THE LABILE FACTOR OF PROTHROMBIN CONVERSION: ITS
CONSUMPTION UNDER NORMAL AND PATHOLOGICAL
CONDITIONS AFFECTING BLOOD COAGULATION 1,2

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INTRODUCTION

The rapid conversion of prothrombin to thrombin by thromboplastin plus calcium requires a
plasma component called "Labile Factor" (2, 3), which is probably identical with plasma Ac-globu-
lin and Factor V (4, 5), and with the prothrombin accelerator of Fantl and Nance (6, 7). 3 Deteriora-
tion of this substance as plasma ages produces an increase in the prothrombin time (2, 3, 8) although
the concentration of prothrombin remains essentially unchanged (9, 10). This defect can be
rectified by the addition of fresh plasma devoid of prothrombin as well as of fibrinogen (11, 8).

Additional information regarding Labile Factor (L.F.) is desirable for several reasons. Its
precise role in prothrombin conversion is obscure, and hemorrhage referable to deficiency of this
entity or of a similar substance has been described (12–14). Furthermore, as will become evident,
the concentration of L.F. must receive due consideration in procedures which measure prothrom-
bin.

In certain hemorrhagic diseases the conversion of prothrombin to thrombin is reduced or retarded
(15, 16). The "prothrombin consumption test," which demonstrates this defect, promises to be-
come important in the investigation and diagnosis

1 Supported by a grant from The Commonwealth Fund.
2 Presented in part before the Federation of American
Societies for Experimental Biology, April, 1950 (1).
3 It should be pointed out that most investigators feel
that Labile Factor, Factor V, plasma Ac-globulin, and the
factor of Fantl and Nance are identical. Thus Quick and
Stefanini (31) use the terms interchangeably to designate one entity. Seegers and his associates (10) are in-
clined to agree that Ac-globulin is the same as L.F. and
Factor V (30, 40). We (24) as well as Flynn and his
coworkers (41) hold the same view. Definite proof re-
garding the identity of all these moieties, however, is
still lacking.

of such disturbances. In order to compute the
consumption of prothrombin during coagulation,
the precise amount of prothrombin in both serum
and its parent plasma must be known. Obviously
changes in the concentration of L.F. in the tran-
sition from plasma to serum may influence con-
siderably the velocity of prothrombin conversion
in serum and, therefore, the determination of se-
rum prothrombin by the one-stage procedure,
since the prothrombin times may reflect the con-
centration of L.F. as well as that of prothrombin
itself. Under such circumstances the computation
of prothrombin consumption from differences be-
tween plasma and serum prothrombic activities
may be erroneous.

Whereas the concentration of L.F. has been de-
termined in the plasma of man and other species
(17), little is known regarding its level in serum.
This information is presented in the present re-
port. Also included is a method proposed for the
determination of serum L.F. The data indicate
that during and after coagulation changes in this
clotting factor occur which are considerably in-
fluenced by conditions, pathological or induced,
which affect clotting.

METHODS

Prothrombin determination: The prothrombic activity
of oxalated plasma (one volume of 0.1 M sodium oxalate
to nine volumes of blood) was determined on undiluted
plasma by the orthodox one-stage procedure (18), and
on diluted plasma by the modification (19) in which the
test plasma is diluted with fresh normal oxalated pooled
plasma rendered prothrombin-free by adsorption with
BaSO4. 4 Prothrombin levels were calculated by inter-
polating the observed prothrombin times on standardiza-
tions obtained on normal plasma similarly diluted.
Serum prothrombin was measured similarly and also

4 Plasma so treated is hereafter referred to as "BaSO4
plasma."
by the modified two-stage method of Ware and Seegers (20).

