Cytolytic replication of echoviruses in colon cancer cell lines

Stina Israelsson¹, Nina Jonsson¹, Maria Gullberg¹,² and A Michael Lindberg¹∗

Abstract

Background: Colorectal cancer is one of the most common cancers in the world, killing nearly 50% of patients afflicted. Though progress is being made within surgery and other complementary treatments, there is still need for new and more effective treatments. Oncolytic virotherapy, meaning that a cancer is cured by viral infection, is a promising field for finding new and improved treatments. We have investigated the oncolytic potential of several low-pathogenic echoviruses with rare clinical occurrence. Echoviruses are members of the enterovirus genus within the family Picornaviridae.

Methods: Six colon cancer cell lines (CaCo-2, HT29, LoVo, SW480, SW620 and T84) were infected by the human enterovirus B species echovirus 12, 15, 17, 26 and 29, and cytopathic effects as well as viral replication efficacy were investigated. Infectivity was also tested in spheroids grown from HT29 cells.

Results: Echovirus 12, 17, 26 and 29 replicated efficiently in almost all cell lines and were considered highly cytolytic. The infectivity of these four viruses was further evaluated in artificial tumors (spheroids), where it was found that echovirus 12, 17 and 26 easily infected the spheroids.

Conclusions: We have found that echovirus 12, 17 and 26 have potential as oncolytic agents against colon cancer, by comparing the cytolytic capacity of five low-pathogenic echoviruses in six colon cancer cell lines and in artificial tumors.

Keywords: enterovirus, echovirus, colon cancer, oncolytic virus, virotherapy

Background

Colorectal cancer is the second and third most commonly diagnosed cancer in the world for women and men respectively. Nearly 50% of those diagnosed with colorectal cancer are killed from it, and in 2008 there were more than 1.2 million new cases estimated of colorectal cancer [1]. Presently the most common treatment of colorectal cancer is surgery, where chemotherapy and radiation may be used as adjuvants [2]. Though progress is made within these therapies, the nearly 50% mortality rate indicates the necessity of new and improved treatments of colorectal cancer.

Viral oncolysis offers a new promising field of cancer treatments. The research area has broadened to include several different mechanisms of antitumoral efficacy; for instance direct tumor cell lysis due to viral replication or viral cytotoxic proteins and transgene expression (reviewed in [3,4]). Many of the oncolytic viruses that have been tested in clinical trials are genetically modified, in particular adenoviruses and herpes simplex virus (reviewed in [5-7]). Several other viruses, including poxviruses, reovirus and Newcastle disease virus have also shown promising antitumor effects in various human malignancies (reviewed in [4,8-10]).

Echoviruses (E) are serotypes belonging in the Human Enterovirus B (HEV-B) species within the enterovirus (EV) genus of Picornaviridae. Picornaviruses are small (diameter about 30 nm), naked, icosahedrically shaped viruses with an RNA genome of about 7.2 to 8.5 kb in length with positive polarity [11,12]. Within the EV genus both E1 and coxsackievirus A21 have been shown to be oncolytic in several human cancers, including malignant melanoma and ovarian cancer [13-18]. In addition, it has been reported that bovine EV shows oncolytic characteristics [19], and attenuated, recombinant poliovirus strains have
been shown to have great oncolytic potential [20-22]. Recently another picornavirus, Seneca valley virus, has also emerged as an oncolytic virotherapy candidate [23,24]. The HEV-B species contains many viruses with rare clinical occurrence and low pathogenicity, making them potentially suitable for oncolytic virotherapy if they show oncolytic characteristics. The serotypes within the HEV-B species that were chosen to be included in this study of cytolytic capacity were E12, E15, E17, E26, and E29. These viruses were all isolated in the 1950's from healthy persons [25-30] and chosen on the basis of their low pathogenicity [31-34]. In this paper, the cytolytic effects of these five serotypes were investigated in six cell lines derived from cancers of the human colon. Colorectal cancer was chosen due to the fecal-oral route of infectivity of enteroviruses [12] and its position as one of the most common forms of cancer with a mortality rate of nearly 50% [1]. Those serotypes that generated the most efficient cytopathic effect (CPE) and highest amount of viral progeny, in cell lines grown as monolayers, were also tested for infectivity in spheroids (reviewed in [35]). Infection of spheroids enables investigation of the capability to spread within an artificial tumor of three-dimensional shape.

