

1 Over 50000 metagenomically assembled draft genomes for the 2 human oral microbiome reveal new taxa

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21 ABSTRACT

22 The oral cavity of each person is home for hundreds of bacterial species. While
23 taxa for oral diseases have been well studied using culture-based as well as
24 amplicon sequencing methods, metagenomic and genomic information remain
25 scarce compared to the fecal microbiome. Here we provide metagenomic shotgun
26 data for 3346 oral metagenomics samples, and together with 808 published
27 samples, assemble 56,213 metagenome-assembled genomes (MAGs). 64% of the
28 3,589 species-level genome bins contained no publicly available genomes, others

with only a handful. The resulting genome collection is representative of samples around the world and across physiological conditions, contained many genomes from Candidate phyla radiation (CPR) which lack monoculture, and enabled discovery of new taxa such as a family within the Achleplasmataceae order. New biomarkers were identified for rheumatoid arthritis or colorectal cancer, which would be more convenient than fecal samples. The large number of metagenomic samples also allowed assembly of many strains from important oral taxa such as *Porphyromonas* and *Neisseria*. Predicted functions enrich in drug metabolism and small molecule synthesis. Thus, these data lay down a genomic framework for future inquiries of the human oral microbiome.

The human microbiome has been implicated in a growing number of diseases. The majority of microbial cells is believed to reside in the large intestine¹ and cohorts with fecal metagenomic data contain over 1000 individuals^{2,3}. For the oral microbiome, hundreds of metagenomic shotgun-sequenced samples have been available from the Human Microbiome Project (HMP) and for rheumatoid arthritis⁴⁻⁶. A number of other diseases studied by Metagenome-wide association studies (MWAS) using gut microbiome data also indicated potential contribution from the oral microbiome in disease etiology⁷⁻¹². Although the MWAS on rheumatoid arthritis was based on a *de novo* assembled reference gene catalog for the oral microbiome⁶, analyses on bacterial genomes would be more desirable. And when oral samples show comparable or better sensitivity and accuracy for disease diagnosis, prognosis or patient stratification than fecal samples, oral samples would be much more convenient as they could be available at any time and taken at a fully controlled setting witnessed by trained professionals. Unlike the anaerobic environment for the gut microbiome, the oral microbiome is believed to be well covered by culturing¹³, and analyses by 16S rRNA gene amplicon sequencing or polymerase chain reaction (PCR) are common. Recently published large-scale metagenomic assembly efforts mostly

included fecal metagenomic data¹⁴⁻¹⁶. It is not clear how much is really missing for the oral microbiome. The saliva, in particular, seems to have more bacterial species per individual than the fecal microbiome¹⁷.

After getting contigs using assembly algorithms suitable for metagenomic data¹⁸, a central idea used by metagenomic binning algorithms is that genes or contigs that co-vary in abundance among many samples belong to the same microbial genome^{8, 19-21}. Large cohorts are therefore prerequisites for high-quality assembly.

Here we present 3346 new oral metagenomic samples, and 56,213 metagenome-assembled genomes (MAGs) which represent 3,589 species-level clades, revealing new taxa as well as substantially complementing the genomic content of known species. This genome reference are highly representative of metagenomic samples not used in assembly, and could facilitating culturing, functional screens as well as disease diagnosis and modulation based on the oral microbiome.

RESULTS

Draft genomes assembled from oral metagenomic data

In order to substantially increase the amount of oral microbiome data, we shotgun sequenced 2284 saliva and 391 tongue dorsum samples from the 4D-SZ cohort^{3, 12, 22}, 671 saliva samples from five ethnic groups of Yunnan province, producing over 43.19 terabytes of sequence data (**Supplementary Table 1**). Together with 808 published samples from 5 studies^{6, 23-26} that have not been used for metagenomic assembly (**Supplementary Table 1**), a total of 4,154 oral samples with metagenomic data were obtained. The data in each sample was single assembled into contigs using SPAdes^{27, 28} (**Fig. 1a, Supplementary Table 1**). Binning was then performed using MetaBAT2²¹ for the 39,458,119 contigs longer than 1.5kb, leading to 56,213 metagenome-assembled genomes (MAGs), 15,013 of which were of high quality according to recently agreed standards²⁹ using CheckM³⁰ (>90% completion, <5% contamination, **Fig. 1a**). The remaining 41,200 also reached the standards for

84 medium-quality MAGs (>50% completion, <10% contamination), while low-quality
85 assemblies were not further analyzed (**Supplementary Table 2**). The median
86 constructed MAGs per sample is 12, with highest from ZhangX_2015
87 (**Supplementary Figure 1a**).

88 **New genomes from the new samples**

89 To evaluate the novelty of the assembled genomes, delineate their taxonomy
90 and potential source, we comprehensively incorporated 190,309 existing isolate or
91 metagenome-assembled genomes from NCBI Refseq, eHOMD³¹ (the expanded
92 Human Oral Database), and recent publications^{14-16, 32} including our culture collection
93 from fecal samples in Shenzhen³³ (**Fig. 1a**).

94 Species-level clusters (SGBs, for species-level genome bins) were computed
95 for the over 0.25 million genomes following multiple steps (**Fig. 1a**, see Methods for
96 details), defined as at least 95% average nucleotide identity (ANI) and at least 30%
97 overlap of the aligned genomes. The clustering well-collapsed the genomes, with
98 about 10-fold reduction in number, i.e. resulting in around thirty thousand species.
99 Besides the 27,936 species that were non-oral according to reference genomes in the
100 cluster (defined in **Fig. 1b**), 2,313 clusters (64% of the total oral species) only
101 contained our MAGs (denoted uSGBs for unknown SGBs), some of which were
102 repeatedly captured in our data, with more than 50 genomes each (**Fig. 1b,c**); the
103 1,276 known oral SGBs (kSGBs) could be further divided according to the percentage
104 of reference genomes in the cluster. Interestingly, kSGBs with over 50% unknown
105 genomes outnumbered kSGBs with 0-50% unknown genomes for clusters containing
106 10 or more genomes (**Fig. 1b,c**), underscoring the discovery power of large
107 metagenomic cohorts. And the top three contributions of uSGBs are 4D_SZ (1441),
108 ZhangX_2015 (445) and Yunnan (334) (**Supplementary Figure 1b**). Comparing the
109 ratio of new MAGs in the samples, we retrieved a greater fraction of previously
110 unknown genomes in dental compared to saliva or tongue samples, even though we

did not take dental samples for this large cohort (**Fig. 1d**). This ratio also appeared to differ between cohorts, with less than 10% unknown for samples from France or the U.S., and more newly matched uSGBs for samples from Fiji, Germany and Luxemburg (**Fig. 1d**). The large cohort available from this study is crucial for the retrieval of novel oral species, contributing over 2000 uSGBs, greatly expanding our knowledge of oral microbiome diversity.

Close to 90% representation of oral metagenomics data by the genomes

We next examined the ability of this species-level genomes set to represent metagenomic shotgun data. We assessed the percentage of reads that could align to cultured genomes only (eHOMD) and cultured complemented by metagenomically assembled genomes. The median was 66.99% mapping with the 1526 genomes from eHOMD (**Fig. 2a**). The 4930 representative human SGBs from a recent large-scale assembly study that included available oral metagenomic samples¹⁶ led to 79.72% mapping, and the representative oral 3589 SGBs from the current study instead led to 88.06% mapping (median for all samples), especially for metagenomes from the U.S. and Germany; and a median of 85.29% mapping even for 81 saliva and subgingival metagenomes from three cohorts that were not used in the assembly process³⁴⁻³⁶ (**Fig. 2a, Supplementary Table 1**). Across physiological states, our SGBs well represented pregnant samples from the U.S. (reaching 92.99% mapping), RA (reaching 90.69% mapping) and diabetes (reaching 83.99% mapping) (**Fig. 2b**). Such a high degree of representation of metagenomic data across geography, ethnicity, age and physiological states suggest that the expanded genomic content of oral SGBs could serve as a starting point for quantitative taxonomic and functional analyses of the human oral microbiome.

