Three Poliovirus Sequences in the Human Genome Associated With Colorectal Cancer

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Abstract. Background/Aim: In a previous study we analyzed poliomyelitis incidence in US states, before the introduction of the polio vaccine. We noted an inverse correlation between polio incidence in US states, between 1937 and 1951, and colorectal cancer incidence (2005-2009). To further elucidate the role that poliovirus could play in colorectal cancer, the full human genome for poliovirus sequences was analyzed using the UCSC Genome Browser. Materials and Methods: BLAT, the Blast-Like Alignment Tool of the UCSC Genome Browser, was used to compare the poliovirus genome to the human genome. Results: BLAT revealed three poliovirus sequences in three chromosomes: Chromosome 20p12.1 (MACROD2 gene), chromosome 1p13.3 (SLC25A24 gene), and chromosome 2p25.1. Conclusion: Poliovirus sequences in the human genome may contribute to programmed cell death and lysis of colorectal cancer cells. A recombinant poliovirus, incapable of reverting to neurovirulence, might be given orally at intervals as a colorectal cancer vaccine to prevent colorectal cancer.

Colorectal cancer rates have dropped steadily for people born between 1890 and 1950, but have been increasing for every generation born since 1950. Adults born in 1990 have twice the risk of colon cancer and four times the risk of rectal cancer compared to adults born in 1950 of comparable age. A total of 16,450 new cases of colon or rectal cancers will be diagnosed this year in Americans under 50, according to the American Cancer Society. In 2014, 43 percent of colorectal cancer cases in those under 50 were in individuals aged 45 to 49, with a rise in colorectal cancers in adults as young as their 20s and 30s (1).

No one knows what may be driving the rising number of cases in young people. Lifestyle, environmental or genetic factors, obesity, cigarette smoking, a diet high in red or processed meats, and lack of physical activity have been tied to increased risk.

In a previous study, we analyzed poliomyelitis incidence in US states, before the introduction of the polio vaccine. We found that polio might protect against colorectal cancer. Polio is predominantly an enteric viral infection that was progressively eradicated in the United States after the introduction of polio vaccine in the early 1950s. We noted an inverse correlation between polio incidence in US states, 1937-1951, and colorectal cancer incidence (2005-2009). The same inverse correlation pertained for colorectal cancer deaths (2).

Oral polio vaccine was given in the US until 2000 when a switch was made to injectable form only, which is still used. The reason for the switch was reversion of the oral vaccine to virulence and vaccine-induced polio. Very few people now in their 20s and 30s have been vaccinated with the oral vaccine, which produces enteric inflammation comparable to wild-type polio virus and could protect against colorectal cancer. The majority of people have been vaccinated with the injectable form, which produces no intestinal inflammation at all and does not protect against colorectal cancer. Even when the oral vaccine was used, between 1960 and 2000, the coverage was 80% (3). So 20% of the population born in this period would be vulnerable to early colorectal cancer.

To further elucidate the role that poliovirus might play in colorectal cancer, the full human genome was analyzed for poliovirus sequences with the UCSC Genome Browser.

Materials and Methods

The UCSC Genome Browser is an on-line genome browser of the University of California, Santa Cruz (UCSC). The browser is an interactive website offering access to genome sequence data from a variety of vertebrate and invertebrate species and major model organisms, integrated with a large collection of aligned annotations.
The graphical view is optimized to support fast interactive performance and is an open-source, web-based tool suite build on top of a MySQL database for rapid visualization, examination, and querying of data at many levels. The Genome Browser Database, browsing tools, downloadable data files, and documentation are all accessible on the UCSC Genome Bioinformatics website (https://genome.ucsc.edu) (4).

To compare the poliovirus genome to the human genome, we used BLAT, the Blast-Like Alignment Tool of the UCSC Genome Browser (4). BLAT can align a user sequence of 25 bases or more to the genome. Because some level of mismatch is tolerated, cross-species alignments may be performed provided the species have not diverged too far from each other; this capability allowed comparison of the poliovirus genome to the human genome. The poliovirus sequence we analyzed with BLAT was FASTA Poliovirus, complete genome NCBI Reference Sequence: NC_002058.3.

