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.

Splean 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 subclonation (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, and IgG, 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 ¹²⁵I-fibrinogen or isolated VIIIR:AG, it reacts with isolated VIII:C and can be used in a two-site immunoradiometric assay for VIIICAG. The epitope against which the antibody is directed is not present on 'serum-VIIICAG'.

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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 glycocalicin no longer respond to ristocetin-human VIII:WF, bovine VIIIR:WF, or to anti-glycocalicin 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-glycocalicin antibodies, IgG and Fab' fragments inhibited ristocetin-human VIIIR:WF- and bovine VIIIR:WF-induced aggregation of fixed, washed platelets and of platelets in plasma while anti-GPs IIb and IIIa antibodies were without effect.

Crossed immunoelectrophorectic studies showed that gly-cocalicin was present on whole platelets in only trace amounts; anti-glycocalicin antibodies, however, recognized a slower migrating component. Platelets incubated in an EDTA-free medium no longer respond to ristocetin-human VIIIR:WF. Membranes isolated from such platelets contained glycocalicin 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-glycocalicin antibodies.

These results show that glycocalicin 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, glycocalicin is released from the membrane and VIIIR:WF-receptor activity is lost.