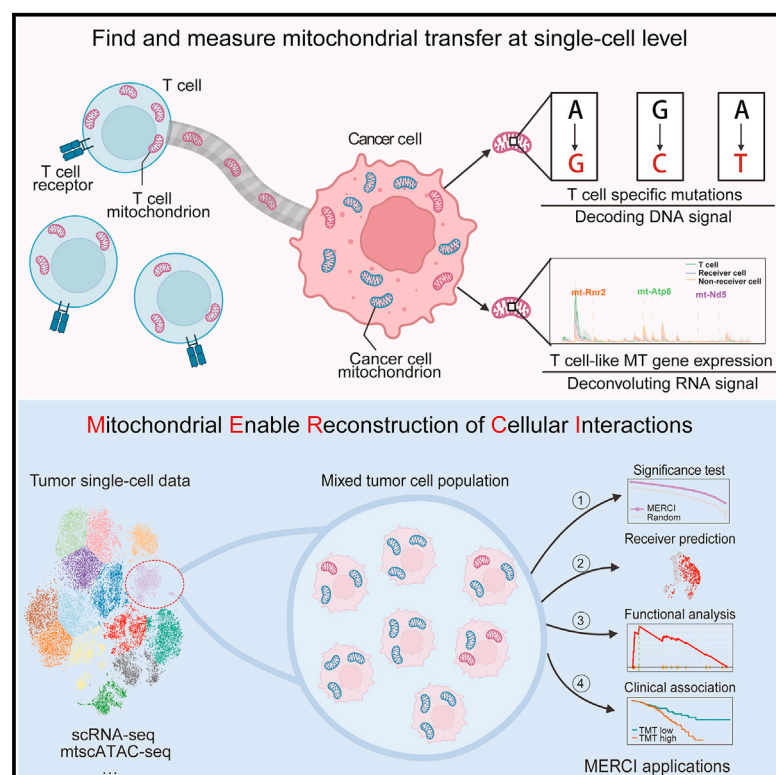


Systematic investigation of mitochondrial transfer between cancer cells and T cells at single-cell resolution

Graphical abstract



Authors

Hongyi Zhang, Xuexin Yu,
Jianfeng Ye, ..., John Minna,
Anli Zhang, Bo Li

Correspondence

anli.zhang@utsouthwestern.edu (A.Z.),
lib3@chop.edu (B.L.)

In brief

Recent work observes mitochondrial transfer from T cells to cancer cells. Zhang et al. systematically investigate this process using single-cell sequencing data from human cancers and identify a distinct phenotype related to mitochondrial transfer. Genes and pathways associated with this phenotype may serve as future therapeutic targets.

Highlights

- Cancer cells “hijack” mitochondria from nearby T cells
- Development of MERCI for tracing intercellular MT transfer
- Genomic inference of receiver cancer cells at single-cell resolution using MERCI
- Receiver cancer cells exhibit a distinct phenotype from the rest



Article

Systematic investigation of mitochondrial transfer between cancer cells and T cells at single-cell resolution

Hongyi Zhang,^{1,5,6,10} Xuexin Yu,^{1,5,6,10} Jianfeng Ye,^{1,10} Huiyu Li,² Jing Hu,^{1,5,6} Yuhao Tan,¹ Yan Fang,³ Esra Akbay,⁴ Fulong Yu,^{7,8} Chen Weng,^{7,8} Vijay G. Sankaran,^{7,8} Robert M. Bachoo,⁹ Elizabeth Maher,⁹ John Minna,² Anli Zhang,^{1,*} and Bo Li^{1,5,6,11,*}

¹Lyda Hill Department of Bioinformatics, UT Southwestern Medical Center, Dallas, TX 75390, USA

²Hamon Center for Therapeutic Oncology Research, UT Southwestern Medical Center, Dallas, TX 75390, USA

³Department of Molecular Biology, UT Southwestern Medical Center, Dallas, TX 75390, USA

⁴Department of Pathology, UT Southwestern Medical Center, Dallas, TX 75390, USA

⁵Center for Computational and Genomic Medicine, The Children's Hospital of Philadelphia, Philadelphia, PA 19104, USA

⁶Department of Pathology and Laboratory Medicine, University of Pennsylvania, Philadelphia, PA 19104, USA

⁷Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA

⁸Department of Pediatric Oncology, Dana-Farber Cancer Institute, Harvard Medical School, Boston, MA 02115, USA

⁹Department of Neurology and Neurotherapeutics, UT Southwestern Medical Center, Dallas, TX 75390, USA

¹⁰These authors contributed equally

¹¹Lead contact

*Correspondence: anli.zhang@utsouthwestern.edu (A.Z.), lib3@chop.edu (B.L.)

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SUMMARY

Mitochondria (MT) participate in most metabolic activities of mammalian cells. A near-unidirectional mitochondrial transfer from T cells to cancer cells was recently observed to “metabolically empower” cancer cells while “depleting immune cells,” providing new insights into tumor-T cell interaction and immune evasion. Here, we leverage single-cell RNA-seq technology and introduce MERCI, a statistical deconvolution method for tracing and quantifying mitochondrial trafficking between cancer and T cells. Through rigorous benchmarking and validation, MERCI accurately predicts the recipient cells and their relative mitochondrial compositions. Application of MERCI to human cancer samples identifies a reproducible MT transfer phenotype, with its signature genes involved in cytoskeleton remodeling, energy production, and TNF- α signaling pathways. Moreover, MT transfer is associated with increased cell cycle activity and poor clinical outcome across different cancer types. In summary, MERCI enables systematic investigation of an understudied aspect of tumor-T cell interactions that may lead to the development of therapeutic opportunities.

INTRODUCTION

The mitochondrion (MT) is a subcellular organelle that participates in critical physiological processes, such as energy production, calcium signaling, and programmed cell death.^{1–3} Mitochondria have their own 16.6Kb genome (16.3Kb for mouse) of circular DNA (mtDNA) with high mutation rate.^{4–6} Each somatic cell contains 10^2 – 10^4 copies of MT genomes,⁷ with a subset of mutations reaching high levels of heteroplasmy⁸ due to cytoplasmic inheritance or random drift.⁹ A few recent studies thus use the SNVs in the mtDNA as endogenous genetic barcodes to trace lineage relationships and clonal dynamics.^{10,11} For example, new methods have been developed to reconstruct shared lineages with cellular heteroplasmy of MT variants from single-cell RNA-sequencing (scRNA-seq) samples.^{12–14} These studies provide evolutionary insights into the fields of developmental biology,^{15,16} stem cell research,¹⁷ and oncology.¹⁸

In the meanwhile, a growing body of literature reports that mitochondria are not only inherited through lineage but also laterally shared between different mammalian cells^{19–21} primarily through tunneling nanotubes (TNTs).^{22–24} This phenomenon was first observed during coculture of mitochondria-deprived A549 lung cancer cells with mesenchymal stem cells, where the latter transfer mitochondria to the former to restore their aerobic respiratory functions.²³ Follow-up studies further demonstrate that this intercellular transfer occurs among diverse mammalian cell types *in vitro* and *in vivo*.^{22,24,25} A recent work examines the nanotube-based communication between cancer and immune cells and reports unidirectional mitochondrial trafficking from T cells to neighboring cancer cells.²⁶ As mitochondria provide both energy and the essential metabolites for T cell activation,^{27,28} this finding may point to a different mechanism of T cell dysfunction.

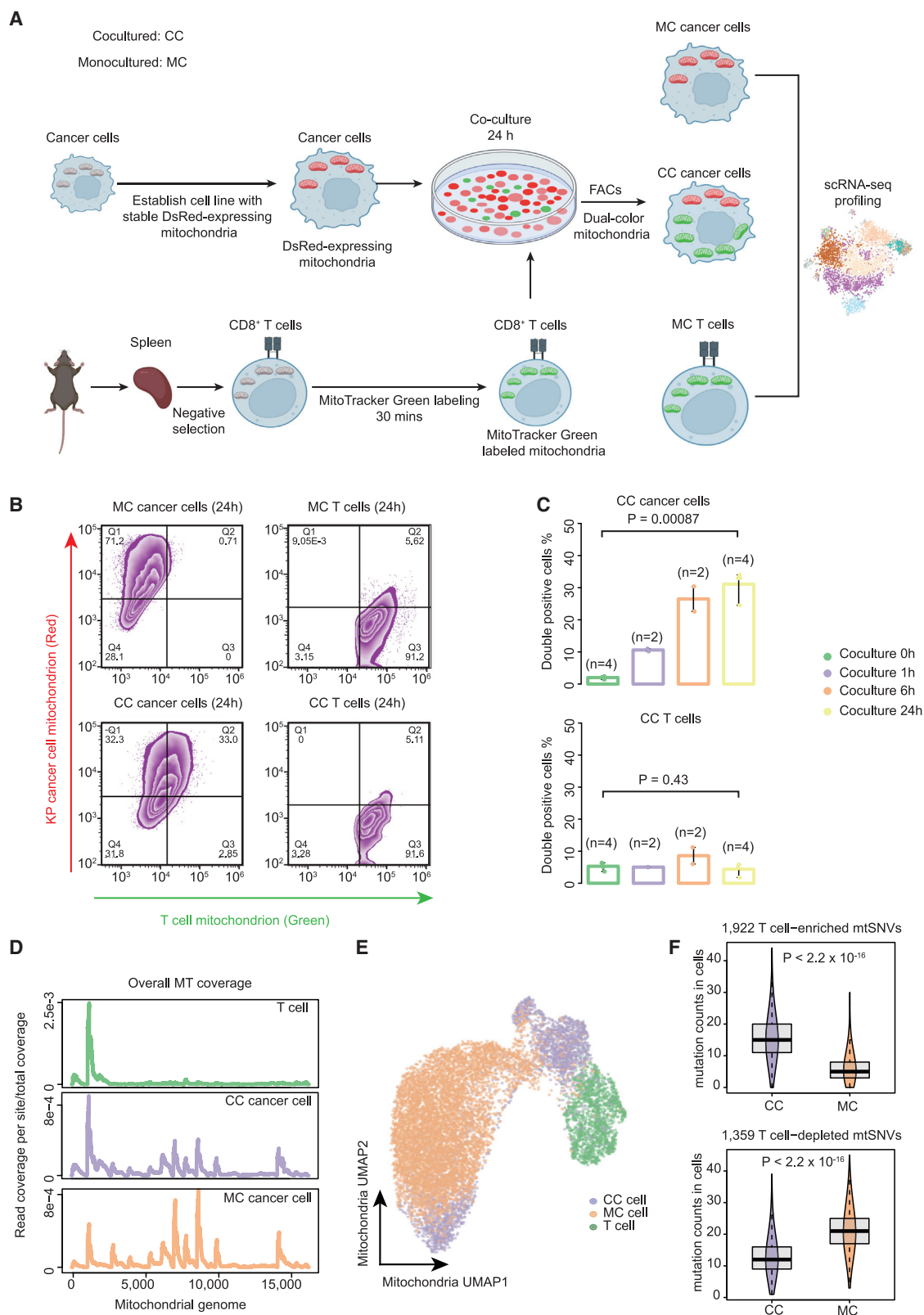


Figure 1. MT transfer signal captured by scRNA-seq data

(A) Cartoon illustrating the process of generating the ground-truth data by coculturing KP cancer cells and CD8⁺ T cells, followed by single-cell RNA-seq library construction and next-generation sequencing.

(legend continued on next page)

Many exciting questions follow this discovery, such as how nanotubes initiate in selected cancer cells, what genes or pathways cause the preferential MT transfer from T cells to cancer cells, and what the consequences of this unidirectional MT transfer are. Currently, most related studies have been conducted *in vitro*,^{20,29} which limited the generalizability of their findings. To better understand the mechanism of tumor mitochondrial interception and to identify precise, druggable targets, it is imperative to investigate this process under physiological conditions. However, *in vivo* tracking of MT transfer relies on genetic engineered mice with photoactivatable mitochondria,³⁰ which cannot be implemented in humans. Consequently, there is no practical approach to investigate this process in the human tumor microenvironment.

To bridge this gap, we developed a single-cell deconvolution method, MERCI (mitochondrial-enabled reconstruction of cellular interactions), to computationally infer the mitochondrial receiver cells in the patient tumor samples. MERCI takes the scRNA-seq data as input, identifies the lineage-specific MT mutations through a verified mutational calling pipeline, and predicts receiver cells based on mutation and mitochondrial gene expression profiles. Application of MERCI to human tumor samples uncovered a reproducible cancer cell phenotype of MT transfer, which is associated with multiple pathways of nanotube formation and energy metabolism. The signature was also predictive of worse clinical outcome for diverse cancer types.

RESULTS

Cancer mitochondrial hijack recapitulated via scRNA-seq data

To reproduce the observation of mitochondrial transfer, we first conducted the *in vitro* experiment described in the recent literature but using a different cancer cell line.^{20,26} Specifically, the murine Kras^{G12D}/p53^{ko} (KP) lung cancer cells³¹ were transfected with pDsRed2-Mito vector to label the mitochondria with red fluorescent signal. Primary autologous CD8⁺ T cells were isolated from mice spleen with mitochondria labeled with Mito Tracker green. Coculture experiments were conducted to visualize and quantify the transfer of mitochondria between cancer and T cells (Figure 1A). Fluorescence imaging revealed a dominant translocation of T cell mitochondria into the cytoplasm of KP cells (Figures 1B, S1A, and S1B). 24 h after coculture, over 30% KP cells carried high level of T cell mitochondria, while very few of T cells were weakly positive for the cancer signal (Figures 1B and S1B). We repeated experiments with different time spans and made similar observations (Figure 1C). We also tested a range of cancer to T cell ratios and observed high level

of mitochondrial transfer with more T cells in the coculture (Figures S1C). These results confirmed the near-unidirectional T cell to cancer mitochondrial flow reported by Saha et al.²⁶ Further, we repeated the coculture experiment using a different cell line, the MC38 colon cancer cells, and observed quantitatively similar MT transfer events (Figures S1D). Our observations suggested that mitochondrial hijacking is a common process initiated by diverse malignant cells.

