

An integrated catalog of reference genes in the human gut microbiome

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Many analyses of the human gut microbiome depend on a catalog of reference genes. Existing catalogs for the human gut microbiome are based on samples from single cohorts or on reference genomes or protein sequences, which limits coverage of global microbiome diversity. Here we combined 249 newly sequenced samples of the Metagenomics of the Human Intestinal Tract (MetaHit) project with 1,018 previously sequenced samples to create a cohort from three continents that is at least threefold larger than cohorts used for previous gene catalogs. From this we established the integrated gene catalog (IGC) comprising 9,879,896 genes. The catalog includes close-to-complete sets of genes for most gut microbes, which are also of considerably higher quality than in previous catalogs. Analyses of a group of samples from Chinese and Danish individuals using the catalog revealed country-specific gut microbial signatures. This expanded catalog should facilitate quantitative characterization of metagenomic, metatranscriptomic and metaproteomic data from the gut microbiome to understand its variation across populations in human health and disease.

The ensemble of microorganisms in our gut, referred to as the human gut microbiota, is known to be important for human physiology and disease in the gut and beyond¹. However, our knowledge of the genetic and functional diversity in gut microbes is far from complete. Increasing numbers of fecal samples are being analyzed by targeted 16S rRNA gene pyrosequencing and to a lesser extent by metagenomic shotgun sequencing, because of the higher costs and more complex data analysis associated with the latter. Metagenomic assembly of short sequencing reads enables functional insights and is a more convenient and unbiased way of obtaining genomic information for environmental microbes, compared to culture-based or single-cell methods. However, data from different studies are scattered (most notably in the MetaHIT² and the Human Microbiome Project (HMP)³ gene catalogs), and there has been no comprehensive and uniformly processed database that can represent the human gut

microbiota around the world. With the increasing amount of sequencing data, it is also not clear at what pace the number of species and genes discovered in the gut microbiome will continue to grow, and to what extent our current sampling and data analyses capture common and rare entities in the gut microbiota.

Catalogs of reference genes in the human gut microbiome are crucial for functional metagenomic analyses². Sequencing reads can be mapped to the catalog to profile the species and gene content of a sample; genes with co-varying abundance levels can be clustered to reveal disease markers in metagenome-wide association studies^{4–7}; analyses of gene content might guide isolation of strains from fecal samples and document the strains' genomic information in the original habitat before possible changes during cultivation; and as metatranscriptomics^{8,9} and metaproteomics¹⁰ become more common, a gene catalog would greatly facilitate analyses of RNA or protein data.

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The MetaHIT² and the HMP³ gene catalogs, based on 124 samples from individuals in European countries (here referred to as 'European samples') and 136 samples from individuals in the United States ('American samples'), respectively, have limited representation and might contain partial or chimeric genes that could be extended or eliminated with more sequencing data and state-of-the-art processing algorithms.

In this study, we established a catalog of the human gut microbial genes by processing 249 newly sequenced samples and 1,018 published samples from MetaHIT^{2,6,7}, HMP³ and a large diabetes study from China⁴, as well as 511 sequenced genomes of gut-related bacteria and archaea. This nonredundant reference catalog of 9,879,896 genes is freely accessible through our website (http://meta.genomics.cn) and the data are deposited in the GigaScience Database¹¹. Beside providing an expanded resource for future analyses, study of the catalog suggests that we may have reached saturated coverage of core gene content and functions, but rare genes will continue to be discovered with increased sampling. We also demonstrate discovery of population-specific characteristics of gut microbiota using the catalog.

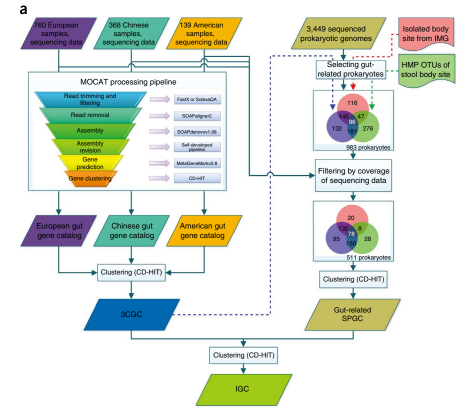
human gut microbiome, we integrated the European-sample microbial gene catalog with data from 368 Chinese samples⁴ and 139 American samples³ (Fig. 1a and Supplementary Table 1). Because we used a standardized and automated workflow that has been shown previously to improve the quality of assembly, gene prediction and redundancy removal^{12,13} (Online Methods), genes from the American samples were 32.8% longer on average and 41.5% fewer in total number compared to the updated HMP catalog (downloaded in August 2013). The updated HMP catalog had more fragmented assemblies compared to ours (despite a slight improvement compared to the original study³) (Fig. 1b). Merging of the cohorts resulted in a catalog of 9,750,788 genes (here called three cohorts nonredundant gene catalog (3CGC)), based on 1,267 gut metagenomes (1,070 individuals) from three continents, amounting to 6.4 Tb of metagenomic sequencing data (Fig. 1 and Supplementary Fig. 1a,b), which is considerably more than for previous cohorts^{2,3,7}.

Abundant gut microbes were well represented in the 3CGC, but some low-abundance yet common microbes were insufficiently covered, probably because of low sequencing depth for these species.

RESULTS Construction of the integrated gene catalog

Here we completed the MetaHIT cohort by sequencing 249 fecal samples from adults in Denmark or Spain, which led to a collection of 760 European samples^{2,6,7} (**Supplementary Table 1**) and a catalog of 8,096,991 nonredundant genes (**Fig. 1**). To create an intercontinental gene catalog for the

Figure 1 Construction of the IGC. (a) Pipeline for data processing and integration (see also Online Methods). Metagenomic sequencing data from the European, Chinese and American cohorts were processed with the MOCAT pipeline¹³ to generate their respective gene catalogs. The three catalogs were merged to form 3CGC. Sequenced prokaryotic genomes or draft genomes regarded as potentially of human gut origin, according to the IMG system¹⁵ (red), 16S rRNA gene sequence (operational taxonomic unit, OTU) from HMP (green), or coverage by 3CGC (blue) (Online Methods). This initial set of 983 genomes was filtered by our metagenomic sequencing data, which resulted in 511 genomes whose genes comprise the SPGC. Finally, 3CGC were merged with SPGC to generate the IGC. (b) General features of the gene catalogs. *, original HMP study reported a gut microbial gene catalog containing 5,183,353 genes³, 13% more than the number shown here from the catalog downloaded from the HMP website in August 2013 (http://www. hmpdacc.org/HMGC/); the sample number is 139 instead of the 136 stated in the original study³. **, original study for the Chinese samples created a gene catalog based on 145 instead of 368 samples⁴. ***, IGC also incorporated 511 prokaryotic genomes. ORF, open reading frame; N50, 50% of the total length at this length or longer; N90, 90% of the total length at this length or longer; NA, not applicable.

