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Recipient Determinants Affecting Conjugational Promiscuity in Enterobacteriaceae

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Recipient Determinants Affecting Conjugational Promiscuity in *Enterobacteriaceae*

Recipient Determinants Affecting Conjugational Promiscuity in *Enterobacteriaceae*

PhD Thesis by Louise Roer December 2015

Submitted: 31.12.2015 Research Group for Genomic Epidemiology National Food Institute Technical University of Denmark Søltofts Plads 221, 2800 Kongens Lyngby, Denmark

PREFACE

The research of this PhD thesis was carried out from July 2012 to December 2015, in the Research Group for Genomic Epidemiology (former Division for Epidemiology and Microbial Genomics) at the National Food Institute, Technical University of Denmark. Part of the research was carried out at the Section of Microbiology, Department of Biology, Faculty of Science, University of Copenhagen. The research was supported by the grant 11-106571 from the Danish Council for Independent Research/Technology and Production Sciences.

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A wise man once told me to follow my dreams. As a total cliché I've always wanted to make a difference. Along this incredible journey, sometimes tough but mostly fantastic, I found my place. I'm sure that my insatiable curiosity and my eager to learn will guide me forward from this point on.

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Above all, I would like to thank my amazing husband Kenneth Roer. You have been the greatest support during this process. Thank you for putting up with me, for believing in me and making me feel that anything is possible. Thank you!

This thesis is dedicated to those who never stop believing in you and encourage you to go follow your dreams.

Skovlunde, December 2015 Louise Roer

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- I. **Roer L**, Aarestrup FM, Hasman H. The *Eco*KI Type I Restriction-Modification System in *Escherichia coli* Affects but Is Not an Absolute Barrier for Conjugation. J Bacteriol 197: 337–342. Available: http://www.ncbi.nlm.nih.gov/pubmed/25384481.
- II. **Roer L**, Hendriksen RS, Leekitcharoenphon P, Lukjancenko O, Kaas RS, Hasman H, Aarestrup FM. Is the evolution of *Salmonella enterica* subspecies *enterica* Linked to Restriction-Modification Systems? *Manuscript for submission to: mSystems.*
- III. **Roer L**, Hasman H, Pamp SJ, Kaas RS, Riber L, Sørensen SJ, Aarestrup FM. *Salmonella enterica* recipient genes affecting plasmid conjugation. *Manuscript in preparation.*

ARTICLES NOT INCLUDED IN THE THESIS

1. Hendriksen RS, Leekitcharoenphon P, Mikoleit M, Jensen JD, Kaas RS, **Roer L**, Joshi HB, Pornruangmong S, Pulsrikarn C, Gonzalez-Aviles GD, Reuland EA, Al Naiemi N, Wester AL, Aarestrup FM, Hasman H. 2015. Genomic dissection of travel-associated extended-spectrum-beta-lactamase-producing Salmonella enterica serovar typhi isolates originating from the Philippines: a one-off occurrence or a threat to effective treatment of typhoid fever? J. Clin. Microbiol. 53:677–80. Available: http://www.ncbi.nlm.nih.gov/pubmed/25428145

SUMMARY

The global emergence and rapid spread of antimicrobial resistance are considered major threats to human health. Horizontal transfer of plasmids is the most important method for transmission of resistance, a method which with the accumulation of sequence data in recent years, was shown to be highly underestimated. Thus, to be able to control and prevent the spread of undesirable genetic traits, it is crucial to understand the factors controlling the conjugational transfer.

With the great attention to whole genome sequencing it is now possible to rapidly predict a variety of different characteristics of pathogens, including antibiotic resistance, plasmid replicons, and the evolutionary relatedness of bacteria. Thus, combinations of the bioinformatic technology and the classical molecular approaches in the laboratory provide new opportunities for research nowadays.

The aim of this PhD study was to improve the knowledge for the implication of restrictionmodification systems in horizontal gene transfer and evolution, and to gain new knowledge of recipient determinants included in regulation of plasmid uptake in bacteria by using a combination of bioinformatics and applied microbiology in the laboratory.

In **Roer I** we addressed the dogma of RM systems being a barrier for horizontal gene transfer, which, to our knowledge never has been proven at a genetic level for conjugation. In this study we constructed an isogenic setup, with single gene-knockout mutants of the type I endonuclease *Eco*KI and the cognate methyltransferase M.*Eco*KI. By utilizing the plasmid pOLA52, with 2 recognitions sites for the RM system, it was possible to investigate the uptake and maintenance of an unmethylated plasmid in the wild-type strain (intact *hsdR* gene), and in a *hsdR* deficient derivate. The inactivation of the restriction enzyme caused a 7-fold increase in conjugational transfer for the unmethylated plasmid compared to the wild-type with intact *hsdR* gene. Interestingly, the RM system did not impose an absolute barrier for conjugational transfer of the wild-type. Thus, these findings are leaving a question of the evolutionary impact of RM systems on organisms which preferably exchange DNA by conjugation.

In **Roer II**, we therefore decided to elucidate on the association between RM systems and the evolution of the 221 genomes of the conjugative bacterial species *Salmonella enterica* ssp. *enterica*. In addition to evaluate the evolution, we decided to investigate the connection between RM systems and the content of some of the evolutionary drivers, plasmids and pathogenicity islands. For the purpose of identifying RM systems and Salmonella Pathogenicity Islands, *RM-Finder* and *SPI-Finder* were constructed. The detected RM systems were compared with the evolution of the

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species, depicted as a network of core- and pan-genome trees. Though, from the study we were able to conclude that in comparison with natural transformable species, there was no evidence for RM systems shaping the evolution of the conjugational species *Salmonella enterica* ssp. *enterica.* In addition, no pattern between the mobile genetic elements and RM systems was observed.

In **Roer III**, we aimed to identify other recipient determinants controlling the conjugational uptake and maintenance. To identify possible gene candidates we used a combination of modern molecular techniques in the laboratory, and bioinformatic gene prediction. The attempt to identify gene candidates was performed on a collection of 93 *Salmonella enterica* ssp. *enterica* isolates, covering 54 different serovars. An initial conjugation experiment was performed to clarify the recipient potential for the 93 isolates, with subsequent grouping into good and poor recipients. From this rather diverse collection, we were not yet able to detect common gene candidates controlling neither increase nor decrease in conjugation abilities. A parallel study, only focusing on *S.* Enteritidis, proved able to detect 33 gene candidates potentially involved in conjugation restriction abilities which are being investigated and awaiting verification. This approach indicates that on species level no common gene controlling conjugation could be detected and other methods should be considered.

Overall, this PhD study has assessed the old dogma of RM systems being a barrier for horizontal gene transfer. This was previously proven in the transfer by transduction and transformation, however in this thesis we provide evidence that the imposed barriers are indefinite in conjugational transfer. Further, the work in this thesis has provided user-friendly tools for easy *in-silico* detection of RM systems (*RM-Finder*) and the transferable genomic islands conferring pathogenicity in *Salmonella* (*SPI-Finder*). It was furthermore demonstrated that, despite the clear correlation between RM systems and evolution in the natural transformable organism *N. meningitidis*, RM systems where not the main driver of evolution when turning to organisms which preferably exchange their DNA material by conjugation. Additionally, the study proved that the possession of mobile genetic elements like plasmids and SPIs cannot be reflected in the RM systems. The PhD study suggests that recipient genes are involved in the control of conjugation; however, different approaches and serovar and strain specific genes should be investigated.

This PhD thesis has altogether improved the knowledge on the influence of RM systems in conjugation and evolution, and placed a potential for further investigation of recipient determinants affecting conjugation.

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RESUMÉ

Den globale udvikling og hurtige spredning af antimikrobiel resistens anses for at være en stor trussel for det menneskelige helbred. Horisontal overførsel af plasmider er en af de mest betydningsfulde mekanismer for overførsel af resistens, en metode som med akkumuleringen af sekvens data de seneste år har vist sig at være særdeles underestimeret. For at kunne kontrollere og forebygge spredningen af uønskede genetiske træk er det afgørende at forstå de faktorer der kontrollerer overførsel ved konjugation. Med den øgede opmærksomhed og udvikling indenfor helgenom sekventering er det nu muligt hurtigt at forudsige forskellige karaktertræk hos patogene bakterier, hvilket inkluderer antibiotika resistens, plasmid replikon og det evolutionære forhold mellem bakterier. Således kan kombinationen af bioinformatik og klassiske molekylære fremgangsmåder i laboratoriet tilføre nye muligheder til vor tids forskning.

Formålet med dette ph.d.-studie var at øge den allerede eksisterende viden indenfor betydningen af RM systemer i horisontal gen overførsel og evolutionen, og at opnå ny viden indenfor recipient faktorer involveret i reguleringen af plasmid optaget i bakterier, ved at benytte en kombination af bioinformatik og anvendt mikrobiologi i laboratoriet.

I **Roer I** håndterede vi dogmet for RM systemer som værende en barriere for horisontal gen overførsel, hvilket ifølge vores viden aldrig har været vist på et genetisk niveau for konjugation. I dette studie konstruerede vi et iso-genetisk opsæt med enkelt gen knockouts af type I restriktionsenzymet *Eco*KI og dens tilhørende methyltransferase M.*Eco*KI. Ved at bruge plasmidet pOLA52, som besidder 2 genkendelses sekvenser for RM systemet, var det muligt at undersøge optaget og opretholdensen af et ikke methyleret plasmid i vild-type stammen (med intakt *hsdR* gen), og i et derivat med ødelagt *hsdR* gen. Inaktiveringen af restriktionsenzymet forårsagede en 7-folds stigning i konjugations overførsel af det ikke methylerede plasmid, sammenlignet med vild-type stammen med intakt *hsdR* gen. RM systemet viste sig overraskende nok ikke at tilføre en fuldstændig barriere for konjugation til vild-typen. Resultaterne efterlader derved spørgsmål om den evolutionære indvirkning af RM systemer på organismer som fortrinsvis udveksler DNA via konjugation.

I **Roer II** besluttede vi derfor at undersøge sammenhængen mellem RM systemer og evolutionen af 221 genomer af *Salmonella enterica* ssp*. enterica,* som benytter konjugation ved overførsel. Ud over at evaluere evolutionen besluttede vi at undersøge sammenhængen mellem RM systemer og evolutionære drivkræfter som plasmider og patogenetiske øer. Til at identificere RM systemer og

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Salmonella Patogenetiske øer (SPI) udviklede vi *RM-Finder* og *SPI-Finder*. De opfangede RM systemer blev sammenlignet med evolutionen af *Salmonella*, gengivet i et netværk af core- og pangenom træer. Dette studie gjorde det muligt at konkludere, at sammenlignet med naturligt kompetente organismer viste RM systemer sig for ikke at forme evolutionen i konjugationsorganismen *Salmonella enterica* ssp. *enterica.* Yderligere var det heller ikke muligt at observere et mønster mellem mobile genetiske elementer og RM systemer.

I **Roer III** forsøgte vi at identificere andre recipient faktorer involveret i kontrollen af konjugations overførsel. Til at identificere mulige gen-kandidater benyttede vi en kombination af moderne molekylære teknikker i laboratoriet og bioinformatisk forudsigelse af gener. Forsøget på at identificere gen-kandidater blev udført på en kollektion af 93 *Salmonella enterica* ssp. *enterica* isolater af 54 forskellige serovars. For at bestemme recipient potentialet af de 93 isolater, blev et indledende konjugations forsøg udført. Herefter blev isolaterne inddelt i grupper af gode og dårlige recipienter. Fra denne meget diverse kollektion har det endnu ikke været muligt at finde nogen fælles gen-kandidater der kontrollerede hverken op- eller nedregulering i evnen til at konjugere. *S.* Enteritidis blev undersøgt i et parallelt studie, hvor det var muligt at identificere 33 gen-kandidater der potentielt kan være involveret i nedregulering af konjugation, hvilket bliver undersøgt nærmere og afventer verificering. Med denne fremgangsmåde er det ikke er muligt at identificere gener der kontrollerer konjugation på organisme niveau, og andre metoder bør overvejes.

Alt i alt har dette ph.d.-studie behandlet det gamle dogme for RM sytemer som værende en barriere i horisontal gen overførsel. Dette har tidligere været vist i overførslen via transduktion og transformation. I denne afhandling leverer vi et bevis for at de pålagte barrierer ikke er fuldkomne i konjugativ overførsel. Yderligere har dette studie leveret brugervenlige redskaber til in-silico påvisning af RM systemer (*RM-Finder*) og de overførbare genomiske øer som tilfører patogenese i *Salmonella* (*SPI-Finder*). Det blev yderligere demonstreret, på trods af en klar sammenhæng mellem RM systemer og evolution i den naturlig overførbare organisme *N. meningitidis*, så var RM systemer ikke en drivende kraft i evolutionen af organismer der fortrinsvis udveksler DNA via konjugation. Studiet viste yderligere at besiddelsen af mobile genetiske elementer, såsom plasmider og SPIs, ikke reflekteres i RM systemerne. Dette ph.d.-studie antyder at organisme specifikke gener kan være involveret i kontrollen af konjugation, men forskellige metoder bør overvejes og serovar og stamme specifikke gener bør undersøges yderligere. Denne ph.d.-afhandling har overordnet øget den eksisterende viden om RM systemers indvirkning på konjugation og evolution, og givet anledning til yderligere undersøgelser af recipient faktorer der kan have indflydelse på konjugation.

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LIST OF ABBREVIATIONS

BACKGROUND

The evolutionary pressure from the increased use of antibiotics in human and veterinary medicine has contributed to the emergence and global spread of antimicrobial resistance, which today is considered one of the major human health threats. Invention of new antimicrobial agents targeting bacterial infections might be an expensive and comprehensive process; consequently it is, as far as possible, important to protect the current agents, through improved knowledge of emergence and spread of the genetic traits.

Horizontal transmission of plasmids harbouring antimicrobial resistance genes is one of the most important methods of resistance transmission between bacterial species. However, little is known about the factors controlling the conjugational transfer of plasmids between different species and clones, and without expanding this knowledge it may be impossible to prevent the spread of genetic traits through conjugation.

It has been observed that host-specific and non-host-specific serovars of *Salmonella* isolated from the same reservoir differs in their antibiotic resistance pattern (VAN DUIJKEREN *et al.* 2002; LAUDERDALE *et al.* 2006). Additionally, preliminary experiments for *Escherichia coli* have suggested that pathogenic clonal lineages serve as better plasmid recipients than non-pathogenic clones (unpublished). Thus it is possible that the ability to promote or reject the uptake of foreign DNA, such as plasmids, is defined by the genetic differences between the bacterial linages. Possible gene candidates include restriction-modification (RM) systems which are thought to act as bacterial host barriers against foreign DNA. For RM systems it is practically common knowledge, hence a dogma that they interfere with horizontal gene transfer, however for conjugation this has not yet been confirmed by a systematic experimental approach. Thus, other recipient genes may be included in the regulation of plasmid uptake. If present, identifying these additional genes acting as barriers or enhancers in conjugation might clarify the spread of plasmid transferred resistance.

PURPOSE

The purpose of the PhD project was to elucidate the implication of restriction-modification systems in horizontal gene transfer between bacterial clones, with the focus on conjugational transfer of plasmid borne antimicrobial resistance in related clones of *Escherichia coli.* Furthermore, to establish the relationship between restriction-modification systems and evolution in *Salmonella* in

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comparison with the content of mobile genetic elements like plasmids and *Salmonella* pathogenicity islands.

Finally, the purpose was to identify additional genes that might block or enhance uptake in the recipient bacteria.

RESEARCH APPROACH

The specific studies conducted during this PhD project focused on the following objectives:

- 1. To quantify the importance of restriction-modification systems in conjugational transfer of plasmids.
- 2. To investigate the influence of restriction-modification systems in long term evolution of *Salmonella enterica* ssp. *enterica.*
- 3. To identify and assess gene candidates from whole genome sequenced bacterial isolates, which promote or resist plasmid uptake.

Chapter 1 **EVOLUTION OF MICROBIAL COMMUNITIES**

The environment of microorganisms can vary considerably, thus bacteria need to be dynamic and constantly capable of adapting to new surroundings. Accumulation of these adaptive changes leads to diversity within the different species. To understand, control and fight the spread of pathogens, we need to understand the processes on how bacteria diversify and the mechanisms that control the evolution. The prokaryotic evolution is a result of three main forces: acquisition of genes, loss of genes and changes within genes, [Figure 1](#page-19-0) (FRASER-LIGGETT 2005; PALLEN and WREN 2007; USSERY *et al.* 2009). Events within these forces happen all the time, leaving evolutionary traces on bacterial genomes.

1.1 Evolution by Single-nucleotide Changes

Single-nucleotide polymorphisms (SNPs) are the simplest and smallest-scale variation within changes of genes (FRASER-LIGGETT 2005). A mutation or base substitution in a genome can be caused by two fundamental processes: incorporation of wrong nucleotides during replication or mutagenesis triggered by physical- or chemical damage or changes. Thus exposure to antimicrobial agents such as rifampicin and ciprofloxacin can cause spontaneous mutations, and in *Helicobacter pylori* frequencies at 1×10^{-8} to 2×10^{-8} per cell per division were detected, conferring rapid adaptation to exposed environments (WANG *et al.* 2001). As spontaneous mutations in general are rare it is unlikely that the same substitution will occur in multiple genomes (unless there is a strong

selective pressure to obtain this given mutation). Hence the presence of the same base SNP in two independent genomes implies that the substitution likely descend from a shared ancestor. However, exceptions do exist e.g. for genes under selective pressure, like resistance genes. Genes under selective pressure are more likely to mutate to make the bacteria capable of adapting to the changed environment, thus for such genes the same base SNP can be found in multiple independent genomes. SNP analysis has been applied to a variety of uniform pathogens with rather conserved genomes (GUTACKER *et al.* 2002; PEARSON *et al.* 2004; MONOT *et al.* 2005; TOUCHMAN *et al.* 2007), thus a problem with the more variable species have been the high proportion of mobile genetic elements (MGE) causing larger insertions, which challenges the one base substitution assumption. However, SNP analysis has proven capable of differentiating between *Escherichia coli* that had diverged for only 200 generations (SHENDURE 2005). Therefore it appears highly suitable for detection of short-term evolution such as antimicrobial adaption, rather than long-term evolution and acquisition of MGEs (ALLARD *et al.* 2012, 2013; GIERALTOWSKI *et al.* 2013; HOFFMANN *et al.* 2013).

Figure 1 | Genome evolutionary events. Genome size changes due to plasmid exchange, insertion sequence elements, phages and duplication and deletions of genes. The genomes in the figure expand (left) or decrease (right) from the light gray to the dark grey backbone. In the events 1, 2 and 3 from the light gray genome (left) a piece of DNA inserts itself into the chromosome, and from the middle genome additional insertion happens in event 4 and 5, the latter located at the same position as event 3. All these insertions could be caused by an IS element, a phage or another mobile genetic element. During genome reduction (right) the inserted region from event 1 was partly removed in event 6, leaving a residue in the genome. Contrary in event 7, more DNA was deleted than initially introduced in event 2, resulting in a genome reduction in comparison with the original genome (light gray genome left). Further, in event 8 a partial deletion of the two individual insertion from event 3 and 5 was resulting in a novel junction, thus in a second step the remaining insertions was deleted. The inserted DNA from event 4 was permanently fixed in the genome. Figure 1 is adapted from (USSERY *et al.* 2009).

1.2 Evolution by Gene Loss and Deletions

As bacterial genomes remain about the same size, a balance of gene duplication or acquisition and gene loss or deletion must exist (MIRA *et al.* 2001). The most severe form of genome reduction is found in intracellular host-associated bacteria such as endosymbionts. These bacteria are segregated from the surroundings in a stable and undemanding niche, which for *Rickettsia prowazekii* has resulted in a substantial downsizing of the genome, leading to renunciation of many of the inherent genes (ANDERSSON *et al.* 1998; WERNEGREEN 2005). With gene loss and deletion of large DNA segments in the reductive evolution, it was initially thought that genes targeted for loss were only redundant genes not necessary for survival in the new host environment. However an increasing number of comparative genomic studies have suggested that a positive selection for gene loss might also be advantageous for the bacteria. In *Salmonella enterica* (*S. enterica*), different deletions appeared beneficial (KOSKINIEMI *et al.* 2012), and for *Bordetella pertussis* and *Shigella* the loss of a cell surface antigen enhanced the ability to avoid the host immune system (NAKATA *et al.* 1993; PARKHILL *et al.* 2003).

1.3 Evolution by Gene Duplication and Acquisition

Acquisition of additional genes through gene duplication can increase the number of key genes responsible for regulation, secondary metabolisms and transport. As much as 50% of larger genomes can be represented by gene paralogs (KONSTANTINIDIS and TIEDJE 2004; FRASER-LIGGETT 2005). However, acquisition of genes through horizontal gene transfer (HGT) is the most effective source of variation within the genome (PALLEN and WREN 2007), providing a possibility for rapid alterations for the constant changing demands on the bacteria (BOUCHER *et al.* 2003). Different genetic elements can be transferred by HGT, among others the virulence determinants on pathogenicity islands (PAIs) and prophages, which appear to play a major role in the evolution of pathogens. In a comparative analysis of *Staphylococcus aureus* and the closely related *Staphylococcus epidermidis*, it was established that the main cause of variation in virulence and pathogenicity was caused by the acquisition of genomic islands, presumably obtained through HGT (GILL *et al.* 2005). Another prominent source for diversification by HGT is mediated by plasmids, which can rapidly confer antimicrobial resistance or numerous of virulence factors and thereby evolve existing clonal lineages. The opportunistic pathogen *Bacillus cereus* is normally differing from the etiologic agent of anthrax, *Bacillus anthracis*, by two plasmids, toxic-encoding pXO1 and capsule gene coding plasmid pXO2. However, whole genome sequence (WGS) analysis revealed that the *B. cereus* G9241 strain was harbouring a plasmid 99.6% identical with the *B. anthracis*

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plasmid pXO1, and a previously unidentified plasmid encoding polysaccharide capsule cluster, resulting in a virulent *B. cereus* strain (HOFFMASTER *et al.* 2004).

