Evaluating the analytical validity of circulating tumor DNA sequencing assays for precision oncology

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Circulating tumor DNA (ctDNA) sequencing is being rapidly adopted in precision oncology, but the accuracy, sensitivity and reproducibility of ctDNA assays is poorly understood. Here we report the findings of a multi-site, cross-platform evaluation of the analytical performance of five industry-leading ctDNA assays. We evaluated each stage of the ctDNA sequencing workflow with simulations, synthetic DNA spike-in experiments and proficiency testing on standardized, cell-line-derived reference samples. Above 0.5% variant allele frequency, ctDNA mutations were detected with high sensitivity, precision and reproducibility by all five assays, whereas, below this limit, detection became unreliable and varied widely between assays, especially when input material was limited. Missed mutations (false negatives) were more common than erroneous candidates (false positives), indicating that the reliable sampling of rare ctDNA fragments is the key challenge for ctDNA assays. This comprehensive evaluation of the analytical performance of ctDNA assays serves to inform best practice guidelines and provides a resource for precision oncology.

Cancer cells undergoing apoptosis or necrosis release fragments of DNA into the circulatory system1-4. These ctDNA fragments might harbor somatic mutations from their tumor of origin, and their abundance correlates with tumor size and stage5-8. Accordingly, ctDNA can act as an accessible biomarker to inform cancer detection, molecular stratification, therapeutic monitoring and post-treatment surveillance9-11.

Assays that measure ctDNA have several advantages over tumor tissue biopsies (Supplementary Table 1). The collection of ctDNA is fast, cheap and minimally invasive and can be performed serially to monitor tumor evolution or response to therapy. In theory, ctDNA can provide a representative cross-section of heterogeneous tumors and multi-focal disease. Moreover, although a tumor tissue biopsy cannot be performed without prior knowledge of the tumor, ctDNA assays can identify evidence of unknown lesions, thereby enabling detection of minimal residual disease (MRD) after treatment or even cancer screening in healthy populations5-9.

These advantages are best realized via the unbiased analysis of ctDNA by next-generation sequencing (NGS), and, on this basis, ctDNA sequencing is being rapidly adopted in precision oncology. However, ctDNA sequencing assays face major technical challenges. Cell-free DNA exists as small fragments (~160 base pairs (bp)) at low concentrations (typically <10 ng or <3,000 genome copies per ml of plasma in patients with cancer)10. Furthermore, only a small fraction of cell-free DNA is tumor derived (commonly <1% of alleles in circulation but sometimes as low as <0.01%)10. The detection

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NATURE BIOTECHNOLOGY | VOL 39 | SEPTEMBER 2021 | 1115-1128 | www.nature.com/naturebiotechnology
of rare somatic mutations from such limited input material is highly challenging.

ctDNA sequencing assays are also affected by a range of experimental variables and artifacts. Extensive polymerase chain reaction (PCR) amplification is typically required to generate an NGS library from the small quantity of cell-free DNA available, as well as further targeted enrichment of informative cancer genes (by hybrid capture or amplicon methods)\(^9\). The small size of cell-free DNA fragments can inhibit target enrichment, reduce alignability to the human reference genome and prevent the resolution of complex loci or mutations. These variables further exacerbate the quantitative biases and sequencing errors that affect all NGS experiments\(^1\).

Despite these challenges, ctDNA assays of increasing resolution have been developed\(^1\). With clinical adoption already underway, it is critical for the community to understand the sensitivity, accuracy and reproducibility of ctDNA assays and the variables that affect analytical performance. Discordant results between alternative assays or parallel ctDNA and tumor biopsy tests have been reported\(^20\). Accordingly, a joint review by the American Society of Clinical Oncology and the College of American Pathologists recently identified the pressing need for proficiency testing using standardized samples to assess the analytical validity of ctDNA assays and enable unbiased comparisons among different technology platforms and laboratories\(^2\).

Here we report the findings of a multi-lab, cross-platform evaluation of analytical performance among NGS-based ctDNA assays carried out as part of the Food and Drug Administration (FDA)-led Sequencing Quality Control Phase 2 (SEQC2) project or the fourth phase of the MicroArray Quality Control consortia. The Oncopanel Sequencing Working Group—comprising academic, industry, government and regulatory stakeholders—tested the performance of five leading ctDNA assays across 12 participating clinical and research facilities. We employed simulated and synthetic experiments, as well as rigorous proficiency testing on contrived human ctDNA reference materials, to measure the effect of variables at each step within the ctDNA sequencing workflow. The study assesses the analytical validity of ctDNA assays for potential clinical applications and informs best practice guidelines (Table 1).

**Results**

**Evaluating ctDNA assays with simulated sequencing data.** The detection of ctDNA fragments occurs by random sampling from a background of non-cancerous cell-free DNA. To investigate the analytical variables that govern this process, in the absence of confounding experimental variables, we initially generated simulated NGS libraries that emulate targeted analysis of cell-free DNA by hybrid capture sequencing (Methods).

We created simulated libraries from 155 cancer genes, covered at ~9,000-fold depth by ~160-bp sequence fragments (Supplementary Fig. 1a). This level of coverage can be theoretically obtained from a routine patient blood draw, given that cell-free DNA occurs at up to ~3,000 genome copies per ml of plasma (Supplementary Table 2). The cancer genes harbored 2,356 simulated somatic mutations (one COSMIC single nucleotide variant (SNV) per exon) that were represented at known variant allele frequencies (VAFs) ranging from 5% to 0.1% (Supplementary Fig. 1a). We then performed an in silico hybrid capture enrichment step to obtain typical convex coverage profiles over targeted exons, resulting in a final 8,252-fold median fragment depth at mutation sites (Supplementary Fig. 1a,b). An example is shown for the oncogene MET (Fig. 1a).

We used these simulated libraries to evaluate the effect of coverage on the detection of ctDNA mutations. Due to random sampling, the number of sequence fragments containing a given mutation follows a Poisson distribution, with a median fragment count that is proportional to the product of VAF and global fragment depth (Supplementary Fig. 1c). At maximum depth, more than 99% of mutations were detected by at least two independent fragments (Supplementary Fig. 1d). However, any decrease in coverage or increase in detection stringency (that is, more than two supporting fragments) caused a reduction in sensitivity.

To model these relationships, we incrementally adjusted the simulated alignment coverage and detection stringency (Methods). For low-frequency mutations (VAF < 0.5%), coverage had a pronounced effect on detection sensitivity, with this relationship modeled by a sigmoidal function (Fig. 1b and Supplementary Fig. 1d). The stringency imposed during mutation detection similarly affected sensitivity for low-frequency mutations (Fig. 1c and Supplementary Fig. 1d). By contrast, mutations at higher frequencies (VAF > 0.5%) were detected with maximum sensitivity even at relatively low fragment depth and high stringencies (Fig. 1b,c and Supplementary Fig. 1d). These analyses illustrate the inherent challenge of reliably detecting low-frequency ctDNA mutations by random sampling.

The enrichment of DNA fragments by hybrid capture results in heterogeneous coverage across targeted exons. Even in the absence of hybridization biases, we found that the detection of ctDNA mutations was similarly heterogeneous: because mutations in the edge regions of exons had lower fragment depth than central mutations, detection sensitivity was up to 10% lower among edge mutations (Fig. 1d and Supplementary Fig. 1e,f). Given that many pathogenic mutations occur within exon edge regions, especially at splice site positions where coverage is lowest\(^7\), this edge effect is a relevant consideration when designing hybrid capture panels for ctDNA sequencing.

The short fragment length of cell-free DNA can cause erroneous or ambiguous alignment to the human reference genome. We found that ~5% of exons analyzed (118/2,356) had suboptimal alignability, with these exons exhibiting lower fragment depth and the mutations they harbored being detected with reduced sensitivity (Fig. 1e and Supplementary Fig. 1f). This effect hindered the detection of mutations in notable gene families, such as the RAS family (KRAS, NRAS and HRAS).\(^7\) The effect of local alignability and exon position was most pronounced when evaluating low-VAF mutations and when depth or stringency was limiting (Fig. 1d,e). These results demonstrate that, even in the absence of experimental variables, genomic context can have an influence on the detection of ctDNA mutations.

**Evaluating ctDNA assays using synthetic DNA controls (sequins).**

We next evaluated the detection of ctDNA mutations using synthetic DNA controls known as ‘sequins’. Sequins are synthetic DNA sequences that emulate natural human genes and mutations and recapitulate many of the technical biases that affect their analysis by NGS\(^11\) (Fig. 2a).

