SEROLOGIC RESPONSES OF KOREAN SOLDIERS SERVING IN MALARIA-ENDEMIC AREAS DURING A RECENT OUTBREAK OF PLASMODIUM VIVAX

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Abstract. Anti-Pv200 antibody levels were assessed in samples from endemic areas of Plasmodium vivax malaria in the Republic of Korea (ROK), using an indirect enzyme-linked immunosorbent assay (ELISA) method. Asymptomatic carriers of P. vivax were detected using nested polymerase chain reaction (PCR) of blood samples. Anti-Pv200 antibody levels in 20 vivax malaria patients (optical density ± standard deviation [OD ± SD] values 1.85 ± 0.29 of IgG isotype and 1.33 ± 1.33 of IgM isotype) were markedly higher than those of uninfected, malaria-naïve controls (0.08 ± 0.16 of IgG isotype and 0.04 ± 0.04 of IgM isotype). Antibody levels for 7 out of 8 soldiers with a recent malaria infection were sustained above the cut-off values for 4 months after successful treatment. Analysis of serum collected from 40 healthy, asymptomatic soldiers who had a P. vivax malaria attack within 3 months after our sampling, revealed 11 antibody-positive samples (27.5%), compared to 5 positive samples (12.5%) collected from a random selection of 40 soldiers. Among a larger pool of 1,713 soldiers who had served in high-risk areas for P. vivax transmission, 15% were antibody positive. Among 1,000 blood samples from asymptomatic soldiers who had served in the high-risk areas, 4 samples (0.4%) were parasite positive, as determined by nested PCR. Our results show that anti-Pv200 antibody levels can provide useful information in the late diagnosis of P. vivax malaria infection in a previously naïve population and also in large seroepidemiologic studies. Furthermore, our results suggest that asymptomatic P. vivax carriers could be important in the current outbreak of malaria in Korea.

INTRODUCTION

Although previously known as an endemic area of Plasmodium vivax, the Republic of Korea (ROK) had been considered free from malaria parasites because no reports of autochthonous transmission had been documented since 1984. However, since re-emergence of a single case of P. vivax malaria in 1993, the number of malaria cases has increased exponentially in the northwestern part of the ROK (northern Kyunggi Province), causing more than 1,700 cases in 1997, and an estimated 4,000 cases in 1998.1–4 Historically, Korean P. vivax malaria has an atypically long incubation period (lasting up to a year or longer) in a large proportion of patients but a typically mild clinical course5—characteristics which appear to apply to the current epidemic. Serologic responses to malaria parasite infections in Korea have not been reported previously, although serology would be a logical approach to detect infection during the long incubation period. The indirect enzyme-linked immunosorbent assay (ELISA) has been developed to detect and monitor antibodies against cell extracts, circumsporozoite (CS) antigen, or merozoite surface protein 1 (MSP 1) of Plasmodium spp.5–10 The indirect ELISA method could be a simple, inexpensive approach for diagnosis of asymptomatic carriers, as well as for the seroepidemiologic study of a large number of populations.

In this study, we measured antibody levels against Pv200 antigen, the MSP 1 homolog in P. vivax (Sal 1 strain), and against P. vivax CS protein by indirect ELISA of serum from infected and uninfected ROK Army soldiers who had served in high-risk areas of P. vivax transmission. We also searched for the presence of circulating P. vivax parasites by performing nested PCR on subjects’ blood. In this report, we demonstrate that the anti-Pv200 antibody levels might provide useful information in diagnosis and prognosis of the population exposed to Korean P. vivax malaria, as well as in large seroepidemiologic studies. We were also able to detect asymptomatic P. vivax carriers by nested PCR.

MATERIALS AND METHODS

Subjects and sample collection. This study conducted by the Korean Armed Forces Central Medical Research Institute was approved and reviewed annually by the Research Review Board in the ROK Armed Forces Medical Command. All participating soldiers were informed of the study objectives and on-going threats of Korean P. vivax malaria; subjects provided written personal histories and signed informed consent for their participation prior to sampling.

