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Natural Transformation of Campylobacter jejuni Occurs Beyond Limits of Growth

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Abstract

Campylobacter jejuni is a human bacterial pathogen. While poultry is considered to be a major source of food borne campylobacteriosis, C. jejuni is frequently found in the external environment, and water is another well-known source of human infections. Natural transformation is considered to be one of the main mechanisms for mediating transfer of genetic material and evolution of the organism. Given the diverse habitats of C. jejuni we set out to examine how environmental conditions and physiological processes affect natural transformation of C. jejuni. We show that the efficiency of transformation is correlated to the growth conditions, but more importantly that transformation occurs at growth-restrictive conditions as well as in the late stationary phase; hence revealing that growth per se is not required for C. jejuni to be competent. Yet, natural transformation of C. jejuni is an energy dependent process, that occurs in the absence of transcription but requires an active translational machinery. Moreover, we show the ATP dependent ClpP protease to be important for transformation, which possibly could be associated with reduced protein glycosylation in the ClpP mutant. In contrast, competence of C. jejuni was neither found to be involved in DNA repair following DNA damage nor to provide a growth benefit. Kinetic studies revealed that several transformation events occur per cell cycle indicating that natural transformation of C. jejuni is a highly efficient process. Thus, our findings suggest that horizontal gene transfer by natural transformation takes place in various habitats occupied by C. jejuni.

Introduction

The Gram-negative organism Campylobacter jejuni is the primary food borne bacterial pathogen in the developed world causing millions of gastroenteritis cases each year. Infections with C. jejuni typically results in acute illness with clinical symptoms ranging from mild to severe mucoid and bloody diarrhoea, and in rare cases the serious neurodegenerative Guillain Barre syndrome [1]. C. jejuni is a zoonotic pathogen and domestic poultry is an important reservoir, as consumption and handling of contaminated meat products is one of the predominant routes of human infections [2]. C. jejuni is highly adapted to the avian gastrointestinal environment and it colonizes the gut of chickens to very high cell numbers without causing significant symptoms in the birds [3,4]. C. jejuni is a fastidious organism, as it only grows under microaerobic conditions in a narrow temperature interval of approximately 30–46°C [5,6]. In addition to a significant degree of host adaptation C. jejuni survives well in the external environment and contaminated water has been the cause of many cases of human gastroenteritis [7,8,9,10].

The adaptive capacity of C. jejuni is in part attributed to a remarkable degree of genetic diversity [11,12]. Studies have revealed substantial variation in genes encoding carbohydrate surface structures, flagella proteins and respiratory enzymes, which contribute to the virulence potential of C. jejuni upon infection but moreover also to survival in the external environment [13,14,15,16,17,18]. The genetic diversity is caused by genomic rearrangements and hyper variable sequences but very importantly also by exchange of DNA between strains [19,20,21,22]. C. jejuni is naturally competent for uptake of DNA from the environment and horizontal gene transfer has been shown to occur efficiently both in vitro and in vivo [21,23].

Competence for natural transformation has been described for bacteria from a wide range of taxonomic groups [24], but the mechanisms of natural transformation have only been investigated in detail for a limited number of species. For most competent bacteria, exemplified by Bacillus subtilis, Streptococcus pneumoniae and Haemophilus influenzae, the development of competence is limited to a transient physiological state, tightly regulated by growth conditions and/or signal peptides [25,26,27], while other organisms, such as Neisseria gonorrhoeae and Helicobacter pylori, seem to be constitutively competent [28,29].

Natural transformation by C. jejuni is far less characterized and improved knowledge of natural transformation of C. jejuni will therefore facilitate in understanding sources of strain diversity and moreover improve the tools for molecular investigations of this organism. It has been shown that the transformation frequency of C. jejuni is highly affected by the carbon dioxide concentration and to a minor extent by the growth phase [23,30], while the levels of oxygen and temperature are reported to influence natural transformation of C. coli [31]. Several genes are required for natural transformation of C. jejuni, such as Cj0011c, ctsD (Cj1474c), ctsE (Cj1471c), ctsF (Cj1470), ctsG (Cj1343c), and Cj1211. They
encode proteins with homology to a DNA receptor, type II secretion systems and/or pseudopilus proteins of other competent bacteria, respectively, and are therefore expected to be involved in uptake and transport of DNA in C. jejuni [32,33,34]. Likewise, the N-linked protein glycosylation (pgl) system is important for competence of C. jejuni [33].

To explore competence of C. jejuni in relation to environmental survival and adaptation, we have investigated the efficacy of natural transformation under conditions both restrictive and permissive for growth. Furthermore, we examined the significance of the Clp protease and central cellular processes on competence of C. jejuni to determine the boundaries of natural transformation. Our findings suggest that natural transformation occurs under a variety of environmental conditions and that the process may be important for the organism in multiple milieus.

