
Complement-fixation Studies of the Specificity of the Interactions between Components of the Virus System of Foot-and-Mouth Disease and its Antibodies

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SUMMARY: The type and subtype specificities of the interactions between fractions of the virus system of foot-and-mouth disease and its antibodies were investigated by complement-fixation methods. Specificity is discussed in terms of a cross-fixation ratio. The 25 mp infective component (D-fraction) combines homotypically with antibody. Thermal degradation of the 25 mp component produces a smaller component of enhanced activity which combines heterotypically with antibody and resembles the naturally occurring 7 mp component (U-fraction). The influence of heating upon the 7 mp component is marked by a sharpened specificity. The specificity and activity of unfractionated starting materials may be interpreted in terms of those of the 25 and 7 mp components present in the separable fractions. Significant differences between the reactions observed in the long and short incubation procedures emphasize the advantages of the latter in specificity studies. The composition and treatment of the initial antigens and the test procedures employed must be carefully defined in such studies of antigen-antibody combination.

As in other systems of multiple antigens and antibodies, viral antigen-antibody interactions reveal affinities which are discussed under the term ‘specificity’. As the component antigens and antibodies become more and more exactly defined it is necessary to refine the definition of specificity to distinguish between the different specificities of the individual components and the ill-defined hybrid specificity of the original material in which these components occur in variable if not unknown proportions. In the particular case of the infective agent in the virus system of foot-and-mouth disease higher and higher specificities are involved in the progression from the recognition of a virus as that of foot-and-mouth disease, to the recognition of the virus as one of the seven distinct types within the disease group, and then to the recognition of that virus as a distinct subtype within the type. The further specificity defined by the adaptation of the virus to a particular animal host is outside the scope of the present study. Superimposed upon this progression of specificities for the infective agent there are the related yet distinct specificities of other antigens within the virus system.

This paper reports a study by complement fixation of the type specificities of the components of the virus system of foot-and-mouth disease. Earlier studies on unfractionated materials and parts of the present work were reviewed by Brooksby (1958, 1958). Recent gel-diffusion studies by Brown & Crick (1958) have confirmed some of the present findings.
METHODS

The antigens and antibodies used in these experiments and the methods of titration of infectivity and complement-fixing activity were as described in previous papers (Brooksby, 1952; Bradish, Brooksby & Tsubahara, 1960).

The starting materials for fractionation were pooled suspensions of the vesicular epithelium and fluid from primary lesions on the plantar pads of groups of 20 or more infected guinea pigs. Original suspensions of 1 g. tissue in 10 ml. phosphate buffer (w/25; pH 7.6) provided clarified starting materials containing about 5 mg. protein/ml. The separation by centrifugation of the components of the virus system proceeded as previously described (Bradish, Brooksby, Dillon & Norambuena, 1952; Bradish, Henderson & Kirkham, 1960). The distribution of infectivity, complement-fixing activity and nitrogen through the various fractions was described in the latter paper. The 25 mp infective component occurs in the D-fraction and the 7 mp complement-fixing component in the U-fraction.

Definition of specificity and cross-fixation ratio

Specificity studies, in general, involve the comparison of two reactions in which one reactant, either antigen or antibody, is common to both. In terms of complement fixation the combination reactions between antigen G₁ and antibody B and between antigen G₂ and the same antibody B are arranged in order of specificity according to the amount of complement fixed in appropriately standardized tests in which the antigen concentrations (G₁) and (G₂) are equal. The more specific reaction fixes the greater amount of complement.

The ratio of the two complement-fixing activities observed when equal samples of an antigen react in standardized tests with antisera A and B may be expressed as

\[
\text{Cross-fixation ratio} = \frac{\text{complement-fixing activity in presence of excess antibody B}}{\text{complement-fixing activity in presence of excess antibody A}}
\]

This ratio is independent of the unknown antigen concentration and provides a measure of the specificity of the combination between antigen and antibody B in terms of that between the same antigen and antibody A (see previous paper; Bradish, Brooksby & Tsubahara, 1960). When the antibody A is defined as that homologous to the antigen then the cross-fixation ratio ranges from zero for the reaction between completely heterologous antigen and antibody, to unity for the reaction between homologous reactants. This convenient scale of cross-fixation is the basis of the interpretation of the present data.

