Spatio-temporal modelling of climate-sensitive disease risk: Towards an early warning system for dengue in Brazil

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1. Introduction and motivation

The early identification of an epidemic of infectious disease is an important first step towards implementing effective interventions to control the disease and reducing mortality and morbidity in human populations (Kuhn et al., 2005). However, an epidemic is often under way before the authorities are notified and control measures are put in place. In this paper we assess the potential for using seasonal climate forecasts to provide early warnings of future increased and geographically specific risk of dengue fever in Brazil.

Dengue fever and its more severe form (dengue hemorrhagic fever) is one of the most important emerging tropical diseases at the beginning of the 21st century in terms of morbidity and mortality (Gubler, 2002; Guzman and Kouri, 2003). Dengue is an acute viral disease characterised by fever, headache, muscle and joint pains, rash, nausea, and vomiting. Dengue haemorrhagic fever is a potentially deadly complication that in severe cases, can cause circulatory failure. Dengue viruses are transmitted by the mosquito with widespread distribution in tropical cities (Hayden et al., 2010). Dengue transmission is influenced by many factors, including climate, which influences mosquito biology and interactions between the mosquito vector and dengue virus (Kuno, 1995; Scott et al., 2000; Sanchez et al., 2006). Dengue is endemic in many tropical and subtropical countries. However, epidemic...
dengue transmission displays a seasonal pattern in response to temperature and rainfall variability (Johansson et al., 2009). There have been recent concerns of a worldwide spread of dengue fever, as a result of climate change, that could favour an expansion of the transmission area (Epstein, 2001; Hales et al., 2002).

In Brazil, the greatest incidence of cases occur from January to May when the climate is warmest and most humid (Braga and Vallee, 2007). Three of the four dengue virus serotypes have spread throughout Brazil, where reported dengue cases in the last decade represent about 60% of dengue cases reported in the Americas as a whole (Nogueira et al., 2007a). Dengue epidemics impact heavily on the national health services. There is no specific treatment for dengue, but appropriate medical care frequently saves the lives of patients with the more serious dengue haemorrhagic fever. A major epidemic occurred in Brazil in 2008, with 787,726 reported cases (January to November) including 4137 cases of hemorrhagic fever and 448 deaths.1 In Rio de Janeiro, military field hospitals were opened during the 2008 outbreak to help to ease the pressure on emergency rooms packed with people suffering from dengue.2

The current monitoring system in Brazil relies on observing dengue incidence in December/January to estimate epidemic potential late in the austral summer. However, this does not provide a quantitative measure or much predictive lead time. The greater the lead time available for forecasting disease risk, the greater the opportunity for effective disease risk intervention, although long term predictions often involve larger errors. Myers et al. (2000) suggested that epidemic forecasting is most useful to health services when case numbers are predicted two to six months ahead. This would allow time for the allocation of resources to interventions such as preparing health care services for increased numbers of dengue patients and educating populations to eliminate mosquito breeding sites, i.e. by regularly emptying water that accumulates in discarded refuse, tyres and domestic water storage containers, commonplace in urban slums/favelas found in some areas of Brazil.

As seasonal climate forecasts predict seasonal or monthly average temperature and precipitation (and other variables) for the forthcoming months/season in both time and space, they could potentially be used in a national dengue early warning system (EWS) for Brazil to aid epidemic planning months in advance. EWS based on seasonal climate forecasts have been developed to predict malaria incidence, for example in Botswana (Thomson et al., 2006), but there has been limited progress in developing EWS for dengue fever. Therefore, the use of seasonal climate forecasts with lead times of one month or more within a dengue EWS is a research area in need of exploration.

Before assessing the viability of using seasonal climate forecasts in a dengue prediction model, a model driven by observed climate variables with time lags relevant to dengue transmission, issued at the same resolution as the climate forecasts, must first be evaluated. If a significant relationship is identified between observed climate and dengue in Brazil, the use of forecast climate for dengue prediction purposes could be valuable. The remainder of this paper focuses on the viability of using observed climate variables in a spatio-temporal dengue prediction model.

