the problem domain and to achieve a faster convergence. In the proposed GA, population generations, cross-over and mutations have been modified.

# Related works

## 4.1 Dengue Epidemic

Increases in disease transmission and the frequency of epidemics have also been the result of circulation of multiple serotypes in Asia [8]. 2.5 billion people around the world living in dengue endemic countries with a risk of contracting DF/DHF, nearly half live in 10 countries of the Southeast Asia region. 75% of the current global disease burden due to DF/ DHF is reported in the Southeast Asia region together with Western Pacific region. In 2002, DF/DHF was ranked as the third most common noticeable disease in Sri Lanka (first and second were malaria and tuberculosis) [9, 10]. In recent years, deaths due to DF/DHF have been greater than those due to malaria, and DF/DHF is becoming the number one killer mosquito-borne infection in Sri Lanka [9]. At present, DF and DHF are prevalent in many urban and semi-urban areas of Sri Lanka with seasonal and periodic epidemics occurring regularly in the island [5]. In recent decades a higher incidence of DF/DHF has occurred in the districts of Colombo, Gampaha, Kalutara, Kurunegala, Kegalle, Ratnapura, and Kandy [11] from 2000 to 2008, the reported number of suspected and serologically positive DF/DHF cases varied from 4749 to 15 643, involving 25–88 deaths, with a major epidemic in 2004 [6]. DF/ DHF have become endemic in northern and eastern Sri Lanka, with a high incidence in the capital cities of the northern and eastern provinces of the island, Jaffna and Batticaloa, respectively [12].

There is a significant relation of the dengue epidemic to the age group of the population. In many age groups, males have been affected. According to a regional study done by the World Health Organization (WHO) in Sri Lanka based on reported cases from 1996 to 2005, there were consistently and significantly larger proportions of males with DF/DHF in those aged 15 years. This male preponderance was reported in every province of the country. Among those aged 1–4 and 5–14 years, there were significantly fewer male cases than expected, although there was some annual variation [13]. The highest incidence occurred in the 5–9 years age-group [3]. Before 2000, one large peak of DF/DHF cases was observed in children and a few cases were observed in adults. After 2000, two reported DF/DHF peaks were observed in children and young adults. Moreover, the mean age of reported DF/DHF cases was shown to have increased from 15 years in 1996 to 25 years in 2006 [4].

Climate change such as temperature, rainfall, and humidity can expand the geographical range of vector mosquitoes. And also, it can extend the disease transmission season, shorten the gonotrophic cycle, and reduce the time taken for ingested viruses to develop to infective stages in mosquitoes, thereby increasing the propagation rates of arboviral diseases transmitted by*A. aegypti and A. albopictus*[7, 14, 15, and 16]. In addition, these works clearly pointed out that there is a strong positive correlation between rainfall and the dengue cases reported. Two DF/DHF peaks occur annually in Sri Lanka. These peaks are association with the monsoon rains. During peak times, the densities of two mosquito vectors (*A. aegypti and A. albopictus*) are high. The first peak appears in June/July, along with the south-western monsoon that commences in late April. The second peak usually occurs in December and it is associated with the north-eastern monsoon rains (October to December) [3].

Temperature is another important factor controlling the seasonality of DF/DHF outbreaks in sub-tropical or temperate regions. It influences vector distribution, the blood feeding activity of the vector, the extrinsic incubation period, and adult longevity. *A. aegypti* has been shown to transmit DENV when the temperature is above 20 8C but not less than 16 8C. A positive correlation has been shown between the temperature and the female vector abundance. In addition, high temperatures may increase the frequency of blood feeding due to a rapid reduction in energy reserves [17]. It is expected that global warming may further facilitate the expanded distribution of DENV mosquito vectors in temperate regions such as northern parts of North America and Europe [18]. This concern has become a far more serious matter with the expanding distribution of *A. albopictus* [19]. Altitude also plays a vital role in limiting the distribution of *A. aegypti*. In India, *A. aegypti* breeding sites range from sea level to 1000 m above sea level. Lower elevations (less than 500 m) have moderate to heavy mosquito populations, while mountainous areas (over 500 m) have low populations [20].

There is couple of researches conducted to study the control measures of the dengue epidemic in Sri Lanka. These studies have revealed couple of strategies to be used in controlling the epidemic. A study conducted in the Kandy District of Sri Lanka showed that the mechanical and biological measures alone are not sufficient to prevent *Aedes* breeding. The prevention of *A. aegypti* and *A. albopictus* breeding in water storage containers would help to control DF/DHF. Therefore, DF/DHF control programs should pay more attention to the control of *Aedes* breeding in domestic water storage containers [5]. More importantly, public education on preventing dengue epidemic will reduce the mosquito breeding sites and hence will be very effective in dengue mitigation [21, 22]. This will call for a system to integrate all these findings and develop a methodology as proposed in this proposal to effectively deal with dengue epidemic in Sri Lanka. We propose to study all the influencing factors and use them in forecasting/predicting system to clearly identify the high-risk areas. And also, the proposed system is capable of estimating the casualties of the upcoming dengue epidemic hence can be better prepared for the event. This information is in great help in educating general public and putting controlling measures into action.

## 4.2 Forecasting /Prediction

### 4.2.1 GIS and statistical models

The authors of [23] and [24] studied the prediction of dengue outbreak in Sarawak and Johor respectively by using statistical models. The work presented in [23] analyzes the interaction between environmental, entomological, socio-demographic factors. GIS technology was used to generate geographic and environmental data on *Aedes albopictus* and dengue transmission. A total of 32,838 *Aedes albopictus* eggs were collected from trapping that spans 56 days. Cluster sampling was conducted to determine whether any of the risk factors (entomological or geographical) were influenced by geographical location. SPSS 10.1 was used on the data collected to perform analysis. Descriptive analysis tools such as frequency, means, and median were used. Two-sample t-test, and Pearson's Chi-Square were used to determine the association between variables and dengue cases reported. Use of differential Global Positioning System in mapping sites of 1m accuracy is also highlighted. Analysis of the data revealed there are major differences in clusters of villages. These differences include number of *A. albopictus* eggs from ovitraps set indoor, outdoor and in dumping sites, container density, house density, and distance of the house from the main road. T-test showed conducted showed that house density, container density, indoor mosquitoes egg count, outdoor mosquitoes egg count, and dumping sites mosquitoes egg count were higher at the roadside villages compared to border villages.

The work presented in [24] considers various environmental factors such as rainfall, temperature, living conditions, demography structure domestic waste management and population distribution in determining the mosquito survival and reproduction. A geostatistical modeling, analysis and mapping approach has been utilized in this research to understand the correlation between dengue fever prevalence, population distribution and meteorological factor, and the characteristics of space-time clusters in the Johor State. Authors showed that combining GIS with geostatistical analysis and space-time permutation scan statistic tools, the spatial variation of dengue incidence can be mapped. They support their claim with the fact that Geographical weighted regression (GWR) analysis gave a strong (R2= 0.87) positive spatial correlation between dengue fever and population distribution. Vaidya A. at el. [61] introduces a mathematical, compartmental model to forecast the population dynamics of a mosquito and its life cycle in relation to seasonal variations of temperature and rainfall. Populations within the compartments were expressed in the form of a system of coupled differential equations, which describe changes in the mosquito population through processes of maturation and mortality. By using regression tools, maturation and mortality rates at various temperatures were estimated.

### 4.2.2 Neural Network

The work presented by authors in [32] had predicted the dengue confirmed cases by using the neural network. The used factors were average temperature, average humidity, total rainfall and the dengue confirmed case number. The authors used an ANN to analyze a set of data consisting of 14,209 dengue reported cases. They reported the results are encouraging. The feature vector was formed by combining mean temperature, mean relative humidity, total rainfall and the total number of dengue confirmed-cases. The authors claim that the proposed prediction model can be used world-wide and in any period of time since the approach does not use time information in building it. The authors proposed an automatic prediction system of Dengue Hemorrhagic Fever outbreak by using entropy and ANN in the research study presented in [25]. In this study, authors mandate the information preprocessing prior feeding into ANN. This step will eliminate redundant data and noises. Temperature, relative humidity, and rainfall were considered in the information extraction phase. Then, a supervised neural network is deployed to predict the possible risk of Dengue Hemorrhagic Fever outbreak. To evaluate the performance of proposed system, the experiments based on the condition of weather data and Dengue Hemorrhagic Fever cases from January 1999 until December 2007 were conducted. Authors claim 85.92% accuracy compared to the actual data.