In late May 1997, samples for serum and whole blood were collected by venipuncture from 1,000 randomly selected Korean soldiers who had served in areas of northern Kyunggi Province, Korea, with high risk of P. vivax transmission during the summer of 1996. These soldiers were still stationed in the P. vivax malaria-endemic areas at the time of sampling in May 1997, but sometime after the autumn of 1996 they had left these high-risk malarious areas to other service locations where the risk of P. vivax transmission was much lower. According to the report by the ROK Armed Forces Medical Command, 40 of the 1,000 soldiers sampled in May 1997, were later diagnosed and treated for P. vivax malaria. The emergence of these cases was within 3 months after our sampling.

For the second round of our large seroepidemiological survey, we collected serum from 1,713 Korean soldiers, beginning in early January 1998, who had served in high-risk areas of northern Kyunggi Province since the summer of 1997. Of these, 8 soldiers who had a previous history of P. vivax malaria could be sampled twice more during the subsequent 5 months.
Negative control sera from malaria-naive subjects were collected from 113 soldiers in the Korean Armed Forces Medical School and from 200 civilians (kindly provided by the Korean Red Cross). None of the 313 subjects who donated negative control sera had a previous history of malaria or temporary residence in malaria-endemic areas.

Positive control sera were obtained from 20 malaria patients who had been admitted to ROK military field hospitals of northern Kyunggi Province and had been confirmed by examination of peripheral blood film smears during the summers of 1997 and 1998.

**Indirect ELISA.** The 19kDa fragment of Pv200, merozoite surface protein 1 (MSP 1) of *P. vivax* (Sal 1 strain), was expressed in the yeast *Saccharomyces cerevisiae*. The expression, purification, and characterization of this antigen have been described in detail elsewhere. V20, CS capture antigen containing 20 tandem repeats of predominant, also known as Pv210, nonapeptide (GDRAA/DGQPA) from *P. vivax* CS protein, was kindly provided by Dr. Robert A. Wirtz (Centers for Disease Control and Prevention). Titers of antibodies against Pv200 and V20 were determined by an indirect ELISA procedure with minor modifications from Kaslow and others, and Wirtz and others. Briefly, 96-well microtiter plates (Costar, Cornning, NY) were coated with 0.1μg of antigen in phosphate buffered saline (PBS) at 4°C overnight. The plates were washed with PBS containing 0.05% Tween20 (Sigma, St. Louis, MO) (PBST), blocked with 5% goat serum (Sigma) in PBST at 37°C for 1 hr, and then washed with PBST to remove unbound proteins. Test sera diluted 1:100 were added to the wells and incubated at 37°C for 1 hr. Then the plates were washed with PBST and incubated with horseradish peroxidase-conjugated, anti-human IgG or IgM antibodies (Sigma) for another 1 hr. The bound antibodies were detected following incubation with 100 μl of O-phenylene diamine (OPD) substrate (Sigma) using a microplate reader at 490 nm wavelength. The results were reported in units of optical density (OD) ± standard deviation (SD). Every anti-Pv200 ELISA experiment included at least 3 negative naïve individuals and 3 positive patient samples. Each serum sample was tested at least 3 times, except in the case of the high numbered seroepidemiologic survey where only positive samples were confirmed. Virtually identical ELISA results were obtained from each sample tested throughout the study.