Materials and methods

Cell lines and viruses

The colon cancer cell lines HT29, SW480, SW620, LoVo, and T84 were purchased from European Collection of Cell Cultures (ECACC, UK) and CaCo-2 was generously provided by C. Gustafsson-Svärd (Linnaeus University, Kalmar, Sweden). A local variant of green monkey kidney (GMK) cells and Chinese hamster ovary (CHO) cells, kindly provided by H.-C. Selinka (University of Tübingen, Tübingen, Germany), were used as positive and negative control for infectivity studies. SW480, SW620, CaCo-2, GMK and CHO were cultured in Dulbecco's modified Eagle's medium (DMEM), HT29 was cultured in McCoy's 5A medium and LoVo and T84 were cultured in Ham's F12 medium. Medias were supplemented with 100 U/ml penicillin, 0.1 mg/ml streptomycin, 2 mM L-glutamine and 10% newborn calf serum (NCS). 5% NCS were used in GMK and CHO media. CaCo-2 media was additionally supplemented with 0.1 mM non-essential amino acids (NEAA). Cell lines were grown in 37°C and 7.5% CO2 (HT29, SW480, SW620, GMK and CHO) or 5.0% CO2 (CaCo-2, LoVo and T84). E15 Charleston was purchased from American Type Culture Collection (ATCC, VR-45), E12 Travis, E17 CHHE-29 and E26 Coral were generously provided by J.-Å. Liljeqvist, (University of Gothenburg, Gothenburg, Sweden) and E29 JV-10, was kindly provided by M. Roivainen (National Institute for Health and Welfare, Helsinki, Finland).

Viral serotype confirmation

Viral serotypes were confirmed genetically by RNA extraction using QIAamp viral RNA MiniSpinKit (Qiagen) according to manufacturer's manual. RNAs were reverse transcribed (RT) by SuperScript III RT enzyme (Invitrogen) using a primer 5'-ATAAGAATGCGGCCGCT27-3' at 50°C for 60 minutes and the enzyme was inactivated at 70°C. cDNAs were amplified by PCR using PuReTaq Ready-To-Go PCR Beads (GE Healthcare) and primers matching the flanking sequences of the VP1 region of each viral strain. Nucleotide sequences were determined using the ABI Prism BigDye terminator cycle sequencing reaction kit (Applied Biosystems) and a 3130 Genetic Analyzer (Applied Biosystems). Sequencer version 4.5 (Gene Codes Corporation) was used for editing of sequences. Obtained viral sequences were determined by strain using basic local alignment search tool (BLAST, National Center for Biotechnology Information, NCBI). Prior to virus infectivity assays all viruses were propagated and titrated in GMK cells using the endpoint titration method of Sperman and Kärber [36].

Virus infectivity assay

Cells were grown in 6 well plates (HT29, SW480, SW620, LoVo, T84, GMK and CHO) or in T25 bottles (CaCo-2) to a confluence of 50-85%. The cells were washed in phosphate buffered saline (PBS) and serum free medias were added to cells: McCoy’s 5A to HT29, DMEM to SW480, SW620, GMK and CHO and OptiMEM to T84, LoVo and CaCo-2. CaCo-2 media was additionally supplemented with 0.1 mM NEAA. Cells were infected with viruses at a multiplicity of infection (MOI) of 1 in double samples and incubated in room temperature for 60 minutes. The virus containing medias were exchanged for new serum free equivalents, and one sample per cell line and virus (E12, E15, E17, E26 or E29) was frozen while the other remaining sample was incubated in 37°C until complete CPE was visible, or for a maximum of seven days. At the time of complete CPE, or after seven days, cells were photographed using microscopy. All infection assays were performed three times. After freezing and thawing three times, all infection samples were titrated in GMK cells.