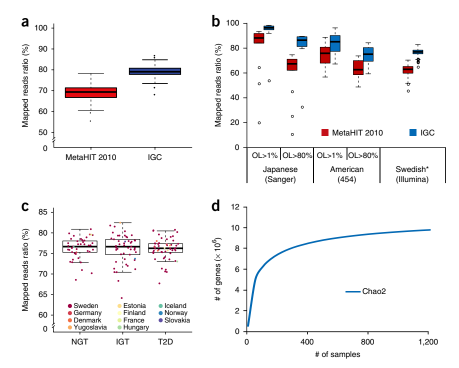


0	Gene catalog	Reference	Sample size	Number of ORFs	Complete ORFs (%)	Total length (bp)	Average length (bp)	N50 (bp)	N90 (bp)	Max length	Min length
	European	Current study	760	8,096,991	56.18	6,039,847,368	746	1,023	381	88,086	102
		MetaHIT 2010 study	124	3,299,822	46.26	2,323,171,095	704	909	378	23,034	102
Ī	American	Current study	139	2,681,342	55.45	1,996,356,219	745	1,005	387	40,011	102
		HMP 2012 study*	139	4,581,984	NA	2,571,088,392	561	765	285	26,109	102
-	Chinese	Current study**	368	3,547,396	60.05	2,750,208,618	775	1,053	405	88,230	102
	3CGC	Current study	1,267	9,750,788	56.34	7,298,407,194	748	1,029	384	88,230	102
	SPGC	Current study	NA	659,492	99.77	612,211,588	928	1,221	513	24,615	100
	IGC	Current study	1,267***	9,879,896	57.74	7,436,156,055	753	1,035	384	88,230	100



gdu

Figure 2 Coverage of the IGC. (a) Percentage of total reads in the MetaHIT 2010 study (n = 124 samples) that could be mapped to MetaHIT 2010 and the IGC. Plotted are interquartile ranges (IQRs; boxes), medians (dark lines in the boxes), the lowest and highest values within 1.5 times IQR from the first and third quartiles (whiskers above and below the boxes), and outliers beyond the whiskers (circles). (b) Percentage of total reads in unrelated studies of Japanese samples (Sanger sequencing, n = 13 samples) and American samples (Roche 454 sequencing, n = 18 samples) that could be mapped to MetaHIT 2010 and IGC with the criterion of identity ≥ 90% and mapped length $\geq 100 \text{ bp}^{16,17}$, and from Swedish samples (Illumina sequencing, n = 145) that could be mapped with identity ≥ 95% (Online Methods)⁵. Results for two overlap cutoffs (>1% and >80%) for queries are shown for Sanger and 454 reads. OL, overlap. *, 130 of the 145 individuals were born in Sweden. (c) Distribution of mapping ratio for the mostly Swedish cohort (shown in **b**) with normal glucose tolerance (NGT), impaired glucose tolerance (IGT) and type II diabetes



(T2D). Each point represents one sample, colored according to nationality at birth. The mapping ratio is not available from the original study⁵. (d) Rarefaction curve based on gene profiles of 1,267 samples using the Chao2 estimator¹⁸ (Online Methods).

For example, the strain labeled Clostridium sp. D5 by the US National Center for Biotechnology Information (NCBI) (but we found it to be classified as Clostridium XlVa in Lachnospiraceae instead of Clostridium in Clostridiaceae, according to the 16S classifier from the Ribosomal Database Project (RDP)¹⁴; Online Methods), was listed by the Integrated Microbial Genomes (IMG) system¹⁵ as a strain isolated from human feces, and we detected it in 53% of stool samples (n = 325) in HMP's 16S rRNA gene data (Online Methods). However, only 4.9% of its genome was covered by 3CGC genes (Supplementary Fig. 1c). To ensure representation of such low-abundance but prevalent organisms, we extracted genes from the genomes of 511 bacterial and archaeal strains that are associated with the human gut and whose genomes were detected in our metagenomic sequencing cohorts (>90% cumulative coverage of the genome by all 1,267 metagenomes; Online Methods). This resulted in a group of 659,492 nonredundant genes, which we refer to as the sequenced prokaryotic gene catalog (SPGC) (Fig. 1 and Supplementary Table 2).

We combined SPGC with 3CGC to form the IGC. The IGC includes 9,879,896 genes, which is nearly three and four times more than the existing MetaHIT and the reassembled HMP gene catalogs, respectively^{2,3} (**Fig. 1**). Each sample contained an average of 762,665 genes and contributed 469 unique genes on average. Any two samples had in common an average of 250,382 genes (32.8% of 762,665 genes).

Quality and completeness of the integrated gene catalog

75.7% and 74.1% of the genes in the IGC were new compared to the MetaHIT 2010 (ref. 2) and the HMP 2012 (ref. 3) gene catalogs, respectively (**Supplementary Fig. 2a,e**). For sequencing reads from the MetaHIT 2010 study², we mapped about 10% more reads to the IGC than to the MetaHIT 2010 catalog, reaching an average mapping rate of 79.24% (**Fig. 2a**). IGC allowed better mapping of sequencing reads (80.54% on average) from the cohorts used for its construction, compared to unintegrated European, Chinese and American gene catalogs (**Supplementary Fig. 3a**). Data from three studies conducted

in America¹⁶, Japan¹⁷ and Sweden⁵ not used in the construction of the IGC had 73.67%, 81.36% and 76.15% of sequencing data represented in the IGC, respectively (**Fig. 2b,c** and Online Methods). Because the percentage of gene-coding regions in all prokaryotic genomes is ~87% (**Supplementary Table 3**), and an estimated 7.25% of sequencing reads with an average length of 77 base pairs could not be mapped reliably as they only partially overlapped with genes (Online Methods), the percentages of mapped reads that we observed with the IGC are close to the maximum achievable mapping rates. In addition, richness estimation based on Chao2 (ref. 18) suggests that the IGC covers 94.5% of the gene content in the sampled gut microbiome (**Fig. 2d**), similar to an estimation of 95.4% using the incidence-based coverage estimator (ICE)¹⁹.

Comparison of the genes assembled in the IGC to the previous catalogs showed that the 12.2% of the MetaHIT 2010 genes not present in the IGC were shorter, more fragmented and often had unknown taxonomy and function compared to the 87.8% MetaHIT 2010 genes present in the IGC (Supplementary Fig. 2c,d). This difference might be due to the approaches used to generate the IGC, including stricter quality control of sequencing reads (using FASTX Toolkit; http:// hannonlab.cshl.edu/fastx_toolkit/), an improved assembler (SOAPdenovo 1.06)²⁰, assembly revision (in the MOCAT pipeline)¹³, more specific gene calling (MetaGeneMark)21, and a standardized and ultrafast clustering algorithm used to merge gene catalogs (CD-HIT)12 (Fig. 1 and Online Methods). Similarly, the 23.6% of HMP 2012 genes that were not present in the IGC were much shorter and aligned with a small portion of sequencing reads compared to the 76.4% HMP 2012 genes present in the IGC (Supplementary Fig. 2g). Of the genes shared among the catalogs, the majority were longer in the IGC (Supplementary Fig. 2b,f).

