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Review Article

Sources of spatial animal and human health data: Casting the net wide to deal more effectively with increasingly complex disease problems

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ABSTRACT

During the last 30 years it has become commonplace for epidemiological studies to collect locational attributes of disease data. Although this advancement was driven largely by the introduction of handheld global positioning systems (GPS), and more recently, smartphones and tablets with built-in GPS, the collection of georeferenced disease data has moved beyond the use of handheld GPS devices and there now exist numerous sources of crowdsourced georeferenced disease data such as that available from georeferencing of Google search queries or Twitter messages. In addition, cartography has moved beyond the realm of professionals to crowdsourced mapping projects that play a crucial role in disease control and surveillance of outbreaks such as the 2014 West Africa Ebola epidemic. This paper provides a comprehensive review of a range of innovative sources of spatial animal and human health data including data warehouses, mHealth, Google Earth, volunteered geographic information and mining of internet-based big data sources such as Google and Twitter. We discuss the advantages, limitations and applications of each, and highlight studies where they have been used effectively.

1. Introduction

Over the last 30 years it has become commonplace for epidemiological studies or surveys to collect locational (spatial) attributes for disease data (Pfeiffer et al., 2008). Although this advancement has been driven largely by the introduction of handheld global positioning systems (GPS), and more recently, smartphones and tablet computers with built-in GPS that facilitate geo-tagged data collection, it also highlights the increased awareness of the importance of the spatial aspect when developing efficacious animal disease surveillance and control strategies (Table 1). Unfortunately, as a result of the particular challenges currently facing health workers and researchers, for spatial disease data to be able to effectively inform innovative surveillance and disease control strategies, it needs to move beyond the fundamentals of collecting georeferenced disease event data in individual studies and instead focus on an inclusive approach that Eysenbach (2001), in his definition of eHealth, referred to as ‘a state-of-mind, a way of thinking, an attitude, and a commitment for networked, global thinking, to improve health care locally, regionally, and worldwide by using information and communication technology’.

This collective, crowdsourced approach was aptly illustrated during the 2014 West Africa Ebola crisis when, faced with only a few rudimentary topographical maps of Guinea but no useful maps upon which to base control and surveillance efforts, Médecins Sans Frontières (MSF) personnel...
Using spatial analysis to inform risk-based animal disease surveillance and control.

**Table 1**

| Mapping disease distribution | Disease distribution maps range from simple dot maps showing the location of disease events to predictive risk maps created using statistical algorithms that combine disease occurrence data with environmental covariates (Pigott et al., 2014). But no matter what form they take, visualizing the spatial pattern of disease – be it at a global, national or local scale – is fundamental for informing risk-based disease surveillance and control strategies in several ways. Simple visualizations allow the extent of the disease to be delineated and disease frequency monitored, and when combined with maps of environmental factors or those highlighting the spatially heterogeneous distribution of at-risk populations, they can also be used to estimate disease burden (Hay et al., 2010; Robinson et al., 2002) and identify target populations for intervention (Temborn et al., 2011; Guerra et al., 2010, 2008, 2006). Visualizing disease distribution can also be fundamental in directing control and elimination efforts. Clements et al. (2013) describe how measures to eliminate malaria from endemic countries have generally adopted a spatially progressive elimination approach referred to as shrinking the malaria map in which eradication efforts initially focus on the geographical perimeter of endemic areas and work inwards, effectively localizing disease distribution which allows for more efficient treatment and control (Feachem et al., 2010). Apart from the key role maps play in informing risk-led decision making, they also serve a more practical purpose such as facilitating integration and synthesis of data from a wide range of diverse sources, each possibly capturing information about disease and relevant risk factors at different scales (Bergquist and Tanner, 2012; Bennema et al., 2014). As a result, cartographers need to decide on the most appropriate scale at which to present the data for it to be useful; data presented at administrative level 1 (province or region) inevitably cannot capture the fine-scale heterogeneity of most infection patterns and so estimates of numbers of individuals requiring treatment tend to be incorrect (Brooker et al., 2010).

Cluster detection

A clustered spatial arrangement of disease events suggests the presence of a contagious process or localised risk factor. Apart from the fact that spatial targeting of interventions at high-risk areas is more cost-effective than uniform resource allocation (Stark et al., 2006) and therefore such identification is essential for informing risk-based disease surveillance and control efforts. Identification of significant disease clusters can also advance our understanding of a disease in several ways including suggesting potential risk factors for further investigation either directly (Calistri et al., 2013; French et al., 2005; Sinkala et al., 2014; Kelen et al., 2012; Ngareda et al., 2015; Poljak et al., 2007; Le et al., 2012; Vigie et al., 2005; Ward and Carpenter, 2000), or indirectly when analysis of model residuals indicate the models’ predictors do not explain fully the spatial heterogeneity in disease distribution (Méroc et al., 2014; Borba et al., 2013), or by defining the scale of disease clustering (French et al., 2005; Le et al., 2012; French et al., 1999; Wilesmith et al., 2003; Picado et al., 2007; Picado et al., 2011; Porphyre et al., 2007; Sanchez et al., 2005; Minh et al., 2009; Minh et al., 2010; Xu et al., 2012; Métras et al., 2012; Abatih and Ersbøll, 2009) and thereby indicate likely transmission mechanisms involved in disease spread (Sinkala et al., 2014; Ward et al., 2013; Looiboyck et al., 2009; Olhson et al., 2014; Rosendal et al., 2014; Poljak et al., 2010). Cluster detection can also be used identify areas where vectors and hosts coincide resulting in potentially increased risk of disease transmission (Shaman, 2007; Hennebelle et al., 2013; Swirski et al., 2007), highlight possible regional differences in disease spread (Kelen et al., 2012), or track the direction and geographical extent of disease spread (Wilesmith et al., 2003; Denzin et al., 2013; Lian et al., 2007).

Spatial modelling

Spatial modelling techniques can be divided into data- and knowledge-driven methods (Stevens and Pfeiffer, 2011), the former characterised by the use of statistical methods for defining relationships between risk factors and disease risk, while knowledge-driven modelling approaches are based on existing knowledge about the causal relationships associated with the disease risk of interest. Statistical analysis is used to generate data-driven models from information collected through surveillance and other means. Such models generate quantitative estimates of risk and the relative weights of risk factors. The results of such models are used for a variety of purposes including targeting areas for disease surveillance, risk management, simulating different control scenarios, or predicting what will happen under different environmental conditions such as those resulting from climate change (i.e. temporal prediction), or identifying new geographical areas suitable for the introduction of diseases (i.e. spatial prediction).

enlisted the help of the Humanitarian OpenStreetMap Team (HOT) to map Guéckédou - the main city in Guinea affected by the outbreak (Hodson, 2014). Within 20 h of receiving the request, online volunteers had mapped three cities in Guinea based on satellite imagery of the area, populating them with over 100,000 buildings; information that proved crucial for door-to-door canvassing of inhabitants and mapping the spread of disease.

