From somatic variants towards precision oncology: Evidence-driven reporting of treatment options in molecular tumor boards

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Abstract

Background: A comprehensive understanding of cancer has been furthered with technological improvements and decreasing costs of next-generation sequencing (NGS). However, the complexity of interpreting genomic data is hindering the implementation of high-throughput technologies in the clinical context: increasing evidence on gene–drug interactions complicates the task of assigning clinical significance to genomic variants.

Methods: Here we present a method that automatically matches patient-specific genomic alterations to treatment options. The method relies entirely on public knowledge of somatic variants with predictive evidence on drug response. The output report is aimed at supporting clinicians in the task of finding the clinical meaning of genomic variants. We applied the method to 1) The Cancer Genome Atlas (TCGA) and Genomics Evidence Neoplasia Information Exchange (GENIE) cohorts and 2) 11 patients from the NCT MASTER trial whose treatment discussions included information on their genomic profiles.

Results: Our reporting strategy showed a substantial number of patients with actionable variants in the analyses of TCGA and GENIE samples. Notably, it was able to reproduce experts’ treatment suggestions in a retrospective study of 11 patients from the NCT MASTER trial. Our results establish a proof of concept for comprehensive, evidence-based reports as a supporting tool for discussing treatment options in tumor boards.

Conclusions: We believe that a standardized method to report actionable somatic variants will smooth the incorporation of NGS in the clinical context. We anticipate that tools like the one we present here will become essential in summarizing for clinicians the growing evidence in the field of precision medicine. The R code of the presented method is provided in Additional file 6 and available at https://github.com/jperera-bel/MTB-Report.

Keywords: Genomic report, Personalized treatment, Cancer genomics, Targeted therapies, Predictive biomarkers, Molecular tumor board, Actionable variants

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Background

Precision medicine (PM) aims at understanding, and thus managing, a patient’s disease based on its molecular profile. In oncology, PM is being implemented through tumor boards (TB), multidisciplinary meetings where patients are discussed from all possible perspectives (diagnostic radiology, surgery, pathology, genetics, oncology, radiotherapy, etc.). Currently, TB include routine testing of several predictive biomarkers which indicate the likelihood to respond to a specific treatment. However, these are restricted to few cancer entities and just a handful of examples exist, for instance EGFR inhibitors in non-small cell lung cancer tumors with EGFR mutations in exons 19/21 (afatinib, erlotinib, gefitinib) or resistance mutation T790 M (osimertinib), HER2 inhibitors (trastuzumab) in breast tumors with HER2 amplification/overexpression, and BCR-ABL inhibitors (bosutinib, desatinib, imatinib, nilotinib, ponatinib) in Philadelphia chromosome-positive hematologic malignancies [1]. Predictive biomarkers are recognized by reference approval and guideline organizations (e.g., The Food and Drug Administration (FDA), The European Medicines Agency, and the National Comprehensive Cancer Network (NCCN)) and are specific for a cancer type. When a tumor progresses after the regulated, evidence-based therapies have been applied, testing for genomic alterations of targets of drugs approved for other cancer entities, or even drugs under clinical trials, becomes a common strategy for rational prescription of drugs. However, even if a drug target harbors a mutation, it does not mean that the variant predicts drug response.Inferring causality between a gene variant and a drug response in an off-label scenario (e.g., another cancer type, unknown mutation in the target gene, and/or mutation in a gene upstream of the target gene) is very challenging [2]. For this reason, the scientific community has already stated the need for a comprehensive knowledge database of variants and affected genes with respect to drug response (i.e., actionable variants) [3]. In the same direction, the clinical community needs decision support platforms for interpreting somatic alterations in regard to clinical action [4–7]. Last, but not least, there is a gap between legislation and health insurance policies with regard to who should pay for these expensive treatments in off-label scenarios. Typically, health insurance companies will only pay for these treatments if the clinician is able to prove causality. Otherwise, the only opportunity to receive an off-label treatment will be under the umbrella of a clinical trial. In turn, clinical trial enrollment can be complex due to strict inclusion criteria (e.g., restricted to some cancer entities, molecular markers, and previous therapies) and location of the clinical facilities.

In addition, the latest advances in next generation sequencing (NGS) will, in the foreseeable future, allow us to sequence the whole genome of patients in a reasonable time. Few worldwide clinical institutions have taken the first steps in this direction by implementing molecular tumor boards (MTBs) to study the feasibility of performing comprehensive genomic profiling (e.g., NGS) in the clinical context [8–16]. However, NGS generates huge volumes of data, ranging between 5 and 200 Gb per patient, depending on coverage, sequencing technique (whole exome, whole genome), etc. Hence, interpreting the clinical implications of single nucleotide variants (SNVs), copy number variations (CNVs), and gene fusions of a tumor biopsy (i.e., somatic variants) becomes increasingly complex. One main difficulty lies in distinguishing the relevant alterations with either diagnostic, prognostic, or predictive meaning from all the others [3, 17]. Compiling a comprehensive list of somatic variants for one patient, scanning through all the variants, and summarizing the clinical implications and evidence based on current literature knowledge is already a huge task. For that, clinicians integrate information from web-based tools such as My Cancer Genome (mycancergenome.org), OncoKB [18], or literature searches (PubMed, Google Scholar). Nevertheless, this strategy is far from being optimal: it requires time, is subject to omission of information, and depends on the clinician’s ability to interpret the clinical implications from genomic research articles. Recent studies have predicted that the use of NGS profiling and drug repurposing have the potential to identify actionable genomic alterations in more than 70% of cancer patients [19, 20]; however, the clinical reality is very far from this.

