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Ho, Jeffery; Yeoh, Yun Kit; Barua, Nilakshi; Chen, Zigui; Lui, Grace; Wong, Sunny H; Yang, Xiao; Chan, Martin Cw; Chan, Paul Ks; Hawkey, Peter M; Ip, Margaret

DOI: 10.1080/19490976.2019.1700755

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Document Version Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Ho, J, Yeoh, YK, Barua, N, Chen, Z, Lúi, G, Wong, SH, Yang, X, Chan, MC, Chan, PK, Hawkey, PM & Ip, M 2020, 'Systematic review of human gut resistome studies revealed variable definitions and approaches', *Gut Microbes*, pp. 1-12. https://doi.org/10.1080/19490976.2019.1700755

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Gut Microbes



ISSN: 1949-0976 (Print) 1949-0984 (Online) Journal homepage: https://www.tandfonline.com/loi/kgmi20

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To cite this article: Jeffery Ho, Yun Kit Yeoh, Nilakshi Barua, Zigui Chen, Grace Lui, Sunny H Wong, Xiao Yang, Martin CW Chan, Paul KS Chan, Peter M Hawkey & Margaret Ip (2020): Systematic review of human gut resistome studies revealed variable definitions and approaches, Gut Microbes, DOI: <u>10.1080/19490976.2019.1700755</u>

To link to this article: https://doi.org/10.1080/19490976.2019.1700755

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REVIEW

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Systematic review of human gut resistome studies revealed variable definitions and approaches

Jeffery Ho^{a,b,c*}, Yun Kit Yeoh ^{a,b,c*}, Nilakshi Barua ^a, Zigui Chen ^{a,b,c}, Grace Lui^{b,c,d}, Sunny H Wong^{b,c,d}, Xiao Yang ^a, Martin CW Chan ^a, Paul KS Chan ^{a,b,c}, Peter M Hawkey^e, and Margaret Ip ^{a,b,c}

^aDepartment of Microbiology, The Chinese University of Hong Kong, Hong Kong SAR, China; ^bCentre for Gut Microbiota Research, The Chinese University of Hong Kong, Hong Kong SAR, China; ^cLi Ka Shing Institute of Health Sciences, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong SAR, China; ^dDepartment of Medicine & Therapeutics, Faculty of Medicine, Prince of Wales Hospital, The Chinese University of Hong Kong, Hong Kong, China; ^eInstitute of Microbiology and Infection, University of Birmingham, Birmingham, UK

ABSTRACT

In this review, we highlight the variations of gut resistome studies, which may preclude comparisons and translational interpretations. Of 22 included studies, a range of 12 to 2000 antibiotic resistance (AR) genes were profiled. Overall, studies defined a healthy gut resistome as subjects who had not taken antibiotics in the last three to 12 months prior to sampling. In studies with *de novo* assembly, AR genes were identified based on variable nucleotide or amino acid sequence similarities. Different marker genes were used for defining resistance to a given antibiotic class. Validation of phenotypic resistance in the laboratory is frequently lacking. Cryptic resistance, collateral sensitivity and the interaction with repressors or promotors were not investigated. International consensus is needed for selecting marker genes to define resistance to a given antibiotic class in addition to uniformity in phenotypic validation and bioinformatics pipelines.

ARTICLE HISTORY

Received 9 July 2019 Accepted 28 November 2019

KEYWORDS Resistome; antibiotic resistance; gut microbiota; meta-genomics; fecal microbiota transplantation

Introduction

Antibiotic resistance is a global public health concern and thus recognition of its reservoirs could facilitate the control of its dissemination. Traditionally, studies of antimicrobial resistance (AR) genes in bacteria started from massive screening of antibiotic resistance phenotypes using macroor micro-broth dilution methods.¹⁻³ For bacterial isolates displaying resistance phenotypes, total genomic DNA is extracted and probed for candidate AR genes using polymerase chain reaction (PCR)-based method or comparative whole-genome sequencing with reference to sensitive type strains.² If a candidate resistance gene could not be identified, genomic DNA is cloned into expression vectors and then transformed into a heterologous sensitive bacterial host for molecular and phenotypic characterization.³ In addition to being timeconsuming, this technique is limited to bacteria that can be cultivated.

