The EB virus in relation to infectious mononucleosis

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Chance favours the prepared mind and it was a fortunate circumstance that the Henles were studying the Epstein-Barr (EB) virus (Epstein, 1970) when one of their technicians, whose serum was being used as a negative control, fell ill with typical infectious mononucleosis. During this illness, she developed not only heterophile antibodies but also fluorescent antibodies to the EB virus (Henle, Henle, and Diehl, 1968). In another example of forethought, sera had been taken at intervals over the years from university students. Examination of these confirmed the association between the development of infectious mononucleosis and the acquisition of antibodies to the EB virus (Evans, Niederman, and McCollum, 1968). Subsequent studies (Niederman, Evans, Subrahmanyan, and McCollum, 1970; Sawyer, Evans, Niederman, and McCollum, 1971; Joint Investigation by University Health Physicians and Public Health Laboratory Service Laboratories, 1971) have shown convincingly that fluorescent antibodies to the EB virus develop during illnesses which are clinically recognizable as infectious mononucleosis and which are associated with the development of a positive Paul-Bunnell test and the appearance of typical abnormal lymphocytes in the peripheral blood. Infectious mononucleosis which is unassociated with the development of a positive Paul-Bunnell test may be associated with cytomegalovirus infection (Klemola and Kääriäinen, 1965; Lamb and Stern, 1966) but is uncommonly associated with EB virus infection (Klemola, von Essen, Henle, and Henle, 1970).

The transmission of EB virus in man has been observed on several occasions. Grace, Blakeslee, and Jones (1969) inoculated serologically negative volunteers with cell extracts which contained EB virus: infectious mononucleosis developed in these individuals and they acquired EB virus antibodies. Accidental infection following perfusion with presumably infected blood has been described (Gerber, Walsh, Rosenblum, and Purcell, 1969) but Stevens and Pry (1971) considered this to be a rare event. Kapsenberg, Langenhuyzen, Nieweg, and Deiss (1970) observed clinical infectious mononucleosis in a girl of 17 which followed a blood transfusion; she developed high levels of antibody to cytomegalovirus and also to EB virus. Spread of EB virus infection, with or without overt illness, has been detected in family and social contacts of children and young adults with infectious mononucleosis (Wahren, Lantorp, Sterner, and Espmark, 1970; Joncas and Mitnyan, 1970); the latter authors remarked upon the apparent low infectivity of this virus. To date, there have been no reports of the production of infectious mononucleosis in animals following the administration of EB virus, although Landon and Malan (1971) observed the decline of (probably) maternal antibodies in three species of macaque monkeys kept in captivity and tested over a period of months. Similarly, Gerber and Rosenblum (1968) have found EB virus antibodies in rhesus monkeys.

Although infectious mononucleosis is commonly associated with EB virus infection, there is much evidence that many infections must be asymptomatic or nearly so. Thus, Lehane (1970) in a prospective study of a military population, demonstrated a ratio of 1:10 in apparent to inapparent cases of infectious mononucleosis associated with EB virus serological conversion. In our own medical students and nurses, tested on entry for antibodies to rubella and to EB viruses, we found a close relationship between a history of past illness and the presence of antibodies to these viruses (Table I). More had rubella antibodies (80%) than EB virus antibodies

<table>
<thead>
<tr>
<th>Rubella (Rubella virus)</th>
<th>Infectious Mononucleosis (EB virus)</th>
<th>Significance of Differences (χ² test)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. tested</td>
<td>273</td>
<td>224</td>
</tr>
<tr>
<td>No. with past history of illness</td>
<td>163/273 (60%)</td>
<td>24/224 (11%)</td>
</tr>
<tr>
<td>No. with antibodies to relevant virus</td>
<td>219/273 (80%)</td>
<td>144/224 (64%)</td>
</tr>
<tr>
<td>No. with past history of illness + antibodies to relevant virus</td>
<td>147/163 (90%)</td>
<td>20/24 (83%)</td>
</tr>
</tbody>
</table>

Table I Rubella and infectious mononucleosis in medical students and nurses
The evidence thus indicates that infectious mononucleosis is not found until early adult life (Newell, 1957).