In this work, we aim to generate an automated tool that produces patient-specific reports comprising a filtered list of patient’s actionable variants with respect to potential treatment options. Although treatment decisions will always require expert knowledge, we hypothesize that such a tool will substantially facilitate the use of NGS in clinical practice by simplifying the task of finding the clinical implications of genomic variants. To this end, we present here a framework that pre-filters, classifies, and reports actionable variants, defined as somatic genomic alterations, which have some evidence of drug response or resistance. We then show the results of applying the tool to different patient cohorts (The Cancer Genome Atlas (TCGA), Genomics Evidence Neoplasia Information Exchange (GENIE), and NCT MASTER). With it, we hope to provide a proof-of-concept of a tool that can filter a list of variants and link them to available evidence for genomic-based treatment decisions (e.g., information from literature, clinical trials, and databases).

Methods

In this article we present a framework to match tumor genomes to targeted therapies. For that, we use public
databases to filter potentially actionable variants from a tumor sample and then classify the results using an evidence-based system. In this section we first detail the databases used. Then, we present the patients’ datasets used for evaluating the feasibility and provide a proof-of-concept of our approach. Finally, we describe all statistical analyses.

**Databases of actionable variants**

Among the main criteria of database selection, we focused on those databases that compile information not only on the drug and actionable gene, but also the actionable variant (SNV, CNV, rearrangement), the type of association (response, resistance), the strength of the evidence (approved, clinical trials, preclinical) and the cancer type. As a result, we have selected the following databases: (1) *Gene Drug Knowledge database* (GDKD) [20] (version 19, downloaded from Synapse syn2370773); (2) *Clinical Interpretation of Variants in Cancer* database (CIViC) [21] (version 01_June_2017); and (3) *Tumor Alterations Relevant for Genomics-driven Therapy* database (TARGET) [12] (version 3). GDKD is a manually curated database of predictive biomarkers updated monthly. It integrates several layers of annotations, comprising cancer type, gene, variant, response/resistance, consensus/emerging, and the corresponding references. CIViC uses very similar layers of annotation, but is a community-driven web resource. For this work, both GDKD and CIViC were modified in such a way that variants in the same gene sharing annotations of disease, drug, evidence, and association levels were aggregated into one single entry. Finally, TARGET was published in 2014 and has not been updated since. Taken together, the three databases compile a comprehensive list of variants comprising a total of 289 actionable genes (conferring either resistance or response to anticancer drugs). The main characteristics of the three databases can be found in Table 1. We also consulted other sources such as NCCN guidelines, mycancergenome.org,

| Table 1 Main characteristics of the public databases of predictive biomarkers | GDKD | CIVIC | TARGET |
|---|---|---|---|
| Number of genes | 170 | 200 | 135 |
| Number of predictive genes | Total 170 | 213 | 111 |
| Exclusive | 46 | 105 | 10 |
| Common | 79 | | |
| Number of variant-drug associations | 618 | 1931 | 111 |
| Number of cancer types | 65 | 177 | Not specified |
| Biomarker types | Predictive | Predictive | Predictive |
| | Prognostic | Prognostic | Prognostic |
| | Diagnostic | Diagnostic | |
| Clinical significance levels | Response | Sensitivity | Resistance or non-response |
| | Sensitivity | | |
| | Increased benefit | | |
| | No response | | |
| | No sensitivity | | |
| | Reduced/decreased sensitivity | | |
| | Resistance | | |
| Evidence levels | NCCN/FDA A: clinical routine | None | |
| | Late trials B: clinical trials | | |
| | Early trials C: case reports | | |
| | Case report D: preclinical | | |
| | Preclinical E: inferential | | |
| Variant specific | Yes | Yes | No |
| References provided | Yes | Yes | No |
| Version | v19 | 1 June 2017 | v3 |
| Source | [20] | [21] | [12] |

Each column summarizes the specificities of each database: GDKD Gene Drug Knowledge Database, CIVIC Clinical Interpretation of Variants in Cancer, TARGET Tumor Alterations Relevant for Genomics-driven Therapy.
and Meric-Bernstam et al. [22] and manually added some expert rules. The complete list of 312 actionable genes can be found in Additional file 1.

Datasets

**TCGA and GENIE datasets**

The Pan Cancer 12 data freeze of TCGA was used as a patient cohort for testing our reporting method. Three data types were downloaded from the Synapse repository: clinical data (syn2325436), somatic SNVs (syn1729383), and somatic CNVs (syn1711454). Data on a total of 5277 samples from 12 different cancer types were collected. Only samples with both mutation and copy number data (3184 samples) were considered for the analyses in this paper. Regarding mutation data, “silent” variants were not studied. For CNVs, the output from GISTIC 2.0 all_thresholded_by_genes was used. This file contains the copy number data of genes discretized into values of the set \{-2, -1, 0, 1, 2\}, where 0 means no deletion or amplification, +/-1 means amplification or deletion above the low noise threshold and +/-2 are amplifications and deletions above the high level threshold. High level thresholds are calculated on a sample basis and are an approximation to homozygous events. Only high level amplifications and deep losses (+/-2) were considered for the analysis, as previously done [23]. The cancer type abbreviations used throughout the article and number of samples are: BRCA (breast cancer, 756), BLCA (bladder cancer, 97), UCEC (uterine cancer, 244), READ (rectal cancer, 69), COAD (colon cancer, 155), OV (ovarian cancer, 313), LUSC (lung squamous carcinoma, 178), LUAD (lung adenocarcinoma, 172), LAML (acute myeloid leukaemia, 190), KIRC (kidney cancer, 417), HNSC (head and neck cancer, 306), and GBM (glioblastoma multiforme, 287).

The GENIE dataset was downloaded from the Synapse repository (SNVs, syn7851250; CNVs, syn7851245; fusions, syn7851249; clinical data, syn7851246). This dataset comprises data of 18,804 patients with more than 50 different cancer entities [24]. Mutation and copy number data were analyzed in the same way as described for TCGA dataset.