Taxonomic representation in the IGC

We taxonomically annotated the IGC using reference genomes of 3,449 bacteria and archaea (Supplementary Table 3 and Online

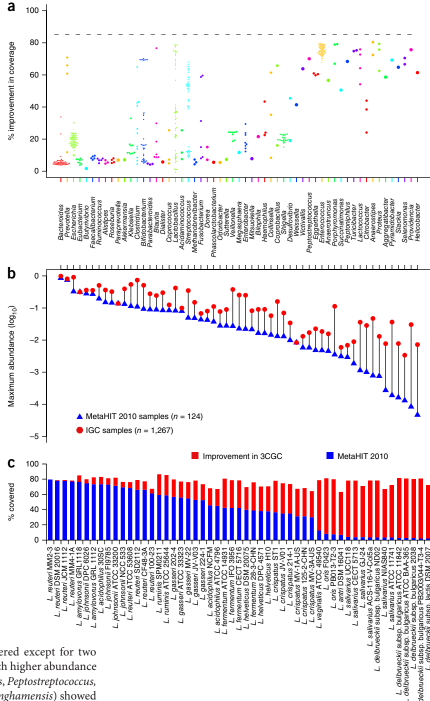
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Figure 3 Improved genome coverage in 3CGC. (a) Improvement in the percentage of each strain's genome covered by 3CGC compared to MetaHIT 2010 genes. Only the 593 strains whose genomes were covered more than 60% by MetaHIT 2010 or 3CGC genes (BLASTN v2.2.24, with criterion of score ≥ 60 and mapped length ≥ 80% for queries) are shown. A complete list of strains is shown in Supplementary Table 4. Strains were grouped by genera. Each dot represents one strain in a genus. Size of the dots scales inversely with the number of strains in the same genus (i.e., a genus with only one strain covered more than 60% had a large dot). The dashed line shows the theoretical maximum of 87% (the average gene content of a bacterial genome). (b) Difference in the highest relative abundance of a genus seen in the MetaHIT 2010 cohort (n = 124) and the IGC cohort (n = 1,267). Genera were ordered according to their relative abundance maxima in MetaHIT 2010 and the resulting x-axis labels are as indicated in a. (c) Genome coverage of different Lactobacillus strains in 3CGC and MetaHIT 2010.

Methods)²². Similar to previous studies^{4,23}, 21.3% of the genes in the IGC could be uniquely and reliably assigned to a phylum and 16.3% to a genus. Genes that could be assigned to genera represented 44.4% of the total sequencing reads (ranging from 5.3% to 78.4% of the sequencing reads in individual samples; Supplementary Fig. 3b).

For 3CGC (IGC without SPGC) compared to MetaHIT 2010, we observed that on average 3CGC had 32.26% higher coverage of individual genomes (the improvement in coverage in 3CGC versus MetaHIT 2010 ranged from -0.71% to +80.44%; average gene content in bacterial genomes is 87%) (Fig. 3a, Supplementary Tables 3 and 4). The improvement in genomic coverage correlated with the increase in maximum abundance of the genera as the cohort size expanded from 124 in MetaHIT 2010 to 1,267 in IGC (Fig. 3a,b). The most abundant genus,

Bacteroides, was no more than 10% better covered except for two strains, whereas genera that were sampled to much higher abundance in the current cohort (e.g., Prevotella, Lactobacillus, Peptostreptococcus, Enterococcus and Helicobacter, specifically, H. winghamensis) showed substantial improvement in their genomic coverage by our gene catalog. At the strain level, pathogenic strains such as *Escherichia coli* O157: H7 and cheese starter strains like L. delbrueckii were substantially better represented in the IGC because of increased sampling (Fig. 3c, Supplementary Fig. 4a and Supplementary Table 4). Analysis of Enterococcus revealed that most samples contained low levels of this genus, but its occasional high abundance in Chinese and European samples, combined with sufficient sequencing depth, enabled 70-80% improvement in the genomic coverage of Enterococcus in the IGC (Supplementary Fig. 4b,c). Thus, increased sampling might be a more effective alternative to deeper sequencing for improved coverage of rare species.



Genera that occurred in large numbers of samples (high occurrence frequency) tended to be those species previously known to inhabit the human gut (Supplementary Fig. 4d, and Supplementary Tables 5 and 6). A notable exception was *Oenococcus* used in wine fermentation, which had not been reported as a gut commensal (**Supplementary Table 6**) but the occurrence frequency of genes annotated to this genus was 13.5% in the current cohort (Online Methods). Although genera not affiliated with the human gut substantially outnumbered genera found in the gut according to the IMG (Supplementary Table 6), they only contributed relatively low-occurrence genes (Supplementary Fig. 4e).



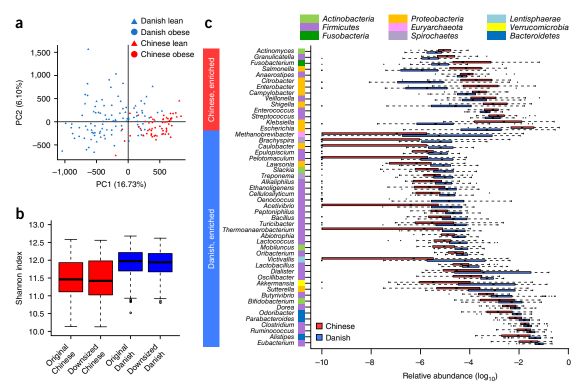


Figure 4 Differences between Chinese and Danish gut microbiota. (a) PCA for gut microbial gene composition in the Chinese (n = 60) and Danish (n = 100) cohorts. The first two principal components are plotted. (b) The within-sample diversity (Shannon index) of the Chinese or Danish cohort, calculated from gene profiles before and after downsizing sequencing data (Online Methods). The difference between cohorts was significant with or without downsizing (Wilcoxon rank-sum test, P < 0.01; two-sided). (c) Top 50 differentially enriched genera in the Chinese and Danish cohorts (P < 0.01, Wilcoxon rank-sum test, two-sided), color-coded by the corresponding phyla. Box-and-whisker plots are as in Figure 2a.

Functional representation of gut microbes

We annotated the genes in the IGC according to the Kyoto Encyclopedia of Genes and Genomes (KEGG) and the evolutionary genealogy of genes nonsupervised orthologous groups (eggNOG) databases^{24,25}. We identified a total of 6,980 KEGG orthologous groups (KOs) and 36,489 eggNOG orthologous groups, which represented 51.6% and 69.3% of the total sequencing reads (**Supplementary Fig. 3b**) and involved 42.1% and 60.4% of the IGC genes, respectively.

