Colorectal Cancer Consensus Molecular Subtypes Translated to Preclinical Models Uncover Potentially Targetable Cancer Cell Dependencies

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Abstract

Purpose: Response to standard oncologic treatment is limited in colorectal cancer. The gene expression–based consensus molecular subtypes (CMS) provide a new paradigm for stratified treatment and drug repurposing; however, drug discovery is currently limited by the lack of translation of CMS to preclinical models.

Experimental Design: We analyzed CMS in primary colorectal cancers, cell lines, and patient-derived xenografts (PDX). For classification of preclinical models, we developed an optimized classifier enriched for cancer cell–intrinsic gene expression signals, and performed high-throughput in vitro drug screening (n = 459 drugs) to analyze subtype-specific drug sensitivities.

Results: The distinct molecular and clinicopathologic characteristics of each CMS group were validated in a single-hospital series of 409 primary colorectal cancers. The new, cancer cell–adapted classifier was found to perform well in primary tumors, and applied to a panel of 148 cell lines and 32 PDXs, these colorectal cancer models were shown to recapitulate the biology of the CMS groups. Drug screening of 33 cell lines demonstrated subtype-dependent response profiles, confirming strong response to EGFR and HER2 inhibitors in the CMS2 epithelial/canonical group, and revealing strong sensitivity to HSP90 inhibitors in cells with the CMS1 microsatellite instability/immune and CMS4 mesenchymal phenotypes. This association was validated in vitro in additional CMS-predicted cell lines. Combination treatment with 5-fluorouracil and luminespib showed potential to alleviate chemoresistance in a CMS4 PDX model, an effect not seen in a chemosensitive CMS2 PDX model.

Conclusions: We provide translation of CMS classification to preclinical models and uncover a potential for targeted treatment repurposing in the chemoresistant CMS4 group.

Introduction

Colorectal cancer is a worldwide health burden, representing the third most common type of cancer and the fourth most common cause of cancer-related deaths (1). Treatment decisions are primarily based on cancer stage and tumor location; however, clinical outcome varies greatly, both with respect to prognosis and treatment response (2). The repertoire of targeted treatments and the number of stratified treatment options based on prognostic and/or predictive factors is limited (3, 4). Colorectal cancer is heterogeneous also at the molecular level (5, 6). This heterogeneity confers primary or secondary resistance to targeted treatments (7) and represents a major challenge for precise interpretation of prognostic and predictive markers (8).

Molecular classification of colorectal cancer has evolved in recent years. Until now, this has been based on the nonoverlapping genomic phenotypes microsatellite instability (MSI) and chromosomal instability, providing both prognostic and predictive information. MSI+ tumors associate with good patient outcome in early stages (9), related to a large mutation burden (10, 11) and cytotoxic immune cell infiltration (12). In the metastatic setting, patients with MSI+ tumors have a poor prognosis (13), but respond well to immune checkpoint inhibition (14). The majority of colorectal cancers have chromosomal instability, and aneuploidy is a predictor of a poor prognosis (15). Recently, more detailed classification of primary colorectal cancer has been proposed based on intrinsic gene expression profiles (16–20), resulting in the four biologically distinct consensus

Note: Supplementary data for this article are available at Clinical Cancer Research Online (http://cincancerres.aacrjournals.org/).

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molecular subtypes (CMS; ref. 21): CMS1 MSI-immune, CMS2 epithelial and canonical, CMS3 epithelial and metabolic, and CMS4 mesenchymal. The CMS classification has prognostic value independent of cancer stage, with dismal survival outcomes for the CMS4 population, even when treated with standard adjuvant chemotherapies (22). A potential predictive value of the CMS classification has not been resolved. Recognizing that the tumor microenvironment is an important contributor to gene expression signals in bulk tumor tissue (24–26), the translation of CMS classification to preclinical models, including cell lines and patient-derived xenografts (PDX) has major challenges. Although CMS labels have previously been assigned to colorectal cancer cell lines (27), development of “adapted” CMS classifiers carefully optimized for preclinical exploration is critical to investigate specific drug sensitivities of subtypes in high-throughput screens. In addition, the question of whether these in vitro models precisely recapitulate the biology of CMS classification has not been resolved.

Here, we studied the distinct molecular and clinicopathologic properties of CMS in an independent, single-hospital series of primary colorectal cancers. Next, we developed a cancer cell–adapted CMS classifier for analysis of preclinical models, and performed high-throughput in vitro drug screening to identify subtype-specific drug sensitivities.

Materials and Methods

Patient material

A consecutive, population-based series of 409 patients treated surgically for stage I–IV colorectal cancer at Oslo University Hospital (Oslo, Norway), between 2005 and 2013 was included (Supplementary Table S1). The study was approved by the Regional Committee for Medical and Health Research Ethics, South Eastern Norway (REC number 1.2005.1629). All patients provided written informed consent, and the study was conducted in accordance with the Declaration of Helsinki. Details of DNA/RNA extraction, as well as MSI status and mutation analyses are included as Supplementary Data.

