THE TOXIC FACTORS IN EXPERIMENTAL TRAUMATIC SHOCK.
IV. THE EFFECTS OF THE INTRAVENOUS INJECTION
OF THE EFFUSION FROM ISCHEMIC MUSCLE

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The fundamental hemodynamic disturbances
which characterize traumatic shock may be ini-
tiated by at least 3 different factors: gross hemorr-
hage or loss of plasma into a traumatized region,
neurogenic and psychogenic elements, and the
absorption of toxic materials from traumatized
regions. Of these etiologic factors, the local loss
of vascular fluid is generally conceded to be of
greatest importance (3, 28, 31, 54), while the
others can be at best but contributory.

Since the original classical experiments of Can-
non and Bayliss (1), the theory of traumatic
toxemia has been furthered by two types of ex-
perimental evidence. In one, a state more or less
resembling clinical shock is produced in animals
by the parenteral administration of various ex-
tracts of normal or traumatized tissues or by the
intraperitoneal implantation of tissue. In the
other, attempts are made in various ways to con-
trol or limit local fluid loss and neurogenic factors
in experimental traumatic shock and to demon-
strate that an additional factor, presumably tra-
matic toxemia, is necessary to explain the results.
This approach has been considered in another
publication (55) and the present discussion will
concern itself only with the first type of
experiment.

Many experiments have been reported in which
extracts of various tissues were found to have
definite vasodepressive or shock-producing prop-
erties (56 to 61). Certain techniques (62, 63)
consist of introducing into the peritoneal cavity of
dogs whole or pulped liver or muscle. Observed
regularly were hemoconcentration, circulatory col-
apse, and death within 24 hours. Furthermore,
at autopsy the viscera consistently showed con-
gestion, edema, and gastrointestinal hemorrhage.
More recently, shock was observed following the
parenteral injection of extract of striated muscle,
and adenosine triphosphate was implicated as the
constituent responsible for these effects (64).

In the experiments cited, there have been no
cardiologic studies, and in only a few has any
attempt been made to prevent bacterial contami-
nation during the course of the experiment. The
inability to reproduce the results of some of these
experiments, when strict aseptic technique was
used (65), raises the question as to how many of
the positive results reported may have been due
to extraneous bacterial contamination.

In many of the cited experiments the criteria
of shock do not appear conclusive. They consist
either of death of the animal, certain autopsy find-
ings, a fall in blood pressure, or combinations of
the three. In some experiments, hematocrit
studies have been done in addition and these have
usually shown a marked hemoconcentration.
Death and hypotension are hardly pathognomonic
of shock, while recent studies in clinical traumatic
shock without evident hemorrhage have revealed
a falling hematocrit (66).

Dosage, rate, and route of administration are of
paramount importance in evaluating the toxicity
of any substance, yet these factors appear to have
received insufficient attention in experiments on
the toxicity of tissue extracts. If one may convert
to clinical terms the experiments of one investi-
gator (64), it would require that a large part of
the muscle tissue of the body be ground, extracted,
and administered in one dose in order for shock
to follow. It would be desirable to determine the
rate at which substances might be absorbed from
a traumatized region and to reproduce that rate

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search, between the Office of Scientific Research and De-
velopment and the Massachusetts General Hospital.
in the administration of tissue extracts. It is not sufficient to demonstrate that substances obtainable from traumatized tissue are toxic; it must be clearly shown that they are toxic in the form, in the dose, and at the rate of absorption which might conceivably occur in clinical traumatic shock.

In view of these considerations, the remarkable thing about the injection of tissue products is not that they are toxic but that they are so often found to be innocuous. In recent experiments with lymph from traumatized limbs, noxious effects were produced in recipient animals in only 30 per cent of the trials (67). In one series of recipients, normal lymph was found to possess toxic properties in 50 per cent of cases (68). The consistently negative results which have been obtained by another group when bacterial contamination is prevented have already been mentioned (65). Extracts of traumatized limbs were found to cause no blood pressure fall or any untoward effects when administered intravenously in excessive dosage (69). This observation is particularly difficult to reconcile with the thesis that significant quantities of shock-producing factors may be liberated from traumatized regions.

