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Meta-analysis of fecal metagenomes reveals global microbial signatures that are specific for colorectal cancer

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Competing Interest

Author Contributions

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Abstract

Association studies have linked microbiome alterations with many human diseases, but not always reported consistent results, which necessitates cross-study comparisons. Here, a meta-analysis of eight geographically and technically diverse fecal shotgun metagenomic studies of colorectal cancer (CRC, N = 768), which was controlled for several confounders, identified a core set of 29 species significantly enriched in CRC metagenomes (FDR < 1E-5). CRC signatures derived from single studies maintained accuracy in other studies. By training on multiple studies we improved detection accuracy and disease specificity for CRC. Functional analysis of CRC metagenomes revealed enriched protein and mucin catabolism genes and depleted carbohydrate degradation genes. Moreover we inferred elevated production of secondary bile acids from CRC metagenomes suggesting a metabolic link between cancer-associated gut microbes and a fat- and meat-rich diet. Through extensive validations, this meta-analysis firmly establishes globally generalizable, predictive taxonomic and functional microbiome CRC signatures as a basis for future diagnostics.

Introduction

Studying microbial communities colonizing the human body in a culture-independent manner has been enabled by metagenomic sequencing technologies [1]. These have yielded glimpses into the complex yet incompletely understood interactions between the gut microbiome – the microbial ecosystem residing primarily in the large intestine – and its host [2]. To explore microbiome-host interactions in a disease context, metagenome-wide association studies (MWAS) have begun to map gut microbiome alterations in diabetes, inflammatory bowel disease, colorectal cancer and many other conditions [3–12]. However, due to the many biological factors possibly influencing gut microbiome composition in addition to the condition studied, a current challenge for MWAS is confounding, which can cause false associations [13, 14]. This issue is further aggravated by a lack of standards in

metagenomic data generation and processing, making it difficult to disentangle technical from biological effects [15].

Robustness of microbiome-disease associations can be assessed through comparisons across multiple metagenomic case-control studies, i.e. meta-analyses. These aim at identifying associations that are consistent across studies and thus less likely attributable to biological or technical confounders. Most informative are meta-analyses of populations from diverse geographic and cultural regions. Previous microbiome meta-analyses based on 16S rRNA gene amplicon data found stark technical differences between studies and the reported taxonomic disease associations were either of low effect size or not well resolved [16–18]. In contrast, shotgun metagenomics enables analyses with higher taxonomic resolution and of gene functions to improve statistical power for fine-mapping disease-associated strains and aid in the interpretation of host-microbial co-metabolism. Thus far however, meta-analyses of shotgun metagenomic data have either reported on features of general dysbiosis in comparisons across multiple diseases [19], or have left it unclear how well microbiome signatures generalize across studies of the same disease when data are rigorously separated to avoid over-optimistic evaluations of their prediction accuracy [20].

Here, we present a meta-analysis of a total of eight studies of CRC including fecal metagenomic data from 386 cancer cases and 392 tumor-free controls. After consistent data reprocessing, we examined an initial set of five studies for CRC-associated changes in the gut microbiome. Firstly, we investigated potential confounders, followed by identifying (univariate) microbial species associations, and inferring species co-occurrence patterns in CRC. Secondly, we trained multivariable classification models for recognition of CRC status, from both taxonomic and functional microbiome profiles and tested how accurately these models generalized to data from studies not used for training. Moreover, we evaluated performance improvements achieved by pooling data across studies and the disease-specificity of the resulting classification models. Thirdly, targeted investigation of virulence and toxicity genes as candidate functional biomarkers for CRC revealed several of these to be enriched in CRC metagenomes indicative of their prevalence and potential relevance in CRC patients. Three additional, more recent studies were finally used to independently validate these taxonomic and functional CRC signatures.

Results

Consistent processing of published and new data for meta-analysis of CRC metagenomes

In this meta-analysis we included four published studies which used fecal shotgun metagenomics to characterize CRC patients compared to healthy controls (referred to by the country codes FR, AT, CN, and US, corresponding to the respective main study population; see Table 1, Supplementary Table S1, and Methods for inclusion criteria). For an additional fifth study population, we generated new fecal metagenomic data from samples collected in Germany (herein abbreviated as DE); a subset of samples from this patient collective were published previously (Table 1, Methods, [8]). These five studies were conducted on three continents and differed in sampling procedures, sample storage, and DNA extraction protocols. Notably, the fecal specimen of the US study were freeze-dried and stored at -80°C for more than 25 years before DNA extraction and sequencing [10]. In all studies, however,

samples were collected prior to treatment, thus excluding cancer therapy as a potential confounding effect [14, 21]. Most samples were even taken before bowel preparation for colonoscopy, with some exceptions in the DE, CN and US studies (Supplementary Table S2). To ensure consistency in bioinformatic analyses, all raw sequencing data were (re-)processed using mOTUs2 for taxonomic profiling [22] and MOCAT2 for functional profiling [23].

Univariate meta-analysis of species associated with CRC

The first aim of the meta-analysis was to determine gut microbial species that are enriched or depleted in CRC metagenomes in a consistent manner across the five study populations. However, as these studies differed from one another in many biological and technical aspects, we first quantified the effect of study-associated heterogeneity on microbiome composition. We contrasted this with other potential confounders ('patient age', 'BMI', 'sex', 'sampling after colonoscopy', and 'library size'; additionally, 'smoking status', 'type II diabetes comorbidity', and 'vegetarian diet' where available Extended Data 1, Supplementary Table S3). This analysis revealed the factor 'study' to have a predominant impact on species composition, which is supported by a recent comparison of DNA extraction protocols, as these typically differ between studies [15]. An analysis of microbial alpha and beta diversity showed study heterogeneity to also have a larger effect on overall microbiome composition than CRC in our data (Extended Data 2).

For the identification of microbial taxa significantly differing in abundance in CRC, parametric effect size measures are not well established, because microbiome data is characterized by non-Gaussian distributions with extreme dispersion; we thus used a generalisation of the fold change (Extended Data 3) and non-parametric significance testing. In this permutation test framework [24] (herein referred to as blocked univariate Wilcoxon tests) differential abundance in CRC can be assessed while accounting for 'study' as a nuisance effect that is treated as a blocking factor; additionally, motivated by our confounder analysis, we also blocked for 'colonoscopy' in all analyses (Methods, Extended Data 1). To rule out spurious associations due to the compositional nature of microbial relative abundance data, we additionally compared the results of this test with a method [25] employing log-ratio transformation (and found highly correlated results, Supplementary Fig. 1, Supplementary Table S4).

At a meta-analysis false discovery rate (FDR) of 0.005, we identified 94 microbial species to be differentially abundant in the CRC microbiome, out of 849 species consistently detected across studies (Supplementary Table S4, Methods). Among these, we focused on a core set of the 29 most significant markers (FDR < 1E-5, Fig. 1a) for further analysis. The latter included members of several genera previously associated with CRC, such as *Fusobacterium*, *Porphyromonas*, *Parvimonas*, *Peptostreptococcus*, *Gemella*, *Prevotella*, and *Solobacterium* (Fig. 1b, [8–11]), and 8 additional species without genomic reference sequences (meta-mOTUs, Methods, [22]) mostly from the *Porphyromonas* and *Dialister* genera and the Clostridiales order (see Extended Data 4 and Supplementary Table S4 for genus-level associations). Collectively, these 29 core CRC-associated species show a previously underappreciated diversity of 11 Clostridiales species to be enriched in CRC (Fig.

