TECHNICAL NOTE

Fusion transcripts and their genomic breakpoints in polyadenylated and ribosomal RNA–minus RNA sequencing data

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Abstract

Background: Fusion genes are typically identified by RNA sequencing (RNA-seq) without elucidating the causal genomic breakpoints. However, non–poly(A)-enriched RNA-seq contains large proportions of intronic reads that also span genomic breakpoints.

Results: We have developed an algorithm, Dr. Disco, that searches for fusion transcripts by taking an entire reference genome into account as search space. This includes exons but also introns, intergenic regions, and sequences that do not meet splice junction motifs. Using 1,275 RNA-seq samples, we investigated to what extent genomic breakpoints can be extracted from RNA-seq data and their implications regarding poly(A)-enriched and ribosomal RNA–minus RNA-seq data. Comparison with whole-genome sequencing data revealed that most genomic breakpoints are not, or minimally, transcribed while, in contrast, the genomic breakpoints of all 32 TMPRSS2-ERG–positive tumours were present at RNA level. We also revealed tumours in which the ERG breakpoint was located before ERG, which co-existed with additional deletions and messenger RNA that incorporated intergenic cryptic exons. In breast cancer we identified rearrangement hot spots near CCND1 and in glioma near CDK4 and MDM2 and could directly associate this with increased expression. Furthermore, in all datasets we find fusions to intergenic regions, often spanning multiple cryptic exons that potentially encode neo-antigens. Thus, fusion transcripts other than classical gene-to-gene fusions are prominently present and can be identified using RNA-seq.

Conclusion: By using the full potential of non–poly(A)-enriched RNA-seq data, sophisticated analysis can reliably identify expressed genomic breakpoints and their transcriptional effects.

Keywords: gene fusion; RNA precursors; RNA-seq; chromosome breakage; genomic structural variation; cryptic exons; TMPRSS2-ERG

Background

Genomic rearrangements are frequently observed in cancer and can drive disease initiation and progression through disruption of tumour suppressor genes and activation of oncogenes [1–3]. Marked examples include TMPRSS2-ERG fusions in prostate adenocarcinoma (PCa) [4] and BCR-ABL in chronic myelogenous leukaemia [5]. DNA rearrangements and their aberrant ligation products are identified as genomic breakpoints by whole-genome sequencing (WGS), but their potential role as driver mutation is mostly unresolved as of yet. The majority of genomic breakpoints intervene intergenic regions and are thus typically not located in messenger RNA (mRNA) and protein-coding sequences [6]. Moreover, genomic breakpoints of fusion genes are mostly located intronically [7]. To reveal their downstream effects, RNA sequencing (RNA-seq) is crucial to investigate changes at the transcriptional level and identify actual (in-frame) fusion transcripts. Conversely, for fusion transcripts, identification of the exact genomic breakpoint(s) can be essential to explain changes in gene expression and to define the origins of alternative promoters and altered splicing or polyadenylation events. Combined genomic and expression data enables further study of the functional consequences of genomic rearrangements and signifies whether an event is merely a passenger or a putative driver mutation [7, 8]. However, for many transcriptome studies, the exact genomic breakpoints of expressed rearrangements have not been resolved because matched WGS, Sanger sequencing, or similar analyses were not performed. Therefore, we set out to determine whether genomic breakpoints could be identified from RNA-seq data.

Next to targeted gene approaches, there are 2 main approaches in preparing RNA-seq libraries [9]. First, the more traditional method includes the positive selection of polyadenylated (poly(A)) mRNA to specifically target mRNA and eliminate abundant ribosomal RNA (rRNA). Because splicing takes place mostly co-transcriptionally, pre-mRNA is typically not polyadenylated and thus not included in this approach. Alternatively, one can extract total RNA and use random hexamer primers to initiate complementary DNA synthesis while removing abundant unwanted RNAs by various additional methods. This approach is referred to as rRNA-minus and is commonly applied when (partially) degraded RNA from formalin-fixed paraffin-embedded (FFPE) samples is sequenced.

rRNA-minus RNA-seq is thus capable of identifying non–poly(A) transcripts such as circular RNAs (circRNAs), specific types of small and long non-coding RNAs, and, importantly, actively transcribed precursor mRNAs (pre-mRNAs) [10]. Although the exact numbers depend on the protocol being used, tissue type, lariats [11], and intron lengths, typically 30–40% of RNA-minus RNA-seq reads map to intronic features, compared with 5–10% in poly(A) RNA-seq [12]. Therefore, rRNA-minus RNA-seq datasets require ≥50% higher sequencing depth to achieve an exon coverage comparable to poly(A) RNA-seq, while being capable of identifying additional RNA classes [9].

Fusion genes such as TMPRSS2-ERG and BCR-ABL are frequently observed as drivers within their respective malignant tissue [13]. Yet, many observed fusion genes are still of unknown consequence and seen in small frequencies in various cancer types. RNA-seq is highly suitable for fusion gene detection [14–16].

Methods to integrate RNA fusions with genomic breakpoints allow further assessment of functional consequences [7, 8, 17]. They are even capable of integrating complex higher order rearrangements but remain dependent on the availability of matching DNA data. State-of-the-art fusion-detection tools such as FusionMap, FusionCatcher, and JAFFA focus on exon regions or splice junctions specifically [18–20], which are the main target of poly(A)+ RNA-seq. Indeed, these tools also work well on RNA-minus RNA-seq because these also include exonic reads. Their efficient search space reduction in turn reduces the overall complexity and processing time. However, using rRNA-minus RNA-seq, typically 30–40% of the aligned reads are intronic and a further 20–25% of all reads are found to be intergenic [12], which are often a priori neglected. This large proportion of intronic and intergenic reads provides an opportunity to identify additional cancer-specific transcripts and exact genomic breakpoints of fusion genes. We have shown in a proof-of-concept that RNA-minus RNA-seq can indeed identify genomic breakpoints [10].
Here, we leverage a novel algorithm named Dr. Disco to report on the presence of genomic breakpoints in RNA-seq data, its implications regarding poly(A)+ and rRNA-minus RNA-seq, and on fused cryptic exons. The algorithm computationally identifies such genomic breakpoints and exon-to-exon junctions in a genome-wide fashion, taking into account the potential of rRNA-minus RNA-seq. It was applied on 6 RNA-seq datasets spanning multiple malignant tissue types (n = 1,275) (Table 1). Indeed, we reveal exact causal genomic breakpoints as derived from RNA-seq alone but limited to regions sufficiently expressed such as fusion gene TMPRSS2-ERG. Furthermore, rRNA-minus RNA-seq data can reveal more transcriptionally active rearrangements than poly(A)+ RNA-seq and results can be useful to supplement WGS. While only large datasets were analysed in this study, the method is developed for single-sample analysis. In summary, rRNA-minus RNA-seq in combination with a suited analysis pipeline gives a more complete view on both the origin and effects of genomic rearrangements and their direct influence on the expression of associated genes.

**Data Description**

RNA-seq data from several types of malignant tissue were used. For the NGS-ProToCol datasets (normal adjacent prostate, n = 41; prostate cancer, n = 51; normal adjacent colon, n = 18; colorectal adenoma, n = 30; and colorectal carcinoma, n = 30), carcinoma, adenoma (only colon), and adjacent normal tissue were rRNA-minus sequenced to study condition-specific molecular differences and further stratify tumour types [21, 22]. The PCA-LINES dataset consists of PCA cell lines PC346C and VCaP and additional PCA patient samples G-089, G-110, G-295, G-316, and G-346, which were WGS, rRNA-minus RNA, and poly(A)+ RNA sequenced. The included VCaP cell line is commonly used as a model system for prostate cancer and is known to contain the TMPRSS2-ERG fusion [23, 24]. The BASIS dataset consists of 560 WGS sequenced breast cancer (BrCa) samples [25] and 289 rRNA-minus RNA samples [26, 27], of which 207 are matching. The Chinese Glioma Atlas (CGGA) is composed of 274 rRNA-minus RNA-seq samples of various types of gliomas [28]. MCF-7 cell line data from ENCODE [29] were used for validation because it is a commonly used gold standard dataset [20]. We made the NGS-ProToCol and PCA-LINES publicly available; other data were taken from the public domain (Table 1).