The theory of traumatic toxemia can, therefore, hardly be called established, and it seems that further experimental work is warranted. The muscle ligation technique previously described (14) affords a method for obtaining in large quantities the fluid which effuses from an area damaged by prolonged ischemia. Chemical examination of this fluid (70) indicates that it represents plasma which passes through the damaged capillaries and muscle tissue carrying with it products of cellular disintegration. It seemed reasonable that toxic factors if they exist should be present in such fluid and that a pharmacodynamic study of the effects of this fluid on intravenous administration might be of value.

METHODS

The operation of muscle ligation, release after 5 hours, and fluid collection as described previously (14) were performed with sterile precautions. The fluid which collected during the 5-hour period following release of the ties was centrifugalized and the supernatant fraction frozen and preserved on carbon dioxide snow until used. Before administration, the fluid was thawed, filtered through a few layers of gauze, warmed to room temperature, and administered intravenously over a period of 15 to 30 minutes. The dose given which approximated 14.7 ml. per kgm. exceeded slightly the average output (12 ml. per kgm.) of the 35 donor animals in this series. The animals used were mongrel dogs in apparently normal health, anesthetized with sodium pentobarbital. Blood pressure was recorded continuously by a mercury manometer connected to a carotid cannula containing dilute heparin or 2.5 per cent sodium citrate solution. Cardiac output was determined at intervals using the Fick principle as first used in shock by two investigators (71). Mixed venous blood was obtained from a catheter passed down one external jugular vein with the tip either in or at the level of the right auricle. Arterial blood was taken from a femoral artery by puncture. The bloods were collected and preserved under oil, and oxygen content was determined by the manometric method of Van Slyke and Neill. Oxygen consumption was measured by a calibrated recording spirometer connected to a tracheal cannula. Carbon dioxide was removed by soda lime in the system. Relative total peripheral resistance was calculated from the relationship:

\[
\text{Relative total peripheral resistance} = \frac{\text{Mean arterial B. P.}}{\text{Cardiac output}}
\]

No attempt was made to calculate the resistance in absolute units since only changes from control values were desired, but in order to permit better comparison between animals of different weights, cardiac outputs were substituted in the formula in terms of liters per minute per kgm. of body weight.

Mixed venous blood hemoglobin concentration was determined photocoelometrically. When used to measure the trend of hemococoncentration in a given experiment this determination possesses the advantage over erythrocyte hematocrits in that it is independent of changes in erythrocyte size and of such variables as speed and duration of centrifugation, plasma viscosity, and specific gravity. Its use as a gross index of plasma loss seems warranted, although, in common with the great vessel hematocrit, it cannot be said to reflect quantitatively the average body hematocrit.

The recipient animals were observed for a period of at least 5 hours after the injection of fluid. If throughout that time no significant change had occurred in the condition of the animal, the experiment was terminated. Where any change had occurred, observations were continued until the death of the animal. Because of the possibility that the plasma proteins present in the injected fluid might augment the blood volume of a normal recipient and either mask any possible toxic effects or cause harm in itself by mere plethora, in 10 experiments the fluid was reinfected at the end of the collection period into the animal which had produced it. It thus merely partially replaced plasma volume already lost. The results were in no way different from those in which the fluid was administered to other recipients. Gross and microscopic examination of the viscera were made in the majority of animals. Bacteriologic studies which were made on the fluids are reported in another communication (11).
RESULTS AND DISCUSSION

The results are presented in Tables I and II. They divide themselves sharply into 2 groups. In 1 (Table I), the injection of the fluid resulted in a progressive decline in blood pressure and cardiac output, death of the animal usually within 5 hours, and consistent autopsy findings. These are the criteria of shock used in our experiments. The characteristic fall of blood pressure was interesting for it was invariably a distinctly delayed reaction. Some fluids had a mild vasodepressor action which caused a prompt and short fall of blood pressure which rapidly disappeared. The blood pressure then usually rose nearly to the control level. After a delay of 30 to 45 minutes, the shock-producing effect became manifest in a gradual progressive fall in blood pressure. Figure 1 illustrates the blood pressure response of an animal following injection of muscle exudate. In a few instances the rate of blood flow through one femoral artery was measured by means of a modification of an air bubble flow meter (72). In animals showing the other criteria of shock following the intravenous injection of muscle exudate, a significant reduction in the peripheral blood flow was observed, consistent with the reduction in cardiac output.

