Patients with high-grade serous ovarian cancer suffer poor prognosis and variable response to treatment. Known prognostic factors for this disease include homologous recombination deficiency status, age, pathological stage and residual disease status after debulking surgery. Recent work has highlighted important prognostic information captured in computed tomography and histopathological specimens, which can be exploited through machine learning. However, little is known about the capacity of combining features from these disparate sources to improve prediction of treatment response. Here, we assembled a multimodal dataset of 444 patients with primarily late-stage high-grade serous ovarian cancer and discovered quantitative features, such as tumor nuclear size on staining with hematoxylin and eosin and omental texture on contrast-enhanced computed tomography, associated with prognosis. We found that these features contributed complementary prognostic information relative to one another and clinicogenomic features. By fusing histopathological, radiologic and clinicogenomic machine-learning models, we demonstrate a promising path toward improved risk stratification of patients with cancer through multimodal data integration.

High-grade serous ovarian cancer (HGSOC) is the most common cause of death from gynecologic malignancies, with a 5-year survival rate of less than 30% for metastatic disease. Initial clinical management relies on either primary debulking surgery (PDS) or neoadjuvant chemotherapy followed by interval debulking surgery (NACT-IDS). Endogenous mutational processes are an established determinant of clinical course, with improved response of homologous recombination-deficient (HRD) disease to platinum-based chemotherapy and poly-ADP ribose polymerase (PARP) inhibitors. More nuanced genomic analyses integrating point mutation and structural variation patterns further refine this stratification into four biologically and prognostically meaningful subtypes including distinct subgroups of HRD, foldback inversion-enriched tumors and those with distinctive accrual of large tandem duplications. Beyond genomic factors, clinical indicators such as patient age, pathological stage and residual disease (RD) status after debulking surgery are also prognostic. However, these clinicogenic factors alone fail to adequately account for the heterogeneity of clinical outcomes. Identifying patients at risk of poor response to standard treatment remains a critical unmet need. Improved risk stratification models would aid gynecologic oncologists in selecting primary treatment, planning surveillance frequency, making decisions about maintenance therapy and counseling patients about clinical trials of investigative agents.

Beyond clinicogenomic features, multiscale clinical imaging is routinely acquired during the course of care, including contrast-enhanced computed tomography (CE-CT) at the mesoscopic scale and hematoxylin and eosin (H&E)-stained slides at the microscopic scale. Digital forms of these diagnostics present opportunities to develop computational models and test whether integrating these data modalities improves identification of risk groups for HGSOC. At the mesoscopic scale, recent radiologic studies have uncovered quantitative CE-CT features that are predictive of early progression, time to recurrence and overall survival in HGSOC. Most studies to date have analyzed the prognostic information captured within adnexal lesions or the whole
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Interpretable features. Apart from stage, HGSOC lacks independent pre-treatment pathological factors by which to stratify patients (Fig. 1c). The models were validated on a pathological model based on pre-treatment tissue samples to obtain during the routine diagnostic workup of patients with ovarian cancer. A quantitative histopathological study of HGSOC involving machine-learning models to improve risk stratification of patients overall survival. Created with BioRender.com. GLSZM-SAE, gray level size zone matrix small area emphasis; GLRLM-GLV, gray level run length matrix gray level variance; Var, variance; Nuc, nuclear; NGS, next-generation sequencing; LSTs, large-scale state transitions; NtAI, number of subchromosomal regions with allelic imbalance extending to the telomere; LOH, loss of heterozygosity.

At the microscopic scale, H&E-stained tissue biopsies enable pathological diagnosis and are routinely acquired before the start of therapy. A quantitative histopathological study of HGSOC identified patterns of immune infiltration on H&E slides that correlate with mutational subtypes. In other cancer types, studies of whole-slide images (WSIs) have advanced our ability to quantify the histopathological architecture of tumors using deep learning and interpretable features. Apart from stage, HGSOC lacks independent pre-treatment pathological factors by which to stratify patients and quantitative approaches thus present an opportunity to systematically develop scaled models that are beyond qualitative human interpretation. Interpretable features are less prone to overfitting in small cohorts and can be more easily interrogated by human pathologists.

Conceptually, genomic sequencing does not account for spatial context and we thus hypothesize that multiscale imaging contains complementary information, rather than merely recapitulating genomic prognostication. We are further motivated by the potential for clinical multimodal machine learning to outperform unimodal systems by combining information from multiple routine data sources. In this work, we set out to study the complementary prognostic information of multimodal features derived from clinical, genomic, histopathological and radiologic data obtained during the routine diagnostic workup of patients with HGSOC (Fig. 1a). We tested the prognostic relevance of ovarian and omental radiomic features derived from CE-CT and developed a model based on omental features (Fig. 1b) and a histopathological model based on pre-treatment tissue samples to risk stratify patients (Fig. 1c). The models were validated on a test cohort and integrated with clinical and genomic information (Fig. 1d) using a late-fusion multimodal statistical framework (Fig. 1e). Our results revealed the empirical advantages of cross-modal integration and demonstrated the ability of multimodal machine-learning models to improve risk stratification of patients with HGSOC.

Results

Cohort and clinical characteristics. We analyzed 444 patients with HGSOC, including 296 patients treated at the Memorial Sloan Kettering Cancer Center (MSKCC) and 148 patients from The Cancer Genome Atlas Ovarian Cancer (TCGA-OV) data. The 40 test cases were randomly sampled from the entire pool of patients with all data modalities available for analysis; data from the remaining 404 patients were used for training. The training set contained 160 patients with stage IV disease, 225 with stage III, 10 with stage II, 8 with stage I and 1 with an unknown stage (Supplementary Table 1). The test cohort contained 31 patients with stage IV and 9 with stage III disease. Median age at diagnosis was 63 (interquartile range (IQR) 55–71) years for the training set and 66 (IQR 59–70) years for the test set. In the training cohort, 175 patients received NACT-IDS and the remaining 82 underwent PDS. In the test cohort, 31 received NACT-IDS and 8 underwent PDS. Overall, 61 patients from MSKCC were known to have received PARP inhibitors (Supplementary Table 1). Treatment regimens are not annotated for 148 TCGA patients. Median overall survival (OS) was 36.3 (IQR 25–55) months for training patients and 37.6 (IQR 26–49) months for testing patients. There were 132 training patients and 17 testing patients with censored OS outcomes (Supplementary Table 2).

