Mosaic DNA Imports with Interspersions of Recipient Sequence after Natural Transformation of *Helicobacter pylori*

Stefan Kulick¹, Claudia Moccia¹, Xavier Didelot², Daniel Falush³, Christian Kraft¹*, Sebastian Suerbaum¹*

¹ Institute of Medical Microbiology and Hospital Epidemiology, Hannover Medical School, Hannover, Germany, ² Department of Statistics, University of Warwick, Coventry, United Kingdom, ³ Environmental Research Institute, University College Cork, Cork, Ireland

**Abstract**

*Helicobacter pylori* colonizes the gastric mucosa of half of the human population, causing gastritis, ulcers, and cancer. *H. pylori* is naturally competent for transformation by exogenous DNA, and recombination during mixed infections of one stomach with multiple *H. pylori* strains generates extensive allelic diversity. We developed an *in vitro* transformation protocol to study genomic imports after natural transformation of *H. pylori*. The mean length of imported fragments was dependent on the combination of donor and recipient strain and varied between 1294 bp and 3853 bp. In about 10% of recombinant clones, the imported fragments of donor DNA were interrupted by short interspersed sequences of the recipient (ISR) with a mean length of 82 bp. 18 candidate genes were inactivated in order to identify genes involved in the control of import length and generation of ISR. Inactivation of the antimutator glycosylase MutY increased the length of imports, but did not have a significant effect on ISR frequency. Overexpression of *mutY* strongly increased the frequency of ISR, indicating that MutY, while not indispensable for ISR formation, is part of at least one ISR-generating pathway. The formation of ISR in *H. pylori* increases allelic diversity, and contributes to the uniquely low linkage disequilibrium characteristic of this pathogen.

---

**Introduction**

*Helicobacter pylori* infects an estimated 50% of the world population, and can result in chronic gastritis, gastric or duodenal ulcers, gastric cancer, and MALT lymphoma [1]. Among the most striking characteristics of *H. pylori* are its amazing allelic diversity and variability. Not only do most all infected persons harbor their own unique strain(s), but *H. pylori* can undergo genetic change during chronic colonization. Such change is driven by the combination of frequent recombination between strains that simultaneously colonize one stomach, and an elevated mutation rate that is thought to be at least partly due to the lack of a MutHLS mismatch repair pathway (*H. pylori* possesses a MutS homolog, which, however, is not involved in mismatch repair [2], for reviews, see [3,4]).

*H. pylori* is naturally competent, and takes up exogenous DNA through an unusual transport system related to type IV secretion systems [5,6]. Population genetic analysis of *H. pylori* nucleotide sequences has provided strong evidence that recombination is far more common in *H. pylori* than in other bacteria [7,8]. More direct information about the effects of recombination on *H. pylori* comes from studies of sequential *H. pylori* isolates, cultured from biopsies taken from the same patient at time intervals of several months to years [9]. Numerous recombination events, many of them spanning only a few hundred base pairs, were detected when 10 gene fragments were sequenced for 24 pairs of such sequential isolates. Bayesian analysis revealed the most likely combination of recombination rate, mutation rate, and length of imported fragments that would generate the dataset. Strikingly, *H. pylori* cells that undergo recombination import unusually short pieces of DNA into their genomes (estimated mean of 417 bp), in contrast to other bacteria, where known lengths range from 2 kb (pneumococci, ref. [10]) to over 10 kb (*E. coli*, *Bacillus subtilis*, refs. [11,12]). However, despite ample evidence of the importance of recombination to the population structure of *H. pylori* [7,13,14], few studies have addressed basic questions of the recombination process. We have developed an *in vitro* transformation protocol that permits to study import events after natural transformation and to assess the role of candidate genes in determining import frequency and length. The data show that import of short fragments of DNA into the chromosome is an intrinsic property of the *H. pylori* DNA recombination/repair machinery. Imports can be interrupted by interspersed sequences of the recipient (ISR) of varying length which can accelerate allelic diversification. Overexpression of the
DNA glycosylase MutY increased the frequency of ISR, suggesting that MutY activity is involved in at least one pathway leading to the formation of ISR.

**Results**

**Short sequence imports after natural transformation of H. pylori**

In order to study the characteristics of import events after natural transformation of H. pylori, we transformed Rif sensitive H. pylori strains 26695, N6, or J99 with genomic DNA purified from Rif resistant H. pylori donor strains, J99-R3, 26695-R1, and N6-R1, and selected Rif resistant clones from these transformation experiments. In a parallel experiment without addition of DNA, the frequency of spontaneous mutations to Rif resistance was determined (Fig. 1A). The spontaneous mutation frequencies were $1.1 \times 10^{-6}$ (26695), $1.6 \times 10^{-6}$ (J99), and $4.6 \times 10^{-7}$ (N6), consistent with the results of earlier studies [15,16]. Addition of DNA from a Rif resistant donor strain to liquid cultures increased the frequency of Rif resistant clones by a factor of 22–199 depending on the recipient/donor combination, due to natural transformation (Table S4).

