Lessons learned in Central Ghana regarding the pandemic influenza a vaccine and how they could be used conserving new ebola virus vaccine

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Abstract

Background: The emergence of influenza A (H1N1) virus prompted the development of influenza A (H1N1) monovalent vaccines (2009-H1N1). The use of the vaccine was recommended by the Center for Disease Control and Prevention Advisory Committee on Immunization Practices. Adverse events after vaccinations occur but are generally rare. The study was carried out to identify reported adverse and serious adverse events after administration of an influenza A (H1N1) vaccine.

Method: This cross-sectional study was carried out between mid of July 2010 to end of August 2010 in Kintampo North Municipality and Offinso South Municipality in Ghana. The study was carried out as part of a pharmacovigilance study in Kintampo Health Research Centre (KHRC). Data were collected from consented participants using questionnaire.

Results: Of the 420 forms that were given out to consented participants in the two regions, 379 (90.2%) were returned with completed information related to the Influenza A (H1N1) vaccine. Participants who took the vaccine reported of adverse events such as fever, headache, chills, stomach ache, diarrhoea, pain in the heart and fast heartbeat.

Of those who received the vaccine, 4.4% (16/366) were hospitalized for the adverse event they reported to have experienced after vaccination.

Of the 4.4% of the vaccinated participants that were hospitalized, 43.8% (7/16) were males and 56.3% (9/16) were females. There was no difference between the proportions (p=0.97) of males and females that were hospitalized after vaccination.

Conclusion: Ghana used Pandemrix- Influenza A (H1N1) vaccine in June 2010. Symptoms reported ranged from what were expected to unexpected symptoms. This survey recorded sixteen (16) hospitalizations due to symptoms reported after vaccination. The study was not controlled and therefore could not make claims of whether the serious adverse events were associated with the vaccine. We recommend continuous and effective post-vaccination and post-marketing monitoring of adverse events after vaccinations.

Introduction

Emerging diseases of animal origin are a growing concern in the public health. About 75% (132/175) of species of pathogens classified as emerging are zoonotic [1]. Severe acute respiratory syndrome (SARS), avian influenza and Ebola Virus Disease (EVD) which have emerged on a global scale are some of the diseases associated with these pathogens.

The novel H1N1 influenza A virus that emerged in humans in Mexico in April 2009 and transmitted efficiently in the human population according to the World Health Organization (http://www.who.int/csr/don/2009_05_25/en/) spread globally to 46 countries with 12,515 reported cases and 91 associated deaths [2]. On 11 June 2009, the WHO raised the pandemic alert from Level 5 to Level 6 and declared the start of an influenza pandemic [3]. Declaration of the pandemic commenced discussion worldwide among public health officials, governments and pharmaceutical companies about development of an H1N1 vaccine [4].

From the sequenced genomes of samples of the new H1N1 taken from human patients diagnosed with the infection, analysis showed that the H1N1 virus had similar antigenic properties to some existing viral strains that made it easier to come up with a new candidate virus for vaccine development [5]. Vaccine was seen as one of the important means to curb the pandemic [6].

Vaccines serve two related though different functions: to protect vaccinated people against infection and severe disease; and to reduce transmission, thereby offering indirect protection to those not vaccinated via herd immunity [7]. Vaccines most effectively reduce transmission if they are given to the groups that are most likely to be infected and most likely to transmit the infection onward [7,8]. The emergence of influenza A (H1N1) virus prompted the development of influenza A (H1N1) monovalent vaccines (2009-H1N1) which its use after it has been licensed by the Food and Drugs Administration, was recommended by the Centers for Disease Control and Prevention (CDC) Advisory Committee on Immunization Practices (ACIP) [9,10]. Influenza A (H1N1) monovalent vaccine (2009-H1N1) was the most effective measure to control the spread of the virus associated with H1N1 Influenza and to reduce its associated morbidity and mortality [10,11]. Vaccines which have side effects and differ in effectiveness are usually administered to healthy individuals trusting in its safety.