Among 404 patients in the training cohort, 243 patients had H&E WSIs, 245 patients had adnexal lesions on pre-treatment CE-CT and 251 patients had omental implants on pre-treatment CE-CT (Fig. 2a). All 40 patients in the test cohort had omental lesions on CE-CT, H&E WSIs and available sequencing by construction; 29 patients had ovarian lesions on CE-CT. Three gynecologic radiologists volumetrically segmented adnexal lesions and representative omental lesions on all sections containing these lesions (Extended Data Fig. 1a). The training and testing data were acquired with similar CT scanners (Extended Data Fig. 1b).
We used clinical sequencing24 to infer HRD status, in particular variants in genes associated with HRD DNA damage response (DDR)26,28 such as BRCA1 and BRCA2 and those specific to disjoint tandem duplicator- and foldback inversion-enriched mutational subtypes (CDK12 and CCNE1 (refs. 5,27), respectively; Figs. 1d and 2b,c). We also examined the genomes of 130 patients with appropriate consent for direct evidence of homologous recombination deficiency, namely COSMIC single-base substitution signature 3, which is associated with defective HRD-DDR. In this subset of MSKCC patients, signature 3 was detected by SigMA28 with high confidence in 48 cases, detected with low confidence in 30 cases and found not to be the dominant signature in 52 cases (Extended Data Fig. 2g,i).

In the TCGA, signature 3 was high in 6 cases and low in 51 cases (Extended Data Fig. 2b). In the TCGA, signature 3 was high in 6 cases and low in 51 cases (Extended Data Fig. 2b). Patients with available sequencing and without evidence for HRD or homologous recombination proficiency (HRP; \(n = 126\)) were treated as HRP. Patients with conflicting evidence (\(n = 6\)) or without sequencing (\(n = 61\)) were assigned a label of ‘ambiguous’ and excluded from all analyses involving HRD status. In total, the training cohort contained 218 HRP and 119 HRD cases (Fig. 2c). The test set contained 12 HRD and 28 HRP cases. HRD status alone (excluding ambiguous) stratified patients by OS with a c-Index of 0.55 in the training cohort and 0.52 in the test set (without fitting any model parameters; Extended Data Fig. 2d,e). Aberrations specific to distinct endogenous mutational processes also stratified patients as expected: patients with HRP disease had worse outcomes than those with HRD disease (\(P = 7 \times 10^{-3}\); Extended Data Fig. 2g,i).

**CE-CT imaging feature selection and stratification.** We began by studying the prognostic relevance of features derived from radiology scans either obtained at our institution (91; 27%) using GE Medical Systems CT scanners or acquired at outside institutions (247; 73%) from a variety of CT scanners (Extended Data Fig. 1 and Supplementary Table 3). The majority of CE-CT scans were acquired with a peak kilovoltage of 120 (median 120 kVp, range: 90–140; Supplementary Table 3) and reconstructed with the standard convolutional kernel using 5-mm slice thickness (median 5 mm; range 2.5–7.5; Supplementary Table 3). Three fellowship-trained radiologists with expertise in gynecologic oncologic imaging manually segmented all adnexal masses and representative omental implants on each pre-treatment CE-CT scan (Figs. 1b and 3a).

We extracted radiomic features from Coif wavelet-transformed images, yielding a 444-dimensional radiomic vector per site per patient after filtering by interquartile range. Using the training cohort, we calculated the hazard ratios (HRs) and prognostic significance of omental and ovarian radiomic features using univariate Cox proportional hazards models (Supplementary Table 4). After correction for multiple hypothesis testing, nine omental features (Fig. 3b) and none of the ovarian features exhibited statistically significant HRs (Fig. 3c). Hence, going forward, we only considered the omental implants. We iteratively fitted and pruned Cox models for multivariable significance on the nine omental features (Algorithm 1), yielding a univariate model based on the autocorrelation of the gray level co-occurrence matrix derived from the high–low–low (HLL) Coif wavelet-transformed images (Fig. 3d). This feature exhibited a log(HR) of 1.68 (corrected \(P < 0.01\); Fig. 3e) and was invariant to CT scanner manufacturers and segmenting radiologists (Extended Data Fig. 3). The model stratified patients in the training and the test sets with concordance indices of 0.55 (95% CI 0.549–0.554) and 0.53 (95% CI 0.517–0.547), respectively.
Fig. 3 | High-autocorrelation omental implants are associated with shorter OS. a, Segmented omental lesion (red) on CE-CT. b, The log HR is depicted for each radiomic feature derived from omental implants (n = 600 features). Features above the line were statistically significant by Cox regression after multiple testing correction of interquartile range-filtered features. c, Adnexal radiomic features (n = 600 features) were not significant by Cox regression after correction of interquartile range-filtered features. d, The hazard ratio with 95% CI as estimated by Cox regression is shown for the feature in the final model, the autocorrelation derived from the gray level co-occurrence matrix for the wavelet-filtered image. e, The value of this feature against OS is plotted for patients in the training set (n = 251 patients). f, Training and test concordance indices for the model are shown; the height of each bar shows the c-Index and the lower and upper points of the respective error bars depict the 95% CI by 100-fold leave-one-out bootstrapping. g, h, Two risk groups based on the model’s predicted risk score are shown for the training and test sets. P values were derived using the log-rank test. glcm, gray level co-occurrence matrix; gldm, gray level dependence matrix; glrlm, gray level run length matrix; glszm, gray level size zone matrix; ngtdm, neighboring gray tone difference matrix.

Histopathological tissue-type classifier for interpretable features. We next trained a tissue-type classifier from histology images using a weakly supervised approach. We annotated tissue types on 60 H&E WSIs, yielding more than 1.4 million partially overlapping tiles, each measuring 128×128 pixels (64×64µm) and containing 4,096µm² of tissue (Fig. 4a). A ResNet-18 convolutional neural network pretrained on ImageNet (Fig. 4b) classified tissue types with an accuracy of 0.88 (range 0.77–0.95) on pathologist-annotated areas labeled as fat, stroma, necrosis and tumor (Fig. 4c) by fourfold slide-wise cross-validation. Notably, the model correctly identified small regions of fat within stromal annotations and necrotic regions within the tumor, supporting the suitability of weakly supervised deep learning for this task and refining annotations into more granular classifications.

