

Method

A virome-wide clonal integration analysis platform for discovering cancer viral etiology

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Oncoviral infection is responsible for 12%–15% of cancer in humans. Convergent evidence from epidemiology, pathology, and oncology suggests that new viral etiologies for cancers remain to be discovered. Oncoviral profiles can be obtained from cancer genome sequencing data; however, widespread viral sequence contamination and noncausal viruses complicate the process of identifying genuine oncoviruses. Here, we propose a novel strategy to address these challenges by performing virome-wide screening of early-stage clonal viral integrations. To implement this strategy, we developed Vcaller, a novel platform for identifying viral integrations that are derived from any characterized viruses and shared by a large proportion of tumor cells using whole-genome sequencing (WGS) data. The sensitivity and precision were confirmed with simulated and benchmark cancer data sets. By applying this platform to cancer WGS data sets with proven or speculated viral etiology, we newly identified or confirmed clonal integrations of hepatitis B virus (HBV), human papillomavirus (HPV), Epstein-Barr virus (EBV), and BK Virus (BKV), suggesting the involvement of these viruses in early stages of tumorigenesis in affected tumors, such as HBV in *TERT* and *KMT2B* (also known as *MLL4*) gene loci in liver cancer, HPV and BKV in bladder cancer, and EBV in non-Hodgkin's lymphoma. We also showed the capacity of Vcaller to identify integrations from some uncharacterized viruses. This is the first study to systematically investigate the strategy and method of virome-wide screening of clonal integrations to identify oncoviruses. Searching clonal viral integrations with our platform has the capacity to identify virus-caused cancers and discover cancer viral etiologies.

[Supplemental material is available for this article.]

Oncoviral infections are responsible for some human cancers (zur Hausen 1991). To date, there are seven well-accepted human oncoviruses, including Epstein-Barr virus (EBV), human T-lymphotropic virus type 1 (HTLV-1), hepatitis B virus (HBV), human papillomavirus (HPV), hepatitis C virus (HCV), Kaposi's sarcoma-associated herpesvirus (KSHV, also known as human herpesvirus 8 or HHV8), and Merkel cell polyomavirus (MCV or MCPyV) (Moore and Chang 2010). These are collectively responsible for 12%–15% of human cancers worldwide (zur Hausen 1991; Bouvard et al. 2009). Additional oncoviruses and virus-caused cancer types have been suggested by convergent evidence from epidemiological, pathological, and oncological studies (Javier and Butel 2008; zur Hausen 2009b; Moore and Chang 2010). An increasing number of cancer types have been newly found to have etiologies partially attributable to known oncoviruses such as HPV and EBV (Young and Rickinson 2004; Woodman et al. 2007) or potentially new oncoviruses. In the past, the discovery of new oncoviruses has led to novel preventative approaches, such as vaccination for HBV and HPV. These approaches have reduced the burden of related cancers, probably more than any single therapeutic treatment (Andre et al. 2008). Thus, it is crucial to uncover new cancer viral etiologies.

The list of known oncoviruses is short, partially due to challenges in identifying them. Virus-caused cancers usually have a long latency after the initial infection, from years to decades, and do not follow the principles of causality (Moore and Chang 2010). Among the seven well-established oncoviruses, three of them (HPV, KSHV, and MCV) were identified by directed searches for viral DNA sequences in tumor tissues (Cao and Li 2018), highlighting the historic successes of nucleotide sequence analysis strategies. In the era of high-throughput sequencing (HTS), the genomes of thousands of tumors have been sequenced, providing exceptional opportunities for discovering new oncoviruses and new cancer types associated with known oncoviruses. Theoretically, one or more reads containing a portion of nucleic acid sequence uniquely mapping to a viral genome can indicate the presence of a virus in a sample. However, two major challenges have prevented previous efforts from successfully identifying new cancer-associated viruses using HTS data: First, contaminating viral sequences are common in HTS data (Moustafa et al. 2017). Viral contaminations here refer to viral sequences that are not a result of infection, but are introduced during sample collection, preparation, and/or

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sequencing procedures. They frequently arise from the introduction of environmental microbes, human resources, synthetic DNA (e.g., vectors), or common cell lines (Jun et al. 2012; Laurence et al. 2014; Salter et al. 2014; Strong et al. 2014; Cantalupo et al. 2015; Friis-Nielsen et al. 2016), as indicated by batch patterns (Laurence et al. 2014), such as sequencing facility-associated virus patterns (Tae et al. 2014). Second, viral infection also may have arisen from an infection occurring after tumor formation, and thus have no contribution to tumorigenesis. The presence alone of viral sequence in tumor genome sequencing data is insufficient to prove an oncogenic role, especially for viruses widely present in the population such as EBV and human herpesvirus 6.

Viral integration is the process by which a virus inserts its DNA or cDNA into its host cell's genome. Integration is a required stage in the life cycle of retroviruses (Goff 1992). Integration may also occur with non-retroviruses via mechanisms such as homologous recombination. The majority of virus-caused human tumors, including most of the tumors caused by HBV, HPV, HTLV-1, and MCV, carry multiple viral integration events in their genomes (Feng et al. 2008; Sung et al. 2012; Hu et al. 2015; Kataoka et al. 2015; Xiao et al. 2016). Thus, identifying viral integrations provides a unique opportunity to overcome the two major challenges in discovering new oncovirus candidates from HTS data: First, identifying integrated viral sequences eliminates the misidentification of contaminating viral sequences since human-virus chimeric sequences flanking integration sites, rather than virus-only sequences, are unlikely to be derived from random viral sequence contamination. Viral integrations derived from cell line contaminations can also be eliminated based on the identical viral integrations already found in cell lines, for example, known HPV-18 integrations in HeLa cells (Cantalupo et al. 2015). Second, by further analyzing the percentage of cells with identical integrations (cellular proportion), we can determine clonal integrations that occurred during the early stages of tumorigenesis. As with somatic mutations, viruses that lead to high cellular proportion "early stage" clonal integrations are potential oncogenic drivers. Some of the integrations, for example, those disrupting tumor suppressor genes, may have direct oncogenic roles. Thus, as a functional

consequence, these integrations may confer selective growth advantages to the cells, leading to increased cellular proportion of these integrations. The presence of high cellular proportion clonal integrations suggests an oncogenic role of the identified virus. Indeed, clonal integration was considered "the strongest evidence" when determining the oncogenic role of MCV in Merkel cell carcinoma (Moore and Chang 2010). We recently compared different methods, including genetic, molecular, and epidemiologic methods, and concluded that identifying clonal integrations using HTS was capable of providing strong genetic evidence for discovering viral etiologies (Cao and Li 2018; Chen et al. 2019; Sulovari and Li 2019). By restricting further analysis exclusively to viruses with clonal integrations, most noncausal viruses can be eliminated in a systematic search for oncoviruses. Therefore, analyzing clonal viral integrations in HTS data is an effective strategy for identifying new oncoviruses and new cancer types associated with known oncoviruses.

