ABSTRACT. The calcium-independent mechanism of cell adhesion was studied in normal and polyoma virus-transformed BHK cells. The degree of Ca\textsuperscript{2+}-independent adhesion was greatly reduced in pyBHK cells, whereas Ca\textsuperscript{2+}-dependent adhesion occurred to the same degree as in BHK cells. This decrease was shown not to be caused by simple masking of the adhesion sites or by their altered sensitivity to trypsin.

Adhesion-blocking antibodies were used to identify molecules responsible for Ca\textsuperscript{2+}-independent adhesion. The antibodies precipitated surface molecules specific for adhesion-competent cells. These have tentatively been named CIDS\textsubscript{BHK} and CIDS\textsubscript{pyBHK}. Both were glycoproteins with respective apparent molecular weights of 120K and 125K. CIDS\textsubscript{pyBHK} incorporated \textsuperscript{3}H-glucosamine more than CIDS\textsubscript{BHK} did. Possible modification of the Ca\textsuperscript{2+}-independent adhesion mechanism in pyBHK cells is discussed.

Studies of cell adhesion are of fundamental importance for the analysis of malignancy as well as for understanding animal morphogenesis. Thus, we have been studying adhesion mechanisms in BHK and pyBHK cells (1, 2). We previously showed that both cell types possess Ca\textsuperscript{2+}-dependent (CD) and Ca\textsuperscript{2+}-independent (CID) mechanisms for cell adhesion and that the degree of CID aggregation is greatly reduced in pyBHK cells, whereas CD aggregation in these cells occurs as in BHK cells (2).

Ozaki et al. reported that microtubule-assembly is not related to the CID adhesion mechanisms of BHK and pyBHK cells (3). Therefore, we need to know what is the molecular basis for the CID adhesion mechanisms of these cells. There have been several reports published on transformation-related changes in cellular products (4–7), and some are thought to be relevant to cell adhesion. For example, fibronectin is known to be markedly reduced after oncogenic transformation (6). Vinculin, a recently found cytoskeletal component (8, 9), is phosphorylated in Rous sarcoma virus (RSV)-transformed cells (7). To our knowledge, however, no alteration of the cell adhesion molecule itself has yet been reported.

The CID mechanism of cell adhesion was studied extensively in our previous research on Chinese hamster V79 cells (10, 11). The molecule responsible for the CID aggregation of V79 cells (CIDS\textsubscript{V79}) was shown by immunochemistry to be a glycopro-
tein with an apparent M.W. of 125,000 daltons (11). In the present study reported here, we found that Fab fragments prepared from the antiserum that detected CIDS inhibited the CID aggregation of BHK and pyBHK cells. Thus, we have tried using this antiserum to identify molecules related to the CID adhesion of these cells.

**MATERIALS AND METHODS**

**Cells.** BHK21 clone 13 (BHK), polyoma virus-transformed BHK (pyBHK) (1) and Chinese hamster V79 cells (12) were used. Cells were grown in Dulbecco’s modified Eagle’s MEM supplemented with 10% fetal calf serum (FCS).

**Cell Dissociation and Reaggregation.** Cells in monolayer cultures were dispersed for the reaggregation experiments as described previously (2). For preparation of cells that had only the CID adhesion mechanisms (LTE-cells), cells were dispersed at 37°C for 15 min with 1 µg/ml crystallized trypsin (Sigma, Type I) in Ca²⁺- and Mg²⁺-free saline buffered with HEPES at pH 7.4 (HCMF) (13) containing 1 mM EDTA. Cells with no adhesion mechanisms (TE-cells) were prepared with 100 µg/ml trypsin in HCMF containing 1 mM EDTA.

Dissociated cells (4 × 10⁵ in 0.5 ml HCMF) were inoculated into one well (15 mm diameter) of a multi-well plate (Falcon 3008) then incubated on a gyratory shaker (New Brunswick, Model G24) at 80 rpm and 37°C. The degree of aggregation was represented by an aggregation index defined as Nt/No, in which Nt and No are the total number of particles per well at times t and 0. The degree of inhibition of aggregation by antiserum or enzymes was represented by the inhibition index Iag defined as Iag=[(Nt(c)—Nt(0))/No —Nt(0)] × 100 (%), in which Nt(c) is the total number of particles at incubation time, t, in the presence of reagents at concentration, c (11). Negative values of Iag indicate the promotion of aggregation.

