THE ADVANTAGES OF HEŁA CELLS FOR ISOLATION OF RHINOVIRUSES

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Isolation of rhinoviruses from clinical specimens is generally performed in semicontinuous lines of human embryo fibroblasts, such as WI-38 cells (Hayflick and Moorhead, 1961) or in primary cultures of human-embryo kidney. It is acknowledged that isolation may be difficult, that strains of fibroblasts vary in their sensitivity to rhinoviruses (Brown and Tyrrell, 1964), and that the development of a cytopathic effect (CPE) may be slow and uncertain (Olson, Willhight and Buescher, 1972). A HeLa cell line particularly sensitive to rhinoviruses was used by Hamparian, Leagus and Hillemann (1964) and a similar line by Fiala and Kenny (1966). Rhinovirus-sensitive HeLa cells have been used to study rhinovirus replication, for example by Stott and Heath (1970), and under suitable conditions at least 55 rhinovirus serotypes are known to produce plaques in them (Conant, Somerson and Hamparian, 1968; Fiala, 1968). Stott and Tyrrell (1968) suggested that HeLa cells could be used for the laboratory diagnosis of rhinovirus infection. For several years in this Unit, nasal washings from volunteers experimentally infected with rhinoviruses have been examined by inoculating in parallel, cultures of WI-38 cells and “rhinovirus-sensitive” HeLa cells, and higher isolation rates have almost always been achieved in the latter. These findings are reported here, together with the results of experiments on the comparative sensitivity of HeLa cells and two semicontinuous lines of diploid human embryo-lung (HEL) cells to three rhinovirus strains representing different serotypes. Two of the viruses were completely unadapted to tissue culture and the third had had only two passages in vitro.

MATERIALS AND METHODS

Studies on volunteers. Volunteers were housed and observed by methods standard in this Unit (Tyrrell, 1963). They were inoculated with rhinovirus in the course of various experimental studies. For re-isolation of virus, 10 ml of Hanks' balanced salt solution was instilled in 1- or 2-ml amounts into each nostril alternately, and the discharged washing fluid was collected in a petri dish. The fluid was mixed with an equal volume of nutrient broth before storage at -70°C. The samples were tested for the presence of rhinovirus simultaneously in WI-38 and HeLa cells; serum designated HB was used in the tissue culture media throughout this part of the work (see below).

Cell cultures. Rhinovirus-sensitive HeLa cells, obtained in 1966 from Dr V. V. Hamparian, were grown and maintained as described by Stott and Tyrrell (1968). Cell stocks

* Visiting worker from Institute of Hygiene and Epidemiology, Prague.
were subcultured once weekly. Cultures for virus titration were prepared in tubes or petri dishes and were used after two days.

The growth medium for the two lines of semicontinuous HEL cells, WI-38 and MRC-5 (Jacobs, Jones and Baillie, 1970) was Eagle’s Basal Medium (BME) for diploid cells (Grand Island Biological Co.) with 0.11% (w/v) sodium bicarbonate, 100 units per ml of penicillin, 100 μg per ml of streptomycin and 10% bovine serum. Cell stocks were subcultured twice weekly, with a dilution of 1 in 2 at each subculture. Cell cultures for virus inoculation were grown in the same medium as that used for the stock cultures, and were maintained in similar medium containing 2% bovine serum.

Bovine sera were obtained from two sources. Harvard bovine serum (HB) was prepared from blood obtained at a local abattoir. Foetal calf serum (FC) was obtained from Flow Laboratories Ltd.

Strains of the rhinovirus (RV) serotypes 1B, 2, 4, 9, 31 and 43 were used. The passage histories since their original isolation from "wild" colds were: RV1B, two passages in volunteers; RV2 as used for inoculation of volunteers, three or four previous passages in volunteers; RV2 as used for experiments in tissue culture, four passages in volunteers and two in HeLa cells; RV4, three passages in volunteers; RV9, four passages in volunteers interspersed with eight passages in human embryo-lung fibroblasts; RV31, one passage in human-embryo nasal-organ culture and one passage in volunteers; RV43, two passages in volunteers. Each serotype was identified with reference antiserum. Viruses for tissue culture experiments were prepared either as pools of nasal washings from volunteers infected with RV4 or RV31, or as fluid from tissue cultures infected with RV2, and were stored in portions at -70°C.

