Effect in vitro on platelet function of two compounds developed from the pyrimido-pyrimidines


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SYNOPSIS

VK 774 and VK 744, two new compounds developed from the pyrimido-pyrimidines, have been found to be powerful inhibitors of platelet function tested in vitro. They inhibit adenosine diphosphate (ADP)-induced platelet aggregation, and the release of platelet factor 3 by kaolin, and VK 774 also reduces platelet adhesiveness and inhibits platelet aggregation ('snowstorm' effect) in the Chandler tube system. Although measured percentage whole blood clot retraction was uninfluenced by these drugs the clot produced with VK 774 was friable and soft. VK 774 appears to be the most powerful of these compounds reported so far, being active in some test systems at $10^{-6}$M, and, if the results of toxicity testing are satisfactory, it should be an important agent for therapeutic trial.

Platelets play a major role in the initiation of thrombus formation and this is particularly true of thrombi arising on the arterial side of the circulation (Mustard and Packham, 1970a). Platelet function may be studied in a number of systems in vitro and the effect of various drugs thereby assessed. The pyrimido-pyrimidine derivatives have been widely studied in this respect but many other compounds interfering with platelet action have been described and the subject has recently been reviewed by Mustard and Packham (1970b). The early pyrimido-pyrimidine derivative diprydiamole (RA8, Persantin, Boehringer Ingelheim, Limited) was found to be effective in vitro in inhibiting platelet aggregation induced by ADP (Emmons, Harrison, Honour, and Mitchell, 1965a; Gray, Wilson, and Douglas, 1968), platelet aggregation in the Chandler tube system, and platelet adhesiveness (Gray et al, 1968). Diprydiamole has also been demonstrated to inhibit thrombus forming at sites of vascular injury in experimental animals (Emmons, Harrison, Honour, and Mitchell, 1965b; McNicol, 1968), and in a controlled trial with random allocation has been shown to be of value when combined with oral anticoagulants in reducing the incidence of embolic episodes in patients with prosthetic heart valves as compared with patients on anticoagulants alone (Sullivan, Harken, and Gorlin, 1969). Since the introduction of diprydiamole, other pyrimido-pyrimidine derivatives have been developed and found in certain in-vitro systems to inhibit platelet function more powerfully. RA 433 was found to be a more potent inhibitor of ADP-induced platelet aggregation in vitro than diprydiamole (Elkeles, Hampton, Honour, Mitchell, and Prichard, 1968; Forbes, McNicol, and Douglas, 1969) but had no effect on thrombus formation in the experimental animal (Elkeles et al, 1968). RA 233 in turn, in a direct comparison with RA 433, has been found to be the more powerful inhibitor of ADP-induced platelet aggregation, of platelet aggregation induced by calcium in a turbidimetric system, and of platelet adhesiveness (Hassanein, Turpie, McNicol, and Douglas, 1970).

This study deals with the effect upon human platelet function in vitro of two new compounds, VK 774 and VK 744, chemically distinct from but historically related to pyrimido-pyrimidine derivatives. Their activity in this respect is directly compared with the pyrimido-pyrimidine derivative RA 233.

Chemistry

VK 774 is 4-morpholino-2-piperazino-thieno-(3, 2-D)pyrimidine-dihydrochloride, and VK 744 is 2-((2 aminoethyl)amino)-4-morpholinothieno (3, 2-D)pyrimidine-dihydrochloride, and their structural formulae together with those of RA 233 and RA 8 (diprydiamole) are set out on page 2. Both VK 774 and VK 744 are soluble in water and their molecular weights are 432-39 and 352-30 respectively. RA 233 is
relatively insoluble in water but soluble in dilute acids; its molecular weight is 421.50.

Materials

**VK 774 (BOEHRINGER INGELHEIM, LIMITED)**
Stock solutions are 10⁻³M and 10⁻⁵M VK 774 in sterile distilled water, stored at 4°C. Dilutions referred to in the text were made in sterile distilled water.