The cross-validation confusion matrix aggregated across folds showed good performance overall (Fig. 4d), with the most significant confusion being necrotic tiles predicted to be tumor and stroma. However, one disadvantage of weakly supervised learning is that neither the training data nor the validation data are exactly labeled. Hence, the cross-validation metrics are not computed against the exact truth. Visual inspection of the predictions were...
qualitatively concordant with only moderate confusion of necrosis with tumor and stroma (Extended Data Fig. 4).

**Histopathological stratification.** We applied the tissue-type classifier to the 243 training H&E WSIs of lesions from pre-treatment specimens (Fig. 1c). We combined these inferred tissue-type maps with detected cellular nuclei, yielding labeled nuclei (Fig. 5a). Subsequently, we extracted cell-type features from these nuclei and tissue-type features from the tissue-type maps based on the methods of Diao et al.\textsuperscript{20}. This yielded a histopathological vector of 216 features. We next identified the HRs of features using univariate Cox models fitted on slides in the training cohort. Several tissue-type features, such as overall tumoral area, were partially determined by specimen sizes and we thus controlled for this during selection. Of the 24 features with a log(HR) found to be significantly different from 0 with 95% confidence, 20 related to tumor nuclear diameter or size, with larger being associated with shorter OS (Extended Data Fig. 5 and Supplementary Table 5). We again iteratively fitted and pruned Cox models as per Algorithm 1, yielding a multivariable model with two features: the mean tumor nuclear area and the major axis length of the stroma (Fig. 5b). This histopathological signature was not confounded by specimen size (Extended Data Fig. 6). This model stratified the training and test sets, with concordance indices of 0.56 (95% CI 0.559–0.564) and 0.54 (95% CI 0.527–0.560), respectively (Fig. 5d; \( P < 0.01 \)). For the test set, the risk groups trended toward (but did not attain) significantly different separation, with median survival of 37 and 50 months (Fig. 5e; \( P = 0.076 \)).

To probe the interpretability of the histopathological features, we investigated the mean tumor nuclear area; we show examples of low (Fig. 5f) and high (Fig. 5g) values, which were associated with better and worse prognosis, respectively.

**Multimodal prognostication.** We tested the prognostic significance of patient age, pathological stage, RD status after debulking surgery, NACT-IDS versus PDS treatment paradigm, receipt of PARP inhibitors in the first 2 years after diagnosis and the presence or absence of adnexal lesions (Supplementary Table 6), ultimately training a model on RD status and PARP inhibitor administration. This model stratified the test set with Harrell’s concordance index, \( c = 0.51 \) (95% CI 0.493–0.528). We then implemented a late-fusion\textsuperscript{8} approach to integrate histopathological, radiomic, genomic and clinical data into multimodal models (Fig. 1e). Specifically, we predicted each patient’s log partial hazard using the Cox model trained using the respective modality, then trained a final Cox model to integrate them (Methods). In the test set, the model combining both imaging modalities (radiomic–histopathological (RH) model) significantly outperformed the HRD status-based model, clinical model and individual imaging models, with a test concordance index of 0.62 (95% CI 0.604–0.638) (Fig. 6a). The model with genomic, radiomic and histopathological (GRH) modalities performed comparably, with a test concordance index of 0.61 (95% CI 0.594–0.625).
The histopathological submodel score remained significant upon addition of HRD status (Fig. 6b). The high- and low-risk groups established by the GRH model were significantly different by log-rank test in the training set (median survival of 34 and 50 months, respectively; \( P = 0.026 \); Fig. 6c). In the test set, the GRH risk groups also showed significantly different OS, with median survival of 30 months for the high-risk group and 50 months for the low-risk group (\( P = 0.023 \); Fig. 6d). At 36 months, 68% and 34% survived for low- and high-risk groups, respectively, in the test set. The separation of the RH model’s risk groups was inferior (Extended Data Fig. 7). Notably, analysis of only training cases with full information (\( n = 114 \)) resulted in poor performance (Extended Data Fig. 8), reinforcing the ability of late-fusion models to learn in the setting of missing data. No robust association was found between modalities to enable interpolation of missing values (Extended Data Fig. 9).

The c-Indices for individual imaging modalities were similar, but identified distinct patient subgroups with good prognosis (Fig. 6e). This is consistent with radiological and histological features containing complementary information content, whereby some patients with good outcomes were identified as high risk by the radiomic submodel but correctly assigned a lower risk score by the histopathological submodel and vice versa. Patients with HRD and HRP disease were distributed relatively evenly, agnostic to unimodal imaging risk scores.

Corroborating this, absolute Kendall rank correlation coefficient values were low between individual modalities (\(< 0.14 \); Fig. 6f), demonstrating that the radiomic and histopathological models ordered patients differently as compared to the genomic model and to one another. The same two risk groups identified by the model in the test set also showed significantly different progression-free survival (PFS) (\( P = 0.040 \); Fig. 6g). Finally, as an orthogonal validation, the inferred risk of all models except the genomic and genomic–histopathological models associated with pathological chemotheraphy response score (CRS) in the training set (Fig. 6h). The test set had only 21 patients with known CRS and only HRD status exhibited statistically significantly different
Discussion

Machine learning in cancer prognostics is a growing field with great potential, but the contribution of common diagnostic modalities to multimodal risk stratification remains poorly understood. Here, we show that integrating multiscale clinical imaging and genomic data increases predictive capacity. These results, in addition to the low correlation between risk scores derived from individual modalities, support the hypothesis that clinical imaging contains complementary prognostic information that is independent of clinicogenomic information. Histopathological and radiological imaging characterize the tumor architecture at microscopic and...
mesoscopic scales, respectively. Therefore, it stands to reason that these data channels complement one another and HRD status, which is derived from spatially agnostic sequencing. The full combined genomic, histopathological, radiological and clinical (GHRC) model did not perform as well as the RH and GRH models, suggesting that multimodality is not a universal guarantee of improved performance\(^1\). In this case, the most likely reason is that the clinical model (based on history of PARP inhibitor administration and RD status after debulking surgery) does not stratify the test cohort, likely due to its small size. Furthermore, the TCGA cohort did not have these informative clinical variables available. Our late-fusion architecture benefits from few parameters to fit—which reduces overfitting\(^4\)—and the ability to learn from partial information cases, but it cannot gate information from noisy modalities. With larger datasets enabling more parameter fitting without overfitting, mechanisms such as attention can be explored to adaptively adjust unimodal contributions.