Culture of spheroids and spheroid infectivity assay

HT29 spheroids were cultured in 24 well plates by inoculating 1 000 cells per well, coated with 1% agarose in PBS, in 1 ml McCoy’s 5A media supplemented with 100 U/ml penicillin, 0.1 mg/ml streptomycin, 2 mM L-glutamine and 10% NCS. Single spheroids were allowed to form for seven days in 37°C, 7.5% CO2. 104 tissue culture infectious dose 50% (TCID50) of each virus (E12, E17, E26 or E29) were added to spheroids, and the infections were followed for nine days. Infections were repeated on at least two
different occasions and each virus was tested for infectivity in spheroids 14 to 18 times in total.

Viability confirmation of spheroids

HT29 spheroids, uninfected or infected by E29, were washed separately three times in PBS to remove any single cells or unbound viral particles and dispersed in 0.25% trypsin-EDTA solution. McCoy’s media supplemented with 100 U/ml penicillin, 0.1 mg/ml streptomycin, 2 mM L-glutamine and 10% NCS was added. Cells were diluted 1:2 in trypan blue solution, incubated for 10 minutes and examined in a Bürker chamber using microscopy. Trypsinated cells from uninfected spheroids were also allowed to proliferate in 6 well plates for two days.

Results

Viral infectivity in colon cancer cell lines

To investigate the oncolytic potential of E12, E15, E17, E26 and E29 against colorectal cancer initially, the cytopathic replication in six colon cancer cell lines (Table 1) was analyzed. Viral ability to cause CPE was investigated (Table 2, Additional file 1) as well as increase in viral titers during infection (Figure 1). GMK and CHO cells were used as positive and negative infectious control respectively (data not shown).

E12 was able to infect all colon cancer cell lines generating an increase in titer of about four logs, 10 \(^{3.4}\) to 10 \(^{7.8}\) TCID\(_{50}/\)ml, in HT29, SW480, SW620 and CaCo-2 (Figure 1). The mean titer of E12 attached to LoVo (10 \(^{2.89}\) TCID\(_{50}/\)ml) and that of the viral progeny (10 \(^{3.94}\) TCID\(_{50}/\)ml) indicate that the virus replicated in the cell line, but no CPE was observed after seven days (Table 2, Figure 1). In all other colon cancer cell lines complete CPE was observed within two to three days, although the viral mean titer decreased during the infection in T84.

E15 generated complete CPE in CaCo-2, HT29, SW620 and T84 within two to seven days, but the mean titers of viral progeny, compared to titers of initially attached virus particles, were markedly higher only in the case of HT29 and CaCo-2 (Table 2, Figure 1). In T84, complete CPE was observed after two days, but no viral progeny was measured by titration. No infection was indicated in LoVo or SW480.

In the case of E17, the virus bound to all colon cancer cell lines (Figure 1). In all cell lines except LoVo, complete CPE was observed within one to seven days (Table 2). CPE was also supported by an increase in mean titer between bound virus and viral progeny of about two to four logs in all cases where CPE was observed, except in SW680, where the mean titer increased with less than one log (Figure 1). Although E17 attached to the cellular surface of LoVo (mean initial titer of attached viral particles; 10 \(^{3.95}\) TCID\(_{50}/\)ml), no CPE was observed, nor was any viral progeny detected by endpoint titration after seven days.

E26 was able to cause complete CPE in CaCo-2, HT29, SW480 and T84 within two to seven days. In SW480 only partial CPE was observed after seven days, while no CPE was observed in LoVo after the same time (Table 2). During infection viral mean titers increased about five to seven logs in all infected colon cancer cell lines except LoVo, where no viral progeny was indicated by titration (Figure 1). Mean titers of viral progeny reached about 10 \(^{5}\) to 10 \(^{8}\) TCID\(_{50}/\)ml in all cell lines except LoVo.

E29 caused complete CPE in two to seven days in CaCo-2, HT29, SW480 and SW620, generating viral mean titers of about 10 \(^{5}\) to 10 \(^{8}\) TCID\(_{50}/\)ml, even though

| Table 1 Cell lines derived from human colorectal cancers used in this study |
|-----------------------------|-----------------|-----------------|
| Cell line | Colorectal disease | Tumor stage | Primary tumor/Derived from metastatic site |
| CaCo-2 | adenocarcinoma | - | Primary tumor |
| HT29 | adenocarcinoma | - | Primary tumor |
| LoVo | adenocarcinoma | Dukes’ type C | Left supraclavicular region |
| SW480 | adenocarcinoma | Dukes’ type B | Primary tumor |
| SW620 | adenocarcinoma | Dukes’ type C | Lymph node |
| T84 | carcinoma | - | Lung |

Data are adapted from the American Type Culture Collection (ATCC) catalogue.