**NCT MASTER dataset**

**Patients**

A proof-of-concept of the method in a clinical context was investigated within a retrospective study. We used the data of 11 patients with advanced tumor diseases who had undergone whole exome and transcriptome sequencing within the so-called NCT MASTER trial, an institutional review board-approved clinical sequencing program for young adults with advanced-stage hematological and oncological diseases across all malignancies. A tumor tissue and a matched normal blood sample for whole-exome sequencing were obtained following written informed consent under an institutional review board-approved protocol.

**Whole-exome sequencing data**

Tissue samples were provided by the NCT Heidelberg Tissue Bank. Whole-exome sequencing of normal and tumor tissue samples was followed by a bioinformatic analysis for detecting SNVs, small insertions and deletions (indels), CNVs, and structural variations that might lead to gene fusions. On average, coverage was 133× and 126× for tumor and normal samples, respectively (sample-wise data can be found in Additional file 2). Reads were mapped to the 1000 Genomes phase 2 assembly of the human reference genome (NCBI build 37.1) using BWA (version 0.6.2) with default parameters and maximum insert size set to 1000 bp [25]. BAM files were sorted with SAMtools (version 0.1.19) [26] and duplicates were marked with Picard tools (version 1.90). For the detection of SNVs, we applied our in-house analysis pipeline based on SAMtools mpileup and bcftools with parameter adjustments to allow the calling of somatic variants with heuristic filtering as previously described [27–29]. We used Platypus [30] version 0.5.2 to identify indels with a similar reliability scoring as for SNVs. All mutations were annotated with ANNOVAR [31] version September 2013 using the RefSeq gene model. From the set of somatic high confidence mutations, we extracted nonsynonymous, stopgain, and stoploss SNVs as well as SNVs at splice sites, and indels that are located in a coding sequence or splice site. CNVs were analyzed by read depth plots and an in-house pipeline using the VarScan2 copy-number and copyCaller modules [32]. Regions were filtered for unmappable genomic stretches and merged by requiring at least 70 markers per called copy number event. We selected regions with a log ratio of tumor coverage over control coverage higher than 0.55 or lower than −0.55 as copy number gains and losses, respectively, and annotated them with RefSeq genes using BEDTools [33]. We searched for structural variants such as translocations that might lead to gene fusions with CREST [34] on the DNA level.

The variant calls were delivered in excel files, and for each patient, a group of expert bioinformaticians and oncologists manually revised the list of somatic alterations looking for actionable alterations that could guide the treatment decision. The genomic somatic calls of the patients, together with the experts’ interpretations, are summarized in Additional file 2.

**Statistical analyses**

We performed an unsupervised clustering using the molecular status of the 312 actionable genes in the 3184 TCGA tumor samples. Four molecular categories were used: wild type, mutated, high-level amplification, and
deep loss. Genes with no mutations or with “silent” variants were considered as wild type. Genes with any other type of mutation were considered as mutated. With respect to CNVs, GISTIC output all_thresholded.by_genes was used; genes with a value of −2 were classified as deep losses and +2 as high-level amplifications. Next, our algorithm to filter variants was applied, and a molecular status matrix (sample x gene) was constructed. Genes without any alteration in any sample were removed. Complete-linkage hierarchical clustering was performed on rows and columns based on the Gower distance metric for nominal data (daisy function from R package cluster). The respective heatmap of all samples and the top 50 genes was plotted with dendrogram on the columns (heatmap.2 from gplots R package).

Results
In this section, we first present the framework to filter, classify, and report actionable genomic alterations to support treatment decisions. We then present the results of applying the tool to three patient cohorts with the aim to: 1) evaluate its feasibility with the TCGA and GENIE cohorts and 2) provide a proof-of-concept of the reports with the NCT MASTER cohort.

Reporting framework
In order to exemplify the steps of the framework (Fig. 1), we will use patient MASTER-04 from the NCT MASTER dataset. This patient presented with metastatic ovarian carcinoma, and, after whole-exome sequencing, the panel of experts of the NCT MASTER suggested everolimus, an mTOR inhibitor, as the best treatment option based on a stopgain mutation in the gene TSC2 (Fig. 2).

Filtering of actionable somatic variants
Two types of information are required as input (Fig. 1a): cancer type (e.g., breast cancer) and the somatic variants. Three kinds of somatic variants can be used as input: SNVs and indels, CNVs, and fusion genes. In patient MASTER-04, 98 missense SNVs, 16 fusion genes, and CNV regions (focal and broad) comprising 3555 genes were identified by the bioinformatic pipeline and used as input for the tool. The method assumes that any quality thresholds are previously applied. Therefore, it assumes that only high quality variants are used as input.

In general, variants are annotated in the databases in two different ways: 1) as hotspot mutations/CNVs/fusions with very well-known therapeutic implications (e.g., T790 M or exon 19 mutations in EGFR, HER2 amplification in breast cancer); or 2) as “any missense mutation” or just “any loss/gain of function” variant in those cases where there is no hotspot mutation known or all variants within a gene are studied (very common in clinical trials and preclinical studies).

To identify actionable variants, the algorithm follows a narrowing down procedure. First, patient variants are queried at gene level. Then, for each altered gene, the patient variant type is matched to database entries (e.g., missense mutations, copy number loss, copy number gain). In case of SNVs, the protein change is further checked. If the patient protein change is present in the database, or if the gene annotation is “any missense mutation”, the variant is retained for further steps. If not, the variant is considered of unknown significance and repurposing rules are applied; a variant of unknown significance will be matched to hotspot mutations on the same gene if all the associations between that gene and the drug on the database are in the same direction (i.e., always in favor of response to a drug, or always indicating resistance). The algorithm will also match a stopgain mutation in a gene “with copy number loss/loss-of-function” entries in the database (Fig. 1b). In patient MASTER-04 we can see an example of such a repurposing rule: TSC2 stopgain mutation (R505X) is identified as a loss-of-function mutation (Fig. 2).