Figure 2 | Concatenated Core-genome tree of 221 *Salmonella enterica* **isolates.** The tree is based on 1,072 core gene clusters, with colour indications of restriction-modification systems present in the isolates. The bootstrap values are indicated at each branch. Figure 2 is adapted from (**Roer II**).

With the accumulation of prokaryotic genome sequences, the challenge of constructing wholegenome phylogenies [\(WGP\)](#page-21-0) began. The extent and influence of HGT on the evolution, as indicated in the two examples with *Staphylococcus* and *Bacillus,* together with disagreements between WGP and rRNA phylogenies indicated that a reconstruction of the Tree of Life would be difficult or even impossible (DOOLITTLE 1999). Consequently, rather than illustrating WGP as a single tree, a consensus appeared that the evolution of prokaryotes was more appropriate depicted as a network of trees covering both horizontal and vertical transferred genes (FRASER-LIGGETT 2005). In Roer *et al.* (**Roer II**), the evolution of *S. enterica* was depicted by the network of the core- and pan-genome

trees, presented in Figure 2 and Figure 3. The pan-genome is defined as any gene family in the bacterial species, including both the core-genome and the dispensable genome. Thus, a pan-genome can either be ope[n or closed](#page-21-0), d[ependent o](#page-22-0)n whether addition of new genomes to the analysis reveals new genes (open), or no new genes (closed) to the pan-genome (MEDINI *et al.* 2005; PALLEN and WREN 2007).

Figure 3 | Pan-genome tree of 221 *Salmonella enterica* **isolates.** The tree is constructed from a presence or absence matrix of gene clusters in the *Salmonella* genomes. The colours indicate the restriction-modification systems present in each isolate. The bootstrap values are shown at each branch. Figure 3 is adapted from (**Roer II**).

The core-genome is the pool of conserved genes shared by all the strains of the bacterial species, which normally are essential for bacterial growth. [The disp](#page-22-0)ensable genome is all the accessory genes present in some but not all strains of the bacterial species, together with the strain-specific genes (MEDINI *et al.* 2005). Further, the accessory genes are contributing to the diversity of the species and consist of genes encoding selective advantage including colonization of new hosts,

adaption to different niches or confer antibiotic resistance. The core-genome tree is constructed from an alignment of the core-genes of the strains, accounting for the vertical evolution. The pangenome is a picture of the dispensable genes as the core-genes are present in all strains, and not contributing to the pan-genome tree. The pan-genome tree is constructed from a presence or absence matrix of the genes, where the relative Manhattan distance between the genomes is used for hierarchical clustering. The pan- and core-genome tree network is valuable in detecting variations within conserved genes but also HGT.

Chapter 2 **PROMISCUITY OF BACTERIA**

It is well-established that considerable horizontal gene transfer can occur between bacterial species. However, along with the accumulation of sequence data it became clear that the genetic and biochemical diversity in the prokaryotic world was highly underestimated, thus horizontal gene transfer was far more extensive in some bacterial species than previously anticipated (PERNA *et al.* 2001; FRASER-LIGGETT 2005).

Some bacterial species like *Mycobacterium tuberculosis*, *Mycobacterium leprae*, *Yersinia pestis*, and *Bacillus anthracis* were found to be genetically uniform pathogens (PALLEN and WREN 2007). In addition, *B. anthracis* was shown to have a closed pan-genome, where the addition of the fourth genome to the analysis did not add new genes to the pan-genome (TETTELIN *et al.* 2005). Furthermore, the pan-genome of nine *B. anthracis* genomes was shown to consist of 2,893 core genes, 85 accessory genes and seven strain-specific genes (ROULI *et al.* 2014). Uniform species are likely to have closed pan-genomes, thus they are more conserved, they have a low ability to acquire foreign DNA, and are normally found in niches with limited access to the global microbial gene pool (MEDINI *et al.* 2005).

In contrast to the bacterial species with closed pan-genomes, bacterial species that normally are capable of colonizing multiple environments and can exchange genetic material in multiple ways, like *Salmonella*, *E. coli*, *Streptococci*, *Meningococci* and *Helicobacter pylori,* are likely to have an open pan-genome, where each new genome sequence adds new genes to the overall gene pool of

the species. Thus, these bacterial species seem more promiscuous in the sense that they are able to acquire DNA outside their own species. With the completion of the genomic sequence of the pathogenic *E. coli* O157:H7 in 2001, Perna *et al.* compared the genome with the non-pathogenic laboratory *E. coli* K-12 strain and revealed 1,387 new genes encoded in strain-specific clusters of *E. coli* O157:H7. These genes encoded protein candidates of virulence factors, numerous prophages and alternative metabolic abilities. This is a practical example of how two members of the same species can differ almost 30% (PERNA *et al.* 2001), which has also been shown for bacterial species such as *H. pylori* and *Staphylococcus aureus,* both with an average of strain specific genes recorded to ~22% (SALAMA *et al.* 2000; FITZGERALD *et al.* 2001). However, these findings were observed in the years 2000 and 2001, where a limited number of sequenced genomes were available. Therefore, assessing the proportion of strain specific genes in the present time could potentially give a different result, with the ongoing exchange of genetic material.

Various studies have been assessing the pan- and core-genomes of *E. coli,* hence estimating the core-genome to be between 1,500-2,000 gene families (WILLENBROCK *et al.* 2007; CHATTOPADHYAY *et al.* 2009; TOUCHON *et al.* 2009; VIEIRA *et al.* 2011; KAAS *et al.* 2012). Additionally, the pan-genome was found in one early study to consist of 11,432 gene families when assessing 20 genomes (TOUCHON *et al.* 2009), while a study only three years later found 16,676 pan-genes when assessing 186 genomes (KAAS *et al.* 2012).

Another highly diverse species is *Salmonella enterica*, which is divided into six subspecies of which *S. enterica* subspecies *enterica* alone is grouped into more than 1,500 individual serovars. Some serovars are defined as host-specific, hence they are only found in a limited and specific number of hosts, compared to the non-host-specific serovars which can be found in a broad range of hosts. Evaluating the pan- and core-genome of *S. enterica*, one study on 73 genomes showed a coregenome of 2,882 gene families with a cognate pan-genome of 10,581 gene families (LEEKITCHAROENPHON *et al.* 2012), where an another study utilizing the same method, estimated the core-genome on 221 genomes to comprise 2,138 gene families with a cognate pan-genome of 16,375 gene families (**Roer II**). The pan-genomes of both *E. coli* and *S. enterica* appear to be open, and follow the rule that addition of more genomes to the analysis causes larger pan-genomes. However, it has been observed for *Salmonella* that host-specific serovars possess considerably less genes related to antimicrobial resistance than serovars with multiple hosts, even though the isolates were isolated from the same reservoir (VAN DUIJKEREN *et al.* 2002; LAUDERDALE *et al.* 2006). This could indicate, that the difference between host- and non-host-specific *S. enterica* could explain

their ability to acquire new genes, as the need for host-specific serovars to adapt to new surroundings is smaller since they are limited to only a few hosts. Thus, some bacterial species and serovars seem to be capable of diversifying by acquiring new genetic material through plasmids or other MGEs more easily than others. The case with *S. enterica* indicates a segregation of promiscuity is possible within a single but yet diverse species.

2.1 Plasmids

Plasmids play a major role in the exchange of genetic material between bacteria, thereby enhancing the genetic diversity and the ability for bacteria to adapt to changing environments. The term plasmid was suggested by Joshua Lederberg in 1952, as a common phrase for all extrachromosomal hereditary determinants, both genetically simple or complex (LEDERBERG 1952). However, now plasmids are described as either linear of circular double-stranded DNA elements capable of autonomous replication in their host, and having a genome size ranging from approximately 750 bp to 2.58 Mbp (SHINTANI *et al.* 2015). Plasmids can be classified by several different criteria, including their transmissibility, incompatibility groups, copy number and host range. Some plasmids are thought of as selfish DNA elements, as they are self-transmissible, and can replicate and survive within a host. However, using these criteria, not all plasmids are to be considered as selfish; in a comparison of 1,730 fully sequenced plasmids, only 15% of the plasmids were predicted to be self-transmissible, compared with 24% mobilizable plasmids which need help to be transmitted and 61% non-transmissible plasmids (SMILLIE *et al.* 2010). In 2015, Shintani *et al.* classified 4,602 complete plasmid genomes by their host (data collected August 2014). They found that 137 of the plasmid genomes were from archaea, 47 were from eukaryotes, and the remaining 4,418 plasmids were from bacteria (SHINTANI *et al.* 2015).

The best studied group of self-transmissible plasmids are the IncP-1 plasmids, which initially were found in clinical bacterial isolates (JOBANPUTRA and DATTA 1974; JACOBY *et al.* 1976). They are still of clinical concern due to their ability to carry and facilitate the spread of antibiotic resistance (INGRAM *et al.* 1973; NOVAIS *et al.* 2006). Additionally, IncP plasmids are present in diverse environments like manure (BINH *et al.* 2008), water streams (SMALLA *et al.* 2006), waste-water treatment plants (SCHLÜTER *et al.* 2007), and agricultural soils (TOP *et al.* 1995; SEN *et al.* 2011), which is also the case for the IncP-1 plasmid pKJK5 isolated from a soil environment (BAHL *et al.*) 2007). A schematic map of a typical IncP-1 plasmid is depicted in [Figure 4.](#page-27-1) The plasmids consist of two regions; a 'plasmid backbone' and the accessory genes. The plasmid backbone includes genes

responsible for replication (*trfA*), transfer and maintenance (*tra* and *ctl*), whereas the accessory regions encode genes that can increase the fitness to the host such as antimicrobial or metal resistance and alternative metabolic pathways (SCHUMANN *et al.* 2008). For the IncP-1 plasmids, the accessory genes are typically found in two different areas, the first located between the *trb* operon that encode mating-pair formation, and the *tra* operon that encode genes responsible for plasmid transfer. Tthe second region is located between the origin of replication (*oriV*) and *trfA* responsible for plasmid replication (SCHUMANN *et al.* 2008; SEN *et al.* 2013). The accessory regions can carry other MGEs such as integrons (Ins), insertion sequences (ISs) and transposons (Tns) that enables the accessory genes to move between the bacterial chromosome and plasmids, contributing to the bacterial evolution (SCHLÜTER *et al.* 2007; SCHUMANN *et al.* 2008). However, plasmids that do not enhance beneficial traits to their hosts also exists, and are named cryptic plasmids (VAN ELSAS *et al.* 1998).

Figure 4 | Genetic map of a typical IncP-1 plasmid. The map displays the different functional elements located on a typical IncP-1 plasmid. Backbone genes are coloured in dark grey, with accessory genes displayed in light grey. The backbone of the plasmid is composed of an origin of replication (*oriV*) and an operon for replication initiation (*trfA*). The *trb* operon is involved in bridge formation and the *tra* operon in DNA processing, both during conjugation. The central control region (*ctl*) comprises regulatory genes involved in maintenance and plasmid stability, whereas the accessory regions can consist of host-beneficial genes, whit potential traits. Figure 4 is adapted from (SEN *et al.* 2013)

2.1.1 Replication

One of the main reasons for the success of plasmids is their ability to self-replicate within their host. In plasmids, the three different well-known mechanisms for replication of circular plasmids includes rolling-circle replication, theta-type replication, and strand displacement-type replication (DEL SOLAR *et al.* 1998).

Rolling-circle replication (RCR) plasmids are found in both Gram-positive and Gram-negative bacteria as well as archaea. RCR is named due to the apparent rolling movement during replication. Plasmids replicating by RCR are generally very small and compact (less than 10 kb) and found in multiple copies (KHAN 2005; SCHUMANN *et al.* 2008). Additionally, RCR plasmids include both conjugative and mobilizable plasmids, often carrying antibiotic or metal resistance genes, however some of the plasmids are cryptic (KHAN 2005). The Rep region of plasmids encodes proteins that are involved in the initiation, elongation and termination of replication (DEL SOLAR *et al.* 1998).

The large plasmids are either replicating through the theta-, or strand displacement-type replication mechanisms (MEIJER *et al.* 1995; GUGLIELMETTI *et al.* 2007). The theta-type replication (TR) is named by the molecule shape of the replication intermediate, which is visible under electron microscopy. The replication initiates with melting of the parental strands and primer RNA synthesis, followed by extension of the primers. In TR the leading strand is synthesized continuously, and the lagging strand discontinuously (DEL SOLAR *et al.* 1998). In contrast, strand displacement-type replication (SDR) synthesizes the two strands bidirectionally. Replication by SDR is initiated by the combination of three proteins, further this mechanism has been associated with mobilizable plasmids of the IncQ family (SAKAI and KOMANO 1996; DEL SOLAR *et al.* 1998).

Besides the size difference between RCR and the two other methods, a substantial difference is the single-stranded DNA intermediate of RCR plasmids (DEL SOLAR *et al.* 1998; KHAN 2005), providing plasmid instability of plasmids replicated by rolling-circle.

Following successful replication of the plasmids, they will segregate to the daughter cells by vertical transfer and establish in the new host cell.

2.1.2 Host Range of Plasmids

The plasmid host range is generally described as the collection of hosts in which a plasmid can replicate. Thus this host range is usually smaller than the transfer range (the range of hosts the plasmid can be transferred to by conjugation), though bigger than the range of host in which the plasmid can be maintained stably without selection (DE GELDER *et al.* 2007). Plasmids can either have a narrow or a broad host range. Narrow host range (NHR) plasmids can be limited in their transfer process, by unsuccessful formation of mating pairs, or in difficulties with plasmid maintenance caused by incorrect expression of the plasmid replication system in the recipient (THOMAS and NIELSEN 2005). Opposite NHR, Broad Host Range (BHR) plasmids are capable of transferring to and replicating within a comprehensive range of hosts. Though, different definitions

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of BHR have been utilized, with some researchers defining BHR as plasmids capable of transferring between different genera, whereas others define BHR as those plasmids that can transfer and replicate within at least two subgroups of a bacterial species (SZPIRER *et al.* 1999). As the transfer range of some BHR plasmids are wider than the replication range, the combination of conjugative transfer followed by integration into the recipient chromosome enables the ability to spread genetic material on a broad taxonomic range. Additionally, mobilization of non-conjugative plasmids with a broader host range increases the spread. For the self-transmittable (conjugative) plasmids, the group of IncP-1 are considered those with the broadest host range (THOMAS 1987; ADAMCZYK and JAGURA-BURDZY 2003; SCHLÜTER *et al.* 2007). Klümper *et al.* showed that the IncP-1 plasmid pKJK5 possessed the ability to transfer to a surprisingly diverse group of recipients extracted from soil, thus showing a rather diverse host range (KLÜMPER *et al.* 2014). However transferring pKJK5 to various *S. enterica* recipients showed differences in the ability to take up and maintain the plasmid (**Roer III**).

Small mobilizable (non-self-transmissible) plasmids, like the IncQ plasmid RSF1010, have showed tendencies of broader host range than large plasmids with the ability to conjugate (GUERRY *et al.* 1974). However, the mobilizable plasmids are deeply dependent on the conjugative plasmids to transfer between hosts.

2.1.3 Incompatibility

Incompatibility has not only been used to classify plasmids since the 1970s (SHINTANI *et al.* 2015), but it still has huge impact on the ability of plasmids to be maintained in a new host. Incompatibility (Inc) is defined as the inability for two plasmids to coexist in the same host without a selective pressure. The failure to coexist can be caused by similarity in replication and partition systems. The partition system is covering the active process that controls a consistent distribution of low-copynumber plasmids to daughter cells (BOUET *et al.* 2007). Hence, if a potential new host already possess an analogous replication mechanism, the new host is unable to distinguish the two plasmids and neither will be replicated frequently enough to be stably maintained (NOVICK 1987; BOUET *et al.* 2007). This will in most cases lead to a loss of the newly transferred plasmid, as the initial copy number is lower.

A PCR-based method was previously used to classify the replicon types within *Enterobacteriaceae*. The limitations to identifying novel and divergent replicons resulted in a more accurate classification based on the full-length sequence (CARATTOLI 2009), enabling grouping of

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unidentified Inc groups (SHINTANI *et al.* 2015). Online WGS tools like *PlasmidFinder* is now available to classify the replicon type based on sequencing data (CARATTOLI *et al.* 2014).

In a review by Shintani *et al.* from 2015, the current number of Inc groups was reported. For *Enterobacteriaceae* 27 different Inc groups were described, 14 Inc groups for *Pseudomonas* and about 18 Inc groups for *Staphylococcus* (SHINTANI *et al.* 2015), indicating a broad member of MGEs.

2.2 Pathogenicity Islands

Another important group of mobile elements that are presumed to be involved in HGT are the Pathogenicity islands (PAIs), which are decisive in the virulence of bacteria (GILL *et al.* 2005; PALLEN and WREN 2007). The HGT facilitates a rapid acquirement of complex virulence functions from different species. PAIs were first described in pathogenic *E. coli*, where Hacker and collegues investigated the genetics behind virulence. By deletion of two large DNA inserts in the chromosome, they found that deletions affected virulence expression, thus mutants with deletions of both inserts resulted in a entirely avirulent strain (KNAPP *et al.* 1986; HACKER *et al.* 1990). These regions responsible for virulence in *E. coli* 536 led to the term pathogenicity islands (BLUM *et al.* 1994).

Figure 5 | General structure of Pathogenicity Islands. (A) shows a schematic model of a bacterial genomic island. The PAIs are frequently inserted at sites with tRNA or tRNA-like genes, flanked by directed repeats (DR) which are used in insertion and deletion processes. (B) The content of guanine and cytosine (G+C) differs from the rest of the genome, which is often used to identify new PAIs. [Figure 5](#page-30-1) is adapted from (SCHMIDT and HENSEL 2004)

PAIs can be diverse in both structure and function, however numerous common genetic features have been observed. For a genomic island to be a PAI it has to carry at least one virulence gene, illustrated by V_1 , V_2 etc. in [Figure 5.](#page-30-1) The majority of the islands can be relatively large, with the majority somewhere between 10 and 200 kb. The PAIs will only be present in pathogens of the same or closely related bacterial species. The guanine and cytosine composistion (base composition) of bacterial DNA can vary from 25 to 75%, however PAI often differ in base composition compared to the rest of the genome in which they are found (HACKER and CARNIEL 2001; HENSEL 2004). As PAI are belived to be acquired by HGT, the reason for this difference in composition is considered to derive from the donor species. However, it has been observed from studying genome evolution that horizontally acquired DNA will tend to shift towards the base composition in the recipient genome, making it difficult to explain why inherent PAIs still differ in their base composition (SCHMIDT and HENSEL 2004).

PAIs are often, like bacteriophages (CAMPBELL 1992), located within an insertion sites such as tRNA, and frequently associated with DNA mobility. Directed repeats (DR) might serve as recognition site for integration, however they can also contribute to the instability of PAIs as DR act as recognition sequences for enzymes that are involved in deletion of MGEs. When PAIs are transferred by transduction (bacteriophages) integrases might mediate the integration into the chromosome of the bacteria, as well as excision when the bacteriophage needs to enter a lytic cycle. Tns are another type of MGEs that are often found in PAIs, thus besides changing the location within the bacterial chromosome, Tns can also jump between chromosome and plasmids. Additionally, the combination of two or more insertion elements (IS) can also result in mobilization of the PAIs, as well as inactivation of genes. Thus, all the MGEs present at PAIs causes unstablility, and deletions of PAIs from chromsomes with distinct frequencies (HACKER *et al.* 1997; HACKER and CARNIEL 2001; SCHMIDT and HENSEL 2004).

The functions encoded by PAIs are dependent on the environment the bacterium lives in. Thus the genetic repertoire encompass several functional groups, with the most common ones being: (i) Iron uptake systems, including siderophores like aerobaction or yersiniabactin used to delivery of iron into microbial cells. (ii) Adhesins counting i.a. type 4 pili, P-pili, S-fimbriae and Sap adhesion. Virulence factors in this group enable bacteria to adhere to the surfaces of the host and promote the infection process in the host. (iii) Pore-forming toxins, exotoxins and enterotoxins such as α hemolysin, listeriolysin, RTX-like exotoxin and pertussis toxin. The toxins can destroy or affect the function of the eukaryotic host cells. (iv) Protein transport through Type III and IV secretion systems. These organelles deliver effector proteins into the host cells, modulating the functions of the host. (v) Invasion genes, such as the *Salmonella* spp. *inv* genes, that facilitate bacterial access to the eukaryotic host cells (SCHMIDT and HENSEL 2004; GAL-MOR and FINLAY 2006).

