Broad host range plasmids can invade an unexpectedly diverse fraction of a soil bacterial community

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Abstract:

Conjugal plasmids can provide microbes with full complements of new genes and constitute potent vehicles for horizontal gene transfer. Conjugal plasmid transfer is deemed responsible for the rapid spread of antibiotic resistance among microbes. While broad host range plasmids are known to transfer to diverse hosts in pure culture, the extent of their ability to transfer in the complex bacterial communities present in most habitats has not been comprehensively studied. Here, we isolated and characterized transconjugants with a degree of sensitivity not previously realized in order to investigate the transfer range of IncP-type broad host range plasmids from three proteobacterial donors to a soil bacterial community. We identified transfer to many different recipients belonging to 11 different bacterial phyla. The prevalence of transconjugants belonging to diverse Gram-positive Firmicutes and Actinobacteria suggests that inter-Gram plasmid transfer of IncP-1 and IncPromA-type plasmids is a frequent phenomenon. While the plasmid receiving fractions of the community were both plasmid- and donor-dependent, we identified a core super-permissive fraction that could take up different plasmids from diverse donor strains. This fraction, comprising 80% of the identified transconjugants, thus has the potential to dominate IncP- and IncPromA-type plasmid transfer in soil. Our results demonstrate that these broad host range plasmids have a hitherto unrecognized potential to readily transfer to very diverse bacteria and can, therefore, directly connect large proportions of the soil bacterial gene pool. This finding reinforces the evolutionary and medical significances of these plasmids.

Keywords: Broad host range/Conjugation/Gene transfer/Plasmid/Transfer range
Introduction

Conjugal plasmid transfer is a process by which bacteria horizontally transfer complete sets of genes to other, potentially distantly related, organisms. Conjugal plasmids frequently carry accessory genes, often encoding antibiotic or metal resistances, catabolic pathways, or virulence factors. They are often implicated in the evolution of pathogenic bacteria and the rapid spread of antibiotic resistance, likely fostering the rise of multiple-resistant microbes in hospitals (Levy & Marshall, 2004) and animal husbandries (Zhu et al., 2013). While the relevance of plasmid transfer has become very acute in this age of massive antibiotic usage, plasmids have been exchanged for much longer and many prokaryotic genomes present signs of intense past horizontal gene transfer (Ochman et al., 2000).

Plasmids present different abilities to transfer into, and be maintained in, distantly related bacterial hosts and are loosely categorized as having a narrow or broad host range. The transfer of narrow host range plasmids is limited at one of the steps required for successful transfer, such as the formation of mating pairs, the avoidance of the recipient’s restriction system, or the correct expression of its replication and maintenance systems in the recipient (Thomas & Nielsen, 2005). Some broad host range plasmids can transfer across bacterial phyla and even across domains of life (Waters, 2001; Heinemann & Sprague, 1989), and several genetic determinants conferring broad host transfer capability have been identified (Jain & Srivastava, 2013).

The host range is thus a key parameter that controls the ecology and fate of plasmids. The evaluation of host range has traditionally been conducted using few individual pure strains as recipients, a situation that contrasts with the fact that most bacteria - and thus most plasmids - exist within complex communities of hundreds to thousands of species (Hong et al., 2006; Brown Kav et al., 2012). Among these diverse communities, all strains are obviously not equally permissive towards plasmid receipt, even for broad host range plasmids. This notion was supported when studying plasmid transfer to a range of strains isolated
from marine water or wastewater treatment bioreactors (Sørensen, 1993; Inoue et al., 2005). With the use of fluorescent reporter genes to track plasmids, which reduces the need for selection and cultivation steps to identify transconjugants, it has become apparent that, in complex communities, broad host range plasmids can indeed be received by bacteria distantly related to the donor, even in the absence of selective pressure for plasmid carriage (Musovic et al., 2014, 2006; Shintani et al., 2014; De Gelder et al., 2005). However these efforts, limited to inspection of a few hundred transconjugants at best, most likely underestimate the true diversity of transconjugal pools and do not accurately describe how plasmid permissiveness may vary across taxa in complex microbial communities.

Horizontal gene transfer between different species has been recognized as a common and major evolutionary process (Zhaxybayeva & Doolittle, 2011), most acutely demonstrated in the heavy interconnection between the resistome of soil dwelling bacteria and human pathogens (Forsberg et al., 2012). The behavior of this environmental resistome may, thus, govern the spread of antibiotic resistance genes to pathogens (Finley et al., 2013). Plasmids serve as main vessels of gene flow in microbial communities, linking distinct genetic pools (Halary et al., 2010; Norman et al., 2009). The in situ host range of plasmids may, then, well govern the taxonomic breadth across which gene flow occurs.

