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Impaired competence in flagellar mutants of *Bacillus subtilis* is connected to the regulatory network governed by DegU

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Summary
The competent state is a developmentally distinct phase, in which bacteria are able to take up and integrate exogenous DNA into their genome. *Bacillus subtilis* is one of the naturally competent bacterial species and the domesticated laboratory strain 168 is easily transformable. In this study, we report a reduced transformation frequency of *B. subtilis* mutants lacking functional and structural flagellar components. This includes *hag*, the gene encoding the flagellin protein forming the filament of the flagellum. We confirm that the observed decrease of the transformation frequency is due to reduced expression of competence genes, particularly of the main competence regulator *comK*. The impaired competence is due to an increase in the phosphorylated form of the response regulator DegU, which is involved in regulation of both flagellar motility and competence. Altogether, our study identified a close link between motility and natural competence in *B. subtilis* suggesting that hindrance in motility has great impact on differentiation of this bacterium not restricted only to the transition towards sessile growth stage.

Originality-Significance statement
Understanding how versatile bacterial phenotypes influence each other is important for our basic understanding of microbial ecology. Our research highlights the novel intertwinement of bacterial differentiation and reveal how lack of single cell motility adjusts DNA exchange among bacterial strains.
Introduction

When facing stressful environmental conditions, bacteria can respond with a variety of post-exponential modifications including secretion of degradative enzymes, sporulation, or genetic competence. *Bacillus subtilis* is one of the bacterial species that are able to take up free DNA from the environment and incorporate it into its own genome, a phenomenon referred to as natural competence (Dubnau, 1991). To import extracellular DNA into *B. subtilis* cells, a pseudopilus formed by proteins encoded by the *comG* operon facilitates binding to the receptor protein ComEA, which is located in the bacterial cell membrane (Inamine and Dubnau, 1995; Chen *et al.*, 2005). As only single stranded DNA is imported, the membrane-associated nuclease NucA catalyzes cleavage of the DNA after successful binding (Provvedi *et al.*, 2001). Subsequent transport of the DNA through a membrane channel formed by the protein ComEC is mediated by the ATPase ComFA that probably requires the transmembrane proton motive force (Maier *et al.*, 2004).

To take up DNA, cells have to be in a developmental state, in which a specific set of genes and regulators are expressed (Dubnau, 1991; Berka *et al.*, 2002). Regulation of the whole apparatus required for competence development is complex. Briefly, entry into the competence state occurs in a bistable manner during the early stationary phase, where a minority of cells produces high level of the competence master regulator ComK above a certain threshold that is required to switch on competence development, the so called ‘K-state’ (van Sinderen *et al.*, 1995; Maamar and Dubnau, 2005; Smits *et al.*, 2005; Dubnau and Losick, 2006). It was demonstrated that noise in the expression of *comK* determines the competent subpopulation and allows a dynamic stress response regarding competence development (Maamar *et al.*, 2007; Mugler *et al.*, 2016). Eventually, ComK activates the expression of late competence operons encoding the DNA-binding and -uptake machinery as well as genes, whose products are responsible for DNA integration (Berka *et al.*, 2002; Ogura *et al.*, 2002; Hamoen *et al.*, 2003). Increase of the ComK level is linked to a quorum sensing-mediated accumulation of the small ComS protein, which interferes with ComK degradation during the exponential phase (Turgay *et al.*, 1998). ComK is able to bind the *comK* promoter, triggering its own transcription, thus creating an auto-stimulatory loop (van Sinderen and Venema, 1994). This binding is further stabilized by the non-phosphorylated form of the regulator DegU to increase the level of ComK above the threshold sufficient for competence development (Hamoen *et al.*, 2000).