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For *S. enterica*, most of the virulence factors are encoded by genes carried in the chromosome, many located within PAIs. Genes needed for invasion of *S. enterica* were found to cluster within a defined region in the chromosome by the group of C. A. Lee (MILLS *et al.* 2006). This lead to the PAI named SPI-1. Thus currently 20 different types of PAIs are recorded in PAIDB for *Salmonella* $(\text{http://www.paidb.re.kr/accessed December 4th 2015).$

2.2.1 SPI-Finder

As whole genome sequencing more or less has become the new standard, a user-friendly web-tool for an easy *in silico* detection of *Salmonella* pathogenicity islands (SPIs) has been developed (**Roer II**). The *SPI-Finder* tool is publicly available as a web-based service hosted by the Center for Genomic Epidemiology (CGE) [\(https://cge.cbs.dtu.dk/services/SPIFinder/\)](https://cge.cbs.dtu.dk/services/SPIFinder/). *SPI-Finder* is built on a BLAST-based methodology for detection of genes from a customized database containing SPI variants.

There have been various initiatives for constructing databases concerning the virulence potential of bacteria, including the database 'Virulence Factors of Pathogenic Bacteria' (VFDB), a comprehensive database which comprises virulence factors form a broad range of bacterial species of medical importance (CHEN *et al.* 2005). However, a drawback with VFDB is the numerous genes present in the database that encodes hypothetical and unknown protein functions. Furthermore, the database only includes 5 of the known *Salmonella* pathogenicity islands. Another initiative is the PAthogenicity Island DataBase (PAIDB) constructed by Yoon *et al.* An initiative with the purpose of providing a database with comprehensive information on both known and potential pathogenicity island regions in prokaryotic genomes (YOON *et al.* 2007). The updated PAIDB version 2.0 contains a total of 223 types of pathogenicity islands, thus the *Salmonella* records have served as inspiration for the customized database behind the *SPI-Finder* (YOON *et al.* 2015). For some of the islands recorded in PAIDB, multiple copies with 100% identity were represented from different host origins. However, in those cases only one representative was included in the *SPI-Finder* database. Different variants within one island were denoted based on their host origin. The SPI database was constructed in FASTA format, with a descriptive header and nucleotide sequence for each record, thus the database can be updated easily by adding new island variants to the file. Another advantage of using FASTA is the uniform format that makes it easy to share.

The BLAST-based program used in *SPI-Finder* was originally developed by Zankari *et al.* for the web-tool *ResFinder* capable of detecting acquired resistance genes from WGS-data. The program

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was developed for detection in both pre-assembled genomes and data of raw reads, with a userselection of minimum percent identity (%ID) between 80 and 100% (ZANKARI *et al.* 2012).

Figure 6 | *SPI-Finder* **output for** *Salmonella* **Choleraesuis str. SC-B67.** For verification of the database, the output for *Salmonella* Choleraesuis str. SC-B67 was correlated with the seven known records in the SPI database for the isolate. Besides the seven records that originated from S. Choleraesuis str. SC-B67, *SPI-Finder* also detected SPI-13 and SPI-14 variants which were first identified in different *S.* Gallinarum isolates.

Subsequently, the original program was updated to only report the best hit within a given position of the data (with allowance of 30% overlap), and to report multiple variants of the same gene in different positions. Further, a second user-selection parameter was added, allowing for selection of a minimum length between 100 and 20%, and the selection interval for %ID was updated to go as low as 30% ID. This updated version of the program was applied to the *SPI-Finder*, allowing only the best hits from the SPI database, which also meet the user-specified parameters, to be reported in the output. The *SPI-Finder* is constructed with an easily understandable output, reporting the

pathogenicity islands detected, the origin of the islands, the %ID of the hit, the length of the hit and database record, the contig followed by the position of the hit in the contig, the insertion location from which the island was originally found, and additionally a function category from which the function of the islands can be interpreted in the category function table located beneath the output table. An output for *S.* Choleraesuis SC-B67 is shown in [Figure 6.](#page-33-0) The hits in the output are colour indicated according to the two parameters; length and identity. Dark green indicates hits of full length and with an identity of 100%, where light green indicates hits with full length but with less than 100% identity. Additionally, results indicated with grey represents results of less than 100% identity that are not of full length (not represented in [Figure 6\)](#page-33-0).

In order to examine the performance and stability of *SPI-Finder*, the two WGS isolates with the most records in the SPI database (*S.* Choleraesuis str. SC-B67 with seven records and *S.* Typhi CT18 with 10 records), were tested against the program. For *S.* Choleraesuis SC-B67 [\(Figure 6\)](#page-33-0), all seven records were found on a 100% ID-level, and in full-length, five additional islands were detected. For S. Typhi CT18 all 10 records were detected with the addition of one island. Thus in both cases, the *SPI-Finder* was capable of detecting all records in the SPI database from the two isolates.

As mentioned, *SPI-Finder* is not the first tool exploring pathogenicity islands; the PAIDB initiative comprises a PAI Finder predicting PAI-like regions. However, a drawback with the PAI Finder is that the user cannot upload WGS data directly to the tool, as it works on open reading frames (ORF). Additionally, the header within the searched file needs to follow a strict pattern: ">ID(integer)|ORF_name|start..end|strand (+ or -)", and the number of query sequences are limited to 1,000. Thus Genebank files of seven different diverse prokaryotic species shows that the number of ORFs are between 1700 and 5566 for a genome, which were also predicted by different gene prediction tools (HYATT *et al.* 2010), indicating that it is impossible to search an entire genome at a time by using PAI Finder.

The attempt with *SPI-Finder* was to design a simple, user-friendly tool to interpret *Salmonella* pathogenicity islands from WGS data. However, different aspects should be considered when using such tool. Notably, searches based on databases are only capable of detecting and reporting records if they are present in the database explored. Consequently, when inquiring a database tool like *SPI-Finder*, the user is to some extend obligated to become acquainted with the content in the database employed. Additionally, it is important to bear in mind that *SPI-Finder* solely determines the

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presence of genes in the genome and does not access any information concerning the expression. *SPI-Finder* provides an assessment of pathogenicity islands in *Salmonella*, but no information is provided about the actual level of pathogenicity.

Another consideration when working with WGS data is the quality of the sequence to be explored, thus identifying the ideal level of %ID to be capable of detecting the presence of genes. In the original program constructed for *ResFinder*, Zankari *et al.* reported an optimal %ID at 98% to circumvent noise signals (ZANKARI *et al.* 2012). However the initial study was performed on Illumina sequencing data which, compared to other technologies, contain less indel errors. For sequence data of lower quality it could be an advantage to decrease the %ID to be capable of detecting genes. As a default *SPI-Finder* do not report hits with an ID of less than 95%. In addition, the minimum length should be considered in comparison with data quality. By default, *SPI-Finder* only report hits that cover at least 60% of the gene length. However, poor genome assembly can result in a false-negative detection of a gene, or the sequence of a present gene might be split on multiple contigs, making it difficult to detect the gene with such BLAST-based tool. Thus, quality control and measures of the input data should be provided for the user to be able to have a critical view on the results. Currently, methods that map reads to databases are tested as an alternative to the BLAST-based method, to circumvent the problems with bad quality assemblies and genes split into multiple contigs.

2.3 The Mechanisms of Horizontal Genetic Transfer

Vertical transfer describes the process from which genetic material is inherent from parental cells to offspring which happens naturally during reproduction. Opposite the vertical transfer process, HGT describes the movement of genetic material between and within different populations of prokaryotes in the community (FRANCINO 2012).

The importance of HGT was first recognized together with the observation of heredity of multiple antibiotic resistances in pathogens in the 1940's (WATANABE 1963; DAVIES and DAVIES 2010). Subsequently, substantial genetic exchange between bacteria was revealed by disparities in base composition and biased codon usage, high gene similarity of distantly related organisms, and variation of gene content within the same species (KOONIN *et al.* 2001; HEUER and SMALLA 2007; SCHUMANN *et al.* 2008). HGT is currently accepted as a dominant driving force in rapid adaptation and evolution of bacterial genomes (BERG and KURLAND 2002; DOOLITTLE *et al.* 2003; HEUER and SMALLA 2007; SCHUMANN *et al.* 2008).

Figure 7 | The three general transfer modes of Horizontal Gene Transfer; Conjugation, Transformation and Transduction. The different mechanisms are explained in the text. [Figure 7](#page-36-0) is adapted from (STEWART 2013).

Several studies have elucidated the extent of HGT in bacterial communities. In a clinical isolate of the gram-positive bacteria *Enterococcus faecalis* more than 25% of the genome potentially consisted of mobile DNA (PAULSEN *et al.* 2003). As previously mentioned, in the case with the non-pathogenic *E. coli* K-12 strain and the pathogenic O157:H7 strain a difference around 30% was observed (PERNA *et al.* 2001), indicating a big intra-species divergence.

The horizontal transfer of genes is mediated by three general mechanisms; conjugation, transformation and transduction, as illustrated in [Figure 7.](#page-36-0)

The individual contribution to HGT is not known for the three transfer mechanisms. However, the transfer of plasmid mediated by conjugation is believed to be one of the most frequent and disseminated mechanisms for transfer of genetic material among bacterial species (JAIN *et al.* 2002; THOMAS and NIELSEN 2005; SCHUMANN *et al.* 2008). This PhD thesis is focusing on conjugational transfer, the mating process between bacteria. However, transduction and transformation will shortly be described in the sections 2.3.2 and 2.3.3.

2.3.1 Conjugation

The conjugational process is highly specific, where the transfer of DNA is specialized by a multiprotein complex, the conjugation apparatus (GROHMANN *et al.* 2003). The active participation of both mating partners, donor and recipient, was already suggested in 1968 by Curtiss *et al*. (CURTISS *et al.* 1968). A physical contact between the cell surfaces of the mating pair is an

important prerequisite for conjugational transfer, which in gram-negative bacteria is mediated by extracellular filaments termed sex pili (GROHMANN *et al.* 2003).

The type IV secretion systems (T4SS) are responsible for the conjugative transfer of plasmids. One of the best studied conjugative transfer systems is the IncP (*tra*) system of RP4, a broad-host range IncP plasmid, of which the most essential mechanism will be described in this section.

The IncP transfer system is comprised in the two regions; Tra1 and Tra2. For DNA to travel across the cell envelopes of the mating cells, two main protein complexes are involved; the multiprotein-DNA complex relaxosome and a mating-pair formation (mpf) complex. The relaxosome complex is formed from genes encoded by both chromosome and plasmid (FÜRSTE *et al.* 1989; LANKA and WILKINS 1995). The mpf complex is encoded in the plasmid *trb* operon (LESSL *et al.* 1993) as illustrated at the IncP plasmid in [Figure 4.](#page-27-0)

Figure 8 | Overview of general mechanisms for plasmid transfer. Conjugative plasmids are carrying an origin of transfer (*oriT*), genes encoding proteins for pili, mating-pair formation complex (mpf) responsible for the T4 envelope spanning channel, a T4CP (yellow), the relaxsosome conjugation initiation complex with the secretion substrate and DNA nicking relaxase (pink). The different steps are described in the text. [Figure 8](#page-37-0) is adapted from (ZECHNER *et al.* 2012)

The conjugational transfer is illustrated step-by-step in [Figure 8,](#page-37-0) and described in the following.

(i) The conjugation is initiated by the assembly of multiple proteins forming the relaxosome

complex at the origin of transfer (*oriT*). (ii) The complex is then preparing the single-stranded DNA intended for transfer (T-strand). This is done by nicking activity of the phosphodiester bond at the *nic* site within *oriT*, which in addition, mediates formation of a covalent tyrosinyl-DNA adduct (ZECHNER *et al.* 2012). (iii) The plasmid-encoded TraG-like T4CP (coupling protein) recognizes the nucleoprotein and docks it to the complex (GRAHN *et al.* 2000). (iv) When the initiation signal is registered, as a result of donor-recipient contact, the T-strand with linked relaxase is actively pumped through the mfp transport apparatus. (v) After the termination of transfer, the original plasmid in the donor is stabilized by conjugative replication, where the missing complementary strand is synthesized. (vi) In the recipient, recircularization of the plasmid is done by joining the free 3'-hydroxyl and the 5'-terminus with the covalently bound relaxase. Finally, the transferred plasmid is stabilized by freeing the relaxase and synthesizing the complementary strand. (GROHMANN *et al.* 2003; ZECHNER *et al.* 2012). The transfer genes at IncP plasmids are not constitutively expressed, but is regulated by local auto-regulators and global regulators, which results in a coordinated expression with other plasmid functions (ZATYKA and THOMAS 1998; DANG *et al.* 1999).

The Type IV secretion system (T4SS) involved in conjugational transfer is capable of connecting a wide variety of organisms (GRAHN *et al.* 2000; THOMAS and NIELSEN 2005), and is thereby one of the most significant machineries in bacterial adaptation and evolution (AMINOV 2011). Additionally, conjugative plasmids facilitate the spread of antibiotic resistance between pathogens, and continue to be a crucial player in the emergence of multidrug resistant pathogens in hospitals (BEOVIĆ 2006) and agriculture (ZHU *et al.* 2013).

2.3.1.1 Methods for Detection of Conjugational Plasmid Transfer

Conjugation abilities of bacterial species and transfer range of plasmids have conventionally been evaluated by using single isolates as recipients (LEDERBERG *et al.* 1952). The most common and still widely used method for detection of plasmid transfer is based on selective plating, where plasmid encoded traits act as markers for conjugational events. This requires for the plasmid to confer resistance to antibiotics or heavy metals, or to possess accessory metabolic pathways, which allow only the transconjugants to grow on the selective media. Nevertheless, the method has allowed for identification of several factors affecting conjugational transfer, including the effect of chemical compounds like biocides in transfer of the transposon Tn*916* (SEIER-PETERSEN *et al.* 2014), conjugational inhibition by unsaturated fatty acids (FERNANDEZ-LOPEZ 2005), the

availability of nutrients (SØRENSEN and JENSEN 1998), changes in temperature and pH values (RICHAUME *et al.* 1989; ROCHELLE *et al.* 1989).

The selective detection method does have other drawbacks than the selective requirements of the plasmids, especially when studying a complex microbial community. The method is restricted to the small fraction of bacteria which is culturable and able to grown on specific growth media; a fraction that probably is below 1% of the total bacterial community (AMANN *et al.* 1995). In addition, this method ignores the fact that most bacteria exist in complex matrices and together in communities with hundreds to thousands of other species, when only focusing on a single recipient strain (HONG *et al.* 2006; BROWN KAV *et al.* 2012).

To overcome some of these drawbacks, *in-situ* reporter genes were introduced on plasmids. The detection of plasmid transfer could then be measured as the expression of the reporter genes in the transconjugants, thus avoiding cultivation with selective plating and the enclosed selective advantage. Different reporter gene systems have been utilized for detecting plasmid transfer, including the *lux* system (HOFFMANN *et al.* 1998), the β-galactosidase assay with the *lacZ* gene (JAENECKE *et al.* 1996), and systems with reporter genes expressing fluorescent proteins (CHRISTENSEN *et al.* 1996; DAHLBERG *et al.* 1998).

To avoid expression of the reporter genes in the donor strain, inducible reporter genes were introduced on the conjugative plasmids. This was achived by introducing the reporter genes behind a *lacZ* promoter subjected to inhibition by constitutive *lacI* repression in the donor (either by insertion of *lacI* on the chromosome or present on a non-transferable plasmid) (DAHLBERG *et al.* 1998; FERNANDEZ-LOPEZ 2005). Thus, expression of the reporter genes would only occur after successful transfer to the recipients. The construction of the inducible *lux* system, measuring plasmid transfer as arbitrary light units (ALU), gave rise to a high-throughput conjugational assay, allowing up to 96 samples at the same time by using microplate luminometers for detection (FERNANDEZ-LOPEZ 2005; PÉREZ-MENDOZA and DE LA CRUZ 2009).

The inducible fluorescent system with *gfp* as reporter gene has been widely used to quantify plasmid transfer using flow cytometers, where the transfer was measured as expression of GFP in the recipients (CHRISTENSEN *et al.* 1996; DAHLBERG *et al.* 1998; SØRENSEN *et al.* 2003). The enhanced ability to study plasmid transfer in microbial communities revealed transfer frequencies up to 1,000 times higher than observed with methods dependent on cultivation (MUSOVIC *et al.* 2006). In 2000, the *gfp* reporter system was further improved by implementing a red-fluorescent

marker gene (*DsRed*) at the donor chromosome. This enabled simultaneously detection and quantification of the donors, recipients and transconjugants (TOLKER-NIELSEN *et al.* 2000).

The improved *gfp* reporter system is still available, thus it provides the opportunity to screen a broad collection of strains or even communities for their ability to conjugate and take up a plasmid. With isolation of single transconjugants by fluorescent activated cell sorting (FACS), and whole genome amplification it is possible to determine the range of recipients in a community, without the need of selection and cultivation (MUSOVIC *et al.* 2006; KLÜMPER *et al.* 2014; SHINTANI *et al.* 2014).

The fluorescent properties of the system have been improved by using the advanced *gfpmut3* variant for the green fluorescent reporter and *mCherry* as the red fluorescent donor marker gene, in the construct of i.a. *E. coli* MG1655::*mCherry-lacI^q* with the plasmid construct pKJK5::*gfp3mut* (KLÜMPER *et al.* 2014)*.*

This latter *E. coli* pKJK5 system, constructed by Klümper *et al.,* was applied in **Roer III**, as a highthroughput screening method, to identify conjugative abilities in a broad collection of *S. enterica* ssp. *enterica*.

Figure 9 | **Quintuple gating for donor, recipient and transconjugant detection by flow cytometry.** Flow cytometry detection of donor, recipients, and transconjugants from a conjugation mixture of a *S. enterica* recipient and *E. coli K-12* MG1655 donor carrying pKJK5. The gating procedure consists of five successive gates in the 3 bivariate plots: Plot I with the Bacterial gate, detects all particles of bacterial size based on front scatter (FSC) and side scatter (SSC); Plot II detects all red fluorescence particles (*mCherry*) representing the donors, and all non-red particles representing either recipients or transconjugants by using side scatter and PE-Texas Red; Plot III divides the recipients and transconjugants by gating non-green and green fluorescence particles (*gfp3mut*) based on side scatter and FITC. [Figure 9](#page-40-0) adapted from (**Roer III)**.

In **Roer III**, the detection of donor, recipients and transconjugants was based on a quintuple gating approach. The five gates are defined in the three bivariate plots in [Figure 9.](#page-40-0) In the first plot, a

combination of side scatter (SSC) and forward scatter (FSC) is used to distinguish bacterial particles from other subpopulations, by using a gate corresponding to particles of bacterial size. In the red fluorescence (PE-Texas Red) vs SSC plot a gate was set to cover all donor cells, with an additional non-red gate covering recipients and transconjugants. In the third plot, green fluorescence (FITC) and SSC was used to distinguish green transconjugants from non-green recipients. This approach allowed us to count 100,000 bacterial events in around 30 seconds, without the need to use selective cultivation.

2.3.2 Transduction

Transduction is another of the three transfer mechanisms comprised by the term HGT. The transfer of non-viral DNA to a new host is mediated by the bacterial viruses, bacteriophages (phage), as illustrated in the upper right corner of [Figure 7](#page-36-0) (STEWART 2013). The mechanism relies on mistakenly packed phage heads where host DNA is integrated in the phage DNA during the reproduction of the phage particle. When host cells are lysed, defective phage particles are released, ready to adsorb to new host cells, where they can inject the carried DNA from the previous host. When delivered to the recipient host cell, the transferred DNA can be integrated into the chromosome (MAZODIER and DAVIES 1991; HEUER and SMALLA 2007). A review by Brüssow *et al.* describes different bacterial species that have acquired pathogenicity determinants through phages (BRÜSSOW *et al.* 2004). Additionally, PAIs in pathogenic strains is believed to have evolved from lysogenic bacteriophages (HACKER *et al.* 2003; DOBRINDT *et al.* 2004).

It was originally assumed that phages was limited to a small range of hosts (BERGH *et al.* 1989; MAZODIER and DAVIES 1991), however phages like P1 and Mu have shown to interconnect with a broad range of bacteria (JENSEN *et al.* 1998; WOMMACK and COLWELL 2000; CHEN and NOVICK 2009). Transfer by transduction has the advantage that DNA is protected from the environment, and the phage particles can persist relatively long under environmental conditions (ZEPH *et al.* 1988; WOMMACK and COLWELL 2000).

The number of bacteriophages in seawater is found to be 10 times higher than the number of bacteria (BRÜSSOW and HENDRIX 2002; DANOVARO *et al.* 2008), and with transduction events of 20 million billions per second in the oceans (CHIBANI-CHENNOUFI *et al.* 2004) marine viruses are believed to be a major contributor in the global ecosystem (SUTTLE 2007; DANOVARO *et al.* 2008; ZHAO *et al.* 2013). However, as the transduction mechanism relies on mistakes in phage packing,

most of the transferred DNA will not be functional genes. Thus, the evolutionary impact of transduction will probably be rather low.

2.3.3 Transformation

The third HGT mechanism is transformation [\(Figure 7\)](#page-36-0). Transformation covers the uptake of free exogenous DNA by competent cells (LORENZ and WACKERNAGEL 1994; DUBNAU 1999). Natural competence is a physiological phase, genetically programmed, where efficient uptake of DNA is permitted. This can either be a persistent phase, as for *N. meningitidis* (HAMILTON and DILLARD 2006), or a phase induced by environmental factors (HANAHAN 1983; NIELSEN and VAN ELSAS 2001). Thus, the prerequisites for transformation are the bacterial change to a competent phase, the availability of exogenous free DNA, followed by the uptake and stable integration of the DNA (HEUER and SMALLA 2007).