**Virus titration methods.** Tube cultures of HEL cells or HeLa cells were used for measurement of 50% tissue-culture infectious doses (TCD50) by the Karber method. Virus dilutions were 3.2-fold (0.5 log10) and three tubes were inoculated with each dilution. Cultures were incubated on a roller drum at 33°C and were observed for characteristic CPE after 4 and 6 days in HeLa cells, or after 4, 7, 10 and usually 14 days in HEL cells, or more frequently than this.

Plaque-forming units (PFU) of RV2 were estimated in HeLa cells by the method of Stott and Heath (1970) with 1 log10 dilution steps and four petri dishes per dilution.

**Experiments in tissue culture.** Each week, HeLa cells and the two lines of HEL cells were subcultured in parallel with single batches of BME and serum, either HB or FC, as required for the various cell types, and cultures were prepared in tubes and petri dishes for virus titrations. The WI-38 cells were used from their 19th to 27th subcultures, and the MRC-5 cells from their 17th to 25th subcultures.

When making comparative virus titrations in the three cell types a single set of virus dilutions, prepared from each virus pool, was used.

**RESULTS**

**Isolation of rhinoviruses from volunteers**

Table I shows the frequency of rhinovirus isolation from infected volunteers, in WI-38 cells and HeLa cells inoculated in parallel. This was almost always higher in HeLa cells, sometimes markedly so.

**Experiments in tissue culture**

The three viruses used for these experiments each produced a characteristic CPE in both HeLa and HEL cells. The M strains RV2 and RV31 produced a more rapidly spreading CPE than did the H strain RV4. The latter virus, when inoculated in low titres, produced recognisable foci of CPE, but in both HeLa
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In fibroblasts the foci frequently did not progress to complete destruction of the cell sheet; however, a typical CPE was reproduced on subculture.

### Table I

Comparison of the sensitivity of WI-38 cells and HeLa cells for the isolation of rhinoviruses from experimentally-infected volunteers

<table>
<thead>
<tr>
<th>Virus</th>
<th>Number of specimens tested</th>
<th>Number of isolations made in WI-38 cells</th>
<th>Number of isolations made in HeLa cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>RV2</td>
<td>73</td>
<td>25</td>
<td>56</td>
</tr>
<tr>
<td>RV2+RV43</td>
<td>22</td>
<td>16</td>
<td>20</td>
</tr>
<tr>
<td>RV9</td>
<td>99</td>
<td>41</td>
<td>52</td>
</tr>
<tr>
<td>RV1B</td>
<td>4*</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

* The mean infectivity titre of these specimens was 10^{1.2} TCD50 per ml in WI-38 cells and 10^{2.1} TCD50 per ml in HeLa cells.

With low-titre inocula of RV4 and RV31 a CPE usually appeared within 5 days in HeLa cells, but was often much more delayed in HEL cells.

**Comparison of HeLa, WI-38 and MRC-5 cells.** This was designed to compare the sensitivities of HeLa, WI-38 and MRC-5 cells, grown and maintained in media containing HB, for infectivity titrations of the three viruses. Table II shows the results of five experiments set up in consecutive weeks. HeLa cells were the most sensitive for all three viruses. The mean titres of RV2 obtained from the five experiments were 10^{6.54} PFU/ml in HeLa cells, 10^{4.28} TCD50/ml in WI-38 cells and 10^{3.97} TCD50/ml in MRC-5 cells (four tests only). It should

### Table II

Infectivity titres of three rhinoviruses pools measured in HeLa, WI-38 and MRC-5 cells, in five successive tests