Here, taking advantage of high throughput cell sorting and next-generation sequencing technologies, we map for the first time the intrinsic diversity of the bacterial recipients of broad host range plasmids in a microbial community extracted from soil, under conditions where cell-to-cell contacts are maximized. We analyzed matings initiated with combinations of three plasmid donors and three plasmids to identify, how permissiveness towards broad host range plasmids is distributed across taxa among the recipient community.
Material & Methods

Donor strain construction

Soil bacterial communities were challenged with various plasmid/donor combinations through solid surface filter matings. The plasmids were marked with a genetic tag encoding conditionally expressible green fluorescent proteins. The used entranceposon (Bahl et al., 2009) carries a lacIq repressible promoter upstream the gfpmut3 gene, encoding for the green fluorescent protein (GFP). Plasmid donor strains were all chromosomally tagged with a gene cassette encoding constitutive red fluorescence and constitutive lacIq production. As a result, there is no gfp expression in the donor strains, but upon plasmid transfer to a soil bacterium, gfp expression is possible, resulting in green fluorescent cells or microcolonies, which can be detected and sorted by fluorescence microscopy or FACS, respectively (Figure 1) (Sørensen et al., 2005).

_Pseudomonas putida_ KT2440, _Escherichia coli_ and _Kluyvera_ sp. served as donor strains, and were each electroporated with the plasmid pGRG36-lacIq-Km-Lpp-mCherry carrying both the transposase genes and the Tn7 lacIq-Lpp-mCherry-KmR region for specific integration of the lacIq-Lpp-mCherry-KmR gene cassette into the chromosomal attTn7 site. Colonies were selected for KmR on LB agar plates at 30 degrees. Colonies were restreaked on selective LB agar plates at 30 °C, incubated in liquid LB overnight culture without antibiotics at 30 °C and finally streaked on LB agar plates without selection at 37 °C for integration of the gene cassette and subsequent loss of the Tn7 helper plasmid. Colonies were tested for successful loss of helper plasmid and chromosomal integration of gene cassette by PCR (McKenzie & Craig, 2006). The same colonies were also phenotypically verified to be bright red fluorescent using stereo microscopy.

Construction of gfpmut3-tagged plasmid pKJK5

Plasmids RP4 and pIPO2tet have been constructed earlier (Musovic et al., 2010, 2014). The 54 kbp IncP-1 plasmid, pKJK5, originally isolated from a soil/manure environment, harbors a tetracycline and a trimethoprim resistance determinant as well as a class 1 integron (Sengeløv et al., 2001).
Entranceposon [Km®\(,\) PA10403-\textit{gfpmut3}], carrying a kanamycin resistance determinant and a LacI\(^q\) repressible promoter upstream the \textit{gfpmut3} gene, encoding the Green Fluorescent Protein (\textit{gfp}), was derived from pEntranceposon [\textit{Km}®] (Finnzymes, F-766) and randomly inserted into the plasmid pKJK5 using the artificial Mu transposon in vitro delivery system as described previously (Bahl et al., 2009). Transformed \textit{E. coli} GeneHogs single colonies were selected for resistance towards trimethoprim and kanamycin and screened for sensitivity towards tetracycline in order to select for plasmid derivatives with an Entranceposon insert location directed to an accessory element (the tetracycline resistance determinant), thereby excluding any potential impacts on conjugation transfer ability. The exact insert location of [\textit{Km}®, PA10403-\textit{gfpmut3}] in the selected pKJK5 derivative of this study was determined by sequencing out from the inserted fragment in one direction using primer Seq_Bw_Ent_gfp: 5’-GCCAGAACCGTTATGATGTCGG-3’. The insertion mapped to position 30.614 bp in the \textit{tetA} gene (30.435-31.634 bp) of plasmid pKJK5 (accession no. AM261282). The selected \textit{gfpmut3}-tagged pKJK5 plasmid was finally introduced into \textit{E. coli} MG1655::\textit{Km}®-Lpp-mCherry, \textit{P. putida} KT2440::\textit{Km}®-Lpp-mCherry and \textit{Kluyvera} spp.::\textit{Km}®-Lpp-mCherry cells by transformation.

**Soil sampling & community extraction**

Soil samples were taken at the annually tilled CRUCIAL (Closing the Rural Urban Nutrient Cycle) agricultural field site (Taastrup, Denmark) from a plot subjected to no further agricultural treatment (Magid et al., 2006). Soil samples were collected in late fall 2012. Samples were taken from three different plots of this treatment. Each plot was sampled for 1 kg of soil at 5 locations. The resulting soil volume was sieved and homogenized to obtain a representative sample. From a total of 30 g of the homogenized chosen soils, indigenous bacterial communities were isolated by Nycodenz®-extraction (Musovic et al., 2010) and used as recipients in the mating assay. Donor strains were grown overnight in LB-medium supplement to the plasmid specific antibiotics (Table 1), harvested by centrifugation.