Taxonomic landscape of the oral microbial genomes

We constructed a phylogenetic tree for the 3,589 oral SGBs, and similar to the gut microbiome, *Firmicutes* took up the largest number of branches (1248 clusters,

12307 genomes, **Fig. 3c**). The other 46276 genomes distributed into 15 phyla, including major human oral phyla such as *Actinobacteria* (490 SGBs, 6477 genomes), *Bacteroidetes* (368 SGBs, 23409 genomes), *Proteobacteria* (364 SGBs, 7570 genomes), *Campylobacteriota* (280 SGBs, 1841 genomes), and *Fusobacteriota* (145 SGBs, 1998 genomes) (**Fig. 3, Supplementary Table 3**). uSGB accounted for 181.27% increase in the reconstructed phylogenetic branch length, with over 80% of the diversity in *Campylobacterota* phylum contributed by the new uSGBs, followed by over 70% for *Patescibacteria* and *Fusobacteriota* (**Fig. 3b**), which seemed overlooked by culturing studies. We estimated there is a median of 210 SGBs with the relative abundance higher than 0.001 per sample (**Supplementary Figure 1c**). Besides uSGBs, there are also very high abundances, explained for 68.10% of richness and 65.23% of relative abundance per sample (**Supplementary Figure 1d,e**). Our MAGs greatly expanded the species or strains diversity within each phylum. As many as 596 SGBs from 4006 genomes belonged to the candidate superphylum of *Patescibacteria* (*Parcubacteria*, also known as OD1), which only have 157 kSGBs with 3115 reference genomes. We note a few not so well studied phyla that were interesting in analogy to the gut microbiome. *Akkermansia* is the only genus from Verrucomicrobiota in the human gut and intensively pursued for its role in health and diseases, and Verrucomicrobiota and Spirochaetota take up a greater fraction in Hadza hunter gatherers compared to developed countries³⁷. Here we identified 6 genomes in 3 SGBs for Verrucomicrobiota, and 900 genomes in 67 SGBs for Spirochaetota. 121 reference genomes were only available for 32 SGBs within Spirochaetota. 198 SGBs with 1169 genomes belong to the candidate division Saccharibacteria (TM7) (**Fig. 3a, Supplementary Table 3**).

At the genera level, *Streptococcus* (460 SGBs), *Campylobacter* (279 SGBs), *Actinomyces* (184 SGBs), *Prevotella* (159 SGBs), *Atopobium* (146 SGBs) were the major genera in the SGBs (**Supplementary Table 3**). 265 of the 2313 uSGBs had taxonomic information until order or family, but cannot be annotated to a known

genus. The top three uSGB classified families were Saccharimonadaceae (17.99%), Streptococcaceae (12.88%) and Campylobacteraceae (9.51%), whereas the most assigned genera were Streptococcus (12.88%), Campylobacter_A (7.65%) and TM7x (5.92%) (**Fig. 3c**).

A new family with small genomes

In the Achleplasmatales order (Mollicutes class) of the *Tenericutes* phylum, a number of our uSGBs with high-quality MAGs formed a clade distinct from *Acheloplasma* and *Candidatus Phytoplasma*, with shallow branches within the clade (**Fig. 4a**). The genome size of this genome-defined family, which we temporarily denoted as *Ca. Bgiplasma*, is 0.69 ± 0.05 Mbp, which is similar to *Candidatus Phytoplasma* (0.64 ± 0.14 Mbp), but much smaller than *Acheloplasma* (1.50 ± 0.20 Mbp). Genomes of such small size were discarded in early efforts of metagenomic assembly¹⁹, but we now know *Ca. Bgiplasma* are complete entities according to single-copy marker genes in CheckM (**Supplementary Table 2**). The GC content of the three clades were also different. *Ca. Bgiplasma* family was more towards normal GC content ($34.57 \pm 0.21\%$), not as low as *Acheloplasma* ($30.99 \pm 1.75\%$) and *Candidatus Phytoplasma* ($25.98 \pm 2.68\%$) (**Supplementary Table 5**). Despite the lack of deep branches, the ANI distribution of uSGB within *Ca. Bgiplasma* family showed two separate groups at genus-level divergence (ANI <85%) (**Fig. 4b**), illustrating diversity within this new family. This 11 uSGBs comprising 29 MAGs contribute more than 0.1% relative abundance in 209 samples, indicating that this family is an potentially important but so far uncharacterized clade in the oral microbiome.

5.53 M genes (87.97% of total) of representatives genome of SGBs can be annotated by EggNOG mapper^{38, 39} with the rate of annotation 89.55% for uSGBs and 81.83% for *Ca. Bgiplasma* family(**Supplementary Table 5**). We found *Ca. Bgiplasma* are gene content dominated by replication, recombination and repair, posttranslational modification, protein turnover, chaperones, and inorganic ion

transport and metabolism, which are reported active up-regulated in *Deinococcus* during gamma-irradiation⁴⁰ (**Supplementary Fig. 2b**).

Distribution of species and strains

The new samples from this study differed in oral microbiome composition compared to published samples across geography/ethnicity (**Fig. 5a,b**). Both the 4D-SZ and Yunnan samples abundantly contained many uSGBs (of the top 10 abundance species in cohort) such as *Neisseria* spp., *Porphyromonas* spp. and kSGBs such as *Haemophilus parainfluenzae* and *Veillonella denticarios*, which were rare in the other cohorts (**Fig. 5b**). Pregnant samples from the U.S. contained *Fannyhessea vaginae* (the vaginal pathogen previously known as *Atopobium vaginae*⁴¹), *Urinacoccus*, etc. that were of much lower abundance in other cohorts (**Fig. 5b**). Samples from Fiji, although not well mapped (**Fig. 2a**), showed high levels of a few SGBs that were also seen in the RA study from Beijing, China, including an SGB from *Saccharibacteria* (TM7) (**Fig. 5b**).

At the strain level, the new samples from the current study greatly expanded the genome collection for common taxa such as *Neisseria* spp., *Porphyromonas* spp., and *Prevotella* spp. (**Fig. 5c,d**). The numbers of publicly available reference genome for the top ten most abundant species in the genera *Porphyromonas* and *Prevotella* were less than 10, and less than 100 for the genus *Neisseria*. Here we obtained more than 1000 genomes for a few of the species, and increased the diversity in all the species in these genera (**Fig. 5c**). Most of the species with a large number of genomes showed strain-level variations (subspecies). The *Prevotella nanceiensis* kSGB, for example, included 3 reference genomes that were similar to a few genomes from developed countries, while our samples contributed two large clusters that were more distantly related (**Fig. 5d**).

New disease markers according to the oral genomes

219 To illustrate the utility of our genome collection in metagenomic studies
 220 including MWAS, we reanalyzed dental and salivary microbiome data from RA
 221 patients and controls⁶. For better confidence in the markers regardless of cohort, we
 222 only analyzed SGBs containing >10 genomes. Similar to the original study, oral
 223 markers selected by a 5x 10-fold cross-validated gradient boosting algorithm
 224 include a number of Gram-negative bacteria e.g. *Haemophilus* spp.,
 225 *Aggregatibacter* spp. enriched in dental samples from healthy volunteers, while
 226 only a *Pseudomonas* SGB and a *Enterococcus* SGB were selected for RA samples
 227 (**Fig. 6a**). Interestingly, the two new RA dental markers appeared more abundant in
 228 control saliva samples. The strongest marker from healthy saliva remained
 229 *Lactococcus lactis*⁵, and *Lactobacillus paracasei*, *Streptococcus infantarius*, were
 230 identified, reminiscent of beneficial effects of *L. casei* gavage in rat model of RA⁴²,
 231 ⁴³. The assembled genomes allowed matching of different species in the *Veillonella*
 232 genus as RA saliva markers. Moreover, *Pauljensenia* spp., a genus recently renamed
 233 from *Actinomyces*⁴⁴, was identified as highly predictive of RA. As *Actinomyces* are
 234 the basis for dental attachment of oral bacteria⁴⁵, potential contribution of
 235 *Pauljensenia* spp. to periodontitis in RA patients remains to be explored; the dental
 236 microbiome was obviously deranged, consistent with epidemiology⁵.

237 A set of saliva samples from colorectal cancer and controls from France are
 238 also available⁴⁶. Here, we found *Pauljensenia* spp., to be control-enriched, along with
 239 *Acinetobacter radioresistens*, *Lachnoanaerobaculum* sp., *Catonella* sp., etc (**Fig. 6b**).
 240 *Streptococcus thermophilus*, a species previously found to be enriched in fecal
 241 samples from control or adenoma compared to CRC patients⁴⁷ was also identified in
 242 control saliva. The markers enriched in CRC oral samples are more unexpected.
 243 Besides *Porphyromonas* spp., *Prevotella maculosa*, we found a *Lachnospiraceae*
 244 SGB (potentially TMA-producing and consistent with gut results^{10, 48-50}),
 245 *Capnocytophaga leadbetteri*, *Cardiobacterium hominis*, etc. (**Fig. 6b**). Thus, the
 246 substantially expanded collection of oral microbial genomes enabled discovery of new

disease markers and genomic representation of previously reported markers,
facilitating the shift from fecal to oral microbiome-based diagnosis and therapeutics.