In addition to this collective approach, for spatial disease data to be effective in the 21st century, it needs to meet certain requirements. Firstly, the increasing number of transboundary disease epidemics has emphasized the need for animal and human health information systems that are no longer circumscribed by regional or national borders; transparent collection and sharing of disease data needs to occur at a global scale. Secondly, globalization has substantially increased the speed and magnitude of disease spread. In the 2001 UK foot and mouth disease (FMD) outbreak it was estimated that at least 57 premises from 16 counties were infected before the first case was reported (Gibbens and Wilesmith, 2002) while in 2007, equine influenza spread rapidly throughout two Australian states as a result of infected horses attending an equestrian event (Cowled et al., 2009); approximately 70,000 horses on over 9000 premises were infected with most of the geographic dissemination occurring within the first ten days of the epidemic. For containment to be effective, reporting of disease events needs to be as rapid as possible. This is of particular concern in developing countries where reporting of animal disease events can be delayed by months (Karimuribo et al., 2012) while lag times for such reports as the Centers for Disease Control and Prevention (CDC) US Influenza Sentinel Provider Surveillance reports are currently in the order of 1–2 weeks (Ginsberg et al., 2009).

During the past decade, collecting spatial disease data has moved beyond the use of handheld GPS devices and there now exist numerous sources of crowdsourced georeferenced disease data such as that available from georeferencing Google search queries or Twitter messages. Not surprisingly, the focus so far has been on human health,
driven to some extent by US President Barack Obama's Global Health Initiative (Initiative GH, 2009) and the World Health Organisation's (WHO) concerted efforts regarding neglected tropical diseases (Cringoli et al., 2013; The Global Network for Neglected Tropical Diseases; Brooker and Utzinger, 2007; van Lieshout and Yazdanbakhsh, 2013; Malone and Bergquist, 2012; Brooker and Smith, 2013; King et al., 2013; Brooker et al., 2006). Also, by their very nature, citizen and internet-based health approaches lend themselves to human rather than animal-health problems and similarly, to developed countries that have the necessary internet infrastructure, rather than developing nations. As a result, internet-based animal-health initiatives currently lag behind those of human health, yet it is to these initiatives that we must look to see how such systems can be best adapted for use in animal health and in developing nations; countries that carry the highest burden of emerging and zoonotic infectious diseases in the world yet frequently have the least capacity for cost-effective risk-led decision making.

This paper reviews a range of sources and features of spatial disease data currently available, discussing their advantages and limitations, and highlighting studies where they have been used. Although the focus is on animal diseases, relevant advancements in human health that could be adopted for animal health purposes are discussed.

2. Sourcing spatial disease data

2.1. Data warehouses

Although animal disease surveillance has traditionally been implemented at national and sub-national levels, the increasing number of transboundary animal disease epidemics has highlighted the need for establishing such systems at broader scales. As a result, data warehouses and disease reporting systems such as World Animal Health Information Database (WAHID) (Jebra et al., 2012) and EMPRES Global Animal Disease Information System (EMPRES-i) (FAO: EMPRES transboundary animal disease bulletin, 2011; Farnsworth et al., 2010; Martin et al., 2007) were launched to encourage and facilitate data collection and sharing at a global level (Table 2). However, in addition to their original role, such data warehouses also provide researchers with cost-effective access to regularly updated spatial disease data, potentially leading to increased knowledge gains, without the need for costly and time-consuming primary research. Moreover, integration of databases from different sources offers researchers a more extensive and comprehensive collection of information than if individual data sources were used with the possibility of better understanding issues at the population level. However, researchers using this data need to remember that although the provenance of national disease surveillance data ensures that specificity is reasonably high, sensitivity is likely to be low and undoubtedly exhibits considerable spatio-temporal heterogeneity with respect to bias and sensitivity (Perez et al., 2011). Users should also bear in mind the limitations of using data that have generally been spatially referenced to administrative centroids rather than exact outbreak locations; in addition to the possibility of ecological fallacy, Stevens et al. (2013) showed that using outbreak data georeferenced to administrative centroids for spatial modelling purposes can be problematic when either constraining the study area or working at relatively low spatial resolutions.

| Data warehouse (URL) | Description |
|----------------------|-------------|
| Disease BioPortal; (http://bioportal.ucdavis.edu) (Perez et al., 2011) | Provides real-time or near real-time access to local, regional and global disease information and data for more than 40 animal diseases and syndromes. Set of techniques for cluster detection and phylogenetic analysis of sequences is available for the user |
| EMPRES Global Animal Disease Information System (EMPRES-i); (http://empres-i.fao.org/eipws3g/) (FAO: EMPRES transboundary animal disease bulletin, 2011; Farnsworth et al., 2010; Martin et al., 2007) | EMPRES-i provides up-to-date information on global animal disease distribution and current threats at national, regional and global level. Disease events can be presented on a map and data may also be exported for further analysis |
| EMPRES-i genetic module (Claes et al., 2014) | EMPRES-i genetic module application combines epidemiological outbreak information (EMPRES-i) with genetic characteristics of influenza viruses (OpenFluDB) |
| FAO GeoNetwork (http://www.fao.org/geonetwork/srv/en/main.home) | FAO GeoNetwork provides access to interactive and downloadable maps, satellite imagery and related spatial databases maintained by the Food and Agricultural Organization of the United Nations (FAO) and its partners |
| Global Livestock Production and Health Atlas (GLiPHA); (http://kids.fao.org/glipha/); (http://kids.fao.org/glipha/); (Clements et al., 2002; Franceschini et al., 2009) | GLiPHA is an interactive, electronic atlas containing global animal production and health statistics. Sub-national statistics relating to the livestock sector can be viewed cartographically, against a backdrop of selected maps such as livestock densities, land-use and topography. Data may either be displayed or exported as tables and charts |
| World Animal Health Information Database (WAHID); (http://www.oie.int/wahis_2/public/wahid.php/WahidHome/Home) (Jebra et al., 2012) | WAHID also provides access to all data held within OIE's World Animal Health Information System (WAHIS). Together with global disease distribution and outbreak maps, WAHID also includes country-level information on exceptional disease events and animal health status together with country-level maps of the prophylactic and control measures in use |
Table 2 contains details of the main animal disease geodata warehouses. EMPRES-i data is obtained from both formal (e.g. reports from the OIE, World Health Organization, national authorities, FAO country or regional projects, field missions and field officers, non-governmental organizations, laboratories and reference centers) and informal sources (e.g. media reports and those disseminated by the Global Public Health Intelligence Network and ProMED) (Farnsworth et al., 2010; Martin et al., 2007), and all outbreaks appearing in the database are followed-up until either confirmed or denied (Farnsworth et al., 2010). WAHID, on the other hand, comprises official information submitted by OIE member countries regarding immediate and follow-up notifications of exceptional disease events, or semester and annual reports on OIE-listed diseases together with background information on animal health and control programs. Alternatively, Disease BioPortal is an unrestricted, public web site created and maintained by the Center for Animal Disease Modeling and Surveillance (CADMS) at the University of California, Davis (Disease BioPortal) incorporating multiple streams of information including WAHID (Jebara et al., 2012), EMPRES-i (FAO: EMPRES transboundary animal disease bulletin, 2011; Farnsworth et al., 2010; Martin et al., 2007), GenBank, the World Reference Laboratory for Foot-and-Mouth Disease (WRLFMD), weekly reports on vesicular-like diseases from Centro Panamericano de Fiebre Aftosa (PANASFOSA) and the Foot-and-Mouth disease (FMD) News and Rift Valley fever (RVF) News produced by CADMS.