**Labile factor**: Pooled oxalated plasma from normal subjects was stored at 4–5°C for variable intervals during which its prothrombin time increased progressively. L.F. activities of test plasma or sera were measured by their restorative effects on the elevated prothrombin time of these aged plasmas in which the factor had largely deteriorated (8, 17). The restorative effect was determined by adding up to one part of plasma or serum (whole or diluted with physiological saline) to four parts of the aged plasma, and by observing the prothrombin times of the mixtures. The values herein reported are the averages of several observations on each mixture. In ranges below 45 seconds the prothrombin times were always duplicable within two seconds; between 45 and 60 seconds, within four seconds.

Except where otherwise indicated, the serum was separated one hour after venous blood was allowed to clot in a glass tube at room temperature. The clot was rimmed and centrifuged, and the serum was removed and incubated at 37°C for 15 minutes in order to inactivate thrombin. The time interval between incubation of the serum and determination of its activity was noted. Experiments were run on non-oxalated as well as oxalated serum (1 oxalate to 4 serum). In some instances citrate (2.5 per cent sodium citrate) was used instead of oxalate. Also studied were sera which were subjected to adsorption with BaSO₄ for 10 minutes at 37°C. Controls for the latter experiments were incubated for the same interval without BaSO₄.

Commercial thromboplastin (Difco) and beef lung thromboplastin were used in the one- and two-stage methods, respectively.

**RESULTS**

The experiments reported below are typical of a series of experiments performed under the same conditions.

**Comparative effect of normal serum and plasma on prothrombic activity of aged plasma**

Oxalated fresh plasma has a profound restorative effect on the retarded prothrombin conversion of stored plasma (Figure 1). This is not due to the prothrombin in the fresh plasma, since removal of the prothrombin by BaSO₄ adsorption does not significantly decrease its restorative activity (Figure 1, Table I). There appears to be remarkably little variation in the L.F. activity of plasma from different normal subjects (Table I). These observations confirm earlier reports (8, 11, 21).

Non-oxalated normal serum tested 75 minutes after coagulation, has as much, and often more, restorative activity than the parent plasma tested simultaneously (Table II). On the other hand,
Table I

Variation of L.F. in fresh normal oxalated plasma before and after adsorption with BaSO₄

<table>
<thead>
<tr>
<th>Subject</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Per cent fresh plasma in fresh plasma-adsorbed plasma mixture</td>
<td>Whole plasma</td>
<td>BaSO₄ plasma</td>
<td>Whole plasma</td>
<td>BaSO₄ plasma</td>
</tr>
<tr>
<td>0</td>
<td>87</td>
<td>87</td>
<td>87</td>
<td>87</td>
</tr>
<tr>
<td>1</td>
<td>43</td>
<td>46</td>
<td>42</td>
<td>46</td>
</tr>
<tr>
<td>2</td>
<td>36</td>
<td>37</td>
<td>35</td>
<td>36</td>
</tr>
<tr>
<td>4</td>
<td>29</td>
<td>29</td>
<td>28</td>
<td>30</td>
</tr>
<tr>
<td>6</td>
<td>27</td>
<td>28</td>
<td>24</td>
<td>27</td>
</tr>
<tr>
<td>10</td>
<td>23</td>
<td>26</td>
<td>23</td>
<td>24</td>
</tr>
<tr>
<td>20</td>
<td>19</td>
<td>21</td>
<td>20</td>
<td>22</td>
</tr>
</tbody>
</table>

BaSO₄ serum is considerably less potent than the control serum (Table III). From interpolation of these data on the curve in Figure 1, it is evident that BaSO₄ removes approximately 80 per cent or more of the restorative activity from serum whereas it does not do so from plasma.

Serum has certain clot-promoting properties not demonstrable in plasma (22). This is due to the serum prothrombin conversion accelerator (spca) which arises during coagulation and which in the presence of L.F. accelerates prothrombin conversion (22–24). Accordingly, the accelerating effect of serum on the prothrombin conversion in aged plasma might be due to both spca and L.F.

