Correspondence

Modification of simple and specific test for measuring lipid peroxides in plasma

In our paper, we used and recommended an isodometric (spectrophotometric) test to measure lipid peroxides in human plasma samples. The use of a commercially available colour reagent (CHO-iodide; Merck, Germany) made the technique very simple and easy to use.

Not long after the paper had been published, we and others have had difficulties performing the test as it was described. Basically, the saline treated (control) sample resulted in higher absorbance than the colour reagent (porex) test sample. This was due to some as yet unidentified changes in the CHO-iodide reagent. The colour reagent presently available somehow quenches the original absorbance of plasma due to yellowish carotenoids. All our efforts to identify and eliminate this interference or change the protocol while measuring the formed I, at 360 nm have failed.

We then adopted a technique, which measures iodine in the presence of starch as a blue colour iodine-starch complex at 560 nm, where the natural yellowish colour of plasma does not interfere with the measurement. This modification has been in use for six months with different batches of CHO-iodide and produced reliable estimate of the overall oxidative capacity of plasma.

Reagents:
CHO-iodide (Merck, Germany).
Butylated hydroxy toluene (BHT): 0.572 mg/ml in ethanol.
Sucrose from soluble from Sigma, No. S-2630: A 1% solution is made fresh in 6-7 mM sodium chloride (39.2 mg/100 ml) by immersing into boiling water for a few minutes. After centrifugation (5 minutes at 12 000 rpm, Eppendorf centrifuge), the clear supernatant fluid is used the same day.
Procedure: Blood is taken into EDTA (1.5 mg/ml), centrifuged, and the plasma samples kept at 4°C for no longer than two days. In a small phial, place 200 µl plasma, 1:0 ml colour reagent (CHOD), 10 µl BHT and 200 µl starch solution. Mix and transfer into a disposable cuvette and read the absorbance immediately at 560 nm against distilled water. Place the cuvette immediately into dark and then read the absorbance again at 60 minutes. The difference in the absorbance represents the peroxide content of plasma. Direct calculation by using molar absorptivity is difficult due to the variation of the absorption with the source of starch. However, for one batch, the system can be calibrated by hydrogen peroxide, according to the stoichiometry of the reaction:

\[(1) \quad H_2O_2 + 2I^- + 2H^+ \rightarrow I_2 + H_2O\]

(2) \[I_2 + I^- \rightarrow \text{I}_3^-\]

Absorption changes are linear between the amounts of peroxide added (0.2-1 nmol) and the \(\text{I}_3^-\) produced.

P GOROG
Pathophysiology Unit, William Harvey Research Institute, St Bartholomew’s Medical College, London EC1M 6BJ

Parathyroid hormone related protein (PTHrP) in hypercalcemia of lymphoproliferative disease

Bolo-Deoku et al have shown that PTHrP was not implicated in a case of hypercalcemia in Hodgkin’s disease, thus expanding the very limited literature on PTHrP in lymphopoietic disorders. In a case of hypercalcemia in chronic lymphoecytic leukaemia (CLL) where the same conclusion was reached.

Eight years after presentation, a patient with stage IV C B-CLL developed acute severe low back pain due to vertebral body collapse. Generalised loss of bone density was apparent radiologically, but no discreet osteolytic lesions. Corrected serum calcium concentration was 2.94 mmol/l (range 2.26–2.60), phosphate activity 1.31 mmol/l (range 0.8–1.4), alkaline phosphatase 84 IU/l (range 30–115), and renal function was normal. Infiltrative bone resorption and increased treatment with corticosteroids failed to control the hypercalcemia and a peak corrected serum calcium of 3.29 mmol/l was reached before effective treatment with pamidronate sodium was started. Prolymphocytoid transformation and myeloma were excluded and there was no evidence of coincidental malignancy. Serum 25-hydroxycholecalciferol was 11.9 ng/ml (range 8–50), PTH 1.7 pg/ml (range 0.9–5.4), and PTH-related protein (PTHrP) was less than 0.23 pmol/l (normal).

While lymphotoksin (tumour necrosis factor \(\beta\), interleukins 1 and 6, and prosta-glandins) has a defined role in the bone disease of myelomatosis, the mechanisms underlying dysfunctional osteoclast activity in other lymphopoietic disorders are unknown. Although larger studies must be performed, it seems unlikely that PTHrP has an important role.

*SV DAVIES
IVOIRA

CAJ WARDROP
Departments of Haematology and Medicine, University of Wales, College of Medicine, Cardiff CF4 4XW


Value of PCNA and Ki67 staining in breast cancer

The failure of PCNA to reflect proliferative activity, as measured by Ki67 or mitotic counting in breast tumours, documented by Leonardi and colleagues, is disappointing. We have found a weak positive correlation between PCNA staining and Ki67 staining in both benign and malignant breast tumours (personal observation).

To some degree such observations may be related to the techniques used. Our practice with both Ki67 and PCNA is to count areas of highest antigen expression after scanning the slide at low magnification. This gives a measure of the maximum proliferative activity in any given tumour and obviates the need to decide how many areas to include or to define a concentration to include (not to mention the intermediate areas).

Using this method, we found no correlation between PCNA and mitotic count in breast epithelial tumours. While Ki67 and PCNA have been observed to be present in paraffin wax sections, with biopsy specimens and small tumours.

These factors tend to reduce the efficacy of PCNA immunostaining, but the main problem with PCNA appears to be its overexpression. It is apparent that when assessing PCNA expression that mitotic figures are frequently unapparent. This is consistent with its described expression in the S-phase under optimal conditions. Yet PCNA expression is frequently higher than Ki67 expression index as we have found, and higher than thymidine labelling in benign breast tissue. As Ki67 and thymidine labelling are restricted to the cell cycle only, PCNA must therefore persist in non-cycling cells but in a variable manner, losing its association with mitotic activity and making its correlation with Ki67 expression tenuous.

This suggests that although PCNA is to some extent comparable with Ki67 it is influenced by factors as yet undefined to such an extent that it loses its usefulness in quantifying proliferative activity.

RP SULLIVAN
M GORTIMER
Department of Pathology, University College Hospital, London, England


Dr Barbarareschi comments:

Dr Sullivan’s letter, which suggests a weak correlation between PCNA and Ki67 immunostaining in benign and malignant breast tumours, agrees with our paper where we concluded that in breast cancer PCNA immunostaining is not a substitute for Ki67...