**Preparation of Antibodies.** Preparation of the antiserum has been described elsewhere. Anti E-V79 antiserum (10) was obtained from rabbits immunized with whole V79 cells that had been dissociated with EDTA and anti TRF antiserum (11) with trypsin-released fragments from V79 cells. Immunoglobulin G(IgG) and Fab fragments of the above antisera and of normal rabbit serum were prepared as described previously (10).

**Radio-labeling of Cell Components.** Surface proteins of the cells were labelled with ¹²⁵I by the lactoperoxidase-mediated method (14). Dispersed cells (1 × 10⁷) were suspended in 2 ml of iodination solution containing 800 µCi Na ¹²⁵I (carrier free, New England Nuclear), 40 µg lactoperoxidase (Miles), 0.2 units glucose oxidase (Miles) and 5 mM glucose in HCMF then incubated for 10 min in an ice bath with occasional shaking.

Polysaccharides of cells were labelled for 24 h with 50 µCi/ml of ³H-glucosamine (32.5 Ci/mM). Cell density was approximately 7 × 10⁴/cm² at the start and increased to about 1.5 × 10⁵/cm² at the end of labelling.

**Immunoprecipitation.** Immunoprecipitation was done by modification of Kessler’s method with formalin-fixed *Staphylococcus aureus* (Calbiochem) (15). The bacteria were heat treated then incubated with 10 µl of rabbit antiserum in a total volume of 1 ml of TBS (10 mM Tris-HCl pH 7.8 containing 0.15 M NaCl) in an ice bath for 30 min. The antiserum-bacteria complex was centrifuged at 3,000 × g for 5 min then suspended in 100 µl TBS. Radio-labelled cells were made soluble with 1 ml of detergent buffer containing 1% Triton X-100 and 0.1% deoxycholate in TBS by incubating them at room temperature for 30 min. Insoluble material was removed by centrifugation at 15,000 × g for 20 min. *S. aureus*, treated as above was added to the lysate, then the mixture was incubated in an ice bath for 30 min. After incubation, the bacteria were washed several times with TBS. Antigens were released by boiling the adsorb-
ents in the SDS sample buffer used for electrophoresis for 3 min.

**Electrophoresis and Autoradiography.** SDS-polyacrylamide gel electrophoresis was done according to the method of Laemmli (16), after which the gels were fixed and stained with Vesterbeg’s stain (450 ml methanol, 1.5 g Coomasie brilliant blue, 180 g TCA, 54 g sulfosalicylic acid and 1.1 liters H₂O) for several hours at room temperature.

After destaining, gels with ³H-labelled samples were impregnated with an autoradiography enhancer (EN³HANCE, New England Nuclear) then soaked in water, after which they were dried and exposed to X-ray film (XAR-5, Kodak). Gels with ¹²⁵I-labelled samples were autoradiographed with the same X-ray film.

**RESULTS**

**Properties of the CID Aggregation of BHK and pyBHK Cells.**

1. **Aggregation kinetics.** Typical curves for the aggregation kinetics of LTE-BHK and LTE-pyBHK cells are shown in Fig. 1. Aggregation of the latter cells is always less than that of the former, and aggregations of both cell types are Ca²⁺-independent. Promotion by Ca²⁺ in the aggregation medium after 1 h of incubation is due to the recovery of the CD mechanism because cycloheximide inhibited this promotion.

2. **Trypsin sensitivity.** To check the possibility that the decreased aggregating ability of pyBHK cells is caused by differences in the trypsin sensitivity of the adhesion mechanisms in those cells, we treated cell monolayers with various concentrations of trypsin in the presence of EDTA then reaggregated the dispersed cells. The degree of the CID aggregation of pyBHK cells was always less than that of the BHK cells throughout the range examined (Fig. 2). Trypsin treatment at concentrations of less than 1 µg/ml somewhat increased the aggregating ability of both cell types. Above this concentration the adhesiveness of the cells decreased with the increase in trypsin concentration and on trypsinization at 100 µg/ml cells became almost non-adhesive. Fifty percent inhibition was achieved ca. 50 µg/ml trypsin for both BHK and pyBHK cells. LTE-cells are represented by (a) and TE-cells by (b) in Fig. 2.