BaSO₄ removes little, if any, L.F. from plasma (8, 21). It is assumed that it likewise does not remove this substance from serum. On the other hand, spca is quantitatively adsorbed (22, 24). Accordingly, BaSO₄ serum is devoid of spca but should retain its full complement of L.F. That BaSO₄ serum has less restorative ability than BaSO₄ plasma indicates, therefore, that the L.F. of normal serum is far less than that of plasma.

Effect of anti-coagulants upon restorative property of normal serum

L.F. is more stable in citrated than in oxalated plasma (8, 25). It is also more stable in native plasma (containing calcium) but not in plasma rendered incoagulable by decalcifying with ion-exchange resins (26). Experiments were performed in order to see whether the L.F. in serum is similarly affected (Table IV). The data indicate that the restorative property of oxalated serum is more labile than that of non-oxalated serum.

The question arises whether oxalate affects L.F., not adsorbable by BaSO₄, or some other serum factor such as spca. Experiments performed on non-oxalated versus oxalated BaSO₄ serum

Table II

Restorative activity of normal plasma and corresponding sera* on retarded prothrombin conversion of aged plasma

<table>
<thead>
<tr>
<th>Interval between clotting and test</th>
<th>Proth-T (sec.) of mixture</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.P.</td>
<td>A.P. plus S</td>
</tr>
<tr>
<td>hrz.</td>
<td></td>
</tr>
<tr>
<td>3 ½</td>
<td>33</td>
</tr>
<tr>
<td>2</td>
<td>45</td>
</tr>
<tr>
<td>3 ½</td>
<td>37</td>
</tr>
<tr>
<td>2</td>
<td>48</td>
</tr>
<tr>
<td>1 ½</td>
<td>66</td>
</tr>
</tbody>
</table>

* All sera separated one hour after clotting at room temperature in glass; both sera and plasma incubated at 37°C for additional 15 minutes and then tested.
† Prothrombin time.
‡ Mixture comprises 0.4 ml. of aged plasma plus 0.1 ml. of material indicated.

Table III

Effect of BaSO₄ adsorption on restorative activity of normal sera (S)* on prothrombin conversion of aged plasma (A.P.)

* Non-oxalated sera, separated one hour after blood was shed. One part serum added to four parts A.P.
† Sera adsorbed with 50 mgm. BaSO₄ per ml.
THE LABILE FACTOR OF PROTHROMBIN CONVERSION

TABLE IV
Deterioration of serum L.F. in presence of oxalate* and citrate*
Prothrombin time (sec.)

<table>
<thead>
<tr>
<th></th>
<th>75 min. after coag.</th>
<th>195 min. after coag.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
<td>II</td>
</tr>
<tr>
<td>Aged plas.</td>
<td>37</td>
<td>37</td>
</tr>
<tr>
<td>Aged plas. plus:‡</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Ser.</td>
<td>17</td>
<td>18</td>
</tr>
<tr>
<td>Cit. ser.</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>Oxal. ser.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Oxalate and citrate added to normal serum 60 min. after coagulation.
‡ 0.1 ml. serum added to 0.4 ml. of aged plasma.

(Table V) indicate that the latter, also, was more labile than its non-oxalated counterpart. On the basis of the assumption that BaSO₄ does not remove L.F. from serum, it is concluded that oxalate accelerates deterioration of this clotting component.

The restorative activity seems more stable in citrate than in oxalate but not as stable as in native serum (Table IV). It appears, therefore, that L.F. deteriorates more rapidly in serum containing oxalate than in native or citrated serum. The same is observed in plasma (8, 17, 25, 26).

CONSUMPTION OF L.F. AND PROTHROMBIN DURING NORMAL COAGULATION

That L.F. disappears from blood during coagulation is clearly established. Its rate of disappearance was studied in relation to the rate of prothrombin consumption. Aliquots of freshly drawn blood were allowed to clot at 37°C in separate dry test tubes. At specified intervals oxalate was added, the mixture was stirred with a glass rod and immediately centrifuged at 2000 r.p.m. for 10 minutes. The sera were separated and subjected to determination of residual prothrombic activity

![Graph](image-url)
There was considerable discrepancy between the one- and two-stage prothrombin values. This is not surprising since one-stage prothrombin times of serum reflect the concentration not only of un-consumed prothrombin but also of residual L.F. together with evolved spca.

