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1. Introduction

The geographic distribution and genetic diversity of dengue virus is deeply rooted in Asia suggesting its origin from this region, with first reported out-break of DHF from Philippine in 1953 (Halstead, 1980). One of the characteristics notable in Asian regions, where the disease is endemic is that dengue hemorrhagic fever outbreaks occur in repetitive cycles of 3-5 years, (Ferguson et al, 1999). The incidence of disease and its severity varies across different dengue virus serotypes and also between primary and secondary infections of same serotypes(Vaughn DW et al).

Due to lack of in-vivo study models, there is little information about factors contributing to disease severity and its variation across dengue virus genotypes and the cyclical nature of dengue outbreaks. It is however critical to study these factors particularly in the South East Asian region where incidence of dengue cases is thought to be associated with variables such as water, sanitation, population density and rate of literacy as opposed to developed countries where ambient temperature, moisture and rainfall perhaps plays the major role. A better understanding of disease epidemiology and pathogenesis will help identify optimum control measures in the region. It will also develop systems for predicting the outcome of mass vaccination when the vaccine becomes available in this region.

The chapter has been divided in three parts: the first part will discuss the historical evolution of the dengue virus in the region its spatial and temporal distribution. It will also look at the effects of covariates such as poverty, water supply, sanitation and global warming on expansion of the dengue endemic regions.

The second part of the chapter will focus on the genetic evolution of the viral isolates circulating in the region. Phylogenetic studies of dengue viruses have uncovered genetic variation within each serotypes, these variations have been organized in discrete clusters on dendograms. Analyses of such studies have broadened our horizon to relate the mutational changes with disease evolution and factors like seasonality and incidence variability. This part of chapter will focus on the common mutational variations that have been reported so far and how these relate with the disease dynamics in the endemic region.

In the third and final part of the chapter an attempt has been made to relate the mutational changes of dengue genotypes with disease severity. Vast array of literature has been published investigating relationship of genetic variation with disease severity. The structure
of virus E-protein that confers the viral infectivity and host immune response of the virus (E.Descloux, 2009) remains the focus of such studies. Sequence variation at different loci such as CprM, E/NS1, preM/E, C/prM/M and untranslated regions etc. have been investigated for its association with disease severity. This part of chapter will throw some light on our current understanding of disease severity and it relation with genetic variation.

2. Historical background of dengue virus in South East Asian

Geographically South East Asia comprises of land south of China to east of India extending as far as to the north of Australia. Although geographically the region is well defined, the list of countries included in this region varies due to political reasons. For the purpose of this review W.H.O based definition has been used. In addition status of dengue virus in further south of the region; including countries like Pakistan and Bangladesh have also been included to encompass the broader spectrum of the region.

2.1 Dengue vector evolution

Evidences suggest that vectors Aedes aegypti and Aedes albopictus originated from darker sylvan forms found in African tropical forests. It is believed to have reached New World from West Africa via slave ships during the 17th century (Gubler, D.J.1998). Aedes aegypti was introduced into the coastal cities of South East Asia from East Africa around nineteenth century via the shipping industry. With the eruption of World War II it became deeply entrenched in many cities (Gubler, D.J.1998). The mitochondrial genetic diversity studies have revealed circulation of two distinct clusters of Aedes aegypti in South East Asia one with strains from French Polynesia, Guinea and Brazil while the other cluster is of strains that migrated from Europa Island in Mozambique and Amazonia (Mousson et al 2005). In contrast; A. albopictus is known to be native to South East Asia. It has spread within past few decades to various countries primarily due to introduction of trade of used tyres worldwide. Using ecological niche modeling Benedict and co-workers have predicted the risk of global invasion by Aedes albopictus secondary to cargo trade and increasing air travel. Although temperate and humid climates are prerequisites for the optimum survival of both the vectors but A. albopictus is known to better acclimatize to the cold and dry weather due to its ability of efficient egg diapause during the extreme conditions, thus favoring its survival in the regions with exotic temperature ranges (Benedict, M.Q, et al 2007).