Transformation is a commonly used method in gene technologies, as artificially constructed changes of environments can trigger the desired bacterial cells to enter a competent phase, ready to act as recipients for uptake of free DNA. However, transformation frequencies among isolates from the same species can differ significantly, indicating that the ability to transform is not shared systematically between isolates from the same bacterial species (SIKORSKI *et al.* 2002; MAAMAR and DUBNAU 2005). Additionally, the efficiency of integrating the newly introduced DNA to the bacterial genome might vary between bacterial species (SIKORSKI *et al.* 2002).

Transformation has been confirmed in various bacterial species and environments (BRÄUTIGAM *et al.* 1997; NIELSEN and VAN ELSAS 2001; AVERHOFF and FRIEDRICH 2003; SØRENSEN *et al.* 2005; HAMILTON and DILLARD 2006), however very limited knowledge exists on the importance of transformation, the subsequent ability of the bacteria to adapt to different environments (HEUER and SMALLA 2007), as well as the contribution to bacterial evolution (JOHNSBORG *et al.* 2007).

Chapter 3 **TRANSFER DEFENCE MECHANISMS**

The evolutionary rate in bacteria is affected by both the ability to exchange DNA as well as the frequencies of transfer, thus any genetic determinant restricting or enhancing this, may influence evolution. A number of well-described mechanisms affecting conjugative transfer include plasmid incompatibility (described in section 2.1.3), entry exclusion, clustered regularly interspaced short palindromic repeats (CRISPR), and restriction-modification systems (RM systems). This PhD thesis is focusing on the barriers of RM systems, and their influence on evolution. However, entry exclusion and CRISPR will shortly be described in the sections 3.2.1 and 3.2.2.

3.1 Restriction-Modification Systems

Restriction-modification systems are transfer defence mechanisms endorsed by the bacterial hosts. Generally all bacteria mark their DNA with an identity signature, thus when DNA is transferred from one bacterium to another, DNA lacking the signature of the recipient strains will be perceived as 'foreign' DNA rather than 'self'. DNA perceived as 'foreign' is generally enzymatically cleaved in the recipient (KING and MURRAY 1994; WILLIAMS 2003; ROBERTS *et al.* 2003).

The discriminatory barrier was first demonstrated in the 1950's with an *E. coli* K-12 strain infected by bacteriophage *λ*, transmitted through either *E. coli* strain C or B (BERTANI and WEIGLE 1953). Around a decade later, it was proven that the establishment of bacteriophages in *E. coli* K-12 strains could be 'restricted' by an enzyme that attacked 'foreign' DNA (MESELSON and YUAN 1968; LINN and ARBER 1968).

Traditionally, the restriction enzyme is found together with a cognate methyltransferase, modifying the DNA by adding a methyl group in a given position. The activities from the two genes constitute the restriction-modification systems. However, there are systems only comprised by a modificationdependent restriction enzyme, which only attack DNA, when the DNA is methylated in a specific pattern (ROBERTS *et al.* 2003). Both variants share the ability to attack DNA recognized as 'foreign' and thereby prevent DNA establishment in the recipient. Thus, while the methyltransferase from a classical RM system is required to protect the cells own DNA from degradation by the cognate restriction enzyme, the modification can act as substrate for restriction activity from a modificationdependent system in another strain (MURRAY 2002). The classical RM systems have been subdivided into three groups (type I-III), based on their complexity, the requirement of cofactors, their recognition sequence, and DNA cleavage site, as illustrated in [Figure 10.](#page-44-0) The modificationdependent systems account for a fourth group (type IV). However, some RM systems do not fit readily into any of the four types (KING and MURRAY 1994).

The key characteristics for the type I RM systems are the hetero-oligomeric complexes which catalyses both restriction and modification, when the recognition sequence is recognized by the specificity subunit (S). They are usually comprised of two R subunits, two M subunits and one S subunit.

Figure 10 | Characteristics and gene organization for the three classical RM systems; Type I-III. RM systems are subdivided according to their complexity, cofactor requirement, recognition sequence and DNA cleavage site. Type I, type II and type III are the classical RM systems. Type IV Modification-dependent systems are not classical systems, as they do not possess a cognate methyltransferase, and only cleave DNA with specific modifications. ENase, endonuclease activity; MTase, methyltransferase activity. [Figure 10 i](#page-44-0)s modified from (MURRAY 2002).

The three type I subunits are encoded by the closely related genes; *hsdR*, *hsdM* and *hsdS*. The genes are transcribed from two promotors; *hsdM* and *hsdS* from the same promotor, and *hsdR* from a separate promotor (LOENEN *et al.* 1987). For methyltransferase activity, only the two subunits M and S are necessary, whereas the third subunit R is required for restriction activity. Thus, the restriction activity will not be active without the M and S subunits. Two functional complexes exists; one with only methyltransferase activity (M_2S_1) , and the RM system which comprises all three subunits $(R_2M_2S_1)$ (MURRAY 2000). In the modification reaction, *S*-adenosyl-methionine (AdoMet) act as the methyl source, whereas AdoMet, ATP, and Mg^{2+} all are required for restriction activity (BICKLE and KRÜGER 1993). In type I RM systems, the recognition sequences are asymmetric and are constructed by two components, the first of 3 or 4 bp and the second of 4 or 5 bp. The two components are separated by a non-specific spacer region of 6-8 bp. When the recognition sequence is hemimethylated the DNA will act as substrate for modification, whereas unmodified recognition sequences will trigger DNA cleavage at sites remote from the recognition sequence (BICKLE and KRÜGER 1993; MURRAY 2002).

Compared to the type I RM systems, the type II systems are less complicated, which is probably why this is also the type of restriction enzymes used in gene technology. The systems are comprised of two separate genes; *Res* encoding the restriction activity and *Mod* encoding the methyltransferase activity. As for the type I systems, the modification requires AdoMet as methyl donor, while the restriction activity for type II systems only is dependent on Mg^{2+} (BICKLE and KRÜGER 1993; ROBERTS *et al.* 2003). The restriction enzyme and the methyltransferase recognize the same recognition sequence, which generally is a palindromic sequence of 4-8 bp. The methyltransferase modify a specific base within the recognition sequence on each strand. The structure of the recognition sequence facilitates cleavage of both DNA strands at the same time, generally within or close to the recognition sequence (ROBERTS *et al.* 2003).

The type III systems are not as simple as type II systems, but are sharing some traits with the type I systems. A hetero-oligomeric complex of the *Res* and *Mod* genes catalyse both restriction and modification activities in the type III systems. As for the type I and II systems, the AdoMet is required for modification, while both Mg^{2+} and ATP are required for restriction activity in type III systems. The recognition sequences for type III systems are asymmetrical, and the cleavage site is located close to the recognition sequence (BICKLE and KRÜGER 1993; ROBERTS *et al.* 2003).

The non-classical RM systems, type IV, restrict modified DNA. However their recognition sequences have usually not been defined (ROBERTS *et al.* 2003).

For decades, the restriction-modification systems have been known to act as barriers, not only on bacteriophages, but for horizontal gene transfer as well (KING and MURRAY 1994; TOCK and DRYDEN 2005; VEIGA and PINHO 2009; VASU *et al.* 2012). Generally, the belief is that DNA lacking the signature of the recipient will act as substrate for restriction, irrespective of whether the 'foreign' DNA would enter the recipient as single- or double-stranded (MURRAY 2002). Plasmid DNA, which enters the recipient as single-stranded DNA through conjugation, will become sensitive to cleavage after stabilization from the synthesis of the complementary strand (THOMAS and NIELSEN 2005).

However, this has, to our knowledge, never been demonstrated at a genotypic level. In **Roer I**, the impact of the type I RM system *Eco*KI, in conjugational transfer, was elucidated by utilizing a combination of isogenic RM variants of the *E. coli* K-12 strain MG1655 and plasmids pOLA52 (SØRENSEN and HANSEN 2003) and pHHA45 (DOLEJSKA *et al.* 2012), respectively harboring either two or zero recognition sequences for the *Eco*KI system.

Each transfer experiment was performed at least in triplicate. The data is visualized as the means of transconjugants per 10⁸ recipients, with standard errors of the mean (SEM). For un-methylated plasmid pOLA52, with two recognition sequences for the *Eco*KI system, statistically significant differences were observed between the restriction deficient recipient MG1655-*hsdR*^{INT} and the remaining two recipients (b*, *P*0.029). No statistically significant differences were observed for the un-methylated pHHA45 plasmid without recognition sequences for *Eco*KI.

In **Roer I** we show that a *hsdR* deficient mutant caused a 7-fold increase in uptake of an unmethylated plasmid with two recognition sites. Though, most importantly we demonstrated that the barrier imposed by the *Eco*KI type I restriction system, had an effect, but that the barrier was not absolute in relation to conjugational plasmid transfer.

3.1.1 Restriction-ModificationFinder

In **Roer II**, the content of RM systems was elucidated in a comprehensive collection of *S. enterica* WGS genomes. For that purpose, another user-friendly web-tool, *Restriction-ModificationFinder* (*RM-Finder*) was developed for an easy *in silico* detection of RM systems. The *RM-Finder* was built on the same BLAST-based methodology as *SPI-Finder*, with a customized database containing restriction enzymes, methyltransferases and specificity subunits for the classical systems, as well as the modification-dependent type IV systems. *RM-Finder* is likewise available as a webbased service hosted by CGE [\(https://cge.cbs.dtu.dk//services/Restriction-ModificationFinder/\)](https://cge.cbs.dtu.dk/services/Restriction-ModificationFinder/).

For databases on RM systems, the Restriction Enzyme database (REBASE) is the only authoritative source, a dedicated work by R. Roberts and D. Macelis (ROBERTS *et al.* 2015). REBASE is a comprehensive database, currently with more than 5,000 complete descriptions of RM systems. The database is containing information about restriction enzymes and related proteins like DNA methyltransferases, specificity subunits, homing endonucleases, helicase domain proteins etc. and contains both published and unpublished references. The data in REBASE is based on genome sequence data subjected to analysis, where predicted components from RM systems are labelled with the suffix 'P' to indicate that they are putative and included in the database (ROBERTS *et al.*) 2015). When the systems become biochemically characterized, the suffix is replaced with a Roman number. Though, with the Single molecule Real Time (SMRT) sequencing, the characterization of especially the methyltransferases are rapidly increasing (FLUSBERG *et al.* 2010; KORLACH *et al.* 2010; KORLACH and TURNER 2012). Impressively, REBASE is continuously updated and their web interface refreshed on a nightly basis. Their ftp access to the data is refreshed once per month, providing data to external sources, like *RM-Finder.* However, the continuous expansion of REBASE indicates that far from all RM systems are known. This can be a problem when searching for RM systems, as it will be impossible to know if all systems are detected in the searched genomes.

Additionally, REBASE is providing various tools for investigating the determinants in RM systems, including a BLAST tool to search sequences against the REBASE database. This tool is capable of using either DNA or protein sequences as input file. However, few major drawbacks from this tool exist; the lack of ability to BLAST either raw data or assembled genomes, and absence of user-

selective parameters. The limitations from REBASE lead to the construction of *RM-Finder* (**Roer II**). REBASE is the source of data behind the database in *RM-Finder*, and includes the relevant restriction enzymes, methyltransferases, and specificity subunits for type I-IV RM systems. The RM database was constructed in the uniform and easily sharable FASTA format, containing a descriptive header, and the nucleotide sequence for each record. The database is automatically updated according to the monthly REBASE update.

The *RM-Finder* is constructed with the updated user-selection parameters, allowing for selection of minimum length and minimum %ID. The default settings for *RM-Finder* are set to minimum ID of 95%, and a minimum length of 60%. Furthermore, it is possible to search only one type of RM systems, and to search all records in the database, including putative genes.

								Type I Restriction enzymes								
Gene	%Identity	HSP/Query length		Contig	Position in contig			Type	Function		Recognition Seq			Accession number		
S.EcoKI	100.00	1395	1395/ NC 000913		45780914579485		Type	specificity subunit		AACNNNNNNGTGC			U00096.			
M.EcoKI	100.00	1590/ 1590		NC 000913		45794824581071		Type	methyltransferase		AACNNNNNNGTGC			U00096,		
EcoKI	100.00	3567/ NC_000913 3567		45812724584838			Type	restriction enzyme		AACNNNNNNGTGC			U00096.			
								Type II Restriction enzymes								
Gene		%Identity	HSP/Query length		Contig			Position in contig		Function Type		Recognition Seq		Accession number		
M.Eco3609Dcm		99.22	1419/ 1419		NC_000913		20289232030341		Type \mathbf{H}	methyltransferase			CCWGG JASV01000004.			
M.EcoKII		100.00	885/ 885	NC 000913		3409675.3410559			Type Ш	methyltransferase			ATGCAT	U00096,		
						Type III Restriction enzymes										
	No restriction enzyme genes found															
						Type IV Restriction enzymes										
	Gene		%Identity	HSP/Query length	Contig			Position in contig		Type	Function		Recognition Seq	Accession number		
EcoKMcrA			100.00	834/ 834	NC_000913		12095691210402			Type IV.	methyl- directed restriction enzyme		YCGR	U00096.		
EcoKMcrBC(EcoKMcrC)			100.00	10471 1047	NC_000913		45749354575981			Type IV	methyl- directed restriction enzyme			U00096,		
EcoKMcrBC(EcoKMcrB)			100.00	1380/ 1380	NC 000913		45759814577360			Type IV	methyl- directed restriction enzyme			U00096.		
EcoKMrr			100.00	915/ 915	NC 000913	45849724585886			Type IV.		methyl- directed restriction enzyme			U00096.		
						extended output										
Results as text			Hit in genome sequences		Restriction enzyme gene sequences											
Results tab separated	Selected %ID threshold: 95.00 %															

Figure 12 | *RM-Finder* **output for** *Escherichia coli* **K-12 strain MG1655.** For demonstration of *RM-Finder*, the complete genome of *E. coli* K-12 strain MG1655 was evaluated.

[Figure 12](#page-48-0) illustrates the output constructed by *RM-Finder*, reporting hits of each of the four systems in an individual section. The output reports the RMS genes detected, the %ID of the hit, the length of the hit compared to the database record, the contig in which the hit was found followed by the position in the contig. Additionally, the type of system, the function of the gene, the recognition sequence (if known), and the accession number are reported in the output. As for *SPI-Finder,* the hits are indicated by colour according to the two user-selection parameters; %ID and minimum length.

To evaluate the performance of *RM-Finder*, the *E. coli* K-12 strain MG1655 was evaluated, and compared with the putative predictions from REBASE [\(http://rebase.neb.com/cgi-](http://rebase.neb.com/cgi-bin/onumget?17068)

[bin/onumget?17068\)](http://rebase.neb.com/cgi-bin/onumget?17068). It can be argued that the MG1655 strain is not the most complicated control strain. However, if *RM-Finder* is not capable of detecting all the known RM systems in a wellcharacterized strain, then the results predicted for uncharacterized strains will be unreliable.

Table 1 | Comparison of REBASE and *RM-Finder*

Predicted by REBASE	Recognition Sequence	Predicted by RM-Finder
M.Eco1655DamP	GATC	Found by including putative
M.Eco1655DcmP	CCWGG	M.Eco3609Dcm
Eco1655McrBP	unknown	EcoKMcrB
Eco1655McrCP	unknown	EcoKMcrC
Eco1655MrrP	unknown	EcoMrr
Eco1655ORF300P	AACNNNNNNGTGC	EcoKI
M.Eco1655ORF300P	AACNNNNNNGTGC	M.EcoKI
S.Eco1655ORF300P	AACNNNNNNGTGC	S.EcoKI
M.Eco1655ORF6040P	ATGCAT	M.EcoKII
Eco1655ORF16910P	YCGR	EcoKMcrA

By only searching the biochemically characterized systems in *RM-Finder*, 90% of the genes were identified in concordance with REBASE. By inspecting the predictions from REBASE M.Eco1655DamP was predicted by the putative M.UbaC1152DamP. When searching the RM-Finder, including putative variants, the M.Eco1655DamP enzyme is predicted from *M.Eco321DcmP* with 100% ID and 100% length.

When using the *RM-Finder tool*, the same precaution should be taken as mentioned for *SPI-Finder*. The WGS data quality should be considered, and even though the data included in *RM-Finder* is comprehensive, database searches can only find what is included in the database.

3.2 Other Defence Mechanisms

In addition to RM systems, other defence mechanisms exist. Some can be encoded or caused by the transferred plasmid to prevent further uptake of MGEs, as for incapability and plasmid entry exclusion, others can, as for RM systems and CRISPR, be encoded on the bacterial chromosome to protect the bacteria. In the following sections, CRISPR and entry exclusion will briefly be introduced.

3.2.1 Host CRISPR Systems

CRISPR is referred to as the bacterial adaptive immune systems, as it has the benefit of continuously adjusting and adapting its reach and defence mechanisms against invasion (GOREN *et al.* 2012). According to the CRISPR database approximately 45% (1,176 out of 2,612) of the bacterial genomes possess CRISPRs (*http://crispr.u-psud.fr/crispr/* Accessed December 22nd, 2015).

The CRISPR systems are combined by two different mechanisms; Immunization of 'foreign' incoming DNA like plasmids or bacteriophages, and secondly subsequent immunity to the same DNA. In the immunization process, 'foreign' plasmid or virus DNA is recognized by a Cas complex, and cleaved into novel spacer unit. The novel spacer unit is inserted at the leader end (5' end) of the CRISPR locus, which induces acquired immunity against subsequent invasion (VAN DER OOST *et al.* 2009; HORVATH and BARRANGOU 2010). As the new spacer units generally are inserted at the leader end of the CRISPR locus, a chronological record of the encountered DNA exists. However, swapping and loos of spacers occur (LILLESTØL *et al.* 2006).

For the immunity process, the repeat-spacer array constructed from the immunization is transcribed into pre-crRNA, which is further processed into mature crRNAs. Subsequently, the crRNAs are guiding a Cas complex to the invading DNA, where inactivation is conducted (HORVATH and BARRANGOU 2010).

CRISPRs are defence mechanisms only recently discovered, however they are now applied to molecular biology research, enabling targeted genome editing (CONG *et al.* 2013; DOUDNA and CHARPENTIER 2014).

3.2.2 Entry Exclusion

Entry exclusion facilitated by plasmids can turn the bacteria into poor recipients to avoid addition of extra genetic elements (GARCILLÁN-BARCIA and DE LA CRUZ 2008). This ability can provide an

evolutionary advantage for the plasmid, as intra-cell competition with other conjugative plasmids or MGEs can be minimized or avoided (THOMAS and NIELSEN 2005).

The exclusion mechanism was shown to operate at both the inner- and outer recipient membrane, and be caused by the two plasmid-encoded genes, *traT* and *traS* (ACHTMAN *et al.* 1977). The protein of *traT* operates at the outer membrane, where modifications of the recipient cell membrane prevent the adhesion of transfer pili encoded by additional plasmids. This results in reduced abilities for the recipient to form mating aggregates with new donor cells (THOMAS and NIELSEN 2005).

As described in section 2.3.1, the initiation of conjugational transfer is dependent on a signal. This signal is proposed to be induced by TraG. After translocation of TraG from the donor to the recipient cell, contact with the inner membrane of the recipient initiates conjugation (AUDETTE *et al.* 2007). The plasmid-encoded *traS* can interfere with this process, thus blocking additional DNA uptake in the recipient (GARCILLÁN-BARCIA and DE LA CRUZ 2008).

The barrier of entry exclusion was shown not to be absolute, and uptake of new plasmids in a host was not completely excluded. However, a decrease of more than 500-fold caused by entry exclusion was observed for conjugation (PÉREZ-MENDOZA and DE LA CRUZ 2009)

3.3 Host Defence and Evolution

Defence mechanisms like RM systems and CRISPR are predicted to have an important influence on the uptake and establishment of DNA. It is therefore reasonable to think that their function can be reflected in the evolution of bacteria.

In was suggested by Budroni *et al.* that the evolution of bacteria was associated with the genomic content of RM systems (BUDRONI *et al.* 2011). The *Neisseria* genus serves as a paradigm for natural transformation, where exchange of genetic material is generally due to the persistent competence in the organism. Budroni *et al.* investigated 20 *Neisseria meningitidis* genomes, covering five serogroups of the recorded 13 for the species (ROSENSTEIN *et al.* 2001; CENTERS FOR DISEASE CONTROL AND PREVENTION 2012). They found that, for the phylogenetic network constructed based on the core genes; genomes from the same clonal complex were forming distinct phylogenetic clades. Further, they provided evidence that the phylogeny, thus evolution, in *N. meningitidis* could be associated with the RM systems identified in the genomes (BUDRONI *et al.* 2011).

In addition to the study by Budroni *et al*., the connection between RM systems and evolution has been elucidated for the organism *S. enterica* ssp. *enterica* (**Roer II**), which is known to generally exchange genetic material through conjugation (FERGUSON *et al.* 2002). The association was investigated for 221 *S. enterica* genomes, covering 97 *S. enterica* ssp. *enterica* serovars, where the evolution was depicted as a phylogenetic network based on a core-genome three [\(Figure 2\)](#page-21-0) and a pan-genome three [\(Figure 3\)](#page-22-0). Assessing the content of RM systems in correlation to the two trees, we did not observe significant linkage, though we observed sub-linage correlation and serovar specific patterns in cases with e.g. *S.* Enteritidis and *S.* Typhymurium (**Roer II**). Additionally, we observed that plasmid replicons, SPIs, and AMR showed a better correlation to serovars than to RM systems. Thus, the study presented in **Roer II** suggests a limited influence of RM systems on the evolution of the conjugative organism *Salmonella enterica* ssp. *enterica* (**Roer II**), which correlates very well with the findings of RM systems not producing an absolute barrier for conjugation (**Roer I**).