To identify exact genomic breakpoints from rRNA-minus RNA-seq, we developed and implemented a novel algorithm, termed Dr. Disco. Briefly, it uses reads with a split alignment or read pairs with an inverted orientation or with a large insert size: discordant reads [30]. It uses reads not only from exons but also intronic and intergenic regions (Fig. 1 and Supplementary Table S1). Indeed, we reveal exact causal genomic breakpoints as derived from RNA-seq alone but limited to regions sufficiently expressed such as fusion gene TMPRSS2-ERG. Furthermore, rRNA-minus RNA-seq data can reveal more transcriptionally active rearrangements than poly(A)+ RNA-seq and results can be useful to supplement WGS. While only large datasets were analysed in this study, the method is developed for single-sample analysis. In summary, rRNA-minus RNA-seq in combination with a suited analysis pipeline gives a more complete view on both the origin and effects of genomic rearrangements and their direct influence on the expression of associated genes.

**Findings**

**Evaluation of poly(A)+ detectors**

The performance of the algorithm identifying mRNA fusions was assessed by analysing the ENCODE MCF-7 dataset, which was subsequently compared to poly(A)+ detector results published earlier [20] (Fig. 2) and with Arriba [31], which also makes use of STAR as aligner. Our method was not superior to JAFFA, and performed rather similarly to FusionCatcher. It was mostly limited in the total number of true-positive results identified, indicating that it is more conservative than JAFFA, SOAPFuse, and Arriba. Although the true-positives ratio for Arriba was considerably lower, the total amount of true-positive results was the highest.

Interestingly, unreported fusions spanning cryptic exons and intergenic regions were observed, including ATXN7-chr1:106,216,304, resulting in a fusion to cryptic intergenic exons supported by a matching genomic breakpoint (chr3:6,394,8014-chr1:106,192,959), and PRPF18-BEND7, resulting in anti-sense transcription of BEND7 spanning cryptic in-gene exons (Supplementary Fig. S2).

Dr. Disco’s performance of detecting only mRNA exon junctions was comparable but not superior to existing tools, while it revealed 27 additional high-confidence junctions using cryptic exons (Supplementary Table S1). The time it took Dr. Disco to complete analysis after the STAR alignment was 949 seconds, 2.07 times slower compared with Arriba (458 seconds). This is in concordance with the expectation that analysing a larger search space requires more conservative filtering and takes more resources to complete.

**Comparison of poly(A)+ and rRNA-minus RNA-seq**

Results from 7 PCa samples with matching rRNA-minus and poly(A)+ RNA-seq (PCA-LINES dataset) were compared (Fig. 3A). Contrary to our initial hypothesis, the poly(A)+ results also revealed intronic junctions, representing genomic breakpoints. Still, rRNA-minus data identified (3.4×) more intronic junctions as compared to poly(A)+ RNA-seq. The intronic junctions identified in poly(A)+ often had lower read counts or were located in untranslated region terminal exons as in-exon located genomic breakpoints (Supplementary Fig. S3). Terminal exons are known for their relative large size as they are ~6–7 times larger than internal exons [32]. The number of exonic junctions, thus predicted mRNA fusions, was nearly identical for rRNA-minus and poly(A)+ RNA-seq (144 vs 155).

**Comparison of RNA-seq with DNA-seq data**

Within the PCA-LINES dataset, the number of WGS-identified genomic breakpoints vastly outnumbered those extracted from the rRNA-minus RNA-seq (6.8%), indicating that only a frac-
Fusion transcripts and genomic breakpoints

We analysed the results of the algorithm on rRNA-minus RNA-seq data (n = 651) from different malignant tissue types (Figs 4B and 5): the BASIS, NGS-ProToCol colon and prostate, and CGGA datasets (Table 1).

Intronic and exonic junctions were identified in each dataset. The different malignant tissue types showed distinct regions enriched with intronic and exonic junctions (Fig. 4B). Known prominent events include TPRRSS2-ERG in PCs, EGFR, CDK4, and MDM2 in glioma, and CCND1 in BrCa. The breakpoints per sample and associated clinical parameters are provided in Fig. 5. The lowest mean number of genomic breakpoints per tissue type was found in normal adjacent samples (colon = 0.5; prostate = 0.9) followed by colorectal adenoma (1.1) (Supplementary Figs S9 and S10). In 2 adjacent normal-looking prostate samples, intronic and exonic junctions were found that were exactly identical to junctions in their matching malignant sample. These adjacent normal-looking tissue samples were most likely contaminated with cancer cells (Supplementary Fig. S9B). Of the different malignant tissue types, colorectal cancer samples were characterized by the lowest mean number of junctions (1.1) followed by combined low- and high-grade glioma (2.1) (Supplementary Fig. S11). Conversely, PCs (4.3) and BrCa (9.3) were characterized with cancer cells (Supplementary Fig. S9B). Of the different malignant tissue types, colorectal cancer samples were characterized by the lowest mean number of junctions (1.1) followed by combined low- and high-grade glioma (2.1) (Supplementary Fig. S11). Conversely, PCs (4.3) and BrCa (9.3) were characterized by relatively high numbers of genomic breakpoints per sample. These mean numbers were not normalized for sequence depth because results are also influenced by dataset-specific differences in read length, stranding, RNA quality, and library preparation. Therefore, comparison of these mean numbers of junctions is confounded by these factors.

Associations between the number of detected intronic junctions per sample and clinical parameters were investigated

Table 1: Dataset overview

| Dataset          | Tissue            | Sequencing              | n   | Depth   | Reference                          | Source  |
|------------------|-------------------|-------------------------|-----|---------|------------------------------------|---------|
| NGS-ProToCol     | Prostate cancer   | rRNA−RNA S              | 41  | ~70M    | EGA00001002816                     | [42]    |
| NGS-ProToCol     | Normal prostate   | rRNA−RNA S              | 51  | ~70M    | EGA00001002816                     | [42]    |
| NGS-ProToCol     | Colon cancer      | rRNA−RNA S              | 30  | ~70M    | EGA00001002854                     | [21, 22, 42] |
| NGS-ProToCol     | Colon adenoma     | rRNA−RNA S              | 30  | ~70M    | EGA00001002854                     | [21, 22, 42] |
| NGS-ProToCol     | Normal colon      | rRNA−RNA S              | 18  | ~70M    | EGA00001002854                     | [21, 22, 42] |
| BASIS            | Breast cancer     | rRNA−RNA S              | 289 | ~150M   | EGA00001001178                     | [26, 27] |
| PCa-LINES        | Prostate cancer   | rRNA−RNA S (+1)*        | 6   | ~37M    | EGA00001001476                     |         |
| PCa-LINES        | Prostate cancer   | poly(A)+ U              | 7   | ~50M    | EGA00001001476                     | [55]    |
| PCMM-FFPE        | Prostate cancer   | rRNA−RNA S              | 529 | ~40M    | GSE48865                            | [28]    |
| CGGA             | Glioma            | rRNA−RNA U              | 274 | ~30M    | GSE48865                            | [28]    |
| ENCODE MCF-7     | Breast cancer     | poly(A)+ S              | 1   | ~138M   | SRR54293                            | [20, 29] |
| BASIS            | Breast cancer     | WGS DNA                 | 560 | ~40×    | EGA00001001178                     | [25]    |
| PCa-LINES        | Prostate cancer   | WGS DNA                 | 7   | ~100×   | EGA00001001476                     | [24]    |
| Weier            | Prostate cancer   | TMPRSS2-ERG targeted DNA| 29  |         | EGA000000000052                     | [56]    |
| Pleasance        | Melanoma          | WGS DNA                 | 9   | ~40×    | EGA000000000052                     | [56]    |

The PCa-LINES dataset consists of 7 samples of which sample G-110 is rRNA-minus RNA-sequenced as part of the NGS-ProToCol study. Therefore the PCa-LINES dataset contributes 7 samples, from 6 unique patients, to the study.

