A METHOD FOR DEPROTEINIZATION OF BLOOD AND OTHER BODY FLUIDS

BY

G. HUNTER

From Cowley Road Hospital, Oxford

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Deproteinization is a necessary step in many procedures for the chemical analysis of body fluids. A variety of reagents have been used for this purpose, especially in blood analysis. Of these perhaps tungstic acid and trichloroacetic acid are the most widely used to-day, and for this reason we may consider them a moment. Folin and Wu (1919) introduced tungstic acid to prepare protein-free filtrates for their system of blood analysis. In this method it is usual to employ 667 m.Eq. of the acid to precipitate the proteins from a litre of plasma or whole blood and twice this concentration to deproteinize the same volume of packed erythrocytes. With the use of sodium tungstate and sulphuric acid equivalent amounts of Na and SO₄ are of course added to the filtrate. Trichloroacetic acid is commonly used in a 5% (w/v) concentration in the filtrate, and it must be used in not less than half of this concentration. If the blood is diluted 1 in 10 this means that the acid used equals 4 m.Eq./litre blood and the filtrate is more than 3.0 N acid.

For such reasons tungstic and trichloroacetic acids are unsuitable deproteinizing agents in many analytical methods, in procedures for the isolation of solutions for chromatography, and other modern techniques. It therefore seems desirable to have other means of deproteinization that will avoid the addition of gross amounts of extraneous matter to the filtrates, that can be rapidly applied, yield filtrates near neutrality, and cause a minimum distortion of the chemical picture of the fluids concerned. The present communication is an attempt in this direction. A method is described for the preparation of protein-free filtrates of blood, which is also applicable to other body fluids. It depends on the adjustment of the pH of the diluted fluid in question to a value suitable for the precipitation by heat of the proteins present. Plasma, cells, and whole blood require about 50, 27, and 40 m.Eq. acetic acid/l. respectively for pH adjustment. The filtrates obtained after heating briefly at 100° C. are clear and colourless and remain clear on addition of sulphosalicylic acid.

Deproteinization of Blood

pH Adjustment.—For this purpose 0.05 N-acetic acid is used routinely, but other acids may be used (see p. 163). One volume of plasma, cells, and whole blood requires 1.0, 0.54, and 0.80 volume respectively of the acetic acid. Water is added to suitable dilution, e.g., 1 in 5 to 1 in 50, and the contents are mixed.

Protein Precipitation.—When the volume of the mixed solution is less than 10 ml it is convenient to have it in a centrifuge tube. This is immersed for three to five minutes in boiling water, then cooled, spun, and the supernatant fluid is transferred to a clean tube.

With larger volumes of course the time of heating must be extended so that the fluid is at 100° C. for two to four minutes.

Deproteinization of Other Fluids

Cerebrospinal fluid (C.S.F.) protein is satisfactorily precipitated with 0.7 volume of the acid at 1 in 10 dilution. In our experience protein is more easily removed from C.S.F. by this method than by either tungstic or trichloroacetic acid.

Oedema fluids, nasal discharges, etc., have been found to require 0.4 to 0.7 volume of the acid, the amount depending largely on the amount of NaHCO₃ and protein present. When the coagulation is not complete the solution should be adjusted to pH 5.3, at which point we have not failed to get good coagulation. When means are not available for pH determination a final adjustment with a drop or two of 1.0 M-acetic acid-sodium acetate buffer pH 5.3 is usually effective.

Deproteinization of Urine

The composition and pH of urine are so variable that no rule can be followed. Our practice is to adjust the pH of a given volume of urine with a measured amount of 1.0 N-acetic acid to 5.2 to 5.3, using a pH meter and glass electrode, then diluting as analytical requirements indicate before heating to precipitate the protein.
Methods

Non-protein nitrogen was determined on the blood filtrates shown in Table I by digestion with H$_2$SO$_4$, addition of K$_2$S$_2$O$_8$ to clear, followed by heating for 30 min., with subsequent nesslerization.

Acid required for the pH shift of Tables II and III was determined by titrating 1 ml. plasma, diluted with 9 ml. water, with 0.05 N-acetic acid to pH 5.3 with the glass electrode; and 1 ml. cells, diluted with 19 ml. water, to pH 7.0 with the same acid.

<table>
<thead>
<tr>
<th>TABLE I NON-PROTEIN NITROGEN OF BLOOD DETERMINED ON CORRESPONDING FILTRATES OBTAINED WITH TUNGSTIC ACID AND THE ACID-HEAT METHOD</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blood No.</strong></td>
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<tr>
<td>1</td>
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<td>4</td>
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<td>5</td>
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</tbody>
</table>

Results are expressed in mg./100 ml.

