HBV vaccine efficacy and detection and genotyping of vaccineé asymptomatic breakthrough HBV infection in Egypt

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Abstract

AIM: To evaluate the impact of mass vaccination against the hepatitis B virus (HBV) in Egypt, and to search for vaccinee asymptomatic breakthrough HBV infection and its genotype.

METHODS: Seven hundred serum samples from vaccinated children and adults (aged 2-47 years) were used for quantitative and qualitative detection of HBsAb by ELISA. Three hundred and sixty serum samples representing undetectable or low or high HBsAb were screened for markers of active HBV infection (HBsAg, HBCab (IgG) and HBeAb by ELISA, plus HBsAg by AxSYM) and HBV-DNA genotyping by nested multiplex PCR and by DNA sequencing.

RESULTS: It was found that 65% of children aged 2-4 years, and 20.5% aged 4-13 years, as well as 45% adults were good responders to HBV vaccination mounting protective level HBsAb. Poor responders were 28%, 59.5% and 34%, and non-responders were 7%, 20% and 21% respectively, in the three studied groups. Markers of asymptomatic HBV infections were HBsAg detected by ELISA in 2.5% vs 11.39% by AxSYM. Other markers were HBcAb (IgG) in 1.38%, HBeAb in 0.83%, and HBV-DNA in 7.8%. All had HBV genotype E infection.

CONCLUSION: It is concluded that HBV vaccine is efficient in controlling HBV infection among children and adults. The vaccine breakthrough infection was by HBV genotype E. A booster dose of vaccine is recommended, probably four years after initial vaccination.

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Key words: HBV vaccine evaluation; Egyptian children; Adults; Genotype E vaccine escape HBV

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INTRODUCTION

The Hepatitis B virus (HBV) is endemic in many developing countries, including Egypt[1-3]. As the majority of chronic carriers of HBV in the world become so as a result of infections which occur prior to the age of 6, the Technical Advisory Group (TAG) of the World Health Organization has recommended that the HB vaccine must be added as a component of the Expanded Program of Immunization (EPI) in countries with a moderate to high prevalence[4]. The recombinant DNA-based vaccine, targeting the HBV major surface protein (r-HBsAg) has been incorporated in the national childhood vaccination program in Egypt since October 1992. Three doses of 10 μg r-HBsAg are given at 2, 4, and 6 mo of age[6]. The vaccination of infants, children and adolescents have produced high rates of seroconversion (95%) and induced adequate levels of HBsAb[5]. The accepted level of seroconversion is 10 or more mIU/L, which provides protection against HBV infection. While the success of the vaccine cannot be denied, it was noticed that vaccine efficiency was improved by the addition of preS1 and preS2 components. Universal vaccination of infants with the HB vaccine has dramatically reduced infection, as well as the hepatitis-B surface antigen (HBsAg) carrier rate of chronic HBV, in addition to a significant decrease in the incidence of childhood hepatocellular carcinoma[8,9]. Protective immunity has been demonstrated in persons and populations up to 5 to 10 years post-vaccination, with associated decrease of asymptomatic HBV breakthrough infection[10-12]. In HBV endemic areas, postnatal or prenatal at the time of delivery mother-to-infant transmission of infection occurs frequently, especially if the maternal serum is hepatitis-Be antigen positive, which is the stage at which 90% of babies acquire HBV infection. Despite r-HBsAg vaccination, some of these infants become persistently infected and increase the worldwide HBV reservoir[13]. Furthermore, 5%-10% of the vaccinees display an inadequate antibody response following primary vaccination with triple doses of either plasma-derived or r-HBsAg vaccine, in addition to 3%-20% of non-responders, and may not be protected from subsequent exposure to HBV infection[13,14]. The introduction of a safe, effective, hepatitis B vaccine has led to universal infant vaccination, resulting in a reduced rate of perinatal HBV infection from infected mothers. Because of appropriate hepatitis B vaccination and passive immunoprophylaxis with hepatitis B immune globulin (HBIG) in infants of mothers with HBV infection, perinatal transmission has been reduced to less than 5% to 10%[14]. The configuration of the HBsAg used in current vaccine formulations contains a determinant which is located between amino acids (aa) 121-149 of the HBsAg immunogenic epitope, that trigger the production of polyclonal antibodies against the HBV major surface protein (HBsAb). The emergence of HBsAg variants, with mutations within aa 121-149 with altered antigenicity, and binding to the HBsAb, has been reported in HBV from vaccinees in several areas of the world[15,16]. Using the currently available diagnostic assays, these variants may go undetected, and could potentially cause breakthrough infections in a vaccinated population, posing a potential threat to the long term success of HBV vaccination programmes[17-19]. Booster doses of Hepatitis B vaccine are recommended only in certain circumstances, for example, for hemodialysis patients, and for those with an ongoing risk of exposure. Annual anti-HBs testing and a booster dose should be administered when anti-HBs levels decline to < 10 mIU/mL. For other immuno-compromised persons, the need for booster doses has not been determined[20]. It has been appreciated that long-term immunity derives from immunological memory, which outlasts the loss of antibody levels. Hence subsequent testing and administration of booster doses is not required in successfully vaccinated immunocompetent individuals. With the passage of time and longer experience, protection has been shown to last for at least 25 years in those who showed an adequate initial response to the primary course of vaccinations[20].