### 4.2.3 Cellular Automata

Cellular automata models began from the concept of John von Neumann to make the machine copies itself. Cellular means "consist of cells". Cellular automata can be multidimensional. If there are two dimensions, it resembles a checkerboard. Each cell has some adjacent cells and called "Neighborhood". Changing the status of a cell in one-time step depends on local rules. The local rules may be the probability [35]. This research uses Moore neighborhood with radius=1 and using the probability in changing status. The main epidemic models are used with a cellular automaton model are SIR and SEIR model. For SIR model, each cell has only one status in one-time step such as 'S' represents susceptible, 'I' represents infected and is able to transmit the disease to the others, 'R' represents recovered. Some diseases have a latent period, a status for this period is 'E' and is called SEIR model [34]. An outbreak of dengue fever is characterized by a SEIR model. Some people are not sick when exposed to the dengue virus. The patient will have an incubation period of about a week after exposure virus and before symptoms to appear. The authors created a model of HBV infection by using cellular automaton [31]. The CA lattice size was 300x5000. The status of a cell in lattice might be "susceptible”, "infected”, "core" or "immune". The local rules were the probability. The work presented in [33] had created a time series model to predict the number of patients with Chickenpox by using probabilistic Cellular Automata. Chromosome of genetic algorithm consists of the state changing probability. Experimental results showed that the bigger number of cells in lattice is better than fewer numbers of cells. The experimental weakness is the forecasting did not use any factor. The proposed model presented in [30] takes a number of people in each status of an epidemic model called SIER into consideration. In this respect, CA take a Genetic Algorithm (GA) to generate the factor weight chromosomes and ANN to determine the probability of state transition 'S' to 'E' at time step t (Pt (s, e)). In addition, other related probabilities are obtained by expert knowledge; P (e, i) = 0.15 and P(i, s)=0.001. P(r, s) is determined by GA. These probabilities were used to calculate the cell number of each state at the next time step of GA. GA compute the fitness for one-time step and repeat every time step finally to compute RMSE. For performance evaluation, 32 factors of dengue causes are used in the model. The dataset collected during 2005 to 2011 consisting of 359 weeks in which 287 and 72 are used to train and test the model, respectively. Authors claim with the results obtained that their method outperforms the artificial neural network approaches.

### 4.2.4 Support Vector Machine

Support vector machine is used in various fields to perform pattern recognition successfully. These areas include face detection/recognition, object detection, image retrieval, information retrieval, speech recognition, and prediction/forecasting. SVM is also used as a regression model in which a value for the dependent parameter is given instead of the class of the parameter. The aim of many nonlinear forecasting methods [26, 27, 28, and 29] is to predict next points of time series. Tay and Cao [29] proposed C-ascending SVMs by increasing the value of C, the relative importance of the empirical risk with respect to the growth of regularization term. This idea assumed that it was better to give more weights on recent data than distant data. Results revealed that C-ascending SVMs produce better results than standard SVM in their problem domain. Fan et al. [28] had adopted SVM approach to the problem of predicting corporate distress from financial statements. In this problem domain, the performance is affected by the choice of input variables. Authors also claimed that selecting suitable input variables has a positive impact on the performance. Input variable must be selected such a way that should maximize the distance of vectors between different classes. And also minimize the distance within the same class. It is suggested that Euclidean distance-based input selection provided a better performance.

In this work we use Support Vector Regression for the prediction/forecasting of the dengue cases in the upcoming epidemic. The main rationale behind the selection of SVM is that it can be easily adopted in the web server. This will enable us to integrate the prediction model in the web user interface.

## 4.3 Response Planning

Dispensing treatments to the public during an emergency is an important task. There are numerous works have been done on various aspects of response planning. Every plan must deploy methods to distribute supplies and dispense medication to the affected population within a specified time frame [36]. There are other important factors such as vulnerabilities of the population that must be taken into consideration when developing a response plan. Each of these factors is considered separately in different works. Routing and scheduling the distribution of supplies have been addressed in different ways, and management of treatment facilities has been studied. Different strategies to distribute medication among the facilities have been introduced in the research work [37]. The challenge of distributing medications and treatment supplies to each local agency is a challenge. The concept of Point Of Dispense (PODs) was introduced by the Center for Disease Control (CDC). PODs strategy is been recognized by the authorities as an effective method of planning an emergency. The CDC maintains a warehouse of treatment supplies and medications and delivers them in accordance with the demand to the local authorities. It is the responsibility of each local authority to develop their own response plan adheres to the guideline setup by the CDC [36]. The PODs concept is well utilized in the RE-PLAN framework, developed at the Center for Computational Epidemiology and Response Analysis (CeCERA) [38]. The framework is capable of producing an effective response plan based on PODs placement where necessary. There are couple of different PODs placement methods introduced in the framework. Each of which is suitable for different scenarios. And hence, the RE-PLAN is a better candidate for the proposed work to make use of by adapting the functionality.

The work presented in [39] has pointed out the additional strain imposed when dealing with mass events involving mass number of people. Public health and policy studies stressed the mandating the managing of limited resources during an emergency [40]. Ethic in allocating resources during a mass casualty event is presented in [36] [41].

The spatial data of the region is a critical component in response planning. This will allow localization of data and provide a visual feedback to the plan designer. The spatial data may include population distribution and road infrastructure, and census data with arbitrary census blocks. For an effective management and manipulation of spatial data, Geographical Information System (GIS) is needed. Integration of GIS data and usage of spatial tools in response planning are widely studied and recommended by various studies [42]. RealOpt [43] proposed a simulation and decision support system created to support planning, designing and placing large-scale emergency dispensing clinics for emergency response. The Centers for Disease Control and Prevention created BioSense, a surveillance system [44], is targeting at early detection of biological emergency events. Coombes [45] and Schneider et al [46] proposed several methods of defining boundaries for the response planning area. The authors proposed algorithms to solve problems associated with discrete and continuous facility PODs allocation [47, 48, 49, 50, 51 and 52]. RE-PLAN framework provides methods to integrate spatial, population and infrastructural data and design and analyze response plans by means of spatial partitioning algorithms and analysis techniques.

## 4.4 Resource allocation

Resource allocation is mostly a subpart of response planning for a natural or manmade disaster. There are several research efforts that focused on optimum allocation of available resources among different entities during disasters. They address different disaster types such as wildfires [64], earthquake [65], and public health emergencies [66].

Research study presented in [64] introduced a minimal stochastic process to represent wildfire progression that nonetheless accurately captures the heavy tailed statistical distribution of fire sizes observed in nature. Authors then couple this model for fire spread to a series of response models that isolate fundamental tradeoffs both in the strength and timing of response and also in distribution of limited resources among multiple competing entities. This framework was used to compute optimal strategies for decision making scenarios that arise in the policy of fire response. There is no computational framework involve in this study that can be used in resource allocation process at ground level. The research study presented in [65] introduced a dynamic optimization model. The model was feed with available resources and a detailed description of the concerning areas to calculate the resource performance. In [66], a solution to overwhelming demand for healthcare resource during a large-scale public health emergency was proposed. This study discusses a resource allocation approach for optimizing regional aid during public health emergencies. They presented a elation between optimal response and delaying the distribution of resources from the central stockpile. And also presented that policy level decisions that alter the objectives of pandemic relief efforts can significantly impact the allocations to affected regions. This study is not presenting a computational framework that can be used to allocate resources among facilities.

Authors of the study [63] proposed an agent-based simulation to be used to determine the allocation of resources for a two-site incident. In this study they try to minimize the latest hospital arrival times for critically injured casualties. Further, the investigate how the optimal resource allocation depends on the distribution of casualties across the two sites. Author is also claimed that this study is tested only on two sites and hence further improvements are needed to apply for larger number of sites.

We propose our solution to analyze and predict the dengue epidemic with a trained model which considers many social economic and geographical factors that positively affect the spread of the epidemic. The main simulating factor for the proposed work is that previous works did not consider, the differences of the administrative regions of Sri Lanka. Each province has its own geographical and social economical characteristics and can be different from each other. Assuming all the units are homogeneous may result in a biased conclusion. We need to overcome this shortcoming with the introduction of local characteristics in the model that are specific for each administrative unit.

# Materials and Methods

The methodology of obtaining an efficient prediction system involves several steps. From data processing to resource allocation, it goes through a distinct set of thorough processing. System flow diagram for the proposed study is shown in the Figure 5.1.