2.2 Factors leading to disease spread in SEA

The factors responsible for the insurmountable expansions of dengue in the region are complex and thought to be intricately linked with vector-host-virus triad, socioeconomic stresses and climatic variations. There are excellent reviews that discuss the impact of these factors in details (Aiken, S.R. 1978, Kendall, C. et al 1991, Halstead, S.B. 1966). Only salient factors in context of SEA will be discussed here. The distribution of DHF outbreaks in SEA correlates with emergence of mosquito A. aegypti in South East Asian countries perhaps due to displacement of indigenous A. albopictus in the region. This is considered to be associated with uncontrolled urbanization leading to shanty towns with inadequate pipe water supply and poor sanitation. A. albopictus is semi domestic species that breeds on natural and man-made breeding sights; it feeds on variety of animals, birds and man. The A. aegypti on the other hand is more
acclimatized to urban set-up, once established the density of this mosquito is directly proportional to density of human population and artificial breeding sites (Merril S.A et al 2005), it feeds almost exclusively on humans. Moreover, A. aegypti is considered to be more competent vector for dengue virus. Genetic traits that determines successful midgut infection by DEN virus have been mapped on several loci on A. aegypti chromosomes (Benedict, M.Q, et al 2007) indicating that vector competence is genetically determined.

The extent to which these mosquitoes compete with each other in the environment is not clear, nonetheless the balance of two species in the region is important, and the socioeconomic factors in SEA appear to be displacing A.albopictus in favour of A.aegypti leaving the population more susceptible. The poor socioeconomic conditions are major contributing factor to sustained vector activity with severe form of disease in the South East Asia. The breeding habitats of A.aegypti have been strongly associated with squatter settlements, inadequate piped water supply and sewage facilities (Halstead, S.B. 1966). In addition, there are impacts of higher environmental temperature in the region. High temperature is inversely related to the mosquito gonotropic cycle and viral extrinsic incubation period; this increases the egg laying episodes resulting in more blood meals and increased risk for viral transmission. In addition shorten extrinsic viral incubation period culminate to increase virus load at time of inoculation (Focks D.A. et al 1993). These effects have been proven for dengue vectors in simulation studies conducted by (Cox J et al 2001) and it has been projected that increase in global temperature would increase the length of transmission season in temperate regions.

2.3 Dengue fever and dengue hemorrhagic fever

The word dengue is believed to have originated from Swahili language “ki denga pepo”, which describes sudden cramp like seizure. The clinical symptoms suggestive of dengue virus infection can be traced back to Chinese Chin Dynasty (265-420 AD) where disease was considered as water poison and was known to be associated with water and insects (anonymous 2006).

Emergence of the disease in the new world can be traced back to the transmigration of the vector in the 17th century. There are reports that suggest possible epidemics of dengue like illness in three major continents (Asia, Africa and North America) as early as 1779 and 1780, within Asia Batavia (now known as Jakarta) was affected by this outbreak (Halstead,S.B. 1966). By early nineteenth century Dengue fever was known to be endemic in the rural areas of South East Asia probably due to the indigenous vector A.albopictus. It manifested as self limiting disease to which native population developed immunity at early age. With the advent of A. aegypti at Asian ports, the disease spread to the main inland cities and towns. It is assumed that unlike rural population, the urban populations of South East Asia remained susceptible to dengue virus and were then infected by newly imported vector. Dengue epidemics progressively became less frequent as urban population became immune to the disease, until 1953 when a new form of dengue fever was reported from Thailand and Manila, where children suffered from fever followed by bleeding diathesis; the disease was then called as Philippine Fever (Aiken, S.R. 1978). By 1960’s the hemorrhagic form of disease had spread to Malaysia, Vietnam, Sri Lanka, Singapore and Indonesia (Halstead,S.B. 1966). The disease epidemiology extended and outbreaks of dengue hemorrhagic fever (DHF) were reported from India (1988) French Polynesia (1990), Pakistan (1992) and Bangladesh (2000).Until recently, DHF was considered to be disease of childhood, especially in South
East Asia where mean age of cases under fifteen, and the modal age of five or slightly higher was reported from countries such as Thailand, Philippines and Malaysia, however, recent reports are now documenting increasing number of DHF and DSS in adult population as well (Khan E et al 2007). The precise cause of DHF/DSS remains elusive despite enormous research in this area. Evidences suggest interplay of multiple factors such as host genetic make-up with unique immune response and viral virulence may play a role in determining the severity of the disease.