Results

Vicaller: a novel bioinformatics tool for virome-wide integration calling

To implement our strategy, we developed a novel bioinformatics platform, *Viral Integration caller* (Vicaller), for detecting virome-wide viral sequences, integration events, and fusion transcripts from HTS data. Specifically, Vicaller determines the cellular proportion for each integration to identify viruses with clonal integrations in the tumor genome as oncovirus candidates. It overcomes the two major challenges for identifying oncoviruses from tumor HTS data because (1) contaminating viral sequences are unlikely to integrate because they are not present in the live cells, and (2) noncausal viruses may integrate; however, integration events derived from noncausal viruses should be independent and sporadic, and thus are unlikely to be clonal. Only viruses that are involved in the early stages of tumorigenesis have the capacity to form clonal viral integrations, and Vicaller is capable of distinguishing them (Fig. 1A). We first generated a comprehensive viral

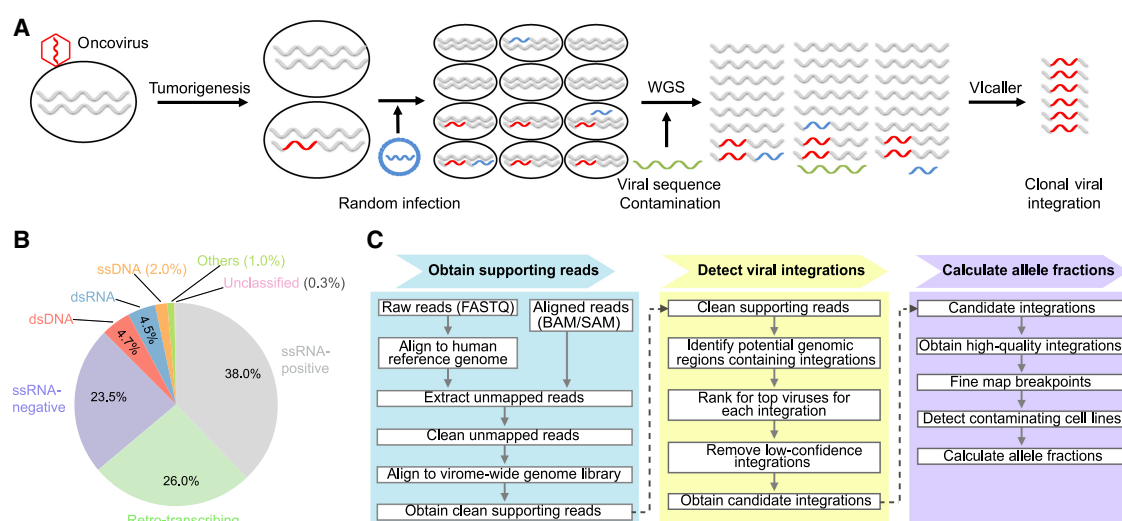


Figure 1. Discovering oncovirus candidates through identification of clonal viral integrations. (A) Identification of clonal viral integrations to eliminate viral sequence contaminations and to prove the involvement of the identified virus in the early stages of tumorigenesis. (B) Composition of Vicaller virome-wide genome reference library. (C) The simplified analytic workflow of Vicaller.

genome reference library containing 411,195 unique whole and partial viral genomes, representing 10,662 distinct viruses (Supplemental Table S1) and covering all six virus taxonomic classes, including dsDNA, ssDNA, dsRNA, ssRNA (–), ssRNA (+), and retroviruses (Fig. 1B). This library incorporates all currently characterized viral genomes and represents the largest existing viral reference genome library.

The Vcaller pipeline consists of three phases: (1) obtaining supporting reads, (2) detecting viral integrations, and (3) calculating integration allele fractions (Fig. 1C; Supplemental Table S2). Three types of supporting reads were identified and used for determining integrated versus nonintegrated viral sequences: (1) viral reads that can be paired-end mapped to a viral genome, (2) chimeric reads, and (3) split reads, the latter two of which can be mapped to both viral and human genomes (for terms and abbreviations, see Supplemental Table S3). Briefly, we first obtained the reads which could not be fully mapped to the human reference genome (unmapped reads), which included paired-end unmapped reads, one-end unmapped reads (one-end mapped reads are also extracted), and soft-clipped sequences. We then aligned these reads to the viral reference genome library to obtain all potential viral, chimeric, and split reads. After excluding reads that aligned to homologs of the human genome or contained primarily repeat sequences, we used the chimeric and split reads to determine viral species, integration status, and the upstream and downstream breakpoints on both the viral and human reference genomes (Supplemental Fig. S1). The integration breakpoints were fine-mapped at nucleotide resolution where possible. Each identified integration was then compared to a list of viral integrations or fusion transcripts present in commonly used cell lines (Supplemental Table S4; Peter et al. 2006; Klijn et al. 2014; Cao et al. 2015; Liu et al. 2016), such as HPV-18 integrations in HeLa cells and HPV-16 integrations in SiHa cells. Any events identical to (or within 10,000 bp of) the known integrations identified in these cell lines were annotated for further evaluation of potential cell line contaminations. The final viral integration events were summarized and reported in a uniform format (Supplemental Table S5). Last, the cellular proportion of each viral integration event was calculated based on the number of reads that supported integration and nonintegration, that is, the integration allele fraction (Supplemental Fig. S2). A 50% integration allele fraction represents 100% cellular proportion in diploid cells. Our comprehensive strategy considers all 12 possible scenarios of supporting reads (Supplemental Fig. S3) and includes all eight possible types of integration events (Supplemental Fig. S4), which increases the chances of discovering integrations.

Identifying viral integrations in simulated data

To evaluate the accuracy of our platform, we applied Vcaller to a series of simulated data sets. We randomly selected and fragmented more than 5000 viral sequences of various lengths from the

virome-wide reference genome library and inserted them randomly into the human genome (Supplemental Fig. S5). An average of 3700 substitution errors per megabase were introduced in all simulated sequence data sets. A series of viral integration-carrying HTS data sets were then simulated with read depths ranging from 1× to 150× (Supplemental Table S6). Vcaller was applied to each data set, and the sensitivity was calculated based on the ratio of correctly identified versus total simulated integration events. The average numbers of chimeric and split reads for these viral integrations are shown in Supplemental Table S7. We found that 5× sequencing depth was sufficient to detect >90% of the simulated viral integrations (Fig. 2A). When the sequencing depth was increased to 20× and 100×, Vcaller was able to capture 95% and 98% of the integrations, respectively (Fig. 2A).