Potential functions in drug metabolism and small molecule synthesis

Many human target drugs are reported to be metabolized to its inactive form by gut human microbiome^{51, 52} or impact the gut bacteria⁵¹⁻⁵³. Gut bacteria genes that metabolite 41 human targeted drugs, 6 non-traditional antibacterial therapeutic and key enzymes experimented validated for 12 human diseases were mapped to our oral SGB genomic contents (**Supplementary Tables 6**). We show that many oral communities share homologous to these gut bacteria encoding enzymes, suggesting the oral microbiome may also play an importance role in medical therapy and disease development (**Fig. 7a**). More specifically, there are total 2696 SGBs contain β -glucuronidase enzyme that can metabolite anti-cancer drug Gemcitabine (2', 2'-difluorodeoxycytidine) into its inactive form⁵². 456 SGBs have agmatine gene for anti T2D drug Metformin and 225 SGBs have tyrosine decarboxylase (TyrDC) for anti-parkinson drug L-dopa⁵¹. There are also 1733 oral SGBs have genes producing small molecule taurine and 5-aminovalerate which are potential drugs for autism spectrum disorder (ASD)⁵¹. Unexpected few SGBs contain CutC/CutD genes which are key enzyme for TMA, a metabolite with high cardiovascular event risk⁵¹.

The inferration of secondary metabolites biosynthetics gene clusters (BGCs) was made by applying antiSMASH⁵⁴ pipeline. The total 12399 BGCs (7804 unknown, 4595 known) have been detected from 91.46%(1167) kSGBs and 66.75%(1544) uSGBs, and the BGCs coding for bacteriocin, arylpolyene, type III **PKS** (polyketide synthase) has appeared more than 500 times on the oral bacterial community(**Fig. 7b**, **Supplementary Table 7**). For each specie's genome, the size percentage (mean: 2.512%) of BGCs was calculated based on the each BGC's location on the each genome. The vast majority of the genome has a BGC range of less than 10% compared to the total genome (**Supplementary Figure 5**), included *Firmicutes*,

274 *Patescibacteria*, *Actinobacteriota*, *Proteobacteria*, etc. Notably, there represented 67%
 275 novel BGCs (2743 known, 5557 unknown) in the kSGBs and 54% novel BGCs (1852
 276 known, 2247 unknown) in the uSGBs. At the phylum level, *Elusimicrobiota*,
 277 *Actinobacteriota*, *Chloroflexota*, *Patescibacteria* contains a higher proportion of
 278 novel clusters (**Fig. 7c**). These unknown BGCs demonstrate the enormous potential
 279 of oral microbes for the synthesis of natural metabolites for drug development and
 280 disease treatment.

281 **DISCUSSION**

282 In summary, we provide the largest set of oral metagenomic shotgun data,
 283 assemble tens of thousands of draft genomes for the human oral microbiome,
 284 including 2,313 new species as well as many new strains of known species. The
 285 results illustrate that culturomics have not even exhausted the microbial complexity in
 286 the more accessible body sites, and that metagenomic data for large cohorts of
 287 non-fecal samples have great potential. A number of taxa with compact genomes were
 288 identified in this study, such as CPR and Mollicutes. Mollicutes such as *Mycoplasma*
 289 and *Ureaplasma* are well known in the female reproductive tract²². Much remains to
 290 be elucidated for the metabolic requirement of small bacteria in the oral microbiome.
 291 Oral bacteria also contributed to discovery of new CRISPR-Cas systems⁵⁵. Species
 292 with thousands of metagenomic and isolated genomes would be amenable to
 293 microbial GWAS⁵⁶ (microbial genome-wide association studies) to discover virulence
 294 factors, drug resistance and more commensal functions, which has so far only be
 295 possible for pathogens.

296 **Accession codes**

297 All the data are available at China National Genebank (CNCB), Shenzhen under the
 298 accession CNP0000687. <https://db.cngb.org/microbiome/>

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 301 Genebank (CNGB), Shenzhen for sample collection, DNA extraction, library
 302 construction, sequencing, and discussions.

303 **AUTHOR CONTRIBUTIONS**

304 J.W. initiated the overall health project, H.J. decided to include oral samples, Z.J., J.Z.,
 305 L.T. worked out the metagenomic assembly approach, with Hadoop support from X.L.
 306 M.H., Z.L., C.L. contributed Yunnan samples. P.C., K.C., X.W., Y.L. contributed
 307 4D-SZ samples, and L.S., X.T. managed the samples and data. J.Z., L.T., Z.J., H.J.
 308 interpreted the data, prepared the display items and wrote the manuscript. All authors
 309 contributed to finalizing the manuscript.

310 **COMPETING FINANCIAL INTERESTS**

311 The authors declare no competing financial interest.
 312

313 **ONLINE METHODS**

314 **The newly cohort and published datasets used in this study.** The 2675(2284 saliva
315 and 391 tongue) oral metagenomics samples from Chinese 4D-shenzhen
316 cohorts(**Supplementary Tables 1 sheet 4**) and the 671 salivary samples from six
317 cities and villages in Yunnan province were collected in this study(**Supplementary**
318 **Tables 1 sheet 5**), and total 706 public oral metagenomics datasets^{6, 23-26} were
319 downloaded from NCBI SRA databases with accession codes SRP029441,
320 ERP006678, SRP133047, ERP110622 and SRP07256, encompassing five different
321 studies (**Supplementary Table 1 sheet 3**) have been reported previously.

322
323 **Sample collection, DNA extraction, sequencing and quality control.** The 2955
324 salivary samples and 391 tongue samples from Shenzhen were self-collected by
325 volunteers, using a kit containing a room temperature stabilizing reagent to preserve
326 the metagenome⁵⁷. DNA extraction of the stored samples within the next few months
327 was performed using the MagPure Stool DNA KF Kit B (MD5115, Magen) from
328 1mL of each sample. Metagenomic sequencing was done on the BGISEQ-500
329 platform⁵⁸ (100bp of paired-end reads for all samples and four libraries were
330 constructed for each lane) and generated 101.4 billions pairs of raw reads. The 671
331 salivary samples from Yunnan province were self-collected using commercial kits
332 (Cat. 401103, Zeesan, China). Collected samples were temporarily stored in -80°C
333 freezers and then transported to CNGB, Shenzhen with dry ice via commercial
334 logistics (SF Express Inc.). DNA was extracted in the same way as above. Sequencing
335 was performed on the BGISEQ-500 machines and generated 26.5 billions single-end
336 100 bp length reads. The raw read length for each end was 100bp. After using the
337 quality control module of metapi pipeline followed by reads filtering and trimming
338 with strict filtration standards(not less than mean quality phred score 20 and not
339 shorter than 51bp read length) using fastp v0.19.4⁵⁹, host sequences contamination

removing using Bowtie2 v2.3.5⁶⁰ (hg38 index) and seqtk⁶¹ v1.3, we totally got 54.9
billions high-quality PE reads and 7.1 billions high-quality SE reads.

Metagenomic De novo assembly, binning and checkm. The high-quality PE and SE
reads was individually assembled using assembly module of metapi pipeline with
different max kmer cutoff by different max read length of each samples applying
SPAdes v3.13.0²⁸ (PE reads with option --meta²⁷). All configuration can see on
<https://github.com/ohmeta/metapi/blob/dev/metapi/config.yaml>. After we
got draft genomes on contig level of each samples, the reads was mapped back to each
assemblies using BWA-MEM v0.7.17⁶² with default parameters and calculate the
contig depth by jgi_summarize_bam_contig_depths²¹, then using MetaBAT2
v2.12.1²¹ to do metagenomic binning individually for each samples. Finally we got
totally 163,718 bins. After MAGs quality assignment by CheckM v1.0.12³⁰ lineages
workflow, 15,013 high-quality (completeness > 90% and contamination < 5%, HQ)
bins and 41,200 medium-quality (completeness > 50% and contamination < 10%, MQ)
bins (**Supplementary Table 2**) have been generated based on MIMAG standard²⁹.
The 16S rRNA sequences in the MAGs were searched by Barrnap v0.9⁶³ with
parameters "--reject 0.01 --evaluate 1e-3" and tRNA sequences in the MAGs were
searched by tRNAscan-SE 2.0.3⁶⁴ with the default parameters.

Public database used. The public bacteria and archaea genomes database used in this
study include (**Supplementary Table 1 sheet 6**):

(a) The NCBI Refseq bacteria and archaea databases
(<ftp://ftp.ncbi.nlm.nih.gov/genomes/refseq/>, accessed in June 2019) contain 155854
microbial genomes.