The comparison between two antigens in complement-fixation tests using the corresponding two homologous antisera requires the further definition: two antigens are distinct when the product of their cross-fixation ratios,
Specificities of components of FMD virus system

tested in excess of the homologous antisera, is significantly different from unity. A similar definition applies to the data of antibody titrations in antigen excess, when the terminology is adjusted accordingly. The application of this criterion to type and subtype differentiation is shown below.

RESULTS

Cross-fixation ratios of U- and D-fractions

Complement-fixation data for the titration in antibody excess of fractions of the virus system are shown in Table 1. Experiments in which the antigen-antibody mixture was incubated with complement for 30 min. at 37° before the introduction of the haemolytic indicator (short test) showed the typical degrees of complement-fixing activity and specificity associated with the U (7 μ component) and D (25 μ component) fractions. The fixation of complement by the clarified starting material in the presence of type-homologous serum was sufficiently in excess of that observed in excess of type-heterologous serum to ensure a correct designation of type in most cases. This

Table 1. Specificity of fractions of the virus system of foot-and-mouth disease separated from guinea-pig pad epithelium and vesicular fluid

Activities are shown in the body of the table as the volumes (μl.) of complement 'fixed' in antibody excess by 1 ml. the undiluted virus fractions. The concentrations of all fractions are the same as those of the clarified starting materials.

<table>
<thead>
<tr>
<th>Type and strain</th>
<th>Starting material</th>
<th>U-fraction</th>
<th>D-fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>fraction of virus titrated</td>
<td>of excess antiserum</td>
<td>Long test</td>
<td>Short test</td>
</tr>
<tr>
<td>O/VI</td>
<td>A/GB</td>
<td>980</td>
<td>440</td>
</tr>
<tr>
<td></td>
<td>C/GC</td>
<td>4010</td>
<td>630</td>
</tr>
<tr>
<td></td>
<td>O/VI</td>
<td>1550</td>
<td>0.63</td>
</tr>
<tr>
<td>Cross-fixation ratio</td>
<td>To A</td>
<td>0.70</td>
<td>0.15</td>
</tr>
<tr>
<td></td>
<td>To C</td>
<td>0.65</td>
<td>0.73</td>
</tr>
<tr>
<td>O/VI</td>
<td>A/85</td>
<td>—</td>
<td>234</td>
</tr>
<tr>
<td></td>
<td>O/87</td>
<td>621</td>
<td>253</td>
</tr>
<tr>
<td>Cross-fixation ratio</td>
<td>To A</td>
<td>—</td>
<td>0.92</td>
</tr>
<tr>
<td>A/GB</td>
<td>A/78</td>
<td>—</td>
<td>206</td>
</tr>
<tr>
<td></td>
<td>C/73</td>
<td>—</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>O/74</td>
<td>—</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>Normal</td>
<td>—</td>
<td>5</td>
</tr>
<tr>
<td>Cross-fixation ratio</td>
<td>To C</td>
<td>0.09</td>
<td>0.25</td>
</tr>
<tr>
<td></td>
<td>To O</td>
<td>—</td>
<td>0.10</td>
</tr>
<tr>
<td>A/GB</td>
<td>A/84</td>
<td>108</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>O/87</td>
<td>—</td>
<td>19</td>
</tr>
<tr>
<td>Cross-fixation ratio</td>
<td>To O</td>
<td>—</td>
<td>0.43</td>
</tr>
<tr>
<td>C/GC</td>
<td>A/GB</td>
<td>310</td>
<td>120</td>
</tr>
<tr>
<td></td>
<td>C/GC</td>
<td>780</td>
<td>240</td>
</tr>
<tr>
<td></td>
<td>O/VI</td>
<td>429</td>
<td>230</td>
</tr>
<tr>
<td>Cross-fixation ratio</td>
<td>To A</td>
<td>0.40</td>
<td>0.50</td>
</tr>
<tr>
<td></td>
<td>To O</td>
<td>0.54</td>
<td>0.96</td>
</tr>
</tbody>
</table>
is equivalent to the standard typing procedure (Brooksby, 1952). Separated U-fractions showed significantly greater cross-fixation than did the clarified starting materials.

The D-fractions, in contrast to the U-fractions, showed cross-fixation ratios which were sufficiently low to permit the conclusion that in the short incubation test the 25 mμ component combined only with antibody of its own type and thereby defined the virus type of the starting material. Uncertainties in typing tests on unfractionated materials may be resolved by typing the separated homotypic D-fractions.