We assembled a mixture of sequin controls representing 134 recurrent and/or clinically actionable somatic mutations within the functional domains of 87 cancer-related genes (Supplementary Data 1). Sequins representing wild-type and mutant alleles for each gene were combined in precise ratios to form a staggered reference ladder spanning a wide range of VAF levels (from 0.1% to 100%; Fig. 2b). The sequin mixture was fragmented and size selected to emulate cell-free DNA fragments (Supplementary Fig. 2a) and then added at ~0.2% fractional abundance to human mock cell-free DNA samples (described below). These combined samples were then analyzed by targeted hybrid capture sequencing (Methods). In total, 119 kb of synthetic sequence was captured and analyzed.

We evaluated the detection of synthetic mutations encoded by the sequin mixture. After calibrating sequins to match the coverage of their accompanying human sample (6,311-fold median fragment depth; Supplementary Fig. 2b), 125/134 synthetic mutations were detected (sensitivity = 0.93). To assess the effect of coverage on sensitivity, we repeated variant detection across a range of downsampld libraries. Decreasing coverage had a strong effect on the detection of low-frequency mutations (VAF < 0.5%), whereas...
In hybridization kinetics, PCR amplification and library conversion

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**Table 1 | Summary of concepts, findings and recommendations**

| Issue                              | Findings                                                                 | Outlook & recommendations                                                                 |
|------------------------------------|---------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|
| Mutation frequency                 | Mutations present above ~0.5% VAF were detected with high sensitivity and reproducibility by all participating ctDNA assays, but performances were generally suboptimal below this level and variable between assays (Fig. 4). | A key challenge in ongoing development of ctDNA sequencing assays is to improve detection sensitivity for low-frequency mutations (VAF < 0.5%). |
| Coverage depth and heterogeneity   | Fragment depth was a critical variable in ctDNA assays, with high coverage essential for sensitive detection of low-frequency mutations (Figs. 3 and 4). In addition to depth, even coverage across target regions was important to ensure high sensitivity and reproducibility (Supplementary Fig. 3). | Improvements to the efficiency and stability of capture enrichment, NGS library conversion and amplification might yield increased coverage depth and decreased heterogeneity, leading to improved performance and robustness. |
| DNA input quantity                 | Increasing DNA input quantity generally improved fragment depth, sensitivity and reproducibility (Fig. 5).                                                                 | Limited availability of cell-free DNA is a challenge for clinical translation of ctDNA assays. Input material might be increased by improvements to the efficiency of plasma DNA extractions, increasing the volume of patient blood draws (when feasible) or obtaining cell-free DNA from other body fluids (for example, urine, stool and cerebrospinal fluid) for relevant cancers. |
| Unique molecular identifiers (UMIs) | UMIs enabled effective consensus error correction, minimizing the detection of false positives (Supplementary Table 5). | Wherever possible, UMIs should be employed for consensus error correction in ctDNA sequencing assays. |
| Inter-laboratory variation         | Participating assays were robust to technical variables among test labs—from plasma extraction to sequencing workflow stages—and were affected largely by random, rather than systematic, variation (Fig. 4 and Supplementary Fig. 6). | Robustness to technical variables is essential for clinical implementation of ctDNA sequencing assays. |
| Random sampling                    | The detection of low-frequency mutations (VAF < 0.5%) by random sampling poses an inherent statistical challenge, even when high fragment depth is available (Fig. 1). | Novel strategies for the enrichment of ctDNA fragments over non-cancerous cell-free DNA (for example, by fragment size selection) and alternative signals, such as ctDNA methylation profiles, might help overcome limitations of random sampling. |
| Targeted enrichment method         | Performance was broadly similar among participating amplicon and hybrid capture assays, with sensitivity and robustness largely determined by the fragment depth achieved, not the method of enrichment (Fig. 6). | Amplicon methods can enable sensitive, cost-effective detection of ctDNA mutations in single genes or mutation hotspots, but small panel sizes limit their suitability for unbiased surveillance (for example, for tumor evolution profiling). |
| Exon edge effect                   | In hybrid capture sequencing, mutations in exon edge regions were detected with lower sensitivity than central regions, due to lower coverage (Figs. 1 and 2). | Increasing the size of captured flanking regions around exons during panel design might alleviate this exon edge effect. |
| Sequence context                   | Mutations in challenging genome sequence contexts, such as high/low GC content, low-sequence complexity or suboptimal alignability, were detected with lower sensitivity (Figs. 1 and 2). | Some of these effects might be alleviated by increasing capture probe density in challenging regions or by improvements to NGS library preparations. |
| Reference standards                | Reproducibility measurements provided a useful but imperfect proxy for analytical performance that is not dependent on the availability of a reference sample and annotation (Fig. 4). | Well-characterized reference standards can directly measure analytic performance characteristics in the absence of confounding biological variables and are a useful tool for comparing ctDNA assays. |

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mutations at intermediate (0.5–5%) and high (>5%) frequencies were detected with high sensitivity even at low fragment depths (0.96–1.00; Fig. 2c). We observed little difference in sensitivity for SNVs and small insertions and deletions (1–16 bp; Supplementary Fig. 2c), suggesting that variant frequency had a larger effect than variant type.

In addition to global coverage depth, performance is also influenced by coverage heterogeneity, resulting from regional variation in hybridization kinetics, PCR amplification and library conversion efficiency. Sequin mutations ranged over 300-fold in fragment depth (12,080-fold to 36-fold), with sequin coverage profiles closely resembling corresponding genes in their accompanying human sample ($r^2 = 0.89$; Fig. 2a and Supplementary Fig. 2d,e). Such heterogeneity had a strong effect on variant detection: mutations in regions of high coverage (>5,000-fold) were detected with up to 30% higher sensitivity than regions of low coverage (<3,000-fold) (Fig. 2d).

To elucidate the underlying determinants of this heterogeneity, synthetic variants were stratified according to genomic context.
two-fold magnitude changes in ctDNA abundance could be accurately measured in patient plasma. At maximum fragment depth, in addition to detecting ctDNA mutations, ctDNA assays can sample (Sample A; Fig. 3a). This pooled sample, as well as each individual cell line, was genotyped to establish a set of ~40,000 ‘known mutations’ and ~10.2 Mb of ‘known negatives’ (positions that matched the human reference genome in every cell line) within the exonic coding regions. Together, these form a reference annotation against which diagnostic performance was subsequently evaluated.

To emulate the range of VAFs typically encountered in ctDNA assays, Sample A was combined at known ratios with DNA extracted from a non-cancer background cell line (Sample B) to create two further reference samples: Lbx-high (20% A/ 80% B) and Lbx-low (4% A/ 96% B) (Fig. 3a). These samples were enzymatically sheared and size selected to form DNA fragment size distributions of ~160–180 bp (Methods).

Changes observed in the abundance of ctDNA fragments in patient plasma might indicate tumor progression, response or resistance to therapy or disease relapse9. It is, therefore, essential that, in addition to detecting ctDNA mutations, ctDNA assays can accurately measure their frequency. At maximum fragment depth, two-fold magnitude changes in ctDNA abundance could be reliably resolved down to a VAF of 0.8% but were inaccurate below this level (Fig. 2b). Resolution was also affected by fragment depth, with decreasing coverage eroding the lower limit of quantitative accuracy (Supplementary Fig. 2g). This demonstrates the difficulty of accurately identifying changes in the abundance of low-frequency ctDNA mutations during patient treatment.

Multi-site, cross-platform proficiency study. We next undertook a large-scale proficiency study using a set of contrived reference DNA samples that are described in detail in a companion article14. Briefly, genomic DNA extracted from ten diverse human cancer cell lines was pooled at equal abundance to create a mock cancer sample (Sample A; Fig. 3a). This pooled sample, as well as each individual cell line, was genotyped to establish a set of ~40,000 ‘known variants’ and ~10.2 Mb of ‘known negatives’ (positions that matched the human reference genome in every cell line) within the exonic coding regions. Together, these form a reference annotation against which diagnostic performance was subsequently evaluated.

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These mock cell-free DNA samples were administered to 12 independent laboratory sites across the United States, United Kingdom, China and Australia (Fig. 3a). Each laboratory performed one or more participating ctDNA sequencing assays (Supplementary Table 3), which included hybrid capture assays from Roche Sequencing Solutions (ROC), Illumina (ILM), Integrated DNA Technologies (IDT) and Burning Rock Dx (BRP) and an amplicon sequencing panel from Thermo Fisher Scientific (TFS). All sequencing was performed using Illumina instruments (NovaSeq or NextSeq), with the exception of the TFS amplicon assay, which was sequenced using Thermo Fisher Scientific’s Ion Torrent instrument (Fig. 3b).

Each participating assay was performed at 2–3 independent test labs, with four technical replicates per lab for each mock ctDNA sample, at a fixed DNA input amount (25 ng). In addition, Lbx-low was analyzed with increased (50 ng) and decreased (10 ng) input amounts to investigate the effect of cell-free DNA input quantity.