Results

Natural Transformation of C. jejuni is Constitutive and Highly Efficient

For studying natural transformation of C. jejuni NCTC11168 we employed a previously described biphasic media system [23] composed of a solid agar layer and liquid medium layer with exponentially growing bacteria placed at 37 °C under microaerobic conditions. As selective markers for DNA uptake and genomic integration, we routinely used genomic DNA carrying either an rpsLSm mutation, providing resistance to streptomycin, or a CamR marker inserted into tlp1, a chromosomal locus not conferring growth defects in vitro [36].

The efficiency of natural transformation was evaluated both relative to the number of bacterial cells and the amount of added genomic DNA. When excess amounts of chromosomal CamR DNA was used in the transformation assay (CFU:DNA genome molecules = ca. 1:10), 0.36% of the total CFUs obtained the resistance marker (Table 1). By limiting the amount of chromosomal DNA (CFU:DNA genome molecules = ca. 10:1) the transformation efficiency could furthermore be evaluated relative to the amount of DNA, which showed 0.54% of the added DNA molecules to give rise to CamR transformants (Table 1). The number of CamR transformants represents cells having obtained a chromosomal region carrying the resistance marker, and these measured CamR transformants are only expected to constitute a minor fraction, while the actual number of transformed cells and the amount of DNA picked up from the environment very well could be substantially higher.

The specificity of natural transformation by C. jejuni NCTC11168 was examined by transforming with DNA of different origin. C. jejuni NCTC11168 was readily transformed with chromosomal DNA from StrepR mutants of two C. jejuni strains, DVI-SC11 and DVI-SC183 isolated from Danish chickens [37] (results not shown). In contrast, the addition of increasing amounts of DNA from the closely related species Helicobacter pylori and Arcobacter butzleri could not outcompete the transformation of isogenic DNA (Table S1). This is not a comprehensive experiment, but these data suggest natural transformation of C. jejuni NCTC11168 to be species- but not strain-specific.

To examine the kinetics of natural transformation in more detail, we investigated if several transformation events occur per cell cycle by transforming simultaneously with chromosomal DNA encoding either CamR or StrepR resistance. Aliquots of an exponentially growing C. jejuni NCTC11168 culture were distributed to 13 samples and each sample equally transformed with 2 μg/ml of both CamR DNA and StrepR DNA. Individual transformations were stopped at various time points (0–120 min.) by the addition of DNase I, and following 2 h of phenotypic expression single and double transformants were scored by plating on selective agar. CamR or StrepR transformants was obtained already after a few minutes of exposure to DNA and a saturated level of transformants, i.e. a stable number of resistant colonies, was obtained after approximately 30 min of co-cultivation with DNA (Figure 1), indicating that single DNA uptake had occurred. The time of transformation saturation was the same for both resistance markers although differences were observed in the number of transformants/ml. The generation time of C. jejuni NCTC11168 is approximately 2 h under the experimental conditions (results not shown), and thus the saturation level of single transformants was obtained already at a quarter of a generation. The appearance of transformants carrying both resistance markers, i.e. bacterial cells have taken up and integrated multiple DNA fragments, occurred after approximately 1.5 h, which is less than a generation, and the fraction of these double transformants increased steadily up to the 2 h time limit of the experiment (Figure 1). These results show that DNA uptake and natural transformation of C. jejuni is a continuous and fast process and that multiple transformation events can occur within a cell cycle.

Quantification of Extracellular DNA in C. jejuni NCTC11168 Cultures

In order to determine the concentration of extracellular DNA in C. jejuni cultures, a fluorescent nucleic acid stain was applied to quantitatively assess the amount of double stranded DNA in the culture supernatant of C. jejuni NCTC11168 Δtlp1. Thus, we found approximately 0.04 μg/ml DNA to be present in the supernatants of early stationary cultures (OD600 >1.0 ≥ ca. 2.5×106 CFU/ml) in MEM without minimal media supplemented with 20 mM sodium pyruvate and 40 μM FeSO4. The result represents the difference between treated and untreated DNase I samples and is corrected

| Table 1. Natural transformation efficiency of C. jejuni NCTC11168. |
|---|
| CFU:DNA | CFU/ml | DNA ng/ml | Transf./CFU | Transf./DNA molecule |
| ca. 1:10 | 3.0×10<sup>2</sup> ± 2.9×10<sup>6</sup> | 6500 ng = 3.5×10<sup>6</sup> DNA molecules | 3.6×10<sup>-3</sup> ± 1.8×10<sup>-3</sup> | -- |
| ca. 1:10 | 3.6×10<sup>2</sup> ± 2.6×10<sup>7</sup> | 65 ng = 3.5×10<sup>6</sup> DNA molecules | -- | 5.4×10<sup>-3</sup> ± 1.0×10<sup>-3</sup> |

*Results are mean of three replicates with standard deviation.

<sup>a</sup>Chromosomal DNA from C. jejuni NCTC11168 Δtlp1::CamR.