Data Acquisition

Data Pre-Analysis

Data Pre-Processing

Model Generation

1. Rainfall, Temperature, Dengue Incidence, Population Data

2. Data Alignment and Cropping

Correlation Analysis

1. Year-wise Data Normalization,

2. Outlier Removal with Iterative Convex Hull Method

1. SVR

2. Least Square

3. K-NN

4. Ensemble (With Best Models)

Model Validation

10-Fold Cross Validation (Prediction)

Resource Allocation

Proposed Genetic Algorithm

Figure 5.1 Proposed Work-Flow for Dengue Epidemic Mitigation

## 5.1 Data Gathering

The number of dengue cases reported depends on various factors such as rainfall, temperature, population density, waste management efficiency, land use, and water body management etc. In this study, only rainfall, temperature, and population densities are considered. These factors are gathered from various sources based on the availability. The following sections explain our strategies of obtaining each factor.

### 5.1.1 Rainfall Data

Rainfall data was obtained from Global Rainfall Map in Near Real Time (GSMaP\_NRT) distributed from JAXA Global Rainfall Watch, which was developed based on activities of the GSMaP (Global Satellite Mapping of Precipitation) project. The GSMaP project is promoted for the study “Production of a high-precision, high-resolution global precipitation map using satellite data," that is sponsored by Core Research for Evolutional Science and Technology (CREST) and it is a part of Japan Science and Technology Agency (JST) [14]. GSMaP\_NRT repository provides hourly rain rate data in 0.1-degree resolution (10km at the equator). Repository divides the globe into 15 distinct regions as shown in Figure 5.2 and provides rainfall data separately for each region as Comma Separated Values (CSV) files. Registered users get free access to the repository. The users can get data using an FTP client that is connected to the repository using credential provided by the repository management. Thailand is included in 02\_AsiaSE area. Table 5.1 lists location specific information for each Asian region. For the model training, data from five consecutive years was used.



Figure 5.2 Definition of text areas of JAXA data repository for text data [14]

Table 5.1 GSMap text area declaration for Asian region [14]

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Area name** | **Lon**  **(W)** | **Lon (E)** | **Lat (S)** | **Lat (N)** | **Description** |
| 01\_AsiaEE | 90 | 155 | 30 | 50 | East Asia |
| 02\_AsiaSE | 90 | 155 | -10 | 30 | South East Asia |
| 04\_AsiaCC | 35 | 90 | 35 | 50 | Central Asia |
| 05\_AsiaSS | 60 | 93 | 5 | 40 | South Asia |
| 06\_AsiaSW | 35 | 65 | 4 | 40 | Arabian Peninsula and East Africa |

### 5.1.2 Temperature Data

Temperature data was obtained from Thai Meteorological Department (TMD) and NASA Earth Observations data archive [69]. Average temperature value for each month for each district was used in the training. Time span of temperature data is five consecutive years.

### 5.1.3 Dengue Case Data

Dengue case data for each district for five consecutive years were obtained from Sri Lanka and Thailand Epidemiology Units. Dengue case data is given in three groups which are Dengue Hemorrhagic Fever (DHF), Dengue Fever (DF) and Dengue Shock Syndrome (DSS). I combine all three categories to form a single entity and used in model training as dengue cases. Dengue case data was obtained from Department of Disease Control, Ministry of Health.

### 5.1.4 Population Data

Population data was obtained from Department od Census from both countries.

## 5.2 Data Processing

### 5.2.1 Extracting Relevant Data and Alignment of Time Resolution

GSMaP\_NRT region 02\_AsiaSE covers a larger area than Thailand geographical region (Figure 5.2). Region 05\_AsiaSS covers a larger area than Sri Lanka. This results in large amount of non-related data being loaded into the spatial database making it heavy for fast computations. To reduce the data load overhead, only the rainfall data that falls inside Thailand and Sri Lankan geographical area were obtained by cropping the dataset using longitude and latitude. Non-relevant data was discarded. As the time resolution of rainfall data is one reading per hour, it is required to compute the monthly rainfall data from hourly data. This matches the time resolution of each factor before use in training process as temperature and population data recorded monthly basis. Further, there are multiple observation points fallen in a single district as shown in Figure 5.3. The average accumulated value of all the points that fall in a district was taken as the monthly rainfall of that district. The unit of recording is mm per hour(mm/hr). Sample data file format for rainfall from GSMaP\_NRT is given in Table 5.2.



Figure 5.3 Rain fall data observation points and geographical boundaries of all four provinces

Table 5.2 Fragment of rainfall data text file from GSMap\_NRT

|  |  |  |
| --- | --- | --- |
| **Lat** | **Lon** | **RainRate** |
| 20.95 | 97.05 | 0.1 |
| 20.85 | 97.05 | 0.06 |
| 20.75 | 97.05 | 0.04 |
| 20.65 | 97.05 | 0.06 |

## 5.3 Pre-analysis of Data

The proposed model used in this study is the SVR [15] and Ensemble. SVR is built on regression analysis. To get a better result from a regression analysis, there must be a positive correlation between explanatory variables (factors) and dependent variable (dengue cases). As the primary model of prediction is SVR, this study needs a data analysis before moving forward with SVR. A separate correlation analysis was conducted for each factor (rain, temperature and population) to determine the suitability of the regression analysis of the proposed factors. Correlation is a statistical relationship between those two sets of data which describes the strength of the relationship between those two data sets in consideration. If the correlation is low there is a weak interdependency between those two sets. If the correlation is high (normally greater than 0.5 negative or positive), there is a considerable relationship between those two sets. Correlation of two data sets is computed as given in the equation below.

(1)

Where is the correlation between datasets X and Y. E is the expected value operator. is the mean of data set X, is the mean of data set Y. and are standard deviation of data sets X and Y respectively. The correlation value is interpreted as shown in Table 5.3.

Table 5.3 Correlation values and their meanings

|  |  |
| --- | --- |
| Correlation Value | Interpretation |
| -1 | A perfect downhill (negative) linear relationship |
| -0.7 | A strong downhill (negative) linear relationship |
| -0.5 | A moderate downhill (negative) relationship |
| -0.3 | A weak downhill (negative) linear relationship |
| 0 | No linear relationship |
| +0.3 | A weak uphill (positive) linear relationship |
| +0.5 | A moderate uphill (positive) relationship |
| +0.7 | A strong uphill (positive) linear relationship |
| +1 | A perfect uphill (positive) linear relationship |

## 5.4 Pre-processing of Data

### 5.4.1 Year-Wise Data Normalization to Eliminate Year Specific Influences

Rainfall pattern for each district is a recurrent pattern that has little variation from the average rainfall data for the year. Dengue incidences are varying from year to year due to various reasons. Case data has a constant and strong correlation to rainfall data. The correlation of the rainfall data is specific to the reference year and does not hold in general due to various other influencing factors. There may be a boost or a decline in the number of cases due to some other influences such as temperature, special environmental events such as flood and droughts, etc. Rainfall pattern along with dengue incidence for the district Amnat Chareon is given in the Figure 5.4 for six years starting from 2012. This variation is not cooperating well with machine learning tools specially with regression tools. Therefore, I use year-wise normalization to eliminate or reduce the impact of the afore mentioned effects from the rainfall and dengue incidences. The normalized dengue incidence pattern along with the rainfall data for the district Amnat Chareon is given in the Figure 5.5.

Figure 5.4 Monthly Rainfall and Incidence Data for Six Years from 2010

Figure 5.5 Normalized Monthly Rainfall and Incidence Data for Six Years from 2010

### 5.4.2 Outlier Removal

Dengue case data is reported partly or fully. If the data reported is partial, then the relationship between influencing factors and dengue cases is not clearly seen. Some elevated numbers have the same effect on the modeling. It is critical to identify these instances in advance and treat them properly to ensure the quality of the generated predictive model. These unusual instances are known as outliers. I proposed a method of outlier removal in which a convex hull is used to determine the outliers. This method uses the fact that the outliers (extreme points) lie further to actual points. As the correlation between influencing factors and the dengue cases reported is strong, the chance of having an observation that position further away from the cluster of points is less. This technique is shown in the Figure 5.6.



Figure 5.6 Outliers in Data Points

Proposed method operates in two stages. Identification of the presence of an outlier and removal of identified outlier. In the first stage, a convex hull generated for the two-dimensional representation of each influencing factor to the dengue cases reported. This stage is shown in the Figure 5.7 with and without outliers. Then the surface area covered by the convex hull is computed. Next, the points that formed the convex hull are removed from the original data set and regenerate the convex hull for the remaining points. The presence of an outlier is determined by a significant difference in surface areas computed. The algorithm advances to the second stage to identify the exact outlier from the set of points that formed the convex hull. Finally, I remove the identified outlier from the original dataset.