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Fig. 2 | Evaluating ctDNA assays with sequins. a, Genome browser view showing fragment coverage within a synthetic sequin control (upper) representing the oncogene FGFR3, harboring multiple synthetic mutations at VAF = 50%. For comparison, coverage is also shown within the natural FGFR3 gene (lower) obtained from the accompanying human sample. b, Scatter box plots show observed versus expected VAFs for synthetic sequin mutations (n=134), which are represented in two-fold VAF increments from 0.1% to 100%. Asterisks indicate significant differences in measured VAFs between increments (two-sided t-test, P < 0.001; n > 8 data points per bin). Boxes show median ± range (whiskers) and IQR (boxes). Colored lines indicate high, mid and low VAF groups used in c–g. Curves modeling the relationship between library depth (median fragment depth) and detection sensitivity for synthetic sequin mutations under various conditions: c shows mutations within different VAF groups, indicated on the lower axis of b; d shows mutations with high fragment depth (>5,000-fold) compared to low fragment depth (<3,000-fold); e shows mutations within exon edge regions (≤20 bp from exon boundary) compared to central regions (>50 bp from exon boundary); f shows mutations in regions of high or low GC content (<40%/>60%) compared to moderate regions; g shows mutations in regions of low sequence entropy (<1.9) compared to typical regions.

To assess the effect of technical variables during plasma DNA extraction, Lbx-low was also analyzed after extraction from a synthetic plasma solution, with extractions performed independently at each test lab (Fig. 3b).

Each sequencing library was then analyzed by the relevant ctDNA assay vendor. Bioinformatic analysis was not standardized across the study, with each vendor, instead, employing an internal analysis pipeline and providing a final set of variant candidates for centralized evaluation by an independent team (Fig. 3a). Together, the proficiency study encompassed 360 ctDNA assays and, to our knowledge, constitutes the most comprehensive evaluation of analytical performance in ctDNA sequencing to date (Supplementary Data 2).

Coverage depth and heterogeneity. We evaluated coverage depth, which is considered a key variable in ctDNA sequencing10. We observed substantial differences in coverage among different assays, with median unique fragment depth ranging from ~4,700-fold (BRP and ROC) to ~1,200-fold (ILM) at 25 ng input (Fig. 3c). Given that DNA input quantities were standardized, these differences reflect the capacity of each assay to exhaustively profile the unique DNA fragments within the input sample and might have a relevant effect on assay performance. Assuming that 25 ng input equates to ~7,500 genome equivalent copies, estimated molecular recovery rates ranged from ~63% in ROC to 17% in ILM (Fig. 3c and Supplementary Tables 2 and 4). The TFS amplicon assay achieved similar fragment depth to participating hybrid capture assays (Fig. 3c).

As observed above, multiple technical variables can result in uneven coverage across target regions. After normalizing for overall depth, assays were distinguished by clear differences in coverage heterogeneity (Supplementary Fig. 3a,b). For example, although the BRP and ROC assays achieved similar median fragment depths,
Fig. 3 | Structure of cross-platform ctDNA sequencing proficiency study. a, Schematic overview of the proficiency study. Briefly, contrived mock cell-free DNA samples (Lbx-high and Lbx-low) were administered to 12 test labs, where they were analyzed by one or more participating ctDNA sequencing assays (ROC, ILM, IDT, BRP and TFS; Supplementary Table 3). Bioinformatic analysis was performed by the relevant assay vendor using their custom pipelines. Results were then submitted for analytical evaluation by an independent team. b, Schematic overview of the proficiency testing scheme. Each participating ctDNA assay was performed at two or three independent test labs, with four technical replicates per lab generated for each test sample. Each of Lbx-high, Lbx-low and Sample b were analyzed at a fixed 25 ng input amount, and Lbx-low was additionally analyzed at 10 ng and 50 ng input amounts and at 25 ng input after extraction from a synthetic plasma solution (Lbx-low-plasma). In total, 360 ctDNA assays were evaluated. c (upper), Violin plots show coverage distributions (unique fragment depth) for Lbx-high and Lbx-low (25 ng input) replicates in each participating assay. c (lower), Distribution of VAF for on-target variant candidates in Lbx-high and Lbx-low (25 ng input). For comparison, expected VAF distributions for known variants in Lbx-high and Lbx-low are also shown (lower left).
Detection of ctDNA mutations in Lbx-low

**Assay:**

- ROC
- ILM
- IDT
- BRP

**Lab:**

- 1
- 2
- 3
- 4

**Replicate:**

- 1
- 2
- 3
- 4

- Known variant: Detected
- Missed
- Missed
- Missed
- Missed
- Missed

**ROC**

- Lbx-high
- Lbx-low

**ILM**

- 2.5%
- 0.5%
- 0.5%
- 0.5%

**IDT**

- 2.5%
- 0.5%
- 0.5%
- 0.5%

**BRP**

- 2.5%
- 0.5%
- 0.5%
- 0.5%

**Sensitivity**

- ROC
- ILM
- IDT
- BRP

**Precision**

- ROC
- ILM
- IDT
- BRP

**VAF**

- 0.1–0.5%
- 0.5–2.5%
- >2.5%

**Reproducibility**

- All comps
- Between-lab
- Within-lab

**Pairwise comparisons**

- ROC
- ILM
- IDT
- BRP

**a** (upper), Ordered heat maps show the detection of known variants (rows) in ctDNA assay replicates (columns). All on-target variants for a given assay are shown. Variants are sorted by expected VAF in descending order, and replicates are arranged hierarchically by assay type, test lab and replicate number. Heat maps show results for Lbx-low at 25 ng input, and equivalent heat maps for Lbx-high are shown in Supplementary Fig. 4a. **a** (lower), Aligned below each heat map column, bar charts indicate the sensitivity of variant detection in each replicate. Sensitivity is reported separately for known variants in the following VAF ranges: 2.5–0.5%, 0.5–0.3%, 0.3–0.2% and 0.2–0.1%, with measurements taken from both Lbx-high (high and mid VAF) and Lbx-low (low VAF). **b**, Precision recall curves compare diagnostic performance of participating ctDNA assays for Lbx-low (25 ng input; VAF range, 2.5–0.1%). Equivalent curves for Lbx-high are shown in Supplementary Fig. 4c. **c, d**, Bar charts show pairwise reproducibility scores for participating assays (n=132 for ROC and ILM and n=56 for IDT and BRP; median ± range). **c**, Reproducibility is reported separately for variant candidates at high, mid and low frequency (as above). **d**, Reproducibility is reported separately for all within-lab and between-lab pairwise comparisons. Note that, due to its smaller panel size and VAF distribution of on-target variants, the TFS amplicon sequencing assay is not included in these analyses.
BRP showed lower heterogeneity than ROC across their respective target regions (normalized interquartile range (IQR) = 0.23 versus 0.35; Supplementary Fig. 3a) and at matched sites present on both hybrid capture panels (normalized IQR = 0.20 versus 0.46; Supplementary Fig. 3b).

**Analytical sensitivity.** We next evaluated the sensitivity of hybrid capture ctDNA assays by measuring the fraction of on-target known variants that were detected in each library (Methods and Supplementary Data 2). Known variants were detected with superior sensitivity in Lbx-high compared to Lbx-low for all assays, reflecting their higher frequency in the former sample (Fig. 3a and Supplementary Figs. 3c,d and 4a,b). Indeed, all assays were highly sensitive for known variants at high (VAF > 2.5%, 0.99–1.00) and intermediate (VAF 0.5–2.5%, 0.96–1.00) frequencies but showed progressively weaker sensitivity for variants at lower frequencies (VAF < 0.5%), with significant variation observed among assays (VAF 0.1–0.5%, 0.39–0.83) (Fig. 4a). The most sensitive assays (IDT and BRP) achieved sensitivity greater than 0.90 for variants with 0.3–0.5% VAF; however, no assays reached this mark for variants with 0.2–0.3% or 0.1–0.2% VAF (Fig. 4a).

As demonstrated above, high coverage is essential for reliable sampling of rare ctDNA mutations. Consistent with this, differences in assay sensitivity partially reflected the differences observed among assays in coverage depth. For example, the high depth achieved by BRP enabled sensitive detection for variants as low as 0.3–0.5% VAF (0.89–0.93), whereas the ILM assay exhibited lower fragment depth and lower sensitivity at this level (0.24–0.77) (Figs. 3c and 4a). However, coverage depth alone was not necessarily a good predictor of sensitivity, with IDT achieving superior sensitivity to ROC despite its lower fragment depth (Figs. 3c and 4a). This result is likely attributed, at least in part, to the lower coverage heterogeneity in the IDT assay (Supplementary Fig. 3a,b), emphasizing the importance of achieving even coverage, in addition to overall depth.