876 KOs were present in the IGC but not in the MetaHIT 2010 catalog, whereas 36 KOs present in the MetaHIT 2010 catalog were absent from IGC because of the increased stringency. Consistent with richness estimation by Chao2 (**Fig. 2d**) and ICE, and as suggested by the local rather than global improvement in the coverage of metabolic pathways from MetaHIT 2010 to IGC (**Supplementary Fig. 4f**), the IGC might provide saturated coverage of the functional capacity of the human gut prokaryotes. Although bacteria are the dominant organisms in the gut microbiota^{26–28}, we obtained 500 more eukaryotic KOs in the IGC compared to MetaHIT 2010, but pathways in higher eukaryotes such as glycosphingolipid biosynthesis, proteoglycan biosynthesis and diterpenoid biosynthesis remained largely absent (**Supplementary Fig. 4f** and **Supplementary Table 7**).

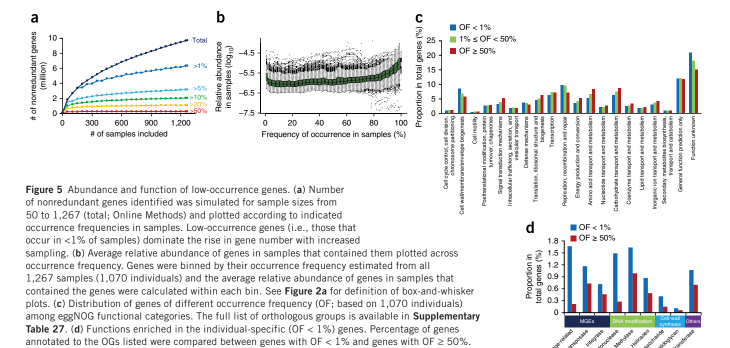
To test the usefulness of the IGC for analyzing metatranscriptomic as well as metagenomic data, we mapped metatranscriptomic sequencing reads from a recent study⁹ to the catalog. After removing genes corresponding to noncoding RNAs such as rRNA, tRNA and signal recognition particle RNA, a higher percentage of the metatranscriptome reads could be mapped to IGC compared to using reference genomes of gut bacteria and archaea only (SPGC) (Online Methods and **Supplementary Fig. 3c**). Despite the stringent alignment criteria (Online Methods), the amount of reads mapping to protein-coding

genes in each sample according to the IGC correlated well with values from the original study (**Supplementary Fig. 3d**). Also, using the IGC instead of the reference genomes (SPGC) allowed identification of more KOs, especially in pathways such as carbohydrate metabolism, cellular processes and signaling, and membrane transport (**Supplementary Fig. 3e**).

Country-specific signatures

To demonstrate the utility of the IGC in quantitative comparisons of the gut microbiome between cohorts, we selected a phenotype-matched group of 60 South Chinese and 100 Danish healthy individuals from the 1,267 samples (**Supplementary Tables 8** and **9**) and profiled their gut metagenomes by comparison to the IGC. Since slightly different DNA extraction methods were used for the two sets of samples^{2–4,6}, before the comparison we randomly selected 11 of the 368 Chinese samples (**Supplementary Table 10**) and extracted the DNA using both protocols to estimate biases resulting from this difference. Metagenomes derived from the same sample using different protocols displayed high self-correlation and the same key features (**Supplementary Fig. 5a–d**). We removed remaining differences before subsequent comparisons (Online Methods).

We could readily separate the Chinese and Danish cohorts by principal component analyses (PCA) based on genes (**Fig. 4a**), KOs or genera profiles (**Supplementary Fig. 5e,f**). Compared to the Danish cohort, the Chinese cohort displayed significantly lower α -diversity in genes and genera but not in KOs ($P = 7.82 \times 10^{-6}$, $P = 1.90 \times 10^{-6}$, P > 0.1, respectively, Wilcoxon rank-sum test), even after normalization of extraction methods and mappable sequencing reads (**Fig. 4b** and **Supplementary Fig. 5c,g,h**). Taxonomically, 151 of the 307 genera showed clear differences between the Chinese and Danish



samples (P < 0.01, false discovery rate (FDR) of 0.0048, power = 0.7, Wilcoxon rank-sum test; **Supplementary Fig. 5i,j** and **Supplementary Table 11**). For example, the Danish samples were generally enriched in the phylum *Firmicutes*, including *Oenococcus* and other lactic acid bacteria, whereas the Chinese samples had greater abundance of *Proteobacteria* (**Fig. 4c**).

The full list of enriched OGs is available in Supplementary Table 28. MGE, mobile genetic elements.

3,491 KOs were significantly different between the two cohorts (P < 0.01, FDR = 0.003, power = 0.7, Wilcoxon rank-sum test; **Supplementary Fig. 5k** and **Supplementary Table 12**). The most prominent differences involved diet-related processes such as energy metabolism, carbohydrate metabolism, amino acid metabolism, and metabolism of cofactors and vitamins, as well as xenobiotic associated functions such as membrane transport and xenobiotic biodegradation and metabolism (**Supplementary Figs. 6** and **7**, **Supplementary Tables 13–25** and **Supplementary Notes**). These differences in metabolic potential of the gut microbiota between healthy Chinese and Danish adults might be influenced by differences in diet (perhaps bread, dairy and vitamins) and environmental factors (perhaps aromatic carcinogens or nitrogen oxides) (**Supplementary Fig. 8** and **Supplementary Notes**).

Individual-specific genes

The increased gene number in the IGC could not be explained by sequencing and assembly error² because such errors in the sequences would have been eliminated as redundant genes (those with >95% identity) during compilation of the gene catalog (**Fig. 1**). To determine the source of the increased number of genes (**Fig. 5**), we simulated the gene content of the catalog when the sample size varied from 50 to 1,267 (**Supplementary Table 26**). The number of genes detected in more than 5% of the samples increased only slightly and approached saturation at about 3.2 million genes; the number of genes present in more than 50% of the subjects remained below 300,000, as in MetaHIT 2010 (ref. 2) (**Fig. 5a**). In contrast, genes found in less than 5% of samples, especially in less than 1% of the samples, continued to increase as sample size increased (**Fig. 5a**). Therefore, genes occurring in a few individuals contributed most to the expanded size of the IGC.

Despite their low occurrence frequency, such genes were abundant in the samples that did contain them (**Fig. 5b**).

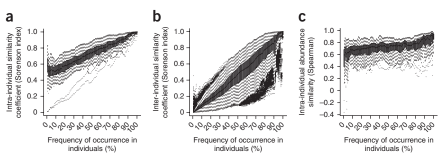
The abundance and repertoire of low-occurrence genes were largely concordant in samples taken from the same HMP individuals at different time points (218 d apart on average), whereas low-occurrence genes from different individuals differed substantially (**Fig. 6**), indicating that these genes were not contamination during sample handling or transient 'passengers' of the gut. Indeed, low-occurrence genes could be more effective than high-occurrence genes when used to distinguish samples from different individuals (**Fig. 6a,b**).

