In recent decades, the emergence and spread of antibiotic resistance among bacterial pathogens has become a major threat to public health. Bacteria can acquire antibiotic resistance genes by the mobilization and transfer of resistance genes from a donor strain. The human gut contains a densely populated microbial ecosystem, termed the gut microbiota, which offers ample opportunities for the horizontal transfer of genetic material, including antibiotic resistance genes. Recent technological advances allow microbiota-wide studies into the diversity and dynamics of the antibiotic resistance genes that are harbouried by the gut microbiota ('the gut resistome'). Genes conferring resistance to antibiotics are ubiquitously present among the gut microbiota of humans and most resistance genes are harboured by strictly anaerobic gut commensals. The horizontal transfer of genetic material, including antibiotic resistance genes, through conjugation and transduction is a frequent event in the gut microbiota, but mostly involves non-pathogenic gut commensals as these dominate the microbiota of healthy individuals. Resistance gene transfer from commensals to gut-dwelling opportunistic pathogens appears to be a relatively rare event but may contribute to the emergence of multi-drug resistant strains, as is illustrated by the vancomycin resistance determinants that are shared by anaerobic gut commensals and the nosocomial pathogen Enterococcus faecium.

1. The gut microbiota: a complex ecosystem and a reservoir for antibiotic resistance genes

Antibiotics have become a cornerstone of medicine in the decades since the Second World War, but even before humans initiated the industrial production and widespread use of antibiotics, they have existed in nature for hundreds of millions of years [1]. In natural environments, antibiotics may incapacitate bacteria that compete for scarce resources and thus provide a selective benefit for the producing strain. In addition, antibiotics could function as signalling molecules which may trigger bacterial developmental processes, such as biofilm formation, that contribute to survival [2,3]. Resistance to antibiotics may arise in a population of susceptible bacteria by the accumulation of mutations (e.g. point mutations in DNA gyrase conferring resistance to quinolones) or by the acquisition of resistance genes that protect the cell against antibiotics. Antibiotic resistance genes can cause phenotypic resistance through a variety of mechanisms, including the enzymatic inactivation of the antibiotic, the modification of the antibiotic target and the prevention of the accumulation of lethal intracellular concentrations of the antibiotic through efflux pumps [4,5]. Just as antibiotics have been present in the environment for aeons, antibiotic resistance genes are ancient too, as illustrated by the estimated emergence of the serine \(\beta\)-lactamases over 2 billion years ago [6]. Horizontal transfer of antibiotic resistance genes also pre-dates the human use of antibiotics, as OXA-type \(\beta\)-lactamases have already been carried on plasmids and have moved between bacterial phyla for millions of years [7]. It should be noted, however, that a gene that confers antibiotic resistance may have an entirely different function in its original bacterial host, as is illustrated by the 2'-N-acetyltransferase encoding gene in the Gammaproteobacterium Providencia stuartii. This enzyme is involved in the modification of peptidoglycan, but because aminoglycoside antibiotics are structurally similar to
Figure 1. Methods for the analysis of the human gut resistome. Starting from a faecal sample, the resistome can be sampled by culture, through targeted detection of resistance genes (by polymerase chain reaction (PCR) or microarray hybridization), by metagenomic sequencing and by functional metagenomics. Further details of these methods are discussed in the text.

The natural substrate of 2'N-acetyltransferase, the enzyme can also inactivate aminoglycosides, providing intrinsic resistance to this class of antibiotics to P. stuartii [8]. Only when such ‘accidental resistance genes’ are mobilized and transferred to other bacterial hosts, can they contribute to the proliferation of antibiotic resistant human pathogens [3]. Currently, antibiotic resistance among human pathogens has become a major threat to modern medicine and there is considerable interest to identify the niches in which bacteria can gain antibiotic resistance genes and the mechanisms by which horizontal transfer of resistance genes occur.

The human body is populated by an estimated 10^{14} bacteria, including harmless symbionts, commensals and opportunistic pathogens [9]. Different body habitats exhibit differences in bacterial composition, presumably reflecting the different micro-environments of the human body [10]. The human gastrointestinal tract harbours a large and diverse bacteria population, which has an important role in human health and disease. The number of bacteria varies along the length of the gastrointestinal tract, ranging from less than 10^3 bacteria ml^{-1} in the stomach and duodenum, increasing to 10^4–10^7 bacteria ml^{-1} in the colon where 10^{11}–10^{12} bacteria ml^{-1} are present [9]. The human gut thus harbours a complex microbial ecosystem, which consists of hundreds of species, collectively termed the gut microbiota. The gut microbiota is relatively stable in healthy adults but the composition of the gut microbiota can change rapidly owing to dietary changes, illness and the use of antibiotics [11,12].