In recent years, the advent of affordable highthroughput sequencing and analysis applied to online antibiotic resistance gene databases enables the avoidance of bacterial culture, facilitating massive resistome-wide studies of potential reservoirs of antibiotic resistance genes.^{4–8} The ubiquity of resistance genes was exemplified by their unanticipated isolation from various environmental habitats waste water treatment plant and soil to food production chain and wild animals.^{9–11}

The term *resistome* is defined as the complete collective assemblage of antibiotic, antiseptic and heavy metal resistance genes in a microbial ecosystem.¹² Studies have applied this definition variably with considerable heterogeneity in candidate genes, methodology and bioinformatics pipelines, precluding direct comparison across studies.^{5–7,13,14} Several reports employed high-throughput microfluidic PCR or customized microarrays by designing probes with reference to existing AR gene databases and targeted

CONTACT Margaret Ip 🔊 margaretip@cuhk.edu.hk 🗈 Department of Microbiology, Chinese University of Hong Kong, Prince of Wales Hospital, Shatin, Hong Kong

*These authors contributed equally to this work

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a selection of resistance genes,¹⁵⁻¹⁷ with the resulting number of AR genes varying from less than 50^{7,14,18} to thousands.^{15,19} Across studies, disparate genes were selected to represent resistance to a given antibiotic class.^{7,14,16,20} Another issue is in comparing gene and/or protein sequence similarities and defining functional conservation. In sequence-based meta-genomic studies, de novo assembled contigs were compared to the existing AR gene databases. Where the assembled contigs displayed sequence similarity beyond a defined threshold, phenotypic resistance was assumed. The degree of similarity was arbitrary, ranging from 80% of amino acid sequence identity⁶ to 95% nucleotide sequence identity.^{21,22} The function of the detected resistance genes was only occasionally validated by expressing in competent bacterial hosts.^{8,19,20,23} It is well-established that a single amino acid substitution could sufficiently change the susceptibility level of a strain by altering the bindaffinity of the drug target site to ing a corresponding antibiotic.²⁴ In cases where up to 20% difference in the amino acid sequence was observed, the resistance phenotype of the bacterial strains concerned would therefore be questionable. In sequence-based meta-genomic studies, the genetic context of the resistant genes, or the flanking regions containing promoter- and repressor-sequences that define the bacterial host origin, was not consistently reported. This should provide important information on possible bacterial hosts, gene expression, and horizontal transferability. The possible bacterial hosts for a given set of antibiotic resistance genes were occasionally predicted using network analysis lacking downstream resistance phenotype validation.²⁵

We reviewed the literature of the human gut resistome determined in healthy populations from various geographical locations. We highlight the characteristics of these studies; the methodologies used in their analyses, and the range of the magnitude of the resistome. We also identify variables that require careful definition and the adoption of a consensus to enable comparisons to be made across studies.

Methods

Bibliographic search and study selection

Search strategy was developed using the MEDLINE Ovid platform (Table S1), which was then translated for use in electronic databases EMBASE, MEDLINE (PUBMED), Web of Science, and Scopus. Briefly, relevant articles were identified using Medical Subject Heading (MeSH) or Title/Abstract keywords from inception to May 2019. All citations were imported into Endnote X8 software by which duplicates were removed. Two researchers (JH and NB) performed article screening and data verification, respectively, according to predefined inclusion and exclusion criteria. Disagreement on study selection was resolved through discussion. This study was reported according to the Preferred Reporting Items for Systematic Reviews and Metaanalysis (PRISMA) statement.²⁶

Inclusion and exclusion criteria

All primary studies with human fecal specimens collected from healthy population or secondary analyses of its kind were included. Studies on population with high risk for carrying antibiotic resistance genes, such as the critically ill, were excluded. Conference abstracts, editorials, systematic reviews, and meta-analyses were excluded. Other exclusion criteria were set in terms of study population, publication type, and study type. The pre-defined inclusion and exclusion criteria are summarized in Table S2.

Data extraction and quality assessment

Full text of the eligible studies were reviewed and the following data extracted: First author, year of publication, study design, number of fecal specimens, specific antibiotic resistance genes, laboratory methods, and bioinformatic pipelines. Whenever appropriate, the corresponding authors would be contacted by e-mail for clarification. The quality of the included studies was rated by subitems of Newcastle–Ottawa scale for nonrandomized observational studies, a tool with established content validity and inter-rater reliability.²⁷ The representativeness of the study cohort and the ascertainment of antibiotic exposure were determined. Two independent reviewers (JH and NB) rated the studies according to the scale. To ensure reliability of the review process, 10% of the articles in the final data set were reviewed by another reviewer (MI). Any discrepancies were resolved by discussion.