It is of interest that within the first 10 minutes, about one-half of the L.F. had disappeared, before any detectable (by the methods used) decrease in prothrombin. Although this must be significant in the role of L.F. in prothrombin conversion, the evidence is too meager to permit any interpretation.

### Effect of thromboplastin on concentration of L.F. in serum

Since thromboplastin added to normal blood yields serum which is always devoid of prothrombin in contrast to normal sera which often show small amounts, it was of interest to study the L.F. content of such sera. Sera from thromboplastin-supplemented blood were far less able to restore prothrombic activity to aged plasma than sera from spontaneously clotted blood (Table VI). This

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**Table VI**

<table>
<thead>
<tr>
<th>Aged plasma plus:</th>
<th>Proth-T (sec.) Exp’1*</th>
<th>Exp’2 †</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>47</td>
<td>63</td>
</tr>
<tr>
<td>S</td>
<td>25</td>
<td>33</td>
</tr>
<tr>
<td>BaSO₄ S</td>
<td>30</td>
<td>44</td>
</tr>
<tr>
<td>TPL S</td>
<td>34</td>
<td>47</td>
</tr>
<tr>
<td>TPL BaSO₄ S</td>
<td>39</td>
<td>64</td>
</tr>
</tbody>
</table>

BaSO₄ S prepared with 25 mgm. BaSO₄ per ml. TPL S obtained from blood (nine vol.) added to TPL solution (one vol.) prepared as for prothrombin determination. Identical results were obtained when the ratio was 19 to 1.

* One volume of material to four of aged plasma.
† One volume of material to one of aged plasma.

by both one- and two-stage methods. Aliquots of the serum were also adsorbed with BaSO₄ (50 mgm. per ml.) and were then tested for L.F. activity. It is evident (Figure 2) that both L.F. and prothrombin disappear during the coagulation process. Whereas L.F. began to decrease promptly after the blood was shed, the two-stage prothrombin values showed no significant change within the first 10 minutes. Within 40 minutes most of both clotting factors had disappeared.

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**Fig. 3.** Restorative Activity of "Dicumarol" Serum on Retarded Prothrombin Conversion of Aged Normal Plasma

Serum was non-oxalated, and was tested 1½ hours after the blood was obtained. Adsorption with BaSO₄ was carried out with 50 mgm. per ml.
difference, evident also in the BaSO₄ adsorbed sera, suggests that more L.F. is consumed when coagulation and prothrombin conversion are accelerated by thromboplastin.

**L.F. in serum from dicumarolized blood**

The purpose of these experiments was to study L.F. depletion during coagulation in relation to the amount of prothrombin available for conversion to thrombin. Dicumarolized blood contains essentially normal amounts of L.F. (17). As is evident in Figure 3, dicumarol serum has a very strong restorative effect on aged plasma, more potent than normal serum (compare with data of Table II; also compare data in Table VII with those in Table V, all obtained on the same aged plasma). The same is true of dicumarol BaSO₄.

**TABLE VII**

<table>
<thead>
<tr>
<th>Aged plasma*</th>
<th>Proth-T (sec.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>61</td>
</tr>
<tr>
<td>Fresh D.S.</td>
<td>11</td>
</tr>
<tr>
<td>Fresh oxal. D.S.</td>
<td>14</td>
</tr>
<tr>
<td>Fresh BaSO₄ D.S.</td>
<td>26</td>
</tr>
<tr>
<td>Fresh BaSO₄ D.S. -oxal.</td>
<td>39</td>
</tr>
<tr>
<td>D.S. kept 3 hrs., room temp.</td>
<td>13</td>
</tr>
<tr>
<td>Oxal. D.S., 3 hrs., room temp.</td>
<td>22</td>
</tr>
<tr>
<td>BaSO₄ D.S., 3 hrs., room temp.</td>
<td>36</td>
</tr>
<tr>
<td>BaSO₄ oxal. D.S., 3 hrs., room temp.</td>
<td>64</td>
</tr>
</tbody>
</table>

The prothrombic activity of the parent plasma was 11 per cent. Serum separated one hour after clotting BaSO₄ S prepared with 100 mgm. BaSO₄ per ml.