2.4 Pathogenesis of severe dengue disease

There are two form of Severe disease, namely dengue shock syndrome (DSS) and DHF without shock. It is proposed that devastating coagulation derangements due to host immune response leads to heamorrhage and shock in severe cases. The concept of original antibody sin leading to immune enhancement is considered to be the main reason whereby infection with one type of dengue virus sensitizes an individual and that subsequent infection with different virus type elicits a hypersensitivity reaction (secondary infection). Various studies have been conducted to show the association of elevated cytokines in patients presenting with DHF and DSS. Elevated serum levels of cytokine and chemokines such as IL-2, IL-8, IL-6, IL-10, IL-13, TNF and INF-γ have been found to be significantly associated with patients presenting with DHF and DSS in clinical setting (Azeredo et al., 2001; Hung, et al., 2004, Clyde. K. et al., 2006). It has been proposed that the pro-inflammatory cytokines released by the cross reactive memory T-cells, induce plasma leakage by its effects on the endothelial cells (Eva.H. et al 2004; Aviruntanan et al., 1998). In fact in-vitro studies have rendered endothelial cell monolayers permeable by the application of chemokine such as IL-1β (Cardier et al.,2005). In vitro-and in-vivo models of studies also suggest role of decreased nitric oxide levels and its relation with IL-10 and raised viral load (Simmons et al., 2007). There is evidence that suggests relation of increased expression of certain cytokines such as IL-1β, TNF-γ, and IL-6 with elevated NO production (Guzik et al., 2003).

With the advances in genomic and bioinformatics tools the scope of genetic studies has greatly expanded particularly in depth data on genomic changes and its association with disease epidemiology, seasonality and severity has been made available. Growing availability of comparative genome sequence data has provided important insights into the molecular evolution of dengue virus. Evidence strongly suggests appearance of new strains correlating with DHF/DSS epidemics. Despite the wealth of genomic data now available the exact cause and effect of viral virulence and clade changes is yet to be proven, however it is quite evident that different serotypes and viral lineage is continually changing with local extinction and emergence of new clade and that the introduction of new clade in the region translates in form of outbreaks of DHF and DSS.

3. Distribution of dengue virus serotypes in SEA

Dengue like other RNA viruses is prone to genetic mutations as it replicates using RNA-Polymerase; enzyme that lacks proof reading mechanism. The mutation rates in the order of 10⁻³ has been reported for dengue (ElodieDes et al 2009) in different host settings. Such mutations often result in variants that become targets of selection; an outcome of underlying genotype and its environment. Despite these facts dengue virus do not evolve as fast as other RNA viruses. The only macro evolutionary divergence is perhaps the radiations in its four serotypes in its primate host (sylvatic strains) around one thousand years ago (ElodieDes
et al 2009). Thereafter genetic mutation in the envelope protein and receptor binding domains resulted in its emergence as infectious pathogen in human population. The divergent forms of these sylvatic strains are often found to be circulating in human habitat, suggesting that enzootic cycles with some spill over in the surrounding human population. This has been shown in Malaysian populations settled near forest and marshy habitats (Wang, E. et al., 2000). The phylogenetic studies conducted based on envelope gene sequences of basal portion of sylvatic lineage, DENV 1,-2,-4 of Malaysian descent suggest that endemic /epidemic strains of these viruses diverged from sylvatic ancestors more than 1000 years ago (Wang, E. et al., 2000). Thereafter, only micro evolutionary change within dengue serotypes have taken place, these changes have nevertheless resulted in substantial genetic diversity with emergence of endemic and epidemic strains in different parts of the region.

Fig. 1. The effects of climatic and social change on vector evolution and disease severity
DEN-3 viruses have undergone independent evolution which has resulted in emergence of four genetic subtypes of which subtype I-III circulate in the South East Asian Region. Subtype I comprises of viruses from Indonesia, Malaysia and the Philippines; subtype II of viruses from Thailand and subtype III includes viruses from Sri Lanka India and Pakistan. The genetic evolution in these subtypes is primarily reported mutations in the prM/M and E structural protein genes. In spite of these mutations, the genomic region has retained greater than 95% amino acid sequence similarity (Lanciotti, R.S et al.,1994), suggesting that these are highly conserved regions responsible for protein architecture and/or biological function. Phylogenetic studies suggest that there are regional foci of virus extinction and selection, one such region is Thailand where the indigenous DEN-3 virus circulating up to 1992 disappeared and was replaced by two new lineages perhaps from a common ancestor (Wittke, V. et al. 2002). The sequence of all Thai DEN-3 isolates recovered after 1992 had T at position 2370 in contrast to the C at this site in the pre-1992 samples (Wittke, V. et al. 2002), and nucleotides difference was observed in at least 45 sites of total 96 sites studied. It appears that the post-1992 strains have replaced the pre-1992 strains. These studies point towards potential of regular extinctions of strains of DEN-3 virus and replacement by new variants in the region (Wittke, V. et al. 2002). Natural selection and/or genetic bottleneck could be the plausible causes for this variation. Since the extinction of pre 1992 strains and appearance of new epidemic strain in Thailand occurred during inter-epidemic period it is therefore hypothesized that the genetic bottleneck is perhaps the cause of regional replacement. This is further supported by studies from India reporting shift and dominance of the dengue virus serotype-3 (subtype III) replacing the earlier circulating serotype-2 (subtype IV) with emergence of increased incidence of DHF and DSS in subsequent outbreaks (Dash, P.K et al. 2006). Strains from the 2005 outbreak in Karachi (Pakistan) were found to be similar to those from Indian strains of dengue serotype 3, and were responsible for deadly outbreak in 2005-06 (Jamil. B. et al. 2007). Thus over the period 1989 and 2000, a new clades of DENV-3 genotype III viruses have replaced older genotype and clades in this region and emergence of new clades coincided with severe epidemics. The epidemiologic data suggests that the DEN-3 virus responsible for recent epidemic outbreaks in Mozambique, Gutamaella, Pakistan and Sri Lanka may have been introduced from India, and changing age structure of dengue patients from 1996–2005 may also be indicative of the selected virus moving into new areas (Kanakaratne, N. et al.2009).