Classification and reporting: evidence-based MTB report
After filtering, the algorithm classifies each variant-drug association according to their evidence. The goal is to include additional options for off-label use that may be applicable for advanced-stage patients who failed all conventional standard therapy. To this end, we have created a six-level system to rank the associations according to their evidence. This system allows a stratification of possible treatments into two axes: strength of clinical evidence (axis 1, 2, 3) and cancer type (axis A, B) (Fig. 1c). Level A means evidence in the same cancer type. Level B means evidence in any other cancer type. On the 1-2-3 axis, level 1 means evidence supported by drug approval organizations or clinical guidelines. Level 2 contains clinical evidence, in which late clinical trials are ranked higher followed by early clinical trials and case reports. Finally, level 3 consists of preclinical evidence.

The output of this algorithm is a report designed as a tool to facilitate genomic-based treatment decisions in MTB (Figs. 1d and 2) by compiling all actionable variants of a patient. The quality of the variants is summarized in two tables (SNVs and CNVs separately). The number of variant-drug associations at each level is depicted in a figure. The results are detailed in a table ranked by 1) drug frequency (number of times the drug appears associated with response), 2) by levels of evidence (A1 > B1 > A2 > B2 > A3 > B3) and 3) by gene. This way of sorting aims at ranking first the therapeutic options with more support, but at the same time with high evidence (in patient MASTER-04, we see that mTOR inhibitor is the treatment with the highest number of results and, hence, it is ranked as the first option).
Also, by grouping the drugs we allow a better visualization of 1) contradicting evidence (resistance and response to the same drug) and 2) different variants supporting the same treatment. To allow a quick interpretation, the type of association (response, resistance) is colored (green, red). The variants of unknown significance are gray.

Scope of the MTB report
To obtain cohort-level information on automatically generated recommendations, we applied our method to 3184 samples of the Pan Cancer 12 dataset from TCGA and to 18,804 samples of GENIE.

Distribution of actionable variants in the TCGA cohort
In order to determine the feasibility and extent of incorporating genomic data into clinical decisions, we first identified the actionable variants of every sample. We performed an unsupervised clustering of the cohort using the molecular status (i.e., wild type, mutated, amplified, deleted) of the variants found by the algorithm (Fig. 3). The clustering did not reveal any histology-specific molecular signature. In other words, we did not observe specific sets of known biomarkers defining cancer types.

Overall, samples with none or few genomic events clustered at the right side of the dendrogram, comprising a subset of acute myeloid leukemia, kidney, and breast cancer samples. On the other extreme of the clustering we found a subset of hypermutated samples of colorectal and endometrial cancers. In between, biomarkers were altered across many histologies in low frequencies rather than being histology-specific. However,
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## PATIENT INFORMATION

| Patient ID  | MASTER-04 | Tissue Type | ? |
|-------------|------------|-------------|---|
| Gender      | Female     | Tumor Content (%) | 20 |
| Disease     | Ovarian carcinoma metastasis | Number SNVs | 98 |
| Previous Therapies | - | Number CNVs | 3555 |
| Tumor Board Decision | TSC2 stopgain: mTOR-Inhibitor (Everolimus) | Number Fusions | 16 |

## GENE-DRUG PREDICTIVE ASSOCIATIONS

**Method:** Somatic variants of the patient (mutations, amplifications, deletions, rearrangements) are searched in curated databases of predictive biomarkers (GRDB®, CIViC®) and reported according to their clinical evidence (see Levels of Evidence). In the following two tables (SNVs and CNVs), basic information of the somatic variants with relevant clinical implications can be found.

### Gene-Drug Predictive Associations (SNVs)

| Gene | Patient’s Variant | Level of Evidence | Seg. Mean | Size (Mb) |
|------|-------------------|-------------------|-----------|-----------|
| AURKA | ampl. | B3 | 0.58 | 39.2 |
| BRCA1 | ampl. | B3 | 0.58 | 39.2 |
| CCNE1 | del. | A2, B2, A3, B3 | -0.84 | 2.6 |
| CDK12 | del. | A1 | -0.87 | 1.9 |
| FANCA | del. | B2 | -0.71 | 59.8 |
| FRS2 | ampl. | B3 | 0.73 | 6.5 |
| MDM2 | ampl. | B2 | 0.73 | 6.5 |
| NF1 | del. | B2, B3 | -0.85 | 2.6 |
| PALB2 | del. | B2, B3 | -0.70 | 17.5 |
| RICTOR | ampl. | B2 | 1.56 | 12.5 |
| SUZ12 | del. | B3 | -0.87 | 4.5 |
| TOP1 | ampl. | B2 | 0.58 | 39.2 |
| TSC2 | R505X | B2, B3 | -0.74 | 11.2 |

**Levels of Evidence:** Findings are classified into 6 levels of evidence combining the axis A-B and the axis 1-2-3. Level A means evidence in the same cancer type. Level B means evidence in any other cancer type. On the 1-2-3 axis, level 1 means evidence supported by drug approval, recommendations or clinical guidelines, level 2 contains clinical evidence (clinical trials, case reports) and level 3 consists of preclinical evidence. The distribution of findings into levels is summarized in the right figure.

**Table of Results:** All the predictive associations are detailed in this table. The results are sorted by 1) drug frequency, 2) levels of evidence (A1-A2,A3-B2-A3-B3). To allow a quick interpretation, the type of association (response, resistance) is colored (green, red) and new variants are gray and underlined.

