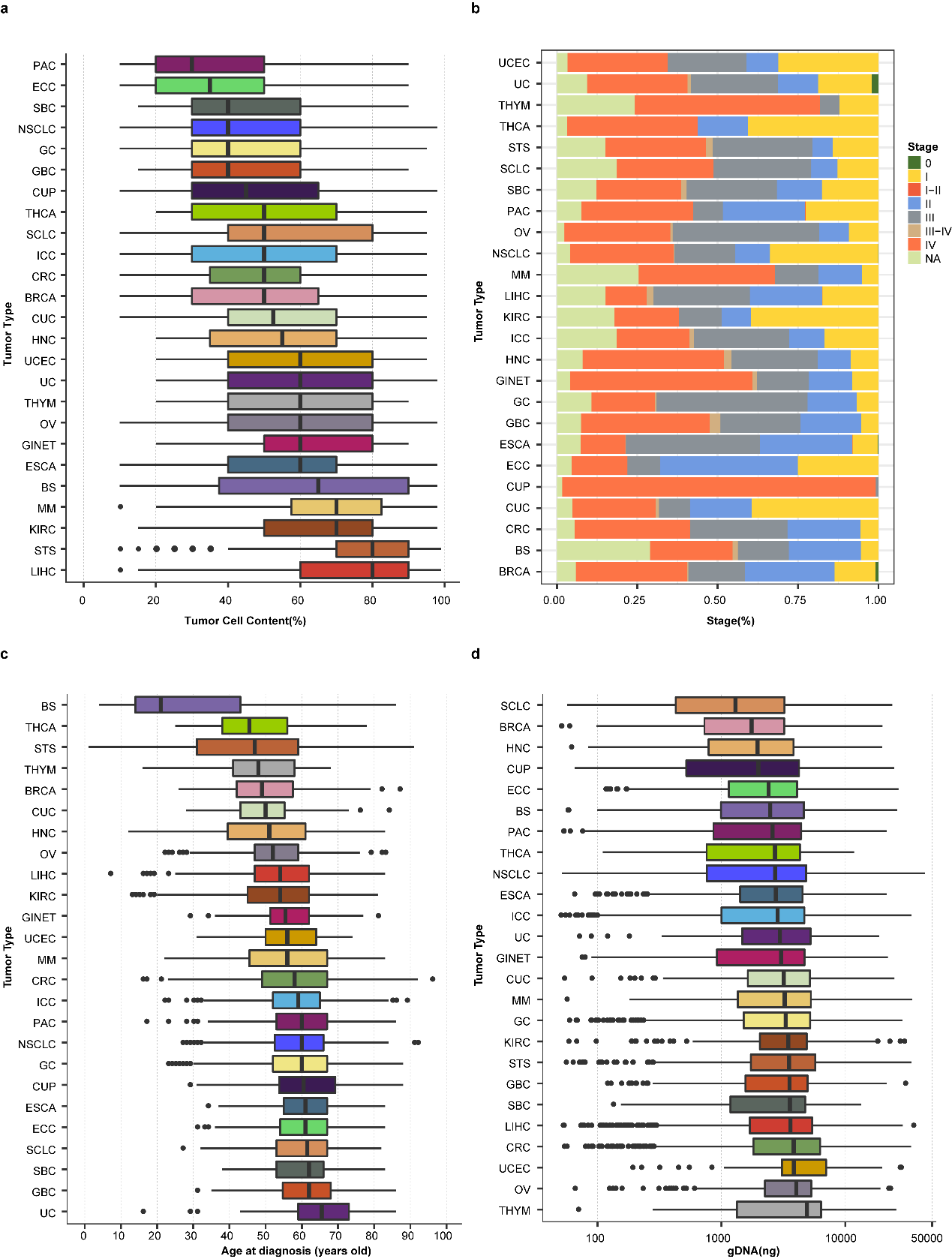
**Supplementary Figures**

Chart, box and whisker chart

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**Supplementary Fig. 1:** (**a**) **Selection processing of OM samples and success rates for analysis.** A total of 11,553 tumor samples were submitted for NGS based cancer assay in the OrigiMed lab. A total of 1,359 samples were excluded in several quality control steps. Finally, 10,194 cases were successfully involved in final analysis with a success rate of 88%. (**b**) **Variations per sample for different sampling methods, treatments and geographic origins.**



**Supplementary Fig. 2: The tumor type-specific distribution of (a) tumor cell purity, (b) stage of disease, (c) age at diagnosis, and (d) genomic DNA (gDNA)** **content extracted from samples.** The median (range) data are shown.