In additional to global information systems, it is also useful for decision makers to have access to national data warehouses. Denmark was the first country to make all animal husbandry data from a range of sources and departments related to Danish production animals available in a single online geodata warehouse (Nielsen, 2011). Disparate databases can be linked at the individual animal level or aggregated at farm, postcode or administrative level. However, the difficulties associated with standardizing and combining a range of data sources – each involving potentially dissimilar unique identifiers, data structures, languages and semantics – limits the development of animal health geodata warehouses and results in bias. Data adapters (applications that convert attributes of one database into attributes compatible with another database) are integral to the creation of such information systems as they can be used to relate diverse databases through the identification of fields containing equivalent information (Perez et al., 2011).

Data quality is of primary concern with data warehouses and assessing the validity of the data is paramount. GLiPHA, which merges livestock health and production data from multiple sources, incorporates a system of error checking rules to help identify inaccuracies and inconsistencies in the data (Clements et al., 2002) while the Danish system uses a number of different tools such as continuous reporting of data, systematic control of entries and irregularities and cross-control of data with other sources (Nielsen, 2011).

Although the logistics involved in linking a range of disparate datasets partly hinders the development of national or global animal health data warehouses, the reluctance of certain agencies and organisations to share disease data freely, transparently and in a timely fashion (Perez et al., 2011) is an additional impediment to the development of a valuable resource that should be integral to 21st century risk-led disease management and decision making. Buy-in from all parties is therefore essential if such information systems are to be successfully established.

2.2. mHealth

While developed nations generally have access to a wide range of good quality georeferenced health-related data, collection, processing and dissemination of such data in resource-poor locations remains challenging owing to lack of the necessary technological infrastructure as well as issues with familiarity and usability (Betjeman et al., 2013). In such countries data is generally still collected via paper-based forms despite associated inefficiencies such as data entry errors and long delays before the processed data is available to decision makers (Anokwa et al., 2009). In fact, a survey of selected human and animal health surveillance systems in different regions of Southern Africa found obvious spatial heterogeneity in the delivery of monthly animal health reports to the central epidemiological department; wards closer to the headquarters submitted reports more regularly than those further away while delivery of all reports could be delayed by as much as six to nine months (Karimuribo et al., 2012).

There is thus an urgent need for resource-poor settings to implement alternate surveillance systems and although lack of technological resources and infrastructure may preclude the use of novel internet-based surveillance approaches, mobile devices such as the now out-of-date personal digital assistants (PDAs) Shirima et al., 2007; Yu et al., 2009; Seebregts et al., 2009; Dale and Hagen, 2007 and more recently mobile phones (Robertson et al., 2010; Jean-Richard et al., 2014; Thinyane et al., 2010), smartphones (Forsell et al., 2011) and tablet computers, are playing an increasingly fundamental role in the collection and processing of animal and human health surveillance data in resource-poor locations (Betjeman et al., 2013; Chretien et al., 2008; Mwabukusi et al., 2014; Istepanian et al., 2004). This is, in part, a result of the extensive penetration of mobile phone use in developing countries over the last decade; estimated to be 63% in sub-Saharan Africa in 2013 and projected to pass 70% by 2015 (Betjeman et al., 2013).

Initially restricted to simple yet effective features such as short messaging service (SMS) and voice calling, the value of mHealth – using mobile devices to collect or distribute health-related information (Istepanian et al., 2004) – for health care workers in developing countries increased immeasurably following the development of smartphones. In addition to the variety of data that can be collected – text, audio, video, photographs and barcode scans – other key benefits of smartphones over non-internet based models include the built-in GPS and accelerometer which allow detailed locational data and changes in movement to be documented (Anokwa et al., 2009). Images, in the form of photographs and videos,
can also be sent which may allow remote diagnoses based on gross pathology.

Furthermore, a range of open-source mobile software and tools such as EpiCollect (Aanensen et al., 2009; EpiCollect) and Open Data Kit (ODK) (Anokwa et al., 2009; Open Data Kit) allows for cheap, efficient and accurate collection and dissemination of data. Both EpiCollect and ODK allow for the generation of mobile-based forms and for a range of data to be collected and stored on the mobile device before being wirelessly transmitted to a central database. Together with a map-based interface, such as that located within spatiotemporal.epidemiology.net (Anokwa et al., 2009), data can be rapidly analysed, mapped and filtered using Google Maps™ via both the web and mobile applications so that both office and field workers can display and analyse data in real-time (Anokwa et al., 2009; Aanensen et al., 2009). Real-time reporting and processing of disease data, followed by rapid transmission of information to decision makers, allows swift action to be taken against possible outbreaks. For example, use of real-time reporting and summarising of surveillance data via mobile devices allowed a potential FMD outbreak in the Ngara district of Tanzania to be rapidly contained (Mwabukusi et al., 2014).

Additional benefits of mobile-based forms include the ability to customize voice-based questionnaires programmed into a smartphone or tablet computer which allows questions to be administered in local languages or dialects (Jantee et al., 2014) while programming the questionnaire to ask questions in sequence significantly reduces the risk of missing data. Similarly, the use of drop-down menus reduces the risk of data entry error (Jantee et al., 2014). Moreover, time spent completing smartphone-based questionnaire surveys is significantly less than when completing paper-based surveys (King et al., 2013; Anokwa et al., 2009; Jantee et al., 2014). However, as availability of open-source mobile software and tools increases – all differing with respect to accessibility, visualization and cost – choosing the most appropriate data-collection tool will depend largely on the type of data being collected (Madder et al., 2012).

Although smart devices have overtaken simple mobile phones restricted to sending voice calls or SMS messages, when combined with applications such as GeoChat, Ushahidi or RapidSMS, the technology can be highly effective. The Cambodian Ministry of Health uses GeoChat for disease reporting and to send staff alerts in response to potential outbreaks (Kamel Boulos et al., 2011). The real-time knowledge at the user end (Karimuribo et al., 2012; Mwabukusi et al., 2014).

Although developing counties are the ideal focus for mHealth data collection and disease control initiatives, there are few published examples of such enterprises, particularly in the field of animal health. Successful mHealth initiatives are more frequently documented in human (Betjeman et al., 2013; Dégélie et al., 2012; Shet and Costa, 2011; Lee et al., 2011; Lozano-Fuentes et al., 2013; Lozano et al., 2012) than in animal health, and in developed (Raja et al., 2014) rather than developing countries. However, practical applications of mobile device use in animal disease control and surveillance in developing countries have been described (Robertson et al., 2010; Jean-Richard et al., 2014; Thinyane et al., 2013) and include enhanced reporting of human rabies exposures at bite treatment centres in Tanzania where mobile phone technologies allowed for rapid communication between human and animal health sectors to ensure follow up of animal cases (Mtma, 2013) and the collection of demographic data and movement tracking of mobile pastoralists and their herds in Chad (Jean-Richard et al., 2014). The real-time knowledge on camp location and populations provided by such a study facilitates health interventions, such as vaccination delivery to both humans and animals, highlighting the potential to develop One Health mHealth approaches. Such an approach provides added value compared to separate animal-human surveillance systems, especially for zoonotic diseases.