Figure 5.7 Outlier Removal Levels

5.4.2.1 Iterative Convex Hull Reduction to Remove Outliers

Outlier removal takes place as an iterative process. First, the convex hull of the two-dimensional dataset is generated. The area of the outer most convex hull is computed. In the next iteration, the points that formed the convex hull are removed from the original data set and regenerate the convex hull for the remaining points. Compute the difference in areas of the convex hull generated from the previous iteration to the current iteration. Iteration stops if the difference in areas is below a predetermined threshold value or preset number of maximum iterations reached. If the difference in area is significant, that indicates the presence of outlier and hence need to determine the outlier. The theory behind the outlier removal process is that the area difference is directly proportional to the number of outliers and the severity of the outlier (how far from the cluster of points).





Figure 5.8 Three Levels of Outlier Removals

Let ΔA is the difference in area between two iterations.

5.4.2.1 Determination of Point to be Removed from Outer Convex Hull in the Presence of Outliers

Once the presence of outliers detected, it is necessary to identify the exact outliers from the points on outer convex hull. This step is important as all the points that make outer convex hull are not outliers. I follow the exhaustive search approach to identify the outliers among points on outer convex hull. I compute the difference in surface area keeping one point of the convex hull out. All the points that generate a difference in area greater than or equal to predetermine threshold value are classified as outliers. All the remaining points will remain in the dataset.

## 5.5 Model Generation

### 5.5.1 Support Vector Machine

Behavior of each factor (rainfall, temperature, and population density) on dengue cases is spatially dependent. That is the effect of rainfall on dengue for each district is different from district to district as presented in [5]. Hence, a separate analysis for each district was conducted and a separate model for each province was generated. Data for five years were combined for each district and fed into the model for training. The proposed arrangement can capture the spatial heterogeneity of each province and hence improve the performance of prediction model.

The SVR model is based on the regression analysis. A regression analysis can estimate the relationship between two data sets (random variable) and fit a cure to the data sets (explanatory variable and dependent variable). This curve can then be used in prediction of unknown cases. The regression curve for this study has three explanatory variable, Rain R, Population P, and Temperature T. The regression model for this study is given in the equation below.

(2)

Where Pi is the population in *i*th region, Ri is the rainfall for *i*th region and is the temperature for *i*th region. The error term is . is the dengue cases for region *i*. Intercept is and it is a constant.

SVR improves the detection speed as it keeps only a subset of training data as support vectors in the model. The SVR uses the same principles as the Support Vector Machine (SVM) for classification, with only a few minor differences. SVR’s output is a real number which makes it difficult to match target output on test dataset. A margin of tolerance (epsilon) is set in approximation to the SVM to address the problem associated with real numbers output.  General construction of SVR is given in the following equations.

SVM regression is constructed by first mapping the input vector X into an m-dimensional feature space using a non-linear mapping function. The linear regression model is then constructed in this feature space. The linear model  is given by equation 3.

   (3)

Where   denotes a set of nonlinear transformations and b is the “bias” term. The bias term can be dropped with the assumption of zero mean data set. is the normal vector.

The quality of estimation is measured by the loss function given in equation 4. The loss function is computed as proposed in [16].

(4)

Then the empirical risk function is given in equation 5.

           (5)

The model generated by minimizing the . This can be achieved by introducing (non-negative) slack variables to measure the deviation of training samples outside http://kernelsvm.tripod.com/index_files/image032.gif-insensitive zone. Thus the SVM regression is formulated by minimization of the function given in equation 6.

(6)

Such that

This optimization problem can be transformed into the dual problem and its solution is given by the equation in 7.

(7)

Such that:

Where is the number of Support Vectors (SVs) and the kernel function is given by the equation in 8.

  (8)

The RBF was used as the kernel function and epsilon was set to 0.001. The cost parameter was kept at 100.

### 5.5.2 k-Nearest Neighbor Regression

The k-nearest neighbor (k-NN) Regression is a conventional non-parametric classifier [67]. The k-NN classifier calculates the distances between the testing point and points in the training data set. There are several distance calculation methods available. Euclidean distance is the most popular metric. K is an integer. The value selected for k has a direct impact of the predicted value and hence must be carefully selected for the problem domain. For this study, Euclidean distance was the distance metric and number of neighbors was three. The final predicted value is the average of k nearest neighbors.

To measure the distance between points A and B in a feature space, various distance functions have been used in the literature, in which the Euclidean distance function is the most widely used one. Let A and B are represented by feature vectors A = (x1, x2, …, xm) and B = (y1, y2, …, ym), where m is the dimensionality of the feature space. To calculate the distance between A and B, the normalized Euclidean metric is used by

### 5.5.3 Least Square Regression

Linear least squares regression is a widely used modeling method. It is also referred to as regression, linear regression or least squares. linear least squares regression can be used to fit the data with a function of the form

in which

1. Each explanatory variable in the function is multiplied by a coefficient
2. There is one constant
3. All terms are summed to form the final function

Error minimization is done as the same way as SVR. The only difference is the way the model is generated. In SVR it uses support vectors to keep the model parameters.

## 5.6 Prediction

A vector of unseen data for rainfall, temperature and population data is fed into the trained model and estimated output is obtained from the SVR, KNN, and LS. This output is not a label as in SVM. It is a real number approximating the number of dengue cases pertaining to the given scenario. The number of cases predicted show cases the severity of the condition that may occur if the given scenario appears in the future.

## 5.7 Model Validation

Conventional regression models are evaluated based on the MSE of the cross validation (mostly 10-fold cross validation). MSE cannot capture the total picture of the behavior of the data set. Several outliers can affect the final outcome of the validation. Another problem of regression analysis is there is no way of computing the accuracy of the prediction with cross validation. Regression gives real values as estimates and there is theoretically infinite number of possibilities with a real number. This fact makes it impossible to compare against target value. Accuracy is computer as per the following equation in SVM like classifiers. SVM like classifiers are based on class labels and hence makes it possible to compute accuracies conveniently.

(9)

In this study a novel yet, simple accuracy calculation method was introduced. A positive confidence boundary parameter α was included in cross validation. If *|actual value – estimated value| > α*, we label the estimated value as a correct prediction and incorrect prediction otherwise. The proposed accuracy calculation is given in the Figure 5.9.

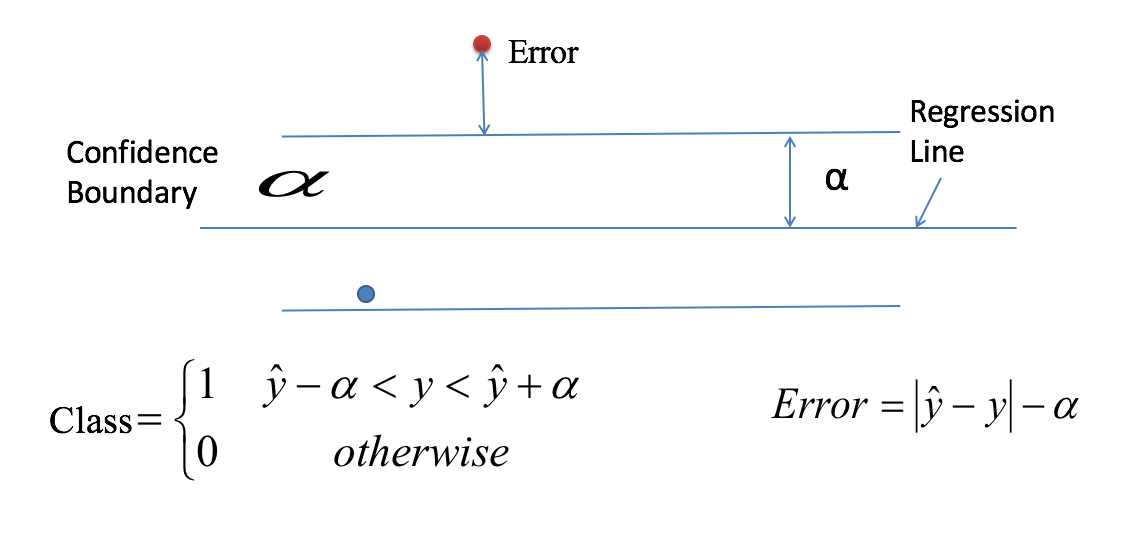


Figure 5.9 Accuracy Calculation of SVR

### 5.7.1 Determination of the Degree of Fit of the Regression Model to the Dataset with Parameter Alpha (α).

The value of *α* is inversely proportional to the model accuracy. If the model generates a higher accuracy value for a lower value of *α,* the regression model fits well to the dataset. If the model accuracy is high only for a large value of *α, the* dataset is loosely correlated to the influencing factors. Higher accuracy for a smaller alpha value indicates that the dataset and the fitted regression is a best fit for the problem domain.