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and efficacy. Most vaccines have contributed substantially to improve human health in the past century.

The timing of vaccine production and release is a crucial factor. Development of the pandemic vaccine for emergency use also required some flexibility in the regulatory pathway because time was not sufficient for full clinical trials [7-12]. The challenge is that between development of a vaccine and its use publicly, there is a multiplicity and heterogeneity of human factors that must be negotiated to realise the public health gains of using the vaccine to immunize [13].

Ideally, vaccines manufactured after pre-clinical trials are to be taken through three phases of clinical trials to assess safety, efficiency, immunogenicity and efficacy prior to marketing approval. This allows the evaluation to be carried out among individuals of varying traits and larger population. The Influenza A H1N1 pandemic vaccine received approval for use after assessment of the convincing results of ongoing clinical trials [14,15]. This can be understood considering the need for a vaccine in the pandemic because the development of the Influenza A H1N1 pandemic vaccine was re-engineered based on an existing vaccine for H5N1 virus strain using the 'Mock-up' approach [10-16]. Hence might not have required the full length of phases in clinical trials. When making decisions about regulatory approval and public health use of vaccines, authorities need to be convinced that the benefits to reduce disease outweigh actual and potential risks of vaccination [17].

Adverse events after vaccinations occur but are generally rare. Some adverse events are unlikely to be detected in pre-licensure clinical trials because of their low frequency, the limited numbers of enrolled subjects, and other study limitations [18,19]. These limitations may hinder the reports of serious adverse events during the clinical trials of vaccines.

Readers' comments and responses on specific stories provided by media outlets and to respond to other readers' comments serve as a gauge of public opinion that is immediate, spontaneous and (presumably) honest [4].

Pandemic Influenza A H1N1 2009 was first recorded in Ghana in October, 2009 [20,21]. Media reportage of the pandemic heightened as there were threats of cases being found and diagnosed in Ghana. In the 2010 half year report of the Ashanti Regional Health Directorate of the Ghana Health Service, an H1N1 outbreak was reported in a in primary school in the capital city [22]. Due to the number of vaccines received, when the Ministry of Health decided to vaccinate Ghanaian against the influenza, it was limited and was targeted at health workers, persons at risk of severe disease, security personnel and schools that were reported to have suspected cases [20,21]. The H1N1 pandemic influenza vaccination which recorded excellent coverage received a lot of negative reports from the media and rumours of severe adverse reactions [22,23].

Ebola Virus Disease (EVD) Outbreak in West Africa (Guinea, Liberia, Sierra Leone, Nigeria, Mali and Senegal) was notified as extraordinary event & Public Health Emergency of International Importance (PHEIC) in August, 2014 by WHO [24]. The situation raised fundamental questions on: how much consideration if experimental treatments or vaccines were deployed; if any interventions proved safe and effective, how can they be made more widely available (Rid and Emanuel, 2014), and the cause of the delay with which the international community reacted to the EVD menace [25].

The plague of the EVD was in an unprecedented manner and killed most of its infected victims since April 2014 [26]. The quest to curb the menace made health regulatory authorities give directives in medical practice that were questioned ethically. This was obvious in the permit given for the Z-MAPP to be used for the treatment of Ebola infected individuals [27,28]. Vaccines have been developed and are already in the processes for clinical trials for possible registration and usage. We report of identified adverse and serious adverse events that were recorded in a survey to assess an influenza A (H1N1) vaccine introduced and administered in the public health system of Ghana to handle the emergence of the pandemic in Ghana from June 2010 to September 2010. This in our view will present lessons that need to be considered in the processes of taking possible Ebola vaccines through clinical trials, approvals and marketing.