365 (b) The eHOMD³¹ database (http://www.homd.org/ftp/HOMD_prokka_genomes)
366 contain 1526 microbial genomes come from human oral environment.

367 (c) The IGGdb¹⁵ (<https://github.com/snayfach/IGGdb>) contain 23790 microbial
368 genomes come from human gut environment.

369 (d) The hSGBRep¹⁶ database contain 4930 representative microbial genomes
370 (http://segatalab.cibio.unitn.it/data/Pasolli_et_al.html) come from human body site
371 include gut, oral, skin, genital tract.

372 (e) The BPUMGs¹⁴ databases
373 (ftp://ftp.ebi.ac.uk/pub/databases/metagenomics/umgs_analyses/) contain 1952
374 microbial genomes come from human gut.

375 (f) The CGR³³ database accession code [PRJNA482748](#) contain 1520 microbial
376 genomes come from human gut bacterial culture collection.

377 (g) The HBC³² database
378 (ftp://ftp.ebi.ac.uk/pub/databases/metagenomics/hgg_mags.tar.gz) contain 737
379 microbial genomes come from human gut bacterial culture collection.

380 **Clustering metagenomic genomes into species-level genome bins.** The 56,213
381 reconstructed genomes and 190,309 reference genomes were grouped into species
382 -level genome bins(SGBs) by a two-step clustering strategy as reported previously¹⁶
383 with a slight modification. In the first step, all-versus-all genetic distance matrix
384 between the 246,522 genomes was carried out using Mash version 2.0⁶⁵ (“-k 21 -s 1e4”
385 for sketching). Then, hierarchical clustering with average linkage and 0.05 genetic
386 distance cutoff on the distance matrix by fastcluster⁶⁶ was resulted to 33008 clusters.
387 Because the Mash will underestimate the distance between the incomplete genomes⁶⁷
388 and split same-species genomes into multiple SGBs, we performed clustering base on
389 average nucleotide identity(ANI) in the second step. First, We divide the SGB into
390 known SGB(kSGB) and unknown SGB(uSGB) according to with or without reference

391 genomes. Then, a representative genome was selected for each SGBs. For the kSGB,
392 the genome which has the largest genome size was selected. For the uSGB, all MAGs
393 were rank by completeness(in decreasing order), contamination(increasing),
394 coverage(decreasing), strain heterogeneity(increasing), N50(decreasing). And
395 representative genome was selected as the one minimizing the sum of the five ranks.
396 We recalculated the more precise genetic distance using pyani v0.2.9⁶⁸(option ‘-m
397 ANIb) for the pairs of representative genomes with mash distances less than 0.95 and
398 only left ANI with genome coverage above 0.3. Following hierarchical clustering
399 with complete linkage based on >95% ANI score, 12,911 representative genome
400 which mash distances less than 0.95 were merged to 11,427 new clusters. Finally, we
401 obtained 31,525 SGBs by two-step clustering strategy. In this dataset, only 3,589
402 SGBs included eHMOD genomes and oral metagenomes MAGs were named Oral
403 SGBs and can be further divided into 2,313 uSGBs and 1,276 kSGBs. The top three
404 contributions of uSGBs are 4D_SZ (1441 uSGBs), ZhangX_2015 (445 uSGBs) and
405 Yunnan (334 uSGBs).The other 27,936 SGBs are non-oral SGBs (Figure 1b).

406 **Reconstruction of the human-oral microbiome phylogenetic structure.** The
407 phylogenetic trees of 3589 representative genomes of SGBs (Figure 3C) and 76
408 genomes of Achleplasmataceae Order were both built using the 400 PhyloPhlAn
409 markers with the parameters “--diversity high --fast --min_num_markers 80” by the
410 PhyloPhlAn2⁶⁹. As input data for PhyloPhlAn2, proteome were predict using Prodigal
411 v2.6.3⁷⁰ with default parameters. Following tools with their set of parameters were
412 used in the configuration files:

413 Diamond v0.9.22.123⁷¹ with parameters: “blastp --quiet --threads 1 --outfmt 6
414 --more-sensitive --id 50 --max-hsps 35 -k 0”;

415 Mafft v7.407⁷² with the “--anysymbol” option;

416 Trimal v1.4.rev15⁷³ with the “-gappyout” option;

417 Iqtree v1.6.12⁷⁴ with parameters: “-quiet -nt AUTO -m LG”.

The phylogenetic trees in figure 3c was generated using GraPhlAn v1.1.3⁷⁵ and the phylogenetic trees in figure 4a, figure s2 were generated using FigTree v1.4.4 (<https://github.com/rambaut/figtree/releases>).

SGBs taxonomic and function analyses. The taxonomic classification of 3589 representative genomes of SGBs was assigned using GTBD-Tk v0.3.2^{70, 76-79} (<https://github.com/Ecogenomics/GTDBTk>) classify workflow with external GTDB database release 89.0(https://data.ace.uq.edu.au/public/gtdbtk/release_89/89.0/). Although some kSGBs already have taxonomy label, we still using GTDB-Tk to classify them because GTDB-Tk has its own taxonomy classification system that is different from the NCBI taxonomy database. Then above the genus level, we manually removed the classification tag with a single letter suffix (**Supplementary Table 3**). Those suffixes used to indicate that taxon needed to be subdivided based on the current GTDB reference tree. We used EggNOG mapper v1.0.3³⁹ to do genome-wide functional annotation through orthology assignment on 3589 SGBs(**Supplementary table 3**) and 29 MAGs in Candidatus bgiplasma (**Supplementary Figure 2b**). The secondary metabolite biosynthesis gene clusters(BGCs) of 3587 oral bacterial genomes was identified respectively by using antiSMASH v5.0.0⁵⁴ with options --fullhmmer --cf-create-clusters --smcog-trees --cb-knownclusters --asf --pfam2go. Then we use the bgctk(<https://github.com/ohmeta/bgctk>) to parse and merge BGCs's results from all json file which was generated by anstiSMASH workflow.

Mapping rate compared between different oral related genomes database. The mapping rates of oral metagenomics reads align to three different oral related genomes databases(eHOMD, hSGB_Rep, oralSGB_Rep) were compared based on the statistics summary of Bowtie2's results(**Supplementary Table 4**). First we randomly

445 selected 100 oral metagenomes samples from each of 4D_SZ and Yunnan cohorts.
 446 With all 808 public samples and 81 additional verify samples which not used to
 447 assembly (**Supplementary table 1 sheet 2**), the total 1089 oral metagenomes samples
 448 were mapped to these databases respectively using Bowtie2 v2.3.5 with default
 449 parameters. The barplot of mapping rate was generated using R package ggplot2
 450 3.1.1⁸⁰ faced with different databases and different country.

451

452 **Metapi for oral SGB metagenomic profiling.** The quantification of species relative
 453 abundance of oral metagenomic samples was performed with the taxonomic profiling
 454 module of metapi pipeline: i) build the oral representative SGBs' index by Bowite2; ii)
 455 align the high-quality reads of each sample to the oral genome index using Bowtie2
 456 with parameters : "--end-to-end --very-sensitive --seed 0 --time -k 2 --no-unal
 457 --no-discordant -X 1200"; iii) The normalized contigs depths were obtained by using
 458 jgi_summarize_bam_contig_depths; vi) base on the correspondence of contigs and
 459 genome, the normalized contig depth were converted to the relative abundance of
 460 each SGB for each samples. Finally we merged all representative SGBs relative
 461 abundance to generate a taxonomic profile.

462

463 **PCOA, heatmap and oral type for metapi profile.** Principal Coordinates Analysis
 464 (Pcoa) of metapi profile was done used dudi.pco function in ade4⁸¹ R package based
 465 on bray distance from vegan2.5.2⁸² R package. The mean top 10 most abundance
 466 SGBs from every study were merged (total 27 SGBs) to visual in pheatmap⁸³ R
 467 package.

468

469 **Pangenome, phylogenetic analysis of kSGB and uSGBs.** From the taxonomic
 470 profiling results of 4820 oral metagenomic samples, the most prevalent eight genus

was selected based the rank of average relative abundance(decreasing), occurrence frequency(decreasing), oral genome number / SGBs size(decreasing), include *Prevotella*, *Neisseria*, *Streptococcus*, *Veillonella*, *Porphyromonas*, *Fusobacterium*, *Pauljensenia*, *Haemophilus*. Then we choose ten most prevalent species for each genus to do pangenome analysis. First each species has a representative genome correspond to each SGBs, so we use prokka v1.13.7⁸⁴ to do genome annotation for all genomes of each SGBs. Then the annotated genomes were used to construct pangenome database for each SGBs via panphlan_pangenome_generation.py (a script come from PanPhlAn v1.2⁸⁵). Finally the gene-family presence / absence profile matrix was transformed to a zero/one matrix for reference genomes and reconstructed genomes of each SGBs to do rarefaction analysis. Accumulation curves (Supplementary Figure 3) based on the number of core gene of each SGBs were bootstrapped ten times at each sampling interval. The observation of intra-SGB phylogenetic structure of *Neisseria* kSGB 3225, *Prevotella* kSGB 3467 and *Porphyromonas* kSGB 3273 was performed by the nonmetric multidimensional scaling analysis using the metaMDS function of R package vegan v2.5.2.