In 1988, genotypes of HBV were proposed by a sequence divergence in the entire genome exceeding 8%, based on a comparison of 18 HBV isolates[21]. Four genotypes were recognized and they were given capital letters of the alphabet from A to D. In 1994, Nordet et al[22] found an additional two HBV genotypes by means of the same criteria, and named them E and F; genotype G was reported in 2006[23], and genotype H was reported in 2002[24]. The vaccine “a” component of HBsAg is stable, due to conserved gene sequences encoding it in all HBV genotypes. On the other hand, the preS1, preS2 containing vaccines, which are more immunogenic, have high specific motifs. Thus there is a need to tailor the vaccine to the HBV genotype prevalent in the geographic areas of vaccination.

The objective of the current study was to evaluate the impact of mass childhood HBV vaccination in Egypt on asymptomatic HBV breakthrough infection in vaccinated individuals, and to determine the causative genotype.

MATERIALS AND METHODS

This study was done in 2004, 12 years after the start of the vaccination programme. Laboratory testing was conducted from October 2004 to January 2007 at the Virology Laboratory, Microbiology Department, Faculty of Medicine, Al-Azhar University, the Virology Laboratory in the Military Central Laboratory and the Molecular Epidemiology Department NAMRU#3 Cairo, Egypt.

Subjects of study

Six hundred serum samples from vaccinated children were collected (after obtaining the legal guardian consent). The six hundred children had received HBV vaccine [Engerix-B (Sankinsline “Sigma” licensed at 1989) from October 1992 until 1996. They received Euvax vaccine (Korea)] at 2, 4, and 6 mo of age, according to the vaccine schedule of the Egyptian Ministry of Health Population (MOHP). Serum samples from children aged 2-4 years of age were obtained from the Pediatric out-patient clinic.
at Al- Zahraa University Hospital, Cairo Governorate, Egypt. Serum samples of children aged from 4-13 years were obtained from the Maternal and Child Health Care Center in Qusena City, Menofya Governorate, or from Al- Zahraa University Hospital, Cairo Governorate, Egypt.

A hundred serum samples from vaccinated adults were collected (after obtaining individual consent) from Benha Teaching Hospital, the Motor Rehabilitation Institute and the Hearing and Speaking Institute, Qualyubia Governorate. All serum samples were collected between October 2004 and August 2005. Sera were divided according to the age of vaccinees into three groups:

**Group I:** two hundred children aged 2-<4 years. These children were considered healthy, with no history of medical or surgical problems, or risk factors for HBV infection, except for males who had been circumcised.

**Group II:** four hundred children aged 4-13 years, some of whom had past surgical history (circumcision, tonsillectomy, para-umbilical & inguinal herniorrhaphy).

**Group III:** one hundred sera from healthy adults, whose age ranged from 20-47 years. This group was without any history of risk for HBV infection, except that some females had a history of previous operations, such as caesarean section, tube dilatation and cervical curettage. All adults had received the 20 μg r- HBsAg at a 0, 1, 6 mo interval schedule. The last dose of vaccine in the 90 adult volunteers was administered within theyear before inclusion in the study, six adults had had their last dose between one and four years previously, and four volunteers had had their last dose six years or more prior to the study. This study did not include another age and gender matched non-vaccinated subject, as almost all children had been vaccinated within the MOHP vaccine schedule.