In addition to multimodal integration, we presented two unimodal models to stratify patients with late-stage HGSOC using routine clinical imaging, validated these models on a test set and studied the relative contributions of each modality to risk stratifying patients with HGSOC. For radiological imaging, we discovered that omental autocorrelation computed from the gray level co-occurrence matrix derived from the HIL Coif wavelet-filtered image was a prognostic feature. This Imaging Biomarker Standardization Initiative-defined feature\(^31,34\) has been found to be strongly or very strongly reproducible in multiple studies\(^25\). It describes the coarseness of the lesion texture and also depends on tissue density. Seven of the other nine omental features with significant log(HR) values were explicitly designed to measure high-density zones and these features did not exhibit log(HR) values significantly different from zero on multivariable regression with the autocorrelation. Hence, the most parsimonious explanation is that higher-density—rather than coarser—omentum implants are an adverse prognostic factor, which could be due to more solid tumors with reduced cystic or fatty components. Omental textures captured by autocorrelation may also reflect differing intratumoral heterogeneity.

To our knowledge, previous HGSOC radiomic models have not explored the prognostic information captured within omental implants, relying instead on more demanding segmentations of adnexal lesions or the entire tumor burden. Notably, we found that none of the radiomic features derived from adnexal masses had log(HR) values significantly different from zero after correction for multiple hypothesis testing, which is possibly due to the late stage of this cohort: the omentum is the most common site of metastasis in HGSOC\(^33\) and may drive further peritoneal seeding. An omental model is advantageous over an adnexal model because omental implants are ubiquitous in advanced-stage disease, even in patients with primary peritoneal high-grade serous cancer that lack adnexal mass(es). Furthermore, an omental implant can be readily segmented even by less experienced observers, whereas adnexal masses can be challenging to distinguish from adjacent loculated ascites, serosal and pouch of Douglas implants and adjacent anatomic structures such as the uterus, especially in the presence of leiomyomas. An omental model is also more practical than a radiomic model based on the whole tumor burden; routine segmentation of the whole tumor volume is impractical in daily practice using current tools due to prohibitively high demand for time and expertise.

For histopathological imaging, we developed an H&E WSI-based model to stratify patients with HGSOC. Although none of the features exhibited log(HR) values significantly different from zero after correction for multiple hypothesis testing, the presence of 20 features highly related to mean tumor nuclear size (such as 60th percentile of tumor nuclear size and 50th percentile of tumor nuclear diameter) with similar HRs in the 24 features with uncorrected significant P values for univariate log(HR) values supports the prognostic relevance of tumor nuclear size. This is further supported by the good stratification of the test set. The larger nuclear size may be associated with events such as whole-genome doubling or cellular fusion and warrants direct study of matched genomes and histopathological sections. The major axis length of stroma is difficult to interpret for a two-dimensional slice of tissue but may reflect distinct patterns of disease infiltration into surrounding stroma. We included the trained weights for our HGSOC model and the source code for extension to other cancer types.

This lack of usable large datasets is one of the main challenges for multimodal machine learning in oncology\(^7\). We have made data from the 296 MSKCC patients with HGSOC available to enable future work toward improving upon the models presented here. Our results demonstrate the benefit of learning from cases with only partial information in multimodal studies: the smaller, full-information subcohort yielded a significantly less-generalizable risk stratification model. Our dataset also offers the advantage of comprising H&E images and CE-CT scans originally acquired at multiple institutions: this improves confidence in the generalizability of the results. Furthermore, we intentionally mined data generated during the standard of care. Using these data instead of specialty research data drastically reduces adoption costs in the clinical workflow for resultant models, but the data were not collected specifically with computational modeling in mind. For example, we included some patients with only germline sequencing of HRD-DDR genes, a clinically relevant but biologically imperfect measure of HRD status: each risk group is enriched for—but not exclusively composed of—the genomic subtype of interest. We expect that clinical whole-genome sequencing will enable more robust genomic analyses.

The improved risk stratification models developed herein show the promise of extracting and integrating quantitative clinical imaging features toward aiding gynecological oncologists in selecting primary treatment, planning surveillance frequency, making decisions about maintenance therapy and counseling patients about clinical trials of investigative agents. The statistical robustness and clinical relevance of the risk groups by both PFS and OS in the test set substantiate the utility of this multimodal machine-learning approach, establishing a proof of principle. Next steps along this line of work include scaled and inter-institutional retrospective cohort assembly for further model training and refinement before prospective validation of clinical benefit in randomized controlled trials\(^6\).

In summary, we have assembled a multimodal dataset of patients with HGSOC and used this to develop and integrate radiological, histopathological and clinicogenomic models to risk stratify patients. We discovered that the autocorrelation of omental implants on CE-CT and average tumor nuclear size on H&E are prognostic factors, that these modalities are demonstrably orthogonal and that their computational integration improves stratification beyond previously known clinicogenomic factors in a test set. Our results motivate further large-scale studies driven by multimodal machine learning to stratify patients with cancer, both in HGSOC and other cancer subtypes.