Disease markers according to the oral genomes. The metagenomics wise association between 3,589 metapi species profiles (SGB) and disease for previously published CRC and RA studies was done using generalized linear model (GLM) with adjust for potential confounders such as gender, age, BMI (Table S1). BMI is only available for RA. Species relative abundances was asin-sqrt transformed as described before⁸⁶. Non-oral SGBs were excluded. Corrected for multiple hypothesis tests was done using FDR. We predicted disease status using gradient boosting model (GBM) in caret⁸⁷ R package, such that 80% of the samples were randomly sampled for each estimator. The depth of the tree at each estimator was not limited, but leaves were restricted to have at least 30 instances. We used 4000 estimators with a learning rate of 0.002. All the FDR <1% oral marker SGBs are included in the model as predictors.

499 To avoid overfitting, 5 repeat ten folds cross validation ROC was used to measure the
500 model performance. VarImp function was used to extract the GBM importance.

501

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507

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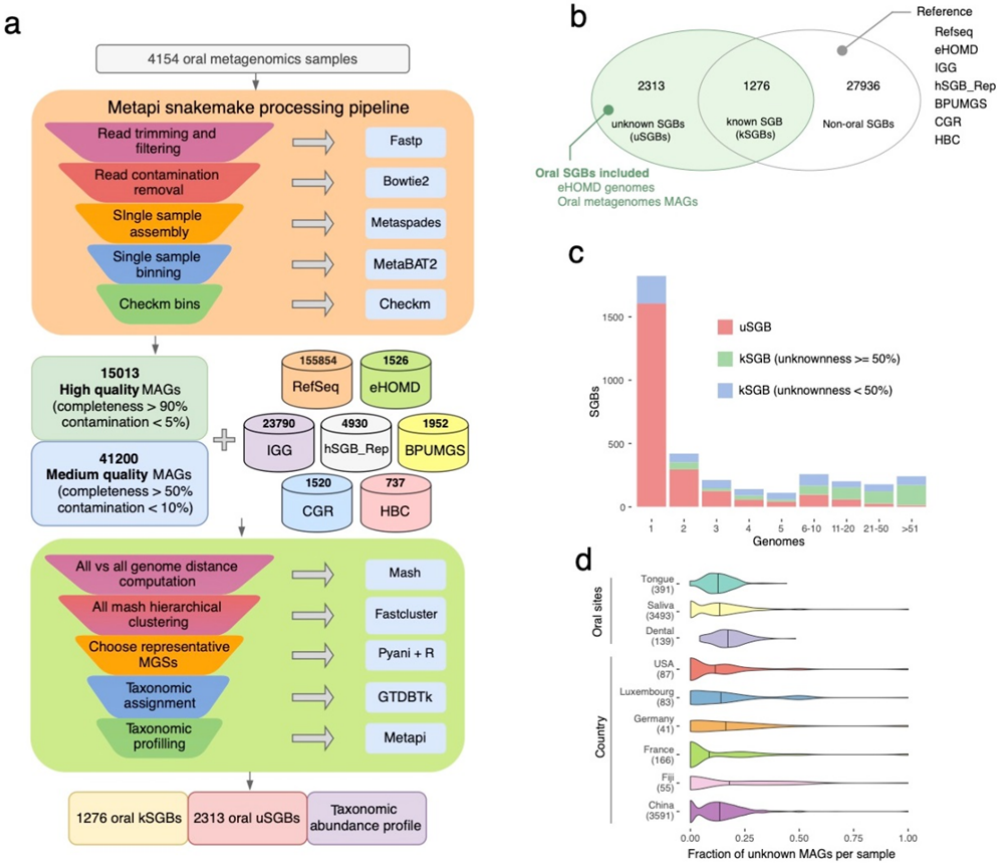
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717

718 **Figures**

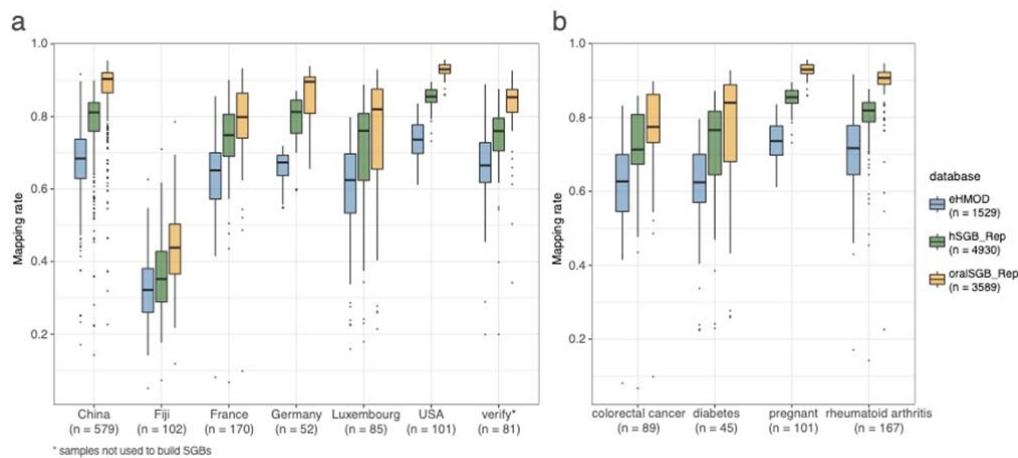


719
720 **Figure 1. 3,589 oral SGBs assembled from 4154 (3346 new sequence)**
721 **meta-analyzed oral-wide metagenomes.**

722 **a**, Species genome bins construction workflow. **b**, Overlap of oral assembled genomes
723 and reference genomes. kSGBs contains both existing microbial genomes (including
724 other metagenomic assemblies) and genomes reconstructed here. uSGBs are only
725 genomes reconstructed here and without existing isolate or metagenomically
726 assembled genomes. Non-oral SGBs contains kSGBs that are not sourced from human
727 oral samples. **c**, Genome numbers distribution of uSGBs and kSGBs. **d**, Distribution
728 of the fraction of uMAGs in each sample by oral sites and country.

729

730



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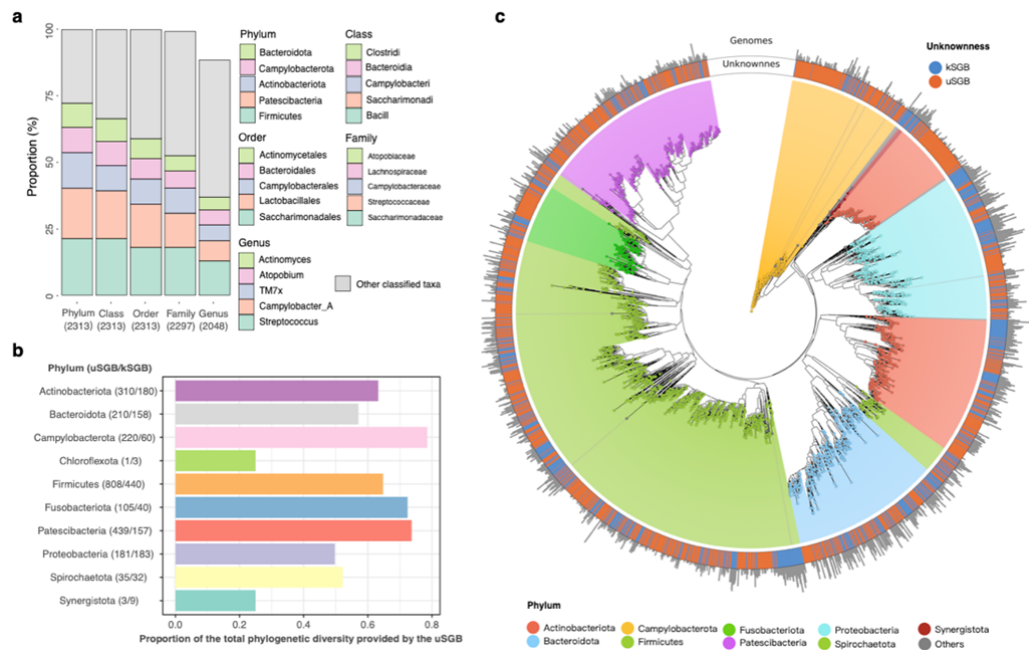
732 **Figure 2. The expanded genome set substantially increases the mappability of**
733 **oral metagenomes.**

734 The raw reads from all the 808 public samples, 100 subsampled of 2674 China
735 Shenzhen and 671 China Yunnan, and 81 additional verify samples which not used to
736 build SGBs were mapped against eHOMD, representative human SGBs (hSGB_Rep)
737 and representative oral SGBs (oralSGB_Rep). Among three databases, our
738 representative oral SGBs have the highest raw-read mappability in all country and
739 verify datasets.