**HBV serological testing**

BIO ELISA (BioKit, Barcelona, Spain) HBsAb kits were used to both quantitatively and qualitatively assess HBsAb in the 700 post-vaccination serum samples. A random selection of 360/585 serum samples that were either negative 82/360 or demonstrated low antibody titer 287/360 (< 10 mIU/L) together with 41/360 serum samples with high antibody titer (> 10 mIU/L), were subsequently screened for HBsAg, using two commercial kits; one to detect both wild and mutant strains (AxSYM from ABBOTT), and the other the ELISA Bioelisa for HBsAg (Bio-Kit, Barcelona, Spain). All serum samples positive for HBsAg were screened for the detection of HBcAb (IgG) (Bio-Kit, Barcelona, Spain) and HBeAb using (Diaisorin, S.P.E Italy).

**HBV-DNA detection and genotyping by nested multiplex PCR**

Serum DNA was extracted using Kaucner and Stinear 1998 heat shock method. In some cases, DNA was extracted using the Qiagen DNA Blood mini kit, according to the manufacturer’s protocol (Qiagen, Valencia, CA, USA). Extracted DNA was subjected to HBV-DNA detection and genotyping using the nested multiplex PCR (nm PCR) method. Briefly, for the first round PCR, the primer pair P1 sense (5’ TCACCATATCTTGGGAAAGA 3’) and S 1-2 antisense (5’ CGAACCACCTGAAATATGGC 3’) were used to amplify the conserved regions of the pre-S1 and S-gene (1063 bases). The reaction mixture contained 5 μL of extracted DNA in 25 μL 1 × PCR buffer containing 1.5MgCl₂, 5 pmol of each primer completed 200 μmol/L of each of the four deoxyribonucleotides, 1U of AmpliTaq Gold DNA polymerase (Perkin-Elmer, Norwalk, Conn.) and completed to 50 μL with DEPEC treated sterile water. The samples were incubated at 95°C for 10 min, followed by 40 amplification cycles of 94°C for 20 sec (denaturation), 55°C for 20 sec (annealing), 72°C for 1 min (extension), and then followed by further extension at 72°C for 10 min. After that the product was kept at 4°C. The second-round PCR was performed for each specimen using inner pair primers in two different combinations, Mix-A: B2:5’ GGCTCMA0TTCMGAGAACAGT 3’ (nt 67-86, types A to E specific, sense), B4R: 5’ CTGCGGGAATGAGATGTG 3’ (nt 113-134, type A specific, anti sense), BBfR: 5’ CAGGTTG0GTGATGAGT GACTGGAGA 3’ (nt 324-345, type B specific, antisense), BCR: 5’ GGTCCTAGATCTGATGTG 3’ (nt 165-186, type C specific, antisense) for genotypes A, B and C using universal common primer B2 (sense) and specific primers A4, B5 and C6. Mix B: BDf: 5’ GCCAAC- AAGTAGGAGCT 3’ (nt 2979-2996, type D specific, sense), BBf: 5’ CACCAGAAATCCAGATGGCACA 3’ (nt 2955-2978, type E specific, sense), BBf: 5’ GTTACAGGGATYTGCTGG 3’ (nt 3032-3051, type F specific, sense), B2R: 5’ GAGCCGCGATYTGCTGCA- CAA 3’ (nt 3078-3097, type D specific, antisense), for genotype D, E, and F using universal primer R10 (antisense) and specific primers D7, E8 and F9. In the second round PCR 5 μL of the first PCR product was added to each mix with the same components of the first round PCR. These were amplified for 40 cycles, with the following parameters: hot start at 95°C for 10 min, followed by 25 cycles of 94°C for 20 sec (denaturation), 58°C for 20 sec (annealing), 72°C for 30 sec (extension), and an additional 20 cycles of 94°C for 20 sec, 60°C for 20 se, 72°C for 30 sec, which was then followed by further extension at 72°C for 10 min, then the product was kept at 4°C. The products of this second round PCR were visualized by electrophoreses on 3% agarose gel, and are differentiated by the size of genotype-specific DNA bands, compared to a 50 base-pair DNA marker (Amersco).