### Methods

This study complies with all relevant ethical regulations and its protocols were approved by MSKCC’s Institutional Review Board. Informed consent was waived for this retrospective study and participants were not compensated. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**MSKCC cohort curation.** Patients were eligible for this retrospective study if they had biopsy-proven newly diagnosed HGSOC and at least one of (1) pre-treatment WSIs of H&E depicting high-grade serous carcinoma or (2) pre-treatment contrast-enhanced abdominal/pelvic computed tomography (CE-CT). Most of the MSKCC cohort was sourced from a retrospective clinical database of patients who underwent diagnostic workup and NACT-IDS at our institution. This database also contained information on the RD status after debulking surgery,
pathological stage, administration of neoadjuvant chemotherapy and patient age at diagnosis from the electronic medical record. To expand the cohort, we also searched the institutional data warehouse for patients with MSK-IMPACT sequence data available pre-treatment CE-CT images. In addition to this retrospective curation, 36 patients were also included from the prospective MSK-SPECTRUM project. Pathological stage was unavailable for 14 patients and we instead recorded the clinical stage as recorded in the institutional database for these patients. We also collected the race for all patients from the institutional data warehouse. OS and PFS were calculated using the date of CT as a start date, when available, or the date of pathological diagnosis otherwise.

To collect H&E imaging, we reviewed the electronic health record to find associated pathology cases with peritoneal lesions (primarily omental) and expert pathologists reviewed the slides to select high-quality specimens for digitization. We also reviewed the institutional data repository for scanned slides associated with the diagnosis that included those containing tumors. All H&E imaging was carried out before treatment.

We subsequently reviewed the associated CE-CT scans for the following the inclusion criteria: (1) intravenous contrast-enhanced images acquired in the portal venous phase, (2) absence of streak artifacts or motion-related image blur obscuring lesion(s) of interest and appropriate signal to noise ratio (Supplementary Table 7). All CE-CT imaging was carried out before treatment. All CT scans were available in the digital imaging and communications in medicine (DICOM) format through our institutional picture archiving and communication system (PACS, Centricity, GE Medical Systems v.7.0).

TCGA cohort selection. From the TCGA-OV project, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35, we selected patients with clinical sequencing data available from the TCGA-OV project35.

In the MSKCC cohort, we used MSK-IMPACT clinical sequencing37, when available, to infer HRD status. Variant calling for these genes and copy number analysis of CNET was performed using the standard MSK-IMPACT clinical pipeline (https://github.com/mskcc/Innovation-IMPACT-Pipeline). For patients with appropriate consent for further genomic re-analysis, we also inferred COSMIC SBS3 activity using SigMA (for cases with at least five mutations across all 505 genes)38 and searched for large-scale state transitions using our own pipeline (https://github.com/jrflab/modules)/39. We used OncoKB to assemble tile predictions into downscaled bitmaps, which were then used for training and omitted vessels from the training data for the models presented in this work. We next used these annotations to generate tissue-type tiles.

Training the histopathological tissue-type classifier. We generated tiles measuring 64×64 (128×128 pixels) with 50% overlap, using the above annotations to delineate regions to be tiled. No other tile sizes were explored; this size was chosen because it offered good resolution while still depicting multiple cells in each tile. Putative tile squares within an annotation but with <20% foreground as assessed by Otus's method were not tiled. Macenko stain normalization was used. We trained a ResNet-18 model (pretrained on ImageNet) for 30 epochs with a learning rate of 5×10−4, 1×10−4 L2 regularization and the Adam optimizer. The objective function was class-balanced cross entropy and we used mini batches of 96 tiles on a single NVIDIA Tesla V100 GPU. We used fourfold, slide-wise cross-validation for model evaluation and hyperparameter tuning. We selected the number of epochs to train the final model using the epoch with the highest lower 95% CI bound estimated using the mean and s.d. of the cross-validation F1 scores. We trained the model on tiles from all 60 slides for 21 epochs.

Histopathological feature extraction and selection. We tiled the WSIs associated with the patients in this cohort without overlap, performing inference using mini batches of 800 across four NVIDIA Tesla V100 GPUs. We used Macenko stain normalization as a result of downsampling, we were then used to calculate tissue-type features in an approach based on previous work40. We included the region properties from scikit-image41 for both the largest connected component and the entirety of each tissue type. We also calculated features such as the area ratio of one tissue type to another and the entropy of tumor and stroma. Using the StarDist method42 for QuPath, we segmented and characterized individual nuclei, using nuclei with a detection probability greater than 0.5. We used a lymphocyte classifier trained iteratively using manual annotations to distinguish lymphocytes from other cells. We assigned a tissue parent type to each CE-CT image feature, classifying features into tissue types based on nearby predicted tissue-type annotations and calculating aggregative statistics by tissue type and cell type of the QuPath-extracted nuclear morphological and staining features, such as variance in eosin staining or circularity. Together, these cell type features and tissue-type features based on tumor, stroma and necrosis constituted the histopathological embedding for each slide.

Clinical data encoding. RD status after debulking surgery was encoded as a binary variable, where patients with ≤1 cm RD (including complete gross resection) were assigned a value of 1 and patients with >1 cm RD were assigned a value of 0. The presence of adnexal lesions on CE-CT was also included as a binary variable. Age at diagnosis was modeled as a continuous variable scaled by the training set range. Tumor stage was encoded as one-hot categorical variables for I, II, III, IV and unknown. Similarly, the primary treatment approach was encoded as a one-hot categorical variable with values NACT-IDS, PDS and unknown.

Feature selection. The same strategy was used to select radiomic, histopathological and clinical features. For each feature, we fitted a univariate Cox proportional model to collect H&E imaging, we reviewed the electronic health record to find associated pathology cases with peritoneal lesions (primarily omental) and expert pathologists reviewed the slides to select high-quality specimens for digitization. We also reviewed the institutional data repository for scanned slides associated with the diagnosis that included those containing tumors. All H&E imaging was carried out before treatment.

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TCGA cohort selection. From the TCGA-OV project, we selected patients with clinical data annotated in the TCGA Clinical Data Resource, pathological grade 3 and at least one of a diagnostic FFPE H&E WSIs or abdominal/pelvic CE-CT scan in the TCGA cohort selection. All clinical and demographic information were extracted from the TCGA CDR. Only diagnostic WSIs of formalin-fixed, paraffin-embedded H&E-stained specimens from the TCGA-OV project were included. All H&E imaging was carried out before treatment.

All CT scans were available in the digital imaging and communications in medicine (DICOM) format through our institutional picture archiving and communication system (PACS, Centricity, GE Medical Systems v.7.0).