We next performed scRNA-seq of three experimental groups: cocultured (CC) cancer cells and T cells as the treatment group and two controls including the monocultured (MC) cancer cells and primary CD8⁺ T cells (Figure 1A). Culture medium was diluted by a factor of 10⁵ prior to loading to 10x chips to mitigate the potential contamination from cell-free DNA/RNA during cell culture (Figure S1E). Half of the CD8⁺ T cells were reserved for scRNA-seq data preparation before coculture with cancer cells to ensure the same mitochondrial genetic background. All the sequenced CC cells were positive for the T cell mitochondrial signal. In total, we obtained 12,145 high-quality cells with over 1,000 reads mapped to mtDNA, including 2,904 CC cells, 7,572 MC cells, and 1,669 T cells (Table S1). Interestingly, the mtDNA read coverage profile of the CC cancer cells already exhibited an “intermediate” pattern between MC cancer and T cells on the overall (Figure 1D) and cellular levels (Figure S2A). For example, CC cells expressed higher protein-coding gene mt-Co3 than the T cells but lower than the MC cells (Figure S2A). Uniform manifold approximation and projection (UMAP)³² analysis using mitochondrial transcriptomic profile also confirmed that most CC cells displayed a mixed phenotype of the two monocultured cell populations (Figure 1E). These results indicated that mitochondria transfer can be manifested on the single-cell gene expression level.

We then investigated SNVs in the mtDNA, which have been used as genetic barcodes to track cell lineage.^{11,12} We customized a variant-calling pipeline (MERCI-mtSNP) for fast retrieving mitochondrial genotypes on cellular level from single-cell sequencing data (Figure S2B, STAR Methods). We tested MERCI-mtSNP by examining the downloaded benchmark scRNA-seq data,³³ which is a modified 10x scRNA-seq data with targeted enrichment to increase the coverage of MT genome. There are 17 *bona fide* mitochondrial variants in this dataset that differentiate two pre-defined cell lineages. MERCI-mtSNP successfully captured all 17 MT mutations in the correct cell populations (Figure S2C).

We then used MERCI-mtSNP to examine scRNA-seq samples derived from KP/T cells and identified 21,341 passed-filtered mitochondrial SNVs (mtSNVs) across all the cells. Next, we sought to trace mitochondrial transfer in the CC cancer cells

(B) The contour plots showing the percentage of double-positive cells (left: KP cancer cells, right: T cells) after 24 h of coculturing (lower) and monoculturing (upper). Double-positive cells are those carrying both endogenous and donor cell-derived mitochondria.

(C) Bar plots showing the percentage of double-positive cells for the KP cancer cells (upper) and the T cells (lower) at different coculture time points. Statistical significance was evaluated using two-sided Student's *t* test. Error bars indicate the range (min to max) of the data.

(D) The distribution of per site read coverage in mitochondrial genome for CC, MC, and T cells. Read depth is normalized by the total reads mapped to the mtDNA region.

(E) UMAP plot showing mitochondrial transcriptional profiles of cells. Cells are colored by different experimental groups.

(F) Boxplots showing the counts of T cell-enriched (upper) and T cell-depleted (lower) mtSNVs in CC and MC cancer cells with the MT read-depth range 1000–2000. Lower and upper box hinges represent 25th to 75th percentiles, central line the median, and the whisker extend to highest and smallest values no greater than 1.5× interquartile range; the violin component refers to the kernel probability density and encompasses all cells. Two-sided Wilcoxon rank-sum test was applied to calculate the *p* values. See also Figures S1 and S2.

using T cell-enriched mutations, which occurred at significantly higher frequencies in the primary T cells compared to the MC cancer cells ($FDR < 0.1$). To be precise, “frequency” refers to the percentage of cells carrying the mutation in a given population. In this analysis, we used half of the MC cells to define these mtSNVs, holding out the other half for testing to avoid data leakage. A total of 1,922 T cell-enriched variants were identified. As expected, the depth-normalized counts of T cell-enriched mtSNVs were significantly higher in the CC cells compared to the held-out MC cells (Figures 1F and S2D). This difference was reversed when using the T cell-depleted mtSNVs (Figures 1F and S2E). Here, the depth-normalized count refers to the number of mutations observed in a cell within a given MT read count range (Methods), which is applied to avoid the bias caused by read coverage difference (Figure S2F). Consistently, 9 out of the top 10 most enriched mtSNVs were significantly more frequent in the CC population (Figure S2G). We provided a table listing all the T cell-enriched variants used in this work (Table S2), which had been selected in the regions with $\geq 5X$ coverage in both cell types by MERCI-mtSNP (Figure S2H). Together, our data indicated that the genetic variants of the transferred mitochondrial genome can be captured by the scRNA-seq data as well.

Tracing MT receivers through single-cell deconvolution

Following the previously described results, we developed MERCI to identify the MT receiver cells through quantifying the donor-derived mtSNVs and estimating the relative abundance of exogenous mitochondria (Figure 2A, STAR Methods). For a given cell, MERCI combines the MT gene expression and mtSNV profiles to predict if it is a receiver and infer the relative fraction of donor-derived mitochondria. Specifically, MERCI first defines T cell-enriched mtSNVs based on the reference (cancer and T cell) populations and calculates an “effective count statistic” (N_{eff}) for each cancer cell to assign each cell with a DNA rank score (Figure S3A). It then uses the averaged MT gene expression profiles of cancer and T cell populations as reference and applies support vector regression (SVR)^{34,35} to estimate the relative abundance of transferred MT in the target cancer cells (Figure S3B). Each cell of interest is assigned with an RNA rank score based on the ordering of the SVR coefficient: the relative proportion of T cell-derived MT. Rank transformation is necessary to make the scores robust to outliers and less sensitive to sequencing depth or cancer type. Cells with scores passing a predefined cutoff for DNA or RNA rank are considered as candidate MT receivers.

We evaluated the performance of MERCI with the ground-truth scRNA-seq data. T cells and half of the MC populations were used to generate the reference MT gene expression and T cell-enriched mtSNVs. CC and the held-out MC cells were mixed to produce an *in silico* sample to test MERCI. After deconvolution, we observed significantly higher proportion of T cell-derived mitochondria in the cocultured cancer cells (Figure 2B), consistent with our previous observation (Figure 1B). We next used the receiver operating characteristic (ROC) and precision-recall curves (PRC) to measure the prediction accuracy of MERCI. Both DNA and RNA rank scores reached a moderate level of area under the curve (AUC) ranging between 0.7 and 0.85 (Figures 2C and 2D), yet neither is accurate enough as a single predictor. As DNA and RNA ranks were esti-

mated from independent sources, it is intuitive to combine them to achieve better performance. One simple approach is to take the averaged rank score as a predictor, which indeed reached an improved AUC of 0.88 for the ROC curve, and 0.85 for the PRC (Figures S3C and S3D). However, this strategy can become problematic when applied to real-world data, because a rank predictor will always make positive calls under a given cutoff even without any true positives in the data. Therefore, in the next section, we assessed a different approach to: 1) test if a sample contains any true MT receivers and 2) isolate the receiver cells based on MERCI rank scores.

Benchmark MERCI rank scores for real-world application

We implemented a simple strategy to call a “positive” only if both ranks are higher than the given cutoff, thus putting an “AND” gate after the DNA and RNA scores. Assume the cutoff is top rank 10%: in a null sample with no true positives, DNA and RNA ranks will be independent, following the same uniform distribution. AND gate selection is expected to make 1% of positive calls. Alternatively, if the sample contains true positives, the two ranks will be correlated by MT receivers, and AND gate will predict higher percentage of positive calls. This rationale can be used to avoid making calls from disqualified samples. We first tested this strategy using the *in silico* mixture sample with $\sim 40\%$ true positive cocultured cancer cells and confirmed that DNA and RNA ranks are indeed correlated, leading to higher-than-expected positive calls across a range of cutoffs (Figure 3A). In contrast, when using the null sample (only monocultured cells) as the input, the reported positives were within the range established by 10,000 randomly permuted ranks (Figure 3B). These results confirmed that the existence of MT receivers in the sample generates statistical dependence between the two MERCI ranks. To quantify this signal, we defined the “ratio of cell number” (R_{cm}) as MERCI reported number of positive calls over the maximum number from 10,000 random permutations. In MERCI, we used $R_{cm} > 1$ (empirical p value < 0.0001) as the criterion to determine if the input sample contained any MT receivers. This criterion performed well as we applied it to the mixture and null samples: at all non-trivial cutoffs, R_{cm} was greater than 1 with $FDR < 0.05$ for the mixture sample, yet insignificant and smaller than 1 for the null sample (Figure 3C).

The power to detect true positives by MERCI is intuitively affected by the fraction of MT receivers in the sample. To assess the limit of MERCI when this fraction is small, we performed an *in silico* titration analysis by gradually lowering the percentage of CC cells from 20% to 0.5% in the mixture sample. We found that with over 7% CC cells, R_{cm} values remained robustly above 1 at different rank cutoffs (10%–50%) (Figures 3D–3F, S3E, and S3F). Further, positive calls made by MERCI were significantly enriched for the true CC cells even with lower fraction (5%–15%) of positive cells (Figure S3G). As the fraction of CC cells dropped below 5%, MERCI did not always detect the true signal, yielding smaller R_{cm} and insignificant p values. We concluded that MERCI is sensitive to determine the existence of MT receivers for samples with over 7% of true positive cells. Depending on the cancer type, the fraction of receivers in the tumor could range between 13% and 45%,²⁶ which is sufficient for MERCI implementation.

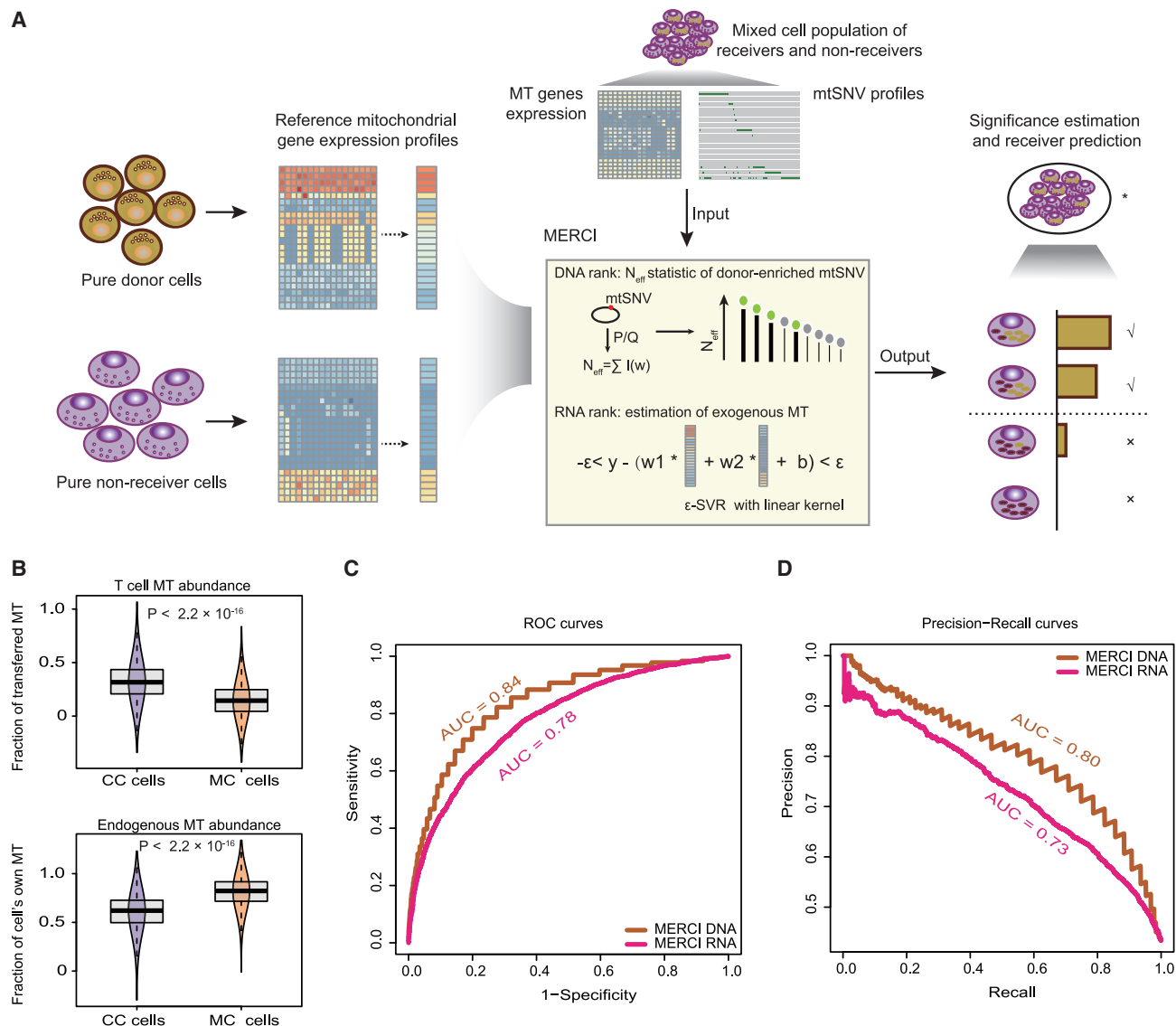


Figure 2. Overview of MERCI and application to the ground-truth data

(A) Schematic illustration of MERCI. Single-cell RNA-seq data from reference donor and non-receiver cells were used to deconvolute MT fractions in the cancer cell population. By combining ranks from DNA and RNA information, MERCI estimates the statistical significance of the existence of true MT receivers, and evaluates each candidate cancer cell as a receiver or not.