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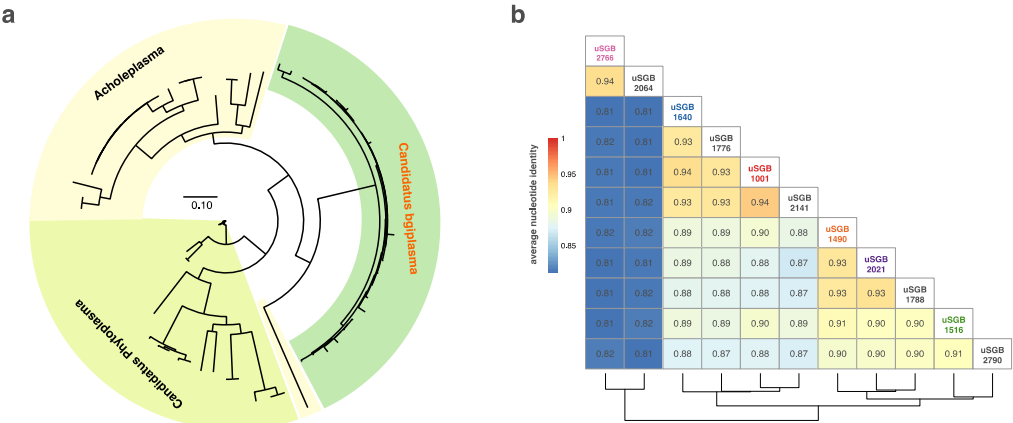
742 **Figure 3. Phylogeny of representative oral SGBs**



743

744 **a**, Taxonomic composition of the 2,313 uSGB, Only the five most frequently
745 observed taxa are shown in the legend, with the remaining lineages grouped as “other
746 classified taxa”. **b**, Proportion of the total phylogenetic diversity provided by the
747 uSGB. **c**, oral-associated microbial phylogenetic tree of representative genomes from
748 3589 species-level genome bin (SGB).

749



750

751 **Figure 4. A new candidatus family is found within the Acholeplasmataceae**

752 **Order.**

753 **a**, phylogenetic tree from all MAGs in the new candidatus family(Candidatus

754 bgiplasma) and known genomes in Acholeplasmataceae Order. Supplementary Figure

755 2a reports the detail of phylogenetic tree in Candidatus bgiplasma. **b**, Average

756 nucleotide identity(ANI) between all uSGB in the *Ca. bgiplasma* represent clearly two

757 genus clades which ANI less than 0.85. Unknown SGBs without HQ MAGs are left

758 black.

759

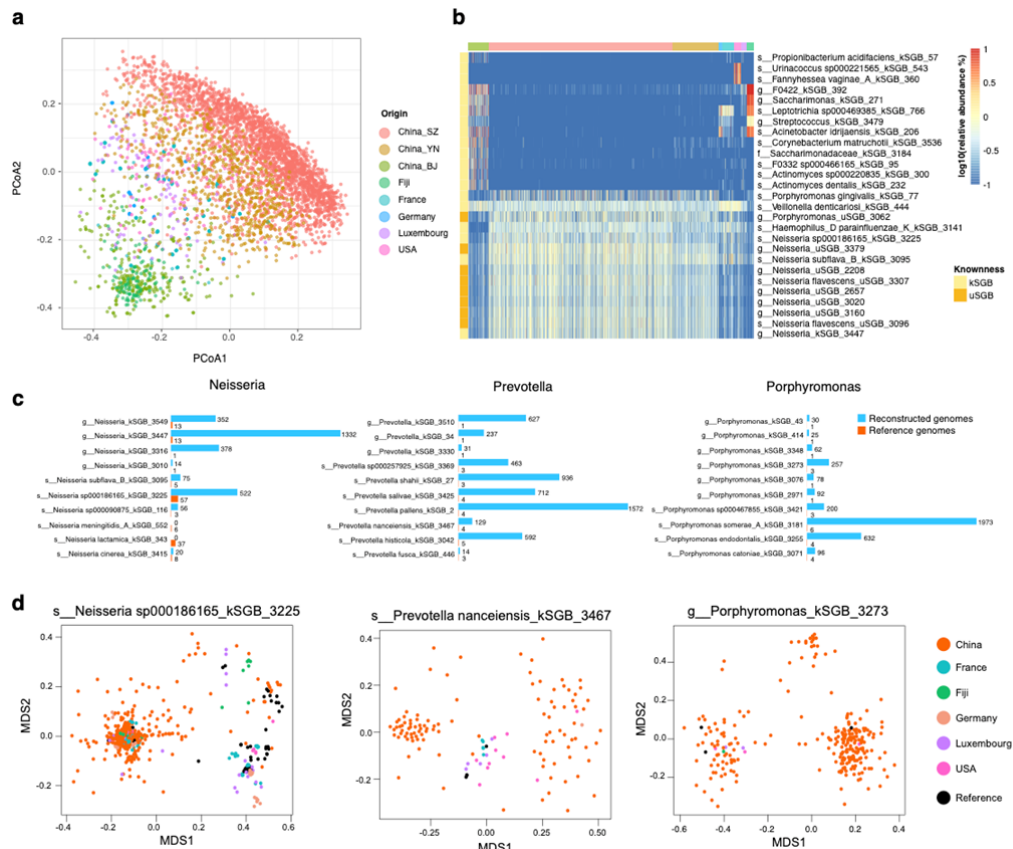


Figure 5. Geographical distribution of oral SGBs and strains.

a, Principal coordinate analysis plot based on Bray-Curtis distances of oral SGB relative abundance profile highlights distinct microbial communities among different origin populations. **b**, The relative abundance of top 10 most abundant SGBs from each origin populations. A large set of reconstructed uSGBs are widely high abundance distribution in our cohort (Shenzhen and Yunnan) and lack of several highly abundant kSGBs in other population. Species are order by hclust with complete linkage and euclidean distance. **c**, Our reconstructed MAGs largely extend the size (genome numbers) of the top 10 most abundance species from common oral genus with few reference genomes. **d**, Multidimensional scaling on average nucleotide identity between MAG and reference genomes in species showed strain variety and that constructed MAGs dominated the sub species. Only HQ MAGs and reference genomes are showed.