The large majority of the bacteria that populate the human gut are strictly anaerobic. Two phyla, the Bacteroidetes and Firmicutes, commonly dominate the gut microbiota of healthy adults. Bacteria from these phyla perform functions that are important for the host, such as the production of vitamins and the degradation of complex carbohydrates from the diet. In addition, several facultatively anaerobic bacteria, like those from the families Enterobacteriaceae and Enterococcaceae are ubiquitous members of the human gut microbiota, but generally at levels that are considerably (at least 100-fold) lower than those of the strictly anaerobic gut commensals [13,14]. The Enterobacteriaceae and Enterococcaceae are of particular interest because organisms from these groups, such as Escherichia coli, Klebsiella pneumoniae, Enterococcus faecalis and Enterococcus faecium have emerged as multi-drug resistant nosocomial pathogens of major importance in the last few decades [15,16]. Evidently, the gut can serve as a reservoir for opportunistic pathogens, which may cause infections in immunocompromised individuals. The gut’s role as a source of opportunistic pathogens is particularly relevant for hospitalised patients, which are at high risk of developing infections. The large quantities of antibiotics that are used in the treatment of this patient group may select for multi-drug resistant opportunistic pathogens in the gut microbiota. Opportunistic pathogens from the gut may cause infections by translocating across the intestinal barrier or, after faecal contamination of skin and other body sites, may cause infections upon placement of a catheter or an intravenous line [15,17].

Because bacterial populations in the human gut are large and share a similar ecology, there is ample opportunity for the transfer of genetic material [18]. Consequently, there is considerable interest to characterize the antibiotic resistance gene reservoir (‘the resistome’) of the human gut microbiota and to understand to what extent the antibiotic resistance genes can spread between different members of the gut microbiota, particularly between commensals and opportunistic pathogens [19]. In this review, I will highlight recent studies that have used metagenomic approaches to identify and quantify antibiotic resistance genes in the bacteria that populate the human gut. The mechanisms and extent by which antibiotic resistance genes that are harboured by anaerobic commensals can transfer to opportunistic pathogens are also discussed.

2. Methods to study the human gut resistome

Methodological approaches for the study of the resistome are outlined in figure 1 and are further discussed below. Notably, all current methods to describe the human gut resistome in terms of type and abundance of antibiotic resistance genes are by themselves inadequate to fully characterize the reservoir of resistance genes and the genetic determinants that are associated with the resistance genes. Therefore, a combination of different methodologies should be used if one wants to fully characterize the human gut resistome.
Bacteria from the human gut can be analysed through laboratory culture and subsequent characterization of the isolated strains. Culture-based studies from the 1970s and 1980s have shown that anaerobic gut commensals from the orders Bacteroidales and Clostridiales can carry antibiotic resistance determinants. These bacteria can not only transfer antibiotic resistance genes in vitro to closely related bacteria, but also to opportunistic pathogens [20–23]. Nevertheless, the isolation and laboratory culture of gut bacteria has long been believed to be practically impossible [24] and consequently there is a lack of information on the antibiotic resistance genes in gut commensals and whether these genes are linked to mobile genetic elements. Recently, important advances have been reported in the laboratory culture of a wide variety of gut commensals [25–27], which has opened up the possibility of performing culture-based analyses of antibiotic resistant bacteria from the gut microbiota. Comparative genomic studies of intestinal bacteria that serve as reservoirs of antibiotic resistance genes have been performed for the Bacteroidales [28] and Enterococcaceae [29,30]. However, culture-based studies to comprehensively map and characterize the bacteria that carry antibiotic resistance genes in the human gut still remain to be performed and it is as yet unclear whether culture-based methodologies can truly capture the entire complement of antibiotic-resistant bacteria in the human gut.