Sensitivity is also influenced by bioinformatic variables. For example, among BRP assays, several known variants were missed by every replicate despite other variants of similar or lower VAF being

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**Fig. 5 | Effect of cell-free DNA input quantity (Lbx-low) on hybrid capture ctDNA assay performance.** a, Violin plots show coverage distributions (unique fragment depth) for Lbx-low replicates at 10 ng, 25 ng and 50 ng input amounts for hybrid capture ctDNA assays. Note that 10 ng ILM assays did not reach minimum coverage requirements, so they were excluded from subsequent analysis. b, Precision recall curves compare diagnostic performance of participating ctDNA assays for Lbx-low at each input amount above (VAF range, 2.5–0.1%). c–e, Curves showing the relationship between cell-free DNA input quantity (Lbx-low) and variant detection sensitivity (c), pairwise reproducibility (d) and FP rates (Fps per kb; e) for each participating ctDNA assay profiling low-frequency variants (VAF range, 0.5–0.1%). Error bars are mean ± 95% confidence interval. Note that, due to its smaller panel size and VAF profiling low-frequency variants (VAF range, 0.5–0.1%), error bars are mean ± 95% confidence interval. Note that, due to its smaller panel size and VAF profiling low-frequency variants (VAF range, 0.5–0.1%).
reproducibly detected (Fig. 4a). This suggests that the BRP analysis pipeline was highly stringent, with strict filtering of variant candidates slightly reducing the sensitivity of variant detection that was achieved (Figs. 3c and 4a).

**Analytical accuracy.** We next identified false-positive (FP) variant candidates that were erroneously detected at known negative positions by each assay (Methods and Supplementary Data 2). With all assays using unique molecular identifiers (UMIs) to correct sequencing errors\(^3\), FPs were relatively rare, ranging from a mean of 1.65 to 5.3 FP candidates per replicate (at 25 ng input; Supplementary Table 5). After accounting for panel sizes, BRP exhibited the lowest FP rate (0.03 FP per kb) and IDT the highest (0.07 FP per kb); however, given the small number of FPs, the differences between assays were not statistically significant (\(P > 0.05\)). Erroneous variant candidates occurred almost exclusively at low frequency (VAF < 0.5%; Supplementary Table 5).

To compare the accuracy of the participating ctDNA assays, we generated precision recall curves\(^4\), ranking known variants and FPs according to their observed VAFs. For Lbx-low samples at 25 ng input, BRP was the most accurate assay, with roughly equivalent sensitivity but superior precision to IDT (Fig. 4b and Supplementary Fig. 4c). Although this analysis enables useful cross-platform comparisons, it should be noted that precision is strongly influenced by the mutational burden of the sample under analysis (that is, the number of positives available for detection) and should not be taken to indicate the inherent precision of the participating ctDNA assays. Overall, despite the differences observed in fragment depth and sensitivity among the different ctDNA assays, FP rates were modest and broadly similar. Therefore, sensitivity, rather than precision, was the major determinant of overall analytical performance.

**Reproducibility among assay replicates.** We next evaluated reproducibility by comparing the outcomes of replicate assays within and between labs (Methods). We defined reproducibility as the fraction of variant candidates shared between any pair of replicates, with all possible pairwise comparisons considered.

Similarly to sensitivity, reproducibility was generally high (0.99–1.00 and 0.95–1.00) for variants at high (>2.5%) and intermediate (0.5–2.5%) VAF, respectively, but was relatively low and differed widely among panels for low-frequency variants (0.1–0.5%, 0.58–0.83; Fig. 4c and Supplementary Fig. 4d). Once again, coverage was a relevant variable but was not alone sufficient to explain differences in reproducibility among assays, with IDT achieving relatively high reproducibility despite its lower fragment depth (Figs. 3c and 4c). The results highlight the difficulty of reproducibly detecting
low-frequency ctDNA mutations and suggest that sensitivity is the major determinant of assay reproducibility. Within known positions, we found that FPs constituted only a small minority (~10%) of the discordant variant candidates between any given pair of replicates (Supplementary Fig. 4e), although we note that this fraction might be larger in samples with lower mutational burden than the reference samples analyzed here.

The concordance between reproducibility and sensitivity indicates that measurements of reproducibility can provide a useful proxy for diagnostic performance that is not dependent on the availability of a reference annotation. However, we noted that, whereas false negatives (FNs) are guaranteed to reduce assay sensitivity, this is not necessarily true for reproducibility, where systematic FNs (that is, known variants missed in every replicate) are not penalized. In this case, reproducibility would slightly overestimate the performance of the BRP assay, in which systematic FNs were most common (Fig. 4a,c). This emphasizes the value of well-characterized reference samples that can directly measure diagnostic performance27.

Finally, despite the wide variation among samples and panels, we observed no significant differences in reproducibility across within-lab and between-lab comparisons (Fig. 4d). This implies that all assays were robust to technical variables among facilities and were affected largely by random, rather than systematic, variation.

### Effect of cell-free DNA input quantity and plasma extraction.

The quantity of cell-free DNA retrieved from a patient blood draw is typically small, and this can be a major limitation for ctDNA assays8. To assess the effect of cell-free DNA input quantity, we next measured the performance of each hybrid capture assay with high (50 ng), medium (25 ng) and low (10 ng) amounts of input DNA (Lbx-low; Methods).

Coverage depth scaled linearly with input quantity for a given assay but varied widely among assays (Fig. 5a). Low-input (10 ng) ILM assays did not reach the minimum coverage requirements for analysis, so they were excluded from subsequent evaluation (Fig. 5a). The ILM assays did not reach the minimum coverage requirements for assay but varied widely among assays (Fig. 5a). Low-input (10 ng) (Lbx-low; Methods).

Coverage depth scaled linearly with input quantity for a given assay but varied widely among assays (Fig. 5a). Low-input (10 ng) ILM assays did not reach the minimum coverage requirements for analysis, so they were excluded from subsequent evaluation (Fig. 5a). The increasing fragment depth afforded by 25 ng input, compared to 10 ng, resulted in substantial improvements in sensitivity, reproducibility and overall diagnostic performance for all assays, particularly for low-frequency variants (Fig. 5b–e and Supplementary Fig. 5a,b). However, some assays (BRP and ROC) showed minimal further improvement with the addition of 50 ng input (Fig. 5b–e and Supplementary Fig. 5a,b). The extent to which performance varied over the range of input quantities tested indicates the robustness of each assay to the variable cell-free DNA input amounts encountered in the clinic. Overall, the greater fragment depth achieved by an assay at a given input level, the more robust that assay was to variation in input quantity, with BRP being the most stable (Fig. 5b–e).

We also evaluated the effect of cell-free DNA extraction, with each laboratory independently performing multiple extractions on DNA from Lbx-low suspended in synthetic plasma solutions (Lbx-low-plasma). Extraction efficiencies ranged from a mean of 33% (TFS) to 55% (BRP) (Supplementary Data 3), with the DNA retrieved being subsequently quantified and analyzed at 25 ng input quantities (Methods). In general, we observed no significant differences in sensitivity, FP rates or overall accuracy between Lbx-low and Lbx-low-plasma (Supplementary Fig. 6a–c). Pairwise reproducibility was equivalent for Lbx-low and Lbx-low-plasma replicates, as well as for the pairwise comparison of Lbx-low to Lbx-low-plasma replicates (Supplementary Fig. 6d). Finally, just as for Lbx-low, there was no difference in pairwise reproducibility across within-lab and between-lab comparisons for Lbx-low-plasma replicates (Supplementary Fig. 6e). These results indicate that all participating ctDNA assays were relatively robust to technical variables between test labs at the plasma DNA extraction stage.

Comparison of TFS amplicon assay to hybrid capture panels.

Amplicon sequencing methods enable targeted analysis of cancer mutation hotspots and can be applied in ctDNA analysis. We next compared the Thermo Fisher Scientific Oncomine ctDNA assay (TFS) to the other participating ctDNA assays, which all use hybrid capture enrichment (Fig. 3a,b). The TFS target regions (~1.9 kb) encompass driver mutation hotspots within 11 cancer genes (Supplementary Table 3), such as BRAF p.V600E, a highly recurrent melanoma mutation that is one of the known variants in our analysis (Fig. 6a). With these target regions almost entirely contained within the ROC, ILM and BRP capture panels, we were able to perform direct comparisons of sensitivity, accuracy and reproducibility within this clinically relevant window (Methods).