Effect sizes and publication bias

Despite multiple reports of the prevalence of antibiotic resistance genes on the same population were identified, the small sample size and inconsistent use of reference genes precluded such an effect size from pooling. In terms of publication bias, a traditional funnel plot using logarithm of effect size against standard error does not apply to proportion data sets due to its inherent nature of asymmetry for extreme proportions. Neither classic fail-safe N nor Orwin's fail-safe N would be applicable for prevalence data because statistical significance or trivial levels do not exist for proportion estimates.

Results and discussion

Study selection, countries of origin, cohort representativeness, and exposure ascertainment

The initial search returned 107 articles. After screening titles and abstracts, 86 full-texts were reviewed. Nineteen of these met the inclusion criteria but were not exempted from the exclusion criteria. Three additional studies were identified by manually checking reference lists of these 19

studies (Figure 1). Of the 22 studies included in this review (Table S3), 15 were primary studies involving subject recruitment and specimen collection.^{4,6,7,14–20,22,23,28–30} These covered populations originated from 18 countries, namely Afghanistan, Canada, China, Denmark, Eritrea, Finland, France, Germany, Iraq, Italy, Norway, Peru, Spain, Syria, Tanzania, The Netherlands, United Kingdom, and the United States. No studies were conducted in 90% of the countries as defined by the United Nations which included more than 193 countries. The remaining seven studies were re-analysis of publicly available datasets.^{5,8,13,21,25,31,32} With reference to the number of population by countries reported by the United Nations,³³ the number of recruited subjects ranged from 0.72 in China to 765 in the Netherlands per 10 million population (median = 19.07, interquartile range 4.56-37.10) (Figure 2). Except for a Dutch study for which a general population-based cohort involving 1,135 individuals derived from 167,000 subjects across three provinces,¹⁸ the remaining studies were either based on convenient samples or did not describe how the cohorts were derived. Antibiotic intake is one of the key factors that may alter the collection of antibiotic resistance genes or resistome.¹⁶ In this connection, ascertainment of antibiotic exposure prior to fecal specimen collection is important. Nonetheless, the period free from antibiotic exposure ranged from 3 months,³ 6 months,^{16,22} to 12 months.^{29,30} Notably, the remaining primary studies did not describe antibiotic exposure in their population. However, they did not report higher number of resistant genes. The details of the included studies are presented in Table 1.



Figure 1. Study selection.



Figure 2. Geographical distribution of studies included in this review. The color intensity indicates the number of participants sampled per 10 million population by countries according to data in the World Data Bank, the United Nations.

Ambiguous definition of a healthy human gut resistome

Human gut resistomes are by large defined as the collection of antibiotic resistance genes present in the gut microbiota.^{14–19} This collection varied from 10 AR genes encoding resistance to five drug classes (*β*-lactams, tetracyclines, macrolides, aminoglycosides, and quinolones)¹⁴ to more than 2,900 AR genes classified into 369 gene types.¹⁵ These 369 AR gene types could be broadly classified into resistance to six drug classes including βlactams and tetracyclines.¹⁵ The number of AR genes surveyed for a given antibiotic class also varied considerably. For instance, more than 30 tetracycline resistance genes were tested in one study¹⁵ but only *tetM* and *tetQ* were investigated in another.¹⁴ Nonetheless, they both reported the detection rate of tetracycline resistance in the gut resistome, which was initially limited by the number of genes surveyed. Interestingly, the highly prevalent tetW gene in healthy subjects¹⁶ was not investigated in another study targeting healthy people after international travel.¹⁴ While the interaction between AR genes per se and their repressors or promoters is well documented, the inclusion of transposons, integrons, and AR geneassociated gene regulators has been variably reported. Several studies did not detect AR geneassociated genetic elements.^{14,16,20} No study has considered collateral sensitivity and silent AR genes in the healthy gut resistome. In metagenomic studies where de novo assembly was used

to detect novel AR determinants, the identity ranged from less than 60% amino acid sequence identity¹⁹ to more than 99% nucleotide sequence similarity to define recently horizontal transferred elements.²¹ A stringent threshold could potentially reduce the number of AR genes identified, and vice versa. Collectively, the difference of the number of AR genes, choice of marker AR gene to represent a functional antibiotic class and the degree of sequence identity to define a putative determinant hinder comparison across studies.