4. Genetic evolution and disease severity in SEA
The micro evolutionary change within dengue serotypes has resulted in substantial genetic diversity with emergence of endemic and epidemic genotypes. With current advances in the field of genetic and molecular techniques scientists are now trying to decipher relation of changing clades with disease severity and epidemic potential. With the availability of complete genomic sequence of the Dengue virus different genetic loci have been investigated to find this relationship. Envelope –gene (E-gene) sequence is the most frequently investigated locus, (Wittke,V.et al., 2002,-Thu, H.M. et al.,2004;Islam, M.A.et al., 2006,27) followed by capsuler C-prM gene (Kukreti, H. 2008;Dash, P.K. 2006;Kanakaratne, N. 2009;Jamil B,2007). In addition non-structural (NS) viral proteins such as NS1 and untranslated genomic region 3’-UTR, 5’ UTR along with complete genomic sequences have been investigated to relate the genetic changes with the disease severity (Mangada, M.N. et al., 1997;Zhou,Y.et al.,2006;Islam, M.A.et al.,2006. Despite the wealth of genomic data
available the exact cause and effect of viral virulence and clade changes is yet to be proven, however, viral lineage is continually changing with local extinction and emergence of new clade. The introduction of new clade in the region translates in form of outbreaks of DHF and DSS. In order to analyze if there is a selection of specific clade in South East Asia that is circulating in the region and causing DHF outbreaks we conducted a meta-analysis. Studies conducted from 1950 to 2009 in South East Asian region that have investigated association of disease severity with specific sequence mutations in the dengue virus genome were retrieved. The objective was to analyze association of disease severity with the specific genomic mutation in the clade circulating and causing periodic epidemics in South East Asia. Since DENV-2 and DENV-3 are more common in this region our study was focused on these two genotypes only. Objectives of the metaanalysis were to identify association of specific genetic mutation in DENV-2 and DENV-3 with clinical severity seen during periodic epidemics in South East Asia. The specific review question was:Is clinical severity of dengue in the South East Asian region associated with emergence of specific mutations in genomes of DENV-2 and DENV-3 genotypes? We hypothesized that there is changing pattern of dengue virus genotypes in South East Asia and these mutations are associated with clinical severity of the disease.

4.1 Methods
4.1.1 Literature search
The literature search was performed from February 2010 to June 2010. Data sources include Medline via Pubmed (1950-February 2010), Cochrane data base of systematic reviews, Google scholar and experts in the field. Secondary references and review articles were scanned for thematic review. Hand search of the journal was also carried out. However, unpublished and ongoing studies could not be explored. Terminologies i.e. dengue type 1-4,
dengue fever, dengue hemorrhagic fever, genetic variation, sequence analysis, south East Asia were used individually as well as in various combinations. Two independent reviewers; reviewed the titles, abstracts and full text articles and selected potentially relevant studies based on inclusion criteria established prior to the literature search. Discrepancy between the reviewers were sought to reach on consensus in consultation with third reviewer. Those potentially irrelevant studies that were ultimately excluded are listed together with the reason for exclusion in Table 1.

4.1.2 Inclusion criteria
Studies which reported dengue virus genotype (mutation / sequencing of viral genetic material) and clinical features of dengue fever patients were included.

4.1.3 Design of the studies
All type of observational studies i.e. case report, case series, surveys and descriptive cross-sectional studies which were focusing on genotype and clinical presentation of dengue patients were included in the review. Population: Population includes patients of dengue fever of all age groups. No age and sex restriction were applied. Outcome of interest: Difference in nucleotide and protein sequences were analyzed and compared according to geographical origin, the sampling period and the clinical presentation. Clinical severity of the disease is defined as presence of DF, DHF or DSS. Language: Only articles in English language were included in the review.