Because of the immunologic basis for colorectal cancer (5), the GISTIC (Genomic Identification of Significant Targets in Cancer) module in cBioportal was used to access data in The Cancer Genome Atlas (TCGA). The GISTIC module identifies regions of the genome that are significantly amplified or deleted across a set of samples. cBioPortal for Cancer Genomics provides visualization, analysis and download of large-scale cancer genomics data sets. The Cancer Genome Atlas is a project that began in 2005 in order to catalogue genetic mutations responsible for cancer, using genome sequencing and bioinformatics.

Results

BLAT revealed three poliovirus sequences in three chromosomes:

- In Chromosome 20p12.1, a poliovirus sequence, with almost 100% identity, consisting of a sense and an antisense strand of 32 nucleotides (6380 to 6411), which codes for the 3Dpol, an RNA dependent RNA polymerase whose function is to make multiple copies of the viral RNA genome. The viral sequence is in an intronic region of the MACROD2 gene, indicated by the dashed lines of Genode. Red line: Genome and poliovirus sequence have different bases at this position. Green line: The poliovirus sequence appears to have a polyA tail that is not aligned to the genome (https://genome.ucsc.edu).

The graphical viewer is optimized to support fast interactive performance and is an open-source, web-based tool suite build on top of a MySQL database for rapid visualization, examination, and querying of data at many levels. The Genome Browser Database, browsing tools, downloadable data files, and documentation are all accessible on the UCSC Genome Bioinformatics website (https://genome.ucsc.edu) (4).

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colon tissue (Figure 2). A similar viral sequence was present in humans and rhesus monkeys.

- In Chromosome 1p13.3, a poliovirus sequence with 89.3% identity, consisting of a sense and an antisense strand of 28 nucleotides, 811 through 838 of the virus, which codes for the VP4 protein of the viral capsid (Figure 3) (7). The sequence is within an intronic region of the solute carrier family 25 member 24 gene (SLC25A24). Increased RNA expression of the sequence was associated with significantly increased survival of colon cancer patients in TCGA (Figure 4).

- In Chromosome 2p25.1, a poliovirus sequence with almost 100% identity, consisting of two sense strands of 20 nucleotides, 7106 through 7125 of the virus, which codes for the 3Dpol. A similar viral sequence was detected in humans and rhesus monkeys. (Figure 5).

With cBioportal, the COAD cohort (adenocarcinoma of the colon) was analyzed, and the relationship of PV R, the poliovirus cellular receptor, to FCAR, a protein coding gene that encodes a receptor for the Fc region of Immunoglobulin A, was examined. A significant correlation in copy number alterations of both genes was found (Figure 6).

As mentioned above, we noted a significant inverse correlation between polio incidence, 1937-1951 and colorectal cancer incidence (2005-2009). The same inverse correlation pertained for colorectal cancer deaths (Figure 7) (2).

To exclude the effect of the quality of reporting by the States (good for some States, mediocre or poor for others), the multivariate general linear model (GLM) of SPSS was used (8). Based on the Annual Supplement Reported Incidence of Notifiable Diseases in the United States, 1960 (9), the relationship of scarlet fever/streptococcal sore throat to male and female colorectal cancer deaths (2005-2009) was analyzed. Any relationship of scarlet fever/streptococcal sore throat to colorectal cancer should be a chance one brought about by variations in State reporting. The relationship of polio incidence to male colorectal cancer deaths ($p<0.0005$) and female colorectal cancer deaths ($p=0.001$) was significant in the GLM. The relationship of scarlet fever/streptococcal sore throat to male colorectal cancer deaths ($p=0.395$) and female colorectal cancer deaths ($p=0.316$) was not significant.

**Discussion**

Eight percent of human DNA comes from human endogenous retroviruses (HERV), and some human diseases have been attributed to this DNA. HERV sequences have occasionally been adopted to serve a useful purpose, such as in the placenta, where they may safeguard fetal-maternal tolerance (10). Some HERV sequences have oncogenic properties; yet treating cancer cells with methyltransferase inhibitors increases HERV RNA and DNA. Like infectious agents, the HERV-derived nucleic acids in the cytoplasm activate innate immune responses that produce tumor cell apoptosis (11, 12).