To mimic tumor purity and cancer cell heterogeneity, we varied the integration allele fractions by mixing the human genomes carrying integrations and those with no integrations at different ratios, for example, 5%, 25%, and 50% for integration-bearing genomes. A series of HTS data sets carrying somatic viral integrations with different allele fractions were then simulated with read depths from 5× to 60× (Supplemental Table S6). At 30× depth, Vcaller was able to correctly detect 80% of the simulated viral integrations when the integration allele fraction was as low as 5%. Integration events with even lower abundance in tumor tissues likely do not support viral involvement in tumorigenesis. When the depth was increased to 60×, nearly all (>98%) integrations with allele fractions >25%, or >90% of integrations with allele fractions as low as 5% could be identified (Fig. 2B; Supplemental Table S8). Precision was defined as the ratio of correctly identified to total identified integrations. In most analyses of the simulated data sets, the precision remained nearly 100%, that is, only four false

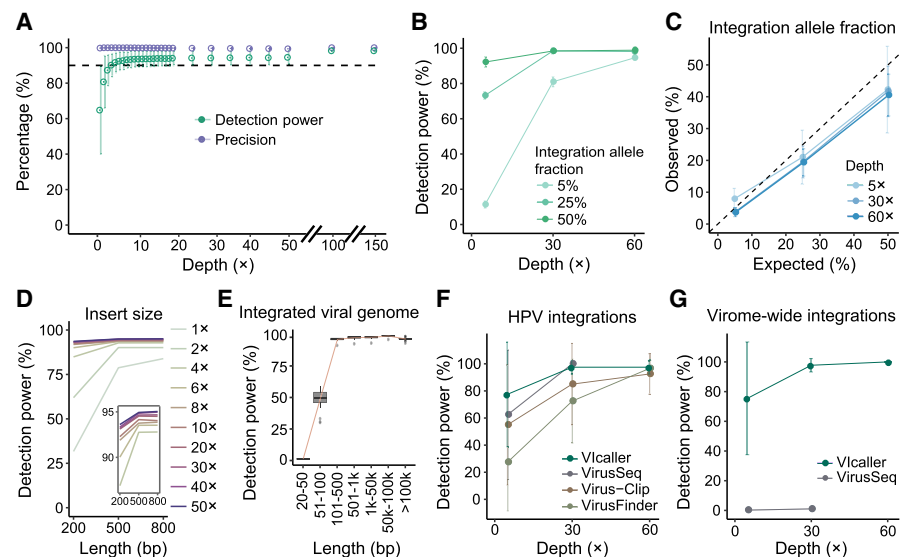
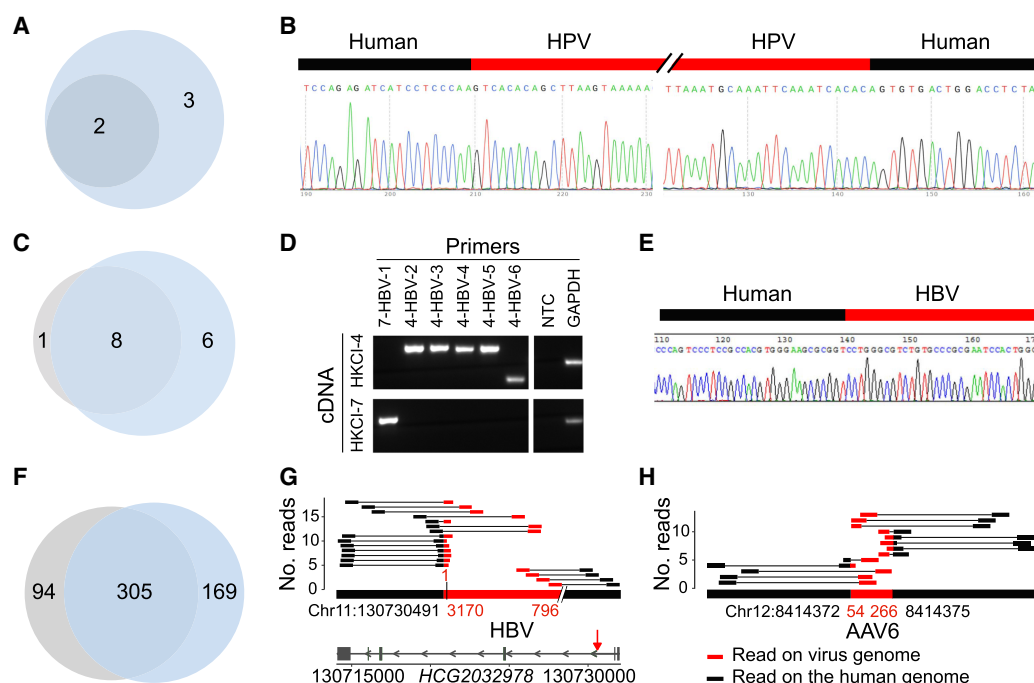


Figure 2. Applying Vcaller to simulation data sets. (A) Detection power and precision were measured by simulated (germline) viral integrations with depths from 1× to 150×. (B) Detection power for integrations with 5%, 25%, and 50% integration allele fractions. On average, 86 viral integrations were used for the calculation. (C) Accuracy of calculated integration allele fractions. (D) Relationship between detection power and insert sizes for paired-end sequence reads at different sequencing depths. (E) Relationship between detection power and lengths of integrated viral sequences. The viral integrations detected under different sequencing depths were combined for the calculation. Comparison of the detection power of Vcaller with existing tools for detecting 10 simulated HPV integrations (F) and 90 simulated virome-wide integrations (G). VirusSeq was only capable of detecting less than 20 human viruses; thus, the detection power was extremely low. It also ran out of server wall time at 60× sequencing depth. VirusFinder and Virus-Clip were not applicable for analyzing data containing the virome-wide integrations.

We also evaluated other factors that may influence the detection power and precision for identifying integration events using HTS data. Overall, a longer insert size for paired-end reads significantly increased the detection power, particularly when it was longer than 500 bp (Fig. 2D). Longer integrated viral sequences also increased the detection power, particularly when the integrated viral sequences were shorter than the insert size (Fig. 2E). When the integrated viral sequences were longer than 1000 bp, the detection power was consistently higher than 98% (Fig. 2E). We further divided the simulated data sets into subgroups by the size of the viral reference genomes and then compared the detection power. Vcaller showed no bias regarding the length of the viral reference genomes (Supplemental Fig. S6). In all cases, the detection power can be enhanced by increasing sequencing depth. Concurrent with the above analyses, we also tested simulated negative controls, that is, a whole-genome sequencing (WGS) data set with no integrations (Supplemental Table S6), and Vcaller was then ap-

We further compared Vcaller to three recently developed tools for detecting viral integrations (Chen et al. 2013; Ho et al. 2015; Wang et al. 2015). We simulated a series of data sets harboring 10 HPV-16 integrations and a series of data sets harboring 90 virome-wide viral integrations, all having varying sequencing depths and integration allele fractions (Supplemental Table S6). Vcaller achieved the highest detection power in identifying the 10 HPV-16 integrations under all conditions (Fig. 2F; Supplemental Table S10). For detecting the virome-wide viral integrations, none of the tools except Vcaller could be used (Fig. 2G; Supplemental Table S10).

To evaluate our Vcaller approach in identifying viral integrations from real HTS data, we first applied it to WGS data of tumor and paired normal tissues from one cervical carcinoma patient. We identified both HPV-18 integration events detected by the HPV-specific approach used in our previous study (Liang et al. 2014). Additionally, we detected three new HPV-18 integration events that were not originally detected (Fig. 3A; Supplemental Tables

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S11A, S12; Liang et al. 2014). All five integrations are present in the tumor, but not the paired normal tissue. All the newly identified integrations were validated via PCR and subsequent Sanger sequencing (Fig. 3B).