However, as with any surveillance system, the success of such efforts depends largely on the extent of local buy-in at all levels, in particular those involved in feeding information back to frontline workers and community organisations; without an efficient two-way flow of information the benefits of mHealth will be limited to having made disease reporting more technologically advanced (Madon et al., 2014).

2.3. Google Earth™ and remote sensing (RS) data

Compared with previous decades when the production of paper-based disease atlases was limited by the expense and inefficiency associated with producing something that was effectively out of date almost before it was published, the advent of interactive digital maps and virtual globes such as Google Maps™ and Google Earth™ allows for easy visualisation of disease data in real time as illustrated by the integration of such digital platforms into an ever-expanding number of animal and human-health projects (Table 3). The value of such technology in creating effective information resources for decision makers is epitomised by Nature’s use of the platform to track the global spatio-temporal spread of highly pathogenic avian influenza (HPAI) H5N1 (Butler, 2006; Google Earth Avian Flu), a project that won the Association of Online Publishers (AOP) Use of a New Digital Platform Award in 2006. A list of all current projects using Google Earth™ and Google Maps™ can be found on the Google Earth Outreach™ website. However, Google Earth™ is not only a visualization
households stratified by distance to the hospital. Using a system software ArcGIS to generate a random sample of combined Google Earth™ and the geographic information.

proximity to a hospital influenced water quality perception and practices in Haiti, Wampler et al. (2013) combined Google Earth™ and the geographic information system software ArcGIS to generate a random sample of households stratified by distance to the hospital. Using a satellite image from EarthExplorer as a basemap in ArcGIS, concentric 1 km buffer zones were created around the hospital. The buffer polygons were then exported to Google Earth™ where the high-resolution imagery allowed individual households within each polygon to be accurately identified and mapped using Google Earth™ pushpins. These point locations were imported into ArcMap where latitude and longitude were added to the dataset and later uploaded into a handheld GPS which was used to locate the households in the field for conducting field surveys (Wampler et al., 2013).

However, remote sensing (RS) data is more often used to provide spatial risk-factor information, particularly for vector borne diseases such as Rift Valley fever (Lacaux et al., 2007; Linthicum et al., 1999; Martin et al., 2007; Pin-Diop et al., 2007; Sallam et al., 2013; Soti et al., 2013; Tourre et al., 2008, 2009; Vignolles et al., 2009, 2010), bluetongue (Klingseisen et al., 2013; Purse et al., 2004, 2013), eastern encephalomyelitis virus (Barrera et al., 2001; Freier, 1993) and African horse sickness (Capela et al., 2003), where the disease vectors are sensitive to changes in specific climatic and vegetation factors that can be captured usefully by satellite technology (Rinaldi et al., 2006; Saxena et al., 2009). To extend the usefulness of RS data for providing risk fact information, image processing software such as ImageJ, can be used to analyse the RS images. For example, to identify suitable refuges for mosquitoes during hot, dry conditions, ImageJ was used to analyse Google Earth™ satellite imagery and the number of plants, total amount of vegetation around a homestead and its percentage of the total area were calculated and related to households that had reported cases of malaria. In this way, ImageJ was used to analyse freely-available Google Earth™ images of malaria-endemic locations to identify potential risk factors associated with vegetation cover (Ricotta et al., 2014).

Table 3
A selection of internet-based animal and human-health projects using Google Maps or Google Earth™ to visualise disease data.

| Project/Organisation (URL) | Description |
|--------------------------|-------------|
| HealthMap and its mobile app Outbreaks Near Me; ([http://www.healthmap.org](http://www.healthmap.org)) | A global disease alert map which aggregates data from a wide range of sources to deliver real-time intelligence on a broad range of emerging infectious diseases. The app includes a participatory surveillance feature that allows users to report outbreaks not yet shown on the map and be credited for their contribution. |
| Predict; ([http://www.vetmed.ucdavis.edu/ohi/predict/index.cfm & http://healthmap.org/predict](http://www.vetmed.ucdavis.edu/ohi/predict/index.cfm & http://healthmap.org/predict)) | Focuses on detection and discovery of zoonotic diseases at the wildlife-human interface and through the HealthMap website provides a dynamic visual display of surveillance data. |
| Animal Disease Reporting System (TSN); ([http://www.fli.bund.de/en/startseite/institutes/institute-of-epidemiology/working-groups/tiereuschennachrichten-tns.html](http://www.fli.bund.de/en/startseite/institutes/institute-of-epidemiology/working-groups/tiereuschennachrichten-tns.html)) | An electronic system for the registration of notifiable and reportable animal diseases in Germany. Disease events can be visualized using Google Earth™ and Google Maps™. |
| CONTRAST ([Utzinger et al., 2013; Stensgaard et al., 2009](http://www.map.ox.ac.uk)) | A multi-disciplinary research platform aimed at investigating control of schistosomiasis. |
| The Malaria Atlas Project (MAP); ([http://www.map.ox.ac.uk](http://www.map.ox.ac.uk)) | MAP uses innovative methods to produce a comprehensive range of malaria maps and estimates to support effective planning of global malaria control at national and international levels. |
| Multi Locus Sequence Typing; ([Databases: http://www.mlst.net Maps: http://maps.mlst.net](http://www.mlst.net)) | Provides basic epidemiological and molecular typing data for a number of bacterial and fungal pathogens and maps the distribution of pathogen genotypes. |
Recently, RS imagery has been used to predict areas of highest risk for schistosomiasis infections in Kenya (http://www.bbc.co.uk/news/science-environment-31483629). Using satellite imagery, the locations of suitable waterbodies for the snail vector have been mapped and compared with satellite imagery of human distribution, to identify area of highest risk of infection.

2.4. Volunteered geographic information (VGI)

Volunteered geographic information (VGI) Goodchild, 2007; Goodchild and Li, 2012, also known as wikification of GIS by the masses (Kamel Boulou et al., 2011) and crowdsourced cartography, refers to ‘the harnessing of tools to create, assemble, and disseminate geographic data provided voluntarily by individuals’ (Goodchild, 2007). A well-known example of VGI is OpenStreetMap (OSM), an open, online, editable map of the world being created by volunteers using a combination of local knowledge, GPS tracks and aerial imagery. As an extension of the basic community mapping effort, volunteers from HOT (Geo-Wiki Project) travel the globe to create collaborative maps of densely packed slums or remote villages, that can be used by aid and development agencies when deciding what infrastructure to build or in the event of a humanitarian crisis. As mentioned in the introduction to this paper, HOT played an important role in the 2014 West Africa Ebola outbreak, rapidly mapping Guéckédou, a city of around 250,000 people in southern Guinea, thereby providing field workers with crucial information they needed to be able to assist with mapping the spatial distribution of the disease and planning and implementing control efforts (Hodson, 2014).