Rif resistant clones were characterized by nucleotide sequence analysis of a 1663 bp fragment of rpoB that included the positions conferring Rif resistance in the donors (Fig. 1B). The rpoB fragments of the three wild type strains differed at 3.7–5.5% of nucleotides. Due to this high degree of sequence divergence, even small imports created a mosaic sequence, and in many cases, both start and end point of DNA import events could be mapped with high accuracy by comparing the mosaic recombinant sequence with both donor and recipient sequences (Fig. 2). In other cases, the import extended beyond the sequenced fragment, providing only a minimum bound for the length of the import. We therefore applied a Bayesian model to the dataset in order to estimate the mean length of imports. Our estimates varied between 1294 bp (J99/26695-R1) and 3853 bp (N6/J99-R3), and the differences observed between some recipient/donor combinations were statistically significant (Table 1). The length of import events was thus longer than previously observed in sequential patient isolates (417 bp, ref. [9]), but still shorter than values reported for transformation in most other bacterial species.

**Interspersed sequences of the recipient (ISR) within imports**

When the rpoB sequences of Rif resistant transformants were compared with those of corresponding donor and recipient, we found that for $\sim$10% of the recombinant clones, the stretch of

![Figure 1. Schematic of the H. pylori in vitro transformation system used to study import events after natural transformation.](image-url)
integrated DNA from the Rif resistant donor was interrupted by one or multiple short fragments where the sequence was identical to that of the recipient. These interruptions of the imported fragment were termed ‘interspersed sequences of the recipient’ (ISR, Fig. 2). Some ISR included only a single polymorphic nucleotide, while others were up to 593 bp long, including up to 21 recipient-specific nucleotides. ISR could be observed in 8 out of 95 clones with DNA import in the recipient/donor combination 26695/J99-R3, 3 out of 26 in 26695/N6-R1, 4 out of 32 in J99/26695-R1, 5 out of 25 in N6/26695-R1, and 0 out of 80 clones in N6/J99-R3 (Table 1). The estimated mean length of ISR for the wild type strains varied between 38 and 121 bp (Table 1 and S7). However, there was no significant difference of ISR lengths between the analyzed recipient/donor combinations.

Short import length and ISR do not depend on DNA uptake via the ComB system

We hypothesized that the short length of imported fragments and/or the generation of ISR might be a consequence of the passage of exogenous DNA via the H. pylori-specific ComB DNA uptake system. To test this hypothesis, a comB10 mutant of H. pylori 26695 was constructed. After verification that this strain was unable to take up DNA by natural transformation, electroporation was used to introduce J99-R3 DNA into the comB10 mutant strain. 25 Rif resistant clones with DNA import were generated by this approach. The mean length of imports was not significantly different from the wild type, and 8 clones contained ISR, indicating that DNA transfer through the ComB system was not responsible for either the short length of imports, or the formation of ISR (Table 2). The frequency of ISR (8/25) was significantly higher in clones obtained by electroporation of the comB10 mutant than in clones isolated after natural transformation of 26695 wild type (8/95; Table 2).

Search for genes determining import length and ISR formation

The results from our in vitro transformation experiments suggested that importing short pieces of donor DNA into the chromosome was an inherent property of the H. pylori DNA recombination or DNA repair machinery. In order to identify genes involved in determining import length and ISR formation, we selected 18 genes with predicted roles in DNA repair or recombination (magIII, magII, mutS, mutL, nth, nucI, recA, recB, recG, recJ, recN, recR, rtaABC, ung, yxeA, and xth) and inactivated these genes in H. pylori 26695 by allelic disruption. Subsequently, all mutant strains were individually transformed with DNA from strain J99-R3. Mutation and recombination frequencies were measured, and the length of imported fragments and the frequency of ISR were determined by sequence analysis of rpoB in Rif resistant clones. Inactivation of mutI, nucI, nth, and ung significantly increased the rate of spontaneous mutations to Rif resistance, while disruption of recA, recB, rtaA, rtaB, and rtaC had the opposite effect. Inactivation of recA, recB, recN, nucI, rtaB, and rtaC significantly reduced the number of recombinant clones.