Pan-cancer analysis

RNA sequencing datasets are given a suffix indicating whether they were performed stranded (S) or unstranded (U); rRNA− indicates rRNA-minus. The PCa-LINES dataset consists of 7 samples of which sample G-110 is rRNA-minus RNA-sequenced as part of the NGS-ProToCol study. Therefore the PCa-LINES dataset contributes 7 samples, from 6 unique patients, to the study.
Figure 1: Overview Dr. Disco pipeline and principle. (A) Schematic representation of fusion-gene RED-BLUE. Owing to relatively large intron sizes, in-gene genomic breakpoints occur most often intronically. The fusion results in different fusion splice isoforms (brown splice junction annotation in fusion gene structure). Fusion splice junction spanning reads form the classical evidence for mature mRNA fusion events (reads marked in green). In rRNA-minus data, intronic pre-mRNA reads (reads marked in pink) may cover causal genomic breakpoints. (B) Pipeline flow chart: RNA-seq data is aligned. Discordant reads are transformed into edges and inserted into a breakpoint graph. In the graph, intronic or exonic derived edges are kept separate. Detection of junctions is performed by analysing the graph for clusters. An additional splice variant correction is applied. Identified junctions are filtered, annotated, and marked intronic or exonic. QC/QA: quality control/quality assurance.

Figure 2: Evaluation on ENCODE MCF-7 dataset [20, 29]. For our algorithm, only non-head-to-tail junctions located on both sides at annotated exons were included. TP: true positive.
Figure 3: Overlap across sequencing types and library size influence. (A) Overlap of cumulative interchromosomal junctions of 7 WGS PCa samples rRNA-minus and poly(A)+ RNA-seq (PCa-LINES dataset). Overlap in only intronic junctions representing genomic breakpoints (left) and only exonic splice junctions (right). Of the 69 exonic junctions only found in rRNA-minus RNA-seq, 40 were detected in the matching poly(A)+ but did not pass filtering. Of the 80 poly(A)+-only exonic junctions, 58 were found in rRNA-minus but did not pass filtering. Not passing filtering mostly occurred because of insufficient discordant reads. (B) The number of predicted junctions as function of sequencing depth (left) and read-length (right) reduction. BrCa samples were selected for high sequencing depth (PR18022 and PR18037) or a high number of junctions (PR4841 and PR8660). Left: The number of predicted junctions per sequencing depth (10–100%) with the full read-length (2 × 75 bp). Reducing the sequencing depth, also for samples with a high sequencing depth, reduces the number of detected junctions. Sample PR4841 reaches a plateau. Right: Each data point represents the number of predicted junctions per given read length, at full sequencing depth. Truncating sequencing reads results in a low number of predicted junctions. However, below 55 nucleotides the number of detected junctions increases.

within datasets (Fig. 5). In BrCa, presence of kataegis \( (P = 1.9 \times 10^{-3}) \) was positively associated with the number of observed intronic junctions whereas estrogen receptor–positive (ER+) tumours were negatively associated \( (P = 0.9 \times 10^{-3}) \) with the number of intronic junctions. In glioma, tumour grade IV is positively associated with the number of intronic junctions per sample \( (P = 1.1 \times 10^{-3}) \), whereas tumour grade II \( (P = 2.9 \times 10^{-3}) \) and presence of IDH1 mutation \( (P = 0.8 \times 10^{-3}) \) were negatively associated. Within PCa no association was observed between the number of intronic junctions and the incidence of high Gleason grade \( (r>0.1; n=24/50) \) and metastasis \( (P = 0.16; n=8/51) \). Within BrCa, the number of intronic junctions correlated positively with the number of WGS-detected genomic breakpoints (Spearman correlation: \( \rho = 0.71, P = 2.2 \times 10^{-16} \), Supplementary Fig. S12). Because of the relative low number of junctions per sample combined with low number of colorectal cancer samples, further in-depth analysis on its recurrent events was not performed.

In the CGGA, BASIS, and NGS-ProToCol datasets ~65% of all intronic and exonic junctions have both sides located within an annotated gene (Fig. 6). Inversely, ~35% of the junctions have ≥1 side located within an intergenic region, regions that are often dismissed a priori by classical fusion gene detection tools [19, 20]. We found transcripts that incorporated cryptic (unannotated) exons, both intergenic as intronic (including anti-sense). For instance, a BrCa sample harboured intergenic junctions in SDC4 transcripts using 5 consecutive cryptic exons (Supplementary Fig. S13). In contrast, a PCa sample had an intergenic rearrangement lacking mRNA-level transcripts, thus only visible by the presence of pre-mRNA (Supplementary Fig. S14).

Genes associated with peaks in breakpoints

There were multiple, cancer type–specific hot spots of junctions located near known oncogenes (Fig. 4) such as KIT, PDGFR, EGFR, CDK4, MDM2 (glioma), TMPRSS2, ERG (PCa), FGFR1, and CCND1 (BrCa). Enrichment analysis was performed using HUGO symbols of genes recurrently hit per dataset, indicating that the pathway “Transcriptional misregulation in cancer [KEGG:05202]” was significantly more frequently hit \( (P = 1.6 \times 10^{-4}) \) within PCa due to TMPRSS2, ERG, ETV1, H3FA3, SLC45A3, and ELK4. Within BrCa, pathways E2F and E5F were significantly enriched \( (P = 6.75 \times 10^{-10}, P = 2.8 \times 10^{-5}) \) in ER+ BrCa and “Proteoglycans in cancer” in ER+ BrCa \( (P = 1.4 \times 10^{-5}) \). Genes that were recurrently hit
in glioma were found more often in pathways “Rap1 signaling pathway” \( (P = 3.2 \times 10^{-4}) \), “Glioma” \( (P = 5.9 \times 10^{-7}) \), and “Ras Signaling” \( (P = 2.6 \times 10^{-3}) \) (Supplementary Table S2).

**Large gene amplifications**

Hot spot regions (20–30 Mb) enriched with RNA-seq–detected junctions were observed in the BrCa (chr11) and glioma (chr12) datasets. These hot spots differed from focal events such as TMPRSS2-ERG in the sense that they were larger, had no consistent fusion partners, and often contained multiple hot spot junctions per sample. If these hot spot region rearrangements are responsible for consistent changes at the transcriptional level, they may provide a selective advantage. In both the BrCa and glioma datasets, transcriptional effects of the hot spot rearrangements were investigated by performing differential gene expression analysis between samples with (BrCa: chr11, glioma: chr12) enrichment with RNA-seq–detected junctions.
Figure 5: Pan-cancer results summary. Intronic and exonic junctions are given per sample for the NGS-ProToCol, BASIS, and CGGA datasets with their associated clinical parameters. For the colon samples, the predicted CMS classes are provided; for the prostate cancer samples, the Gleason grade and metastatic progression are provided; for the breast cancer samples, the ER, BRCA1, BRCA2, kataegis, and Dr. Disco–detected chr11-hot spot status are provided; and for the glioma samples, the grading, recurrence, IDH1 mutation status, sex, and the Dr. Disco-detected EGFR and chr12 hot spot status are provided.

Figure 6: (Inter)genic junction status. Frequency of intronic and exonic junctions and their gene/intergenic status. Because the glioma dataset was sequenced un-stranded, junctions with 1 intergenic side are grouped together. In all datasets, ∼3/8 of the junctions have ≥1 intergenic side. Inter- and intrachromosomal junctions were included, suspected circRNAs were discarded, and unlocalized and unplaced sequences (chrUn...) and alternate loci (chr...alt) spanning junctions were discarded. Intronic and exonic junctions corresponding to the same event were treated as a single entry.

and 13 samples, respectively), yet were not among the strongest up-regulated genes of the overall region. Instead, genes with a strong increase in logFC were FGF4 and CCND1, the cluster KCTD21, ALG8 and GAB2, and genes downstream of TENM4. Up-regulation of the overall region indicated amplifications of CCND1 and/or the gene cluster, which is in concordance with previous reports [33]. We presume that selection of breakpoints near SHANK2 is influenced by being adjacent to CTTN, a gene containing an enhancer often co-amplified with CCND1 [34]. The high frequency of junctions in the relatively large, yet not heavily up-regulated SHANK2 (785 kb) and TENM4 (788 kb) suggests that they are "collateral damage" of the amplifications, a hypothesis that has been described in glioma [35]. This hypothesis is further supported by the lack of consistent fusion partners, consistency in acting as acceptor or donor, and the absence of a clear spike in cumulative breakpoints (Fig. 7A and B; Supplementary Table S3).