<table>
<thead>
<tr>
<th>TABLE II ESTIMATE OF ACID* REQUIRED TO SHIFT pH OF 1 LITRE PLASMA TO 5-3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein, 70 x 0.243 (Peters and van Slyke, 1931) = 17 m.Eq. NaHCO$_3$, at pH 5.3 (4 m.Eq. as NaHCO$_3$) = 24 m.Eq. Total by estimate = 41 m.Eq. Found in practice = 40 m.Eq.</td>
</tr>
</tbody>
</table>

* It is assumed that plasma contains 70 g. protein and 28 m.Eq. NaHCO$_3$, 40 m.Eq. strong acid is equal to about 50 m.Eq. acetic acid at pH 5.3.

<table>
<thead>
<tr>
<th>TABLE III ESTIMATE OF ACID REQUIRED TO SHIFT pH OF 1 LITRE LAKED CELLS FROM 7.5 TO 7.0</th>
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</thead>
<tbody>
<tr>
<td>About 2.5 m.Eq. acid required to shift 17 g. Hb through 1 pH unit (Peters and van Slyke, 1931). Hence acid required here is 2.5 x 20 = 50 m.Eq. To shift 16 m.Eq. NaHCO$_3$ from pH 7.5 to pH 7.0 requires 320 x 5 = 1600 m.Eq. 320 x 2.5 x 0.5 = 23.5 m.Eq. Total by estimate = 17 m.Eq. Found in practice = 24 m.Eq.</td>
</tr>
</tbody>
</table>

* It is assumed that cells contain 320 g. haemoglobin and 16 m.Eq. NaHCO$_3$.

Results

The filtrates obtained are clear and colourless and remain clear on addition of sulphosalicylic acid. This is conclusive evidence that there is less than about 2 mg. protein/100 ml. filtrate. The xanthoproteic and biuret tests are faintly positive in the filtrates as they are also in corresponding tungstic acid filtrates and do not necessarily signify the presence of protein. For practical purposes the filtrates are therefore free from protein.

Though free from protein, the non-protein nitrogen (N.P.N.) content of the filtrates from plasma, cells, and whole blood is uniformly higher than the N.P.N. values obtained on corresponding tungstic acid filtrates, as shown in Table I.

The pH of the filtrates is somewhat higher, 0.1 to 0.2 pH unit, than that of the solutions before heating, due to loss of CO$_2$.

Jaundiced sera yield colourless filtrates indicating that, of the pigments, at least bilirubin is adsorbed on the protein precipitate.

Discussion

This method of deproteinization has proved practicable for the determination of ergothioneine in blood (Hunter, 1949), polyvinyl pyrrolidone (plasmosan) in blood serum (Campbell and Hunter, 1953), and for the estimation of isonicotinic acid hydrazide in blood serum (Hunter, 1955). It has now been found that filtrates from blood and other body fluids prepared in this way are suitable for the determination of Mg and Ca (Hunter, 1956).

The "heat and acetic acid test" has of course long been used as a qualitative clinical test for protein in urine, but it is usually performed rather haphazardly, without control of pH, and to differentiate protein from precipitated phosphates.

The method has not been suggested as a general and quantitative means for deproteinizing blood and other body fluids, though there are a few observations in the literature on the controlled use of acetic acid, e.g., by Rimington (1940) to prepare filtrates of plasma containing mucoprotein, by Solomon, Johnson, Sheffner, and Bergeim (1951) to prepare tissue filtrates for chromatography, and by Ramsay (1953), who used an acetic buffer at pH 5.0 to remove protein in the course of the determination of Fe in plasma and serum. As far as I am aware there has been no study of this method of deproteinization.

The arbitrary finding that plasma requires the addition of almost twice as much acid as an equal volume of packed cells was at first surprising in view of the much greater concentration of protein and base in the cells. It was observed, however, that the protein-free filtrates from plasma or serum had a pH about 5.3, and the protein-free filtrates from packed cells had a pH about 7.0 at room temperature, pHs not far removed from the isoelectric points of denatured serum albumin and globulin and the isoelectric point of haemoglobin respectively. Postulating a normal plasma as containing 70 g. protein and 28 m.Eq. NaHCO$_3$, the amount of strong acid necessary to shift its pH to 5.3 is calculable with fair accuracy, according to Table II, and agrees in a satisfactory manner with the amount found in practice. The NaHCO$_3$ present in plasma accounts for more than half the acid required.
As might be expected the pH value of the acid used has some effect on the amount required in different pH regions. For example, there is little difference in the range pH 7.5–7.0, whether hydrochloric or acetic acid is used, but when the pH has to be shifted to about 5.3, more of the weak acid is required. The following amounts (in ml.) of different acids were required to shift 1 ml. of a diluted plasma to pH 5.3: Acetic 1.00, benzoic 0.88, lactic 0.80, perchloric, trichloroacetic and hydrochloric 0.75, tartaric 0.85, sulphuric 0.80, and citric 1.05. Similar values are obtained on titration of bicarbonate solutions. On heating the plasma solutions clear filtrates were obtained. It should be noted, however, that there is a marked rise in the pH of trichloroacetic acid filtrates owing to the decomposition of the acid on heating.

