Abstract

Background: In Turkey, varicella vaccine was introduced into routine childhood immunization in 2013, with a single dose administered to children aged 12 months. However, there is limited information on the morbidity (incidence and seroprevalence), mortality and burden of disease of varicella in the overall Turkish population.

Aim: To determine varicella seroprevalence and its social determinants in Manisa Province, Turkey in children aged > 2 years before single-dose varicella vaccination was introduced in 2013.
Methods: The presence of anti varicella-zoster virus IgG antibodies was determined using enzyme-linked immunosorbent assay in serum samples collected from 1250 participants.

Results: The overall seroprevalence was 92.8% and the seroprevalence was > 90% among all age groups except 2–9 years (55.7%). Seroprevalence was significantly associated with family size, annual per capita equivalent income, number of people per room and education level. After adjusting by age, only education level remained significantly associated with seroprevalence, reflecting the early age effect.

Conclusion: High seroprevalance depends on natural exposure to the infectious agent itself and is not associated with social determinants. High vaccine coverage should be maintained for effective varicella control and switching to a 2-dose schedule may also be considered to reduce the number and size of outbreaks in the Turkish population.

Keywords: seroprevalence, social factors, Turkey, vaccination, varicella

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disease caused by varicella-zoster virus (VZV), which has no animal reservoir (1,2). While varicella occurs predominantly in childhood in temperate climate regions, it is also a disease of people aged 20–25 years in subtropical and tropical climate regions, with a lower overall incidence (2–4). Although varicella is often mild and self-limiting in healthy children, some complications such as secondary bacterial infection of the skin, viral or bacterial pneumonia, encephalitis and cerebellitis can be life threatening especially in adolescents, adults and immunosuppressive individuals (2,5–7).

In 1998, the World Health Organization (WHO) recommended routine vaccination for countries where varicella is an important public health and socioeconomic problem, if high vaccine coverage could be provided and it were economically viable (8). In some countries such as the United States of America (USA), Greece, Republic of Korea, Canada and Saudi Arabia, varicella is included in the routine childhood immunization programme. In some countries such as England and Belgium, the vaccine is administered to seronegative individuals after childhood and at high risk for varicella (9,10). In developing countries, as the burden of other diseases is higher than that of varicella, introduction of varicella vaccination into the national immunization programme is not a high priority. For these countries, a targeted vaccination strategy may be an option (2,8).

In Turkey, varicella vaccine was introduced into the routine childhood immunization programme in 2013, with a single dose administered to children aged 12 months. However, there is limited information on the morbidity (incidence and seroprevalence), mortality and disease burden of varicella in the overall Turkish population (11–22). Furthermore, varicella was only included in the notifiable diseases list in Turkey in 2011. Therefore, we aimed to determine varicella seroprevalence in the population aged > 2 years in Manisa Province and to examine some of the social determinants for varicella seroprevalence. This reflects the population before the introduction of varicella vaccination into the routine childhood immunization programme in Turkey in 2013.

**Methods**

This cross-sectional study was conducted as part of the population survey titled “Determination of the seroprevalence of some vaccine preventable diseases in Manisa, 2014”. Manisa is located in Western Anatolia and has a population of ~1.3 million. In terms of socioeconomic development, Manisa was ranked 23rd among 81 provinces of Turkey in 2011 (23).
using the EpilInfo computer programme (Centers for Disease Control and Prevention, Atlanta, USA), assuming anticipated seronegativity of 2.0% and absolute precision of 0.75%. The estimated sample size was then inflated by 30% to reach 1740 to overcome nonresponse bias. The sample was selected using a simple random sampling method from the Manisa Province Family Medicine Information System. Individuals who were selected for the study were invited to the family health centres by physicians or midwives. Data were collected from participants in family health centres between 18 March and 22 June 2014.

Ten interviewers were trained and standardized for data collection. Written informed consent was obtained from the participants, and questionnaires were completed by the interviewers during face-to-face interviews. Serum samples were collected in the family health centres, transported to a centre in Manisa at 4–8°C and stored at −20°C. Once weekly, samples were sent to the laboratory at −20°C to be stored at −80°C until they were tested. The study protocol was approved by the Dokuz Eylul University Clinical Research Ethics Board.