1b). In contrast to the majority of species that are more strongly affected by study heterogeneity than by CRC status, 26 out of the 29 CRC-associated species varied more by disease status (Fig. 1d).

All of the core CRC-associated species were enriched in patients and were often undetectable in metagenomes from non-neoplastic controls. While previous studies were contradictory in the reported proportion of positive versus negative associations [8, 9, 17, 20], our meta-analysis results are more easily reconciled with a model in which – potentially many – gut microbes contribute to or benefit from tumorigenesis than with the opposing model in which a lack of protective microbes contributes to CRC development (Fig. 1b). Although these core taxonomic CRC associations were highly significant and consistent, individual studies showed marked discrepancies in the species identified as significant (Fig. 1a). Retrospective examination of the precision and sensitivity with which individual studies detected this core of CRC-associated species showed relatively low sensitivity for the US study (consistent with the original report [10]) and low precision of the AT study due to associations that were not replicated in other studies (Supplementary Fig. 2).

Analyzing patient metagenomes for co-occurrences among the core set of 29 species that are strongly enriched in the CRC microbiome revealed four species clusters with distinct taxonomic composition (Fig. 2a, Extended Data 5, Methods). Two of them showed strong taxonomic consistency: Cluster 1 exclusively comprised *Porphyromonas* spp., and cluster 4 only contained members of the Clostridiales order. In contrast, the other two clusters were taxonomically more heterogeneous with cluster 3 grouping together the species with highest prevalence in CRC cases (all among the ten most highly significant markers), consistent with a co-occurrence analysis of one of the data sets included here [11]. Cluster 2 contained species with intermediate prevalence.

Investigating whether these four clusters were associated with different tumor characteristics, we found the *Porphyromonas* cluster 1 to be significantly enriched in rectal tumors (Fig. 2b), consistent with the presence of superoxide dismutase genes in *Porphyromonas* genomes possibly conferring tolerance to a more aerobic milieu in the rectum (Extended Data 5). The Clostridiales cluster 4 was significantly more prevalent in female CRC patients. All species clusters showed a slight tendency towards late-stage CRC (i.e. AJCC stages III and IV), but this was only significant for cluster 3. Associations with patient age and BMI were weaker and not significant (Extended Data 5). To rule out secondary effects due to differences in patient composition among studies, all of these tests were corrected for study effects (by blocking for 'study' and 'colonoscopy', see Methods). At the level of individual species, significant stage-specific enrichments could not be detected suggesting CRC-associated microbiome changes to be less dynamic during cancer progression than previously postulated [26], although fecal material may be less suitable to address this question than tissue samples.

Metagenomic CRC classification models

To establish metagenomic signatures for CRC detection across studies in face of geographic and technical heterogeneity, we developed multivariable statistical modeling workflows with rigorous external validation to avoid prevailing issues of overfitting and over-optimistic

reports of model accuracy [19]. As a precaution against over-optimistic evaluation, these workflows are independent of the above-described differential abundance analysis. Instead, LASSO (Least Absolute Shrinkage and Selection Operator) logistic regression classifiers were employed to select predictive microbial features and eliminated uninformative ones (Methods).

In a first step, we used abundance profiles from five studies including the 849 most abundant microbial species and assessed how well classifiers trained in cross validation (CV) on one study generalize in evaluations on the other four studies (study-to-study transfer of classifiers) (Fig. 3a). Within-study cross-validation performance, as quantified by the Area Under the Receiver Operating Characteristics (AUROC) curve, ranged between 0.69 and 0.92 and was generally maintained in study-to-study transfer (AUROC dropping by 0.07 ± 0.12 on average) with two notable exceptions. First, in line with the univariate analysis of species associations, CRC detection accuracy on the US study was lower than for the other studies, both in cross-validation and in study-to-study transfer. This could potentially be explained by the US fecal specimen, unlike in the other studies, being freeze-archived for >25 years before metagenomic sequencing [10]. Second, classifiers trained on the AT study did not generalize as well to the other studies, consistent with low study precision seen in univariate meta-analysis (Supplementary Fig. 2). Given the microbial co-occurrence clusters described above, we wondered whether species-species interactions would provide additional information relevant for CRC recognition that is not contained in species abundance profiles. However, nonlinear classifiers able to exploit such interactions did not yield significantly better accuracies (Supplementary Fig. 3, see also [27]), suggesting that the linear model based on few biomarkers (on average 17 species account for more than 80% of the classifier weight, Extended Data 6) is near optimal for CRC prediction.

We further assessed if including data from all but one study in model training improves prediction on the remaining held-out study (leave-one-study-out validation, LOSO). LOSO performance of species-level models ranged between 0.71 and 0.91, and when disregarding the US study as an outlier was 0.83 (Fig. 3b). This corresponds to a LOSO accuracy increase of 0.076 ± 0.03 compared to study-to-study transfer. These results suggest that one can expect a CRC detection accuracy 0.8 (AUROC) for any new CRC study using similarly generated metagenomic data. We moreover verified that metagenomic CRC classification models trained on species composition were not biased for clinical subgroups. With the exception of slightly more sensitive detection of late stage CRC (P = 0.03, mostly originating from the US study, Extended Data 7), we did not observe any classification bias by patient age, sex, BMI, or localization. Together this suggests that these metagenomic classifiers are unlikely to be strongly confounded by the clinical parameters recorded.

Several previous studies comparing microbiome changes across multiple diseases reported primarily general dysbiotic alterations and highlighted the need to examine the disease specificity of microbiome signatures [17, 19]. Therefore, we assessed false positive (FP) predictions of our metagenomic CRC classifiers on fecal metagenomes of type 2 diabetes [4, 5], Parkinson's disease [12], ulcerative colitis and Crohn's disease [6, 7] patients, reasoning that classifiers relying on biomarkers for general dysbiosis would yield an excess of FPs on these cohorts. However, our LOSO classification models calibrated to have a false-positive

rate (FPR) of 0.1 on CRC datasets in fact maintained similarly low FPRs on other disease datasets ranging from 0.09 to 0.13 (Fig. 3c). Interestingly, disease specificity of LOSO models was significantly improved over that observed for classifiers trained on a single study, indicating that inclusion of multiple studies in the training set of a classifier can substantially improve its specificity for a given disease.