**Solid surface filter mating assay**
The extracted recipient community was challenged with exogenous plasmids via solid-surface filter matings (Musovic et al., 2010) modified to an initial ratio of donor to recipient bacteria of 1:1 at a density of approximately 30,000 bacteria/mm² on the filter. As a growth medium we used a 10% soil extract medium as described by Musovic et al. (2010) buffered at pH 7.2 with 5mM MOPS and supplemented with 20 µg/mL Nystatin to avoid fungal growth. Unlike in Musovic et al (2010), we did not use additional nutrient additions, but only relied on soil extracted nutrients to support activity during the mating incubations. Successful conjugation was checked after 48 hours by epifluorescence stereo microscopy and confocal laser scanning microscopy (CLSM) (Figure 1) (Musovic et al., 2010).

**Cell collection and triple gated fluorescence activated cell sorting of transconjugants**

Cells from 5 filters per mating combination replicate were harvested in 2mL 0.9% NaCl-solution by vortexing for 3 minutes. Flow cytometric detection of cells was carried out using a FACSARia IIIu (Becton Dickinson Biosciences, San Jose, CA). The following settings and voltages were used during analysis: forward scatter (FSC) = 505 V, side scatter (SSC) = 308 V, and detectors for green (BP filter 530/30 nm) and red fluorescence (BP filter 610/20 nm) were set at 508 V and 500 V, respectively. A 70 µm nozzle was used at a sheath fluid pressure of 70 psi. The BD FACSDiva™ software v6.1.3 was used for both operating and analyzing results. Sorting was performed using a 488 nm (20 mW) laser connected to the green fluorescence detector at 515-545 nm and a 561 nm (50 mW) laser connected to the red fluorescence detector at 600-620 nm. Three gates were defined in bivariate plots to sort for transconjugants. On the SSC-A vs FSC-A plot a gate for only particles of bacterial size was used. On the FITC-A vs SSC-A plot a gate was set that covered all green fluorescent particles, while using an additional non-red gate on the PE-Texas Red-A vs SSC-A plot excluded all small autofluorescent particles from soil or leaking donors (Figure 2) to sort out only transconjugants. All samples were diluted in 0.9% NaCl to approximately 2000 counting events s⁻¹ before fluorescent activated cell sorting to assure for optimal sorting. Transconjugants that originally made up for less than 0.1% of the total community in the filter matings and were enriched to up
to 82% in a first fast sorting step, before isolating over 10,000 transconjugants per sample in a second purification step, leading to 100% purity of green cells as observed by fluorescent counting in the flow cytometer. Plating of more than 200 isolated transconjugants on 10% soil extract medium (Musovic et al., 2010) resulted in detection of green fluorescence in all colonies, additionally verifying purification of gfp-expressing transconjugants. Of the isolated transconjugants, twenty were subject to 16S rRNA gene sequences; the recovery of proteobacterial, sphingobacterial and actinobacterial phylotypes indicated diversity among transconjugants.

**Bacterial cell lysis, amplification and sequencing**

Bacterial transconjugal cells from the second sort, initially collected in 5 mL sterile polystyrene round-bottom Falcon™ tubes (BD Biosciences, San Jose, CA) with 0.5 mL 0.9% NaCl solution, were transferred to 1.5 mL Eppendorf tubes and centrifuged at 10,000 x g for 30 min to collect the cell pellets. The supernatant was carefully removed, the cell pellet suspended in 20 μL of Lyse and Go PCR Reagent (Thermo Scientific, Waltham, MA, USA) and the lysis mixtures transferred to 0.2 mL amplification tubes. Cell lysis was subsequently performed in an Arktik™ Thermal Cycler (Thermo Scientific, Waltham, MA, USA) using the program: one initial cycle at 57 °C for 30 s, a second cycle at 8 °C for 30 s, a third cycle at 65 °C for 90 s, a fourth cycle with heating to 97 °C for 3 min, a fifth cycle with cooling to 8 °C for 60 sec, a sixth cycle with heating to 65 °C for 3 min followed by additional heating to 97 °C for 60 s and cooling to 65 °C for 60 s with a final end-step at 80 °C. DNA-containing cell lysis products were immediately put on ice and used directly for subsequent PCR. Then, 5 μL of the cell lysis product from the previous step was used directly for sequencing library preparation. Tag-encoded 16S rRNA gene pyrosequencing was carried out after amplification of the V3 and V4 region (Primers: 341F: 5´-CCTAYGGGRBGCASCAG-3 and 806R 5´-GGACTACNGGGGTATCTAAT-3) using the PCR procedures and GS FLX Titanium chemistry as described previously (Hansen et al., 2012).

