Clotting of Citrated Plasma and Citrate Utilization by Intestinal Gram-negative Bacilli

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SUMMARY: Salmonella, shigella, alkalescens-dispar, proteus, paracolon and coliform strains were tested for the presence of a coagulase enzyme as manifested by the ability to clot plasma. The shigella and proteus strains examined gave negative results; a high percentage of the remaining strains clotted citrated plasma but this reaction was due to metabolism of citrate and subsequent liberation of calcium ions and not to a coagulase enzyme. Results depended on the bacterial strain and on the kind and dilution of plasma used. Utilization of citrate in citrated plasma was compared with the ability of the tested strains to attack citrate in Koser’s ammonium citrate medium and in sodium citrate peptone medium. The Shigella and Proteus species examined failed to utilize citrate in any of the media under the conditions of the tests, Salmonella species gave fairly uniform results in the three media, while paracolon, coliform and alkalescens-dispar strains differed in their activity, the highest percentage of positive reactions being obtained in plasma-containing media.

In the course of a study of Gram-negative bacilli these organisms were examined to see whether they possessed coagulases capable of clotting plasma. A survey of literature showed that numerous bacterial species have been examined for plasma-coagulating ability. Conflicting results have been obtained, due possibly to different experimental conditions. In the present communication only intestinal Gram-negative bacilli will be considered.

Loeb (1903) used aseptically drawn goose-blood, which was centrifuged and diluted 1/10. He noticed the strong coagulating power of Staphylococcus aureus and that Salmonella typhi had no apparent effect, while Pseudomonas aeruginosa and Escherichia coli had some coagulating ability. Much (1908), in experiments with human plasma and citrate, fluoride or hirudin as anticoagulants, reported negative results with Sal. typhi, E. coli, Friedländer’s bacillus and Proteus spp. Kemkes (1928) published negative findings in tests with oxalated plasma and the following organisms: Ps. aeruginosa, Ps. fluorescens, Sal. typhi, Shigella flexneri and Sh. dysenteriae. Lusena (1929) noted that some Sal. typhi cultures coagulated decalcified plasma. Gross (1931), found various bacteria, Sal. typhi, Sal. paratyphi, Ps. aeruginosa, shigella and proteus strains, negative in their action on rabbit plasma. Fisher (1936) investigated the coagulating properties of staphylococci on oxalated and citrated plasma and observed that contaminants such as some strains of Bacillus subtilis and Ps. aeruginosa caused clotting of rabbit but not human plasma, while Friedländer’s bacillus, Alcaligenes sp., E. coli and Aerobacter aerogenes did not clot either. Reimer (1936) used various kinds of plasma without addition of anticoagulant and compared the clotting time of plasma mixed with bacterial cultures with that of control plasma. He found no
analogy between activity on human and other animal plasmas; *E. coli* accelerated clotting of human plasma, while *Sal. typhi* and *Sal. paratyphi-B* inhibited clotting. Fredericq (1942) made experiments with oxalated plasma and recorded that *Ps. aeruginosa*, *Ps. fluorescens* and *B. subtilis* had a coagulating effect, that of *Ps. aeruginosa* being the strongest.

Rita (1945a, b) recorded clotting of citrated plasma by *Sal. typhi* and other species of *Salmonella* but found no action on plasma to which oxalate or fluoride had been added. This writer concluded that strains which utilized citrate released calcium ion and thus caused plasma clotting. Rita (1945c), in tube experiments, noticed also the slow (8–10 hr.) clotting caused by salmonellinas as compared with the quick (1–3 hr.) reaction caused by staphylococci.

Harper & Conway (1948) investigated 8 strains of coliform and 5 of paracolon bacilli and found that 11 of the 18 strains clotted human citrated plasma at 37° within 18 hr.; the clotting reaction was proved to be due to the breakdown of citrate with liberation of calcium ion and not to a coagulase. However, Krech (1952) demonstrated an active agent in filtrates of *E. coli*. He used citrated plasma of man, horse, cow and guinea-pig and found that horse plasma was most actively coagulated and that a temperature of c. 16° was the important factor in the test.

The present communication describes the investigation of a collection of Gram-negative intestinal bacilli, including salmonella, shigella, alkalescens-dispar, proteus, coliform and paracolon strains, with regard to clotting action on citrated and heparinized plasma and citrate utilization.