In the MSKCC cohort, we used MSK-IMPACT clinical sequencing, when available, to infer HRD status. Variant calling for these genes and copy number analysis of CNET was performed using the standard MSK-IMPACT clinical pipeline (https://github.com/mskcc/Innovation-IMPACT-Pipeline). For patients with appropriate consent for further genomic re-analysis, we also inferred COSMIC SBS3 activity using SigMA (for cases with at least five mutations across all 505 genes) and searched for large-scale state transitions using our own pipeline (https://github.com/jrflab/modules). We used OncoKB to assemble tile predictions into downscaled bitmaps, which were then used for training and omitted vessels from the training data for the models presented in this work. We next used these annotations to generate tissue-type tiles.

Training the histopathological tissue-type classifier. We generated tiles measuring 64×64 (128×128 pixels) with 50% overlap, using the above annotations to delineate regions to be tiled. No other tile sizes were explored; this size was chosen because it offered good resolution while still depicting multiple cells in each tile. Putative tile squares within an annotation but with <20% foreground as assessed by Otus’s method were not tiled. Macenko stain normalization was used. We trained a ResNet-18 model (pretrained on ImageNet) for 30 epochs with a learning rate of 5×10−4, 1×10−4 L2 regularization and the Adam optimizer. The objective function was class-balanced cross entropy and we used mini batches of 96 tiles on a single NVIDIA Tesla V100 GPU. We used fourfold, slide-wise cross-validation for model evaluation and hyperparameter tuning. We selected the number of epochs to train the final model using the epoch with the highest lower 95% CI bound estimated using the mean and s.d. of the cross-validation F1 scores. We trained the model on tiles from all 60 slides for 21 epochs.

Histopathological feature extraction and selection. We tiled the WSIs associated with the patients in this cohort without overlap, performing inference using mini batches of 800 across four NVIDIA Tesla V100 GPUs. We used Macenko stain normalization as a result of downsampling, we were then used to calculate tissue-type features in an approach based on previous work. We included the region properties from scikit-image for both the largest connected component and the entirety of each tissue type. We also calculated features such as the area ratio of one tissue type to another and the entropy of tumor and stroma. Using the StarDist method for QuPath, we segmented and characterized individual nuclei, using nuclei with a detection probability greater than 0.5. We used a lymphocyte classifier trained iteratively using manual annotations to distinguish lymphocytes from other cells. We assigned a tissue parent type to each CE-CT image feature, classifying features into tissue types based on nearby predicted tissue-type annotations and calculating aggregative statistics by tissue type and cell type of the QuPath-extracted nuclear morphological and staining features, such as variance in eosin staining or circularity. Together, these cell type features and tissue-type features based on tumor, stroma and necrosis constituted the histopathological embedding for each slide.

Clinical data encoding. RD status after debulking surgery was encoded as a binary variable, where patients with ≤1 cm RD (including complete gross resection) were assigned a value of 1 and patients with >1 cm RD were assigned a value of 0. The presence of adnexal lesions on CE-CT was also included as a binary variable. Age at diagnosis was modeled as a continuous variable scaled by the training set range. Tumor stage was encoded as one-hot categorical variables for I, II, III, IV and unknown. Similarly, the primary treatment approach was encoded as a one-hot categorical variable with values NACT-IDS, PDS and unknown.

Feature selection. The same strategy was used to select radiomic, histopathological and clinical features. For each feature, we fitted a univariate Cox proportional model.
hazards model to the full training set using the Python Lifelines package without regularization and we plotted the univariate coefficient and significance confidence. For features whose model failed to converge, we re-attempted fitting with L2 regularization of 0.2 and any model still failing to converge was assigned a log(HR) of 0 and P value of 1. For histopathology, we controlled for relative specimen size by including it in each Cox model. We next removed features with scaled IQR below 0.1. Subsequently, for radiomics, which is the largest feature space, we used the Benjamini–Hochberg method to correct for multiple hypothesis testing 29. Taking the ordered list of features significant with 95% confidence, we next applied Algorithm 1 to select features, yielding modality signatures with low multicollinearity.

Algorithm 1. Multivariable model selection procedure. Input: A list of unique candidate features ordered by P value of the Cox model for each candidate feature. Output: A list of features significant with confidence α on a multivariable regression model g, where j ∈ [1, k] and i ≤ k. Require: k ≥ 1, i → 1 while i ≤ k do g ← P−significance(g) ≥ significance assessed by Cox regression if P < α then j ← i+1 end if i ← i+1 end while

The only modification to this procedure occurred for the ablation experiment to test the importance of learning from the partial information cases: we used a threshold of 0.31 for each model as none was significant with P < 0.05 and we did not correct for multiple hypothesis testing in the omental radiomic features during the ablation experiment as none would be significant by this metric.

Survival modeling. We used linear Cox proportional hazards models with L2 regularization (α = 0.5) and no L1 regularization for all multimodal and unimodal models. No substitution was fitted for the genomic modality; patients assigned to the HRP subtype were designated high risk (risk score 1.0) and patients assigned to the HRD subtype were designated low risk (risk score 0). No interaction terms were used. We used Kaplan–Meier analysis to determine whether each model stratified patients into clinically significant groups. To delineate group membership, we tested percentile thresholds in [0.33, 0.34, …, 0.64, 0.65, 0.66], choosing the value that maximized significance of the separation in the training set by the log-rank test. This was performed individually for OS and PFS, where relevant. P values for concordance indices were calculated using 1000-fold permutation tests. The 95% CI for c-Indices were calculated using 100-fold leave-one-out bootstrapping. All P values for Kaplan–Meier analysis were calculated by the multivariate log-rank test. P values for covariate significance in Cox proportional hazards models are reported for models fitted with α = 0.5. The fraction surviving was estimated using linear interpolation.

Multimodal integration. We chose a late-fusion approach to increase unimodal sample sizes available for parameter estimation 30. Parameters for unimodal submodels were estimated using all available unimodal data (radiomic parameters were estimated across the 251 training CT cases with omental lesions and histopathological parameters were estimated across the 243 training H&E cases), where each submodel inferred a partial hazard for each patient. The negative partial hazard was used to enable compatibility with the concordance index as implemented in the lifelines Python package 31. For the second-stage late-fusion model, we estimated parameters for a multivariate Cox model integrating the negative log partial hazards inferred by each modality using only the intersection set of patients.