< Tumor stage is not specified.
the virus attached to the cellular surfaces of all cell lines in very low amounts (Table 2, Figure 1). E29 also caused an infection in T84, generating a viral progeny mean titer of $10^{4.88}$ TCID$_{50}$/ml, but no CPE was observed. LoVo was uninfected by E29.

To summarize these results, E12, E17, E26 and E29 are able to proliferate efficiently and cause cytolytic infection in most of the cell lines included in this study. E15 was excluded at this stage, due to its relatively poor ability to proliferate and cause CPE in the colon cancer cell lines investigated.

**Viral infectivity in HT29 spheroids**

Three-dimensionally in vitro growing cells, referred to as spheroids (reviewed in [35]), allow viruses to infect cells in a more tumor-like environment rather than the infection of a bi-dimensional monolayer. Spheroids of the human colon cancer cell line HT29 were infected by E12, E17, E26 and E29. The results show that E12, E17 and E26 destroyed the spheroids within five to nine days (Figure 2). E29 was not able to destroy HT29 spheroids within nine days. Viability of spheroids, both uninfected and infected by E29, was confirmed by individualization of spheroid cells using trypsin treatment, followed by trypan blue staining (data not shown). Viability of uninfected spheroids was additionally confirmed by allowing the individualized cells to proliferate as monolayers (data not shown).

**Discussion**

Though improvements are being made, patient prognosis of colorectal cancer is still poor [1]. Presently surgical restriction, chemotherapy and radiation are the most common therapeutic approaches [2]. A promising new field in the battle against cancer is oncolytic virotherapy, which is undergoing extensive investigation. In this study, we investigated the cytolytic capacity of low- or apathogenic echoviruses within the HEV-B species in colon cancer cell lines, grown both as monolayers and as three-dimensional spheroids.

Wild type enteroviruses may be suitable as oncolytic viruses for several reasons. (i) Enteroviruses have been studied epidemiologically for over 50 years and many of these viruses are strictly human pathogens causing low- or even asymptomatic infections [12]. It is therefore well known that there are viruses within this genus with a relatively safe profile from an apathological point of view. This is a favorable characteristic of an oncolytic virus. It should be mentioned though, that low-pathogenic enteroviruses might be associated with more severe symptoms occasionally [37,38]. (ii) Wild type enteroviruses with low pathogenicity can be oncolytic without genetic engineering [16,17]. This eliminates the risks of potential reversion to virulent phenotypes of attenuated oncolytic viruses or loss of malignant tissue specificity as a consequence of genetic engineering. (iii) There are antipicornavirus drugs such as Pleconaril. This drug has been used successfully on a compassionate-release basis [39,40]. (iv) Low-pathogenic enteroviruses may be rare in the human population [16,31]. This decreases the risk of patients having an acquired immunity towards the virus prior to deliberate infection for oncolytic virotherapy purposes. (v) Possibly several different serotypes with
similar oncolytic properties within an EV species could be identified. Serotypes could then be altered successively if administering more than one oncolytic virus to a patient. This would provide a strategy to avoid obstacles due to acquired immunity towards already administered viruses. (vi) The small virion size (about 30 nm in diameter) of picornaviruses [11] could be an oncolytic advantage. Fibrillar collagen has proven to inhibit relatively large herpes simplex virus from reaching malignant cells within a tumor, while nanoparticles spread more easily [41]. (vii) Picornaviruses have a relatively fast replication cycle, ranging from five to ten hours [11]. This could also be an advantage for viral oncolysis, since a faster replicating virus may reduce a tumor in size more effectively than a slower replicating virus [42].