In the case of cells the situation is quite different, as seen from Table III. The effect of NaHCO₃ is almost negligible over the relevant pH range, but to shift the haemoglobin over 0.5 pH takes more acid than that required to shift the plasma protein over 2.0 pH. Again the acid calculated as required is near that found in practice, especially when a small amount of plasma with the cells is accounted for.

What remains surprising is that the proteins in whole blood are effectively precipitated at about pH 6.4. When a 1 in 10 dilution of plasma is heated at this pH it will not coagulate well; the supernatant will be opaque or very milky in appearance. Likewise solutions of cells heated at pH 6.4 give dirty red suspensions. Yet when both cells and plasma are present in about equal volume the supernatant is clear and colourless. At pH 6.4 the plasma proteins will be charged negatively and the haemoglobin positively, and it is suggested on those grounds that coprecipitation is a major factor in the coagulation of the denatured protein from solutions of whole blood.

**Range of pH Tolerance for Deproteinization.**—It may be observed that pH 5.3 has been chosen in Table II to calculate the buffering capacity of plasma. This point was chosen as a mean of a range pH 5.7–5.0, in which crystal clear filtrates of plasma are usually obtained. This range is of considerable importance, as the present method would not otherwise be practicable for different bloods. Expressed in another way, it is found that 1 ml. diluted normal plasma requires 1.0 ml. 0.05 N-acetic acid for adjustment to the pH suitable for deproteinization by heat. Were 0.9 ml. acid added, the pH in such a diluted plasma would be near 5.7, and were 1.1 ml. acid added the pH would be near 5.0. We thus have a range of tolerance of ±10% of the amount of acid recommended for use. This covers all but quite exceptional plasmas—more than a 20% change in plasma protein or more than a 15% change in plasma bicarbonate.

Like arguments apply to the deproteinization of cells and whole blood.

It may be noted that the loss of CO₂ from blood has no influence on the amount of acid required for pH adjustment of the whole blood, as it is a volatile acid. However, plasma separated from blood with a low CO₂ tension will have more chloride present and thus require less acid for pH adjustment, and vice versa for plasma separated from blood with a high CO₂ tension. Only under extreme conditions does this chloride shift exceed the range stated above.

Despite the range of tolerance recognized here it should be emphasized that the amounts of acid used must be measured precisely.

The general directions are only applicable to fresh bloods. On long standing, acid will frequently develop in blood.

**Fluids Other Than Blood.**—The principles discussed above for the precipitation of the proteins from blood plasma would appear to apply to body fluids in general. In the case of C.S.F. the protein is normally a negligible factor as a buffer, but the total CO₂ is somewhat higher than that of plasma. Hence we might expect that unit volume will require about 0.7 volume of 0.05 N-acetic acid. Cerebrospinal fluid also serves to show the sensitivity of the process. For example, 1 ml. of a C.S.F. containing 22 mg. protein/100 ml., with 8.3 ml. water and 0.7 ml. acetic acid, showed a well-coagulated precipitate after heating.

When the acid requirement of the fluid is unknown and its volume perhaps small, or pH determinations are not readily made, it is suggested that 0.6 volume of acid be tried, and if this fails to give a good coagulation a drop or two of acetate buffer at pH 5.3 may be used. As noted above, Ramsay has suggested the use of a similar buffer at pH 5.0, but it may be calculated from his data that he is adding some 25 times more acetate than used in the present method.

In a case of nephrosis studied with Dr. L. C. A. Nunn of the Pathology Laboratory, Stoke Mandeville Hospital, Aylesbury, the present method was found useful in the precipitation of urinary protein without the precipitation of polyvinyl pyrrolidone ("plasmosan") which had been administered to the patient to maintain oncoltic pressure. It was thus possible to determine both the protein and plasmosan excreted in the urine.
Non-protein Nitrogen.—The non-protein nitrogen values of the filtrates obtained by the present method present a point of some interest. Its elucidation awaits further investigation. However, it was stated by Benedict and Newton (1929) that tungstic acid precipitates ergothioneine in blood, and from similar findings the present method of deproteinizing (Hunter, 1949) was developed. It would appear that non-protein substances other than ergothioneine are involved, and it would no doubt be of some interest to find out what they are.

It appears unlikely that any proteolysis occurs, even in plasma at pH 5.3, in the short heating time. Extension of the heating time to 30 min. was found to have no effect on the non-protein nitrogen.

Summary

A method is described for the deproteinization of blood and other body fluids, at the usual dilutions of 1 in 5 to 1 in 50, which depends on the addition of small amounts of acid followed by a brief period of heat at 100° C.

The requisite amount of acid is calculable from the buffer capacity, due mainly to NaHCO₃ and protein, and the pH shift involved.

The filtrates obtained are clear and colourless and show no clouding on the addition of sulphosalicylic acid, but they contain more non-protein nitrogen than corresponding filtrates obtained with tungstic acid.

Some of the factors in the deproteinization process are discussed and the advantages of such filtrates for analytical and other purposes are indicated.

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G. Hunter

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