**MATERIALS AND METHODS**

*Organisms.* The following strains were examined: 52 salmonellas, 34 shigellas (7 *Sh. dysenteriae*, 18 *Sh. flexneri*, 7 *Sh. boydii*, 2 *Sh. sonnei*), 4 alkalescens-dispar group, 53 proteus (23 *Pr. mirabilis*, 25 *Pr. morganii*, 5 *Pr. vulgaris*), 140 paracolons (91 *Paracolobactrum coliforme* type I, 9 *P. coliforme* type II, 12 *P. intermedium* type I, 5 *P. intermedium* type II, 10 *P. aerogenoides* type I, 11 irregular citrate-negative and 2 irregular citrate-positive) and 14 coliforms (*E. coli* type I). Salmonella strains originated from the National Collection of Type Cultures, shigella and alkalescens-dispar strains were sent by Dr Ewing, U.S.A., while proteus, paracolon and coliform strains were from our own collection isolated from normal and abnormal faecal specimens and classified according to *Bergey’s Manual of Determinative Bacteriology* (1948).

*Methods.* Rabbit and human plasmas were tested. Rabbit blood was collected aseptically by cardiac puncture and mixed in the proportion 9 ml. blood with 1 ml. of 10% (w/v) sodium citrate (final concentration 1%). Human plasma was obtained from a blood bank and contained 4 vol. blood to 1 vol. of 2% (w/v) sodium citrate (final concentration 0.4%). In tests with heparinized plasma 4 drops of the anticoagulant were added to 20 ml. citrated human plasma. The citrate media used included Koser’s (1924) ammonium citrate solution and citrate in Bacto-peptone basal medium (Kauffmann, 1951).

Slide coagulase tests were performed with undiluted citrated rabbit plasma. For tube coagulation experiments rabbit plasma was diluted 1/10 or 1/5
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with normal saline; the human plasma was diluted 1/4 or 1/2 or used undiluted. In tube tests 0.5 ml. of appropriate plasma was added to 0.5 ml. of an 18 hr. culture of the required organism in Hartley’s tryptic broth. The results were recorded after 24 hr. incubation at 37°C; negative tubes were kept under observation for 3 days.

Each batch of plasma was tested for clotting capacity by adding 1 drop of 5% calcium chloride to 0.5 ml. plasma. Control slide and tube coagulase tests were included in which known coagulase-positive and coagulase-negative strains of staphylococci were used.

Utilization of citrate in Koser’s medium at 37°C was considered to be positive when growth was evident after a third successive subculture by inoculation with a straight wire at 48 hr. intervals. Utilization of citrate in the citrate peptone medium was assessed with lead acetate indicator after incubation of the cultures in triplicate sets at 37°C for 2, 7 and 14 days.

RESULTS

Clotting of citrated plasma. The slide coagulase tests were uniformly negative. Occasionally granularity slowly took place which at first glance could have been mistaken for clotting, but after some practice it became evident that late and doubtful reactions were non-specific and should be recorded as negative.

Table 1. Clotting reactions with citrated and heparinized plasma caused by Gram-negative organisms

<table>
<thead>
<tr>
<th>Citrated plasma</th>
<th>Heparinized plasma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rabbit*</td>
<td>Human†</td>
</tr>
<tr>
<td>Plasma dilution</td>
<td></td>
</tr>
<tr>
<td>1/10</td>
<td>1/5</td>
</tr>
<tr>
<td>1/4</td>
<td>1/2</td>
</tr>
<tr>
<td>Undiluted</td>
<td>1/2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Strains</th>
<th>No. of strains tested</th>
<th>0.1</th>
<th>0.2</th>
<th>0.1</th>
<th>0.2</th>
<th>0.4</th>
<th>0.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salmonella</td>
<td>52</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>Shigella</td>
<td>34</td>
<td>45</td>
<td>7</td>
<td>46</td>
<td>6</td>
<td>45</td>
<td>46</td>
</tr>
<tr>
<td>Alkalescens-dispar</td>
<td>4</td>
<td>4</td>
<td>0</td>
<td>4</td>
<td>0</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Proteus</td>
<td>53</td>
<td>0</td>
<td>53</td>
<td>0</td>
<td>53</td>
<td>0</td>
<td>53</td>
</tr>
<tr>
<td>Paracolon</td>
<td>140</td>
<td>92</td>
<td>48</td>
<td>101</td>
<td>39</td>
<td>2</td>
<td>138</td>
</tr>
<tr>
<td>Coliform</td>
<td>14</td>
<td>14</td>
<td>0</td>
<td>14</td>
<td>0</td>
<td>14</td>
<td>0</td>
</tr>
</tbody>
</table>

* Diluted 1/10 in 10% (w/v) sodium citrate.
† Diluted 1/5 in 2% (w/v) sodium citrate.
+ = clot; − = no clot.