In addition to laboratory culture, several culture-independent approaches exist that can be used to probe the antibiotic resistance gene reservoir of the gut microbiota by using DNA that is isolated from faecal samples (figure 1). Resistance genes from the gut microbiota can be detected and quantified by quantitative PCR [31,32] or by microarray hybridisation [33,34]. These methods provide a relatively fast overview of the presence and abundance of the targeted antibiotic resistance genes, but they are limited in their ability to detect genes of which the sequence is not fully complementary to the primers and probes and they will, by definition, not provide information on the presence of antibiotic resistance genes that are not targeted by the primer pairs or probes. In addition, PCR and microarray methodologies do not provide information on the genetic context of the resistance genes or the bacterial hosts of the resistance genes.

In an approach termed metagenomic sequencing, DNA that is purified from faeces is sequenced using modern sequencing technologies. The resulting sequencing datasets can be analysed by assembly of the short reads into larger contiguous DNA fragments or by mapping the sequence reads to reference sequences. This method allows the determination of the phylogenetic composition of the microbiota and can be used to simultaneously detect and quantify antibiotic resistance genes in the microbiota [35].

In an alternative approach, termed functional metagenomics, faecal DNA is randomly cloned into an E. coli vector, either as less than 5 kbp fragments in a routine cloning vector or as fragments of approximately 40 kbp in a fosmid vector. The libraries of randomly cloned DNA are then plated on media that contain antibiotics, resulting in the isolation of antibiotic-resistant clones from the libraries. The vector inserts of resistant clones are then sequenced to identify the genes that confer resistance to the antibiotic of interest [4,36]. In the case of fosmid libraries, the antibiotic resistance gene that is harboured by a particular fosmid clone can be identified by in vitro transposon mutagenesis of the fosmid. Subsequent screening for transposon mutants that have lost the antibiotic-resistant phenotype and sequencing of the transposon insertion sites results in the experimental validation of a gene’s function in antibiotic resistance [32,37]. Functional metagenomic analyses of the gut resistome are considerably more labour-intensive than the previously discussed methods but have the advantage that novel resistance genes (i.e. those that are currently not included in antibiotic resistance gene databases) are identified and that, in the case of large-insert fosmid libraries, information can be obtained about the genetic context of antibiotic resistance genes.

3. Metagenomics of the human gut resistome

Several studies have recently applied metagenomic sequencing and functional metagenomics to probe the resistome in the gut of healthy [38–41] and hospitalized individuals [32,42].

These studies have revealed that the human gut microbiota forms a large reservoir of antibiotic resistance genes, with Forslund et al. [38] finding resistance genes for 50 of 68 classes of antibiotics in 252 faecal metagenomes, at an average of 21 antibiotic resistance genes per sample. These samples were collected from individuals in Spain, Italy, Denmark, France, Malawi, the USA and Japan. Hu et al. [39] identified a total of 1093 antibiotic resistance genes in 162 individuals from China, Denmark and Spain, in a dataset that partially overlapped with that analysed by Forslund et al. [38]. Genes providing resistance to the antibiotic tetracycline (tet32, tet40, tetO, tetQ and tetW) are present in the microbiota of all individuals and are also the most abundant family of resistance genes [38,39]. The tetO gene is the most abundant resistance gene in Chinese, Danish and Spanish individuals, which may be explained by the high prevalence of this resistance gene in Bacteroides isolates, which has increased from 30% in the early 1970s to more than 80% at the start of the twenty-first century [43]. Other resistance genes that were ubiquitously present putatively confer resistance to aminoglycosides (ant(6’)-Ia), bacitracin (bacA) and the glycopeptid vancomycin (vanRA and vanRG) [39]. However, it should be noted that the Antibiotic Resistance Genes Database (ARDB) [44], which was employed by the authors of both studies, contains many genes that have an unclear role in antibiotic resistance. Specifically, the bacA gene appears to be present in most bacterial genomes, where it has a role in peptidoglycan synthesis [45]. In addition, the vanRA and vanRG genes are involved in the transcriptional regulation of vancomycin resistance genes in enterococci [46]. These regulatory genes are therefore not directly involved in the remodelling of peptidoglycan cross-links, leading to vancomycin resistance, but may be broadly present, regulating genes that do not have a role in vancomycin resistance [47]. The large number of genes with housekeeping or regulatory functions in ARDB should lead researchers to use more recently developed tools and resistance gene databases to characterise the resistome in natural environments, such as CARD [48], RED-DB (http://www.fibim.unisi.it/REDDB), ResFinder [49], ARG-ANNOT [50] or Resfams [51].