We have investigated the cytolytic capacity of E12, E15, E17, E26 and E29 by infection of six colon cancer cell lines derived from both primary tumors and metastases (Table 1). To be able to address the infectious ability of the viruses in these cell lines as explicitly as possible, no serum was added to the medias in these viral infectivity experiments. The results were recorded both by the ability to cause CPE (Table 2, Additional file 1), and by proliferation efficacy (Figure 1). Highly cytolitic viruses with an efficient propagation were also allowed to infect spheroids (Figure 2), to investigate their ability to spread within and destroy an artificial tumor.

When analyzing the results of the monolayer studies (Table 2, Figure 1) CaCo-2, HT29 and SW620 were cytolically infected by all viruses, while LoVo was not cytolically infected by any of the five viruses. However, our results suggest that E12 causes a persistent infection in LoVo. We have not been able to find a likely cause to this infectious disability, and it may be due to both difficulties of viral entry as well as intracellular proliferation factors. A cytolytic infectious cycle within the tumor cells is a favorable characteristic of an oncolytic virus. The release of viral progeny enables spreading to neighboring malignant cells as well as metastatic sites that are distant from primary site of administration. Interestingly, it has been shown that the genetic difference between an EV causing a cytolytic infection and one causing a non-cytolytic infection, in a particular cell line, may be as little as one amino acid change [43].

The viral infections of SW480 and SW620 are interesting, due to the fact that these cell lines originate from the
same cancer patient. SW480 was established from a primary adenocarcinoma of the colon, while SW620 was established from a lymph node metastasis one year later (Table 1). The cell lines seem to respond differently to infection of the same virus. E26 does not cause cytolysis in SW480 to the same extent that it does in SW620 (Table 2). The mean viral progeny of E17 in SW620 is about $10^4$ TCID$_{50}$/ml, while the mean viral progeny in SW480 is more than $10^6$ TCID$_{50}$/ml (Figure 1). E12 and E29 cause CPE faster in SW480 than in SW620, and even though E15 is not able to infect SW480, the virus causes a cytolytic infection in SW620 (Table 2, Figure 1). This indicates that the same viral serotype may have a different impact when infecting cancer cells derived from the same original cancer. Hence, there may be a “sliding therapeutic window” in the progression of a cancer, where a specific oncolytic virus has the most effective anti-cancer effect. In this perspective, several oncolytic viral serotypes with similar oncolytic properties may be very beneficial, not only from an immunological point of view.

All viruses cause a cytolytic infection in T84 except E29, which seems to cause a non-cytolytic infection (Table 2, Figure 1). Strangely, the viral progeny titers of E12 and E15 are lower than the initial titers in T84 (Figure 1). This could to some extent be explained by viral progeny neutralization caused by neutralizing substances released by lysed cells, or less viable progeny production of E12 and E15 in T84. More viral particles adsorbing to the T84 cellular surface than the number of particles entering and infecting the cells could also contribute to this effect. In the case of E12, it is known that decay-accelerating factor (DAF or CD55) is a cellular receptor [44,45]. This receptor may not be enough for cellular entry, hence infection, for DAF-binding picornaviruses [46]. It is possible that unknown entry-receptors of E12 and E15 are not expressed on the surface of T84 or expressed in very low amounts. In addition, viral attachment-induced apoptosis could contribute to the CPE observed in E12 and E15 infected T84 cells. It has been shown that high MOIs of UV-inactivated reovirus virions can induce apoptosis in the absence of viral replication by interacting with cell surface receptors [47]. Due to the inefficient cytolytic replication of E15, this virus was excluded from further studies.

In addition to confirmation of successful lytic infections and measurement of viral progeny following infection, an interesting pattern emerges when comparing binding titers of the different viruses in Figure 1. Several serotypes generate similar binding titers independent of cell line infected. A MOI of 1 was used in the experiment and the time required for complete CPE is not connected to initial binding titers (Table 2). The titers of attached E12 and E17 are relatively high, while the amount of bound E29 (and in some cases E26 and E15) are not detected in all cell lines. This could be the result of different entry strategies. If viruses with very low or undetected binding titers use a highly specific attachment and entry process, infection could be successful even at undetected initial amount of bound virions using the TCID$_{50}$ method. This method does not tell how many infectious particles are present in the original sample, but what dilution of virus will generate CPE in 50% of the cells inoculated [36]. A high titer of bound viruses indicates that relatively many virions interact with the cell surface initially. This could be an effect of a more virion dense attachment and entry strategy, possibly by the use of more than one receptor.