Table 1. Maximum likelihood estimation (MLE) of the mean length of donor DNA imports in the rpoB gene, number of clones with ISR in the imported DNA fragments and MLE of ISR length of H. pylori wild type strains.

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Donor</th>
<th>Num clones</th>
<th>MLE of import length (bp)</th>
<th>Bayes Factor</th>
<th>Clones with ISR</th>
<th>Bayes Factor</th>
<th>MLE of ISR (bp)</th>
<th>Bayes Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>26695</td>
<td>J99-R3</td>
<td>95</td>
<td>1681</td>
<td>39</td>
<td>8</td>
<td></td>
<td>39</td>
<td></td>
</tr>
<tr>
<td>J99</td>
<td>26695-R1</td>
<td>26</td>
<td>2434</td>
<td>0.29</td>
<td>3</td>
<td>0.19</td>
<td>121</td>
<td>1.20</td>
</tr>
<tr>
<td>N6</td>
<td>26695-R1</td>
<td>32</td>
<td>1294</td>
<td>0.19</td>
<td>4</td>
<td>0.20</td>
<td>38</td>
<td>0.24</td>
</tr>
<tr>
<td>J99-R3</td>
<td>N6</td>
<td>25</td>
<td>3819*</td>
<td>28.21</td>
<td>5</td>
<td>0.66</td>
<td>110</td>
<td>1.62</td>
</tr>
<tr>
<td>80</td>
<td>3853*</td>
<td>8</td>
<td>4.43 × 10^6</td>
<td>8</td>
<td>0.12</td>
<td>114</td>
<td>2.26</td>
<td></td>
</tr>
</tbody>
</table>

See Methods for interpretation of Bayes Factor values. **Strongly significant results (Bayes Factor ≥10)** are marked with an asterisk.

1 Number of clones with DNA imports in rpoB.

2 Approximated using the Bayesian Information Criterion (cf. Methods).

doi:10.1371/journal.pone.0003797.t001
while disruption of nth, recG, recR, and xseA increased it (Table S4).
However, out of 18 genes investigated, only one, mutY, had a
significant effect on import length and ISR formation.

Role of the DNA glycosylase mutY in import length and
ISR formation

The H. pylori antimutator gene mutY (HP0142) encodes a DNA
glycosylase involved in preventing specific C to A transversions
[16–19]. Due to this antimutator function of mutY, a 26695 mutY
mutant had a strongly increased frequency of spontaneous
mutation to Rif resistance. In order to test the impact of an
inactivation of mutY on length of import and ISR, a prescreening
procedure was applied to select recombinants from the Rif
resistant clones isolated from the transformation experiment. 43
clones with import were selected from a total of 250 colonies
screened, and characterized by p6B sequence analysis.

The mean length estimate of import events in the mutY mutant
was 3268 bp, almost twice that of observed in wild type bacteria.
This difference was statistically significant (Table 3). ISR were
detected in 10/43 (23%) of recombinant clones; this ISR
frequency was, however, not significantly different from wild type.

To rule out that the increased import length was caused by
polar effects of the insertion of the resistance cassette on genes
downstream of mutY, we next performed a functional complemen-
tation experiment by introducing into the chromosome of the
mutY mutant an intact copy of mutY while disruption of
ISR formation

Table 2. Import length and ISR formation are independent of the DNA uptake system ComB.

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Num clones</th>
<th>MLE of import length (bp)</th>
<th>Bayes Factor$^2$</th>
<th>Clones with ISR</th>
<th>Bayes Factor$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>26695</td>
<td>95</td>
<td>1681</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>26695comB10</td>
<td>0</td>
<td>-</td>
<td></td>
<td>8</td>
<td>1.00</td>
</tr>
<tr>
<td>26695comB10 EP$^3$</td>
<td>25</td>
<td>2228</td>
<td>0.15</td>
<td>8*</td>
<td>11.45</td>
</tr>
</tbody>
</table>

J99-R3 DNA was used as donor DNA for all transformations. See Methods for interpretation of Bayes Factor values. Strongly significant results (Bayes Factor ≥10) are marked with an asterisk.