In Section 2 we outline some of the key processes involved in dengue transmission and describe the data used in the statistical modelling. Section 3 documents the exploratory data analysis and model selection process to find which monthly climate variables and time lags are statistically significant for modelling dengue incidence in Brazil, using a negative binomial generalised linear model (GLM) to allow for overdispersion. In Section 4 we focus on the South East of Brazil, where dengue predominates, and refine the previously selected Brazilian global model by reverting to a Poisson formulation and explicitly including spatially unstructured and spatio-temporal structured random effects via a Bayesian framework to account for unobserved/confounding factors. Section 5 then assesses the ability of the refined model to issue dengue epidemic warnings for the peak dengue season in 2008 when a serious epidemic occurred. The final section discusses future ideas for research and summarises the main findings of the paper.

2. Dengue transmission

A number of complex factors are related to dengue transmission. In particular population growth and unplanned urbanization, resulting in substandard housing, inadequate water, sewerage and waste management systems, which allow mosquito reproduction. Poverty and health inequality are behind almost all of these factors (Gubler, 2002). Given favourable climatic conditions for development of the dengue-carrying mosquito, the urban environment plays a major role in determining transmission rates. Rainfall may influence the filling of containers out in the open (e.g. old tyres) which create potential breeding sites for the mosquito. More importantly, the breeding of mosquitoes depends on temperature, humidity, the mosquitoes' life expectancy, life-long fecundity, biting activity and virus incubation (Favier et al., 2005). Several previous studies have examined the link between climate and dengue. In many tropical countries, a positive association between rainfall and dengue incidence have been documented (Li et al., 1985; Moore et al., 1978; Gould et al., 1970). However, a significant relationship was not found for other regions (Eamchan et al., 1989; Goth et al., 1987; Kuno, 1995). Some authors have found that time-lagged climate variables of up to two or three months have a statistically significant association with dengue (Li et al., 1985; Schreiber, 2001; Wu et al., 2007). Precipitation and temperature oscillations over large parts of Latin America and the Caribbean are strongly influenced by changes in Pacific sea surface temperatures (SST) as part of the El Niño Southern Oscillation (ENSO) (Glantz, 2001) and these in turn can influence vector competence and surviviorship. Several studies have also used some index of ENSO to model dengue (Brunkard et al., 2008; Cazelles et al., 2005; Gagnon et al., 2001; Hales et al., 1999). Therefore, the inclusion of covariates based on the urban environment, climate (e.g. temperature, precipitation, Pacific SST) and their lagged effects appear to be potentially important components of a climate informed dengue prediction model.

Dengue fever data (counts of confirmed cases per month) from January 2001 to December 2008 (96 months) were obtained at municipality level (5651 municipalities) from SINAN DATASUS—an Information System for Notifiable Diseases, established by the Brazilian Ministry of Health.2 A network of laboratories, capable of diagnosing dengue infections, has been implemented in all states. The network is responsible for confirmation of cases to support epidemiological surveillance (Nogueira et al., 2007b). However, this network is not accessible to all municipalities. Dengue counts were aggregated to the microregion level (558 microregions), where a microregion typically consists of one large city and

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several smaller municipalities. This alleviates problems of misreporting due to variation in the availability of health services and epidemiological facilities at the municipality level. Fig. 1a shows dengue counts for this period grouped into the five main regions of Brazil (Fig. 1b) and Fig. 1c shows the total dengue cases in each microregion for the period January 2001–December 2008. Dengue is most prevalent in the South East. Two major epidemics occurred in the late austral summer of 2002 and 2008, while considerably less dengue occurred in 2004 and 2005. There is very little dengue in South Brazil and the North West Amazon.