(B) Boxplots showing the estimated abundance (SVR coefficients) of T cell transferred (upper) and endogenous mitochondria (lower) in CC and MC cancer cells. Lower and upper box limits represent 25th to 75th percentiles, central line the median, and the whiskers extend to highest and lowest values no greater than $1.5 \times$ interquartile range; the violin component refers to the kernel probability density and encompasses all cells. Statistical significance was evaluated using two-sided Wilcoxon rank-sum test.

(C and D) Receiver operating characteristic (ROC) (C) and precision-recall (PR) (D) curves using MERCI-derived rank scores as predictors of mitochondrial receivers, i.e., CC cells. Area under the ROC and PR curves (AUC) was labeled. See also [Figure S3](#).

Next, we assessed how rank cutoffs influence the prediction of MERCI using the criteria of sensitivity, specificity, and precision. The benchmark *in silico* sample contained all the CC cells ($n = 2,904$) as true positives and held-out MC cells ($n = 3,786$) as true negatives, which was used to evaluate the performance of MERCI under different cutoffs ([Figures 3G–3I](#)). MERCI achieved high specificity and precision with more rigorous cutoffs, at the cost of a reduced sensitivity. When using a less strict cutoff, for example, at the threshold of top rank 50%, MERCI reached

approximately 90% specificity and 80% precision, with a moderate sensitivity of 56.3%. Therefore, in our downstream analysis, we consistently used a “lenient” cutoff up to 50% given $R_{\text{cm}} > 1$ to balance the sensitivity and precision.

Finally, in a real-world scenario, cancer samples are comprised of both “receiver” and “non-receiver” cells without known class labels. Thus, it is challenging to find the “pure” non-receiver cancer cells to infer the reference MT gene expression and the T cell-enriched mtSNVs. To solve this problem, we

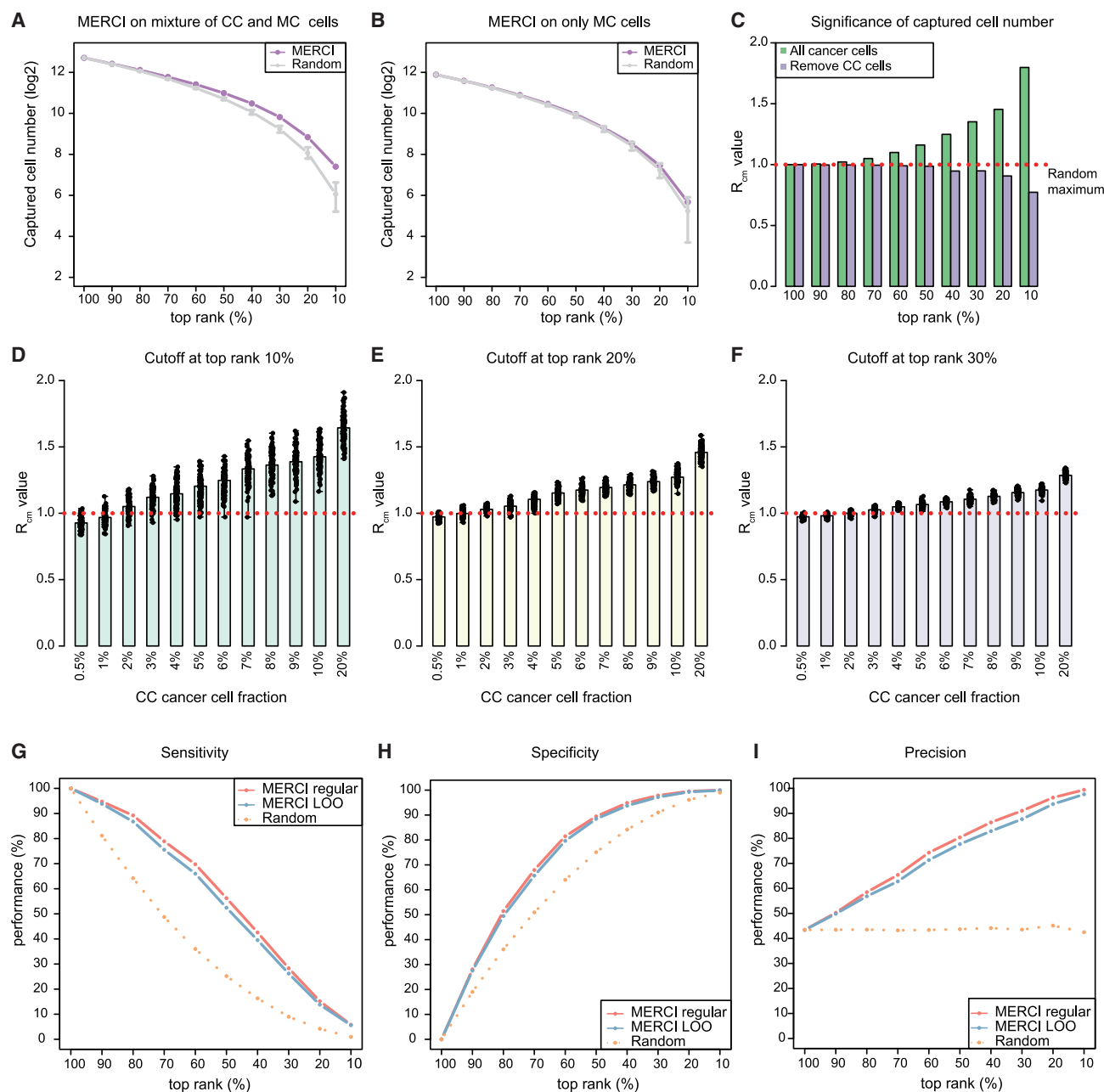


Figure 3. Benchmarking of MERCI for real-world application

(A and B) Dot-line plots showing the number of positive calls captured by using AND gate to MERCI DNA and RNA rank scores across a range of cutoffs. Purple dots represent the results of *in silico* mixture sample and gray intervals indicate the ranges established by 10,000 randomly permuted ranks. Error bars indicate the range (min to max) of the data.

(C) Bar plots showing the R_{cm} values at different rank cutoffs. Red dotted line indicates $R_{cm} = 1$.

(D–F) Significant estimation of positive calls when different fractions of true receivers are included. Bar plots showing the averaged R_{cm} values reported by MERCI for downsampled datasets at rank cutoffs at top 10% (D), 20% (E), and 30% (F) respectively. Black dots indicate the R_{cm} values of downsampled datasets. Error bars indicate the range of (min to max) of the data.

(G–I) The sensitivity, specificity, and precision of MERCI when using different rank cutoffs to predict the MT receivers. See also Figure S3.

implemented a “Leave One Out” (LOO) strategy: for a given cancer cell of interest, we considered all the cancer cells except the given one as the non-receiver reference to estimate the rank scores. This approach is reasonable because majority of tumor

cells in the cancer population do not receive mitochondria from the T cells.²⁶ We assessed the performance of MERCI LOO using the same *in silico* sample described previously (Figures 3G–3I). The slightly lower sensitivity (Figure 3G) is

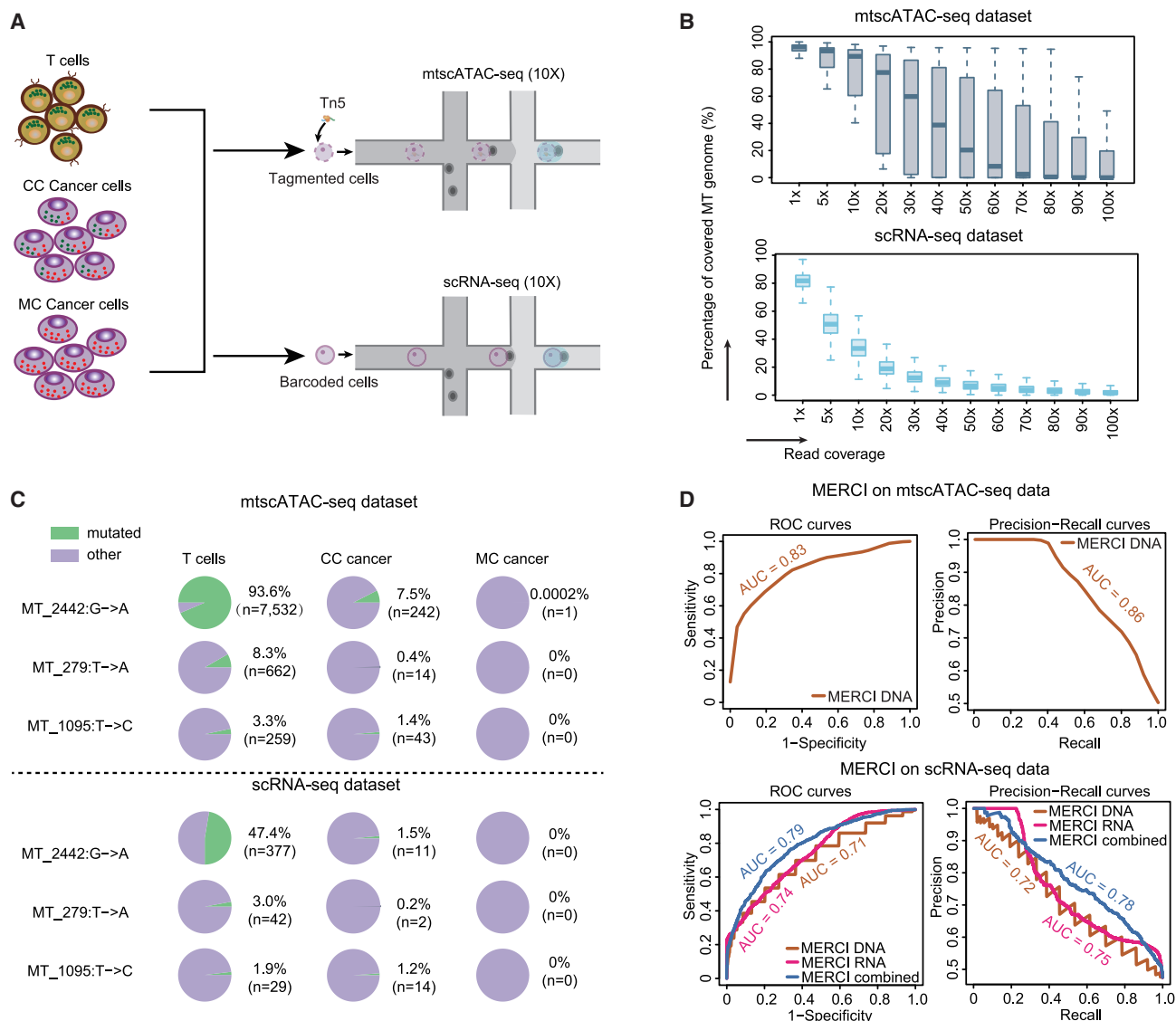


Figure 4. Independent validation using mtscATAC-seq technique

(A) Diagram showing the process of generating mtscATAC-seq and matched scRNA-seq datasets.

(B) Boxplots showing the distribution of read coverage per cell in the mtscATAC-seq dataset (upper) and the matched scRNA-seq dataset (lower). Box center line: median; box limits: upper and lower quartiles; box whiskers: 1.5× interquartile range (IQR).

(C) The cell frequencies of three T cell-specific variants in T cells, CC, and MC cancer cell populations. Label of the variant marks the 0-index coordinate, followed by the nucleotide change.

(D) Receiver operating characteristic (ROC) and precision-recall (PR) curves using MERCI-derived rank scores as predictors of mitochondrial receivers. As it is not feasible to estimate the RNA rank scores from mtscATAC-seq dataset, the upper panel only shows the performance DNA rank scores. See also Figure S3, Tables S1, and S2.

expected as the presence of the true receivers in the reference may diminish the cancer-T cell difference, which does not influence the specificity (Figure 3H). Together, our analysis provided a practical way to apply MERCI to real-world samples.

Evaluation of MT transfer and MERCI performance with mtscATAC-seq

Detection of mitochondrial transfer in our analysis so far is primarily based on 10x Genomics scRNA-seq platform, which, although heavily used in recent clinical studies, is not optimized

for mtDNA profiling due to its limited and uneven coverage in this region.^{11,36} Therefore, we repeated the coculture experiments (Figure 1A) and implemented a recently developed, cutting-edge technology for mtDNA single-cell profiling (mtscATAC-seq) (Table S1, Figure 4A)¹⁰ to profile the MT regions for CC, MC, and primary T cells. In brief, mtscATAC-seq repurposed scATAC-seq to deeply sequence mtDNA (Figure S3H), offering significantly improved coverage (per-site, per-cell average 48X) within the mitochondrial region in our analysis (Figure 4B). In the same batch, we also generated matched