Identifying HBV integrations in liver cancer data sets

We further applied Vcaller to an RNA-seq data set of hepatocellular carcinoma (HCC) cell lines (Lau et al. 2014). Human HBV commonly integrates into the genomes of liver tumor cells (Bréchet et al. 2000). When inserted into genic regions, the HBV genes may be expressed as host–virus fusion transcripts and contribute to tumorigenesis. By performing virome-wide analyses of three cell lines, we detected eight of the nine fusion transcripts reported by the HBV-specific approach used in the original study (Fig. 3C; Lau et al. 2014). In addition, we identified six new fusion transcripts that were not detected by the previous method (Supplemental Table S11B; Lau et al. 2014). All six newly identified fusion transcripts were validated via reverse transcription PCR (RT-PCR) (Fig. 3D) followed by Sanger sequencing (Fig. 3E).

We then applied Vcaller to a larger WGS data set derived from 99 HBV-associated HCC patients (Sung et al. 2012). We detected a total of 474 integration breakpoints (derived from 388 unique HBV integration events) in the 99 pairs of tumor and matched normal tissues (Supplemental Table S11C). Most of these integrations were derived from partial HBV genomes. For example, the average length of these integrated HBV sequences was 1645 bp, with the minimum and maximum lengths being 59 bp and 3201 bp, respectively (Supplemental Fig. S7; Supplemental Table S11C). We randomly selected a total of 425 chimeric and split reads that supported these integrations and aligned them to the NCBI Nucleotide database using BLASTN (Camacho et al. 2009). All reads (100%) were verified (Supplemental Table S13). Among the 99 samples, 88 were also previously analyzed with an HBV-specific approach (Supplemental Table S14; Sung et al. 2012). In comparison, among the 88 samples our virome-wide approach detected the majority (305 of 399, 76.4%) of the 399 HBV integration breakpoints reported previously (Fig. 3F; Supplemental Fig. S8; Sung et al. 2012). A total of 94 of these previously identified breakpoints were not detected by Vcaller. However, these integration breakpoints were not fully validated, that is, only 32 of the 399 breakpoints were selected for PCR/Sanger sequencing validation in the original study (Sung et al. 2012). Additionally, Vcaller identified 169 new HBV integration breakpoints that were not detected in the original study (Fig. 3F; Sung et al. 2012). For example, we found strong evidence of 19 read pairs supporting a new HBV integration event of at least 808 bp in the *HCG2032978* gene (Fig. 3G). In support of the newly identified HBV integration events, we compared our findings to the results of an HBV sequence enrichment-based experimental analysis named HIVID (Li et al. 2013b), which was applied to 28 of the 88 tumors. Among the 28 tumors, Vcaller found 15 new integration events not detected in the original analysis, 13 of which (87%) were detected by HIVID. Seven of the 13 integrations identified by both Vcaller and HIVID, but missed in the original study, were analyzed by PCR and Sanger sequencing (Li et al. 2013b), and all were successfully validated (Supplemental Table S15). On the other hand, Vcaller found none of the five false-positive HBV integrations that were detected by HIVID but not successfully validated by PCR and Sanger sequencing (Supplemental Table S16; Li et al. 2013b).

Because the Vcaller approach is virome-wide and not limited to specific virus types, we also identified integrations of adeno-

associated virus (AAV) in two of the 99 tumor samples (Supplemental Table S17). One of the non-HBV integrations was derived from AAV-6 (AF028704) and was supported by 13 chimeric and split reads. It was 212 bp in length with a 2-bp deletion on the human genome at the breakpoint (Fig. 3H). Our analysis revealed that the other non-HBV integration had very low abundance (Supplemental Table S17). These results collectively demonstrate that Vcaller can detect new viral integrations and fusion transcripts from WGS and RNA-seq data, respectively. In all, our virome-wide approach obtained significantly higher sensitivity and accuracy than the currently available virus-specific approaches.

Determining clonal HBV integrations in the *TERT* and *MLL4* genes

Because of the high sensitivity and precision of Vcaller, we have identified many additional HBV integrations not detected in the original analysis, allowing us to perform a comprehensive characterization of HBV integrations in the HCC samples. Consistent with the previous HBV-specific study (Sung et al. 2012), the HBV integrations identified by Vcaller support the presence of integration hotspots, for example, clusters of breakpoint locations in the telomerase reverse transcriptase (*TERT*) and lysine methyltransferase 2B (*KMT2B*, also known as *MLL4*) genes. In the 99 liver cancer samples, we detected most of the reported *TERT* integrations (20 of 24), plus nine new *TERT* integrations (Fig. 4A). Among the 24 integrations, nine were experimentally validated (Sung et al. 2012), all of which were detected by Vcaller. We also identified all nine reported *MLL4* integrations (100%), plus two new *MLL4* integrations (Fig. 4B). We found that most of the *TERT* integrations (26 of 29) were located in the *TERT* promoter region (Fig. 4A), and almost all of the integrations contained at least one viral gene enhancer (*Enh1* or *Enh2*) or promoter (*XP*, *CP*, *SP1*, or *SP2*), particularly *Enh2*, *CP*, and *SP2* (Fig. 4C; Supplemental Table S18). The *TERT* gene expression levels were enhanced in the patients with integrations in *TERT* relative to those without integrations in *TERT* (Lau et al. 2014). We also found that most of the *MLL4* integrations were located between exons 3 and 6 (Fig. 4B). The HBV integrations enhanced the *MLL4* expression levels of the exons downstream from the integration breakpoint relative to the exons upstream of the breakpoint (Dong et al. 2015), indicating a potentially truncated *MLL4*, presumably serving as an oncoprotein. These results suggest oncogenic functions of up-regulation of *TERT* and truncation of *MLL4* in HCC. None of the samples had integrations in both *TERT* and *MLL4*, indicating that integrations in these two genes may be mutually exclusive (Fisher's exact test, $P=0.01$). Such mutual exclusivity usually implies functional redundancy or synthetic lethality of the two oncogenic variants.

By comparing the integration allele fractions of different integration events (Supplemental Fig. S2), we found that integrations in *TERT* and *MLL4* gene loci had significantly higher allele fractions compared to integrations in other parts of the genome (Fig. 4D). Even in the same individual, the *TERT* integration had higher allele fraction relative to other integrations (paired *t*-test $P=0.0006$ and fraction difference = 21.86 [10.56–33.17]) (Fig. 4E). A similar trend was found for *MLL4* (Supplemental Fig. S9). These observations indicate that the integrations in *TERT* and *MLL4* likely occurred in the early stages of liver tumorigenesis and/or conferred selective growth advantages to these cells. In both cases, the high cellular proportion implicates an oncogenic role for the virus.