Glioma samples having a junction harbouring the chr12 hot spot region (Fig. 7D–F) were analysed similarly and also showed up-regulation of genes in the hot spot locus, with an increased logFC of CDK4, MDM2, and neighbouring genes. Both CDK4 and MDM2 are known to be hyper-amplified in glioblastoma [36], often by double minute chromosomes [37]. The junctions showed a sharp increase in close proximity of CDK4 (Fig. 7D and E), likely indicating a common start of the amplification event. These breakpoints and up-regulated genes ceased just prior to LRIG3. Similarly, glioma samples harbouring rearrangements near the commonly hyper-amplified EGFR showed up-regulation of the surrounding locus (Supplementary Fig. S16).

Using RNA-seq data only, genomic rearrangements can be identified that can thereafter be used to reveal associated over-expression of oncogenes that have resulted from high-copy gene amplifications.
Figure 7: Differential gene expression in junction hot spot regions. (A–C) Overview of chr11 junctions, junction positions, and hot spot–associated differential gene expression in BrCa, using RNA-seq data only. (A) Intrachromosomal junctions not marked as putative circRNA, indicated by horizontal lines. (B) Junction end positions from intronic and exonic, inter- and intrachromosomal junctions not marked as putative circRNA. (C) Chromosomal differential expression plot for locus chr11:60,000,000–90,000,000 (grey square) with a q-value threshold of 0.001. Genes with the highest number of rearrangements, SHANK2 and TENM4, are illustrated with coloured boxes. Peaks in logFC were observed surrounding ORAOV1, CCND1 and FGF4, and surrounding TENM4. (D–F) Overview of chr12 junctions, junction positions, and hot spot–associated differential gene expression in glioma. (D) Intrachromosomal junctions not marked as putative circRNA are indicated with lines. (E) Junction end positions from intronic and exonic, inter- and intrachromosomal junctions not marked as putative circRNA. The junction enriched region chr12:40,000,000–75,000,000 is indicated with a grey square. (F) Chromosomal differential expression plot for locus chr12:40,000,000–75,000,000 with a q-value threshold of 0.01. Peaks in logFC from up-regulated genes are found near CDK4 and MDM2.

Chromothripsis

In VCaP, the q-arm of chr5 has been subjected to chromothripsis as revealed by 468 intrachromosomal WGS-detected breakpoints [24]. Seventeen intronic and exonic junctions were identified in rRNA-minus RNA-seq; thus evidence for chromothripsis events was identified at the (pre-)mRNA level (Supplementary Fig. S17). In 3 BrCa samples, high numbers of WGS-detected genomic breakpoints were identified on the q-arm of chr17. RNA-seq analyses revealed intrachromosomal and exonic junctions concordant with WGS data, which recurrently involved the genes BCAS3, APPBP2, MED13, USP32, and VMP1 (Supplementary Fig. S18). Taken together, this demonstrates the possibility of observing chromothripsis-derived junctions in RNA-seq.

TMPRSS2-ERG

From previous analyses it emerged that TMPRSS2-ERG was the most prominent focal event identified. Therefore, we leveraged NGS-ProToCol prostate tissue sample data to study this fusion in detail. TMPRSS2-ERG is a highly prevalent fusion gene in prostate cancer (∼50% of the diagnosed patients) [4], resulting in TMPRSS2-driven up-regulation of ERG. In 32 of the 51 samples Dr. Disco identified mRNA fusion transcripts of TMPRSS2-ERG, including genomic breakpoints in 27 of 32 samples (Fig. 8). These fusions were in concordance with high ERG expression in those samples exclusively. The detection rate for genomic breakpoints for this oncogenic fusion gene is thus markedly higher than for the overall number of genomic breakpoints. The genomic breakpoint did not pass filtering in sample 072, was marked exonic in sample 027, and was merged with closely adjacent (<450 bp; insert size) exonic junctions in 3 samples (053, 050, and 065), indicating that breakpoint-spanning reads were present in all 32 TMPRSS2-ERG–positive RNA-seq samples.

Three samples had their ERG-flanking genomic breakpoint located in an intergenic region upstream to ERG’s first exon.
Fusion transcripts and genomic breakpoints

Figure 8: TMPRSS2-ERG junction map. TMPRSS2 and ERG loci with endpoints of the junctions in NGS-ProToCol RNA-seq and non-matching targeted DNA-seq (Weier dataset). Gene structures are indicated at the bottom. Intronic junctions (representing genomic breakpoints) and genomic breakpoints (Weier dataset) are indicated in blue and exonic junctions in red. (A) For TMPRSS2, most breakpoints are detected after exon 1, up to exon 3. At mRNA level, apart from the first exons (1a and 1b), also exon 0 and exon 2 were commonly included in fusion transcripts. Two novel recurrent cryptic exons (*1 and *2) were common in fusion transcripts. (B) Three NGS-ProToCol samples (048, 05, and 075) have their genomic breakpoint before ERG and result in transcripts with additional, novel, intergenic cryptic exons. (Supplementary Figs S19 and S20). In 2 of these 3 samples, TMPRSS2-ERG fusion transcripts were identified containing cryptic intergenic exons (chr21:38,692,521–38,692,797 and chr21:38,701,593–38,701,947; hg38). In the same 2 samples, ERG had additional deletions, removing exon 2 (Supplementary Fig. S19).

The most abundant exonic junctions were T1-E4 and T1-E5 (Fig. 8, Supplementary Figs S20 and S21), which is in concordance with previous reports [38]. Genomic breakpoints were indeed located in hot spot regions within the first 2 introns of TMPRSS2 and the last half of ERG intron 3 [39]. Subsequent analysis of shallow sequenced FFPE RNA-seq samples revealed TMPRSS2-ERG in 181 samples (Supplementary Figs S20 and S21) and confirmed this remarkable breakpoint preference region within ERG intron-3 more precisely.

Two novel exons in TMPRSS2 were observed in both fusion and wild-type transcripts (Fig. 8). These cryptic exons were both expressed at a low level as they represented 3% of all TMPRSS2-ERG reads in samples having the splice variant. Furthermore, intragenic TMPRSS2 exon-0 [40] was detected in fusion mRNA-transcripts within 18 of 32 TMPRSS2-ERG–positive samples.

One sample contained an exonic junction originating in ERG and spanning to TMPRSS2 in which the gene order and included exons indicated that this ERG-TMPRSS2 fusion was caused by a reciprocal translocation instead of the common 3-Mb deletion (Supplementary Fig. S22). Other TMPRSS2-related fusions were TMPRSS2-RERE, SERINC5-TMPRSS2, TMPRSS2-TBX3, TMPRSS2-PADI4, MGA-TMPRSS2, and TMPRSS2-CATSPER2 (Supplementary Table S4).

VCaP

PCA cell line VCaP has been subjected to intensive research, revealing that it is TMPRSS2-ERG positive and contains 2 additional related genomic rearrangements (breakpoint A and breakpoint B) [24, 39]. TMPRSS2-ERG in VCaP was analysed using both rRNA-minus and poly(A)+ RNA-seq data.

Poly(A)+ RNA-seq shows that only the first exon of TMPRSS2 splices to ERG, even though the genomic breakpoint to ERG is located in the fifth intron (Supplementary Fig. S23A). The rRNA-minus data not only confirm this splice junction but also reveal all TMPRSS2- and ERG-spanning genomic breakpoints, concordant with the WGS results. Moreover, read stranding indicates that a region containing the fourth and fifth exon is inverted and
that its breakpoint A is an inversion. Breakpoint B is an amplification, and the junction from TMPRSS2 to ERG is again inverted such that ERG is transcribed in its original orientation. The junction from TMPRSS2 to ERG deletes the genomic region containing ERG’s exons 2 and 3. Thus, only TMPRSS2 exon 1 splices to ERG because exons 2 and 3 are deleted and exons 4 and 5 are inverted (Supplementary Fig. S23B). The small proportion of reads still present within the deleted TMPRSS2 exons 2 and 3 in the rRNA-minus data originate from the non-fusion allele(s). The rRNA-minus RNA-seq data not only revealed both intronic and exonic junctions but also showed the complex downstream effects on transcription and splicing.