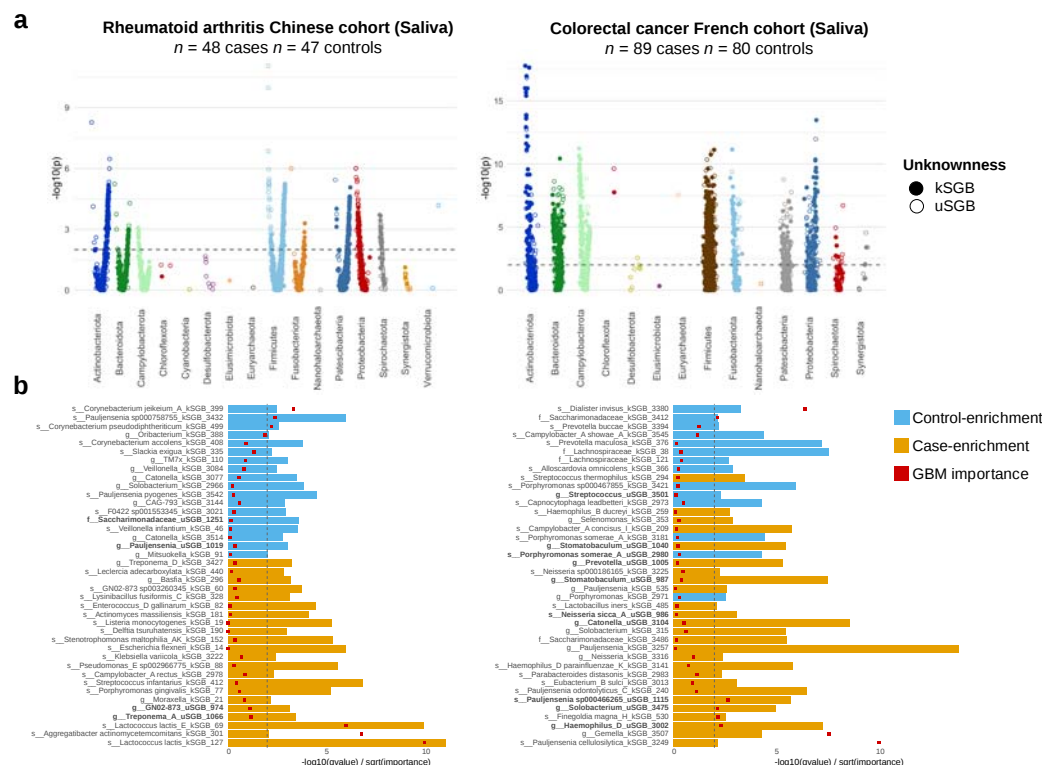
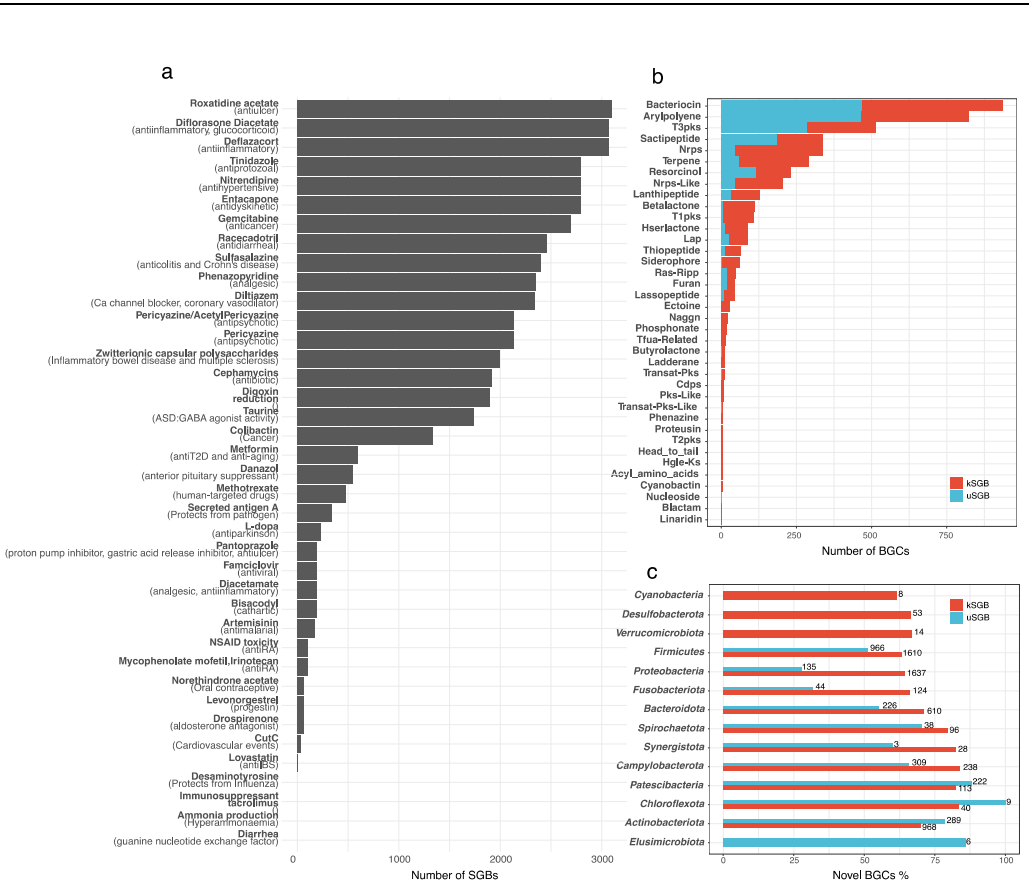


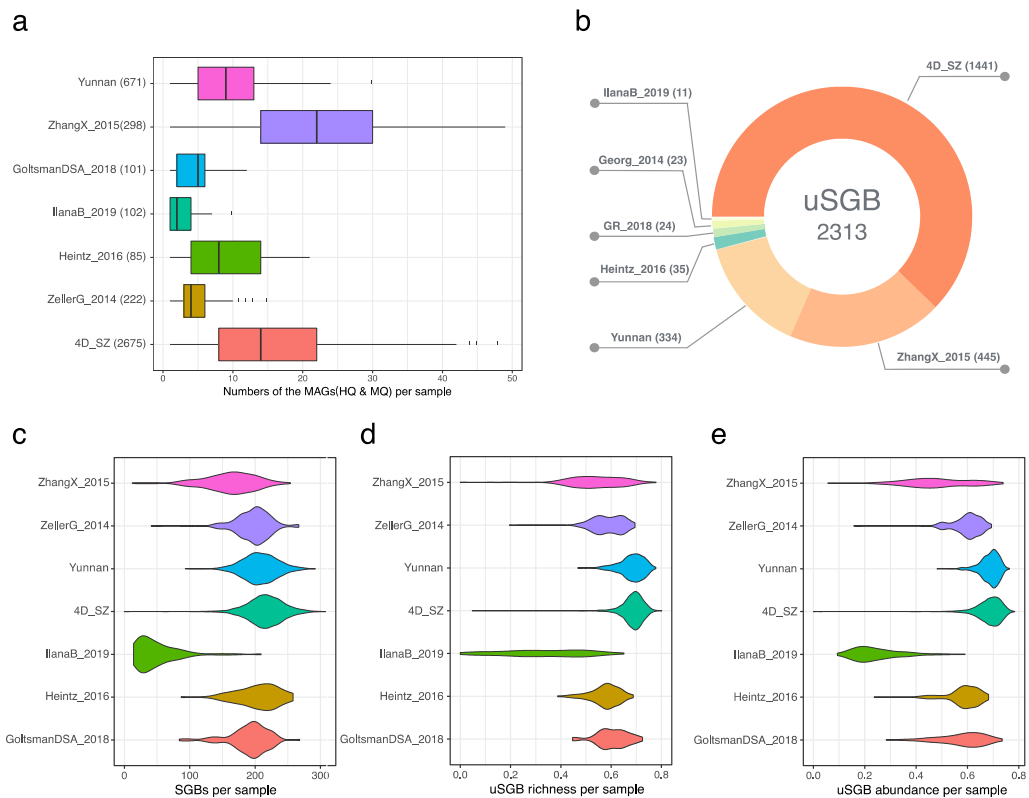
Figure 6. Disease markers according to the oral genomes.

a, The Manhattan plot shows metagenomic wise association of oral SGBs for RA and CRC studies. The species are ordered according to their phylogeny (bottom) and the association direction (positive or negative). Each point is one SGB and point height indicates the FDR value correction for multiple hypothesis tests from a generalized linear model (GLM) test between diseased and healthy species abundance after adjusting Age, Gender, BMI. The dotted line indicates a false discovery rate (FDR) of 1%. **b**, SGBs association with RA and CRC. We select 40 large and most importance oral SGBs (>10 genomes) for disease prediction using Gradient Boosting Machine (GBM). The species are order according to their partial spearman correlation adjusted age, gender, BMI and GBM importance. The bar length indicated the FDR value between groups as described above. The dotted line indicates a false discovery rate (FDR) of 1%. The red square in bar is the sqrt GBM importance. uSGBs are highlight in bold label text.