3. **Inhibition of aggregation by anti E-V79 Fab.** Anti E-V79 Fab inhibited the aggregation of LTE-BHK and LTE-pyBHK cells as well as that of LTE-V79 cells.

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Fig. 1. Aggregation kinetics of LTE-BHK and LTE-pyBHK cells. LTE-BHK (—) and LTE-pyBHK cells (—) were aggregated in the presence (○) and absence (□) of 1 mM Ca²⁺ in the aggregation medium.
Table 1. Aggregation of these cells was inhibited completely at high concentrations (2 mg/ml) of Fab. As judged by the Fab concentrations needed for 50% inhibition, anti E-V79 Fab was least effective on BHK cells. Fab from normal rabbit serum promoted, rather than inhibited, the aggregation of all the cells, the lag being around −5%. Anti-TRF Fab inhibited neither the aggregation of LTE-BHK nor LTE-pyBHK cells.

4 Effect of hyaluronidase. Because hyaluronidase synergistically inhibited the aggregation of LTE-V79 cells with anti-TRF Fab (11), we examined the involvement of hyaluronic residues in the Ca2+-independent adhesion in BHK and pyBHK cells by treating LTE-cells with hyaluronidase from Streptomyces hyalurolyticus (Seikagaku Kogyo, Japan). The enzyme had opposite effects on LTE-BHK and LTE-pyBHK cells. The former cells became slightly less adhesive on enzyme treatment, whereas LTE-pyBHK cells became more adhesive, but the degree of aggregation for pyBHK cells was lower than that for BHK cells even after the most effective treatment.

Table 1. Inhibition of Aggregation by Anti E-V79 Fab

<table>
<thead>
<tr>
<th>Cell</th>
<th>Inhibition* by Fab at 0.1 mg/ml</th>
<th>Inhibition* by Fab at 0.2 mg/ml</th>
<th>Inhibition* by Fab at 2 mg/ml</th>
<th>Fab concentration for 50% inhibition</th>
</tr>
</thead>
<tbody>
<tr>
<td>LTE-BHK</td>
<td>19.2</td>
<td>64.6</td>
<td>100</td>
<td>0.16 mg/ml</td>
</tr>
<tr>
<td>LTE-pyBHK</td>
<td>29.2</td>
<td>83.7</td>
<td>100</td>
<td>0.11</td>
</tr>
<tr>
<td>LTE-V79</td>
<td>45.8</td>
<td>66.7</td>
<td>100</td>
<td>0.11</td>
</tr>
</tbody>
</table>

LTE-cells were aggregated in the presence and absence of anti E-V79 Fab for 1 h, then the inhibition index (lag) was calculated.

* Indicated as the lag (%) defined in the text.
Immunoprecipitation of Adhesion Molecules. For the identification of molecules related to the CID adhesion of BHK and pyBHK cells, LTE- and TE-cells were prepared from BHK, pyBHK and V79 cells, and their surface proteins were radioiodinated. Lysates of the labelled cells were immunoprecipitated with anti E-V79 antiserum and these precipitates were analyzed by SDS-polyacrylamide gel electrophoresis followed by autoradiography. Several bands were detected in each lane of the autoradiograms (Fig. 3), but a comparison of the LTE- and TE-cells of each cell type showed that only one band predominates in the LTE-cells (asterisks in Fig. 3(A) and (B)). The LTE-specific band in V79 cells coincides with the position of CIDS\textsubscript{v79}

**TABLE 2. EFFECT OF HYALURONIDASE TREATMENT ON AGGREGATION OF LTE-BHK AND LTE-pyBHK CELLS**

<table>
<thead>
<tr>
<th>Hyaluronidase concentration (U/ml)</th>
<th>LTE-BHK</th>
<th>LTE-pyBHK</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Aggregation (N\textsubscript{t}/N\textsubscript{a})</td>
<td>Inhibition (lag(%))</td>
</tr>
<tr>
<td>0\textsuperscript{a}</td>
<td>.149</td>
<td>0</td>
</tr>
<tr>
<td>20</td>
<td>.168</td>
<td>2.2</td>
</tr>
<tr>
<td>40</td>
<td>.183</td>
<td>4.0</td>
</tr>
<tr>
<td>80</td>
<td>.214</td>
<td>7.6</td>
</tr>
<tr>
<td>100</td>
<td>.196</td>
<td>5.5</td>
</tr>
<tr>
<td>No treatment\textsuperscript{b}</td>
<td>.096</td>
<td>---</td>
</tr>
</tbody>
</table>

LTE-BHK and LTE-pyBHK cells were treated with hyaluronidase and reaggregated for 1 h. For the enzyme treatment, 3 x 10⁶ cells were incubated with appropriate concentrations of hyaluronidase in 0.5 ml HCMF, pH 6.6, at 36°C for 60 min with vigorous shaking. For explanation of aggregation and inhibition indices, see text.