Expression of the human endogenous retrovirus (HERV)-H family is associated with colorectal carcinomas. One study identified colorectal cancer specific HERV-H sequences in all stages, along with correlations between HERV-H expression, lymph node involvement, and microsatellite instability (13).

Poliovirus is a human enterovirus and member of the *Picornaviridae* family. Human rhinovirus type 2 (HRV2) is another member of this family. Poliovirus infects human cells by binding to an immunoglobulin-like receptor, CD155, also known as the poliovirus receptor (PVR), on the cell surface. Interaction of poliovirus and CD155 facilitates an irreversible conformational change of the viral particle necessary for viral entry. The 5' end of poliovirus RNA is extremely long – over 700 nucleotides – and highly structured. This region of the viral genome is called internal ribosome entry site (IRES), and directs translation of the viral RNA. Genetic mutations in IRES prevent viral protein synthesis (14).

The three HERV poliovirus sequences that were identified in this study in the human genome on chromosomes 20p12.1, 1p13.3 and 2p25.1 relate to colorectal cancer as follows:

- The MACRO Domain Containing 2 (MACROD2) gene on chromosome 20p12.1 has chromosome breaks in 41% of colorectal cancers (15). The gene codes for a deacetylase that removes ADP ribose from mono-ADP ribosylated proteins. The encoded protein translocates from the nucleus to the cytoplasm upon DNA damage. MACROD2 expression predicts response to 5-FU-based chemotherapy in stage III
colon cancer (16). The poly A tail of the poliovirus is not aligned to the genome. A 3’ poly(A) tail is a common feature of picornavirus RNA genomes and the RNA genomes of many other positive-strand RNA viruses (17).

- The gene SLC25A24 on chromosome 1p13.3 is part of the SLC family. Proteins in the SLC family transport various compounds across cell membranes and component parts. The protein produced by the SLC25A24 gene carries molecules across the inner membrane of the mitochondria. The protein is an ATP-Mg/Pi carrier, transporting ATP molecules that are bound to magnesium (Mg) through the mitochondrial inner membrane in exchange for adding or removing phosphate atoms from the mitochondria. The exchange is essential for normal energy production, the metabolism of various molecules, and protein production within cells. SLC25A24 is involved in carcinogenesis (18) and the Fontaine progeroid syndrome (19).

- The lipopolysaccharide binding protein 32 (LBP-32) gene on chromosome 2p25.1 is up-regulated by a factor of 2.57 in colorectal cancer (20). This gene encodes a transcription factor that functions during development.

It is not surprising that poliovirus might reduce the risk of colorectal cancer. A genetically modified poliovirus, PVRSRIPO, infused directly into the brain, significantly improves long-term survival in patients with recurrent glioblastoma (21). PVRSRIPO is a recombinant, live, attenuated, nonpathogenic form of the oral poliovirus Sabin type 1, in which the IRES has been replaced with the IRES from human rhinovirus type 2 (HRV2). Because of the replacement, PVRSRIPO cannot revert to virulence, as can Sabin type 1 virus. Most tumor cells express the PVR gene and its protein CD155, which allows tumor cells to selectively take up polio virus. The virus then replicates and produces tumor cell lysis, with an end result comparable to apoptosis (programmed cell death).
Induction of apoptosis in malignant cells is a major goal of cancer therapy. Numerous apoptosis-regulating genes have been identified (22). The results of this analysis suggest that poliovirus sequences in the human genome may contribute to programmed cell death of colorectal cancer cells. In combination with an enteric poliovirus to prime the immune system, the risk of colorectal cancer may be reduced in susceptible individuals. In the case of the MACROD2 gene, enteric poliovirus might insert sequences into chromosome breaks, making colorectal tumor cells more vulnerable to apoptosis and tumor cell lysis.

A recombinant poliovirus, incapable of reverting to neurovirulence, might be given orally at intervals as a colorectal cancer vaccine to prevent colorectal cancer. Further study would be worthwhile.

**Conflicts of Interest**

None of the Authors have any conflicts of interest to disclose regarding this study.

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