Both previously discussed large-scale metagenomic sequencing studies of the gut resistome [38,39] analysed datasets of healthy individuals from different countries, which had very different practices in terms of antibiotic use in human and veterinary medicine. Interestingly, individuals from countries with relatively reticent policies of antibiotic use
use in humans and animals (specifically Denmark in these studies) have lower levels of antibiotic resistance genes in their gut microbiota than people from countries where antibiotic use is considerably higher, like Spain and China [38]. This finding suggests that policies concerning antibiotic use in human and veterinary medicine can have a major effect on the relative abundance of antibiotic resistance genes in the gut microbiota of the inhabitants of the different countries [38,39]. However, it should be noted that the difference in antibiotic resistance gene abundance in the microbiota of individuals from these countries is relatively small (approx. 1.5 to twofold) and it remains to be determined whether this difference in abundance contributes to the burden of antibiotic resistant infections in these countries. Interestingly, analysis of single nucleotide polymorphisms (SNPs) in the antibiotic resistance genes indicated that the resistance genes have a specific geographical signature, with the sequences of resistance genes from Chinese individuals forming a cluster that is distinct from resistance genes of Danish and Spanish individuals, which are intermingled in a phylogenetic analysis [39]. A similar specific clustering of Chinese individuals was observed when the abundance of resistance genes in different populations was analysed [52]. The regional differentiation in the sequence and abundance of resistance genes again may be linked by the differences in antibiotic use and exposure of the microbiota to antibiotics in China and European countries, but further analyses with additional data from individuals in other, particularly non-European, countries, will need to be performed to confirm these observations.

Recently, metagenomic sequencing has been used to study the gut resistome of patients throughout hospitalization [32,42]. As in the studies discussed above, many antibiotic resistance genes were found to be present in the microbiota of the studied patients. The relative abundance of antibiotic resistance genes generally appeared to increase in response to antibiotic therapy. For example, Buelow et al. [32] found that the relative abundance of genes that confer resistance to aminoglycosides expanded during hospital stay, particularly during hospitalization in an intensive care unit (ICU) (figure 2). The expansion of the resistome may have been linked to the use of tobramycin (an aminoglycoside antibiotic) that is used as part of a prophylactic antibiotic therapy, termed selective decontamination of the digestive tract, which is used in some European countries to lower the risk of infections with opportunistic pathogens during ICU hospitalization [53]. Similar effects on the resistome were observed in a metagenomic study in which four patients received different courses of antibiotics [42]. However, the use of antibiotics does not always lead to an expansion of the gut resistome in patients and resistance genes from the human microbiota may even be eradicated during antibiotic therapy [32,42]. This can occur during combination therapy, when two (or more) antibiotics are used simultaneously: when a bacterium carries a resistance gene for one of the antibiotics, but is still susceptible to the other antibiotic that is used during therapy, the resistance gene will be lost from the microbial population.

Even though some information on the genetic context of resistance genes can be gleaned by assembling metagenomic sequencing data [52], the bacterial host of antibiotic resistance genes identified by metagenomic sequencing can frequently not be determined with certainty as resistance genes are often associated with promiscuous genetic elements that can be located in a wide variety of organisms. The identification and quantification of antibiotic resistance genes by metagenomic sequencing will also fail to identify genes that have not previously been linked to antibiotic resistance and are therefore not present in antibiotic resistance gene databases that are used to analyse metagenomic sequencing data. Several functional metagenomic studies have been performed to study the human gut resistome and these have provided...
an important complement to metagenomic sequencing studies. Sommer et al. [41] describe a functional metagenomic study of the resistome in faeces and saliva of two healthy individuals. DNA fragments (1–3 kb in size) were randomly cloned into an E. coli cloning vector after which the resulting library was screened for phenotypic resistance to thirteen antibiotics, leading to the isolation of 290 antibiotic resistant clones. Subsequently, the plasmid inserts of resistant clones were sequenced to identify the genes that conferred resistance. Interestingly, most of the antibiotic resistance genes that were identified in this way had relatively low nucleotide identity (69.5% on average) to genes that had been deposited at GenBank at the time of analysis. Sequencing of the inserts that originated from faecal material indicated that most of the genes originated from strictly anaerobic gut commensals, such as bacteria from the genera Bifidobacterium and Bacteroides. Notably, these resistance genes were distinct (on average 60.7% nucleotide identity) from the resistance genes that were found in pathogenic bacteria that can grow aerobically, suggesting that resistance genes from human gut commensals are unlikely to transfer frequently to pathogens. A recent, methodologically similar study found that the resistome of 22 healthy infants and children is already highly diverse and contains genes conferring resistance to 14 of the 18 tested classes of antibiotics [54]. Both novel and previously described resistance genes were identified in the infant gut microbiota and clinically relevant antibiotic resistance genes could be identified, even in the absence of selective pressure in the form of antibiotic therapy.