Using the eggNOG database²⁵ we compared the functions of genes seen in less than 1% of the individuals with genes found in more than 50% of the individuals, referred to here as 'individual-specific' and 'common' genes, respectively. The individual-specific genes were modestly enriched in the categories cell wall/membrane/envelope biogenesis and DNA replication, recombination and repair. The common genes were enriched in functions such as signal transduction mechanism, energy production, carbohydrate transport and metabolism, and amino acid transport and metabolism (Fig. 5c and Supplementary Table 27).

When we looked at the exact orthologous groups (groups of genes with homologous sequence and function in different organisms) in the eggNOG resource, genes responsible for the synthesis of cell wall components, especially peptidoglycans and lipopolysaccharides, were overrepresented in the individual-specific set (Fig. 5d and Supplementary Table 28). We also observed an eightfold higher fraction of phage-related proteins, including tail proteins, phage repressors and terminases, among these individual-specific genes. DNA-related functions such as transposases, endonucleases and DNA methylases were enriched in the individual-specific genes (Fig. 5d and Supplementary Table 28), possibly linked to exposure of the gut microbes to foreign DNA. In addition, the individualspecific genes encoded more acetyltransferases, such as GCN5-related N-acetyltransferases that inactivate aminoglycoside-type antibiotics. These results suggest that common genes supply functions essential for survival, whereas individual-specific genes likely reflect adaptation



Figure 6 Temporal stability of low-occurrence genes. (a,b) Genes were binned by their occurrence frequency estimated from all 1,070 individuals. Within each bin, the Sorenson index based on gene content was estimated for pairwise comparisons of multiple samples taken from 43 HMP individuals (a), and first time point (stool 1) samples from 94 different HMP individuals (b). The two sets of dots in a that showed substantially lower Sorenson indices in all occurrence frequencies than the rest of the data originated from comparison between



763536994-stool 2 sample with the same individual's stool 1 and stool 3 samples; this sample seems to be an outlier. (c) Relative abundance of genes in common among samples from 43 HMP individuals sampled at two time points were compared to calculate the Spearman's correlation coefficient. See **Figure 2a** for definition of box-and-whisker plot and Online Methods for computation.

to host immune system, viral infection, antibiotic treatment and other challenges experienced by the gut microbiome.

DISCUSSION

The IGC is a comprehensive resource for further investigations of the gut microbiome, covering strains with a diverse range of occurrence frequencies, abundance and transit durations in the human gut. Future efforts to enhance this catalog could be more targeted to samples with high abundance of a particular strain of interest, which might indicate deviation from a healthy status or relate to a particular environmental factor. As the gut could be seeded by microbes present in food and drinks⁹, quantitative information on the intake and excretion of microbes, the half-life of a strain in the gut²⁹ and so forth would be necessary to define a gut commensal reliably. It is also possible that invasive techniques such as colonoscopy would identify more mucosal-associated microbes than fecal sampling.

Our analysis of two phenotype-matched cohorts of healthy Chinese and Danish adults based on the IGC revealed differences in their gut microbiota regarding many aspects of nutrient metabolism as well as xenobiotic detoxification, which might have been shaped by diet and environment (**Supplementary Fig. 8** and **Supplementary Note**). However, other influences, such as host genetics, remain possible.

Low-occurrence genes contributed overwhelmingly to the increased total gene number in the IGC and might reflect the distinct combination of genetic, nutritional and medical factors in a host. Although the individuals had no recent antibiotic treatment, we observed enrichment in possible antibiotic resistance genes both at the population level^{30,31} (penicillin resistance in Danes and multidrug resistance in Chinese; **Supplementary Table 21**) and in the individual-specific genes (e.g., acetyltransferases and peptidoglycan synthesis), which highlights the need for close monitoring of direct and indirect exposure to antibiotics.

Gut bacteriophages are believed to be mostly temperate but can be induced to enter the lytic cycle^{32,33}. We identified genes for maintenance of lysogeny, such as phage repressors, as well as various genes involved in replication and infection. Other individual-specific genes might also be carried by phages, which are known to alter the metabolism of and confer stress resistance to their bacterial host^{33–36}, and appear stably associated with a given individual³². It remains to be explored whether rare genes in the non-gut microbiome are also enriched for phages or adaptive functions possibly carried by phages³⁶.

Similar to the field of human genetics, where the search for new alleles has progressed from common to rare, our data indicate that cataloging of our 'other genome', the human gut microbiome, is also entering the stage for identification of rare or individual-specific genes instead of common and shared genes. It is also reaching the stage for quantitative comparisons between populations around the world. A reference gene catalog such as the IGC allows rapid and

multi-omic profiling of the genetic and functional repertoire of a given gut metagenome, and facilitates investigations of its geographical, genetic, temporal and physiological characteristics.

A website (http://meta.genomics.cn, optimized for Safari) has been set up to better visualize the annotation information of the gene catalog and guide researchers who are interested in using our data set and downloading specific sets of data.

METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. European Bioinformatics Institute Sequence Read Archive: ERP004605 (metagenomic sequencing data of the 249 European samples and 11 Chinese samples).

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS

J.L., Q.F., S.D.E., P.B. and Jun W. managed the project. T.N., T.H., F.G. and O.P. performed clinical sampling. C.M., W.Z., F.L. and Jua.W. performed DNA extraction. J.L., M.A., K.K., P.B. and Jun W. designed the analyses. J.L., H.J., X.C., H. Zhong, Q.F., E.P., A.S.J., B.C., L.X., S.L., D.Z., Z.Z., W.C., H. Zhao, S.E. and H.B.N. performed the data analyses. J.L., X.C., S.S., J.R.K., Z.Z. and W.C. constructed the integrated gene catalog and performed the functional and taxonomic annotation analyses. J.L., X.C., H. Zhong, B.C. and S.L. performed the country-specific signature analyses. J.L., H.J. and H. Zhong wrote the paper. S.S., M.A., X.X., J.Y.A.-A., H.Y., Ji.W., S.B., K.K., O.P., J.D., S.D.E., P.B. and Jun W. revised the paper. The MetaHIT Consortium members contributed to design and execution of the study.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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ONLINE METHODS

Sample collection and transfer. Under the MetaHIT consortium, 249 fecal samples were collected in a container provided for this purpose at homes of the participating individuals, immediately transferred to a $-20\,^{\circ}\mathrm{C}$ freezer and brought frozen in a cold box to the clinic on the next day. The samples were transferred then to $-80\,^{\circ}\mathrm{C}$ and kept at that temperature or on dry ice. Informed consent was obtained from all Danish volunteers from the Ethical Committees of the Capital Region of Denmark and from all Spanish volunteers from Hospital Universitari Vall d'Hebron. All other samples have been reported previously²–4,6,7.

Sample DNA extraction. DNA extraction from the 249 new MetaHIT samples was performed as previously described³⁷.