To assess changes in the HBV S gene, the following primers were designed to amplify the whole S gene, and used in a semi-nested PCR, HBsP1f (forward): 5’ GGAGYKG- GCGCATC CGGS 3’, HBsP2f (forward): 5’ GTTACAGGCGGTTTTCGTG 3’ and HBsP4r (reverse): 5’TCA­CACATCATC ATGATG TG 3’. The first round PCR was done using HBsP1f and HBsP4r, the reaction

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mixture consists of 5 μL of extracted DNA in 50 μL of a reaction buffer made of 30 pmol of each primer, 10 μmol/L of each of the four deoxynucleotides, 1 U of AmpliTaq Gold DNA polymerase, and 5 × PCR buffer containing 25 mol/L MgCl₂. The samples were incubated at 95°C for 10 min, followed by 35 cycles of 94°C for 30 sec (denaturation), 60°C for 1 min (annealing), 72°C for 1 min (extension), followed by further extension at 72°C for 10 min. The second round PCR was done in the same way as the first round, using HBSP2f and HBSP4r primers. To confirm the obtained genotype, and to determine the existence of any mutations, DNA sequencing of the S gene product was conducted using BigDye terminator technology and an ABI 377 fluorescence automated sequencer. DNA sequence was manually edited, using the software program Bioedit v7.0.5 [28]. Sequence comparisons of the obtained Egyptian strain (a positive control chronic hepatitis patient EGYAZC P1P2P4 bankit 1229997 GQ253-10c) were made using the program Clustal X [30] and HBV gene sequences retrieved from the Gene Bank: HBV genotype E (AB1832AB274977, LAGOS58S81AJ604967, CAR194AM494753, X75657, EU239226 PW6, 235-01DQ-060830, MAL136AJ604992, CMR936 ABI194948) [34] and HBV genotype A X75669, HBV genotype B X75660, HBV genotype C X75656, HBV genotype F X75658 [33]. In addition, alignment of 87 nucleotide sequence obtained from preS1 genes from Egyptian HBV isolate (a sample from a vaccinated child EGYAZV B2 265R) and Gene Bank strains as indicated: HBV genotype E PW6 EU239226 [31], HBV genotype A DQ788725 [33], HBV genotype B FM211360 [18], HBV genotype Cx75656 and HBV genotype D x75658 [33] was done, and Phylogenetic analysis and distance calculations for molecular evolutionary analyses were conducted using MEGA version 3.1 [34]. Phylogenies were constructed using the Neighbor-joining method and substitutions were modeled using the Kimura 2-parameter model, and phylogenetic analysis of selected sequences based onPreS1 and S fragments was done according to Felsetten [1988] [35].

RESULTS

**Egyptian children and adults' immune response to r-HBsAg vaccination**

The r-HBsAg vaccine used in Egypt appears to be efficient in inducing HBsAb immune response, which may, in turn, be efficient in controlling HBV infection in children and adults (≥ 10 mIU/mL), as 93% and 80% of children aged 2-<4 years, 4-13 years respectively, and 79% adults acquired protective HBsAb. Having low HBsAb titre (less than 10 mIU/mL) were 28%, 59.5% and 34% in the three studied groups respectively (Table 1).

### Table 1 Percent of positive cases, for HBs antibodies (IgG) by ELISA in the three studied population

| Group | Children (2-< 4 years) | Group II | Children (4-13 years) | Group III | Adults (20-47 years) | Total No |
|-------|------------------------|----------|-----------------------|-----------|----------------------|---------|
|       | N=200                  | N=400    | N=100                 |           |                      | N=700   |
| Anti-HBs IgG |       |          |                       |           |                      |         |
| High +ve samples (10-99.9 mIU/mL) | 130 (65) | 82 (20.5) | 45 (45) | 257(36.7) |         |
| Low +ve samples (<10 mIU/mL) | 56 (28) | 238 (59.5) | 34 (34) | 328(46.9) |         |
| Negative samples (Non-responders) | 14 (7) | 80 (20) | 21 (21) | 115 (16.4) |         |
| Total positive samples | 186 (93) | 320 (80) | 79 (79) | 585 (83.6) |         |

**Breakthrough HBV infection in vaccinated children and adults**

The infection determined by detection of HBsAg by BioELISA test was 2.5% vs 11.39% by AxSYM. Active HBV replication by DNA amplification procedures was 6.11%, and other serological markers of infectivity were HBeAb (IgG) 1.38% and HBeAb 0.83% (Tables 2, 3) respectively.

Among serum samples from children aged 2-<4 years old, 65% had a high HBsAb titer, 28% had low titer, and 7% did not have detectable HBsAb. The majority of these children had no markers of HBV infection by BioELISA test for HBsAg. Only 5/5 (9.09%) were infected as determined by detection of the HBsAg by AxSYM (Tables 1-3).