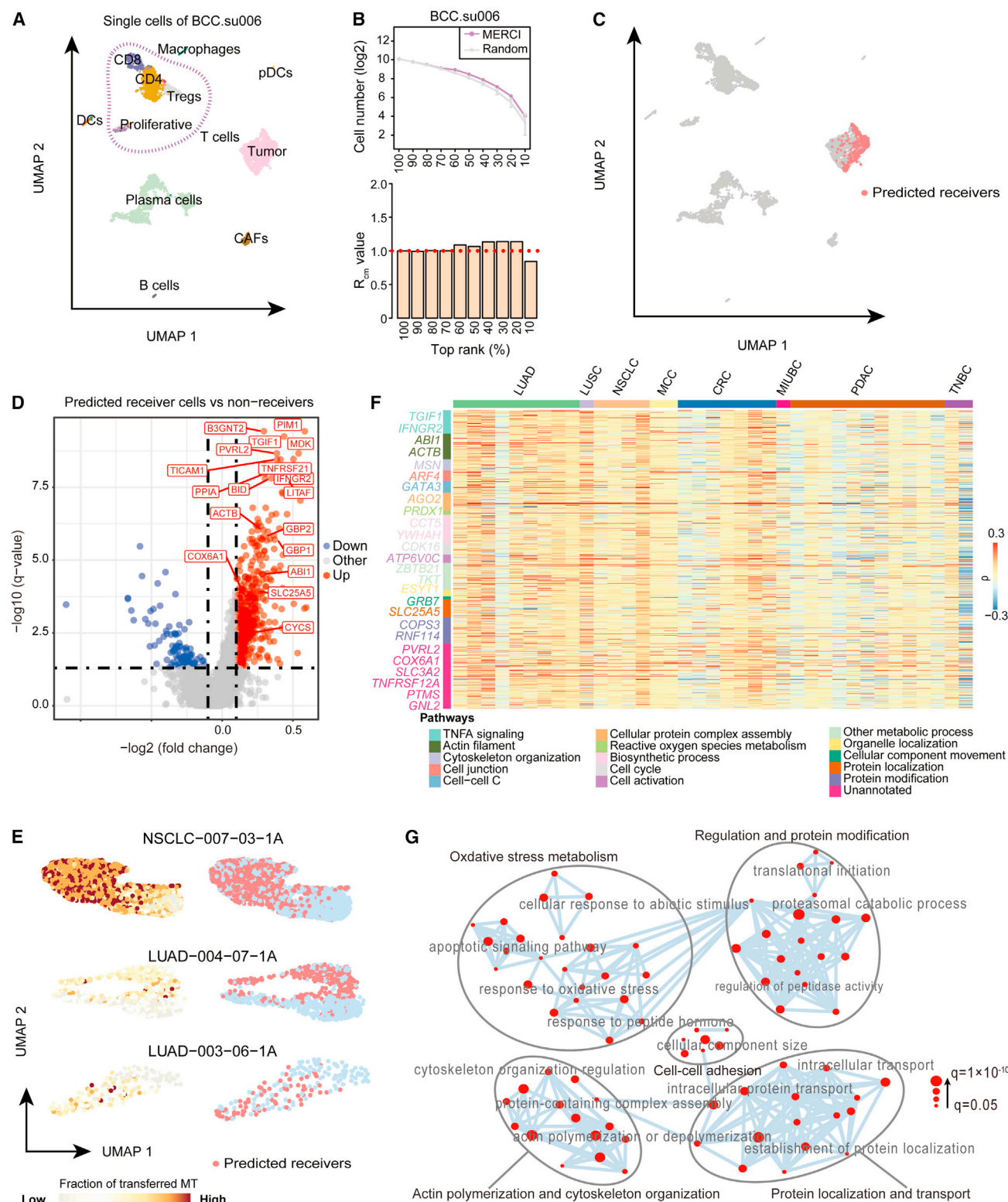


Figure 5. Distinct MT receiver phenotype predicted by MERCI in human tumor samples

(A) UMAP plot showing the cell clusters and distribution of single cells from BCC patient “su006.”
(B) Significance estimation of the number of positive calls reported by MERCI for scRNA-seq data of BCC patient “su006.”
(C) UMAP plots showing the projection of MERCI predicted receiver cells.

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scRNA-seq data (15X coverage) for each sample to evaluate MERCI.

We next called mtSNVs for each sample through the MERCI-mtSNP pipeline and identified 1,989 T cell-enriched variants from mtscATAC-seq data (Table S2). As expected, CC cancer cells, the *bona fide* receivers, harbored significantly more T cell-enriched mtSNVs compared to the negative control (MC cancer cells), a finding also confirmed using matched scRNA-seq data (Figures S3I and S3J). A scrutiny over the variants called from mtscATAC-seq revealed several interesting mtSNVs. For example, MT_2442: G→A, among others, occurred in 94% of the T cells, while almost completely absent in the MC population. We observed that a sizable fraction (7.5%) of cancer cells cocultured with T cells picked up this mutation (Figure 4C). These variants provided compelling genetic evidence of mitochondrial transfer from T cell to the cancer cell population. In contrast, cocultured T cells (Figure S3K) did not show higher amount of cancer-enriched mtSNVs (Figures S3L and S3M), supporting the nearly unidirectional mitochondrial transfer from T cells to cancer cells.

To further evaluate MERCI's performance in identifying receiver cells, we applied MERCI to the mtscATAC-seq data from the mixed population of CC and MC cancer cells. The outcomes revealed that MERCI is capable of differentiating CC cancer cells from a blend of cancer cells with both ROC and PRC AUC values exceeding 0.8 (Figure 4D). We repeated this analysis using the scRNA-seq data matched to the mtscATAC-seq samples and confirmed that MERCI reached similar prediction accuracy as in the discovery cohort. These confirmed the ability of MERCI to uncover true MT receivers from scRNA-seq samples.

Distinct MT receiver phenotype in human solid tumor uncovered by MERCI

We next implemented MERCI to scRNA-seq samples in the public domain, which were derived from patients with basal cell carcinoma (BCC)³⁷ or esophageal squamous cell carcinoma (ESCC).³⁸ Both cohorts provided sufficient cancer and T cells that were ideal for discovery of gene markers related to MT transfer. We first analyzed the samples from three BCC patients and tested the existence of receivers in each sample using the R_{cm} criterion. These samples were selected as they contained sufficient numbers ($n > 500$) of both cancer and T cells. While two samples showed no evidence of MT receivers ($R_{cm} < 1$, Figures S4A and S4B), positive signals were detected in one sample, "su006" (Figures 5A and 5B). Specifically, its R_{cm} values were significantly greater than 1 at the rank cutoffs of 20%–60% ($FDR < 0.05$, Figure 5B), thus qualified the next step implementation of MERCI. We then used the LOO approach to predict MT receivers using the recommended parameters (rank cutoff =

50%), and called a total of 309 tumor cells (~30% of all tumor cells). Interestingly, the predicted receiver cells formed a tight subcluster within the original cancer cell cluster, which was not observed for the predicted non-receivers (Figures 5C and S4C), indicating a converged cancer cell phenotype associated with MT transfer.

We proceeded to investigate this phenotype through differentially expressed genes (DEGs) of the MT receivers against the other cancer cells. The genes associated with ATP synthesis and oxidative phosphorylation, such as *BID*, *COX6A1*, *CYCS*, and *SLC25A5*, were significantly upregulated in the predicted receivers (Table S3, Figure 5D), supporting that these cells carried more mitochondrial-mediated energy production. As expected, genes involved in cytoskeleton regulation and actin polymerization, such as *PIM1*, *ACTB*, *PFN1*, *PVRL2*, and *GBP1/2* etc., were also expressed at higher levels, consistent with the mechanism of nanotube formation. Interestingly, essential components of the tumor necrosis factor alpha (TNF- α) pathway, including *TGIF1*, *IFNGR2*, and *LITAF*, were among the top upregulated genes. Functional enrichment analysis of the 608 DEGs also revealed that "TNF- α signaling via NF- κ B" was the most enriched pathway (Table S4, Figure S4D). Previous studies have reported that TNF- α pathway is closely involved in nanotube initiation and intercellular nanotube connectivity,^{39,40} which is in line with our observations. This phenotype was further reproduced when we analyzed the scRNA-seq data of another three patients with ESCC, where MERCI-predicted receivers also showed distinct gene expression pattern from non-receivers (Figure S4E). The signature genes of the MT receivers from the ESCC patients significantly overlapped with those from the BCC sample (Figure S4F). Specifically, genes involved in cytoskeleton remodeling, cell junction and TNF- α pathways, such as *ACTB*, *PFN1*, *PVRL2*, and *TGIF1*, were also upregulated (Figures S4G–S4J).

To verify the previously described findings, we extended MERCI analysis to incorporate more cancer samples from the public domain by releasing the requirements for sample inclusion: ≥ 100 counts for both T cells and cancer cells. We assessed 156 samples with both cancer and T cells from recent clinical studies, as cataloged in CancerSCEM database,⁴¹ and subsequently filtered in 61 samples from lung, gastric, skin, pancreatic, bladder, colorectal, and breast cancers (Table S1). We applied MERCI on each sample and found that the fraction of T cell-derived mitochondria in cancer cells was significantly associated the gene expression variations on UMAP plot in 61% (37/61) of these samples ($|p| > 0.2$, $p < 0.001$, Table S5, Figures 5E and S5A). Further, we observed that the 608 DEGs identified in BCC were also positively correlated with the estimated fraction of T cell-derived mitochondria in most of these

(D) Volcano plot showing DEGs of the predicted receiver cells versus non-receivers, with fold change calculated using the mean values of the two groups. Statistical significance was evaluated using two-sided Wilcoxon rank-sum test, with FDR corrected using Benjamini-Hochberg procedure.

(E) UMAP plots illustrating the estimated fraction of transferred MT in cancer cells (left) and the distribution of predicted receivers (right) for three selected cancer patients. These patients were chosen based on the highest Pearson correlations between the fraction of transferred mitochondria (Fr.T-Mito) and the gene expression phenotype measured with UMAP (either umap1 or umap2).

(F) Heatmap showing the Spearman correlation coefficients between the expression of 608 DEGs identified in BCC dataset and the Fr.T-Mito in cancer cells across 37 samples of different cancer types. LUAD: lung adenocarcinoma, LUSC: lung squamous cell carcinoma, NSCLC: non-small-cell lung cancer, MCC: merkel cell carcinoma, CRC: colorectal cancer, MIUBC: muscle-invasive urothelial bladder cancer, PDAC: pancreatic ductal adenocarcinoma, TNBC: triple-negative breast cancer. The selected representative genes of different pathways were marked with different colors.

(G) Gene ontology network based on the commonly enriched GO terms of the 95 MT transfer-related genes. Each node represents a gene ontology. Node size corresponds to gene ratio of each GO term vs. total analyzed genes in human BCC sample. See also Figures S4 and S5, and Tables S3, S4, and S5.

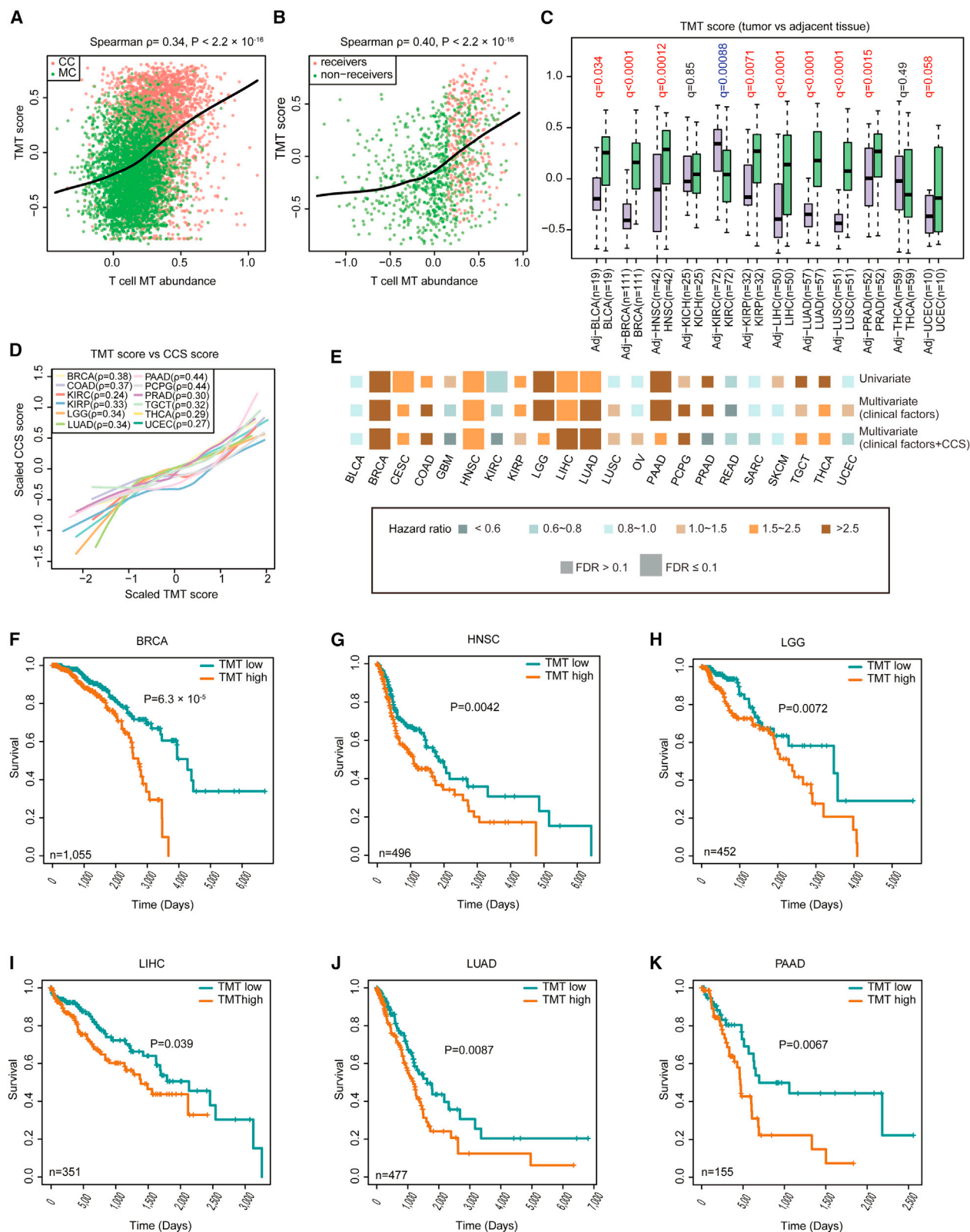


Figure 6. Functional and clinical impact of TMT score across different cancer types

(A and B) TMT scores of cancer cells showing high correlation with the estimated foreign mitochondrial abundance for both murine training (A) and human BCC (B) data. The Spearman correlation test was used to calculate the p values.

(legend continued on next page)

37 samples. Specifically, genes in pathways related to biological processes such as biosynthesis, cell activation, TNF- α signaling, and cell cytoskeleton remodeling were also significantly correlated with this fraction (Figure 5F). These findings suggested that the gene markers identified by MERCI from human samples are reproducible across multiple cancer types.