<sup>b</sup>Limits of range tested: 10<sup>4</sup>–10<sup>10</sup> transformants/CFU.

<sup>c</sup>Limits of range tested: 10<sup>-2</sup>–10<sup>-1</sup> transformants/DNA molecule.

<sup>d</sup>Excess amounts of DNA.

<sup>e</sup>Limited amounts of DNA.

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for background fluorescence from the media and DNase I. The measured extracellular DNA concentration was furthermore confirmed in natural transformation assays using the culture supernatants as DNA source (results not shown). The measured extracellular DNA concentration corresponds to approximately one molecule of a \textit{C. jejuni} genome per 100 CFU under the investigated conditions.

\textbf{C. jejuni is Competent Under Growth-restrictive Conditions}

We observed \textit{C. jejuni} NCTC11168 to be competent in both exponential and late stationary growth phase, although the number of transformants per CFU was approximately 400 times lower for bacterial cells in late stationary phase compared to cells in the exponential growth phase (Table 2). This observation prompted us to investigate if \textit{C. jejuni} is competent under conditions not supporting growth. Growth of \textit{C. jejuni} is restricted to conditions mimicking the environment of the gastrointestinal tract, i.e. microaerobic conditions and temperatures between 30–46°C [5,6]. Natural transformations were carried out and compared for various conditions inside and outside this window of growth (Table 3). The level of natural transformation was highest under microaerobic conditions at 42°C, which reflects the conditions of the avian gut. However, transformation was found to occur at all tested conditions although at highly variable degree. Generally, the temperature had a much greater influence on the transformation than the concentration of oxygen, since decreased temperatures caused a more severe reduction of the transformation than did changing the oxygen concentration from microaerobic to aerobic conditions. Noteworthy, the level of natural transformation of \textit{C. jejuni} was almost equal at aerobic and microaerobic conditions at 20, 30 and 37°C, respectively. On the other hand, an anaerobic environment was significantly less supportive of natural transformation as compared to microaerobic conditions (Table 3).

To further explore the effect of growth restrictive conditions we investigated natural transformation and survival of \textit{C. jejuni} at variable pH (Figure 2). From pH 5 to ca. 8.5 more than 50% of bacterial cells survived within the 4 h experiment, while the fraction of viable cells rapidly dropped outside this pH interval. In contrast, natural transformation occurred very efficiently in the pH interval 7 to 11 with a relatively constant level of viable CFU’s obtaining the resistance marker, while at pH conditions outside this interval the number of transformants per CFU was reduced by 10–10^5 fold (Figure 2). From these results it is clear that the pH interval for natural transformation is different from the pH interval supporting growth, since higher pH correlates with lower survival but higher number of transformants.

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|}
\hline
\textbf{Growth phase} & \textbf{OD600} & \textbf{Aggregates}\(^a\) & \textbf{Transformants/CFU}\(^b\) \\
\hline
\textbf{Exponential} & 0.05 & \begin{tabular}{c}
4.3 \times 10^8 \pm 7.4 \times 10^6
\end{tabular} & \begin{tabular}{c}
2.3 \times 10^2 \pm 3.2 \times 10^{-4}
\end{tabular} \\
\textbf{Late stationary} & 2.2 & \begin{tabular}{c}
4.3 \times 10^8 \pm 7.6 \times 10^7
\end{tabular} & \begin{tabular}{c}
5.9 \times 10^5 \pm 2.6 \times 10^{-4}
\end{tabular} \\
\hline
\end{tabular}
\caption{Natural transformation in exponential and stationary growth phases.}
\end{table}

\(^a\)Aggregates of bacterial cells resuspended before transformation.
\(^b\)Mean of four replicates with standard deviations.

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Figure 1. Kinetics of natural transformation. \textit{C. jejuni} transformed (t = 0) with equal amounts of isogenic chromosomal DNA carrying a Strep\(^b\) or a Cam\(^b\) resistance marker, respectively. At various time points (0–120 min.) following addition of DNA the transformation was terminated by adding DNase I. Single and double transformants were scored by plating on selective agar. Results are representative from three independent experiments. Diamonds, Cam\(^b\) transformants, Squares, Strep\(^b\) transformants; Triangles, Cam\(^b\) Strep\(^b\) double transformants.

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Table 2. Natural transformation in exponential and stationary growth phases.
Taken together these results show that natural transformation of C. jejuni can take place under several conditions not supporting growth. Moreover, we found no indications that stress enhances or induces competence of C. jejuni, since the levels of transformation were found to peak at optimal growth conditions.

**Competence is Dependent on the Level of Cellular Energy and Protein Synthesis**

In order to explore the regulation of natural transformation, we investigated the impact of cell density and the contribution of central cellular processes, such as respiration, translation and transcription to transformation.