National cartographic data such as altitude and biome were obtained from the Brazilian Institute for Geography and Statistics (IBGE). Census data at the microregion level such as population, percentage of urban population and the percentage of households with a water supply provided by a network, refuse collection and at least one bathroom, was obtained from an aggregated database SIDRA maintained by IBGE. Each microregion belongs to an administrative main region (1. North, 2. North East, 3. South, 4. South East, 5. Central West) and a biome (1. Amazon Rainforest, 2. Caatinga, 3. Cerrado, 4. Atlantic Rainforest, 5. Pampa, 6. Pantanal). A spatial variable named zone was defined according to the six biomes but by also subdividing the Atlantic Rainforest biome into three areas (North, South East and South) according to different climatic regimes. For example, south of the Tropic of Capricorn (23.5°S) the climate is more temperate and humid, while in the North East portion of the Atlantic Rainforest the climate is relatively warmer. Therefore, eight zones are defined for which climatic, geographical and ecological conditions are approximately homogeneous. In a modelling context, zone is treated as a categorical variable, or factor. Fig. 2 shows the spatial distribution of altitude and urban population in Brazil and the location of the geographical zones (Figs. 2a–c). Figs. 2d–f illustrate the relationship between these covariates and standardised morbidity ratio (SMR) for the given time period where, for a microregion i, the SMR is defined as the ratio of observed \( y_i \) to expected \( e_i \) dengue cases in the time period. The expected cases \( e_i \) in each microregion are calculated as the population at risk \( p_i \) multiplied by the global dengue detection rate over the whole of Brazil for the time period \( e_i = p_i \times \sum y_i / \sum p_i \). Altitude has a statistically significant negative relationship with dengue SMR (as altitude increases, dengue counts decrease) and percentage of urban population had a statistically significant positive relationship, given a microregion with excess risk of dengue fever (SMR > 1), as urban areas are ideal environments for mosquitoes and many people living in close proximity create a human virus reservoir.

Fig. 3 illustrates that dengue has a strong annual cycle which differs between geographical zones. The spatially varying dengue annual cycle is included in the model specified in Section 3, as an interaction between the categorical variables zone and month. As only part of the cycle may be attributable to climatic conditions, the inclusion of this interaction could account for other confounding variables, such as seasonal population movements, leading to differences in the annual cycle across zones.

Observed gridded (2.5° × 2.5° latitude–longitude grid) monthly mean precipitation data was obtained from the Global Precipitation Climatology Project (GPCP) (Adler et al., 2003). Reanalysis monthly mean surface air temperature data was obtained from the NCAR/NCEP Reanalysis (Kalnay et al., 1996). These climatic variables are referred to as ‘observed’ climate for the remainder of the text. Niño 3.4 is an index used to measure the strength of El Niño and La Niña events (Barnston et al., 1997) and is defined as the departure in monthly sea surface temperature from its long-term mean averaged over the region (120°W–170°W and 5°S–5°N). A positive (negative) index indicates El Niño (La Niña) conditions. A time series of the monthly Niño 3.4 index was obtained from NOAA Climate Prediction Center.

Microregion and gridded data were combined by assigning a grid point to each microregion on the basis that the microregion is contained within the grid square (see Fig. 4).

3. Model selection using a generalised linear model

Poisson models are widely used in the analysis of count data. However, it is well established that observed count data, e.g. disease cases, often display substantial extra-Poisson variation, or overdispersion (Lawless, 1987). Overdispersion was evident in this dengue dataset. Fitting a Poisson generalised linear model (GLM) involving the full set of explanatory variables described earlier results in a residual deviance more than a hundred times larger than the residual degrees of freedom, implying that as the mean dengue count increases, the variance increases at a much greater rate. In Section 4 we will consider making explicit allowance for this overdispersion within the Poisson framework via the inclusion of appropriate random effects, but for model selection purposes within this section we accommodate overdispersion implicitly by using the negative binomial distribution
for the observed counts, viz:

$$f(y; \mu, \theta) = \frac{\Gamma(y+\theta)}{\Gamma(\theta)^y} \left( \frac{\mu}{\mu+\theta} \right)^\theta \left( \frac{\mu+\theta}{\mu} \right)^y,$$

with mean $\mu$, scale parameter $\theta$ and variance function $V(\mu) = \mu + \mu^2/\theta$. The log-link $g(\mu_i) = \log \mu_i$ is utilised to allow comparison of point estimates to a Poisson model.

