No evidence for integrated viral DNA in the genome sequence of cutaneous squamous cell carcinoma

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To the Editors:

Cutaneous squamous cell carcinoma (cSCC) is the second most common type of non-melanoma skin cancer and is dramatically increased in the setting of immunosuppression such as organ transplantation (Grulich et al., 2007; Hartveilt et al., 1990; Jensen et al., 1999; Lindelof et al., 2000). This magnitude of increase is similar to that seen in virus-related cancers, such as human herpesvirus-8 mediated Kaposi’s sarcoma and Epstein-Barr virus related lymphoma (Vajdic et al., 2006). There have been multiple attempts to define a viral etiology for cSCC and to link cSCC with human papillomavirus (HPV). There is a clear epidemiologic association between HPV and cSCC (Aldabagh et al., 2012), but detection has been limited to viral DNA. Two studies have demonstrated the absence of viral transcription in cSCC RNAseq data, confirming that direct viral transformation of keratinocytes is not required for maintenance of tumor (Arron et al., 2011; Ganzenmueller et al., 2012). These data suggest that, if there is a viral pathogen causing cSCC, the mechanism of carcinogenesis is not simply through expression of viral oncoproteins.

This raises the possibility of alternate mechanisms of oncogenesis, including insertional mutagenesis, in which viral integration into the genome disrupts human genes of cell cycle regulation (Leib-Mosch et al., 1990). This would be unlikely for a papillomavirus, but high-risk α-papillomaviridae do aberrantly integrate as part of cervical oncogenesis (Wentzensen et al., 2004), and it is possible that cutaneous β-papillomaviridae do the same. Alternately, DNA from a previously undetected virus may be integrated into the genome or persist as extragenomic DNA. We used a newly published software package, Integrated Metagenomic Sequence Analysis (IMSA) (Dimon et al., 2013), to address the hypothesis that HPV or another virus may be integrated into the cSCC genome or otherwise present in cancer DNA.
Exome sequence from 12 cSCC and patient-matched normal skin were obtained from a previously reported study (Durinck et al., 2011). Exome sequence from the MCF7 breast cancer cell line was used as a negative control. Previously described WGS data from SiHa and CaSki cell lines were used as positive controls for viral analysis (Conway et al., 2012).

Pathogen analysis was performed with IMSA (Dimon et al., 2013). Low quality and human reads were filtered against the human genome (hg19) and non-human reads were aligned to the NCBI nucleotide database. An average of 9.4x10^7 reads were obtained from each sample; 2.4x10^6 non-human reads remained after IMSA filtering (0.25%). Non-human and intronic reads in exome datasets result from incomplete capture efficiency. In order to confirm our ability to detect potential pathogen reads in exome datasets, we measured the percent of aligned reads on target using Picard. Picard is a set of standard bioinformatics software tools for high throughput sequence analysis. The tools are open source and can be found at [http://picard.sourceforge.net/](http://picard.sourceforge.net/). The specific tool that we used was CalculateHsMetrics, which is designed to measure the efficiency of targeted sequencing such as exome sequencing. Capture efficiencies ranged between 66.1% and 78.1%. On average 27.3% of the reads in each dataset were non-exome. These libraries were sequenced to great depth for the purpose of cancer mutation analysis, therefore ample sequence was available for metagenomics.

IMSA generates a quantitative score for each taxonomic node, reflective of the level of nonhost DNA. To compare between datasets, we normalized the taxonomy score per million reads. cSCC had more bacterial reads on average than normal skin (65% vs 33% of nonhuman reads, Figure 1). 16% of cSCC and 30% of normal nonhuman reads mapped to vertebrates, most commonly nonhuman primate. These likely represent human DNA with sufficient divergence or sequencing errors to pass through the filters.

1% or fewer reads mapped to virus (Figure 2). The median normalized score for cSCC samples was 0.26 (IQR 0.1-0.7), no different from normal tissue (0.24, IQR 0.06-0.9, paired t-test p=0.3). In contrast SiHa had a score of 6.25 and CaSki had a score of 1272.6, derived exclusively from HPV16 reads.

One exome set contained significantly higher viral read scores, 6.2 in tumor and 36.3 in normal skin. These reads were derived from torque teno virus (TTV). TTV is a ubiquitous 3.8Kb circular single stranded DNA virus that has not been associated with disease. TTV viremia reflects immune function (Moen et al., 2003), and this tumor-normal pair was from an immunosuppressed lung transplant recipient. Two of these reads indicated integration into the human genome.

To maximize the probability of detecting pathogen, we also examined WGS from three tumor-normal pairs. Whole genome sequence (WGS) libraries were prepared with the NuGEN Ovation Ultralow Library System and sequenced with the Illumina sequencing-by-synthesis platform. 3x10^8 reads were obtained from each WGS readset; with over 2x10^6 non-human reads remaining after IMSA filtering. The tumor-normal pairs had normalized viral read scores within the background range; 0.04/0.06, 0.24/0.08, and 0.01/0.05, (p=0.9, 95% CI -0.6-0.6) (Figure 2).
These data demonstrate the absence of viral DNA, including HPV DNA, in the exome and WGS of cSCC. This is consistent with prior qPCR data demonstrating extraordinarily low HPV DNA copy number in cSCC (typically less than 1 copy/100 cells) (Arron et al., 2011) and with the newly developing consensus that viral infection is not required for tumor maintenance. In contrast, viral DNA was readily detected in two cervical cancer cell lines, CaSki and SiHa.

Despite these findings, the consistent association of HPV DNA with cSCC remains unclear. One remaining possibility is a hit and run mechanism, by which the virus destabilizes the keratinocyte until sufficient UV-mediated DNA mutations accumulate for transformation. This proposed role for viral tumor induction would be compatible with existing sequencing data, which has been limited to invasive tumor tissue.

It is possible that virus plays no role in cSCC but is rather a marker of cutaneous immunosuppression. This would explain the predilection of HPV for sun-exposed skin and the increased prevalence of HPV in the skin of immunosuppressed patients (Aldabagh et al., 2012). Our finding of TTV in the tumor and adjacent normal skin of a lung transplant recipient suggests that immunosuppression predisposes to unrelated viral carriage.

It remains imperative to determine whether cSCC is a viral cancer, as this would have major implications for patient risk assessment and screening. Identifying a causative virus would open avenues for improved detection, targeted molecular therapies, and a potential vaccine for cancer prophylaxis. Additional studies may continue to shed light on this controversial and important subject.

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Abbreviations

| Abbreviation | Description                        |
|--------------|------------------------------------|
| cSCC         | cutaneous squamous cell carcinoma  |
| HPV          | human papillomavirus               |
| IMSA         | integrated metagenomic sequence analysis |
| TTV          | Torque teno virus                  |
| WGS          | Whole genome sequencing             |

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Figure 1.
Taxonomic classification of nonhuman reads in cSCC and normal skin. The percents shown are rounded to the nearest integer. In viral samples, the average percent viral reads is 0.145% while in normal skin the average percent viral reads is 1.139%.
Figure 2.
Viral sequence scores per million sequence reads.