Functional metagenomic signatures for CRC

As shotgun metagenomics data, in contrast to 16S rRNA gene amplicon data, allow for a direct analysis of the functional potential of the gut microbiome, we examined how predictive metabolic pathways and orthologous gene families differing in abundance between CRC patients and controls would be of CRC status. When applying the same classification workflow as above to eggNOG orthologous gene family abundances [28], CRC detection accuracy was very similar to that observed for taxonomic models (Fig. 3de). AUROC values ranged from 0.70 to 0.81 for study-to-study transfer (per-study averages, Fig. 3e) and from 0.78 to 0.89 in LOSO validation with a pattern of generalization across studies resembling that for taxonomic classifiers. The accuracy of functional signatures did not strongly depend on eggNOG as an annotation source, but was similar when based on other comprehensive functional databases, such as KEGG [29] (Extended Data 8). When using individual gene abundances from metagenomic gene catalogues as a classifier input [30], we observed higher within-study cross-validation AUROC values of 0.96 in all studies, but lower generalization to other studies (AUROC between 0.60 and 0.79) (Extended Data 8).

To explore changes in metabolic capacity of gut microbiomes from CRC patients more broadly, we quantified gut metabolic modules (defined in [31]) and subjected these to the same differential abundance analysis developed for microbial species. Gut metabolic modules with significantly higher abundance (FDR < 0.01, Wilcoxon test blocked for study and colonoscopy) in CRC metagenomes predominantly belonged to pathways for the degradation of amino acids, mucins (glycoproteins) and organic acids. This clear trend was accompanied by a depletion of genes from carbohydrate degradation modules (Fig. 4ab). Differences in all four high-level categories were highly significant (P < 1E-6 in all cases, blocked Wilcoxon tests) and consistent across studies (Fig. 4b). Overall these results establish a clear shift from dietary carbohydrate utilization in a healthy gut microbiome to amino acid degradation in CRC consistent with an earlier report based on a subset of the data [8]. Correlation analysis suggests that increased capacity for amino acid degradation is mostly contributed by CRC-associated Clostridiales (cf. cluster 4 in Fig. 2, Supplementary Fig. 4). About one half of these metagenomic pathway enrichments are also in agreement with independent metabolomics data suggesting increased availability of amino acids in epithelial cells or feces of CRC patients (Supplementary Table S5, [32-36]). While the observed pathway enrichments could potentially result from many factors, including unmeasured ones [13], they are consistent with established dietary risk factors for CRC, which include red and processed meat consumption [37] and low fiber intake [38].

The large metagenomic data set analyzed here allowed us to quantify the prevalence of gut microbial virulence and toxicity mechanisms thought to play a role in colorectal

carcinogenesis. Prominent examples include the Fusobacterium nucleatum adhesion protein A (encoded by the fadA gene), the Bacteroides fragilis enterotoxin (bft gene) and colibactin produced by some *Escherichia coli* strains (*pks* genomic island) [39, 40]. Moreover, intestinal *Clostridium* spp. are known to contribute to the conversion of primary to secondary bile acids using several metabolic pathways including 7a-dehydroxylation, encoded in the bai operon [41]. The products of this 7a-dehydroxylation pathway, deoxycholate and lithocholate, are known hepatotoxins associated with liver cancer [42] and hypothesized to also promote CRC [43]. Although intensely studied at a mechanistic level, these factors are not (well) represented in general databases that can be used for metagenome annotation (Supplementary Fig. 5). Thus, we built a targeted metagenome annotation workflow based on Hidden Markov Models to identify and quantify virulence factors and toxicity pathways of interest in CRC. Additionally, we used co-abundance clustering to infer operon completeness for factors encoded by multiple genes (Methods, Extended Data 9, Supplementary Fig. 5). While fadA, bft, the pks island and the bai operon were clearly detectable in deeply sequenced fecal metagenomes, they varied broadly with respect to abundance, significance and cross-study consistency of enrichment (Fig. 4c): fadA and *pks* were significantly enriched in CRC metagenomes (P = 5.3E-10 and 4.1E-4 respectively), whereas no significant abundance difference could be detected for *bft* in fecal metagenomes, despite reports on its enrichment in the mucosa of CRC patients [44], its carcinogenic effect in mouse models [45], and synergistic action with pks [46]. Our quantification of the bai operon showed a highly significant enrichment in CRC metagenomes (P = 1.6E-9) observed across all five studies (Fig. 4d) at an average abundance that exceeded fadA and pks copy numbers (Fig. 4c). Metagenome analysis indicated that at least four Clostridiales species (including the well characterized C. scindens and C. *hylemonae* [47, 48]) have a (near) complete 7α -dehydroxylation pathway contributing to the observed enrichment of bai operon copies (Extended Data 9). To validate this finding and further explore its value towards diagnostic application, we developed a targeted quantification assay for the *baiF* gene based on quantitative PCR (qPCR, see Methods). Quantification of *baiF* by qPCR using genomic DNA from 47 fecal samples of the DE study population was found to be similar to, yet more sensitive than by metagenomics (Fig. 4e). Gut microbial *baiF* copy numbers clearly distinguished CRC patients from controls (P =0.001) at an AUROC of 0.77, which in this subset of samples is surpassed by only a single species marker for CRC (Extended Data 9). Although consistent with increased deoxycholate metabolite levels reported for serum and stool samples of CRC patients [49], this finding does not imply 7a-dehydroxylation pathway activity. We therefore quantified *baiF* expression using RNA extracts from the same set of fecal samples, and found also transcript levels to be elevated in CRC patients (Fig. 4f). The observed weak correlation of baiF expression with genomic abundance (Fig. 4f) might be explained by dynamic transcriptional regulation [47] and bai expression in feces might not accurately reflect the tumor microenvironment. Taken together, these data suggest gut microbial metabolic markers to be meaningful and highly predictive of CRC status.

Validation of CRC signatures in independent study populations

Even though CRC classification accuracy for both species and functions were evaluated on independent data, we nonetheless sought to confirm it using two additional study

populations from Italy (IT1 and IT2, combined N = 61 CRC, N = 62 CTR, [27], see Methods, Table 1) and one from Japan (JP, N = 40 CRC, N = 40 CTR, see Methods, Table 1). The overlap of single species associations detected in the IT2 study and those from the meta-analysis was found to vary within the range seen for the other studies, whereas for IT1 and JP the overlap was slightly lower (cf. study precision in Supplementary Fig. 2, Extended Data 10). Nonetheless, the AUROC of LOSO classification models based on species ranged between 0.79 and 0.81 and that for the classifiers based on eggNOG from 0.71 to 0.92 (Fig. 5ab). We also validated CRC enrichment of *fadA*, *pks* and *bai* genes in these three study populations (Fig. 5c). Altogether these results highlight consistent alterations in the gut microbiome of CRC patients across eight study populations from seven countries in three continents.

Discussion

Through extensive and statistically rigorous validation, in which data from studies used for training is strictly separated from that for testing, our meta-analysis firmly establishes that gut microbial signatures are highly predictive of CRC (see also [27]). In particular metagenomic classifiers trained on species profiles from multiple studies maintained an AUROC of at least 0.8 in seven out of eight data sets and achieved an accuracy similar to the fecal occult blood test, a standard non-invasive clinical test for CRC (Supplementary Fig. 6, cf. [8]). These results thus suggest that polymicrobial CRC classifiers are globally applicable and can overcome technical and geographical study differences, which we found to generally impact observed microbiome composition more than the disease itself (Fig. 1c, Extended Data 1, 2). The generalization accuracy of classifiers across studies seen here is higher than that reported in 16S rRNA gene amplicon sequencing studies, which are characterized by even larger heterogeneity across studies [16, 18] (Supplementary Fig. 7).