In a recent study by Fricke *et al.* the evolutionary influence of CRISPR was investigated in *S. enterica* (FRICKE *et al.* 2011). From the 28 *S. enterica* genomes included in the study, the phylogeny was only partially reflected in the composition of the CRISPR arrays. However matches between CRISPR spacers and the content of plasmids and prophages could indicate a potential for CRISPR to be involved in sub-linage evolution (FRICKE *et al.* 2011).

Even though, barriers like RM systems seem to have an influence on the evolution of transformable organisms (BUDRONI *et al.* 2011), the indefinite barrier of RM systems in conjugation (**Roer I),** the lack of association between evolution and RM systems (**Roer II),** and the only partial association between CRISPR and evolution (FRICKE *et al.* 2011), implies that for conjugal organisms, other factors might be involved in shaping the evolution and importantly the spread of genetic traits like AMR.

Chapter 4 **OTHER RECIPIENT DETERMINANTS AFFECTING CONJUGATION**

Numerous studies have elucidated on environmental factors or exogenous components affecting conjugations (RICHAUME *et al.* 1989; ROCHELLE *et al.* 1989; FERNANDEZ-LOPEZ 2005; SEIER-PETERSEN *et al.* 2014). However, only few studies have investigated the contribution and ability of bacteria to act as recipients.

In 1974 Skurray *et al.* showed that some *E. coli* K-12 mutants were defective in conjugation, due to the lack of a major outer membrane protein (SKURRAY *et al.* 1974). The missing protein in the recipient precluded mating pair formation. Thus, the defective recipient inspired to the term "Conmutant" (conjugation negative mutant). Further, evidence was provided, that different recipient cell surface components like LPS and the outer membrane protein OmpA were involved in Conrecipients for conjugation (HAVEKES *et al.* 1977; MANOIL and ROSENBUSCH 1982). However, mutants generated at that time, especially cell envelope mutants, were generated by random mutagens like ethylmethanesulfonate (EMS), which made it difficult to identify the exact site of mutation (HAVEKES *et al.* 1977; MANOIL and ROSENBUSCH 1982).

In a recent study, Pérez-Mendoza and de la Cruz tried to identify recipient genes in *E. coli* affecting the ability in plasmid conjugation, by using a more systematic approach (PÉREZ-MENDOZA and DE LA CRUZ 2009). By using the inducible *lux* system for conjugational detection (described in 2.3.1.1), they were capable of performing high-throughput screening in the attempt to identify recipient factors involved in conjugation. The *E. coli* mutant collection, the Keio collection (BABA

et al. 2006), enabled systematic screening of more than 99% of the non-essential genes from the *E. coli* K-12 genome. However, none of the 3,908 recipients tested, showed a dramatic drop in conjugation frequency compared to the wild-type *E. coli* K-12 strain BW25113. As a second attempt to identify recipient genes involved in conjugation, Pérez-Mendoza and de la Cruz constructed a transposon insertion mutant library of the DH5α *E. coli* strain. From this, they confirmed the previous findings of LPS being involved in conjugational transfer (PÉREZ-MENDOZA and DE LA CRUZ 2009). Therefore, they concluded that recipient bacterial cells cannot evade from being used as recipients in conjugation. However, their study was only based on two *E. coli* strains, both with wild-types showing reasonable conjugation abilities. Thus, these two isolates could potentially lack the restricting or enhancing abilities for conjugation.

As described in Chapter 3 with transfer defence mechanisms, recipient systems like RM and CRISPR do exist. However, RM systems was both shown not to be an absolute barrier in conjugation between isogenic *E. coli* strains (**Roer I**), and not to shape the evolution of *S. enterica* ssp. *enterica* (**Roer II)**. Thus, in **Roer III**, we aimed to identify other recipient determinants either enhancing or restricting plasmid uptake in *S. enterica* ssp. *enterica*. To overcome the drawback of not knowing if the isolates investigated, potentially could possess determinants either restricting or enhancing conjugation, the study was initiated with identifying the recipient potential of all isolates included. 93 isolates, covering 54 serovars were included in the study, to provide a broad genetic pool for investigation. From the initial conjugation, the isolates were subdivided into two groups of good (10 isolates) and poor (83 isolates) recipients. However, the comparative bioinformatic approaches applied could not yet identify gene candidates for neither enhancing nor restricting plasmid uptake. Thus other approaches should be considered in the identification of genes in *Salmonella* subspecies *enterica* that might control conjugation (**Roer III**). A parallel study on *S.* Enteritidis identified 33 gene candidates potentially restricting conjugation, which are being investigated and awaiting verification (**Roer III**).

Though, little is still known about recipient determinants affecting conjugational transfer. So far, membrane proteins have demonstrated to be involved in conjugation (SKURRAY *et al.* 1974; HAVEKES *et al.* 1977; MANOIL and ROSENBUSCH 1982). The findings in **Roer III**, with transfer frequencies from the same donor construct, varying from 0.0 to 2.5×10^{-1} transconjugants per potential recipients indicates that recipient determinants must exist (**Roer III**). This contradicts the conclusion by Pérez-Mendoza and de la Cruz, which states that in conjugation, bacteria cannot avoid being used as recipients (PÉREZ-MENDOZA and DE LA CRUZ 2009).

Chapter 5 **CONCLUSION AND FUTURE PERSPECTIVES**

For decades, it has been a dogma that RM systems act as a barrier for horizontal gene transfer. This has previously been demonstrated for transfer of DNA by transduction and transformation. The studies within this PhD thesis have illustrated that, the barriers imposed by the RM system *Eco*KI, are not absolute in conjugational transfer of an IncP-1 plasmid (**Roer I**). However, an inactivation of the restriction enzyme could cause a 7-fold increase in plasmid uptake. As the barriers of RM systems was demonstrated to be indefinite, it was interesting to investigate if the observed reduction in transfer after all could be reflected in the evolution of an organisms using conjugation as the preferred method in DNA exchange (**Roer II**). By developing and using the easily interpreted and user-friendly tools, *RM-Finder* and *SPI-Finder*, the PhD study demonstrated that the evolution of the conjugative organisms *S. enterica* ssp. *enterica*, was not associated with RM systems, and that, plasmid replicons and SPIs could not be reflected in the content of RM systems either. This led to the thought of other recipient determinants being involved in the control of plasmid uptake and in the evolution of bacteria (**Roer III**). By combining results obtained in the laboratory with bioinformatic approaches, we were able to predict 33 gene candidates restricting plasmid uptake in *S.* Enteritidis, which are being investigated and awaiting verification. However, we were not yet able to detect common genes on species level that potentially could control conjugation.

There is still an enormous lack of knowledge regarding the recipient determinants that control the uptake and establishment of plasmids. Thus, the global emergence and rapid spread of undesirable genetic traits are of significant importance, with antimicrobial resistance being an obvious example of such traits. The usage of antimicrobials in treatment of both human and animals has created a reservoir of resistant organisms and a pool of resistance genes available for further transmission.

One of the most striking and current examples is the rent observations of the plasmid-mediated *mcr-1* gene, which is conferring resistance towards colistin. A resistance which previously was mediated by chromosomal mutations with subsequent vertical heredity to daughter cells (HASMAN *et al.* 2015; LIU *et al.* 2015). However, the plasmid-mediated gene now enables rapid horizontal spread between bacteria. In Denmark, the gene was detected in an *E. coli* isolate only susceptible to a narrow range of antimicrobial classes (HASMAN *et al.* 2015). Thus, acquisition of additional resistances will leave a limited number of (if any) suitable options for treatment of bacterial infections. This emphasizes the importance of understanding the mechanisms and genes involved in controlling the uptake and maintenance of plasmids. Through improved knowledge we can hopefully predict and control the conjugational spread of generic threats in the future.

When elucidating on conjugational transfer, a big effort has been made on explaining the conjugative apparatus encoded on plasmids, and environmental settings affecting the transfer. However, little is known about the ability for bacteria to act as recipients. Different membrane proteins and barriers in the bacteria have been proposed to be involved in regulation of horizontal gene transfer. With new knowledge of RM systems not providing an absolute barrier, it would be interesting to elucidate on the mechanisms which allow some plasmids to evade the digestion of the restriction enzymes. Since most restriction enzymes works on double stranded DNA, one theory could be that the methyltransferase methylates some of the plasmids as the bacterial 'self', simultaneously with the synthesis of the stabilizing complementary strand right after transfer to the recipient, thus protecting the DNA from the restriction enzyme. This theory should be investigated to further improve the knowledge of the effect of RM systems in conjugation.

The development of benchtop sequencing technologies in the recent years has made whole-genome sequencing applicable in modern research, which also enabled a comprehensive analysis of a diverse dataset comprising 221 genomes in **Roer II**. With the continuous decrease in both purchasing and sequencing cost and the increased knowledge within this field, sequencing will most likely be standard in most research laboratories in the future. However, to enable practical usage of

the new technology, the knowledge obtained in the laboratory needs to be converted into bioinformatic tools like the *RM-Finder* and *SPI-Finder*. Currently, tools to predict plasmid replicons, antimicrobial resistance genes, virulence factors, serotypes etc. are developed (ZANKARI *et al.* 2012; CARATTOLI *et al.* 2014; JOENSEN *et al.* 2014, 2015), and over the next years we will probably see a great increase in the development of bioinformatic tools and usage. However, everything cannot be predicted by *in-silico* analysis alone, to improve our knowledge of mechanisms and bacterial behaviour we still need to do research in the laboratory. But the technology definitely has the potential in diagnostics and routine surveillance settings, where bacteria are screen for known characteristics.

Combining molecular technology and advanced bioinformatics is thus a new strategy in contemporary research, an approach pursued in **Roer III**. However, for now we were not capable of linking genes to neither enhanced or restricted conjugational abilities on species level, but a comprehensive investigation and characterization of the 93 isolates included in the study could potentially reveal a connection between isolates within the two groups of recipients. Another interesting analysis could be a SNP comparison of the genomes, thus unique SNPs within genes could explain a functional inactivation of proteins involved in conjugation. If the combination with bioinformatics cannot predict or answer why differences are observed in conjugational abilities, alternative approaches would be to turn the focus back to functional analysis in the laboratory. One such approach could be to screen the bacterial genes by performing random gene-knockouts of the genes in the recipient. However, the genes responsible for the recipient potential could as well be an essential gene for the bacteria, in which case a knock-out of the gene would be fatal for the bacteria. The knock-out construct should be accompanied by the construction of a transposon cloning library of the recipient. Different approaches combining the technologies, as well as sticking to molecular microbiology can be followed in the attempt to identify the recipient determinants that are involved in the control of conjugational uptake of plasmids.

In summary, this PhD study is improving the knowledge of RM systems and their impact on conjugation and evolution. Moreover, this PhD implies that recipient genes could be involved in controlling conjugation, and are leaves great opportunities for revealing of genes involved in plasmid uptake.

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ARTICLES

Roer I

The *Eco***KI Type I Restriction-Modification System in** *Escherichia coli* **Affects but Is Not an Absolute Barrier for Conjugation**

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The EcoKI Type I Restriction-Modification System in *Escherichia coli* **Affects but Is Not an Absolute Barrier for Conjugation**

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The rapid evolution of bacteria is crucial to their survival and is caused by exchange, transfer, and uptake of DNA, among other things. Conjugation is one of the main mechanisms by which bacteria share their DNA, and it is thought to be controlled by varied bacterial immune systems. Contradictory results about restriction-modification systems based on phenotypic studies have been presented as reasons for a barrier to conjugation with and other means of uptake of exogenous DNA. In this study, we show that inactivation of the R.EcoKI restriction enzyme in strain *Escherichia coli* **K-12 strain MG1655 increases the conjugational transfer of plasmid pOLA52, which carriers two EcoKI recognition sites. Interestingly, the results were not absolute, and uptake of unmethylated pOLA52 was still observed in the wild-type strain (with an intact** *hsdR* **gene) but at a reduction of 85% compared to the uptake of the mutant recipient with a disrupted** *hsdR* **gene. This leads to the conclusion that EcoKI restriction-modification affects the uptake of DNA by conjugation but is not a major barrier to plasmid transfer.**

The exchange of chromosomal and/or extrachromosomal DNA, such as plasmids, viruses, and transposons, is crucial for the evolution of bacteria and their ability to adapt to new environments. Exchange of genetic material occurs among both related and unrelated species of bacteria and is driven by the three horizontal gene transfer (HGT) mechanisms: conjugation, transformation, and transduction [\(1](#page-78-0)[–](#page-78-1)[3\)](#page-78-2). Restriction-modification (RM) systems are described as major barriers to HGT [\(4](#page-78-3)[–](#page-78-4)[6\)](#page-78-5) and comprise restriction endonucleases with a cognate methyltransferase. These recognize and cleave DNA not modified by the methyltransferase, thereby making the bacterium able to distinguish between its own (methylated) DNA and incoming nonmethylated DNA.

Based on their protein-complex subunit composition and functionality, RM systems can be divided into four types; this study focuses on type I. Type I systems require products of the three genes *hsdR* (restriction), *hsdM* (methylation), and *hsdS* (sequence specificity) and cleave randomly at a remote distance from the recognition sequence. Restriction occurs only when a protein complex of all three gene products (R_2M_2S) is formed, whereas methylation of the DNA requires formation of a complex of only the HsdM and HsdS proteins $(M₂S)$ [\(7\)](#page-78-6).

Some studies have indicated that transfer by conjugation is unaffected by RM systems but that unmodified phage or free DNA in transformation is readily degraded [\(8](#page-78-7)[–](#page-78-8)[11\)](#page-78-9). This has led to the view that the conjugational transfer of plasmids through a singlestranded DNA intermediate is immune to restriction by RM systems, as the great majority of these recognize only nonmethylated double-stranded DNA [\(12](#page-78-10)[–](#page-78-11)[17\)](#page-78-12). Other studies have, however, contradicted this. In 1964, Arber and Morse [\(18\)](#page-78-13) proposed that host specificity (RM systems) might play a role in the acceptance or rejection of DNA transferred by conjugation in *Escherichia coli*. In many studies from the 1960s, the transfer was measured with recombinants of Hfr strains [\(18](#page-78-13)[–](#page-78-14)[20\)](#page-78-15), but Arber and Morse made a phenotypic study showing that the conjugational transfer of episomes (with the ability to express genes without integration into the bacterial chromosome) was affected in the same manner as in phages [\(18\)](#page-78-13). Other experiments have shown reduced conjugational transfer between different bacterial species with diverse restriction-modification systems, indicating that they may be the cause of this reduction in transfer $(21, 22)$ $(21, 22)$ $(21, 22)$, but none of these observations have been confirmed with isogenic strains by modern molecular techniques. Recent studies indicated that SauI, a type I RM system for *Staphylococcus aureus*, may be a barrier to transfer into and between *S. aureus* isolates [\(23\)](#page-78-18), but Veiga and Phino showed that inactivation of the SauI system was not sufficient for producing strains that efficiently take up foreign DNA [\(6\)](#page-78-5), again questioning the importance of RM systems as barriers to conjugational transfer.

In the current study, we aimed to clarify the impact of a restriction-modification system in the conjugational gene transfer of single-stranded plasmidic DNA [\(24,](#page-78-19) [25\)](#page-78-20) at the genotypic level. We focused on the impact of the type I RM system EcoKI, with the recognition sequence AACN_6GTGC , in the transfer of conjugative plasmids between RM variants of the *E. coli* K-12 strain MG1655. In addition to possessing the type I RM system, which we examine in the current study, MG1655 possess three different methylationrequiring type IV systems, EcoKMcrA, EcoKMcrBC, and EcoKMrr. All three systems are sequence specific and will not interfere with plasmid transfer between isogenic strains used in the current study [\(26](#page-78-21)[–](#page-78-22)[29\)](#page-78-23). We provide evidence that the RM system EcoKI has a significant and negative effect on conjugation but also that this is not a major barrier to conjugation.

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^a For bacterial strains and plasmids, the relevant genotype, phenotype, and other characteristics are shown. Abbreviations: INT, interruption; COMPL, complementation. *b* The genes targeted by the primers used in the construction of knockout strains or complementary plasmids are shown in parentheses at the end of the entry.

MATERIALS AND METHODS

Media and reagents. *E. coli* cells were cultured in brain heart infusion (BHI) broth at 37°C. For growth on agar, Luria-Bertani (LB) or BHI agar plates were used. The following antibiotics and concentrations were used: ampicillin (Amp), 50 µg/ml for cloning or 100 µg/ml in HGT assays; chloramphenicol (Cam), 25 µg/ml; kanamycin (Kan), 25 µg/ml; tetracycline (Tet), 5 μ g/ml or 10 μ g/ml; rifampin (Rif), 25 μ g/ml; and nalidixic $acid (Nal), 25 µg/ml.$

Strains and plasmids. Bacterial strains, plasmids, and oligonucleo-tides used in this study are listed in [Table 1.](#page-75-0) 32 The conjugation experiments were performed from donors with $(hsdM^+)$ and without ($\Delta hsdM$) the ability to methylate the DNA and with functional, disrupted, and complemented restriction abilities. Two plasmids with a published DNA sequence, pOLA52 and pHHA45 (GenBank accession numbers [EU370913.1](http://www.ncbi.nlm.nih.gov/nuccore?term=EU370913.1) and [JX065630.1,](http://www.ncbi.nlm.nih.gov/nuccore?term=JX065630.1) respectively), were used to assess the influence of the RM systems on conjugational transfer. pOLA52, belonging to the IncXI incompatibility group, has two recognition sites for the EcoKI system, while pHHA45, belonging to the IncN incompatibility group, does not contain any sites. Both IncXI and IncN plasmids are known to transfer in the single-stranded form [\(24,](#page-78-19) [25,](#page-78-20) [30,](#page-78-24) [31\)](#page-78-25).

Construction of TargeTron insertion mutants of *E. coli* **MG1655.** Two TargeTron mutants with interruptions, MG1655-*hsdR*INT and MG1655-*hsdM*INT, were created by following the guidelines from SigmaAldrich [\(32\)](#page-78-26) for insertion mutations in *E. coli* strains, with the plasmid pAR1219 as the source of T7 RNA polymerase. Plasmid pACD4K-C-loxP was used as the donor for the group II intron, retargeted by PCR with primers designed for position 1740|1741 in *hsdR* (primers LR1, LR2, and LR3) and position 720|721 in *hsdM* (primers LR5, LR6, and LR7). Gene disruptions were induced by the addition of 20 μ l of a 10 mM stock solution of isopropyl- β -D-thiogalactopyranoside (IPTG) to 2 ml of culture.

The plasmids (retargeted pACD4K-C-loxP and pAR1219) were cured by overnight growth in broth, followed by plating on BHI, and patched on BHI with 50 μ g/ml Amp and BHI plus 25 μ g/ml Cam to identify plasmid-free isolates. MG1655-hsdR^{INT} was made electrocompetent as described by D. O'Callaghan and A. Charbit [\(33\)](#page-78-27), but the glycerol washing step was performed with the full-strength original volume (100 ml). MG1655-*hsdR*INT was further transformed with the plasmid 706-Cre to remove the kanamycin resistance marker (Gene Bridges GmbH). Single colonies of both MG1655-*hsdM*INT and MG1655-*hsdR*INT were tested by PCR for the TargeTron insert and removal of the kanamycin gene (MG1655-*hsdR*INT only) by using *Taq* polymerase (Fermentas) and internal and external primers LR3 and LR4 (*hsdR*) or LR7 and LR8 (*hsdM*). For further verification, the PCR products were purified using a GFX purification kit (GE Healthcare) and sequenced by Macrogen Korea.

FIG 1 Single-stranded horizontal gene transfer with isogenic donor and recipient strains of *E. coli* K-12 MG1655. All experiments were performed at least in triplicate, and data are means of transconjugants per 10⁸ recipients, with standard errors of the mean (SEM). The levels of transfer of the plasmids for the three recipients were compared individually for each donor (strain and plasmid). Statistically significant differences were observed between the recipients MG1655-*RN* and MG1655-*hsdR^{COMPL}* with the donor MG1655-*A* (a*, *P* = 0.021) and between MG1655-*hsdR*^{INT} and the remaining two recipients with the donor MG1655*hsdM*^{INT} (b*, *P* = 0.029) with the plasmid pOLA52, which harbors two recognition sites for EcoKI. No statistically significant differences were observed with the two donors with the plasmid without recognition sites for EcoKI, pHHA45.

Complementation of *hsdR* **with p***hsdR***.** For complementation in the *hsdR* mutant, we amplified a 3,633-bp fragment encompassing the *hsdR* gene with the primers LR9 and LR10 by PCR and cloned the resulting fragment into the PvuII site of pMSC83. For a detailed description, see the supplemental material.