4.1.4 Exclusion criteria
All those studies focusing on dengue vector control, clinical trials on vaccines, clinical trials on drugs, pure prevalence or incidence, unusual case report or case series without genotype and studies conducted in countries other than south East Asian region were excluded.

4.1.5 Data extraction
Data extraction of the included studies was done by using structured data extraction form specifically made for the review. Data was extracted for country of origin, year of publication, clarification of objectives, type of study, its duration and setting, results on both genotype and clinical severity etc.

4.1.6 Data synthesis
A narrative data synthesis was carried out to show result summary of all included studies which include description of clinical features and genotype of dengue virus. However, meta-analysis could not be performed due to non availability of required data i.e. measure of strength of association. Hence, pooled effect of genetic variation on clinical severities among dengue patients could not be provided.

4.1.7 Quality assessment
According to Cochrane Collaboration’s recommendation, the quality of included studies have been assessed by using criterion which asses the quality of studies by focusing on study type, sample size calculation, clarity of objective, selection of cases, and internal validity of selected studies.
Fourteen studies were finally selected based on inclusion criteria i.e., association of dengue genotype and clinical severity of the diseases in the patients and were conducted in different countries of South East Asia. Setting of these studies were; Thailand 6 (Zhang, C. et al 2006, Rico-Hesse R. et al. 1998, Wittke, V. et al., 2002), Myanmar (Thu H.M. et al., 2004), India (Kukreti, H. et al., 2008; Dash, P.K. et al., 2006), Bangladesh (Islam, M.A. et al., 2006), Sri Lanka (Kanakaratne, N. et al., 2009), Taiwan (King, C.C. et al., 2008) and Pakistan (Jamil B, et al 2007). These studies were published from 1997 to 2009. Since the focus of our study was on DEN-2 and DEN3 viruses 12 studies out of these 14 were finally included in this study.
4.1.8 Study sample characteristics


Age ranges for dengue patients in these studies varied from 1 year to 70 years. The total numbers of dengue patients were 7663 in these studies. Characteristics of the studies included in this review have been summarized in table 2. A total of 285 virus isolates were subjected to genotyping/ sequence analysis in these studies. All four genotypes were studied in three studies (Zhou Y, et al 2006, Jarman RG, et al. 2008, Rico-Hesse R. et al. 1998); only DEN 3 in five studies (Wittke, V. et al. 2002, Kukreti, H. et al. 2008; Islam, M.A. et al. 2006), only DEN 2 in three studies (Zhou Y, et al. 2006, Mangada MNN et al. 1997, Zhang, C. et al. 2006), only DEN 4 in one studies (Klungthong C, et al. 2004), whereas DEN 1 and DEN 3 in one study (Kukerti H, et al. 2008) and DEN 1, DEN 2 and DEN 3 studied in one study (Jarman RG et al. 2008).

4.1.9 Clinical definition

Dengue case was defined on the bases of presence of IgM, IgG, or fourfold or greater rise in hemagglutination inhibiting (HI) antibody titer against dengue virus, and presence of dengue virus specific nucleic acids in RT-PCR. Clinical severity was defined as presence of hemorrhagic manifestation and DHF related symptoms such as thrombocytopenia, skin rash, gum bleeding, gastrointestinal bleeding, hemorrhagic sclera, epistaxis, edema and ascitis. Where as other studies simply defined as presence of DF, DHF grade I, II and III and DSS as per WHO criteria.

5. Nucleotide sequencing and phylogenetic analysis

Envelope -gene (E-gene) sequence was most frequently investigated loci, nine studies were focused on this region followed by C-prM gene, in three studies both genetic loci studied in one study (9) and one study included NS1 along with PrM and E loci. The 3’-UTR, 5’ and 3’ UTR and complete genomic sequences were studied in one each.

Homology search and comparisons of most obtained sequences were performed using commercially available software systems such as DNASIS, DNASTar, 3’ –UTR secondary structures were estimated using MFOLD package, while nucleotide sequence alignments (Phylogenetic analysis) were performed using CLUSTAL X, MEGA version, and maximum likely hood methods available e.g. PAUP PROGRAM.

5.1 Mutations observed

5.1.1 E-gene mutations

In case of DEN-2 virus, maximum numbers of viral isolates have been analyzed in studies from Thailand. The E-NS1 region of 77 different variants of DEN-2 studied using Maximum Parsimony analysis of 240 nucleotide sequence, showed 11 of 240 nucleotide variation; 4.6% divergence but did not reveal significant segregation of virus according to geographic location (Rico-Hesse R. et al. 1998). Similarly, Phylogenetic analysis of 120 E gene of DEN-2 by another group from Thailand has confirmed existence of six genotypes of this virus; however evolutionary relationships among the genotypes is difficult to determine (Zhang, C. et al 2006). In terms of dengue pathogenesis these studies failed to show segregation of DF versus DHF-associated viruses on the evolutionary tree. There are no clear-cut evolutionary divergence or branching of DF versus DHF isolates, suggesting that nucleotides from this region of the genome encode amino acids that are apparently not under immune selection (Rico-Hesse R. et al. 1998 and Zhang, C. et al 2006).