Colorectal cancer cell lines

Totally 169 colorectal cancer cell lines were analyzed (Supplementary Table S2), including 38 cell lines in-house (details of growth conditions in Supplementary Data) and publicly available gene expression data from 136 cell lines [five overlapping with the in-house dataset; obtained from Gene Expression Omnibus (GEO) accession numbers GSE36133 (28), GSE57083, and GSE59857 (29)]. The number of cell lines derived from unique patients was 148. Cell line identities were verified by fingerprinting according to the AmpFlSTR Identifier PCR Amplification Kit (Life Technologies by Thermo Fisher Scientific), and matched to the profiles reported by the ATCC.

Cell lines were regularly tested for mycoplasma contamination according to the MycoAlert Mycoplasma Detection Assay (Lonza Walkersville Inc.).

Gene expression analysis

The primary colorectal cancers were analyzed for gene expression using Affymetrix GeneChip Human Exon 1.0 ST Arrays (HuEx; n = 201 colorectal cancers) or Human Transcriptome 2.0 Arrays (HTA; n = 208 colorectal cancers) according to the manufacturer’s instructions (Affymetrix Inc.). The in-house cell lines were analyzed on HTA arrays. The data have partly been published previously (GEO accession numbers GSE24550, GSE29638, GSE69182, GSE79959, and GSE97023) and the remaining samples (n = 174 colorectal cancers) have been deposited to GEO with accession number GSE65288. Details of data preprocessing of the in-house and public datasets, as well as CMS classification of the primary colorectal cancers are included as Supplementary Data. Gene set expression enrichment analyses were performed using the R package GSA (30) and a customized collection of 51 colorectal cancer-related gene sets. Sample-wise gene set expression enrichment scores were calculated using the R package GSVA (31).

Development of the cancer cell–adapted CMS classifier

A CMS classifier enriched for cancer cell–intrinsic gene expression signals was developed based on RNA sequencing data from primary colorectal cancers in The Cancer Genome Atlas (TCGA; n = 560) and colorectal cancer cell lines (n = 37 unique; ref. 32), as well as a public microarray dataset of PDX tumors and primary colorectal cancers (n = 40 and 30, respectively; ref. 33). For the TCGA data, preprocessed gene-level RSEM expression values were downloaded from the Broad GDAC Firehouse (level 3; doi:10.7908/C11G0KM9) and CMS assignments from the Colorectal Cancer Subtyping Consortium web site at SAGE Synapse (21). The samples were randomly assigned to a training (75%, n = 417) and a test (25%, n = 143) dataset.

Genes with subtype-specific expression were identified as genes with high relative expression in each CMS group in the TCGA training set. Differential expression analysis was done by comparing each subtype with the rest using the voom approach with quantile normalization in the R package limma, and genes with a log2 fold-change >1 and adjusted P value <0.1 in each subtype were retained. To enrich for genes likely to be informative in cell lines and PDX models, and to exclude genes with high expression in the tumor microenvironment, two
additional filters were applied. First, only genes with high expression in colorectal cancer cell lines (top 25% expressed genes in at least three samples) and high expression variation (top 25% interpercentile range (10th–90th) among the samples) in the RNA sequencing cell line dataset were retained. Second, genes with high expression in primary colorectal cancers compared with PDX tumors were filtered out, retaining only genes with a mean log₂ fold-change below 2 in the primary colorectal cancer versus PDX dataset.

On the basis of this filtered template gene set representing cancer cell–adapted expression signatures of each CMS group, a collection of 148 colorectal cancer cell lines derived from unique patients (totally 169 cell lines) was classified using the nearest template prediction (NTP) algorithm (34) with cosine correlation distances to predict the proximity of each sample to the four template signatures. P-value and false discovery rates (FDR) were calculated on the basis of random resampling (n = 1,000) of the template genes. Sensitivity analysis of the gene expression thresholds applied during filtering of the template gene set is described in the Supplementary Data.

CMS classification of PDX models

PDX models of primary colorectal cancers or liver metastases (n = 32) were established as previously described (35). One tumor from each mouse and samples from four matching primary colorectal cancers were analyzed for gene expression on Affymetrix Human Gene 2.0 ST arrays (details of data preprocessing in the Supplementary Data). Sample classification was performed using the adapted CMS classifier.

Drug screening in colorectal cancer cell lines

An in-house collection of 33 cell lines (Supplementary Table S2B) was analyzed for drug sensitivities in an in vitro screen using an established high-throughput platform (36) including 459 clinically approved or investigational drugs representing different molecular target classes. A drug sensitivity score (DSS) was calculated per drug and cell line relative to a negative and a positive control, based on cell viability after drug treatment at five different concentrations over a 10,000-fold concentration range. Drugs (n = 218) with low efficacy (DSS values above 7 in less than three cell lines) and low variation in DSS values (cross-sample range below 7) were excluded from further analyses. Differential drug sensitivity among sample groups was analyzed by independent samples t tests.

Transcriptional profiling and Western blotting in cells treated with luminespib

Three CMS4 cell lines with varying levels of sensitivity to HSP90 inhibition (Caco2, LIM2099, and SW480) were seeded in 60 mm dishes 24 hours prior to exposure to DMSO (control) or 50 mmol/L luminespib. RNA was isolated after treatment for 6 hours (Qiagen Allprep DNA/RNA/miRNA Universal Kit) and analyzed on Affymetrix HTA microarrays. Differential gene expression analysis was performed by paired samples t-tests comparing treated and control cells using limma. Protein expression of HSP70 and Ki67 was analyzed for drug sensitivities in an in vitro setting and Western blotting in cells treated with luminespib.