CircRNA detection

Head-to-tail aligned reads (Supplementary Fig. S24) are marked as chimeric (discordant) by STAR and are used as input for our method. Such reads are not only observed in transcripts from genomic tandem duplications but also from circular mitochondrial DNA and circRNAs. Using the PCA-LINES rRNA-minus samples, we found that 88.6% of the junctions with a head-to-tail orientation were located exactly on exon-junctions corresponding to annotated circRNAs from circBase v31 (Supplementary Fig. S25).

Discussion

RNA-seq is generally performed as poly(A)+ RNA-seq, and fusion gene detection algorithms are in particular focused on annotated exons or splice junctions. It has become common practice to sequence ribosome-depleted total RNA (rRNA-minus) [12], especially for partially degraded (FFPE) RNA samples. rRNA-minus RNA-seq is interesting because it also yields non-polyadenylated transcripts, including pre-mRNA-derived intronic sequences. As a result, there is more genomic coverage in rRNA-minus RNA-seq alignments compared to poly(A)+ RNA-seq (Supplementary Fig. S26), which provides more opportunity to reveal fusion transcripts and a broader understanding of the transcriptome. Because genomic breakpoints are often harboured within introns [6] and intergenic regions (Supplementary Fig. S27), we interrogated to what extent rRNA-minus RNA-seq can be used to reveal genomic breakpoints because this also captures intronic (pre-mRNA) reads. Addressing this required analysing the genome without regional restrictions.

Here, we confirm, by using Dr. Disco, that RNA-seq data can be used to reveal genomic breakpoints of expressed transcripts in an automated fashion. Detection was limited to ~7% of WGS-detected breakpoints but markedly higher for the driver TMPRSS2-ERG fusion gene (85% detected; 100% presence). Because the algorithm was conservative in detecting mRNA fusions, it is likely that genomic breakpoints were missed and the actual percentage is somewhat higher. Conversely, estimation of this percentage implies that WGS results offer the ground truth, but these are also affected by noise, coverage, and filter cutoffs, indicating that this percentage is an approximation. The results commonly included intergenic junctions. For instance, 3 TMPRSS2-ERG fusions had their breakpoint located before ERG, supported by cryptic intergenic splice junctions and intergenic pre-mRNA coverage (Supplementary Fig. S19). Furthermore, intronic intergenic junctions in chromothripsis regions in 3 BrCa samples were in 14 of 18 cases validated by WGS junctions (Supplementary Fig. S18A).

That both Arriba and Dr. Disco make use of STAR and that Arriba finds a higher number of true-positive results indicates that improving the filtering is an important future step, but care must be taken not to compromise Dr. Disco’s true-positive ratio. The large search space combined with graph analysis was an effective solution as shown by providing a unique view on 1,275 transcriptomes. Whereas our initial hypothesis was that only rRNA-minus RNA-seq would reveal genomic breakpoints, this also accounts for poly(A)+. In poly(A)+ RNA genomic breakpoints are observed less frequently, with lower confidence, and often in long untranslated regions. The VCaP TMPRSS2-ERG analysis underlined the differences: on the basis of rRNA-minus RNA-seq, each genomic rearrangement with resulting splice variants and their strand as well as the order of events could be deduced, while both the poly(A)+ and WGS alone were insufficient.

CircRNAs are a relatively new group of non-polyadenylated transcripts with >90,000 different human circRNAs identified so far [41, 42]. The distinctive signature of proximate exonic head-to-tail junctions sets them apart from other junctions, except for small tandem duplications. A useful addition to the algorithm could be annotation of the junctions using a circRNA database such as circBase [41]. The proposed method is not specifically designed to identify circRNAs because it has stringent cut-off levels, merges splice variants into subgraphs, and requires >1 read. The number of circRNAs identified is therefore lower as compared with dedicated detection tools such as CIRI [43, 44].

The number of RNA-seq–detected intronic junctions representing genomic breakpoints varied widely between the 4 different cancer types (PCa, BrCa, colorectal cancer, and glioma). This variation is in line with the omics-reported number of structural variants—low in colorectal cancer [45] while high in BrCa [46, 47]—but these differences are confounded by the influence of sequencing depth, length, and library preparation, which vary per dataset.

While only a fraction of all genomic rearrangements is transcribed, an even smaller fraction is causal for fusion genes, which are currently of high interest in RNA research. Transcribed genomic breakpoints more often involve driver events than non-transcribed genomic breakpoints, as seen with TMPRSS2-ERG. Known exceptions that can be considered driver events include promoter and enhancer rearrangements such as known for AR and FOXP1 [48], but also tumour suppressor gene deletions [34, 49]. The reasoning that RNA-seq is not a replacement for WGS is valid. However, we show that by also looking at intronic and intergenic regions, more cancer-specific transcripts are identified and context is provided than being restricted to classic mRNA fusion genes only.

Although WGS depth surpasses 40× coverage, Dr. Disco showed that 26% and 48% of all RNA-seq intronic junctions in PCa and BrCa, respectively, were not identified by WGS. While these will contain false-positive results for sure, 100 of the 357 were found in 8 of the 207 samples all characterized by high number of genomic breakpoints. We suspect that this discrepancy is partially due to actual genomic breakpoints missed by WGS because of the following factors: high RNA-seq coverage of highly expressed genes (up to 1000×), clonality because this difference was in particular high for a small subset of samples, low local coverage in WGS, and selection criteria in software such as cut-offs and read mapping rules. In the PCMM-FFPE dataset, samples with low insert sizes or short read lengths often resulted in insufficient split-reads whilst resulting in many false positive read-pairs in full transcriptome analysis. However, FFPE data could still be used effectively in identifying the targeted, highly expressed, TMPRSS2-ERG fusion events.
Fusion transcripts and genomic breakpoints

The large number of PCa samples allowed TMPRSS2-ERG to be analysed in-depth, revealing additional cryptic and intergenic exons including TMPRSS2 exon 0 [40], a detailed map of the genomic breakpoints, genomic breakpoints located before ERG that combined with cryptic intergenic exons co-exist with exon 2 deletions, and a tumour that harbours TMPRSS2-ERG that originated from a reciprocal translocation rather than a deletion. In VCaP, stranded RNA-seq provided an advantage in deciphering the chronological order of complex genomic events. Moreover, this underlined the importance of automatic resolution of complex genomic rearrangements or poly-fusions. The current implementation does not offer such top-level integration for poly-fusions, while techniques exist with this purpose in mind [7, 50]. Integration of such techniques prompts future work. The current algorithm uses discordant reads exclusively. It would be interesting to investigate the added value of extending the detection with regions enriched with discordant opposite stranded reads, to strengthen detection of junctions having insufficient coverage of discordant reads.

In both BrCa and glioma, RNA-seq data alone revealed hot spot regions of junctions with subsequent up-regulation of known amplified oncogenes and neighbouring genes within these regions. The inconsistent transcriptional direction of the junctions combined with the lack of consistent acceptor/donor genes provides additional context that distinguishes these events from focal fusion genes. In BrCa, this combined analysis indicated that the events are related to CCND1 amplifications, despite the frequent events in sizeable genes TEMN4 and SHANK2 of which their fusion transcripts are not driving cancer.

Chromothripsis-derived junctions matching WGS-detected genomic breakpoints were present at the RNA level, in VCaP and 3 BrCa samples. As with most genomic rearrangements, the majority of the chromothripsis rearrangements were not detected on the RNA level. Solely on the basis of RNA-seq data, it will be difficult to prove the presence of chromothripsis because not all parameters that define this specific process can readily be extracted (e.g., copy-number variations, short insertions, loss of heterozygosity) [51, 52]. However, potential indicators for oncogenic chromothripsis events can be present in RNA.

While our preliminary aim was to study to which extent genomic breakpoints are present in RNA, we were surprised by how common both intergenic events and cryptic exons are. That ∼35% of the junctions in rRNA-minus datasets were full or partial intergenic events is an under-representation because, as for example with TMPRSS2-ERG exon-0, intergenic splice variants are merged with gene-spanning splice variants and will be considered in-gene as-whole. Intergenic mRNA fusions are characterized by incorporation of (typically multiple) cryptic exons. But cryptic exons are not limited to intergenic events, as we reported that cryptic exons in fusion transcripts transcribed in the antisense direction of a gene are common.