DEN-3 has replaced DEN-2 as most frequently isolated virus in Thailand since late 1980’s (Wittke,V.et al.,2002). The evolutionary history of Thai DEN-3 viruses, has been studied by comparative analysis of the nucleotide sequence of E protein genes of currently prevailing isolates with those from all previously published E gene sequences of DEN-3 virus available in Gen Bank (Wittke,V et al. 200218), this study has shown E-gene of DEN-3 to be relatively conserved at amino acid level, however, four amino acid changes have been identified within genotype II of Thai strains. The amino acid changes observed at positions (E172 I-V) and (E479 A-V) are the only difference found between pre and post-1992 viruses. Similarly, there is little evidence to support in-situ evolution among the virus samples that were studied over prolong period ranging from days to months in a selected community in Thailand(Jarman, R.G.et al.,2008) very few mutational changes were noted, and association of these mutations with disease severity could not be delineated either. Analysis by E-sequence of eight DEN-3 strains from Bangladesh (2002 out-break strains) were found to be very closely related to Thai isolates that caused out-break in 1998 in Thailand. The multiple alignment of amino acid (aa) sequence revealed that Bangladeshi isolates and Thai isolates shared common aa changes at position E127 (I-V), suggesting that 2002 outbreak in Bangladesh was due to introduction of Thai isolates (Islam, M.A.et al.,2006), however the statistical association of aa changes with disease severity could not be delineated. In case of Sri Lankan DEN-3, type III is the most frequent strain with two distinct clades IIA and IIB linked to mild and severe disease epidemics on the island respectively (Kanakaratne, N. et al.,2009). Phylogenetic studies of E-NS1 junction of DEN-2 isolates from Sri Lanka has categorized the isolates into 4 genotypes designated as Malaysian/Indian subcontinent, Southeast Asian, American, and West African (Sylvatic) and Sri Lankan isolates are closely related to Indian / Malaysian genotype.

5.2 C-prM mutations

The phylogenetic analysis of 433 base pair region (nucleotides 180—612) of the DEN-3 CprM gene junction showed that sequences of Delhi isolates (2006 outbreak) were closely related to sequences from Guatemala (1998) and presented a nucleotide identity of 95.9—98.2% (mean 97.05%). On comparison of Delhi 2006 sequences with other Indian sequences from years 2003, 2004, and 2005, mean sequence divergence of 2.85%, 2.15%, and 1.6%, respectively, were observed (Kukreti, H. et al.,2008. Common amino acid mutations observed in 2006 DENV-3 sequences are given in table 3. Similar study performed on DEN-3 isolates of 2003-04 outbreaks in New Delhi, found them to be closely related and belonged to subtype III from Sri Lanka (Dash, P.K.et al.,2006). Moreover, Phylogenetic analysis of C/PrM/M region of
DEN-3 isolates from Pakistan (2004-05 outbreak isolates) also showed sequence homology with 2003-04 New Delhi outbreak strains suggesting that circulation of common isolates of DEN-3 subtype III in the region. There was no clear statistical association of disease severity with the specific isolates or geographical origin.

<table>
<thead>
<tr>
<th>DENV-Protein</th>
<th>Geographical Origin</th>
<th>year</th>
<th>aa change (positions)</th>
<th>Relation to Disease severity</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Envelope (E) Den 3</td>
<td>Thailand</td>
<td>2008</td>
<td>Phylogenetic analysis reveals multiple genetic variants (mutational positions not mentioned)</td>
<td>Could not be ascertained</td>
<td>Jarman RG</td>
</tr>
<tr>
<td>DEN 3 E region</td>
<td>Taiwan</td>
<td>2008</td>
<td>E301 (L to T)</td>
<td>Distinct clade causing epidemic outbreak</td>
<td>King, C.C. et al., 2008</td>
</tr>
<tr>
<td>Envelope (E) Den 3</td>
<td>Bangladesh</td>
<td>2005/6</td>
<td>E81 (I-T) E140 (I-T) E127 (I-V)</td>
<td>No association with disease severity could be determined</td>
<td>Kukreti H</td>
</tr>
<tr>
<td>Den 3 C-preM/E</td>
<td>Srilanka</td>
<td>2003-06</td>
<td>Phylogenetic analysis reveals multiple genetic variants (mutational positions not mentioned)</td>
<td>2 distinct clades linked to mild (IIIA) and severe (IIIB) disease epidemics</td>
<td>Kanakaratne et al</td>
</tr>
<tr>
<td>C-prM</td>
<td>India</td>
<td>2006</td>
<td>C-prM108(M-I) C-prM112(T-A)</td>
<td>May be attributed to increased incidence of DHF &amp; DSS in India</td>
<td>Jamil B, et al</td>
</tr>
<tr>
<td>DEN 3 prM region</td>
<td>Taiwan</td>
<td>2008</td>
<td>CprM 55 (L-H) PrM 57 (T-A)</td>
<td>No association with disease severity could be determined</td>
<td>King CC</td>
</tr>
</tbody>
</table>