Method

Descriptions: Vaccine: Pandemrix vaccine is a monovalent, split virion inactivated influenza vaccine containing antigen from A/California/07/2009 (H1N1) v strain [29]. It contains an immunologic adjuvant AS03 which consists of DL-a-tocopherol (vitamin E), squalene and polysorbate 80, with Thiomersal (thimerosal) as preservative with additional important non-medicinal ingredients such as formaldehyde, sodium deoxycholate, and sucrose. The final formulation contains 3.75 μg haemagglutinin (HA) per 0.5 ml dose adjuvanted with AS03. The vaccine is given as a single dose for most people, with a two-dose schedule for children under the age of 10 years and immunocompromised adults [29,30].

Adverse event: Any symptom a participant experienced after the administration of the pandemic influenza vaccine.

Serious Adverse event: In the period of the survey, we considered any symptom a participant experienced after the administration of the pandemic influenza vaccine, reported and was hospitalized to stabilise.

Study design and site: The cross-sectional study was carried out between mid of July 2010 and 31st of August 2010 in the Kintampo North and Offinso South Municipality in the Brong Ahafo and Ashanti Regions of Ghana respectively.

Kintampo North Municipal cover in total, an area of 5,108km2 with a resident population of approximately 96,358 (Ghana Districts). The study area is located within the forest-savannah transitional ecological zone in Ghana where subsistent farming is the predominant occupation. Mean monthly temperatures range between 18°C to 38°C and rainfall averages 1250 mm per annum. Since 2010 the study area has records of average vaccination coverage of at least 90% of the various vaccines in the Expanded Programme on Immunization (EPI).

Offinso South Municipal is located in the extreme north-western part of the Ashanti Region. Offinso Municipal lies between longitude 1° 65W and 1° 45E and latitudes 6° 45N and 7° 25 S. The municipal covers an area of 1255km2 with a resident population of approximately 138,190. The municipality had an average EPI coverage of at least 80% in 2010 and also an H1N1 vaccination coverage of about 62% of the target population during the period of vaccination [31].

Study Population: The study team distributed questionnaire to consented participants who could read and understand English with minimum instructions and introduction since the questionnaire were written in English. Aside this criterion, individuals targeted to be interviewed were to be more than fifteen (15) years of age. Questionnaires were therefore distributed to prospective participants.
assumed to be at the level of Senior High School and beyond (Midwifery/ Nursing Training and Teacher Training Institutions) based on age for interviewing. The selection of the population was made to reflect the group of individuals that were considered by the Ministry of Health to be at high risk and were considered for the vaccination.

**Procedures**

Members of the study team piloted the administration of the questionnaire which was adopted from the tools use for the Safety and Pharmacovigilance component of a study in the Kintampo Health Research Centre from the Food and Drugs Authority, Ghana with information from the CDC website on known symptoms associated with influenza A H1N1 infection. Information from the media reports were also used to modify the questionnaire to find out if any unsuspected adverse event could be found among individuals vaccinated with Pandemrix.

Mode of administration of the questionnaire was regularised in order to obtain data that would be comparable since we were collecting and collating data from two different regions of the country.

The study was introduced and explained to individuals, groups of people who were considered eligible. Inform consents were given by individuals who signed to respond to the questionnaires administered.

**Ethical Issues:** Informed consent was sought from all participants in the study after a detailed explanation of the purpose of the study. Permissions were also sought from the heads of the various institutions that were used for the study. The survey was approved by the Institutional Ethics Review Committee of Kintampo Health Research Centre, Ghana.

**Data Management and Analysis:** Microsoft Access data management software version 2007 was used for data entry. Stata software version 12 and GraphPad Prism software version 5.0 were used for statistical analyses of the data. All data were double entered on computers, and consistency checks were performed and discrepancies resolved with reference to the original form. The proportions of participants who reported adverse events after vaccination were tabulated. Differences in symptoms reported among hospitalized and those not hospitalized after vaccination were considered significant with 95% confidence interval.

**Results**

Of a total of 420 forms that were given out to consented participants, 379 (90.2%) were returned with completed questionnaires. The selection of the population was made to reflect the group of individuals that were considered by the Ministry of Health to be at high risk and were considered for the vaccination.