Cryptic exons are of importance because they may encode nonsense proteins with completely novel neo-antigens that are more divergent than point mutation–based neo-antigens and could therefore be more immunogenic [53]. We want to emphasize the importance of fusions beyond those incorporating annotated exons because we show that they can be transcribed into stable mRNA, thereafter be translated into protein, and potentially be oncogenic and/or immunogenic. This, deciphering the consequence of rearrangements, annotation of cryptic exons, and their coding potential for nonsense protein sequences is relevant for therapeutic interventions using tumour-specific antigens [54].

Conclusion

Facilitated by Dr. Disco, we set out to extract both intronic and exonic junctions from comprehensive rRNA-minus RNA-seq datasets and identified novel genomic breakpoints, circRNAs, novel gene and intergenic fusions, cryptic exons, and chromothripsis events and were able to link expressed rearrangements to transcriptional outcome. Discovering both genomic breakpoints and exonic junctions from only RNA-seq data requires an analysis strategy keeping these 2 levels of information separated. The number of breakpoints detected is limited to ∼7% because most are not within expressed regions. These results indicate that this analysis is not a replacement for WGS, but performing analysis like this will result in considerably more cancer-specific transcripts than interrogating classical fusion genes only. This holds in particular for rRNA-minus RNA-seq, which harbours more intronic reads. Furthermore, combined WGS and RNA analysis showed that RNA can function as informative supplement to WGS analysis because of stranding, expression, and resolution of the fusion gene structure(s). rRNA-minus RNA-seq provides more unique and complete information on non-polyadenylated and aberrant transcripts and, if the pre-mRNA is sequenced, the genomic breakpoints that underline transcriptional changes.

Thus, RNA-seq data can reveal genomic breakpoints, (cryptic and/or intergenic) splicing, and gene expression information, which together can reveal consequences and their selective advantage for cancer development and progression and be a useful supplement to DNA-seq.

Methods

Sequencing and datasets

The sequencing datasets used are given in Table 1. For NGS-ProToCol and the rRNA-minus RNA-seq of PCa-LINES, RNA was extracted using RNA-Bee (Campro Scientific, Berlin, Germany), and the library prepared for RNA-seq used the NEBNext Ultra Directional RNA Library Prep Kit for Illumina with rRNA reduction (New England BioLabs, Ipswich, Massachusetts, United States of America). The sample preparation was performed according to the protocol “NEBNext Ultra Directional RNA Library Prep Kit for Illumina” (NEB, Cat. No. E7420S/L and E6310S/L/X). Briefly, rRNA was reduced using the RNase H-based method. Then, fragmentation of the rRNA-reduced RNA and a complementary DNA synthesis was performed. This was used for ligation with the sequencing adapters and PCR amplification of the resulting product. The quality and yield after sample preparation were measured with the Fragment Analyzer (Advanced Analytical). Clustering and DNA sequencing using the Illumina cBot and HiSeq 2500 was performed according to manufacturer’s protocols (ServiceXS, Leiden, The Netherlands). A concentration of 16.0 pM of DNA was used as input. HiSeq control software HCS (v2.2.58) was used. Image analysis, base calling, and quality check were performed with the Illumina data analysis pipeline RTA (v1.18.64) and Bcl2fastq (v2.17). The 126-bp stranded Illumina HiSeq 2500 paired-end reads have a peak in fragment size of 300–600 bp and the samples have a mean depth of 70 million paired-end reads.

Of the PCa-LINES samples, each sample was WGS DNA sequenced and processed using the Complete Genomics platform [24, 57] (CompleteGenomics, San Jose, California, United States of America). The matching poly(A)- RNA-seq samples were taken from the TraIT-Cell Line Use Case study [55, 58]. rRNA-minus RNA-seq sample G-110 was not sequenced.
within PCa-LINES but sequenced in NGS-ProToCol as sample 7046-004-052.

In the BASIS RNA-seq dataset [26], total RNA was extracted and cleaned from abundant RNAs such as rRNA and transfer RNA as described elsewhere [27]. The BASIS DNA-seq data preparation and analysis is described elsewhere [25] and coordinates were converted to hg38 using pyliftover (v0.4) where needed.

The detection of genomic breakpoints from additional TMPRSS2-ERG fusions determined by targeted DNA-seq was described elsewhere [39], and genomic coordinates were obtained from this study accordingly. Genomic breakpoints of TMPRSS2-ERG and chromothripsis on chr5 in VCaP were described elsewhere [24, 39]. Predicted CMS classes for NGS-ProToCol colon samples were described elsewhere [22]. CGGA metadata were described elsewhere [28].

**Computational analysis**

RNA-seq data were aligned with STAR [59] (v2.4.2) using fusion settings and hg38 as reference genome. More details are given in the Supplementary Methods. Dr. Disco (v0.17.8, git commit 2a9f32950b71029b124f4d16544b2953c57d8e) was used for analysis. Arriba [31] (v2.1.0, git commit 349d24c28917f6c9320b1caab73afbb93f7bfbf) was used for evaluation analysis. For Supplementary Fig. S26, we designed and used our free software package to generate Lorenz and coverage plots and statistics (bam-lorenz-coverage, v2.3.0). Processed bam files used to estimate general genome coverage statistics were obtained from EGAS00000000052 [56]. Pathway enrichment was performed with gProfiler web (https://biit.cs.ut.ee/gprofiler/gost [60]), using gene identifiers as non-ordered query. For differential expression analysis, the annotation of the results of Dr. Disco, and further integration with gene sets for determining intergenic status, Ensembl 89 was used.

Plots were made with base R (3.6.3), ggplot2 (3.3.5), plotrix (3.8.1), and circlize (0.4.13), and illustrations, with Inkscape (0.92.4). Differential gene expression analysis was performed using edgeR (3.28.1) [61]. Associations between the frequency of breakpoints per sample and clinical parameters were tested using edgeR. Every gene located on the chromosome on which the locus is located is plotted with its genomic center as defined by Ensembl 89 on the x-axis and with edgeR’s logFC on the y-axis. A gene that is up-regulated in the mutant group has a positive logFC change, and a gene that is down-regulated, a negative logFC. When the gene is not significantly differentially expressed across the wild-type and mutant group (q-value below predetermined threshold) the gene will be coloured grey. If the difference is significant, it will be coloured green (up) or red (down).

Snapshots of discordant alignments were made in IGV (2.8.0) using the split view and with “Color alignments by” set to “read group.”

**Data Availability**

Supporting data, including reference data, final results tables, and chimeric alignments, and an archival copy of the code are available in the GigaDB database [62]. The results of the evaluation on the ENCODE MCF-7 dataset are given as Supplementary Table S1 [62].

**Availability of Source Code and Requirements**

- Project name: Dr. Disco
- Project home page: https://github.com/yhoogstrate/dr-disco
- bio.tools: https://bio.tools/dr_disco
- RRID:SCR_021739
- Operating system(s): GNU/Linux
- Programming language: Python
- Other requirements: STAR (aligner)
- License: GNU GPL 3.0
- Project name: bam-lorenz-coverage
- Project home page: https://github.com/yhoogstrate/bam-lor-enze-coverage
- bio.tools: https://bio.tools/bam-lorenz-coverage
- RRID:SCR_021837
- Operating system(s): GNU/Linux
- Programming language: Python
- Other requirements: Python libraries: numpy, pysam, tqdm, matplotlib & click