**Sequence analysis and tree construction**
Results and Discussion

High throughput isolation and sequencing of transconjugants

We explored the ability of a bacterial community extracted from soil to engage in horizontal gene transfer and receive one of three gfp-tagged broad host-range plasmids from three different red fluorescent-tagged donor strains in which plasmid-mediated gfp expression is repressed (Table 1). In soil, physical barriers limit contact between freshly introduced plasmid donors and potential recipients (Dechesne et al., 2005); here we maximized cell-to-cell contact in a gene transfer assay (Musovic et al., 2010) to study the intrinsic permissiveness of the recipient community. All three plasmids (RP4, pIPO2tet, and pKJK5) were introduced to the soil community in matings with a Pseudomonas putida donor strain, while plasmid pKJK5 was also introduced via Escherichia coli and Kluyvera spp. donors (Supplementary Table 1). After mating, the gfp-expressing transconjugant cells (Figure 1) were isolated from the mixed community by fluorescent activated cell sorting (FACS). A novel triple gated FACS approach based on size, green fluorescence, and lack of red fluorescence, allowed specific isolation of large numbers of transconjugant cells, in spite of their low relative abundance (less than 0.1%) in the mating mixture (Figure 2). At least 14,000 transconjugant cells were obtained for each mating, corresponding to 28,000 – 116,500 transconjugants per donor/plasmid combination, depending on the number of replicate matings. The eleven pools of sorted transconjugants as well as the total soil recipient community were then subjected to deep amplicon sequencing of 16S rRNA genes, resulting in 29,894 to 50,398 sequences per sample after processing with the mothur pipeline.
(Schloss *et al.*, 2009). This corresponds to more sequences than sorted transconjugants for most samples (Supplementary Table 1), providing an adequate picture of the observed plasmid transfer range.

**Transconjugal pools are plasmid- and donor-specific**

The phylogenetic structure of the transconjugal pools was compared after clustering the partial 16S rRNA gene sequences in OTUs at 97% similarity. The eleven transconjugal pools clustered clearly and significantly apart from the recipient community, as shown by PCoA (Figure 3) and AMOVA (Excoffier *et al.*, 1992) ($p=0.028$). Mating plates contained soil extracts as nutrient sources and growth on filter did not significantly modify the soil community structure ($p=0.797$) based on UNIFRAC comparisons (Lozupone *et al.*, 2011), in spite of a diversity reduction by 72%. The transconjugal pools were clearly distinct from the recipient community, but also differed from each other based on plasmid or donor. Considering different plasmids in an identical donor strain (*P. putida*) and providing the same plasmid (pKJK5) in different donor strains revealed phylogenetically distinct transconjugal pools (AMOVA, $p<0.001$). Hence, plasmid acquisition is not a stochastic process, even for broad host range plasmids. While replicates of the same donor/plasmid combinations differed based on weighted UNIFRAC comparisons ($p<0.05$), the average interreplicate dissimilarity ($W=0.36$) was clearly less than dissimilarity between different plasmid/donor combinations ($W=0.49$) or between transconjugal pools and the soil community ($W=0.60$). Slight differences between the replicates can also be seen in their phylum level distribution of transconjugants (Supplementary Figure 1). This dissimilarity between replicates can most likely be decreased through sorting of higher numbers of transconjugants per replicate, since replicates from the same donor-plasmid combinations grouped significantly together in PCoA ($p<0.01$) (Figure 3). Based on this PCoA grouping and because the numbers of replicates per combination differed (Supplementary Table 1), replicates were pooled for subsequent phylogenetic analysis.

