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Microbiome analyses of blood and tissues suggest cancer diagnostic approach

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Summary

Systematic characterization of the cancer microbiome provides a unique opportunity to develop cancer diagnostics that exploit non-human, microbial-derived molecules in a major human disease. Based on recent studies showing significant microbial contributions in select cancer types^{1–10}, we re-examined *treatment-naïve* whole genome and whole transcriptome sequencing studies

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Contributions

The research topic was developed by E.K., G.D.P., T.K., S.J., J.M., S.J.S., S.M.M., A.D.S., S.P.P., and R.K.. The TCGA microbial-detection pipeline was co-developed by E.K., S.J.S., J.M., J.K., and G.D.P.. The supervised normalization pipeline was developed by G.D.P.; the decontamination pipeline by G.D.P., A.D.S., and S.P.P.; and the ML pipeline by G.D.P., A.D.S., T.K., and S.J.. SourceTracker2 analyses, including re-running HMP2 shotgun metagenomic data through the microbial-detection pipeline were completed by E.K., Q.Z., and G.D.P.. Samples for the prospective validation study were collected by R.H., R.M., and S.P.P., then processed for sequencing by C.C., S.F., and G.D.P., then bioinformatically analyzed by E.K., S.W., and A.D.S., and then put through normalization and ML pipelines by G.D.P. and A.D.S.. The cell-free microbial DNA extraction protocol was originally designed and refined by C.C., S.F., S.M.M., and A.D.S.. The original version of the manuscript was written by G.D.P., A.D.S., S.P.P., and R.K.. All authors contributed to the final version of the manuscript.

Competing interests

Clarity Genomics, the employer of E.K., did not provide funding for this study. Both G.D.P. and R.K. have jointly filed U.S. Provisional Patent Application Serial No. 62/754,696 and International Application No. PCT/US19/59647 on the basis of this work. G.D.P., R.K., and S.M.M. have started a company to commercialize the intellectual property. R.K. is a member of the SAB for GenCirq, Inc., holds an equity interest in GenCirq, and can receive reimbursements for expenses up to \$5,000/yr. R.K., A.D.S., and S.M.M. are directors at the Center for Microbiome Innovation at UC San Diego, which receives industry research funding for various microbiome initiatives, but no industry funding was provided for this cancer microbiome project.

Supplementary Information is linked to the online version of the paper at www.nature.com/nature. This includes **Tables S1–S8**.

(n=18,116 samples) from 33 cancer types in The Cancer Genome Atlas¹¹ (TCGA) for microbial reads, and found unique microbial signatures in tissue and blood within and between most major cancer types. These TCGA blood signatures remain predictive when applied to stage Ia-IIc cancer patients and cancers lacking any genomic alterations currently measured on two commercial-grade ctDNA platforms, despite very stringent decontamination analyses that discard up to 92.3% of total sequence data. We then independently show the ability to discriminate between and among non-cancer, HIV-, healthy controls (n=69) and multiple cancer types (prostate, lung, and melanoma, 100 total samples) solely using plasma-derived, cell-free microbial nucleic acids. We thus propose a new class of microbial-based oncology diagnostics, warranting further exploration.

Background

Cancer is classically considered a human genome disease^{12,13}. However, recent studies show significant microbial contributions in select cancer types, primarily fecal microbiome contributions to gastrointestinal cancers¹⁻¹⁰. However, the extent and diagnostic implications of microbial contributions across diverse cancer types remain unknown. Possible sample contamination during collection, processing, and sequencing limits these investigations: procedural controls have rarely been implemented in cancer genomics projects. Employing recently-developed tools¹⁴⁻¹⁸ to minimize contributions of contaminants to microbial signatures could enable rational development of microbially-based diagnostics.

To characterize the cancer-associated microbiome, we re-examined microbial reads from 18,116 samples across 10,481 patients and 33 cancer types from The Cancer Genome Atlas (TCGA) compendium of whole genome sequencing (WGS; n=4,831) and whole transcriptome sequencing (RNA-Seq; n=13,285) studies. Microbial reads were previously identified in *ad hoc* analyses — Epstein-Barr Virus (EBV) in stomach adenocarcinoma¹⁹, Human Papillomavirus (HPV) in cervical cancer²⁰ — and systematically on small subsets of samples, e.g. viromes of 4,433 TCGA samples across 19 cancer types²¹ and bacteriomes of 1,880 TCGA samples across 9 cancer types¹⁷. Most TCGA sequencing data remains unexplored for microbes. Here we present the most comprehensive cancer microbiome dataset yet created using two orthogonal microbial-detection pipelines, systematically measuring and mitigating technical variation and contamination. We use machine learning (ML) to identify microbial signatures discriminating cancer types, and compare their performance.

Because TCGA processing did not control for microbial contamination and excluded healthy individuals, we performed an additional analysis on the most contentious TCGA sample type analyzed (blood) using gold-standard microbiology protocols^{18,22}. We focused on plasma-derived microbial DNA to commensurably benchmark against clinically available ctDNA assays. Deep metagenomic sequencing on plasma samples from cancer individuals with prostate, lung, and skin cancers (n=100), and non-cancer, HIV-, healthy controls (n=69) suggested that healthy-versus-cancer and cancer-versus-cancer discriminations were possible from cell-free microbial profiles. These findings suggest a new class of microbial-based

cancer diagnostics that may complement existing ctDNA assays for cancer detection and monitoring.

Results

TCGA cancer microbiome and its normalization

Of 6.4×10^{12} sequencing reads in TCGA, 7.2% were classified as non-human, of which 35.2% were assigned to bacteria, archaea, or viruses with 12.6% resolved at the genus level by Kraken²³, which matches short genomic substrings (k-mers) to taxa in a reference database, representing 2.5% and 0.9% of total reads respectively (Fig. 1a; Tables S1–2; Extended Data Fig. 1a shows TCGA study abbreviations). After filtering samples for quality-controlled metadata (Fig. 1b) and normalizing by sample number within a cancer type and sample type (Extended Data Figs. 1f–g), WGS provided significantly more microbial reads than RNA-Seq experiments for primary tumor (p-value= 2.08×10^{-9}), solid-tissue normal (p-value= 1.26×10^{-7}), metastatic (p-value=0.0396), and recurrent tumor samples (p-value=0.0336; all two-sided Mann-Whitney). Fast kmer-matching approaches are prone to false positives, so we performed slower, but potentially more specific, genome alignments of Kraken-positive, genus-level microbial reads on which our findings are based for four TCGA cancer types (CESC, STAD, LUAD, OV) with known microbial relationships^{5,19,20} and/or with paired proteomic data²⁴ and found a low estimated false positive rate of 1.09% (Table S3), suggesting the Kraken data was valid for downstream analyses.

Substantial batch effects are known in TCGA expression and human genomic data^{25,26} and were replicated in metagenomic data (Fig. 1c, Extended Data Figs. 1b,d). Therefore, we implemented a pipeline that converted discrete taxonomical counts into log-counts per million (log-cpm) per sample using Voom²⁷, and performed supervised normalization (SNM) (Methods)²⁸. Principal variance components analysis^{29,30} showed that normalization reduced batch effects while increasing biological signal, including “disease type” (i.e. cancer type), above the individual technical variables (Figs. 1d–e; Extended Data Figs. 1c,e).

Predicting among and within cancer types

Using normalized data, stochastic gradient boosting ML models were trained to discriminate between and within cancer types and stages. The performance of these models was strong for discriminating (i) one-cancer-type-versus-all-others (n=33 cancer types) and (ii) tumor-versus-normal (n=15 cancer types) (Figs. 1f–g, Extended Data Figs. 2a–f; all performance metrics online: http://cancermicrobiome.ucsd.edu/CancerMicrobiome_DataBrowser/). Differences in sensitivities and specificities between cancer types may be partially attributed to different class sizes, as a significant linear relationship was evident in one-cancer-type-versus-all-others comparisons between the minority class size and AUROC (p-value=0.0231) and AUPR (p-value=0.0089) values (two-sided hypothesis tests of slope; Extended Data Figs. 2g–h). Cancer microbial heterogeneity may also contribute to this differential performance, although spatial examination of these historic tissue samples is beyond the scope of this study. Tissue-based microbial models performed well for discriminating stage I versus IV tumors (n=8 cancer types) for COAD, STAD, and KIRC,

but not the other five cancers tested (Fig. 1h), nor for discriminating intermediate stages (not shown). These results suggest that microbial community structure dynamics may not correlate with cancer stages as defined by host tissue for all cancer types.

To evaluate the generalizability of our approach across datasets, we randomly sorted raw TCGA microbial counts into two batches, repeated all procedures on each independently, tested each independently-trained model on the other half's data, and found highly similar performance (Extended Data Fig. 3a). Discriminatory microbial signatures held when examining singular methodologies (WGS or RNA-Seq), one sequencing center that performed either WGS (Harvard Medical School; HMS) or RNA-Seq (UNC) (Extended Data Fig. 3b–i), or using only genomic alignment-filtered Kraken data (Table S3; Extended Data Fig. 4a–h).

For further validation, we applied SHOGUN³¹, an alignment-based microbial taxonomic pipeline using a reduced, phylogenetically-based, bacteria-only database on 13,517 TCGA samples (WGS: n=3434; RNA-Seq: n=10,083), covering every analyzed cancer type (n=32), sample type (n=7), sequencing platform (n=6), and sequencing center (n=8) in the Kraken-based analysis. The SHOGUN-derived data replicated batch effects identified in Kraken-derived data despite a smaller, non-identical underlying database (Extended Data Figs. 4j–l). We input it, and a corresponding subset of Kraken-derived data (Methods), independently into our normalization and ML pipelines and found no major differences in discriminatory performances between the datasets (Extended Data Figs. 4m–t). Together, the results imply that microbial communities are unique to each cancer type and that our approach of normalization and model training to distinguish cancers based on microbial profiles alone can be applied more broadly.

Biological relevance of microbe profiles

Given the strong discrimination of microbial signatures, we sought evidence of biological relevance using ecologically-expected and/or clinically-tested outcomes. To assess whether cancer-associated microbes are ecologically-expected, i.e. part of the 'native' organ-specific commensal community, we trained a Bayesian microbial-source tracking algorithm³² on data from 217 samples across 8 body sites in the HMP2 project³³ processed with our microbial detection and normalization pipeline to estimate the body-site contribution from 70 solid-tissue normal samples in the COAD cohort and 122 SKCM primary tumors (Methods). Stool was the primary known body-site contributor only to COAD profiles (average mean fractional contribution=20.17%; SE=2.55%; Fig. 2a), but not SKCM profiles (one-tailed Mann-Whitney: p-value=0.0014; Extended Data Fig. 5b), suggesting a local source for part of the community.

Fusobacterium spp. is important in the development and progression of gastrointestinal tumors^{1,19,34,35} and *Fusobacterium* genus was overabundant in primary tumors compared to solid-tissue normals (all p-values 8.5×10^{-3}) and especially to blood-derived normals (all p-values 3.3×10^{-11}) (Fig. 2b). Pan-cancer analyses also showed an overabundance of *Fusobacterium* when comparing all broadly-defined gastrointestinal (GI) cancers in TCGA (n=8) against non-GI cancers (n=24) in both primary tumor tissue (p-value $<2.2 \times 10^{-16}$) and adjacent solid-tissue normal (p-value=0.031) (Fig. 2c; Extended Data Fig. 5a). Similar to

previous investigations in TCGA STAD¹⁹, we did not identify differences in *Helicobacter pylori* between primary tumors versus adjacent solid-tissue normals (p-value=0.72, not shown; all tests two-sided Mann-Whitney).

We then confirmed clinically annotated TCGA viral infections and compared our microbial-detection pipeline to studies that examined the TCGA virome with two different bioinformatic pipelines: (i) *de novo* metagenome assembly methods and (ii) read-based methods (PathSeq³⁶ algorithm)^{19,21}. There was differential abundance of the *Alphapapillomavirus* genus between primary tumors in individuals clinically tested as ‘positive’ or ‘negative’ for HPV infection in CESC and HNSCC samples (all p-values 3×10^{-9} ; two-sided Mann-Whitney; Figs. 2d–e). Blood-derived normal samples from CESC patients were used as negative controls and were not statistically different (p-value=0.99; two-sided Mann-Whitney), and selective overabundance for *Alphapapillomavirus* held when comparing across all other cancer types and sample types (Extended Data Fig. 5c). LIHC individuals with a prior history of hepatitis B had selective overabundance of the HBV genus (*Orthohepadnavirus*) in both primary tumors and adjacent solid-tissue normals compared to LIHC patients with a prior history of alcohol consumption and hepatitis C (*Hepacivirus* genus) (Fig. 2f; primary tumor p-values 2.8×10^{-7} ; solid-tissue normal p-values 0.011); blood-derived normals were used as negative controls and were not statistically different (p-values 0.44; all tests two-sided Mann-Whitney). Also in agreement with the previous reports¹⁹, the genus for EBV (*Lymphocryptovirus*) was selectively overabundant in EBV-infected primary tumors compared to patients assigned to other STAD molecular subtypes (Fig. 2g; p 2.2×10^{-16}). Solid-tissue normals and blood-derived normals were used as negative controls and were not statistically different (blood: p-values 0.52; tissue: p-values 0.096; all tests two-sided Mann-Whitney).

These data are consistent with the feature importance information found in our models predicting one-cancer-type-versus-all-others. Namely, cancers with known microbial “drivers” or “commensals” provided initial evidence that the models were ecologically-relevant; for example, *Alphapapillomavirus* genus was the most important feature for identifying CESC tumors; for COAD tumors, the *Faecalibacterium* genus; for LIHC tumors, the *Orthohepadnavirus* genus was the second most important feature (after the hepatotoxic *Microcystis* genus³⁷). For additional hypothesis generation, we created an interactive website to enable exploration of normalized microbial abundances found in TCGA cancers and major sample types (Extended Data Fig. 5d–e; http://cancermicrobiome.ucsd.edu/CancerMicrobiome_DataBrowser/). We provide raw and normalized microbial abundance datasets for public reuse, and anticipate the opportunity to integrate these with host multi-omic data to generate additional mechanistic hypotheses. Collectively, the findings provide ecological-validation of our bioinformatic and normalization approaches for viral and bacterial data while extending the results to many more samples and microbes.

Measuring and mitigating contamination

We recognize the importance of measuring and mitigating the potential effects of contamination to best characterize putative cancer-associated microbes^{14–18}. Previous work identified just six contaminants in TCGA (*Staphylococcus epidermidis*, *Propionibacterium*

acnes, *Ralstonia* spp., *Mycobacterium*, *Pseudomonas*, *Acinetobacter*) based on common low-read abundances across cancer types¹⁷, but recent literature demonstrated that external contaminants more consistently have frequencies inversely correlated with sample analyte concentration and can be detected using a robust statistical framework^{16,38}.

Based on the latter approach, we used (i) DNA and RNA concentrations calculated during TCGA sample processing (n=17,625) and taxon read fractions (n=1993) to identify putative contaminants and (ii) additionally removed genera typically found in “negative blank” reagents¹⁴ (n=94 genera) (Methods). Extended Data Fig. 6a outlines the approaches taken from surgical resection through bioinformatic processing; we also spiked five types of pseudo-contaminants into the raw dataset to track through decontamination, supervised normalization, and ML. Given known technical variation (Figs. 1c–e), we processed samples in batches by sequencing center (n=8) and removed taxa found to be a contaminant amongst *any* center. This identified 283 putative contaminants, including 19.1% (n=18 genera) of the reagent “blacklist”¹⁴. After combining these two lists (n=377 genera), we manually reviewed the literature (Table S6–7) to re-allow pathobiont genera or mixed-evidence genera (both a pathogen and common contaminant, e.g. *Mycobacterium*). This resulted in two datasets, one with “likely contaminants removed” and another with “all putative contaminants removed.” We also created a third “most stringent filtering” dataset that discarded ~92% of the total reads using a stricter filtering schema (Methods; Extended Data Fig. 6b). Finally, we grouped samples into individual sequencing plates at each center and removed all putative contaminants identified in any one “plate-center” batch (n=351; Table S8; Methods), in addition to the aforementioned reagent “blacklist” (497 total genera). Decontamination did not appear to differentially affect the samples types or cancer types under study (Extended Data Fig. 7).