In contrast, only 20.5% of children aged 4-13 years had a high HBsAb titer, 59.5% had a low HBsAb titer, and 20% did not have detectable HBsAb. Screening for HBV infection in this age group revealed that 2.04% and 12.24% were positive for HBsAg by BioELISA and AxSYM respectively. It was noticed that 7.14%, 9.55% and 20.75% of high, low and undetectable HBsAb were positive for HBsAg, 7.34% (2.25% in low HBsAb and 26.4% in undetectable HBsAb) for HBV-DNA, 0.81% for HBeAb and 0.4% for HBeAb (Tables 1-3).

Finally, within the adult age group 45% of the serum samples had a high antibody titer, 34% of samples had a low antibody titer and 21% were negative. In this age group, 6.66% and 11.66% were positive for HBsAg by ELISA and AxSYM, respectively, 5% were positive for HBeAb, and 3.33% were positive for HBeAb; HBV-DNA was detected in 6.66% of the serum samples (Tables 1-3).

On examining 41 high HBsAb positive samples, regardless of the assay, they were negative for all markers tested, except for one sample which was positive for HBsAg by AxSYM assay (Table 2).

Out of 22 cases positive for HBV DNA; 4 cases of group III who were also HBsAg positive by both methods,
and 18 cases of group II (5 of them were HBsAg positive by BioELISA 11 by AxYM and two were HbcAb positive), while 12 participants had HBV-DNA as the only marker for HBV infection. Considering HBV-DNA detection by nmPCR as a reference test, it was found that BioELISA specificity (100%), BioELISA sensitivity (96.29%), AxYM specificity (50%), AxYM sensitivity (96.65%).

Genotyping and sequencing of the S gene
In all the HBV-DNA positive samples, genotype E positive control and samples were detected at 167bp specific for type E. The result was confirmed by DNA sequencing in the available PCR products (Figure 1 and 2). A correlation between HBsAg detection by BioELISA and HBV genotyping by nm PCR revealed that all samples positive by BioELISA (9 samples) and the 11 samples positive by AxYM were also positive by nm PCR. HBV DNA was detected in 12 samples that were HBsAg negative by both techniques.

It was also found that tested subjects, either HbcAb positive or negative, were infected with the same genotype. Sequence comparisons of 565 nucleotide (genome position: ~155-720) obtained from HBV S gene from Egyptian strain (a positive control chronic hepatitis patient) EGYAZC P1P2P4 and HBV/E gene sequences retrieved from the Gene Bank revealed only five nucleotide sequences (No 186, 201, 473, 515, 521) between them. The closest sequence was from CMR936 AB194948 (Figure 1). Furthermore, alignments of 87 nucleotide sequences (genome position: ~3048-3135) obtained from preS1 genes from Egyptian HBV isolate (a sample from a vaccinated child) EGYAZV B2 265R and the Gene Bank strains indicated that there is 100% similarity with HBV genotype E PW6 EU239226 (Figure 2).

**DISCUSSION**

**Evaluation of r-HBsAg vaccination in Egypt**
Studies on vaccinated children during infancy and early childhood in countries with a high endemicity of chronic HBV infection have shown that more than 50% of participants had measurable anti-HBs levels of at least 10 mIU/L from 4 to 10 years after vaccination.[36,37]

The main finding in this study was that 93% of children aged 2-4 years had responded to the vaccine, compared with 80% of children aged 4-13 years, and 79% of adults. In the present study, adults had a higher rate of high-titre antibodies (45%) than children (age 4-13 years) (20.5%), because the last vaccine dose in 90
adult volunteers was administered within a year before their inclusion in the study, six adults had their last dose between one and four years previously, and four volunteers had had their last dose six years or greater prior to the study. However, the children had received their doses since their birth, consequently they had had the vaccine administered between the ages of 4 to 13 years. It is well known that HBsAb level declines by time. The percentage of undetectable HBsAb level was lowest (7%) in 2–4 years old vaccinees, and rose to 20% and 21% as the age of the vaccinee increased by the time of vaccination. This is comparable to McMahon et al., 2005[38] who found HBsAb response in 89% of the study population with 19% between 2 and 9.9 mIU/L and 70% had levels greater than 10 mIU/L. In the current study, 83.6% had detectable HBsAb, but only 36.7% of them had levels of $\geq$ 10 mIU/L. In the study of Zanetti et al., 2005[39], more than 60% of children and nearly 90% of recruits maintained protective HBsAb, and recorded undetectable concentrations in about 9% of children and 4% of recruits and detectable amounts lower than 10 mIU/L in 27% and 7% respectively, more than 10 years after vaccination. In Egypt, El-Sawy and Mohamed (1999)[40] tested the post-vaccination seroprotection rate in sera collected during one month (93.3%) and 5 years (53.3%). In Taiwan, the HBsAb was detected post-vaccination in 100% of 2 year-olds, and in 75% of 6 year old children[41]. In a study from Taiwan, they noted that a single dose of vaccine boosted the immune response in almost all individuals. The results led to the suggestion that booster doses may be necessary.
in seronegative subjects for at least 15 years after neonatal immunization. They believed that this applies to both hyper-endemic and low endemic areas of the world[12].