Previous microbiome meta-analyses suggested that the majority of gut microbial taxa differing in any given case-control study reflect general dysbiosis rather than disease-specific alterations illustrating the difficulty of establishing disease-specific microbiome signatures [17, 19]. Here, by combining data across studies for training (LOSO), we were able to develop disease-specific signatures that maintained false positive control on diabetes and IBD metagenomes at a very similar level as for CRC (Fig. 3c) despite these diseases having shared effects on the gut microbiome [17, 50] and an increased comorbidity risk [51].

Although for diagnostic purposes, unresolved causality between microbial and host processes during CRC development are not a central concern, elucidating the underlying mechanisms would greatly enhance our understanding of colorectal tumorigenesis. Towards this goal, we developed both broad and targeted annotation workflows for functional metagenome analysis. First, we found functional signatures based on the abundances of orthologous groups of microbial genes to yield accuracies as high as taxonomic signatures (Fig. 3), which raises the hope for future improvements in metagenome annotation to translate into microbiome signature refinements. Second, by investigating potentially carcinogenic bacterial virulence and toxicity mechanisms taking a targeted metagenome annotation approach, we confirmed highly significant enrichments of the colibactin-producing *pks* gene cluster and the *Fusobacterium nucleatum* adhesin *FadA* in CRC

metagenomes (Fig. 4c). Our results support the clinical relevance of these factors adding to the experimental evidence for their carcinogenic potential [46, 52–54]. We further examined the bai operon, encoding enzymes that produce secondary bile acids via 7adehydroxylation, as an example of toxic host-microbial co-metabolism (see [27] for another intriguing example). While a dehydroxylated bile acids are established liver carcinogens [42], their contribution to CRC is less clear [43]. Here, we have, for the first time, shown bai to be highly enriched in stool from CRC patients (Fig. 4cd) and confirmed this finding at both the genomic and the transcriptomic level using qPCR (Fig. 4ef). As bai enrichment (and expression) is likely a consequence of a diet rich in fat and meat [55], it is intriguing to explore whether bai could be used as a surrogate microbiome marker for such difficult-tomeasure dietary CRC risk factors. To further unravel the molecular underpinning of these dietary CRC risk factors, molecular pathological epidemiology studies that investigate the mucosal microbiome as part of the tumor microenvironment, hold great potential [56, 57]. However, they will require more comprehensive diet questionnaires, medical records, and molecular tumor characterizations than are available for the study populations analyzed here. In this context, carcinogens possibly contained in the virome also warrant further investigation [58, 59], but for this goal, metagenomic data needs to be generated with protocols optimized for virus enrichment [60].

Taken together, our results and those by Thomas, Manghi et al. [27], strongly support the promise of microbiome-based CRC diagnostics. Both taxonomic and metabolic gut microbial marker genes established in these meta-analyses could form the basis of future diagnostic assays that are sufficiently robust, sensitive, and cost-effective for clinical application. The targeted qPCR-based quantification of the *baiF* gene is a first step in this direction. Our metagenomic analysis of this and other virulence and toxicity markers bridge to existing mechanistic work in preclinical models and could enable future work aiming to precisely determine the contribution of gut microbiota to CRC development.

Data and Code Availability

The raw sequencing data for the samples in the DE study that had not been published before (see Methods), are made available in the European Nucleotide Archive (ENA) under the study identifier PRJEB27928. Metadata for these samples are available as Supplementary Table S6.

For the other studies included here, the raw sequencing data can be found under the following ENA identifiers: PRJEB10878 for [11], PRJEB12449 for [10], ERP008729 for [9], and ERP005534 for [8]. The independent validation cohorts can be found in SRA under the identifier SRP136711 for [27] and in the DDBJ database under the ID DRA006684.

Filtered taxonomic and functional profiles used as input for the statistical modeling pipeline are available in Supplementary Data 1.

The code and all analysis results can be found under https://github.com/zellerlab/crc_meta.

Methods

Study inclusion and data acquisition

We used PubMed to search for studies that published fecal shotgun metagenomic data of human colorectal cancer patients and healthy controls. The search term, all hits, and the justification for exclusion or inclusion are available in Supplementary Table S1. Raw fastq files were downloaded for the four included studies from the European Nucleotide Archive, using the following ENA identifiers: PRJEB10878 for [11], PRJEB12449 for [10], ERP008729 for [9], and ERP005534 for [8].

DE study recruitment and sequencing

The German (DE) study population data consist of 60 fecal CRC metagenomes, 38 of which were sequenced and published in [8] under ENA accession ERP005534. The fecal metagenomes from additional 22 CRC patients recruited for the same ColoCare study (DKFZ, Heidelberg, [61, 62]) were sequenced later as part of this work. All fecal samples were collected after colonoscopy. Sixty gender- and age-matched participants of the PRÄVENT study run by the same clinical investigators were included as healthy controls; as these were not subjected to colonoscopy, the presence of undiagnosed colorectal carcinomas cannot be completely ruled out but is expected to be unlikely due to low prevalence of preclinical CRC in the general population [63].

Written informed consent was obtained from all additional 22 CRC patients and 60 controls. The study protocol was approved by the institutional review board (EMBL Bioethics Internal Advisory Board) and the ethics committee of the Medical Faculty at the University of Heidelberg. The study is in agreement with the WMA Declaration of Helsinki and the Department of Health and Human Services Belmont Report.

Genomic DNA was extracted from the fecal samples (preserved in RNALater) and libraries were prepared as previously described [8]. Whole-genome shotgun sequencing was performed by using Illumina HiSeq 2000 / 2500 / 4000 (Illumina, San Diego, USA) platforms at the Genomics Core Facility, European Molecular Biology Laboratory, Heidelberg.

Independent validation cohorts

During the revision of this manuscript, we included three independent study populations for external validation. Two of them were recruited in Italy (IT1 and IT2) with informed consent from all participants and ethical approval by the Ethics committee of Azienda Ospedaliera of Alessandria and that of the European Institute of Oncology of Milan. Shotgun fecal metagenomic data was generated as described in [27].

The third study population was recruited in Japan (JP) with informed consent and ethical approval of the institutional review boards of the National Cancer Center Japan - Research Institute and the Tokyo Institute of Technology. DNA was extracted from frozen fecal samples using a GNOME DNA Isolation Kit (MP Biomedicals, Santa Ana, CA) with an additional bead-beating step as previously described [64]. DNA quality was assessed with an

Agilent 4200 TapeStation (Agilent Technologies, Santa Clara CA). After final precipitation, the DNA samples were resuspended in TE buffer and stored at -80°C before further analysis. Sequencing libraries were generated with the Nextera XT DNA Sample Preparation Kit (Illumina, San Diego, CA). Library quality was confirmed with an Agilent 4200 TapeStation. Whole-genome shotgun sequencing was carried out on the HiSeq2500 platform (Illumina). All samples were paired-end sequenced with a 150-bp read length to a targeted data set size of 5.0 Gb.

