

A MONOCLONAL ANTIBODY TO VIII:C PRODUCED BY A MOUSE HYBRIDOMA. H.P. Muller, N.H. van Tilburg, R.M. Bertina, J. Derks and E. Klein-Breteler. Haemostasis and Thrombosis Research Unit, Department of Medicine and Department of Human Genetics, Leiden University Hospital, Leiden, The Netherlands.

Spleen cells of a Balb-c mouse immunized with VIII:C (isolated by affinity chromatography) were fused with mouse myeloma cells (MOPC-21 derivative). After the fusion 12/32 wells produced an inhibitor to VIII:C. After subcloning (3 x) a stable hybridoma line was obtained. The antibody in the supernatant was detected with a modified VIII:C inhibitor technique. The supernatant of in vitro cell cultures of the hybridoma cells contained anti-VIII:C titers (Bethesda) of about 0.3-1.0 units/ml. Injection of the hybridoma cells in pristane pretreated Balb-c mice results in anti-VIII:C titers of 5,000-10,000 units/ml ascites.

Analysis of the produced immunoglobulin demonstrated the presence of one band after isoelectric focussing, which contained heavy chains both of IgG<sub>1</sub> and IgG<sub>2B</sub> subclass. Because of the unusual kinetics of the monoclonal antibody with VIII:C extensive characterisation of the nature of its VIII:C neutralising properties was necessary.

The monoclonal antibody does not bind <sup>125</sup>I-fibrinogen or isolated VIIIIR:AG, it reacts with isolated VIII:C and can be used in a two-site immunoradiometric assay for VIIIICAG. The epitope against which the antibody is directed is not present on 'serum-VIIIICAG'.

ANTIBODIES AGAINST PLATELET MEMBRANE GLYCOPROTEINS: EFFECT ON RISTOCETIN-INDUCED PLATELET AGGREGATION. E.F. Ali-Briggs, C.S.P. Jenkins and K.J. Clemetson. Departments of Hematology, Wilhelmina Gasthuis, Amsterdam, and Montefiore Hospital, Bronx, NYC, and Theodor Kocher Institute, Berne, Switzerland.

Some membrane glycoproteins (GPs) have been isolated by lectin-affinity chromatography and antibodies towards them have been raised. Platelets that have lost glycofocalicin no longer respond to ristocetin-human VIII:WF, bovine VIIIIR:WF, or to anti-glycofocalicin or anti-GPs Ia and Ib antibodies but are still agglutinated by anti-GPs IIb and IIIa antibodies. Anti-GPs Ia and Ib and anti-glycofocalicin antibodies, IgG and Fab' fragments inhibited ristocetin-human VIIIIR:WF- and bovine VIIIIR:WF-induced aggregation of fixed, washed platelets and of platelets in plasma while anti-GPs IIb and IIIa antibodies were without effect.

Crossed immunoelectrophoretic studies showed that glycofocalicin was present on whole platelets in only trace amounts; anti-glycofocalicin antibodies, however, recognized a slower migrating component. Platelets incubated in an EDTA-free medium no longer respond to ristocetin-human VIIIIR:WF. Membranes isolated from such platelets contained glycofocalicin which cross-reacted with a remnant of the slower migrating component. Anti-GPs Ia and Ib antibodies gave more complex patterns but it was possible to identify the slower moving component recognized by the anti-glycofocalicin antibodies.

These results show that glycofocalicin is not normally found as such on whole platelets but is present as a precursor which is most likely GP Ib. On degradation of this precursor, glycofocalicin is released from the membrane and VIIIIR:WF-receptor activity is lost.