\(\textsuperscript{a}\) LTE-cells were incubated without hyaluronidase at 37°C for 1 h.

\(\textsuperscript{b}\) LTE-cells were reaggregated for 1 h without incubation with or without hyaluronidase.

Fig. 3. Immunological detection of CIDS's in BHK, pyBHK and V79 cells. Radio-iodinated LTE-(a) and TE-cells (b) were lysed and immunoprecipitated with anti E-V79 antiserum. The immunoprecipitates were separated by SDS-polyacrylamide gel electrophoresis. Autoradiograms of 7.5% (A) and 15% (B) gels are shown. Asterisks show bands specific for LTE-cell precipitates.
detected with anti TRF. These LTE-cell specific bands probably represent CID adhesion molecules as they exist on the surface of CID adhesion-competent cells but not on adhesion-incompetent cells, and they react with adhesion-blocking antibodies. These bands have tentatively been named CIDS<sub>BHK</sub> and CIDS<sub>pyBHK</sub>. Some bands also are specific to TE-cells. They probably represent inactivated forms of CIDS's.

Close examination of the autoradiogram showed that CIDS<sub>pyBHK</sub> migrated more slowly than did CIDS<sub>BHK</sub>. This difference was reproducible by repeated immunoprecipitation. The molecular weights were estimated as ca. 120K for CIDS<sub>BHK</sub> and 125K for CIDS<sub>pyBHK</sub>. There was no detectable anti TRF precipitation of CIDS's in BHK and pyBHK cells.

*<sup>3</sup>H-glucosamine Labelling of CIDS's. When LTE-BHK and LTE-pyBHK cells were prepared from *<sup>3</sup>H-glucosamine-labelled BHK and pyBHK cells then lysed and immunoprecipitated with anti E-V79 antiserum, components with molecular weights corresponding to those of CIDS<sub>BHK</sub> and CIDS<sub>pyBHK</sub> were detected on a fluorogram (arrowheads, Fig. 4). Both types of components were absent from precipitates of TE-BHK and TE-pyBHK cells. There were some other bands developed less intensively in TE-cells than in LTE-cells in both the BHK and pyBHK cells. Although the conditions for radiolabelling, immunoprecipitation and electrophoresis were exactly the same for all four types of cells, the radioactivity of samples from pyBHK cells was higher than that from BHK cells. Differences between the two cell types are summarized in Table 3.

**DISCUSSION**

Our previous research has shown that there are Ca<sup>2+</sup>-dependent (CD) and Ca<sup>2+</sup>-
Modified Adhesion in Transformed Cells

One million cells were cultured in the presence of 50 μCi/ml 3H-glucosamine for 24 h. LTE- and TE-cells were prepared from these labelled cells, then lysed and immunoprecipitated with anti E-V79 antiserum, after which they were electrophoresed. Samples of cell lysates and precipitates were used to measure radioactivities of the whole cell lysates and of the total precipitates. For measurement of the radioactivity of CIDS’s, the gel was superimposed on its fluorogram and corresponding regions of the gel were cut out. All radioactivities shown were adjusted to $2 \times 10^6$ cells.

independent (CID) mechanisms for cell-to-cell adhesion in BHK and pyBHK cells and that the degree of CID adhesion is much less in pyBHK cells than in their normal counterparts (2). The results described above suggest that this decrease is caused by an alteration(s) in the adhesion molecule itself rather than being caused by different trypsin sensitivity or simple masking of adhesion sites by proteinaceous substances.

