Platelet antigens in varicella associated thrombocytopenia

J Winiarski

Abstract

Serum IgG, or predominantly, IgM antibody binding to electrophoretically separated normal platelet membrane protein antigens were detected by immunoblotting in five children with thrombocytopenia associated with varicella. Glycoproteins GPIb, GPIIb, GPIIIa, and other 25-260 kilodalton (kDa) proteins were identified as target antigens, suggesting a transient autoimmune mechanism causing the thrombocytopenia.

Purpura is a well known complication of varicella but the mechanisms causing thrombocytopenia have not been fully elucidated. Thrombocytopenia may be caused by either immune mediated platelet destruction or direct viral interaction with megakaryocytes or platelets. Circulating platelet binding antibodies and increased platelet associated immunoglobulins have been described, but the antigenic specificity of the putative platelet binding antibodies has remained unknown. Specific antibodies to defined platelet antigens in thrombocytopenia associated with varicella are reported here.

Patients and methods

Blood was drawn from three boys and two girls, aged 1 to 11 years, within a week after onset of purpura, that presented three to seven days after the eruption of a varicella exanthem. All children were previously untreated and in good health. Nadir platelet counts ranged from 2-7 x 10^9/1. A bone marrow examination was performed only in case 3 who, after seven days of purpura, had slightly decreased megakaryocytes. Platelet counts remained at 1 x 10^9/1. Three children who had a CD 45+ cell count of > 150 x 10^9/l within 20 days. Serum samples were obtained from four of the children one to seven years later. Sera from 24 healthy blood donors were used as negative controls and anti-PIIIA sera as positive controls. Separation of platelet membrane glycoproteins using sodium dodecylsulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting were performed as described elsewhere. Briefly, pooled normal platelet membranes were dissolved in SDS and applied to a slab gel together with molecular weight standards. SDS-PAGE was performed under non-reducing conditions using the discontinuous buffer system of Laemmli. The separated proteins were electrophoretically transferred from gels to nitrocellulose membranes and blocked in fat free milk. Membrane strips were incubated with patient or control sera, diluted 1:25. After several washes the blots were then stained with alkaline phosphatase conjugated rabbit anti-human IgG and then labeled with protein A or rabbit anti-human IgM and then stained after addition of substrate for the detection of glycoprotein bound antibodies. Glycoproteins were identified on parallel blots by alkaline phosphatase conjugated Lens culinaris lectin which binds to platelet glycoproteins Ib, IIb, and IIIa. To confirm the specificity for platelet surface antigens, sera were absorbed with fresh platelets and used in parallel with unabsorbed sera. Sera were also screened with a solid phase platelet membrane enzyme linked immunosorbent assay (ELISA). All positive results were confirmed on repeat testing.

Results

By immunoblotting, platelet glycoprotein binding IgG was detected in sera from three patients and IgM in five patients. IgM corresponded in part to the IgG bands, but additional reactions were noted. The bands comigrated with GPIb (170 kDa), GPIIb (140 kDa), and GPIIIa (95 kDa) in four patients. Other uncharacterized 25-260 kDa bands were also seen (figure and table). Absorption of sera with packed platelets eliminated antibody binding to glycoproteins Ib, IIb, IIIa and to the other proteins indicating specificity for native surface antigens. By ELISA, four children were shown to have membrane binding antibodies. In sera collected one to seven years later, the earlier positive reactions were mostly weak or no longer detected (table). In case 4, IgG platelet
ELISA and immunoblot assays for serum antibodies to platelet antigens in four children with varicella associated purpura and in controls

<table>
<thead>
<tr>
<th>Case No</th>
<th>Serum sample taken*</th>
<th>Presence (+) or absence (-) of platelet membrane binding antibodies in serum</th>
<th>Antibodies to SDS-PAGE separated platelet membrane proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>During</td>
<td>-</td>
<td>IgG: 38, 52 (25)</td>
</tr>
<tr>
<td></td>
<td>After (1)</td>
<td>+</td>
<td>IgM: 95</td>
</tr>
<tr>
<td>2</td>
<td>During</td>
<td>-</td>
<td>IgG: 25, 38, 52 (25), (38)</td>
</tr>
<tr>
<td></td>
<td>After (3)</td>
<td>-</td>
<td>IgM: 95</td>
</tr>
<tr>
<td>3</td>
<td>During</td>
<td>ND</td>
<td>IgG: 25, 38, 52, 170, 260</td>
</tr>
<tr>
<td></td>
<td>After (4)</td>
<td>-</td>
<td>IgM: 95</td>
</tr>
<tr>
<td>4</td>
<td>During</td>
<td>+</td>
<td>IgG: 25, 38, 52, 170, 260</td>
</tr>
<tr>
<td></td>
<td>After (7)</td>
<td>-</td>
<td>IgM: 95</td>
</tr>
<tr>
<td>5</td>
<td>During</td>
<td>+</td>
<td>IgG: 95, 140, 38, 52, 95</td>
</tr>
<tr>
<td>Controls (n = 24)</td>
<td>-</td>
<td>-</td>
<td>IgM: 38</td>
</tr>
</tbody>
</table>

* Sera were collected during thrombocytopenia and in the convalescent period (No of years after thrombocytopenia).
†ELISA.
‡Bands detected by immunoblotting; molecular weights of target antigens in kDa. Parentheses denote a weak reaction.
ND = not done.

