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PLATELET SIZE, NOT PLATELET MASS, DETERMINES INTRINSIC KINETIC DIFFERENCES IN PLATELET RECRUITMENT INTO AGGREGATES FOR ADP, U46619, AND PAF, BUT NOT FOR RISTOCETIN. Truman Wong and Mony M. Frojmovic. Department of Physiology, McGill University, M. Frojmovic. Department of Ph Montreal, Quebec, Canada H3G 1Y6.

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Previous studies of platelet aggregation using resistive counting methods (PA) have suggested a dependence on platelet size (\bar{v}), but have not been evaluated for varying platelet number (N_0) and associated total platelet mass. Here, the relationship between \bar{v} , N_0 and function was examined in size dependent human subpopulations fractionated by counterflow centrifugation. The original platelet population and three size dependent platelet fractions were concentrated and resuspended into autologous citrated platelet poor plasma at varying N_0 for 5 donors. The initial rate and sensitivity of PA were determined generally at 3-5 seconds following ADP/ristocetin addition. Extent of PA was determined at 10 seconds. At similar N_0 (180 \pm 50 x 10 3 μ 1), large platelets (L; \bar{v} = 7.4 \pm 0.3 fl; 16 \pm 4% of total population) were two-fold more sensitive and more rapidly recruited into both PA and turbidometrically measured macroaggregates (TA) in response to ADP than the smallest platelets (S; \bar{v} = 4.6 \pm 0.4; 16 \pm 5%). Aggregation kinetics and sensitivity for the mid-sized platelets (\bar{v} = 5.9 \pm 0.3; 31 \pm 7%) were intermediate between the large (L) and small (S) platelet fractions. When platelet counts were adjusted to yield similar total platelet mass (N_0 x \bar{v}), these differences persisted for PA, but not for TA. Subsequent studies were all made for platelet suspensions at similar mass. Maximal rates of ADP-induced shape change were comparable for L vs. S platelets. Significant differences in the initial rate and maximal extent of PA between the size-dependent fractions were also seen for a stable PGH analogue (146619) and platelet activating factor PA between the size-dependent fractions were also seen for a of PA between the size-dependent fractions were also seen for a stable PGH, analogue (U46619) and platelet activating factor (PAF). Most platelets were maximally recruited into micro-aggregates (60-80% PA) for all sized fractions. Kinetics and sensitivity for ristocetin-induced agglutination were comparable between the different sized fractions. The above size-dependent differences in aggregation for physiological activators appear to arise from intrinsic membrane/cell biochemical differences, not observed for ristocetin-von Willebrand (Factor VIII)-induced agglutination.

AGGREGATION RESPONSE OF PLATELETS DURING INHIBITION OF PHOSPHOLIPASE A₂. R.S. Labow (1). E. Meek (1). G.A. Adams (1.2).G. Rock (1.2.3). Ottawa Centre, Canadian Red Cross, Blood Transfusion Service (1), Depts. of Biochemistry (2) and Medicine (3), University of Ottawa, Ottawa, Ontario, Canada

Arachidonic acid (AA) is liberated from platelet membrane phospholipids during stimulation and promotes cellular aggregation phospholipase during simulation and pholocos centular aggregation through the synthesis of thromboxane A₂. Two pathways; phospholipase A₂ (PLA₂) or phospholipase C (PLC) followed by the action of acylglycerolipases, are thought to be activated during platelet stimulation and supply the necessary AA. We have platelet stimulation and supply the necessary AA. We have reported that mono (2-ethylhexyl)phthalate (MEHP), a physiological metabolite of the plasticizer di(2-ethylhexyl)-phthalate (DEHP), commonly used in a variety of medical devices and storage containers, inhibits PLA₂, but not PLC in platelet lysates. The effects of MEHP on intact platelets were studied. PLA2 activity in intact platelets or lysates was assayed by incubating them with 2-l⁴C-arachidonyl-phosphatidylcholine and measuring formation of free l⁴C-arachidonic acid in 10 min. Platelet lysates hydrolyzed 10% of the substrate while 2.6% was hydrolyzed by intact platelets. The amount of MEHP needed to inhibit ¹⁴C-AA liberation was 0.35 mM for platelet lysates and 0.7 mM for intact platelets. Platelet aggregation induced by collagen was inhibited by MEHP (1 mM), although responses to adenosine diphosphate, AA and ionophore were unaffected. Identical effects on platelet aggregation were found when indomethacin (0.1 mM) was added. Higher concentrations of MEHP blocked platelet aggregation induced by adenosine diphosphate or AA but not ionophore or synergistic pairs of these stimuli, indicating a more generalized membrane disruption at higher MEHP concentrations. These results suggest that MEHP acts in a similar manner to indomethacin to block PLA2-mediated liberation of arachidonate during platelet aggregation (supported by MRC and NHRDP, Canada)