Taxonomic profiling and data preprocessing

The metagenomic samples were quality controlled using MOCAT2's -rtf procedure, which is based on the 'solexaqa' algorithm [23]. In particular, reads that map with at least 95% sequence identity and alignment length of at least 45 bp to the human genome hg19 were removed. In a second step, taxonomic profiles were generated with the mOTU profiler version 2.0.0 ([22, 65, 66] – see motu-tool.org and GitHub version tag 2.0.0) using the following parameters: -1 75, -g 2 and -c. Briefly, this profiler is based on ten universal single-copy marker-gene families (COG0012, COG0016, COG0018, COG0172, COG0215, COG0495, COG0525, COG0533, COG0541 and COG0552) [66]. These marker-genes were extracted from >25,000 reference genomes and >3,000 metagenomic samples allowing to profile prokaryotic species with a sequenced reference genome (ref-mOTUs) and ones without (meta-mOTUs). The read count for a mOTU was calculated as median of the read count of the genes that belonged to that mOTU.

mOTU profiles were first converted to relative abundances to account for library size. Then, profiles were filtered to focus on a set of species that are confidently detectable in multiple studies. Specifically, microbial species that did not exceed a maximum relative abundance of 1E-03 in at least 3 of the studies were excluded from further analysis, together with the fraction of unmapped metagenomic reads.

Functional metagenome profiling and data preprocessing

High-quality reads (same quality filtering as for taxonomic profiling) were aligned against a combined database (IGChg38 hereafter) consisting of the hg38 release of the human reference genome and the integrated gene catalog (IGC) containing 9.9 million nonredundant microbial genes [30] using BWA mem [67] (Version: 0.7.15-r1140) with default parameters. The purpose of adding the human genome to the reference database was to filter out reads that mapped as well or better to some human sequence than to any bacterial gene. Alignments were computed separately for paired-end and single read libraries (single reads could result from read pairs where one read was filtered out in the quality filtering procedure described above). Alignments were then filtered to only retain those longer than 50bp with >95% sequence identity. Then the highest scoring alignment(s) was/were kept for each read. As IGChg38 is a database of predominantly genes and not genomes, there will be a substantial proportion of read-pairs where one end maps within the gene while the other end does not – it either maps to an adjacent gene or remains unmapped due to intergenic regions not contained in the database. Therefore, we counted a whole read-pair aligning to a gene when (i) both ends from a read pair map to the same gene, (ii) only one end from a read-pair maps to the gene, or (iii) a read from the single read library maps to the gene. We then

counted only the read-pairs that map uniquely to one gene in the IGC, thus excluding ambiguous read pairs mapping with similarly high scores to multiple genes in the database. For a given metagenomic sample, we further normalized the abundance of each IGC gene by the length of that gene. We then estimated relative abundance of IGC genes by dividing gene abundances by the total abundance of all genes in IGC (excluding the human chromosomes).

Because metagenomes from CRC patients were not included when the IGC was constructed, we analyzed how well CRC-associated species as identified in this meta-analysis were represented in the IGC. Using a phylogenetic marker gene (COG0533), which is also used by the species profiling workflow on which the meta-analysis is based, for 24 out of the 29 core CRC-associated species we found a match in the IGC with at least 90% nucleotide identity, indicating that a sequence from the same species (above 93.1% identity) or a slightly more distant relative is present in the IGC (Supplementary Fig. 8).

The relative abundance of eggNOG orthologous groups [28] was estimated by summing relative abundances of genes annotated to belong to the same eggNOG orthologous group as of the most recent annotations provided by MOCAT2 [23]. To obtain KEGG orthologous groups (KO) and pathway abundances, we applied the same procedure, but using KEGG annotations for IGC provided by MOCAT2 [29].

Overview over statistical analyses

For univariate association testing between the abundances of microbial taxa or gene functions we used nonparametric tests throughout; all of these were two-sided Wilcoxon tests except were otherwise noted. To account for potential confounding and heterogeneity between data sets we employed a stratified version of the Wilcoxon test [24] (see below for details). ANOVA was conducted on rank-transformed data. Significance of binary cooccurrence patterns was assessed using (stratified) Cochrane-Mantel-Haenszel tests.

Multivariable analysis was done with strict separation between training and test data. This importantly also pertained to feature selection, which was either done via the LASSO [68] or by nested cross-validation procedures to avoid overoptimistic performance assessment [69] (see below for details). All samples included in this meta-analysis came from distinct individuals to ensure that generalization across subjects – rather than across timepoints within a given subject – is assessed.

Confounder analysis

To quantify the effect of potential confounding factors relative to that of CRC on single microbial species, we used an ANOVA-type analysis. The total variance within the abundance of a given microbial species was compared to the variance explained by disease status and the variance explained by the confounding factor akin to a linear model including both CRC status and confounding factor as explanatory variables for species abundance. Variance calculations were performed on ranks in order to account for non-Gaussian distribution of microbiome abundance data. Potential confounders with continuous values were transformed into categorical data either as quartiles or for the case of body mass index (BMI) into lean/obese/overweight according to conventional cutoffs (lean: < 25, obese: 25 - 30, overweight: > 30).

Univariate meta-analysis for the identification of CRC-associated gut microbial species

Significance of differential abundance was tested on a per-species basis using a blocked Wilcoxon test implemented in the R coin package [24]. Informed by the results of the preceding confounder analysis, we blocked for `study` and additionally `colonoscopy` in the CN study. Within this framework, significance is tested against a conditional null distribution derived from permutations of the observed data. Notably, permutations are performed within each block in order to control for variations in block size and composition. To adjust for multiple hypothesis testing, P-values were adjusted using the false-discovery rate (FDR) method [70].

As nonparametric effect size measures we used the area under the ROC curve (AUROC) with permutation-based confidence intervals computed using the pROC package in R [71]. We further developed a generalization of the (logarithmic) fold change that is widely used for other types of read abundance data. This generalization is designed to have better resolution for sparse microbiome profiles (where 0 entries can render median-based fold change estimates uninformative for the large portion of species with a prevalence below 0.5). The generalized fold change (gFC) is computed as mean difference in a set of pre-defined quantiles of the logarithmic CTR and CRC distributions (see Extended Data 3 for further details; we used quantiles ranging from 0.1 to 0.9 in increments of 0.1).

For the retrospective analysis of study precision and recall for detecting microbial species associations from the meta-analysis, the true set was defined as the species which were associated at a given FDR in the meta-analysis. Then, we checked how well this set of species would be recovered using the single-study significance as determined by the Wilcoxon test. Study precision corresponds to the proportion of meta-analysis significant species among those detected as significant in a single study. Similarly, recall (or sensitivity) corresponds to the proportion of species out of the true set of meta-analysis significant species that were recovered in a given study.