**Transconjugal pools span most of the major bacterial phyla**
More than 300 transconjugant OTUs were detected across all plasmid/donor combinations (Figure 4, Figure 5), a large expansion over the low number of distinct bacterial isolates identified previously from matings in complex environmental communities (De Gelder et al., 2005; Musovic et al., 2014, 2010; Shintani et al., 2014). As expected, Proteobacteria, known to be the main hosts for the studied broad-host-range plasmids (Suzuki et al., 2010), were represented. Unlike in previous studies (Musovic et al., 2010; Shintani et al., 2014), all five classes (α-ε) of Proteobacteria were identified among the transconjugants. More strikingly, the diversity of transconjugants extended much beyond the proteobacterial phylum, and included diverse members of ten additional phyla including Verrucomicrobia, Bacteroidetes and Actinobacteria, some of which are known as poorly cultivable (Joseph et al., 2003). The IncP transfer apparatus is known to build conjugative bridges between a huge variety of organisms (Grahn et al., 2000; Thomas & Nielsen, 2005). Shuttle vectors for gene transfer from Proteobacteria to distantly related recipients such as Cyanobacteria (Wolk et al., 1984) or gram-positive bacteria and yeast (Heinemann & Sprague, 1989; Samuels et al., 2000) have, indeed, been built using the RP4 transfer system, a IncP-1α subgroup plasmid. While the wide transfer potential of the RP4 conjugation system has therefore been known in artificial constructs under laboratory conditions, we are the first to prove that a large proportion of this transfer potential can be realized in nature. A similarly huge transfer potential is demonstrated for pKJK5, a plasmid closely related to RP4 belonging to the IncP-1ε subgroup and for pIPO2tet, a currently unclassified, phylogenetically more distant, cryptic plasmid that merely seems to provide plasmid mobilization capability to its host (Figure 4). We thus show that a variety of broad host range plasmids can effectively be transferred to, and encoded genes can be expressed in large proportions of the bacterial tree of life, much beyond the limited transfer range identified so far (Shintani et al., 2014). The realized transfer range in the soil community under natural conditions might be even higher taking into account that Nycodenz extraction might not be able to recover all bacterial phyla from the soil sample (Holmsgaard et al., 2011). Of the total extractable soil microbial community only the phyla Chloroflexi, Deinococcus-Thermus, Nitrospira and SR1, were not represented in the transconjugal pools in our experiments.
In particular, we identified transfer from the used Gram-negative donor strains to a wide variety of gram positive bacteria (Figure 4, Figure 5). Over 15 OTUs within the Actinobacteria phylum and more than 10 OTUs belonging to 6 different orders of Bacilli and Clostridia within the Firmicutes phylum were identified as transconjugants. Inter Gram conjugal gene transfer has been shown with vectors consisting partly of the broad host range transfer machinery of RP4 recombined with the $sacB$ gene from Gram-positive *Bacillus subtilis* (Schäfer et al., 1994), but has only exceptionally been identified in natural habitats (Musovic et al., 2006). Our observations suggest that it may be a more common process than previously considered.

**Abundance in recipient community and phylogenetic distance to the donor do not explain the composition of transconjugal pools**

In spite of the large diversity within the transconjugal pools, not all OTUs of the recipient community were represented in each pool and the relative abundance of OTUs in transconjugal pools was very heterogeneous. Our method cannot distinguish between original horizontal plasmid transfer events from subsequent vertical plasmid transfer through growth of transconjugants on the mating filter. Therefore, relative abundance in the transconjugal pools can be influenced by the relative growth rate of recipients. However, the fact that OTU abundance in the transconjugal pools is not explained by their abundance in the reference soil recipient community (Figure 4, Figure 5, and Supplementary Table 2) indicates that plasmid transfer occurs preferentially to some recipients and that transconjugal pools are not simply determined by the recipient’s growth ability.

Next, we tested whether phylogenetic distance between donor and recipient, calculated based on the Sogin distance algorithm (Sogin et al., 2006), influenced the abundance of individual OTUs among the transconjugal pools. We found no significant correlation between phylogenetic distance to the donor and recipient frequency in the transconjugal pools ($p=0.09-0.94$) for any of the donor plasmid combinations (Supplementary Figure 2). For example, the most abundant OTUs in soil that do not appear in the transconjugal pools (Supplementary Table 2) are Gammaproteobacteria; they display more than 90% 16S
rRNA gene sequence similarity to the donor strains, while other OTUs with less than 70% sequence
similarity to donor cells, such as several members of the Flavobacterium phylum, did receive at least one of
the plasmids. Transfer of an IncP-1 plasmid from *E. coli* to phylogenetically distant Flavobacteria was
detected in soil microcosms (Pukall *et al.*, 1996), indicating that transfer to distant nodes of the
phylogenetic tree is not only possible, but also realized in undisturbed soil environments. In pure culture,
permisiveness towards broad host range plasmids of isolates that are indistinguishable by 16S rRNA gene
analysis can differ by more than 100-fold (Heuer *et al.*, 2010). Here we confirm that inferring plasmid
uptake and transfer frequency cannot be predicted based on the phylogenetic identity of an OTU.

However, we confirm the role of donors in defining the plasmid transfer host range (De Gelder *et al.*, 2005),
and show that this effect is significant even for two donors belonging to the same family of
Enterobacteriales (*E. coli* & *Kluyvera* sp.) and thus sharing a high genomic similarity. The reasons behind this
are uncertain, but certain strains might have distinct abilities to achieve efficient cell-to-cell contact with a
specific recipient, for example through specific mating mediating pheromones (Hirt, 2002). Earlier studies
have shown that plasmid exchange between two taxonomically different species can exceed intraspecies
transfer frequencies (Bingle *et al.*, 2003), proving that the regulatory interactions of donor, recipient, and
plasmid can influence transfer efficiency.