First, we observed the transformation frequency of C. jejuni to increase at reduced cell density. The cell densities of exponentially growing cultures (OD$_{600}$ 0.3–0.4) were reduced 20 fold by dilution with growth media and transformed with rpsLS$_{m}$ DNA in the standard assay. This caused a 50 fold increase in the number of transformants per CFU for the diluted cultures compared to the original undiluted cultures (Diluted cultures: $4.1 \times 10^{-2} \pm 5.7 \times 10^{-2}$ transf./CFU and undiluted cultures: $8.2 \times 10^{-2} \pm 2.1 \times 10^{-2}$ transf./CFU. The results are mean of two independent experiments).

C. jejuni perform oxidative phosphorylation to obtain ATP, and an active electron transport chain is therefore crucial for energy generation. The cellular energy level of C. jejuni can be reduced by respiratory inhibitors that inhibit the transfer of electrons in central enzymes of the electron transport chain [36,38]. When we examined natural transformation in the presence of the respiratory inhibitors HQNO and sodium azide we found that the transformation mechanism of C. jejuni is an energy requiring process strongly depending on an active electron transport chain, as transformation was strongly inhibited by these compounds (Table 4). In addition, the energy level prior to natural transformation is important for transformation, since the number of transformants per CFU was also greatly reduced by the presence of the respiratory inhibitors just before the addition of DNA (Table 4).

To determine if the level of gene and protein expression per se affects competence of C. jejuni, natural transformation was investigated while inhibiting the transcriptional and translational processes, respectively. We found natural transformation of C. jejuni was inhibited by the translation inhibitors erythromycin and tetracycline, while largely being unaffected by the transcriptional inhibitor, rifampicin (Figure 3). C. jejuni is intrinsically resistant to rifampicin [39]. The inhibitory effect on transformation by the translation inhibitors was not caused by lethal effects of these antibiotics, since the reduction in viability was less than 50% and therefore much less than the reduction in transformation frequency (Figure 3). Natural transformation of C. jejuni thus seems to be modulated at the level of protein synthesis.

**Natural Transformation of C. jejuni is Dependent on the ClpP Protease**

The Clp proteolytic complex is involved in regulation of competence in B. subtilis and S. pneumoniae and to investigate if this

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**Table 3. Natural transformation of C. jejuni at varying temperature and atmospheric conditions$^{a,b}$.**

| Temperature | Oxygen level | 
|-------------|--------------|
|             | Aeroic       | Microaerobic | Anaerobic |
| 20°C        | 0.11 ± 0.13  | 0.16 ± 0.27  | Nd. $^c$  |
| 30°C        | 1.15 ± 0.92  | 1.96 ± 7.37  | Nd. $^c$  |
| 37°C        | 97.3 ± 10.3  | Index 100$^b$ ± 55.5 | 28.1 ± 12.9 |
| 42°C        | 130 ± 50.8   | 353 ± 37.8   | 16.2 ± 6.49 |

$^a$Exponentially growing cells were supplied with isogenic chromosomal DNA at 1:3 ratio (CFU$/$DNA) and transformation was allowed to proceed for 4 h before plating on selective agar. Results are the average of three independent experiments with two replicates.

$^b$Results are normalized to the number of transformants per ml at 37°C under microaerophilic conditions.

$^c$Not determined.

![Figure 2. Natural transformation at variable pH.](image)

Natural transformation of C. jejuni in BH broth with pH values from 4 to 12. Values of CFU$/$ml are normalized to the value at pH 7, while transformants$/$CFU are exact values. Results are representative from three independent experiments. Squares, transformants$/$CFU. Circles, survival, CFU$/$ml normalized to pH 7.

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complex also is involved competence of *C. jejuni*, mutants of the ClpP protease and the ClpX and ClpA ATPases were analyzed in the standard transformation assay. This showed the Δ*clpP* mutant of *C. jejuni* to be practically untransformable, while mutants lacking either the ClpX or the ClpA ATPases showed approximately 93 and 80% reduced transformation, respectively, when compared to the wild-type (wt) strain (Figure 4). Competence of the Δ*clpP* strain could be restored by inserting a copy of the *clpP* gene into a nonessential chromosomal locus of the mutant (resulting in strain Δ*clpP*+*clpP*), although the obtained level of transformation was lower than that of the wt strain (Figure 4). These experiments show that competence of *C. jejuni* is dependent on ClpP.

Natural Transformation of *C. jejuni* is Neither Involved in DNA Repair Nor Provides a Growth Benefit

*C. jejuni* lacks a typical SOS response and many of the DNA repair genes [20], and therefore we speculated that this organism may use external DNA to repair damaged DNA. However, the efficiency of transformation was unaltered following exposure to DNA damaging agents such as UV light or Mitomycin C (Table S2). Also, survival following UV light exposure did not increase when the bacteria were allowed to recover in the presence of excess amounts of DNA (Table S2). Furthermore, external DNA did not significantly alter the frequency of spontaneous nalidixic acid resistant colonies within three generations after the addition

| Respiratory inhibitor | Time of addition | CFU/ml | Transformants/CFU |
|-----------------------|------------------|--------|-------------------|
| Negative control      | –                | Index 100 | Index 100 |
| HQNOd                 | During transformation | **88** | **0.71** |
| HQNOd                 | Prior to transformation | **57** | **25** |
| Sodium azidee         | During transformation | **65** | **0.61** |
| Sodium azidee         | Prior to transformation | **64** | **4.5** |

*Exponentially growing cells were incubated with the respiratory inhibitor for 2 h prior to a wash out of compound and addition of DNA.