**Hospitalization, a serious adverse event, of 4.4% (16/366) was recorded among the participants who received the vaccine for the adverse events reported after vaccination. Majority, 56.3% (9/16), of the hospitalized individuals stayed on admission for three (3) days with only one person who had a prolonged stay of seven (7) days in hospital (Figure 1).

Of the 4.4% of the vaccinated participants that were hospitalized, 43.8% (7/16) were males and 56.3% (9/16) were females. There was no significant difference ($p=0.97$) between the proportions of males and females that were hospitalized after vaccination.

In total, 68.3% (250/379) of the participants responded to the question on the duration it took for an adverse event experienced after vaccination to resolve. The duration a participant endured an adverse event ranged from a minimum of one day to a maximum of twenty days (Table 3).

**Discussions**

Ghana started use of Pandemrix- Influenza A (H1N1) vaccine in June 2010. Upon its introduction, there was a lot of public outcry about the adverse events experienced after vaccination. The heightened media attention increased the awareness of the public to the potential vaccine side effects. The symptoms observed from the participants in multiple responses after vaccination that were heightened ranged from "severe pain at injection site" (12.3%), "headache" (11.7%), "fever" (10.3%) and "General body pains" (9.0%) to unexpected "fast-heart beat" (2.0%) and "pain in heart" (0.7%) adverse events (Table 2). Notably, with the emerging suspicion that Pandemrix could be linked to narcolepsy, the survey recorded 4.7% of drowsiness (feeling sleepy) and 4.7% of sleeplessness [32,33].

This survey of 379 people considered to be in a high-risk group to receive the pandemrix recorded sixteen (16) hospitalizations due to symptoms reported after vaccination. The study was not controlled and therefore could not make claims of whether the serious adverse events were associated with the vaccine. However, it is interesting to note the prolong hospitalization was recorded of individuals that received the vaccine and were hospitalized for the adverse events they developed.

The study was not designed to do proper systematic causal assessments of the serious adverse events and therefore does not have...
Table 2. Frequently reported adverse event associated with influenza A (H1N1) 2009 vaccine in Central Ghana

<table>
<thead>
<tr>
<th>Adverse event</th>
<th>N (% of responses)</th>
<th>% of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fever</td>
<td>148 (10.3)</td>
<td>40.7</td>
</tr>
<tr>
<td>Headache</td>
<td>168 (11.7)</td>
<td>46.2</td>
</tr>
<tr>
<td>Fatigue</td>
<td>116 (8.1)</td>
<td>31.9</td>
</tr>
<tr>
<td>Vomiting</td>
<td>13 (0.9)</td>
<td>3.6</td>
</tr>
<tr>
<td>Nausea</td>
<td>34 (2.4)</td>
<td>9.3</td>
</tr>
<tr>
<td>Stomach ache</td>
<td>23 (1.6)</td>
<td>6.3</td>
</tr>
<tr>
<td>General body pains</td>
<td>139 (9.7)</td>
<td>38.2</td>
</tr>
<tr>
<td>Swelling at injection site</td>
<td>28 (2.0)</td>
<td>7.7</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>20 (1.4)</td>
<td>5.5</td>
</tr>
<tr>
<td>Inflammation at site</td>
<td>12 (0.8)</td>
<td>3.3</td>
</tr>
<tr>
<td>Itching</td>
<td>22 (1.5)</td>
<td>6.0</td>
</tr>
<tr>
<td>Dizziness</td>
<td>48 (3.4)</td>
<td>13.2</td>
</tr>
<tr>
<td>Drowsiness</td>
<td>68 (4.7)</td>
<td>18.7</td>
</tr>
<tr>
<td>Sleeplessness</td>
<td>68 (4.7)</td>
<td>18.7</td>
</tr>
<tr>
<td>Loss of appetite</td>
<td>56 (3.9)</td>
<td>15.4</td>
</tr>
<tr>
<td>Chills</td>
<td>77 (5.4)</td>
<td>21.2</td>
</tr>
<tr>
<td>Restlessness</td>
<td>48 (3.4)</td>
<td>13.2</td>
</tr>
<tr>
<td>Stuffy nose</td>
<td>21 (1.5)</td>
<td>5.8</td>
</tr>
<tr>
<td>Fast heart-beat</td>
<td>28 (2.0)</td>
<td>7.7</td>
</tr>
<tr>
<td>Pain in heart</td>
<td>10 (0.7)</td>
<td>2.8</td>
</tr>
<tr>
<td>Severe pain at injection site</td>
<td>176 (12.3)</td>
<td>48.4</td>
</tr>
<tr>
<td>Runny nose</td>
<td>34 (2.4)</td>
<td>9.3</td>
</tr>
<tr>
<td>Others</td>
<td>8 (0.6)</td>
<td>2.2</td>
</tr>
<tr>
<td>None</td>
<td>69 (4.8)</td>
<td>19.0</td>
</tr>
<tr>
<td>Total</td>
<td>1434 (100)</td>
<td>394.0</td>
</tr>
</tbody>
</table>