In the other group (Table II), none of these effects were observed. Even though the results described in the previous paper of this series indicate the dominance of fluid loss, this is good evidence for an additional inconstant toxic factor in muscle exudate which has the effect of producing shock, in spite of increasing the plasma volume by roughly 25 per cent. The effect was obtained consistently in the pooled fluids (number 9) accumulated from 9 dogs, indicating that the effect does not depend on individual sensitivity of certain animals to the toxin.

Examination of the possible sequence of events in the first group of animals indicates that the rapid and progressive decrease in cardiac output precedes, and is the probable cause of, the drop in blood pressure. The progressive increase in total peripheral resistance may be the result of a compensatory peripheral vasconstriction. The reduction in cardiac output, however, is not specifically explained by the data available. It could result from either an intrinsic myocardial change or a decrease in venous return to the heart. The

### TABLE I

**Positive muscle exudate assays**

<table>
<thead>
<tr>
<th>Donor dog number</th>
<th>Fluid number</th>
<th>Recipient dog number</th>
<th>Weight (kgm.)</th>
<th>Dose (ml. per kgm.)</th>
<th>Blood pressure (mm. Hg)</th>
<th>Cardiac output (liters per kgm. per minute)</th>
<th>Relative total peripheral resistance</th>
<th>Blood hemoglobin (grams per 100 ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>I</td>
<td>II</td>
<td>III</td>
<td>I</td>
</tr>
<tr>
<td>Pool*</td>
<td>9**</td>
<td>122</td>
<td>11.5</td>
<td>15.0</td>
<td>134</td>
<td>110</td>
<td>60</td>
<td>0.13</td>
</tr>
<tr>
<td>Pool</td>
<td>9</td>
<td>123</td>
<td>5.0</td>
<td>15.0</td>
<td>120</td>
<td>75</td>
<td>60</td>
<td>0.10</td>
</tr>
<tr>
<td>Pool (dial.)</td>
<td>9</td>
<td>128</td>
<td>7.5</td>
<td>25.0</td>
<td>130</td>
<td>100</td>
<td>34</td>
<td>0.13</td>
</tr>
<tr>
<td>Pool (dial.)</td>
<td>9</td>
<td>129</td>
<td>6.5</td>
<td>27.0</td>
<td>138</td>
<td>124</td>
<td>45</td>
<td>0.12</td>
</tr>
<tr>
<td>Pool (prot. frac.)</td>
<td>9</td>
<td>132</td>
<td>8.0</td>
<td>45.0</td>
<td>140</td>
<td>72</td>
<td>40</td>
<td>0.15</td>
</tr>
<tr>
<td>Pool</td>
<td>9</td>
<td>136</td>
<td>9.0</td>
<td>24.0</td>
<td>160</td>
<td>138</td>
<td>104</td>
<td>0.13</td>
</tr>
<tr>
<td>148*</td>
<td>17</td>
<td>149</td>
<td>9.8</td>
<td>5.0</td>
<td>145</td>
<td>50</td>
<td>65</td>
<td>0.03</td>
</tr>
<tr>
<td>151*</td>
<td>19</td>
<td>151</td>
<td>9.7</td>
<td>29.0</td>
<td>130</td>
<td>130</td>
<td>32</td>
<td>0.16</td>
</tr>
<tr>
<td>157*</td>
<td>24</td>
<td>173</td>
<td>12.0</td>
<td>15.0</td>
<td>142</td>
<td>135</td>
<td>65</td>
<td>0.16</td>
</tr>
<tr>
<td>180*</td>
<td>28</td>
<td>182</td>
<td>6.0</td>
<td>15.0</td>
<td>110</td>
<td>78</td>
<td>30</td>
<td>0.34</td>
</tr>
<tr>
<td>181</td>
<td>29</td>
<td>183</td>
<td>5.8</td>
<td>15.0</td>
<td>108</td>
<td>55</td>
<td>35</td>
<td>0.12</td>
</tr>
<tr>
<td>Average</td>
<td>154</td>
<td>154</td>
<td>23.5</td>
<td>22.0</td>
<td>120</td>
<td>120</td>
<td>80</td>
<td>1300</td>
</tr>
</tbody>
</table>