However, DegU is not only crucial for competence initiation, but also involved in the regulation of many other processes including protease production, biofilm development, and, particularly, flagellar motility (Murray *et al.*, 2009; Mukherjee and Kearns, 2014). The main components of the hook and basal body of the flagellum are encoded by the large *fia/che* operon. Transcription of this operon is activated by a complex formed by the regulator SwrA and phosphorylated DegU (DegU~P),
which binds to one of the fla/che promoters (Mordini et al., 2013; Mukherjee and Kearns, 2014). Amongst other genes, the operon contains the gene encoding the sigma factor $\sigma^D$ that activates transcription of motility genes outside the fla/che operon like the hag gene (encoding flagellin), motA and motB (encoding flagellar stator proteins), as well as transcription of lytF, which is necessary for separation of motile cells after cell division (Serizawa et al., 2004; Chen et al., 2009). The level of $\sigma^D$ and its position in the fla/che operon determines the cell fate, i.e. subpopulations of motile single cells or non-motile chains (Cozy and Kearns, 2010). The function of DegU~P changes in the absence of SwrA. In this case, DegU~P seems to inhibit motility via the same promoter of the fla/che operon (Amati et al., 2004). Additionally, DegU~P can activate the anti-sigma factor FlgM by binding to its promoter region in the absence of SwrA (Hsueh et al., 2011) allowing FlgM to antagonize $\sigma^D$ (Caramori et al., 1996). Consequently, DegU~P indirectly suppresses transcription of $\sigma^D$-dependent genes (Hsueh et al., 2011). It was suggested that a completion of flagellum assembly can be sensed by the DegSU two component system: FlgM, which is activated by DegU~P, causes inhibition of $\sigma^D$-dependent genes, when the assembly of the flagellum is impeded (Cozy and Kearns, 2010; Hsueh et al., 2011).

In addition to its role on modulating the expression of flagellum-related genes in B. subtilis, the phosphorylation and therefore the activity of DegU, has been shown to be influenced by a mechanical signal transmitted by the flagellum (Cairns et al., 2013). Inhibition of flagellar rotation by the flagellar clutch or by tethering the flagella results in an increased DegU~P level in the cell.

In this study, we report a correlation between motility function and competence development, which in B. subtilis is connected by the multifunctional response regulator DegU. We show that mutants lacking a functional flagellum such as Δhag, ΔmotA, and ΔflgE exhibited a reduced transformation frequency. This was due to a decrease in competence gene expression, particularly reduced levels of the competence master regulator ComK, which can be reverted by overexpressing comK in the hag mutant. Finally, we suggest that the reduced transformation frequency was likely due to an imbalance in the phosphorylation level of DegU.
Results

Lack of active flagella impairs competence for DNA uptake in *B. subtilis*.

While genetically modifying various *B. subtilis* strains, a striking difference in transformation frequency was observed between the wild type and a non-motile mutant lacking the gene encoding flagellin, *hag*. To explore this phenomenon, we tested the transformability of wild type (strain 168) and *hag*-mutant in competence medium (see Experimental Procedures), where the *hag*-mutant showed a more than 100-fold reduced transformation frequency relative to the wild type (Fig. 1A, B): while the transformation frequency of the wild type ranged between $3 \cdot 10^{-5}$ and $5 \cdot 10^{-5}$, that of the *hag*-mutant was reduced to values below $3 \cdot 10^{-7}$. Similarly, the undomesticated *B. subtilis* strains DK1042 (transformable derivative of NCIB 3610) and PS216 showed reduced transformation efficiency when the *hag* gene of these strains was disrupted (Fig. S1). To investigate whether this difference in transformation frequency between the two strains resulted from a lower growth rate of the *hag*-mutant, the growth behavior of wild type and *hag*-mutant grown in competence medium was evaluated over time. As depicted in Fig. 1B, the *hag*-mutant showed a clear growth advantage and reached a higher OD compared to the wild type (unpaired two-sample t-test with Welch Correction: $P = 0.001$, $n = 5$), thus supporting our previous observations (Hölscher *et al.*, 2015).