### 5.7.2 10—Fold Cross Validation

Cross-validation is a model validation technique for assessing how the results of an analysis will generalize to the underlying population of the extracted data set. It is used in predictions where one needs to evaluate the outcome of the model. In a prediction problem, a model is usually given a training dataset with known outcome values, and a dataset of unknown outcome data. The goal of cross-validation is to test the model’s ability to predict new data that was not used in training stage of the model. In this study, I used 10-fold cross validation where dataset is divided into 10 subsets. One portion is kept as the testing data set and 9 portions are used in training. This process is repeated 10 times alternating the testing portion from the initial division. Final outcome of the model is obtained averaging the 10 outputs of the process.

## 5.8 Resource Allocation

A successful mitigation plan includes an effective resource allocation scheme. An effective resource allocation should eliminate wastage, over allocation and under allocation of resources while satisfying the demand of each facility for a given resource. Over allocating may result in run out of available resources before every demand is satisfied. Under allocating may result in failure of mitigation plan. Hence, an effective resource allocation is needed to ensure a proper distribution of available resources among facilities to satisfy their demands.

### 5.8.1 Problem Definition

Let F1,F2….Fn is the set of facilities demanding for resources R1, R2,……Rm with a demand for each resource W00, W01,…….Wnm. with the constraint . As the available resources may not be sufficient enough to fulfill the demand of each facility, the allocation algorithm must generate an optimum allocation that minimizes the penalty of allocating less than the demand. The final allocated scheme is represented as A00, A01,…..Anm. This problem can be modeled as a bi-partite matching between facilities and resources with a weight o each mapping as the demand. I represent this problem using a bi-partite grape as show in figure.

### 5.8.2 Problem Representation

The problem of resource allocation with demand can be successfully represented with a bi-partite graph. The graph is constructed as resources and facilities as two sides and demand as weight of associations. The graphical representation can be seen in Figure 5.10. In the figure resources are represented with R0…Rm, facilities are represented with F0,…Fn and corresponding demands are represented with W00,…..Wnm.

F1

Fn

Rm

Rm

W02

Wij

Figure 5.10 Bi-Partite Graph of Resource Mapping

### 5.8.3 Digital Representation

The resource allocation problem is represented as an adjacency matrix that is algorithm friendly. Resources are arranged along x axis and facilities are arranged along y axis. Weight (demand) of the matching is the corresponding element of the matrix.

Demand of resources

F1

.

.

Fn

R1 . … … … Rm

Allocation of resources

R1 . … … … Rm

F1

.

.

Fn

### 5.8.4 Finding the Optimum Solution for Resource Allocation Problem

The best allocation is to deliver each facility resources they demanded. The scarcity of resources prevents us from delivering the demanded amount of resources for all the facilities. It is required to find the optimum resource allocation that will minimize the impact of being delivered below the demanding amount. It is required to determine an effective method of calculating the impact of resource allocation on each facility. I suggest calculating a penalty score for each facility based on its important properties. The severity of allocating fewer resources than demanded is depended on the population that needed to be served and surface area of the serving region. The penalty score is calculated according to the equation given below.

The goal of the minimization algorithm is to minimize the total penalty score which is given in the equation below.

### 5.8.5 Weight Adjustments

The best approach is to use exhaustive search to look for the best match for weight for allocated matrix. The number of candidates in the exhaustive search space is increased exponentially with the number of facilities and number of available resource categories. The exhaustive search is then become resource intensive and is not an ideal candidate to find weights of the allocated matrix. I propose to use Genetic Algorithm approach with several modifications to match the problem domain.

Genetic Algorithm can be successfully applied in searching for a best fit from large number of candidate solutions. GA performs best when the available number of candidates is high. State of the art GA is defined as follows.

Genetic Algorithm is an optimization technique built on the principle of Genetics and Natural Selection. GA comes under the category machine learning. Given a set of possible solutions to the given problem, GA tries to select the best match based on the criteria set in advance. The GA selects a subset of population that meets the given criteria. This subset undergoes two steps, namely crossover and mutation, producing new children similar to natural process of selecting best fit candidates. This process is repeated over several iterations. Each candidate solution is assigned a fitness value and the fitter individuals are given a higher chance to mate and yield more “fitter” individuals.

GAs is randomized in nature when selecting crossover points and mutation points. The randomness in crossover and mutation is not applicable in my problem domain. I introduce a variant of crossover and mutation such that the resulting off springs are compliance with the requirements of the resource allocation.

I propose several major modifications to the standard GA for achieving optimum results for the resource allocation problem. These modifications are generalized and can be applied to any resource allocation problem. The modifications are listed below.

1. Introducing chromosome representation for resource allocation
2. Change the way the standard GA generates the initial population.
3. Chane the way the standard GA does crossover operation
4. Change the way the standard GA does mutation
   1. Introduction of resource lock chromosome
   2. Introducing sliding mutation scheme to accelerate the convergence
5. Change the iteration of the GA to accommodate sliding mutation scheme

All the above stages are detailed in the following sections.

### 5.8.6 GA Representation of the Problem

The problem of allocating set of resources among set of facilities must be represented in a single string of genes called a chromosome. I select decimal numbers in each location of gene as oppose to binary digits in standard GA. The binary chromosome and the proposed chromosome are given in the Figure 5.10 below.

Standard binary chromosome

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| 1 | 1 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 0 |

Proposed chromosome

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| W11 | W12 | .. | W1m | .. | .. | Wn1 | Wn2 | .. | Wnm |

Gene Boundary for Facility 1

Figure 5.11 Modified Chromosome for Genetic Algorithm

As shown in the figure, resources for a given facility must be placed in adjacent genes of the chromosome. Gene boundary is used here after throughout the document as to represent the set of resources for a given facility.

Resource allocation requirements that must be embedded in GA are given below.

* Total number of allocated resources must match the total number of available resources when performing mutations.
* The Gene Boundary must not be broken apart when performing crossovers.

### 5.8.7 Proposed Population Generation Procedure

As opposed to the standard GA I use a customized population generation technique. In the proposed scheme, a constraint is set forth and eliminates chromosomes that violate the constraint. All the resulting chromosomes allocate resources that are under the limits of availability of resources. The constraint is modeled as shown in equation below. The number of facilities is n and number of resources is m.

Constraint -> Available amount of resources (i)

### 5.8.8 Proposed Crossover Operation

Crossovers are performed such that the gene boundaries are preserved. A gene boundary is treated as a single entity in crossover and hence the problem reduced to binary crossover in standard GA. A sample crossover is shown in Figure 5.11.

Chromosome 1

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| W11 | W12 | .. | W1m | .. | .. | Wn1 | Wn2 | .. | Wnm |

Crossover point

Chromosome 2

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| W11 | W12 | .. | W1m | .. | .. | Wn1 | Wn2 | .. | Wnm |

Resulting Chromosomes

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| W11 | W12 | .. | W1m | .. | .. | Wn1 | Wn2 | .. | Wnm |

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| W11 | W12 | .. | W1m | .. | .. | Wn1 | Wn2 | .. | Wnm |

Figure 5.12 Proposed Cross-Over Operation

### 5.8.9 Proposed Mutation Operation

The proposed mutation operation selects number of mutation points based on the mutation rate as like standard GA. It is required to make sure that the resulting chromosome is adhering to the requirements set forth by the problem domain as opposed to standard GA. In standard GA it is simply select random points based on the mutation rate and flip the selected bit. In the proposed mutation operation, if the resulting chromosome demands more resources than available amount of resources after mutation it is not a good fit. A random mutation is not suitable in the proposed problem domain. A constraint must be set in mutation operation to consider the total amount of allocation for a given resource must not exceed the total amount of availability of the resource.

I randomly pick a resource type from the list of available resources. Next, set of facilities is randomly picked to apply the mutation. A resource exchange operation takes place between selected facilities for randomly selected resource type. This will make sure that the allocated amount of resources is not exceeding the available amount of resources. The proposed mutation operation is shown in Figure 5.12. Let the mutation operation randomly picked facility 1 and facility n for resource exchanging. All the other facilities remain unchanged and resource type 1(Wi1) was randomly selected for mutation.