In order to select which explanatory variables are important for modelling dengue counts in Brazil for the 96 month period (January 2001–December 2008), the negative binomial GLM described above was fitted using the MASS package (Venables and Ripley, 2002) in R (R Development Core Team, 2008), starting with a maximal model based on all of the covariates described in the previous section, i.e. spatial covariates related to the urban environment, altitude, the annual cycle and interactions with geographical zone, observed climate variables with associated time lags (0–3 months) and the Niño 3.4 index with time lags of up to 6 months. Exploratory analyses were then carried out using different subsets of variables to select an...
appropriate prediction model (e.g. examining model fit with and without climate information and with different interactions). These analyses were assisted by the use of stepwise model selection algorithms based on the Akaike information criterion (AIC), which not only rewards goodness of fit, but also includes a penalty that discourages overfitting. The final most parsimonious model which emerged from the investigation is as follows:

\[ y_{it} \sim \text{NegBin}(\mu_{it}, \theta) \]

\[
\log(\mu_{it}) = \log(e_i) + \sum_j \beta_j x_{jit} + \sum_j \gamma_j w_{jit} + \sum_j \delta_j z_{jit},
\]

where \( y_{it} \) is dengue count for microregion \( i=1, \ldots, 558 \) and time \( t=1, \ldots, 96 \), \( \mu_{it} \) is the corresponding mean dengue count and \( \theta \) is the scale parameter. The expected cases \( e_i = p_i \lambda \) are treated as an offset in the model based on the population \( p_i \) in microregion \( i \) and the overall average dengue rate per month \( \lambda \). The variables \( x_{jit} \) represent the selected climate influences: precipitation one month previous \( (j=1) \), precipitation two months previous \( (j=2) \), temperature one month previous \( (j=3) \), temperature two months previous \( (j=4) \) and Niño 3.4 six months previous \( (j=5) \). The variables \( w_{jit} \) are: altitude \( (j=1) \) and percentage of urban population \( (j=2) \). Finally, \( z_{jit} \) is a series of factors reflecting zone, month and interaction between zone and month.

All covariate coefficients were found to be significantly different from zero at \( p=0.001 \) level. The estimated parameters and standard errors for the climate variables included in the final model are listed in Table 1. Precipitation and temperature with time lags of 1 and 2 months were found to be the most statistically significant and are positively related to dengue. This may be the result of warm/humid conditions promoting mosquito development and rain water filling discarded containers outdoors to create mosquito breeding sites. Therefore, an epidemic could be more likely if the temperature and/or precipitation in the preceding months are above average. The Niño 3.4 index is negatively related to dengue. This is because the major dengue epidemics in 2002 and 2008, in particular, were preceded by negative SST anomalies in the Niño 3.4 region. The scale parameter \( \theta \) was estimated to be 0.32 with standard error 0.002, confirming a mean variance relationship considerably different from that of the Poisson (equal mean and variance), hence justifying the use of a negative binomial rather than a Poisson GLM for model selection purposes.

