Cells transformed by murine herpesvirus 68 (MHV-68) release compounds with transforming and transformed phenotype suppressing activity resembling growth factors

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Summary. – In this study, we investigated the medium of three cell lines transformed with murine herpesvirus 68 (MHV-68) in vitro and in vivo, 68/HDF, 68/NIH3T3, and S11E, for the presence of compounds resembling growth factors of some herpesviruses which have displayed transforming and transformed phenotype suppressing activity in normal and tumor cells. When any of spent medium was added to cell culture we observed the onset of transformed phenotype in baby hamster kidney cells (BHK-21) cells and transformed phenotype suppressing activity in tumor human epithelial cells (HeLa). In media tested, we have identified the presence of putative growth factor related to MHV-68 (MHGF-68). Its bivalent properties have been blocked entirely by antisera against MHV-68 and two monoclonal antibodies against glycoprotein B (gB) of MHV-68 suggesting viral origin of MHGF-68. The results of initial efforts to separate MHGF-68 on FPLC Sephadex G15 column in the absence of salts revealed the loss of its transforming activity but transformed phenotype suppressing activity retained. On the other hand, the use of methanol-water mobile phase on RP-HPLC C18 column allowed separation of MHGF-68 to two compounds. Both separated fractions, had only the transforming activity to normal cells. Further experiments exploring the nature and the structure of hitherto unknown MHGF-68 are now in the progress to characterize its molecular and biological properties.

Keywords: murine herpesvirus; MHV-68 transformed cell line; putative growth factor MHGF-68; separation techniques

In nineties of the 20th century, a new class of compounds encoded by some herpesviruses (pseudorabies and herpes simplex virus (HSV) 2) resembling growth factors has been found in medium of mammalian cells cultured under conditions non-permissive for virus replication. These compounds designed as virus related growth factors (PRGF and HSGF) displayed unique effects on cell culture either infected or transformed with corresponding virus (Golais et al., 1990, 1992a). Non-transformed cells cultivated in the presence of these factors acquired the transformed phenotype (“criss-cross” pattern of growth), while the phenotype of transformed cells became changed towards the normal ones (Gašperík et al., 1994; Konvalina et al., 2002). Both activities could be neutralized by antisera to corresponding virus and also by some monoclonal antibodies directed against viral glycoprotein B (gB) (Golais et al., 1992b). Furthermore, our previous studies on HSV-1 strains that differ in gB sequence (e.g in syn1 locus) revealed, that gB might be involved at least in the function of these factors (Golais et al., 1992b; Rajčáni et al., 1996).

Recently, two cell lines transformed by MHV-68 derived from human dermal fibroblasts and mouse embryonal cells were prepared, in this paper designed as 68/HDF and 68/
NIH3T3. In both, the presence of viral antigen and genome but the absence of infectious virus release have been found (Mrázová et al., 2015). In this study, we demonstrated the presence of putative MHV-68 related growth factor (MHGF-68) in the medium of both MHV-68 transformed cell lines following its biological activities that have arisen after its addition to cultivation medium of normal baby hamster kidney cells (BHK-21) and transformed human epithelial cells (HeLa). We have found that BHK-21 cells cultivated with medium of MHV-68-transformed cells, either with 68/HDF or 68/NIH3T3 cells (Fig. 1a), acquired changed morphology resembling that observed with similar herpesvirus-related growth factors (Dušinská et al., 1994; Konvalina et al., 2002). On the other hand, HeLa cells cultivated with the same media displayed the phenotype of non-transformed cells (Fig. 1c). We have found that compounds present in the media of cell lines transformed with MHV-68 in vitro exhibited similar bivalent properties described earlier for growth factors of herpesviruses. When these media were incubated with rabbit antiserum against MHV-68 (30 min at 37°C) before its testing on BHK-21 or HeLa cell cultures, both activities disappeared. This finding indicated that the observed activities are related to compounds in the medium to be of viral origin. In following experiments we treated these media with several monoclonal antibodies against gB of MHV-68 (Gillet et al., 2006; Glauser et al., 2011). As summarized in Table 1, two out of seven monoclonal antibodies tested, MG-1A12 (recognizing conformational epitope and requir-
Table 1. Neutralization of biological activity of MHGF-68 (present in the medium of 68/HDF and 68/NIH3T3 cells) with monoclonal antibodies against glycoprotein B of MHV-68