Overall, performance was similar between the TFS amplicon assay and hybrid capture assays. TFS showed perfect detection sensitivity for on-target known variants in Lbx-high and achieved equivalent sensitivity to ROC and BRP for Lbx-low, when high input was used (25 ng or 50 ng; Fig. 6a,b). Due to its lower fragment depth (Fig. 3c), TFS suffered a larger reduction in sensitivity than ROC and BRP when input quantity was restricted (10 ng) but still outperformed ILM, which was the hybrid capture assay with the lowest fragment depth (Fig. 6a,b). With this loss of sensitivity, BRAF p.V600E (VAF=0.33%) was missed in half of all TFS replicates at 10 ng, whereas it was detected with perfect reliability at 25 ng and 50 ng (Fig. 6a). TFS and ILM also showed poor reproducibility at low input levels compared to ROC and BRP (Fig. 6c). Finally, as we observed for hybrid capture panels (Fig. 4d), TFS showed no difference in reproducibility between within-lab and between-lab comparisons (Fig. 6d).

To further evaluate the detection of low-frequency mutations, TFS test sites also analyzed synthetic DNA control (AcroMetrix Oncology Hotspot Control) containing 15 known cancer mutations that overlapped TFS hotspot regions (out of 521 in total; 50 ng input; Supplementary Methods). This enabled more robust measurement of diagnostic performance for low-frequency variants, with all 15 mutations being present at ~0.1% VAF. Low-frequency mutations were detected with relatively high sensitivity (0.86–1.0; Supplementary Fig. 7a). Several FPs were also detected; however, these were almost entirely excluded by applying a minimum detection threshold of VAF>0.05% (Supplementary Fig. 7a,b). Accordingly, a strong improvement in reproducibility (median, 0.65 versus 0.94) was observed when applying this filter, at a relatively small cost to sensitivity (median, 0.96 versus 0.90) (Supplementary Fig. 7c).

Overall, these results indicate similar performance between the TFS amplicon sequencing assay and participating hybrid capture-based assays for the detection of SNVs. Indeed, we found that the performance of a given assay, and especially its robustness to reductions in input quantity, was largely determined by the fragment depth achieved, not the method of target enrichment or sequencing.

### Discussion

The ability to diagnose and monitor cancer through ctDNA sequencing promises to revolutionize clinical oncology6–9. Accordingly, there is considerable interest and investment in the ongoing development of NGS-based ctDNA assays89. However, the reliable detection of trace amounts of fragmented ctDNA from a routine blood draw remains a major technical challenge11. Government, regulatory and clinical organizations have, therefore, called for thorough analytical evaluation of ctDNA assays, to define diagnostic limits, assess reproducibility and identify key experimental variables that affect performance84.

This study begins to address these unmet needs and, to our knowledge, provides the first large-scale assessment of analytical performance among industry-leading ctDNA assays. We report that mutations represented above ~0.5% VAF could be detected with
high sensitivity, accuracy and reproducibility by all participating assays. However, variant detection was generally unreliable and variable among assays for mutations lower than ~0.5% VAF. This was primarily driven by the proportion of known variants that missed detection (that is, lack of sensitivity) due to stochastic sampling, in agreement with our initial simulated experiments. FPs were a less significant source of discordant results, with UMIIs used effectively to minimize errors in all assays. Cell-free DNA input quantity was a key variable, with increasing input leading to improved fragment depth, sensitivity and reproducibility.

Previous studies reported discordant results among alternative assays or parallel ctDNA and tumor biopsy tests, although the underlying causes and extent to which this resulted from biological rather than technical factors were unclear26–27 (see below). We also observed discordant results among vendors, labs and assay replicates. However, this was limited to low-frequency mutations (VAF < 0.5%) and largely reflected the limitations of stochastic sampling rather than technical biases or errors. In fact, we found that participating assays were generally robust to technical variables among test labs, from plasma extraction to sequencing workflow stages.

The performance characteristics of the assays evaluated here were broadly similar to what has been reported by several ctDNA sequencing providers (based on internal testing) that did not participate in this study. During validation of the Guardant360 CDx hybrid capture assay, variants were detected with high sensitivity (~94%) at VAF ≥ 0.4%, declining to ~64% among variants with VAF ranging from 0.05% to 0.25%28. FoundationACT showed ~99% sensitivity for SNVs with VAF > 0.5%, ~95% for 0.25%–0.5% VAF and ~70% for 0.125–0.25% VAF29. MSK-ACCESS showed ~98% sensitivity for SNVs with VAF > 0.5%, declining to ~74% for 0.1%–0.5% VAF30. Validation of the amplicon-based InVisionFirst assay suggested that this might have superior limit of detection (LOD) to the hybrid capture assays mentioned above, with ~99% sensitivity for SNVs as low as 0.25–0.35% VAF31. Consistent with our findings, all of these providers also reported low FP rates. Although direct comparisons among studies that used different test samples and DNA input quantities must be treated with caution, it appears generally true that the sensitive detection of ctDNA mutations below ~0.5% VAF is a major challenge.

Although the accurate detection of mutations >0.5% VAF and robustness to technical variables among ctDNA assays is cause for optimism, data arising during early clinical implementation of several tests highlight the necessity for reliable detection of low-frequency mutations. A survey of more than 1,000 plasma samples from patients with cancer tested with Guardant360 CDx found that half of all detected SNVs occurred below ~0.5% VAF and a quarter below ~0.2% VAF32. Among 859 patients tested with FoundationACT, half of all variants detected had VAFs below ~1.3% and a third below ~0.5%33. Among 435 patients tested with MSK-ACCESS, more than 5% of all mutations detected were missed by NGS but identified by more sensitive genotyping methods, with these having a median VAF of ~0.08%29. These results, which are generally based on the analysis of patients with advanced disease, demonstrate the tendency for ctDNA mutations to be represented at very low frequencies. Moreover, given the likelihood that many low-frequency variants missed detection, the median VAFs reported in these studies represent upper-bound estimates.

The analytical performance characteristics of a given ctDNA assay determine its potential suitability for specific applications in research and clinical oncology, with different assays being suited for different purposes. For example, higher sensitivity and precision and lower LOD are required for molecular characterization of early-stage versus late-stage cancer, due to the lower mutational burden and abundance of ctDNA fragments in circulation34.

That mutations above ~0.5% VAF were accurately detected indicates that the participating ctDNA assays might be suitable for molecular stratification and profiling tumor evolution in patients with advanced cancer, where informative mutations are commonly detected with VAFs ranging from ~1% to 10%35,18–20. Given that variants were also accurately quantified above a limit of quantification of ~0.8% VAF, ctDNA sequencing appears suitable to monitor frequencies over time and in response to therapeutic intervention14. Characterization of early-stage, localized disease with ctDNA sequencing requires accurate detection of mutations with VAFs ranging from ~0.1% to 1% VAF, although we note that this is likely to vary between cancer types and individual patients3,4,41,42. In patients with non-small-cell lung cancer, for example, a tumor ~10 cm in diameter yields ctDNA fragments at ~0.1% VAF, on average, in the plasma1. Therefore, although further improvement is required to ensure reliable detection of low-frequency mutations, suitability for early-stage cancer appears within reach of current ctDNA sequencing assays.

The ability to detect and monitor post-surgical MRD by ctDNA sequencing is an application with great potential utility37. However, MRD monitoring demands highly sensitive detection of mutations with VAFs ranging from ~0.1% to 0.01% to reliably predict disease relapse3,13,45. Given that the participating assays in our study were generally unreliable for mutations with VAFs ~ten-fold higher than this, substantial further development is required for use in monitoring MRD, and targeted analysis of known mutations by droplet digital PCR (ddPCR) (or related approaches) remains the most promising strategy for this application.

The breadth of a ctDNA assay’s target regions and the types of mutations detected are also relevant considerations. For example, large hybrid capture panels such as the ILM panel tested here (154 target genes, ~500 kb) are ideal for unbiased genomic characterization in advanced metastatic disease but are not cost-effective for targeted monitoring of known driver mutations during and after therapy. Amplicon methods, such as the TFS panel tested here (11 target genes, ~1.9 kb), enable more affordable, focused analysis of mutation hotspots, but their small panel sizes limit suitability for unbiased genomic surveillance. Limited ability to detect mutations types beyond SNVs and small indels is a further drawback of amplicon-based approaches.

The use of ctDNA sequencing for early cancer detection demands unbiased surveillance of broad target regions with high sensitivity and low LOD. Moreover, given the low prior probabilities involved in screening healthy individuals, FP rates must be exquisitely low for an assay to achieve clinical utility. Given these requirements, considerable improvements to existing assays, as well as continued depreciation of sequencing costs, will be needed for this much-anticipated application to approach feasibility.

Given the discussions above, improved sensitivity for mutations below ~0.5% VAF should be a priority for the ongoing development of ctDNA assays. Optimizations to the efficiency of plasma DNA extractions, target capture and NGS library preparations might yield incremental improvements in coverage and, thereby, in the detection of low-frequency mutations. Such advances are critical to ensure the robustness of ctDNA assays to the variable cell-free DNA input quantities encountered in the clinic36. However, participating assays showed diminishing returns with increasing sample input and/or fragment depth and remained unable to exhaustively identify low-frequency mutations.

