Platelet functions in relation to diet and serum lipids in British farmers*

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SUMMARY Coagulation and platelet aggregation induced by thrombin, ADP, adrenaline, and collagen were studied in three contrasted groups, each of 20 to 22 middle-aged male farmers. Serum lipids were similar in the three groups. In the west of Scotland group, however, platelet reactivity was significantly greater than in the east of Scotland. This was associated with a dietary intake, evaluated by three different techniques, higher in saturated fat but also lower in polyunsaturated fat and alcohol. Platelet function in the southern England group also correlated with dietary fats and in addition inversely with calcium intake. On an individual basis in the 63 farmers, all the platelet function tests were significantly correlated with the intake of saturated fat regulated by that of calcium and alcohol.

The dietary effects on platelets appear to be mediated by the fatty acid composition of plasma lipids and of platelet phospholipids. In that fraction, the fatty acids 20:3ω9, 22:3ω9 and 20:4 were the most closely related to the platelet function tests. The trienoic acid 20:3ω9, identified with essential fatty acid deficiency, was also correlated with the intake of saturated fat and calcium. In this study, platelet functions were more dependent upon the dietary factors associated with coronary heart disease such as saturated fats, calcium, and alcohol than upon serum lipids.

Animal studies have shown that prolonged feeding with saturated fats predisposes to thrombosis initiated in various ways. In man, platelet function can be modified by dietary changes and studies performed on French farmers have shown that platelet responses were more closely related than serum lipids to the intake of saturated fats and the known incidence of coronary heart disease in the region.

Farmers were chosen for these studies because of their comparable life style. In addition, farmers take every meal at home and, consequently, precise evaluation of the food intake can be more easily achieved particularly since their diet does not appear to change much from day to day. The aim of the present pilot study performed in 1978 in the United Kingdom was to investigate in healthy subjects the relation between diet, serum lipids and platelet function in geographical areas with differing incidences of coronary heart disease. Farmers were chosen from three areas: south west Scotland (coronary heart disease death rate per 100 000 in men aged 35 to 44: 95); east Scotland (coronary heart disease death rate: 65), and south east England (coronary heart disease death rate: 54).

Recommendations made by the Scottish Home and Health Department Working Party were that careful dietary studies were desirable within defined populations. This study attempts to parallel epidemiological work with detailed laboratory investigations.

Subjects and Methods

Subjects
Healthy male farmers who were actively farming, and were born between 1 January 1933 and 31 December 1938, in the region, were selected. All the farmers fulfilling the above criteria, identified through Health Centre records, within a radius of approximately 10 miles from the following small towns: Kirriemuir and Friockheim in east Scotland; Stranraer, in south west
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Scotland; Denmead, in south England, were asked to participate.

Sixty-nine farmers met the criteria and were contacted by their local Health Centre. Sixty-three agreed to participate and informed consent was obtained from all subjects, 52 of whom were also owners of their farm.

The project was approved by the British Medical Association Ethical Committee.

Dietary survey
The study was undertaken using a mobile laboratory with the same personnel and instrumentation as in previous studies. Each subject, was assessed in respect of his dietary habits and his state of health and nutrition in the following ways.

(a) The 24-hour recall of diet
This was performed one or several days before the weighing technique. Both the farmer and the lady of the house were present for the interview which also included questions about smoking habits.

(b) Weighing technique and diet composite
This was performed as in previous studies. The dietitian was present on one week day, at each of the subject’s two main meals (midday and evening) and each of the food items and drinks was weighed before being eaten. An equal portion of this food was placed in a closed refrigerated container and similarly for the breakfast and any other food intake, and the subject’s spouse was asked to keep an identical portion refrigerated. Each subject’s daily food collection was mixed, ground (Waring Blender), and weighed, with two 180 g samples being taken and frozen for subsequent analysis by the Municipal Laboratory in Bordeaux (France), with the techniques already used. The food items were then coded by the dietitian before computer processing. The food table composition used was essentially the British table recently published, but with a number of meat products and fats such as margarine and dripping, being analysed for their fatty acid composition, in our laboratory.

(c) Physical examination
This was performed by a local physician, except for the anthropometric determinations (weight, height, 12 skinfolds) which were done by the dietitian. The blood pressure was determined by both the nurse of the team and the local physician.