**VK 744 (BOEHRINGER INGELHEIM, LIMITED)**
Stock solutions are 10⁻³M and 10⁻⁵M VK 744 in sterile distilled water, stored at 4°C. Dilutions referred to in the text were made in sterile distilled water.

**RA 233 (BOEHRINGER INGELHEIM, LIMITED)**
Stock solutions are 10⁻³M and 10⁻⁵M RA 233 in 0.025 N HCl, stored at 4°C. Dilutions referred to in the text were made in 0.025 N HCl.

**ADENOSINE 5-DIPHOSPHATE (ADP) (SIGMA CHEMICAL COMPANY, ST LOUIS)**
Stock solution of 100 µg/ml ADP in barbitone/saline buffer, pH 7.2; stored at −20°C. Dilutions were made in barbitone/saline buffer.

**TUBING**
In Chandler tube experiments transparent vinyl tubing (N/17, Portland Plastics Limited, Kent) and plastic adaptors (10M/634, Portex) were used.

**GLASS BEAD COLUMNS**
Transparent vinyl tubing (NT/13, Portland Plastics Limited, Kent) and translucent silicone tubing (Esco Rubber Limited, London) were used.

Ballotini glass beads, 0.57 mm diameter; calcium chloride, 0.25M, 0.025M; kaolin, 5% kaolin in imidazole buffered saline pH 7.2; Russell viper venom (Stypven, Burroughs Wellcome Company).

Citrated blood was collected by clean venupuncture in plastic syringes with 21-gauge needles, 9 volumes of blood being mixed with 1 volume of 3.8% trisodium citrate in siliconized graduated centrifuge tubes maintained at room temperature. Platelet-rich plasma was obtained by centrifugation of citrated whole blood at 600 g for five minutes at room temperature. Siliconized glassware was used throughout (Siliclad, Clay-Adams, Inc, New Jersey).

Methods

The following parameters of platelet function were studied and the methods are described in detail by Forbes et al (1969).

**ADP-INDUCED PLATELET AGGREGATION**
This was studied by the method of Born (1962) at room temperature. Platelet-rich plasma (1.8 ml) was incubated at room temperature for five minutes with 0.2 ml dilutions of test solution or with solvent control, and challenged by 0.5 µg of ADP (final concentration 0.24 µg ADP/ml solution). The optical density was recorded every 10 seconds for 1 minute after addition of ADP.

Indices of platelet aggregation and disaggregation are expressed as follows: 30-60 platelet aggregation is fall in optical density between 30 and 60 seconds after the addition of ADP; maximum aggregation is the difference between the lowest recorded optical density and the constant arbitrary baseline of 0-600; percentage disaggregation is the ratio between the increment in optical density occurring in five minutes from the point of maximum aggregation and the maximum platelet aggregation.

**PLATELET ADHESIVENESS**
The glass bead column technique of Hellem (1960) as modified by Hirsh, McBrade, and Dacie (1966) was used. Citrated whole blood (4.0 ml) was incubated at room temperature for 10 minutes with 0.04 ml of test solution or control.

**RELEASE OF PLATELET FACTOR 3 BY KAOLIN**
A modification of the technique of Spaet and Cintron (1965) was used. Platelet-rich plasma (0.9 ml) was preincubated at 37°C for 10 minutes with test dilutions or solvent control (0.1 ml).

**CHANDLER TUBE TECHNIQUE**
The method used was that described by Chandler (1958) as modified by Cunningham, McNicol, and...
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Douglas (1965). Small volumes were used: platelet-rich plasma (1.0 ml), 0.877% saline (2.86 ml), and test solution or solvent control (0.04 ml) were incubated in the Chandler tube at 37°C for five minutes before recalcification with 0.25M calcium chloride (0.1 ml).

Whole Blood Clot Retraction
A modification of the method described by Dacie and Lewis (1968) was used; whole blood (4.95 ml) was mixed in siliconized centrifuge tubes containing copper wire spirals with test solution or solvent control (0.05 ml) before clot formation.