This deficit in sensitivity reflects the difficulty of reproducibly detecting rare ctDNA fragments by random sampling of amplified fragments—an inherent statistical challenge that might not be fully overcome simply by increasing global fragment depth. Substantial additional improvements to ctDNA assays might, therefore, require further innovations, such as the selective enrichment of ctDNA fragments over the background of non-cancerous cell-free DNA fragments, potentially by exploiting discrepancies in fragment size35,46 or methylation status37.
ctDNA assay performance will also benefit from ongoing improvement in the detection of cancer mutations beyond SNVs and small indels. Translocations and other structural variants can be informative cancer biomarkers, but their detection is challenging due to the small fragment sizes and amplification biases in ctDNA assays. Further assay development and proficiency testing on these features are, therefore, required.

The reliability of ctDNA sequencing assays is commonly assessed by measuring concordance among alternative assays or assay replicates, parallel ctDNA and tumor biopsy tests or orthogonal analysis of plasma samples with non-NGS-based techniques, such as ddPCR. These approaches have been applied across large cohorts of clinical specimens to inform assay development and validation.

Although highly useful, several caveats must be acknowledged. As noted earlier, concordance measurements among assays do not penalize systematic errors, such as mutations that are missed by multiple assays or replicates, and might, therefore, overestimate performance. The comparison of plasma and tumor tissue biopsies is also confounded by an array of biological factors, such as tumor type, stage, morphology and heterogeneity. For example, a mutation detected in a tumor tissue biopsy but not with matched ctDNA sequencing might be truly absent from the plasma, due to restricted local circulation at its site of origin. Therefore, discordance is not necessarily indicative of poor analytical performance for either test. Although orthogonal analysis of a plasma sample with ddPCR can reliably determine whether detected mutations are true, it is not practical with this targeted approach to assess all invariant sites across a large hybrid capture sequencing panel to rule out potential FNs.

Arguably, the most relevant limitation in using bona fide cell-free DNA samples for analytical validation experiments is the inability to standardize across different assays, sites and replicates. Patient plasma samples vary widely in mutational burden and cell-free DNA yields, with no single sample representing the full diversity of mutation types and frequencies that should be considered in rigorous proficiency testing. The use of different clinical cohorts, cell-free DNA input quantities and test materials makes it difficult to draw reliable comparisons among validation studies performed by different ctDNA sequencing providers.

To avoid these caveats, we used a combination of synthetic DNA controls (sequins, AcroMetrix) and cell-line-derived reference samples (Lhx-high and Lhx-low) to evaluate the analytical performance of ctDNA assays. This approach allows 1) appropriate numbers, types and frequencies of known mutations to be analyzed; 2) unambiguous classification or true/false positives/negatives across large target regions; 3) standardization of samples and input quantities among sites, assays and assay replicates; and 4) absence of confounding biological variables.

However, there are also limitations to this approach. The enzymatic process by which samples were fragmented does not perfectly emulate the fragmentation of natural cell-free DNA, on which the participating assays have been optimized. It is, therefore, possible that the efficiency of fragment capture and library conversion for these contrived samples might be lower or higher for any given kit than for natural cell-free DNA. Moreover, we cannot be certain that the kinetics of fragmentation are equivalent among all genome regions, potentially resulting in some sites being over-represented or under-represented or represented by fragments of atypical sizes. Although we deliberately devised reference samples with a high mutational burden to improve the power of performance measurements, this means that measures of assay precision are inflated and should not be interpreted as realistic measures of performance on clinical samples (instead, they are useful for making technical comparisons among assays, given that common reference samples were tested). Contrived reference samples can never fully recapitulate the many nuanced biological factors that influence the potential utility of ctDNA sequencing assays in real clinical contexts. For example, the ability to distinguish informative ctDNA mutations from a background of benign variants in cell-free DNA, generated during clonal hematopoiesis, is a challenge that cannot be evaluated using synthetic reference materials. Therefore, although contrived samples are an ideal substrate on which to assess the analytical performance characteristics of ctDNA assays in the absence of confounding biological variables, they alone cannot be used to determine clinical thresholds (for example, limit of detection and limit of blank), which must be accounted for, rather than excluded, such variables. Importantly, clinical performance cannot exceed analytical performance, and analytical validity is a prerequisite for clinical validity and clinical utility, with these properties requiring demonstration in clinical trials employing assays that have achieved analytical validity.

This study advances the community’s understanding of analytical performance characteristics in ctDNA sequencing, outlines a set of best practice guidelines (Table 1) and constitutes a step toward the ultimate goal of establishing clinical utility for precision oncology. Moreover, this study establishes a unique set of reference materials and annotations and an analytic framework for standardized proficiency testing on ctDNA assays. Although ongoing studies are required to establish the potential clinical validity and utility of ctDNA assays, the SEQC2 Oncopanel Sequencing Working Group has helped lay the foundation for such future work.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41587-021-00857-z.

Received: 20 July 2020; Accepted: 15 February 2021; Published online: 12 April 2021

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Methods

Simulated ctDNA sequencing assays. To model the parameters of ctDNA sequencing assays, we generated simulated NGS libraries that emulate targeted analysis of cell-free DNA by hybrid capture sequencing. Simulated reads were created using seqsimulator42 [https://github.com/mbomman], with uniform and error rates set to zero (<0.0 - 0.0 ± 0.0 ±). Paired-end (2 × 150-bp) read fragments were generated, with a mean fragment size of 160 bp and a standard deviation of 15 bp (<150 – 250 d 160 s 15>). Read fragments were simulated uniformly over 155 cancer-related loci on a generic gene panel for oncology applications (Roche NimbleGen; for research use only, not for diagnostic procedures), based either on the hg38 reference sequence or a modified hg38 sequence containing one Cosmic SNV per exon (n = 2,356; selected randomly and inserted using gatk FastAAlternateReferenceMaker (v3.8)31).

Reads simulated from reference and mutant sequences were combined in precise ratios to create eight independent simulated libraries in which all SNVs were represented at a 5%, 1%, 0.5%, 0.4%, 0.3%, 0.1%, with genes covered uniformly at ~9,000-fold fragment depth. Simulated read fragments were aligned to hg38 using bwa mem (v0.7.16)54. In silico capture enrichment was performed by intersecting aligned read fragments with the capture targets BED file and retaining only fragments with ≥60 bp overlap to a target region. This process creates convex coverage profiles over targeted exons that resemble typical coverage profiles obtained during hybrid capture sequencing (Supplementary Fig. 1a). These libraries were then downsampled (gatk DownsampleSam) to create additional libraries with incremental reductions in fragment depth.

Simulated SNVs were then detected using VarScan (v2.4.3)39, and detection sensitivity (TPs/TPs + FNs) was calculated for SNVs within each library, across a range of detection stringency levels (that is, the minimum number of supporting fragments for an SNV to be called). To measure the effect of variant position, simulated SNVs were parsed based on their distance to the nearest exon boundary (edge regions <20 bp; central regions >50 bp). To measure the effect of local alikability, SNVs were binned based on the alikability of their occupied exon sequence. Exons were considered to have 'suboptimal' alikability if 2% of overlying alignments had a mapping quality score = 0.

Manufacture and sequencing of synthetic DNA controls (sequins). We evaluated the detection of ctDNA mutations using synthetic DNA controls known as 'sequins' (www.sequinstandards.com). Synthetic controls were manufactured in a purpose-built facility at the Garvan Institute of Medical Research. Detailed descriptions of the design, manufacture and experimental validation of sequins were published previously55.

For this study a custom sequin mixture for ctDNA sequencing experiments was created, encompassing 354 individual sequins ranging in size from ~1 kb to 6 kb for a combined ~757 kb of synthetic DNA sequence in total. This mixture provides synthetic 'circular' representations of relevant exons and domains within 87 cancer-related genes and includes 134 synthetic mutations (Supplementary Data 1). As described previously, sequin sequences were initially synthesized and validated by a commercial vendor (Thermo Fisher Scientific, GenArt). Within a purpose-built manufacturing facility, synthetic sequences were amplified by bacterial culture, excised by restriction enzyme digest, quantified by ultraviolet fluorometry (Thermo Fisher Scientific Qubit) and combined using a liquid-handling robot (Eppendorf epMotion). By combining synthetic molecules representing each mutant allele in precise ratios, synthetic mixtures were represented across a wide range of VAF levels, ranging from 100% to 0.1% in two-fold increments. This staggered reference ladder allows detection sensitivity and quantitative accuracy to be assessed at different VAF levels.