**DNA template preparation and nested PCR.** To detect blood stage parasites of *P. vivax*, nested PCR amplification was applied according to the method of Snounou and others. To prepare the *P. vivax* DNA template, frozen whole blood samples containing EDTA were thawed at room temperature and diluted in 10 volumes of PBS with 0.05% saponin (Sigma). The parasites were released from erythrocytes at room temperature for 30 min and collected by centrifugation (6000 x g for 5 min). The supernatant was discarded; the parasite and leukocyte pellet was resuspended in 25 μl of PCR buffer without MgCl2 and the mixture was incubated at 95°C for 10 min. The first round of nested PCR amplification was then performed with a 5 μl aliquot of supernatant from this mixture solution. For the first round of PCR amplification, two *Plasmodium* genus-specific primers, rPLU6 (5’-TAA AAA TTG TTG CAG TTA AAA CG-3’) and rPLU5 (5’-CCT GTT GCC TTA AAC TTC-3’) were used for amplification of small subunit ribosomal RNA (ssrRNA) genes. The second round of amplification was performed with a 1 μl aliquot of product from the first round of PCR, using species-specific primers of rVIV1 (5’-CGC TTC TAG CTT AAT CCA CAT AAC TGA TAC-3’) and rVIV2 (5’-ACT TCC AAG CCG AAG CAA AGA AAG TTC TTA-3’) to produce a *P. vivax*-specific 120 bp fragment of ssrRNA genes. All PCR reactions were performed in 2 mM MgCl2, 50 mM KCl, 10 mM TrisCl (pH 8.3), 0.1 mg/ml gelatin, 125 μM of each of the four dNTPs, 250 nM of each oligonucleotide primer, and 0.4 unit of AmpliTaq polymerase (Perkin Elmer Cetus).

**RESULTS**

**Antibody levels in control group and malaria patients.** To determine the cut-off values of anti-Pv200 antibody levels and to evaluate the usefulness of the indirect ELISA method for anti-Pv200 antigen in diagnosing *P. vivax* malaria in Korea, anti-Pv200 antibody levels were measured in sera of 313 normal individuals who had never been exposed to malaria parasites and 20 patients ill with active malaria. As shown in Figure 1, mean anti-Pv200 levels of IgG and IgM isotype antibodies in normal control sera were OD ± SD of 0.08 ± 0.16 and 0.04 ± 0.04, respectively. Based on these results, OD values of 0.5 and 0.2 (mean + 3 × SD) were chosen as the lower limits of positive reactions for IgG and IgM ELISA results, respectively (these cut-off values exclude >97% of the control group). Anti-Pv200 antibody levels of 20 Korean *P. vivax* malaria patients had mean values of 1.85 ± 0.29 for the IgG isotype and 1.33 ± 0.29 for the IgM isotype (Figure 2). All of the patients’ sera were above the cut-off values, except for one case that was below the cut-off for IgM.

**Changes in antibody levels with a history of malaria.** Serum samples from 8 individual soldiers with recent illness
FIGURE 2. Anti-Pv200 antibody levels in *Plasmodium vivax* malaria patients determined by indirect enzyme-linked immunosorbent assay (ELISA). The thick solid horizontal line and dotted horizontal line represent the cut-off values of optical densities (OD) in IgG isotype (0.5) and IgM isotype (0.2) antibodies, respectively. The thin horizontal lines indicate the mean values of OD in each anti-Pv200 antibody titer.

from *P. vivax* infection were collected at 4, 6, and 9 months after recovery following treatment with chloroquine/primaquine. The anti-Pv200 IgG antibody levels were sustained above the limit of positive values for 4 months in all but one of the soldiers tested, and for 6 months in 3 of the soldiers tested (Figure 3).

**Elevated antibody levels before malaria attack.** According to the report by the ROK Armed Forces Medical Command, 40 of the 1,000 soldiers sampled in May 1997, had an attack of *P. vivax* malaria within 3 months after our sampling. Since these soldiers left the high-risk malarious area after autumn 1996, we assumed that their illnesses were caused by infection acquired during the summer of 1996. Anti-Pv200 antibody levels of these 40 soldiers were compared to the levels of 40 randomly selected soldiers from the pool of 1,000 (Figure 4). Eleven (27.5%) of the serum samples were positive for anti-Pv200 IgG and IgM in the 40 soldiers who subsequently experienced a malaria attack, while only five (12.5%) sera had positive values among the 40 randomly selected soldiers. The value distribution of anti-Pv200 antibody levels for the remaining 920 soldiers was virtually identical (12.7% positive for anti-Pv200 IgG and IgM, data not shown) to that of the 40 randomly selected soldiers.

**Antibody levels in high risk areas.** In late January 1998,
we collected 1,713 serum samples from soldiers who had been serving since the summer of 1997 in areas with high malaria transmission rates. Of all the samples tested, 15% were positive for anti-Pv200 IgG or IgM (Figure 5).