Table 1 shows that none of the strains investigated clotted heparinized plasma, thus showing the absence of coagulase under the conditions of the test. This was confirmed by tests with oxalated rabbit plasma; in no instance was a positive reaction obtained.
All shigella and proteus strains were uniform in their inability to clot citrated plasma. All the *E. coli* and alkalescens-dispar strains and the majority of salmonella and paracolon strains clotted citrated plasma within 24 hr. With citrated human plasma an increase in the number of positive readings was obtained with 1/2 dilution of plasma (citrate concentration 0·2%) as compared with 1/4 dilution, but with undiluted plasma only one paracolon strain gave an additional positive reaction. Citrated rabbit plasma was clotted by most strains at 1/10 dilution (citrate concentration 0·1%) and only a few more strains clotted plasma at 1/5 dilution. With a few exceptions strains clotted both citrated human and rabbit plasma; under the conditions of test 1 salmonella and 10 paracolon strains gave positive readings with human plasma only, while 1 salmonella and 3 paracolon strains acted only on rabbit plasma. Thus the ability to clot citrated plasma depended on the bacterial strain and also on the dilution and kind of plasma used, the concentration of citrate being the important factor. Harper & Conway (1948) showed that clotting takes place when the concentration of citrate is less than that necessary for binding free calcium ions. The clot produced in our tests was soft, occasionally with a slight precipitate and unlike the fibrin-thread type of staphylococcal coagulum which retracts from the walls of the tube.

Additional tests were performed to find out whether a change of pH value was responsible for plasma clotting. Control plasma + sterile broth had a pH of 7·2. Clotted plasma cultures were shaken to break up the clot and the pH measured by means of a glass-electrode. Various cultures showed an increase of pH value to 7·9-8·4; no clotting occurred in control media when the pH was adjusted to 8·4.

Of 140 paracolon strains tested, 16 liquefied gelatin; of these 9 did so within 1–2 days, but the remaining 7 strains acted more slowly. Eight active gelatin-liquefying strains did not clot plasma. Tests with these strains were repeated and the tubes examined every hour, but clot formation followed by solution was not observed.

Table 2. Utilization of citrate in various media

<table>
<thead>
<tr>
<th>Strains</th>
<th>No. of strains tested</th>
<th>Clotting of plasma</th>
<th>Koser media</th>
<th>Citrate peptone medium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>No.</td>
<td>%</td>
<td>No.</td>
</tr>
<tr>
<td>Salmonella</td>
<td>52</td>
<td>47</td>
<td>90</td>
<td>46</td>
</tr>
<tr>
<td>Shigella</td>
<td>34</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Alkalescens-dispar</td>
<td>4</td>
<td>4</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>Proteus</td>
<td>53</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Paracolon</td>
<td>140</td>
<td>111</td>
<td>80</td>
<td>31</td>
</tr>
<tr>
<td>Coliform</td>
<td>14</td>
<td>14</td>
<td>100</td>
<td>0</td>
</tr>
</tbody>
</table>

Citrate utilization. Table 2 shows the results of experiments on citrate utilization in various media: human and rabbit citrated plasma, Koser's ammonium citrate medium and citrate peptone medium. All the *Shigella* and
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three of the *Proteus* species (*Pr. rettgeri* was not tested) gave negative results in all media. Some proteus strains (*Pr. mirabilis* and *Pr. vulgaris*) showed growth in Koser's medium on first subculture, but growth was not maintained on the second transfer. Readings for the salmonellas ran parallel in all three media, with two exceptions: of 52 strains tested, 45 utilized citrate in all three media and 5 strains (*Sal. paratyphi-A* var. *durazzo*, *Sal. abortus-equi*, *Sal. sendai* and 2 strains of *Sal. typhi*), gave negative results; 1 strain of *Sal. typhi* attacked citrate in rabbit plasma only; *Sal. carrau* utilized citrate in all media except rabbit plasma. Paracolon, coliform and alkalescens-dispar strains gave a much higher percentage of positive reactions with citrated plasma than with Koser's or citrate peptone media. Only 2 paracolon strains utilized citrate in Koser's or citrate peptone media without being able to do so in plasma. Of the 29 paracolon strains which did not clot plasma, 20 were *Paracolobactrum coliforme* type I, one was *P. coliforme* type II, one was *P. aerogenoides* type I, one was *P. intermedium* type II, and 6 were irregular citrate-negative types.