Table 2. Genetic Characteristics and relation to disease severity in patients with DEN-3 infections reports from South East Asian Region
with specific serotype, as viruses isolated from DHF patients fell at different locations on the phylogenetic tree (Kukreti, H. et al., 2008; Dash, P.K. et al., 2006). Using maximum likelihood and Bayesian approaches, phylogenetic analysis of Taiwan's indigenous DENV-3 isolated from 1994 and 1998 dengue/DHF epidemics were found to be of three different genotypes - I, II and III each associated with DEN-3 circulating in Indonesia, Thailand and Sri Lanka, respectively (King, C.C. et al., 2008). The authors of this study analyzed complete nucleotide sequence of DEN-3 for its mutation and its relation with regional evolution. The highest level of nucleotide sequence diversity, and the positive selection site was detected at position 178 of the NS1 gene. Although the authors have identified the NS1 gene as the positive selection site and the envelope protein site for purifying selection pressure, however direct association of these changes with disease severity was not determined. Study from Bangkok Thailand performed sequence analysis on E/NS-1 region of Thai isolates to determine if viral strains from less severe DENV infections had distinct evolutionary nucleotide pattern then those with more severe form (Rico-Hesse R. et al., 1998). This study found that two distinct genotypes were identifiable from both DF and DHF cases, suggesting its evolution from common progenitor that perhaps shares the potential to cause severe disease.

<table>
<thead>
<tr>
<th>DENV-Protein</th>
<th>Geographical Origin</th>
<th>year</th>
<th>aa change (positions)</th>
<th>Relation to Disease severity</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>E/NS1 (77 DEN-2 virus strains studied)</td>
<td>Thailand</td>
<td>1998 from 1980</td>
<td>11 nucleotides (4.6% divergence) between Strain PUO-218-280, 22 nt or 9.2% divergence PUO-218-D80141</td>
<td>No specific association with disease severity</td>
<td>(Klunthong C)</td>
</tr>
<tr>
<td>E/NS1 junction Den 2</td>
<td>Sri Lanka</td>
<td>2003-06</td>
<td>239-nt (from positions 2311–2550)</td>
<td>Could not be ascertained</td>
<td>(Kanakaratne et al)</td>
</tr>
<tr>
<td>3' and 5' UTR Den 2</td>
<td>Thailand</td>
<td>1996-97</td>
<td>5' NCR homologus 3' UTR trinucleotide change 297±299 (two transversions and one transition)</td>
<td>Trinucleotide change may alter the functional characteristic of Secondary structure</td>
<td>(Mangada MNM)</td>
</tr>
<tr>
<td>Den 2 E/C/NS2A</td>
<td>Thailand</td>
<td>2006</td>
<td>approx 10^{3} substitutions</td>
<td>no apparent association</td>
<td>(Zhang C)</td>
</tr>
<tr>
<td>3'-UTR</td>
<td>Thailand</td>
<td>1973 to 2003</td>
<td>Variable secondary structures were detected</td>
<td>No clear association</td>
<td>(Zhou Y)</td>
</tr>
</tbody>
</table>

Table 3. Genetic Characteristics and relation to disease severity in patients with DEN-2 infections reports from South East Asian Region
5.3 Untranslated Region (UTR) mutations
The 3’ UTR region is thought to play a pivotal role in the DENV biology; it contains several conserved regions as well as 3’ long Stable Hair Pin structure which is conserved among all the members of the family Flaviviridae. It has been proposed that this structure interacts with viral and host nucleic acid and protein factors to form a complex to regulate transcription and replication (Zhou, Y. et al., 2006). Therefore it appears to play a significant role in the efficiency of RNA- translation, and virus ability to cause infection, hence the role of 3-UTR in determining the severity of dengue disease seems plausible. The literature reviewed under this study did show considerable intra-serotype diversity at 3-UTR region with greatest variability seen in DEN-4 followed by DEN-1.