Coagulation and aggregation tests
Before venesection (130 ml) each subject fasted and refrained from smoking for 10 hours. The following tests were performed as in previous studies: (a) The recalcification plasma clotting time of platelet rich plasma without or with (Styphen time) the addition of Russell’s viper venom (1/100,000) (Sigma Chemical Co, Saint-Louis, Mo, USA); (b) the cephalin recalcification clotting time of platelet poor plasma; (c) the platelet factor 3 recalcification clotting time which is the clotting time of washed platelets resuspended in a standard platelet poor plasma; (d) the aggregation of platelet rich plasma with a platelet count of 300,000 per μl, in response to the following agents: thrombin, adenosine diphosphate, collagen, adrenaline bitartrate as in previous studies and in addition arachidonic acid. The arachidonate solution was prepared by diluting 50 mg arachidonic acid (Sigma chemical Co) with 1:65 ml NaOH, 0:1 N, sonicating for 10 seconds, and adding 14:85 ml of complete Tyrode. The solution was distributed in small containers, frozen and before use diluted to half with Tyrode to give a final concentration in platelet rich plasma of 70 and 35 μg/ml.

The final determinations for both coagulation and aggregation were performed in a Rubel-Renaud recording aggro-coagulometer (US patent no 4.116.564), a modification of Born’s original turbidimetric technique.

Lipid analysis
Fatty acid analysis of the diet, plasma total lipids, and platelet total phospholipids were performed after lipid extraction and methylation as previously reported, by gas liquid chromatography except that the 6 foot glass columns were packed with 10% Supelco 2340 on 100/120 chromosorb W-AW and with temperature programming (150°-210° C, 2°/min). The platelet total phospholipids were separated by thin layer chromatography in one dimension on silica gel 60F–254 aluminium foils (Merck, E., Darmstadt, Germany) with the following solvent mixture: petroleum ether-diethyl ether-acetic acid (90:30:2, v/v). Total serum cholesterol and triglycerides were determined as in previous studies and the high density lipoprotein-cholesterol by the Technicon automated technique after heparin-manganese precipitation.

The statistical significance of the results was evaluated by Student’s t test, and the regression analysis, uni- or multivariate, performed by computer, according to the method of Efroyzman.

Results
In Fig. 1 are reported different variables studied in the farmers from the three regions. None of them except the cigarette consumption is significantly different from one region to another, in particular
serum cholesterol, HDL-cholesterol, and triglycerides.

From the haematological results (Fig. 2) it can be seen that the plasma clotting times and the platelet factor 3 recalcification clotting time (evaluating the platelet clotting activity) are similar at Denmead and at Kirriemuir. By contrast at Stranraer, both tests are significantly shorter (accelerated) \((p<0.001)\) than in the two other regions. The cephalin recalcification clotting time, which evaluates the activity of the plasma clotting factors, does not show any significant difference between the three regions. The Stypven-time of platelet rich plasma was 26-6±1-1 s at Denmead, 28.4±0.8 s at Kirriemuir, and 23.9±0.7 s at Stranraer (mean ± SE). The values obtained at Stranraer are significantly accelerated when compared with those of Kirriemuir \((p<0.001)\) and of Denmead \((p<0.005)\). All the tests of aggregation are significantly lower at Kirriemuir than those at Stranraer and Denmead. Except for thrombin induced aggregation which is lower at Denmead, the results of the other aggregation tests are similar at Denmead and at Stranraer.

The correlations on an individual basis between the various haematological tests are given in Table 1. The italics demonstrate two series of highly significant correlations.

(a) Those between the clotting tests and thrombin aggregation: these two different tests were performed by two different technicians.

(b) The correlations between the other tests of aggregation: none of these tests, however, are significantly correlated with serum cholesterol, HDL-cholesterol, or triglycerides.

The fatty acid composition of plasma total lipids is shown in Fig. 3. The following ratio saturated + monounsaturated/linoleic + linolenic acids appears to be related to platelet factor 3 recalcification clotting time and to thrombin-induced aggregation. The correlation coefficient between that ratio and thrombin induced aggregation is \(r=0.26\) \((p<0.05)\). It rises to \(r=0.32\) \((p<0.01)\) when the interference of plasma triglyceride concentration is removed.