**Breakthrough HBV infection**

In the present study, detection of markers of HBV infection in the form of HBsAg was 0% among children aged 2< 4 years, 2% among children aged 4 -13 years and 6.66% among adults, as established by the ELISA method. Similarly Alam et al (2007)[13] found in their study that the frequency of HBV infection in the Pakistani population was higher in individuals aged from 20-40 years. In the present study, none of the high HBsAb- positive individuals were positive for HBsAg, supporting a previous observation that an HBsAb titer greater than 10 mIU/mL can be considered protective[14-15].

Population-based studies of HB immunization after 10-15 years follow-up showed a reduction in chronic HBsAg carrier prevalence from high (8% or greater) to low (< 2%) endemicity in immunized cohorts of infants[18]. The current results are comparable to the Shatat et al (2000)[17] study in Alexandria, Egypt, in which only one child out of 184 vaccinated 5 year old children who had received the full course of EPI r- vaccine was HBsAg positive, while El-Sawy and Mohamed (1999)[19] did not find HBsAg positive sera among 180 children with a one month to 5 year time lapse since their last dose of vaccination. Moreover, in a serosurvey in Alexandria, Reda et al (2003)[20] revealed that the number of HBsAg carriers is significantly lower among the vaccinated (0.8%), compared to the unvaccinated in 6 year old children (2.2%). All these findings, as well as the present study reflect the impact of HB vaccination in lowering the HBsAg carriage rate in Egypt, but it also raises several questions. Is the time schedule for the 2nd and 3rd dose of r- HB vaccine appropriate? How frequent is post-vaccination breakthrough HBV infection, and which genotype is associated with it?

The same observations have been recorded all over the world, in Gambia[21], in Taiwan[22], in Indonesia[23], in Senegal[24], in a hyper-endemic area in Southern Italy[25], in Chinese children[26] and in Saudi Arabia[27]. However, the current results have revealed a decrease in the titer of HBsAb as subjects grow older, associated with an increased probability of becoming infected over time. A higher rate of HBV-positive cases was observed among the non-responders, when compared to subjects who mounted an elevated level of anti-HBs IgG antibodies.

By comparison, the current results showed that the AxSYM system yielded significantly more positive results than the ELISA test with respect to the detection of HBsAg 11.39% by AxSYM in comparison to 2.5% by ELISA. This finding may be due to the emergence of mutant HBV that could not be detected by ELISA or false positive AxSYM results as noted previously[28].

It was found that 1.38% of the studied groups had HBeAb, and 0.83% had HBeAb. Since serological data were not obtained either before or after vaccination, it is impossible to conclude whether these individuals were already infected at the time of vaccination or whether they had been subsequently infected with the hepatitis B virus.

Considering viremia, in the present study, 22 (6.11%) of all participants were positive for HBV DNA; 4 cases of group III were HBsAg positive by both methods, and 18 cases of group II (5 of them were HBsAg positive by BioELISA 11 by AxSYM and two were HBeAb positive), so 12 participants had occult HBV infection. Similarly, McMahon et al (2005)[30] found that all detected cases of HBV DNA were HBsAb negative. In China, a higher result of HBV viremia (36% of the vaccinated one year old children) was reported[31].

In the current study, HBV-DNA/HBsAg positive children may be either born to an HBV-positive mother or infected with an HBV mutant. It was noticed in study of Karthigesu et al (1999)[32] that vaccinated children may show serological evidence of breakthrough infections, particularly if they had a low HBsAb titer. They recorded that single-point mutation at nucleotide 421 of the S gene is associated with such breakthrough infections. It was recorded also by Coleman et al (2006)[33] that a child remained both DNA/HBsAg positive for > 12 years, despite having a protective HBsAb titer against the wild type virus that had a substitution mutation of glycine to arginine at HBsAg aa position 145.