To further investigate cross-species conservation of MT transfer, ^{13,26} we searched for all the genes positively correlated with the fraction of T cell-transferred mitochondria in cancer cells (Spearman $\rho > 0$ and FDR < 0.05) in the human BCC sample and the murine benchmark sample. We observed 95 overlapped genes as MT transfer related. The number of overlapped genes between human and mouse data is significantly higher than expected ($p = 1.28 \times 10^{-8}$, Figure S5B), suggesting that these overlapped genes might be evolutionary conserved. Gene set enrichment analysis confirmed that MT transfer-related genes are mainly enriched in the pathways related to nanotube formation, such as cell-cell adhesion, actin polymerization, and cytoskeleton organization (Figures 5G, S5C, and S5D). These results indicated that nanotube-related genes could potentially serve as a marker of MT transfer.

Pan-cancer analysis of MT transfer in large human patient cohorts

To quantify the MT transfer activity in cancer cells or bulk tumors, we generated a 17-gene signature from the 95 evolutionary conserved MT transfer-related genes (Table S6). Selection of the 17 signature genes was guided by prior knowledge of gene function, as these genes are directly involved in the pathways associated with nanotube formation or mitochondrial activity (Figure S6A). We excluded genes involved in various metabolic pathways, such as “oxidative phosphorylation” and “regulation of ATP synthesis”, as they are more likely the consequences, not the cause of mitochondrial transfer.⁴² As our goal is to find markers related to the initiation of this process, these genes were not suitable to serve as signatures. We defined the gene set enrichment score⁴³ based on the mRNA expression of these 17 genes as tumor MT transfer (TMT) score, which served as a measure of MT transfer in tumor. To access its performance, we first calculated TMT score for each cancer cell in the murine benchmark and human BCC sample, and confirmed that the MT receivers showed significantly higher TMT scores than the non-receivers (Figures 6A and 6B). Next, the TMT score was highly correlated with the exogenous MT abundance in the cancer cells

with a Spearman correlation coefficient higher than any of the 17 genes alone ($\rho = 0.34$ for benchmark data and $\rho = 0.40$ for BCC data, Figures S6B and S6C).

To further investigate the functional relevance of MT transfer in cancer, we estimated the TMT score of 10,628 human cancer or adjacent normal samples in The Cancer Genome Atlas (TCGA) database. To ensure statistical power, we analyzed the 22 major cancer types with over 100 patients. In 9 out of the 12 cancers with at least 10 adjacent normal samples, we observed significantly higher TMT scores in the tumors than in the matched normal tissues (Figure 6C), consistent with the enhanced ability of mitochondrial capture in cancer cells.^{29,42,44} Further, as MT transfer is reported to promote recipient cell proliferation and survival through improved energy production,^{45,46} we assessed the association of TMT score with cell cycle activity.⁴⁷ To get an unbiased estimation, we applied partial Spearman correlation corrected for tumor purity for the related variables. Consistently, the TMT score is positively correlated with cell cycle score (CCS) in almost all cancer types (Figure S6D). In 12 out of 22 cancers, the TMT scores are highly correlated with CCS (partial Spearman $\rho > 0.2$, FDR < 0.05, Figure 6D), suggesting increased cancer cell proliferation induced by MT trafficking.

Hypoxia is a common signature of cancer,⁴⁸ and therefore, oxygen deficiency in the tumor microenvironment could influence the activity of MT transfer. We next investigated this potential connection using a recently published estimation of hypoxia status in the TCGA samples.⁴⁹ Interestingly, in non-small-cell lung and pancreatic cancers, tumor hypoxia level is significantly correlated with the TMT score (partial Spearman $\rho > 0.2$ adjusted for tumor purity, FDR < 0.05, Figures S6E and S6F). One possible explanation to this result is the previously reported mechanism that limited oxygen supply promotes cancer cells to hijack foreign mitochondria to keep up with energy production.⁵⁰

Finally, to investigate the potential role of MT transfer in prognosis, we performed univariate and multivariate Cox regression with TMT score as the primary predictor, while controlled for potential clinical confounders including age, tumor grade, clinical stage, tumor purity, etc. We observed that high TMT scores were consistently associated with poor survival in multiple cancer types (Figure 6E), including breast, head and neck, brain, liver, lung, and pancreatic cancers (Figures 6F–6K). Given the high correlation between TMT score and cell cycle activity, we speculated that TMT score's impact over patient outcome might be through CCS, which has also been reported to influence

(C) The TMT scores of primary tumor and adjacent (Adj) samples in 12 cancer types. BLCA: bladder urothelial carcinoma, BRCA: breast invasive carcinoma, HNSC: head and neck cancer, KICH: kidney chromophobe, KIRC: kidney renal clear cell carcinoma, KIRP: kidney renal papillary cell carcinoma, LIHC: liver hepatocellular carcinoma, PRAD: prostate adenocarcinoma, THCA: thyroid carcinoma, UCEC: uterine corpus endometrial carcinoma. Statistical significance was estimated using non-parametric paired Wilcoxon signed-rank test, with FDR adjusted using Benjamini-Hochberg procedure. Box center line: median; box limits: upper and lower quantiles; box whiskers: $1.5 \times \text{IQR}$.

(D) Lowess smooth curves showing positive correlations between TMT scores and CCS in selected cancer types.

(E) Association of TMT score with patient overall survival based on both univariate and multivariate Cox proportional hazards models in different cancer types. CESC: cervical squamous cell carcinoma and endocervical adenocarcinoma, COAD: colon adenocarcinoma, GBM: glioblastoma multiforme, LGG: lower-grade glioma, OV: ovarian serous cystadenocarcinoma, PAAD: pancreatic adenocarcinoma, PCPG: pheochromocytoma and paraganglioma, READ: rectum adenocarcinoma, SARC: sarcoma, SKCM: skin cutaneous melanoma, TGCT: testicular germ cell tumors. We applied two multivariate Cox models: one with just well-known clinical confounders as covariates and the other with clinical confounders plus CCS as a covariate. Size denotes statistical significance at the cutoff of FDR = 0.1; color denotes the hazard ratio.

(F–K) Kaplan-Meier estimates of overall survival, according to TMT score calculated from RNA-seq data of BRCA (F), HNSC (G), LGG (H), LIHC (I), LUAD (J), and PAAD (K). The patients were stratified into two groups (TMT high and low) based on median value of TMT scores. Statistical significance was evaluated using log rank test. See also Figure S6 and Table S6.

survival.⁴⁷ Interestingly, when adding CCS as a covariate in Cox model, the hazard ratio of TMT score became non-significant in brain (LGG) and pancreatic (PAAD) cancers, while the other cancer types remain unaffected (Figure 6E). These results provided evidence that MT acquisition by cancer cells might promote proliferation, which ultimately caused shortened survival in selected cancers. Meanwhile, TMT score still serves as an independent prognostic predictor for other cancer types, which is in line with the fact that mitochondrial transfer metabolically empowers the aggressive behavior of cancer cells and causes immune cell dysfunction.²⁶

DISCUSSION

In this work, we described MERCI for tracing MT transfer from T cells to cancer through combined use of MT mutation and gene expression profiles from scRNA-seq samples. MERCI was validated and benchmarked using experimentally generated ground-truth data, with demonstrated utilities in the real-world settings. Application of MERCI to the human cancer samples uncovered a previously unreported cancer phenotype related to MT transfer. Analysis of large human patient cohorts through TMT score further revealed that MT transfer might negatively influence patient outcome, partially through promoting cancer cell proliferation.

Nuclear mitochondrial DNA segments (NUMT)^{51,52} generated via the translocation of mtDNA fragments into the nuclear genome have been presented as a confounder in studies using MT variants as cell lineage markers. This is because variants derived from the NUMT regions usually satisfy the inclusion criteria designed for lineage tracing, such as high cellular heteroplasmy. NUMT were not explicitly handled our study, though, as they usually occur in all cell types and will unlikely contribute T cell-enriched variants for the downstream inference in MERCI.

Horizontal MT transfer between mammalian cells is mediated by TNTs, gap junctions, microvesicles, or direct extrusion and internalization,^{24,53,54} with TNT formation being the most prevalent mechanism for intercellular mitochondrial transport.^{20,29} TNT-mediated MT hijack from T cells might serve as another immune-evasive approach *in vivo*.²⁶ Thus, selectively blocking the formation of TNTs can be a promising antitumor therapeutic opportunity. However, due to the lack of specific TNT markers, the current pan-inhibitors (such as farnesyltransferase inhibitors and cytochalasin) only partially prohibit TNT formation,^{26,55} leading to limited tumor growth control. We discovered that MT transfer is highly associated with many putative regulators of the actin filament network (Tables S3 and S4), such as *PIM1*, *MYO1B*, *PFN1*, and *ABI1*, which might be candidates for TNT disruption.^{24,56,57} Another interesting target is cell adhesion molecule *PVRL2* (encoding Nectin-2), a top predicted marker of MT transfer (Figure 5D). *PVRL2* might participate in MT transport, as its disruption results in morphologically aberrant spermatozoa of mice with defects in mitochondrial localization.⁵⁸

The dysfunction of tumor-infiltrating lymphocytes (TILs) is one of the main reasons for uncontrolled tumor progression.^{59,60} In parallel to the heavily investigated immune checkpoint molecules (such as PD-1, LAG3, TIM-3, and CTLA-4), the newly discovered mitochondrial drainage by cancer cells provided another possible mechanism for the metabolic dysfunction and

exhaustion of TILs. As reported in previous work, partial loss of mitochondria in T cells may lead to shortage of energy production and dysfunction.^{61,62} Capturing the signatures of T cell exhaustion resulted from MT transfer may reveal novel genes or pathways for T cell revitalization. Although MERCI was optimized to detect gain-of-MT signals in the cancer cells, its framework could be reconfigured to detect donor T cells with proper benchmark experiments and training data. We thus anticipate a better understanding of tumor-immune interactions to unfold with the future investigation of the donor T cells in the process of MT transfer. Related to this point, in fact, both this study and the study by Saha et al.²⁶ observed a small fraction (3%–5%) of T cells that received some mitochondria from cancer cells. This signal is much weaker compared to the transfer from T cells to cancer cells, both in receiver cell percentage as well as the amount of mitochondrial translocation. Although weak, this effect might be useful to detect antigen-specific T cells closely attached to the cancer cells.

Limitations of the study

While our analyses provided more insights into this process, there are several important aspects that remain unknown. First, what is the driving mechanism that regulates mitochondria uptake by the cancer cells? GTPase *Miro1* has been reported to mediate mitochondrial trafficking,^{63,64} yet it did not emerge as a top hit in the unbiased genomic analysis. Our data suggest that MT receivers upregulated multiple genes involved in cell junction and cytoskeleton remodeling pathways (*ACTB*, *GBPI/2*, *PIM1*, *PFN1*, etc) that might directly or indirectly influence MT trafficking.^{65,66} Selective knockout of these top targets will be useful to illustrate their roles in mitochondrial transfer. Second, can non-malignant cells acquire mitochondria from T cells under physiological conditions? Mitochondrial transfer to damaged or stressed epithelial cells has been repeatedly observed in previous studies as a way of quick tissue repair.^{67,68} So far, mitochondrial drainage targeting T cells from any non-malignant cell types has not been reported. We posit that due to their small size and relatively smaller mitochondrial reservoir, T cells are not ideal donors of mitochondria in damaged tissues. Last, why different cancers (lung, breast, pancreatic, skin, etc) all demonstrated the same phenotype? Despite high level of heterogeneity, cancer cells exhibited common hallmarks, such as sustained growth signaling, metastasis, immune evasion, etc.⁶⁹ It is possible that drainage of mitochondria from tumor-infiltrating T cells delivers selective advantages that caused the convergent evolution of cancer cells with diverse genetic backgrounds. While immune evasion is an apparent mechanism for survival benefit,⁷⁰ further investigations will be needed to confirm: 1) if tumor-associated T cells are preferentially targeted in this process and 2) if and how T cells become dysfunctional after mitochondrial transfer.

Low detection sensitivity is a potential limitation of MERCI because per cell MT read coverage could be low for some 10x libraries, leading to compromised detection power of MT mutations. Using libraries sequenced at 100K reads per cell (2.3K reads in the MT region), MERCI reached 56.3% sensitivity in our benchmark. Further increase of coverage of the mitochondrial genome is expected to improve its performance. Recently, Tyler et al. introduced the MAESTER scRNA-seq platform, which combined

standard 10x protocol with mitochondrial transcriptome enrichment to increase per site and per allele MT coverage for each cell, enabling high-confidence MT variant detection in scRNA-seq dataset.³³ mtscATAC-seq implemented in this study is another example of successful mtDNA enrichment for single-cell genomic analysis. Adaptation of these new technologies for future data generation will likely increase the prediction accuracy of MERCI and broaden its application. In addition, although we defined TMT score to investigate MT transfer in the bulk tumor samples, it should be noted that this score is derived from a composite gene signature that might not be exclusive to this biological process. Conclusions regarding this analysis require future validations with tailored clinical samples.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.ccell.2023.09.003>.