* Data incomplete.
** Pooled fluid 9 is discussed in detail in the following paper of this series (23). Following dialysis (dial.), the non-dialyzable portion retained its activity, and the protein fraction (prot. frac.) which salted out from 0.2 to 0.7 saturation with ammonium sulfate was the most highly active. The dosage of these fluids has been calculated from the original fluid involved, and does not take into account losses in toxicity which must have occurred during dialysis and fractionation.
*** Average dose omitting fractionation experiments, 14.7 ml. per kgm.
1 Before injection.
II Immediately after injection.
III Final or 5 hours.
latter change could in turn be due either to a loss in blood volume through capillary leakage or a pooling of the blood in dilated vascular beds. The absence of any significant hemoconcentration, however, is evidence against generalized capillary leakage of plasma as an initiating factor in these experiments. In this respect alone do these animals differ from those described by Moon (63); there is, however, a likely explanation for the hemoconcentration which he observed in a large part of his experiments. His experiments consisted of introducing a weighed amount of muscle pulp plus 100 ml. of saline into the peritoneal cavity of dogs and subsequently measuring the fluid recovered from the peritoneum at autopsy. While the volume of exudate did not increase much in his observations, it is highly likely that the injected saline solution was replaced by a solution high in plasma proteins. The loss of plasma proteins into the peritoneum may well account for the hemoconcentration. In our observations, the reverse situation holds for we were diluting plasma with an intravenous injection of fluid high in proteins.

The post-mortem appearance of the animals in Group I was similar to that in Moon's experi-

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**TABLE II**

**Negative muscle exudate assays**

<table>
<thead>
<tr>
<th>Donor dog</th>
<th>Fluid number</th>
<th>Recipient dog number</th>
<th>Weight</th>
<th>Dose</th>
<th>Blood pressure</th>
<th>Cardiac output</th>
<th>Relative total peripheral resistance</th>
<th>Blood hemoglobin</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>km/h</td>
<td>liters/km/min</td>
<td>grams/hour</td>
<td>grams per 100 ml</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ml per km</td>
<td>per minute</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>mm Hg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>II</td>
<td>III</td>
<td></td>
<td></td>
<td>I</td>
<td>II</td>
<td>III</td>
<td></td>
</tr>
<tr>
<td>135</td>
<td>10</td>
<td>135</td>
<td>20.0</td>
<td>14.8</td>
<td>94</td>
<td>118</td>
<td>92</td>
<td>0.07</td>
</tr>
<tr>
<td>137</td>
<td>11</td>
<td>137</td>
<td>17.5</td>
<td>11.4</td>
<td>108</td>
<td>124</td>
<td>130</td>
<td>0.08</td>
</tr>
<tr>
<td>138</td>
<td>12</td>
<td>138</td>
<td>20.0</td>
<td>15.0</td>
<td>127</td>
<td>123</td>
<td>132</td>
<td>0.10</td>
</tr>
<tr>
<td>142</td>
<td>13</td>
<td>142</td>
<td>20.5</td>
<td>7.0</td>
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<td>87</td>
<td>0.12</td>
</tr>
<tr>
<td>143</td>
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<td>143</td>
<td>16.0</td>
<td>5.0</td>
<td>102</td>
<td>90</td>
<td>82</td>
<td>0.14</td>
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<tr>
<td>145</td>
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<td>0.16</td>
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<td>16</td>
<td>147</td>
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<td>6.7</td>
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<td>0.18</td>
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<td>151</td>
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<td>15.0</td>
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<tr>
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<tr>
<td>166</td>
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<td>167</td>
<td>6.0</td>
<td>15.0</td>
<td>160</td>
<td>95</td>
<td>140</td>
<td>0.24</td>
</tr>
<tr>
<td>168</td>
<td>23</td>
<td>169</td>
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<td>15.0</td>
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<td>130</td>
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<td>174</td>
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<td>3.0</td>
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<td>60</td>
<td>125</td>
<td>0.28</td>
</tr>
<tr>
<td>171</td>
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<td>175</td>
<td>6.8</td>
<td>13.0</td>
<td>160</td>
<td>132</td>
<td>142</td>
<td>0.30</td>
</tr>
<tr>
<td>176</td>
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<td>178</td>
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<td>9.0</td>
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<td>125</td>
<td>118</td>
<td>0.32</td>
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<tr>
<td>177</td>
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<td>179</td>
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<td>15.0</td>
<td>95</td>
<td>93</td>
<td>130</td>
<td>0.34</td>
</tr>
<tr>
<td>214-215</td>
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<td>216</td>
<td>6.8</td>
<td>13.5</td>
<td>144</td>
<td>90</td>
<td>125</td>
<td>0.36</td>
</tr>
<tr>
<td>217-218</td>
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<tr>
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<td>33</td>
<td>223</td>
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<td>120</td>
<td>135</td>
<td>0.40</td>
</tr>
<tr>
<td>226-227*</td>
<td>34</td>
<td>228</td>
<td>5.8</td>
<td>14.0</td>
<td>130</td>
<td>130</td>
<td>120</td>
<td>0.42</td>
</tr>
</tbody>
</table>