**Antibody levels against *P. vivax* CS protein.** Antibody levels against *P. vivax* CS protein were tested with V20 antigen in four groups shown in Figure 6: A) 20 malaria patients; B) 12 soldiers within 6 months after malaria attack; C) 40 soldiers who had malaria within 3 months after sampling; and D) 40 soldiers who did not have a malaria attack within 3 months following our sampling. The mean IgG antibody titer of group A patients was 0.40 ± 0.76, which was not significantly higher than those for the other groups (B, 0.15 ± 0.37; C, 0.24 ± 0.09; and D, 0.06 ± 0.09) (Figure 6A). Likewise, the mean IgM antibody level for group A was not significantly higher (0.37 ± 0.53) compared to those for the other groups (B, 0.03 ± 0.53; C, OD 0.01 ± 0.02; and D, 0.02 ± 0.02) (Figure 6B).

**Detection of *P. vivax* in blood.** Among 1,000 whole blood samples collected in late May 1997, *P. vivax* parasites were detected in 4 soldiers by nested PCR (Figure 7). These soldiers had no clinical signs or symptoms of malaria at the time of our sampling.

**DISCUSSION**

The present serologic study clearly demonstrated that distinct anti-Pv200 antibody responses occur in the course of *P. vivax* infection experienced during the recent malaria epidemic in the northwestern part of the ROK. There were clear differences in anti-Pv200 antibody responses between patient and non-patient groups. These results suggest that anti-Pv200 antibody detection by ELISA may be useful in diagnosing Korean *P. vivax* malaria in suspected patients. During the course of our study, we demonstrated the existence of distinct anti-Pv200 antibody responses in Korean soldiers serving in high-risk *P. vivax* malaria areas.
cence of elevated levels of anti-Pv200 antibodies from several suspected patients who had not yet shown *P. vivax* on blood film smears, but who later showed positive results of *P. vivax* on blood film smears as malaria symptoms progressed. Further studies using larger sample sizes may be required to firmly establish the sensitivity and specificity of the method. One possible confusion in the interpretation of anti-Pv200 antibody titers is that some individuals may maintain high anti-Pv200 antibody titers for an extended period after recovery from malaria, and even after complete clearance of the parasite. According to the data from the prospective longitudinal study of 8 soldiers, most individuals with Korean *P. vivax* malaria maintain high anti-Pv200 antibody titers of the IgG isotype for 4 to 6 months after drug cure (Figure 3). It is possible that some individuals maintain high anti-Pv200 antibody titers for longer periods. The IgG isotype of anti-Pv200 antibodies had higher normal background levels (Figure 1) but a more consistent high response to infection (Figure 2) than did the IgM isotype. Although there is a danger of false negatives, the combination of both IgG and IgM isotype levels against Pv200 antigen could detect Korean patients infected with *P. vivax*, either in addition to, or as a replacement for traditional laboratory diagnosis.