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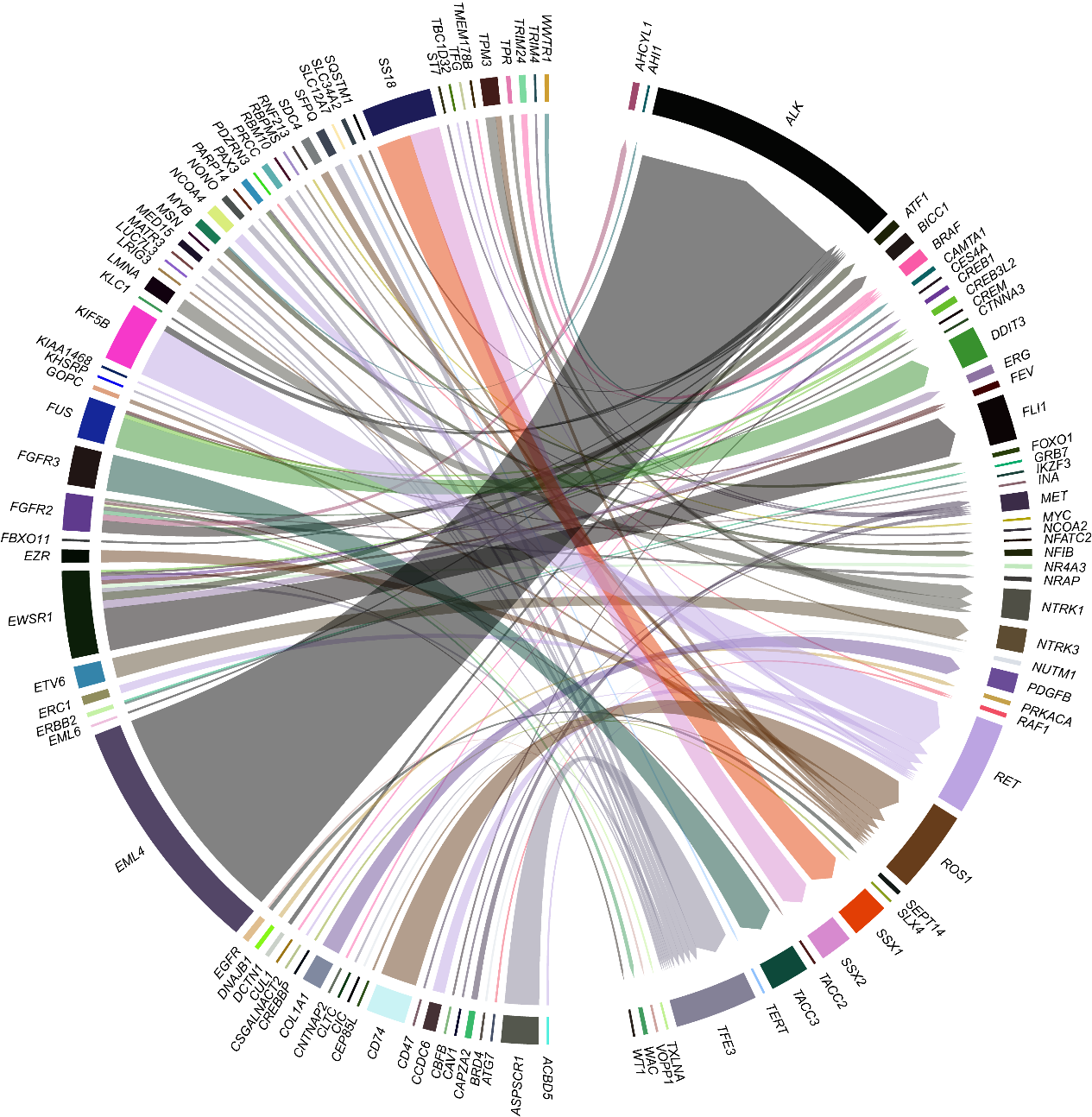
**Supplementary Fig. 3: Distribution of variant allele frequency (VAF) and mean unique coverage.** Distribution of VAF (≥ 2%) for mutations detected and reported in the OM dataset is shown above. Distribution of mean unique sequence coverage of 10,194 samples successfully sequenced is shown below.