To identify the molecules responsible for the CID aggregation of BHK and pyBHK cells, we used anti E-V79 antiserum, an antiserum originally raised for the study of adhesion molecules in Chinese hamster V79 cells (10). This use was based on the fact that the CIDS’s in various cells can cross-react to a degree (17). In fact, Fab fragments prepared from anti E-V79 antiserum inhibited the aggregation of LTE-BHK and LTE-pyBHK cells. Among the target molecules of the anti E-V79 antibodies we detected surface molecules specific to LTE-BHK or to LTE-pyBHK cells. These we have tentatively named CIDSBHK and CIDSpyBHK. Although further confirmation is necessary, these probably are adhesion molecules because they are the only molecules found that satisfy the characteristics of adhesion molecules. i.e., CIDSBHK and CIDSpyBHK exist on the surface of CID-adhesion competent cells (LTE-cells) but not on the surface of CID-adhesion incompetent cells (TE-cells), and they react with adhesion-blocking antibodies.

The detection of CIDSBHK and CIDSpyBHK revealed a striking similarity among the CIDS’s of BHK, pyBHK and V79 cells. All are glycoproteins with apparent molecular weights of ca. 120,000 daltons and are immunologically cross-reactive. Details of their natures, however, differ. CIDSBHK and CIDSpyBHK did not react with anti TRF antibodies in either the inhibition-of-aggregation assay or in immunoprecipitation. This probably was because the antiserum mainly detects antigenic determinants specific for Chinese hamster cells. This phenomenon is interesting because of the generality and specificity of the CID mechanism of cell adhesion in different cell types.

The molecular weights of CIDSBHK and CIDSpyBHK are very close, but the weight of the latter is slightly larger. Although there was no quantitative difference between iodinated CIDSBHK and CIDSpyBHK that would explain the decreased adhesiveness of pyBHK cells, the incorporation of 3H-glucosamine differed markedly. As increased

<table>
<thead>
<tr>
<th></th>
<th>BHK</th>
<th>pyBHK</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LTE</td>
<td>TE</td>
</tr>
<tr>
<td>Whole cell lysate</td>
<td>$1.67 \times 10^7$</td>
<td>$1.18 \times 10^7$</td>
</tr>
<tr>
<td>Whole precipitate</td>
<td>$1.3 \times 10^4$</td>
<td>$0.9 \times 10^4$</td>
</tr>
<tr>
<td>CIDS</td>
<td>$13 \times 10^2$</td>
<td>$0.4 \times 10^2$</td>
</tr>
<tr>
<td>CIDS/TOTAL</td>
<td>$8.1 \times 10^{-5}$</td>
<td>$0.4 \times 10^{-5}$</td>
</tr>
</tbody>
</table>

One million cells were cultured in the presence of 50 μCi/ml 3H-glucosamine for 24 h. LTE- and TE-cells were prepared from these labelled cells, then lysed and immunoprecipitated with anti E-V79 antiserum, after which they were electrophoresed. Samples of cell lysates and precipitates were used to measure radioactivities of the whole cell lysates and of the total precipitates. For measurement of the radioactivity of CIDS’s, the gel was superimposed on its fluorogram and corresponding regions of the gel were cut out. All radioactivities shown were adjusted to $2 \times 10^6$ cells.

TABLE 3. INCORPORATION OF RADIOACTIVITY TO BHK AND pyBHK CELLS METABOLICALLY LABELLED WITH 3H-GLUCOSAMINE
incorporation into pyBHK cells is not solely a phenomenon of CIDS<sub>pyBHK</sub> but is common to the components of all the pyBHK cells, it seems to confirm other findings that sugar chains in transformed cells are generally larger than their normal counterparts (18). Overglycosylation of cellular constituents can be assumed to be reflected in the larger molecular weight of CIDS pyBHK and to be the cause of the decrease in its ability to bind cells. This assumption could explain why hyaluronidase treatment of LTE-pyBHK cells partially restored the aggregating ability of pyBHK cells.

Recent findings have shown that a product of the src gene of RSV, (pp60<sup>src</sup>), is associated with protein kinase activity (19, 20) and that many products in RSV-transformed cells are phosphorylated (21). Similar alterations in enzymatic activities have been found in other transformed cells (22). Thus, we need to determine the relationship between the expression of polyoma T-antigens (23) and the modification of adhesion molecules in pyBHK cells. Further studies are necessary to show by what mechanism adhesion molecules are altered during the malignant transformation of cells.

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