Most of the above epidemiological studies on the relationship between infectious mononucleosis and EB virus infection have depended upon the demonstration of antibodies by indirect fluorescence (Henle and Henle, 1966). Knowledge of EB virus replication is now rather fuller and has disclosed a range of antigens which arise during this process. These antigens and the antibodies which may be produced against them may be detected in a variety of ways.

**Antibodies to the EB Virus**

**FLUORESCENT ANTIBODIES (VIRAL CAPSID ANTIGEN ANTIBODIES)**

The indirect fluorescent antibody test, using cells fixed in acetone (Henle and Henle, 1966), was the first method for detecting EB virus antibodies to be described and is usually the one which is implied when no other indication is given. These antibodies are directed towards the virus capsid (Zur Hausen, Henle, Hummeler, Diehl, and Henle, 1967; Epstein and Achong, 1968; Henle, Henle, Zajac, Pearson, Waubke, and Scriba, 1970) and are now termed viral capsid antigen (VCA) antibodies. This antigen develops late during virus replication and, as it is associated with the virus particle, is not detectable in cell lines which are substantially ‘virus-free’ (as determined by electron microscopy), such as the Raji cell line or probably the RM cell line which we derived from a patient with infectious mononucleosis. During infectious mononucleosis, VCA antibodies develop rapidly and may be present shortly before the onset of frank illness. As they are present in over half of the age group in which infectious mononucleosis is commonly observed, their presence or absence is of little use in the diagnosis of this disease unless, of course, sera taken before the onset of illness are available. Viral capsid antigen antibodies decline gradually (Niederman et al, 1970) but may show considerable fluctuations (Sohier, 1970).

**ANTIBODIES TO CELL MEMBRANE ANTIGENS**

Cell membrane antigens were initially detected by indirect immunofluorescence in living Burkitt tumour cells (Klein, Clifford, Klein, and Stjernsward, 1966) but difficulties in standardization led to the use of established cell culture lines and a blocking technique with a standard serum that was shown to be free from iso-antibodies (Klein, Clifford, Klein, Smith, Minowada, Kourilsky, and Burchenal, 1967). The membrane antigen is associated with the virus capsid but differs from the VCA antigen and is probably an early product of virus synthesis, as its formation is not inhibited by cytosine arabinoside or IUDR, whereas puromycin inhibits it completely (Gergely, Klein, and Ernberg, 1971). It can be detected in cultured cells derived from patients with infectious mononucleosis (Klein, Pearson, Henle, Henle, Diehl, and Niederman, 1968) and these patients develop antibodies to cell membrane antigens.

**ANTIBODIES TO EARLY ANTIGEN**

During the course of experimental infections of foetal lymphoblastoid cells with EB virus (Henle, Diehl, Kohn, Zur Hausen, and Henle, 1967; Pope, Horne, and Scott, 1968; Dunkel and Ziegel, 1970), a further antigen, the early antigen (EA), was detected (Henle, Henle, Ho, Burtin, Cachin, Clifford, de Schryver, de The, Diehl, and Klein, 1970; Hinuma, Sairenji, and Ohta-Hatano, 1971). This antigen is another early product of viral replication and is independent of viral DNA synthesis (Gergely et al, 1971). As in the case of the cell membrane antigens it has some resemblance to antigens observed in cell cultures of tumours induced experimentally by oncogenic DNA viruses (Rapp, Butel, Feldman, Kitahara, and Melnick, 1965; Rapp, 1968).

**COMPLEMENT-FIXING ANTIBODIES**

Armstrong, Henle, and Henle (1966) described
complement-fixation tests with an antigen prepared by the disruption of cultured Burkitt tumour cells. Antibodies to this antigen were present in many individuals but they differed clearly from those directed against the VCA antigen. Gerber and Birch (1967) prepared a similar antigen and purified it by sucrose gradient centrifugation. They selected the fractions which contained a high proportion of virus, as assessed by electron microscopy, and observed in their serological tests a close concordance with VCA antibodies: an antigen prepared in the same way from the 'virus-free' RAJI cell line did not react with known positive sera. However, Pope, Horne, and Winters (1969), using a crude antigen, observed complement-fixing activity in preparations of RAJI cells, as well as in preparations of virus-containing cells. It has now become clear that both soluble and sedimentable complement-fixing antigens are present in cultured Burkitt tumour cells and in cultured human leucocyte cultures (Gerber and Deal, 1970; Vonka, Benyesh-Melnick, Lewis, and Wimberly, 1970; Vonka, Benyesh-Melnick, and McCombs, 1970; Hollinshead, Lee, and Alford, 1971). Walters and Pope (1971) have separated three components of their complement-fixing antigen: these are a virion-associated component, and two soluble components, one heat-labile and one heat-resistant. Preparations from RAJI cells contain only the third of these antigenic components. These soluble antigens may also be analogous to the T antigens induced by oncogenic DNA viruses (Shigeta, Minowada, and Moore, 1971).