For comparison of DNA extraction methods, BGI's protocol⁴ was identical to the MetaHIT protocol³⁷ except that for each fecal sample (up to \sim 1 g), 25 mg of lysozyme and 12.5 mg of proteinase K was added after the initial centrifugation to facilitate cell lysis. Incubation was performed at 37 °C for 1 h to conform to the optimal reaction temperature of lysozyme.

To assess the influence of different DNA extraction protocols, we randomly selected 11 fecal samples from the Chinese cohort⁴ and sent them to Institut National de la Recherche Agronomique (INRA). Our MetaHIT collaborators in INRA extracted the DNA from these 11 samples again following the MetaHIT protocol.

HMP uses PowerSoil DNA isolation kit (MO BIO Laboratories)³, which gives a low DNA yield according to assessments by us (data not shown) and others³8. Combined with the lower $\alpha\text{--}diversity$ in the HMP samples (Supplementary Fig. 51)³9 and the non-overlapping ages (<40 for HMP versus >40 for MetaHIT), we did not include HMP data in our intercontinental comparison.

DNA library construction and sequencing. DNA library construction was performed following the manufacturer's instructions (Illumina). We used the same workflow as described elsewhere² to perform cluster generation, template hybridization, isothermal amplification, linearization, blocking and denaturation, and hybridization of the sequencing primers.

We constructed Illumina libraries for 249 new MetaHIT samples from the European cohort with insert size of 350 bp, followed by high-throughput sequencing to obtain around 36 million paired-end (PE) reads. The read length for each end was 90 bp. High-quality reads were extracted by the MOCAT pipeline from the Illumina raw data ¹³. The proportion of high-quality data in these samples was 89.5% on average.

We constructed Illumina libraries for 11 randomly selected samples from the Chinese cohort, followed by high-throughput sequencing to obtain around 14 million PE reads or 15 million single-end (SE) reads. The read length for each end was 90 bp. High-quality reads were extracted by the MOCAT pipeline from the Illumina raw data¹³. On average, the proportion of high-quality data in these samples was 87.9%, and the actual insert size of our PE library ranged from 311 bp to 326 bp.

The Illumina libraries of 511 European fecal samples from the MetaHIT project and libraries of 368 Chinese fecal samples were constructed and sequenced at BGI using the same protocol as the 249 MetaHIT samples^{2,4,6}. 139 HMP samples were processed by HMP sequencing centers using a similar protocol and platform³.

Public data used. The public gut microbial metagenomes used in this IGC include: (i) 139 HMP samples from stool body site³, which were downloaded from http://www.hmpdacc.org/HMASM/; (ii) 368 Chinese fecal samples⁴, which were downloaded from NCBI (accession codes SRA045646 and SRA050230); (iii) 511 European fecal samples from the MetaHIT project, which were downloaded from the European Bioinformatics Institute (EBI) with accession codes ERA000116, ERP003612 and ERP002061 (refs. 2,6,7), and shared within the MetaHIT consortium. All of these public metagenomic sequencing samples were processed using the MOCAT pipeline to extract high-quality reads¹³.

Other gut metagenomic data used to validate representativeness of IGC include: (i) data from US individuals¹⁶, which were downloaded from NCBI with the accession code SRA002775; (ii) data from Japanese individuals¹⁷,

which were downloaded from EBI with the accession code PRJNA28117; and (iii) data from European individuals⁵, which was downloaded from NCBI with the accession code ERP002469.

Two previously published gene catalogs for the human gut microbiome used in this project include: (i) a gene catalog established from 124 Europeans by MetaHIT², which was downloaded from http://gutmeta.genomics.org.cn/; (ii) a gene catalog established by HMP³, which was downloaded from http://www.hmpdacc.org/HMGC/ in August 2013.

Gut metatranscriptomic data from 59 samples were downloaded from the Gene Expression Omnibus under accession GSE46761 (ref. 9). All of these public metatranscriptomic sequencing samples were processed by the MOCAT pipeline to extract high-quality reads¹³.

Collection and quality control of 3,449 sequenced prokaryotic genomes or draft genomes. Prokaryotic genomes were collected and filtered as described 22 . Briefly, all prokaryotic genomes available at NCBI and EMBL Bank on 23 February 2012 were downloaded and genomes with more than 300 contigs and N50 < 10 kbp were removed. In addition, we removed genomes for which less than 30 of 40 universal single-copy marker genes were identified 40,41 . Finally, for genomes with the same taxonomy identifier, but different project identifiers, one genome was randomly chosen, which resulted in a set of 3,449 genomes used in this study.

Construction of the integrated gene catalog (IGC). Illumina sequencing reads for fecal samples from European, Chinese and American adults were independently processed (quality control, removal of human sequences, assembling, assembly revision and gene prediction) using MOCAT¹³, which could process metagenomes in a standardized and automated way while improving the quality of assembly and gene prediction compared to using default parameters for the supported programs based on parameter exploration and data-driven parameter optimization at run time¹³. We chose FASTX Toolkit (http://hannonlab.cshl.edu/fastx_toolkit/) for quality control, SOAPaligner2 (ref. 42) for identifying human sequences, SOAPdenovo v1.06 (ref. 20) for assembling and MetaGeneMark²¹ for gene prediction in the MOCAT pipeline. The configuration file we used in MOCAT has been deposited on GigaScience Database¹¹. Genes in each cohort were clustered using CD-HIT¹². The gene catalogs were then merged to generate a human gut microbial gene catalog based on all 1,267 samples, referred to as 3CGC.

3,449 sequenced bacteria and archaea genomes or draft genomes were gathered²², and human gut-related prokaryotes were selected in two steps. First, strains that satisfied any one of these three criteria were included: (i) the strain's habitat is "human gastrointestinal tract" according to IMG (http://img. jgi.doe.gov/cgi-bin/w/main.cgi) (downloaded on 24 July 2012), i.e., the strain's "Body Site" is "Gastrointestinal tract" or "Isolation" is "human feces." (ii) 16S rRNA sequence of the strain is identical to that of an OTU reported by HMP as from stool body site⁴³. 485 nonchimeric HMP OTUs from stool body site were aligned to the 16S rRNA gene of each strain using mothur (version 1.23.1)44, with a global identity cutoff of ≥99.5%. (iii) Ratio of genes covered by 3CGC with a weak criterion is high. Genes from each strain were aligned to 3CGC using BLAT⁴⁵ with the criterion of overlap \geq 10% and identity \geq 95%. Strains with over 80% of their genes covered by 3CGC were selected. We obtained 983 gut-related prokaryotic strains following these three criteria (Fig. 1a and Supplementary Table 3). Second, these 983 prokaryotic genomes were filtered by the cumulative coverage by our metagenomic sequencing data (more than 90% of genome by 1,267 samples) to confirm that they are part of the human gut microbiome. The genomes or draft genomes of each strain were initially aligned with sequencing reads from 100 samples using SOAP2 (ref. 42) with identity ≥ 90%. Strains whose genome had not yet been covered over 90% were aligned with data from all 1,267 samples for further selection. After such filtering, 511 prokaryotes remained and were used to construct the gut-related SPGC (Fig. 1 and Supplementary Table 3).