To emulate the fragmentation of cell-free DNA, the synthetic sequin mixture was enzymatically sheared using NEDNext dsDNA Fragmentase in 10X fragmentation reaction buffer for 30 min at 37°C. The reaction was terminated by the addition of 0.5 M EDTA, and the resulting fragments were purified using double-sided SPIR size selection. To exclude fragments >250 bp, 0.65X Agencourt AMPure XP beads were used, whereas 1.8X Agencourt AMPure XP beads were used to enrich fragments <250 bp, and the purified DNA was visualized with an Agilent TapeStation. This fragment-sheared sequin mixture was validated by NGS, using a Nextera XT DNA Library Prep Kit and sequenced on an Illumina MiSeq.

After validation, the fragmented sequin mixture was spiked into mock human ctDNA reference samples (Lbx-high and Lbx-low; described below) at ~0.2% fractional concentration. These combined samples were analyzed by hybrid capture sequencing, using a custom oncology panel (Roche NimbleGen; for research use only, not for diagnostic procedures) targeting the 87 human cancer genes that were represented by sequin controls, as well as the sequin controls themselves (this approach is described in detail elsewhere)55. In total, 119 kb of synthetic sequence was captured and analyzed.

NGS libraries were prepared with a KAPA LTP Library Preparation Kit (Illumina platform KR0453, v6.17), in conjunction with IDT xGen dual index adaptors, according to the manufacturer's protocol, with ten cycles of PCR amplification. Capture enrichment was performed according to an established protocol (Roche Double Capture Technical Note, August 2012). Purified libraries were quantified on an Agilent TapeStation and sequenced on an Illumina NovaSeq (S1 flow cell).

Sequin bioinformatics analysis. Targeted NGS libraries containing reads from mock human ctDNA samples spiked with synthetic sequin controls were initially trimmed using TrimGalore [https://github.com/FelixKrueger/TrimGalore] and then processed using the purpose-built anaquin toolkit for sequin analysis, via a workflow that is described in detail elsewhere55. Briefly, sample-derived and sequin-derived reads were not separated, and sequin reads were reversed in orientation (anaquin split). Sample and sequin reads were aligned separately to the hg38 reference genome using bwa mem (v0.7.16). Off-target reads were excluded and PCR duplicates collapsed using gatk MarkDuplicates (v4.0). Sequin-derived alignments were then calibrated to equivalent coverage depth to accompanying sample alignments within matched genome regions (anaquin calibrate). VarScan (v4.0) was then used to call variants (SNVs and indels) within target sequin regions, with a minimum of three supporting read fragments required for detection. anaquin somatic was used to evaluate the detection of sequin variants.

Sequin libraries were then incrementally downsampled (gatk DownsampleSam), and variant detection was repeated across a range of deprecimating fragment depths. Detection sensitivity (the fraction of known sequin variants detected) was calculated in each downsamplied library to generate curves that model the relationship between sensitivity and fragment depth, with sequin variants parsed into pre-defined VAF bins (<0.5%, 0.5%–5% and >5%). Sequin variants were also parsed according to 1) fragment depth (high fragment depth, >60X fold; low-fold: 2); 2) distance to the closest exon boundary ('edge regions', <20 bp; 'central regions' >50 bp); 3) GC content within a 120-bp local window (high, >60%; low, <40%); and 4) sequence complexity within a local 120-bp window. Complexity was calculated using SeqComplex [https://github.com/caballero/SeqComplex], with windows showing entropy scores <1.9 considered to have low complexity.

Preparation of human ctDNA reference samples. The cross-platform ctDNA sequencing proficiency study from the present manuscript used a set of contrived reference DNA samples that are described in detail in a companion article55. Briefly, Sample A comprised genomic DNA extracted from ten diverse cancer cell lines (Agilent UHR cell lines), and Sample B is non-cancerous genomic DNA (Agilent Male Control DNA). Sample A and Sample B were combined to create two further reference samples: Lbx-high (20% A / 80% B) and Lbx-low (4% A / 96% B). Note that Lbx-high and Lbx-low are also referred to as Sample D and Sample E (before fragmentation) in the accompanying article describing the preparation of reference samples53.

Aliquots of these samples, as well as Sample A and Sample B, were enzymatically fragmented using the KAPA Frag Kit (KAPA Biosystems), according to the manufacturer's instructions. Samples dissolved in TE buffer were first purified with the Agencourt AMPure XP Kit at a 3:1 bead-to-sample volumetric ratio to remove EDTA. Purified DNA samples (5 μg) in 10 mM Tris-HCl, pH 8.0 (35 μl) were mixed with KAPA Frag Enzyme (10 μl) and 10x KAPA Frag Buffer (5 μl) on ice and incubated at 37°C for 25 min. After fragmentation, DNA samples were purified with Agencourt AMPure XP beads again, as described above.

Size selection of fragmented DNA (5 μg per well) was performed on a Pippin Prep instrument (Sage Science) using 3% agarose gel cassette (Sage Science, No. IP250) with the range from 110 to the BP calibration ladder (Agilent Technologies) to achieve an average fragment length of ~165 bp. After size selection, DNA samples were characterized using an Agilent Bioanalyzer 2100 with DNA high sensitivity kit (Agilent Technologies). Typically, a yield of 6–8% was obtained. Samples were quantified and dissolved in Tris buffer (10 mM Tris, pH 8.0) at 5 ng μl−1 for storage and distribution.

The plasma DNA sample (Lbx-low-plasma) was prepared with a DNA concentration at 40 ng ml−1 in synthetic plasma (Horizon Discovery), and two aliquots of 8 ml were shipped to each test site in two 10-ml tubes. Test labs did not assess the concentration of DNA samples, except the plasma sample, to avoid any site-to-site variation that could be introduced by independent quantification. Qubit dsDNA HS (High Sensitivity) 1.0 μl (6,000 copies μl−1 for the Qubit dsDNA HS assay calibration with distributed ctDNA samples as standards.

Multi-site ctDNA proficiency testing. Each testing laboratory performed one or more participating ctDNA sequencing assays (five in total; Supplementary Table 3), according to the vendor's instructions. Sequencing was performed using Illumina (NovaSeq 6000 and NextSeq 500) or Thermo Fisher Scientific (IonTorrent) instruments (Supplementary Table 3). Sequencing information is supplied in Supplementary Table 6. Details for all experimental procedures for each assay are provided in the Supplementary Methods.

Each participating assay was performed in 2–3 independent labs, with four technical replicates per lab for each mock ctDNA sample, at a fixed DNA input amount (25 ng). In addition, Lbx-low was analyzed with increased (50 ng) and
decreased (10 ng) input amounts. Each test lab also performed four independent plasma ctDNA extractions on the provided Lbx-low plasma sample (2.5 ml per replicate) and analyzed the extracted DNA at a fixed DNA input amount (25 ng per replicate). All sequencing libraries were then administered to the relevant ctDNA assay vendor for blinded analysis. Bioinformatics analysis was not standardized across the study, with each vendor, instead, employing an internal analysis pipeline and providing a final set of variant candidates for centralized evaluation by an independent team at the Garvan Institute of Medical Research and the FDA National Center for Toxicological Research. All participating assays used UMIs to collapse duplicate read pairs into consensus fragments. Comparisons of assay yields and depths throughout the study are based on unique fragment depth rather than raw read or alignment counts. Detailed information about the bioinformatics pipeline employed by each vendor is provided in the Supplementary Methods.

Evaluation of results. Pre-processing variant candidates. Assay vendors provided a single independent VCF file containing candidate variants called in each assay replicate, at each test lab. The following pre-processing steps were used to ensure direct comparability among all call sets. Where relevant, candidates marked with filter flags by the vendor bioinformatics pipeline, or indicated as VAF = 0, were excluded. Multi-allelic variant sites were broken into multiple individual variants using bcftools norm (v1.9). Complex and/or multi-nucleotide variants were broken into their simplest individual components using RTG-tools (https://github.com/RealTimeGenomics/rtg-tools) vcfdecompose (v3.10.1) with the--break-mnps-break-indels parameters set to TRUE, and gatk LendAlignAndTrimVariants (v4.0.11) was used to ensure consistent representation of indels.

Sensitivity and accuracy. To measure the sensitivity of participating ctDNA assays, we compared each set of variant candidates to the reference annotation described in ref. 2. RTG-tools vcfdeval (v3.10.1) was used to compare the VCF file for each assay replicate to the set of ‘known variants’ in Lbx-high/Lbx-low and the set of known germline variants in the non-cancer background Sample B. These comparisons were restricted to the intersection of all ‘known positions’, with the on-target reportable regions provided by the relevant assay vendor. All candidates within known positions were classified as TP or FP. Sensitivity was defined as the number of TPs in a given replicate divided by the number of on-target known variants for the relevant panel and was calculated both globally and within pre-defined VAF bins (0.1–0.5%, 0.5–2.5% and >2.5%). FP rates were defined as the number of FPs in a given replicate divided by the size of the on-target known negative positions for the relevant panel, thereby accounting for differences in panel size. Precision recall curves were generated by incrementally varying the minimum VAF threshold (below which candidates are excluded) from 0% to 100% and recalculating sensitivity (TP/(TP + FN)) and precision (TP/(TP + FP)) at each increment.