Horizontal gene transfer of single-stranded DNA by conjugation. The strains MG1655-A and MG1655-hsdM^{INT} were transformed with the plasmid pOLA52, containing two recognition sites for EcoKI, or pHHA45, lacking EcoKI recognition sites, to act as donors in the conjugation experiments. The strains MG1655-*RN*, with a functional *hsdR* gene on the chromosome, MG1655-*hsdR*INT, with an interrupted restriction gene, and MG1655-*hsdR*^{COMPL}, with a complemented restriction gene, were used as recipients.

Two individual conjugation experiments were performed, one with transfer of pOLA52 and one with pHHA45. In both cases, overnight cultures of donor and recipients were reinoculated into fresh preheated BHI medium and grown to an optical density at 600 nm $OD₆₀₀$ of 0.5. Then, 1-ml samples of each donor and recipient were mixed in 24-well microtiter plates and incubated at 37°C for 5 h. Conjugation mixtures were diluted and plated on selective plates for CFU counting.

RESULTS

Horizontal gene transfer of single-stranded DNA by conjugation. The results of the conjugational transfers are presented in [Fig. 1.](#page-76-0)

All three recipients, MG1655-*RN*, MG1655-*hsdR*INT, and MG1655-hsdR^{COMPL}, accepted the methylated plasmid pOLA52 from donor MG1655-*A* at high ratios (transconjugants per 10⁸ recipients) of 2.05 \times 10³, 1.04 \times 10³, and 0.67 \times 10³, respectively. The level of conjugational transfer into the MG1655-hsdRCOMPL strain complemented with the *hsdR* gene in *trans* was significantly lower (*P* = 0.021) than in the MG1655-*RN* strain, with a wild-type functional *hsdR* gene on the chromosome, possibly due to the higher copy number, stronger promoter, or both.

For the transfer of the unmethylated plasmid pOLA52, with two recognition sites, from the donor MG1655-*hsdM*INT to the three recipients, MG1655-*RN*, MG1655-*hsdR*^{INT}, and MG1655-

hsdR^{COMPL}, the ratios were 0.42×10^3 , 2.79 \times 10³, and 0.17×10^3 , respectively. The recipient MG1655-*hsdR*INT showed a statistically significantly ($P = 0.029$) higher level of transfer, which was more than 6.5 times higher than those of the two recipients with functional *hsdR* genes.

In the transfer of pHHA45, no significant difference was observed between the ratios of 8.45 \times 10⁴, 7.73 \times 10⁴, and 3.58 \times 10⁴ from the donor MG1655-A or between the ratios of 9.47×10^4 , 6.25×10^4 , and 7.66×10^4 from the donor MG1655-*hsdM*^{INT} to the recipients MG1655-*RN*, MG1655-*hsdR*INT, and MG1655 *hsdR*^{COMPL}, respectively (see Table S1 in the supplemental material).

Complementation of *hsdR* **restores restriction activity.** The restriction gene *hsdR* was cloned into the expression vector pMSC83 under the control of the arabinose promoter pBAD, which is known to be leaky in rich media (34) . The conjugation experiment was therefore performed without addition of arabinose to avoid overexpression of the *hsdR* gene, which could potentially be harmful to the cell. As a control for sufficient *hsdR* expression and to verify that the decrease in transfer observed in [Fig. 1](#page-76-0) was caused by expression of the *hsdR* gene alone and not the vector pMSC83, conjugation with the complemented strain and a control strain with the pMSC83 vector was performed. For the control experiment, the methylation-deficient donor MG1655 hsdM^{INT} was used with each plasmid (pHHA45 and pOLA52). The results of the conjugative control experiment are presented in [Fig. 2.](#page-77-0)

In experiments with the unmethylated plasmids pHHA45 and pOLA52, the conjugational transfer of pOLA52 to MG1655 h sdR^{COMPL} was significantly decreased ($P = 0.00031$) compared with that to MG1655-hsdR^{CONTROL}, with transfer ratios of 0.21 \times 10^3 and 5.89×10^3 , respectively. With the transfer of pHHA45, no significant difference was observed between the transfer ratios of 5.70×10^4 and 3.37×10^4 for MG1655- $\emph{hsdR}^{\rm COMPL}$ and MG1655hsdR^{CONTROL}, respectively.

FIG 2 Conjugative effect of *hsdR* from the unmethylated donor strain MG1655-*hsdM*INT inserted into the complemented strain MG1655 *hsdR*COMPL compared to that in a control strain harboring the empty vector pMSC83. For conjugation with the unmethylated plasmid pHHA45, which lacks recognition sites for MG1655-*hsdR*, no significant difference was observed between the complementation and control strains. For conjugation with the plasmid pOLA52, which has two recognition sites, a significant difference was observed between the two recipients $(a^*, P = 0.00031)$.

DISCUSSION

Previous studies of the influence of RM systems in conjugation have generated conflicting conclusions in relation to the effect of RM systems on plasmid transfer [\(8](#page-78-7)[–](#page-78-29)[15\)](#page-78-30). This has led to some controversy on how restriction-modification systems act on the uptake of single-stranded DNA, such as plasmids transferred by conjugation. An obvious driver of the conflicting conclusions from these studies is the fact that many of these studies were carried out in an era before the emergence of molecular techniques in microbiology and before the genetic determinants responsible for the RM phenotypes were identified.

The current study aimed to utilize isogenic strains and defined knockout genetic constructs to study how RM systems influence plasmidic transfer and showed that the type I restriction-modification system EcoKI in *E. coli* K-12 MG1655 affects conjugational transfer if the transferred DNA includes nonmethylated recognition sites.

Transfer of the methylated plasmid pOLA52 from the wildtype host to the three different recipients, with different restriction abilities, was not expected to have any significant influence on its uptake, as the plasmid was modified as "self" and in all three cases should have been immune from degradation. Transfer of the nonmethylated plasmid into the complemented strain was significantly lower than into the wild-type strain; more surprisingly, the same was true for the methylated plasmid. One explanation for this decrease might be incompatibility between pOLA52 and the vector carrying the complementation gene *hsdR*, but this was not expected, as the two replicons belong to IncXI and ColE10, respectively. This was also verified by the control experiment [\(Fig. 2\)](#page-77-0), where the results clearly showed that the decrease in transfer was caused by the restriction gene alone and not the vector pMSC83. Therefore, a more likely explanation may be that overexpression of EcoKI, resulting in the formation of R_2M_2S complexes rather than $M₂S$ complexes, leads to degradation of hemi-methylated DNA, as in the type II systems described by Nelson et al. [\(35\)](#page-78-31).

The unmethylated plasmid pOLA52 was taken up significantly less in the wild-type strain than in strain MG1655-hsd $R^{\rm INT}$, which has an interrupted restriction gene. The plasmid without recognition sites for EcoKI can transfer efficiently between donor and recipients independently of methylation and restriction abilities. This was shown by the transfer of the plasmid pHHA45 from both the methylation-deficient donor MG1655-*hsdM*INT and the MG1655-*A*donor with the functional *hsdM* gene to three different recipients with functional, disrupted, and complemented restriction genes.

Even though the transfer efficiencies of the two plasmids, pOLA52 and pHHA45, differ by nearly 2 logs in the *hsdR*-disrupted recipient, we find pHHA45 an appropriate control plasmid, as both plasmids are narrow-host-range plasmids and about the same size. More plasmids could be investigated to confirm our observation that the conjugational transfer or uptake in *E. coli* K-12 strain MG1655 is in fact dependent not only on the donor's methylation and the recipient's restriction abilities but also on the presence of recognition sequences on the plasmid. The results of this genotypic study are in agreement with the study performed by Arber and Morse [\(18\)](#page-78-13), where the RM-deficient strains were selected based on phenotypes by testing their ability to restrict different phages, but they were not further characterized.

From the present study, we have shown that a type I RM system can act as a barrier to the conjugational transfer of plasmids, but in many of the previous studies, the restriction is described only phenotypically [\(10,](#page-78-8) [18\)](#page-78-13) or as a transfer between unrelated species [\(8,](#page-78-7) [11,](#page-78-9) [12\)](#page-78-10) without our knowing the type of RM system involved. There is evidence that a type-III-like RM system may act as barrier to transformation in *S. aureus* strains [\(36\)](#page-78-32) but not to conjugational transfer. Further, the present study focuses on only a single system, in a single isolate, with only two different plasmids. This leads to the questions of whether all type I systems influence conjugational transfer and whether the three remaining systems have the same ability to protect hosts from invading foreign DNA. Murray and colleagues suggest that the protection from foreign DNA might be altered by alleviation of chromosomal restriction genes, which might lead to uptake [\(37](#page-78-33)[–](#page-78-34)[39\)](#page-79-1).

Pérez-Mendoza and de la Cruz [\(40\)](#page-79-2) investigated two different knockout libraries, the Keio collection of single knockouts and a random insertion library, to determine how recipient cells contribute to bacterial conjugation [\(40\)](#page-79-2). Their only finding was that the lipopolysaccharide (LPS) showed strong conjugation inhibition when conjugation was performed in liquid, but with filter mating, the reduction was restored. The Keio collection is based on a restriction-deficient K-12 strain variant (BW25113), which explains why they did not find EcoKI as a contributing factor in conjugation, as we did in this study.

Even though Pérez-Mendoza and de la Cruz did not find any genes responsible for the conjugational uptake in the recipient, this does not preclude the possibility that such genes exists. The two parental *E. coli* strains used in their study may be missing regulatory genes responsible for conjugational transfer. To identify possible barriers to conjugational uptake, as well as uptake by the other HGT mechanisms, good and poor recipients must be identified by phenotype and compared at the genetic level.

In summary, the EcoKI RM system found in *E. coli* K-12 strain MG1655 affects the conjugational transfer of plasmid pOLA52,

harboring two recognition sites, but the results imply that this effect is not absolute and that uptake is still possible, though at a lower level. The results showed 4.88-times-higher uptake of methylated pOLA52 than of the unmethylated plasmid in wild-type MG1655. This leads to the conclusion that plasmids with the same methylation pattern as the recipient can have a competitive advantage when entering a new host.

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Roer II

Is the Evolution of *Salmonella enterica* **subspecies** *enterica* **Linked to Restriction-Modification Systems?**

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For Submission to mSystems

Supplemental material can be accessed at: https://www.dropbox.com/sh/wz4v9rzem87eccn/AAAPJqt4m_TWX-Bhv7M_1PI_a?dl=0

ABSTRACT

Salmonella enterica ssp. *enterica* is a highly diverse foodborne pathogen that is sub-divided into

more than 1,500 serovars. The diversity is believed result from mutational evolution as well as

intra- and interspecies recombination that potentially could be influenced by Restriction-

Modification systems (RM systems). The aim of this study was to investigate if RM systems were

linked to the evolution of *Salmonella enterica* ssp. *enterica*.

The study included 221 *Salmonella enterica* genomes of which 68 were *de novo* sequenced, in

addition to 153 public available genomes from ENA. The dataset covered 97 different serovars of

Salmonella enterica ssp. *enterica,* and additional five genomes from four other *Salmonella*

subspecies as an outgroup for constructing the phylogenetic trees. The phylogenetic trees were

constructed based on similarity in core-genes as well as the presence or absence of pan-genes. The

topology of the trees was compared to the presence of RM systems, antimicrobial resistance (AMR)

genes, *Salmonella* Pathogenicity Islands (SPIs), and plasmid replicons.

We did not observe any significant correlation between the evolution and the RM systems in *S.*

enterica ssp. *enterica*. However, sub-lineage correlations and serovar specific patterns was observed

in few cases. Additionally, the results implied that plasmids replicons, SPIs, and AMR all were

better correlated to serovars than to RM systems.

This study suggests a limited influence of RM systems on evolution of *Salmonella enterica* ssp.

enterica, very likely due to the conjugational mode of horizontal gene transfer in *Salmonella*. Thus,

other factors must be involved in shaping the evolution of bacteria.

IMPORTANCE

The evolution of bacterial pathogens, their plasticity and ability to constantly change and adapt to

new surroundings are crucial for understanding the epidemiology and public health. Along with the

 application of genomics, it became clear that horizontal gene transfer played a key role in evolution. Thus, to understand, prevent, fight, and control the spread of undesirable pathogens, we need to elucidate the processes that drive the horizontal gene transfer, the evolution and diversification. Restriction-modification systems are thought to cause rearrangements within the chromosome, as well as act as a barrier to horizontal gene transfer. However, here we show that the very persuasive correlation between restriction-modification systems and evolution in other bacterial species is not present in *Salmonella enterica* ssp*. enterica.* In summary, this work illustrates that other mechanisms must be involved in controlling the evolution.

INTRODUCTION

 The *Salmonella* genus is highly diverse and comprised by the two species; *Salmonella enterica* and *Salmonella bongori*. The species *S. enterica* contains six subspecies of which the highly diverse *S. enterica* ssp. *enterica* can be sub-divided into more than 1,500 serovars. The subspecies is a common cause of diseases in humans and domestic animals (1, 2) and one of the leading causes of foodborne illness worldwide (3).

 Recombination between genomes is thought to be a major driver in evolution (4), and to contribute to the diversity within the *Salmonella* genus (5, 6). In *Neisseria meningitidis* it was suggested that the phylogenetic relationship is associated with the content of Restriction-Modification systems (RM systems) (7). Further, rearrangements of genomes caused by RM systems are described as factors that could influence the evolution of pathogens (4). In addition to their role in rearrangements, RM systems are also considered to be a barrier for horizontal gene transfer between bacteria, thus serving as an immune defence system for uptake of foreign(4) DNA (8–10). The precise contribution have to our knowledge never been quantified, but we have recently shown that for conjugational transfer between isogenic *Escherichia coli isolates,* RM systems are not absolutely barrier (11).

 RM systems are comprised of a restriction enzyme (RE) and a cognate methyltransferase. The restriction enzyme recognizes and digests foreign incoming DNA, whereas the methyltransferase perform methylation of the bacterium's own DNA to protect itself from degradation from the cognate restriction enzyme (12, 13). This enables the bacteria to distinguish between own (methylated) DNA and incoming non-methylated DNA.

The RM systems are divided into four types (I-IV), based on their protein-complexes, the sub-unit

composition, and the functionality of the system (14). The Type I systems are complexes of three

gene products; *hsd*R (R: restriction), *hsd*M (M: methylation), and *hsd*S (S: sequence specificity).

RESULTS

Genomes

- A total of 221 *Salmonella* genomes were included in the analysis represented by 153 genomes previously described by Timme *et al.* (1) and retrieved from The European Nucleotide Archive. This collection was merged with 68 genomes sequenced as part of the 100K Foodborne Pathogen Genome Project [\(http://100kgenome.vetmed.ucdavis.edu/,](http://100kgenome.vetmed.ucdavis.edu/) NCBI BioProject PRJNA186441). The final collection consisted of 216 *S. enterica* ssp. *enterica* genomes and five genomes of other subspecies. The 221 *Salmonella* genomes were summarized in the supplemental material (Table S1). **Characterization of Restriction-Modification systems** To characterize the RM systems of the 221 genomes, a WGS analysis was performed using the newly developed tool Restriction-ModificationFinder 1.0. Partial systems were completed by individual BLAST analysis of up- and downstream sequences against the REBASE database (19). In total, we identified 113 putative RM systems i.e.; 58 Type I RM systems (TI) with 43 of unknown recognition sequence, 23 Type II RM systems (TII), two Type III RM system (TIII), and 30 Type IV RM systems (TIV), respectively. In addition, numerous methyltransferases outside RM systems were identified, including Type I, Type, II and Type III methyltransferases (Table S2). The additional methyltransferases (without associated restriction enzyme) identified were not include in the analysis, as the barrier for horizontal gene transfer (HGT) is thought to be caused by the cleavage from RE. Two of the genomes only contained one RM system, where the other genomes contained between two and seven systems. One of the Type III RM systems were shared by 191 genomes, the type I
- system TI-1 was shared by 203 of the genomes and 37 RM systems were specific to a single

124 genome. The remaining systems were shared by two to 38 genomes. The distribution of the RM systems was illustrated in the presence/absence matrix in [Fig. 1](#page-106-0), and in details in [Fig. S1.](#page-111-0) The analysis revealed very diverse content of RM systems, and assessing the highest level of discrimination in the cladogram, 120 distinct clusters were formed with 77 clusters containing a single genome.

The Salmonella Pan- and Core-Genomes

 The pan- and core-genomes were estimated based on the 221 *S. enterica* genomes. The progression of the pan- and core-genomes is shown in [Fig. S2,](#page-111-1) as increasing numbers of the *S. enterica* genomes were added to the analysis. Analysing the pan-genome, consisting of any gene families found, the gene families increased gradually with the one by one addition of the different *S. enterica* ssp. *enterica* serovars compared to a distinct increase in the number of gene families with the inclusion of *S. enterica* ssp. *diarizonae*. In contrast, the number of conserved gene families across the *S. enterica* genomes in the core-genome analysis seemed relative consistent but with similarly drops when introducing other *S. enterica* subspecies to the analysis. The final analysis compressing all 221 *S. enterica* genomes contained 16,375 gene families in the pan-genome (File S1) and 2,138 gene families in the core-genome (File S2), respectively. Analysing the total number of gene families in the pan-genome, each *S. enterica* genome contribute in average with 65 new gene families, increasing the diversity within the *S. enterica* species.

The link between evolution and Restriction-Modification systems

 To study the genomic evolution of *S. enterica* ssp. *enterica,* differences within the core-genes were examined for all 221 genomes and illustrated by the phylogenetic core-genome tree in [Fig. 2.](#page-107-0) This evolution of *S. enterica* ssp. *enterica* is not only formed by the differences in the genes shared by

 the genomes, but also the loss of genes leading to differences in gene content. [Fig. 3](#page-108-0) displays the pan-genome tree based on the presence and absence of genes across all the genomes included in the study.

 For both the core- and pan-genome trees, the high bootstrap values of 1 near the root of the trees reflected a general trustable structure for the phylogenetic trees of the *Salmonella* genus, whereas the low bootstrap values found in some of the branches likely indicated that some of the genomes in those branches were highly similar in their core- or pan-gene content, thus it was difficult to determine the definite arrangement within the branch. However, both the core- and pan-genome trees separated well the serovars amongst one another with relatively high bootstrap values at the tips of the trees.

 To examine the hypothesis of RM systems being linked to the evolution of *S. enterica* ssp*. enterica*, parallel analyses were performed to identify groups of genomes with highly similar RM pattern that were also forming discrete phylogenetic clusters on the core- and pan-genome trees. Convergence was indicated by colours in the matrix of RM systems ([Fig. 1](#page-106-0)), and the core- and pan-genome trees ([Fig. 2](#page-107-0) and [Fig. 3\)](#page-108-0). A few small clusters with almost identical RM content, partly clustering together in the core- and pan-genome trees were observed. In most cases the genomes within a cluster belonged to the same serovar, as for *S.* Bareilly or *S.* Agona (dark green and light blue). However, three larger clades, with 21, 8 and 7 genomes, were identified in the RM matrix. Of the 21 genomes in the first clade, containing serovars of Typhimurium, Saintpaul, Paratyphi B and Heidelberg, 90% of the genomes were located together in one distinct cluster in the core-genome tree and the remaining 10% were also located together. For the pan-genome tree, the 21 genomes were identified at 4 different locations with the distribution of 67%, 19%, 9% and 5% of the genomes, respectively. The two larger clades of 8 and 7 genomes were located adjacent to each other in the RM matrix, the first containing serovars of Stanleyville, Gallinarum, Pullorum and Dublin, and the

 second containing serovars of Enteritidis and Berta. Comparing to the core- and pan-genome trees, in both cases the two clades were located together with all 100% of the genomes.

 Plasmid Replicons, Antimicrobial resistance and Pathogenicity Islands in *Salmonella enterica* All 221 *Salmonella* genomes were analysed for the content of plasmid replicons by using the Center for Genomic Epidemiology (CGE) web-tool PlasmidFinder 1.2 (20), of which 118 of the genomes did not contain any replicons present in the PlasmidFinder database. In the remaining 103 genomes, 40 different plasmid replicons were identified with each genome containing up to seven different replicons.

 Assessing the replicons in comparison with the RM systems observed in the *S. enterica* ssp. *enterica* serovars, no convergence was observed [\(Fig. 4\)](#page-109-0) and RM systems and plasmid replicons were never observed on the same contigs. Evaluating the plasmid replicons in correlation to serovars, no correlation was observed between serovars and quantity of replicons; the 11 genomes with the highest replicon content represented 10 different serovars. However, for multiple isolates with the same serovar common replicons are observed, such as with the *inc*FII and *inc*X1 replicons present in all the *S*. Dublin genomes. Additionally, *inc*FII was observed in 50% of the *S*. Enteritidis genomes, and together with the *inc*FIB replicon in 50% of the *S*. Typhimurium and in both 4,[5],12:i:- genomes. Interestingly, even though the numbers of replicons are not equal in identical serovars, the results imply a better association between replicons and serovars compared to the association between replicons and RM systems..

 AMR genes were found in 220 of the genomes, varying from one to 19 different genes per genome, with *aac(6')-Iy* present as the only resistance gene in 140 of the genomes. The correlation of AMR and RM systems was examined [\(Fig. S3\)](#page-111-2), however no congruency was observed most likely due to

 the in-built bias of the strain collection. Despite this potential bias, the assessment of correlation between AMR and plasmid replicons revealed different resistance genes located on the same contigs as plasmid replicons. Few genomes contained more than four plasmid replicons and between 10 and 16 resistance genes, correlating high resistance to the number of replicons present in the genomes.

