The Value Of Clinical Parameters In Cytomegalovirus Infection Among Antibody Sero- Positive Blood Donors In A Nigerian Teaching Hospital

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INTRODUCTION

Cytomegalovirus (CMV), otherwise called human herpes virus type 5 (HHV5), is one of the human herpes viruses that may present with diverse signs and symptoms.

Available literature suggest that such systemic viral infections may exhibit a spectrum of clinical manifestations that range from completely asymptomatic to most severe forms of clinical presentation. Clinical features attributable to CMV can be congenital or acquired. Congenital infections manifest with intrauterine growth retardation (IUGR), jaundice, hepatosplenomegaly, thrombocytopenia and encephalitis with or without microcephaly. In the neonate, CMV is now known to be the commonest cause of congenital infections, affecting around 0.3-1% of all live births. About 5-10% of congenitally infected infants have symptoms at birth, while fatal disease occurs in 20% of these infants. With postnatal infections and other forms of acquired infections, the incubation period (IP) is thought to be 3-12 weeks. Primary CMV infection in the postnatal period is usually mild or asymptomatic but can occasionally be accompanied by the syndrome of infectious mononucleosis that is symptomatically indistinguishable from EBV- induced infectious mononucleosis with malaise, fever of up to 39-40°C, chills, sore throat and headache as the typical presenting features. Lymphadenopathy is however uncommon in CMV induced mononucleosis. Primary CMV infection in the immunocompromised patients is far less likely to be asymptomatic and commonly presents with spiking pyrexia, gut and retinal involvement and pneumonitis which cause grave prognosis with 80-90% mortality. Diagnosis of CMV should be suspected if a patient has symptoms of infectious mononucleosis but has negative test results for Infectious Mononucleosis and EBV or has negative heterophile antibody test. Hence, like any other clinical case, history and physical examination should serve as relevant tools in accurate diagnosis of CMV infection. Given that CMV is a transfusion transmissible viral infection, laboratory physicians should be interested in the clinical profile of CMV antibody seropositive blood donors. More so because clinical evaluation by way of
history taking and physical examination constitute a critical part of the initial sorting and assessment of prospective blood donors. It is against this backdrop that this study emanated.

MATERIALS AND METHODS

Two hundred consenting prospective blood donors who reported to the blood bank of the Jos University Teaching Hospital, (JUTH) Jos from October 2006 to December 2006, were recruited into this prospective study. Their blood samples were screened for the presence of CMV antibodies. Excluded from the study were pregnant females and donors with language barrier. All tests were done using kits manufactured by “Dialab” Austria (www.dialab.at). The kit is based on ELISA principles and technique. The manufacturer’s instructions as well as procedures were strictly adhered to or followed. All donors who tested positive for CMV antibodies proceeded with the study while negative donors were excluded at this stage. Detailed history and thorough physical examination were carried out on all CMV seropositive donors with a view to eliciting signs and symptoms that would normally indicate the presence of an infectious and systemic disease. Standard thermometers used to measure both buccal cavity and armpit or axillary body temperatures in order to objectively determine clients’ febrility or otherwise. This stage of the study was observational and results were described as numbers and percentages. Ethical approval was obtained from the Research and Ethical Committee of JUTH. Informed consent was obtained from all participants. The data were analyzed using Epi info computer software version 3.3.2.

RESULTS

A total of 200 prospective blood donors who presented for bleeding were screened. One hundred and eighty four donors tested positive for Cytomegalovirus antibodies, representing a prevalence rate of 92 %.( table I).The age range of the study population was between 19 and 55 years, with a mean of 37 years. The age distribution and CMV antibody seropositivity among donors is shown in table I. Of the 184 donors who are positive for CMV, 97.8 % (=180 donors) were males while 2.2% (=4 donors) were females, the two groups were however merged due to the number of females which was rather small for a separate statistical consideration. CMV antibody prevalence was highest in the 25-29 years age group (30.4%) and lowest in the extreme age groups (1.6%). Figure 1 shows a standard (calibration) curve, plotted to determine the CMV antibody status of donors. Donors with antibody levels in excess of 0.5 iu/ml were regarded as seropositive. The clinical findings in CMV antibody seropositive donors were recorded in table II. One hundred and sixty five donors (90%) were asymptomatic while 5% had fever with the average temperature of 38.3°C. Five seropositive donors (2.7%) presented with sore throat, while 4 donors (2.2%) had fatigue. Only one donor (0.5%) had a history of previous blood transfusion. With respect to physical signs on examination, One hundred and seventy two donors (94%) had no demonstrable physical signs. Pallor was found in 5.4% of donors. None of the donors presented with skin rash. There was no evidence of significant lymphadenopathy or hepatosplenomegaly in any of the donors.

DISCUSSION

The high proportion of donors (90%) who were asymptomatic and who revealed no clinical abnormality on complete physical examination (94%) confirms the common impression that CMV infection is usually asymptomatic especially among immune competent persons. The results are therefore not surprising. Studies conducted among immunocompromised patients indicate that it is such patients that are more likely to present with the classic symptoms of cytomegalovirus infection. Other authors have documented similar patterns of asymptomatic presentation of CMV infection although not necessarily among blood donors. These observations indicate the opportunistic tendency of cytomegalovirus. Most of the other papers cited by us showed that other herpes and non-herpes viral infections may present with diverse signs and symptoms. The possible reasons for this difference in clinical behavior may be difficult to clearly highlight now. However, bearing in mind that clinical manifestations emanate from underlying pathophysiology, differences in virus induced pathophysiological mechanisms may be a plausible factor. The results of this study show that most seropositive blood donors are asymptomatic. These donors were predominantly aged between 20 and 55 years and represent a largely “well” segment of the general population and should mirror closely the overall pattern of CMV presentation in the general population. The implication of this is that the general population will need
appropriate health education and public enlightenment on this seemingly clinically benign virus. More especially, immunocompromised individuals will need to be informed that they are particularly at risk of fatal complications and outcomes from CMV. There is therefore a need for this group of the population to know their CMV status, and insist on CMV negative blood units whenever transfusion is indicated. In analyzing the results of this study, it will be noted that there was no control group. The study design was essentially an observational study, with a descriptive design. Further analytical studies utilizing suitably selected control groups are needed to better highlight the spectrum of clinical presentation and tendencies of Cytomegalovirus.

In conclusion, this study has shown that clinical manifestations in CMV seropositive blood donors are non-specific and are therefore not of any diagnostic significance. Most seropositive donors are asymptomatic. These findings indicate that diagnosis of CMV infection anchors on laboratory parameters. Consequently, management of health institutions in general, and laboratories in particular are encouraged to put in place adequate facilities for CMV diagnosis.

References

18. “Dialab” Austria http://www.dialab.at
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