Epidemiological factors shaping the healthy gut resistome

At an individual level, the gut resistome landscape is largely shaped by age,¹⁵ sex,¹⁸ co-morbidities,¹⁶ gut microbiota composition,25 living conditions such as household environment,¹⁹ and daily life exposure.⁶ For instance, the Hadza hunter-gatherer had a gut resistome profile similar to that in soil.⁶ While antibiotic exposure was anticipated to alter the human gut resistome in people with existing co-morbidities,¹⁶ the gut resistome appears to be highly resilient in healthy subjects following short-term experimental exposure to antibiotics.^{17,22} At the population-wide level, a single ecological study suggested that the gut resistotype was associated with the country-level consumption of antibiotics.²⁸ In the 21st century, the mobility of humans across countries also accounts for the variation of gut resistome. Despite an initially low prevalence of $bla_{\text{CTX-M}}$ (9%) and qnrB (6.6%)

Ref.	. Country(ies)	Sample size	Method(s)	ARG (N) ^a	Pipeline(s)	Reference Database(s)	Key findings	Limitations
4	Canada	24	SeqMeta	13,218	RAY Meta	MERGEM	Culture-enrichment in prior to sequencing allows	Enrichment step has eliminated otherwise
					2.2.1		identification of core and accessory ARGs in rare gut microbial	prevalent Akkermansia and Prevotella.
5	Demark, Spain	663	SeqMeta	6,095	Prodigal, CD-HIT v4 5 7	Resfinder, ARG- ANNOT, Lahey Clinic RFD-DR	species. The predicted ARGs had 29.8% amino acid identity to known ARGs, supporting ARG transfer between microbes is rare.	ARGs in the subdominant microbial species and that non-homologous to the known ARG classes may be misced
9	Tanzania, Italy	38	SeqMeta	2,570	TBLASTN	ARDB; ResFams	The gut resistome of the Hadza hunter-gatherer resembles the soil, with less syntenic mobile elements than that of the Italian.	Survey of ARGs was limited to known genes. Functional validation was not performed.
~	Syria, Iraq, Iran, Eritrea, Afghanistan, Germany	606	Mf-PCR	42	NA	AN	Refugees had more than five ARGs, with high prevalence of bla_{CX-M} , bla_{SHV} , bla_{oxa-1} , $vanC1$, and $qnrB$. German had three or less ARGs.	Mobility of the ARGs were not investigated and thus their transferability was unclear.
∞	China, Denmark, Spain	162	SeqMeta; FunMeta	1,093	BLASTx	ARDB	Chinese had the highest abundance of ARGs, which are phylogenetically distant from others.	The limited samples may not reflect population-wide profile.
13	USA, China, Spain, Denmark	1,267	SeqMeta	112	BLASTx, SOAP	ARDB, ARG- ANNOT	Fig. 5 and the similar among all population studied. Chinese had the hichest ARG shundare than others	Insufficient epidemiologic data that supported out ABG being affected by antibiotic policy
14	The Netherlands	122	PCR	10	NA	NCBI	The prevalence of <i>blactry</i> , <i>qnrB</i> and <i>qnrS</i> increased from 6.6% to 9% before travel to 33.6% to 55.7% after	Small study population size precluded risk analysis.
15	China	77	Microarray	2,915	NA	ARDB	international travel. ARG diversity is related to age.	False-positive or -negative hybridization signals may exist
16	The Netherlands	10	Mf-PCR	81	NA	ResFinder, NCBI	The catA and tetW genes were more abundant in healthy subjects than the critically ill.	The findings may not apply to countries where resistant bacteria is prevalent.
17	Canada	70	Microarray	254	NA	CARD, ResFinder, ARG-ANNOT	Gut resistome in healthy adults showed prompt recovery to baseline in one week after antibiotic perturbation.	The microarray did not include all known ARGs.
18	The Netherlands	1,179	SeqMeta	11	ShortBRED	CARD	Females had higher diversity of ARGs and prevalence of <i>Inu</i>	Small number of male subjects may be
19	Peru	263	SeqMeta	1,100	ShortBRED	CARD, Lahey	The resistome from human gut and their associated animals and environment are shared	No functional validation of the ARGs identified.
20	China, Europe, USA	2,768	SeqMeta	19,569	SOAP- denovo	ResFinder	Mobile resistomes are mainly found, in order of frequency, in <i>Proteobacteria</i> (399 genes), <i>Firmicutes</i> (86), <i>Bacteroidetes</i> (46), and <i>Actinobacteria</i> (40).	Nucleotide identity cutoff was set at 95% against ResFinder.
21	Caucasian male	12	SeqMeta	NA	CARD-RGI	CARD, ResFams,	Healthy subjects with glycopeptide or aminoglycoside resistance genes were prone to <i>de novo</i> colonization upon antibiotic exposure.	The annotation algorithm was conservative.
22	UK, France, Italy, Finland, Norway, Scotland	22	Microarray; FunMeta	NA	Chromas, DNAMAN	NCBI	The <i>tet(W)</i> gene and transposons Tn4451 and Tn1549 were highly prevalent in fecal metagenomes.	Only tetracycline and erythromycin resistance genes were investigated.
23	Finland, France, Italy, Norway, UK	100	PCR; Microarray; FunMeta	97	NA	NA	ARGs identified by microarray were not recovered by functional screen. Combining these methods increase the identifiable ARG repertoire.	The cloned ARG in functional screen needs to be expressive in the heterologous host.
25	11 countries	180	SeqMeta	1,209	ARGs-OAP	SARG	The <i>ermF</i> was the representative macrolide resistant gene in the Chinese population.	Bacterial hosts of ARGs were predicted by network analysis.
								(Continued)