<table>
<thead>
<tr>
<th>Monoclonal antibody</th>
<th>Characteristics of monoclonal antibody</th>
<th>Neutralization of biological activity of MHGF-68</th>
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<tr>
<td>MG-4D11</td>
<td>recognizes C-terminal half of gB (55 K), works on western blots</td>
<td>–</td>
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<tr>
<td>MG-1C12</td>
<td>recognizes conformational gB epitope requiring the whole protein</td>
<td>–</td>
</tr>
<tr>
<td>BN-2B11</td>
<td>recognizes conformational epitope in N-terminal half of gB</td>
<td>–</td>
</tr>
<tr>
<td>MG-1A12</td>
<td>recognizes conformational epitope requiring the whole protein</td>
<td>+</td>
</tr>
<tr>
<td>SC-9E8</td>
<td>IgM, recognizes N-terminus of gB, works on western blot, sensitive to cell type-specific O-glycosylation</td>
<td>–</td>
</tr>
<tr>
<td>MG-2H4-D9</td>
<td>recognizes conformational epitope in N-terminal half of gB, neutralization</td>
<td>–</td>
</tr>
<tr>
<td>BN-1A7</td>
<td>recognizes conformational epitope in N-terminal half of gB</td>
<td>–</td>
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</tbody>
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Fig. 2

Determination of MHGF-68 transforming activity to BHK-21 cells by limiting dilution of medium obtained from cultivation of 68/NIH3T3 cells and separated to two components, MHGF_A (fractions 6–12) and MHGF_B (fractions 15–21) on RP-HPLC C18 column using methanol-water mobile phase

Better separation of MHGF-68 obtained from medium of both cell lines was achieved by FPLC on Sephadex G15 column (data not shown). An interesting fact observed was that the choice of mobile phase used determinates MHGF-68 activity retention. While phosphate buffered solution (PBS) (pH 7.2) retained both activities of MHGF-68 (Fig. 3a) (the highest in the fraction 4), the use of redistilled water caused...
the loss of transforming activity of MHGF-68 while suppressing transformed phenotype activity in HeLa cells was retained (Fig. 3b). The latter activity of MHGF-68 (fraction 4) seems to be even better than of any MHGF-68 fraction eluted with PBS. Based on results obtained from both types of chromatography, we assume that the absence of salts or presence of organic solvents might influence the structure important for biological properties of this newly found growth factor.

In the following experiments we compared the results obtained for MHGF-68 present in medium of \textit{in vitro} transformed cells with that of tumor B cell line S11E, derived from an MHV-68 infected mouse suffering from lymphoproliferative disease. The cells of S11E line contain MHV-68 genome in both linear and episomal form and spontaneously release infectious virus (Usherwood \textit{et al.}, 1996). When we cultivated BHK-21 or HeLa cells with the medium of S11E cells we observed transforming or trans-

![Diagram](image-url)
formation suppressing activities in relevant cell culture identical with those observed for MHGF-68 described above (data not shown). Further experiments have shown that both activities present in the medium of S11E cells could be neutralized with antiserum against MHV-68 as well as with the same monoclonal antibodies (MG-1A12 and SC-9E8) described above. Thus, in the medium of S11E cells we very likely identified the same compounds, MHGF-68, found in the media of cell lines transformed by MHV-68 in vitro. The evaluation of its biological properties after separation on the C18 or Sephadex G15 column demonstrated similar biological activities described for MHGF-68 present in the cultivation medium of 68/HDF and 68/NIH3T3 cells (data not shown).

Finally, we identified MHGF-68, a growth factor related to MHV-68, in the medium of in vitro and in vivo transformed with MHV-68 cell lines, 68/HDF, 68/NIH3T3, and S11E. MHGF-68 displayed transforming and transformed phenotype suppressing activity in BHK-21 and HeLa cell culture resembling growth factors of some herpesviruses. We identified MHGF-68 to be vacant of transforming activity in the absence of salts (e.g. by treatment with redistilled water during the separation on the column), which might help to find its potential application in the treatment of tumors in the future. The attempts are now in the progress aimed at uncovering the nature, the structure as well as molecular and other biological properties of MHGF-68, hitherto unknown compounds related to MHV-68.

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References