**Additional Files**

Supplementary Dr. Disco Technical Specification
Supplementary Figure S1. Exonic and intronic junctions
Supplementary Figure S2. Snapshot of fusion using cryptic exons in ENCODE MCF-7 dataset
Supplementary Figure S3. IGV Screeshot of in-exon genomic breakpoint
Supplementary Figure S4. Junctions in PCa-LINES dataset
Supplementary Figure S5. rRNA-minus RNA-seq and WGS data intersection BASIS dataset
Supplementary Figure S6. Correlation binned density WGS breakpoints and rRNA-minus junctions
Supplementary Figure S7. Intrachromosomal junctions in BASIS dataset
Supplementary Figure S8. Intrachromosomal junctions in BASIS dataset
Supplementary Figure S9. Intrachromosomal junctions in NGS-ProToCol prostate dataset
Supplementary Figure S10. Intrachromosomal junctions in NGS-ProToCol colon dataset
Supplementary Figure S11. Intrachromosomal junctions in CGGA dataset
Supplementary Figure S12. Correlation RNA-seq depth, WGS breakpoints, and RNA-seq junctions in BASIS dataset
Supplementary Figure S13. Snapshot cryptic fusion in BrCa sample PR9608a
Supplementary Figure S14. Snapshot of intergenic breakpoints in PCa sample 7046-004-134
Supplementary Figure S15. Chromosomal differential expression plot of chr11 in BASIS dataset
Supplementary Figure S16. Chromosomal differential expression plot of EGFR in CGGA dataset
Supplementary Figure S17. Chromothripsis on chr5q in VCaP
Supplementary Figure S18. Catastrophic events on chr17 in BASIS dataset
Supplementary Figure S19. Snapshot of intergenic breakpoints in TMPRSS2-ERG
Supplementary Figure S20. Detailed overview of junctions in ERG
Supplementary Figure S21. Detailed overview of junctions in TMPRSS2
Supplementary Figure S22. Schematic representation of TMPRSS2-ERG as reciprocal translocation
Supplementary Figure S23. TMPRSS2-ERG in VCaP
Supplementary Figure S24. Schematic overview discordant read orientations
Supplementary Figure S25. Overlap head-to-tail junctions and circBase in PCA-LINES dataset
Supplementary Figure S26. Lorenz and coverage plots of rRNA-minus, poly(A)+, and WGS data
Supplementary Figure S27. (Inter)genic events in WGS Supplementary Methods
Supplementary Table S1. Results SRR534293
Supplementary Table S2. gProfiler
Supplementary Table S3. SHANK2 and TENM4 related junctions
Supplementary Table S4. TMPRSS2-ERG and related junctions

Abbreviations
bp: base pairs; BrCa: breast cancer; circRNA: circular RNA; edgeR: empirical analysis of DGE in R; ER+: estrogen receptor–positive; FFPE: formalin-fixed paraffin-embedded; IGV: Integrative Genomics Viewer; logFC: logarithmic fold change; Mb: megabase pairs; mRNA: messenger RNA (‘- capped, polyadenylated, and spliced); PCA: prostate cancer; poly(A)+: polyadenylated; premRNA: RNA that is actively being transcribed by polymerase (not polyadenylated); rRNA-minus RNA-seq: RNA-seq prepared such that there is no specific positive selection for poly(A)-tails while reducing the amount of ribosomal RNA; poly(A)+ RNA-seq: RNA-seq prepared with a positive selection for poly(A)-tails; STAR: Spliced Transcripts Alignment to a Reference; WGS: whole-genome sequencing.

Ethical Approval
For the CTMM NGS-ProToCol study (NGS-ProToCol, Next Generation Sequencing from Prostate to Colorectal Cancer - Center for Translational Molecular Medicine (2014-2015); https://www.lygature.org/ctmm-portfolio), 51 prostate cancers from the Erasmus MC were snap-frozen and stored in liquid nitrogen as previously described [63]. Use of the samples for research purposes was approved by the Erasmus MC Medical Ethics Committee according to the Medical Research Involving Human Subjects Act (MEC 2004-261; MEC 2010-176).

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

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Authors’ Contributions
Y.H. and G.W.J. designed most of the experiments. Y.H. carried out most of the experiments and analysis and wrote the manuscript and software. M.A.K., R.B., J.v.R., H.J.G.v.d.W., R.H., A.P.S., and G.W.J. contributed to the methodology. N.D., D.S., D.v.d.M., F.P., C.H.B., G.J.L.H.v.L., M.S., J.W.M.M., W.v.W., B.J., E.C., M.d.J., G.A.M., E.v.d.B., A.S.B., R.H., R.J.A.F., and G.W.J. acquired data. H.J.G.v.d.W., J.v.R., S.v.L., C.R., R.B., M.S., A.P.S., and M.A.K. analysed and prepared data. P.J.F., R.B., M.S., G.W.J., B.J., R.J.A.F., and P.J.v.d.S. made large contributions to writing the manuscript. G.J., A.P.S., P.J.F., C.H.B., R.J.A.F., and P.J.v.d.S. acquired funding.

References
1. Weinhold N, Jacobsen A, Schultz N, et al. Genome-wide analysis of noncoding regulatory mutations in cancer. Nat Genet 2014;46(11):1160–5.
2. Calabrese C, Davidson NR, Demircio˘glu D, et al. Genomic basis for RNA alterations in cancer. Nature 2020;578(7793):129–36.
3. Li Y, Roberts ND, Wala JA, et al. Patterns of somatic structural variation in human cancer genomes. Nature 2020;578(7793):112–21.
4. Tomlins SA, Rhodes DR, Perner S, et al. Recurrent fusion of TMPRSS2 and ETS transcription factor genes in prostate cancer. Science 2005;309(5748):644–8.
5. Burmeister T, Schwartz S, Bartram CR, et al. Patients’ age and BCR-ABL frequency in adult B-precursor ALL: A retrospective analysis from the GMALL study group. Blood 2008;112(3):918–9.
6. Annala MJ, Parker BC, Zhang W, et al. Fusion genes and their discovery using high throughput sequencing. Cancer Lett 2013;340(2):192–200.
7. McPherson A, Wu C, Wyatt AW, et al. NFuse: Discovery of complex genomic rearrangements in cancer using high-throughput sequencing. Genome Res 2012;22(11):2250–61.
8. Zhang J, White NM, Schmidt HK, et al. INTEGRATE: Gene fusion discovery using whole genome and transcriptome data. Genome Res 2016;26(1):108–18.
9. Zhao S, Zhang Y, Gamini R, et al. Evaluation of two main RNA-seq approaches for gene quantification in clinical RNA sequencing: PolyA+ selection versus rRNA depletion. Sci Rep 2018;8(1):4781.
10. Erdem-Eraslan I, Van Den Bent MJ, Hoogstrate Y, et al. Identification of patients with recurrent glioblastoma who may benefit from combined bevacizumab and CCNU therapy: A report from the BELOB trial. Cancer Res 2016;76(3):525–34.

11. Taggart AJ, Fairbrother WG. ShapeShifter: A novel approach for identifying and quantifying stable lariat intronic species in RNAseq data. Quant Biol 2018;6(3):267–74.

12. Zhao W, He X, Hoadley KA, et al. Comparison of RNA-Seq by poly (A) capture, ribosomal RNA depletion, and DNA microarray for expression profiling. BMC Genomics 2014;15(1):419.

13. Heyer EE, Deveson IW, Wooi D, et al. Diagnosis of fusion genes using targeted RNA sequencing. Nat Commun 2019;10(1):1388.

14. Edgern H, Murumagi A, Kangaspeska S, et al. Identification of fusion genes in breast cancer by paired-end RNA-sequencing. Genome Biol 2011;12(1):R6.

15. Liu S, Tsai WH, Ding Y, et al. Comprehensive evaluation of fusion transcript detection algorithms and a meta-caller to combine top performing methods in paired-end RNA-seq data. Nucleic Acids Res 2015;44(5):e47.

16. Kim D, Salzberg SL. TopHat-Fusion: An algorithm for discovery of novel fusion transcripts. Genome Biol 2011;12(8):R72.

17. McPherson A, Wu C, Hajirasouliha I, et al. Conrad: Detection of expressed rearrangements by integrated analysis of RNA-Seq and low coverage genome sequence data. Bioinformatics 2011;27(11):1481–8.

18. Ge H, Liu K, Juan T, et al. FusionMap: Detecting fusion genes from next-generation sequencing data at base-pair resolution. Bioinformatics 2011;27(14):1992–8.

19. Nicolici D, Satalan M, Edgren H et al. FusionCatcher - a tool for finding somatic fusion genes in paired-end RNA-sequencing data. 2014. https://www.biorxiv.org/content/10.1011650v1. Accessed: 19 November 2014.