Table 1. Summary of studies included in this review.

GUT MICROBES 😔 5

Tab	e 1. (Continued).							
		Sample		ARG		Reference		
Ref.	Country(ies)	size	Method(s)	(N) ^a	Pipeline(s)	Database(s)	Key findings	Limitations
28	Denmark, Spain, USA	252	SeqMeta	3,496	MOCAT	ARDB	The most prevalent human gut ARGs were those that encoded resistance to antibiotics used in animals in	Only three developed countries were involved.
29	USA	2	FunMeta	NA	NA	NCBI	a country-specific manner. Half of the resistance genes recovered from aerobic gut icolates were identical to that in nathorous	The majority of gut commensals was
30	Germany	2	SeqMeta	NA	DIAMOND	CARD	The gut resistome in two persons changed differently in response to ciprofloxacin treatment.	Given sequence reads did not span the whole ARG length, its frequency could not be
31	USA	29	SeqMeta	21	Prodigal, BLACTD	DeepARG	Sixty four ARGs were shared amongst the gut resistomes from	identified. The genetic contexts of the ARGs were unclear.
32	Malawi, USA, Venezuela	110	SeqMeta	70	BLAST+	ARDB, ResFams	Antibiotic resistance and virulence genes are correlated and may be co-selected.	Results were based on sequence analysis without downstream laboratory validation.
^a Nur Mc SO	hber of antibiotic re: bile Elements and Re AP, short oligonuclec	sistance gé sistance G stide analy	enes (ARG) sun enes Enhanced sis package.	veyed o for Met.	r annotated; :agenomics; N	FunMeta, functiona Af-PCR, microfluidic	al metagenomics; Marilyn, Marilyn Robert's website for macrolic PCR; PATRIC, Pathosystems Resource Integration Center; Ref, refe	des and tetracycline resistance genes; MERGEM, arence; SeqMeta, sequence-based metagenomics;

before departure from the Netherlands, a country with low prevalence of antibiotic resistance, as many as 33.6% and 36.9% of the Dutch travelers acquired *bla*_{CTX-M} and *qnrB* genes, respectively, following travel to Southeast Asian and the Indian subcontinents.¹⁴ Cross-country migrations of refugees and asylum seekers also potentially altered the gut resistome amongst the local population due to higher prevalence of resistance genes in such developing countries as Afghanistan, Iraq, and Syria.⁷ A recent study in the UK has shown that the two major risk factors for carriage of CTX-M in the healthy community are travel to an area of high occurrence and birth-origin in such countries which is associated with frequent travel to those areas of the world.^{34,35}

Contemporary approaches to investigate human intestinal resistome

Essentially, three approaches of studying the gut resistome have been identified. Classically (and historically), a selected pool of AR genes was detected by either PCR or customized microarrays.^{7,14–17s} Different marker AR genes were chosen to represent resistance phenotype to a given antibiotic class.^{14–17} These methods were also limited to the small number of AR genes possibly be detected, reducing the representativeness of the AR gene repertoire. The second approach employed total bulk DNA directly isolated from samples, for which only known AR or related genes with similar sequence identity were detected. In this method, DNA was extracted from fecal specimens and sequenced with a depth of 1.2Gb to 12.6Gb per sample.^{4,19} Contigs were then assembled based on user-defined settings, the sequence of which was compared against existing antibiotic resistance gene databases such as Comprehensive Antibiotic Resistance Database (CARD),^{18,19,22,24} Lahey beta-lactamase database,¹⁹ Antibiotic Resistance Gene-ANNOTation (ARG-ANNOT),¹³ Antibiotic Resistance Database (ARDB),^{13,25,28} and Resfinder.²¹ A combination of these databases could also be included in a study to increase the detection rate of AR genes.^{13,19,30} Based on local AR gene distribution, additional augmentation could also be possible.²⁸ The third approach entails direct fragmentation of DNA extracted from fecal samples and cloned into heterologous hosts such as E. coli using shuttle vectors including plasmids,¹⁹ fosmids⁸ or bacterial artificial chromosomes,