1Number of analyzed clones with DNA imports.
2Approximated using the Bayesian Information Criterion (cf. Methods).
3EP: electroporation.
doi:10.1371/journal.pone.0003797.t002

Discussion

The extent of allelic diversity within H. pylori has puzzled
investigators since the late 1980s. Studies of population-wide
nucleotide sequence variation, sequence comparisons of multiple
strains isolated from a single biopsy, as well as comparisons of genetic
relationships of sequential H. pylori isolates provided evidence that
frequent recombination plays a central role in creating allelic
diversity in H. pylori [7,9,20]. Studies of sequential isolates have also
indicated that H. pylori imports short pieces of DNA with an
estimated mean length of 417 bp into its chromosome after natural
transformation [9]. However, systematic analyses of chromosomal
import events after natural transformation have not been performed in
H. pylori. In this study, we present extensive sequence data from a
total of 1090 recombinant H. pylori clones resulting from transfor-
mation experiments with different combinations of recipients and
donors, and investigated the roles of a large set of DNA
recombination and repair genes in determining the length of
imports after natural transformation.

Length of import events

The estimated mean length of imports after transformation of
H. pylori wild type strains ranged from 1294 to 3853 base pairs.

Table 3. Import length and clones with ISR in selected mutants of H. pylori 26695 transformed with DNA from J99-R3.

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Num clones</th>
<th>MLE of import length (bp)</th>
<th>Bayes Factor$^2$</th>
<th>Clones with ISR</th>
<th>Bayes Factor$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>26695</td>
<td>95</td>
<td>1681</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>26695magIII</td>
<td>51</td>
<td>1729</td>
<td>0.06</td>
<td>9</td>
<td>0.53</td>
</tr>
<tr>
<td>26695mutY</td>
<td>43</td>
<td>3268*</td>
<td>38.88</td>
<td>9</td>
<td>1.15</td>
</tr>
<tr>
<td>26695mutY comp</td>
<td>40</td>
<td>1882</td>
<td>0.08</td>
<td>25*</td>
<td>2.28 × 10$^4$</td>
</tr>
<tr>
<td>26695nth</td>
<td>53</td>
<td>1919</td>
<td>0.08</td>
<td>9</td>
<td>0.45</td>
</tr>
<tr>
<td>26695ung</td>
<td>56</td>
<td>2390</td>
<td>0.60</td>
<td>7</td>
<td>0.18</td>
</tr>
</tbody>
</table>

Full data for all 18 mutants examined are contained in Tables S5–6. See Methods for interpretation of Bayes Factor values. Strongly significant results (Bayes factor ≥10) are marked with an asterisk.

1Approximated using the Bayesian Information Criterion (cf. Methods).
doi:10.1371/journal.pone.0003797.t003

PLOS ONE | www.plosone.org | 4 November 2008 | Volume 3 | Issue 11 | e3797
While these values are still short in comparison with import length data reported from most other bacterial species, they exceeded those obtained by model-based analysis of sequences from sequential H. pylori isolates (417 bp, ref. [9]). These results indicate that import of short fragments after natural transformation is due to intrinsic properties of the H. pylori recombination/repair machinery. However, they also point at the existence of additional mechanisms contributing to the even shorter import events observed in vivo. For example, the moderately acidic environment of the natural ecological niche of H. pylori, a thin zone within the gastric mucus layer with a pH of ~5.5–6 [21], may damage naked double stranded DNA (e.g. by depurination) before it is taken up by the recipient cell. This and other hypotheses are currently under investigation in our laboratory.

The length of imported fragments varied significantly between different combinations of donor and recipient. We first hypothesized that these differences might have been a consequence of varying levels of sequence similarity between the rpoB alleles (sequence identity ranged between 96.2% and 94.6% for the combinations of rpoB tested). However, our results argue strongly against this hypothesis, because in some cases, import lengths were significantly different when combinations of donor and recipient were reversed (e.g. N6/26695-R1 and 26695/N6-R1). Therefore, the mechanisms responsible for the differences of import length remain unknown, although our data suggest that import length is mainly determined by properties inherent to the recipient strains, such as their content of restriction modification systems. The distance of the import border to the central point mutation followed a geometric distribution, yielding no evidence that the location of the starting/end point of the import depended on specific sequence motifs.

Since H. pylori imports DNA through an unusual type IV secretion system (ComB, ref. [5]), we hypothesized that this machinery might be involved in generating short import events. However, this also seems unlikely, because the import length did not change when DNA was introduced into a ComB-deficient mutant by electroporation.

Interspersed sequences of the recipient (ISR)

One striking feature of the imports we observed in H. pylori was the occurrence of short stretches of recipient sequence interspersed within the imported regions. The mean length of ISR was 82 bp. While these disruptions of the imported alleles were observed only in ~10% of clones, ISR are likely to have an impact on allelic diversity, because they have the potential to significantly decrease linkage between individual polymorphic sites. H. pylori is characterized by a uniquely low linkage disequilibrium [7], and ISR are likely to contribute strongly to this allelic diversity, together with an overall high rate of recombination and a relatively short length of imported fragments.