Results

Validation of clinicopathologic and biological associations of CMS in primary colorectal cancer

A prospective, single-hospital series of primary colorectal cancers (n = 409; Supplementary Table S1) was classified according to CMS based on gene expression profiles using the random forest (RF) predictor implemented in the R package CMSclassifier (ref. 21; Fig. 1A). The previously described molecular (MSI status, BRAF, KRAS, and TP53 mutations) and clinicopathologic (patient gender, tumor localization, tumor differentiation grade, and cancer stage) associations of each subtype were confirmed (Fig. 1B and C; Supplementary Tables S3 and S4; Supplementary Data). In particular, patients with CMS4 tumors had a poorer 5-year RFS and OS rate than patients with CMS1–3 tumors [HR = 1.8 (patient ID 1)] were selected for drug treatment. Experiments were conducted following the European Union's animal care directive (2010/63/EU) and were approved by the Ethical Committee of Animal Experimentation of Vall d’Hebron Institute of Research (VHIB)/Vall d’Hebron Institute of Oncology (VHIO; ID: 18/15 CEA). NOD-SCID (NOD.CB17-Prkdcscid/ NcrCrj) mice were purchased from Charles River Laboratories (Wilmington, MA). One hundred thousand patient-derived cells suspended in PBS were mixed with Matrigel (1:1 v/v ratio; BD Biosciences) and injected subcutaneously into both flanks of NOD-SCID mice. When the tumor reached 0.5 cm³ in volume, mice (n = 34 for both models) were randomized to each of four different treatment arms, including a control arm (empty vehicle), 5-fluorouracil (5-FU) monotherapy, luminespib (HSP90 inhibitor) monotherapy, and 5-FU + luminespib combination therapy. Luminespib (25 mg/kg in PBS; MCE) was administered by intraperitoneal injection three times per week. 5-FU (40 mg/kg in PBS; Sigma-Aldrich) was administered by intraperitoneal injection twice per week. When matching endpoint criteria, mice were euthanized and complete necropsies were performed. Protein expression of HSP70 and Ki67 was analyzed in posttreatment tissue samples by immunohistochemistry (Supplementary Data).

Statistical analyses

Statistical tests were conducted in R (v.3.3.3), including Fisher exact test of contingency tables with the function fisher.test, t tests with equal or unequal variances (Welch t test) using the function t.test, prediction accuracy using the confusionMatrix-function in the package caret, and two one-sided t tests for equivalence using the t.test-function in the package equivalent, with the magnitude of similarity determined by the parameter epsilon. Unsupervised principal components analysis (PCA) was done using the prComp function. Univariable and multivariable survival analyses were conducted with Cox's proportional hazards regression, with calculation of P-values from Wald tests for predictive potential using the SPSS software version 21 (IBM Corporation). Kaplan–Meier survival curves were compared with the log-rank test. Five-year relapse-free survival (RFS, considering relapse after complete resection or death from any cause as events) and overall survival (OS, considering death from any cause as events) were used as endpoints. Antitumor activity in PDX models was analyzed using a generalized linear mixed model of tumor volume fold changes, with random effects and treatment arm and time as covariates.
(95% confidence interval, CI, 1.2–2.7) and 2.0 (95% CI, 1.3–3.1); $P = 0.005$ and 0.001 for RFS and OS, respectively; see also Fig. 1D and Supplementary Fig. S1A]. This was independent of known clinicopathologic prognostic factors and MSI status in multivariable analyses [HR = 1.4 (95% CI, 0.9–2.2) and 1.6 (95% CI, 1.04–2.6), $P = 0.1$ and 0.03 for 5-year RFS and OS, respectively; Supplementary Table S5]. The distinct biological properties of each CMS group, including infiltration patterns of immune and stromal cells, were also validated by gene set expression enrichment analyses (Fig. 1E; Supplementary Fig. S1B; Supplementary Table S6 and Supplementary Data).

CMS classification of preclinical models

Our main interest was to study CMS-specific drug sensitivities in cell line models and particularly in the poor prognostic CMS4 group. As confirmed in our clinical cohort, the transcriptome of CMS4 primary colorectal cancers is greatly influenced by signals from the tumor microenvironment, and application of the original RF CMSclassifier to a collection of 148 unique colorectal cancer cell lines showed that it failed to accurately identify this subtype in the in vitro models. Using default settings, 82 cell lines (55%) were unclassified and among the classified, 41 (62%) were CMS2, and only 3 (5%) were CMS4 (Supplementary Fig. S2A). Gene set analyses showed that the three CMS4 cell lines indeed had clear CMS4 characteristics, including epithelial to mesenchymal transition (EMT) and TGFβ responses, but this was true also for additional, unclassified samples (Supplementary Fig. S2b).