We stress that these *in silico* decontamination methods are not substitutes for implementing gold-standard microbiology practices on cancer samples, including sterile processing, sterile-certified reagents, ‘negative blanks’ of reagents processed from start to finish, and multiple-sample pooling as “positive” controls^{18,22}. The *in silico* tools described here reflect the state of the art, but are not designed to detect abundant ‘spikes’ of contaminants or cross-contaminants. These latter contaminants should not drive uniform discriminatory signals between and within cancer types collected over many centers and years, but may limit biological conclusions, particularly in small studies, if not controlled.

Another risk with stringent decontamination is that real signals reflective of commensal, tissue-specific microbial communities and concomitant cancer-predictive microbial profiles may be discarded. To evaluate this concern, we re-calculated the body-site attribution percentages for COAD solid-tissue normals (n=70), and found that successively-stringent decontamination improves recognition of concomitant tissues before becoming unrecognizable (Extended Data Figs. 6c–f).

We recalculated all ML models shown in Figs. 1f–h and compared their performances before and after each decontamination approach (Extended Data Figs. 6g–i). Most models did not rely on spiked pseudo-contaminants (Extended Data Fig. 8a), though DLBC and MESO models (with very few available samples; Table S4) appear to be exceptions and may be

unreliable. As expected, comparisons where knowledge about the tissue type is informative (e.g. COAD-versus-all-other-cancer-types) generally have poorer performance with stringent decontamination, but within-tissue comparisons (e.g. tumor-versus-normal) often have equivalent or increased performance. These results suggest that stringent filtering may be desirable in certain comparisons, but a universal approach to decontamination may preclude biologically informative results.

Predictions using microbial DNA in blood

There is mounting evidence that blood-based microbial DNA (hereafter “mbDNA”) can be clinically informative in cancer^{39–44}, including those featuring blood-barrier or lymphatic disruptions (e.g. COAD)³⁹, but it is unknown how broadly this applies. Using WGS data from TCGA blood samples, we employed our ML strategies on the full and four decontaminated datasets and found that blood-borne mbDNA could discriminate between numerous cancer types (Fig. 3a), regardless of the microbial taxonomic algorithm and database used for classification or when using only genomic-alignment-filtered Kraken data (Extended Data Figs. 4g–h,s–t). Retrospective analysis showed that few models included spiked pseudo-contaminants for predictions (Extended Data Fig. 8b); models that did (CESC, KIRP, LIHC) may be less trustworthy.

Spurred by these findings, we sought to benchmark our ML models against existing cell-free tumor DNA (ctDNA) assays, focusing on circumstances where ctDNA assays fail: stage Ia–Ic cancers and tumors without detectable genomic alterations. After removing all blood-derived normal samples from patients harboring stage III or IV cancers, we built new ML models and found high cancer type discrimination using blood mbDNA (Fig. 3b). We further used gene-lists from the Guardant360[®] and FoundationOne[®] Liquid assays^{45,46} to filter out TCGA patients with 1 targeted modification (~70%; Extended Data Figs. 8c–e) and found high discriminatory performances using the same ML approach for most remaining cancer types (Figs. 3c–d).

These analyses are limited by the fact that ctDNA assays are plasma-based rather than whole-blood derived^{45,46}, and the distribution of mbDNA among blood compartments is unknown. It is impossible to assess if mbDNA was coming from live or dead microbes, as RNA data was unavailable, nor whether mbDNA is cell-free or in host leukocytes, as TCGA SOPs allowed whole blood or buffy coat extraction (Methods). It is also impossible to know the origin of blood mbDNA without examining primary specimens and, possibly, matched gut epithelia, as certain cancer types may ‘leak’ mbDNA non-intuitively (e.g. gut bacterial translocation in leukemia)^{8,10}. There is likely a continuum of ideal decontamination as the effect of decontamination on model performance varied across cancer types, but our filtering was limited by (i) not having access to the primary specimens, (ii) genus-level taxonomic resolution, and (iii) not knowing which non-TCGA samples were concurrently processed.

Validating microbial signatures in blood

To demonstrate the real-world utility of these results while commensurably benchmarking against plasma-based ctDNA assays^{45,46}, we evaluated the use of plasma-derived, cell-free mbDNA signatures to discriminate among healthy individuals and multiple cancer types in a

validation study while implementing gold-standard microbiology controls for low-biomass studies^{18,22}. Although plasma represents a distinct subset of whole blood not studied in TCGA, limiting direct comparability, it carries major advantages in archival stability (e.g. freezability), biorepository availability, and biological interpretation (i.e. non-living material). Our cohort included 69 non-cancer, HIV- individuals and 100 patients from three types of high-grade (stage III-IV) cancers: prostate cancer (n=59; “PC”); lung cancer (n=25; “LC”), and melanoma (n=16; “SKCM”) (Fig. 4a). Without prior literature to estimate effect sizes, we used independent simulations on TCGA blood samples from matched cancer types at The Broad Institute and HMS to estimate minimum sample sizes (Extended Data Fig. 9a; Methods). Cell-free DNA was extracted from these plasma samples with extensive controls^{18,22} (Extended Data Figs. 9b–c), and processed for whole metagenomic sequencing by a limited set of users, using a single library preparation method⁴⁷, in a single batch, in one deep-sequencing run. We performed human-read removal, classification of remaining reads by Kraken, stringent decontamination using both DNA concentrations and negative blanks¹⁶, and Voom-SNM. Demographic comparisons and permutation analyses suggested necessary normalization for age and sex (Extended Data Figs. 9d–e,h–j; Methods), and direct age regression performance showed mean absolute errors similar to the gut microbiome (unpublished results, Extended Data Fig. 9g). ‘Bootstrapping’ the same ML protocol used in the TCGA analyses showed strong, generalizable discrimination between healthy controls and grouped cancer patients (Fig. 4b; Methods). Due to small sample sizes, we performed leave-one-out (LOO) iterative ML on the normalized data and found high discriminatory performance in pairwise- and multiclass-comparisons between and among healthy controls and cancer types except for the smallest SKCM cohort (Figs. 4c–k). Therefore, we iteratively subsampled PC and LC groups to match SKCM cohort size and performed pairwise LOO discrimination of each cancer type against subsampled healthy controls (Extended Data Fig. 9k; Methods). PC and LC cohorts were still separable at SKCM cohort size (Mean_AUROC=0.891, 95% CI:[0.879,0.903]; Mean_AUPR=0.827, 95% CI:[0.815,0.839]; 100-iterations), revealing universal deficits in SKCM performance. This deficit may have a biological basis as SKCM was the second-worst performer in TCGA blood discriminations for four of five datasets tested (Fig. 3a), although this warrants further confirmation. To ensure that the microbial assignments by Kraken were valid, we repeated all bioinformatic, normalization, and ML steps using bacterial assignments from SHOGUN³¹ and its separate database⁴⁸, which showed highly concordant performances (Extended Data Fig. 10). We anticipate refinement of the taxonomic assignments for cfDNA signatures as concomitant microbial databases improve⁴⁹. Plasma microbial abundances detected are also explorable in our web interface (Extended Data Fig. 5d–e; http://cancermicrobiome.ucsd.edu/CancerMicrobiome_DataBrowser/).

Discussion

Collectively, our data suggest widespread associations between cancer and specific microbiota across diverse cancer types. These microbial profiles appear to discriminate within and between most cancer types, including with blood-based mbDNA at low-grade tumor stages and in patients without any detectable genomic alterations on commercial ctDNA assays. These results often remain valid even after extensive internal validation

checks and decontamination, at times discarding >90% of the total data. High discriminatory performance among healthy controls and multiple cancer types using only cell-free mbDNA in plasma, while adopting more extensive internal and external contamination controls than TCGA, suggests the feasibility and generalizability for clinically-relevant and retrospective testing with widely-available samples. More work is needed to evaluate whether the observed nucleic acids are coming from live microbes, host cells, or lysed bacteria in the tumor microenvironment and blood. Notably, many technical and biological factors limit analysis of retrospective cancer sequencing data for low-biomass microbes, and advancing this field will require collaborations between cancer biologists and microbiologists. Nonetheless, our results suggest that a new class of microbial-based cancer diagnostics may provide significant future value to patients.

Methods

TCGA data accession

All TCGA sequence data (Tables S4–5) were accessed via the Cancer Genomics Cloud (CGC) as sponsored by SevenBridges (<https://cgc.sbgenomics.com>)⁵¹. Details of how these samples were acquired and processed are comprehensively described elsewhere⁵². Standard operating procedures (SOPs) for TCGA were accessed via the NCI Biospecimen Research Database (<https://brd.nci.nih.gov/brd/sop-compendium/show/701>). Matched patient metadata, including molecular subtypes, were accessed via the CGC through both SevenBridges and the Institute for Systems Biology (ISB; <https://isb-cgc.appspot.com/>)⁵³, via the TCGAMutations R package⁵⁴, or were taken directly from their respective TCGA publication's supplemental data^{19,55}. Genomic alteration statuses for all TCGA patients were queried and downloaded via the cBioPortal^{56,57}. Gene panels for commercial ctDNA assays were accessed from company white papers for the Guardant360® assay (https://www.therapysselect.de/sites/default/files/downloads/guardant360/guardant360_specification-sheet_en.pdf) and the FoundationOne® Liquid assay (https://assets.ctfassets.net/vhribv12lmne/3SPYAcBgdqAeMsOqMyKUog/d0eb51659e08d733bf39971e85ed940d/FIL_TechnicalInformation_MKT-0061-04.pdf). For TCGA metadata accession and transformation from hierarchical formats to flat tables, custom Python scripts (available on Github; <https://github.com/biocore/tcga>) were written to query SevenBridges's metadata ontology and organize the data where possible; for information not stored in that ontology, we used the ISB's CGC R programming language API⁵³ to access its recent metadata release (tcga_201607_beta.Clinical_data).

Kraken TCGA microbial-detection pipeline

The SevenBridges CGC interface enabled rapid development of the bioinformatic pipeline for this project while ensuring its future reproducibility⁵¹. Bioinformatic tools were either loaded directly from the CGC platform (e.g. samtools, BWA) or uploaded and run as separate Docker containers (e.g. QIIME, Kraken) in order to create customized 'app' workflows. These 'app' workflows take sample BAM files as inputs and labeled which DNA or RNA reads within each sample were "microbial". These 'app' workflows can be publicly shared for reproducibility purposes, as needed. The computational analyses themselves were hosted on Amazon Web Services (AWS; <https://aws.amazon.com/>) through the CGC

interface and most often used AWS's "x1.16" EC2 compute instance, comprised of the following specifications: 64 vCPU, 174.5 ECU, 976 GB of Memory, and 1920 GB of Instance Storage. The computational wall-time was approximately 6-months using these specifications.

Sequencing reads that did not align to known human reference genomes (based on mapping information in the raw BAM files) were mapped against all known bacterial, archaeal, and viral microbial genomes using the ultrafast Kraken algorithm²³. A total of 71,782 microbial genomes were downloaded using RepoPhlan (<https://bitbucket.org/nsegata/repophlan>) on 14 June 2016, of which 5,503 were viral and 66,279 were bacterial or archaeal. Based on prior literature, bacterial and archaeal genomes were filtered for quality scores of 0.8 or better⁵⁸, which left 54,471 of them for subsequent analysis, or a total of 59,974 microbial genomes.

As previously described in detail²³, the Kraken algorithm breaks each sequencing read into k-mers (we used default 31-mers) and exactly matches each k-mer against a database of microbial k-mers, which was built from the 59,974 microbial genomes described above prior to running the algorithm. The set of exact k-mer matches for a given read, in turn, provides a putative taxonomy assignment of the lowest common ancestor for that read, most accurately to the genus level, to which we summarized our data. The matching and classification operations are orders of magnitude faster than performing direct genome alignments. As a safeguard against false positives and to properly benchmark our pipeline, we took four cancer types (COAD, CESC, OV, LUAD) and aligned the reads Kraken classified as "microbial" to the 59,974 microbial genomes using BWA⁵⁰, which is computationally more expensive but yields a result with higher specificity and taxonomic resolution (i.e. to species and strain level). The four cancer types that were directly aligned included CESC as a putative positive viral control (i.e. HPV), STAD as a putative positive bacterial control (i.e. *H. pylori*), and two others (LUAD, OV) based on microbial signatures in the literature and/or available mass-spectrometry proteomic information (to look for microbial proteins; data not shown)^{5,24,59-61}. We found that 98.91% of reads classified to genus level or lower by Kraken (on which our main findings are based) also aligned with BWA to the microbial database (bacteria, archaea, viruses; see Table S3), or a false positive rate of 1.09%, suggesting that the genus-level, Kraken-labeled, pan-cancer microbial reads were sufficiently usable for further analyses.

SHOGUN TCGA bioinformatic processing

To evaluate the robustness of cancer type discriminations using different taxonomic identification algorithms, we utilized a previously published shallow shotgun taxonomic assignment approach (co-developed by Q.Z. & R.K.)³¹ and a separate phylogeny-centric database called 'Web of Life' (WoL; developed by Q.Z. & R.K.; n=10,575 bacterial and archaeal genomes; <https://biocore.github.io/wol/data/genomes>)⁴⁸ on TCGA samples. SHOGUN utilizes computationally intensive direct genomic alignments for taxonomy assignments rather than an ultrafast kmer-based approach like Kraken. To reduce processing time for TCGA samples, reads classified as microbial in origin by Kraken were used as input for the SHOGUN³¹ align function, which used Bowtie2⁶² to map reads against the 'Web Of Life' database⁴⁸ to generate taxonomy profiles. 13,517 total samples (WGS: n=3434; RNA-

Seq; n=10,083) were processed, covering all TCGA cancer types (n=32), sample types (n=7), sequencing centers (n=8), and sequencing platforms (n=6) under study in the Kraken analysis including 21 TCGA cancer types (n=9444 samples) that had all samples in the Kraken analysis re-analyzed by SHOGUN. Profiles were then collapsed to the genus level using QIIME 2⁶³. Analyses were run on a local compute cluster comprised of 1024 Intel Ivy-bridge compute cores, as well as 384 AMD compute cores, and 12TB of total RAM with a 10GbE compute network for approximately 5-months of computational wall-time; typical job submissions for a single cancer type utilized ~30 cores and ~250GB of RAM.