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depending on the insert size.^{20,23} This method has the advantage of identifying novel AR genes and confirming the role of the AR genes albeit its expression may differ in the heterologous host. Collectively, targeted AR gene detection will be inexpensive, allowing surveys be conducted in resource-limited settings such as low- and middle-income countries. In places where affordability is not a concern, an initial sequencebased metagenomic survey can be conducted to identify putative AR genes, followed by confirming its biological role in an expressive bacterium such as E. coli. All in all, while metagenomic shotgun sequencing can sidestep cultivability of gut commensals in which anaerobic bacteria predominate, its small size of contig assembly renders the characterization of antibiotic-resistant gene harboring transposons or other mobile elements difficult.²¹ This can be overcome by constructing and screening of fosmid libraries using longer DNA fragments isolated from pulsed-field gel electrophoresis assuming that the gene encodes the same phenotype in both the heterologous host and the native bacterium.⁸ The bioinformatic pipelines and reference databases for gut resistomes are summarized in Tables 2 and 3, respectively.

Common antibiotic resistance determinants found in healthy human gut

Of 180 fecal metagenome datasets tested, the AR genes *aadE*, *bacA*, *acrB*, *tetM*, *tetW*, *vanR*, and *vanS* were present in all people.²⁵ The healthy human gut is expected to carry AR genes which are associated with intestinal anaerobic commensals such as *catA* and *tetW* encoding chloramphenicol

and tetracycline resistances, respectively.¹⁶ Interestingly, functional metagenomic studies have recovered clones with AR genes encoding resistance to β -lactamases (*bla_{TEM}*), macrolides (*ermB*, *ereA*), aminoglycosides (*strA*, *strB*), trimethoprim (*dfrA14*) and sulfonamides (*sul2*) other than bacitracin and tetracyclines.²⁰ To date, genes conferring resistance to isoniazid, rifampicin, oxazolidinones have not yet been reported in the healthy human gut microbiome.

Quality assessment

Of 22 included studies, seven were re-analysis of publicly available metagenomic datasets whereas 15 were primary studies of which only one was population based.¹⁸ The remaining used either convenient samples or less than 10 samples with limited representation of the target population. Only six studies reported prior exposure to antibiotics amongst the participants.^{5,16,20,22,29,30} The scores for each study as evaluated by Newcastle–Ottawa scale for non-randomized cohort studies are summarized in Table S4.

Discussion

The abundance and diversity of human gut microbiota and its association with physiological functions are well recognized. Nonetheless, their roles as potential carriers of AR genes have received unequal attention. With the increasing prevalence of multiple drug-resistant bacteria worldwide, the potential reservoir of AR genes in the gut

Table 2. Common bioinformatic pipelines for gut resistome analysis.

		Latest		
Pipeline	Comments	Update	Website	Ref.
fARGene	ARGs can be identified and reconstructed directly from metagenomic data, suitable for unknown resistomes.	May 2019	https://github.com/fannyhb/ fargene	36
ResFinder	Allows the identification of acquired AMRs and chromosomal mutations.	Apr 2019	https://cge.cbs.dtu.dk/services/ ResFinder	37
ARG-OAP	Online pipeline for fast annotation of ARGs.	June 2018	http://smile.hku.hk/SARGs	38
GROOT	Combination of a variation graph representation of genes with a Forest indexing scheme.	April 2019	https://github.com/will-rowe /groot	39
ARIBA	Efficient for identifying acquired AMR genes and variants.	June 2019	https://github.com/sanger- pathogens/ariba	40
DeepARG	This is a deep machine learning algorithm for ARGs.	June 2017	https://bench.cs.vt.edu/deeparg	41
ShortBRED	Profiling protein families of ARGs from shotgun metagenomics data.	Unclear	http://huttenhower.sph.garvard. edu/shortbred	42

ARDB, Antibiotic Resistance Gene Database; CARD, The Comprehensive Antibiotic Resistance Database; MERGEM, Mobile Elements and Resistance Genes database Enhanced for Metagenomics; RED-DB, Resistance Determinants Database; SARG, Structured Antibiotic Resistance Genes.

microbiota and the possibility of their transfer to human pathogens are of concern. In this article, we systematically reviewed studies on healthy human digestive tract resistome and highlight the methodological heterogeneity across studies which may potentially generate biased results.