Statistics and reproducibility. No statistical method was used to predetermine sample size. Data were excluded from the analyses only for the reasons detailed above and before any machine-learning modeling. The training and test sets were chosen at random from the patients with all four data modalities available. The investigators were not blinded to allocation during outcome assessment. Data distributions were not assumed to be normal for any tests. The hazards were assumed to be proportional for survival modeling, but this was not formally tested. Analysis was conducted in QuPath v.0.2.3 (with the StarDist extension), ITK SNAP v.3.8.0 and custom code written in Python v.3.9.4 (using Pandas v.1.2.4, NumPy v.1.20.2, PyTorch v.1.5.1, TorchVision v.0.8, OpenSlide v.1.1.1, Seaborn v.0.11.1, Matplotlib v.3.2.2, SciPy v.1.6.3, scikit-learn v.0.24.0, PyRadiomics v.3.0 and Lifelines v.2.25.2).

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

Data availability. DNA sequencing, H&E WSI and CT data that support the findings of this study have been deposited at Synapse (Sage Bionetworks) under accession code syn25946117. Additional H&E WSI, CT imaging and genomic data were derived from the TCGA Research Network: https://cancergenome.nih.gov/ and The Cancer Imaging Archive: https://www.cancerimagingarchive.net/. Raw data from MSK-IMPACT performed in the CLIA laboratory in the Department of Pathology is not currently permitted in public repositories because ethical and legal implications are still being discussed at an institutional level; thus, the derivative features related to HRD status are shared in the repository. Source data have been provided as Source Data files. All other data supporting the findings of this study are available from the corresponding author on reasonable request. Source data are provided with this paper.

Code availability. All code necessary for the analyses is shared without access restrictions in a public repository (https://github.com/kmboehm/onco-fusion).
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Author contributions

K.M.B. reviewed the healthcare record for cases, wrote the manuscript and conducted the radiological, histopathological and clinical data analysis and modeling. E.A.A., J.S.K.-F. controlled and segmented CT studies and reviewed the healthcare record for pathological cases. L.E. annotated CRNs, reviewed slides for digitization and annotated tissue types on H&E slides. I.N. quality controlled and segmented CT studies. M.A. annotated the chemotherpay response score for histopathological specimens. I.V.G. selected slides for tissue-type annotation and provided input on the genomic and histopathological analyses. D.Z., K.L.R. and Y.L. reviewed the healthcare record for outcomes and treatment modalities. D.P., A.A. and D.R. produced data engineering infrastructure for H&E slide annotation. A.P. also facilitated data and code sharing. P.S. analyzed the mutational signatures in the MSKCC cohort. P.I.C. segmented CT studies under supervision. C.F. provided data engineering support. M.C. provided biostatistical guidance regarding outcome definitions. J.R.F. supervised the analysis of mutational signatures. R.V. and H.V. provided input on radiomic modeling, N.G. coordinated protocols and cohort lists. R.S. provided support for CT study de-identification and segmentation. S.L. helped generate Extended Data Fig. 1. A.M. provided input on study direction. J.G. oversaw the data engineering efforts supporting the project. Y.L. and S.P.S. conceived of and supervised the study. K.M.B. and S.P.S. wrote the manuscript with input from Y.L., E.A.A. and all other authors, who approved the manuscript.

Competing interests

S.P.S. is a shareholder and consultant to Imagia Canexia Health Inc. Y.L. is a shareholder of Y-mabs Therapeutics Inc. and a consultant to Calyx. J.S.K.-F. reports receiving personal/consultancy fees from Goldman Sachs, REPARE Therapeutics, Paige.AI and Eli Lilly, membership of the scientific advisory boards of VolitionRx, REPARE Therapeutics and Paige.AI, membership of the Board of Directors of DecipherBio, GenomicPath, and a Medical Scientist Training Program Grant from the National Institute of General Medical Sciences, Novartis, Genentech and InVicro. J.S.K.-F. owns Paige.AI stock options. The other authors declare no competing interests.

