Next-Gen Sequencing Analysis and Algorithms for PDX and CDX Models

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Running Title

Improved PDX, CDX Data Processing

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Abstract

Patient-derived xenograft (PDX) and CTC-derived explant (CDX) models are powerful methods for the study of human disease. In cancer research, these methods have been applied to multiple questions including the study of metastatic progression, genetic evolution and therapeutic drug responses. Since PDX and CDX models can recapitulate the highly heterogeneous characteristics of a patient tumor, as well as their response to chemotherapy, there is considerable interest in combining them with next-generation sequencing (NGS) in order to monitor the genomic, transcriptional, and epigenetic changes that accompany oncogenesis. When used for this purpose, their reliability is highly dependent on being able to accurately distinguish between sequencing reads that originate from the host, and those that arise from the xenograft itself. Here we demonstrate that failure to correctly identify contaminating host reads, when analyzing DNA- and RNA-sequencing (DNA-Seq and RNA-Seq) data from PDX and CDX models is a major confounding factor that can lead to incorrect mutation calls and a failure to identify canonical mutation signatures associated with tumorigenicity. In addition, a highly sensitive algorithm and open source software tool for identifying and removing contaminating host sequences is described. Importantly, when applied to PDX and CDX models of melanoma, these data demonstrate its utility as a sensitive and selective tool for the correction of PDX- and CDX-derived whole exome and RNA-Seq data.

Implications: This study describes a sensitive method to identify contaminating host reads in xenograft and explant DNA and RNA sequencing data, and is applicable to other forms of deep sequencing.

Introduction
Human xenograft models have been widely used to study cancer. They provide an excellent tool with which to investigate the dynamics of oncogenesis, tumour heterogeneity, evolution and responses to therapy (1-8). This has led to considerable interest in combining them with NGS. This is challenging because downstream analyses are highly dependent on the quality and purity of the samples (9), leading to poor mutation calling accuracy and poor estimates of gene expression. While efforts can be made to mitigate these effects experimentally, high levels of infiltrating stromal cells often render this impractical. Consequently, levels of contamination as high as 73% have been observed in pancreatic cancer PDX models (10), and data are often variable (11). Instead, studies have typically addressed read-heterogeneity \textit{in silico} (9,12). Although the precise filtering strategy differs between studies, these studies all compare reads to both the mouse and human genomes and then eliminate those that match strongly to the mouse genome.

Despite the importance of reliable read filtering, only one method, Xenome, is implemented and readily available as a software tool (13). It is a computationally efficient approach that works by identifying 25-mer matches between the experimental data and the two candidate genomes, and using these to partition the data into host, graft, and ambiguous sets.

Here we describe a new algorithm for de-convolving host and graft reads. Unlike Xenome our algorithm makes use of full-length alignments and their scores, and can use values extracted from the CIGAR string or mapping quality scores when alignment scores are unavailable. With paired end data, in which two reads are generated for each DNA or RNA fragment, corresponding to its 5' and 3' ends, the algorithm resolves conflicts at the individual read level, not the fragment level, allowing more data to be retained. These approaches allow a weak but significant match to one organism to be ignored in favour of a stronger match to the other.
Together these enhance its discriminatory power. We demonstrate its utility for the analysis of human melanoma CDX models. The algorithm is freely available and released as an open source tool at https://github.com/CRUKMI-ComputationalBiology/bamcmp.git.

Materials & Methods

DNA/RNA from xenografts is always contaminated, and while an assay has been published (10) to quantify the proportion of human/mouse DNA in the samples from pancreatic cancer, a generalised method is still lacking. Various studies have reported the need to pre-process xenograft data before performing downstream analysis (9,13).

This method was developed to address the issues involving the analysis of both DNA/RNA xenograft data with a high accuracy. The model was designed so as to be generally applicable to any type of genomic data and also to a subset of common aligners. The method is based on filtering the host reads from the graft reads after aligning the reads to both host and graft genome using pre-existing alignment methods such as Burrows Wheel Aligner (BWA) (14), Bowtie2 (15) for DNA-Seq or Mapsplice2.0 (16), Tophat2 (17), STAR (18), and others, for RNA-Seq data.

Full experimental details for the BRAF<sup>V600E</sup> mutated cutaneous melanoma sample, patient information, ethics approval, and animal procedures, are available in reference (3).