Chromosome 1

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| W11 | W12 | .. | W1m | .. | .. | Wn1 | Wn2 | .. | Wnm |

Facility 1 Facility n

Mutation -> W11 = W11 – a Mutation -> Wn1 = Wn1 + a

Figure 5.13 Proposed Mutation Operation

### 5.8.10 Fitness function

The fitness function is modeled as a function of risk associated with the given facility based on the location of the facility, population count of the facility, amount of area to be served by the facility. The proposed method of fitness calculation will make sure that resources are allocated appropriately. Final fitness value is computed for all the facilities (from facility Fi=0 to n) for all the resources (from resource Rj=0 to m) together as shown in the equation below.

Where the risk is obtained from the ensemble model for each district. The higher the predicted dengue incidence for a given district the higher the risk. The predicted value is obtained for a future feature set containing future rainfall, future temperature and population.

5.8.11 The Proposed Concept of Locked Genes

In the real-world settings, the possibility of having more of a certain resource than the total demand from all the facilities is not rare. This will bring up a concern that the amount of time we spend on mutating on that resource is a waste of computational time. We can improve the performance of the GA by eliminating these resources in the iterations. I proposed a concept of locked genes in to the standard GA in which genes that has more resources than requested are locked. The GA does not mutate the locked genes. There will be a in-memory locked chromosome as a reference for GA to use in each iteration. Sample lock chromosome is given in Figure 5.13. The sample chromosome is generated for two facilities requesting five resources. The first resource is abundant and hence needed to be locked. Other four resources are not abundant and hence needed to be allocated efficiently via GA.

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| 1 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 |

Figure 5.14 Lock Chromosome for two facilities with five resources. First resource is abundant and hence locked.

### 5.8.12 Proposed Sliding Mutation Scheme

Standard GA uses random mutation and crossover operations. This will inherently produce random children and hence the fitness of the generated child will be varying. The algorithm selects chromosomes randomly to apply the two operations. This will lead to lower the fitness of the selected best fit chromosome as well. To reduce the fluctuation of fitness, I introduce a new method of mutation in each iteration. The proposed method will make sure that the next generation will be generated based on a sliding mutation scheme where the amount of mutation applied to a given gene is proportional to the risk associated with the chromosome. This method will mutate chromosomes with an amount that proportional to the risk associated with the chromosome. This will drive each chromosome gradually towards the best fit. The cross over operation will ensure that the GA will not end up in a local minimum. The experimental results proved that the proposed scheme perform very well and resulted in a higher bet fit value than standard GA in all the trials that performed. The proposed sliding mutation is given in equation below.

randomMutation = rand (0, (CurrentAllocation \* Risk));

The Figure 5.14 shows the pseudocodes of the proposed GA with modifications and standard GA. Added steps are bolded in the proposed GA.

|  |
| --- |
| **Algorithm 1**: Standard GA |
| 1: k ← 0;  2: PK ← InitPopulation(n) {Generating Initial Population with n individuals}  3: Compute fitness(i) for each i ∈ Pk; {Evaluate everyone in the population Pk: }  **4: While** not termination **do**  5: Select (1 − ChurnEntropy) × n members of Pk and insert into Pk+1; { Select Subset of PK:}  6: Select ChurnEntropy × n members of Pk; pair them up; produce offspring; insert the offspring  into Pk+1; { Crossover:}  7: Select MutationEntropy × n members of Pk+1; invert a randomly-selected bit in each;  {Mutate}  8: Compute fitness(i) for each i ∈ Pk; { Evaluate Pk+1:}  9: k ← k + 1;  **10: end while**  11: return the fittest individual from Pk; |

|  |
| --- |
| **Algorithm 2**: Proposed GA |
| 1: k ← 0;  2: PK ← InitPopulation(n, **resourcesMap**) {Generating Initial Population with n individuals and  constraints}  3: Compute fitness(I, facilityInfo) for each i ∈ Pk; {Evaluate everyone in the population Pk: }  **4: While** not termination **do**  5: Select (1 − ChurnEntropy) × n members of Pk and insert into Pk+1; { Select Subset of PK:}  6: Select ChurnEntropy × n members of Pk; **generate crossover with constraints**; insert the  offspring into Pk+1; { Crossover:}  7: Select MutationEntropy × n members of Pk+1; **perform constrained and sliding mutation**  **based on risk**; {Mutate}  applyRiskBasedPenalty for each individual; {applying over and under allocation penalty}  8: Compute fitness(i) for each i ∈ Pk; { Evaluate Pk+1:}  9: k ← k + 1;  **10: end while**  11: return the fittest individual from Pk; |

Figure 5.15 Standard and Proposed GA for resource allocation

### 5.8.13 Time and Space Complexity Analysis of the Proposed GA

5.8.13.1 Space Complexity

Space complexity is a constant for the proposed GA. Let L be the number of genes in the chromosomes, N is the number of chromosomes in the population, and m is the number of generations it produces. As it replaces the existing least fit chromosomes with newly generated best fit chromosomes resulting in the same number of individuals in the population, the population will not grow along with the iterations. Proposed GA uses two times the space required to store the generated population as the same way it does in the standard GA.

Hence, the space complexity of the proposed GA is O(2\*L\*N) and it is a constant complexity.

5.8.13.2 Time Complexity

It is not an easy task to formulate an exact equation for the calculation of time complexity of the GA. GA uses evolutionary algorithm making it hard to come up with an exact complexity value. But it is possible to formulate an estimated complexity for the GA. In this section, I formulate a complexity estimator for the proposed GA.

Let Gk is the kth generation, N is the number of individuals in each generation, L is the length of the chromosome, O(crossover) is the complexity of the crossover operation, O(fitness) is the complexity of the fitness function which depends on the implementation of the fitness function and O(mutation) is the complexity of the mutation operation. The proposed GA can adapt fitness function and mutation scheme according to the problem domain. And pm and pc are the entropy of mutation and crossover operations. In this study, O(fitness), O(crossover) and O(mutation) are O(L) operations with is the time complexity is a constant.

Hence the total time complexity is the summation of computations required by the fitness function, crossover operation and mutation operation. At each iteration there will be sorting operation performed to get the best fit chromosome. Hence, it is adding to the computation complexity.

Let’s consider Gk generation,

Number of fitness calculations (Fk ) = NL

Number of crossover operations (Ck ) = pc\* N

Number of mutation operations (Mk ) = pm\*N\*L

Hence, the number of total computations needed at kth iteration

= Fk + Ck + Mk

Time complexity O(GA) = O(

O(GA) = O(Nm)

The proposed GA has a constant time complexity as the standard GA does.

# Results:

## 6.1 Pre-analysis of Data

I conducted correlation analysis of the dataset used in this study. It is vital to have a strong correlation between dependent and independent variables to build a better SVR model. The initial correlation analysis did not show a strong correlation between rainfall and case incidences. The literature strongly pointed out that there is a strong correlation between rainfall and dengue incidence. This finding strongly suggested to have a preprocessing of data to eliminate noise in dataset which is the main influencing factor to have a very low correlation value. And also, I had to find out that is there any other factors affecting the correlation between rainfall and dengue incidence. Therefore, I performed several data preprocessing steps to eliminate external influences on rainfall and dengue incidence. The initial correlation analysis is given in the Figure 6.1. The correlation for the global model containing all 76 districts is 0.523.

Figure 6.1 Correlation between Rainfall and Dengue Incidence for Raw Data

## 6.2 Preprocessing of Data

### 6.2.1 Data Normalization

Initial correlation analysis did not produce a desirable result. There were several external influencing factors contributing to the low correlation values. The main reason for that to happen was the effect of year specific event happened during the period of data collection. I used a dataset which combines multiple years’ worth dengue incidences and weather data. There is a possibility of having a special event impacts such as flooding on reported data and hence the effect can be elevated or declined. To minimize the effect of special event in individual year, I normalize the data set year wise. The improvement is visible and reflected well in the correlation analysis of the dataset before and after normalizing. The complete correlation analysis for all districts with and without data normalization is given in Figure 6.2.