<table>
<thead>
<tr>
<th>Virus</th>
<th>Cell line</th>
<th>Infectivity titres (log_{10} PFU per ml or TCD50 per ml)* obtained in test number</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>RV2</td>
<td>HeLa</td>
<td>7.0</td>
</tr>
<tr>
<td></td>
<td>WI-38</td>
<td>4.7</td>
</tr>
<tr>
<td></td>
<td>MRC-5</td>
<td>NT</td>
</tr>
<tr>
<td>RV4</td>
<td>HeLa</td>
<td>2.4</td>
</tr>
<tr>
<td></td>
<td>WI-38</td>
<td>&lt;1.9</td>
</tr>
<tr>
<td></td>
<td>MRC-5</td>
<td>NT</td>
</tr>
<tr>
<td>RV31</td>
<td>HeLa</td>
<td>NT</td>
</tr>
<tr>
<td></td>
<td>WI-38</td>
<td>NT</td>
</tr>
<tr>
<td></td>
<td>MRC-5</td>
<td>NT</td>
</tr>
</tbody>
</table>

* RV2 was titrated in HeLa cells as plaque-forming units per ml (PFU per ml); all other titres are expressed as 50% tissue-culture doses per ml (TCD50 per ml). The titres given for RV2 in HeLa cells would be about 0.4 log_{10} units lower if measured as TCD50 per ml.

NT = Not tested.
be noted, however, that titres of RV2 in HeLa cells were measured by an
enumerative (plaque) method, as PFU/ml, whereas all the other titrations were
quantal (50% endpoint) assays measuring TCD50/ml. The latter method
theoretically, though not necessarily in practice, gives titres 1.44 times higher
than plaque assays carried out in the same cells. To compare the sensitivity
of the two techniques in HeLa cells under the conditions of our tests, a pool of
RV2 was titrated five times by both methods. The plaque method actually gave
infectivity titres (mean, 10⁷.08 PFU per ml) which were about 0.4 log10 units

**TABLE III**

*Differences in infectivity titre obtained with three rhinoviruses, in several replicate experiments in parallel sublines of HeLa, WI-38 or MRC-5 cells, grown in medium containing either foetal or non-foetal bovine serum*

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Virus</th>
<th>Differences* between the log10 TCD50 values obtained in FC-containing and HB-containing cultures</th>
<th>Means of values shown in preceding column for each cell line</th>
</tr>
</thead>
<tbody>
<tr>
<td>HeLa</td>
<td>RV2</td>
<td>−0.5, +0.5, 0</td>
<td>−0.055</td>
</tr>
<tr>
<td>HeLa</td>
<td>RV4</td>
<td>0, 0, +0.2</td>
<td></td>
</tr>
<tr>
<td>HeLa</td>
<td>RV31</td>
<td>+0.2, −0.5, −0.4</td>
<td></td>
</tr>
<tr>
<td>WI-38</td>
<td>RV2</td>
<td>+0.5, 0, +1.3</td>
<td>+0.56 (p&lt;0.1)†</td>
</tr>
<tr>
<td>WI-38</td>
<td>RV4</td>
<td>+0.5, +0.5</td>
<td></td>
</tr>
<tr>
<td>MRC-5</td>
<td>RV2</td>
<td>+0.3, +0.2, −0.1, +0.7, +1.3</td>
<td>&gt; +0.36 (p&lt;0.02)†</td>
</tr>
<tr>
<td>MRC-5</td>
<td>RV4</td>
<td>+0.1 (in 3 experiments), +0.8, &gt; +0.3</td>
<td></td>
</tr>
<tr>
<td>MRC-5</td>
<td>RV31</td>
<td>&gt; +0.2</td>
<td></td>
</tr>
</tbody>
</table>

* The differences are those observed in replicate experiments.
† Significance of difference from zero (t-test, method of paired comparisons).
FC = Foetal calf serum. HB = Harvard bovine serum (non-foetal).