*Valid cases: 364 participants vaccinated with H1N1 influenza vaccine with multiple responses to symptoms experienced.

Table 3. Indication of the number of days participants experienced the side effect after vaccination

<table>
<thead>
<tr>
<th>Duration for AE to clear</th>
<th>N (% of responses)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-3 days</td>
<td>128 (51.2)</td>
</tr>
<tr>
<td>4-6 days</td>
<td>75 (30.0)</td>
</tr>
<tr>
<td>7-9 days</td>
<td>29 (11.6)</td>
</tr>
<tr>
<td>10-20 days</td>
<td>18 (7.2)</td>
</tr>
<tr>
<td>Total</td>
<td>250 (100)</td>
</tr>
</tbody>
</table>

Figure 1. Number of days of hospitalization of vaccinated participants

Participants were selected to minimize bias by simply selecting participants who were ready to consent and respond to our questionnaire, providing a control group for better analysis would have minimized speculations and assumptions.

The study had a limitation of selection bias. Since the study was interested to source information on adverse and serious adverse events after vaccine administration, participants’ selection was from high risk individuals who had been considered for vaccination with the H1N1 Influenza Pandemrix vaccine.

Also, there was the possibility of the effect of recall bias and the ability of participants to understand the various checklist of symptoms presented on the questionnaire. Though the symptoms were presented in very simplified English language, the study did not assess the level of participants understanding of the symptoms which could also affect what was chosen as response.

Lessons to EBV vaccines

Effectiveness data if available would be necessary to perform a clear and fair risk-benefit balance assessment; this assessment is important taking into account the characteristics of the H1N1 influenza pandemic [9-34]. Pharmaceutical companies have produced vaccines to EVD and the desire to take it through clinical trials and further use for protection if found efficacious. These gave rise to ethical, safety and scientific concerns in the manner in which the processes to conduct the clinical trials were being hastened [35-36]. It is of good motive these processes were being done to satisfy the quest for vaccine to the plagues, many of which were of zoonotic origin.

We recommend continuous and effective post-vaccination and post-marketing monitoring of adverse events after the introduction of any of the Ebola vaccines to monitor safety and review spontaneously reported adverse events which will add to safety data that will emerge from other sources.

Authors’ contributions

DAG, DKG and KP designed the study. DAG, DKG, KP, LA, AK, SA performed the experiments and co-wrote the manuscript. KB supported with the data collection. EM, AM and SOA contributed to the study design, carried out the statistical analysis, performance of experiments and wrote the manuscript. DAG, KP, SOA designed the experiments, supervised the study and co-wrote the manuscript.

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Conflict of Interest

The authors declare no conflict of interest.

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