Further, it was tested whether the addition of DNA at different time points would increase the transformation frequency of the *hag*-mutant. However, the mutant showed a consistently low transformation frequency over the course of several hours, indicating that a shifted timing of the initiation of the competence state is unlikely to be the reason for the observed decrease in transformation frequency (Fig. 1C). To test whether this phenomenon is restricted to the *hag*-mutant or connected to the lack of an active motility apparatus in general, mutants lacking other functional flagellum-related genes were investigated. The transformation frequencies of mutants lacking the gene encoding one of the flagellar motor units, *motA*, and the gene encoding the hook protein, *flgE*, were decreased in both cases compared to the wild type (Fig. 2; unpaired two-sample t-test with Welch Correction: $P = 0.01$ for WT - *ΔmotA*, $P = 0.039$ for WT - *ΔflgE*, $n = 9$ for both).

Although the wild type transformation frequency was slightly different, the transformation frequencies of both *ΔmotA* and *ΔflgE* were around 10-times lower than that of the wild type (Fig. 2). In contrast, a *cheA*-mutant lacking the main chemotaxis sensor kinase showed a similar transformation frequency than the wild type (Fig. S2; unpaired two-sample t-test with Welch Correction: $P = 0.232$, $n = 3$), suggesting that the presence of an active flagellum, but not directed motility per se is required for full competence development. In sum, these results demonstrate that the observed impaired competence is linked to a loss of flagellar function.
Lack of competence in flagellar mutants is due to the reduced expression of competence genes.

To determine if the detected diminished transformation frequency of flagellar mutants was due to altered competence gene expression, the fluorescent reporter \( P_{\text{comG}^\text{gfp}} \) was introduced into these strains. This reporter allows the detection of cells expressing the \( \text{comG} \) operon-encoding genes required for pseudopilus formation and DNA uptake. In addition, this reporter provides a proxy on the activity of the ComK protein, the master regulator of competence. Qualitative microscopy analyses of cultures harboring the reporter, and which were grown in competence medium for 5 h, showed indeed a decreased number of fluorescent (i.e. \( \text{comG} \) expressing) cells in the \( hag \) mutant compared to the wild type, whereas a control strain lacking \( \text{comK} \) showed no fluorescence (Fig. 3A).

For quantitative determination of competence gene expression within the population, flow cytometric measurements were performed that revealed 24.7% of fluorescent cells in wild type cultures (mean value), but only 4.6% of fluorescent cells for the \( hag \)-mutant (Fig. 3B, C; unpaired two-sample t-test with Welch Correction: \( P = 0.004, n = 3 \)), thus confirming the microscopy results. Similarly, the \( \text{motA} \) and \( \text{flgE} \) mutants were analyzed microscopically as well as by using flow cytometry. Both methods revealed fewer cells activated transcription of competence genes in these mutants compared to the wild type (Fig. 4; unpaired two-sample t-test with Welch Correction: \( P = 0.017 \) for WT - \( \Delta \text{motA} \), \( P = 1.3 \times 10^{-9} \) for WT - \( \Delta \text{flgE} \), \( n = 3 \) for both; mean percentage of fluorescent cells: 16.7% for wild type, 4.5% for \( \Delta \text{motA} \), 4.2% for \( \Delta \text{flgE} \)). Flow cytometry measurements at different time points during growth in competence medium confirmed a similarly reduced fraction of competent cells in the \( hag \) mutant compared to the wild type strain (Fig. S3).