Quantitative measurement and normalization of TCGA technical variation

Cognizant of how technical variation between TCGA sequencing centers (n=8), sequencing platforms (n=6), experimental strategy (WGS vs. RNA-Seq), and possible contamination could confound our results, we developed a pipeline to quantify and remove batch effects while maintaining or increasing the signal attributed to biological variables. Briefly, we filtered out samples with poor metadata quality (i.e. missing race or ethnicity, ICD10 codes, DNA/RNA analyte amounts, or FFPE status information); transformed our discrete taxonomical count data to approximately normally-distributed, log-count per million (log-cpm) data using the Voom algorithm²⁷, which models and removes the data's heteroskedasticity; and lastly performed supervised normalization (SNM) on the data to remove all significant batch effects while preserving biological effects²⁸. Voom is traditionally used in combination with limma⁶⁴ for differential expression (or abundance) analysis of discrete count data, but we only used it for the algorithmic transformation to 'microarray-like' data, which permitted subsequent SNM. The Voom and SNM model matrices were equivalent and built using "sample type" as the target biological variable (n=7; e.g. primary tumor tissue) due to expected biological differences between them, for which signal should be preserved during the SNM; conversely, the following were modeled as technical covariates to be mitigated during SNM: sequencing center (n=8), sequencing platform (n=6), experimental strategy (n=2), tissue source site (n=191), and fixed-formalin paraffin-embedded (FFPE) status (n=2; 'YES' or 'NO'). It was not possible to model "disease type" as the target biological variable due to complete confounding between certain cancer types and sequencing centers (i.e. some cancer types were only sequenced at one TCGA site). During the Voom transformation, weighted trimmed mean of M-values (TMM) normalization from the edgeR package⁶⁵ was employed for most data ("full dataset", "likely contaminants removed" data, "plate-center decontaminated" data, and "all putative contaminants removed" data) while dropping unvarying features (filterByExpr() function; edgeR), as shown by limma's user guide (<https://www.bioconductor.org/packages/devel/bioc/vignettes/limma/inst/doc/usersguide.pdf>; Chapter 15: RNA-Seq Data, p. 70). In other cases ("most stringently filtered" data, "SHOGUN TCGA data", "Kraken TCGA data matched to SHOGUN TCGA data", and both plasma microbiome datasets), quantile normalization was utilized since downstream SNM correction was not compatible with stringently filtered TMM normalized, feature-dropped data, as these datasets already had significantly reduced or low feature counts. With the exception of "most stringently filtered" data, all quantile normalized datasets were compared only to other quantile normalized datasets. Principal components were calculated before and after SNM correction of the Voom-adjusted data, and principal variance components analysis (PVCA)^{29,30} quantified

these changes between raw count data, Voom-adjusted data, and Voom-SNM normalized data. The mathematical basis for PVCA is well described by NIEHS (<https://www.niehs.nih.gov/research/resources/software/biostatistics/pvca/index.cfm>), and we set the one tunable parameter to 80%, based on their recommendation of 60–90%.

Using SourceTracker2 as a validation analysis to address contamination concerns

Shotgun sequencing data from the NIH's Integrative Human Microbiome Project³³ ("HMP2"), which swabbed eight different body sites among 217 total samples, were downloaded and run using the same TCGA Kraken microbial-detection pipeline as described above, including against the same microbial database (n=59,974 bacterial, archaeal, and viral metagenomes) for taxonomy assignments. HMP2 data were summarized at the genus-level, as per our TCGA cancer microbiome data, and then were used to train a Bayesian source tracking model (SourceTracker2;<https://github.com/biota/sourcetracker2>)³². Details of the Bayesian model have been previously described by our lab³². Using SourceTracker parlance, these HMP2 samples served as "sources" while the Voom-SNM normalized samples acted as "sinks," and the SourceTracker algorithm was used to calculate the proportion of each "source" attributable each "sink." In lay terms, we estimated the proportion of body site from HMP2 data attributable to each Voom-SNM normalized cancer microbiome sample using the Bayesian model. After (i) intersecting the genera in our cancer microbiome dataset with those in HMP2, (ii) converting the log₂-cpm normalized values to scaled relative abundances (scaled by 10⁶ to give approximately 1 million total reads, as HMP2 data had 917,450 reads), and (iii) converting the data to BIOM table format⁶⁶, we applied the model on solid-tissue normal samples from the TCGA COAD cohort (n=70) and on primary tumor SKCM samples (n=122). SKCM primary tumor samples were chosen instead of solid-tissue normal samples as the best proxy of skin flora, since only one adjacent solid-tissue normal sample for SKCM was available (see Table S4). SourceTracker2 default settings (alpha1=0.001, alpha2=0.1, beta=10, restarts=10, draws_per_restart=1, burnin=100, delay=1) were utilized for both runs. The outputs were calculated in terms of mean fractional contributions of each source to each sink; averages and standard errors of these values were subsequently calculated. Statistical differences between fecal contribution to COAD and SKCM samples (Extended Data Fig. 5b) were calculated using a one-sided Mann-Whitney test. The above protocol was repeated for the four decontaminated datasets to generate Extended Data Figs. 6c–f.

TCGA ML methods

Stochastic gradient boosting machine (GBM) learning models were trained, automatically tuned, and tested using the R programming language (<https://www.r-project.org/>), GBM package^{67,68}, and Caret package⁶⁹. Training and testing occurred on separate, randomly selected, stratified sampling splits of 70% and 30% of the data, respectively, and a fixed random number seed was used to ensure reproducibility of the model results and comparability among models. During model training, the data were first centered and scaled for each sample to have mean zero and unit standard deviation; four-fold cross validation was employed to create multiple subsets of the training data and to perform a basic grid search optimization of GBM parameters, including interaction depth (1, 2, or 3) and number of trees (50, 100, or 150), while maximizing AUROC of the final, fully trained model.

Learning rate (“shrinkage”) was held constant at 0.1 and the number of minimum observations per node was fixed at 5. In cases of class imbalance, we up-sampled the minority class during training to help the model generalize after observing that other methods (i.e., differential class weighting, downsampling the majority class, minority class interpolation) did not consistently improve performance (data not shown). Comparisons were not made when the minority class contained fewer than 20 samples total due to the inability of the classifiers to be adequately trained and tested with so few samples. Final model performances, including ROC curves, PR curves, and confusion matrices (generated based on a probability cutoff of 50% for class #1 versus class #2 discrimination), were generated by applying the final model on the unseen 30% holdout test set. ROC and PR curves as well as AUROC and AUPR values were calculated using the PRROC package⁷⁰ while confusion matrices were calculated using the Caret package⁶⁹. Variable importance scores of the resultant, non-zero model features were estimated using the GBM and Caret packages^{67–69}. Percent contribution of a particular feature to the model’s predictions was estimated by dividing that feature’s variable importance score by the sum of all variable importance scores for a given model (cf. Extended Data Figs. 8a–b).

TCGA ML benchmarking and generalizability

As a benchmarking and generalizability assessment, we split TCGA in two stratified data halves (across sequencing center, sample type, and disease type) of raw Kraken-derived, genus-level microbial count data (split #1: n=8814; split #2: n=8811 samples), ran them both separately through our Voom-SNM protocol, built separate ML models on each normalized half, and then tested these tuned ML models on each other’s normalized data. We then compared these model performances against a third ML model that was built on the full Voom-SNM normalized dataset (n=17,625 samples) and used 50%–50% training and testing splits. Final performance was compared across all three approaches using their respective 50% holdout test set AUROC and AUPR. For additional internal validation, we built models predicting one-cancer-type-versus-all-others using just (i) RNA samples or (ii) DNA samples, as well as on (iii) samples from one sequencing center that only did RNA-Seq (UNC) or (iv) DNA-Seq (HMS) (Extended Data Fig. 3).

TCGA decontamination analyses

Broadly speaking, there are two classes of possible contamination that affect next-generation sequencing data: external contamination (e.g. reagents, investigators’ or subjects’ bodies, environmental contributions) and internal contamination (i.e. cross-contamination between samples during processing or sequencing)^{14,16}. Our overall decontamination approach attempts to (i) simulate contamination to estimate its contribution to predictive performance and/or model unreliability, (ii) mitigate external contamination as much as possible, and (iii) measure the degree of internal contamination using sensible positive and negative controls. External contaminants were identified and removed using sample analyte concentrations for all TCGA samples (n=17,625), as recently described in the literature^{16,38}, and by using a “blacklist” of microbes identified from reagents in sequencing kits similar to those used in TCGA¹⁴. Internal contaminants are particularly difficult to identify without having access to the primary samples or knowing which other samples (especially non-cancer samples) were run at the same time. As such, the only internal contaminants that were identified and

removed as clear cross-contaminants were 4 reads assigned to the *Ebolavirus* genus (2 reads from 1 TCGA-LGG sample at The Broad Institute and 2 reads from 1 TCGA-HNSC sample at HMS), almost certainly from concurrent studies on the 2014 West Africa outbreak at these same sequencing centers during the TCGA study collection period (2006–2016)^{71,72}, and 4 reads assigned to the *Marburgvirus* genus (from 2 TCGA-OV samples at The Broad Institute), also likely of similar origin or as false positives (i.e. *Ebolavirus* and *Marburgvirus* are both of the *Filoviridae* family). Doing so is in line with our previously published work that removes microbial assignments that cannot be related to the biology at hand⁷³. It is further unlikely that such cross-contaminants, especially of extremely low abundance, would drive uniform discriminatory signals between and within cancer types collected over many centers and years. For other possible cross-contaminants, we relied on estimating their contribution using Bayesian analyses (described above) of ecologically expected communities rather than their identification and removal.

Firstly, we spiked five pseudo-contaminants into the raw dataset (Extended Data Fig. 6a top right) to track them through decontamination, SNM, and ML. This included the following: (1) 1000 reads across all samples from Harvard Medical School (HMS); (2) 1000 reads across all samples from HMS, Baylor College of Medicine, Washington University School of Medicine, and Canada's Michael Smith Genome Sciences Centre; (3) 1000 reads across all samples from all sequencing centers; (4) 10^6 reads spiked across 100 randomly selected samples from HMS; (5) 10^6 reads spiked across 1000 randomly selected samples from all sequencing centers. The mean raw read count across all samples and taxa was 1481.20, so pseudo-contaminants containing 1000 reads can be considered 'low-level' background while those with 10^6 reads are considered 'high-abundance' spikes. If pseudo-contaminants are present in downstream ML models after training, three interpretations are available: Evaluate the percent predictive contribution of the pseudo-contaminants via feature importance scores and decide if it is negligible or not; eliminate any ranked model features below the pseudo-contaminant; or, most conservatively, flag the entire model as being unreliable.

Since TCGA did not include any "negative blank" reagent tubes during sample processing, we next attempted to pair a microbial "blacklist" at the genus-level that used similar reagents and/or library prep kits. TCGA SOPs mainly employed QIAGEN products (Qiagen, Valencia, CA) for extracting DNA and RNA in tissues (DNA/RNA AllPrep kit) and DNA in blood (QiaAmp Blood Midi Kit)⁵². Salter and colleagues¹⁴ described such a list (n=94 genera) for DNA extraction kits in metagenomic experiments, including from QiaAmp kits that used the same silica membrane-based DNA purification as those employed in TCGA blood extractions, obtained across four years of "negative blank" sequencing and three high-throughput sequencing centers. Additional putative external contamination was identified on the basis that sequences from contaminants generally have frequencies that are inversely correlated with sample analyte concentration^{16,38}. A robust statistical framework recently validated this principle¹⁶, providing the opportunity to exploit sample DNA/RNA concentrations recorded by TCGA as a means to identify putative contaminants. The two main assumptions of this framework are (i) the contaminants are added in uniform amounts across samples and (ii) the amount of contaminant DNA/RNA is small relative to the true sample DNA/RNA (microbial or host). Filtering was then conducted using the associated *decontam* R package (<https://github.com/benjjneb/decontam>)¹⁶ using the recommended

hyperparameter threshold ($P^*=0.1$) and a more stringent approach ($P^*=0.5$). Note, $P^*=0.5$ means that taxonomies are classified as “contaminant” or “not” if the contaminant model or non-contaminant model fit the distribution better. Since we found sequencing center to contribute significant variation to the raw count data, we processed the data in batches corresponding to them, whereby a taxon identified as a contaminant at *any* center was subsequently discarded for all centers (i.e. `batch.combine=“minimum”` in *decontam*). Putative lists of contaminants ($P^*=0.1$: $n=283$ genera; $P^*=0.5$: $n=1818$ genera) were then combined/intersected with the microbial “blacklist” ($n=94$ genera) and subtracted from the full dataset. Manual literature inspection of the smaller combined contaminant list ($n=377$) re-allowed 89 genera that were potentially pathogens or commensals (Table S6). This resulted in three new datasets: “likely contaminants removed,” “all putative contaminants removed,” “most stringent filtering.” As a further conservative measure, we took all TCGA sample barcodes (e.g. TCGA-02-0001-01C-01D-0182-01; as shown on NCI’s documentation https://docs.gdc.cancer.gov/Encyclopedia/pages/TCGA_Barcode/) and extracted all sequencing plate-sequencing center combinations, as named by the barcode’s last two sets of integers (i.e. plate 0182 from center 01, or 0182–01, in this example). Since *decontam* calculates the equivalent of a linear regression between taxon read fractions and analyte concentrations for all samples in a batch to determine if a given taxon is classified as a “contaminant,” we required more than 10 samples per plate-center combination to qualify as a batch, giving 351 total plate-center batches. A $P^*=0.1$ was used (default value), and, as before, if a taxon was identified as a contaminant in any one of the 351 batches (`batch.combine=“minimum”`), it was removed from the dataset ($n=421$ taxa removed; Table S8). After intersecting with the microbial “blacklist,” a total of 497 genera were removed. This provided the fourth decontaminated dataset, and all of them were then processed through the same SNM and ML pipelines described above.

Comparing ML performances between BWA, SHOGUN, and Kraken data

BWA filtering occurred against the same database used to generate the Kraken-based assignments ($n=59,974$ microbial genomes [bacteria, archaea, viruses]). Then, filtered BWA microbial count data were batch corrected the same way as the Kraken data via Voom-SNM, except that DNA and RNA data were normalized separately due to confounding between experimental strategy and sequencing center of the reduced sample count. Samples from the raw Kraken-derived data were then matched against samples processed by BWA and normalized the exact same way as the BWA data. This resulted in a total of four normalized datasets: DNA BWA data, RNA BWA data, DNA Kraken-subsetted data, and RNA Kraken-subsetted data. All four normalized datasets were then inputted for ML and their performances were compared to each other (Extended Data Fig. 4a–h).

The ‘Web of Life’ database⁴⁸ used for SHOGUN taxonomy assignments did not contain viruses, and SHOGUN processed a subset of all TCGA samples evaluated by Kraken (13,517 vs 17,625 samples). Thus, to make a fair comparison between their downstream ML performances, raw Kraken count data were subsetted to remove all identified viruses and to match the same samples processed by SHOGUN. Both datasets were then identically normalized by Voom (using quantile normalization) and SNM algorithms (using the same

biological and technical variables as in the main TCGA analysis described above) before being fed into the ML pipelines for discrimination between and within cancer types.

Complementary diagnostic analyses

When evaluating the applicability of blood mbDNA to low-grade cancers, all patients with stage Ia-c and IIa-c classified tumors were grouped together while discarding all others. For comparisons against the Guardant360® and FoundationOne® Liquid ctDNA assays, all TCGA patients containing *at least one* genomic alteration evaluated on their coding gene panels were filtered out; this included whether mutations were considered to be “passengers” or “drivers.” Remaining patients were used for ML analyses as described above.

TCGA simulations to estimate required sample sizes for validation study

To estimate the number of required samples from prostate, lung, and skin cancer (melanoma) for discrimination, we performed empirical simulations on TCGA blood samples at two different sequencing centers (Broad, HMS) that were all sequenced on one type of platform (Illumina HiSeq). We first used Kraken-derived microbial count data and then repeated the simulations with SHOGUN-derived microbial count data. This most closely mimicked expected real-world experimental conditions of the validation study.

First, all TCGA PRAD, LUAD, LUSC, and SKCM blood samples at Broad and HMS that were sequenced on Illumina HiSeq machines were subsetted from the raw Kraken data of microbial counts (Broad: n=99; HMS: n=288). Our lung cancer samples from author S.P.P. were of mixed origin so we combined LUAD and LUSC blood samples into a single non-small-cell-lung cancer (NSCLC) umbrella disease type; however, this only applied to Broad samples, as all blood-derived lung cancer samples at HMS were LUAD in origin. This left a breakdown of samples as follows: {HMS: 66 LUAD, 104 PRAD, 118 SKCM; Broad: 42 NSCLC [24 LUAD, 18 LUSC], 17 PRAD, 40 SKCM}. Then, each raw count dataset for HMS and Broad was independently normalized through Voom (using quantile normalization) and SNM algorithms, using “disease type” as the biological variable of interest and “tissue source site” as the technical variable, as all other technical factors were precluded by picking a single sequencing center, data type, and platform.