Figure 6.2 Correlation comparison with and without Normalization

### 6.2.2 Outlier Removal with Convex Hull Iterative Approach

I suggested to use an iterative outlier removal method. The experimental analysis revealed that each level of outlier removal process increased the correlation between rainfall and dengue incidence. The concept behind the approach is to eliminate outliers and hence increase the correlation. The following set of figures show three levels of outlier removal process and their corresponding correlation values. Figure 6.3 shows set of districts undergone outlier removal and Figure 6.4 shows correlation values for each district in each level of removals. The remainder of the outlier removal graphs are attached in Appendix A.

|  |  |
| --- | --- |
| Amnat Charoen | Ang Thong |
| Bangkok | Bueng Kan |

Figure 6.3 Multi Level Outlier Removal with Convex Hulls (v1-rainfall, v2-case incidence)

Figure 6.4 Correlation for all Districts at each Outlier Removal Level

It is clear from the Figure 6.4 that correlation between rainfall and case incidence is increased with the outlier removal. For a handful of districts, the correlation is not improving as expected. There can be other influencing factors or reporting errors for those districts. For most of the districts there is a good or considerable improvement which is helping SVR to perform better.

The remainder of the results are organized as follows. Geostationary behavior of the dengue epidemic is studied in Sri Lanka and results are presented. The generation of prediction models was done for Thailand and results are presented. As per the fact that there is no real data available for resources, a thorough testing was conducted on synthetic data for resource allocations with the proposed Genetic Algorithm approach.

I conducted a thorough analysis I Sri Lanka which gave a clear understanding on how the dengue epidemic is geographically varying. The study conducted in Sri Lanka revealed the usage of district specific prediction models will outperform global prediction models. The results are listed in the following section.

## 6.3 Results of GWR and Least Square Analysis of Dengue Epidemics in Sri Lanka

### 6.3.1 Least Square Analysis

In OLS results VIF values shows whether the predictor variables are multicollinear. VIF < 10 means the variables are not multicollinear. In this study, rainfall and population density were used as explanatory variables. VIF value determined these two factors are not correlated and hence can be used in regression analysis together. Every explanatory variable used is unique and contributing to the variation in dengue incidence. OLS regression result also shows that the Adjusted R-Squared value is 0.332054 for the year 2014. This indicates the model built with a combination of population density and rainfall data explains 33.2% of the variation in dengue incidences. According to the OLS regression results, all explanatory variables (rainfall and population density) are statistically significant but the value for Jarque-Bera statistics is also significant. Significance in Jarque-Bera statistics indicates the model is biased and hence undesirable. Also, the Koenker test is statistically significant (P value < 0.01 for both rainfall and population). This implies non-stationary relationship between the dependent and some or all of the explanatory variables. That reveals the explanatory variables (rainfall and population density) behave differently in different spatial regions.



Figure 6.5 Spatial distribution of dengue incidence from 2011 to 2015 in Sri Lanka.

GWR analysis

For GWR, adaptive kernel was used as kernel type and AIC was selected as bandwidth. In GWR AIC value determines the performance of the model. AIC can be used to compare two different models generated with regression analyses. AIC value for OLS is greater than in GWR. Hence GWR is a better analysis tool for dengue incidence with rainfall and population density as explanatory variables.

The GWR model results show that the Adjusted R-Squared values is 0.5632 (R2= 0.621). This indicates the model generated with population density and rainfall as explanatory variables can explain 56.3% of the variance in dengue incidences in 2014. These results also reveal that there are other variables besides population density and rainfall data that has stronger relationships with dengue incidence. These variables are not included in the model. But the model cannot provide a clue of those variables so that they cannot be included in the model. They have to be identified by experimenting with various candidate explanatory variables.



Figure 6.6 GWR standard residual map for dengue incidence with rainfall and population density for the year 2014.

The standard residual map for the model developed for dengue incidence is shown in the Figure 3. The red areas indicate under predictions where the actual number of dengue cases is higher than the model predicted values. The blue areas indicate over predictions where actual dengue cases are lower than predicted values. Random locations of red or blue areas indicate the model performs fairly well. Red or blue clustered areas indicate under/over prediction of the model and hence the performance of the model is poor. Spatial clustering of over/under prediction indicates missing one or more key explanatory variables in the model. The standard residual map in Figure 3 shows clustered over and under predicted areas.

It is required to find how well each explanatory variable predicts the dengue incidence for each administrative region. It is revealed from previous sections that there are no global explanatory variables that hold consistent relationship across administrative regions. An analysis was conducted to reveal the variation in strength of explanatory variables in each administrative region in explaining the relationship between the variable and dengue incidence. Results of the analysis are shown in Figure 6.7 (a) and (b). Figure 6.7 (a) provides the spatial distribution of regression coefficients for rainfall and Figure 6.7 (b) provides the same for population density. Lighter colors represent lower coefficients and darker colors represent higher coefficients.

Mapping these coefficients shows the relationship between each explanatory variable and the dependent variable that how they change across the study area. The darker areas in figures indicate the explanatory variables, rainfall and population density, are strong predictors of the dengue incidence, whereas, the lighter areas are locations where they are comparatively weak.

GWR regression results show that relationship of incidence with rainfall and population density is spatially varying across districts of Sri Lanka. Figure 6.7 (a) shows that spatial distribution of regression coefficient of population density is a strong predictor in eastern coastal areas mainly in Trincomalee district, and a weak predictor in Mannar. Figure 6.7 (b) also shows that spatial distribution of regression coefficient of rainfall is a strong predictor in northern areas including Mannar and in eastern coast it is a weak predictor. There is an inverse effect of rainfall and population density on dengue incidence. When rain becomes a strong predictor in some areas population density is a weak predictor and vice versa. It is very important to understand this variation for making local policies to mitigate dengue. GWR model can also be used to predict values of dependent variables for locations within the study area with unseen explanatory variables values. This will give an estimate of the dependent variable (dengue incidence) using the regression model generated.



Figure 6.7 spatial distribution of regression coefficients for (a) population density (b) rainfall.

## 6.4 Generating Prediction Models for Dengue Epidemic in Thailand

Multiple phases of testing were conducted on the same data set with multiple level of noise removals. A separate SVR model was trained for each district. Each model was validated using 10-fold cross validation. SVR models are assessed using MSE and proposed accuracy calculation method. Finally, multiple classifiers are combined and generated a micro ensemble to eliminate bias in each classifier for a given district.

### 6.4.1 Prediction Results for Global Model

Global model contains data from all 76 districts. Each data point is treated the same way as all the other and geographical variations re not considered. There is a single SVR is trained on the entire data set. The plot of rainfall vs dengue incidence is depicted in Figure 6.8. The results for various scenarios are listed below.

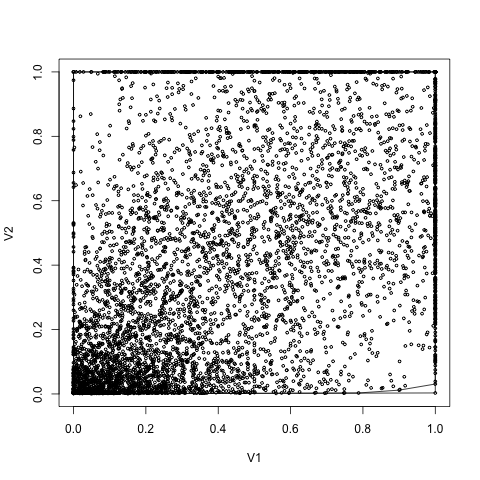


Figure 6.8 Plot of Rainfall vs Dengue Incidence for the Global Dataset (v1-rainfall, v2-dengue incidence)

Correlation of dengue incidence to rainfall for global dataset is 0.523. As per the figure, it is clear that the global model has many outliers that make it to have lower correlation coefficient. It is hard to determine an optimal value for the level of outlier removals. Correlation coefficients for 3 levels of outlier removal is given in the Table 6.1. SVR model validation results from 10-fold cross validation with no outlier removal is given in the Table 6.2. Average accuracy and MSE for level1 and level2 outlier removal on global dataset are given in Table 6.1.

Table 6.1 Correlation Coefficients for Three Levels of Outlier Removals on Global Dataset

|  |  |
| --- | --- |
| Level of outlier removal | Correlation Coefficient |
| Level 0 | 0.523 |
| Level 1 | 0.524 |
| Level 2 | 0.525 |

As per the table, it is clear that the increase in correlation coefficient in each level of outlier removal is very small. It is advisable to have many levels of outlier removals to get a higher coefficient for correlation. There is no means by which the optimum number of outlier removals are estimated. The results obtained from the global model using 10-fold cross validation is listed in Table 6.2. Note that the accuracy is calculated based on the method proposed in this study.