The environmental resistome has been extensively studied. It was speculated that AR genes have been selected because of human use of antibiotics. Isolation of AR genes in ancient permafrost and in communities without anthropogenic activity disproved this hypothesis, suggesting that antibiotic resistance genes may pre-date the widespread use of naturally occurring antibiotics.¹² The rapid emergence of resistance to sulfonamides in the 1930s demonstrates that a non-natural chemical entity can readily select for genes encoding enzymes with preexisting functionality in bacteria to confer resistance to these synthetic agents. The presence of relevant antibiotic resistance

mechanisms in organism such as *Streptomyces* from which the antibiotics was isolated for clinical use was assumed to be a part of self-protection mechanism for survival.¹² This finding led to numerous studies on disparate microbial communities, predominantly in the soil resistome which is reviewed elsewhere.⁹

Defining an antibiotic resistance gene could be difficult, particularly in cases where sequencebased metagenomic studies have identified a gene with considerable similarity to a known AR gene but without knowledge of its phenotype.³¹ This is further complicated by using *de novo* assembly which identifies thousands of potential AR genes with variable degree of similarity to the existing known AR genes.²¹ The number of candidate AR genes thus identified may actually reflect the cutoff of similarity to define an AR gene *per se*, without reference to the functional significance of the difference in DNA sequence, such as that based on

Table 3. Refe	erence database	s for a	antibiotic	resistance	genes.
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		Latest			
Database	Comments	Update	Information contained	Website	Ref.
ARDB	No longer being maintained. All information has moved to CARD.	July 2009	13,293 genes, 377 types, 257 antibiotics, 632 genomes, 933 species and 124 genera	https://ardb.cbcb.umd. edu/	43
ARGO	Data was restricted to β -lactamase, tetracycline, and vancomycin genes.	Unclear	555 β-lactamases and 115 vancomycin resistance genes reported before 2004.	http://www.argodb.org/	44
ARG-ANNOT	Based on a local BLAST program in Bio-Edit software, allowing offline analysis.	Unclear	1,689 antibiotic resistance genes	No longer available	45
CARD	This is one of the most extensive AMR sequence databases, covering intrinsic, mutation-driven, and acquired resistance, including mobile genetic elements.	April 2019	2570 reference sequences, 1233 SNPs, 7305 plasmids, 5524 chromosomes	https://card.mcmaster. ca/	46
Lahey Clinic database	This database provides comprehensive information on β -lactamase genes.	May 2019	β -lactamase classification and amino acid sequences for TEM, SHV, and OXA genes.	https://www.lahey.org/	47
SARG	The redundant sequences were removed and query sequence detection optimized.	June 2018	9080 sequence belonged to 189 ARGs subtypes.	http://smile.hku.hk/ SARGs	38
MERGEM	This database contains resistance genes suitable for k-mer filtering.	Unclear	The website is no longer maintained	http://www.mergem.gen ome.ulaval.ca/	48
RED-DB	Both partial and complete coding sequences are available.	May 2019	10,742 non-redundant genes covering ten antibiotic classes.	http://www.fibim.unisi. it/REDDB	-
ResFams	Database of proteins encoding drug resistance, organized by ontology.	Feb 2018	2,454 unique sequences, presenting 54 Resfams protein families.	http://www.dantaslab. org/resfams	49
AMRFinder	Database of AMR protein and gene nomenclature with hidden Markov models	June 2019	4,579 AMR protein sequences and more than 560 hidden Markov models.	https:/www.ncbi.nlm. nih.gov/pathogens/anti microbial-resistance /AMRFinder/	50

ARDB, Antibiotic Resistance Gene Database; CARD, The Comprehensive Antibiotic Resistance Database; MERGEM, Mobile Elements and Resistance Genes database Enhanced for Metagenomics; RED-DB, Resistance Determinants Database; SARG, Structured Antibiotic Resistance Genes.

the active sites that determine resistance. In addition, sequence-based metagenomic approach is limited by the variability of expression of the AR genes in heterologous hosts. Some of the AR genes identified by sequence similarity may be silent or nonfunctional in the native hosts.