Other examples of crowdsourced cartography include Geo-Wiki (Geo-Wiki Project), a global network of volunteers working to improve the quality of global land-cover maps. The website allows volunteers to view land-cover maps in real time, with Google Earth™ as a backdrop, and to apply their local-area knowledge to determine whether or not the classification is correct, and to amend or update it if necessary. Within the main project are subsidiaries such as Risk Geo-Wiki, AusCover Geo-Wiki and Livestock Geo-Wiki, which also hosts the updated global livestock distribution maps (Robinson et al., 2014). The Global Geo-Referenced Field PhotoLibrary is a global repository of georeferenced field photos which allow land cover changes to be tracked over time and provide ground truthing for satellite imagery (Xiao et al., 2011).

Despite their usefulness in producing general maps, the area in which VGI has so far proved most valuable is that of crowdsourced disaster surveys where online volunteers work from satellite imagery to identify buildings which appear to be damaged or destroyed, and to create maps of the disaster area by which aid workers can navigate. This is an excellent example of ‘collective intelligence’ (Spielman, 2014), the premise being that, under the right circumstances, collections of individuals are smarter than even the smartest individuals in the group. Similarly, if the collectively generated end-product is better than the best individual contribution, then the aggregated is incredibly valuable (Spielman, 2014); one of the reasons why the most successful mapping projects often address a pressing need (e.g. Haiti post-earthquake or Ebola outbreak in Guinea) or concentrate on areas with poor geographic coverage (e.g. slums) (Spielman, 2014).

However, these disaster surveys highlighted an important limitation associated with using untrained volunteers as, although the maps they created proved to be invaluable, damage assessments were poor (Zastrow, 2014) with satellite judgements by HOT personnel corresponding with a later ground survey only 36% of the time (Zastrow, 2014). The question of VGI accuracy extends beyond that of disaster situations and is particularly important when deciding whether citizen scientists can provide information that is of high enough quality to be used in formal scientific investigations. See et al. (2013) found that when using satellite imagery to describe land cover and human impact on the Geo-Wiki website, although there was little difference between expert and non-expert responses when categorising degree of human impact, experts were better than non-experts at identifying a range of land-cover types (See et al., 2013). However, accuracy of VGI can be improved by providing targeted training materials for volunteers (e.g. providing volunteers with pre-disaster imagery against which to compare current images (Zastrow, 2014), guidance on what features to look for (Zastrow, 2014; See et al., 2013) and instituting a continual learning process by providing volunteers with feedback on their contributions (See et al., 2013).

Unsurprisingly, not all crowdsourced information is of equal quality; some data are of higher quality than others just as some contributors are consistently better than others (Haklay, 2010). Given that crowd sourced data are of varying quality, when aggregating such data one has to guard against regression to the mean. That is, a few highly accurate or highly credible contributions should not be degraded by being combined with many contributions of low quality, even if these exceptional contributions are outliers (Spielman, 2014). This is aggravated by the fact that participation in internet-based mapping systems is highly skewed with a few contributors accounting for a large proportion of contributions (Goodchild and Li, 2012; Sieber and Rahemtulla, 2010; Elwood et al., 2012). Without effective means of aggregation maps will either be shaped by the most active contributors, or map features will simply reflect the average of contributions.

OSM works by assembling all volunteer data into a patchwork map (Goodchild, 2007) which is in turn converted into a single map by aggregating the data using the following three review process: crowdsourced (other users check contributions), social (a set of elite users adjudicate problems) and geographic (features are validated based upon geographic context) (Goodchild and Li, 2012). OSM contributions are aggregated primarily through crowdsourced review following a last-in, first-out model; users see only the most recent edit. Although this form of aggregation relies heavily on trust and does not directly leverage prior contributions, it has been shown that more edits of an OSM feature generally leads to greater positional accuracy (Haklay, 2010). If conflicts arise, OSM uses the social review process whereby a set of elite users adjudicate problems.
As quality of VGI contributions vary, the addition of robust measures of quality would be useful to indicate the level of confidence associated with each piece of information, and although traditional statistical concepts of uncertainty and bias are hard to apply to VGI, other options are available. For example, See et al. (2013) found that when classifying land-cover, volunteer accuracy appeared to be higher when responses for a given location were more consistent and when the volunteers indicated higher confidence in their responses, suggesting that these additional pieces of information could be used to develop associated robust measures of quality (See et al., 2013). Additional possibilities include the application of Bayesian probability or Dempster-Shafer theory (Eastman, 2009) to provide a measure of confidence.

2.5. Internet-based epidemic intelligence

Identification of outbreaks at their earliest stages – followed by a rapid response – can substantially reduce the impact of epidemics yet surveillance capacity for such detection can be costly. The Internet however is revolutionizing how epidemic intelligence is gathered, particularly in developed countries, allowing us to detect disease outbreaks earlier than when using traditional surveillance approaches, with the added bonuses of reduced costs and increased reporting transparency. For obvious reasons these approaches have, so far, focused on important human diseases but there is potential for the development of similar tools for surveillance of key animal diseases.

2.5.1. Mining primary internet-based data sources: big data

A huge volume of real-time information about infectious disease outbreaks is to be found in various forms of internet-based data streams (Brownstein et al., 2008; Grein et al., 2000; Heymann and Rodier, 2001). Known as internet-based big data, the term refers only partly to the volume of data available for analysis and mainly to the fact that access to the almost limitless body of internet data – Google processes, on average, over 40,000 search queries every second (Google Search Statistics) - allows us to learn things that we could not when using smaller, limited datasets (Cukier and Mayer-Schoenberger, 2013). Big data are generally characterized by 3Vs: volume (relative magnitude of dataset), velocity (rate at which new data are generated) and variety (heterogenous structure of dataset [e.g. text, video, audio]) (Gandomi and Haider, 2015). In addition, big data are also characterized by their ability to convert many aspects of the world that have never previously been quantified, into valuable data that can be analysed; a process known as datafication (Cukier and Mayer-Schoenberger, 2013).

However, there are several limitations associated with big data, of which researchers accustomed to working with smaller, conventional datasets, need to be aware (Cukier and Mayer-Schoenberger, 2013). Firstly, when working with big data researchers need to accept that the data will not be pristine; however, working with vast quantities of data of slightly questionable quality invariably trumps using small amounts of very exact data. Secondly, big data requires a move from causation to correlation, so rather than trying to identify why something happens, big data allows us to search mammoth amounts of information about an event and anything associated with it, in order to identify patterns that might help predict future occurrences. In the words of Cukier and Mayer-Schoenberger (2013), ‘big data helps answer what, not why, and often that’s good enough’.

If mined using internet-based tools, these big data are often capable of detecting the first evidence of a disease outbreak. Such systems are based on the assumption that changes in information and communication patterns on the Internet can act as early warning of changes in population health (Wilson and Brownstein, 2009) and comprise automated biosecurity intelligence text-mining systems that continuously query, filter, integrate and visualise infectious disease data from myriad primary or secondary data sources. Two such sites that have received a lot of attention are Google and the social-media platform, Twitter.