The respiratory inhibitor were added to the exponentially growing cells together with the DNA.

Normalized to the negative control.

100 μg/ml.

500 μM.

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Figure 3. The importance of transcription and translation for competence. Natural transformation in the presence of transcriptional or translational inhibitors. CFU/ml (light grey) and transformants/CFU (dark grey). The results represent a mean of three replicates with standard deviations. Data are normalized to the value of the negative control (no additive). Rif, 500 μg/ml rifampicin; Erm, 2 μg/ml erythromycin; Tet, 20 μg/ml tetracycline. White bars, CFU/ml; grey bars, transformants/CFU.

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of DNA (results not shown). These data suggest that natural transformation does not serve to repair DNA following DNA damage in \textit{C. jejuni}.

To investigate if \textit{C. jejuni} utilizes DNA as a nutritional source, we compared the growth of the wt strain in minimal media (MEMx without nucleosides) in the presence and absence of excess amounts of extracellular DNA. However, the addition of 10 \(\mu\)g/ml chromosomal DNA did not result in higher cell numbers than the cultures without added DNA (Figure S1). Likewise, cocultivation of a competent strain (\textit{rpsL}) with a competent-deficient strain (\textit{DclpP}) in minimal media with and without excess amounts of extracellular DNA did not reveal any growth or survival advantages for the competent strain from early exponential to late stationary phase (results not shown).

**Discussion**

In this study, we have explored natural transformation of \textit{C. jejuni} in order to determine how the transformation process is affected by environmental conditions and to assess the physiological impact of this process. We find that natural transformation in \textit{C. jejuni} occurs with a high efficiency (Table 1), and that the uptake of DNA from the extracellular environment must be a continuous and fast process resulting in multiple transformation events per cell cycle (Figure 1).

Horizontal gene transfer occurs actively when strains of \textit{C. jejuni} are co-cultivated either \textit{in vitro} or \textit{in vivo} [21]. This gene transfer is prevented by DNase and culture supernatants may serve as DNA source for transformation thus indicating that the exchange of genetic material is independent of cell-to-cell contact (results not shown) [32]. Therefore, genomic DNA of \textit{C. jejuni} must be present in the extracellular environment either via release or by cell lysis. We estimated the extracellular DNA concentration of \textit{C. jejuni} NCCTC11168 to be ca. 0.04 \(\mu\)g/ml, which corresponds to approximately one \textit{C. jejuni} genome per 100 CFU under the investigated conditions. This is less than what have been described for \textit{N. gonorrhoeae}, which actively releases DNA via a type IV secretion system [40,41], but comparable to what have been observed for \textit{N. meningitides} where DNA is released in a process relying on autolysins [42]. However, such autolysins have not been described for \textit{C. jejuni}.

\textit{C. jejuni} was most efficiently transformed at optimal growth conditions, i.e. in the exponential growth phase, at 37–42°C in a microaerobic atmosphere and pH ca. 7 (Table 2–3, Figure 2), which is consistent with previous observations from \textit{C. jejuni} and \textit{C. coli} [23,30,31]. The level of transformation was significantly reduced by inhibition of the electron transport chain by respiratory inhibitors (Table 4), while the addition of fresh nutrients by culture dilution had a positive effect on the transformation frequency. Taken together these results indicate the efficiency of natural transformation by \textit{C. jejuni} is connected to growth rate and thus may be correlated with the physiological fitness of the organism.

Competence development of \textit{B. subtilis}, \textit{H. influenzae} and \textit{S. pneumoniae} is induced upon nutrient limitation, pheromones and/or by environmental stress [25,26,27,43]. In contrast, we find \textit{C. jejuni} to be competent in both the exponential and late stationary growth phase as well as at growth limiting conditions such as low temperature, at aerobic and anaerobic levels of oxygen and in alkaline environments, that are conditions restrictive for growth but not for physiological activity of \textit{C. jejuni} [5]. In this aspect, \textit{C. jejuni} resembles \textit{N. gonorrhoeae} in being competent during all phases of growth [44]. However, natural transformation of \textit{N. gonorrhoeae} is dependent on the expression of essential pili proteins and specific DNA uptake sequences [45], which so far have not been identified in \textit{C. jejuni}. Based on our findings we consider natural transformation of \textit{C. jejuni} to be a constitutive process that operates...
regardless of environmental conditions and beyond the conditional boundaries supporting growth.