In multiple studies, AR genes appear to be exclusively present in gut commensals.^{5,16,20} Longitudinal analysis of fecal metagenomic assembly from critically ill subjects surprisingly identified that all antibiotic-resistant genes including $aph(2^{"})$ -Ib and aadE-like gene were carried by mobile genetic elements such as plasmids and insertion sequences in anaerobic gut commensals among Firmicutes, Bacteroidetes, and Actinobacteria.¹⁶ The aminoglycoside resistance genes were harbored by butyrate-producing Clostridium cluster IV commensal which are present in healthy humans.¹⁶ These genes encoding clinically relevant drug resistance, if transferred to pathogens, will pose a threat to the patients.

In recent decades, the increasing use of disinfectants in the food industry and clinical settings led to research on reduced susceptibility to disinfectants. Surprisingly, studies on gut resistome did not consider the presence of genes encoding reduced susceptibility to biocides such as quaternary ammonium compounds and triclosan. It has been shown that environments with high concentrations of quaternary ammonium biocides select for high carriage rates in bacteria of Class 1 integrons and therefore might be expected to co-select for AR genes.⁵¹ Numerous studies employed quantitative polymerase chain reaction and commercially available microarray assays to study gut resistome. These findings would therefore be limited to known resistance genes. Coupled with functional screen of large DNA inserts in appropriate vectors and expression hosts, functional metagenomics appears to be a promising robust molecular tool to identify novel antimicrobial resistance genes and their origins. For instance, functional screens of specimens with known plasmid-borne resistance genes detected in microarray-hybridization did not recover any transformed clones with plasmid-borne resistant determinants but instead detected a clone with resistant gene carried in chromosome.²⁰ This discrepancy may indicate that the genes detected by microarray may be silent and that both determinants may be present in the microbial community.

The use of metagenomic shotgun sequencing circumvents the problem of non-culturable gut commensals.⁵²⁻⁵⁴ The results are often limited by the bioinformatics pipelines and ARG database(s) selected for analyses. Recently, the National Center for Biotechnology Information (NCBI) produced a high-quality, curated, AMR gene reference database consisting of up-to-date protein and gene nomenclature. A comparison of the susceptibilities of three common Gram-negative foodborne pathogens against the database gave high consistent predictions of 98.4%. This database is designated as AMRFinder with more than 390,000 entries, which is one of the most comprehensive databases.⁵⁰ However, the small size of the contigs which are assembled renders the characterization of the resistance-gene harboring transposons or other mobile elements difficult.²¹ This can be overcome by constructing and screening of fosmid libraries using longer DNA fragments isolated from pulsed-field gel electrophoresis assuming that the gene encodes the same phenotype in both the heterologous host and the native bacterium.8

Strengths and limitations of this review

To our knowledge, this is the first systematic review on studies investigating healthy human gut resistome. Using pre-defined inclusion and exclusion criteria, we comprehensively searched the available databases for relevant studies. Our review highlights the methodological heterogeneity across studies that demands harmonization. However, our review is limited by the small sample size in most of the included studies and that most of them were based on convenient samples or republicly analysis of available datasets, a shortcoming which will be addressed as newer technologies are developed and deployed.

Conclusions

In conclusion, there are considerable methodological heterogeneities across studies. The choice of marker genes to represent resistance phenotype to a given antibiotic class varied. Uniformity in phenotypic validation and bioinformatics pipelines will need to be addressed to facilitate inter-study comparisons.

Disclosure of potential conflicts of interest

None to declare.

Funding

We thank the support of funding from Health and Medical Research Fund (Project number 18170082, PI: MI), Food and Health Bureau of the Hong Kong Special Administrative Region, People's Republic of China. The study was partially supported by a seed fund for gut microbiota research provided by the Faculty of Medicine, The Chinese University of Hong Kong. The funding bodies did not involve in the design of the study and collection, analysis, and interpretation of data and in writing the manuscript.

ORCID

Yun Kit Yeoh D http://orcid.org/0000-0003-0241-6117 Nilakshi Barua D http://orcid.org/0000-0002-7007-5433 Zigui Chen D http://orcid.org/0000-0002-8577-1298 Xiao Yang D http://orcid.org/0000-0002-3071-6531 Martin CW Chan D http://orcid.org/0000-0002-1568-1596 Paul KS Chan D http://orcid.org/0000-0002-6360-4608 Margaret Ip D http://orcid.org/0000-0003-1291-6537

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