(2.5-fold) higher than the 50% endpoint method (mean, 10⁶.66 TCD50 per ml),
presumably because of the more efficient virus absorption of the plaque tech-
technique. Therefore, if the titrations of RV2 in HeLa cells, recorded in table II as
PFU per ml, had instead been carried out by the 50% endpoint method, the
readings obtained would probably have been about 0.4 log10 units lower.
However, the RV2 titres (TCD50 per ml) in the HEL cells were too far below
the titres (actual PFU per ml or estimated TCD50 per ml) obtained in the HeLa
cells for the differences to be attributable solely to the change in titration tech-
nique. The low sensitivity of HEL cells, compared with that of HeLa cells, was
confirmed by the results obtained with RV4 and RV31; their titres, measured in
TDC50 per ml in all three cell lines, were higher in HeLa cells than in either of
the HEL cell lines. When RV4 and RV31 were used, a CPE often did not occur
at all in HEL cells within the 2-week observation period, whereas in HeLa cells,
a CPE was detectable within 5 days. The two HEL cell lines were similar in
their sensitivity to the three rhinoviruses; the WI-38 cells were possibly slightly
more sensitive than the MRC-5 cells. There was no evidence that the passage
level of the HEL cells influenced their virus sensitivity.
Comparison of the effects of foetal and non-foetal bovine serum. In the preceding series of experiments all media contained non-foetal HB. A second series of experiments was carried out to compare the effects of foetal and non-foetal sera on the sensitivity of cells to RV infection. Preliminary passages of the three cell lines in media containing different batches of FC were necessary, to select serum which gave good cell growth. Each cell line was then maintained as two parallel sublines in medium containing either HB or FC for between one and seven subcultures. In a series of replicate experiments the three rhinoviruses were titrated in parallel in the three pairs of sublines. Table III shows the differences between the titres (TCD50) obtained with each virus in the parallel sublines. The MRC-5 and WI-38 cells maintained in medium containing FC generally gave slightly higher titres than in medium containing HB, and the difference was statistically significant for MRC-5 cells, with which a larger number of tests was made. The kind of serum used had no obvious effect on the virus titres obtained with HeLa cells.

**DISCUSSION**

Our results show that a suitable line of HeLa cells can be more sensitive for detecting rhinoviruses in nasal secretions than the semicontinuous HEL cells customarily used for this purpose. Only six serotypes were used but it seems likely that the findings will apply generally. Although small inocula of poorly-growing serotypes often produced only focal and non-progressive CPE in HeLa cells, the lesions were clearly recognisable and appeared more rapidly than in HEL cells.

Repeated experiments with nasal washings from infected volunteers, over a period of many months, indicated that HeLa cells were better than WI-38 cells for virus isolation when both lines were grown in medium containing non-foetal bovine serum. Infectivity titrations with rhinoviruses which were minimally adapted or entirely unadapted to tissue culture confirmed that HeLa cells were more sensitive than WI-38 cells to "wild" viruses. Although the sensitivity of both WI-38 and MRC-5 cells increased somewhat when a suitable foetal bovine serum was used instead of non-foetal serum, HeLa cells remained superior. The sensitivity of HeLa cells was unaffected by the type of serum used, and it is interesting to speculate as to the mechanism by which non-foetal serum improved the sensitivity of one cell type but not another. The fact that both the HEL and HeLa cells grew equally well in media containing FC or HB suggests that a viral inhibitor active only in HEL cells may have been present in the non-foetal serum.

**SUMMARY**

A sensitive line of HeLa cells was found to be better than two lines of diploid human embryo-lung (HEL) cells, WI-38 and MRC-5, for the isolation of several serotypes of rhinovirus from experimentally-infected volunteers. Comparative infectivity titrations of three rhinoviruses which were either unadapted or only partly adapted to tissue culture confirmed the greater sensitivity of the HeLa cells. The substitution of foetal calf serum for non-foetal bovine
serum in the tissue culture medium slightly enhanced the sensitivity of the HEL cells to rhinoviruses but had no such effect on HeLa cells; HeLa cells nevertheless remained more sensitive than HEL cells. The line of HeLa cells studied merits more general use for the isolation of rhinoviruses from clinical specimens.

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REFERENCE