The formation of complex mosaic alleles has previously been observed after transformation or transduction of homologous DNA into E. coli. Abastado et al. obtained multiple different sequences with “patchwork structure” after transformation of E. coli with plasmids containing a heteroduplex [22,23]. These mosaics were obtained in wild type as well as in recA-deficient recipients, suggesting that their formation resulted from repair processes rather than recombination. Since mosaicism was detected with oligonucleotide probes, detailed analysis of the imported sequences could not be performed at that time. McKane and Milkmann used restriction fragment polymorphisms to characterize import events after transduction of ~100 kb of DNA into E. coli, reporting that import frequently occurred in multiple fragments with a mean size of 8–14 kb. Similar to our data, fragment lengths were dependent upon the recipient strain used, and this was thought to be due to differences in the restriction modification systems [11]. To our knowledge, the present study is the first to use nucleotide sequencing of a large number of recombinant clones to analyze import patterns resulting from transformation. It is thus not possible to conclude with certainty whether ISR similar to those observed with H. pylori occur in other species, and at which frequency.

Formation of ISR might occur randomly at any position within the transformed fragment, or depend on the occurrence of specific sequence motifs, such as restriction sites. We tested the existence of such associations for 36 restriction sites within the rpoB sequence, and found no significant association between restriction sites in the rpoB fragment and the occurrence of ISR (data not shown). However, due to the relatively short length of the sequenced fragment, this analysis does not rule out the existence of such associations. Many restriction sites are present only once or twice in the sequenced fragment, making our dataset lowly informative about the association of ISR with particular sequence motifs.
We note that there was a significant increase in the number of clones with ISR when DNA was introduced by electroporation. Whether this is due to an increase of DNA repair activities after electroporation, damage to DNA generated by the pulse, the non-selective process by which the DNA is introduced into the cell by electroporation, or yet other mechanisms remains to be investigated.

To address the question whether ISR occur in vivo, we reanalyzed mosaic nucleotide sequences from sequential H. pylori isolates that were previously shown to have undergone recombination in vivo [9]. These paired sequences contain multiple events where blocks of sequence polymorphisms are interrupted by stretches of identical sequence (e.g., the atpA alleles of the pair NQ352). These patterns likely represent ISR, but it is impossible to prove this with certainty, because only recipient and recombinant sequences are known, whereas the donor sequence is not available, which would also be required to establish that the mosaic patterns are indeed ISR.

A screen for genes controlling import length and ISR: the role of mutY

In order to identify genes that are involved in the control of import length and ISR formation, 18 candidate genes were selected. Mutants in these genes were constructed by allelic disruption and characterized with respect to import length and ISR, as well as mutation and recombinant frequencies (Tables S4, S5, S6, S7). This screen identified only one gene, mutY, whose inactivation affected import length and whose overexpression increased ISR frequency.

The DNA glycosylase MutY is part of the base excision repair (BER) pathway and has a strong antimutator function in H. pylori [16,18,19,24]. Unexpectedly, a mutY mutant exhibited a significantly increased import length compared with the wild type strain, while the frequency of ISR was not significantly affected. A functional complementation experiment was performed to rule out polar effects. Since the intact copy of mutY introduced in this strain is under control of a strong promoter [19], the complemented strain contains more mutY transcript than the wild type strain (Fig. S1). This complementation reduced the import length back to levels not distinguishable from wild type. Strikingly, it also caused a strong and highly significant increase in the number of ISR, suggesting that the MutY enzyme may be involved in initiating ISR formation after transformation. An involvement of MutY in recombination has been demonstrated before, albeit in a very different model system [25]. A hypothetical model of MutY-mediated formation of a short ISR is shown in Fig. 4. The inactivation of three other predicted BER glycosylase genes, magIII, ung, and nth, had no significant effect on the import length, and ISR were still observed in these mutants, indicating that neither of them is essential for ISR formation. We note that all single and double mutants studied were still capable of generating ISR. This may suggest that ISR formation can occur via multiple partially redundant repair pathways, and complete abrogation of ISR might require inactivation of multiple repair genes.

Conclusion

The combination in H. pylori of competence for natural transformation, and high degree of sequence divergence between naturally occurring alleles of spoB has enabled us to perform an in depth study of import events after transformation, and to identify the formation of mosaic sequences with interspersed recipient fragments as a mechanism likely contributing to the uniquely high allelic diversity in H. pylori. The experimental setup has also permitted to identify the glycosylase MutY as one gene involved in the control of import length and initiation of ISR formation. Our system will provide a powerful tool for further studies of recombination and possibly gene conversion in H. pylori, and should also be applicable to other transformable bacteria where many fundamental questions about the effect of recombination on the chromosome remain to be addressed.