* One volume of material to four of aged plasma (60 days at 4–5°C).

serum compared with its normal counterpart. Again, as with normal serum, oxalate accelerated L.F. deterioration (Table VII).

It was of interest to study the stability of L.F. in dicumarol serum as contrasted with normal serum. Accordingly, experiments were performed on the same sera recorded in Tables V and VII, after three days' storage at 4–5°C. Determinations were made with the same stored plasma which now had a prothrombin time of 80 seconds compared with 61 seconds three days earlier. The three-day-old normal serum lowered the prothrombin time from 80 to 64 seconds, whereas the three-day-old dicumarol serum reduced it to 32.

Whether this difference, however, reflected a greater lability, or a lesser initial concentration, of L.F. in the normal serum is not clear.

The oxalated dicumarol serum became inert within three days.

**Effect of thromboplastin on L.F. of dicumarol serum**

As already shown, thromboplastin added to normal blood reduced serum L.F. activity. On dicumarol blood this effect was not observed. Serum derived from thromboplastin supplemented dicumarol blood contained the same L.F. activity as dicumarol serum derived without the thromboplastin. This was true whether non-oxalated was compared with oxalated serum, and BaSO₄ compared with non-BaSO₄ serum under the same conditions. It appears, therefore, that the effect of thromboplastin on the L.F. concentration of normal serum was related to the velocity and/or the amount of prothrombin conversion.

Again, in dicumarol-thromboplastin serum L.F. was much less stable in the presence of oxalate.

In these experiments 0.3 ml. of thromboplastin extract was added to 10 ml. of dicumarol blood containing 3 per cent prothrombin. The aged plasma on which the sera were tested had a prothrombin time of 37 seconds.

**L.F. in hemophilic serum**

Hemophilic serum has more restorative activity on aged plasma than normal serum (Table VIII), despite the fact that the interval between the shedding of the blood and the testing of the serum was

**TABLE VIII**

<table>
<thead>
<tr>
<th>Aged plasma*</th>
<th>Proth-T (sec.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Saline</td>
<td>Exp't 1 80</td>
</tr>
<tr>
<td>Norm. S</td>
<td>26</td>
</tr>
<tr>
<td>Hem. S</td>
<td>17</td>
</tr>
<tr>
<td>Norm. BaSO₄ S</td>
<td>34</td>
</tr>
<tr>
<td>Hem. BaSO₄ S</td>
<td>Exp't 2 28</td>
</tr>
<tr>
<td>Saline</td>
<td>80</td>
</tr>
<tr>
<td>Hem. S</td>
<td>14</td>
</tr>
<tr>
<td>Hem. BaSO₄ S</td>
<td>28</td>
</tr>
<tr>
<td>Hem. Tpl.† S</td>
<td>38</td>
</tr>
<tr>
<td>Hem. Tpl. BaSO₄ S</td>
<td>53</td>
</tr>
</tbody>
</table>

* One volume of material added to four of aged plasma.
† Sera, non-oxalated, separated one hour after clotting. BaSO₄ S obtained with 30 mgm. BaSO₄ per ml.

5 Obtained from patients with myocardial infarct treated with dicumarol for the prevention of thromboembolic disease.
inevitably greater as a result of the prolonged clotting time of the hemophilic blood. Also, BaSO₄ removes some restorative activity.

As in all the other experiments, oxalated serum from hemophilic blood had far less restorative ability than non-oxalated sera. This was true with whole as well as BaSO₄-adsorbed serum.