By contrast, the other tests of aggregation seem to be related to the fatty acid ratio arachidonic/linoleic on a group basis though no significant correlation could be obtained on an individual basis.

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**Table 1: Correlations \((r)\) between certain blood variables in 63 farmers**

<table>
<thead>
<tr>
<th>Variable</th>
<th>PCT</th>
<th>THR</th>
<th>ADP(P)</th>
<th>EPI(P)</th>
<th>COLL</th>
<th>F&lt;sub&gt;CT&lt;/sub&gt;</th>
<th>STYPVEN</th>
<th>ADP(S)</th>
<th>EPI(S)</th>
<th>AA</th>
<th>CHO</th>
<th>HDL-CHO</th>
<th>Trig</th>
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<td>0.03</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.03</td>
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<td>0.00</td>
<td>0.03</td>
<td>0.02</td>
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Abbreviations as in Fig. 1 and 2.

STYPVEN = Stypven-time of platelet rich plasma.

Highly significant \((p<0.001)\) correlations are in italics \((r=0.25, p<0.05; r=0.32, p<0.01; r=0.41, p<0.001)\).
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Denmead, Kirriemuir, Stranraer (20 subjects, 22 subjects, 21 subjects)

Mean ± S.E. **p<.05 **p<.01 *p<.001

Fig. 2  Haematological results. PCT, recalcification plasma clotting time of platelet rich plasma (evaluates the whole blood clotting activity); CEP-CT, cephalin clotting time of platelet poor plasma (evaluates the global activity of the plasmatic clotting factors); F3-CT, recalcification clotting time of platelets washed and resuspended in a standard platelet poor plasma (evaluates solely the platelet clotting activity, that is PF3). Aggregation (performed in citrated platelet rich plasma, 300 000 platelets/µl). THR, thrombin (final concentration in platelet rich plasma, 0.12 NIH Units per ml); ADP, adenosine diphosphate (0.92 x 10^-6 mol in platelet rich plasma); P, primary aggregation; S, secondary aggregation; EPIN, adrenaline (0.6 x 10^-6 mol in platelet rich plasma); COLL, collagen (126 µg/ml platelet rich plasma); AA, arachidonic acid (70 µg/ml platelet rich plasma). Extent of aggregation for all tests as the maximal deflection in cm.

Results of the dietary habit evaluation are reported in Fig. 4 (diet composite chemical analysis of a 24-hour period) and in Fig. 5 (past recall and weighing technique based on two different days). Fig. 6 shows that the three methods yield closely similar results.

The intake of polyunsaturated fat, with both techniques, was significantly lower at Stranraer than at Kirriemuir and also than at Denmead as evaluated by the chemical analysis. At Kirriemuir, the intake of saturated fat was significantly lower whether alcohol was included or not included in the calculation of calories (Table 2). In practical terms, the mean intake of visible fat in the form of butter, cream, and vegetable fat (margarine + oil) was in g/day (mean ± SE) of 40±3.5, 24±9.0 and 10±2.7, respectively, at

Fig. 3  Fatty acid composition of the plasma total lipids. Saturated (Sat) = myristic (14:0) + palmitic (16:0) + stearic (18:0); Monounsaturated (Mono) = palmitoleic (16:1) + oleic (18:1) + eicosenoic (20:1)
Stranraer, whereas it was of 14.5±5.0, 3.1±2.7 and 44±6.1 at Kirriemuir.

Several univariate correlations between diet components and the blood variables examined in the 63 farmers were highly significant as shown in Table 3. Of particular interest are the correlations between: (a) dietary cholesterol, the alcohol intake, and platelet functions as well as high density lipoprotein cholesterol; (b) dietary cholesterol and platelet factor 3 recalcification time and serum cholesterol; (c) the saturated fat intake evaluated by two different techniques and the platelet function tests.