 The presences of SPIs were assessed in all 221 *Salmonella* genomes utilizing the newly developed web-tool SPI-Finder 1.0, and visualized in [Fig. 5.](#page-110-0) SPIs were found in all 221 genomes, and the number of SPI's in each genome varied from one to 14 islands/genes of islands.

The comparison of the SPIs and the content of RM systems revealed no clear association [\(Fig. 5\)](#page-110-0).

However, associating the SPIs to individual *Salmonella* serovars such as *S. enterica ssp. enterica*

 serovars Typhimurium, Paratyphi A, Choleraesuis, Heidelberg, and Saintpaul, some association was apparent within a serovar.

DISCUSSION

 bacteria, determining whether or not foreign DNA was established in the cell. For *N. meningitidis* the impact of RM systems in evolution was recently elucidated, and the purpose of the current study was to clarify if a similar association could be identified in *S. enterica* ssp. *enterica*, by investigating a large subset of different whole genome sequenced *S. enterica* ssp. *enterica* serovars and an outgroup of five genomes from other subspecies. However, it was not possible to draw any significant association between RM systems and the overall evolution of *Salmonella enterica* ssp. *enterica,* even though we did observe genomes from the same RM clades in discrete phylogenetic clusters of both the core- and pan-genome trees. The *Neisseria* genus, including *N. meningitidis,* is known to serve as a paradigm for natural transformation, where genetic exchange happens frequently due to their persistent competence independent of the phase of their life cycle (21). As RM systems have been recognized as barriers for transformation in multiple species (22–24), the link between a natural transformable species and the RM systems, as shown for *N. meningitidis,* seems reasonable. However transformation in *Salmonella* is unlikely as they are not natural transformable, and the transfer of genetic material

For decades, RM systems have been recognized for their ability to act as "immune systems" for

happens mainly through conjugation (25). In addition, a recent study performed in *E. coli* (11)

indicated that the barriers imposed by the RM systems in conjugational plasmid transfer were not

absolute, which could explain the lack of significant association in our study when trying to

associate RM systems to the plasmid replicons, AMR, SPI's, and their influence on the evolution.

The core- and pan-genome trees were constructed with a method previously described by

Leekitcharoenphon *et al.* (26), where 73 genomes were evaluated. They found a core-genome of

2,882 genes and a cognate pan-genome of 10,581 genes. In our study, the core-genome was found

to comprise 2,138 genes and the cognate pan-genome to contain 16,375 genes of the 221 genomes

 assessed. This indicates an open pan-genome for *S. enterica* ssp. *enterica*, where addition of genomes to the analysis increases the total gene pool in the pan-genome, compared to the very uniform species *Bacillus anthracis,* having a closed genome, where the addition of genomes to the analysis will not increase the gene pool (27). This also supports the knowledge of *S. enterica* ssp. *enterica* being a highly diverse bacterial species.

Assessing the RM systems, we identified 113 RM systems and numerous individual

 methyltransferases, with each genome harbouring two to seven RM systems. Vasu and Nagaraja (4) recently described how changes in specificity or acquisition of new RM systems could alter the strains genetically from the original clonal population, as the methyltransferase modifies the genome of its new host, and the RE prevents genetic exchange between closely related strains. Thus, mutations accumulate in the "new strain", leading to genetic diversity. We found that 191 of the 221 genomes analysed shared a Type-III RM recognition sequence. Taking this result into consideration it is plausible that the Type-III system was the first RM system introduced in *S. enterica* ssp. *enterica* with subsequently a larger diversity following the later acquisition of new RM systems as described by Vasu and Nagaraja. However the same RM systems could also have been introduced in different branches at different time points, resulting in identical RM systems clade across the trees.

 Assessing the pan- and core-genome trees, there are indications of some clustering of genomes with similar RM systems i.e. the cluster of *S.* Enteritidis and *S.* Dublin as well as the red RM systems clade consisting of *S.* Typhimurium, and *S.* Heidelberg located together in both trees. However, the influence is not significant, indicating that the evolution could be driven by several factors. For instance, a previous study compared 28 *Salmonella enterica* isolates and provided evidence that clustered regularly interspaced short palindromic repeats (CRISPR) mediated sub-lineage evolution (28). Other drivers in evolution are host and environmental adaptations, which besides gene

 acquisition can be caused by gene loss and deletions, gene duplication and changes within genes by e.g. mutations (29, 30).

 In the study on *N. meningitidis* they investigated the association of RM systems, homologous recombination and the phylogenetic network (7). The main study was performed on 20 genomes, covering five serogroups out of the recorded 13 serogroups for *N. meningitidis* (31, 32). Budroni *et al*. found that genomes from the same clonal complex (CC) were located together in phylogenetic clades based on their core genes. In addition, the clades could be associated with the RM systems identified (7). In our study, we investigated 217 genomes of *S. enterica* spp*. enterica* containing 97 different serovars with an outgroup of five genomes from four other subspecies. Considering the highly diverse dataset investigated in this study, compared to the one of *N. meningitidis*, we observed small trends of sub-lineage association of RM systems and evolution. This could indicate that even though the dataset investigated in this study was comprehensive, more genomes of each serovar should be included to cover the complete picture of the influence of RM systems in evolution of *S. enterica* ssp. *enterica*. Thus, with the current speed in WGS, this might be realistic in the nearest future. Thus, even in the ideal data scenario the lack of association is very likely due to the incomplete barrier of RM systems in conjugation.

 As for all database dependent approaches, the methods are only capable of detecting and reporting records present in the database explored. Our analysis for detecting the RM systems was limited to the current knowledge presented in the REBASE (33), where the recognition sequences of various number of Type I specificity subunits were not yet determined. Thus it is likely that some strains have acquired RM systems with identical recognition pattern, however this is presently unknown. Thus, this is why the all database dependent analyses should be interpreted with care. The plasmid replicons which potentially could have an effect on the bacterial diversity due to

horizontal gene transfer were also identified, but no clear correlation between the RM systems and

 the content of harboured plasmid replicons was observed. However, this approach might be complicated by the fact that plasmids are transferable and affected by factors such as fitness costs, selective pressure (34–37), and again the fact that the RM barrier is not absolute (11). Thus, the analysis performed on the plasmid replicons illustrates the current status of the time of isolation in contrast to what plasmids potentially could be acquired. AMR can be encoded by genes located on transferable plasmids, why this potentially could reflect the promiscuity of the genomes reflected in current time which could explain the lack of association between AMR and RM systems. This might also be explained by the possible biased dataset in respect of selective criteria e.g. susceptibility to antimicrobial agents, plasmid content or virulence (SPI) for the isolates. Though, the content of the RM systems is not believed to be affected by the possible biases. The mechanism behind the acquisition of SPIs is horizontal gene transfer (17, 18). Nevertheless, the maintenance of SPIs within the genomes are considered stable (38), and is therefore a good measure of the barriers of RM systems, compared to plasmids replicons and AMR which can easily be lost if they do not confer any beneficial traits to the host. Despite this speculation, the influence of RM systems on the distribution of the SPI's was not supported by our analysis - on the contrary there were indications of some SPI's being serovar specific, which corresponds to previous findings (38). In conclusion, recombination and rearrangement events caused by RM systems are, in several cases, described as driven factors for evolution, contributing to the diversity within a species (4, 39–44). However, high recombination between two distantly related lineages of *S. enterica* is exceptional (6, 45), thus explaining the difficulties of linking the RM systems to the evolution of *S. enterica* ssp. *enterica*. Thus, recombination occurs within and between closely related serovars (6). In this study, we showed that RM systems could not be linked to the evolution of *S. enterica* ssp. *enterica,* very likely due to the incomplete barriers of RM systems in conjugation. However, we provided evidence of closely related serovars with identical RM systems; i.e. *S.* Dublin and *S.*

- could have stronger links between their genomic evolution and the presence of RM systems
- compared to the lack of association for the entire subspecies *enterica*.

MATERIAL AND METHODS

Salmonella **Strains**

From an in-house strain collection at the Technical University of Denmark, National Food Institute

(DTU FOOD), a sub-collection of 68 *S. enterica* ssp. *enterica* isolates with a global origin and a

focus on multidrug resistance was submitted to the 100K Food Pathogen Genome Project

[\(http://100kgenome.vetmed.ucdavis.edu/,](http://100kgenome.vetmed.ucdavis.edu/) NCBI BioProject PRJNA186441) for WGS.

Subsequently, the genomes from that project were merged with a genomic collection consisting of

105 *Salmonella* strains; with a majority originating from the American Type Culture Collection –

often pan-susceptible, sequenced by the Center for Food Safety and Applied Nutrition (FDA-

CFSAN) and US Department of Argiculture (USDA) (1) and with 48 public available *Salmonella*

genomes retrieved from the European Nucleotide Archive and included this study. The final dataset

of 216 *S.* ssp. *enterica* genomes was constructed with focus on diversity including a total of 97

different *Salmonella* serovars. Additionally, five genomes of four other subspecies were included in

the dataset forming an outgroup. This dataset might have an in-build bias in respect of the selective

 criteria e.g. susceptibility to antimicrobial agents. Full genomic information is shown in Table S1 in the supplemental material.

Whole genome sequencing

Genomic DNA was extracted from the 68 *Salmonella* isolates using an Invitrogen Easy-DNA kit

(Invitrogen, Carlsbad, CA, USA), and DNA concentrations were determined using a Qubit double-

stranded DNA (dsDNA) BR assay kit (Invitrogen). The genomic DNA was prepared for Illumina

pair-end sequencing using Illumina (Illumina, Inc., San Diego, CA) NexteraXT Guide

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[\(http://support.illumina.com/downloads/nextera_xt_sample _preparation_guide_15031942.html\)](http://support.illumina.com/downloads/nextera_xt_sample%20_preparation_guide_15031942.html).

 The DNA was dispatched for Illumina HiSeq whole genome sequencing at the School of Veterinary Medicine, UC Davis, USA in relation to the 100K Foodborne Pathogen Genome Project. The raw reads of the 68 sequenced genomes, received from UC Davis, were assembled using the Assembler 1.0 pipeline from the CGE available on [http://cge.cbs.dtu.dk/services/all.php,](http://cge.cbs.dtu.dk/services/all.php) which is based on the Velvet algorithms for *de novo* short-read assembly. A complete list of genomic sequence data is available in Table S1 in the supplemental material.

Construction of core-pan-genome plot and pan- and core-genome trees

 Open reading frames (ORFs) were predicted on the contigs using Prodigal software (46) and same gene predictor was subsequently used to eliminate biases in annotation quality and to standardize the genes found in all genomes (47).

 The predicted genes were translated into amino acid sequences and aligned all-against-all using BLASTP (48). Genes were in this study considered to belong to the same gene family if the alignment length was at least 50% of the longest sequence and more than 50% in similarity ("the 50/50 rule") (27).

Pan*-* **and Core-Genome Plot**

 The core-pan-genome plot was constructed by comparing the gene families from all genomes. The pan-genome was constructed from the union of the genes from the genomes under consideration, while the core-genome was built from the intersection of genes families shared by every genome under analysis (26, 27).

Pan-genome Tree

 The pan-genome tree was reconstructed from a matrix consisting of gene families (rows) and genomes (columns). In the matrix, the absence and presence of genes across the genomes were represented by 0's and 1's, respectively. The genomes were clustered using hierarchical clustering of the relative Manhattan distance between genomes and the bootstrap values were calculated to represent the confidence of branches (26, 49).

Core-genome Tree

The core genes were aligned in a blast-like manner using BLAT v. 35 (50) to the predicted genes of

each genome. The genes found in all genomes were then aligned using MUSCLE v. 3.8.31 (51) and

concatenated to a single alignment. 500 resamples of the alignment were created with Seqboot

version 3.67 (part of the PHYLIP package (52)).

A gene was considered "identified" according to the "50/50" rule. DNADist (52) was employed to

calculate the genomic distances from the initial alignment as well as each of the 500 resamples.

FastMe (53) was used to calculate trees from the distance matrices. The tree from the original

alignment was compared to the 500 trees created from the resamples using CompareToBootstrap

(54). The final tree was visualized with FigTree [\(http://tree.bio.ed.ac.uk/software/figtree/\)](http://tree.bio.ed.ac.uk/software/figtree/).

Construction of RM-Finder and SPI-Finder

 To be able to analyse the genomes for their content of RM systems and SPI's, two public available online tools were developed. Both tools were build on a BLAST-based methodology for detection of genes from customized databases, originally developed by Zankari *et al.* for in silico detection of acquired resistance genes (55). The tools were developed to process both pre-assembled genomes and data of raw reads from different sequencing platforms, with user-selections parameters for minimum percent identity (%ID) and minimum length. The default settings were chosen as minimum ID at 95% to avoid noise and fragments of the genes, and a minimum length of 60% to be able to detect genes in the start or end of contigs from bad assemblies.

The database behind RM-Finder originate from the authoritative source REBASE (19, 33), and

includes Type I-IV restriction genes, methyltransferases and specificity units. The database is

categorised into two groups, one only including genes with confirmed function, and one where

- putative genes are included. The RM-Finder database is monthly updated.
- The *Salmonella enterica* records from the PAthogenicity Island DataBase (PAIDB) served as
- inspiration for the customized database behind SPI-Finder (56, 57).
- Extensive explanations of the output can be assessed as separate tabs at the online tools;
- <https://cge.cbs.dtu.dk/services/Restriction-ModificationFinder/> and
- [https://cge.cbs.dtu.dk/services/SPIFinder/.](https://cge.cbs.dtu.dk/services/SPIFinder/)

Identification of RMS-genes, Plasmid Replicons, SPI's and Antimicrobial resistance

- To analyse the content of RM systems in the 221 genomes, all ORFs were submitted to the
- Restriction-ModificationFinder version 1.1 available on the CGE website
- [\(https://cge.cbs.dtu.dk/services/Restriction-ModificationFinder/\)](https://cge.cbs.dtu.dk/services/Restriction-ModificationFinder/), to identify restriction- (R),
- methyltransferase- (M) and specificity (S) genes. Subsequently, the R, M and S genes identified
- were individually inspected to form RM systems, and putative systems were assigned when all
- genes required were present and adjacent on the contig, even if truncated or frame-shifted. For
- systems with unknown specificity, systems were assigned according to the specificity subunit
- present. However, incomplete systems were investigated for truncated genes. Additionally, contigs
- with incomplete systems were inspected by BLAST against REBASE, and putative genes for
- completion were revealed. Based on the predicted recognition sequences, the systems were merged
- and named according to the type of system. A RM systems presence/absence matrix was
- constructed in R-2.14 [\(http://cran.r-project.org/bin/windows/base/old/2.14.0/\)](http://cran.r-project.org/bin/windows/base/old/2.14.0/) with hierarchical
- clustering, and euclidean distance [\(Fig. 1](#page-106-0) and Fig. S1).
- The 221 draft genomes were analyzed for the content of plasmid replicons, pathogenicity islands
- and antimicrobial resistance genes, by using the CGE web-tools; PlasmidFinder 1.1 (20), SPIFinder
- 1.0 [\(https://cge.cbs.dtu.dk/services/SPIFinder/\)](https://cge.cbs.dtu.dk/services/SPIFinder/) and ResFinder 2.0 (55) with %ID = 80.00 and
- minimum length at 60.00%. The content of plasmid replicons and pathogenicity islands was
- 408 interpreted by Circos plots [\(http://circos.ca/\)](http://circos.ca/). For the antimicrobial resistance genes, a
- presence/absence matrix was constructed as described for the RM systems (Fig. S2).

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- **Fig. 1. Presence or absence matrix of the 113 restriction-modification systems.**
- In the matrix, each row represents one genome analysed, and each column represents one of 113 RM systems. The
- cladogram is a hierarchical clustering of the genomes based on the Euclidean algorithm. Convergence of genomes with
- highly similar RM system content and discrete phylogenetic clades on the core- and pan-genome trees was indicated by
- the colours in the cladogram.

- Phylogenetic tree constructed on core genes. Discrete phylogenetic clades with highly identical RM system content
- were indicated by the colours defined i[n Fig. 1](#page-106-0)**.**

constructed from presence/absence matrix of genes across genomes. The colours represent the different groups of RM

systems defined in [Fig. 1](#page-106-0)**.**

 Fig. 4. Map of plasmid replicons in *Salmonella enterica.* The genomes in the map are ordered according to their RM systems, and coloured as in [Fig. 1.](#page-106-0) The bright gray colour indicates absence of replicon in the given genomes, where presence is indicated by a colour specific for the replicon. From the outside to the middle the order of the replicons are:

- *inc*A/C, *inc*A/C2, ColMGD2, Col156, Col8282, ColE10, ColpVC, ColRNAI, *inc*FIA, *inc*FIB, *inc*FIC, *inc*FII, *inc*HI1A,
- *inc*HI1B, *inc*HI2, *inc*HI2A, *inc*I1, *inc*I2, *inc*N, *inc*P, p0111, *inc*Q1, *inc*R, *inc*X1, *inc*X4.

 Fig. 5. Map of *Salmonella* **Pathogenicity Islands found in** *Salmonella enterica* **genomes.** The genomes are ordered by their RM system profile, indicated by the colours from [Fig. 1.](#page-106-0) Each SPI is defined by a circle with different variants

- indicated by colour in the given circle.
- From the inner circle and outwards, the order of the SPI's is: C63PI, CS54-island, SGI1, SPI-1, SPI-2, SPI-3, SPI-4,
- SPI-5, SPI-7, SPI-8, SPI-9, SPI-10, SPI-11, SPI-12, genes of SPI-13, genes of SPI-14.
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- File S2: The 2,138 Salmonella Core genes in FASTA format

Roer III

Identification of *Salmonella enterica* **recipient genes affecting plasmid conjugation**

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Supplemental material can be accessed at: <https://www.dropbox.com/sh/src29dglfq2h0pu/AADsmz81L9M5VzfKpVNORf0Ha?dl=0>

1 **Title: Identification of** *Salmonella enterica* **recipient genes affecting plasmid**

2 **conjugation**

3 Running title: Recipient genes affecting conjugation

Abstract

 The spread of genetic threats through horizontal gene transfer is of very high importance. By understanding the mechanisms and genes associated with conjugational ability in recipients we might be able to predict, delay or even control the spread of antimicrobial resistance. The aim of this study was to identify species-specific gene family candidates from whole genome sequenced *Salmonella enterica*, which are enhancing or restricting plasmid uptake and maintenance in recipient cells.

 The study included 93 *Salmonella enterica* isolates, covering 54 different serovars. To evaluate the ability of the 93 isolates to take up a plasmid, an initial conjugation experiment was performed with the *mCherry* marked *Escherichia coli* K-12 strain MG1655 donor, and the *gfp* reporter constructed IncP-1β plasmid pKJK5. The conjugation frequencies, ranging from 0.0 to 2.5×10^{-1} transconjugants per potential recipients, facilitated a subdivision of the genomes by using the kmeans algorithm. The predicted gene families of the resulting groups of ten good and 83 poor recipients were compared by different approaches to identify gene family that might be involved in either enhancing or restricting conjugation. In addition, a parallel analysis was performed, on 11 *S.* Enteritidis isolates, to elucidate if serotype specific genes could be involved in conjugation. For the entire collection of 93 *Salmonella enterica* isolates, we were not able to detect unique gene family candidates associated with the either high or low transfer frequencies. However, when assessing the 11 *S.* Enteritidis genomes separately, 33 gene family candidates were identified that were associated with restricting conjugational abilities.

 This study suggests that no common genes exist for *Salmonella enterica*, which neither enhancing nor restricting conjugation. However, serovar specific gene family candidates should be

- investigated to clarify their impact on conjugation. Additionally, the results obtained in this study
- indicate that bacteria can avoid being used as recipients.

Introduction

 Horizontal gene transfer has played a major role in both long- and short term bacterial evolution. Mobile genetic elements such as plasmids and transposons are widespread in all bacterial species [1]. With the global emergence and rapid spread of multiple antibiotic resistance in recent years, it has become clear that the importance of horizontal gene transfer in evolution is of very high significance, perhaps even higher than previously expected [2–4], especially with the global transmission of resistance plasmids into several bacterial species [5]. We also observe seemingly a difference in the ability of specific clonal lineages and bacterial species to acquire resistance plasmid, which cannot just be explained by differences in reservoirs [6, 7]. If we can identify genetic traits associated with reduced or increased ability to acquire resistance genes we might better predict where resistance will emerge.

 In conjugation, there is a need for active participation of both mating partners [8]. In a recent study, Pérez-Mendoza and de la Cruz tried to identify recipient genes in *Escherichia coli* affecting the ability in plasmid conjugation [9], but the results indicated that none of the non-essential genes from the *E. coli* recipient played an important role in conjugation. Different host defense mechanisms have been suggested to impose barriers for conjugation, including the RM systems. However a study performed on *E. coli* revealed that these barriers were not absolute [10], and in addition could not be reflected in the evolution of *Salmonella enterica* ssp*. enterica* (unpublished data from Roer *et al*., 2015). It has been observed that host-specific *Salmonella* serovars are carrying considerably less resistance genes compared to the serovars with multiple hosts, although they have been isolated from the same reservoir [6, 7]. This phenomenon has so far not been explained, thus it is likely that a varying ability to receive DNA could be caused by genetic difference among the clones.