The immediacy of Twitter offers health officials an enormous advantage as both a surveillance and research tool. For example, emergency departments in Boston learned about the 2013 marathon bombings through Twitter before announcements from conventional sources such as the media or established emergency service communication channels (Cassa et al., 2013). While terrorist attacks are an extreme case, the general principle also holds true for early warning of disease epidemics. Similarly, in addition to posting information about their health on social-media sites such as Twitter, data from search-queries have been found to be highly predictive of a wide range of population-level health events. For example, trends in Google and Yahoo search-queries have been used to predict influenza and dengue fever outbreaks (Chan et al., 2011) and estimate the prevalence of Lyme disease (Seifter et al., 2010). In addition, the relative immediacy of internet-based surveillance systems also allows for much quicker targeting of infection hot-spots in pandemic situations, as was done by companies such as Google during the 2009 swine-flu pandemic (Chew and Eysenbach, 2010; Signorini et al., 2011; St Louis and Zorlu, 2012).

2.5.1.1. Search-term surveillance. Google’s Flu Trends (GFT) (Ginsberg et al., 2009; Google Flu Trends) is perhaps the most well-known of the search-term surveillance systems. Combining data-mining of Google search queries and statistical modelling to provide a baseline indicator of the trend or changes in the rate of influenza, GFT provides estimates of weekly regional US influenza activity with a reporting lag of only one day compared with the 1–2 week delays associated with the CDC Influenza Sentinel Provider Surveillance reports (Ginsberg et al., 2009). GFT has been extended to include surveillance for dengue – Google Dengue Trends (GDT) (Chan et al., 2011; Google Dengue Trends) – and also been used to develop early warning systems for influenza epidemics (Pervaiz et al., 2012; Dugas et al., 2013; Cook et al., 2011).

Implemented in 29 countries, with a focus on Europe (Google Flu Trends; Eurosurveillance Editorial Team, 2009; Valdivia et al., 2009), GFT is currently best suited...
Crowdsourced tracking systems. Crowdsourced tracking surveillance systems, such as data-mining of Twitter posts, apply algorithms to filter tweets by specific keywords, assess their relevance and accuracy, geo-tag tweets and compare this information to other surveillance data. For example, NowTrending uses Twitter to track disease trends at both regional and national levels, presenting the most commonly tweeted diseases in a WordCloud. These metrics are intended to serve as an indicator of potential emerging health issues to spur further investigation and collection of direct measures of disease. In addition, recent studies demonstrate the value of combining social media with routine epidemiological data to detect or predict disease outbreaks, including influenza and cholera (Chew and Eysenbach, 2010; St Louis and Zorlu, 2012; Broniatowski et al., 2013; Chunara et al., 2012; Abrams et al., 2013) and to estimate weekly levels of influenza-like illness (Signorini et al., 2011).

Although one of the main advantages of crowd-sourced tracking surveillance systems is that of timeliness through the availability of real-time, georeferenced data (Stoové and Pedrana, 2014), a major limitation is the large amount of unrelated ‘noise’ (Chew and Eysenbach, 2010; Broniatowski et al., 2013; Denecke et al., 2013), although Broniatowski et al. (2013) appear to have developed an algorithm that can successfully distinguish relevant tweets from noise. The lack of specificity caused by noise may be less of an obstacle if the analysis is supported by trained public health officials who can investigate signals as they develop; Barboza et al. (2014) showed that systems including human moderation were found to have a 53% higher specificity after adjustment for other variables.

An additional weakness of Twitter is that its users do not represent a random sample of the population; the majority of Twitter users are aged between 18 and 50 (http://www.pewinternet.org/fact-sheets/social-networking-fact-sheet/) and therefore, drawing conclusions without considering the primary population demographic can be problematic. Furthermore, these surveillance tools appear to be most effective in developed countries; in Turkey a comparison of flu-related tweets with real world records showed no strong correlation (Bilge et al., 2012). However, language does not appear to be an issue as attested by a Portuguese study that successfully trained a Naïve Bayes classifier to identify tweets mentioning flu or flu-like illness or symptoms (Santos and Matos, 2014).

Despite limited evidence of internet-based surveillance systems to detect emerging threats before more traditional systems (Heymann and Rodier, 2001; Zeldenrust et al., 2008; Cowen et al., 2006), their primary value currently lies in their ability to act as an early-warning system thereby lessening the consequences of an outbreak (Wilson and Brownstein, 2009; Hartley et al., 2013). As such, although novel surveillance systems are still a long way from replacing traditional surveillance methods, they can usefully complement conventional approaches (Milinovich et al., 2014), to the extent that they have become an important component of the influenza surveillance scene. For example, WHO’s Global Outbreak Alert and Response Network uses such data as part of its day-to-day surveillance activities (Grein et al., 2000; Heymann and Rodier, 2001) and is authorized to act on this information (Wilson et al., 2008). In addition, internet-based data sources exist outside traditional reporting channels and as such, are invaluable to public-health agencies that rely on the timely flow of information across administrative borders.

However, search-term surveillance and crowdsourcing tracking systems clearly require in-depth evaluation, especially with respect to false positives and gaps in coverage and further work is necessary to determine how much of a change from baseline warrants further investigation.

2.5.2. Mining secondary internet-based data sources

In contrast to the surveillance systems that mine the primary data available through tweets and Google searches, there are also a number of surveillance systems that mine secondary data systems such as internet-based media sites; for example BioCaster (Collier et al., 2008; Collier, 2006; BioCaster), EpisPIDER (Keller et al., 2009; Tolentino et al., 2007), HealthMap (Brownstein et al., 2008; Wilson et al., 2013).
The value of such systems for flagging potential health threats is evidenced by the fact that GPHIN identified the 2002 severe acute respiratory syndrome (SARS) outbreak in Guangdong Province, China, more than two months before the World Health Organisation’s (WHO) official announcement (Mykhalsovskiy and Weir, 2006). Similarly, HealthMap identified news stories reporting a strange fever in Guinea nine days before official notification of the 2014 West Africa Ebola outbreak (Milinovich et al., 2015). Developing country initiatives include India’s Media Scanning & Verification Cell (MSVC) which scans global and national media sources and flags unusual health events, and has successfully flagged a number of outbreaks before they were identified by traditional surveillance systems (Sharma et al., 2012).

Although a comparison of BioCaster, EpiSPIDER and HealthMap identified significant differences in their ability to obtain relevant disease information (Lyon et al., 2012) – owing mainly to differences in sources searched, languages read, regions of occurrence and types of cases (Lyon et al., 2012; Barboza et al., 2014) – running the three surveillance systems in parallel has been shown to enhance early detection of disease anomalies over traditional surveillance approaches (Barboza et al., 2014). However, such automated systems are not without problems; for example, the location detection tool of all three systems assumed that the number of articles plotted for a country reflected the number of articles found about that country, which was not necessarily true (Lyon et al., 2012).