One important aspect of such a model to a public health decision maker is its ability to predict dengue during the peak dengue season from February–April (FMA). In Fig. 5, scatter plots with fitted loess curves show the relationship between observed and predicted dengue using the GLM model for the FMA season 2001–2008 for Brazil (Fig. 5a) and the South East region where dengue predominates (Fig. 5b). Although the model clearly fails to capture much of the variability in dengue counts in this season, there is an overall positive association between observed and predicted counts at both the national and regional level. The influence of the climate variables in the model predictions is demonstrated in Fig. 6a, which shows the time series of total observed dengue cases for the FMA season, predicted dengue using a GLM without any climate information (dotted line) and with climate information (dashed line). The climate variables are the only source of temporal information in the model. Therefore, by not including them the same prediction is produced for every month/season of each year. By including climate information, some of the temporal variability is captured albeit with limited skill. Fig. 6c illustrates how the GLM predicts dengue for the FMA season in 2008. In some areas, the predicted dengue level corresponds to the observed level, for example, in coastal margins of the South East region (see Fig. 6b). However, low levels of dengue are overestimated in the South and the model fails to reproduce the variability in dengue cases across the Amazon. When we focus at the region level (South East) and microregion level (Río de Janeiro) for which dengue early warnings would be most useful, time series of dengue for the FMA season 2001–2008 show that the climate informed GLM fails to reproduce the dengue epidemic in 2002 and the increase in dengue from 2006–2007 (Fig. 7a and b).

This GLM clearly fails to capture much of the temporal variability in dengue counts, which may be attributable to factors such as population immunity to the dominant circulating serotype or specific health interventions and vector control measures. However, information regarding these aspects of the disease system are not readily available. Therefore, the use of unstructured random effects may be valuable to allow for unobserved latent structures in the model (McCulloch and Searle, 2001), for example, to capture the impact of unknown/unobserved confounding factors, such as the introduction of a new dengue serotype in a certain area of Brazil. Also, by using a GLM independence is assumed in both time and space and neither of these assumptions may be valid. There could be strong temporal correlation effects within some areas and there could also be spatial clustering effects in neighbouring microregions. To allow for such latent effects and correlation structures, the GLM is refined in the next section by reverting to a Poisson framework but using a generalised linear mixed model (GLMM) which includes spatially unstructured and spatio-temporal structured random effects in the linear predictor. This explicitly models the extra-Poisson variation or overdispersion previously allowed for using the negative binomial.

| Observed climate | Coefficient estimate | Standard error | Prob > |z| |
|------------------|----------------------|----------------|--------|
| Precipitation lag 1 | 0.018 | 0.0037 | 5.12 \times 10^{-4} |
| Precipitation lag 2 | 0.022 | 0.0036 | 6.45 \times 10^{-11} |
| Temperature lag 1 | 0.091 | 0.0093 | 2 \times 10^{-16} |
| Temperature lag 2 | 0.161 | 0.0093 | 2 \times 10^{-16} |
| Niño 3.4 lag 6 | -0.204 | 0.0119 | 2 \times 10^{-16} |
We focus our analysis on the South East region of Brazil (see Fig. 1a) where dengue is most prevalent and there are a large number of densely populated urban centres, which could benefit from a climate informed dengue EWS. This is also the region where the previously reported GLM predictions did appear to capture some of the observed spatial variability in dengue counts (see Fig. 6c).

4. Development of a generalised linear mixed model

As described above, we now focus on the 160 microregions in South East Brazil and return to a Poisson model for the dengue count data to develop a GLMM that includes random effects in the linear predictor. One approach to fitting such a model is to use a Bayesian framework. Markov Chain Monte Carlo (MCMC) methods make Bayesian modelling of complex situations involving many parameters a practical feasibility (see Gilks et al., 1996; Brooks, 1998 for more details). One further advantage of the Bayesian approach is that the associated MCMC sampling yields full posterior predictive distributions which automatically incorporate all components of variance at the different levels in the model. A full assessment of prediction uncertainty is therefore more easily obtained with Bayesian estimation than with the more traditional maximum likelihood approach.