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**Supplementary Fig. 4: Location of *EGFR*, *ERBB2*, *PIK3CA* and *KRAS* gene mutations which were concomitant with amplification of corresponding genes**.

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**Supplementary Fig. 5: Distribution of CNVs in primary/ metastatic tumors and across tumor types.** (**a**) Pan-cancer chromosome distribution of CNVs in primary and metastatic tumors. (**b**) The tumor type-specific distribution of recurrent CNVs. Red represents gene amplification and blue represents deletion. Frequencies are displayed in color gradient.

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**Supplementary Fig. 6: Spectrum of known fusions identified in OM cohort.** Known recurrent gene fusion driver-partner relationships across tumor types are profiled. A total of 94 known relationships reported previously were detected spanning 115 genes. The thickness of the line between two genes implies the relative count.

Chart, scatter chart

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**Supplementary Fig. 7: Fusion subtypes between *ALK* and *KIF5B* or *EML4*.** The middle line represents the fusing exons of *ALK* gene; fusing exons of *KIF5B* and EML4 are displayed at the left and right side, respectively. The exact fusion subtype relationships can be read according to the corresponding exons from *ALK* and *KIF5B* or *EML4*. Blue dots represent known fusions detected in both the Quiver dataset (http://quiver.archerdx.com/) and the OM dataset; red dots represent novel fusions only detected in the OM dataset; green dots represent fusions only found in the Quiver dataset.

**W:\综合项目数据\引力项目\review_comments\202003for check\figures\CRCmsi_group and smoke-4.tif Supplementary Fig. 8: (a) Distribution of hypermutated subtypes of CRC and (b) correlation between gender and smoke feature in lung cancers.** The median initial diagnosis age (58) of all CRC patients is used to separate into younger and older patients. *POLE*-associated CRC is defined as microsatellite stable (MSS) tumors with high mutation burden and at least one inactive POLE mutation. The “smoker” includes current smokers and former smokers, while “nonsmoker” means never smoker.

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**Supplementary Fig. 9: Comparative analysis of median TMB across various tumor types between the aOM cohort and MSK cohort.** The median value of TMB of each tumor type (aOM: MSK) is shown. TMB values that were statistically significantly (by Wilcoxon-test, *P* < 0.01) different between the aOM cohort (x-axis) and the MSK cohort (y-axis) are labeled.

Chart, bubble chart

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**Supplementary Fig. 10: (a)** **Tumor type-specific distribution of 891 samples which were at least one of the following: MSI-H, TMB-H, PD-L1 positive, and (b) the association between PD-L1 positive and MSI-H and TMB-H in ECC, GBC, BS, OV, SCLC and BRCA.**

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**Supplementary Fig. 11:** **Significance of *CD274* amplification in Chinese patients with solid tumors**. **(a)** The tumor type-specific distribution of tumors with *CD274* amplification is presented as the total or stage-specific percent of patients (%) across tumor types; stages are shown in different colors. **(b)** Expression of PD-L1 protein in tumors with *CD274* amplification. Thirty of 85 tumors with *CD274* amplification were tested for PD-L1 expression by IHC staining assay with Abcam 28-8 antibody. PD-L1 strong positive is defined as TPS > 50%. Among 30 tumors with tested PD-L1 protein, 70% showed positive expression. **(c)** The variation spectrum of 85 tumors with *CD274* amplification was profiled. The top 20 mutated genes are displayed in descending order. The upper panel represents the TMB value and the right panel indicates the frequencies of mutated genes.

Chart

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**Supplementary Fig. 12: (a)** **Number of patients with clinical actionable variants** **and (b) the distribution of TMB-H and PD-L1 positive in patients without clinical actionable variants**. The exact number of patients with clinical actionable variants according to four variant subtypes including substitution/indel, amplification, fusion/rearrangement, and truncation.

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**Supplementary Fig. 13:** **Overview of the NGS based cancer assay (CSYS) workflow.**