Species co-occurrence and cluster analysis in CRC metagenomes

For the analysis of gut bacterial species co-occurring in CRC microbiomes, relative abundances of the core set of associated species (excluding the CRC-depleted *Clostridiales* meta-mOTU [1296]) were discretized into binary values to determine whether a CRC (metagenomic) sample is "positive" or "negative" for a given microbial marker. To normalize for differences in prevalence (and therefore specificity) of these markers we adjusted the threshold value, above which a sample is labeled "positive" based on the abundance in healthy controls. For each microbial species, the 95th percentile in healthy controls was used as threshold, which effectively results in adjusting the per-marker false positive rate to 0.05. Based on the binarized species-by-sample matrix, species were then clustered using the Jaccard dissimilarity as implemented in the vegan package in R [72]. Associations between species clusters and meta-variables were tested as 2-by-n (where n is the number of categories in the meta-variable tested) contingency tables using a Cochrane-Mantel-Haenszel test with study as blocking factor as implemented in the coin package [24].

Multivariable statistical modeling workflow and model evaluation

As a main goal of our work is to assess the generalization accuracy of microbiome-based CRC classifiers across technical and geographic differences in patient populations, we extensively validated classification models across studies taking the following two approaches.

In *study-to-study transfer* validation, metagenomic classifiers were trained on a single study and their performance externally assessed on all other studies (off-diagonal cells in Fig. 3ac). Effectively we implemented a nested cross validation procedure on the training study to compute within-study accuracy (cells on the diagonal in Fig. 3ac) and tune the model hyperparameters.

In *leave-one-study-out* (LOSO) validation, data from one study was set aside as an external validation set, while the data from the remaining 4 studies was pooled as a training set on which we implemented the same nested cross validation procedure as for study-to-study transfer (see [19] for a more detailed description of LOSO).

Data preprocessing, model building, and model evaluation was performed using the SIAMCAT R package (https://bioconductor.org/packages/SIAMCAT, version 1.1.0).

Preprocessing of taxonomic abundance profiles for statistical modeling

Relative abundances were first filtered to remove markers with low overall abundance and no variance (an artifact for single-study data arising from the joint data filtering described above), log-transformed (after adding a pseudo-count of 1E-05 to avoid non-finite values resulting from log(0), [73]) and finally standardized as z-scores. Data were split into training and test set for 10 times repeated 10-fold stratified cross validation (balancing class proportions across folds). For each split, a L1-regularized (LASSO) logistic regression model [68] was trained on the training set, which was then used to predict the test set. The lambda parameter, i.e. regularization strength was selected for each model to maximize the area under the precision recall curve under the constraint that the model contained at least 5 non-zero coefficients. Models were then evaluated by calculating the area under the Receiver Operating Characteristics curve (AUROC) based on the posterior probability for the CRC class.

In model transfer to a hold-out study, the holdout data were normalized for comparability in the same way as the training dataset by using the frozen normalization function in SIAMCAT, which retains the same features and re-uses the same normalization parameters (e.g. the mean of a feature for z-score standardization). Then, all 100 models derived from the cross validation on the training dataset (10 times repeated 10-fold CV) were applied to the holdout dataset and predictions were averaged across all models.

In the LOSO setting, data from the four training studies were jointly processed as a single dataset in the same way as described above using 10 times repeated 10-fold stratified cross validation.

Preprocessing of functional abundance profiles

Functional profiles, such as eggNOG gene family or KEGG module abundance profiles were preprocessed as described above for species profiles, but using 1E-06 as maximum abundance cutoff and 1E-09 as a pseudo-count during log transformation. Since these abundance tables contained several thousand input features we implemented an additional feature selection step, which was nested properly into the cross-validation procedures as described above. This nested approach is crucial to avoid over-optimistically biased performance estimates ([74], Chapter 7.10). Specifically, features were filtered inside each training fold (without using any information from the test fold) by selecting the 1600 features with highest single-feature AUROC values (for features depleted in CRC, 1 - AUROC was used for feature selection).

Preprocessing of gene abundance profiles

To ascertain the predictive power of a classifiers based on IGC gene abundances [30] we applied a series of filters to the abundance tables to reduce the number of genes that would be the input of the LASSO modelling. These filters where applied once on a per-study level and once in a leave-one-study-out (LOSO) mode, where they were applied jointly to all studies in the training set, with the remaining one being held out for external validation.

The following filters were applied in this order:

- 1. All genes with 0 abundance in 15% of samples (regardless of CRC status) were discarded.
- 2. The remaining data was discretized using the equal frequencies method implemented in the 'discretize' function of the sideChannelAttack R package (version 1.0-6) as a preparation to the minimal-redundancy-maximal-relevance (mRMR) algorithm [75].
- **3.** As a feature selection procedure, mRMR (code version from 20 April 2009 downloaded from http://home.penglab.com/proj/mRMR/ on 3 Dec 2016) was run on the gene abundance table to retain the 100 top genes as output.

LASSO models were then built on log10-transformed abundances (pseudo-count of 10E-09, centered and scaled) of the sets of 100 top genes returned by mRMR. The whole process was repeated 10 times in a 5-fold stratified cross-validation scheme to allow for an estimation of the confidence of the AUROCs of the resulting models. We used the LiblineaR package (version 2.10-8) to build the LASSO models in R and tested a sequence of 20 cost parameters (equivalent or the lambda parameter controlling regularization strength) evenly spaced from 0.001^2 to 0.2^2 . The cost parameter was selected to maximize the AUROC within the training set.

External evaluation of disease-specificity of the metagenomic classifiers

To assess how disease-specific the predictions of the CRC models are, we applied these to data from case-control studies investigating other human diseases. Fecal metagenomic data of patients with Parkinson's disease [12], type 2 diabetes [4, 5], and inflammatory bowel disease [6, 7] were taxonomically profiled as described above. The parameters for quality

control with MOCAT2 and for the mOTU profiler were the same as described above, except for the data from [6], where we used -1 50 (to set the threshold for minimum alignment length to 50) as the read length is shorter (average read length 71) compared to the other more recently generated Illumina shotgun metagenomic data.

Relative abundance data were treated exactly as another holdout dataset for each model, i.e. applying the frozen normalization prediction routines as described above. For each CRC model applied to the external datasets, a cutoff on its prediction output was adjusted to yield a false positive rate (FPR) of 0.1 on the controls of its respective (CRC) training set. Subsequently its FPR on metagenomes from patients suffering from the above-mentioned (non-CRC) conditions was assessed to evaluate its disease specificity. The rationale behind this is that a metagenomic classifier recognizing general features of dysbiosis would be expected to predict CRC patients and those suffering from other conditions at a similar rate; such a classifier would thus in the above-described evaluation display a much higher FPR than on the controls of its training set. In contrast maintaining a low FPR in this evaluation indicates that the classification model is based on CRC-specific features rather than hallmarks of general dysbiosis or nonspecific inflammation.

Functional profiling of gut metabolic modules (GMMs)

Gut metabolic modules were computed as originally proposed [31], using the KEGG KO profiles based on the IGC (see Functional metagenome profiling above) as input. Statistical analysis and generalized fold change calculations were performed analogously to species profiles (see above). Gut metabolic modules were summarized across functional groups (e.g. amino acid degradation) as geometric mean of all modules within the respective group.