Furthermore, this classification failed to accurately distinguish between the two epithelial subtypes CMS2-canonical and CMS3-metabolic (Supplementary Fig. S2C). To improve the classification of preclinical models, we therefore generated a novel CMS classifier enriched for cancer cell-intrinsic gene expression signals (Fig. 2A; details of the public expression datasets and analysis thresholds used are included in the Materials and Methods). First, potential template genes were identified as genes with high relative expression in each CMS group in primary colorectal cancers ($n = 1,994$ unique genes; Supplementary Table S7). Next, this gene set was filtered to exclude (i) genes with a low expression level or expression variation in colorectal cancer cell lines ($n = 1,454$ genes) and (ii) genes expressed in the tumor microenvironment, identified as genes with a high expression in primary colorectal cancers compared with PDX tumors ($n = 57$ additional genes; Supplementary Table S7). The resulting list of genes ($n = 483$; Supplementary Table S8) were used as templates for CMS.
classification based on the NTP algorithm (34). This new classifier is publicly available as the R package CMScaller and can be downloaded from https://github.com/Lothelab/CMScaller (Eide and colleagues; submitted for publication).

To assess prediction accuracy in patient samples, the adapted CMS classifier was applied to four independent series of primary colorectal cancers (total n = 709) analyzed on four different gene expression platforms. Classification concordance compared with

Figure 2.
CMS classification of colorectal cancer cell lines. A, Flowchart of development of the cancer cell–adapted CMS classifier. Candidate template genes with high relative expression in each CMS group were identified in colorectal cancers from TCGA (left). Prior to CMS classification using Nearest Template Prediction (right), genes with low expression levels and/or expression variation in colorectal cancer cell lines and genes with high expression in the tumor microenvironment were filtered out (blue background). B, Confident CMS classification was obtained for 126 (85%) of 148 colorectal cancer cell lines from unique patients using the adapted CMS classifier, with similar distribution among the subtypes as for the consecutive patient series. The molecular and biological characteristics of the CMS groups were also recapitulated among the cell lines, as shown in C, for MSI status, BRAF mutations, KRAS mutations, and TP53 mutations, as well as (D) by gene set expression enrichment analyses.
Drug Responses Among Consensus Molecular Subtypes of Colorectal Cancer

Table 1. Prediction accuracy of the cancer cell-adapted CMS classifier in primary colorectal cancers

<table>
<thead>
<tr>
<th>Patient series</th>
<th>Analysis platform</th>
<th>Samples classified by both CMS classifiers</th>
<th>Reference subtype</th>
<th>Cancer cell-adapted CMS classifier</th>
<th>Prediction accuracy (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TCGA test-set</td>
<td>RNA sequencing</td>
<td>91 (64%)</td>
<td>CMS1</td>
<td>CMS1</td>
<td>85% (76%–91%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CMS2</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CMS3</td>
<td>0</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>CMS4</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>GSE14333 (n = 157)*</td>
<td>Affymetrix HG U133 Plus 2.0 arrays</td>
<td>116 (74%)</td>
<td>CMS1</td>
<td>15</td>
<td>86% (79%–92%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CMS2</td>
<td>1</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>CMS3</td>
<td>3</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>CMS4</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>In-house patients (n = 208)*</td>
<td>Affymetrix HTA 2.0 arrays</td>
<td>165 (78%)</td>
<td>CMS1</td>
<td>36</td>
<td>92% (87%–96%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CMS2</td>
<td>0</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>CMS3</td>
<td>1</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>CMS4</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>In-house patients (n = 201)*</td>
<td>Affymetrix Human Exon 1.0 ST arrays</td>
<td>138 (69%)</td>
<td>CMS1</td>
<td>22</td>
<td>87% (80%–92%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>CMS2</td>
<td>0</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>CMS3</td>
<td>1</td>
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<td></td>
<td></td>
<td></td>
<td>CMS4</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

*Reference CMS classes obtained from Guinney and colleagues (21).

*Reference CMS classes obtained using the RF predictor implemented in the R package CMSclassifier.

the original RF CMSclassifier ranged from 85% to 92%, demonstrating robust performance independent of analysis platform (Table 1).

Colorectal cancer cell lines. CMS classification was obtained for 126 (85%) of the 148 unique colorectal cancer cell lines using the adapted classifier and an FDR threshold from NTP of 0.2 (Supplementary Table S2A). The CMS distribution across the cell lines was similar to the in-house patient series (Fig. 2B; \( P < 0.05 \) from paired test of equivalence with magnitude of similarity above 0.8).

In comparison with the original RF CMSclassifier, the concordance in subtype assignments for cell lines classified by both approaches was high (88%), and the added value of the adapted classifier was primarily the higher classification rate, in particular in CMS3 and CMS4 (Supplementary Table S9). To determine whether key characteristics of the CMS groups were recapitulated in the cell lines, we explored associations between CMS and other molecular data. Similarly to primary colorectal cancers, CMS1 cell lines showed strong enrichment for MSI (\( P = 2 \times 10^{-4} \)) and BRAF mutations (\( P = 6 \times 10^{-4} \); Fig. 2C and Supplementary Table S10). CMS3 cell lines were frequently MSI+ and KRAS mutated, whereas TP53 mutations were enriched in CMS2, although not statistically significant. Gene expression-based PCA indicated that CMS1/4 versus CMS2/3 represented the primary sample split (\( P = 2 \times 10^{-26} \) from comparison of principal component 1 (PC1) between the two sample groups; Supplementary Fig. S3), and gene set analyses confirmed that CMS1 and CMS4 cell lines were undifferentiated, whereas CMS2 and CMS3 showed clear epithelial characteristics (Fig. 2D and Supplementary Table S11). CMS2 and CMS3 additionally had upregulation of HNF4A targets, whereas CMS3 was particularly enriched for metabolic pathways. CMS4 was specifically characterized by EMT activation, extracellular matrix organization, and TGFβ responses.