![Gene-Drug Associations Diagram](image)

**Gene-Drug Associations (CNVs)**

| Gene | Patient’s Variant | Level of Evidence | Drugs | Association | Evidence | PMID |
|------|-------------------|-------------------|-------|-------------|----------|------|
| FANCA | del. | prostate | any (LpF) | response | PARP inhibitors | early trials | 26510020 |
| PALB2 | del. | prostate | any (LpF) | response | PARP inhibitors | early trials | AACR 2015 (abstr CT322), 26510020 |
| BRCA1 | del. | ovarian | del. (LpF) | sensitivity | PARP inhibitors | preclinical | 22342242 |
| CDK12 | del. | ovarian | any (LpF) | sensitivity | PARP inhibitors | preclinical | 2440700 |
| PALB2 | del. | Pancreatic | any (LpF) | sensitivity | PARP inhibitors | preclinical | 22565339 |
| TSC2 | R505X | Angiomyolipoma | any (LoF) | response | mTOR inhibitors | early trials | NCT01858805 |
| RICTOR | ampl. | Lung | ampl. (GpF) | response | mTORC1/2 inhibitors | case reports | 21325172, 20048774 |
| NF1 | del. | Neurosarcoma | any (LoF) | sensitivity | mTOR inhibitors | preclinical | 18483311, 30505189, 24504877 |

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1. Demestmann et al., Cancer Discov (2015), v19
2. Greif et al., Nat Genet (2017), version 01 June 2017

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Fig. 2 (See legend on next page.)
some exceptions could be observed: TP53 and PIK3CA were mutated in ~30–40% of the samples, FGFR3/4 were almost exclusively amplified in a subgroup of breast and squamous malignancies (head and neck and lung squamous cancers), and CDKN2A/B deletions clearly defined a subgroup of glioblastoma samples mixed with lung and head and neck samples. The two latter examples are interesting since clinical trials for these biomarkers exist for breast and glioblastoma tumors, respectively. However, according to these results, FGFR3/4 and CDKN2A/B could be biomarker candidates to be studied on lung and head and neck cancers.
Finally, for this cluster analysis 156 genes (out of 312) were removed due to the lack of alterations in any samples. The fact that half of the biomarkers are not present in a cohort of 3184 samples highlights the low frequency of some genomic events and the problem that this poses for biomarker validation. Of course, the tissue representation of this cohort has also an important impact on the alterations found. All in all, the results support the principle of performing comprehensive molecular characterization in clinical practice instead of only testing a histology-specific panel of biomarkers.

**Impact of genomic-guided treatment options on TCGA cohort**

Following the identification of actionable variants, the variant–drug pairs of each patient were classified into one of the six levels of evidence, as described in the “Reporting framework” section and Fig. 1. The results of this analysis allowed us to evaluate the performance of our method at suggesting treatment options. We identified the fraction of patients that received treatment suggestions at each level of evidence independently (Fig. 4a), and cumulatively from the highest evidence to the lowest (A1 > B1 > A2 > B2 > A3 > B3) (Fig. 4b). We could also determine the number of genes with actionable variants identified at each level (Fig. 4c). Finally, these three measurements are depicted in Fig. 4d for the entire cohort.

The algorithm found actionable variants with supporting evidence for drugs approved for the underlying entity (level A1) in 9.9% of the investigated cancer patients. A1 actionable variants were found in only half of the cancer types, and, among these, two main groups could be differentiated: 1) colon, rectal, and leukemia (> 30% of patients) and 2) lung squamous, lung adenocarcinoma, breast and ovarian cancer (< 15% of patients) (Fig. 4a, b). Of course, there may be further A1 actionable variants in the cohort that we cannot identify at the DNA level (e.g., Her2neu or CD20 protein expression) and because we did not include fusion data in this analysis (e.g., ALK fusion in lung adenocarcinoma).

Nevertheless, the percentage of patients with actionable variants doubled (22.7%) when we considered also level B1—drugs approved for the treatment of other cancer types (Fig. 4d). This rise was due to alterations at a low percentage in genes with proven predictive value in the context of approved anti-cancer drugs (e.g., EGFR, BRAF, ERBB2, BRCA1, BRCA2, RET, ALK) in all cancer types (Additional file 3). Kidney cancer as well as acute myeloid leukemia overall tended to harbor fewer actionable mutations (Fig. 4c), in which the incidence of B1 actionable variants reached less than 10% of patients (Fig. 4a), pointing towards different molecular mechanisms driving these two cancer types. As for breast cancer, there was almost 80% overlap of patients between A1 and B2 levels (Fig. 2b). The reason for it is that they are based on the same alteration: HER2 amplification is an approved biomarker for trastuzumab, lapatinib, pertuzumab in breast cancer (A1) as well as for trastuzumab in gastric cancer (B2).

The most significant increase in the ratio of patients with actionable alterations happened when considering drugs in clinical trials (A2), which reached 64.1% of patients (Fig. 4d). In this context, the number of patients with variants studied in clinical trials was significantly lower in glioblastoma and kidney cancer when compared to other entities (Fig. 4a, A2 level). Clinical trials on other cancer entities (B2) further increased the coverage to 89% of patients. Whereas at the B2 level the increase seems to be correlated to a higher number of different biomarkers included in clinical trials, this is not the case for the A2 level (Fig. 4c, d). Additionally, kidney cancer and acute myeloid leukemia still remained different at the B2 level (43.6 and 38.4%, respectively, vs > 80%; Fig. 4a). Finally, including preclinical evidence (A3, B3) did not have a further impact on the range of patients with tumors harboring actionable alterations (Fig. 4b, d).

We compared our results with previous in silico efforts to characterize the actionable landscape of TCGA cohort (Table 2). These studies used different subsets of TCGA cohort and also different databases to identify actionable variants. For these reasons, the results are not directly comparable. However, reporting lower-evidence biomarkers (i.e., off-label scenarios and on substances in clinical trials) undeniably increased treatment recommendations for cancer patients in all studies. Also, there were clear similarities in the different classification systems used. The study from Dienstmann and colleagues [20] presented the most similar results to ours, most likely because our method uses, among others, their database.