Materials and Methods

Bacterial strains and culture conditions

Bacterial strains used in this study are listed in supporting information [SI] table S1 (references cited in supporting tables S1, S2, S3 are listed in Supplementary References S1). Helicobacter pylori wild type strains 26695 [26], J99 [27], and N6 [28] were cultured from frozen stocks on blood agar plates (Columbia agar base II, Oxoid, Wesel, Germany) containing 10% horse blood and antibiotics (vancomycin [10 μg/ml], polymyxin B [2500 U/ml], amphotericin B [4 μg/ml], and trimethoprim [5 μg/ml]). Plates were kept in an incubator with 5% O2, 10% CO2 and 85% N2 at 37°C for 24 h - 48 h. Mutant strains were cultivated on blood agar plates containing kanamycin (20 μg/ml), chloramphenicol (20 μg/ml), or both antibiotics as required. For selection of rifampicin (Rif) resistant H. pylori donor strains, 106 wild type bacteria were plated
on blood agar plates supplemented with rifampicin (10 µg/ml). Liquid cultures were performed in brain heart infusion [BHI, Oxoid] media with yeast extract (2.5 g/l), 10% heat inactivated horse serum and antibiotics (see above) in microaerobic atmosphere using anaerobic jars (Oxoid) and Anaerochult® C gas generating bags (Merck).

E. coli strains DH5α [29] and MC1061 [30] were used for the DNA cloning experiments and were grown in LB broth or on LB plates (Lennox L Broth, Invitrogen GmbH, Karlsruhe, Germany) supplemented with ampicillin (200 µg/ml), chloramphenicol (20 µg/ml) and/or kanamycin (20 µg/ml) as required.

DNA techniques
All standard cloning and DNA amplification, purification and manipulation procedures were done according to standard protocols [31]. Preparation of genomic DNA was performed using the QIAamp DNA Minikit (QIAGEN, Hilden, Germany).

Large-scale purification of DNA was performed using QIAGEN Genomic-tip 100/G columns. Plasmid DNA from E. coli strains were isolated using QIAGEN tip 100 columns.

Insertion mutagenesis in H. pylori
Construction of mutants by natural transformation mediated allelic exchange was performed as described previously [28,32]. A list of the oligonucleotides used for mutagenesis, with introduced restriction sites is provided in Table S2. Briefly, the target genes were amplified by PCR and cloned into pUC18. The resulting plasmids (Table S3) were used for inverse PCR amplification with primers containing restriction sites. Inverse PCR reactions were designed in order to delete a part of the target gene and to introduce a unique restriction site. The PCR products were subsequently digested with the appropriate restriction enzyme, and ligated with an antibiotic resistance cassette (aphA3-III or cat, refs. [33,34] flanked by compatible restriction sites. The direction of transcription of the antibiotic resistance gene was the same as that of the target gene to avoid possible polar effects caused by the strong promoters of the antibiotic resistance cassettes.

Plasmids containing the interrupted gene were used as suicide plasmids in natural transformations of the H. pylori strains 26695, J99, or N6. The successful chromosomal replacement of the intact target gene with the disrupted gene construct via allelic exchange (double crossover) was checked by PCR using suitable primer combinations. For mutT, whose inactivation caused significant changes of the length of imported fragments, we performed a functional complementation experiment.

In vitro transformation system of H. pylori
The transformation of H. pylori recipient strains with genomic DNA from Rif resistant donors was performed in 10 ml BHI including yeast extract (2.5 g/l), 10% horse serum and antibiotics (vancomycin [10 µg/l], polymyxin B [2500 U/ml], amphotericin B [4 µg/ml] and trimethoprim [5 mg/ml]).