The inclusion of random effects introduces an extra source of variability (a latent effect) into the model to capture the impact of unknown/unobserved confounding factors. For example, serotype introduction, which can vary spatially and temporally. Unstructured random effects can help account for overdispersion in the distribution of dengue counts $y_i$. However, this does not allow for explicit spatial dependence between $y_i$. This dependence can be included by adding a spatially structured random effect. A typical choice for a spatially structured prior is a conditional intrinsic Gaussian autoregressive model (CAR) (see Besag et al., 1995):

$$y_i \mid y \sim N \left( \sum_{j \neq i} a_{ij} y_j, \sigma^2 \sum_{j \neq i} a_{ij} \right).$$

where $a_{ij}$ are adjacency weights for the microregions, here taken to be simple binary values: $a_{ij} = 1$ if microregion $i$ has a common

**Fig. 5.** Scatter plot and loess curve (solid line) to show observed and predicted dengue fever, using GLM model for 3 month season FMA 2001–2008 for (a) Brazil and (b) South East region.

**Fig. 6.** (a) Time series of total observed dengue (solid line), GLM predicted dengue with climate (dashed line) and GLM predicted dengue without climate (dotted line) for FMA season 2001–2008 in Brazil, maps to show sum of (b) observed and (c) predicted dengue cases for microregions of Brazil, FMA season 2008. Categories defined by quintiles of observed dengue for FMA season 2008.
boundary with microregion $j$, $a_{ij} = 0$ otherwise. The hyperparameter $\sigma_j$ controls the strength of the local spatial dependence. As the CAR is improper, a 'sum to zero' constraint is applied to $v_i$ and it is then advisable to take a uniform flat prior for the intercept $\alpha$ (see model specification below).

Models to predict vector-borne disease may include an autoregressive time series component (e.g. Gomez-Elipe et al., 2007), based on the idea that the current value of the time series $y_{it}$ can be explained as a function of past values. Accordingly, a first order autoregressive temporal effect $\omega_t$ was included in the model, where $t$ is calendar month and $\omega_1$ (August) is set equal to zero in the model specification to avoid identifiability problems.

Therefore, the spatio-temporal GLMM adopted is given by:

$$y_{it} \sim \text{Pois}(\mu_{it})$$

$$\log(\mu_{it}) = \log(e_{it}) + \alpha + \sum_j \beta_j x_{ijt} + \sum_j \gamma_j w_{jt} + \phi_i + v_i + \omega_t$$

$$\alpha \sim U(-\infty, +\infty)$$

$$\phi_i \sim \text{N}(0, \sigma_{\phi}^2)$$

$$v_i \sim \text{CAR}(\sigma_v^2)$$

$$\omega_0 = 0, \omega_t \sim \text{N}(0, \sigma_{\omega}^2), \quad t = 2, \ldots, 12.$$

Independent diffuse Gaussian priors (mean 0, precision $1 \times 10^{-6}$) were taken for $\beta_j$ ($j = 1, \ldots, 5$) and $\gamma_j$ ($j = 1, 2$), whilst independent gamma hyperpriors with equal shape and inverse scale parameter (0.01) were used for the precisions ($\omega_0 = 1/\sigma_{\phi}^2, \omega_1 = 1/\sigma_v^2, \omega_0 = 1/\sigma_{\omega}^2$) of the priors for the spatially unstructured $\phi_i$ and spatially structured $v_i$ random effects, ($i = 1, \ldots, 160$), and the temporally autocorrelated random effects $\omega_t$ ($t = 2, \ldots, 12$).

This model was fitted with WinBUGS software (Lunn et al., 2000), using two parallel MCMC chains, each of length 25,000 with a burn-in of 20,000 and thinning of 10 to obtain samples from $P(\alpha, \beta, \gamma, \phi, v, o, \omega_0, \omega_1, \omega_2 | y)$. As mentioned earlier, the explanatory variables $x_{ijt}$ ($j = 1, \ldots, 5$) and $w_{jt}$ ($j = 1, 2$) are as before. However, all covariates are now standardised to zero mean and unit variance to aid convergence. This model is fit at the region level, therefore, the zone factor is omitted as there is little variation in zone type in the South East region, and any geographic differences between microregions are captured by the spatial random effects. Satisfactory convergence was confirmed using a range of standard criteria (Gelman et al., 2004). Posterior distributions for each parameter associated with the climate covariates in the model are given in Fig. 8 (with posterior means in parentheses). The climate parameters are all significantly different from zero and the sign of the association between dengue and each climate variable is consistent with the results from the GLM fit (Table 1).