Targeted functional analysis of virulence and toxicity pathways of potential relevance in CRC

To investigate toxins and virulence mechanisms that have previously been implicated with CRC [40], we constructed for each gene belonging to the respective virulence or toxicity pathway a hidden Markov model (HMM). Each HMM was built from a multiple sequence alignment generated by MUSCLE [76], containing the respective reference sequences and close homologs identified using PSI-Blast [77]. Multiple sequence alignments are available together with the code for this paper (https://github.com/zellerlab/crc_meta). Then, we screened the IGC metagenomic gene catalogue [30] with each HMM using the HMMER software (version 3.1b2) [78]. Genes with an E-value below 1E-10 were filtered for uniqueness, since in some cases the HMMs would call different regions in the same gene. For single gene virulence factors (i.e. *fadA* and *bft*), potential IGC hits were aligned against the reference sequence using the Needleman-Wunsch algorithm in the EMBOSS package [79]. Hits were then filtered based on percentage of sequence identity (cutoff: 40%) and sequence similarly to the species relative abundance profiles based on maximum relative abundance (cutoff: 1E-07) in order to exclude genes with limited relevance. Statistical analysis was performed on the sum of all genes.

For virulence pathways containing more than one gene, the IGC hits of each functional group within the pathway were aligned against the respective reference sequence and filtered

for percentage of sequence identity and maximum abundance. Then, all hits were clustered based on the Pearson correlation of the log-abundances across all samples using the Ward algorithm as implemented in the *hclust* function in R. The gene clusters were filtered based on operon completeness (how many genes of the operon were present in the cluster) and average correlation within the cluster (Extended Data 9). For statistical analysis, the genes in the selected gene clusters were summed up within each group or all together for the overall analysis.

Quantitative PCR for baiF

Real-time quantitative PCR to quantify the abundance and expression of *baiF* was performed on a subset of samples in the DE cohort (20 control and 24 colorectal cancer samples, see Supplementary Table S6). For these samples, DNA and RNA extraction was done with the Allprep PowerFecal DNA/RNA kit (Qiagen, Cat No: 80244) with additional RNAse and DNAse digestion steps, respectively, as described by the manufacturer. DNA and RNA concentrations were determined by Qubit Fluorometer (Invitrogen) and quality control of all RNA samples was done using an Agilent 2100 Bioanalyzer in combination with RNA 6000 Nano and Pico LabChip kits.

First-strand cDNA was synthesized by SuperScript IV VILO Master Mix with ezDNAse enzyme and random hexamer primers (Invitrogen, catalogue number 11766500) as recommended by the manufacturer. Reaction were performed as described in the protocol with one minor change of temperature (incubation for the reverse transcription step at 55°C).

To quantify *baiF* relative to the total bacterial RNA/DNA in a sample, qPCR was performed in triplicates for 16S rRNA and the *baiF* genes, using both cDNA and genomic DNA (gDNA) as template. We used the following primers for *baiF*: TTCAGYTTCTACACCTG (forward), GGTTRTCCATRCCGAACAGCG (reverse), and standard primers F515 and R806 for 16S [80]. RT-PCR reactions were prepared with a final primer concentration of 0.5 μ M, including 5 ng of genomic DNA or 10 ng of cDNA in 20 μ l final reaction volume, and reactions were performed with SYBR Green qPCR mix on StepOne Real-Time PCR system (Thermo Fisler Scientific). Cycling conditions were as follows; initial denaturation of 95°C for 10 min, then 40 cycles of denaturing at 95°C for 15 s, annealing at 60°C for 60 s followed by melt curve analysis.

Delta-Ct values were calculated as difference between *baiF* and 16S Ct values. Significance of the comparison between control and colorectal cancer samples was tested on the delta-Ct values using a one-sided Wilcoxon test as a confirmation of metagenomic enrichment.



Extended Data 1.

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Extended Data 2.

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Extended Data 3.

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Extended Data 5.

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Extended Data 6.

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Extended Data 7.



Extended Data 8.

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Extended Data 9.

a meta	FR	AT	CN	US	DE	IT1	IT2	JP	b	
Parvimonas micra [1145]	20	9	1	70	1	84	2	1	1 00	
Gemella morbillorum [4513]	22	1	2	110	10	131	1	5	— IT1	
Peptostreptococcus stomatis [4614]	45	37	3	19	3	191	27	2	— IT2	
F. nucleatum subsp. animalis [0776]	3	9	13	7	7	84	23	24	JP	
unknown <i>Dialister</i> [5867] 1	1	17	8	6	4	677	11	36	0.75	2
unknown <i>Porphyromonas</i> [7656] 6	30	4	283	18	4	213	618	377		leta
Solobacterium moorei [0531] 7	79	204	5	303	2	109	15	37	<u>.</u>	님
Porphyromonas uenonis [4616] 8	35	14	557	8	10	459	67	27		RO
Clostridium symbiosum [1475] 8	30	125	8	20	14	45	267	82		.00
unknown <i>Clostridiales</i> [6105] 10	63	119	4	279	8	292	761	219	" "hurl	01
Hungatella hathewayi [0882] 11	2	132	17	114	32	80	365	74	0.25	
Prevotella intermedia [0515] 12	5	32	15	204	121	526	67	27	Ma manager and	
Porphyromonas somerae [2101] 13	5	7	113	438	64	68	142	690		
Porphyromonas asaccharolytica [1517] 14	22	78	128	119	8	20	156	60	0.00 -	
F. nucleatum subsp. nucleatum [0777] 15	5	138	34	29	72	311	372	32		
Parvimonas sp. [4961] 16	63	50	23	169	57	311	67	9	1.00	
Prevotella nigrescens [0276] 17	83	138	71	84	14	601	123	87		
unknown <i>Porphyromonas</i> [5431] 18	97	23	113	68	26	42	209	96		
Ruminococcus torques [1376] 19	184	24	33	123	54	137	530	121	0.75	
F. nucleatum subsp. vincentii [0754] 20	5	300	66	62	20	550	156	26		Met
Fusobacterium sp. oral taxon 370 [1403] 21	5	181	19	141	121	217	372	96		a-F
unknown <i>Peptostreptococcaceae</i> [5742] 22	223	167	6	349	52	68	19	17		DR
Anaerococcus obesiensis/vaginalis [0429] 23	105	7	266	204	98	311	51	225		Ē
unknown <i>Anaerotruncus</i> [6835] 24	22	38	283	313	33	489	348	187		05
Porphyromonas uenonis [2102] 25	146	50	146	49	62	304	688	27		
unknown <i>Clostridiales</i> [7553] 26	35	682	34	1	46	213	574	89	0.25 MM https://www.	
unknown <i>Porphyromonas</i> [6490] 27	191	108	229	8	36	141	372	60		
Clostridium boltae/clostridioforme [0886] 28	109	220	24	119	40	548	595	487	and the second s	
Subdoligranulum sp. [4738] 29	43	817	11	178	26	399	179	10	0.00	
									0.00 0.25 0.50 0.75 1.00	

Extended Data 10.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Despite study differences, meta-analysis identifies a core set of gut microbes strongly associated with CRC.