As the alignment to both the host and graft is performed using the same aligner, the alignment scores from the aligner can be easily utilized to differentiate the origin of the reads. The reads are filtered on the basis of any of the four different parameters
described below, ordered on the basis of stringency (parameter names are in parentheses). In each case reads are assigned to the genome with the highest score. Consequently, no explicit thresholds are required:

1. Alignment scores (AS) if generated by the software (as).
2. CIGAR string values along with NM and MD tags discerned by the aligner (match).
3. Mapping Quality (MAPQ) scores of the alignments (mapq).
4. Remove everything that matches the host genome (balwayswins).

The reads are categorised into human only, mouse only and both. The latter category is further categorised into align better to human or align better to mouse after filtering. The reads that align only to human as well as those that align better to human (from the both category) are merged and returned as human reads; those that remain are assigned to mouse. The method can be used as a standalone application for filtering the contaminated reads or incorporated in the pipelines of routine NGS analysis. It has been implemented in C++ utilizing the htslib library from SAMtools (19).

**Filtering process:**

1. Align the fastq files to both human and mouse genomes.
2. Filter the mouse reads on any of the four filtering parameters.
3. Downstream processing as applicable (Mutation calling/read count generation/peak calling).

**Usage:**

All analyses for this study were performed with default parameters for Mapsplice2 (version 2.1.6), BWA-mem (version 0.7.11), Picard (version 1.96), GATK (version 3.3), Samtools (version 1.3.1) and Mutect (version 1.1.7). Output files from Xenome required minor additional processing in order to format them correctly for subsequent use by BWA and Mapsplice; results from Xenome were calculated from the graft reads only.

Results & Discussion

The data utilized in this study were derived from a cutaneous melanoma (20) patient (Patient 10) with a BRAF<sup>V600E</sup> mutation (3). The patient presented with primary melanoma on the back and bilateral axillary nodal metastasis. A PDX was derived from the bilateral axillary nodal metastasis. The patient relapsed after 3 months with liver, spleen and lymph node metastases. A CDX (CDXF1) was established from the patient’s CTCs taken at the time of relapse and grown in subsequent passage (CDXF2) that developed macro-metastases in liver, lymph nodes, kidneys, lungs, brain and distant subcutaneous tissue (3). Whole exome sequencing (WES) and RNA-sequencing (RNA-Seq) were used to profile the lymph node tumour (Tumour/Primary Tumour), PDX, CDXF1, CDXF2, CDXF2 Liver metastasis (Liver Met) and CDXF2 Lymph Node metastasis (LN Met). WES was also performed for patient whole blood (Germline) and Mouse kidney.

Here and throughout, all data were processed through the same pipeline (Supplementary Figure 1), with summary statistics computed in R (21) and Bioconductor (22). WES data were first aligned to human (hg19) and mouse (mm10)
genomes separately using BWA (14) with default parameters. While 99.81% human germline (i.e. never in mouse) reads aligned to the human genome, 42.68% of these mapped also to mouse; similar patterns were also observed for the mouse germline data (99.66%; 40.08%). Similar proportions of cross-species matches were also observed for the mouse xenograft material (Figure 1A). Together these data illustrate how a naïve filtering strategy that simply discards reads that map significantly to the mouse genome will be driven largely by orthology between human and mouse, and will thus discard substantial proportions of the data. We therefore sought to develop a filtering strategy better able to distinguish between host and graft reads.

WES data were processed using the default GATK framework (23) with mutations called using Mutect (24). Following filtering, using our new algorithm a minimum of 99.5% of human and mouse germline reads were correctly assigned to the right organism, while at most 0.20% reads could not be reliably mapped to either genome (Figure 1A). Similar improvements were observed for the xenograft material.

We next asked what effect the software had on mutation calling. Somatic mutations were called relative to the human germline control using Mutect. Without filtering, data were highly variable (631 to 8465 SNVs/sample), and concordance poor, despite the fact that all samples were derived from the same patient (Figure 1B). The number of Single Nucleotide Variants (SNV) predicted for each sample was also correlated with the level of host read contamination \( r = 0.98 \) in the xenograft samples (Figure 1C). Consistency increased dramatically after filtering (Figure 1A, B, D, E). Importantly, this was achieved with minimal effect on sensitivity: only one SNV called in the human primary tumour was lost and no false positives were obtained when the data was passed through the filtering pipeline (Figure 1F). On performing the same analysis using Xenome, fewer SNVs were detected; 4 SNVs were lost after filtering and an additional 6 false positives were obtained in the primary tumour...
We also calculated the variant allele frequency (VAF) using primary tumour before and after filtering. Since in the primary tumour data, no reads should be removed, optimal performance would result in no changes to the VAF. This analysis revealed higher agreement before and after filtering using our algorithm (Figure 1H), than with Xenome (Figure 1I). Similar analysis of CDX data reveals a similar trend, as expected (Supplementary Figure 2).