A posterior predictive distribution can be obtained for each microregion by drawing random samples from a Poisson distribution with mean equal to the MCMC samples from the model fit. The mean of the posterior predictive distribution for all microregions in the South-East region were obtained for the peak dengue season FMA. In Fig. 9 a scatter plot with fitted loess curve shows the relationship between observed and predicted dengue using the GLMM model for the FMA season 2001–2008. When compared to predicted values from the GLM (see Fig. 5b), the loess curve has shifted towards the 45° line and more of the variability in dengue cases has been captured by the prediction model. Fig. 10 illustrates the spatial distribution of observed (Fig. 10a) and predicted dengue for FMA season 2008 using both the GLM (Fig. 10b) and GLMM (Fig. 10c). While the GLM predicted medium levels of dengue across much of the region, the GLMM captures more of the observed variation. When compared with Fig. 7, Fig. 11 shows how the addition of random effects to the model has improved dengue predictions for both the South East as a whole (Fig. 11a) and for the microregion Rio de Janeiro (Fig. 11b), particularly for the 2008 epidemic.

5. Probabilistic epidemic prediction

The specified Bayesian hierarchical model can also be used to predict the probability of dengue exceeding a pre-defined epidemic threshold in each microregion. As the GLMM here provides a posterior predictive distribution for each microregion (rather than a point estimate), the probability of exceeding an epidemic threshold can be calculated and the decision to trigger an alert can be based on the probability of exceeding the

threshold being greater than a specified alert level, (e.g. probability of exceedance $Z_{90\%}$). Many epidemic detection algorithms have been investigated to detect epidemics (Cullen et al., 1984; Hay et al., 2002; Teklehaimanot et al., 2004). As an example, we consider the event of dengue exceeding an epidemic threshold of the mean plus one standard deviation for each microregion in South East Brazil in FMA 2008. The epidemic threshold is based on the dengue counts in the FMA season for the previous seven years (FMA 2001–2007). We can assess the ability of the GLMM to predict ‘dengue epidemics’ across South East Brazil during the FMA season in 2008 using a contingency table (see Table 2). Observed dengue counts for the 3-month season were compared with model predictions where the probability of an epidemic exceeded an alert threshold chosen to be 90%.

The contingency table provides information on the overall predictive skill of the warning system. The proportion correct, defined as the proportion of the 160 microregions for which the prediction correctly anticipated the subsequent epidemic or non-epidemic ($a + d / a + b + c + d$), was 83%. The hit rate (the proportion of epidemics that were correctly predicted $a / a + c$) was 60%. Conversely, the false alarm rate (the proportion of epidemics that were predicted but did not occur $b / b + d$) was 8%.

Fig. 12 shows the posterior predictive distribution for the FMA season 2008 for the microregion Linhares, found on the coastal region of Espírito Santo. Here, the probability of exceeding the epidemic threshold was found to be $Z_{90\%}$, based on the epidemic threshold of mean plus one standard deviation derived from the distribution of dengue for the season FMA 2001–2007. A successful epidemic alert would have been issued for this microregion using the GLMM with the given epidemic threshold and alert level. By lowering the alert level below 90% the hit rate for the region increases but so does the false alarm rate. In practice, the choice of epidemic threshold and alert level should be selected by decision makers based on expert opinion and available resources.