Optimal performance of the classifier is dependent on unbiased representation of all CMS groups in the query sample set, and to estimate stability, cell line classification was repeated after random resampling of cell line subsets (\( n = 1,000 \) resamplings of 50% of the cell lines). The majority of cell lines (82% of the 148 unique) retained their CMS group in more than 95% of the resamplings (Supplementary Table S12). The classification uncertainty was highest in CMS1 (Supplementary Fig. S4A), which may be associated with an enrichment of MSI+ samples in the cell line collection (38% versus 18% in our patient series). However, gene set expression analysis specifically among MSI+ cell lines showed expected CMS-associations, also for CMS1 (Supplementary Fig. S4B).

To assess the independence of the adapted classifier from tumor stroma, the stromal and epithelial compartments of laser microdissected primary colorectal cancers (GSE35602; ref. 38) were analyzed. Some template genes had high relative expression in the stromal samples (Supplementary Fig. S5), and an additional template gene filter was therefore tested by excluding these genes (Supplementary Table S7). Cell line classification with the reduced template gene set was highly concordant with the initial adapted classifier [90% accuracy (95% CI, 83–95%)] among the confidently classified cell lines; Supplementary Table S12), indicating that the influence of stromal gene expression signals on sample classification was low. Furthermore, gene set analyses and resampling of the cell lines (as above) indicated that the reduced template gene set did not improve the subtype assignment or classification stability (Supplementary Fig. S5).

Patient-derived xenografts. The adapted classifier was also applied to a set of 32 PDX models of colorectal cancer (22 derived from primary tumors and 10 from liver metastases), and the matching primary tumor from four patients. Subtype assignment was obtained for 28 (88%) of the PDX tumors (FDR from NTP lower than 0.2), including 7 (25%) to CMS1, 13 (46%) to CMS2, 5 (18%) to CMS3, and 3 (11%) to CMS4. Concordant subtypes were assigned for three of the four matching PDX-patient tumor pairs (Supplementary Table S13). Although with a lower sample number, the in vivo models also recapitulated important features...
of the CMS groups. In concordance with results from the patient series and cell lines, CMS1/4 versus CMS2/3 represented the primary sample split based on gene expression PCA (Supplementary Fig. S6). Furthermore, gene set expression analyses showed that CMS1 was enriched for "MSI-like" and "BRAF-mutant-like" PDX tumors, whereas CMS2 and CMS3 had epithelial characteristics, with enrichment of colonic differentiation signatures. CMS2 additionally had high WNT signaling and CMS3 had enrichment of metabolic signatures. CMS4 showed enrichment for angiogenesis. For comparison, PDX classification based on the reduced template gene set (additionally filtered for stromal gene expression) was highly concordant [93% accuracy (95% CI, 77–99%)].

CMS defines subgroups of cell lines with distinct drug response profiles
To explore subtype-specific drug responses, 33 cell lines established from 29 patients (Supplementary Table S2b) were selected for in vitro pharmacogenomic profiling using an established high-throughput drug screening platform (n = 459 drugs; Supplementary Table S14; ref. 36). DSS values (37) were calculated for each drug based on cell viability after treatment at five different concentrations, and quality control showed strong reproducibility of the DSS values between independent drug screens of the same cell line (RKO; Pearson correlation 0.99, standard deviation of difference between repeated screens 1.36). Furthermore, drug screen reproducibility between paired cell lines from each of four patients was associated with their pair-wise similarity in gene expression (Supplementary Fig. S7A). For subgroup comparisons, paired cell lines were excluded (HCT15, WIDR, SW620, and IS1), and the final set (n = 29) represented all four CMS groups (n = 7, 9, 5, and 8 predicted CMS1, 2, 3, and 4, respectively, not restricted by the FDR from CMS prediction; Supplementary Fig. S7B).

Principal component analysis based on DSS values from the drug screen indicated a separation of the cell lines into two response groups by MSI status (Supplementary Fig. S8A; P = 7 × 10^{-7} by Welch t-test comparing PC1 between MSI+ and microsatellite stable, MSS, samples). Comparisons of individual drug responses between the two sample groups confirmed that this distinction was primarily caused by a strong relative sensitivity to chemotherapeutic drugs in MSI+ cell lines, in particular topoisomerase inhibitors and gemcitabine (Supplementary Fig. S8B; Supplementary Table S15). CMS accounted for additional variation in DSS values (Supplementary Fig. S8C) and to explore subtype-specific sensitivities, drug response comparisons were made between all the individual CMS groups (Fig. 3; Supplementary Table S16). Consistent with the high prevalence of MSI in CMS1, CMS1 cell lines were more sensitive to antimetabolites and