The HBV genotype E recorded in this study has not previously been reported in Egypt; the most prevalent genotype in Mediterranean, Middle East and Egypt is the genotype D[34-35]. Sequence comparisons of the obtained Egyptian strain EGYAZV P1P2P4 and HBV/E gene sequences retrieved from the Gene Bank revealed that there were only five nucleotide sequence differences between them, the closest sequence was from CMR936 AB194948. Also alignments of 87 nucleotide sequence obtained from preS1 genes from the Egyptian HBV isolate EGYAZV B2 265R and the Gene Bankstrains indicated that there is a 100% similarity with HBV genotype E. The closest sequence was from CMR936 AB194948. The same conclusion was arrived at by Mulders et al (2003)[36], who reported that HBV genotype/E has low sequence diversity throughout the expanses of the HBV/E crescent, which covers almost 6000 km from Senegal to Angola. This suggest that it has a short evolutionary history in humans, and is incompatible with the evolution from the closest human virus genotype D. Transmission during childhood is supposed to be the most common mode of infection in Africa, and most children infected before the age of 6 mo become chronic carriers[37]. Early age of infection and high probability of chronic carrier status results in a high rate of transmission[38]. It was speculated from this study that the presence of this genotype in Egypt for the first time may be due to virus mutation in the “a” determinant that causes this vaccine escape mutant infection. Similarly, in Argentina they found that HBV genotype E was detected in two Argentinean sisters; one of them had been vaccinated against HBV[39].

**Conclusions and recommendations**

HBV breakthrough infection was induced by a novel HBV genotype (E) with respect to that reported in Egypt (geno-
type D). The Hepatitis-B vaccine appears to be efficient in controlling HBV infection in children and adults. It was noticed that the HBsAb level decreases by age, with increased liability to get infected, and that those with undetectable HBsAb also had a higher rate of infection. Further studies are needed to evaluate the spread of this genotype in Egypt. Furthermore, the need for and timing of a booster dose should be studied (by whom and when?).

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COMMENTS

Background

The apparent prevalence of hepatitis B virus (HBV) infection in Egypt has decreased after the Expanded Program of Immunization (EPI) vaccination program; however, the frequency of asymptomatic HBV carriers in response to the vaccination program needs to be determined.

Research frontier

Studies on vaccinated children during infancy and early childhood in countries having high endemicity of chronic HBV infection have shown that more than 50% of participants had measurable anti-HBs levels of at least 10 mIU/L 4 to 10 years after vaccination. Population-based studies in Alexandria, Egypt, of HBV immunization after 10-15 years follow-up, showed a reduction in chronic HBsAg carrier prevalence from high (8%) or greater) to low (< 2%) endemicity in immunized cohorts of infants. These same observations were recorded all over the world, in Gambia, in Taiwan, in Indonesia, in Senegal, in a hyper endemic area in Southern Italy, in Chinese children and in Saudi Arabia.

Innovations and breakthroughs

Considering viraeemia and breakthrough infections, it was observed that there is a decrease in the titer of HBsAb as age progresses, with an increased probability to become infected over time. Similar results were recorded by McManus et al. in China. Kato et al and Karthiagesu et al all detected cases of positive HBV DNA in those who were HBsAb negative than in those who had HBsAb. The genotype of HBV in this study was genotype E, which has not previously been reported in Egypt: genotype D is the most prevalent in Mediterranean and Middle East and Egypt. Similarly in Argentina they found the HBV genotype E in two Argentinean sisters; one of them had been vaccinated against HBV.

Applications

Further studies are needed to evaluate the spread of the HBV genotype E in Egypt. The Hepatitis-B vaccine appears to be efficient in controlling HBV infection in both children and adults, so it is recommended that it should be given to the high risk groups all over the country. It was noticed that HBsAb level decreases with age leading to increased liability to get infected, and that those with undetectable HBsAb have a higher rate of infection, so the need for a booster dose should be studied (to whom and when?). The number of HBsAg positive samples by AxSYM was higher than that of the BioELIS test. This raises the question of whether they were true or false positives? Further studies using different kinds of ELISA tests are needed to confirm or deny this observation.

Peer review

This is an interesting study that investigated HBV vaccine efficacy in Egypt. It is readable and publishable.
Abusahdy EAE et al. HBV genotype E among vaccinated Egyptians

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