In multivariate analysis, considering the effect of linoleic acid added to that of the saturated fatty acids, only slightly improved (particularly with the chemical analysis evaluations) the correlations with platelet function tests. By contrast, the correlation with total serum cholesterol became significant (chemical analysis) with a negative sign for linoleic acid. The inclusion of calcium together with saturated fats made no substantial change to the correlations with the serum lipids (cholesterol, triglycerides, high density lipoprotein cholesterol) whereas it significantly (p<0.05) improved correlations with the platelet factor 3 recalcification time and thrombin-aggregation, with the nutrient intake determined by chemical analysis.

Concerning platelet phospholipid fatty acid composition (Fig. 7), only a few minor differences could be observed between the regions, in the 20:3ω9, 20:4,
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22:5o,3, the ratio 20:3o9 + 22:3o9/18:2, and in the 18:0 dimethyl acetal. Of special interest appears to be the fatty acids 20:3o9, 22:3o9 (which runs parallel to 20:3o9) and 20:4, since they are significantly correlated with the platelet functional tests on an individual basis (Table 4). Moreover, 20:3o9 in platelet phospholipids was also significantly correlated with the intake of saturated fat (r = 0.27) (Table 5). When in addition to saturated fat, the calcium intake was considered in trivariate analysis, the coefficient correlation increased from 0.27 to 0.33, with a negative sign for calcium.

Discussion

These results on serum lipids and platelet function in Scottish farmers follow the same pattern as in the French groups in areas of high and low risk for coronary heart disease. While there was no difference in the lipaemia (including HDL-choesterol) between the two regions despite a much higher intake of saturated and a lower intake of polyunsaturated fatty acids in west Scotland, the platelet functions were much lower in east Scotland, in the subjects examined. A second study (to be reported elsewhere) performed in 1979, again in Scottish farmers, but only on smokers this time, confirmed completely the present study on the lower platelet function and lower fat intake in east Scotland, even though the areas investigated were different (Inverurie and Girvan).

The difference noted with the diet surveys is further substantiated by the fatty acid composition of the plasma total lipids, which shows a significantly lower level of linoleic acid in west Scotland (the region with the higher coronary heart disease risk) and a slightly higher level of oleic acid, which is the usual pattern observed in atherosclerotic patients with or without demonstrated platelet hyperactivity.

Similar results were also reported recently in a comparison between men from Edinburgh and from Stockholm. Though serum cholesterol was identical in the two cities, a striking difference in the level of linoleic acid in plasma lipids and adipose tissue was
observed. This decrease in linoleic acid appears to be associated with the three times greater mortality rate from coronary heart disease in Edinburgh.14 The correlation between the plasma fatty acid composition and the platelet function tests is of interest; it confirms our previous observations,12 and emphasises the obvious dependence of platelet functions on the diet and plasma fatty acid composition.

The difference in the dietary habits between the groups in east and west Scotland appears to be mostly the result of the following.

(1) The visible fats which are essentially dairy products in the west, and vegetable fats (oil and polyunsaturated margarines) in the east, at least in the subjects studied. Dairy farming is the main activity in south Scotland and ease of access to dairy products may be of importance in the type of diet consumed. It seems that a change from dairy to beef farming has taken place in east Scotland in the past decade.

The prominent influence of saturated fats (and eventually of dietary cholesterol) on platelet functions is substantiated by the correlations observed in the present study between these nutrients and the platelet function tests. Similar correlations have been found in French farmers.4 5 15

(2) The higher intake of alcohol in the east Scotland group. Alcohol, in the present study, was inversely correlated with some of the platelet function tests and

Table 3  Correlations (r) between blood parameters and diet components in 63 British Farmers studied in three regions

<table>
<thead>
<tr>
<th>Diet</th>
<th>Method</th>
<th>F3-CT</th>
<th>THR</th>
<th>ADP</th>
<th>EPI</th>
<th>COLL</th>
<th>Cho</th>
<th>HDL</th>
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<tr>
<td>Alco</td>
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<tr>
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<td>PW</td>
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<td>SAT</td>
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<tr>
<td>SAT and</td>
<td>C</td>
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<td>0.32</td>
<td>0.22</td>
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* Trivariate analysis. Abbreviations as in Fig. 1. Cho, cholesterol; ALCO, alcohol; PW, past recall + weighing technique; C, chemical analysis.
Significant correlations are italicised (r=0.25, p<0.05; r=0.32, p<0.01; r=0.41, p<0.001). In parentheses, the sign of 18:2 or Ca²⁺ in the multivariate analysis.