Similarly, three broad host range plasmids, all carried by the same *P. putida* strain, were transferred to
distinct pools of recipients. Yano *et al.* (2013) hypothesized that, genetic differences appearing among
closely related IncP-1 plasmids through plasmid backbone evolution can result in significant diversities in
host range efficiency without affecting their broad host range nature. Such backbone alterations exist
between the IncP-1α (RP4) and IncP-1ε (pKJK5) core regulatory proteins such as *KorB*, *TrfA*, *TrbA* and *Ssb*
(Bahl *et al.*, 2007). Although these two plasmids are incompatible (both IncP-1), differences in gene
silencing and expression of the different core proteins could explain the different transconjugal patterns.
Since already minor differences in regulation between two IncP-1 plasmids lead to distinct transconjugal
pools, it is coherent that the unrelated transfer machinery of plasmid pIPO2tet caused significantly (p<0.05) dissimilar transconjugal pools when compared to the IncP ones.

A core super-permissive community fraction dominates gene transfer

Out of 281 OTUs identified in the transconjugal pools with the three different broad host range plasmids and *P. putida* as donor, 74 OTUs were common to all three pools (Figure 6A). A similar observation (46 out of 279 OTUs shared) held when comparing the transconjugal pools for plasmid pKJK5 introduced via three different donors (Figure 6B). Therefore, the majority of transconjugant OTUs were only identified in single donor/plasmid combinations. This might result from mating pair combinations that each favor or reduce gene transfer abilities (Bingle *et al.*, 2003; Thomas & Nielsen, 2005; Yano *et al.*, 2013).

While only 74 and 46 OTUs are shared among the compared transconjugal pools, these OTUs represent over 80% of the transconjugal sequences (Figure 6C&D). This core super-permissive community fraction shared by all five transconjugal pools is able to take up diverse broad host range plasmids from diverse donor strains at high frequencies. The presence of this shared core in each analyzed transconjugal pool is the crucial discriminant that groups transconjugal pools apart from the original soil community (Figure 3). The core super-permissive community consists mainly of diverse Proteobacteria like Enterobacteriales (γ), Burkholderiales (β), Pseudomonadales (γ) and Rhizobiales (α) (Figure 4). In addition, within this core super-permissive fraction, several OTUs that are rare in the recipient community (<0.001%) are more than 20-fold overrepresented in transconjugal pools (Figure 4). The participation of these rare community members in gene transfer might play a crucial role in increasing the communal gene pool through rapid recombination with plasmids, since the rare biosphere can harbor a great reservoir of genes (Sogin *et al.*, 2006).

Medical relevance

The large realized transfer potential of newly introduced plasmids in soil may be of medical importance. In recent EAHEC outbreaks in Germany, recombination of a pathogenic with the plasmid of a non-pathogenic *E.coli* strain increased the pathogenic potential to cause a deadly combination (Brzuszkiewicz *et al.*, 2011).
Soil borne antibiotic resistance has been found to be shared with human pathogens (Benveniste & Davies, 1973; Forsberg et al., 2012). Several organisms among the identified transconjugants belong to groups known to contain opportunistic human pathogens, providing a direct link between plasmid encoded soil resistome and opportunistic pathogens. These groups include the proteobacterial Enterobacteria, Pseudomonas or Campylobacter but also groups from other phyla such as Fusobacterium, Streptococcus and Staphylococcus, most of which are treated with antibiotic therapy. Especially the acquisition of new antibiotic resistance genes through plasmid mediated gene transfer may push the pathogenic potential of Staphylococcus, originating from rapid evolution of virulence and drug resistance (Holden et al., 2004), even further.

The observed transfer of broad-host-range IncP-1 type plasmids between Gram-negative and Gram-positive bacteria might lead to a reassessment of the potential of soil bacterial communities to spread antibiotic resistance genes. Indeed, Actinobacteria, the origin of many soil-borne resistance genes (D’Costa et al., 2006) which are sometimes identified in clinical isolates of Gram negative antibiotic-resistant bacteria (Benveniste & Davies, 1973), are frequent among the transconjugants we identified. Broad host range plasmids of the IncP-1 and IncPromA group can thus provide a direct link between diverse bacterial groups. Especially IncP-1ε plasmids such as pKJK5 have been identified as vectors of antibiotic resistance genes transfer among Proteobacteria by additionally hosting Class 1 integron gene cassettes (Heuer et al., 2012).

These Class 1 integrons may not only spread in their originally identified Gram-negative Enterobacteriaceae hosts but can also be found among many Gram-positive bacteria (Nandi et al., 2004). Here we demonstrated a possible direct way of accession of these Class 1 integrons in Gram-positive bacteria through IncP-1ε plasmid transfer from Proteobacteria.

**Ecological & Evolutionary Relevance**

Plasmid host range can be defined in several ways depending on the duration and intimacy of the considered plasmid-host relationship, including the transfer host range, the replication and maintenance
host range, or the evolutionary host range (Suzuki et al., 2010). We show here that the immediate transfer range for IncP plasmids is much wider than previously reported, proving that in absence of physical barriers to cell-to-cell contact, broad host range plasmids have a high likelihood to be, hosted by very diverse bacteria, at least transiently.