2.5.3. Active participatory online surveillance

In addition to the passive internet-based surveillance systems there are a number of active participatory surveillance systems that capture voluntarily submitted symptom data from the general public and can aggregate and communicate that data in near real-time thereby providing unique disease information that is not available through traditional surveillance sources. To date, such systems exist only for human diseases with a focus on influenza. Examples include Influenzanet, (FluTracking), Reporta, Flu Near You, Dengue na Web and SaludBoricu.

These systems show a high degree of accuracy and increased sensitivity and timeliness relative to traditional healthcare-based systems (Wojcik et al., 2014) and have proven useful for identifying risk groups, assessing disease burden, evaluating vaccination coverage and effectiveness, and informing disease transmission models (Marquet et al., 2006; Paolotti et al., 2010, 2014; Van Noort et al., 2012; Van Noort et al., 2007; Parrella et al., 2009; Friesema et al., 2009; Brooks-Pollock et al., 2011). In addition, they are cheaper and more flexible than traditional systems. Nevertheless, they present important challenges including, biases associated with the population that chooses to participate, difficulty in adjusting for confounders due to patients’ unwillingness to complete long surveys, and limited specificity because of reliance on syndromic disease definitions (Wojcik et al., 2014).

3. Conclusion

As a result of 21st century challenges, such as globalization and global warming, health officials and researchers are faced with increasingly complex and challenging disease problems which demand access to new and different types of data in order to inform effective risk-based disease surveillance and control strategies. The increasingly wide range of available spatial disease data may allow us to meet those challenges, assuming we see them as opportunities rather than problems; opportunities to convert aspects of the world that have never previously been quantified into valuable data that can shine a new light on health problems. Opportunities to develop timely and cost-effective online disease surveillance systems for developing nations that lack the necessary resources and infrastructure to implement traditional surveillance systems. Opportunities to transpose novel human health surveillance systems for use in animal health situations. And opportunities to respond to the increasingly complex disease problems facing us with state-of-the-art and spatially-explicit, risk-based disease surveillance and control strategies.

Conflict of interest

The authors report no conflict of interest.

References

Aanensen DM, Huntley DM, Feil EJ, Al-Own FA, Spratt BG. EpiCollect: linking smartphones to web applications for epidemiology, ecology and community data collection. PLoS ONE 2009;4(9):e6986.
Abath EN, Erssbøll AK, Lo Fo Wong DMA, Emborg HD. Space–time clustering of ampicillin resistant Escherichia coli isolated from Danish pigs at slaughter between 1997 and 2005. Prev Vet Med 2009;80(1–2):90–101.
Abrams JY, Copeland JR, Tauxe RV, Dale KA, Belay ED, Mody RK, et al. Real-time modelling used for outbreak management during a cholera epidemic, Haiti, 2010–2011. Epidemiol Infect 2013;141(06):1276–85.
Althouse BM, Ng YY, Cummings DAT. Prediction of dengue incidence using search query surveillance. PLoS Negl Trop Dis 2011;5(8):e1258.
Anokwa Y, Hartung C, Brunette W, Borriello G, Lerer A. Open source data collection in the developing world. Computer 2009;42(10):97–9.
AusCover Geo-Wiki. <http://www.geo-wiki.org/branches/auscover> [accessed January 2015].
Baker S, Holt KE, Clements ACA, Karkey A, Aryal A, Boni MF, et al. Combined high-resolution genotyping and geospatial analysis reveals modes of endemic urban typhoid fever transmission. Open Biol 2011;1(2).
Barboza P, Vaillant L, Le Strat Y, Hartley DM, Nelson NP, Mawudeku A, et al. Factors influencing performance of internet-based biosurveillance systems used in epidemic intelligence for early detection of infectious diseases outbreaks. PLoS ONE 2014;9(9):e90536.
Barrera R, Torres N, Freier J, Navarro J, García C, Salas R, et al. Characterization of enzootic foci of Venezuelan equine encephalitis virus in Western Venezuela. Vect Born Zoon Dis 2001;13(3):219–30.
Benemsa SC, Scholte RGC, Molento MB, Medeiros C, Carvalho OdS: Fasciola hepatica in bovines in Brazil: data availability and spatial distribution. Revista do Instituto de Medicina Tropical de São Paulo 2014;56:35–41.
Bergquist N, Tanner M. Visual approaches for strengthening research, science communication and public health impact. Geospat Health 2012;6(2):155–6.
Loobuyck M, Froséling J, Lindberg A, Björkman C. Seroreivalence and spatial distribution of Neospora caninum in a population of beef cattle. Prev Vet Med 2009;92(1–2):116–22.

Lozano R, Naghavi M, Foreman K, Lim S, Shibuya K, Aboyans V, et al. Global and regional mortality from 235 causes of death for 20 age groups in 1990 and 2010: a systematic analysis for the Global Burden of Disease Study 2010. Lancet 2012;380(9859):2095–128.

Lozano-Fuentes S, Weydan F, Hernandez-Garcia E, Sadhu D, Ghosh S, Biemans J. Cell phone-based system (Chaaq) for surveillance of immatures of dengue virus mosquitoes. J Med Entomol 2013;50(4):879–89.

Lyon A, Nunn M, Grossel G, Burgman C. Comparison of Web-Based Biosecurity Intelligence Systems: BioCaster, EpiSPIDER and HealthMap. Transbound Emerg Dis 2009;56(3):223–32.

Madder M, Walker JC, van Kruyen J, Koelewijn E, Berkers D, et al. e-Surveillance in Animal Health: use and evaluation of mobile tools. Parasitology 2012;139(Special Issue 14):1831–42.

Madon S, Amaguru JVO, Malecena MN, Michael E. Can mobile phones help control neglected tropical diseases? Experiences from Tanzania. Soc Sci Med 2014;102:103–10.

Malone J, Bergquist N. Mapping and modelling neglected tropical diseases and poverty in Latin America and the Caribbean. Geospat Health 2012;6(3):51–5.

Margaret R, Bartels RA, Van Noort S, Koppeschaar C, Paget J, Schellevis F, van der Zee J. Internet-based monitoring of influenza-like illness (ILI) in the general population of the Netherlands during the 2003–2004 influenza season. BMC Public Health 2006;6:242.

Martin V, De Simone I, Lubroth J. Geographic information systems as tools to the international surveillance and control of transboundary animal diseases, a focus on highly pathogenic avian influenza. Vet Ital 2007;43(3):437–50.

Martin V, De Simone I, Lubroth J, Ceccato P, Chevalier V. Perspectives on using remotely-sensed imagery in predictive veterinary epidemiology and global early warning systems. Geospat Health 2007;2(1):3–14.

McVey DJ, Brownstein JS. Wikipedia usage estimates prevalence of influenza-like illness in the United States in near real-time. PLoS Comput Biol 2014;10(4):e1003581.

Méro E, De Regge N, Rioucreux F, Caij AB, van den Berg T, van der Stede Y. Distribution of Schmallenberg Virus and Seroprevalence in Belgian Sheep and Goats. Transbound Emerg Dis 2014;61(5):425–31.

Métrás R, Porphyre T, Pfeiffer DU, Kemp A, Thompson PN, Collins LM, et al. Exploratory time-space analyses of rift valley fever in South Africa in 2008–2011. PLoS Negl Trop Dis 2012;6(8):e1808.