Finally, the gene catalog based on metagenomic sequencing data (3CGC) and the gene catalog based on sequenced prokaryotic genomes (SPGC) were combined using CD-HIT to generate the IGC (Fig. 1).

The gene catalogs, annotation information, abundance profile, assemblies and predicted open reading frames of the 1,267 samples have been deposited into the GigaScience database¹¹.



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Investigation on the representation of a low-abundance but prevalent human gut bacterium. Genome of NCBI 556261.HMPREF0240_10201 (Clo stridiaceae | Clostridium | Clostridium sp. D5 in NCBI, Lachnospiraceae | Clostrid ium XlVa according to the RDP database¹⁴, Feburary 2014) originally isolated from human feces was chosen as a reference. Genes from 3CGC were aligned to the NCBI 556261 genome by BLAST with the criterion of more than 95% identity and 90% overlap of query. Only 4.9% the genome was represented by 3CGC. The occurrence frequency of the strain was assessed by sequencing data of 325 stool samples from HMP in 16S rRNA gene variable regions³. Tags of each sample with length more than 150 bp were aligned to the 16S rRNA gene from the NCBI 556261 genome by mothur (version 1.23.1)⁴⁴ with more than 97% identity and more than 90% overlap of query. 53% of the HMP stool samples carried this species (2.1 tags on average), which indicated that it is a universally present but low-abundance species in the human gut environment. Cumulative coverage of its genome by metagenomic sequencing reads from 1,267 samples was assessed by SOAP2 (ref. 42) with more than 95% identity. The best covered sample was chosen according to the maximum number of NCBI 556261 genes covered by ORFs assembled from the sample.

Phylogenetic annotation based on reference genomes. Phylogenetic annotation was performed using an in-house pipeline. (i) We aligned 9.7 million genes of 3CGC onto the database of 3,449 prokaryotic genomes using BLASTN (v2.2.24, default parameters except that -e0.01 -b100 -K 1 -F T). (ii) For each gene, only the top 10% highest-scoring alignments covering \geq 80% of gene length and identity \geq 65% were retained. (iii) Each gene was assigned the taxonomy of the alignment(s) with 50% or higher consensus above the similarity threshold for taxonomic rank (>65% for phylum, >85% for genus and >95% for species). (iv) The 0.7 million genes of SPGC were assigned the taxonomy they came from.

As explained previously^{4,23}, our phylogenetic annotation method ensures unique assignment and minimizes ambiguity. The false positive rates at phylum level and genus level were 0.77% and 1.84%, respectively²³.

Functional annotation (KEGG and eggNOG). We aligned putative amino acid sequences translated from the integrated gene catalog against the proteins or domains in eggNOG (v3.0) and KEGG databases (release 59.0, genes from animals or plants were excluded) using BLASTP (v2.2.24, default parameter except that -e 0.01 -b 100 -K 1 -F T). KEGG annotation was performed using an in-house pipeline, where each protein was assigned to a KO when the highest-scoring annotated hit(s) contained at least one alignment over 60 bits. eggNOG annotation was performed using Smash Community (v1.6, find_best_hit.pl&og_mapping.py, with default parameters)⁴⁶.

Comparison between MetaHIT 2010, HMP 2012 and IGC genes. 9.9 million genes from IGC and 3.3 million genes from the MetaHIT 2010 catalog² were pooled together, and redundant genes were identified using CD-HIT¹¹ with \geq 95% identity and \geq 90% overlap. For the shared (overlapped) genes, the length was compared and discrepancies greater than 10% were regarded as significantly longer or shorter in IGC. 9.9 million genes from IGC and 4.6 million genes from the updated HMP 2012 catalog³ were compared using the same workflow.

Aligning public human microbial sequencing data onto gene catalogs. Roche 454 reads from 18 US twins and their mother 16 and Sanger reads from 13 Japanese individuals 17 were aligned to MetaHIT 2010 and IGC using BLASTN (v2.2.24), with the criterion of mapped length \geq 100 bp. The ratio of reads that could be aligned to MetaHIT2010 or IGC was filtered by two overlap thresholds (the proportion of a read aligned to the gene catalog), 1% and 80% (Fig. 2b).

Illumina reads from 145 European individuals (130 of them were born in Sweden)⁵ were aligned to MetaHIT 2010 and IGC using SOAP2 with the criterion of identity \geq 95%⁴².

Aligning public human microbial metatranscriptomic data onto gene catalogs. With our gene catalog constructed directly from the gut microbiome, we were able to allocate transcript sequences from 59 metatranscriptomic sequencing samples⁹ onto the gene catalog (IGC) using SOAP2 (≥95% identity)⁴² to identify expressed genes, and retrieve their annotated functions.

SPGC, the gene catalog compiled from 511 gut-related bacterial or archaeal genomes or draft genomes present in the 1,267 metagenomes, was used to compare with IGC, because the set of 539 human-associated microbial reference genomes used in the original study is not known (the human microbiome database used⁴⁷ now contains 2673 genomes, as of 31 March 2014, http://www.hmpdacc.org/catalog/). The hashing-based software SSAHA2 (ref. 48) used in the original study has a very loose alignment criterion (parameters: '-best 1 -score 20 -solexa') and likely does not support unequivocal identification of gene functions. Besides, SSAHA2 is substantially slower than short-read aligners such as SOAP2 in terms of aligned bases per unit time, and is too slow to handle our gene catalogs.

A few noncoding RNAs were involved while integrating 511 human gut-associated reference genomes into our catalog. In order to calculate the ratio of reads mapped to protein-coding genes (coding sequences; CDS), we eliminated genes annotated to non-coding RNAs in SPGC and IGC, with the keywords 'RNA' but no '-ase', 'enzyme' or 'protein' from the original genome annotations²². SPGC and IGC contained 923 and 866 noncoding RNA genes (rRNA, tRNA, SRP RNA, etc.), respectively, according to this search criteria.

Computation of relative gene abundance. High-quality reads from each sample were aligned against the gene catalog by SOAP2 using the criterion of identity $\geq 95\%^{42}$. Sequence-based abundance profiling was performed as previously described⁴.

Construction of genus, KO and enzyme profiles. For the genus profile, we used phylogenetic assignment of each gene from the original gene catalog and summed the relative abundance of genes from the same genus to yield the abundance of that genus. Relative abundance of each genus in a sample constituted the genus profile of that sample. The KO profile was constructed using the same method. The relative abundance of an enzyme was calculated from summation of the relative abundance of its corresponding KOs.

Estimating loss of mappable reads at gene boundaries. When mapped against the gene catalog, a portion of short reads would be lost at the boundary regions of a gene (Supplementary Fig. 9a).