Supplementing hemophilic blood with thromboplastin, which accelerates coagulation, yields serum containing far less L.F. activity, practically the same as that observed in serum from normal blood similarly supplemented (Table VIII).

**Serum L.F. in thrombocytopenia and thrombasthenia**

These findings led to experiments in other pathological conditions where prothrombin conversion is reduced as a consequence of inadequate elaboration of thromboplastic material. It is evident (Table IX) that in thrombocytopenic purpura as well as in thrombasthenia, two disorders where prothrombin consumption is poor (15, 27), serum L.F. activity was abnormally elevated. In these cases, the BaSO₄ sera also were more potent in rectifying the retarded prothrombin conversion of aged plasma than BaSO₄ normal serum. In all instances, the BaSO₄ removed the large amounts of residual prothrombic activity characteristically present in serum in these disorders. Recovery in one patient with acute thrombocytopenia was associated with restoration of serum L.F. and prothrombin to normal values.

**L.F. in “serum” from afibrinogenemic subject**

Experiments performed on blood from a subject with afibrinogenemia exclude the possibility that L.F. disappears during coagulation as a result of adsorption on the fibrin clot. Blood was drawn with and without added oxalate; the latter remained completely fluid indefinitely. One hour after the blood was shed, oxalate was added to the non-oxalated sample. The bloods were centrifuged. The plasma and “serum” were then incubated at 37°C for 15 minutes. Plasma prothrombin was normal by both one- and two-stage methods. The L.F. activity of the plasma also was normal. The “serum” contained no prothrombin by the two-stage procedure, and the L.F. activity of the BaSO₄ adsorbed “serum” was approximately 10 per cent of that of the parent plasma. Also of interest was the finding that the unadsorbed serum contained a normal amount of sPCA. These observations indicate that the disappearance of L.F. activity during coagulation is not related to formation of a fibrin clot. It may be concluded, furthermore, that sPCA elaboration is also independent of fibrinogen and its conversion to fibrin.

**DISCUSSION**

Although the importance of a non-prothrombin plasma constituent in the conversion of prothrombin by thromboplastin plus calcium is now definitely established, its role in this reaction is obscure. Quick and his collaborators, who have designated the entity by the term “Labile Factor” because of its lability, claim that it enters into the reaction stoichiometrically along with thromboplastin and calcium (28). On the other hand, the Wayne University group (Ware, Seegers, and colleagues), calling their substance “plasma Ac-globulin,” consider it to be a relatively inert precursor of “serum Ac-globulin” to which it becomes converted by thrombin. Serum Ac-globulin then functions as an accelerator of the activation of

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*We are indebted to Dr. Louis K. Diamond of the Children's Medical Center, Boston, for making blood, from this patient, available.*
prothrombin \((4, 29, 30)\). Owren, who holds similar views, terms his substance Factor V which is converted to Factor VI during coagulation, and which accelerates thrombin elaboration \((12, 5)\).

Regardless of whether these moieties are identical and it is becoming increasingly evident that they are), certain facts are clear. When L.F. is progressively reduced below 50 per cent of that in normal plasma, the prothrombin time increases \((8)\). Prothrombin consumption is also curtailed \((31)\). Thus, a given prothrombin time reflects the amount of L.F. \((\text{or plasma Ac-globulin and Factor V, if we assume that they are identical})\) as well as the concentration of prothrombin itself. Accordingly, in order to measure prothrombin as such, adequate quantities of this other factor must be assured. This point has received scant consideration in reports on serum prothrombin and on prothrombin consumption during coagulation, especially those based upon one-stage prothrombin determinations, where the velocity of prothrombin conversion is measured.

The foregoing data, in accordance with recent reports \((28, 32)\), show that substantial amounts of L.F. disappear during normal coagulation. Depletion of L.F. begins before prothrombin falls off, but soon the rates with which both are consumed seem parallel. Accelerating and increasing prothrombin conversion with supplements of thromboplastin causes greater losses. Conversely, in hemophilia, thrombocytopenia, and thrombasthenia where prothrombin conversion is retarded or reduced, L.F. consumption appears reduced. Furthermore, curtailing the amount of available prothrombin also decreases the amount of L.F. consumed.