H. pylori wild type or mutant strains were grown for 24 h on non selective (for wild type strains) or selective (kanamycin and/or chloramphenicol as required for each knockout mutant) blood agar plates, as described above and harvested in 1.5 ml BHI broth. For strains J99, 26695 and N6, a suspension with an OD600 of 1 contains approx. 3 x 10⁹ bacteria [16]. The suspension was diluted to a start concentration of 2.1 x 10⁶ bacteria/ml in a final volume of 10 ml. For cultures containing knockout mutant strains, kanamycin and/or chloramphenicol (each at 20 µg/ml) were added as required. The samples were then incubated for 20–24 h at 37°C in a rotary shaker (175 rpm) under microaerobic conditions using Anaerochult C bags (Merck). The bacterial concentration of the preparatory culture was measured and the suspension was used to start the experiment. Three 10 ml cultures per strain were adjusted to 1.5 x 10⁷ bacteria/ml and cultivated for 16 h as described above to achieve a concentration of approx. 5 x 10⁹ bacteria/ml. At this point, donor DNA (1 µg/ml) was added to two of the three cultures. All three cultures were then further incubated for 8 h. The OD₆₀₀ was measured and for each culture incubated with donor DNA, inocula corresponding to 10⁸, 10⁹, and 10¹⁰ bacteria were plated on selective blood agar plates containing rifampicin (10 µg/ml) and cultivated for 5 days at 37°C. In parallel, a volume corresponding to 10⁸ bacteria from the recipient cultures cultivated without donor DNA were plated on selective Rif containing blood agar plates to determine the frequency of spontaneous Rif resistant mutants.

From each experiment, at least 16 clones were expanded in order to sequence a fragment (1663 bp) of the rpoB gene (see below). These experiments were reproduced at least 10 times for each H. pylori wild type strain and 3 times for each H. pylori mutant strain.

Electroporation of H. pylori
For electroporation experiments, bacteria were grown to a density of 3 x 10⁶ cells/ml in 10 ml liquid cultures as described above, centrifuged and suspended in 10% glycerol (1 ml). 90 µl of the suspension was mixed with 10 µl of DNA in prechilled 0.2 cm electroporation cuvettes. Electroporation was performed at 2.5 kV, 25 µF and 400 Ω in a Gene Pulser (Biorad). After electroporation, the bacteria were added to 2 ml BHI media with 10% horse serum and yeast extract and incubated for 8 h. Plating and counting were done as described above.

Partial sequence analysis of rpoB from H. pylori transformants
To determine the position of point mutations or import events in rpoB, a 2370 bp PCR fragment of rpoB was amplified with primers HPrpoB-1 and HPrpoB-6. This PCR product was used as the template for sequencing reactions with the primers HPrpoB-3, -4, -5, -6, -9w, and -10. Sequence reactions were done using the BigDye Terminator Cycle Sequencing Kit v1.1 (Applied Biosystems, Foster City, USA), and analyzed with the ABI 3130xl genetic analyzer (Applied Biosystems). The six sequences from each rifampicin resistant clone were assembled using the software Bionumerics V 4.5 (Applied Maths, Sint-Martens-Latem, Belgium), yielding a continuous, double-stranded 1663 bp fragment of rpoB that included the Rif resistance mediating point mutation of the donor strains.

PCR-based prescreening for clones with DNA imports in the strains 26695mutT and 26695rufV
Since it was difficult to find DNA imports in the strain 26695mutT due to the high background of spontaneous mutations and extraordinary low recombination levels in 26695rufV, it was necessary to screen the Rif resistant clones after transformation by an import specific PCR using the primers HPrpoB-IsrX and HPrpoB-I. Primer HPrpoB-IsrX, specific for the Rif resistance mediating point mutation in strain J99-R3, was designed according to the method described by Furuta et al. [35]. PCR positive clones were subsequently used for sequencing as described above.

Semi-quantitative reverse transcriptase PCR
Semi-quantitative RT-PCR was performed essentially as described previously [36]. Briefly, RNA was prepared from H. pylori strains grown in liquid culture to a density of 3 x 10⁹ bacteria/ml using the Qiagen RNeasy kit. Reverse transcription was performed using 2 µg of DNase I treated RNA, a random
hexamer primer mix and superscript III™ reverse transcriptase (Invitrogen, Karlsruhe, Germany) at 42°C for 1 h. Semi-
quantitative PCR of mutF was done with 2.5 μl cDNA in a total volume of 25 μl using the primers HPmutY1979 and HPmutY1980. To compare total amounts of RNA for different samples, a fragment of the 16S rRNA was amplified with the primer pair C05 and C97.

**Statistical methodology**

Statistical analysis was performed by Bayesian model comparison, where two competing models are compared using the Bayes Factor (BF) which is equal to the ratio of probabilities of observing the data under each model (see refs. [37,38] for reviews). This has the advantage to directly weight the evidence for two competing hypothesis, taking into account their relative complexity, so that over-complicated models are naturally penalized. Interpretation of the Bayes Factor was done following the guidelines of Jeffreys [39]: Negative ($<1$); Barely worth mentioning ($1<3$); Substantial ($3-10$); Strong ($10-30$); Very strong ($30-<100$); Decisive ($\geq 100$).