6. Discussion and conclusion

The preliminary modelling results in this paper indicate that climatic covariates play a statistically significant role in the transmission of dengue fever. Although climate information alone does not account for a large proportion of the overall variation in dengue cases in Brazil, spatio-temporal climate information with the addition of spatio-temporal random effects do account for some of this variability, particularly for the 2008 peak dengue season, when a serious epidemic occurred. Therefore, the inclusion of seasonal climate forecasts in a dengue EWS for Brazil is worth investigating. The next step would be to assess the predictive validity of the model when replacing ‘observed’ with ‘hindcast’ (i.e. retrospective forecasts made for a historical period in pseudo-operational mode) climate variables. ‘Hindcast’ precipitation, temperature and Niño 3.4 data are available from forecasting systems such as the UK Met Office seasonal forecasting system (Graham et al., 2005) and the European Centre for Medium Range Forecasts (ECMWF) System 3 (Anderson et al., 2007). These systems typically produce ensemble predictions with lead times up to 6 months. By replacing ‘observed’ with ‘hindcast’ climate variables in the above GLMM, a dengue prediction could be made 5 months ahead of the dengue season of interest. For example, to predict dengue incidence for March 2010, the model could be run in October 2009 using the observed Niño 3.4 index for September 2009 (6 month lag), and precipitation and temperature forecasts for January and February 2010 issued in October 2009. The reliability of a climate-based EWS will depend on the skill of the forecasting system or multi-model

Table 2

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<th>Predicted probability ≥ 90%</th>
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combined and calibrated system such as EUROBRISA (Coelho et al., 2006), in predicting seasonal climate conditions for the region of interest.

Previous sections have highlighted the potential for incorporating climate information into a spatio-temporal EWS for dengue in Brazil. However, before implementing such an operational system several technical issues need to be considered. For example, the definition of epidemic thresholds by public health decision makers. Thresholds should be designed to minimise false alarms and false negatives (i.e. failing to predict that an epidemic will occur) and should correspond with the epidemic response capabilities in specific locations. The spatial scale of the system affects the type of response activity that could be implemented. For example, at the microregion level interventions such as health care provisions may be possible. However, vector-control efforts may be more difficult to target. Predictive output from an EWS needs to be continuously monitored and evaluated over time and models should be refigured as new dengue/climate data becomes available.

Spatial demographic data from the census (and interim projections) should also be updated when necessary. In order to issue the most reliable epidemic predictions forecast climate should be replaced with observed climate as time progresses towards the peak epidemic season, so that updated epidemic alerts can be reissued to public health decision makers. However, time delays in obtaining and collating real-time information for both confirmed dengue cases from SINAN and climate forecasts and observations could hinder the ability to provide warnings far enough in advance. Another important consideration is the dissemination and visualisation of early warnings of increased level of dengue risk. It is vital to train public health decision makers on how to interpret and use dengue risk forecasts, including awareness about forecast limitations to avoid misinterpretation and/or over interpretation.

Developing a climate-based EWS for dengue using climate and disease information over such short time periods remains a major challenge. During the study period, the Niño 3.4 index strongly influences the temporal signal of predicted dengue. From June 2007, a moderate La Niña event developed, which strengthened in early 2008. Was the dengue epidemic in 2008 influenced by this La Niña event or was this a coincidence? ENSO may play a role in synchronizing epidemics, however, periods between epidemics may also be a function of herd immunity from previous epidemics, and these two cycles (ENSO and herd immunity) may have coincided during the 2001–2008 study period. Further investigation is needed to understand temperature and precipitation patterns associated with warm phase and cold phase ENSO for this region in Brazil and to consider the possibility of a non-linear relationship between precipitation/temperature and dengue. The model parameterisation would benefit from the inclusion of one or more past epidemics to address these problems.

Despite this, it is hoped that this spatio-temporal dengue prediction model is a step towards the development of a useful decision making tool for the Brazilian health services. Such spatio-temporal models offer an opportunity to balance global climate variables and local responses, e.g. the influence of ENSO on dengue incidence is likely to occur unequally across the region due to particular socio-economic local conditions. Another advantage of the GLMM is the ability to address specific public health issues in terms of probabilities. This model could be extended to other regions in the world where climate-sensitive infectious diseases (e.g. cholera, malaria, leptospirosis, plague) present a burden to public health infrastructure, particularly in developing countries.

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