Accelerating coagulation and increasing prothrombin consumption in hemophilia by additions of thromboplastin induces greater L.F. consumption. It may, therefore, be concluded that this clotting factor is utilized in relation to the amount and/or velocity of prothrombin consumption. Furthermore, critically curtailing the concentration of L.F. limits the velocity as well as the amount of thrombin evolution \((33)\). These findings lend support to the idea that L.F. enters into the reaction as a real participant rather than the alternative view that it is merely an "accelerator" acting outside the thrombin-forming system proper. This is in accordance with the interpretation of Quick and Stefanini \((28)\) regarding L.F. and Lewis and Ferguson concerning Ac-globulin \((34)\).

The amount of L.F. in serum is considerably influenced by the presence of decalcifying anticoagu- lancs and the interval of time elapsing after coagulation. These facts must be considered in studies where these agents may be added to blood at various intervals after blood is shed in order to stop prothrombin conversion at various stages of the coagulation process. It is noteworthy that L.F. deteriorates more rapidly in oxalated than in citrated serum, as in plasma under the same condition.

All of these facts are pertinent to the determination of serum prothrombin and prothrombin consumption by the one-stage method. The orthodox procedure \((18)\), modified by the addition of fibrinogen, will give grossly inaccurate results for two reasons: \((a)\) During coagulation the L.F. disappears to a variable extent, which is considerably influenced by conditions of coagulation. Consumption of L.F. may reduce its concentration in serum to critically low levels, thus causing elevated prothrombin times. \((b)\) Also, the prothrombin conversion accelerator \((\text{spca})\) which evolves during coagulation varies in its concentration in serum, again depending upon various conditions of coagulation \((23)\). This clot-promoting factor increases the velocity of prothrombin conversion by thromboplastin and calcium, resulting in shorter prothrombin times than can be attributed to the prothrombin itself. It has already been shown that for this reason, one-stage determinations of the prothrombin in serum are inaccurate \((15)\). Now there is additional basis for this conclusion.

Difficulties also arise in the determination of serum L.F. by the one-stage procedure. Clearly, the substance is measurable by its restorative activity on aged plasma in which this clotting factor has largely deteriorated. Thus, the restorative effect of serum might validly be considered a measure of L.F. if we could be sure that during coagulation alterations did not take place in components other than prothrombin or L.F., which might influence the velocity of prothrombin conversion in aged plasma. That such alterations do occur, however, has been shown conclusively. Spca evolves which, in the presence of L.F., will accelerate thrombin evolution in stored plasma \((24)\). Seegers and his group had shown earlier that serum and serum
fractions activate and accelerate prothrombin conversion to a greater degree than plasma, and Owren has reported similar observations (12). Also, Milstone found that in the course of coagulation, "prothrombokinase" is converted to "thrombokinase" which is a potent accelerator of prothrombin conversion (35).

It is accordingly evident that some consideration must be given to clot-promoting agents which arise as a consequence of the coagulation process, and which may accelerate prothrombin conversion in aged plasma, simulating or enhancing L.F. activity. This is substantiated by the observations that BaSO₄ removes considerable restorative activity from serum despite the fact that this agent does not appreciably adsorb L.F. Since BaSO₄ removes spca (23) and thrombokinase (35), the effect on the restorative activity of serum is attributed to removal of this (these) substance(s). It should also be noted that BaSO₄ also adsorbs residual serum prothrombin. The restorative activity of BaSO₄ treated serum is, therefore, a more accurate index of serum L.F. than untreated serum.

The question arises whether serum Ac-globulin and Factor VI can be similarly excluded. According to Murphy and Seegers (36) and as has been confirmed by us (24), serum Ac-globulin is extremely unstable in normal human serum, virtually disappearing shortly after coagulation. The same is said to be true of Factor VI (12). This does not necessarily obviate the need for considering these factors because under abnormal conditions of coagulation they may be more stable, persisting in the serum. Indeed, substantial Ac-globulin activity is demonstrable in human serum prepared from blood allowed to clot slowly in siliconized tubes (36).