E12, E17, E26 and E29 were further investigated in spheroids, cells grown in three-dimensional clusters, hence more resembling tumors than cells grown as monolayers. The results (Figure 2) show that E12, E17 and E26 were able to dissolve spheroids of HT29 cells grown for seven days, starting with 1 000 cells. E29 was not able to efficiently kill HT29 cells when grown as spheroids. These cells were only readily infected and lysed by E29 when grown as monolayers (Table 2, Figure 1). The viability of E29 infected and uninfected spheroids were confirmed after seven days by trypsinization and trypan blue staining. Trypan blue staining showed that E29 had some cytolytic effect on the spheroids, but not enough to dissolve them. Almost all cells from uninfected spheroids were viable after seven days (data not shown). The viability of uninfected spheroids was also confirmed by letting individualized spheroid cells proliferate as monolayers again (data not shown). These results show that even though a virus may cause cytolysis in a certain cell population when grown as a monolayer, the same virus may not be as infectious when the cells are grown in three-dimensional clusters.

A major concern regarding all oncolytic viruses is the existence of neutralizing antibodies from prior natural or deliberate exposure. Employing low- or apathogenic viruses for oncolytic purposes raises the question whether these viruses are frequently circulating in the human population without causing any symptoms although inducing immunity. Because they are not frequently associated with disease there are very little data published regarding circulation of these viruses in the population. The seroprevalence of the oncolytic candidates we present here needs to be addressed in coming studies. However, seroprevalence of E1 and coxsackievirus A21, two other low-pathogenic oncolytic enteroviruses, seems to be less than 10% in preliminary studies [16,48]. Besides investigation of the potential risk of preexisting neutralizing antibodies, this study will need to be complemented with infection studies in primary cell cultures and animal models. This would further elucidate the oncolytic capacity of these viruses. Knowledge of infection preferences, for example receptor usage, of these viruses would also be a great advantage when investigating the ability to utilize these viruses for
oncolytic purposes. E12 and E29 have been shown to utilize DAF (CD55) as cellular receptor [44,45,49]. DAF is a vital part of directing the immune system away from self-tissue [50], and has been shown to be upregulated in colon cancer [51,52]. In addition to in vivo studies of the oncolytic capacity, the receptor usage of these viruses needs to be further elucidated while bearing in mind that enteroviruses may utilize more than one receptor [46,53]. Our study pinpoints E12, E15 and E26 as new potential candidates for oncolytic virotherapy. It also puts focus on a group of entroviral serotypes that have not been extensively studied before, due to the fact that they are not causing serious diseases or financial losses to the human population. However, seen in the light of oncolytic virotherapy, these viruses may be of great interest as new treatments in the battle against cancer.

Conclusions
In conclusion, the echoviruses 12, 17 and 26 show potential as oncolytic agents against colon cancer. This was found by investigating the cytolytic capacity and replicative efficiency of E12, E15, E17, E26 and E29 in six colon cancer cell lines as well as in artificial HT29 tumors in vitro.

Additional material

Additional file 1: Pictures of infected colon cancer cell lines CaCo-2, HT29, LoVo, SW480, SW620 and T84 infected with echovirus 12, 15, 17, 26, 29 (at a multiplicity of infection of 1) and uninfected (Control). Pictures were taken at the time of complete cytopathic effect or seven days post infection at 4 000 or 10 000 x magnification.

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Author details
1School of Natural Sciences, Linnaeus University, SE-391 82 Kalmar, Sweden.
2Current address: National Veterinary Institute, Division of Virology, Technical University of Denmark, Lindholm, 4771 Kalvehave, Denmark.

Authors’ contributions
SI planned the experimental setup, prepared viral stocks, carried out viral infectivity assays and cultured and infected spheroids. SI wrote the manuscript. NJ participated in preparation of virus stocks and viral infectivity assays. MG participated in titrating viral stocks. AML was involved in the study design and revision of the manuscript. All authors have read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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