In agreement with the notion that *C. jejuni* is constitutively competent we found no evidence of natural transformation being directly dependent on cell density. Again, this is in contrast to the regulation of competence in *B. subtilis* and *S. pneumoniae* in which extracellular competence-stimulating peptides initiate the transcription of competence genes in the stationary and exponential growth states, respectively [43]. Central regulators of competence in these organisms are the key transcriptional regulator ComK and the alternative sigma factor ComX, respectively, as well as the ClpP protease, which degrades ComK and ComX [43,46,47]. Neither ComK nor ComX homologues are found in *C. jejuni*, but we found an interesting common characteristic of competence between *C. jejuni* and these organisms, since the ClpP protease strongly affect natural transformation of *C. jejuni* (Figure 4). Likewise, *C. jejuni* mutants of the complementary ClpX and ClpA ATPases showed decreased ability to be naturally transformed (Figure 4) suggesting that each of the two Clp ATPases could function together with ClpP in this process. A polar effect on the downstream deformylase (def) gene has previously been reported for the clpP mutant [48]. However, competence of the clpP mutant could partly be restored by reintroducing the wt clpP allele, and we therefore conclude that the competence deficiency of the mutant was indeed attributed to the ClpP protease. The function of the Clp proteolytic complex in competence of *C. jejuni*, however, remains unknown, and can be either directly by processing of competence proteins or indirectly by processing of other proteins with downstream effects on natural transformation. A previous proteomic analysis showed the ΔclpP mutant to carry altered levels of several proteins with as various functions such as motility and glycosylation [48]. Protein glycosylation is important for competence of *C. jejuni* [35], and since the ΔclpP mutant was shown to contain reduced amounts of the UDP-GlcNac/Glc4-epimerase (gna/gale) [48], which is a central enzyme of the glycosylation pathway [49], and a gnu mutant furthermore displays reduced level of natural transformation [50], we hypothesize that the competence deficiency of the ΔclpP mutant is, at least partly, caused by reduced protein glycosylation. However, it is likely that other factors are also involved in the lacking competence of the ΔclpP mutant.

Natural transformation in *C. jejuni* is dependent on protein synthesis, since inhibition of translation caused a severe reduction in the transformation efficiency, while transcriptional inhibition did not reveal the same effect (Figure 3). This finding points to a translational control of competence in *C. jejuni*, which can be caused either by a requirement of de novo synthesis of essential competence proteins or physical instability/rapid turnover of such proteins. This consideration is in agreement with observations of reduced transformation frequency of *H. pylori* following gentamicin treatment [51], and a correlation between trans-translationally tagging and competence of *H. pylori* [52]. Moreover, natural transformation of *N. gonorrhoeae* requires active protein synthesis in addition to a type IV pilus assembly apparatus [53,54].

DNA damaging agents cause an induction of essential competence genes in *H. pylori* and *S. pneumoniae*, and the level of natural transformation increase following DNA damage in these organisms [51,55]. In this study, it was explored if a similar correlation between DNA damage and natural transformation exists in *C. jejuni*. However, we found the level of natural transformation to be unaffected by DNA damage caused by Mitomycin C and UV light and moreover, high concentrations of available DNA in the extracellular environment did not increase either the recovery from DNA damage by UV light or the spontaneous mutation rate (Table S2 and results not shown). Likewise, *E. coli* has been shown to utilize DNA as a nutritional source [56,57], but a similar benefit of the constitutive competence of *C. jejuni* could not be identified (Figure S1).

In conclusion, this study shows that competence of *C. jejuni* is a constitutive process dependent on protein synthesis and the ClpP protease. The efficiency of natural transformation seems correlated to growth, but nevertheless *C. jejuni* is able to take up DNA even under conditions not supportive of proliferation. We therefore hypothesize that horizontal gene transfer by natural transformation is important for milieu adaptation of *C. jejuni* both inside the gastrointestinal tract and in the external environment.

### Experimental Procedures

#### Cultivation of *C. jejuni*

The original clinical isolate of *C. jejuni* NCTC11168 [11168-O] [58] was routinely cultivated on Blood agar Base II (Oxoid) supplemented with 5% bovine blood or in Brain Heart Infusion broth (BHI) (Oxoid). The bacteria were incubated at 37°C in a controlled atmosphere (8% O2, 6% CO2, 4% H2, and 84% N2) in a MACS VA1000 microaerophil workstation (Don Witley Scientific) unless otherwise stated. When cultivated in broth, serial dilutions of bacteria in 8–12 ml media were made in 90 mm Petri dishes and incubated overnight as described above. Preceding all experiments, *C. jejuni* cultures were investigated for contamination by phase contrast microscopy and cultures of the appropriate growth phase selected by optical density measurements.