Additional information

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Extended Data Fig. 1 | Segmenting radiologist and CT vendor in training and test sets. a, Three fellowship-trained radiologists segmented the training (N = 298 patients) and test cases (N = 40 patients). b, Scanner vendors (same number as in preceding panel).
Extended Data Fig. 2 | Genomic features of the training and test sets. **a**, Distribution of large-scale state transitions and threshold. **b**, Signature 3 detections by SigMA with high confidence (HC; N = 48 patients) and low confidence (LC; N = 30 patients), Clock signature (N = 50 patients), Signature 18 (N = 1 patient), and MSI (N = 1 patient). (c) Signature 3 frequencies for all TCGA-OV cases with sequencing from⁴³ are shown. 338 patients with low Sig. 3 and 47 with high Sig. 3. **d–e**, Kaplan–Meier analyses of patients by genomic subtype in the training and test sets (p-value by log-rank test). **f**, Incorporating thresholded LST counts as indicators of HRD status did not increase the difference in OS of the HRD and HRP curves (p-value by log-rank test). **g**, Stratification by PFS using specific mutational subtypes: HRD-Deletion (HRD-DEL), HRD-Duplication (HRD-DUP), Foldback Inversion (FBI), and Tandem Duplications (TD) (p-value by multivariate log-rank test). **h**, Stratification by OS using the same mutational subtypes (p-value by log-rank test). **i**, Kaplan–Meier analysis by OS for only patients with explicit evidence of HRD or HRP, excluding presumed HRP (p-value by multivariate log-rank test).
Extended Data Fig. 3 | Radiomic feature values by segmenting radiologist, CT scanner, and site. The radiomic feature chosen for the model by a segmenting radiologist, b CT vendor, and c whether the scan was acquired at our institution (MSKCC) or elsewhere.
Extended Data Fig. 4 | Example cross-validation histopathologic tissue-type classifications. Three samples (a–c) chosen at random from all slides used for cross-validation.
Extended Data Fig. 5 | Histopathologic feature discovery. The logarithm of the univariate hazard ratio is depicted for each histopathologic feature (N = 281 features) before interquartile range-based filtering, with the cluster in the upper right quadrant comprising primarily features describing tumor nuclear diameter and size.
Extended Data Fig. 6 | Histopathologic embeddings by specimen size. UMAP embeddings of the two-feature histopathologic signature (for N = 283 patients), with each slide's point colorized by the relative specimen size (here, the quantile of the number of foreground tiles detected).
Extended Data Fig. 7 | Test performance of histopathologic-radiomic model. a, Kaplan–Meier analysis of OS for the RH model’s risk scores (p-value by log-rank test). b, Kaplan–Meier analysis of PFS for the RH model’s risk scores (p-value by log-rank test).
Extended Data Fig. 8 | Learning only from cases with full information (N = 114) worsens performance. Log hazard ratios for radiomic features derived from a omental implants (N = 600 features) and b adnexal masses (N = 600 features; uncorrected p-values shown). c, Log hazard ratios for histopathologic features (N = 281 features; uncorrected p-values shown). d, Concordance indices for the test set by overall survival. The height of each bar shows the c-Index, and the lower and upper points of the respective error bars depict the 95% C.I. by leave-one-out bootstrapping. Asterisks denote 95% confidence of significant ordering of the test set by 1000-fold permutation test. e, KM analysis of OS for the GRH model in the test set. P-value calculated using the log-rank test. f, KM analysis of PFS for the GRH model in the test set. P-value calculated using the log-rank test.
Extended Data Fig. 9 | No robust association exists between individual modalities in the training set. a, The maximal magnitude of the Pearson correlation between individual modalities is 0.191. b, The maximal magnitude of the Spearman correlation between individual modalities is 0.192.
Extended Data Fig. 10 | Chemotherapy response scores for all models on the test set. a–o, for C, G, GC, GH, GHC, GR, GRC, GRH, GRHC, H, HC, R, RC, RH, and RHC models, respectively. The box of each plot depicts the 25th, 50th, and 75th percentiles, and the whiskers depict the entire range except for outlier points beyond 1.5 times the interquartile range past from the median 50% of data. Significance was assessed by a one-sided Mann-Whitney U test without correction for multiple tests. * denotes $p < 0.05$, ** denotes $p < 0.01$, ns denotes $p > 0.05$. P-value in b is 0.012. Each plot depicts $N=9$ patients with CRS 3/NET and $N=12$ patients with CRS 1/2.
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

The electronic healthcare record was used to acquire clinical outcomes, and PACS (GE Centricity v 7.0) was used to review CT scans for quality. H&E WSIs were reviewed and segmented in the institutional slide viewer web application. cBioPortal was used to acquire genomic information. For the TCGA, the TClin was used to acquire CT images, the GDC Portal was used to acquire H&E WSIs, and cBioPortal was used to acquire genomic information.

Data analysis

Analysis was conducted in QuPath 0.2.3 (with the StarDist extension), ITK SNAP 3.8.0, and custom code written in Python 3.9.4 (using Pandas 1.2.4, NumPy 1.20.2, PyTorch 1.5.1, TorchVision 0.6, OpenSlide 1.1.1, Seaborn 0.11.1, Matplotlib 3.4.2, SciPy 1.6.3, scikit-learn 0.24.0, PyRadiomics 3.0, and Lifelines 0.25.7). Conda environments are provided to streamline reproducibility.

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DNA sequencing, H&E WSI, and CT data that support the findings of this study have been deposited at Synapse (Sage Bionetworks) under the accession code syn25946517. Additional H&E WSI, CT imaging, and genomic data were derived from the TCGA Research Network: http://cancergenome.nih.gov/ and The Cancer
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**Life sciences study design**

All studies must disclose on these points even when the disclosure is negative.

- **Sample size**
  
  No statistical method was used to predetermine sample size. After reviewing patients with biopsy-proven HGSC and at least one of the following two conditions: (1) CE-CT available in PACS at our institution or (2) H&E WSIs digitized at our institution or source tissue available before chemotherapy, we assembled a dataset of 444 patients. Data were excluded from the analyses only for the reasons detailed in the methods and prior to any machine learning modeling.

- **Data exclusions**
  
  Exclusion criteria were pre-established and included conflicting evidence for HRD status (without high-confidence higher-order feature such as signature 3 detected by SigMA), no tumor, excessive chattering, or intraparenchymal (non-solid tissue) lesions on H&E WSIs, and artifacts, low signal-to-noise ratio, or poor intravenous contrast bolus timing on CT. Every effort was made to include H&E WSIs with minimal tumor and CT images with imperfect quality to test potential clinical feasibility.

- **Replication**
  
  A test set were used to control for overfitting. Our cohort comprised both internal (VSKCC) and external H&E and CT images. Given the data science nature of the project, true replication (i.e., generating new data and testing the same relationships) will require further test set curation. Four-fold cross-validation was used to ensure generalizable performance of the H&E tissue type classifier.

- **Randomization**
  
  The test set was sampled randomly from patients with available CT imaging, H&E imaging, and HRD status.

- **Blinding**
  
  The investigators were not blinded to allocation during outcome assessment. Radiologists and pathologists did not have outcomes readily available during data annotation.

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**Human research participants**

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| Population characteristics | Recruitment |
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| The median age was 63 years [IQR 55-71] for the training set and 66 years [IQR 59-70] for the test set. All patients were female. All patients had biopsy-proven high-grade serous ovarian cancer. In the training set, 175 received neoadjuvant chemotherapy, 82 underwent primary debulking surgery, and 147 had unknown treatment. In the test set, 31 received neoadjuvant chemotherapy, 8 underwent primary debulking surgery, and 1 had unknown treatment. The training set contained 218 homologous recombination proficient and 119 homologous recombination deficient cases, and the test set contained 12 homologous recombination deficient and 28 homologous recombination proficient cases. | Participants were not recruited, but rather retrospectively identified from the institutional data warehouse as per the outlined criteria for data availability and biopsy-proven HGSC diagnosis. |
Ethics oversight

Memorial Sloan Kettering Cancer Center’s Institutional Review Board

Note that full information on the approval of the study protocol must also be provided in the manuscript.