Because libraries with relatively small inserts were generated in the studies discussed above, only limited information can be obtained on the genetic context of the resistance gene. This limitation can be circumvented by the generation of long-insert (fosmid) libraries of which the inserts can be sequenced in their entirety to allow reliable determination of the origin of the cloned resistance genes and to assess its potential association with mobile genetic elements. Compared to small-insert libraries, fosmid libraries may result in lower numbers of antibiotic resistant clones, because the antibiotic resistance genes have to be expressed from their native promoters rather than a strong, E. coli-optimized, promoter in small-insert libraries [55]. Reflecting the high abundance of tetracycline resistance genes in the gut resistome, fosmid clones that are resistant to this antibiotic can be readily isolated from fosmid libraries constructed with faecal DNA. In a study that used fosmid libraries to determine the diversity of tetracycline resistance genes and the bacteria that carry these genes in a mother and her infant child, tetM and tetL genes were found to be carried by streptococci in the infant metagenomic library [40]. The tetracycline resistance genes tetO and tetW could only be detected by PCR in the uncloned DNA that was purified from the infant faecal sample. By contrast, tetM and tetL were not found in the metagenomic library of the mother. Instead, this library contained the tetracycline resistance genes tetO, tetW and tetX, which were carried by strictly anaerobic Bacteroidetes and Firmicutes [40]. This observation suggests that resistance gene transfer can occur from the maternal microbiota to the child. However, skin bacteria (including streptococci and staphylococci) which colonize the infant intestinal tract in the first days and weeks of life are likely to be more important sources of antibiotic resistance genes [56].

Buelow et al. [32] have applied fosmid-based functional metagenomics, in addition to metagenomic sequencing, to map the gut resistome during and after patient hospitalization. The authors were able to isolate five clones that acquired resistance to the β-lactam ampicillin, the macrolide erythromycin and tetracycline. Antibiotic resistance genes in the fosmid were identified by in vitro transposon mutagenesis and all fosmid inserts were sequenced to completion, allowing the analysis of the genetic context of the resistance gene. All resistance genes were predicted to originate from anaerobic gut commensals (Bacteroidetes, Firmicutes and Actinobacteria). Notably, the aminoglycoside resistance aph(2")-Ib, which was predicted to originate from the Firmicute Subdoligranulum, was genetically linked to genes with putative roles in plasmid conjugation and replication. This gene has previously been described in the opportunistic pathogen E. faecium and may contribute to high-level aminoglycoside resistance in this organism [57].

Metagenomic studies have mapped the human gut resistome in great detail and have shown that anaerobic gut commensals form the main reservoir of antibiotic resistance genes. However, many questions remain on the potential mechanisms by which resistance genes can circulate among the bacteria in the gut microbiota and the extent by which these can transfer from mostly harmless gut commensals to opportunistic pathogens.

4. Gut commensals as hubs for resistance gene transfer in the gut

Horizontal transfer of antibiotic resistance genes can occur through several different mechanisms (figure 3). During transformation, naked DNA, which may contain antibiotic resistance genes, is taken up by a bacterium. During conjugative transfer, resistance genes can spread after the formation of a mating bridge between two cells, via which plasmids or conjugative transposons can move from a donor cell to a recipient. In transduction, resistance genes that are encoded by bacteriophages can transfer from one cell to another and be integrated into the chromosome of the recipient cell, in a process called lysogeny [58]. Of these three processes, transformation does not appear to contribute appreciably to horizontal gene transfer in the mammalian intestinal tract [59], but both conjugation and transduction importantly contribute to the spread of antibiotic resistance genes in the gut microbiota.