One of the intriguing findings in the present study was that 27.5% of 40 soldiers who later developed symptomatic malaria had elevated anti-Pv200 antibody levels up to 3 months prior to illness (Figure 4). This implies that the antibody titers against Pv200, or MSP 1 antigen of *P. vivax* might have increased in the very early erythrocytic stage of infection, if not in the hepatic stage. All 40 soldiers were probably carrying *P. vivax* during a prolonged incubation period at the time of our sampling and could have been infected with *P. vivax* during the previous summer since all of the subjects had served in areas of high risk *P. vivax* transmission in the summer previous to our sampling, but were stationed in low-risk *P. vivax* areas thereafter. In support of this conclusion, the North Korean (NK) strain of *P. vivax*, characteristically, has a long incubation period in many patients. This phenomenon of a long dormancy has been widely and consistently observed in the course of the *P. vivax* malaria outbreak in the ROK. According to the epidemiologic survey conducted by the ROK Armed Forces Medical Command from 1996 through 1998 (Roh C-S and others, unpublished data), more than 60% of malaria patients in the ROK Armed Forces became infected with *P. vivax* during the summer of the previous year and became ill sometime after the winter and spring that followed. During these prolonged incubation periods of up to 1 year or longer, no apparent clinical signs and symptoms were observed. On the other hand, the rest of the malaria patients had short incubation periods typical of *P. vivax* malaria in many other parts of the world. Those malaria patients with a prolonged incubation period might play a very important role in maintaining infectious sources of *P. vivax* malaria for the following summer. Hence, the early detection of malaria patients in a prolonged incubation period could be crucial in trying to eradicate malaria from Korea. Anti-Pv200 antibody ELISA could be useful in the early detection of Korean *P. vivax* malaria in a certain portion of the population. Anti-CS antibody levels in the various groups showed a broad range of values. Since the antibody titers in each group had diverse distributions, no clear cut-off values could be defined. CS protein can be classified into two groups according to the central 20 nonapeptide repeats: predominant type I repeats with a GDRADQQPA sequence and variant type II repeats with an ANGAGNQPG sequence. The use of different *P. vivax* CS proteins appears to be regionally biased. Therefore, the CS protein of *P. vivax* in the recent outbreak of malaria in the ROK is likely to have the sequence homologous to the NK strain (predominant type I repeats). In our results, however, antibody responses against CS protein with predominant type I repeats appeared to show no significant differences between patients and other groups (Figure 6). More than 50% of patients showed markedly lower anti-CS protein antibody titers (OD < 0.10). In addition to the poor antibody responses against CS protein of *P. vivax* after a prolonged incubation period, the possibility that those patients with low antibody titers against CS protein with type I repeats might have been infected with *P. vivax* strain carrying CS protein with variant type II repeats cannot be excluded.

The survey of 1,713 soldiers in high-risk malaria areas by anti-Pv200 antibody ELISA demonstrated that at least 10% of soldiers appeared to have been exposed to the malaria antigen. This ELISA test could be used in seroepidemiologic studies as an efficient tool for monitoring malaria prevalence and incidence in large cohorts.

As a result of the nested PCR of blood samples, we detected 4 soldiers (0.4%) carrying *P. vivax* parasites who had no clinical signs and symptoms associated with *P. vivax* malaria at the time of our sampling. A few weeks later, 1 of those 4 soldiers was admitted to the field hospital for *P. vivax* malaria. The remaining 3 soldiers were also ordered to be treated with curative drugs for *P. vivax* malaria. Although anecdotal, these results indicate that asymptomatic blood carriers of *P. vivax* parasites are present in the Korean population.

The current epidemic of *P. vivax* malaria in Korea had an insidious beginning with only a single case detected in 1993. The medical community initially suggested that transmission in the ROK was due to mosquitoes flying across the border from the Democratic People’s Republic of Korea (DPRK). However, although the majority of infection had occurred in soldiers stationed along the DPRK border, it was soon apparent that there was extensive transmission in the ROK civilian community. Without active surveillance, some civilian cases will inevitably go undetected because some people are either asymptomatic or suffer only mild symptoms. Though parasitemic, these infected people will not seek treatment and will therefore contribute to expansion of malaria parasite foci in Korea. Given the extremely long incubation period of *P. vivax* in some individuals, infected people are more likely to travel throughout the peninsula before seeking treatment. Extensive geographic movement of the parasite has already occurred in American soldiers who carried the infection from Korea to the United States. Since the resurgence of epidemic malaria after a virtual absence of *P. vivax* transmission in the ROK for the last 20 years, we have a unique opportunity to study the serology of Korean *P. vivax* malaria in young subjects, most of whom have never experienced the threat of malaria before. To stop
the current epidemic, a combination of vector control (Strickman D and others, unpublished data), personal protection, active case detection, and treatment are likely to be necessary. Considering the challenge of case detection caused by a long incubation period and the sometimes mild symptomatology of *P. vivax*, the anti-Pv200 ELISA and nested PCR could be valuable tools in a program to systematically eradicate malaria from the Republic of Korea once again. These techniques could have been useful in assessing the extent of malarial resurgence earlier in the epidemic.

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