**Impact of genomic-guided treatment options on the GENIE cohort**

The same analysis was performed on 18,804 samples of the GENIE cohort (Additional file 4). This cohort is especially interesting because of a higher proportion of advanced tumors—compared to TCGA—and because of the targeted screening for cancer-relevant genes—panels covering from ~ 50 up to ~ 450 genes. Our method identified A1 actionable variants in 15.3% of the samples. Compared to TCGA cohort, main differences could be observed in non-small cell lung cancer and leukemia in the A1 level. In B1, GENIE cohort showed a clearly higher percentage of patients with actionable alterations (38.4%). This is not surprising since these patients have acquired mutations during the course of the disease (EGFR T790 M) and the fact that this dataset includes fusions (ROSI, ALK in lung cancer, ABL1 in leukemia).
However, the remaining levels of evidence shared similar ranges in both datasets, being even a bit superior in TCGA cohort. This is most likely due to the fact that TCGA covered exome-wide analysis whereas GENIE is restricted to a panel of genes.

Retrospective evaluation of the MTB report on 11 advanced cancer patients
As a proof-of-concept of our reporting method (MTB report), we performed a retrospective study on 11 patients with advanced malignancies. Whole exome sequencing of normal and tumor tissue samples was performed within the NCT MASTER program, a clinical sequencing program including young adult cancer patients, followed by bioinformatic analysis for detecting SNVs, indels, CNVs, and fusions (see “Methods”). For each patient, a group of experts (bioinformaticians and oncologists) manually revised the list of somatic alterations looking for variants that could guide the treatment decision. The selected actionable alterations were discussed in the NCT MASTER program molecular tumor board. In case of many variants in the same

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![Fig. 4 Heatmap representation of the distribution of identified actionable variants in TCGA cohort. Somatic alterations of each sample (3184 samples) were analyzed as depicted in Fig. 1 and the resulting biomarker–drug associations were assigned to one of the six levels of evidence. Evidence in wild-type variants and resistances are not included in this representation (unless if the evidence is in level A1, e.g., NRAS, KRAS wild type in colorectal cancer).](image)
tumor, only the one to three most relevant findings were discussed. We have compared the alterations discussed by the experts in the NCT MASTER program (MASTER report) with the results reported by our method (MTB report). The 11 MTB reports generated with our tool can be found in Additional file 5.

The MTB reports showed a very good performance in terms of sensitivity in the detection of actionable variants: in 10 out of 11 patients, mostly all variant–drug associations in the MASTER report matched our MTB report (Table 3). Overall, only four genes suggested in the MASTER reports were not detected by our tool: \textit{PTPRJ}, \textit{PTPN12}, \textit{LCK}, and \textit{NTRK3}. The first three are not present in any of the databases used by our method. As for \textit{NTRK3}, it is present in GDKD as a gene fusion biomarker. However, the patient presented a missense mutation in this gene, and our tool does not allow repurposing of fusions into missense mutations to avoid too many false positive results.

The majority of actionable variants in the MASTER reports were found in low evidence levels of the MTB report, mostly B2 and B3 levels. This agrees with the distribution of actionable events in TCGA cohort. Furthermore, the patients enrolled in this program are advanced cancer patients, meaning that they were heavily pretreated and did not have standard therapies left; hence, we would not expect A1 findings.

The actionable variants reported in the MTB report were redundant in terms of gene–drug associations; yet, they provided unique information regarding cancer type or clinical evidence (A, B, C). Indeed, the more times a drug was reported, the more evidence supported the eligibility of that drug (see “Drug support” percentages in Table 3). However, mutations in controversial genes and pathways may also generate conflicting results (e.g., \textit{KRAS} mutation, see report MASTER-02 in Additional file 5), demonstrating the need for a multidisciplinary discussion of NGS data in the clinical context within MTBs.

Alterations of DNA repair pathway genes (\textit{BRCA1}, \textit{BRCA2}, \textit{RAD51}, \textit{PALB2}, \textit{CDK12}) predicted response to PARP inhibitors in several MTB reports. Based on the concept of synthetic lethality upon an initial disruption on DNA repair, MASTER reports included platin-based chemotherapies whenever this pathway was mutated. Therefore, these two therapies were considered equivalents and assessed as a “match” in Table 3.

### Discussion

Despite rapid advances in high-throughput screening technologies, only a handful of predictive biomarkers are routinely tested in TB to guide treatment decisions (e.g., \textit{EGFR} in non-small cell lung cancer or \textit{KRAS} in colorectal cancer). Some clinical institutions are now starting to implement panel sequencing for some tumor entities (e.g., lung cancer and myelodysplastic diseases). Moreover, several clinical studies have anticipated the feasibility of using NGS for treatment decisions, although its impact on patient outcome is still under discussion [8–11, 13, 16]. However, the high dimensionality of NGS data together with the growing evidence of biomarkers for targeted therapies makes the task of identifying actionable somatic variants time-consuming. In this work, we present a tool for matching tumor genomes to treatment options. In brief, this tool filters actionable somatic variants and delivers the results in evidence-based reports. This filtered list of actionable variants is aimed to assist bioinformaticians and oncologists in their task of interpreting somatic variants prior to MTB discussions. In our analysis, the presented framework was able to identify potentially actionable variants in 94% of cancer patients according to TCGA cohort, and, most importantly, it was able to include experts’ treatment suggestions in a retrospective analysis.