Milinovich GJ, Williams GM, Clements ACA, Hu W. Internet-based surveillance systems for monitoring emerging infectious diseases. Lancet Infect Dis 2014;14(2):160–8.

Milinovich GJ, Magalhães RJS, Hu W. Role of big data in the early detection of Ebola and other emerging infectious diseases. Lancet Global Health 2015;3(1):e26–1.

Minh PQ, Morris RS, Schauer B, Stevenson M, Benschop J, Nam HV, et al. Spatio-temporal epidemiology of highly pathogenic avian influenza outbreaks in the two deltas of Vietnam during 2003–2007. Prev Vet Med 2009;89(1–2):1–13.

Minh PQ, Stevenson MA, Jewell C, French N, Schauer B, Stevenson M, Benschop J, Nam HV, et al. Spatio-temporal analyses of highly pathogenic avian influenza H5N1 outbreaks in the Mekong River Delta, Vietnam, 2009. Spatial Spatiotemporal Epidemiol 2010;1:1:S3.

Müllner A, Robertson C, Sawford K, Daniel SLA, Nelson TA, Stephen C. Mobile phone–based infectious disease surveillance system Sri Lanka. Emerg Infect Dis 2010;16(10):1524–31.

Ortiz JR, Zhou H, Shao DK, Neuhiel KM, Fowlkes AL, Goss CH. Monitoring influenza activity in the United States: a Comparison of Traditional Surveillance Systems with Google Flu Trends. PLoS ONE 2011;6(4):e18137.

Paolotti D, Gioannini C, Colizza V, Vespignani A: Internet-based monitoring system for influenza-like illness: H1N1 surveillance in Italy. 3rd International ICST Conference on Electronic Healthcare for the 21st century 2010.

Picado A, Carahasan M, Colizza V, Eames K, Edmunds J, Gomes G, et al. Web-based participatory surveillance of infectious diseases: the Influenzanet participatory surveillance experience. Clin Microbiol Infect 2014;20:17–21.

Pfarrlla A, Dalton C, Pearce R, Litt J, Stocks N. ASPREN surveillance system for influenza-like illness – a comparison with FluTracking and the National Notifiable Diseases Surveillance System. Aust Fam Physician 2009;38:932–6.

Perez A, Alkhams M, Carlsson U, Brito B, Carrasco-Medanic R, Whedbee Z, et al. Global animal disease surveillance. Spat Statiotemporal Epidemiol 2011;2(3):349–54.

Pervaz A, Pervaiz M, Abdur Rehman N, Saif U. FluBreaks: early epidemic detection from Google Flu Trends. J Med Inform Res 2012;14(5):e125.

Pfeiffer DU, Robinson TP, Stevenson M, Stevens KB, Rogers DJ, Clements ACA. Spatial analysis in epidemiology. Oxford: Oxford University Press; 2008.

Picado A, Speybroeck N, Kivaria F, Mosha RM, Sumaye RD, Casal J, Berkers D. Foot-and-Mouth Disease in Tanzania from 2001 to 2006. Transbound Emerg Dis 2011;58(1):44–52.

Pigott DM, Bhatt S, Golding N, Duda KA, Battle KE, Brady OJ, Messina JP, Balard Y, Bastien P, Prattong F et al.: Global distribution maps of the arbovirauses; vol. 2014.

Pin-DioP R, Toure I, Lancelot R, Ndiaye M, Chavernac D. Remote sensing and geographic information systems to predict the density of ruminants, hosts of Rift Valley Fever virus in the Sahel. Vet Ital 2007;42(3):675–86.

Polak Z, Dewey CE, Martin SW, Christensen J, Carman S, Friendship RM. Spatial clustering of swine influenza in Ontario on the basis of herd-level disease status with different misclassification errors. Prev Vet Med 2007;81(4):236–49.

Polak Z, Dewey C, Rosendal T, Friendship R, Young B, Berke O. Spread of porcine circovirus associated disease (PCVAD) in Canada (Canada): swine herds: part I, Exploratory spatial analysis. BMC Vet Res 2010;6:59.

Porphyre T, McKenzie J, Stevenson M. A descriptive spatial analysis of tuberculosis on intensively controlled cattle farms in New Zealand. Vet Res 2007;38(3):465–79.

PromED-mail. <http://www.promedmail.org/> [accessed January 2015].

Purse BV, Tatem AJ, Caracappa S, Rogers DJ, Mellor PS, Baylis M, et al. Modelling the distributions of Culicoides bluetongue virus vectors in the United Kingdom using remotely-sensed imagery in predictive veterinary epidemiology. Comput Biol 2013;9(10):e1003256.

RapidSMS. <http://www.rapidsms.org/> [accessed January 2015].

Rapoport. <http://reporta.3.org.mx/> [accessed January 2015].

Ricotta E, Frese S, Choobwe C, Louis T, Shiff C. Evaluating local vegetable cover as a risk factor for malaria transmission: a new analytical approach using Image. Malar J 2014;13(1):94.

Rinaldi L, Musella V, Riggeri A, Cringoli G. New insights into the application of geographical information systems and remote sensing in veterinary parasitology. Geospat Health 2006;1(1):33–47.

Risk Geo-Wiki. <http://www.geo-wiki.org/branches/risks/> [accessed January 2015].

Robinson C, Sawford K, Daniel SLA, Nelson TA, Stephen C. Mobile phone-based infectious disease surveillance system Sri Lanka. Emerg Infect Dis 2010;16(10):1524–31.

Robinson TP, Harris RS, Hopkins JS, Williams BG. An example of decision support for trypanosomiasis control using a geographical information system in eastern Zambia. Int J Geogr Inf Sci 2002;16(4):345–60.
system in malaria epidemic management and surveillance. [Article in Chinese]. Zhongguo Xue Xi Chong Bing Fang Zhi Za Zhi 2013, 25(2):172–176.

Ward MP, Carpenter TE. Techniques for analysis of disease clustering in space and in time in veterinary epidemiology. Prev Vet Med 2000;45(3–4):257–84.

Ward MP, Cowled BD, Galea F, Garner MG, Laffan SW, Marsh I, et al. Salmonella infection in a remote, isolated wild pig population. Vet Microbiol 2013;162(2–4):521–9.

Wilesmith JW, Stevenson MA, King CB, Morris RS. Spatio-temporal epidemiology of foot-and-mouth disease in two counties of Great Britain in 2001. Prev Vet Med 2003;61(3):157–70.

Wilson K, Brownstein JS. Early detection of disease outbreaks using the Internet. Can Med Assoc J 2009;180(8):829–31.

Wilson K, von Tigerstrom B, McDougall C. Protecting global health security through the International Health Regulations: requirements and challenges. Can Med Assoc J 2008;179(1):44–8.

Yu P, de Courten M, Pan E, Galea G, Pryor J. The development and evaluation of a PDA-based method for public health surveillance data collection in developing countries. Int J Med Informatics 2009;78(8):532–42.

Zastrow M. Crisis mappers find an ally. Nature 2014;515:321.