Table 6.2 Results of 10-fold Cross Validation of SVR on Global Dataset

|  |  |  |
| --- | --- | --- |
| Fold | Accuracy | MSE |
| 1 | 63.25 | 0.19322216 |
| 2 | 56.85 | 0.20835374 |
| 3 | 59.23 | 0.20811646 |
| 4 | 55.94 | 0.21453015 |
| 5 | 63.07 | 0.19367864 |
| 6 | 59.41 | 0.20107433 |
| 7 | 58.68 | 0.20458908 |
| 8 | 60.51 | 0.2005876 |
| 9 | 60.14 | 0.20600364 |
| 10 | 59.04 | 0.20545753 |
| Average | 59.61 | 0.20356133 |

### 6.4.1 Prediction Results for Local Models

Each district is considered as a separate entity and having unique characteristics. Hence, the models generated for individual district is specific to that particular district. The results of local models trained for 76 districts are given below.

There are three models generated for the same data set and validated based on 10-fold cross validation. The best models are then used in ensemble generation and achieve better output from the ensemble model. The ensemble model then be used in prediction and risk estimation. The models are generated based on SVR, Least Square and K-NN tools. Performance of each model in each outlier removal level along with the performance of ensemble are given in Figure 6.9, Figure 6.10 and Figure 6.11 for level of outlier removals 0, 1 and 2 respectively. It is very clear from the figures that the accuracy improvement with each level of outlier removal.

Figure 6.9 Model Performance without Outlier Removal

Figure 6.10 Model Performance with Level1 Outlier Removal

Figure 6.11 Model Performance with Level2 Outlier Removal

## 6.5 Resource Allocation

I performed resource allocation on four different synthetic datasets confirming to the setting listed in the Table 6.3. I listed the complete setup for the Trial 1 and list only best fit value graph and the allocation results for the remaining trials. All the other results are found under appendix A.

Table 6.3 Experimental Trial Setup

|  |  |  |  |
| --- | --- | --- | --- |
| Trial No | Number of facilities | Number of Resources | Facility Properties |
| 1 | 10 | 10 | 4 |
| 2 | 50 | 5 | 4 |
| 3 | 100 | 10 | 4 |
| 4 | 500 | 10 | 4 |

Performance Comparison of GA with random and sliding mutation with lock chromosome

### 6.5.1 Trial 1

Table 6.4 Facility Information (High risk facility is highlighted in red)

|  |  |  |  |
| --- | --- | --- | --- |
| **Population** | **Risk** | **Area** | **put** |
| 293931 | 4435 | 259 | 655 |
| 424873 | 2722 | 619 | 233 |
| 300904 | 1930 | 743 | 264 |
| 222133 | 4372 | 195 | 954 |
| 185651 | 2426 | 328 | 185 |
| 257318 | 2368 | 354 | 742 |
| 453762 | 2720 | 522 | 419 |
| 141650 | 1956 | 493 | 174 |
| 402820 | 1555 | 128 | 720 |
| 259771 | 1944 | 840 | 466 |

Table 6.5 Resource Availability

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **R1** | **R2** | **R3** | **R4** | **R5** | **R6** | **R7** | **R8** | **R9** | **R10** |
| 1304 | 7441 | 7297 | 980 | 6276 | 9691 | 5735 | 6476 | 2754 | 146 |

Table 6.6 Requested Resources from each facility

|  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Resource**  **Facility** | **R1** | **R2** | **R3** | **R4** | **R5** | **R6** | **R7** | **R8** | **R9** | **R10** |
| F1 | 43 | 369 | 362 | 49 | 208 | 481 | 190 | 214 | 91 | 5 |
| F2 | 66 | 564 | 553 | 74 | 317 | 734 | 290 | 327 | 139 | 7 |
| F3 | 55 | 467 | 458 | 61 | 262 | 608 | 240 | 271 | 115 | 6 |
| F4 | 207 | 1770 | 1735 | 233 | 995 | 2305 | 909 | 1027 | 437 | 23 |
| F5 | 102 | 875 | 858 | 115 | 492 | 1140 | 450 | 508 | 216 | 11 |
| F6 | 207 | 1770 | 1735 | 233 | 995 | 2305 | 909 | 1027 | 437 | 23 |
| F7 | 129 | 1108 | 1087 | 146 | 623 | 1444 | 570 | 643 | 273 | 14 |
| F8 | 164 | 1400 | 1373 | 184 | 787 | 1823 | 719 | 812 | 345 | 18 |
| F9 | 189 | 1614 | 1583 | 213 | 908 | 2102 | 829 | 936 | 398 | 21 |
| F10 | 143 | 1225 | 1201 | 161 | 689 | 1595 | 629 | 711 | 302 | 16 |

Table 6.7 Lock Chromosome

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 |

Performance of the GA for random, no-sliding with constraints and sliding with constraints GAs. There are 10 facilities with four properties. Each facility is requesting 10 different resources. The performance of each category is shown in Figure 6.12.

Figure 6.12 Performance of Proposed GA for 10 Facilities Requesting 10 Resources

Resource Allocation Results of Trial 1 for high risk facility and low risk facility are given in Figure 6.13 and Figure 6.14.

Figure 6.13 High risk facility

Figure 6.14 Lowest risk facility

### 6.5.2 Trial 2

Performance of the GA for random, no-sliding with constraints and sliding with constraints GAs. There are 50 facilities with four properties. Each facility is requesting 5 resources. The performance of the trial 2 is given in Figure 6.15.

Figure 6.15 Performance of the Proposed GA for 50 facilities and 5 Resources

The high-risk facility is the facility with id 27. The following graphs (Figure 6.16) shows the resource allocation for the facility 27 with random GA and sliding with constrained GA.

Figure 6.16 Resource Allocation for the High-risk Facility of Trail 2

### 6.5.3 Trial 3

Performance of the GA for random, no-sliding with constraints and sliding with constraints GAs. There are 100 facilities with 4 properties that request for 10 resources. Performance is given in Figure 6.17

Figure 6.17 performance of the Proposed GA for 100 Facilities with 10 Resources

The high-risk facility is the facility with id 49. The following graphs (Figure 6.18) shows the resource allocation for the facility 49 with random GA and sliding with constrained GA.

Figure 6.18 Resource Allocation for the Trail 3

### 6.5.4 Trial 4

Performance of the GA for random, no-sliding with constraints and sliding with constraints GAs. There are 500 facilities with 4 properties that request for 10 resources. Performance is given in Figure 6.19.

Figure 6.19 Performance of the Proposed GA for 500 Facilities with 10 Resources

The high-risk facility is the facility with id 377. The following graphs (Figure 6.20) shows the resource allocation for the facility 377with random GA and sliding with constrained GA.

Figure 6.20 Resource Allocation for the Trail 4

## 6.6 Comparison of proposed GA with sliding mutation against standard GA with random allocation and mutation

Standard GA with random population generation and mutation is always start with a lower fitness value. The variation in fitness value among each individual is very high giving a very lower value of average fitness value compared to best fit value. In contrast, proposed GA is always starts with a higher fitness value for both best fit values and average fitness value. The difference between best fitness and average fitness is small for the proposed GA as the constrained based population generation always produces offspring that are closer to the target chromosome. These observations are clearly shown in Figure 6.21.

Figure 6.21 Comparison of Standard GA and the Proposed GA

# Conclusion

A computational approach for dengue epidemic prediction and mitigation was proposed. The framework consists of preprocessing of data where noises and inherent influences from other factors removed from data sets. Different behavior of dengue disease in each year is identified and regularized using year-wise data normalization. Noises were removed from data sets with the proposed outlier removal process using iterative convex hull approach. The behavior of the disease was modelled using an ensemble of regressors to overcome the geographical dependency of the dengue disease.

The proposed regressor ensemble was used to predict the dengue epidemic and identified the high-risk areas. This information is then used in proposed resource allocation method to efficiently allocate limited resources. The proposed resource allocation methodology was modelled based on the state-of-the-art Genetic Algorithm with major modifications to the algorithm. The mutation, crossover and initial population generations are modified to introduce constraints and to match the problem domain.

The proposed data cleaning, problem modelling, and resource allocation are promising and proved by the generated results.

## 7.1 Future Directions

A web-based system is proposed to manage every aspect of the dengue mitigation from data cleaning to resource allocation. This system will enable administrative staff to deploy an effective response plan for the upcoming dengue epidemic.

A mobile app is proposed to report case data in real time. Target audience of the mobile app is the general public. This will enable us to gather near real-time data from general public. A moderation mechanism needs to be implemented to validate the case data reported.

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