To assess the clotting time of rabbit plasma (diluted 1/10), tests were carried out with a number of paracolon and coliform strains (Table 3). Only one paracolon strain produced a soft clot after 2 hr. incubation, the majority of paracolon and coliform strains gave positive reactions after 10 hr.; no additional positive results were recorded after 21 hr. In contrast to the slow action of the majority of paracolon and coliform strains, our control coagulase-positive staphylococci coagulated plasma in tube tests after 2 hr.

Table 3. The effect of incubation time on clotting reactions

<table>
<thead>
<tr>
<th>Strains</th>
<th>No. of strains tested</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>10</th>
<th>21</th>
<th>24</th>
<th>48</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paracolon</td>
<td>78</td>
<td>0</td>
<td>1</td>
<td>8</td>
<td>14</td>
<td>51</td>
<td>66</td>
<td>66</td>
<td>66</td>
</tr>
<tr>
<td>Coliform</td>
<td>13</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>10</td>
<td>13</td>
<td>13</td>
<td>13</td>
</tr>
</tbody>
</table>

DISCUSSION

False positive coagulase reactions with citrated plasma by some salmonella strains (*Rita, 1945a, b*) and by some paracolon and coliform strains (*Harper & Conway, 1948*) were shown to be due to utilization of citrate with liberation of calcium ion. Evans, Buettner & Niven (1952) observed that a similar reaction accounted for the clotting of plasma by certain streptococci.

Our experiments with Gram-negative intestinal bacilli show that the majority of salmonella, paracolon, coliform and alkalescens-dispar strains which clotted human or rabbit citrated plasma did so because of citrate breakdown. On the other hand, all *Shigella* species and three *Proteus* species did not cause clotting. Lominski, Conway, Harper & Rennie (1947) pointed out that some *E. coli* strains which failed to attack citrate in Koser’s medium (ammonia as source of nitrogen) attacked citrate in the presence of a different source of nitrogen, e.g. peptone. The *Shigella* and *Proteus* species examined
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here gave negative results in the three citrate media; this may be helpful in
diagnostic work, e.g. when it is difficult to classify an organism as shigella or
paracolon, a positive citrate test would exclude shigella. *Pr. mirabilis* and
*Pr. vulgaris* strains were reported (Rustigan & Stuart, 1945; Cook, 1948;
*Bergey's Manual, 1948*) to utilize citrate, but Kauffmann (1951) listed these
species as citrate-negative. It is possible that proteus strains of faecal or
non-faecal origin may differ in citrate utilization; our cultures were of faecal
origin. It is clear from our findings that attention should be given to the
conditions and details of these tests; in our experiments with Koser's medium
a positive reading was recorded when growth appeared after three successive
subcultures from inoculation with a straight needle at 48 hr. intervals.
Rustigan & Stuart (1945) observed that proteus strains grew at different rates
on citrate medium. Cook (1948) mentioned that he inoculated Koser's citrate
media with a straight wire. The published results, therefore, emphasize the
need to standardize citrate utilization tests.

Rita (1945c) noticed that *Sal. typhi* strains varied in ability to clot plasma,
and suggested that the division of *Sal. typhi* into strains which clotted or did
not clot citrated plasma might be useful in epidemiological investigations.
Corda (1947) disagreed with this basis for differentiation because of the
instability of the character. Our experiments also showed the unreliability of
this criterion since results occasionally varied with the batch of plasma used.
Krech (1952) described some *E. coli* strains which possessed an active agent
in bacteria-free filtrates which was capable of coagulating plasma at room
temperature but not at 37°. This, apparently, is a different reaction from that
investigated in our study. The ability of *Ps. aeruginosa, Ps. fluorescens,
B. subtilis, Chromobacterium prodigiosum* and actinomycetes to clot plasma is
attributed by Fredericq (1942, 1946a, b) to a different factor, namely a
secretion of coagulase at 37°. Our work supports the suggestion of Harper &
Conway (1948) and of Evans et al. (1952) that, to avoid false positive coagulase
reactions, tests should be performed with plasmas which contain anticoagu-
lants other than citrate, e.g. oxalate or heparin.

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