Figure 3.
Differential drug responses among CMS groups. High-throughput drug screening (filtered list of 241 of totally 459 drugs) of colorectal cancer cell lines (n = 29) revealed differential drug responses among the CMS groups. Each plot represents a comparison of two subtypes, as indicated, and each dot represents one drug. Selected drugs are colored according to molecular targets, as indicated.
Inhibitors of topoisomerases and mitosis than CMS2. In addition, CMS1 showed stronger sensitivity to heat shock protein 90 (HSP90) inhibitors than both CMS2 and CMS3. There were few drugs with differential sensitivity between CMS1 and CMS4, or between CMS2 and CMS3. However, CMS2 cell lines were more sensitive to EGFR and HER2 inhibitors than both CMS3 and CMS4. CMS4 cell lines showed strong sensitivity to HSP90 inhibitors, atorvastatin (HMG-CoA reductase inhibitor), 2-methoxyestradiol (2ME; combined angiogenesis and tubulin inhibitor), and disulfiram (inhibitor of alcohol dehydrogenase) compared with both CMS2 and CMS3 (these selected drug screen data are available in Supplementary Table S17).

Summarized, these comparisons indicated that EGFR and HER2 inhibitors had particularly strong activity in CMS2, which was confirmed in a direct comparison of CMS2 versus CMS1/3/4 cell lines (Fig. 4A; Supplementary Table S18). Strong relative response to anti-EGFR treatment in CMS2 was also validated in published data of cetuximab treatment in 130 unique cell lines (29), independent of KRAS and BRAF mutation status (Supplementary Fig. S9). In addition, CMS1 and CMS4 appeared to be sensitive to similar classes of agents, in line with the major distinction observed in the gene expression data between the undifferentiated CMS1/4 and epithelial-like CMS2/3 cell lines. Indeed, correlation analyses between PC1 of the DSS values and sample-wise gene set expression enrichment scores (calculated using the R package GSVA; ref. 31; gene sets listed in Supplementary Table S6), showed that the overall drug response pattern among the cell lines was most strongly correlated to a colonic differentiation signature (‘colonic crypt, top’; Spearman correlation: 0.7, \( P = 2 \times 10^{-7} \); Supplementary Fig. S8D). Accordingly, CMS1/4 cell lines were compared with CMS2/3, and a strong relative response to several HSP90 inhibitors (luminespib, ganetespib, and radicicol), 2ME, indibulin (another tubulin-inhibitor), atorvastatin, and tipifarnib (farnesyltransferase inhibitor) in CMS1/4 was confirmed (Fig. 4B; Table 2). These same drugs had stronger relative activity in CMS1/4 also when analyzing MSS cell lines only, when including only cell lines with FDR from CMS assignment below 0.2, when including the opposite set of the paired cell lines, and based on CMS classification using the reduced template gene set (additional filtered for stromal gene expression; Supplementary Fig. S10).

**Strong relative activity of HSP90 inhibitors in CMS1 and CMS4 is validated in vitro**

For independent biological validation of differential drug activity in CMS1/4 compared with CMS2/3 cell lines, five additional cell lines were predicted to belong to either the CMS1 (LIM2405) or CMS4 (CAR1, HCA7, LIM2099, and OUMS23) subtypes based on their gene expression profiles, and subsequently screened for drug sensitivities with the same experimental setup as in the initial discovery screen. Two CMS3 cell lines (HT29 and LS174T) were included as controls in the validation drug screen. Clear differential sensitivity for all three HSP90 inhibitors (luminespib, ganetespib, and radicicol), 2ME, atorvastatin, and disulfiram was validated in CMS1 and CMS4 compared with CMS3 (Fig. 4C).

Furthermore, strong sensitivity to HSP90 inhibition in CMS1 and CMS4 was validated in public drug response data from 15 colorectal cancer cell lines (nine overlapping with our drug screen) treated with ganetespib (39). The cell lines were classified using the adapted classifier and the CMS1/4 group was found to have lower IC\(_{50}\)-values for ganetespib (mean 24 nmol/L) than CMS2/3 (mean 52 nmol/L), indicating higher sensitivity in the first group (Supplementary Fig. S11A). Similarly, among 32 colorectal cancer cell lines from the Genomics of Drug Sensitivity in Cancer Project (16 overlapping with our drug screen), higher sensitivity to the HSP90 inhibitor CCT018159 was confirmed in CMS1/4 [average log\(_{10}\)IC\(_{50}\) in nmol/L] 3.4] compared with CMS2/3 [average log\(_{10}\)IC\(_{50}\) in nmol/L] 5.6, \( P = 0.0004 \) by Welch t-test. Here, stronger relative sensitivity in CMS1/4 was found also among MSS cell lines only (Supplementary Fig. S11B).