Platelet counts were performed with formal-citrate as the diluting fluid (Dacie and Lewis, 1968).

Results
ADP-Induced Platelet Aggregation
In 14 experiments both VK 774 and VK 744 produced statistically significant inhibition of 30 to 60 sec and maximal platelet aggregation at a final concentration of 10^-4M, though VK 774 was the more powerful (p < 0.001) and almost completely inhibited aggregation. VK 744 also produced a significant increase in percentage disaggregation (p < 0.001). RA 233 at this concentration (10^-3M) had no significant effect on 30 to 60 sec or maximum platelet aggregation but significantly increased percentage disaggregation (p < 0.001), although VK 744 was more powerful in this respect (p < 0.001). In seven experiments at a concentration of 10^-4M, of the three drugs, only VK 774 had a statistically significant effect in the system, producing inhibition of 30 to 60 sec aggregation (0.01 > p > 0.001) and of maximum platelet aggregation (p < 0.001). Figure 1 shows the mean results of the three drugs in the 14 experiments at 10^-4M concentration, and of the effect of VK 774 at 10^-4M concentration compared with solvent controls.

Platelet Adhesiveness
In seven experiments at a final concentration of 10^-4M VK 774 significantly reduced retention of platelets by glass beads (p < 0.001), but VK 744 and RA 233 had no statistically significant effect at this concentration. These results are shown in Figure 2.

Release of Platelet Factor 3 by Kaolin
In seven experiments at a concentration of 10^-5M all three drugs produced statistically significant reduc-

Fig. 1 ADP-induced platelet aggregation. Results of 14 experiments using VK 774, VK 744, and RA 233 at a concentration of 10^-5M, and of seven experiments at a concentration of 10^-4M (VK 774 only shown) compared with solvent controls. ADP concentration 0.24 μg/ml.

Fig. 2 Effect of VK 774, VK 744, and RA 233 (concentration 10^-5M) on platelet retention by glass beads (VK 774 p < 0.001; VK 744 0.1 > p > 0.05; RA 233 p > 0.1).
Effect of two concentrations of VK 774, VK 744, and of RA 233 on the release of platelet factor 3 by kaolin. (Above, concentration $10^{-5}$M, VK 774, $p < 0.001$; VK 744 $0.01 > p > 0.001$; RA 233 $0.05 > p > 0.02$. Below, concentration $10^{-6}$M, VK 774 $p < 0.001$; VK 744 $p > 0.01$; RA 233 $p > 0.1$.)

Concentration in platelet factor 3 release. VK 774 was more potent in this respect than VK 744 ($p = 0.001$), which in turn was more powerful than RA 233 ($0.01 > p > 0.001$). At a concentration of $10^{-4}$M in seven experiments only VK 774 had a statistically significant effect ($p < 0.001$). These results are shown in Figure 3.

**CHANDLER TUBE TECHNIQUE**

The results obtained in seven experiments using VK 774, VK 744, and RA 233 at a final concentration of $10^{-4}$M compared with solvent controls are shown in Figure 4. VK 774 and RA 233 produced significant prolongation of platelet aggregation time ('snowstorm' effect). They also considerably reduced the magnitude of the snowstorm effect as judged by inspection; this action was particularly noticeable with VK 774. VK 744 had no significant effect in this system.

**WHOLE BLOOD CLOT RETRACTION**

In seven experiments at $10^{-4}$M concentration VK 774, VK 744, and RA 233 had no statistically significant effect on percentage clot retraction although some of the clots with VK 774 were noted to be friable and soft.

Fig. 3 Effect of two concentrations of VK 774, VK 744, and of RA 233 on the release of platelet factor 3 by kaolin. (Above, concentration $10^{-5}$M, VK 774, $p < 0.001$; VK 744 $0.01 > p > 0.001$; RA 233 $0.05 > p > 0.02$. Below, concentration $10^{-6}$M, VK 774 $p < 0.001$; VK 744 $p > 0.01$; RA 233 $p > 0.1$.)