Conjugation of plasmids or conjugative transposon between related organisms appears to be a relatively common event in the gut [28,30,60] and conjugation may therefore contribute importantly to the spread of resistance genes. Analyses of genome and plasmid sequences have indicated that conjugation between remotely related bacteria, while rarer than between closely related bacteria, contributes to the dissemination of antibiotic resistance genes [61,62]. Further evidence for in vivo conjugative spread of resistance genes from anaerobic gut commensals to opportunistic pathogens is illustrated by the observation that the vanB-type vancomycin resistance transposon is commonly carried by anaerobic gut commensals of the phylum Firmicutes [63,64], which can serve as a continuous source for vancomycin resistance genes for the nosocomial pathogen E. faecium [65]. Among Gram-negative bacteria, the Bacteroidetes appear to play an important role in the in vivo transfer of resistance genes as they are able to successfully acquire DNA from a wide variety of bacteria, including the Gram-positives E. faecalis.
and *Clostridium perfringens*, which can then be transferred to other gut dwelling bacteria [43]. A specific bacteroidal conjugative transposon, termed CTnDOT, is particularly important for the ability of this group of gut commensals to spread erythromycin and tetracycline resistance. Interestingly, transfer of CTnDOT is triggered by exposure to low levels of tetracycline [66], thereby directly linking antibiotic use and the spread of antibiotic resistance determinants. In addition to the strictly anaerobic gut commensals discussed above, facultatively anaerobic bacteria, in particular the lactic acid bacteria (enterococci, streptococci and lactobacilli), appear to be important conduits of horizontal gene transfer in the intestinal tract [67]. Enterococci appear to be exceptionally well suited to function as ‘drug resistance gene traffickers’ in the human gut [68]. In addition, Enterobacteriaceae can readily exchange plasmids encoding antibiotic resistance and virulence genes during colonization of the intestinal tract [69,70].

The interaction between the human host and the microbiota and its effect on conjugative gene transfer in the gut deserves further attention as there appears to be a role for the mammalian host in determining the extent of horizontal gene transfer in the microbiota as human intestinal epithelial cells can produce a protease-encoding compound that can lower conjugational transfer of an antibiotic resistance plasmid between *E. coli* strains. [71]. By contrast, inflammation of the gut boosts conjugative gene transfer between pathogenic and commensal *E. coli* strains [72].

Phages are present in numbers that equal those of bacteria in the intestinal tract [73–75] and are hypothesized to importantly shape the composition of the gut microbiota [76]. Analyses of metagenomic sequencing datasets have further expanded our understanding of the diversity and abundance of bacteriophages in the gut microbiota. Integrated prophages, which can carry antibiotic resistance genes, frequently enter a lytic cycle in the gut and could thereby transfer antibiotic resistance genes between gut bacteria [77]. Supplementing these findings, a recent study using quantitative PCR on phage DNA that was isolated from faeces of 80 healthy humans found that more than 70% of samples were positive for an antibiotic resistance gene, with the β-lactamases *bla*TEM and *bla*CTX-M-1 and the quinolone resistance gene *qnrA* being most abundant [78]. Experimental evidence for the role of bacteriophages in the horizontal transfer of resistance genes has been obtained in a study where mice were treated with the β-lactam antibiotic ampicillin or the fluoroquinolone ciprofloxacin [79]. Both antibiotic treatments led to an increase of antibiotic resistance genes in the phage metagenome. Remarkably, when phages from antibiotic-treated mice were isolated and used to infect an aerobic culture of the microbiota of non-treated mice *ex vivo*, the frequency of resistant isolates was significantly higher compared with the culture that was infected with phages isolated from the microbiota of untreated mice. These data show that phage transduction can contribute importantly to the expansion of the gut resistome during antibiotic therapy [79].

5. Discussion and future perspectives

Antibiotic resistance genes are widely distributed in the environment and the studies that are discussed in this review have provided convincing evidence that the gut microbiota
form a large reservoir for antibiotic resistance genes. Conjuga-
tive transfer of plasmids and transposons, and transduction are
two important mechanisms by which antibiotic resistance
genes can spread through the microbiota. However, there
remain many open questions on the relative contribution of the
resistome in donating antibiotic resistance genes to
opportunistic pathogens.