UV-related melanoma is strongly associated with a UV mutation signature comprising a disproportionate number of G>A/C>T transitions (25). Although detected in the primary tumour, this signature was not evident in the xenograft samples prior to filtering. After filtering it emerged strongly (Figure 1J).

RNA-Seq data from the same study were then aligned using MapSplice2.0 (16), and filtered using values extracted from the CIGAR string to provide mapping scores. As with the WES data, cross-species mappings were substantially reduced following filtering (Figure 2A), with levels of mouse contamination concordant to those of the WES data, but at an overall higher level (~15%). In order to investigate the effect of filtering on expression changes, we calculated fold-changes between the human primary and the mouse CDX model CDXF1, and compared those to the fold changes calculated after filtering. Fold changes for majority of the loci remained consistent, with 17 protein coding genes differing more than 2-fold between filtered and unfiltered sets (Figure 2B). When mouse filtering was applied to the human tumour data 202 protein coding genes were removed due to sequence homology. 1394 protein-coding genes exhibited greater than 4-fold difference between the unfiltered CDX and the filtered CDX data (values were computed for the mean of CDXF1 and CDXF2). Over-enrichment analysis of this set using gProfiler (26) found significant enrichment for genes associated with the extracellular matrix (Figure 2C), indicating that the reads filtered from the data set are of mouse stromal cell origin. Broadly,
similar results were obtained with Xenome (Supplementary Table 1), although a small portion of reads that mapped better to the mouse genome remained, even after filtering. 20 protein coding genes exhibited more than 2-fold change between filtered and unfiltered sets (Supplementary Figure 3), and a similar number of protein-coding genes (1405) displayed greater than 4-fold difference between the unfiltered CDX and the filtered CDX data. However, 988 protein-coding genes were absent from the filtered tumour dataset, vs. 202 with our algorithm, again confirming the improved selectivity of the bamcmp-based pipeline.

Conclusion
In conclusion, we present a sensitive and selective tool for identifying contaminating host reads in deep sequencing data from xenograft and explant models. While the results we present here focus on WES and RNA-seq data, the approach is equally applicable to other deep sequencing analyses including ChIP-seq and WGS.

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References


**Figure Legends**

**Figure 1.** Filtering reads of mouse origin improves sensitivity and selectivity of mutation calling from CDX models

A. Proportion of reads mapping to human and mouse genomes before and after filtering. Mouse: mouse germline sequenced from kidney; LN Met: lymph node metastasis; Liver Met: liver metastasis; CDXF1, CDXF2: CTC derived xenografts;
PDX: patient derived xenograft; Tumour: Patient primary tumour; Germline: patient whole blood.

B. Number of Single Nucleotide Variants (SNVs) called relative to human germline sequence, before and after filtering.

C. Number of SNVs called increases linearly with the number of mouse reads detected.

D. Correspondence in SNVs before filtering.

E. Correspondence in SNVs after filtering.

F. Filtering patient primary tumour against mouse removes only one SNV erroneously, and does not lead to others being detected.

G. As F, but filtering using Xenome.

H. Comparison of Variant Allele Frequency (VAF) before and after filtering for the primary tumour data.

I. As H, but filtering using Xenome.

J. The canonical C>T transition signature of UV damage is only detectable with correct read processing.

Figure 2. Filtering removes mouse reads from RNA-sequencing data without systematically disrupting expression levels.

A. Substantial reduction in cross-species mappings following filtering of RNA-sequencing data.

B. High correspondence in fold changes between human primary and CDX model before and after filtering. Loci no longer detected following filtering have been removed from the figure.

C. Over-representation analysis of loci no longer detected in xenograft data following filtering (BP: Biological Process; CC: Cellular Component; MF: Molecular Function).
Figure 1
Figure 2
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