#### Natural Transformation Assays

For a standard natural transformation assay, 200 μl of an exponentially growing *C. jejuni* in BHI (optical density 600 nm: 0.1–0.3) was transferred to a 2.0 ml cryo tube containing 1 ml BHI-agar. This biphasic culture was preincubated for 2 h under microaerobic conditions at 37°C followed by the addition of 10–400 ng DNA and transformation for 2–4 h under the same conditions. The transformed culture was subsequently diluted in isotonic sodium chloride solution and CFUs were scored by plating on Blood agar Base II, while transformants, which had obtained resistance marker(s), were scored by plating on Blood agar Base II supplemented with 20 μg/ml chloramphenicol, 50 μg/ml streptomycin or 30 μg/ml kanamycin when appropriate. Control experiments confirmed that the transformation process was terminated by plating on selective media. Chromosomal DNA purified by the DNeasy Blood & Tissue Kit (Qiagen) from *C. jejuni* NCTC11168 ΔclpP::CamR or *C. jejuni* NCTC11168 rpsLΔR [36,59] were routinely used for natural transformation assays. For test of DNA specificity, chromosomal DNA from *C. jejuni* DVI-SC11 StrepR or *C. jejuni* DVI-SC1181 StrepR [37] were applied. For competition experiments with foreign DNA, increasing amounts (0–200 ng) of chromosomal DNA from *Helicobacter pylori* ATCC700392 or *Arcobacter butzleri* ATCC4916 were supplied together with 20 ng of isogenic chromosomal DNA. For investigations on cell density and transformation efficiency, exponentially growing bacteria were diluted 20 fold in BHI prior to transformation with the standard transformation assay.

#### Kinetics of Natural Transformation

300 μl aliquots of an exponentially growing culture of *C. jejuni* NCTC11168 were dispensed to 13 tubes and each sample transformed with 2 μg/ml isogenic chromosomal DNA carrying a StrepR or a CamR resistance marker, respectively. The transformations were terminated at various time points (0–120 min.) by the addition of 0.017 units of DNase I (Fermentas),
and following 2 h of phenotypic expression, single and double transformants were scored by plating on selective agar.

**Quantification of Extracellular DNA**

*C. jejuni* NCTC11168 ΔclpP was cultivated to early stationary phase in MEM-sex (Life Technologies) supplemented with 20 mM sodium pyruvate and 40 μM FeSO₄. Bacterial cells were removed by centrifugation and the concentration of double stranded DNA in the sterile filtered supernatants were determined by fluorescence spectrometry and the Quant-iT™ Picogreen® dsDNA assay kit (Life Technologies) using the manufacturer’s instructions. Background fluorescence levels were disregarded by determining the difference between untreated and DNase I treated samples and correcting for fluorescent from the media and DNase I. The extracellular DNA concentration was further confirmed by natural transformation assays of *C. jejuni* wt with the *C. jejuni* ΔclpP supernatants in comparison with a standard row of transforming DNA. An estimate on the number of extracellular DNA genomes per CFU was made using an estimated molecular weight of the *C. jejuni* NCTC11168 genome (1.6 × 10⁶ bp * 660 g/mol *bp*), the Avogadro constant (6 × 10²³ molecules/mole), the measured DNA concentration and CFU/ml.

**Natural Transformation Under Stress Conditions**

Natural transformations under environmental stress conditions was determined with the standard natural transformation assay described above with the following exceptions: The effect of temperature and oxygen was analyzed by adding 400 ng of DNA to transform *C. jejuni* NCTC11168 wt for four hours at 20, 30, 37 or 42° C under aerobic conditions, or in sealed containers with microaerobic or anaerobic environments provided by CampyGen or AnaeroGen sachets (Oxoid), respectively. The effect of pH was analyzed by harvesting by centrifugation exponentially growing *C. jejuni* NCTC11168 bacteria, resuspending in sodium chloride solution, and transferring to BHI broth with pH values from 4–12. Biphatic media was not applied for these samples. Following two hours of preincubation, 30 ng of isogenic CamR chromosomal DNA was added and the transformation allowed to proceed for two hours before plating on selective and non-selective solid media.

**Inhibition of Electron Transport Chain, Transcription and Translation**

Various compounds were added to the standard natural transformation assay carried out without biphatic media. The respiration was inhibited by the addition of 100 μg/ml HQNO (2-n-Heptyl-4-hydroxyquinoline-N-oxide) (Alexis) or 500 μM sodium azide (Fluka) either only during the preincubation step or during the transformation step. For inhibition of transcription and translation, the preincubation step was omitted and 500 μg/ml rifampicin, 2 μg/ml erythromycin, or 20 μg/ml tetracycline, respectively, were added together with the DNA. Following two hours of transformation the inhibitory compounds were washed away by exchange of media, the transformation processes terminated by addition of 0.015 units DNase I (Fermentas) and phenotypic expression allowed to proceed for two hours before plating on selective and non-selective solid media.