Average 12.9 12.0 128 116 120 0.16 0.14 0.13 1320 1180 1315 14.1 13.3 13.8

* Data incomplete. I Before injection. II Immediately after injection. III Final or 5 hours.

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**Fig. 1. Kymograph Record Illustrating the Blood Pressure of an Animal Following Injection of Muscle Exudates**

The white blocks indicate cardiac outputs taken at various times during the course of the experiment. The first cardiac output taken after administration of the toxin (Number 2) is (in this case) increased, although such was not usually the case. The blood pressure response is a typical one. Cardiac output values are as follows, expressed in liters per minute: Number 1—1.47, 2—2.22, 3—0.69, 4—0.60, 5—0.35.
ments. There was intense visceral congestion, especially of the liver, together with moderate to marked hemorrhage into the intestinal lumen. This began abruptly at the pylorus, was most marked in the duodenum, and diminished distally.

A consideration of the proportion of positive to negative results in these experiments is illuminating. Of the total of 32 fluids studied, only 9 (or 28 per cent) were found to possess significant toxic properties, while the remaining 72 per cent were apparently innocuous.²

If the 9 toxic fluids derived their properties from noxious products of tissue breakdown, it is difficult to see how the other 23 specimens could have failed to contain these substances. All the muscles were subjected to the same degree and duration of ischemia and all the fluids were high in protein content indicating the degree of local capillary damage. Moreover both toxic and nontoxic fluids revealed similar electrolyte composition and the same concentration of proteolytic enzymes (70) indicating similar degrees of cellular breakdown. These considerations and the very inconstancy of positive results from fairly rapid intravenous injection suggest that the positive effects are due to inconstant factors which are not present in all muscles, but are probably an extraneous complication introduced into some experiments.

The possibility presents itself that an explanation for the positive effects in one-fourth of these fluids lies in bacterial contamination. The results of a detailed study of the number and types of organisms present in these fluids appears elsewhere (11), and the evidence there presented clearly demonstrates a correlation between the toxicity of a fluid and its bacterial content.

Since the majority of these fluids yielded no evidence of a shock producing factor, it appears warranted to conclude that the usual metabolic breakdown products in anoxic muscles do not produce the vascular changes which precede shock. In our muscle ligation technique, the circulation in the anoxic muscles is usually well reestablished, so that diffusible toxic elements should be present in the circulation as a whole, as well as in the muscle exudate. However, this factor is controlled by the similar results which were obtained when fluids were reinjected into the original donor, as when given to recipient dogs.

**SUMMARY**

1. The exudate which accumulates after muscle anoxia was collected and injected into the same or recipient dogs.

2. In three-quarters of such observations, no evidence of shock was observed, while in one-quarter of the animals shock was produced by this technique and caused death.

3. The inconstancy of the presence of this toxic factor suggests it is an extraneous agent, not present in the usual cellular constituents and metabolic products found in all muscle exudates. Since the exudates are invariably contaminated with bacteria, it is thought that the toxin may be bacterial in origin.

Bibliography follows Paper VI of this series.

² Table II does not include the earlier experiments in which the cardiac output was not measured.