We have been studying (Sutton, Marston, and Emond, 1971) antibodies to a soluble complement-fixing antigen prepared from virus-containing EB3 cells by the method of Armstrong, Henle, and Henle (1966). A similar antigen prepared from the apparently 'virus-free' RM cell line reacted in the same way with test sera and it is thus possible that our antigen resembles the soluble antigen of Vonka et al (1970).

Over half (64%) of medical students and nurses born in Europe or in North America possessed antibodies; almost all (95%) of those born in tropical areas had similar evidence of previous infection (Table II). In 224 medical students and nurses born in Europe or North America there was a significant association between a history of infectious mononucleosis and the presence of complement-fixing antibodies (Table III) and, in 73 patients admitted to hospital with infectious mononucleosis, compared with 88 patients admitted to the same hospital with other conditions, there was a significant association between low levels of complement-fixing antibodies and active infectious mononucleosis (Table IV). These patients slowly developed antibodies and, when tested some months later, almost all had acquired high levels (Table V).

### Table II EB complement-fixing antibodies in nurses and medical students

<table>
<thead>
<tr>
<th>Origin of Subjects</th>
<th>EB Virus Complement-fixing Antibody Titres</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4 or less</td>
</tr>
<tr>
<td>Africa, West Indies, Asia</td>
<td>23 (95%)</td>
</tr>
<tr>
<td>Europe, North America</td>
<td>144 (64%)</td>
</tr>
</tbody>
</table>

### Table III Relation of EB complement-fixing antibodies to past history of infectious mononucleosis in medical students and nurses

<table>
<thead>
<tr>
<th>History of Infectious Mononucleosis</th>
<th>EB Virus Complement-fixing Antibody Titres</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yes</td>
<td>4</td>
</tr>
<tr>
<td>No</td>
<td>124</td>
</tr>
</tbody>
</table>

\[ p = 0.02 \] (exact test)

### Table IV EB complement-fixing antibodies in patients with infectious mononucleosis and with other conditions

<table>
<thead>
<tr>
<th>Time of Taking Sera</th>
<th>EB Virus Complement-fixing Antibody Titres</th>
</tr>
</thead>
<tbody>
<tr>
<td>During active phase of infectious mononucleosis (mean 0.4 months after onset)</td>
<td>11</td>
</tr>
<tr>
<td>After recovery from infectious mononucleosis (mean 1.4 months after onset)</td>
<td>1</td>
</tr>
</tbody>
</table>

\[ p = 0.008 \] (exact test)

### Table V EB Complement-fixing antibodies in 23 patients with infectious mononucleosis

Unlike the VCA antibodies, soluble complement-fixing antibodies to the EB virus develop slowly after the onset of illness and there is some evidence (Sutton, Almond, Marston, and Emond, 1971) that the combined use of these tests could form the basis for a differential test for recent or remote infection (Table VI).
The EB virus in relation to infectious mononucleosis

<table>
<thead>
<tr>
<th>Period when Sera were Collected</th>
<th>0-2 Weeks</th>
<th>2-24 Weeks</th>
<th>Over 25 Weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sera with Fluorescent Antibody</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Titres &gt; 8</td>
<td>0/4</td>
<td>18/27 (67%)</td>
<td>23/28 (82%)</td>
</tr>
<tr>
<td>Total number of sera tested</td>
<td>20/36 (56%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sera with Complement-fixing Antibody</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Titres &gt; 16</td>
<td>0/7</td>
<td>10/42 (24%)</td>
<td>40/53 (76%)</td>
</tr>
<tr>
<td>Significance of differences</td>
<td>NS</td>
<td>p &lt; 0.05</td>
<td>p &lt; 0.0005 NS</td>
</tr>
</tbody>
</table>