The fixation of complement by antigen-fractions in the presence of normal, inactivated guinea-pig serum was usually less than 2% of that in homologous serum.

Table 1 also shows results by the long-incubation test. Values of both complement-fixation and cross-fixation ratio were significantly greater in the long- than in the short-incubation test. This loss of specificity by the U-fraction in the long test was coupled with the disproportionate enhancement of activity demonstrated in the previous paper. It is apparent that the short-incubation test offers considerable advantages in sensitivity to virus type, particularly in typing studies of unfractionated starting materials.

In both long- and short-incubation tests the cross-fixation ratios for the D-fractions was significantly less, and for the U-fractions significantly more, than those for the starting materials. These data generally are consistent with the recognition of the starting material as a mixture of a homotypic 25 mμ component and a heterotypic 7 mμ component which respond differently to incubation procedures in the complement-fixation test.

Specificity in relation to sub-types of virus and corresponding antisera

In the previous section the cross-fixation ratio was shown to be determined by the virus fraction and incubation method used and to differentiate clearly between reactants of distinct virus type. Lower order variations in the properties of antigens and antibodies within a single virus type have been reported (Traub & Möllmann, 1946; Galloway, Henderson & Brooksby, 1948). Such strains are called variants or subtypes.

Table 2 shows the complement-fixation data relating to two strains of virus of A type and three strains of C type. Titrations of antibody in antigen excess and of antigen in antibody excess are shown for the U- and D-fractions of the virus systems. The cross-fixation ratio, now the ratio of the complement-fixing activity in excess of strain-heterologous reactant to that in excess of strain-homologous reactant, is shown in Table 2 in brackets beside the corresponding complement-fixing activities. In all experiments, and as reported in the previous paper (Bradish, Brooksby & Tsubahara, 1960), the titration of antibody in antigen excess allowed a more sensitive detection of differences than the alternative titration of antigen in antibody excess.

In the titrations of either antibody or U-fraction antigen of type A, strain Ca 1, the fixation with antigen or antibody of type A, strain GB, was greater than that with the strain-homologous reactants. A similar situation appears
in the data for type C, strain 149, in Table 2 and for type A, strain HK. The remaining antigen-antibody combinations show greatest activities with strain-homologous reactants. These observations of U-fraction activities in opposition to the subtype differentiation of the parent material emphasize the generally heterologous character of the 7 M component of the virus system and the caution which must be exercised in the subtyping of unfractonated samples.

Table 2. Influence of variations within type on antigen-antibody combination reactions

<table>
<thead>
<tr>
<th>Antigen: type/strain/fraction</th>
<th>Titration of antibody with excess antigen</th>
<th>Type and strain of serum used as antibody</th>
<th>Titration of antigen with excess antibody</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A/GB</td>
<td>A/Ca1</td>
<td>A/GB</td>
</tr>
<tr>
<td>A/Ca1/U</td>
<td>48 (0.15)</td>
<td>16 (1)</td>
<td>0-140 (1-6)</td>
</tr>
<tr>
<td>A/Ca1/D</td>
<td>41 (1)</td>
<td>0-049 (0-95)</td>
<td>0-052 (1)</td>
</tr>
<tr>
<td>A/GB/U</td>
<td>330 (1)</td>
<td>31 (2)</td>
<td>0-260 (1)</td>
</tr>
<tr>
<td>A/GB/D</td>
<td>390 (1)</td>
<td>0-072 (1)</td>
<td>0-038 (0-53)</td>
</tr>
<tr>
<td>C/149</td>
<td>38 (1)</td>
<td>44 (0-5)</td>
<td>75 (1-2)</td>
</tr>
<tr>
<td>C/149/D</td>
<td>38 (1)</td>
<td>0-69 (1)</td>
<td>0-67 (0-97)</td>
</tr>
<tr>
<td>C/733/U</td>
<td>70 (1-8)</td>
<td>88 (1)</td>
<td>75 (1-2)</td>
</tr>
<tr>
<td>C/733/D</td>
<td>130 (1)</td>
<td>0-40 (0-85)</td>
<td>0-47 (1)</td>
</tr>
<tr>
<td>C/U3/U</td>
<td>44 (1-2)</td>
<td>40 (0-46)</td>
<td>63 (1)</td>
</tr>
<tr>
<td>C/U3/D</td>
<td>100 (1)</td>
<td>0-70 (0-69)</td>
<td>0-85 (0-83)</td>
</tr>
</tbody>
</table>

In terms of the product of cross-fixation ratios (Table 3) the virus strains GB and Ca1 of type A are distinct subtypes. This is shown also by the more sensitive titration of the antisera in antigen excess. The type C strain U3 is probably distinct from strains 733 and 149, whereas the latter show no significant differentiation. In earlier studies of unfractonated materials (Skinner, Henderson & Brooksby, 1952) strains 733 and 149 were identified as subtypes; this discrepancy has not been investigated further. Table 3 also shows the cross-fixation ratios appropriate to type differentiation studies and demonstrates the relatively small difference between subtypes as compared with that between types.