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AUTHOR CONTRIBUTIONS

B.L. and H.Z. conceived of this project. H.Z. and X.Y. wrote the MERCI-mtSNP and MERCI R package. H.Z. and X.Y. performed the benchmark analysis. A.Z., J.Y., and H.L. carried out the cell culture research, animal care, and single-cell experiments. A.Z. and Y.F. provided the mice and reagents. J.H. and Y.T. helped the data analysis. H.Z. wrote the manuscript together with X.Y. and B.L. F.Y., C.W., and V.G.S. helped with mtscATAC-seq protocol and manuscript revision. J.Y. generated the used sequencing data. E.A. provided the KP cancer cells. J.M., R.B., and E.M. provided experimental reagents. B.L. and A.Z. supervised the whole project.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Alexa Fluor® 700 anti-mCD8 α (clone53-6.7)	Biolegend	Cat# 100730; RRID: AB_493703
Ultra-LEAF™ Purified anti-mouse CD3 Antibody	Biolegend	Cat# 100239; RRID: AB_2810313
Ultra-LEAF™ Purified anti-mouse CD28 Antibody	Biolegend	Cat# 102115; RRID: AB_11150408
Chemicals, peptides, and recombinant proteins		
MitoTracker™ green FM	ThermoFisher	Cat# M7514
MitoTracker™ red CMXRos	ThermoFisher	Cat# M7512
PEI MAX	Polysciences	Cat# 24765-1
Fixable viability Dye eFluor 506	eBioscience	Cat# 65-0866-18
Critical commercial assays		
CD8 $^{+}$ T cell isolation kit	Stem Cell Technologies	Cat# 19853
Chromium Next GEM Single Cell ATAC Kit v2	10x Genomics	PN-1000406
Chromium Single Cell V(D)J Reagent Kit v1.1	10x Genomics	PN-1000167
Chromium Next GEM Chip H Single Cell Kit	10x Genomics	PN-1000162
Agilent High Sensitivity DNA Kit	Agilent Technologies	Cat# 5067-4626
Mitochondria Isolation Kit for Cultured Cells	Thermo Fisher Scientific	Cat# 89874
Deposited data		
Raw and processed data	This paper	GEO: GSE235675
Processed data	This paper	Zenodo: https://doi.org/10.5281/zenodo.8065206
Human reference genome UCSC, GRCh38; Murine reference genome UCSC, mm10	University of California Santa Cruz	https://genome.ucsc.edu/cgi-bin/hgGateway
Full list of public single cell datasets used in this study	N/A	Table S1
scRNA-seq data for BCC samples	Yost et al. ³⁷	GEO: GSE123814
scRNA-seq data for ESCC samples	Zhang et al. ³⁸	GEO: GSE160269
scRNA-seq data for NSCLC samples	Zeng et al. ⁴¹	EBI: E-MTAB-6149
scRNA-seq data for CRC samples	Zeng et al. ⁴¹	EBI: E-MTAB-8410
scRNA-seq data for MCC samples	Zeng et al. ⁴¹	GEO: GSE117988
scRNA-seq data for MCC samples	Zeng et al. ⁴¹	GEO: GSE118056
scRNA-seq data for LUAD samples	Zeng et al. ⁴¹	GEO: GSE123904
scRNA-seq data for STAD samples	Zeng et al. ⁴¹	GEO: GSE134520
scRNA-seq data for NSCLC samples	Zeng et al. ⁴¹	GEO: GSE143423
scRNA-seq data for TNBC samples	Zeng et al. ⁴¹	GEO: GSE148673
scRNA-seq data for MIUBC samples	Zeng et al. ⁴¹	GEO: GSE145137
scRNA-seq data for PDAC samples	Zeng et al. ⁴¹	GSA: PRJCA001063
bulk RNA sequencing data for cancer samples	TCGA	https://portal.gdc.cancer.gov/
Experimental models: Cell lines		
Kras ^{G12D} /p53 ^{ko} (KP) lung cancer cells	Dr. Esra Akbay	N/A
KP-mito-DsRed	This paper	N/A
MC38 colon cancer cells	ATCC	N/A

(Continued on next page)

Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Experimental models: Organisms/strains		
C57BL/6J	Jackson Laboratory	Cat# 000664
Recombinant DNA		
pDsRed2-Mito	Takara	Cat# 632421
Software and algorithms		
MERCI-mtSNP and MERCI R package code	This paper	https://github.com/shyhihihi/MERCI
CytExpert	Beckman Coulter, Inc	https://www.beckman.com/coulter-flow-cytometers/cytoflex/cytextpert
FlowJo	Tree Star Inc.	https://www.flowjo.com/solutions/flowjo
GraphPad Prism software 7.0	GraphPad Software, Inc.	https://graphpad.com/scientific-software/prism/
Python version 3.8	Python Software Foundation	https://www.python.org/downloads/
R version 4.0.2	The R Foundation	https://www.r-project.org/
Cell Ranger 3.1.0	10x Genomics	https://10xgenomics.com/
Cell Ranger ATAC 2.0.0	10x Genomics	https://10xgenomics.com/
Seurat 3.1.2	Seurat developers	https://satijalab.org/seurat/articles/get_started.html
pysam	pysam developers	https://github.com/pysam-developers/pysam
e1071(version 1.7.4)	N/A	https://cran.r-project.org/web/packages/e1071/index.html
fgsea(version 1.16.0)	N/A	https://bioconductor.org/packages/release/bioc/html/fgsea.html
msigdb (version 7.2.1)	N/A	https://cran.r-project.org/web/packages/msigdb/index.html
simplifyEnrichment (version 1.0.0)	N/A	https://bioconductor.org/packages/release/bioc/html/simplifyEnrichment.html
Cytoscape (version 3.9.1)	Cytoscape developers	https://cytoscape.org/
GSVA	Hänzelmann et al. ⁴³	https://bioconductor.org/packages/release/bioc/html/GSVA.html
BioRender	biorender.com	https://www.biorender.com/

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources, reagents and codes should be directed to and will be fulfilled by the lead contact, Dr. Bo Li (liib3@chop.edu).

Materials availability

This study did not generate any new unique reagents or models.

Data and code availability

scRNA-seq datasets generated in this study, as well as the MT variants and coverage information are available at Zenodo <https://doi.org/10.5281/zenodo.8065206>, and Gene Expression Omnibus (GEO) database with accession number GEO: GSE235675. The other scRNA-seq datasets of human cancer are available under the GEO accession codes provided in Table S1 and key resources table. The TCGA data is available at <https://portal.gdc.cancer.gov/>. The source codes of MERCI-mtSNP and MERCI R package, and the codes to run example data are available at: <https://github.com/shyhihihi/MERCI>. Any additional information is also available from the corresponding author (Dr. Bo Li) upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Cancer cell line and culture

The KP lung cancer cells were provided initially by Dr. Esra Akbay. The KP-mito-DsRed stable cell line was generated in the lab with pDsRed2-Mito vector (Takara, Cat# 632421) transfection using PEI MAX® (Polysciences, Cat# 24765-1) with long-term selection using G418 following manufacture's protocol. KP cell lines were grown in 5% FBS (ThermoFisher Scientific, Cat# 26140079) supplemented RPMI 1640 (Millipore Sigma, Cat# R8758). Cell lines were routinely tested using a mycoplasma contamination kit (E-myc, Cat# 25233, Bulldog Bio) and cultured under 5% CO₂ at 37 °C. The MC38 cell lines were purchased from the American Type Culture Collection and cultured in DMEM supplemented with 10% heat-inactivated FBS, 100 U ml⁻¹ penicillin and 100 U ml⁻¹ streptomycin under 5% CO₂ at 37 °C.

Mice

6-8 weeks old female C57BL/6J mice were purchased from Jackson Laboratory and housed at UTSW Animal Resource Center. All mice were maintained at 68-79 °F, 30-70% humidity, in individually ventilated cages, with no more than five mice per cage on 12-hour on:off light:dark cycles. Mice had unrestricted access to RO chlorinated water and irradiated 2916 Teklab global diet (Envigo, Cat# 2916). The mice were housed at least for a week before starting the downstream process. Animal care and experiments were performed to comply with institutional and National Institutes of Health protocols and guidelines. This study has been approved by the Institutional Animal Care and Use Committee of University of Texas Southwestern Medical Center.

METHOD DETAILS

T cell isolation, activation and mitochondria labeling

Splenocytes were isolated by harvesting the spleen from 6-8 weeks old mice following red blood cell lysis with ACK buffer. CD8⁺ T cells were isolated by immunomagnetic negative selection using EasySep™ mouse CD8⁺ T cell isolation kit (Stem Cell Technologies, Cat# 19853). Isolated T cells were activated with 1 µg/mL anti-CD3/CD28 (Biolegend, Cat#100239 and Cat#102115) antibodies for 24h. Activated CD8⁺ T cells were then stained with 100 nM MitoTracker™ green FM (ThermoFisher, Cat# M7514) at 37 °C for 30 min following the manufacturer's instruction. MitoTracker™ green FM (ThermoFisher, Cat# M7514) at indicated concentration would not stain all T cell mitochondria positive, to avoid non-mitochondrial labeling. Stained CD8⁺ T cells were then washed with medium, centrifuged down, and then resuspended to remove exceeded dye in the medium. Isolated CD8⁺ T cells were maintained in 10% heat deactivated FBS supplemented RPMI 1640 (Millipore Sigma, Cat# R8758). In the control experiment, KP-mito-DsRed cancer cells were labeled with MitoTracker™ red CMXRos (ThermoFisher, Cat# M7512) using the same procedure.

Coculture assay and flow cytometry analysis

The KP-mito-DsRed cancer cells were plated at 2x10⁶ cells/well of 6 well plate overnight to allow cell attachment before co-culturing. MitoTracker™ green FM stained CD8⁺ T cells were then added to the tumor cells at 2x10⁵ cells/well with a final volume of 2ml of RPMI 1640 supplemented with 10% heat deactivated FBS in each well. Cancer cells and CD8⁺ T cells were then cocultured under 5% CO₂ at 37 °C for 24h before mitochondria horizontal transfer evaluation. Co-cultured supernatant containing CD8⁺ T cells were then collected, as well as 0.25% trypsinized co-cultured KP-mito-DsRed cells. All the cells were spined down, and a single cell suspension was used to perform staining and analysis. Samples were incubated with anti-FcγIII/II receptor (clone 2.4G2) for 15 min at 4 °C to block potential non-specific binding of conjugated antibodies. Anti-mCD8α (Biolegend, Cat# 100730) were incubated with single-cell suspension for 30 min at 4 °C. Fixable viability Dye eFluor 506 (eBioscience, Cat# 65-0866-18) was used to exclude the dead cells. Subsequently, the dissociated cells were filtered through a 70 µm cell strainer and resuspended in FACS buffer before analysis. Data were collected and analyzed on CytoFlex flow cytometer (Beckman Coulter, Inc). To control the effect of different co-culture ratios on mitochondrial transfer. A wide range of co-culture ratios (10:1, 5:1 and 2:1) were also tested in our control experiments for KP and MC38 cancer cells.

Single cell suspension preparation and flow sorting

CD8⁺ T cells successfully labeled with MitoTracker™ were sorted as FITC positive through BD FACSMelody™ sorter. Single-cell suspension of co-cultured KP cells (either DsRed or mitoTracker Red labeled cells) and CD8⁺ T cells were prepared and stained as described during flow cytometry analysis. KP cells containing mitochondria derived from T cells (CD8 negative, FITC and DsRed/ MitoTracker Red positive) and CD8⁺ T cells were sorted with BD Melody flow sorter. Monocultured cancer cells and CD8⁺ T cells were prepared as single-cell suspension simultaneously following the same protocol as the cocultured cancer cells.

mtscATAC-seq libraries preparation and sequencing

Mitochondrial single-cell assay for transposase-accessible chromatin with sequencing (mtscATAC-seq) technique was performed strictly following the revised 10x scATAC-seq protocol developed by Lareau et al.^{10,71} Briefly, after washing, the sorted cells were fixed in 1% formaldehyde (FA; ThermoFisher #28906) in PBS for 10 min at RT, quenched with glycine solution to a final concentration of 0.125 M before washing cells twice in PBS via centrifugation at 400 g, 5 min, 4 °C. Cells were subsequently treated with lysis buffer (10mM Tris-HCL pH 7.4, 10mM NaCl, 3mM MgCl₂, 0.1% NP40, 1% BSA) for 3 min for CD8⁺ T cells and 5 min for KP cells on ice,

followed by adding 1 ml of chilled wash buffer and inversion (10mM Tris-HCL pH 7.4, 10mM NaCl, 3mM MgCl₂, 1% BSA) before centrifugation at 500 g, 5 min, 4°C. The supernatant was discarded, and cells were diluted in 1x Diluted Nuclei buffer (10x Genomics) before counting using Trypan Blue and a Countess II FL Automated Cell Counter (Invitrogen). mtscATAC-seq libraries were generated using the 10x Chromium Controller and the Chromium Next GEM Single Cell ATAC Kit v2 (PN-1000406) according to the manufacturer's instructions (CG000496 Rev-B), and sequencing was run on Illumina NovaSeq 6000 using paired-end sequencing of 150-bp read length, aiming to allocate 35,000 paired-end reads per cell in a mtscATAC-seq library.

scRNA-seq libraries preparation and sequencing

scRNA-seq was performed by the 10x Genomics single-cell 5' V(D)J library platform. Single-cell suspension of negative controls and sorted cells were counted manually under a microscope stained with trypan blue. The concentration of single-cell suspensions was adjusted to 900-1000 live cell/μl. Cells were loaded at 10,000 cells/chip position. Single-cell libraries were generated with Chromium Single Cell V(D)J Reagent Kit (10x Genomics, PN-10000006, PN-10000009) following per manufacturer's instruction. Purified libraries were analyzed by Illumina NovaSeq with 150-bp paired-end reads at a targeted median read depth of 100,000 reads per cell.

Data processing and read alignment

For each scRNA-seq library used in this study, fastq files were processed using the Cell Ranger (v6.0.0) Software Suite. Cellranger count was used to align reads to mouse mm10 or human GRCh38 reference genome, as well as to filter and quantify. The generated files of Cell Ranger for each library included a cell barcode file, a feature file and a gene expression matrix (in unique molecular identifier (UMI) count). Seurat pipeline (v3.1.2) was applied to the raw UMI count table, following log-normalization. A normalized gene expression matrix was generated for each sample. The cell barcode file was used for subsequent mitochondrial variant calling, and only MT reads associated with cell barcodes listed in this file were retained. The raw data from mtscATAC-seq libraries underwent preprocessing with the Cell Ranger ATAC pipeline (version 2.0.0) and alignment with the mm10 reference via the cellranger-atac count command. The cell barcodes listed in 'filtered_peak_bc_matrix' of the output files were utilized for subsequent mitochondrial variants calling.