In a previous paper (24) evidence was presented which indicated that "serum" Ac-globulin probably comprises a mixture of plasma Ac-globulin (L.F.) plus spca, which together account for its greater prothrombin-converting activity than plasma Ac-globulin alone. The relatively low initial concentration of Ac-globulin in human plasma and its rapid depletion during coagulation explains the apparent extreme lability of "serum" Ac-globulin in human, in contrast to bovine, serum. On the basis of this interpretation, and assuming Factor VI to be identical with "serum" Ac-globulin, removal of spca by adsorption with BaSO₄ adequately takes care of clot-promoting activity of serum over and above that of its parent plasma as far as the restorative effect of serum on aged plasma is concerned.

In evaluating the restorative effect of serum as a measure of its L.F. content, consideration should also be given to the products of platelet disintegration which are said to accelerate prothrombin conversion in aged plasma (37, 38). It appears that platelets contain at least two clot-promoting substances, one which resembles Ac-globulin, and another which accelerates the interaction of thrombin with fibrinogen (39, 30). Whether these appear, and persist, in serum as a consequence of clotting is not clear. Pending further investigation, it may be concluded that the restorative effect of BaSO₄ serum on aged plasma provides a suitable method for measuring serum L.F. By comparing the prothrombin times of mixtures containing various dilutions of BaSO₄ serum in aged plasma with similar mixtures containing the BaSO₄ parent plasma, the L.F. of the serum in per cent of that originally present in the blood can be determined.

CONCLUSIONS

1. Normal serum has as much, and frequently more, restorative activity than the parent plasma on the retarded prothrombin conversion of aged plasma.

2. This is largely due to spca which evolves during coagulation and which, together with whatever Labile Factor remains unconsumed, accelerates prothrombin conversion.

3. Removal of spca by adsorption with BaSO₄ yields a serum which still has some restorative activity. This activity is a measure of residual L.F. in serum, which under normal conditions is small.

4. L.F. begins to disappear from blood shortly after its withdrawal, even before detectable changes in the prothrombin concentration have occurred. Within 40 minutes consumption of both factors is virtually complete. Disappearance of L.F. during coagulation is unrelated to the conversion of fibrinogen to fibrin.

*It has previously been shown that spca does not act on this phase of the coagulation process (22, 24).
5. Accelerating coagulation and prothrombin conversion by thromboplastin supplements increases the amount of L.F. consumed.

6. Conversely, curtailing the amount of prothrombin converted to thrombin, by dicumarol or by pathological conditions such as hemophilia, thrombocytopenia or thrombathen results in sera containing abnormally large amounts of L.F. This could be corrected in hemophilia by thromboplastin supplements, but not in dicumarol blood.

7. These observations lend support to the concept that L.F. is an active participant in the prothrombin conversion reaction.

8. Serum L.F. seems more stable in citrate than in oxalate, but not as stable as in native serum. This is in accordance with its behavior in plasma under similar conditions.

9. These findings are discussed insofar as they relate to methods for the determination of serum prothrombin, serum L.F. and prothrombin consumption under normal and pathological conditions.

REFERENCES


ASSOCIATION ANNOUNCEMENT

The 43rd annual meeting of the American Society for Clinical Investigation will be held at the Chalfonte-Haddon Hall, Atlantic City, N. J., on Monday, April 30, 1951, at 9 a.m.

Other meetings to be held also at the Chalfonte-Haddon Hall are:
- The 8th annual meeting of the American Psychosomatic Society, Saturday, April 28, 1951.
- The National Meeting of the American Federation for Clinical Research, Tuesday, May 1, 1951.
- The 64th meeting of the Association of American Physicians, Tuesday, May 1, and Wednesday, May 2, 1951.