IgG and IgM antibodies to SDS-PAGE separated platelet membrane glycoprotein antigens as detected by immunoblotting in patient and control sera. Apparent molecular weights are given in kDa. Lanes a, b, and c show platelet antibodies of IgG class and lanes d, e, and f of the IgM class. Lane a was incubated with PPV antisera. The PFU alloantigen is situated on GPIIIa (95 kDa). Lanes b and e, normal sera; c and d, case 5. The 140 kDa band in lane c comigrated with GPIIb. Lane f, case 2.

membrane antibody was detected in the ELISA but not by immunoblotting. IgG-F(ab')2 binding to GPIIb/IIIa could be confirmed as previously described.

Discussion

Various mechanisms underlying the virus associated thrombocytopenias have been proposed. Non-immunological factors such as viral interaction with platelets or megakaryocytes have been suggested when purpura presents in the viraemic phase of varicella, while the 'postinfectious' purpura would more likely be immune mediated. Increased megakaryocyte counts may, however, be found in patients with thrombocytopenia presenting early during the exanthematous phase. Feusner et al reported shortened platelet survival, increased megakaryocyte counts, and platelet binding antibodies in serum and on platelets in four children with varicella associated thrombocytopenia, indicating an immune mediated platelet destruction.

IgG antibodies directed to platelet membrane glycoprotein antigens have been shown in chronic idiopathic thrombocytopenic purpura, confirming the autoimmune nature of that disease. Few data have been available about the pathogenesis of acute idiopathic thrombocytopenic purpura. Recently antibodies to platelet membrane glycoproteins have been observed, and also binding via the IgG F(ab')2 part to the GPIIb/GPIIIa complex in patients with acute idiopathic thrombocytopenic purpura. The present finding of IgG and predominantly IgM antibodies directed to platelet surface glycoproteins, including GPIb, GPIIb, and GPIIIa, during active disease suggests that they may have a role as a thrombocytopenic factor in these varicella patients. The presence of HLA or alloantibodies is unlikely in the untransfused children. In ELISA, with a different antigen presentation, platelet membrane antibodies were detected in four patients. The course of thrombocytopenia might be connected with the primary IgM antibody response to an infection. Varicella zoster virus appears in the blood five days before the onset of a rash. Elicited antiviral IgG is detectable in serum three days after exanthem, by which time the viraemic phase is concluded. An early thrombocytopenia could be mediated by IgM class platelet cross reactive antibodies or antibodies produced by unspecifically stimulated autoreactive B lymphocytes. It is concluded that the detection of platelet antigen reactive autoantibodies strengthens the concept of a transient autoimmune platelet destruction in children with chicken pox induced thrombocytopenia.
Is a specialist paediatric diabetic clinic better?

S Bloomfield, J W Farquhar

Abstract
Diabetic control in 88 children attending three
general paediatric clinics was compared
prospectively over one year with that of 89
children attending a specialist paediatric
diabetic clinic. Glycated haemoglobin (HbA1)
concentration and days admitted were signifi-
cantly lower in the group attending the
specialist clinic. This has implications for the
organisation of paediatric diabetic services.

Diabetic control in many children is not as good
as it could be. It has been suggested that better
diabetic control is achieved in children attend-
ing specialist paediatric diabetic clinics, but
few objective studies have been done. We have
collected, over a one year period, data about
diabetic control in children attending a special-
ist paediatric diabetic clinic in a children's
hospital compared with those attending paedia-
tric clinics in three district general hospitals.

Methods
The diabetic clinic at the Royal Hospital for
Sick Children (RHSC) acts as a primary referral
centre for children under 16 years in south east
Scotland. It is staffed by three paediatricians
two consultants and one senior registrar), one
adult diabetologist who facilitates gradual trans-
fer of adolescents to the adult clinics, a dietitian,
a full time nurse specialist, and a dental
hygienist. There is 24 hour access by telephone
to medical or nursing staff for advice and the
specialist nurse visits at home as required.

Three general paediatric clinics (A, B, and C)
in district general hospitals in central Scotland
provide care for diabetic children who are seen
by a consultant paediatrician or paediatric
registrar. A dietitian is available in all these
clinics, and a diabetes nurse specialist who is
shared part time with the local adult clinics is
sometimes available.

Information about children under 13 years of
age on 1 October 1985, and with diabetes of
more than three months duration, was obtained
prospectively for one year as part of a research
project concerning diabetes education. Data
included a medical and social profile, methods
of diabetic care, diabetic events, anthropo-
metric measurements, and measurement of
glycated haemoglobin (HbA1) at each visit. (All
blood samples for HbA1 were analysed at RHSC
by a Corning electrophoretic method; the
normal reference range is 4.7–7.9%.) Samples
were analysed within one week and remained
stable.

Data from the three district general hospitals
and the RHSC clinic were compared by χ2 or
Kruskal-Wallis tests as appropriate, and asso-
ciations between quantitative and ordinal
variables were tested by Kendall rank correla-
tion.

Results
Comparisons were made at the end of one year
(a) between the three general paediatric clinic
populations, and (b) for all these three clinics
combined (if there was no significant difference
between them) with the RHSC paediatric
diabetic clinic. Age, duration of disease, age at
diagnosis, the number of boys, and social class
distribution were similar (table 1). The average
time spent with the paediatrician at each clinic
visit was 25 minutes at RHSC and approxi-
mately 15 minutes in the general clinics.

Children attending the clinic at RHSC were
admitted to hospital for significantly fewer days

<table>
<thead>
<tr>
<th>Table 1 Characteristics of diabetic children &lt;13 years of age attending paediatric clinics in district general hospitals (A, B, C) and a specialist paediatric diabetic clinic (RHSC)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Mean (SD) age (years)</td>
</tr>
<tr>
<td>Mean (SD) duration diabetes (years)</td>
</tr>
<tr>
<td>Male/female</td>
</tr>
<tr>
<td>No in social class I and II</td>
</tr>
</tbody>
</table>

There was no significant difference between groups for any variable.
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