Analysis of mouse scRNA-seq data

All normalized expression matrices were merged into one file by the standard Seurat v3 integration workflow⁷² to remove potential technical variabilities. Since our purpose is to analyze the mitochondrial genomic feature of every single cell, we performed an additional filtering operation to remove those cells with low read depths in the mitochondrial genome region. The aligned and processed .bam files generated from Cell Ranger were further processed using customized Python scripts. Cells with at least 1,000 reads uniquely mapping to the mitochondrial genome were considered for downstream analysis.

UMAP projection using mitochondrial reads

To visualize the mitochondrial transcriptional data in a 2-dimensional space, we performed a uniform manifold approximation and projection (UMAP) embedding of the cells by their mitochondrial read coverage. First, we extracted all the high-quality reads (mapping score=255) that uniquely mapped to the mitochondrial genome, and calculated the basewise coverage for each cell. When counting the coverage for each base, we only considered the reads with base quality > 15. The 16.3-kb long genome of mouse MT provided a read-depth matrix, with 16.3 k rows and columns corresponding to each cell. Raw read counts were then log-normalized for each cell using Seurat. The ScaleData function was used to calculate the scaled z-scores for mitochondrial genome sites, which were used as input of the principal component analysis (PCA). With the first 10 principal components as features, the UMAP projections were generated using the RunUMAP function.

Overview of MERCI

Mitochondrial variant calling by MERCI-mtSNP

To efficiently retrieve the cell-level mitochondrial variants from single-cell sequencing data, we developed a mutation calling pipeline called MERCI-mtSNP (Figure S2B). MERCI-mtSNP was designed to call single-nucleotide variants (SNV) in MT genomics data generated from popular bulk or single-cell sequencing technologies, such as 10x Genomics scRNA-seq, smart-seq2 and ATAC-seq. More specifically, the aligned bam file was used as the input of MERCI-mtSNP, and all reads uniquely aligned to the mitochondrial genome were extracted. For 10x scRNA-seq and mtscATAC-seq data, the reads were further separated by cell barcodes. This step generated a new mitochondrial bam (MT bam) file only containing those extracted mitochondrial reads. We performed variant calling for each cell independently, keeping cells with at least K mitochondrial reads for the scRNA-seq (K=1,000) and mtscATAC-seq (K=2,000) data. The imported MT bam file using the pysam module determined per base and per allele read counts. We removed those reads marked with 'duplicate', 'secondary', 'supplementary' or 'unmapped', and only kept the reads with the high mapping score (255 for scRNA-seq, >=5 for mtscATAC-seq). In mtscATAC-seq data, we observed that some regions in the mtDNA regions are similar to the mouse genome, resulting in multiple mapping locations low mapping score. To include reads in these regions, we lowered the mapping quality cutoff while restricted the primary mapping location to mtDNA. Per-based coverage statistics were then calculated using the remaining mitochondrial reads. For each altered base at a given locus, the variant allele frequency (VAF) was computed by using the number of the supporting reads divided by the total read depth. To get high-quality variants, only the aligned reads with base-quality score >15 (>25 for mtscATAC-seq) were used to calculate the VAF values. In the scRNA-seq data, we noticed

the existence of ‘mutation clusters’, which are consecutive variants occurring within a small genomic region, which are likely to be some false positives due to technical artifact. Therefore, variants observed in the clustered genome loci (within 5 bp) with only supported by one altered read were removed. A csv file and txt file would be generated as the output of MERCI-mtSNP, representing the information of MT coverage and MT variants, respectively.

Since MERCI-mtSNP ignores the mutations in the nuclear genome, it has a favorable computational efficiency with 52s for running out a single cell smart-seq2 data (450,358 MT reads) or 2.5h for a normal-sized 10x scRNA-seq data (8,721 cells, 1,147 MT reads per cell) with 256G memory and 12 CPU nodes.

Binomial model-based filtering of MT variants

Assuming a mitochondrial single nucleotide variant (SNV) with a true heteroplasmy level p , the statistical power for detecting this mutation depends on the local sequencing coverage n at the site of this variant. It should be noted that heteroplasmy indicates the proportion of mitochondrial genomes containing a specific mutation, which is different with observed VAF. VAF is usually determined by multiple factors including heteroplasmy, sequencing error and read depth. Assuming the sequencing error to be e ($e=0.001$), the probability of observing at least one alternate read can be written as:

$$P(p, n) = 1 - p_w^n$$

in which

$$p_w = 1 - p_{mut}$$

$$p_{mut} = p \times (1 - e) + (1 - p) \times \frac{e}{3}$$

Where p_{mut} and p_w represent the probabilities of a read being altered and wild type, respectively. The probability of detecting mutation at a heteroplasmy of 0.2 is greater than 60% when read coverage is 5 or higher (Figure S2F). Therefore, only variants with at least 5 total reads (wild-type and variant) reads were kept. For a given cell and a given variant, if the coverage at the locus is ≥ 5 , we define this variant “observable” in this cell. Variants observable in less than 10% of cells were removed. Unobservable positions are represented as missing values. We implemented additional filtering for mtscATAC-seq data because its comparatively higher coverage per site per cell can result in more sequencing errors. Specifically, the probability of observing at least m identical alternate reads due to sequencing error can be expressed as follows:

$$P(m) = \begin{cases} 1 & \text{if } m = 0 \\ 1 - \sum_{i=0}^{m-1} \text{Binom}\left(i|n, \frac{e}{3}\right) & \text{if } m > 0 \end{cases}$$

The minimum number of alternate reads k supporting that $P(k)$ is less than a given false-positive rate (FPR) is:

$$k = \min(m) | P(m) \leq FPR$$

With FPR set to 5×10^{-4} , we can compute a k value for each variant. All variants with supporting reads fewer than k were considered as potential sequencing errors ($FPR > 5 \times 10^{-4}$), and removed from the mtSNV profile. Finally, we obtained a matrix of VAF with rows being SNPs and columns being cells, which was further transformed into a binary matrix: with 0 being VAF=0 and 1 VAF>0.

Defining T cell enriched/depleted MT variants

The monocultured cancer cells were randomly split into a training and a validation set. T cell enriched mtSNVs were defined as those more frequently occurring in T cells than in cancer cells of the training set ($FDR < 0.1$, $OR > 1$). Similarly, T cell depleted variants represented the mutations less observed in T cells ($FDR < 0.1$, $OR < 1$). Statistical significance was estimated using Chi-square test followed by FDR correction with the Benjamini–Hochberg (BH) procedure.

‘Effective count statistic’ of T cell enriched mtSNV

The observation of T cell enriched mtSNVs in cancer cells provides key evidence of MT transfer. In this study, we consider each cancer cell as a potential receiver of mitochondria from T cells. Let S denote the set of mutations that are enriched in the T cell population. For a given cancer cell and mutation i in S , we observe the coverage of this site n_i and the count of mutated reads m_i ($m_i > 0$). Although the ratio m_i/n_i is an unbiased estimation of the heteroplasmy of the mutation i , given the relatively shallow per-cell coverage of the single cell RNA-seq data, this estimate is noisy and can be inaccurate. Therefore, we used the following hierarchical Bayesian approach to determine if mutation i is resulted from MT transfer from T cells or otherwise. We define a weight w_i associated with mutation i as the ratio of marginal likelihood:

$$w_i = \frac{P_i}{Q_i}, i \in S \quad (\text{Equation 1})$$

where

$$P_i = \Pr(m_i | n_i, \text{MT transfer})$$

$$Q_i = \Pr(m_i | n_i, \text{no MT transfer})$$

Intuitively, if MT transfer occurs, we anticipate $w_i > 1$ for mutation i . Next, we calculate the likelihood given the probability p_i^0 of observing a mutated read:

$$\Pr(m_i | n_i, p) = \binom{n_i}{m_i} \times (p_i^0)^{m_i} \times (1 - p_i^0)^{n_i - m_i} \quad (\text{Equation 2})$$

Under the null scenario, i.e. variant coming from either sequencing error or as a result of the background population heteroplasmy p_i^c :

$$p_i^0 = p_i^c \times (1 - e) + (1 - p_i^c) \times \frac{e}{3} \quad (\text{Equation 3})$$

The variable p_i^c in Equation 3 can be estimated by pooling all the single cells from the given cell population. Assuming that a total of M_i altered reads support the mutation i , with sequencing depth N_i , then $E(p_i^c) = \frac{M_i}{N_i}$. According to the Central Limited Theorem, when N_i is a sufficiently large number, p_i^c is asymptotically Normal with estimated deviation:

$$\sigma_i^c = \sqrt{\frac{M_i \times (N_i - M_i)}{N_i^3}}$$

Then the standard deviation of p_i^0 should be:

$$\sigma_i^o = \left(1 - \frac{4}{3}e\right) \times \sigma_i^c$$

We hypothesized that the condition of with or without MT transfer will alter the range of the true heteroplasmy of the variant allele in the expressions of P_i and Q_i in Equation 1. Specifically, when there is no MT transfer, it is reasonable to believe that the mutation allele frequency p_i is unlikely to take a value that is 'a lot' higher than p_i^0 , with 'a lot' quantified by σ_i^o . Given the property of Normal distribution, we set the upper limit of p_i in the case of no MT transfer to be $s_1 = \min(p_i^0 + 1.96\sigma_i^o, 1)$, such that p_i will be smaller than this limit 97.5% of the chance. As it is not possible to learn which T cells had been in contact with the cancer cell of interest, when MT transfer occurs, we assume that the heteroplasmy of the variant in the T cell population can uniformly take value somewhere between p_i^c to 1. Considering the statistical uncertainty, we anticipate p_i to be higher than $s_2 = \max(p_i^0 - 1.96\sigma_i^o, 0)$. Accordingly,

$$P_i = \frac{n_i!}{m_i! \times (n_i - m_i)!} \times \int_{s_2}^1 (p_i)^{m_i} \times (1 - p_i)^{(n_i - m_i)} dp_i$$

$$Q_i = \frac{n_i!}{m_i! \times (n_i - m_i)!} \times \int_0^{s_1} (p_i)^{m_i} \times (1 - p_i)^{(n_i - m_i)} dp_i$$

The integral can be written as a form of incomplete Beta function to get the expression of w_i :

$$w_i = \frac{P_i}{Q_i} = \frac{1 - I_{s_2}(m_i+1, n_i - m_i+1)}{I_{s_1}(m_i+1, n_i - m_i+1)} \quad (\text{Equation 4})$$

where

$$I_x(m_i + 1, n_i - m_i + 1) = \frac{\int_0^x t^{m_i} \times (1 - t)^{(n_i - m_i)} dt}{\text{Beta}(m_i+1, n_i - m_i+1)}$$

$$= \sum_{j=m_i+1}^{n_i+1} \binom{n_i+1}{j} x^j (1 - x)^{n_i+1-j}$$

Therefore, the final expression of w_i is:

$$w_i = \frac{1 - \sum_{j=m_i+1}^{n_i+1} \binom{n_i+1}{j} s_2^j (1 - s_2)^{n_i+1-j}}{\sum_{j=m_i+1}^{n_i+1} \binom{n_i+1}{j} s_1^j (1 - s_1)^{n_i+1-j}} \quad (\text{Equation 5})$$

All the variables in the Equation 5 can be estimated from the single cell RNA-seq data. Based on the value of w_i , we define the 'effective count statistic' (N_{eff}) of each cancer cell as follows:

$$N_{\text{eff}} = \sum_{i \in S} I(w_i)$$

in which $I()$ is indicator function:

$$I(x) = \begin{cases} 1 & \text{if } x > 1 \\ 0 & \text{otherwise} \end{cases}$$

Effective count N_{eff} is an estimate of MT transfer occurrence for each cancer cell of interest based on T cell enriched MT mutations, while controlled for sequencing error and read depth. We use N_{eff} in MERCI as the DNA source of information, which is further taken the rank transformation to reduce the impact of outliers.

Deconvolution model with support vector regression

The mixed mitochondrial transcriptome expressed in receiver cells allowed us to propose a deconvolution model on mitochondrial mRNA mixture \mathbf{m} for imputing an unknown fraction of mitochondrial composition for each cell.^{34,73,74} The mitochondrial genome contains 13 protein-coding genes (MT genes) that provide essential functions for making enzymes involved in oxidative phosphorylation. The mRNA expression of each MT gene in a receiver cell is expected to be contributed by both intrinsic (endogenous) MT and donor-transferred (exogenous) MT. Thus, the deconvolution problem can be represented by the equation:

$$\mathbf{m} = \mathbf{f} * \mathbf{g} + \delta$$

Where \mathbf{f} denotes a vector of the mitochondrial fraction of different source of origins in the mixture and \mathbf{g} is the signature matrix of MT gene expression profiles (MGEP) in purified donor and receiver cell types. δ is the random noise following a normal distribution with zero mean. In this work, MERCI adopted epsilon-support vector regression (ϵ -SVR) with linear kernel application³⁵ to estimate the fraction of external mitochondria by decomposing the relative RNA transcripts of MT genes. ϵ -SVR is an extended use of support vector machine (SVM) in regression problems. Unlike ordinary linear regression, the objective of ϵ -SVR is to find an appropriate line (or hyperplane in higher dimensions) to fit the data with acceptable error (ϵ) instead of minimizing the squared error sum. Therefore, ϵ -SVR can largely reduce overfitting and be robust to noisy data, making it suitable for deconvolving the mixture profile of gene expression.^{34,75}

For a given receiver cell, the regression line of ϵ -SVR can be written in the equation:

$$\hat{y} = w * x + b \quad (\text{Equation 6})$$

Where $w = (w_1, w_2)$ denotes the weights of the exogenous (w_1) and endogenous (w_2) MT within the cytoplasm of this cell, and x is the vectorized reference MGEP of potential donor cells and receiver cells. This formula represents the linear mixture of different cell types' gene expression pattern with trainable weights. The goal is to minimize the difference between the mixed expression pattern and the observed data at given constraints. Let y denote the observed expression of a MT gene. To solve the weights, the following objective function of ϵ -SVR (L_2 -norm penalty function) and constraints are used:

$$\min \left(\frac{1}{2} \|w\|^2 + C \sum_{i=1}^N (\xi_i + \xi_i^*) \right) \quad (\text{Equation 7})$$

With Constraints:

$$y_i - \hat{y}_i \leq \epsilon + \xi_i \quad (\text{Equation 8})$$

$$\hat{y}_i - y_i \leq \epsilon + \xi_i^* \quad (\text{Equation 9})$$

where N is the number of data points, $C > 0$ is the input penalty coefficient and ξ_i, ξ_i^* are positive slack variables ensuring equality. ϵ is set as 0.1 in this study.