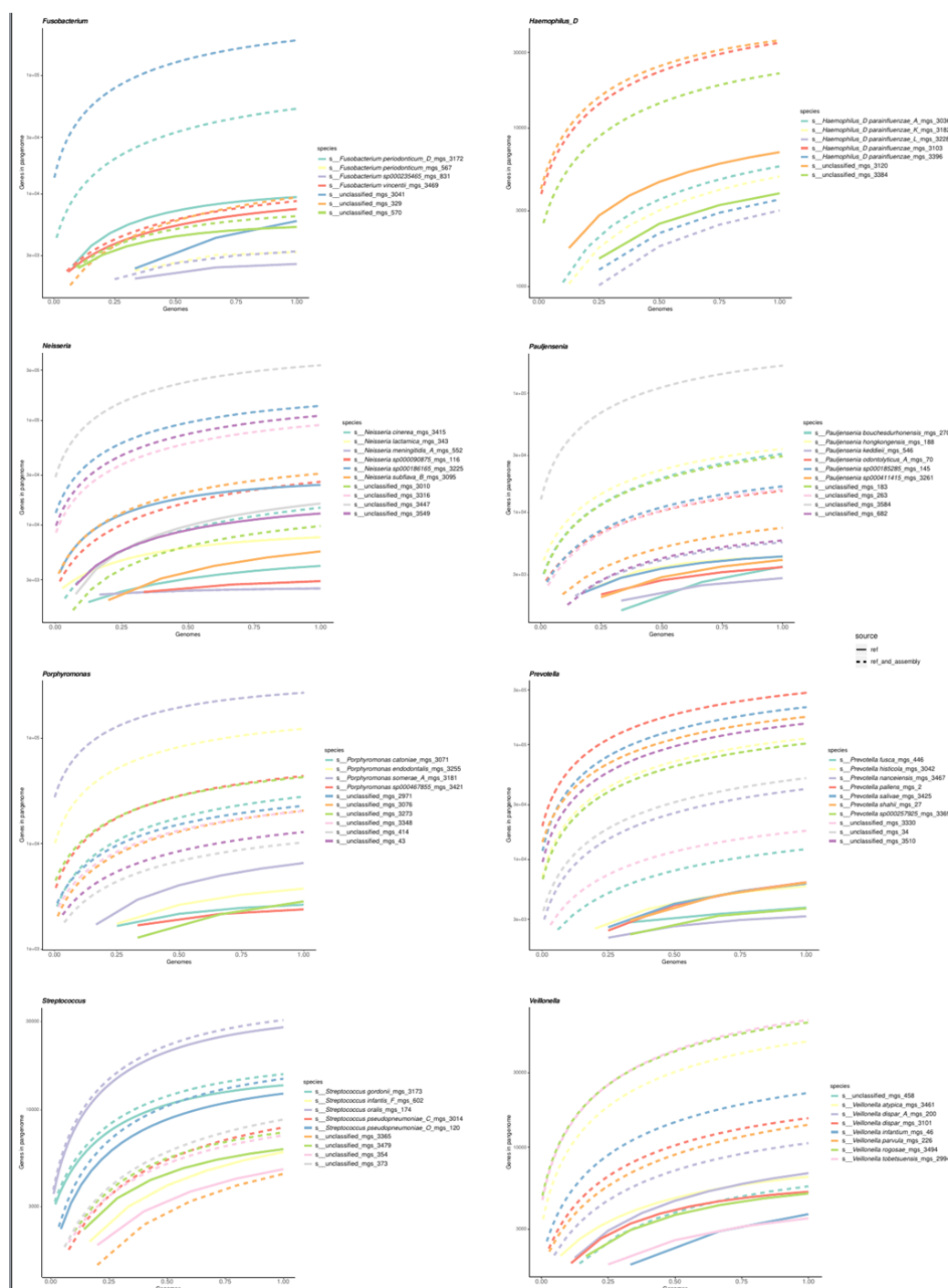
791
792 **Figure 7. A comprehensive mapping of the function repertoire of the human oral**
793 **microbiome.**
794 **a**, Number of oral SGBs who share homologous to gut microbiome enzyme coding
795 drug metabolite or healthy related function. Details see Supplementary Table 6. **b**,
796 Summary of predicted BGCs in oral microbiome. Numbers of BGC of 38 different
797 types detected in the 2711 oral bacterial genomes were grouped by kSGB and uSGB.
798 **c**, Fraction of novel BGCs across phylum levels. The numbers of bar show the novel
799 BGC count while bar length represent kSGBs/uSGBs ratio on the phylum levels.
800

801 **Supplementary Figures**



802
803 **Supplementary Figure 1. Summary of assembly quality, SGBs distribution**
804 **across 7 studies.**
805 **a**, Numbers of medium quality and high quality MAGs in samples from 7 studies. **b**,
806 3,589 uSGBs origin distribution across studies. **c**, The number of all SGBs (>0.001
807 abundance) for each samples. **d**, uSGB richness (number of uSGB/number of all SGB)
808 for each sample. **e**, Sum of all uSGB abundance for each samples.

809

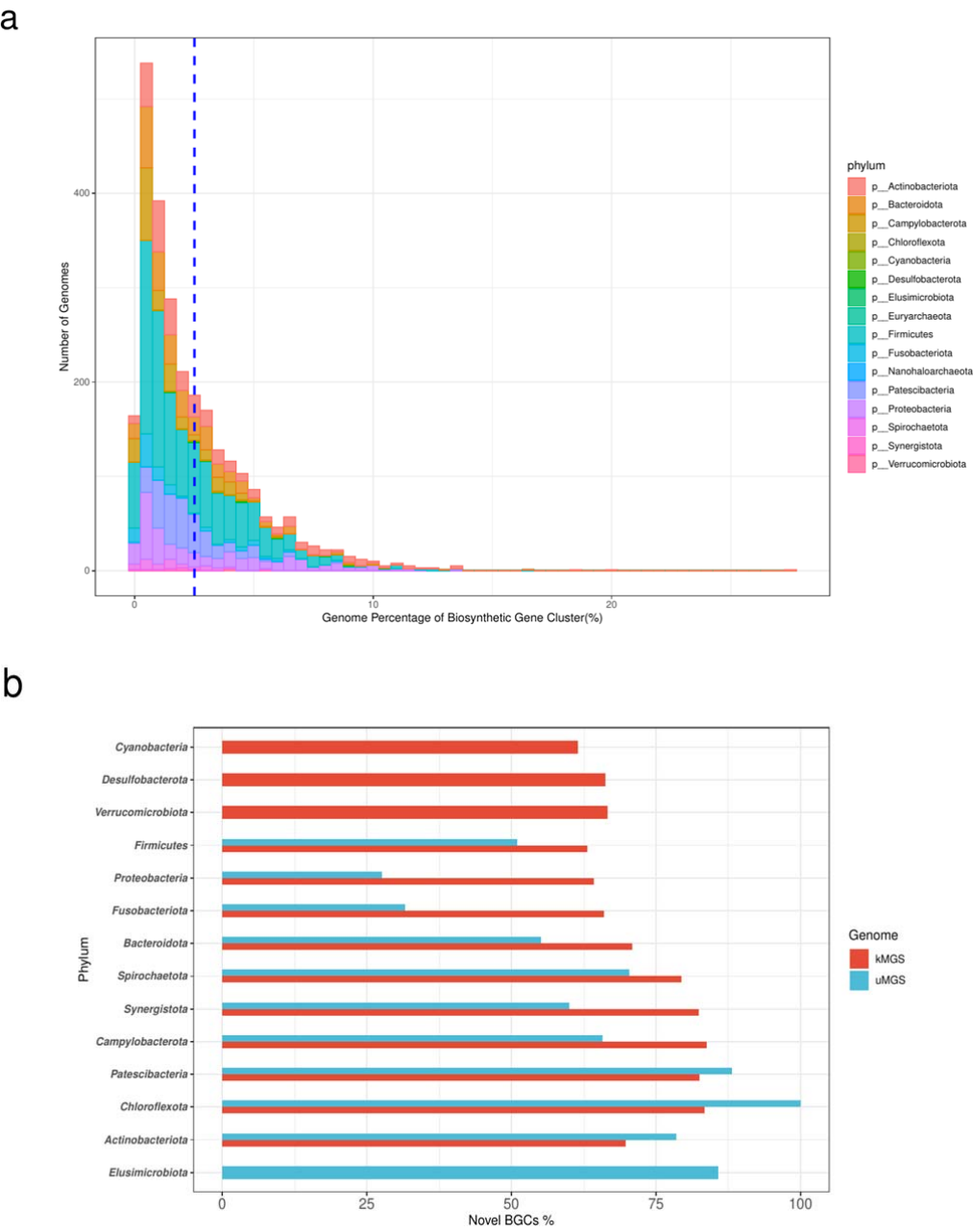


819

820 **Supplementary Figure 3. The pangenome genetic diversity.**

821 The gene number rarefaction curve indicated that the genetic diversity have been
822 increased through more genomes metagenomically assembled, included 71 species
823 genomes come from eight most prevalent genus.

824



830

831 **Supplementary Figure 5. The detection of BGCs on the human oral microbiome**

832 **a**, The distribution of genome percentage of biosynthetic gene cluster. From oral
833 microbiome included 2713 genomes form 16 different phylum. The x coordinate
834 corresponds to the proportion of BGC size, and the y coordinate corresponds to the
835 number of genomes. The blue vertical line indicates the average proportion of BGC
836 size: 2.512%. **b**, Novel BGCs proportion on phylum level of oral microbiome. The x

837 coordinate corresponds to the proportion of the number of the novel BGCs, the y

838 coordinate corresponds to the 14 different phylum grouped by uSGB and kSGB.

839