However, comparative analysis of plasmid sequences has indicated that the evolutionary host range of IncP plasmids seems to be mostly limited to Proteobacterial classes (Suzuki et al., 2010). This suggests that these plasmids are not maintained long enough outside of this phylum to be significantly affected by non-Proteobacterial genomes. Long-term evolutionary adaptation of the plasmid backbone to the new host, as known for IncP plasmids (Norberg et al., 2011), might therefore also not take place. Poor maintenance of these plasmids in non-Proteobacterial hosts is the likely bottleneck explaining the difference between the very wide realized transfer range and the narrower evolutionary range. Mating pair formation and conjugation systems in these plasmids are evolutionary adapted to connect and span Gram-negative membranes. The observed transfer to Gram-positive bacteria might therefore become a dead end in many cases for Gram-negative associated plasmids if the Type IV coupling and secretion system cannot efficiently spread the plasmid to other neighboring bacteria. However, an actinobacterial Mycobacterium strain has been shown to host and transfer a IncP type plasmid indicating that maintenance and transfer is possible across the Gram border (Leão et al., 2013). Also, the transient presence of a plasmid can provide the new host with a punctual adaptive gene pool and result in a short-term, but highly significant, fitness gain.

Accessory genes on plasmids are mostly arranged in transposons flanked by insertion sequence (IS) elements, which can recombine with the recipient bacterial chromosomes (e.g. Class 1 integron of pKJK5) delivering packages of fitness altering DNA without the need for plasmid replication. Additionally, transient hosts can increase the transfer range further by allowing transfer to organisms that had a lower transfer potential from the original donor strain (Yano et al., 2013).
We show within a bacterial community that there is a high variability in permissiveness to broad-host range plasmids that cannot be explained by the phylogeny of the potential recipient. The ability to take up diverse broad host range plasmids from different hosts at high frequencies as represented by the super permissive fraction of the community has not previously been described. We do not know if it is a strain-specific trait and how environmental conditions affect its manifestation. Also, we do not know to what extent the employed mating conditions might have biased the observed pattern of super permissive plasmid recipients. But, if strain-specific, these super-permissive strains would be expected to play a disproportionate role as central nodes in networks of lateral gene acquisitions (Popa et al., 2011). Most gene acquisitions occur between donors and recipients residing in the same habitat (Popa & Dagan, 2011) and while gene acquisition in nature mainly occurs within taxonomically homogenous groups, the heterogeneous soil community provides a hot-spot for gene acquisition from phylogenetically distant groups (Popa et al., 2011). In soil a few strains build the core nodes of a heavily connected network of lateral gene acquisition (Popa et al., 2011), which could be a possible indication of being part of the super-permissive fraction. These species are mainly found within Enterobacteriales (Gammaproteobacteria), Burkholderiales (Betaproteobacteria), and Staphylococci (Bacilli), groups that contain most of our super-permissive OTUs. Finding the same group of bacteria as central nodes in lateral gene transfer networks (Popa et al., 2011) and as main contributors to plasmid flow in soil suggests that there is a indeed a link between increased plasmid uptake ability and long-term gene acquisition potential.

Supplementary information is available at ISMEJ’s website

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Competing Financial Interest Statement
All authors declare no competing financial interest.

References


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**Figure 1:** Typical transconjugal microcolonies for plasmid pKJK5::gfp introduced through *E. coli* MG1655::Km\(^R\).-Lpp-mCherry. Observation was carried out with the confocal laser scanning microscope (CLSM). Transconjugants are green fluorescent, due to gfp-expression. Gfp-repressing donor strains are red fluorescent through chromosomal mCherry tagging. Black background represents soil bacteria not taking part in gene transfer events.

**Figure 2:** FACS sorting of transconjugal cells from a mating mixture initiated with soil bacteria and *E. coli* carrying pKJK5. The procedure consists in three successive gates (marked by pink stars in line A): Gate I sorts for bacterial size based on front and side scatter; Gate sorts for green fluorescent cells; Gate III selects only those green cells that possess no red fluorescence. Line A shows the sorting of the initial soil bacterial recipient community in absence of any donor strain and proves that the presence of green autofluorescent particles (A-II) does not yield false positive as they are excluded at the third gate, due to their red fluorescence (A-III). The sorting of a pure culture of the donor strain is shown in line B, where, again, no false positive events are recorded at the final gate. Line C represents the analysis of the mating mixture before sorting. Line D shows the enrichment of transconjugants after the first fast enrichment sorting step to over 80% transconjugal cells, with minor contamination by donor or soil particles. Line E shows how only pure transconjugants are obtained after the second purification sorting step.