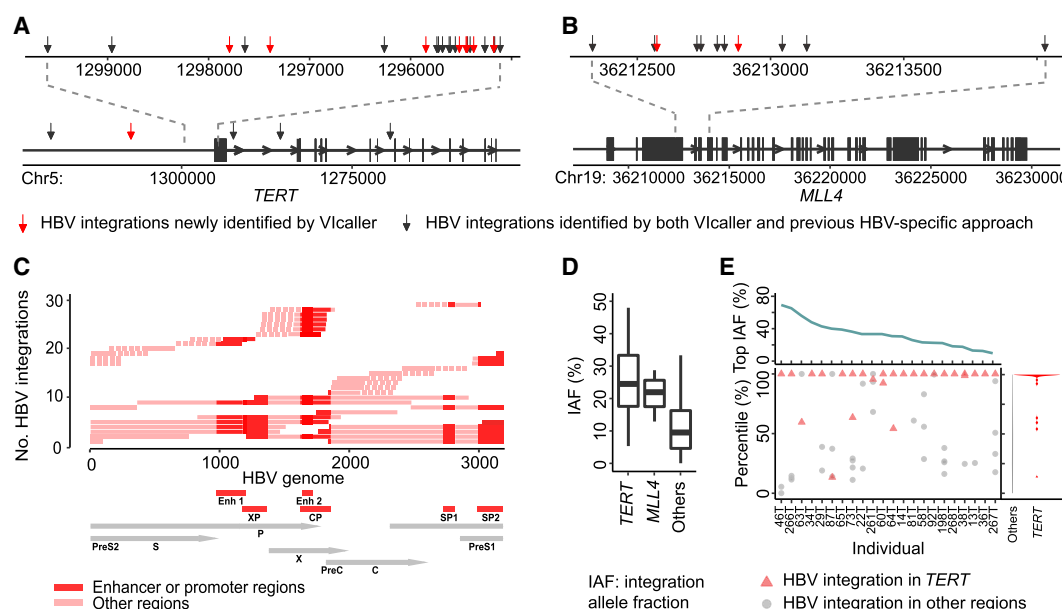


Figure 4. Characteristics of HBV integrations identified in tumors. (A, B) Sites of HBV integrations in two oncogenes: (A) *TERT*; (B) *MLL4*. (C) The integrated HBV sequences in *TERT*. The solid red lines above the HBV genome represent the integrated sequences with both breakpoints identified, whereas the dotted lines represent those with only one breakpoint identified. The HBV genes are in gray, and the promoters and enhancers are in red. (D) Comparison of integration allele fractions among HBV integrations in *TERT*, *MLL4*, and other chromosomal regions. (E) Integration allele fraction comparison of all HBV integrations in the samples with integrations in *TERT*. The top shows the highest integration allele fraction in each sample. The bottom left shows all HBV integrations in each sample, including those in *TERT*, and other regions (except *MLL4*). The bottom right shows the violin plot distributions of allele fractions of integrations in *TERT* compared to those in other regions (except *MLL4*). The result for *MLL4* is shown in Supplemental Figure S9.

Identifying HPV and BKV clonal integrations in bladder cancer

High-risk HPV and BKV have been linked to bladder cancer. The genomic sequences and fusion transcripts of the two viruses have been found in a small percentage of bladder tumors (Tang et al. 2013; The Cancer Genome Atlas Research Network 2014b). A recent paper reported genomic integrations of HPV and BKV in four bladder cancer samples (Cantalupo et al. 2018). To further evaluate our strategy and determine whether the integrations were clonal, we applied Vcaller to the WGS data of these four bladder cancer samples as well as an additional 102 randomly selected bladder cancer samples (Supplemental Table S19). Using our virome-wide integration screening, we found 10 integration events in the four samples, but no integration events in the additional samples. One sample carried five integrations of BKV (Fig. 5A; Supplemental Table S11D), and the other three carried two HPV-16, two HPV-56, and one HPV-45 integration, respectively (Fig. 5A). These integrations were only observed in tumor tissues, but not in paired normal tissues. The integration allele fractions for the most abundant integration event in each sample were 64.7%, 48.3%, 39.0%, and 30.0%, respectively (Fig. 5A), indicating that these integrations were shared by most cancer cells. The fact that we observed clonal integrations suggests the involvement of these viruses in the early stages of bladder tumorigenesis. All three observed HPV strains are considered high-risk HPV, capable of inducing cancers in a wide range of tissues (zur Hausen 2002). The oncogenic role of BKV has also been suggested (Abend et al. 2009), but not yet established. Figure 5, B and C, show a BKV integration and an HPV-45 integration, respectively. These results show that our novel strategy and platform can be used for the detection of oncoviruses present in additional cancer types using WGS data.

Identifying EBV integrations in non-Hodgkin's lymphoma and gastric adenocarcinoma

EBV was the first identified human oncovirus and associated with cancers such as non-Hodgkin's lymphoma (Heslop 2005) and gastric cancers (Iizasa et al. 2013). EBV genomes present largely as episomes, and to date, integrations of EBV into tumor genomes were mostly observed in established cell lines, such as Raji cells (Luo et al. 2004; Cao et al. 2015; Xiao et al. 2016). By screening a WGS data set (Supplemental Table S20) of 12 diffuse large B-cell lymphoma tumors (Morin et al. 2011), we detected an EBV (AB828191.1) integration in a patient (09-33003/DLBCL-PatientM). This patient had the shortest overall survival (1.31 yr) among the 12 patients (the average was 4.38 yr), and the difference was statistically significant (Z -test $P=0.008$). This EBV integration was only detected in the tumor, but not in paired blood cells. The integration was 20,941 bp in length and located in an intron of the *EHD1* gene (Fig. 5D). The allele fraction of this integration was 18%, supported by 22 chimeric and split reads. Additionally, we applied Vcaller to gastric adenocarcinomas, of which ~10% are EBV positive (Iizasa et al. 2013; The Cancer Genome Atlas Research Network 2014a). In 29 stomach adenocarcinoma samples (Supplemental Table S21), we found an EBV integration in one sample that was detected in the tumor but not in the paired normal tissue (Supplemental Fig. S10; Supplemental Table S11D). These results demonstrate that our approach can identify integrations derived from a wide range of viruses in cancer WGS data.

Determining capacity to identify integrations from uncharacterized viruses

KSHV and MCV were first isolated from tumors when searching for oncoviruses. It is likely that additional unknown viruses capable