The simulations were performed as follows on the normalized datasets: (1) Random stratified sampling picked equal numbers of samples from the three classes; (2) one sample of the three class subsample was left out; (3) a ML model was built on all the remaining samples in the subsample and applied on the left out sample to make a prediction with a certain probability; (4) steps 2–3 were repeated until all samples had been iterated through; (5) using the list of observed classes and list of predicted classes along with their probabilities, multi-class performance metrics were estimated; (6) another stratified random sample was selected of the same sample size and steps 2–5 were repeated 9 more times (i.e. total of 10 times) to estimate standard errors of the multi-class performance metrics; (7) steps 1–6 were repeated for individual class sample sizes of 5 through 40 with a step size of 5 samples. In cases where the stratified sampling size was larger than the number of samples in a class, all samples in that class were used. Collectively, this provided an estimate of the number of samples to perform multi-cancer discrimination well (Extended Data Fig. 9a).

The empirical performance estimates (mean AUROC, mean AUPR) suggest that having at least 15 samples per cancer class should be sufficient. Note that it was not possible to estimate an ideal sample size for healthy controls since TCGA did not include them.

Clinical cohort selection and IRB protocols numbers

A total of 169 patient biobanked, frozen plasma samples were analyzed as part of this study, all from UC San Diego. All studies were approved by the Institutional Review Board (IRB) at UC San Diego, and under their respective IRB-approved protocols, patients provided written informed consent for sample donation and study. All prostate cancer plasma samples (n=59) came from R.M. under IRB protocol 131550. All lung cancer and melanoma plasma samples came from S.P.P. under IRB protocol 150348. All non-cancer, HIV-, healthy control subjects (n=69) came from a group led by R.H. under the following IRB protocol numbers: 130296, 091054, 172092, 151057, and 182064.

Plasma-derived, cell-free microbial DNA sample processing, and sequencing

Total circulating DNA was extracted from a volume of 250 μ L of plasma from each sample using the QIAamp Circulating Nucleic Acid Kit (QIAGEN) according to the manufacturer's instructions. And purified with AMPure XP SPRI paramagnetic beads (Beckman Coulter). Sequencing libraries were prepared from purified cfDNA using the KAPA HyperPlus Kit (Kapa Biosystems) with standard Illumina indexed adapters (IDT) as detailed in Sanders J. *et al.* (2019)⁴⁷. Sample libraries were characterized using the Agilent 4200 TapeStation System (High Sensitivity DNA Kit) and quantified by qPCR using the NEBNext® Library Quant Kit for Illumina® (New England Biolabs). Paired-end 2 \times 150 bp sequencing was performed on a NovaSeq 6000 instrument (Illumina), and samples were pooled across all four lanes during sequencing.

Bioinformatic processing for plasma microbiome samples

A total of 21,600,141,264 reads were generated on the single NovaSeq 6000 sequencing run across all samples. 19,046,611,360 reads were assigned to human samples (i.e. negative and positive controls removed), of which 2.186% of the total reads were classified as “non-human”. Raw sequencing data were demultiplexed and adapter-trimmed using Atropos⁷⁴. Additional quality filtering was done using Trimmomatic with the following settings—(ILLUMINACLIP:TruSeq3-PE-2.fa:2:30:7, MINLEN:50, TRAILING:20, AVGQUAL:20, SLIDINGWINDOW:20:20)⁷⁵. An additional adapter sequence consisting of a string of only G was added to the standard TruSeq3 adapters to remove trailing G stretches from the 5' end of reads. Read pairs were discarded if either mate mapped to the human genome (major-allele-SNP reference from 1000 Genomes Project) using Bowtie2 with the fast-local parameter set^{62,76}. Paired-end reads were then merged using FLASH with the following parameters—(minimum overlap: 20, maximum overlap: 150, mismatch ratio: 0.01)⁷⁷.

The filtered, merged reads were then processed either by Kraken, using the same workflow and database (n=59,974 microbial genomes) detailed above, or with SHOGUN as detailed here. Samples were processed on individual plasma microbiome samples (i.e. on a per-sample-per-lane basis since samples were pooled across all four sequencing flow cells during the run). After per-sample-per-lane taxonomy assignment by Kraken or SHOGUN,

microbial counts across lanes were aggregated for each sample after hierarchical clustering procedures showed consistent grouping by sample IDs rather than by flow cell lane. For SHOGUN-derived data, both successfully merged and unmerged reads were used as input for the SHOGUN align function, using Bowtie2 to map reads against the Web Of Life database to generate taxonomy profiles (<https://biocore.github.io/wol/data/genomes/#>)³¹, which were then collapsed to the genus level using QIIME 2⁶³. Taxonomy profiles of each sample were then filtered to remove all taxa whose relative abundance was less than 0.01%.

Plasma microbiome technical validation and data decontamination

To evaluate the performance of the sequencing run and bioinformatic microbial-detection pipelines, spiked wells and experimental serial dilutions of *Aliivibrio fischeri* (genus: *Aliivibrio*) included on the sequencing plate were examined against other sample types for differential abundance and in isolation for log-fold changes in abundance across dilutions. These technical “positive” controls are plotted in Extended Data Figs. 9b–c for both Kraken and SHOGUN-derived taxonomy assignments.

Three kinds of “negative” blank controls were included on the sequencing plate: (1) DNA extraction blanks, which had reagents from the DNA extraction stage through sequencing; and (2) DNA library preparation blanks, which had reagents from the library preparation stage through sequencing; and (3) empty control wells, which had water added to them and then reagents during library preparation and would contain splashed and/or aerosolized microbial nucleic acids. As used in the TCGA analysis, *decontam*¹⁶ was again used to decontaminate the plasma microbial data, except that it had access to both “negative” blank controls and DNA concentrations for all samples (excluding empty control wells for the latter). As a conservative measure, we used a $P^*=0.5$ hyperparameter value for *decontam* for both “prevalence” (i.e. blank-based) and “frequency” (i.e. concentration-based) modes of decontamination; this hyperparameter value is equivalent to the “most stringent decontamination” in TCGA that discarded >90% of the total data. For “prevalence” mode, a $P^*=0.5$ will flag any taxon that is more prevalent in “negative” controls than biological ones as a contaminant; for “frequency” mode, a $P^*=0.5$ will flag any taxon whose model (i.e. a regression model) fits a contaminant distribution better than a non-contaminant distribution using read fractions and DNA concentrations¹⁶ For Kraken count data, “prevalence” mode discarded 21 taxa and “frequency” mode discarded 1261 taxa (out of 1753 original assignments); for SHOGUN count data, “prevalence” mode discarded 57 taxa and “frequency” mode discarded 244 taxa (out of 1181 original assignments). Decontaminated data for both Kraken and SHOGUN were fed into downstream normalization and ML pipelines.

Plasma microbiome data normalization, permutation testing, and ML

An attempt to predict age using raw microbial count data was performed using gbm ML models (architectures same as those described above for TCGA) and leave-one-out (LOO) iterative ML (Extended Data Fig. 9g).

To confirm the importance of normalizing for age and gender in this cohort, we performed a permutation analysis with 100 iterations for each factor and then simultaneously for both

factors (Extended Data Figs. 9h–j). Briefly, the following four steps were performed: (1) Randomly swap age and/or sex labels among all samples; (2) run Voom-SNM on the raw data, using disease type as the biological variable of interest and permuted age and/or sex as the technical factors; (3) perform a ML analysis to discriminate grouped cancer samples from healthy controls using 70%/30% training/testing splits with a fixed random number seed and internal 4-fold cross validation to obtain a two-class performance estimate (AUROC, AUPR); (4) repeat steps 1–3 for a total of 100 times to create a null performance distribution. Next, using correct, fixed age and/or sex assignments, we ran steps 2–3 for a total of 100 times while randomly selecting the random number seed in step 3. Lastly, this performance distribution was directly compared to its null distribution for significance using a two-sided Mann-Whitney test. Since all of these tests were extremely significant (all p-values 1.5×10^{-13}), we incorporated age and sex as technical factors in the Voom-SNM while holding disease type as the biological variable of interest. Note, all lung cancer samples were labeled with a consolidated disease type label during normalization regardless of pathological subtype, as done in the TCGA cancer simulations (described above). All “negative blank” and “positive monoculture” controls were removed prior to Voom-SNM.

ML on the Voom-SNM normalized plasma microbiome samples was done exactly as previously described for TCGA samples, except for the sampling schema, due to the orders of magnitude smaller sample sizes. First, to estimate generalization of healthy versus grouped cancer discriminations, we ‘bootstrapped’ 70%/30% training/testing splits with 4-fold cross validation during training for 500 iterations. ‘Sampling with replacement’ was allowed in that every training/testing split (i.e. every iteration) was unique; however, in no case was a sample allowed to be both a training case and a testing case. Summary statistics on the resultant performance metrics from all 500 iterations estimated the AUROC and AUPR distributions and confidence intervals (Fig. 4b; Extended Data Fig. 10a). Second, pairwise and multi-class discriminations between and among healthy controls and individual cancer types were done with leave-one-out (LOO) ML. In other words, one sample was iteratively left out, a model was iteratively trained on the remaining samples with 4-fold cross validation for hyperparameter tuning, and a prediction was iteratively made on the left out sample with a probability given by the model. The final list of actual classes for all samples were compared to the list of predicted classes and their probabilities to estimate AUROC and AUPR metrics, as described previously using the PRROC R package⁷⁰. Multi-class performance was estimated by taking the mean of all 1-versus-all-others comparisons, as reported by the multiClassSummary() function in the caret R package⁶⁹.

Iterative subsampling to evaluate the contribution of smaller sample sizes on the melanoma cohort performance (Extended Data Fig. 9k) was done as such: (1) Perform random stratified sampling of a single cancer type and healthy controls of 16 samples each (32 total); (2) perform LOO iterative ML and evaluate performance on those 32 samples for healthy vs. cancer discrimination; (3) repeat steps 1–2 one-hundred times to estimate performance standard errors; (4) repeat steps 1–3 for each of the three cancer types. The same process was also done for iterative subsampling of PC and LC cohorts to study the impact of decreased sample size on their discrimination. Note that the entire melanoma cohort was used during each stratified subsampling, since the goal was to compare its cohort size to the other sample sizes.

Statistical analyses

All statistical analyses were done using R version 3.4.3. The ggpubr package (<https://github.com/kassambara/ggpubr>) performed nonparametric statistical testing between groups and accounted for multiple hypothesis testing correction when necessary. Note that p-values less than 2.2×10^{-16} cannot be accurately calculated by R, so p-values less than this are listed as “ $p < 2.2 \times 10^{-16}$ ”; it is not a range of p-values. Measurements were taken from distinct samples and not by repeatedly measuring samples. Sample size estimates for the prospective validation study came from empirical simulations with TCGA blood samples and relied on the GBM package^{67,68}, Caret package⁶⁹, and MLmetrics package (<https://github.com/yanyachen/MLmetrics>) for performing ML and multi-class performance estimation. All other multi-class performance estimates were calculated using the Caret⁶⁹ and MLmetrics packages.

Reporting summary

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

Website

The website referenced in this manuscript can be accessed at http://cancermicrobiome.ucsd.edu/CancerMicrobiome_DataBrowser/. Links are directly available on the website to the data repository as well as the code hosted on GitHub (upper right). Additionally, there are five separate tabs (left-hand side) for interactively plotting Kraken and SHOGUN-derived normalized microbial abundances in TCGA and the plasma microbiome data (raw counts or normalized data), as well as for interactively examining all model performances and ranked feature lists shown in this paper (approximately 600 models).

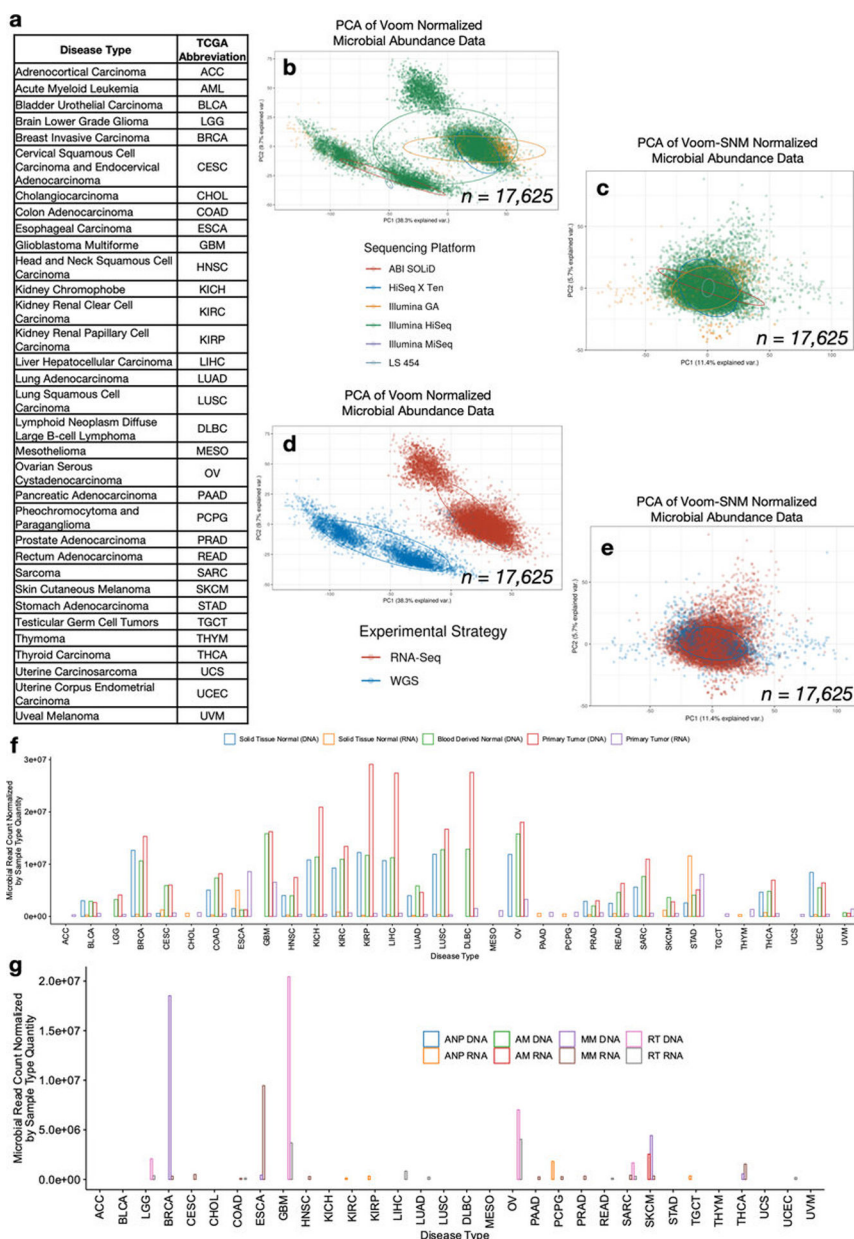
Data availability

Pre-processed cancer microbiome data generated and analyzed in this study (i.e. summarized read counts at the genus taxonomic level) as well as the metadata are available at ftp://ftp.microbio.me/pub/cancer_microbiome_analysis/. Raw outputs of Kraken- or SHOGUN-processed TCGA sequencing data comprise hundreds of terabytes of files and are not directly available unless otherwise coordinated with the corresponding author. However, all raw TCGA data and the bioinformatics pipeline necessary to generate such raw outputs from Kraken can be accessed through SevenBridge’s CGC. Each of the hundreds of ML models in this work generated a list of ranked features used to make predictions, and we provide the code to generate these lists below, in addition to showing them on our website. Raw data for the plasma validation study are available through the European Nucleotide Archive (accession IDs ERP119598: HIV-; ERP119596: PC; ERP119597: LC and SKCM); these data and the SHOGUN-processed data for the plasma validation study are available in Qiita (<https://qiita.ucsd.edu/>) under study IDs (12667: HIV-; 12691: PC; 12692: LC and SKCM).