 In this study we aimed to identify genetic traits associated with uptake of IncP-1 plasmids. IncP-1 plasmids were originally found in clinical bacterial isolates [11, 12], and are still of clinical concern because of their ability to carry and facilitate the spread of antibiotic resistance [13, 14] and the IncP transfer (*tra*) system of the broad-host-range plasmid RP4 is one of the best-studied conjugative systems [15]. We investigated several *Salmonella* recipients of various serovars, and divided them into groups of good and poor recipients based on their transconjugant frequencies using an *E. coli* donor. Different bioinformatical approaches were taken in the attempt to identify genes responsible for promoting or resisting conjugational transfer.

Material and Methods

Bacterial donor strain and conjugative plasmid

The mCherry tagged donor *E. coli* MG1655::*lacI^q-pLpp-mCherry-Km*^{*R*} with the conjugative *gfpmut3*-tagged pKJK5 plasmid was kindly provided by Uli Klümper *et. al* [16]. Essentially, the plasmid was genetic marked with a gfpmut3 gene, encoding green fluorescent protein (GFP), 76 conditionally expressible by a lacI_q repressible promoter located upstream of gfpmut3 gene [17]. Further, the plasmid donor strain *E. coli* MG1655 was chromosomally marked with a gene cassette encoding constitutive expression of both red fluorescence (mCherry gene) and lacIq. As a result, the 79 constitutive lac I_q expression ensures a gfp repression on the plasmid in the donor strain, but upon plasmid transfer to the recipients, gfp expression is possible resulting in green fluorescent transconjugants detectable by flow cytometry. Donor strain and conjugative plasmid are listed in

[Table 1.](#page-133-0)

Salmonella enterica **Recipients**

 From an in-house strain collection at the Technical University of Denmark, National Food Institute (DTU FOOD), a collection of 35 *S. enterica* spp. *enterica* isolates with a global origin and a focus on serovar diversity was merged with a sub-set of 58 genomes previously sequenced as part of the so-called 100K project (NCBI BioProject PRJNA186441). The final recipient collection consisted of 93 *Salmonella* spp. *enterica* genomes covering 54 different serovars. A complete list with genomic information is avalible in S1 Table in the supplemental material.

Whole genome sequencing

From the in-house collection, genomic DNA was extracted from the 35 *Salmonella* isolates using

Invitrogen Easy-DNA kit (Invitrogen, Carlsbad, CA, USA). DNA concentrations were determined

using Qubit double-stranded DNA (dsDNA) BR assay kit (Invitrogen). The genomic DNA was

- prepared for Illumina pair-end sequencing using Illumina (Illumina, Inc., San Diego, CA)
- NexteraXT Guide 150319425031942 and following protocol revision C
- [\(http://support.illumina.com/downloads/nextera_xt_sample _preparation_guide_15031942.html\)](http://support.illumina.com/downloads/nextera_xt_sample%20_preparation_guide_15031942.html),
- and submitted for whole genome sequencing at the in-house Illumina MiSeq sequencer.
- The raw reads of both in-house 35 genomes and the 58 genomes from 100K project were assembled
- using the Assembler v1.0, which is based on the Velvet algorithms for *de novo* short-read assembly,
- available from the Center for Genomic Epidemiology (CGE), [http://cge.cbs.dtu.dk/services/all.php.](http://cge.cbs.dtu.dk/services/all.php)

Conjugation with pKJK5::*gfp*

102 A single colony of the donor strain was grown in LB with TMP at 37° C overnight. The individual 103 recipients were inoculated in 24 well plates (Nunc) and grown overnight at 37^oC. The growth was stopped by incubating the overnight cultures on ice for 3 hours. The optical density (OD) was measured at 600 nm and donor and recipients were diluted to a final OD at 0.5. The donor was mixed with the individual recipients in a ratio of 1:1 in 24 well plates, and incubated 18 hours for 107 conjugation at 37°C. The conjugations were interrupted by properly mixing the liquid, and in the initial screening plates were stored on ice for 30 minutes. Donor, recipients and transconjugants were detected and counted by flow cytometry. All samples were diluted in 0.9% NaCl to reach approximately 2,000 counting events per second before running on flow cytometer.

Flow cytometric detection

Detection of transconjugants, donor, and recipient cells was carried out by using a FACSAria IIIu

(Becton Dickinson Biosciences, San Jose, CA, USA). The following settings and voltages were

used for detection: forward scatter (FSC) = 505V, side scatter (SSC) = 308V, the bandpass filter

530/30 nm and 508V were used for detection of green fluorescence (GFP) whereas the bandpass

filter 610/20 nm and 500V were used for red fluorescence detection. A 70 mm nozzle was used with

 a sheath fluid pressure of 70 psi. For operating the system the BD FACSDiva software v.6.1.3 was used, whereas FlowJo v.10 was used for analyzing the data. Five gates were defined in three bivariate plots to distinguish donors, recipients and transconjugants. On the SSC-A vs. FSC-A plot, one gate was defined for particles of bacterial size. On the PE-Texas Red-A vs. SSC-A plot two gates were defined, a gate covering all red fluorescent particles defining the donor bacterial cells, and a non-red gate for the recipient- and transconjugant cells. From the non-red gate, on a FITC-A vs. SSC-A plot, a gate was set to cover all green fluorescent particles, while an additional non-green gate was set to account for the recipient cells [\(Figure 1\)](#page-134-0). Each sample was set to analyze a total of 100,000 events with the size of bacteria.

Defining good and poor recipients

 From the conjugation experiments, transfer frequencies were defined as number of transconjugants per number of potential recipients, and the mean transfer frequencies (TFs) were calculated for each isolate. The algorithm kmeans in R-2.14 [\(http://cran.r-project.org/bin/windows/base/old/2.14.0/\)](http://cran.r-project.org/bin/windows/base/old/2.14.0/) was used to divide the isolates into two groups (good and poor recipients, respectively) based on their mean TF. The algorithm partitions each point into the group where the sum of squares from the point to the center of the cluster gives the lowest value. To illustrate that it was reasonable to group by the TF mean, a clusplot was drawn based on data from donor, recipient, transconjugant, TF and transconjugant/donor for each isolate.

Identification of recipient determinants involved in conjugation

For identification of recipient determinant of the 93 genomes that might be involved in conjugation,

two different approaches were taken. In the first attempt all 93 genomes were analysed together.

- The Prodigal software [18] was used to predict open reading frames (ORFs) on the scaffolds to
- eliminate biases in annotation quality and to standardize the genes found [19]. Further, the predicted
- genes were translated into amino acid sequences and aligned all-against-all using BLASTP [20].

 Genes were considered as part of a gene family if the alignment length was at least 80% of the longest sequence and had more than 80% in similarity (80/80 rule) [21]. A common core-pan- genome plot was constructed by comparing the gene families from all the genomes. The pan- genomes were constructed by the union of all the genes, while the core-genomes were assembled from the intersection of gene families shared by at least 90% of genomes [21, 22]. By using the perl script for comparative genomic analysis from the CMG-biotool [23], unique genes was determined by comparing the intersection of genes in one group with the ORFs of the other group.

 In the second attempt the genomes were, prior to the subjected analysis, divided into the two groups of good and poor recipients. ORFs, gene families, pan- and core-genomes were predicted for each group, as described above. To identify unique genes, the core-genome of one group was compared with pan-genome of the other group by using BLASTP and the 80/80 rule. This method will ignore orthologues. The resulting gene family candidates were further inspected by using BLASTP against the predicted ORF from each genome from the opposite group, to ensure that the predicted gene families would be unique.

 In addition to the analyses of all 93 *S. enterica* genomes, the 11 *S.* Enteritidis genomes spanning both groups (good and poor) were analysed separately, to search for serovar specific gene family candidates. For this analysis, the first approach was used, where ORFs, gene families, and core- and pan-genomes were constructed based on all 11 genomes.

Results

High throughput conjugational screening and grouping of isolates

 The high throughput conjugational approach was conducted to screen a high number of varying *S. enterica* isolates for their conjugational ability. This was done to identify recipients with poor versus enhanced ability to take up a plasmid. The plasmid pKJK5, encoding the green fluorescence gene *gfp,* was introduced to the *S. enterica* recipients via the red fluorescent tagged donor *E. coli* K- 12 MG1655 which represses *gfp*. Flow cytometry detection revealed transfer frequencies (TF) of 166 the 93 *S. enterica* recipients varying from 0.0 to 2.5×10^{-1} transconjugants per potential recipients,

illustrated in the boxplot in [Figure 2A](#page-135-0).

 From the TF mean for each isolate, the kmean algorithm was used to divide the isolates into two groups, resulting in a group of 10 good recipients, and a group of 83 poor recipients. The clustering based on the TF mean for each isolate was represented by a principal component analysis (PCA), illustrated in [Figure 3.](#page-136-0) Isolates from the group of good recipients were illustrated by the red triangles, and isolates from the poor group of recipients by blue circles. The two groups were located apart without overlap in the 2D representation in [Figure 3.](#page-136-0)

 Comparative genomics and identification of gene family candidates involved in conjugation By investigating the gene families found in the 83 bad recipients, absent in the 10 good recipients, and vice versa, we attempted to identify gene families involved in enhancing or restricting plasmid uptake. The pan- and core-genome of the 93 recipients were predicted to 22,161 gene families and 1,933 gene families, respectively. However, in the attempt to identify group dependent gene families, no families were identified by this approach.

 In the second attempt, separate core- and pan-genomes were constructed for the two groups. The core-genome (conserved genes) from one group of recipients was compared with the pan-genome

 (entire gene-pool including core-genome) from the other groups of recipients, to identify unique gene families only shared within one of the group. The pan- and core-genomes of the good recipient group were predicted to 8,733 and 3,249 gene families by prodigal with following all-against- alignment. For the group of poor recipients, the pan- and core-genomes were predicted to 20,458 and 2,052 gene families.

 By using the second approach to compare the gene families between the two groups of recipients, we identified 6 gene families from the good recipients, potentially encoding genes facilitating plasmid transfer. Additionally, 4 gene families were identified in the group of poor recipients, as gene candidates for resisting plasmid transfer. However surprisingly, when manually searching for the identified ORFs in the opposite group by using BLASTP, none of the gene family candidates were found to be unique for the groups.

 Additionally, to narrow down the diversity within the dataset and to look for serovar-specific gene candidates, the analysis was performed on a single serovar as well. For the 11 *S*. Enteritidis genomes included in the study, four genomes were predicted to belong to the good recipient group and seven genomes was predicted to belong to the poor recipient group. The TF for the 11 genomes were illustrated in [Figure 2B](#page-135-0), with the TF values for the 18 S. Typhimurim genomes in [Figure 2C](#page-135-0) for comparison. The common pan- and core-genomes were predicted to 6,149 and 3,736 gene families, respectively. The interception of gene families from one group, totally absent from the other group, revealed that the bad group was sharing 34 gene families not identified in the group of good recipients. Contrary, no unique gene families were identified from in the good group. To verify the findings, the 34 predicted proteins from the poor recipient group were manually searched against the four genomes from the good recipients. This revealed that one of the genes was found in one of the four genomes with an identity of 99.6%, and the remaining 33 was found with less than 66% identity at amino acid level. Thus, 33 gene family candidates, possible restricting conjugation,

 were identified in *S.* Enteritidis. The protein sequences of the 33 unique genes can be found in supplemental material (S9 File), together with the sequences of all the predicted pan- and core-genomes (S1 File - S8 File).

Discussion

 important in evolution of bacteria [2–4]. However, it is equally important in global emergence and rapid spread of undesirable genetic traits, like the spread of antibiotic resistance. A striking example is the recent finding of the plasmid-mediated colistin resistance gene, *mcr-1,* a resistance which until now only was mediated by chromosomal mutations with following vertical transmission to daughter cells [24, 25]. Hasman *et al.* detected the *mcr*-1 gene in a highly resistant *E. coli* isolate from a human bloodstream infection. The isolate was only susceptible to a limited number of antimicrobial classes, including the carbapenems. Additional acquisition of resistance will leave a 220 limited number of suitable treatment options [25], which underlines why understanding how genes are controlling the uptake and establishment of the plasmids in the recipients is of great importance. The objective of this study was to identify genetic traits of recipient strains, associated with uptake of IncP-1 plasmids, a mediator for spread of antimicrobial resistance. Based on transconjugant frequencies from a conjugational setup, the 93 *S. enterica* recipients clustered into two groups based on high and low transconjugant frequencies. By using different approaches for comparative genomics of the two groups, we initially identified 6 gene family candidates for enhancing conjugation, in addition to 4 gene family candidates restricting conjugation. However, further analysis with BLASTP against all predicted ORFs from the opposite group, revealed that none of the candidates were in fact unique to any of the two defined groups. The main dataset of 93 recipients was covering 54 different serovars, which is arguably a very diverse dataset and could have contributed to the lack of success to identify unique gene candidates. Thus, to decrease the genetic diversity, a single serovar was assessed separately. Assessing only the *S.* Enteritidis serovar, it was possible to identify 33 unique gene families in the genomes belonging to the low transfer

In recent time, it has become clear that the genetic spread through horizontal gene transfer is highly

frequency group, indicating the potential for serovar specific genes involved in the restriction of

 conjugational transfer. Adding additional *S.* Enteritidis serovars to the analysis could increase the serovar specific diversity and perhaps decrease the number of significant genes found in the analysis.

 The attempt to identify genes affecting recipient abilities in plasmid conjugation has previously been done by Pérez-Mendoza and de la Cruz [9]. Their study was performed with *E. coli* as recipient, and their strategy was based on high-throughput screening for systematic evaluation of individual *E. coli* genes in bacterial conjugation, where 20,000 random Tn-insertion mutants as well as the Keio collection of 3,908 individual deletion mutants, were screened. The attempt to identify recipient genes was focusing on two different K-12 strains, without knowing if genes affecting conjugation would be present at all [9]. They concluded that conjugation occur with little regard to the components of the recipient cells, which cannot avoid being used as recipient [9].

 In the present study, we tried to deal with the drawback of not knowing the level of conjugational ability of the recipients investigated, by performing an initial conjugation experiment of 93 *S. enterica* ssp. *enterica* isolates, with following grouping of good and poor recipients. Unfortunately, no gene families could be associated with either the good recipient- or poor recipient groups when assessing the entire *enterica* subtype of *Salmonella*. However, other analysis identifying unique SNPs could perhaps reveal an association not recorded with the utilized methods. Further, a comprehensive characterization including plasmid replicons, antimicrobial resistance, MLST profiles, etc. could as well guide the analysis in a direction or give an impression of where to search for differences that could cause the different recipient abilities.

 As mentioned above, an explanation could be the high complexity within the dataset, as the initial study was performed on a variety of *S. enterica* ssp. *enterica* isolates. To reduce the diversity, the 11 genomes of *S.* Enteritidis were assessed in an individual analysis, with the grouping defined

 from the analysis of the 93 genomes. However, when assessing the conjugational frequencies of *S.* Enteritidis, it became clear that it can be difficult to define an arbitrary threshold separating good recipients from poor recipients. It can be questioned if we need the threshold to identify genes involved in conjugation, or if a strain by strain comparison, taking the TF into account, would be more appropriate. Comparing the conjugational frequencies of *S.* Enteritidis and *S.* Typhimurium, the frequencies for the two serovars are not in the same range, however the same pattern of increase in frequencies can be observed within each of the two serovars [\(Figure 2\)](#page-135-0). Thus, a comparison of the gene content of the two serovars could be a subject for identification of gene-candidates. From the kmean algorithm none of the *S.* Typhimurium but four out of 11 *S.* Enteritidis isolates were determined as good recipients. Assessing the continuously increase between of the isolates illustrated in [Figure 2,](#page-135-0) this classification into good and poor recipients do not seem reasonable, as no distinct grouping is observed within neither of the two serovars. Additionally, the similar pattern for the two serovars, but different range of transfer, indicates that the ratio of conjugational transfer is very serovar specific. An explanation for this difference in range could be the individual sub- linage content of RM systems for the two serovars (unpublished data from Roer *et al*., 2015). However, this does not explain the difference observed within each serovar. Even though there is no clear grouping of good and poor recipients within a serovar, differences in recipient abilities can be observed which even appear significant between the two isolates with the highest and lowest frequencies. Comparing these results to the conclusion by Pérez-Mendoza and de la Cruz [9], it is not obvious that the bacteria cannot avoid being used as recipients, especially as some of the transfer frequencies in this study are recorded to zero and some are as high as 2.5×10^{-1} transconjugants per potential recipients, from the same donor and conjugation apparatus. The very high TF observed for some of the isolates is very unusual in conjugation, indicating that these isolates indeed have good recipient abilities. Thus, some recipient components must be involved in

 the recipient conjugational ability. As RM systems did not impose absolute barriers, other factors must be involved in the control of conjugation.

 The continuously increase in *S.* Enteritidis and *S.* Typhimurium could indicate that multiple genes are involved in determining the ability to take up plasmids, causing a synergy effect increasing or decreasing of the transfer frequency. However, it could also be different genes accounting for the differences in recipient abilities for each isolate, which probably could be answered by protein annotation and sequence alignment of the predicted proteins. Changing the arbitrary threshold for *S.* Enteritids could potentially reveal more or less gene family candidates. An interesting approach with this continuously increase in frequency could be to do multiple comparisons, where each "jump" is investigated in correspondence to the remaining genomes. Thus, for each

increase/decrease the responding gene could be predicted.

 Interestingly, only gene families predicted to be involved in reducing plasmid uptake were observed from this analysis. However, considering the fitness costs of harbouring plasmids, it is reasonable to think that bacteria in general develop methods for restricting plasmid uptake as a protection or defence mechanism, and not develop enhancing abilities.

 Playing the devil's advocate it could be argued that it is possible that the 33 genes only are predicted as unique due to the size difference between the two groups, as the poor recipient group contain seven genomes compared to the four in the good group. Thus, maybe the 33 gene family candidates are found due to natural diversity between the strains, and do not possess any impact on conjugational transfer. Therefore, the 33 gene family candidates predicted in this study should be further investigated, both by function prediction, prevalence of genes, and in molecular techniques in the laboratory.

 The current study was constructed by using a combination of molecular technologies and bioinformatics to predict gene family candidates involved in regulation of recipient abilities in

 conjugation. Though, some drawbacks from using bioinformatics in prediction of bacterial behaviour do exist. The prediction of gene families is based on the "80/80" rule. However a single nucleotide difference can cause alteration of the protein which in some cases can lead to functional inactivation without truncating the protein. An alternative approach could be to screen the bacterial genes for examining enhanced or restricted abilities to uptake plasmids. One example would be to perform random gene-knockout of a good recipient, with following identification of genes either decreasing or increasing the conjugational abilities, as performed in by Pérez-Mendoza and de la Cruz in *E. coli* [9]. However, in this case the conjugational ability would be known in advance. Another approach could be a transposon-library of the bacterial genome, which can be introduce to a recipient with the opposite conjugation ability, than the wildtype.

 In summary, the findings from this study indicate that differences in conjugation ability can be observed, both at species level and serovar level for *Salmonella,* with differences varying from 0.0 318 to 2.5×10^{-1} transconjugants per potential recipients. No common genes could yet be linked to enhanced or restricted conjugational abilities, when comparing a broad community of *Salmonella enterica* ssp. *enterica* isolates, but additional bioinformatic approaches are ongoing to explore this dataset further. With regards to *S.* Enteritidis, 33 gene family candidates could be predicted, which are being investigated and awaiting verification. Thus, the collection of genomes used in this study, together with the results from the initial conjugation experiment leaves great opportunities for revealing of genes involved in plasmid uptake.

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405 **Table 1 | Bacterial donor strain and plasmid used in this study.**

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407

408
409 409 **Figure 1** | **Conjugational screening detected by flow cytometry.** Flow cytometry detection of Donor, Recipients, and Transconjugants from a conjugation mixture of *S. enterica* and *E. coli K-12* MG1655 carrying pKJK5 410 Transconjugants from a conjugation mixture of *S. enterica* and *E. coli K-12* MG1655 carrying pKJK5. The gating procedure consists of five successive gates in the 3 different plots: Plot I with the Bacterial gate, det

411 procedure consists of five successive gates in the 3 different plots: Plot I with the Bacterial gate, detects all particles of bacterial size based on front and side scatter; Plot II detects all red fluorescence partic

412 bacterial size based on front and side scatter; Plot II detects all red fluorescence particles representing the donors, and all non-red particles representing either recipients or transconjugants by using side scatter

413 all non-red particles representing either recipients or transconjugants by using side scatter and PE-Texas Red; Plot III divides the Recipients and Transconjugants by gating non-green and green fluorescence particles b 414 divides the Recipients and Transconjugants by gating non-green and green fluorescence particles based on side scatter and FITC.

and FITC.

416

417 **Figure 2 | Conjugation frequencies for** *S. enterica* **ssp.** *enterica***. A,** Transconjugant frequencies of the initial 418 screening of recipients with good or poor uptake. The frequencies are represented as a box for each isolate, illustrating the results of 419 the results of the three repetitions. The frequencies are represented as a bo 419 the results of the three repetitions. The frequencies are represented as a box for each isolate, illustrating the results of 420 the three repetitions. **B**. Conjugation frequencies for the 11 S. Enteritidis isolates in the three repetitions. **B**, Conjugation frequencies for the 11 *S*. Enteritidis isolates included in the study. **C**, Conjugation frequencies for the 18 *S*. Typhimurium isolates included in the study. The isolates in B and 421 frequencies for the 18 *S*. Typhimurium isolates included in the study. The isolates in B and C are ordered according to their mean transfer frequency. their mean transfer frequency.

2D Representation of the Clustering

viability of the data.

Supporting information

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