(a) Meta-analysis significance of gut microbial species derived from blocked Wilcoxon tests (n=574 independent observations) is given by bar height (false discovery rate, FDR, of 0.05). (b) Underneath, species-level significance as computed by two-sided Wilcoxon test (FDR-corrected P-value) and generalized fold change (Methods) within individual studies are displayed as heatmaps in gray and color, respectively (see color bars and Table 1 for details on studies included). Species are ordered by meta-analysis significance and direction

of change. (c) For a core of highly significant species (meta-analysis FDR 1E-5), association strength is quantified by the area under the Receiver Operating Characteristics curve (AUROC) across individual studies (color coded diamonds) and 95% confidence intervals are indicated by gray lines. Family-level taxonomic information is color-coded above species names (numbers in brackets are mOTU species identifiers, see Methods). (d) Variance explained by disease status (CRC vs controls) is plotted against variance explained by study effects for individual microbial species with dot size proportional to abundance (Methods); core microbial markers are highlighted in red. *F. nucleatum – Fusobacterium nucleatum*.





(a) The heatmap shows for all CRC patients (n=285 independent samples) if the respective sample is positive for each of the core set of microbial marker species (see Methods for adjustment of positivity threshold). Samples are ordered according to the sum of positive markers and marker species are clustered based on Jaccard similarity of positive samples, resulting in four clusters (Methods). Barplots in (b), (c), and (d) show the fraction of CRC samples that are positive for marker species clusters (defined as the union of positive marker

species) broken down by patient subgroups based on differences in tumor location, sex, or CRC stage, respectively. Statistically significant associations between CRC subgroups and marker species clusters were identified using the Cochran–Mantel–Haenszel test blocked for study effects and are indicated above bars (P < 0.1).



Figure 3. Both taxonomic and functional metagenomic classification models generalize across studies in particular when trained on data from multiple studies.

CRC classification accuracy resulting from cross validation within each study (gray boxes along diagonal) and study-to-study model transfer (external validations off diagonal) as measured by AUROC for classifiers trained on (**a**) species and (**d**) eggNOG gene family abundance profiles. The last column depicts the average AUROC across external validations. Classification accuracy, as evaluated by AUROC on a held-out study, improves if taxonomic (**b**) or functional (**e**) data from all other studies are combined for training (leave-one-study-

out, LOSO validation) relative to models trained on data from a single study (study-to-study transfer, average and standard deviation shown). Bar height for study-to-study transfer corresponds to the average of four classifiers (error bars indicate standard deviation, n=4). (c) Combining training data across studies substantially improves CRC specificity of the (LOSO) classification models relative to models trained on data from a single study (depicted by bar color, as in (c) and (d)) as assessed by the false positive rate (FPR) on fecal samples from patients with other conditions (see legend). Bar height for study-to-study transfer corresponds to the average FPR across classifiers (n=5) with error bars indicating the standard deviation of FPR values observed.

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Figure 4. Meta-analysis identifies consistent functional changes in CRC metagenomes.

(a) Meta-analysis significance of gut metabolic modules derived from blocked Wilcoxon tests (n=574 independent samples) is indicated by bar height (top panel, FDR of 0.01). Underneath, the generalized fold change (Methods) for gut metabolic modules [31] within individual studies is displayed as heatmap (see color key below (b)). Metabolic modules are ordered by significance and direction of change. A higher-level classification of the modules is color-coded below the heatmap for the four most common categories (colors as in (b), white indicating other classes). (b) Normalized log abundances for these selected functional

categories is compared between controls (CTR) and colorectal cancer cases (CRC). Abundances are summarized as geometric mean of all modules in the respective category and statistical significance determined using blocked Wilcoxon tests (n=574 independent samples, see Methods). (c) Normalized log abundances for virulence factors and toxins compared between metagenomes of controls (CTR) and colorectal cancer cases (CRC) (significant differences P < 0.05 were determined by blocked Wilcoxon test, n=574 independent samples, see Methods for gene identification and quantification in metagenomes; fadA: gene encoding Fusobacterium nucleatum adhesion protein A, bft: gene encoding Bacteroides fragilis enterotoxin, pks: genomic island in Escherichia coli encoding enzymes for the production of genotoxic colibactin, and bai: bile acid inducible operon present in some Clostridiales species encoding bile acid converting enzymes). (d) Metaanalysis significance (uncorrected P-value) as determined by blocked Wilcoxon tests (n=574 independent samples) and generalized fold change within individual studies are displayed as bars and heatmap, respectively, for the genes contained in the *bai* operon. Due to high sequence similarity to *baiF*, *baiK* was not independently detectable with our approach. (e) Metagenomic quantification of *baiF*(metag. ab. – normalized relative abundance) is plotted against qPCR quantification in genomic DNA (gDNA) extracted from a subset of DE samples (n=47), with Pearson correlation (r) indicated (see Methods). (f) Expression of *baiF* determined via qPCR on reverse-transcribed RNA from the same samples in contrast to genomic DNA (as in e). The boxplots on the side of (e), (f) show the difference between cancer (CRC) and control (CTR) samples in the respective qPCR quantification (P-values on top were computed using a one-sided Wilcoxon test). All boxplots show interquartile ranges (IQR) as boxes with the median as a black horizontal line and whiskers extending up to the most extreme points within 1.5-fold IQR.

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Figure 5. Meta-analysis results are validated in three independent study populations

CRC classification accuracy for independent datasets, two from Italy and one from Japan (see Supplementary Table S2), is indicated by bar height for single study (white) and leaveone-study-out (grey) models using either (**a**) species or (**b**) eggNOG gene family abundance profiles (cf. Fig. 3). Bar height for single study models corresponds to the average of five classifiers (error bars indicate standard deviation, n=5). (**c**) Normalized log abundances for virulence factors and toxins (cf. Figure 4c) compared between controls (CTR) and colorectal cancer cases (CRC). P-values were determined by blocked, one-sided Wilcoxon tests (n=193

independent samples). Boxes represent interquartile ranges (IQR) with the median as a black horizontal line and whiskers extending up to the most extreme points within 1.5-fold IQR.

Table 1

Fecal metagenomic studies of colorectal cancer included in this meta-analysis.

See Methods for inclusion criteria and Supplementary Table S2 for extended meta-data. For a detailed description of patient recruitment and data generation for the DE study, see Methods. The data for 38 samples from the DE study had been published previously as part of an independent validation cohort in [8].

Country Code	Reference	No. of cases	No. of controls					
FR	Zeller et al., 2014 [8]	53	61					
AT	Feng et al., 2015 [9]	46	63					
CN	Yu et al., 2017 [11]	74	54					
US	Vogtmann et al., 2016 [10]	52	52					
DE	this study	60	60					
External validation cohorts								
IT1	[27]	29	24					
IT2	[27]	32	28					
JP	Courtesy of T. Yamada et al.	40	40					