Change of antigen activity produced by heat treatment

Since the extent and specificity of antigen-antibody combination, as indicated by the complemen-fixation test, is dependent upon the time-temperature conditions of incubation a number of experiments were made to examine...
the stability of the antigens during heating. Clarified starting materials and the derived U- and D-fractions were heated in water baths at temperatures between 85° and 56° for periods up to 24 hr. Control samples were held at 4°. After treatment samples were cooled to 4° until titrated by complement fixation in antibody excess. Typical results are shown in Fig. 1. When incubated at 85°-40° the complement-fixing activities of the D-fractions increased steadily until at 16–24 hr. the activities were about twice those of the control samples. Incubation for up to 30 min. at 56° or for 1–4 hr. at 45° was sufficient to produce the same approximate doubling of the initial activity. Continued heating at 45° for up to 24 hr. produced no significant loss of activity. In eleven such experiments at temperatures between 35° and 56° the complement-fixing activities of D-fractions increased on the average to 190% (150–230%) of the initial values. Parallel incubation of the corresponding U-fractions

Table 3. Differentiation of foot-and-mouth disease virus strains as types and subtypes

<table>
<thead>
<tr>
<th>Virus type of strains used</th>
<th>Strains and fractions under comparison</th>
<th>Cross-fixation ratio in long and short tests</th>
<th>Product of cross-fixation ratios</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Ca1-fraction</td>
<td>1.6, 1.2</td>
<td>0.8, 0.55</td>
</tr>
<tr>
<td></td>
<td>GB-U fraction</td>
<td>0.50, 0.46</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ca1-U fraction</td>
<td>0.95, 0.97</td>
<td>0.50, 0.58</td>
</tr>
<tr>
<td></td>
<td>GB-D fraction</td>
<td>0.53, 0.60</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ca1-antiserum</td>
<td>2.0</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>GB-antiserum</td>
<td>0.15</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>149-U fraction</td>
<td>1.2</td>
<td>1.08</td>
</tr>
<tr>
<td></td>
<td>733-U fraction</td>
<td>0.90</td>
<td></td>
</tr>
<tr>
<td></td>
<td>149-U fraction</td>
<td>1.5</td>
<td>1.05</td>
</tr>
<tr>
<td></td>
<td>U3-U fraction</td>
<td>0.70</td>
<td></td>
</tr>
<tr>
<td></td>
<td>733-U fraction</td>
<td>0.86</td>
<td>0.83</td>
</tr>
<tr>
<td></td>
<td>U3-U fraction</td>
<td>0.97</td>
<td></td>
</tr>
<tr>
<td></td>
<td>149-U fraction</td>
<td>0.97</td>
<td>0.82</td>
</tr>
<tr>
<td></td>
<td>733-D fraction</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td></td>
<td>149-D fraction</td>
<td>0.85</td>
<td>0.59</td>
</tr>
<tr>
<td></td>
<td>U3-D fraction</td>
<td>0.69</td>
<td></td>
</tr>
<tr>
<td></td>
<td>733-D-fraction</td>
<td>0.87</td>
<td>0.72</td>
</tr>
<tr>
<td></td>
<td>U3-D fraction</td>
<td>0.83</td>
<td></td>
</tr>
<tr>
<td></td>
<td>149-antiserum</td>
<td>1.8</td>
<td>0.9</td>
</tr>
<tr>
<td></td>
<td>733-antiserum</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>149-antiserum</td>
<td>1.2</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td>U3-antiserum</td>
<td>1.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>733-antiserum</td>
<td>0.46</td>
<td>0.55</td>
</tr>
<tr>
<td></td>
<td>U3-antiserum</td>
<td>1.2</td>
<td></td>
</tr>
</tbody>
</table>

Comparative type-differentiation

| A and O                    | A/GB-U fraction                        | 0.43, 0.22                                  | 0.40, 0.06                       |
|                           | A/OVI-U fraction                       | 0.92, 0.27                                  |                                |
|                           | A/GB-D fraction                        | 0.04, 0.03                                  | 0.006, 0.001                     |
|                           | A/OVI-D fraction                       | 0.16, 0.03                                  |                                |
Specificities of components of FMD virus system

(Fig. 1) showed a distinct behaviour since now the complement-fixing activities decreased in all cases. The activities fell to about 75% of the initial values after 16 hr. at 35°-40° or 1 hr. at 50°-56°.