**Table VI Development of fluorescent antibodies and complement-fixing antibodies to EB virus infectious mononucleosis**

**Precipitating Antibodies**

Precipitating antibodies to EB virus antigens have been described by several groups (Old, Boyse, Oettgen, De Harven, Geering, Williamson, and Clifford, 1966; Fink and Cowles, 1968; Stevens, Pry, and Blackham, 1970). Old and his colleagues, using an antigen prepared from sonicated cells and a standard Ouchterlony technique, detected such antibodies in about 50% of patients with Burkitt tumour and in a low proportion (1 of 17) of patients with infectious mononucleosis. Using this method, our results have been similar. Of 55 patients admitted to hospital with infectious mononucleosis, three developed precipitating antibodies from two to nine months after the onset of illness; these sera showed reactions of identity with each other and with a serum from a patient with a Burkitt tumour. Of 199 controls tested, one possessed precipitating antibody: this was a woman of 45 years, admitted with an unrelated complaint, who had an unusually high EB complement-fixing antibody level and who had a considerably (four-fold) raised level of gamma globulins.

Stevens et al (1970) and Fink and Cowles (1968) have used a micro-technique and claim a much greater sensitivity, the former authors finding antibodies in 121 of 261 healthy controls and in 27 of 42 patients with infectious mononucleosis. The antibodies are probably IgG (Oettgen, Aoki, Geering, Boyse, and Old, 1967) and the precipitating antigens are probably associated with EB virus particle (Stevens et al, 1970). The nature of this test lends itself to antigenic comparison, and Fink, King, and Mizell (1968) have claimed to have thus demonstrated the identity of the EB virus with the Lueke frog adenocarcinoma and Ono, Tanabe, Naito, Doi, and Kato (1970) have reported that a herpes-type virus recovered from a chicken with Marek’s disease shared at least one antigen with the EB virus.

**The Response to Infection with EB Virus**

**Virological**

We have seen that a variety of antibodies, directed against the EB virus capsid, against the cell membrane and against several antigens which arise early in the replicative cycle, can be detected in patients with infectious mononucleosis, as well as in those with Burkitt’s tumour or with nasopharyngeal carcinoma. These antigens appear to develop sequentially following cellular infection with the EB virus and in character they are very reminiscent of the antigens specified by some oncogenic DNA viruses, such as SV40 (Rapp, 1968). The presence of DNA virus nucleic acid homology in a virus-free line of Burkitt tumour cells (Zur Hausen and Schulte-Holthausen, 1970), and the recent demonstration by Epstein (1971) that human embryo fibroblasts can be transformed by the EB virus, are also relevant.

Some variation occurs in response to infection and deserves remark. Henle et al (1970) observed that antibodies to early antigens were rarely seen in sera other than those from patients with Burkitt’s tumour, nasopharyngeal carcinoma, or infectious mononucleosis, and Pearson, Henle, and Henle (1971) noted that three lymphoblastoid cell lines responded in differing ways to experimental infection with EB virus, one line producing early antigen but not membrane or viral capsid antigens, another responding with high titres of viral capsid and membrane antigens. Such differences might well influence the expression of virus infection, acting (by analogy with herpes simplex virus) via the glycoprotein specified by the virus (Spear, Keller, and Roizman, 1970), which is detectable by an altered immunological specificity on the cell membrane (Roizman and Spring, 1967) and which is accompanied by alterations in the ‘social behaviour’ of the cell (Roizman, 1971). Whether these differences are due to intratypic variation in the virus or to host variation (possibly genetic in nature) is not known; in the system described by Pearson et al (1971), host variation is clearly important.

These observations chime well with our present knowledge linking the EB virus with Burkitt’s lymphoma and with nasopharyngeal carcinoma, and the connexion with infectious mononucleosis is thought-provoking. Clinical infectious mononucleosis bears no patent relationship with malignant disease although, rarely, it may be followed by Burkitt’s tumour (Cohen, Hirshaut, Stevens, Hull, Davis, and Carbone, 1970) or by Hodgkin’s disease (English, 1970). In the later condition, elevated EB
virus antibody titres have been observed (Levine, Ablashi, Berard, Carbone, and Waggoner, 1970; Johansson, Klein, Henle, and Henle, 1970) and Reed-Sternberg-like cells have been observed in infectious mononucleosis (Lukes, Tindle, and Parker, 1969; McMahon, Gordon, and Rosen, 1970). Although there has been speculation that infectious mononucleosis is an atypical form of acute leukaemia with a self-limiting course (Dameshek and Gunz, 1964), EB virus studies have not supported this and concurrent infectious mononucleosis and acute leukaemia is not uncommon (Lampkin, Canales and Mauer, 1967; Deardorff, Gerber, and Vogler, 1970; Ragab and Vietti, 1969; Stevens, Levine, Lee, Sonley, and Waggoner, 1971).