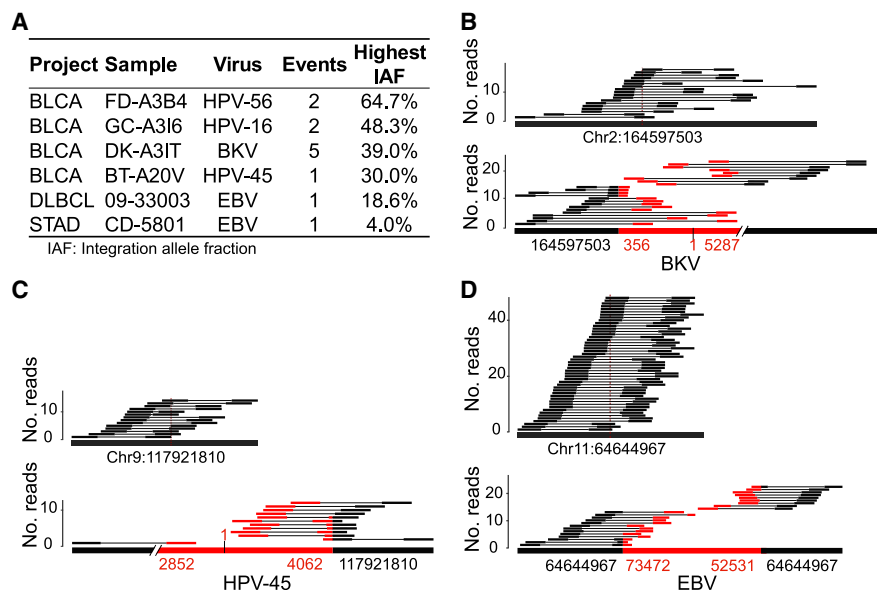


Figure 5. Identifying oncovirus candidates with integrations in bladder cancer, diffuse large B-cell lymphoma, and gastric adenocarcinoma samples. (A) Summary of identified integrations: The Cancer Genome Atlas (TCGA) Urothelial Bladder Carcinoma (BLCA); TCGA Stomach Adenocarcinoma (STAD); The Cancer Genome Characterization Initiative Diffuse Large B-Cell Lymphoma (DLBCL). (B) Sequence read alignment of a BKV (AB485698.1) integration event with 39% integration allele fraction detected in a bladder cancer sample TCGA-DK-A3IT. A total of 23 supporting read pairs, including 19 chimeric and four split reads, were found crossing the two breakpoints, supporting an integration, and 18 read pairs that support no integration were fully mapped to the human reference genome. (C) Sequence read alignment of an HPV-45 (EF202163.1) integration event with 30% integration allele fraction detected in a bladder cancer sample TCGA-BT-A20V. A total of 12 supporting read pairs, including 11 chimeric reads and one split read, were found crossing the two breakpoints, supporting an integration, and 14 read pairs were fully mapped to the human reference genome, supporting no integration. (D) Sequence read alignment of an EBV (AB828191.1) integration event with 18.6% integration allele fraction detected in a diffuse large B-cell lymphoma sample 09-33003. A total of 22 supporting read pairs, including 18 chimeric and four split reads, were found crossing the two breakpoints, supporting an integration, and 48 read pairs that support no integration were fully mapped to the human reference genome.

of causing cancers remain undiscovered. Viral integration screenings should not be limited to characterized viruses. Thus, we evaluated whether our platform had the capacity to identify some integrations derived from uncharacterized viruses. We first simulated a data set containing 97 random integrations of MCV (NC_010277.2) with 50% integration allele fraction in Chr 22 at 30x sequencing depth. To mimic MCV as an uncharacterized virus, we removed the MCV genome from the Vcalleer virome-wide reference library. By running Vcalleer on the simulated data with the MCV-depleted library, we still found 90% of the simulated MCV integration events (87 of 97). As expected, these events were annotated as arising from viruses having the highest sequence similarity with MCV, including *Gorilla gorilla gorilla* polyomavirus (GgorgPyV1) and *Pan troglodytes* virus polyomaviruses (PtvPyV) (Fig. 6A; Supplemental Table S22). The former is known to be closely related to MCV (Leendertz et al. 2011). Similarly, we ran Vcalleer on a simulated data set containing 97 HPV-18 integrations using an HPV-18-depleted library. We detected 94% of the simulated HPV-18 integration events (91 of 97). As expected, they were annotated as arising from other HPV strains (Fig. 6A; Supplemental Table S22). Most of the detected integration allele fractions were as high as 20%–30% (Fig. 6B), allowing for most integrations to be identified as clonal or subclonal.

We applied a similar strategy to the four bladder cancer samples carrying HPV or BKV integrations. We aligned all supporting

reads of the 10 integration events to the target virus-depleted library and still found integrations in two of the four samples (three of the 10 integration events) using the default parameters. As expected, they were annotated as from closely related viruses. For example, the two BKV integrations were annotated as from Vervet monkey polyomavirus 2 and JC polyomavirus, respectively. However, when we applied a less stringent threshold (two or more supporting reads with a minimum alignment score of 25), we found integrations in all four samples (seven of the 10 integrations). The allele fractions of all seven identified integrations remained clonal or subclonal (Supplemental Table S23).

These results suggest that our strategy and platform have the potential capacity to capture integrations from some uncharacterized viruses, especially clonal and subclonal integrations, benefiting from sequence similarity with related viruses. In future oncovirus discovery analyses, if certain nonhuman viruses are repeatedly identified with clonal or subclonal integrations, the possibility of the presence of an uncharacterized human virus should be considered because some novel oncoviruses may be missed if analyses are limited to characterized viruses.

Discussion

In this study, we first developed a novel strategy to discover oncoviruses and virus-caused tumors. Analyzing clonal integrations overcomes two major challenges: noncausal viruses and viral contaminations. Identification of clonal viral integrations was considered “the strongest evidence” in the discovery of MCV’s role in Merkel cell carcinoma (Moore and Chang 2010). However, the strategy of high-throughput screening

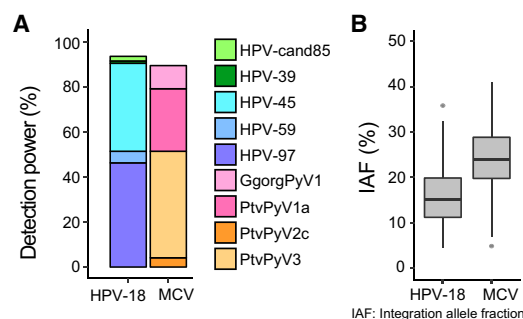


Figure 6. Viruses and integration events detected after removing the target viral genomes from our virome-wide database. (A) Percentage of simulated integrations detected after removing the HPV-18 (left) or MCV (right) references. (B) Integration allele fraction detected after removing the HPV-18 (left) or MCV (right) references. Three detected integration events that had >50% fractions are not shown in the figure, including two among the 91 HPV-18 events, and one among the 87 MCV events.

for clonal integrations has not been previously proposed or used to identify new oncovirus candidates. Second, we developed the first virome-wide clonal integration detection platform. Virome-wide analysis is necessary for identifying novel cancer viral etiologies. Our platform is also innovative because it includes integration allele fraction calculation, which provides additional supportive information to study the roles of viruses in tumorigenesis; yet it has not been utilized in previous bioinformatics analyses. Third, we have demonstrated that the strategy and platform are capable of identifying virus-caused tumors for a broad range of cancer types. By performing virome-wide screens using WGS data in several cancer types, we identified early-stage clonal integrations of HBV, HPV, BKV, and EBV, which were known or speculated to be oncoviruses in related tumor types. Fourth, we demonstrated the potential capability of our platform to identify integrations derived from uncharacterized viruses based on their sequence similarity with characterized viruses, allowing for further investigation. For example, if the MCV references were removed from our virome-wide reference library, most of the simulated MCV integration events were still found as clonal or subclonal integrations derived from other viruses having a high degree of sequence similarity to MCV. It is worth noting that MCV's oncoviral role was discovered because a piece of its genome that had high sequence similarity with African green monkey lymphotropic polyomavirus was "fished out" in Merkel cell carcinoma (Feng et al. 2008). Thus, for future cancer genomic analyses, if there are any new integration-capable oncoviruses or new cancer types associated with known oncoviruses, application of our strategy and platform to large collections of WGS data sets will be able to identify them.