The results obtained in these experiments by the short and long incubation tests were generally similar, although differences in detail were apparent under certain conditions. Figure 2 shows the comparison of the results by the two methods, expressed as the ratio of the complement-fixing activity in the long incubation test to that in the short incubation test. This ratio of the test results is given for the U- and D-fractions of the virus system after different thermal treatments. For the U-fraction the test ratio was initially 3.5 and fell steadily on heating to 2.3 after 16 hr. at 50°. For the D-fractions the test ratio was initially 1.8 and fell steadily with increasing time and temperatures above 40° to values as low as 0.6. Treatment at 35° for 1-16 hr., however, caused the test ratio to increase to about 2.2. This distinction of
response between treatment at 35° and 40° or above is particularly interesting since the disintegration of the 25 mμ component (D-fraction) at the lower temperature produced a component with test ratio akin to that of the 7 mμ component (U-fraction). Degradation of either component at 40° or above decreased the distinction between the tests. This suggests that part of the distinction between the tests is related to the degradation of the antigens during continued incubation with antibody.

Change of specificity of antibody-combining activity following heat treatment of antigen fractions

The data shown in Fig. 8 demonstrate the changes in specificity observed following heat treatment of the antigens for 4 hr. at 45° or 56°. Heated and control fractions were titrated in antibody excess by complement fixation in the long-incubation test. The cross-fixation ratios show that the D-fractions were initially almost completely type-specific in their capacity to combine with antibody but became strongly heterotypic after heat treatment had doubled their homotypic capacity. In contrast to the D-fractions, heat treatment of the U-fractions improved the type-specificity of their combination with antibody. In some cases the heated U-fractions showed cross-fixation ratios approaching those of the highly specific initial D-fractions. The starting materials, regarded as natural mixtures of the D- and U-fractions, showed properties intermediate between those of these components. Heat treatment of starting materials produced an increase of up to c. 50 % in the value of complement-fixing activity in the presence of excess type-homologous antibody. This was accompanied by a slight increase in the degree of cross-fixation with antibody of heterologous type.
Change of sedimentation properties following heat treatment or storage of antigen fractions

Sedimentation analyses of control or heated U-fractions by angle or radial tube sampling methods showed, in confirmation of earlier data (Bradish et al. 1952; Bradish, Brooksby & Dillon, 1956), a complement-fixing component of sedimentation coefficient $8 \pm 1$ Svedberg units. This corresponds with an equivalent protein particle diameter of about 7 mμ, which was little changed by heat treatment under the conditions defined. Parallel experiments with the initial D-fractions indicated a particle size of about 25 mμ by electron microscopy (Bradish, Henderson & Kirkham, 1960) and a consistent minimum sedimentation coefficient of about 70 Svedberg units by angle or radial tube sampling methods, with assay by infectivity.

Table 4. Influence of heat treatment upon sedimentation properties of complement-fixing components of foot-and-mouth disease virus fractions

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Initial activity (%) relative to clarified starting material (c.s.m.)</th>
<th>Activity after heating at $56^\circ$ (% initial c.s.m.)</th>
<th>Percentage of activity of heated sample remaining in supernatant fluid after centrifugation at 12G to remove 25 mμ component</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clarified starting material</td>
<td>100</td>
<td>115</td>
<td>95</td>
</tr>
<tr>
<td>U-fraction (7 mμ)</td>
<td>41</td>
<td>25</td>
<td>80</td>
</tr>
<tr>
<td>D-fraction (25 mμ)</td>
<td>50</td>
<td>76</td>
<td>90</td>
</tr>
<tr>
<td>U-fraction (7 mμ)</td>
<td>50</td>
<td>80</td>
<td>75, 85</td>
</tr>
<tr>
<td>D-fraction (25 mμ)</td>
<td>50</td>
<td>76</td>
<td>90, 78</td>
</tr>
</tbody>
</table>

Investigation of the sedimentation properties of thermally disintegrated D-fractions regularly showed a sedimentation coefficient of only 7 to 11 Svedberg units. Electron micrographs of such degraded D-fractions showed in agreement with the sedimentation data, the absence of the 25 mμ particle and the presence of aggregates and strings of 5–10 mμ particles. Table 4 shows that after heat treatment the activity of the initially infective D-fractions sedimented as a 7 mμ component, similar to that in the U-fractions.