**IMMUNOLOGICAL**

The EB virus, by virtue of its habitat, is in a favourable position to distort humoral and cellular immune responses. Wollheim and Williams (1966) and our own group (Table VII) have observed considerable increases in immunoglobulin levels, especially in the IgM fraction, during the acute phase of infectious mononucleosis; these high levels normally decline over the ensuing weeks or months. Wollheim and Williams estimated that, in the acute phase of illness, about 5% of the IgM immunoglobulin was attributable to heterophile antibody and, as this declines rapidly, in the later stages almost all remains unaccounted for. Part of this is, presumably, virus antibody, i.e., EB virus antibody, but some of this IgM immunoglobulin may be 'inappropriate antibody' (Waldenström, 1968). Thus, antinuclear factor has been observed in patients with infectious mononucleosis (Wollheim and Williams, 1966, Holborow, Asherson, Johnson, Barnes, and Carmichael, 1963; Carter, 1966), as has anti 'I' (Wollheim and Williams, 1966), rheumatoid factor (Dresner and Trombly, 1959; Holborow et al, 1963), and cryoproteins (Kaplan, 1968). In general, these abnormal antibodies are seen infrequently and, in our series, we have observed a positive latex test in one of 79 patients with infectious mononucleosis, many of whom were tested at various stages during their illness.

There has been at least one report of transient depression of cell-mediated hypersensitivity, as assessed by the tuberculin reaction. Jones (1970) observed that six of 13 patients with acute infectious mononucleosis exhibited negative reactions to tuberculin (100 TU) early in their illnesses which reverted to positive in convalescence. Of our own patients, de Coutinho (personal communication) has observed two who were given BCG at unspecified times before the illness (possibly in childhood) and were tuberculin negative during the acute phase of infectious mononucleosis, reverting to a positive state during convalescence.

**Conclusions**

There is good evidence that much of the population is infected with the EB virus in childhood or later. Most infections are, apparently, asymptomatic but, in some, infection with the EB virus is associated with 'Paul-Bunnell positive' (but not 'Paul-Bunnell negative') infectious mononucleosis. Following EB virus infection, whether accompanied by overt illness or not, a variety of specific antibodies develop. These are directed against the EB virus particle and against antigens specified during early stages of viral synthesis; the latter resemble antigens which are formed during experimental infections with oncogenic DNA viruses.

Infectious mononucleosis, unlike the other conditions for whose aetiology the EB virus is a candidate, is not a malignant condition. Nevertheless, the suspicion of oncogenicity attached to the EB virus and its wide prevalence in the community make closer observation of its host-virus reactions an important object for further investigations. Such work as has been done suggests that the EB virus may be responsible for abnormal humoral and cellular

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**Table VII**  Geometric mean values and Student's t test analyses for age and immunoglobulins in patients with infectious mononucleosis, with other conditions, and healthy medical students and nurses

<table>
<thead>
<tr>
<th>Immunoglobulins (%)</th>
<th>Age (years)</th>
<th>Number Tested</th>
</tr>
</thead>
<tbody>
<tr>
<td>IgM</td>
<td>IgA</td>
<td>IgG</td>
</tr>
<tr>
<td>Patients with infectious mononucleosis</td>
<td>269.5</td>
<td>197.2</td>
</tr>
<tr>
<td>Patients with other conditions</td>
<td>151.6</td>
<td>122.8</td>
</tr>
<tr>
<td>t</td>
<td>5.9</td>
<td>4.6</td>
</tr>
<tr>
<td>P</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Medical students and nurses</td>
<td>123.4</td>
<td>107.6</td>
</tr>
<tr>
<td>t</td>
<td>13.2</td>
<td>7.3</td>
</tr>
<tr>
<td>P</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

1 GMTs of logarithmically transformed data.
responses in man and indicates that further studies in this direction would be fruitful.

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References