Code availability

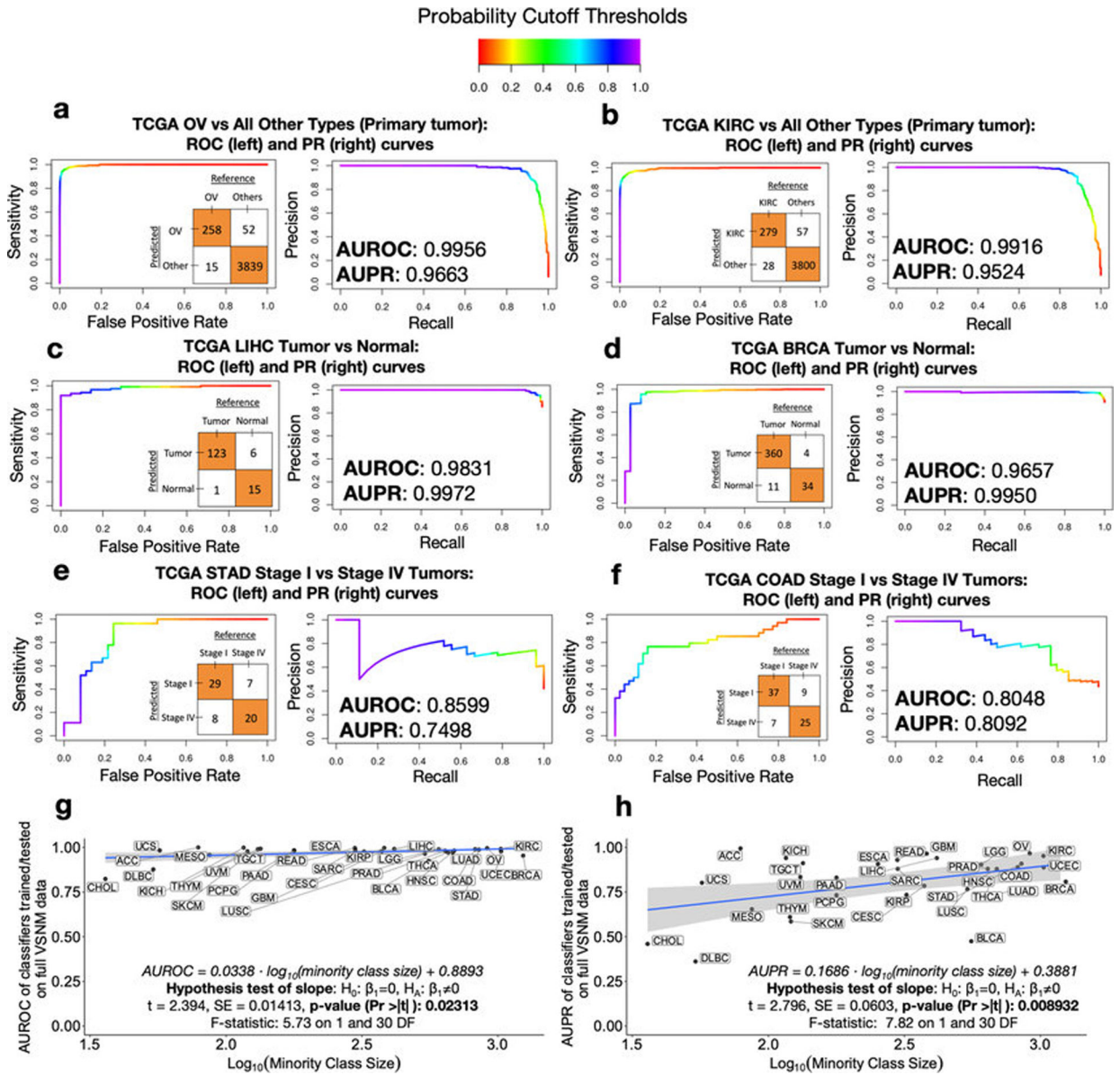
All programming scripts used to access, manage, and run data on the CGC as well as development of the supervised normalization, decontamination, ML pipelines, and so forth can be found at our GitHub repository link: <https://github.com/biocore/tcga>. These can be applied directly on the summarized, genus-level count data given above. Our CGC pipeline is also publicly shareable and available upon reasonable request to the corresponding author.

Extended Data



Extended Data Figure 1: Continued overview of the TCGA cancer microbiome.

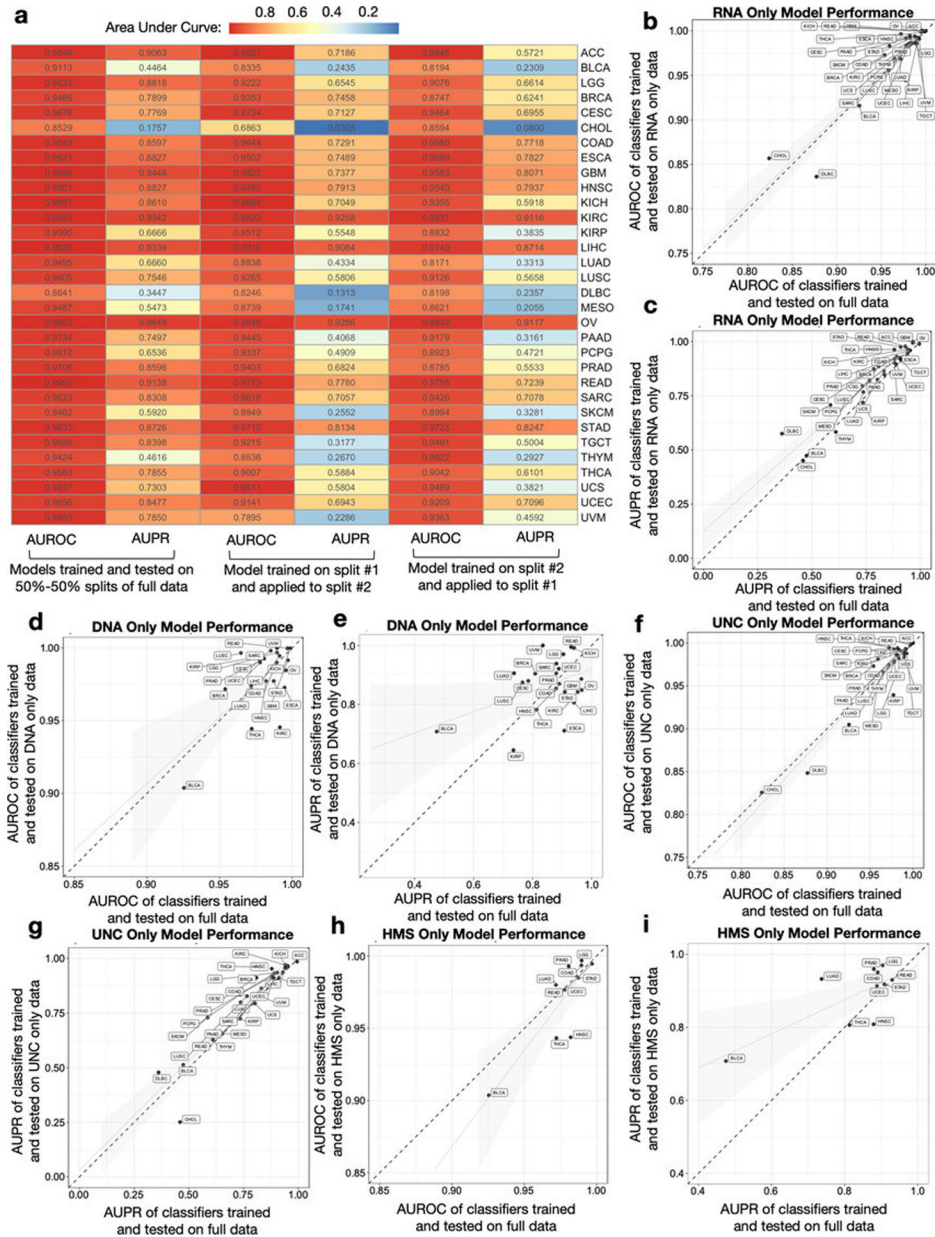
a, TCGA study abbreviations. **b**, Principal components analysis (PCA) of Voom normalized data, where colors represent sequencing platform of the sample and each dot denotes a cancer microbiome sample. **c**, PCA of the data following consecutive Voom-SNM supervised normalization, as labeled by sequencing platform. **d**, PCA of Voom normalized data, where colors represent experimental strategy of the sample and each dot denotes a cancer microbiome sample. **e**, PCA of the data following consecutive Voom-SNM supervised normalization, as labeled by experimental strategy. **f-g**, Microbial reads counts as normalized by the quantity of samples within a given sample type across all cancer types in TCGA after metadata quality control (Fig. 1b), including the three major sample types analyzed in the paper (**f**) and the remaining sample types (**g**). Note the following abbreviations: ANP = Additional - New Primary; AM = Additional Metastatic; MM = Metastatic; RT = Recurrent Tumor.



Extended Data Figure 2: Performance metrics details discriminating between and within TCGA cancer types using microbial abundances.

a-f, Expanded examples from the heatmaps in Figs. 1f-h. A color gradient shown at the top denotes the probability threshold at any point along the ROC and PR curves. An inset confusion matrix is shown using a 50% probability threshold cutoff, which can be used to calculate sensitivity, specificity, precision, recall, positive predictive value, negative predictive values, and so forth at the corresponding point on the ROC and PR curves. **g-h**, Linear regressions of model performance, specifically AUROC (**g**) and AUPR (**h**), for discriminating between cancer types in a one-cancer-type-versus-all-others manner, as a function of minority class size. Performances are shown for models using microbes detected in primary tumors, which had the greater number of samples (n=13,883) and cancer types (n=32) to compare. Since AUROC and AUPR have domains of [0,1] and the minority class size varied from 20 to 1238 samples, the latter is regressed on a log₁₀ scale. Inset hypothesis

tests and associated p-values are based on the null hypothesis of there being no relationship between the dependent and independent variables (two-sided test).



Extended Data Figure 3: Internal validation of ML model pipeline.

a, Two independent halves of TCGA raw microbial count data were normalized and used for model training to predict one-cancer-type-versus-all-others using tumor microbial DNA and RNA; each model was then applied on the other half’s normalized data. This heatmap compares the performances of these models as compared to training and testing on 50%–50% splits of the full dataset. **b-c**, Model performance comparison when subsetting the full Voom-SNM data by primary tumor RNA samples (n=11,741) across multiple sequencing centers to predict one-cancer-type-versus-all-others. **d-e**, Model performance comparison

Extended Data Figure 4: Orthogonal validation of Kraken-derived TCGA cancer microbiome profiles and their ML performances.

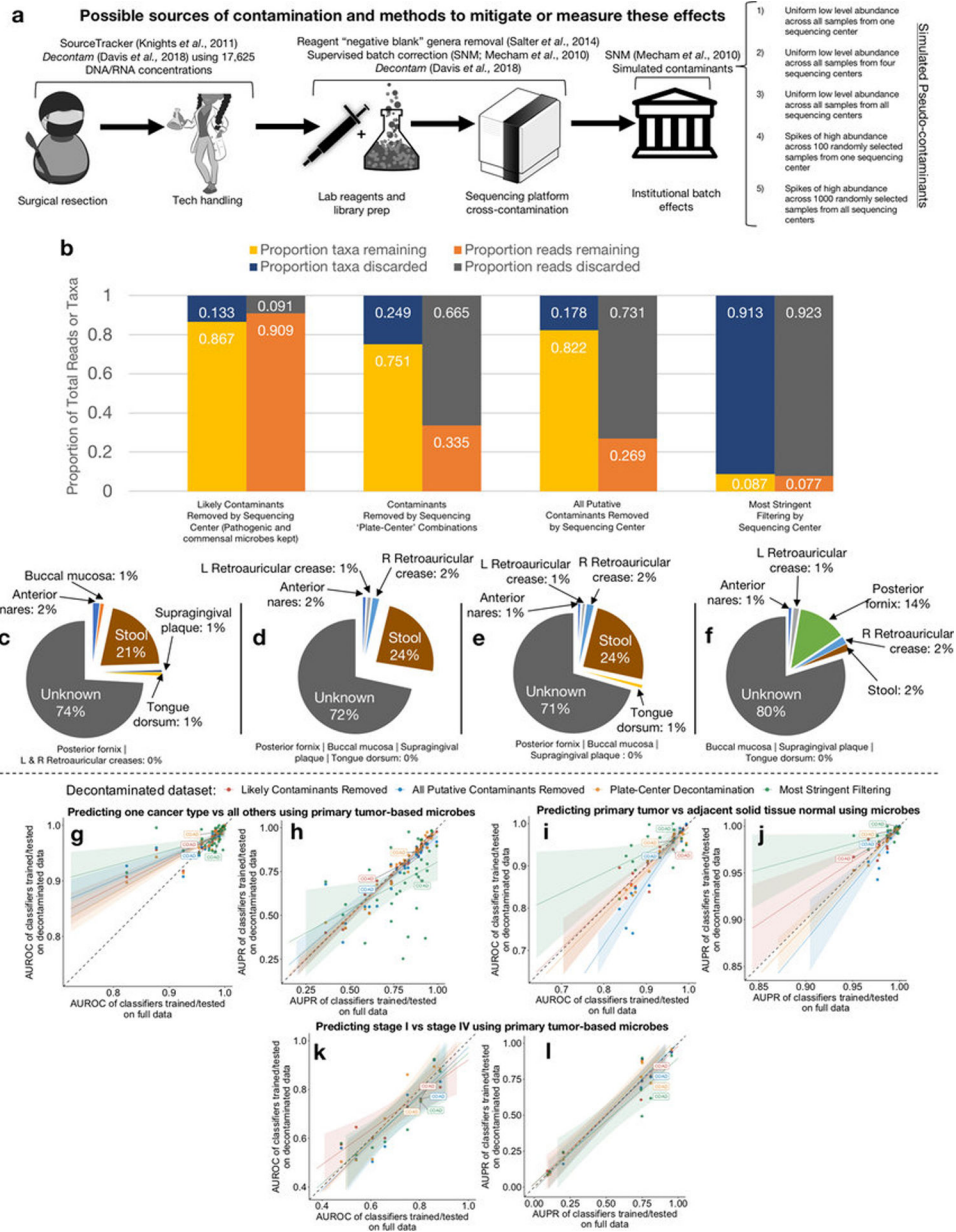
a-h, Four TCGA cancer types (CESC, STAD, LUAD, OV) underwent additional filtering after Kraken-based taxonomy assignments via direct genome alignments (Burrows-Wheeler Aligner⁵⁰, BWA). ML performances are compared between the normalized, BWA filtered data and matched, independently normalized Kraken data for one-cancer-type-versus-all-others using primary tumor microbes (**a-b**), tumor-versus-normal discriminations (**c-d**), stage I versus stage IV tumor discriminations using primary tumor microbes (**e-f**), and one-cancer-type-versus-all-others using blood-derived microbes (**g-h**) (Methods). **i**, Venn diagram of the taxa count between the BWA filtered data and the Kraken “full” data. **j-t**, An orthogonal microbial-detection pipeline called SHOGUN³¹ (“SHallow shOtGUN sequencing”) that uses direct genome alignments and a separate database (‘Web of Life’ [WoL]; n=10,575 microbial genomes [bacteria, archaea]; <https://biocore.github.io/wol/>)⁴⁸ was run on a subset of TCGA samples covering every analyzed cancer type (n=32), sample type (n=7), sequencing platform (n=6), and sequencing center (n=8) in the Kraken-based analysis (n=13,517 total samples). SHOGUN-derived microbial count data were normalized via Voom-SNM, analogous to its Kraken counterpart, and utilized for downstream ML analyses. **j**, Venn diagram of the SHOGUN-derived microbial taxa and the Kraken-derived microbial taxa. Note the use of separate databases and the fact that WoL does not include viruses while the Kraken database does. **k-l**, PCA of Voom (**k**) and Voom-SNM (**l**) normalized SHOGUN data, colored by sequencing center. **m-t**, ML performance comparisons between models trained and tested on SHOGUN data and matched Kraken data, using the same 70%/30% splits, for one-cancer-type-versus-all-others using primary tumor microbes (**m-n**), tumor-versus-normal discriminations (**o-p**), stage I versus stage IV tumor discriminations using primary tumor microbes (**q-r**), and one-cancer-type-versus-all-others using blood-derived microbes (**s-t**). For fair comparison, matched Kraken data were derived by removing all virus assignments in the raw Kraken count data and subsetting to the same 13,517 TCGA samples analyzed by SHOGUN; these matched Kraken data were then normalized independently via Voom-SNM the exact same way as the SHOGUN data (Methods) and fed into downstream ML pipelines. For all ML performances: A minimum minority class sample size of 20 was required to be eligible. For regression subfigures: The dotted diagonal line denotes perfect performance correspondence; generalized linear models with standard error ribbons are shown.