EBV infects 90% of the world population (Cohen 2000); however, only a small proportion develop EBV-associated cancers (Young et al. 2016). The oncogenic role of EBV was established based on epidemiological, pathological, and oncological evidence, such as increased risks of EBV-associated cancers in EBV-infected individuals; EBV's capacity to cause infectious mononucleosis, a self-limiting lymphoproliferative disease; and its capacity to transform human normal lymphocytes (Cao and Li 2018). Monoclonal amplification of the EBV genome in cancer cells, indicated by loss of tandem repeat polymorphism, was also used to prove EBV's oncogenic role (Raab-Traub and Flynn 1986; Weiss et al. 1989). Our clonal integration analysis has a similar concept, and it can be applied directly to HTS data. EBV was largely considered incapable of integrating into the human genome. Our findings of clonal EBV integrations in human cancer further support ERV's role as an oncovirus (at a lower frequency compared to HBV and HPV).

Vcaller, the platform that we developed for implementing this strategy, is capable of correctly detecting virome-wide integrations in WGS and fusion transcripts in RNA-seq data. Vcaller incorporates the majority of the useful functions and features from existing virus and viral integration detection software (Supplemental Table S24; Hawkins et al. 2011; Kostic et al. 2011; Bhaduri et al. 2012; Borozan et al. 2012; Chen et al. 2013; Li et al. 2013a; Naeem et al. 2013; Wang et al. 2013, 2015; Katz and Pipas 2014; Naccache et al. 2014; Forster et al. 2015; Ho et al. 2015; Liang et al. 2017; Tennakoon and Sung 2017) as well as existing transposable element detection tools (Bourque et al. 2018; Goerner-Potvin and Bourque 2018). Vcaller implements many features, such as accurate estimation of integration allele fraction and fine-mapping of breakpoints (Supplemental Table S25). Because it considers all possible combinations of supporting read (Supplemental Fig. S3) and integration types (Supplemental Fig. S4), and applies stringent and comprehensive quality controls, Vcaller achieves high detec-

tion power and precision. We compared Vcaller with three recently developed software programs for candidate viral integration detection, and Vcaller consistently obtained the highest detection power and precision (Supplemental Table S10; Supplemental Fig. S11). Vcaller includes a step to annotate integration events that have been previously reported in commonly used cell lines (such as the Hela and SiHa cell lines). Vcaller accepts either raw FASTQ reads or aligned BAM files as its input and supports both single-end and paired-end sequences. Vcaller is open source and has the flexibility to be customized. All major software and utilities used by Vcaller can be easily replaced with alternative tools or updated to different versions. Vcaller is user-friendly with minimal bioinformatics skills required for users. Taken together, Vcaller can accurately detect virome-wide integrations in various HTS data sets, even for integrations with low cellular abundance and in cancer genomes with high mutation rates. The sensitivity and precision of our platform demonstrate the effectiveness of Vcaller for capturing clonal integrations present in tumor-derived WGS data. Therefore, Vcaller is a powerful tool for identifying new cancer viral etiologies. In the future, we will continue adding new viral reference genomes to our virome-wide library and will incorporate de novo assembly and viral sequence taxonomy prediction for the discovery of uncharacterized viruses.

In a proof-of-principle viral integration analysis, we found recurrent clonal integrations of high-risk HPV strains in bladder cancer samples. Although HPV transcripts, integrations, and fusion transcripts were previously detected in bladder cancer (Tang et al. 2013; The Cancer Genome Atlas Research Network 2014b; Cantalupo et al. 2018), the casual relationship between HPV exposure and bladder cancer has not been well-established (zur Hausen 2009a). As shown in this study, finding a high proportion of cells carrying identical HPV integrations provides strong genetic evidence to support the involvement of HPVs in the early stages of bladder tumorigenesis or the positive selection of the infected cells. Therefore, this study provides new genetic evidence in support of high-risk HPVs as oncovirus candidates in a subgroup of bladder carcinomas. Finding new oncoviruses is likely to be achieved by applying this strategy to cancer types with speculated, but undiscovered, viral etiologies. The most recently discovered human oncovirus was identified 10 yr ago from a rare skin cancer (Moore and Chang 2010). With decreasing sequencing costs, an increasing number of tumors, especially in understudied cancer types, will be sequenced, providing unique opportunities to identify novel viral etiologies. Our method will provide one solution that can be easily adapted for future cancer genome sequencing studies. Adding clonal integration analysis to standard analysis pipelines will aid in maximizing gains from sequencing investments.

In addition to discovering oncoviruses and virus-induced tumors, Vcaller has other applications. Viral integration analysis can lead to a more in-depth cancer genome profiling. For instance, HBV integrations at certain locations, such as the promoter of *TERT* and the 5' coding region of *MLL4*, may contribute to liver tumorigenesis and play a role in clonal selection. Our analysis found that the allele fractions of integrations in five other genes: *GAS7*, *PRDM16*, *ARID1B*, *AFF1*, and *NRG1*, were also among the highest in the corresponding tumor. Three of these five genes have been reported to be associated with cancer (So et al. 2003; Huang et al. 2004; Helming et al. 2014), indicating that the other two genes might also be cancer related. Given the potential roles of viral integration in tumorigenesis, this study has shown that viral integration should be included in standard WGS data analyses along with point mutations, Indels, copy number alterations/

aberrations, and chromosomal rearrangements. Furthermore, tumor growth follows a pattern of clonal evolution comprised of continuous clonal expansion, genetic diversification, and clonal selection (Greaves and Maley 2012). Viral integration sites are largely random, and most of the events are likely neutral. Continuous integrations at random locations contribute to the genomic diversity of tumor cells, and the integration sites are inherited during clonal expansion. Therefore, these integrations can also be used to study cancer heterogeneity and to track cell lineages during metastasis and drug resistance, especially when combined with single-cell sequencing techniques. Cell line contamination is relatively common in HTS studies (Jun et al. 2012; Laurence et al. 2014; Salter et al. 2014; Strong et al. 2014; Cantalupo et al. 2015; Friis-Nielsen et al. 2016). Identifying viral integrations identical to the events in an established cell line at nucleotide resolution suggests contamination by this cell line. This strategy was used by Vcanner to discover and filter out this type of contamination. Currently, we include a total of 25 cell lines (Supplemental Table S4; Peter et al. 2006; Klijn et al. 2014; Cao et al. 2015; Liu et al. 2016). By incorporating a comprehensive database of fine-mapped viral integrations and fusion transcripts derived from all commonly used cell lines in the future, our platform can also be used to detect cell line contaminations. Therefore, Vcanner can be easily adapted for use in a broad range of new applications.