Reproducibility. To measure the reproducibility of participating ctDNA assays, we performed reciprocal pairwise comparisons among variant call sets for all replicates of a given assay/sample/input. Comparisons were performed using RTG-tools vcfdeval (v3.10.1) and were restricted to the capture target regions provided by the relevant assay vendor. For a given pair of replicates, reproducibility was defined as the fraction of total variant candidates that were concordant among call sets, with all possible pairwise comparisons being performed. Reproducibility was calculated globally, across within-lab and between-lab comparisons, and within pre-defined VAF bins (0.1–0.5%, 0.5–2.5% and >2.5%). When calculating reproducibility within VAF bins, candidates in a given bin in the first sample were compared to the whole call set for the second sample, and vice versa, to avoid bin edge effects.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Descriptive data about individual ctDNA assays are provided in Supplementary Data 2. Descriptive data about individual variants, including their detection status in each ctDNA assay, are provided in variant classification tables within the Source Data Excel file. These tables were used to generate variant detection heat maps and other data plots. Raw sequencing data have been deposited to the National Center for Biotechnology Information Bioproject PRJNA677999. Variant calls generated by each assay vendor (in VCF format) and panel region files (in BED format) can be accessed at the following link: https://figshare.com/projects/SEQC2_Onco-panel_Sequencing_Working_Group__-__Liquid_Biopsy_Study/94523. Source data are provided with this paper.

Code availability

Variant call sets for each ctDNA sequencing assay were generated by internal bioinformatics pipelines by each assay vendor. Although these pipelines are not open source, detailed descriptions and relevant software version numbers are provided in the Supplementary Methods. All data plots were generated using R (v3.5 or later) or GraphPad Prism (v8).

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Acknowledgements

All SEQC2 participants freely donated their time, reagents and computing resources for the completion and analysis of this project. We thank our expert colleague, S. J. Dawson, for providing useful feedback during manuscript preparation. We acknowledge the following funding sources: NHMRC grants APP1108254 and APP1114016 (to T.R.M.), BAA grant HHSF223201510172C (to D.J.), Shanghai Municipal Science and Technology Major Project grant 2017SHZDX01 (to L.S.), the National Natural Science Foundation of China grant 31720103909 (to L.S.), MRFF grant MRF1713594, Cancer Institute NSW Early Career Fellowship 2018/EC013 and philanthropic support from the Kinghorn Foundation (to I.W.D.). The contents of the published materials are solely the responsibility of the administering institution, a participating institution or individual authors, and they do not reflect the views of any funding body listed above. This research includes contributions from, and was reviewed by, the FDA and the NIH. This work has been approved for publication by these agencies, but it does not necessarily reflect official agency policy. Certain commercial materials and equipment are identified to adequately specify experimental procedures. In no case does such identification imply recommendation or endorsement by the FDA or the NIH, nor does it imply that the items identified are necessarily the best available for the purpose.

Author contributions

W.T., D.J. and J.X. conceived the project. I.W.D., J.W., W.J., D.J., T.R.M. and J.X. devised the experiments. I.W.D. performed simulated experiments. B.S.M., J.B., I.S., A.B., D.C., J.C., M.H., N.M., P.M., R.S., D.S., P.S., H.T., D.T., H.A., H.B., B.B., D.D., A.G., S.G., K.H., C.M., A.R., P.R., R.R., R.S., M.S., P.S., M.S., V.T. and S.V. performed and/or coordinated laboratory experiments. I.W.D. performed data analysis. I.W.D. and T.R.M. prepared the figures. I.W.D., J.W., T.R.M. and J.X. devised the project. I.W.D., J.W., W.J., D.J., T.R.M. and J.X. conceived the project. I.W.D., J.W., W.J., D.J., T.R.M. and J.X. devised the experiments. I.W.D. performed simulated experiments. B.S.M., J.B., I.S., A.B., D.C., J.C., M.H., N.M., P.M., R.S., D.S., P.S., H.T., D.T., H.A., H.B., B.B., D.D., A.G., S.G., K.H., C.M., A.R., P.R., R.R., R.S., M.S., P.S., M.S., V.T. and S.V. performed and/or coordinated laboratory experiments. I.W.D. and B.S.M. performed data analysis. I.W.D. and T.R.M. prepared the figures. I.W.D., D.J., T.R.M. and J.X. prepared the manuscript, with support from all co-authors.

Competing interests

The Garvan Institute of Medical Research has filed patent applications on synthetic controls for genomics.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41587-021-00857-z.

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Peer review information Nature Biotechnology thanks Michael Berger and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

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Software and code

Policy information about availability of computer code

Data collection

Variant callsets for each ctdNA sequencing assay were generated by internal bioinformatics pipelines by each assay vendor. While these pipelines are not open source, detailed descriptions and relevant software version numbers are provided in the Supplementary Methods section. The following software was used:

Roche AENE/O ctdNA Analysis Server v1.1
Roche Oncology Analysis Server v1.1
Picves v5.2.10.49 (https://github.com/Illumina/Picves)
Picard v2.18.9 IlluminaBasecallsToSam
fgbio v0.7.0 ExtractUmisFromBam
Picard v2.18.9 SamToFastq
Picard v2.18.9 MergeBamAlignment
Picard v2.18.9 CollectHsMetrics
fgbio v0.7.0 ExtractUmisFromBam.
fgbio v0.7.0 GroupReadsByUmi
fgbio v0.7.0 CallDuplexConsensusReads
fgbio v0.7.0 FilterConsensusRead
fgbio v0.7.0 ClipBam
bwa-mem v0.7.15
samtools v1.5
AstraZeneca VarDict v1.5.8
bc2fastq v2.20
Trimomatic 0.36
BWA aligner 0.7.10
VarScan v2.4.3
ANNOVAR 20160201
Data analysis

All software used used to evaluate results and produce figures are publicly available and/or open source. Full descriptions and version numbering are found in the Materials & Methods section. The following software was used:

- wgsim [v1.9: https://github.com/lh3/wgsim]
- gatk FastAAlternateReferenceMaker [v3.8]
- bwa mem [v0.7.16]
- VarScan2 [v2.4.3]
- TrimGalore [https://github.com/FelixKrueger/TrimGalore]
- anaquin toolkit [v3.22.0]
- gatk MarkDuplicates [v4.0]
- gatk DownsampleSam [v4.0]
- gatk leftAlignAndTrimVariants [v4.0]
- bedtools [v2.25.0]
- SeqComplex version 1 [https://github.com/caballero/SeqComplex]
- bcftools norm [v1.9]
- RTG-tools vcfdecompose [v3.10.1]
- RTG-tools vcfeval [v3.10.1]
- R [v3.5]
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All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
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- A list of figures that have associated raw data
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Descriptive data about individual ctDNA assays are provided in Supplementary Data 2. Descriptive data about individual variants, including their detection status in each ctDNA assay, are provided in variant classification tables within the Source Data Excel file. These tables were used to generate variant detection heatmaps and other data plots. Raw sequencing data has been deposited to the NCBI Bioproject PRJNA677999. Variant calls generated by each assay vendor [in VCF format and panel region files (in BED format) can be accessed at the following link: https://figshare.com/projects/SEQC2_Onco-panel_Sequencing_Working_Group_-_Liquid_Biopsy_Studv94523

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Sample size
As a proficiency study, this study necessarily employed large sample sizes and high replication. This is explained clearly in section "Multi-site, cross-platform proficiency study" (p10) and presented schematically in Figure 3.
For every condition/assay/sample, 8-12 replicates were performed across 2-3 independent test sites and 360 independent ctDNA sequencing assays were performed overall.

Data exclusions
Low input (10ng) samples from the Illumina panel were excluded from performance comparisons as they did not reach the minimum depth requirements for Illumina’s analysis pipeline. This is clarified in Figure 5 and within the relevant main text section (p14):
"Low input (10 ng) ILM assays did not reach the minimum coverage requirements for analysis, so were excluded from subsequent evaluation (Fig. 5a)."
Some samples from the IDT panel were excluded from performance comparisons as they did not reach the minimum depth requirements for IDT's analysis pipeline. This is documented in the Supplementary Methods (p16):
"10ng samples with < 500x median target coverage, and other samples with <2000x median target coverage was flagged as outliers."

Replication
For every condition/assay/sample, 8-12 replicates were performed across 2-3 independent test sites and 360 independent ctDNA sequencing assays were performed overall.

Randomization
The study design did not involve any randomisation.
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