Extended Data Figure 5: Pan-cancer microbial abundances and an interactive website for TCGA cancer microbiome profiling and ML model inspection.

a, Pan-cancer *Fusobacterium* normalized abundances with a one-way ANOVA (Kruskal-Wallis) test for microbial abundances across cancer types for each sample type. Sample sizes are inset in blue, and TCGA study names are listed at the bottom. **b**, SourceTracker2 results for fecal contribution, as based on HMP2 data, for TCGA-COAD solid-tissue normal samples and TCGA-SKCM primary tumor samples. Only 1 solid tissue normal sample was available for TCGA-SKCM (Table S4), so primary tumors were used instead as the best proxy of expected skin flora. It is expected that colon samples should have higher fecal contribution than skin, so a one-sided Mann-Whitney test was employed. Since SourceTracker2 outputs mean fractional contributions of each source (i.e. HMP2) to each sink (i.e. COAD, SKCM samples), the center value of each bar plot is the mean of these values and the error bars denote the standard error. The sample sizes are inset below the bars in blue. **c**, Pan-cancer *Alphapapillomavirus* normalized abundances with a one-way ANOVA (Kruskal-Wallis) test for microbial abundances across cancer types for each sample type. Sample sizes are inset in blue, and TCGA study names are listed at the bottom. TCGA studies that clinically tested patients for HPV infection have “negative” or “positive” appended depending on the result of the test. **d**, Interactive website screenshot showing plotting of *Alphapapillomavirus* normalized microbial abundances using Kraken-derived data. Plotting using SHOGUN-derived normalized microbial abundances is available on another tab of the website (left-hand side). **e**, Interactive website screenshot of ML model inspection. Selecting the data type (e.g. all likely contaminants removed), cancer type (e.g. invasive breast carcinoma), and comparison of interest (e.g. tumor vs normal) will

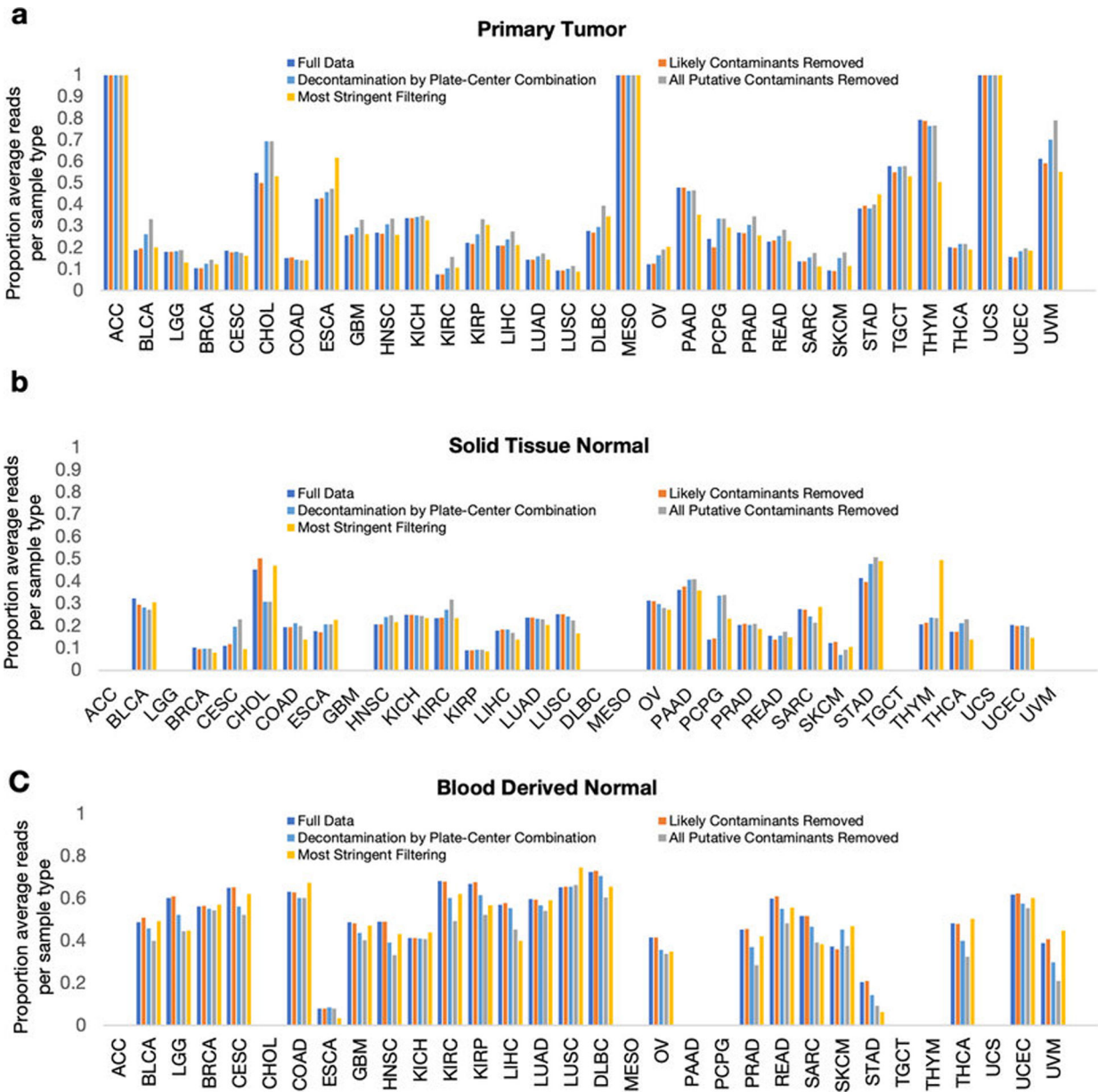
automatically update the ROC and PR curves, as well as the confusion matrix (using a probability cutoff threshold of 50%) and the ranked model feature list. Website is accessible at http://cancermicrobiome.ucsd.edu/CancerMicrobiome_DataBrowser/. All box plots show median, 25th and 75th percentiles, and whiskers that extend to 1.5× the interquartile range.



Extended Data Figure 6: The decontamination approach along with its results, benefits, and limitations on cancer microbiome data.

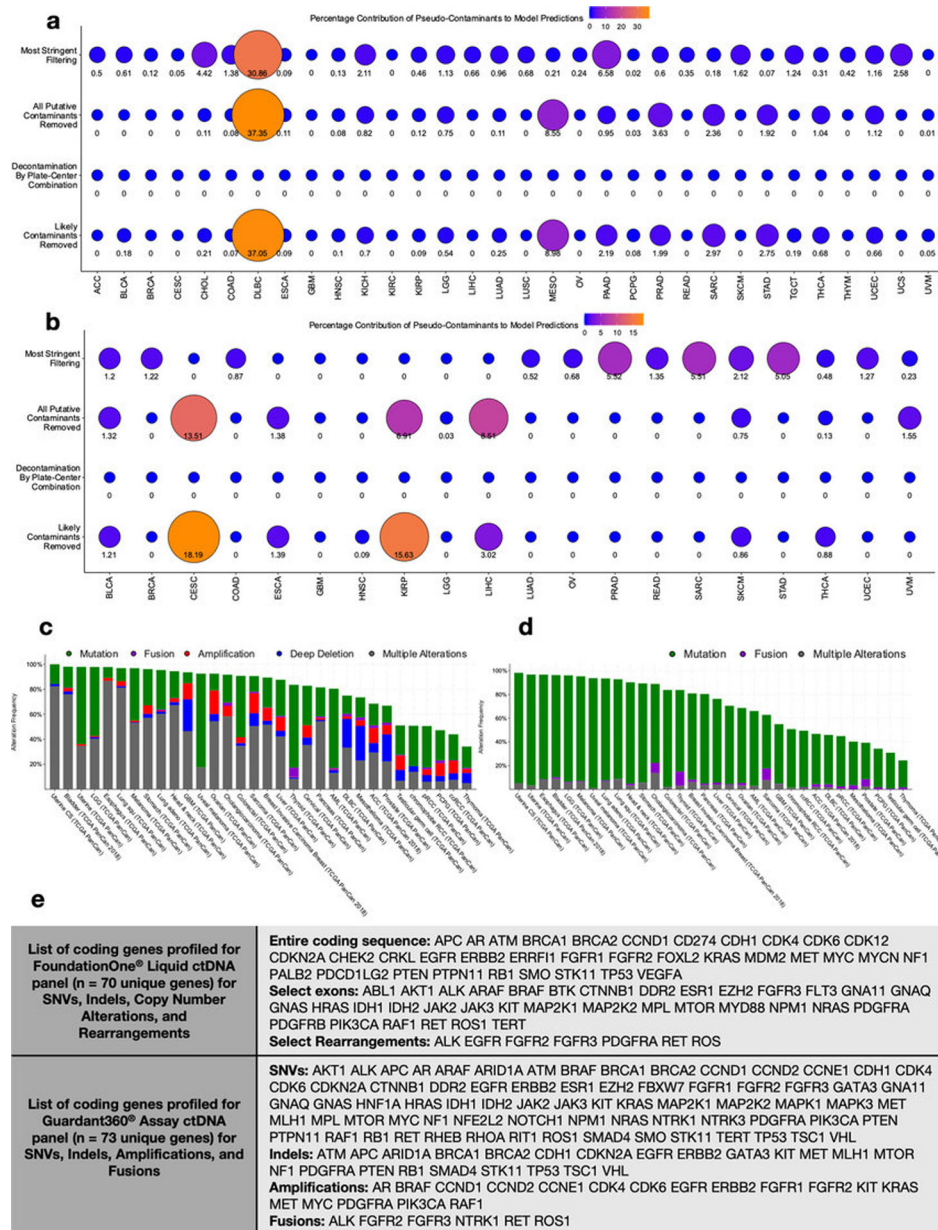
a, Various approaches used to either evaluate, mitigate, remove and/or simulate sources of contamination. **b**, The proportion of remaining taxa or microbial reads in TCGA after varying levels of decontamination. Decontamination by “sequencing center” removed all taxa identified as a contaminant at any one sequencing center (n=8 ‘batches’); decontamination by “plate-center” combinations removed all taxa identified as a

contaminant on any one single sequencing plate having more than 10 TCGA samples on it (n=351 ‘batches’). **c-f**, Body-site attribution prediction on the “likely contaminants removed” dataset (**c**), the “plate-center decontaminated” dataset (**d**), the “all putative contaminants removed” dataset (**e**), and the “most stringent filtering” dataset (**f**). **g-l**, All of the models and concomitant performance values (AUROC and AUPR) were re-generated using the four decontaminated datasets described above (each labeled with a different color; see legend located above plots). The AUROC and AUPR values obtained from models trained and tested on the decontaminated datasets are plotted against the AUROC or AUPR values from the “full” dataset (shown in Figs. 1f–h). The dashed diagonal line denotes a perfect linear relationship. Generalized linear models have been fitted to the corresponding datasets’ AUROC and AUPR values; standard errors of the linear fits are shown by the associated shaded regions. COAD model performances are identified throughout the figures.



Extended data figure 7: Decontamination effects on proportion of average reads per sample type. **a-c**, The total read count (i.e. DNA and RNA) of each major sample type (primary tumor [a], solid-tissue normal [b], blood-derived normal [c]) was summed and divided by the total number of samples within each sample type. This normalized read count (per sample type) was then divided by the summed normalized read count across all sample types for each cancer type, thereby providing an estimate of the proportion of average reads per sample type per cancer type. This was repeated for all five datasets, as shown by the legend, to assess if decontamination differentially impacts certain sample types and/or certain cancer types; relative stability in the percentages shown would suggest a lack of differential contamination. Minor sample types that were not further analyzed in this paper by decontamination or ML (e.g. additional metastatic lesions; n=4 sample types; Extended Data Fig. 1g) are not shown and comprised only 3.80% of total TCGA samples. Note, in the

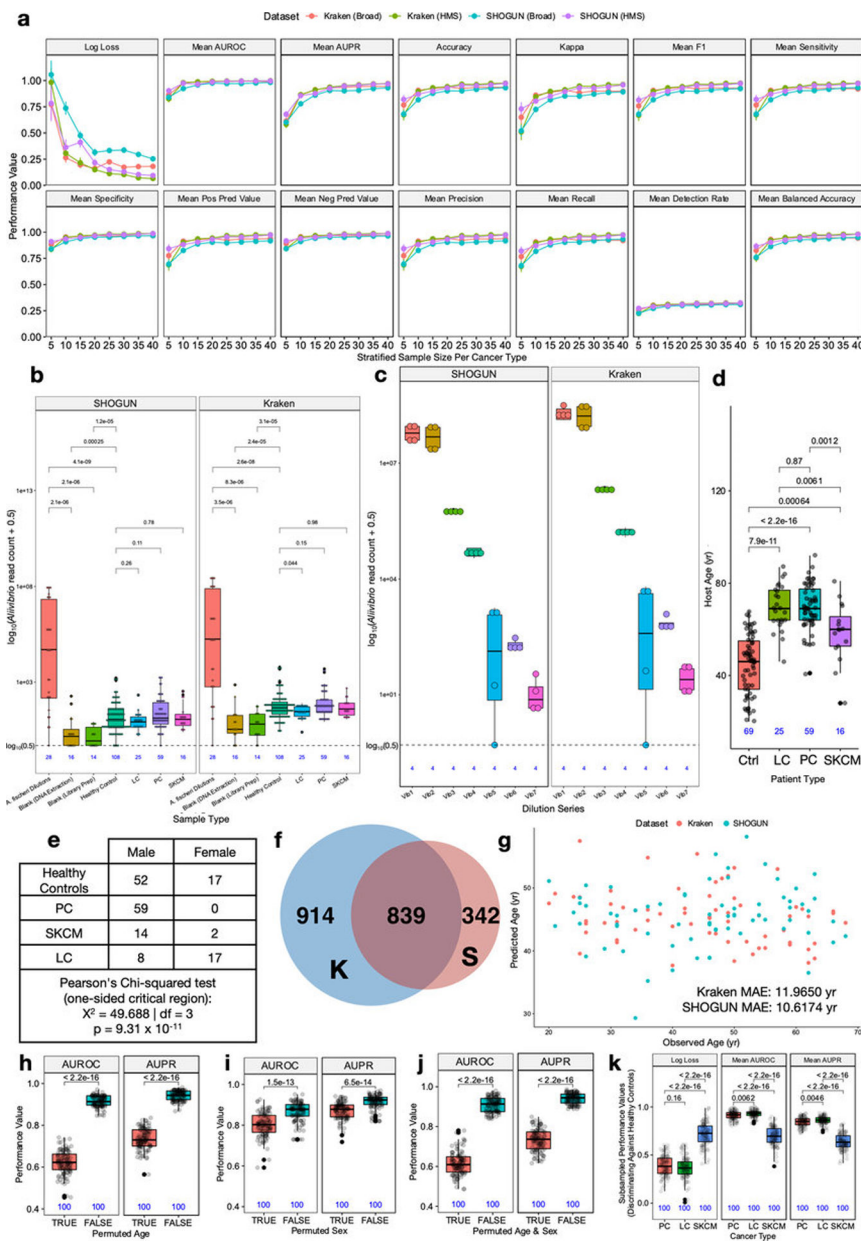
special case that only one sample type exists for a given cancer type (i.e. primary tumor in ACC, MESO, UCS), then all bars will show that 100% of the normalized reads came from that one sample type.