Methods

Vcanner pipeline

Vcanner aligns FASTQ reads to the human reference genome, using BWA-MEM (Li 2013) for WGS data and TopHat2 (Kim et al. 2013) for RNA-seq data, with default parameters, or directly uses aligned BAM/SAM files. SAMtools (Li et al. 2009) is then used to extract paired-end unmapped reads with the parameters “-f 12 -F 256,” reads having one end mapped with the parameters “-f 8 -F 260,” and reads having one end unmapped with the parameters “-f 4 -F 264.” HYDRA’s bamToFastq (Quinlan et al. 2010) and in-house Perl scripts are then used to convert these reads to paired-end files (Supplemental Code). SE-MEI (<https://github.com/dpryan79/SE-MEI>) is used to extract the soft-clipped sequences, that is, unmapped sequences in mapped reads. All soft-clipped sequences that are 20 bp or longer are extracted with the parameter “-l 20,” and only the obtained soft-clipped sequences from paired-end reads mapped in proper pairs are kept for subsequent analyses (Supplemental Methods). To eliminate low-quality data, Vcanner uses NGS QC Toolkit (Patel and Jain 2012) and Perl scripts (Supplemental Code) to remove low-quality nucleotides (Q20) from the end of each read, or the entire read based on its quality score (i.e., the read is shorter than 20 bp or high-quality nucleotides account for <80% of the read). The clean reads are then aligned to our virome-wide reference genome library using BWA-MEM with the parameters “-k 19 -c 100000 -m 50 -T 20 -h 10000 -Y -M,” which are less stringent than the defaults. The putative supporting reads, including viral, chimeric, and split reads, as well as their physical locations on the human and viral genomes are obtained using Perl scripts (Fig. 1C; Supplemental Figs. S1, S3; Supplemental Code). Vcanner further realigns all resulting reads that are fully or partially mappable to viruses back to the human reference genome using BWA-MEM with the same (less stringent) parameters used for viral genome alignment (Supplemental Fig. S12). As human repetitive sequences, such as tandem and complex repeats, and other low-complexity regions, influence viral integration detection (Supplemental Fig. S13), as an optional step, Vcanner can use three different tools, including RepeatMasker

(Smit et al. 2015), TRF (Benson 1999), and DUST (Morgulis et al. 2006) to screen all supporting reads for repeat sequences. Vcanner removes the reads that map to the human genome, which are potential virus-human homologs, with these less stringent parameters, and the reads containing shorter than 20 bp nonrepetitive (uniquely mapped) sequence if there are no other supporting reads surrounding the breakpoint (i.e., within the insert size length). Last, the chimeric and split reads that do not follow the insert size distribution are also removed (Supplemental Fig. S14). The remaining reads constitute the clean supporting reads.

Viral integration status is determined primarily by the presence of chimeric and split reads (Supplemental Fig. S15). For each viral integration event, Vcanner identifies the physical locations of the upstream and downstream breakpoints on both the human and viral reference genomes. For integration events that are matched to multiple viruses or chromosomal locations, we determine the most likely candidate based on the number of supporting reads and their alignment scores. If the chimeric and split reads are mapped to more than one virus, this usually reflects homologous sequences present among these viruses, not random sequence matches. Considering that multiple levels of evidence are used to detect an integration, including (1) at least 20 bp mapped sequence required for each read, (2) multiple chimeric/split reads required for each breakpoint, and (3) at least one breakpoint (rather than viral sequences) required for each integration, it is extremely unlikely that random sequence matches make a significant contribution to the ranking of viral candidates. Viruses with a larger genome may have increased random sequence matches; however, except for extreme situations, such ranking metrics should not be biased for larger viruses.

The details are described in Supplemental Figure S1. When only one integration event exists in a genome, the viral read depth distribution will also be used as a confirmation of the detected event (Supplemental Fig. S16). To report a viral integration event, Vcanner requires a minimum of two chimeric and/or split read pairs with at least one uniquely mapped chimeric read pair (i.e., alignment score ≥ 50 if two supporting reads, and alignment score ≥ 30 if three or more supporting reads). These criteria were proven reliable based on Sanger sequencing experiments from this study and others (Sung et al. 2012).

Quality control

To eliminate potential false positives and low-quality viral integrations, stringent quality control metrics were developed and implemented in Vcanner, as described in detail in Supplemental Table S26. Furthermore, to achieve highly confident results, we used four different bioinformatics methods or settings to verify whether each of the supporting chimeric and split reads was uniquely mapped to expected human and viral genomic locations, and whether the mapped locations were consistent across different mapping tools (in silico validation). Only the integrations having reads uniquely mapped to both human and viral genomes, and that were consistently aligned by different alignment tools, were kept for further analyses. Last, we annotated the results with a list of integrations detected in commonly used cell lines. Similarly, we also annotated the results for potential vector contaminations with the VecScreen database.

Fine-mapping chromosomal locations of breakpoints

Both chimeric and split reads are used to fine map the physical locations of each breakpoint on the human and viral genomes (Supplemental Fig. S17; Supplemental Methods). The length of the integrated viral sequence is calculated for events with both

upstream (5') and downstream (3') breakpoints detected on the viral genome.

Vicaller output

The identified viral integration events are summarized in a uniform format. Each row of this output file contains a unique viral integration event. Each event contains 27 characteristics (Supplemental Table S5), such as genomic location, number of supporting reads, and alignment scores.

Integration allele fraction and cellular proportion

For each viral integration event, we calculated its integration allele fraction based on the combined number of chimeric and split reads versus the number of human reads crossing the two integration breakpoints (Supplemental Fig. S2). The human reads were extracted from the sorted BAM files using SAMtools. Specifically, we calculate the total number of chimeric and split reads from both upstream and downstream breakpoints (denoted as "a"), which support integration; and that of the human reads crossing the breakpoint (denoted as "b"), which support nonintegration at this chromosomal location. The ratio $a/(a+2b)$ is used as the integration allele fraction (Supplemental Fig. S2). When only one breakpoint is identified, the ratio $a/(a+b)$ is applied instead. A 50% integration allele fraction represents 100% cellular proportion in diploid cells.

Early-stage and growth-selected clonal integrations

An integration is considered "early stage" clonal and growth selected when its cellular proportion is reasonably high (such as 25% of the cells, considering various adjacent normal cells in the tumor) in the same specimen. By comparing the cellular proportions of clonal integrations with those of known tumorigenic somatic mutations, we can identify integrations that occurred in the early stages of tumorigenesis and conferred selective growth advantages to these cells. Viruses occurring in multiple samples with clonal integrations are considered oncovirus candidates for further evaluation.

Software availability

Vicaller is an open-source software. Vicaller v1.1 source code, documentation, and example data are available (www.uvm.edu/genomics/software/Vicaller.html; and <https://github.com/daweili-lab/Vicaller>). The source code is also available in the Supplemental Code.

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Author contributions: D.L. and J.C. initiated and designed the project. X.C., D.L., J.K., and A.S. analyzed the results. X.C., W.S.L., and N.W. performed the experimental work. X.C. developed the software code. D.L., J.C., and X.C. wrote the manuscript.

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