The lost ratio of abundance, rate lost could be calculated as (Supplementary Fig. 9b),

$$\text{if } (2 \times L_{\text{boundary}}) \leq L_g, \text{rate}_{\text{lost}} = \frac{L_{\text{boundary}} - 1}{L_g}$$

if $(2 \times L_{\text{boundary}}) > L_g \ge L_{\text{boundary}}$,

$$\text{rate}_{\text{lost}} = \begin{cases} 1 - \frac{L_g + 2}{4 \times L_{\text{boundary}}}, & \text{if } L_g \text{ is even} \\ 1 - \frac{(L_g + 1)^2}{4 \times L_g \times L_{\text{boundary}}}, & \text{if } L_g \text{ is odd} \end{cases}$$

if
$$L_{\text{boundary}} > L_{\sigma}$$
, rate_{lost} = 1

where L_g is gene length; $L_{
m boundary}$ is length of boundary region which equals read length in our situation.

We used all the genes from 511 human gut-associated prokaryotes (Supplementary Table 2) to estimate the proportion of lost sequencing data from prokaryotic gene coding region for individual samples (Supplementary Fig. 9c). The proportion of lost sequencing data from prokaryotic gene coding region,

$$Rate_{T,lost} = \sum_{i=1}^{n} rate_{lost,i} \times \frac{Lg_i}{L_{511,genome}}$$

where i (1,2, ... n) refers to each gene from the 511 human gut-associated prokaryotes; Lg_i is the length of gene i; rate_{lost, i} is the lost abundance for gene i; $L_{511, \text{genome}}$ is the total genome size of 511 human gut-associated prokaryotes.

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For a given read length of 77 bp (the average read length of 1,267 samples in this study) (**Supplementary Table 1**), the proportion of lost sequencing data from prokaryotic gene coding regions is estimated to be 7.25%.

BMI criteria used for European and Chinese cohorts. A number of reports indicated that the BMI criterion for Asians is lower than for Europeans, and that Asians tend to accumulate abdominal fat and develop obesity-related diseases without overall obesity^{49–51}. Accordingly, we used a lower BMI cutoff to define obesity status in Chinese. For Chinese, we used BMI values < 21 kg/m² for being lean and $\geq 25 \text{ kg/m}^2$ for obesity. For Danish we used BMI values < 25 kg/m² for being lean and $\geq 30 \text{ kg/m}^2$ for obesity.

Biodiversity and richness analysis: α -diversity. Based on the gene, genus or KO profile, we calculated the α -diversity (within-sample diversity) to estimate the gene, genus or KO diversities of a sample using the Shannon index:

$$H' = -\sum_{i=1}^{S} a_i ln a_i$$

where S is the number of genes and a_i is the relative abundance of gene i. A high α -diversity indicates a high evenness or many types of genes present in the sample.

Adjustment by linear regression. A linear regression equation was generated based on the 11 randomly selected samples whose DNA was extracted twice using both the BGI and MetaHIT protocols (**Supplementary Fig. 5c**). For comparison with the Danish cohort, the within-sample diversity index of the Chinese cohort (n = 60) was adjusted according to the linear regression equation to eliminate possible biases introduced by different DNA extraction protocols.

Read downsizing. To eliminate the influence of fluctuations in sequencing amount, we sampled the alignment results and downsized the number of mapped pairs to 11 million for each sample.

Rarefaction curve analysis. To assess the gene richness in our Chinese or Danish cohort, we generated a rarefaction curve. For a given number of individual samples, we performed random sampling 1,000 times in the cohort with replacement and estimated the total number of genes that could be identified from these samples by the Chao2 richness estimator 18 . To minimize erroneous identification, only the genes with ≥ 1 pair of mapped reads were determined to be present in a sample.

Statistical analysis of the gut metagenome. To identify associations between metagenomic profiles and populations, a two-tailed Wilcoxon rank-sum test was used in the profiles. We identified a genus marker if its P value was <0.01 and occurrence frequency >10% in at least one cohort, and identified a KO/enzyme marker if its P < 0.01 and occurrence frequency > 30% in at least one cohort.

The statistical method used to detect biases in extraction methods was similar to the above-mentioned method, but we did not consider occurrence frequency because the sample size was only 11.

Estimating the false discovery rate and statistical power. Instead of a sequential P-value rejection method, we applied the 'qvalue' method proposed in a previous study⁵² to estimate the FDR. Statistical hypothesis tests were performed on a large number of features of the genus profiles and KO profiles. Given that a FDR was obtained by the q value method⁵³, we estimated the power P_e for a given P-value threshold as

$$P_e = \frac{N_e(1 - \text{FDR}_e)}{N(1 - \pi_0)}$$

where π_0 is the proportion of null distribution P values among all tested hypotheses; N_e is the number of P values that were less than the P-value threshold;

N is the total number of all tested hypotheses; ${\rm FDR}_e$ is the estimated false discovery rate under the $P\text{-}{\rm value}$ threshold.

Simulation for the dependence of gene catalog size on sampling scale.

Two data tables were prepared before simulation. The first one was a 'gene profile' table, containing information about the relative abundance of each gene for each sample. The table was generated using the method in ref. 4. The second one was a 'genes assembled' table, containing information about whether a gene was assembled due to the presence of an individual sample. The table was generated from the clustering output file from CD-HIT, which traced genes corresponding to the same cluster (representative gene) to the original sample.

Simulation was performed by random sampling without replacing the selected samples, with sample size from 50 to 1,267 samples, 50 samples per step. For each simulated set, the estimated size of the nonredundant gene catalog was calculated from the 'genes assembled' table, and the distribution of genes in a range of occurrence frequencies was calculated through the 'gene profile' table. For each sample size, the simulation was performed 1,000 times, and the averages were plotted.

Concordance of low-occurrence genes between samples. Sorenson index, also known as Sørensen-Dice index, was used to measure the presence/absence similarity of genes of all occurrence frequencies (according to all 1,070 individuals) in HMP samples.

Sørensen's original formula was applied to the presence/absence data, in the form:

$$QS = \frac{2C}{A+B} = \frac{2|A \cap B|}{|A|+|B|}$$

where A and B are the number of genes in samples A and B, respectively, and C is the number of species shared by the two samples; QS is the quotient of similarity and ranges from 0 to 1.

Spearman's rank correlation coefficient was used to measure the abundance similarity of genes of all occurrence frequencies in HMP samples.

Intraindividual similarity was based on the samples taken from the same 43 HMP individuals at different times (218 d apart on average). And the interindividual similarity was based on any two stool samples (the first time point) from 94 different HMP individuals.

Source genera for genes of diverse occurrence frequencies. The occurrence frequency of each IGC gene was rounded to the nearest integer, for example, 1% represents $1\% \pm 0.5\%$. For each source genus, the numbers of genes in each occurrence frequency percentile were counted (Supplementary Table 5). Supplementary Figure 4e was derived from this table with a gene number cutoff of ≥ 10 .

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