Remarkably, the application of the report strategy on TCGA dataset successfully found actionable variants in 9.9% of patients considering only approved drugs (A1), increasing to 22.7% when considering \textit{off-label} use (B1), and to 89% with evidence from clinical trials (B2). These results were shown to be in line with previous in silico efforts to characterize the therapeutic landscape of

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**Table 2** TCGA actionable landscape in different publications

|                  | Standard therapy | Clinical trials | Preclinical | Total | Number of TCGA samples | Databases used by the study |
|------------------|------------------|-----------------|-------------|-------|------------------------|-----------------------------|
|                  | Label            | Off-label       | Label       | Off-label | Label       | Off-label |                   |                               |
| MTB              | 9.9 (A1)         | 22.7 (B1)       | 64.1 (A2)   | 89 (B2)  | 90.6 (A3)   | 94.1 (B3) | 94                  | 3184 GDKD, CIViC, TARGET      |
| Dienstmann et al. 2015 [20] | 11 (5)           | –               | 39 (4)      | 75 (3)   | 93 (1-2)    | –          | 93                  | 4392 GDKD                     |
| Rubio-Perez et al. 2015 [19] | 5.9              | 40.2            | 73.3        | –        | –          | 73.3       | 4068                | Rubio-Perez et al. 2015       |
| Chakravarty et al. 2017 [18] | 7.5 (1–2A)       | 16 (2B)         | 26 (3A)     | 41 (3B)  | –          | –          | 41                  | 5983 OncoKB                    |

The table shows the cumulative percentages of patients with actionable variants identified at different levels. The name of the levels used in each publication are specified in parentheses. The studies being compared are: MTB (this publication), Chakravarty et al. 2017 [18], Dienstmann et al. 2015 [20], and Rubio-Perez et al. 2015 [19].
public datasets [18–20, 35]. Even though these studies slightly differ in the percentages, the overall message remains the same: reporting lower-evidence variants (i.e., off-label scenarios and on substances in clinical trials) undeniably increases treatment recommendations for cancer patients. Nevertheless, these results do not mean that 89% of patients can be treated based on a genomic rationale, but rather highlights a trend of increasing opportunities for personalized cancer treatment.

Recent prospective trials using NGS to guide treatment decisions have reported actionable variants in more than 80% of patients (in the most inclusive sense of the word, as used in the current study), or around 50% of patients (attending to a narrower definition, defined as actionable variants for approved drugs and accessible clinical trials) [10–16, 35]. Altogether these studies confirmed the extent of reporting methods; unfortunately, they also showed a gap between identifying actionable alterations and the actual numbers of treated patients (5–13%). The inability to prescribe treatments based on NGS results was in general due to the impossibility to prescribe drugs in off-label scenarios, restricted access to clinical trials, and clinical deterioration. Whereas continuous efforts in molecular characterization of tumor diseases and successful targeted treatment approaches keep raising the expectations on precision medicine, the effect of individual NGS-based treatment decisions on patient outcome is still under discussion in several open clinical trials (SHIVA02, IMPACT, MOSCATO, NCI-MATCH, NCT MASTER).

In the same line, it is important to emphasize that clinical interpretation of somatic variants is subject to many challenges, comprehensively reviewed by [2, 5, 22]. Among them, determining the clinical relevance of a biomarker–drug association is one of the main difficulties. A common approach is to classify the findings into categories or tiers, what we refer to here as "levels of evidence". Previously proposed classification systems

| New ID | Cancer | Number of SNVs | Number of CNVs | MASTER report | MTB report |
|--------|--------|----------------|----------------|---------------|------------|
|        |        | Gene           | Drug           | Number of results | Match level | Drug support |
| MASTER-01 | Breast cancer metastasis | 104 | 3410 | BRCA1/2 deletions, RAF1, PDXGFR amplifications, FGFR1 (T88N) | PARP inhibitors, Sorafenib, FGFR inhibitor | 43 | Yes | B2 | 11.6% |
| MASTER-02 | Pancreatic adenocarcinoma | 49 | 1528 | KRAS (G12D) | – | 92 | – | – | – |
| MASTER-03 | Leiomyosarcoma of the retroperitoneum | 31 | 3181 | PTPRJ deletion, CDK12 + BRCA2 deletions | Pazopanib, Cisplatin | 12 | No | – | – |
| MASTER-04 | Ovarian carcinoma | 98 | 3385 | TSC2 (R505X) | mTOR inhibitor | 56 | Yes | B2 | 12.5% |
| MASTER-05 | Myxoid liposarcoma | 11 | 72 | PIK3CA (C420R) + PTEN (R130G) | mTOR inhibitor | 75 | Yes | B2 | 9.3% |
| MASTER-06 | Neuroendocrine tumor | 2703 | 2060 | MTOR (P2490L, G332R), PTPN12 (S509 N, G523S), KIT (A837T), LCK (P74L) | mTOR inhibitor, Lapatinib, erlotinib, imatinib, desatinib | 89 | Yes | B2 | 16.8% |
| MASTER-07 | Neuroendocrine tumor | 645 | 941 | ERBB3 (V104 M), RAF1 (S259P), MTOR (E1485G) | mTOR inhibitor | 40 | Yes | A2 | 12.5% |
| MASTER-08 | Cholangiocarcinoma | 28 | 1001 | ERRFI1 (R199X) | Erlotinib | 17 | Yes | – | 0.58% |
| MASTER-09 | Clear cell sarcoma | 11 | 1 | NTRK2 (R116W) | Lestaustinib, midostaurin | 3 | No | – | – |
| MASTER-10 | Histiocytic sarcoma | 7 | 5 | BRAF (F595 L) + HRAS (Q61R) | MEK inhibitor, sorafenib (multi TKi) | 11 | Yes | B2 | 33.3% |
| MASTER-11 | Pulmonary adenocarcinoma | 70 | 146 | EGFR (p.745-750del) | Erlotinib | 54 | Yes | B2 | 22.2% |
have between three and six levels, and consider informative variables such as validation stage of the drug–biomarker predictive association (FDA approved, clinical trials, case studies, preclinical) [10, 12, 22], histology type [12, 22, 36], gene variant [36], and biological significance [10, 12]. Our six levels of evidence system emphasizes the validation stage of the biomarker–drug predictive association (axis 1–3) because drug availability has been identified as one of the main reasons of the low impact of genomic-driven cancer therapies [10, 11]. In addition, it accounts for histology type (axis A–B) to better inform off-label use. That being said, we do not aim to substitute the other classifications, but rather to ease the assessment of the variants in the reports. In a broader perspective, the same is applicable to the main goal of the MTB reports. They aim at reducing the work load that represents searching for actionable variants rather than suggesting just the best treatment option: this is a task for clinicians. We designed our method to include the maximum knowledge important for clinical decisions, but keeping the presentation as simple and compressed as possible. As a consequence, one should be very careful with all recommendations since, although backed by literature, many actionable variants can be far from real clinical practice.