A complication in the interpretation of these experiments arose in some cases when analyses in terms of complement-fixing activity of unheated D-fractions gave sedimentation coefficients of about 8 Svedberg units. This indicated the lability of the 25 mμ infective component, which may disintegrate more readily when partially purified than when present as a component of the starting material. It appears that the 25 mμ component disintegrates continuously to produce an accumulating concentration of a smaller component of distinct properties. This process is accelerated by heating and also, to a lesser extent, by the manipulation inherent in experimental procedures. For this reason periods of manipulation at room temperature were restricted to a
total of less than 2 hr. Final assays of the complement-fixing activity or infectivity of fractions were complete within 4–6 days of the collection of the initial infective materials.

That disintegration occurred even under the nominally favourable conditions of storage at $-20^\circ$ is indicated by the following experiment. Vesicular epithelium stripped from the tongue of a steer infected with virus of type O, strain 39, was divided into 1 cm. squares and distributed in four 5 ml. bottles. An immediate partition analysis (Bradish et al. 1952) of the suspension prepared from the contents of one bottle indicated that 38% of total complement-fixing activity was due to the 7 m$\mu$ component. The remaining bottles were stored at $-20^\circ$ and suspensions were prepared for partition analysis after 1, 4 and 14 weeks. After these times the complement-fixing activity due to the 7 m$\mu$ component contributed, respectively, 49, 64 and almost 100% of the total activity. Thus even at $-20^\circ$ the proportion of total activity due to the extracted 25 m$\mu$ component diminished from 67% initially to 51, 36 and 0% at 1, 4 and 14 weeks, respectively.

DISCUSSION

It had been shown earlier that the virus system of foot-and-mouth disease comprises at least two distinct components. The present study indicates, in terms of complement-fixation data, the serological specificity and inter-relationship of these components. The 25 m$\mu$ infective component which occurs in the separated D-fraction is a labile antigen which combines only with antibody of its own type. This component is readily degraded by heating to produce an essentially non-infective 7–11 m$\mu$ component which combines with antibody of any type within the disease group. Such degradation is accompanied by an approximate doubling of complement-fixing activity in the presence of homologous antibody. The disintegration of the 25 m$\mu$ infective component to produce a component which is similar in many respects to the naturally occurring 7 m$\mu$ component suggests that this latter component may arise either as a precursor or as a product of the larger particle, or perhaps in both ways. In most starting materials collected from guinea pig or ox about half of the initial complement-fixing activity is due to the 7 m$\mu$ component. If this arises through the disintegration of intact infective particles it might be anticipated that under certain conditions starting materials would be collected in which all complement-fixing activity was contributed by intact 25 m$\mu$ particles. That such preparations are encountered only as separated D-fractions and not as starting materials implies either that disintegration proceeds as virus accumulates at the particular site in the host tissue, or that some 7 m$\mu$ component is formed independently. The infectivity and complement-fixing activity of the material collected from the host may thus be dependent not only upon the course of virus multiplication but also upon superimposed decay processes.

In addition to the infective 25 m$\mu$ component of complete type-specificity and the non-infective (probably only weakly antigenic) 7 m$\mu$ particles pos-
Specifities of components of FMD virus system

Sessing only disease-group specificity, it is necessary to consider the presence of a non-infective yet fully antigenic component. The presence of such a 25 μm component is indicated by the 100- to 1000-fold disparity between particle counts by the electron microscope and titrations of numbers of infective units. The potency of vaccines also presumes a non-infective yet fully antigenic component. It is probable that non-infective 25 μm particles are responsible for the major part of the antigenicity and homotypic antibody-combining activity of virus samples. Such intermediate, non-infective components, which retain antigenicity or type-specificity, are of importance to the understanding of the relationships which exist between infectivity, antigenicity, antibody-combining capacity and physical-chemical structure.

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