**HSP90 inhibition is associated with upregulation of heat shock response**

To identify the transcriptional changes associated with response to HSP90 inhibition, three CMS4 cell lines (CAC02, LIM2099, and SW480) were treated with luminespib. Differential gene expression analysis of treated compared with control cells (DSMO) showed that upregulation of heat shock response was the dominant response mechanism, with upregulation of several members of the HSP family (Fig. 4D; Supplementary Table S19). Upregulation of two main HSP90 co-chaperones, HSP70 and HSP40, was confirmed at the protein level (Supplementary Fig. S12A). Heat shock transcription factor 1 (HSF1) and its transcriptional activity has previously been described to be a resistance mechanism against HSP90 inhibition, and concordantly, we selected a CMS4 colorectal cancer cell line treated with HSP90 inhibition may alleviate chemoresistance in CMS4 in vivo

In our drug screen panel, CMS4 had a particularly poor response to fluoropyrimidines (\( P \leq 0.05 \) among MSS cell lines; Supplementary Fig. S13). Previous studies have suggested that HSP90 inhibition may sensitize colorectal cancer cell lines to chemotherapy, and although monotherapy with HSP90 inhibitors has shown low efficacy in metastatic colorectal cancer (42), response has been obtained by combination therapy with HSP90 inhibitors and capcitabine (5-FU pro-drug) in patients who have progressed on fluoropyrimidines (43). Accordingly, to analyze a potential effect of HSP90 inhibition in vivo, we selected a CMS4 PDX model (MSS, KRAS/NRAS wild type, BRAF/V600E mutated) for treatment in a randomized and controlled set-up. Immunodeficient NOD-SCID mice (n = 34) were injected with cells derived from a liver metastasis of a chemotherapy-naïve colorectal cancer patient and randomized to four treatment arms: (i) control arm with vehicle, (ii) single agent 5-FU, (iii) single agent luminespib, and (iv) combination therapy with 5-FU + luminespib. Consistent with the cell line data, this CMS4 model showed poor response to chemotherapy (Fig. 4E). Chemoresistance was confirmed by staining for the proliferation marker Ki67 in posttreatment samples, and there were no significant changes in Ki67 expression in mice receiving 5-FU compared with vehicle-treated controls. Furthermore, monotherapy with luminespib did not impact on tumor growth, but combined administration of 5-FU + luminespib resulted in significantly greater antitumor activity.
Discussion

Response to standard oncologic treatment is limited in colorectal cancer and there is great potential to improve treatment efficacy by molecularly-guided repurposing of targeted drugs. We identify strong relative activity of HSP90 inhibitors in in vitro models of the transcriptomic CMS1 and CMS4 groups of colorectal cancer by high-throughput drug screening, using a new and cancer cell-adapted CMS classifier. HSP90 inhibition has previously been extensively investigated in cancer and has demonstrated antitumor activity in several solid tumor types, mainly as combination therapies (41). However, low response rates are observed in unstratified patient populations. In the only phase

Table 2. Differential drug sensitivity between CMS1/4 and CMS2/3 cell lines

<table>
<thead>
<tr>
<th>Drug</th>
<th>Average difference in DSS*</th>
<th>P</th>
<th>FDR</th>
<th>Molecular targets/mechanisms</th>
</tr>
</thead>
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<tr>
<td>PF-03758309</td>
<td>10.8</td>
<td>$8 \times 10^{-4}$</td>
<td>$8 \times 10^{-3}$</td>
<td>PAK inhibitor</td>
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<tr>
<td>Rigosertib</td>
<td>10.5</td>
<td>$4 \times 10^{-4}$</td>
<td>$6 \times 10^{-3}$</td>
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<tr>
<td>Disulfiram</td>
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<td>$1 \times 10^{-2}$</td>
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<tr>
<td>YM155</td>
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<td>$1 \times 10^{-2}$</td>
<td>Survivin inhibitor</td>
</tr>
<tr>
<td>Tipifarnib</td>
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<td>$9 \times 10^{-3}$</td>
<td>Farnesyltransferase inhibitor</td>
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<td>Luminespib</td>
<td>8.3</td>
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<td>$4 \times 10^{-3}$</td>
<td>HSP90 inhibitor</td>
</tr>
<tr>
<td>Ganetespib</td>
<td>8.1</td>
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<td>$2 \times 10^{-3}$</td>
<td>HSP90 inhibitor</td>
</tr>
<tr>
<td>Idarubicin</td>
<td>7.9</td>
<td>$4 \times 10^{-4}$</td>
<td>$6 \times 10^{-3}$</td>
<td>Topoisomerase II inhibitor</td>
</tr>
<tr>
<td>Teniposide</td>
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<td>$1 \times 10^{-2}$</td>
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<tr>
<td>Indiniblin</td>
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<td>$8 \times 10^{-3}$</td>
<td>Mitotic inhibitor; microtubule depolymerizer</td>
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<td>$2 \times 10^{-2}$</td>
<td>RNA and DNA synthesis inhibitor</td>
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<tr>
<td>Clofarabine</td>
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<td>$3 \times 10^{-4}$</td>
<td>$1 \times 10^{-2}$</td>
<td>Antimetabolite; Purine analog</td>
</tr>
<tr>
<td>Danusertib</td>
<td>7.1</td>
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<td>$2 \times 10^{-3}$</td>
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<td>$4 \times 10^{-3}$</td>
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<tr>
<td>Radicicol</td>
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<td>Cytarabine</td>
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<td>Antimetabolite, interferes with DNA synthesis</td>
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<td>$9 \times 10^{-4}$</td>
<td>$8 \times 10^{-3}$</td>
<td>Nucleoside analog; RNA synthesis inhibitor</td>
</tr>
</tbody>
</table>

*Top 20 drugs (FDR from independent samples t-tests below 0.05) sorted by average difference in DSS values between CMS1/CMS4 (n = 15) and CMS2/CMS3 (n = 14) cell lines.
*Positive values indicate drugs with strongest effect in CMS1/CMS4 cell lines.