Fig. 4 Effect of VK 774, VK 744, and RA 233 in concentrations of $10^{-4}$M on platelet aggregation time in the Chandler tube system (VK 774 $0.02 > p > 0.01$; VK 744 $p > 0.1$; RA 233 $0.01 > p > 0.001$).
Discussion

The first step in the initiation of thrombus formation is the adherence and aggregation of platelets to each other and to the vessel wall (Mustard, Murphy, Rowsell, and Downie, 1962), and ADP released into the blood stream by various mechanisms appears to play a major part in stimulating this process (Mustard and Packham, 1970a). Subsequent release of platelet factor 3 (phospholipoprotein), promoted by platelet aggregation (Mustard, Hegardt, Rowsell, and MacMillan, 1964; Hardisty and Hutton, 1966), accelerates the clotting mechanism and leads to stabilization of the thrombus with fibrin production, and eventual clot retraction reduces the friability of the clot. Substances capable of inhibiting platelet adhesion and aggregation, release of platelet factor 3, and clot retraction, may be expected to be of possible therapeutic value. Which particular platelet function test best reflects the thrombus-forming properties of a platelet is, however, unknown, and tests in vitro using the injured artery model are not necessarily more valid than tests in vivo (Elkeles et al., 1968). A useful review of platelet function tests and their relevance to thrombosis has been given by Hampton (1967).

The place of anticoagulant therapy in arterial thrombosis remains controversial. Platelet behaviour, of central importance in the initiation of thrombi, especially arterial, is largely uninfluenced by anticoagulant drugs. Drugs such as the pyrimido-pyrimidine derivatives which interfere with platelet function, however, offer a potential therapeutic advance in the management of arterial thrombosis, and there is already clinical and experimental evidence of the usefulness of dipyridamole in the prevention of thromboembolic complications of prosthetic heart valves (Sullivan et al., 1969; Harker and Slichter, 1970). So far neither RA 433 nor RA 233 have been evaluated clinically though they are well absorbed orally and appear to be relatively non-toxic in experimental animals (Elkeles et al., 1968; Hassanein et al., 1970). VK 774 and VK 744 are two new compounds developed from the pyrimido-pyrimidines and their effect on platelet function in vitro is reported and compared directly with that of RA 233. VK 774 and VK 744 inhibit ADP-induced platelet aggregation and VK 744 is considerably more powerful in this respect, almost completely inhibiting aggregation at 10^{-5}M and being the only drug still significantly effective at 10^{-6}M. VK 744, and to a significantly lesser degree RA 233, increase platelet disaggregation though there is no significant effect on aggregation by RA 233 at the concentrations used. Platelet adhesiveness is significantly reduced only by VK 774, VK 744, and RA 233 having no significant effect at the concentration used.

All three drugs impair the release of platelet factor 3 by kaolin, the VK compounds proving more potent in this respect than RA 233, and VK 744 having statistically the most powerful effect. At a concentration of 10^{-4}M again only VK 744 has a statistically significant inhibitory effect on platelet factor 3 release. Platelet aggregation in the Chandler tube system, as estimated by the time of appearance of the snowstorm effect of aggregation, is significantly inhibited by VK 774 and RA 233 but not by VK 744. There is no significant difference noted between VK 774 and RA 233. They both also considerably reduce the magnitude of the snowstorm effect and this is particularly noticeable with VK 774 with which clotting occurs with little or no recognizable prior snowstorm effect of platelet aggregation. In the concentration used there is no effect on percentage whole blood clot retraction by any of the three drugs although the final clot with VK 774 is noted in some cases to be friable and soft.

These results show that VK 774 and VK 744 have anti-platelet properties similar to the previously tested pyrimido-pyrimidine derivatives RA 233, RA 433, and dipyridamole, and in some of the in-vitro systems tested are significantly more powerful. Of the two VK compounds tested, VK 774 stands out clearly in this study to be considerably the more potent drug and therefore the most powerful of these compounds so far reported. Initial toxicity studies with VK 774 are promising, and if these are satisfactorily completed it should be an important agent for therapeutic trial.

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