The macrolide resistance genes *ermB*, *ermF* and *ermG* and
the tetracycline resistance genes *tetM* and *tetQ* can spread
among phylogenetically diverse Gram-negative and Gram-
positive gut bacteria [60], but these may be exceptions as
many of the resistance genes that are detected in gut commen-
sals do not occur in opportunistic pathogens. Indeed, resistance
genes from gut commensals appear to mostly spread to related
bacteria. For example, the β-lactamase *cblA* is frequently pre-
sent in *Bacteroides* and is one of the most abundant resistance
genes in the microbiota of healthy individuals and patients
[38,39]. However, this gene does not appear to transfer to
opportunistic pathogens, like the Enterobacteriaceae, even
though functional metagenomic selections have shown that
this gene confers β-lactam resistance to *E. coli* [32,41]. The
observation that the gut microbiota of two healthy volunteers
contained nine β-lactamase sequence families (eight of which
have highest identity to genes in *Bacteroides*) that had not pre-
viously been shown to occur in opportunistic pathogens [41],
also indicates that major barriers exist for the transfer of resist-
ance genes between anaerobic gut commensals (in particular,
*Bacteroides*) and Gram-negative facultative anaerobic opportunis-
tic pathogens, like the Enterobacteriaceae. Conjugal transfer of
resistance genes from *Bacteroides* to *E. coli* is possible under labora-
tory conditions [20,21], but, apparently, the conditions in the
intestinal tract preclude the efficient mobilization and transfer
of antibiotic resistance genes from Bacteroidetes to Enterobacter-
icaeae. The bacteroidal β-lactamases are therefore unlikely to
contribute importantly to the burden of antibiotic resistance
among Enterobacteriaceae.

It is less clear to what extent gut commensals from the
phylum Firmicutes can transfer their resistance genes to
Gram-positive, gut-dwelling opportunistic pathogens, such as
enterococci and streptococci. A lack of data on the plasmids
and conjugal transposons that are carried by the strictly
anaerobic Gram-positive bacteria in the gut complicates the
assessment of the transferability of these genes. However, con-
jugative transposons of the Tn916/Tn1545 family (carrying
tetracycline, macrolide and aminoglycoside resistance genes)
are widely disseminated in clostridia, enterococci and strepto-
cocci, suggesting that resistance gene transfer occurs between

these groups of bacteria [80]. Similarly, aminoglycoside resist-
ance genes are carried by bacteria from the class Clostridia
[32,81], even though anaerobic gut commensals are thought to
be intrinsically resistant to aminoglycosides [82]. The role
(if any) of these genes in aminoglycoside resistance in Clostri-
dia is therefore unclear. Nevertheless, the aminoglycoside
resistance genes from Clostridia are also found in enterococci
and Enterobacteriaceae [83], again suggesting that Gram-positi-
ve gut commensals serve as hubs for the transfer of resistance
genes. Arguably, the *vanB* transposon is clinically the most rele-
vant antibiotic resistance determinant that can be acquired
from the microbiota. Clinical *E. faecium* strains are able to
acquire the *vanB* transposon, and gain resistance to vancomy-
cin at a minimal fitness cost [84]. The recent, and largely
unexplained, increase of *vanB*-positive *E. faecium* strains in Europe [85,86] may possibly be caused by a further spread of
*vanB*-type transposons which are then repeatedly and indepen-
dently acquired by vancomycin-susceptible *E. faecium* isolates.

Currently, it is already feasible to use metagenomic sequenc-
ing as a diagnostic tool for the detection of pathogens in faecal
samples in outbreak situations [87] and fluctuations of the
resistome have been tracked in individual patients [32,42].

The routine implementation of metagenomic sequencing of
the microbiota in clinical practice is some way off as costs for
sequencing and analysis are still high. Perhaps more impor-
tantly, the diagnostic relevance of resistome profiling by
sequencing is currently limited as it takes at least several days
to perform sequencing and analyses. Future advances in
DNA sequencing technologies are likely to facilitate high-
throughput characterization of the resistome by metagenomic
sequencing in hospitalized patients. The choice of antibiotics
for therapy might then be guided, at least partially, by the com-
position and relative abundance of the antibiotic resistance gene
reservoir in patients. To fully assess the risks that are associated
with the selection for antibiotic resistance genes in gut commen-
sals, further studies are needed to profile the antibiotic
resistance genes from the gut microbiota, particularly with
respect to their potential for horizontal gene transfer.

Acknowledgements. The author thanks Rob Willems for helpful discussions.

Funding statement. W.V.S. is supported by the European Union Seventh
Framework Programme (FP7-HEALTH-2011-single-stage) ‘Evolution
and Transfer of Antibiotic Resistance’ (EvoTAR) under grant agreement
number 282804, The Netherlands Organisation for Scientific Research
(VIDI grant 917.13.357) and The Netherlands Organisation for Health
Research and Development ZonMw (Priority Medicine Antimicrobial
Resistance; grant 205100015).

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