**Construction of Complemented ClpP Mutant and Natural Transformation of Clp Mutants**

Deletion mutants of *C. jejuni* NCTC11168 clpP, clpA and clpX have been described previously [48]. For complementation of the ΔclpP mutant we applied a pKfdxA vector [60] kindly donated by Dr Duncan Gaskin. This vector contains a BsmBI restriction site located downstream from fdaA promoter. A divergent promoter controls expression from a gene conferring Kanamycin resistance. Both promoters are located between flanking sequences of pseudogene c70466 that allow for recombination of any gene cloned into BsmBI site. The clpP gene was amplified by with primers clpP-for (5’ ATGGTTATTCATGTGCTAATGTT) and clpP-rev (5’ ATTCCTCCACATGTCATTTAAGCCTTCTTG) containing BsmBI restriction sites. The resulting PCR product and the pKfdxA vector were then cut with the Es3fl enzyme (an isochizomer of BsmBI, Fermentas) and ligated, placing the gene under the control of the fdaA promoter. Ligated constructs were sequenced to confirm the presence of the insert, its correct orientation, as well as lack of point mutations. Vectors with successfully cloned clpP gene were then introduced by electroporation into strain *C. jejuni* clpPΔcam [48] and the transformants were selected for Chloramphenicol and Kanamycin resistance.

Natural transformation of *C. jejuni* NCTC11168 ΔclpP, ΔclpA, ΔclpX and ΔclpP+clpP was investigated in the standard natural transformation assays using 400 ng of isogenic chromosomal StrepR DNA.

**DNA Damage, Mutation Rate and Natural Transformation**

Natural transformation of *C. jejuni* wt was monitored during DNA damage caused by 10–500 ng/ml Mitomycin C (Sigma-Aldrich) in standard transformation assays without preincubation and biphatic media. Natural transformation and recovery following DNA damage caused by UV irradiation was examined as follows: Exponentially growing *C. jejuni* NCTC11168 bacteria were harvested by centrifugation, resuspended in isotonic salt solution to OD₆₀₀ ca. 0.3 and treated with 0–80 J/m² UV in an ultraviolet crosslinker (UVP). Following UV exposure, 2×100 μl aliquots of bacterial suspension from each UV treatment were added either 100 μl BHI + 400 ng isogenic CamR chromosomal DNA or just 100 μl BHI. Following 2 h recovery at 37°C in a microaerobic environment, samples were added 3 μl DNase I (Fermentas) and plated on non-selective and selective agar for scoring of transformants and CFU/ml. Transformants/CFU were compared between UV treated and untreated samples, while CFU/ml were compared for samples recovered with or without added DNA. The spontaneous mutation rate during natural transformation of *C. jejuni* was analyzed by plating and enumerating NalR colonies on Blood agar Base II supplemented with 50 μg/ml nalidixic acid at various time points following natural transformation with excess amounts of isogenic StrepR chromosomal DNA.

**Competence and Long-term Growth Competition with Excess Amounts of DNA**

Bacteria of an exponentially growing culture of *C. jejuni* NCTC11168 in BHI were harvested by centrifugation, washed in minimal media and resuspended to OD₆₀₀ 0.03 in minimal media MEM-sex without nucleosides (Life Technologies, catalog no 32561) supplemented with 40 μM FeSO₄. The culture was subsequently divided in two, and one fraction supplemented with 10 μg/ml of isogenic chromosomal CamR DNA. CFU/ml was determined at the start of the experiment and after 24 h of incubation at 37°C under microaerobic conditions. Growth competition experiments with *C. jejuni* NCTC11168 pfdA⁺ and ΔclpP:KanR [48] were carried out at 37°C under microaerobic conditions in the defined minimal medium MEM-sex (Life Technologies) supplemented with 40 μM FeSO₄ and 10 μg/ml chromosomal DNA isolated from *C. jejuni* ΔclpP. Cultures were
inoculated with approximately $10^8$ CFU/ml of each strain and subpopulation titers were determined periodically for four days by plating on selective media. The experiments were performed in duplicates.

**Supporting Information**

Figure S1 Growth of *C. jejuni* NCTC11168 is not supported by extracellular DNA. Growth of *C. jejuni* NCTC11168 in minimal media (MEM) without (white) and with 10 ng/ml of isogenic chromosomal DNA (grey). The results represent a mean of four replicates with standard deviations. (TIF)

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3. Jacobs-Reitsma W, Lyhs U, Wagenaar J (2008) Comparative studies of *Campylobacter jejuni* distant lineages with 32561, Life Technologies) without (white) and with 10 ng/ml of isogenic chromosomal DNA (grey). The results represent a mean of four replicates with standard deviations. (TIF)

**Author Contributions**

Conceived and designed the experiments: CSV. Performed the experiments: CSV ML. Analyzed the data: CSV LB HI. Contributed reagents/materials/analysis tools: CSV ML. Wrote the paper: CSV LB ML. HI.

**Table S1** DNA from *A. butleri* and *H. pylori* cannot outcompete natural transformation of isogenic DNA. (PDF)

**Table S2** Natural transformation and survival of *C. jejuni* following DNA damage. (PDF)
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