CHANGES IN PLATELET HALF-LIFE, SENSITIVITY PROSTANOIDS AND AGGREGATION INDUCED IN THE DOG BY BODY HYPOTHERMIA. L.M. Cunha-Ribeiro, S. Cunha, T. Brandão, F. S. Gonçalves, A. Almeida-Dias and J.M. Pina-Cabral. Haemostasis Center (INIC) and Dept. of Physiology, Porto Medical School, Porto, PORTUGAL.

During body cooling, there is a progressive thrombocytopenia, which is reversible after rewarming and is not prevented by previous treatement with aspirin, ticlopidine or prostacyclin. In this work in order to evaluate if hypothermia induces alterations of platelet function we studied, in the dog, the platelet aggregation (PA) and the inhibitory action of PGE1 and of a stable prostacyclin before analog, Hoprost, hypothermia(37°C.) rewarming(37°C.).Platelet half-life was also studied in another group of dogs before induction of hypothermia and after rewarming and recuperation of the animals. PA has been evaluated by platelet counting in whole blood.Platelet half-life was estimated by serial determinations of MDA following administration of aspirin.PA induced by ADP (30µM) decreased 40% after rewarming(n=8). Platelet sensitivity to PGE1 (35 nM -1.4 µM f.c.) and Iloprost (7nM-172nM f.c.) was also decreased after rewarming: inhibition index 2.08±1.082 versus 1.19±0.362 (n=8; p<0.01) and 2.48±1.250 versus 1.10±0.227 (n=8; p<0.005) respectively.Platelet half-life increased after hypothermia from 3.99±0.730 days to 4.48±0.846 days(n=8 p<0.05).In the control group(n=6) platelet half-life determined twice with one week interval, did not change significantly.

We conclude that body hypothermia decreases platelet reactivity to ADP; renders platelets less sensitive to the inibitory effect of prostanoids and increases platelet half-life. These results are probably due to alterations in platelet membrane induced by cooling and rewarming the animals.

ABNORMALITIES OF PLATELET AGGREGATION AND ENHANCED FACTOR X AC-TIVATOR ACTIVITY OF WASHED PLATELETS IN SICKLE CELL D.A.F. Chamone (1), A.Y. Hoshikawa-Fujimura (2), C. Massumoto (1), G. Bellotti (3), F. Arashiro (2) and M. Jamra (1). Department of Internal Medicine, Division of Hematology (1), Faculty of Pharmaceutical Sciences (2) and Institute of Heart Disease Massumoto (3), São Paulo, Brazil

The occurence of microvascular occlusion is one of the most prominent pathologic features of sickle cell anemia. The mechanism of vaso occlusion has generally been attributed to the ab-normal shape and reduced deformability of the sickled erithrocy tes. However, the involvement of vascular endothelium, plate lets and their interactions with coagulation factors may also be of pathogenic significance in microvascular occlusive crises.

We investigated the interaction between vascular endothelium, platelets and blood coagulation factors in 23 with Sickle Cell Disease (SCD) and in normal volunteers.

Factor X activator activity in washed platelets was performed according to Semeraro and Vermylen (1977), thromboxane B₂ (TXB2) and 6-keto-PGF1α were determined using specific radioimmunoassays..PAF-acether from platelets was determined according to Chignard et al (Nature, 1979, 279:799). Platelet aggregation was performed with a Chrono-Log Aggregometer (Model 440) on platelet rich plasma (PRP) using the Born method. Prostacy clin release from endothelium was performed according to Mon-

cada et al (Lancet i:18, 1977).

Our results showed that platelets from patients with SCD ha we enhanced factor X activator activity (p < 0.0001), produce more PAF-acether than controls (p < 0.02) and showed hyperaggregability in these patients as compared to normal volunteers (p < 0.00001).

We concluded that platelets from homozygous sicklers have enhanced factor X activator activity as well as increased capacity for PAF-acether production. These abnormalities may contri bute to the incidence of vaso occlusive crises in these pati-