Among 1740 people who were sampled, 168 could not be reached in their place of residence, 312 refused to participate in the study and 10 participants’ blood samples were excluded from the analyses due to haemolysis or insufficient quantity. We analysed data from 1250 people (response rate: 71.8%).

The dependent variable of the study was varicella seropositivity. The independent variables were gender, age groups, area of residence, family size, number of people per room, annual per capita equivalent income, education level and employment status. The presence of specific varicella antibodies in serum samples was determined using Euroimmun (Medizinische Labordiagnostika AG, Lübeck, Germany) anti-VZV IgG ELISA test. Test results with categorical variables were summarized with count, percentage and 95% confidence interval (CI) and compared using the χ² test in univariate analysis. Continuous variables were summarized with median, minimum and maximum values. Crude and age-adjusted odds ratios and 95% CIs for social determinants of varicella seropositivity were calculated using logistic regression analysis. All statistical analyses were performed using SPSS version 15.0 (SPSS Inc., Chicago, IL, USA).

Results

The study group was 52.1% female and 47.9% male, with a median age of 36 (2–89) years. Varicella seroprevalence was 92.8% (95% CI: 91.2–94.1%) among all the participants. The seroprevalence changed between 92.6 and 100.0% in children aged > 9 years but it was 37.1 and 61.9% for the 2–4- and 5–9-year age groups, respectively (P Figure 1). There was no
significant difference in seroprevalence according to gender (P = 0.37), area of residence (P = 0.57) and employment status (P = 0.83) (Table 1). Seroprevalence was significantly increased in participants whose family size was ≤ 4 (OR = 1.75, 95% CI: 1.12–2.72); participants who had ≤ 1 person per room (OR = 2.32, 95% CI: 1.49–3.62); and participants in the 3rd and 4th quartiles of annual per capita equivalent income (OR = 1.94, 95% CI: 1.03–3.65 and OR = 2.21, 95% CI: 1.14–4.27). These associations disappeared after age adjustment. There was also a significant difference in varicella seroprevalence according to education level. Seroprevalence was > 95% for participants at all education levels, apart from those who were not of school age (38.1%) or attending primary school (74.2%) (P

Discussion

Varicella seroprevalence was 92.8% among people aged > 2 years in Manisa in 2014. The study population did not receive varicella vaccine in the national immunization programme since it was not introduced for children aged 12 months until 2013. Therefore, our findings reflect the seroepidemiology of varicella before the vaccine was introduced as part of the national immunization programme. The seroprevalence was > 90% among all age groups except 2–9 years, which was 55.7%. This suggests that varicella exists in the population and almost 90% of the population contract the virus by age 10 years.

In most temperate climates, > 90% of the population become seropositive for varicella before adolescence and the incidence ranges from 13 to 16/1000 per year (2). In a comparative seroepidemiology study in 11 countries in Europe between 1996 and 2003, varicella seroprevalence was > 90% in children aged 10–14 years, except in Italy (81.7%) and England and Wales (89.7%) (24). There have been several previous epidemiological studies on varicella in Turkey that included different age groups (11–22). In 9 provinces of Turkey in 2002, the seroprevalence was 77.8% in the population aged

It is known that the most important risk factors associated with severity and mortality of varicella are older age and a compromised immune system. Varicella case fatality rates are about 1 per 100 000 people in children and the risk of death is 4 times higher in infants than in children and 23–29 times higher in adults. Average crude varicella mortality rates range from 0.3 to 0.5 per million people annually. It is estimated that varicella leads to 4.2 million severe complications annually that require hospitalization and 4200 deaths globally (26). In developed countries, overall case fatality rates are 2–4 per 100 000 cases compared to 1–3 per 1000 cases for measles (2,26). Despite the routine use of measles and pertussis vaccination, the age-standardized death rate per 100 000 cases in 2010 was lower for varicella (0.1) than measles (1.7) and pertussis (1.1) (26). According to WHO recommendations, varicella should be considered an important public health problem that requires routine vaccination (8). A few studies in Turkey have evaluated the financial burden and mortality of varicella in children and
adults (18,19). In a multicentre study from 2008 to 2010 that included 824 children aged

