NEW TRENDS IN CURRENT IMPLEMENTATION OF RHEOLOGICAL TECHNOLOGIES FOR THE HUMAN BLOOD

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During the last ten years the number of the in vivo ex vivo techniques in haemorheology have considerably increased and in spite of a consensus meeting held in 1986 (International Committee for Standardization in Haemorheology, ICSH, 1986) little improvement can be noticed at the present time.

It appears that the number of proposed tests remains high, as many of them are obsolete and/or no longer commercially available. Cataloguing the different tests, coming out of all rheological laboratories all around the world, is extremely difficult. One can however distinguish the public techniques (viscosity, viscoelasticity, red cell aggregation, deformability), which can be obtained from manufacturers and the private or "wild" techniques only settled inside research laboratories.

Of importance is: viscosity, filtration, ektacytometry, micropipettes and in vitro flow systems (Tableau I).

TABLE I
THE METHODS IN HAEMORHEOLOGY

1- Public Techniques

Viscosity
Plasma Viscosity
Coulter Capillary Viscosimeter, Coulter Electronics
Capillary tube Viscosimeter, Schott A G
Whole Blood Viscosity
Contraves LS 30 (Couette type Viscosimeter)
Brookfield microcone plate Viscosimeter, model LVT (Brookfield Engineering Lab. Inc., USA)

- Viscoelasticity
OCR-D: oscillating capillary rheometer, PAAR, Graz (Viscoelasticity and flexibility)

- Red cell aggregation
Mini-erythrocyte aggregometer : MA1, Myrenne GmbH
Erythroaggregometer Affibio, Regulex
Laser-assisted Optical Rotational Cell Analyser (Mechatronics, Hoorn, NL)

- **Deformability**
  Hanss Hemorheometer
  St George's Filtrometer, Carrimed, UK
  CTA: Cell Transit Analyser, ABX Company
  Ektacytometer Technicon
  Laser-assisted Optical Rotational Cell Analyser (Mechatronics, Hoorn, NL)

2-Private (wild) Techniques

- **Viscosity**
  Cui Xizhong Cone Plate Viscosimeter Tianjin
  Capillary Viscosimeter, Shangai Medical University

- **Filtration**
  Selecting-erythrocyte Rigidometer (H Kiesewetter 1983)
  Filtrometer S JA Evans 1990
  Filtrometer AM Ehrly 1973
  Filtrometer G Fisher 1993 (WH Reinhardt 1990)
  Filtrometer Y Isogai 1981
  CPA: Cell Passate Analyzer, Hoechst AG, Wiesbaden

- **Erythrodeformeter**
  R J Rasia & G Schültz 1993, A Luquita, 1994

- **Micropipette**
  Micropipette System, Paulishke & Nash, 1993
  Micropipette System, JC Lelèvre, 1992

- **In vitro - flow systems**

The current objectives, in haemorheology, are to be precised in order that some techniques can survive, knowing that manufacturers stop rather quickly the providing of machines...

1- Monitoring and calibration

One particular point in rheology is the lack of mastered methods of calibration when the tests are let to be running on. In France only two attempts have been made in this way. The first is the comparison of Contraves LS in 5 laboratories after dispatching control viscous samples which showed slight differences. The second is a current research study about 5 Affibio erythroaggregometers in the same laboratories, which exhibit important differences for the same samples, thus constraining to perform an immediate adjustment of both the machinery and the soft system... However more extensive and subsequent studies have shown that this method is stable, well defined, related to haematocrit, fibrinogen and ESR, but nevertheless giving more accurate information (1)

It must be assessed that, up to the present, no other device, system and or method are going to be calibrated or controled in any way... The concept of "normal" plasma should be rejected, as there is no normal plasma when are considered the variations of plasma factors according to the age, the state of inflammation etc... Only a strategy where the patient appears like his own control may be acceptable, before and after a treatment, for
example, or when the measurement is plotted against another proper signal or test (see below)
Particularly important is to consider that the so called "filtration techniques" are uncalibrated, submitted to unmastered factors and therefore obviously without any signification. Only very precise experiments performed with elaborated devices as the St George's filtrometer and the CTA may be taken into some account.

2 Clinical applications
At the patient bed side, tests related to the thyxotropy phenomenon are useful, for they have been studied in relevant clinical studies. Both plasma and blood viscosity, red cell aggregation and some viscoelasticity methods enter this group. The obtained results are part of the prognosis and able to evaluate the effect of treatments (haemodilution, drugs).

One particularly interest is that they indicate the final effect of blood closed factors: i.e.: haemoconcentration, high density of the plasma, dysglobulinaemia, polycythaemia, shape of cells, increased cohesion of red cell aggregates.
Among those former factors the level of the fibrinogen is particularly important and may be different along with the age of patients, the season and the country ...(2)(3)(4)(5)(6).

The filtration techniques which are largely undetermined, do not appear useful in the literature, on the clinical point of view. A lot of changes in patients: i.e.: diabetes, sickle cell anaemia, atherosclerosis have been refered, but nothing remains in practice.

3 Pharmacological studies
Again the effects of drugs on cells has been studied using the filtration methods. These have to be carefully selected for many of them are undetermined. The more recent are better: St George's, acting upon a cell suspension or the CTA and CPA analysing the cells through a single calibrated pore. In such very cautious experiments the cells may improve their "deformability" or their "filterability" under the influence of pharmacological products.

4 New trends for new devices
The micropipette technique either for white or red cells is promising and under development, but will certainly remain a specialized research test.

The in vitro flow systems, mimicking microcirculation, are interesting: they allow studies dealing with the behaviour not only of the circulating cells, but also of the endothelial wall barrier (cultured human or animal endothelium). Recent advances in this field have shown that the properties of the microvessels are, by many aspects, shear-dependent at the level of the wall and submitted to the up-regulation of receptors and ligands recently defined. The use of inhibitors and monoclonal antibodies allow experiments to be performed, creating a better understanding of the action of rheological factors at the level of the microcirculation.

5 What could be the future in haemorheology?

a Physiology & pharmacology
The haemorheological methods will remain useful in animal experiments and in human, in order to reach a better knowledge of the microcirculation for a given function of organ. Furthermore the influence of drugs can be analyzed in such techniques, using blood, plasma and cells.

b At the patient bedside
When an interest in rheology is to be found out for patients, only the thyxotropic related methods are able to provide this. In pre-operative periods, before and after haemodilution, haemorheology is useful, as well as in neonates.
One can also investigate the importance of rheological data in performing the comparison with other biological or functional parameters. For example red cell aggregation correlated with TC-PO2 investigation in ischemic patients (7). This latter result appears extremely interesting and incentive for a future development of haemorheological investigation at the patients bed side.

REFERENCES