A comparative analysis of 3’ UTR conducted for DENV isolates from Bangkok, Thailand compared Thai sequences with 61 globally sampled isolates of DENV taken from patients with varying disease severity. Although some genetic variations were found both within and among the serotypes notably at 3’ Long Hairpin Stable structure, however these mutations did not show consistent association with the clinical outcome of the DENV infection (Zhou, Y. et al., 2006). Study focusing on terminal 3’ 5’UTR sequences of four DEN-2 from Thailand 1998 outbreak strains, showed complete homology for sequences at 5’ UTR (highly conserved region) when compared with the prototype virus New Guinea C strain.

Fig. 2. The Geographical distribution of mutations in DEN-2 and DEN-3 viruses detected at different genomic loci of isolates from South East Asia

6. Conclusion
Ades aegypti was introduced into the coastal cities of South East Asia from East Africa around nineteenth century via the shipping industry. With the eruption of World War II it
Genetic Diversity of Dengue Virus and Associated Clinical Severity During Periodic Epidemics in South East Asia

deeply entrenched in many cities. The distribution of DHF outbreaks in SEA correlates with emergence of mosquito *A. egypti* in South East Asian countries due to uncontrolled urbanization leading to displacement of indigenous *A. albopictus* from the region. Phylogenetic analysis suggests that there are foci of virus extinction and selection in South East Asian region, one such region is Thailand where the indigenous DEN-3 virus circulating up to 1992 has disappeared and replaced by two new lineages perhaps from a common ancestor. These studies point towards potential of regular extinctions of strains of dengue virus particularly DEN-3 virus and replacement by new variants in the region. Natural selection and / or genetic bottleneck are plausible causes for this variation. Since the extinction of pre 1992 strains and appearance of new epidemic strain in Thailand occurred during inter-epidemic period we therefore hypothesize that the genetic bottleneck is perhaps major cause of regional replacement. This is further supported by studies from India reporting shifting and dominance of the dengue virus serotype-3 (subtype III) replacing the earlier circulating serotype-2 (subtype IV) with emergence of increased incidence of DHF and DSS in subsequent outbreaks. Strains from the 2005 outbreak in Karachi (Pakistan) were found to be similar to those from Indian strains of dengue serotype 3, and were responsible for deadly outbreak in 2005-06.

Despite the growing genomic data base in the gene bank there are fundamental gaps in our understanding of epidemiological and evolutionary dynamics and its relation with disease severity. There are two possibilities that explain the association between clade replacement and increased viral virulence. The first is the possibility of these viruses to be better fit and therefore produce high viremia in infected humans, consequently with better transmission of virus by the vector. The other hypothesis to explain the possible virulence of emerging clades in the region is its improved ability to avoid neutralization by serotypes cross reactive antibodies (Kochel et al., 2005). Thus there is relative abundance of different serotypes and viral lineage is continually changing in South East Asia. In face changing threshold of host immunity, periodic epidemics of DHF and DSS is due to local extinction and emergence of new clades. Over the period 1989 and 2000, a new genotype of DENV-1 and new clades of DENV-3 genotype III viruses have replaced older genotype and clades in this region and emergence of new clades coincided with severe epidemics. Thus South East Asia displays greatest degree of genetic diversity, suggesting that it is the hub for the evolution of new epidemic strain. However, selection of specific clade and association of specific sequence variation with disease severity at various genomic levels reported in the literature reviewed in this study lacks strength of association i.e. reporting Relative Risk (RR)/ Odds Ratio (OR) limits our interpretation regarding causality or pin pointing specific clade with virus virulence, and therefore further studies are recommended.

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8. References


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Tropical Medicine has emerged and remained as an important discipline for the study of diseases endemic in the tropic, particularly those of infectious etiology. Emergence and reemergence of many tropical pathologies have recently aroused the interest of many fields of the study of tropical medicine, even including new infectious agents. Then evidence-based information in the field and regular updates are necessary. Current Topics in Tropical Medicine presents an updated information on multiple diseases and conditions of interest in the field. It includes pathologies caused by bacteria, viruses and parasites, protozoans and helminths, as well as tropical non-infectious conditions. Many of them are considering not only epidemiological aspects, but also diagnostic, therapeutical, preventive, social, genetic, bioinformatic and molecular ones. With participation of authors from various countries, many from proper endemic areas, this book has a wide geographical perspective. Finally, all of these characteristics, make an excellent update on many aspects of tropical medicine in the world.

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