Extended data figure 8: Measuring spiked pseudo-contaminant contribution in downstream ML models and theoretical sensitivities of commercially available, host-based, cell-free DNA (ctDNA) assays in TCGA patients.

a-b, Feature importance scores were calculated for all taxa used in models trained to discriminate one-cancer-type-versus-all-others in all four decontaminated datasets (Extended Data Fig. 6b) using primary tumor microbial DNA or RNA (**a**), or using blood-derived mbdDNA (**b**). These decontaminated datasets were spiked with pseudo-contaminants prior to the decontamination and normalization pipelines to evaluate their performance (Methods),

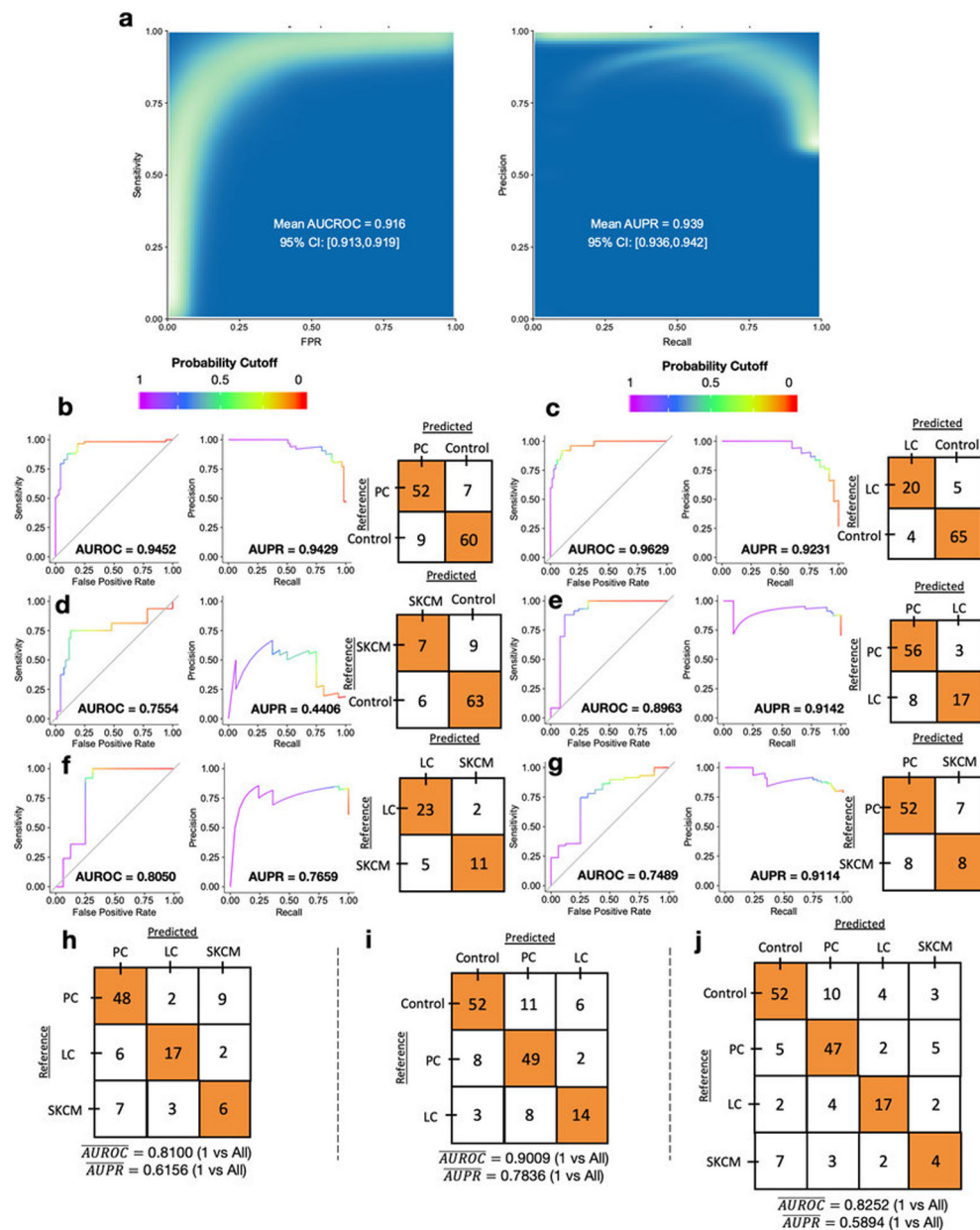
and the test set performances of the models shown are given in Extended Data Figs. 6g–h and Fig. 3a, respectively. Any spiked pseudo-contaminant(s) used by a model had their feature importance score(s) divided by the sum total of all feature importance scores in that model to estimate a percentage contribution of them towards making accurate predictions; the higher the score (out of 100), the less biologically reliable the model is. Note, “0” means that no spiked pseudo-contaminants were used for making predictions by the model; none of the models generated on the “plate-center decontaminated” data included spiked pseudo-contaminants as features. **c-d**, Percent distribution among TCGA studies with patients having one or more genomic alterations on FoundationOne® Liquid ctDNA coding genes (**c**) or on Guardant360® ctDNA coding genes (**d**). Data are downloaded from <https://www.cbioportal.org/>. **e**, The specific list of coding genes for the FoundationOne® and Guardant360® ctDNA assays and their examined alterations (source listed in Methods).



Extended Data Figure 9: Supporting analysis for real-world, plasma-derived, cell-free microbial DNA analysis between and among healthy individuals and multiple cancer types.

a. Discriminatory simulations in TCGA used to empirically power the real-world validation study (Fig. 4; Methods for details) and the theoretical performance metrics for each stratified sample size (per cancer type) using blood samples from the three cancer types of interest (prostate cancer (PC), lung cancer (LC), melanoma (SKCM)). Center values for each stratified sample size are the means of the performances and error bars denote the standard errors. Stratified sampling means that a sample size of five per cancer type would be a total of 15 blood samples under study for three-class discrimination (Methods). TCGA blood-derived normal samples were subsetted from The Broad Institute (Broad) and Harvard Medical School (HMS) such that they came from one sequencing center (i.e. Broad or HMS), one sequencing platform (Illumina HiSeq), and one experimental strategy (WGS).

The two kinds of LC in TCGA (LUAD, LUSC) were combined to reflect the samples available for the validation study. The resultant Broad and HMS datasets, as raw microbial counts, were then normalized separately via Voom-SNM, as would be in the validation study, and fed into a multi-class ($n=3$), leave-one-out (LOO) ML pipeline. Ten permuted iterations per stratified sample size per cancer type were used to estimate standard errors of theoretical performance estimates; for example, a stratified sample size of 40 would involve training and testing 1200 ML models ($= 40 \text{ samples} * 3 \text{ cancer types} * 10 \text{ iterations}$), from which 10 performance estimates would be made on 120 samples each to estimate standard errors (see Methods for details). All of this was repeated for SHOGUN-derived data as well. **b**, Evaluation of *Aliivibrio* genus abundance values (raw read counts) among positive control bacterial (*Aliivibrio*) monocultures, negative control blanks, and human sample types using both Kraken and SHOGUN-derived taxonomy assignments. Note the \log_{10} scale and 0.5 pseudo-count lower limit, shown with a dotted line. **c**, Evaluation of *Aliivibrio* genus abundance (raw read counts) across bacterial monoculture dilutions. Note the \log_{10} scale and 0.5 pseudo-count lower limit, shown with a dotted line. **d**, Distribution of ages among non-cancer healthy controls (“Ctrl”), grouped lung cancer (LC), prostate cancer (PC), and melanoma (SKCM) patients. **e**, Distribution of gender among non-cancer healthy controls, LC, PC, and SKCM patients with inset Pearson’s chi-squared testing (one-sided critical region). **f**, Venn diagram of taxa assignments between Kraken, which used the same database built for TCGA ($n=59,974$ microbial genomes [bacteria, archaea, viruses]), and SHOGUN, which used the ‘Web of Life’ database ($n=10,575$ microbial genomes [bacteria, archaea]; <https://biocore.github.io/wol/>)⁴⁸. **g**, Iterative leave-one-out (LOO) ML regression of host age using raw microbial count data from either Kraken (pink) or SHOGUN (aqua) derived assignments in healthy non-cancer patients. Mean absolute errors (MAE) evaluated across all samples are shown in the plot for Kraken and SHOGUN data. **h-j**, The effects of permuted age (**h**), sex (**i**), and age and sex (**j**) prior to Voom-SNM on ML performance to discriminate healthy versus grouped cancer patients using cell-free microbial DNA. One-hundred permutations were used for each comparison (Methods). **k**, Iterative subsampling of PC, LC, SKCM, and healthy control groups to match SKCM cohort size ($n=16$ samples), followed by LOO pairwise ML of each subsampled cancer type against subsampled healthy controls. One-hundred permuted iterations were used to estimate discriminatory performance distributions and standard errors (Methods). For subfigures **b**, **d** and **h-k**: Significance testing was performed using a two-sided Mann-Whitney test for all comparisons with multiple testing correction when testing >2 comparisons; all box plots show median, 25th and 75th percentiles, and whiskers that extend to $1.5\times$ the interquartile range. For all box plots and bar plots, sample sizes are inset in blue below them.



Extended Data Figure 10: SHOGUN-derived ML performances to discriminate between cancer types and healthy, non-cancer subjects using cell-free microbial DNA.

a, ‘Bootstrapped’ performance estimates for distinguishing grouped cancer (n=100) from non-cancer healthy controls (n=69). ROC and PR curve data from 500 iterations of with different training/testing splits (70%/30%) are shown on the rasterized density plot; mean values and 95% confidence interval estimates are inset on the plot. **b-g**, Leave-one-out (LOO) iterative ML performance between two classes: PC vs. controls (**b**), LC vs. controls (**c**), SKCM vs. controls (**d**), PC vs. LC patients (**e**), LC vs. SKCM patients (**f**), and PC vs. SKCM patients (**g**). **h-j**, Multi-class (n=3 or 4), LOO iterative ML performances to distinguish between cancer types, as well as between cancer patients and healthy non-cancer controls. Mean AUROC and AUPR, as calculated on one-versus-all-others AUROC and AUPR values, are shown on the bottom of the confusion matrices. **h**, LOO ML performance

between the three cancer types under study. **i**, LOO ML performance between the three sample types with 20 samples in the minority class (i.e. the cutoff used in the TCGA analysis, Figs. 1f–h). **j**, LOO ML performance between all four sample types under study. For all subfigures with confusion matrix plots: LOO ML was employed instead of single or ‘bootstrapped’ training/testing splits due to small sample sizes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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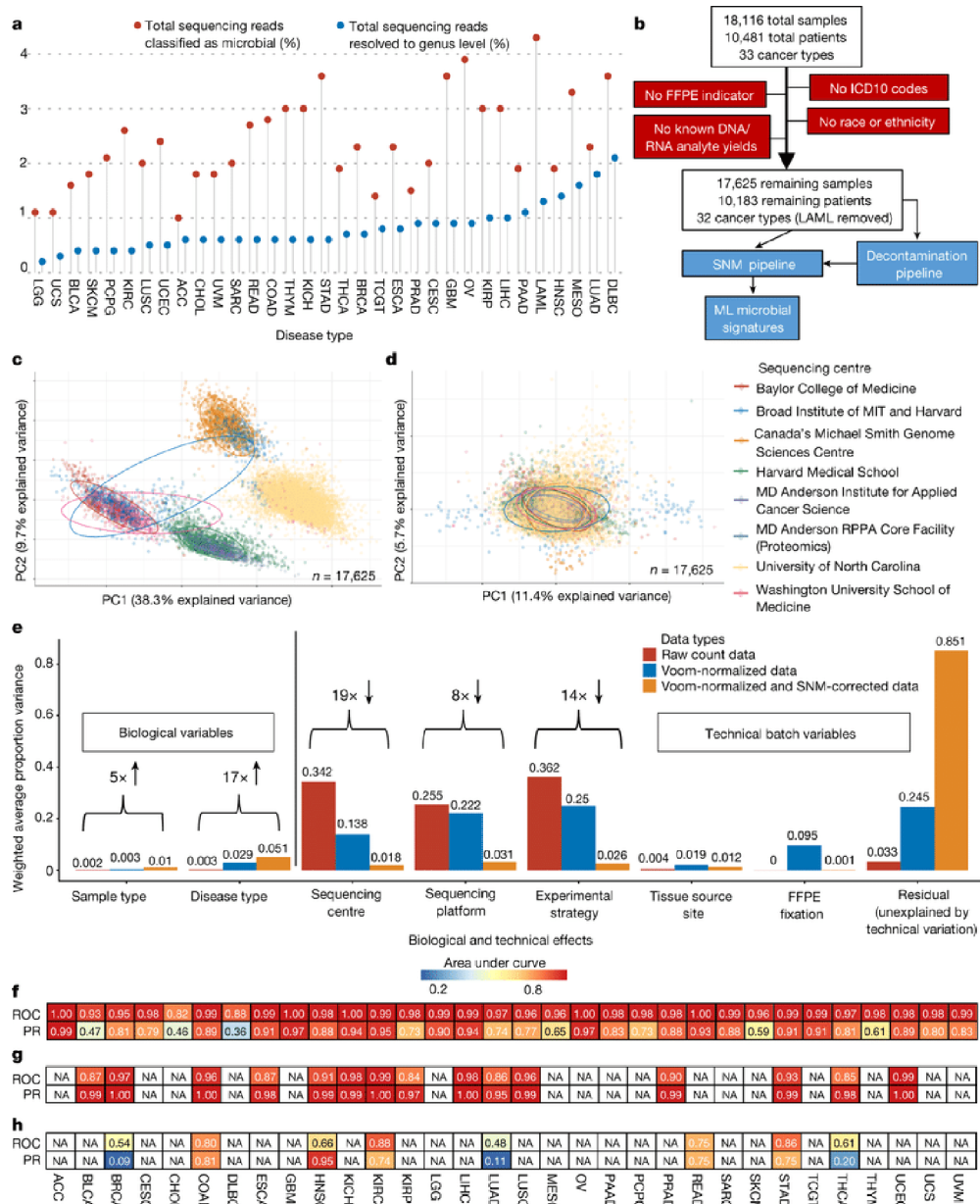


Figure 1: Approach and overall findings of the cancer microbiome analysis of The Cancer Genome Atlas (TCGA).

a, Lollipop plot showing the percentage of sequencing reads identified by the microbial-detection pipeline in TCGA dataset by Kraken, and the number of reads resolved at the genus level. **b**, CONSORT-style diagram showing quality control processing and the number of remaining samples. **c**, Principal components analysis (PCA) of Voom normalized data, with cancer microbiome samples colored by sequencing center. **d**, PCA of Voom-SNM data. **e**, PCA of Voom-SNM data. **f**, Principal variance components analysis of raw taxonomical count data, Voom normalized data, and Voom-SNM data. **f-h**, Heatmaps of classifier performance metrics (area under the ROC curve, “AUPRC”, or PR curve, “AUPR”) from red (high) to blue (low) for distinguishing between TCGA primary tumors (**f**), tumor-versus-normal (**g**), and stage I

versus IV cancers (**h**) with “NA” denoting <20 samples available in any ML class for model training. Column names are TCGA study IDs (Extended Data Fig. 1a).