20. Davidson NM, Majewski IJ, Oshlack A. JAFFA: High sensitivity transcriptome-focused fusion gene detection. Genome Med 2015;7(1):43.

21. Komor MA, Bosch LJW, Bounova G, et al. Consensus molecular subtype classification of colorectal adenomas. J Pathol 2018;246(3):266–76.

22. Komor MA, de Wit M, van den Berg J, et al. Molecular characterization of colorectal adenomas reveals POFUT1 as a candidate driver of tumor progression. Int J Cancer 2020;146(7):1979–92.

23. Korenchuk S, Lehr JE, McLean L, et al. VCaP, a cell-based model system of human prostate cancer. In Vivo 2001;15(2):163–8.

24. Teles Alves I, Hiltemann S, Hartjes T, et al. Gene fusions by chromothripsis of chromosome 5q in the VCaP prostate cancer cell line. Hum Genet 2013;132(6):709–13.

25. Nik-Zainal S, Davies H, Stahl J, et al. Landscape of somatic mutations in 560 breast cancer whole-genome sequences. Nature 2016;534(7605):47–54.

26. Smid M, Rodríguez-González FG, Sieuwerts AM, et al. Breast cancer genome and transcriptome integration implicates specific mutational signatures with immune cell infiltration. Nat Commun 2016;7:12910.

27. Smid M, Wilting SM, Uhr K, et al. The circular RNAome of primary breast cancer. Genome Res 2019;29(3):356–66.

28. Bao ZS, Chen HM, Yang MY, et al. RNA-seq of 272 gliomas revealed a novel, recurrent PTPRZ1-MET fusion transcript in secondary glioblastomas. Genome Res 2014;24(11):1765–73.

29. Dunham I, Kundaje A, Aldred SF, et al. An integrated encyclopedia of DNA elements in the human genome. Nature 2012;489(7414):57–74.

30. McPherson A, Hormozdiari F, Zayed A, et al. Defuse: An algorithm for gene fusion discovery in tumor rna-seq data. PLoS Comput Biol 2011;7(5):e1001138.

31. Uhrig S, Ellermann J, Walther T, et al. Accurate and efficient detection of gene fusions from RNA sequencing data. Genome Respir 2021;31(3):448–60.

32. Bolisetty MT, Beemon KL. Splicing of internal large exons is defined by novel cis-acting sequence elements. Nuclear Acids Res 2012;40(18):9244–54.

33. Elsheikh S, Green AR, Aleksandarany MA, et al. CCND1 amplification and cyclin D1 expression in breast cancer and their relation with proteomic subgroups and patient outcome. Breast Cancer Res Treat 2008;109(2):325–35.

34. Morton AR, Dogan-Artun N, Faber ZJ, et al. Functional enhancers shape extrachromosomal oncogene amplifications. Cell 2019;179(6):1330–41.

35. Nikolaev S, Santoni F, Garieri M, et al. Extrachromosomal driver mutations in glioblastoma and low-grade glioma. Nature Commun 2014;5(1):5690.

36. Rollbrocker B, Waha A, Louis DN, et al. Amplification of the cyclin-dependent kinase 4 (CDK4) gene is associated with high cdk4 protein levels in glioblastoma multiforme. Acta Neuropathol 1996;92(1):70–4.

37. Decarvalho AC, Kim H, Poisson LM, et al. Discordant inheritance of chromosomal and extrachromosomal DNA elements contributes to dynamic disease evolution in glioblastoma. Nat Genet 2018;50(5):708–17.

38. Clark J, Merson S, Jhavar S, et al. Diversity of TMPRSS2-ERG fusion transcripts in the human prostate. Oncogene 2007;26(18):2667–73.

39. Weier C, Haffner MC, Mosbruger T, et al. Nucleotide resolution analysis of TMPRSS2 and ERG rearrangements in prostate cancer. J Pathol 2013;230(2):174–83.

40. Hermans KG, Boormans JL, Gasi D, et al. Overexpression of prostate-specific TMPRSS2(exon 0)-ERG fusion transcripts corresponds with favorable prognosis of prostate cancer. Clin Cancer Res 2009;15(20):6398–403.

41. Gläzar P, Papavasileiou P, Rajewsky N. CircBase: A database for circular RNAs. RNA 2014;20(11):1666–70.

42. Chen S, Huang V, Xu X, et al. Widespread and functional RNA circularization in localized prostate cancer. Cell 2019;176(4):831–43.

43. Zeng X, Lin W, Guo M, et al. A comprehensive overview and evaluation of circular RNA detection tools. PLoS Comput Biol 2017;13(6):e1005420.

44. Gao Y, Zhang J, Zhao F. Circular RNA identification based on multiple seed matching. Brief Bioinform 2018;19(5):803–10.

45. Priestley P, Baber J, Lolkema MP, et al. Pan-cancer whole-genome analyses of metastatic solid tumours. Nature 2019;575(7781):210–6.

46. Alaei-Mahabadi B, Bhadury B, Karlsson JW, et al. Global analysis of somatic structural genomic alterations and their impact on gene expression in diverse human cancers. Proc Natl Acad Sci U S A 2016;113(48):13768–73.

47. Yoshihara K, Wang Q, Torres-Garcia W, et al. The landscape and therapeutic relevance of cancer-associated transcript fusions. Oncogene 2015;34(37):4845–54.

48. van Dessel LF, van Riet J, Smits M, et al. The genomic landscape of metastatic castration-resistant
prostate cancers reveals multiple distinct genotypes with potential clinical impact. Nat Commun 2019;10(1):5251.

49. Abdulaziz H, Petreaca B, Petreaca R. Frequent homozygous deletions of the CDKN2A locus in somatic cancer tissues. Mutat Res 2019;815:30–40.

50. Tian L, Li Y, Edmonson MN, et al. CICERO: A versatile method for detecting complex and diverse driver fusions using cancer RNA sequencing data. Genome Biol 2020;21(1):126.

51. Korbel JO, Campbell PJ. Criteria for inference of chromothripsis in cancer genomes. Cell 2013;152(6):1226–36.

52. Govind SK, Zia A, Hennings-Yeomans PH, et al. ShatterProof: Operational detection and quantification of chromothripsis. BMC Bioinformatics 2014;15(1):78.

53. Menez-Jamet J, Gallou C, Rougeot A, et al. Optimized tumor cryptic peptides: The basis for universal neoantigen-like tumor vaccines. Ann Transl Med 2016;4(14):266.

54. Gubin MM, Artyomov MN, Mardis ER, et al. Tumor neoantigens: Building a framework for personalized cancer immunotherapy. J Clin Invest 2015;125(9):3413–21.

55. Hoogstrate Y, Zhang C, Senf A, et al. Integration of EGA secure data access into Galaxy. F1000Res 2016;5:doi:10.12688/f1000research.10221.1.

56. Pleasance ED, Cheetham RK, Stephens PJ, et al. A comprehensive catalogue of somatic mutations from a human cancer genome. Nature 2010;463(7278):191–6.

57. Hiltemann S, Jenster G, Trapman J, et al. Discriminating somatic and germline mutations in tumor DNA samples without matching normals. Genome Res 2015;25(9):1382–90.

58. Zhang C, Bijlard J, Staiger C, et al. Systematically linking transSMART, Galaxy and EGA for reusing human translational research data. F1000Res 2017;6:doi:10.12688/f1000research.12168.1.

59. Dobin A, Davis CA, Schlesinger F, et al. STAR: Ultrafast universal RNA-seq aligner. Bioinformatics 2013;29(1):15–21.

60. Reimand J, Arak T, Adler P, et al. gProfiler—a web server for functional interpretation of gene lists (2016 update). Nucleic Acids Res 2016;44(W1):W83–9.

61. Robinson MD, McCarthy DJ, Smyth GK. edgeR: A Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 2009;26(1):139–40.

62. Hoogstrate Y, Komor MA, Böttcher R, et al. Supporting data for “Fusion transcripts and their genomic breakpoints in poly(A)+ and rRNA-minus RNA sequencing data.” Gigascience Database 2021; http://dx.doi.org/10.5224/100939.

63. Hendriksen PJM, Dits NJF, Kokame K, et al. Evolution of the androgen receptor pathway during progression of prostate cancer. Cancer Res 2006;66(10):5012–20.