Figure 4.

Selective activity of HSP90 inhibitors in CMS1 and CMS4. A, High-throughput drug screening of colorectal cancer cell lines (n = 29) showed that CMS2 was more sensitive to EGFR and HER2 inhibitors than the other three CMS groups. B, Compared with CMS2 and CMS3, CMS1 and CMS4 cell lines were more sensitive to three HSP90 inhibitors (red: luminespib, ganetespib and radicicol), 2ME (green: combined angiogenesis and tubulin inhibitor), atorvastatin (dark blue; HMG-CoA reductase inhibitor), Indiniblin (pale blue; tubulin-inhibitor), and disulfiram (pink inhibitor of alcohol dehydrogenase). C, A validation drug screen of five additional cell lines predicted to belong to the CMS1 or CMS4 groups (green and black cell lines are MSI+ and MSS, respectively) confirmed strong sensitivity (red) to HSP90 inhibitors, 2ME, atorvastatin and disulfiram in comparison to two CMS3 cell lines included in the validation screen, as well as in comparison to the mean sensitivity in CMS2 and CMS3 cell lines in the initial screen. D, Three CMS4 cell lines with response to HSP90 inhibition (Caco2, Lm2099, and SW480) were treated with luminespib or DMSO (control). Paired differential gene expression analysis showed upregulation of several members of the HSP family after HSP90 inhibition. E, In CMS4 PDX models (n = 34) of a liver metastasis from a chemotherapy-naive colorectal cancer patient, combined administration of 5-FU and luminespib showed stronger antitumor activity than single agent treatment with 5-FU or luminespib, or in vehicle-treated controls. Tumor growth is plotted as the mean ± SE of tumor volume fold changes of all mice per treatment arm at the indicated time points. No significant changes in Ki67 protein expression in posttreatment samples (relative to -treatment samples) confirmed that the CMS4 model was chemoresistant, whereas increased expression of HSP90 treatment showed a targeted effect of HSP90 inhibition (P-values were calculated by Welch t-test; sample numbers vary due to availability of high-quality samples or data). F, CMS2 PDX models (n = 34) were highly chemosensitive, as shown by a strong antitumor activity of 5-FU monotherapy and reduced Ki67 expression in posttreatment samples, and there was no synergistic effect of combining 5-FU with luminespib. In contrast with the in vitro data, luminespib monotherapy had a moderately stronger antitumor activity in CMS2 (relative to vehicle-treated controls) than in CMS4, but this was not associated with changes in HSP70 expression in CMS2 posttreatment samples.
This is the case also for CMS classification. Although devoid of tumor stroma, the cell lines recapitulated the individual CMS groups and their biological properties. The adapted classifier is enriched for cancer cell-intrinsic gene expression signals, although not completely independent of the tumor microenvironment. Still, additional filtering of the template gene set to further reduce the potential influence of stromal gene expression had little impact on sample classification, indicating robustness. Importantly, the classifier performed well also in tumor samples, confirming reproducibility of the classification in primary colorectal cancers, and showing translation of the classification to PDX models, where contamination of gene expression signals from murine stroma may be a challenge. It has recently been recognized also by others that the original CMS classifier fails to identify some of the CMS groups not only in cell lines, but also in patient-derived organoids and xenografts (50). We argue that this may be alleviated by our adapted classifier.

Important features such as the level of intra- and/or intertumor heterogeneity of CMS, as well as the stability of the subtypes during metastatic progression, are still unknown. However, we validated the clinicopathologic and biological properties of the CMS groups in a single-hospital series of primary colorectal cancers. We also identified strong relative response to EGFR and HER2 inhibitors in cell lines of the CMS2 subtype. This is consistent with the high relative frequency of EGFR and ERBB2 (encoding the HER2 protein) amplification in CMS2 (21), and with the strong sensitivity to cetuximab demonstrated in cell lines of the late transient-amplifying gene expression-based subtype (29) and in PDXs of a subtype with high WNT signaling (50), both of which are largely overlapping with CMS2. Altogether, this reinforces the potential of CMS as a framework for stratified treatment in colorectal cancer.

In conclusion, we show reproducibility of the CMS groups in primary colorectal cancer and provide translation of the classification to preclinical models. Drug screening of cell lines identified CMS1 and CMS4 as potential predictive biomarkers for response to HSP90 inhibition. In vivo, this targeted treatment may alleviate chemoresistance in CMS4. The poor patient prognosis associated with CMS4 warrants additional studies to pursue the potential for clinical testing of HSP90 inhibitor repositioning and combination therapy in colorectal cancer.

Disclosure of Potential Conflicts of Interest
J. Tabernero is a consultant/advisory board member for Bayer, Boehringer Ingelheim, Genentech/Roche, Lilly, MSD, Merck Serono, Novartis, Roche, Sanofi, Symphogen, and Taiho. No potential conflicts of interest were disclosed by the other authors.

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Administrative, technical, or material support (i.e., reporting or organizing data, constructing databases): J. Bruun, A. Munumagi, M. Arjama, H.G. Palmer, A. Nesbakken, R. Dienstmann, R.A. Lothe

Study supervision: R. Dienstmann, R.A. Lothe

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