Fig. 7  Fatty acid composition of the platelet total phospholipids. DMA, Dimethyl acetal.
positively correlated with the level of high density lipoprotein cholesterol, a result similar to that obtained in French farmers. Alcohol is known to increase the level of high density lipoprotein cholesterol and inhibits platelet functions both in man and in animals. Consequently, the higher intake of alcohol in the east Scotland group might be partly responsible for the lower response of platelets to aggregation by ADP, adrenaline, and collagen observed in this region. In southern England (Denmead), a region in which the incidence of coronary heart disease does not appear to be much different from that of east Scotland, at least in the general population, only the clotting activity of platelets and, to some extent, the response to thrombin, were comparable, among the tests performed, to the results obtained in East Scotland. It has to be emphasised that it is only these two tests that we have repeatedly found to be closely related to thrombosis in animals and to coronary heart disease in man.

The lower response in Denmead, of the platelet clotting activity and of the thrombin aggregation could be the result of the intake of calcium or of linoleic acid, both being significantly higher when determined by chemical analysis in this region as compared with west Scotland. Chemical analysis is the only technique which can precisely evaluate the intake of: (a) minerals such as calcium which depend on the local source of water; (b) polyunsaturated fats since in margarines and prepared food, the amount of polyunsaturated can change conspicuously in a short time; (c) lipids which vary according to the way the meat is cooked.

Multivariate analysis of the results from the diet composite chemical analysis suggests that the saturated fat induced hyperactivity of some platelet functions (platelet factor 3 recalcification time, thrombin) was inhibited mainly by dietary calcium. This has been confirmed in more extensive studies, in French farmers as well as in animal studies. Thus, the protective effect of dietary calcium on platelet functions, probably by promoting the excretion of long chain saturated fats as reported in animals and man, a result that we have confirmed recently in animal studies (to be published elsewhere) might be consistent with the known protective effect associated with hard water on coronary heart disease.

Concerning the mechanisms involved in the platelet hyperactivity induced by dietary fats, we have previously shown that it seems to depend, in animals and man, mainly on the increase in the platelet phospholipids of oleic acid and its trienoic derivative 20:3o9. In the present study, 20:3o9, 22:3o9, and 20:4 were correlated on an individual basis with the platelet function tests. In addition, the percentage of 20:3o9 in the platelets was also directly dependent upon the intake of saturated fat, and inversely related to that of calcium, as for the platelet function tests.

While the role of 20:4 in the platelet phospholipids seems to be obvious as the precursor of endoperoxide for aggregations dependent upon prostaglandins (adrenaline, collagen), we have no definite explanation of the effect of 20:3o9 and 22:3o9 on platelet functions. Nevertheless the potential detrimental effect of 20:3o9 (and probably of 22:3o9, its derivative) which is specifically increased in essential fatty acid deficiency, has been consistently observed in human and animal studies in relation to thrombosis, atherosclerosis, and coronary heart disease. Deficiency in essential fatty acids has been recently emphasised in relation to atheroma in man, but attention has been focused, until now, on the deficiency in 18:2 or its derivatives rather than on the possible noxious effects of 20:3o9.

Finally, in the present investigation, in which the smokers studied had not smoked for 10 hours before blood removal, no relation could be found between cigarette smoking and any of the platelet functions examined. In the additional studies conducted in 1979 on smokers from two Scottish areas, however, we have observed in confirmation of Levine that most

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Prophospholipid fatty acids most closely related to platelet function tests in 63 British farmers, univariate analysis (r).</th>
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<tr>
<td></td>
<td>20:3o9</td>
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<td>FCT</td>
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<td>EPI</td>
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Abbreviations and significance as in Fig. 2 and Table 3.

Table 5: Correlations (r) between certain platelet phospholipid fatty acids, diet components and platelet function tests in 63 British farmers studied in 1978.

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<td>ADP</td>
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* Trivariate analysis

Abbreviations and significance as in Fig. 2 and Table 3.
of the platelet function tests were much increased when performed 10 minutes after smoking one cigarette. This effect was additive to the hyperactivity resulting from the saturated fat diet.

References


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