The retrospective analysis established a proof-of-concept of the reports, but also shed light on ways to improve upon the framework. For instance, our reports did not capture indirect targeting unless they were present in the databases, which led to an omission of some experts’ suggestions. For that, we believe it is important to provide the possibility of visualizing somatic alterations in their pathway context. Also, our reports missed target–drug associations such as LCK–dasatinib because LCK mutations have not been shown to predict drug response. This could be improved by including drugs and their targets in the aforementioned pathway visualization. Nevertheless, as we state throughout this article, inferring causality between target gene and drug response is not straightforward. This is the reason why we have so far focused the tool only towards variants with proven evidence of drug response, yet this evidence is not always strong enough and needs to be assessed by a clinician with regard to the clinical course of each patient. For all these reasons, it is important to stress that the reports are aimed at reducing the time-consuming task of linking genomic variants to clinical evidence but they always need a reevaluation by a clinician. Finally, other interesting information that could be included are germline variants. Besides non-cancer syndromes and pharmacodynamic responses, these variants are being demonstrated also to have therapeutic utility in some cases, such as in prostate cancer and childhood leukemias [37, 38].

Taken together, the results of this study have further strengthened the hypothesis that an automated reporting of potential therapeutic options is crucial to make the clinical interpretation of NGS a task with a clinically acceptable turn-around time. We have shown that our method is able to reach a large number of cancer patients. Also, we have demonstrated the reproducibility of experts’ suggestions. This work establishes a proof-of-concept of MTB reports, which provide a structured picture of the actionable variants of a tumor with the already pre-filtered information necessary for deeper treatment discussions.

Conclusions

We have shown here a method that relies entirely on public knowledge, hence encouraging transparency and joint public efforts for a curated database of actionable variants [3]. We strictly report evidence-based variants, so that the clinician can easily verify with reference the causality between variant and drug response. We encourage the design and enrollment of patients into biomarker-driven clinical trials in order to generate more evidence. Nevertheless, this effort must be accompanied by a systematic recording of treatment decisions and outcomes of patients, also of single case studies [39]. We believe that if NGS is undeniably entering the clinical context, a standardized method to annotate public knowledge of actionable variants is crucial to accomplish a successful and appropriate use of the data.

Additional files

| Additional file 1: | List of predictive biomarkers. (XLSX 14 kb) |
| Additional file 2: | Summary of NCT MASTER cohort. (XLSX 334 kb) |
| Additional file 3: | Percentage of patients of the top ten biomarkers at each level of evidence in each tumor type. Level 2 is divided into three further groups: 2a (late clinical trials), 2b (early clinical trials), and 2c (case reports). (PDF 43 kb) |
| Additional file 4: | Heatmap representation of the distribution of identified predictive biomarkers in the GENIE cohort. (EPS 87 kb) |
| Additional file 5: | Eleven reports of NCT MASTER patients. (PDF 175 kb) |
| Additional file 6: | R code of the presented method. The user can generate MTB reports for TCGA samples. (RAR 237 kb) |

Abbreviations

CIVC: Clinical Interpretation of Variants in Cancer database; CNV: Copy number variation; FDA: The Food and Drug Administration; GDKD: Gene Drug Knowledge Database; GENIE: Genomics Evidence Neoplasia Information Exchange; MAST: Molecularly Aided Stratification for Tumor Eradication Research; MTB: Molecular tumor board; NCCN: National Comprehensive Cancer Network; NCT: Nationale Centrum för Tumorerkrankungen; NGS: Next-generation sequencing; PM: Precision medicine; SNV: Single nucleotide variation; TARGET: Tumor Alterations Relevant for Genomics-driven Therapy Database; TB: Tumor board; TCGA: The Cancer Genome Atlas

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Availability of data and materials
TCGA dataset analyzed during the current study is available in the Synapse repository. The NCT MASTER dataset is included in this published article (Additional file 2). The R code of the method presented in this article is provided in Additional file 6 and available in https://github.com/perera-bel/MITB-Report.

Authors’ contributions
JPB and TB designed the study and wrote the manuscript. BB provided major contributions to the study design. JPB performed TCGA data analysis. BH accomplished the bioinformatic analysis of the NCT MASTER dataset. MF, CH, AB, SF, and HG provided important clinical insights. All authors contributed at all stages and critically reviewed the content. All authors read and approved the final manuscript.

Ethics approval and consent to participate
All patient data were obtained following written informed consent under the NCT MASTER trial, an institutional review board-approved clinical sequencing program for younger patients with advanced-stage hematological and oncological diseases across all malignancies. The trial protocol was approved by the Ethics Committee of Heidelberg University. Research within NCT MASTER was conducted in accordance with the Declaration of Helsinki.

Consent for publication
Written informed consent for publication was obtained under the NCT MASTER trial.

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

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