When the Bayes Factor could not be explicitly calculated, a conservative estimate was computed using the Bayes Information Criterion (BIC; refs. [37,40])

$$BF \approx \exp \left( \frac{BIC_1 - BIC_2}{2} \right) = \exp \left( l_2 - l_1 + \log(n) \frac{k_1 - k_2}{2} \right)$$

(1)

where $l_1$ and $l_2$ are the maximized value of the log-likelihood under the two models, $k_1$ and $k_2$ the number of free parameters in the two models, and $n$ the number of observations. Comparisons of frequency data between any two recipient/donor combinations were done using the BIC for two models: one where the data from the two combinations comes from the same Normal distribution and one where it comes from two distinct Normal distributions.

Import length is often modelled using a geometric distribution [9,13,41]. Using the BIC, we showed that this is a better model than a negative binomial model, despite its additional free parameter. We do not observe exactly where recombination starts and ends for an observation $i$, but instead an interval $[m_i; M_i]$, where $m_i$ might happen with $M_i = \infty$ if the beginning or end is out of the sequenced region), so that the likelihood of $N$ observations under the geometric model with mean length $\delta$ is equal to:

$$L(\delta) = \prod_{i=1}^{N} \left( \frac{1 - \frac{1}{\delta}}{\frac{1}{\delta}} \right)^{m_i} \left( 1 - \frac{1}{\delta} \right)^{M_i}$$

(2)

Lengths of imports on either side of the resistance points for each recipient/donor combination were compared using the BIC with one model where left and right come from the same geometric distribution and one model where they do not. We found no evidence for a difference in length on left and right. The effect of gene knock-outs on the lengths of import was evaluated in the same way (comparing results between two combinations rather than between left and right of a single combination).

Let $p$ denote the probability of occurrence of ISR in a sequenced clone. The number $m$ of clones containing ISR amongst $n$ sequenced clones is thus distributed as Binomial($n,p$). Assuming a Jeffrey’s prior on $p$ (i.e. Beta($\frac{1}{2}, \frac{1}{2}$)), we compared the value of $p$ between two recipient/donor combinations ($m_1, n_1$ and $m_2, n_2$) using the Bayes Factor:

$$BF = \frac{B(m_1 + 1/2, n_1 - m_1 + 1/2) B(m_2 + 1/2, n_2 - m_2 + 1/2)}{B(m_1 + m_2 + 1/2, n_1 + n_2 - m_1 - m_2 + 1/2)}$$

(3)

where $B(.,.)$ denotes the Euler Beta function.

**Supporting Information**

- **Table S1** Bacterial strains used in this study
  Found at: doi:10.1371/journal.pone.0003797.s001 (0.05 MB PDF)
- **Table S2** Oligonucleotide primers and PCR products used in this study
  Found at: doi:10.1371/journal.pone.0003797.s002 (0.06 MB PDF)
- **Table S3** Plasmids used in this study
  Found at: doi:10.1371/journal.pone.0003797.s003 (0.02 MB PDF)
- **Table S4** Mutation and recombination frequencies leading to Rif resistance in *H. pylori* wild type strains and mutants
  Found at: doi:10.1371/journal.pone.0003797.s004 (0.03 MB PDF)
- **Table S5** Maximum likelihood estimation (MLE) of the mean length of DNA imports left and right from the Rif resistance mediating mutation.
  Found at: doi:10.1371/journal.pone.0003797.s005 (0.02 MB PDF)
- **Table S6** Clones with ISR observed in the analyzed *H. pylori* wild type and mutant strains.
  Found at: doi:10.1371/journal.pone.0003797.s006 (0.02 MB PDF)
- **Table S7** Maximum likelihood estimation (MLE) of the mean length of ISR in the analyzed *H. pylori* wild type and mutant strains
  Found at: doi:10.1371/journal.pone.0003797.s007 (0.02 MB PDF)

**Supplementary References S1** References for supplementary material
Found at: doi:10.1371/journal.pone.0003797.s008 (0.04 MB DOC)

**Figure S1** Semi-quantitative RT-PCR analyses of the abundance of mutF mRNA in the *H. pylori* strains 26695 and 26695mutYcomp.
Found at: doi:10.1371/journal.pone.0003797.s009 (0.12 MB PDF)

**Acknowledgments**

The authors thank Christine Josenhans for helpful discussions, Martin Blaser for plasmid pADC-HpMutY, and Kerstin Ellrott for excellent technical assistance.

**Author Contributions**

Conceived and designed the experiments: SK CM CK SS. Performed the experiments: SK CM XD DF CK SS. Wrote the paper: SK CM XD SS.
References