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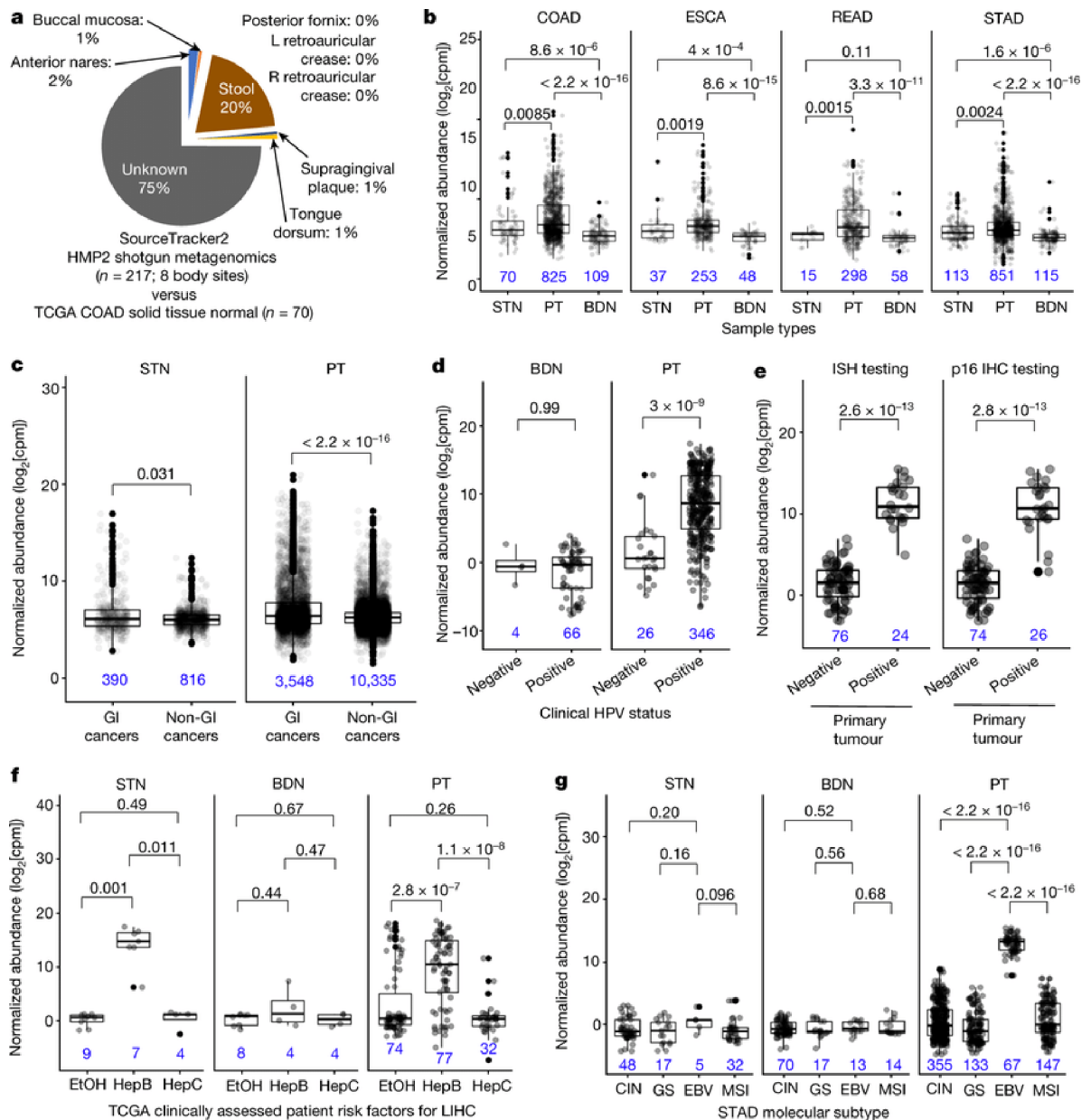


Figure 2: Ecological-validation of viral and bacterial reads within the TCGA cancer microbiome dataset.

a. Average body site attribution for solid-tissue normal samples from COAD ($n=70$) using SourceTracker2³² trained on the Human Microbiome Project 2 (“HMP2”) dataset. **b.** Differential abundances of the *Fusobacterium* genus for common gastrointestinal (GI) cancers associated with *Fusobacterium* spp.^{1,19,34,35}. **c.** Differential abundances of *Fusobacterium* among grouped GI cancers ($n=8$) and non-GI cancers ($n=24$) (Methods). **d-e.** Normalized HPV abundances for HPV-infected CESC patients (**d**) or HPV-infected HNSCC (**e**), as denoted in TCGA. **f.** Normalized *Orthohepadnavirus* abundance in LIHC patients with clinically adjudicated risk factors: prior hepatitis B infection (Hep B); heavy alcohol consumption (EtOH); or prior hepatitis C infection (Hep C). **g.** Normalized EBV abundance

in STAD integrative molecular subtypes: chromosomal instability (CIN), genome stable (GS), microsatellite unstable (MSI), or EBV-infected samples (EBV). All sub-figures: blood-derived normals and/or solid-tissue normals are shown as comparative negative controls; two-sided Mann-Whitney tests were used with multiple testing correction for >2 comparisons; all box plots show median, 25th and 75th percentiles, and whiskers extending to 1.5× the interquartile range with sample sizes inset in blue.

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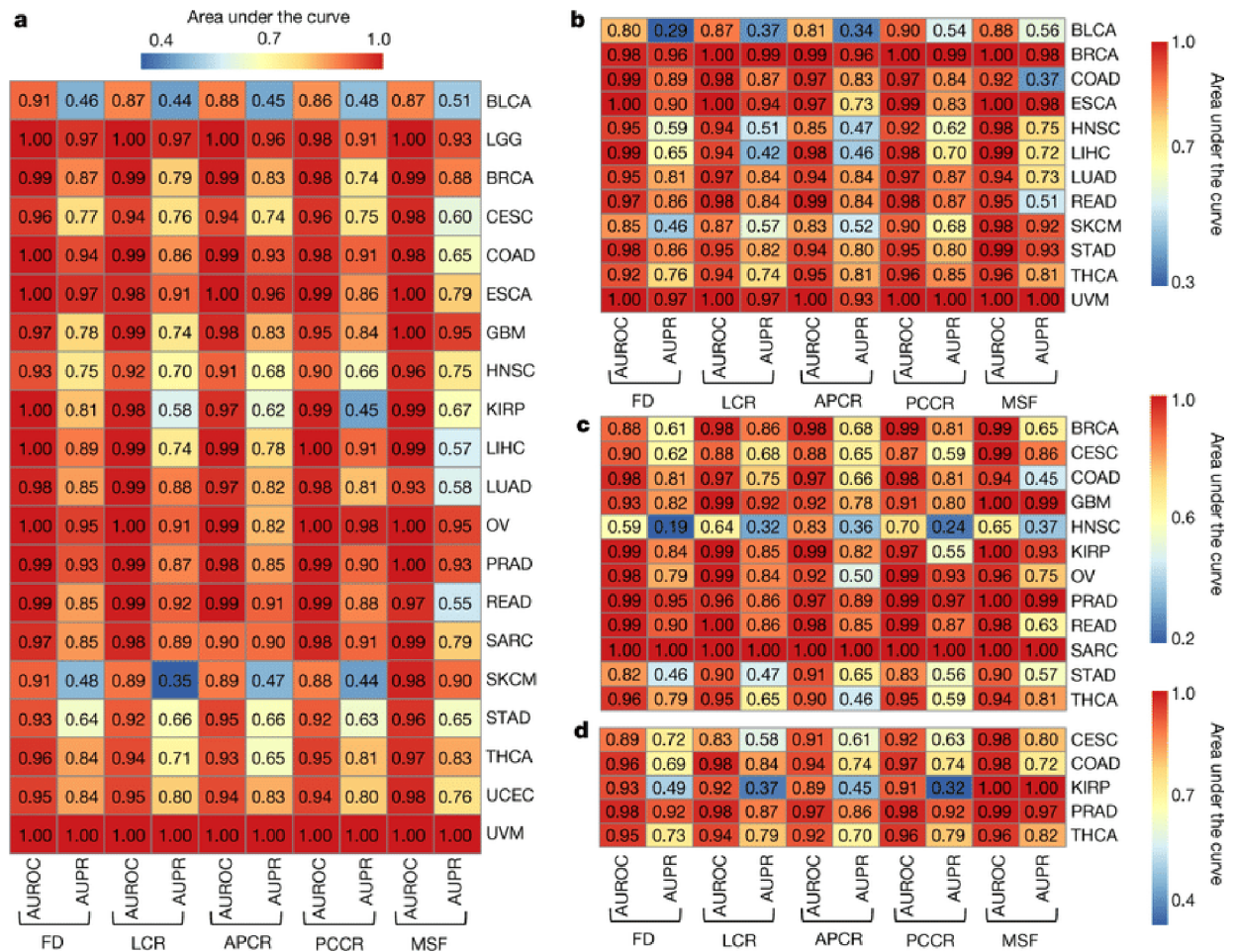


Figure 3: Classifier performance for cancer discrimination using microbial DNA (mbDNA) in blood and as a complementary diagnostic for cancer 'liquid' biopsies.

a, Model performance heatmap analogous to Figs. 1f–h to predict one-cancer-type-versus-all-others using blood mbDNA with TCGA study IDs on the right (Extended Data Fig. 1a);

20 samples were required in each ML minority class to be eligible. **b**, ML model performances predicting one-cancer-type-versus-all-others using blood mbDNA for stage Ia–IIc cancers. **c–d**, ML model performances using blood mbDNA from patients without detectable primary tumor genomic alterations, per Guardant360® (**c**) and FoundationOne® Liquid (**d**) ctDNA assays.

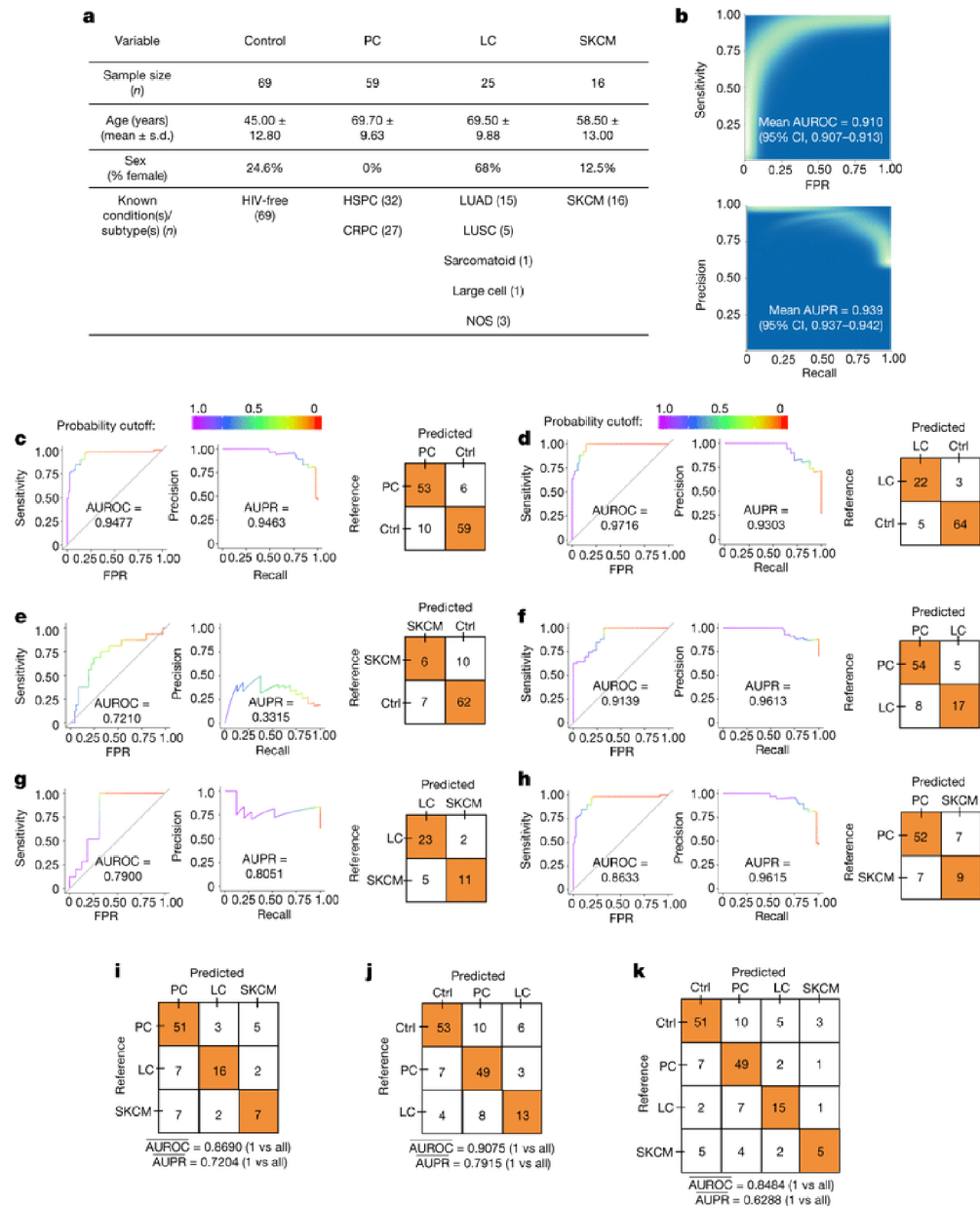


Figure 4: Performance of machine learning (ML) models to discriminate between cancer types and healthy non-cancer subjects using plasma-derived, cell-free mbdNA.

a. Demographics of samples analyzed in the validation study. All cancer patients had high-grade (stage III-IV) cancers of multiple subtypes and were aggregated into prostate cancer (PC), lung cancer (LC), and melanoma (SKCM) groups. **b.** ‘Bootstrapped’ performance estimates for distinguishing grouped cancer (n=100) from non-cancer healthy controls (n=69). Rasterized density plot of ROC and PR curve data from 500 iterations of with different training/testing splits (70%/30%); mean values and 95% confidence interval estimates inset. **c-h.** Leave-one-out (LOO) iterative ML performances between two classes: PC vs. controls (**c**), LC vs. controls (**d**), SKCM vs. controls (**e**), PC vs. LC patients (**f**), LC vs. SKCM patients (**g**), and PC vs. SKCM patients (**h**). **i-k.** Multi-class (n=3 or 4), LOO iterative ML performances to distinguish cancer types (**i**) and between mixed cancer patients

and healthy non-cancer controls (**j,k**). Overall LOO ML performance was calculated as the mean of one-versus-all-others comparisons' performances (area under the ROC curve "AUROC", or PR curve; "AUPR").

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