Varicella is one of the most contagious diseases. The basic reproduction number (R0) estimates for varicella ranged between 3.3 and 16.9 in serum banks that were collected between 1996 and 2003 in 11 European countries, and herd immunity thresholds varied from 70 to 94% (24). This means that if the varicella-susceptible proportion in a population is over this threshold, an outbreak is not expected to occur in that population (24–27). However, we currently have no R0 and herd immunity threshold estimation for the Turkish population. WHO recommends that resources should be sufficient to ensure reaching and sustaining varicella vaccine coverage for ≥ 80% of the population. If vaccine coverage remains

We found that varicella seroprevalence was not associated with the social factors that were explored, except for education level, which actually reflects the early age effect. Our expectation was to demonstrate higher exposure to VZV and higher varicella seroprevalence in disadvantaged groups, for example, people with lower education, lower income per capita or crowded housing. However, only the group who was not of school age or at primary school had low varicella seroprevalence, suggesting that exposure to varicella increased with schooling. Some studies have reported that living in urban areas, living in a large family (≥ 4 people) and children with more siblings are factors associated with higher varicella seroprevalence. Many studies have found no gender difference for varicella seroprevalence (2, 3, 16, 17, 28–30). Consistent with the current study, varicella seroprevalence was not associated with household size, income, occupation and education in a population-based study in Izmir in 2009–2010 (16).

The current study had some strengths. First, the study group was randomly sampled from a general population and had a high response rate, hence it was representative of all age groups > 2 years in Manisa Province. The study reflected the epidemiology of varicella before the introduction of routine varicella vaccination, which can provide invaluable baseline information for evaluating the impact of the vaccination programme. Although some socioeconomic and primary healthcare quality variations existed between the provinces, the findings may give some indicators about the whole country since our study group was a random sample from Manisa.

Our study also had some limitations. First, vaccination histories were not taken from the participants, and some may have received varicella vaccine in private clinics before the vaccination programme started. However the number of such people should not be a large proportion of the study participants. Second, the overall response rate was 72% and we evaluated the age and gender structure of responders and nonresponders. There was no significant difference between the 2 groups by age but there were more female than male participants in the study. However, there was no significant gender difference in seroprevalence; therefore it can be considered that the overall seroprevalence was not greatly affected by this limitation. Third, since there were fewer participants in the subcategories according to social determinant variables, the statistical power may have been lower to determine the difference in seroprevalence between those subcategories. For example, the literate group had only 42
participants and the unemployed group 68.

In conclusion, since the study included age groups before starting routine varicella vaccination in Turkey, the observed varicella seroprevalence depended on natural exposure to VZV and was not associated with social determinants. For successful varicella control by routine vaccination programme, vaccine coverage and effectiveness, burden of varicella and change in seroepidemiology in Turkey should carefully be followed up. High vaccine coverage should be targeted and maintained. Otherwise, varicella outbreaks affecting older age groups, with more severe clinical features, are inevitable. To reduce the number and size of varicella outbreaks, as well as severe morbidity and mortality, switching to a 2-dose vaccination schedule may also be considered for the national immunization programme in Turkey.

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Séroprévalence et déterminants sociaux de la varicelle en Turquie
Résumé

Contexte: En Turquie, le vaccin contre la varicelle a été introduit dans le programme de vaccination systématique des enfants en 2013, avec une seule dose administrée aux enfants de 12 mois. Toutefois, les informations concernant la morbidité (incidence et séroprévalence), la mortalité et la charge de morbidité de la varicelle dans l’ensemble de la population turque sont limitées.
**Objectif**: Déterminer la séroprévalence de la varicelle et ses déterminants sociaux dans la province de Manisa (Turquie) chez des enfants de plus de deux ans avant l’introduction de la vaccination en une seule dose en 2013.

**Méthodes** : La présence d’anticorps IgG dirigés contre le virus varicelle-zona (VZV) a été déterminée à l’aide de la méthode immuno-enzymatique (ELISA) sur des échantillons de sérum collectés auprès de 1250 participants.

**Résultats** : La séroprévalence globale de la varicelle était de 92,8 % et la séroprévalence était supérieure à 90 % au sein de tous les groupes d’âge, excepté chez les 2-9 ans (55,7 %). La séroprévalence de la varicelle était associée à la taille de la famille, au revenu annuel équivalent par habitant, au nombre d’individus par classe et au niveau d’éducation.