To solve this constrained optimization problem, Lagrange multipliers $\alpha = (\alpha_1, \alpha_2, \dots, \alpha_N)$ and $\alpha^* = (\alpha_1^*, \alpha_2^*, \dots, \alpha_N^*)$ are introduced^{76,77} and the Lagrangian function L can be obtained to be:

$$L(w, b, \xi, \xi^*, \alpha, \alpha^*) = \frac{1}{2} \|w\|^2 + C \sum_{i=1}^N (\xi_i + \xi_i^*) + \sum_{i=1}^N \alpha_i (y_i - wx_i - b - \epsilon - \xi_i) + \sum_{i=1}^N \alpha_i^* (wx_i + b - y_i - \epsilon - \xi_i^*) \quad (\text{Equation 10})$$

By using the Lagrange multiplier techniques, the optimization problem in support vector regression is transformed to:

$$\min_{w,b} \left(\frac{1}{2} \|w\|^2 + C \sum_{i=1}^N (\xi_i + \xi_i^*) \right) = \min_{w,b} \max_{\alpha, \alpha^*} L(w, b, \xi, \xi^*, \alpha, \alpha^*) \quad (\text{Equation 11})$$

This leads to the following dual optimization problem.³⁵

$$\min_{w,b} \max_{\alpha, \alpha^*} L(w, b, \xi, \xi^*, \alpha, \alpha^*) = \max_{\alpha, \alpha^*} \min_{w,b} L(w, b, \xi, \xi^*, \alpha, \alpha^*) \quad (\text{Equation 12})$$

Using the condition that the partial derivatives of L with respect to the primal variables will be zero at optimality. The dual problem can be written as following⁷⁷:

Maximize

$$\max_{\alpha, \alpha^*} \left(-\varepsilon \sum_i^N (\alpha_i^* + a_i) + \sum_i^N y_i (\alpha_i^* - a_i) - \frac{1}{2} \sum_{i,j=1}^N (\alpha_i^* - a_i) (\alpha_j^* - a_j) \langle x_i, x_j \rangle \right) \quad (\text{Equation 13})$$

s.t.

$$\sum_{i=1}^N (a_i - \alpha_i^*) = 0$$

$$0 \leq a_i, \alpha_i^* \leq C$$

w in the primal formulation can be expressed easily in terms of α and α^* .

$$w = \sum_{i=1}^N (a_i - \alpha_i^*) x_i \quad (\text{Equation 14})$$

Thus, the linear ε -SVR hyperplane can also be represented by

$$y = \sum_{i=1}^N (a_i - \alpha_i^*) \langle x_i, x \rangle + b \quad (\text{Equation 15})$$

where b is computed using the fact that Equation 8 becomes an equality with $\xi_i = 0$ if $0 < \alpha_i < C$ and (9) becomes an equality with $\xi_i^* = 0$ if $0 < \alpha_i^* < C$. The current implementation of ε -SVR in MERCI is based on the R package e1071. The estimated regression coefficients (w_1 and w_2) in MERCI denote the relative, not absolute, quantity of intrinsic and foreign mitochondria in a receiver cell.

Rank score calculation

MERCI generated two rank scores (DNA rank, RNA rank) for each input candidate cell. Based on the N_{eff} statistic of donor cell (T cell in this study) enriched MT variants (\mathbf{N}_o), the DNA rank score is calculated as the descending order of \mathbf{N}_o ($R_{\mathbf{N}_o}$). The RNA rank score is obtained from decreasingly ranking the estimated abundance of donor cell derived MT calculated from the Equation 14 (R_{w_1}). In Figures S3C and S3D, the combined rank score of each cell was computed by averaging the DNA rank score and RNA rank score. MERCI considers both DNA and RNA rank scores to predict receiver cells using an AND gate. A candidate cell will be assigned as a receiver only if its DNA and RNA rank scores are both within the user-defined threshold (R_c):

$$\max (R_{\mathbf{N}_o}, R_{w_1}) \leq R_c$$

Depth-normalized counts of T cell enriched mtSNVs

The probabilities of observing a mtSNV are influenced by the read depth (Figure S2F). Therefore, we compared the observed counts of T cell enriched/depleted mtSNVs between CC cells and MC cells only within the same read depth range. We explored different ranges (1000-2000, 2000-3000, 3000-4000, 4000-5000) and showed the comparison using cells with 1000-2000 MT reads (Figure 1F) because this range covered the most of cells. We did not consider cells with MT reads > 5000 because this range contains very few (<100) MC cells.

MERCI LOO strategy for real-world application

In case the reference data of true non-receivers is not available, we used a leave-one-out strategy to get the reference MGEP and donor-enriched MT variants. When MERCI calculates the rank scores for a given cell, we left this cell in the input cell set and used remaining cells as reference non-receiver cells. The reference MGEP was then obtained by averaging the expression of MT genes in these cells. Donor cell enriched MT variants are derived from comparing the mitochondrial mutation profile of these non-reference receiver cells to the potential donor cells. The other steps are the same as the regular MERCI pipeline.

Significance estimation

Bona fide mitochondrial receivers are expected to have higher DNA and RNA rank scores. This effect causes an association between the two scores, enabled us to test if the sample contains true receivers. For example, if no receiver cell is included, the two rank scores are expected to be independent. When we set the rank threshold at top rank 10%, theoretically $\sim 1\%$ of input cells will be randomly captured By MERCI as the intersection of two cell sets: one with top 10% DNA rank scores and the other with top 10% RNA rank scores. However, if the input cells include a certain proportion of true receivers, the fraction of captured cells will be significantly higher than 1%. Thus, we used randomly permuted ranks to produce a null distribution (\mathbf{R}) of captured cell numbers. At a given rank cutoff, MERCI predicts N receivers based on two lists of random rank scores. This process is repeated 10,000 times to generate the empirical estimate of \mathbf{R} (with a list of N , or \mathbf{N}). Let N_c denotes count of predicted receivers using the original ranks, we define a statistic R_{cm} as below:

$$R_{cm} = \frac{N_c}{\max(\mathbf{N})}$$

We used a strict criterion to determine whether the receiver cells are included in the input cells or not. The captured cell number is considered significant only if $R_{cm} > 1$, with empirical p value < 0.0001 , i.e. all the predicted cell counts by random permutations are smaller than the real value.

Down-sampling analysis

In the ground-truth data, the fraction of cocultured cancer cell accounts for 43% of total input cancer cells. To analyze the dependence of the number of MERCI captured cells on the fraction of true receiver cells in the input cell set, down-sampling was performed to test the influence of signal/noise ratio on R_{cm} . Down-sampling of cocultured cancer cells generated simulated data sets with different fraction (from 0.5%~20%) of true receivers (Figures 3D–3F). There were 100 simulated data sets produced for each fixed fraction. Significance estimation and R_{cm} calculation was then repeated for these subsets of cells from ground-truth data, which enabled us to determine whether the number of captured cells is stably significant when analyzing each subset with smaller positive signal. We then calculated the R_{cm} values for each simulation to determine the power of MERCI at lower receiver cell percentage.

Analysis of BCC scRNA-seq data

We downloaded the expression data of 11 patients with advanced basal cell carcinoma (BCC) from the GEO database (GEO: GSE123814).³⁷ Of these patients, only three of them, ‘su005’, ‘su006’ and ‘su008’ contained both T cells and tumor cells, with each subset comprising more than 500 cells, and were kept for the downstream analysis. We accessed the raw fastq files of these patients from the SRA database (PRJNA509907), which were further processed with Cell Ranger and Seurat as described above. We directly used the cell type labels provided by the cell information table.³⁷ Different subpopulations of T cells such as CD8⁺ memory, CD8⁺ active and CD4⁺ T cells were merged into one T cell cluster. After obtaining the MT mutation profile for each cell, MERCI significance estimation was then performed for each patient. A significant signal was observed for the tumor cells from samples of patient ‘su006’. Thus, the MERCI LOO pipeline was applied to predict the mitochondrial receiver tumor cells and quantify the MT composition.

Analysis of DEGs and MT transfer related genes

The DEGs of the predicted receiver cells in BCC sample ‘su006’ were identified using the FindMarkers function with a Wilcoxon rank-sum test. A total of 608 DEGs were called with an adjusted FDR < 0.05 (Benjamini–Hochberg method) (Table S3). Candidate genes of MT transfer were defined as those with a significantly positive correlation with the estimated abundance of exogenous MT across cancer cells (Spearman $\rho > 0$, FDR <0.05 , two-sided Spearman’s correlation coefficient test following BH correction). Totally, 398 and 1,678 MT transfer related genes were obtained for murine CC cells and BCC tumor cells, respectively. The two lists overlapped by 95 genes, which were defined as MT transfer related genes.

Analysis of ESCC scRNA-seq data

The scRNA-seq data sets of 58 patients with esophageal squamous-cell carcinoma (ESCC) were downloaded from the GEO (GEO: GSE160269) and SRA (SRA: PRJNA672851) databases.³⁸ Of them, we only analyzed the raw fastq files for twenty-seven patients because they contained both sufficient T cells and tumor cells ($n>500$). The preprocess of single cell data is the same as the BCC data. Using T cells as potential donors, we calculated R_{cm} statistics for tumor cells (epithelial cells) of all samples. Significant signals of receiver existence were detected in samples of three patients ‘P16T’, ‘P52T’ and ‘P39T’. Next, we used MERCI LOO pipeline to predict the MT receivers from tumor cells of these three patients using the rank parameter cutoff 50%.

Analysis of other single cell transcriptome data

We investigated the single cell RNA-seq datasets in the CancerSCEM database⁴¹ and downloaded the scRNA-seq data of 156 samples with both cancer and T cells reported in the original publications. We used the same data processing workflow as described in CancerSCEM to process the downloaded data and reproduced the T cell and cancer clusters for each sample. We then applied MERCI LOO pipeline to the 61 samples that contained sufficient number (>100) of T cells and cancer cells. Subsequently, we calculated Pearson’s correlation coefficient between the estimated fraction of T cell-derived mitochondria and two UMAP coordinates

(umap1, umap2) for each sample to measure the phenotypic impact of MT transfer (Table S5). We further evaluated if a gene is associated with MT transfer by calculating the Spearman's correlation coefficient between the estimated T cell mitochondrial fraction and the gene expression level and corrected the FDR using the Benjamini-Hochberg method.

Gene set enrichment analysis

The R packages *fgesa* (v1.16.0) and *msigdbR* (v7.2.1) were used to perform the gene ontology and pathway enrichment analysis. The pathway gene sets used in this work were extracted from the online databases Kyoto Encyclopedia of Genes and Genomes (KEGG), Gene Ontology, BioCarta, Reactome and curated Hallmark gene sets.⁷⁸ Function terms or pathways were considered significant if Benjamini-Hochberg adjusted FDR < 0.05. To investigate the programs associated with MT transfer, we extracted the common GO biological processes enriched by the candidate genes of MT transfer using the benchmark data and human BCC sample, respectively. An enrichment map was produced based on these ontologies using *simplifyEnrichment* (v1.0.0) and *Cytoscape* (v3.9.1).

Analysis of MT transfer score for TCGA samples

From 95 MT transfer related genes, we selected a 17 gene signatures (*ABI1*, *ARF6*, *F11R*, *BAIAP2L1*, *MYADM*, *ACTR3*, *ARHGEF5*, *SRC*, *CAPZA2*, *GOLPH3*, *RAB10*, *ARF4*, *CAP1*, *PSEN1*, *YWHAG*, *YWHAZ*, *HSPA4*) that are involved in pathways of mitochondrial transportation or nanotube formation (Table S6). We excluded genes with direct association with oxygen-dependent metabolism. The tumor MT transfer (TMT) score for each cell or tumor sample was calculated by using gene set variation analysis⁴³ based on the mRNA expression of these 17 genes. The Wilcoxon paired signed-rank test was used to assess the statistical difference between TMT scores of tumoral and match normal samples in different cancer types. The cell cycle scores (CCS) and hypoxia scores of TCGA cancer samples were obtained from Lundberg et al.⁴⁷ and Ye et al.⁴⁹ Partial Spearman's rank correlation coefficient was used to assess the correlation of TMT scores with CCS and hypoxia scores in TCGA cancer types by adjusting for tumor purity.

QUANTIFICATION AND STATISTICAL ANALYSIS

All sequencing data were analyzed using R v4.0.2 and Python v3.8. The statistical analyses in this work, including two-sided Wilcoxon rank-sum or signed-rank, chi-squared, odds ratio, correlation tests and downsampling analysis were implemented with the R programming language. All P values from multiple tests were corrected using the Benjamini-Hochberg method and *p.adjust* function. Survival analyses were conducted with Kaplan-Meier and Cox proportional hazards regression methods using R package *survival* v3.3.1. Cox regression was adjusted for age, race, gender (excluding cancer types specific to one gender), tumor purity, molecular subtypes (for BRCA), clinical stage (for CESC and HNSC), histologic diagnosis (for CESC, HNSC, LGG, LIHC and PAAD), tobacco smoking history (for CESC, HNSC, LUAD and PAAD), tumor grade (for HNSC, LGG, LIHC and PAAD) and clinical TNM stage subgroups (for HNSC) as clinical covariates. Survival curves were compared across groups with log-rank test. The other method details (such as sequencing data processing and gene set enrichment analysis etc.) and the related software or packages could be found in [method details](#).