**Figure 3:** Principal Coordinate Analysis of individual transconjugal pools, as well as of the extracted soil community (Soil_Extract) and the reference soil community as grown on filters (Soil_Filter) based on the ThetaYC algorithm (Yue & Clayton, 2005). Each axis explains a certain fraction of dissimilarity according to the axis loading given in brackets. The three different plasmids are represented by color. The three different donor strains are named next to the data points.

**Figure 4:** Phylogenetic tree showing all identified transconjugal OTUs for three different plasmids (pKJK5, RP4, pIPO2tet) from the same donor (*P. putida*). The colors of the branches mark different phylogenetic
groups. The three donor strains are shown in white letters in the trees. Green heatmap-circle around the
tree represents the log transformed relative OTU abundance in the soil reference recipient community.

Three heatmap-circles in blue and red display the x-fold over- and underrepresentation of the OTU in the
respective transconjugal pool in comparison to the abundance in the reference soil sample. Stars mark the
shared (present in all 3 transconjugal pools) and abundant (present at more than 1% relative sequence
abundance) transconjugant OTUs, which constitute the core super-permissive community fraction. Sample
size was normalized to 30000 sequences per transconjugal pool.

**Figure 5:** Phylogenetic tree showing all identified transconjugant OTUs for the same plasmid (pKJK5)
introduced through 3 different donor strains (*P. putida*; *Kluyvera* sp.; *E. coli*) (B). Colors of the branches mark
different phylogenetic groups. The three donor strains are shown in white letters in the trees. Green
heatmap-circle around the tree represents the log transformed relative OTU abundance in the soil
reference recipient community. Three heatmap-circles in blue and red display the x-fold over- and
underrepresentation of the OTU in the respective transconjugal pool in comparison to the abundance in
the reference soil sample. Stars mark the shared (present in all 3 transconjugal pools) and abundant
(present at more than 1% relative sequence abundance) transconjugant OTUs, which constitute
constituting to the core super-permissive community fraction. Sample size was normalized to 30000
sequences per transconjugal pool.

**Figure 6:** Venn diagram of transconjugal pools for plasmid pKJK5 transferred from three different donor
strains (*E. coli*, *P. putida* & *Kluyvera* sp.) (A&B) and for three different plasmids (pKJK5, RP4, pIPO2tet)
introduced through *P. putida* into the soil community. Venn diagrams are presented for OTU incidence
(C&D) and for OTU relative abundance (right, 100% represents the total number of transconjugal
sequences). OTUs were defined at 97% sequence similarity and sequence sample size was normalized to
30000 per transconjugal pool.
Figure 1:
Figure 2:
Figure 3:
Figure 4:
Figure 5:
Figure 6:
Title:
Deep sequencing of transconjugal pools reveals unexpectedly diverse bacterial community fraction receiving broad host range plasmids

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**Supplementary Information content:**

**Supplementary Table 1:** Number of FACS sorted transconjugant cells; raw and quality checked sequences for each mating combination and its replicates.

**Supplementary Table 2:** OTUs with more than 0.01% sequence representation in the soil reference community that were not represented in the transconjugal pools. Sequences were classified using the RDP classifier. Numbers behind the name indicate % sequence similarity to the closest representative.

**Supplementary Figure 1:** Phylum level distribution of the isolated and sequenced transconjugants in each of the 11 samples including all replicates per donor and plasmid combination.

**Supplementary Figure 2:** Relative over-/underrepresentation of OTUs in transconjugal pools compared with the soil reference recipient community as a function of the OTU’s Sogin phylogenetic dissimilarity to the respective plasmid donor strain. Overrepresentation is calculated as relative abundance of the OTU in the transconjugal pool divided by relative abundance in soil reference community subtracted by 1. Underrepresentation is calculated as the inverse of overrepresentation. Values given in the figure display the square root of the absolute over or underrepresentation.
### Supplementary Table 1:

<table>
<thead>
<tr>
<th>Donor</th>
<th>Plasmid</th>
<th>Replicate</th>
<th>Sorted Transconjugants</th>
<th>Raw reads</th>
<th>Sequences after mothur processing</th>
<th>Sequences/Transconjugants sorted</th>
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Number of FACS sorted transconjugant cells, raw and quality checked sequences for each mating combination and its replicates.
Supplementary Table 2:

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<th>Phylum</th>
<th>Class</th>
<th>Order</th>
<th>Family</th>
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<td>100 unclassified</td>
<td>100</td>
</tr>
</tbody>
</table>

OTUs with more than 0.01% sequence representation in the soil reference community that were not able to take up any of the introduced plasmids. Sequences were classified using the RDP classifier. Numbers behind the name displays percent of sequence similarity to the closest representative.