**Reduced competence in \( hag \) mutant can be rescued by overexpression of \( \text{comK} \).**

The reduced competence gene expression in the tested flagellar mutants suggested a regulatory link between flagellar motility and competence. To investigate if regulatory elements upstream of \( \text{comK} \) were responsible for our observations and if a bypass of those could therefore rescue transformation frequency in the flagellar mutant, we examined a strain with an additional copy of \( \text{comK} \) under the control of a xylose-inducible promoter (\( P_{xyl}^\text{comK} \)). Indeed, in combination with \( P_{xyl}^\text{comK} \), the transformation level of the \( hag \) mutant increased back to a level that was statistically indistinguishable from wild type levels (mean transformation frequency of \( 5.3 \times 10^{-6} \) for the wild type and \( 8.5 \times 10^{-6} \) for \( \Delta \text{hag} P_{xyl}^\text{comK} \); Kruskal-Wallis test: \( P = 0.453, n = 9 \), Fig. 5A). Despite this observed increase in the \( hag \) strain upon \( \text{comK} \) overexpression, the wild type strain, which contained an inducible copy of \( \text{comK} \) showed a higher transformation frequency (Fig. 5A, Kruskal-Wallis test: \( P = 3.4 \times 10^{-4} \) for WT - WT \( P_{xyl}^\text{comK} \), \( P = 3.4 \times 10^{-4} \) for WT \( P_{xyl}^\text{comK} - \Delta \text{hag} P_{xyl}^\text{comK} \), \( n = 9 \) for both), which
was probably due to higher levels of *comK* transcription at the native locus as previously observed (Hahn *et al.*, 1996).

*Reduced competence in flagellar mutants is likely connected to unbalanced DegU phosphorylation.*

As the above results suggested that regulatory elements in response to impaired flagellar motility are responsible for the decreased *comK* expression, we investigated DegU as a likely candidate causing the reduced competence in flagellar mutants. As non-phosphorylated DegU was implicated to be required for *comK* transcription (Dahl *et al.*, 1992; Hamoen *et al.*, 2000), two variants of *degU* were tested: *degU*32, which harbors a mutation resulting in an extended half-life and thus higher stability of the phosphorylated form of the DegU protein (DegU~P), and *degU*146, which is cannot be phosphorylated (Dahl *et al.*, 1991; Dahl *et al.*, 1992; Kunst *et al.*, 1994). Both variants were tested in wild type as well as the Δ*hag* background to observe differences in transformability compared to the wild type strain. The results of this experiment indicated that the transformation frequency of the *degU*32 strain was slightly decreased (Figure 5B), which is consistent with previous publications, suggesting that non-phosphorylated DegU is required for priming *comK* transcription. The observed difference, however, was only marginally significant in our experimental setup (Figure 5B; Kruskal-Wallis test: *P* = 0.078, *n* = 6). Surprisingly, when combined with the Δ*hag* mutation, the transformability of *degU*32 increased significantly far above wild type levels, despite presumably possessing low levels of non-phosphorylated DegU to induce the ComK auto-stimulatory loop (Fig. 5B; Kruskal-Wallis test: *P* = 0.004, *n* = 6). Furthermore, we observed a tendency towards a reduced albeit non-significant transformation frequency in the *degU*146 strain compared to the wild type (Fig. 5B; Kruskal-Wallis test: *P*>0.05, *n* = 6). This result was similar to the one observed for *degU*32, although no negative impact on transformability was expected in strain *degU*146 due to the abolished phosphorylation of DegU. Interestingly, the *degU*146 strain combined with the Δ*hag* mutation exhibited transformation frequencies at the same level than the wild type strain (Fig. 5B; Kruskal-Wallis test: *P*>0.05, *n* = 6) that was significantly higher than the transformation frequency of the single *degU*146 mutant (Fig. 5B; Kruskal-Wallis test, *P* = 0.007, *n* = 6). These results suggest that altering the phosphorylation state of DegU in flagellar mutants can revert the negative impact on competence, which was caused by a lack of motility.

*Increased viscosity enhances competence in B. subtilis*

A recent study showed that restricting the flagellar rotation by viscous medium results in induction of flagellar gene transcription and activation of the DegSU two-component system in *Paenibacillus* sp. NAIST15-1 (Kobayashi *et al.*, 2017). Accordingly, we tested whether an increased viscosity of the
medium changes the transformability in *B. subtilis*. Indeed, the average transformation frequency of the wild type strain was three-fold higher in a medium of increased viscosity. The corresponding statistical test, however, indicated only a trend towards a statistically significant difference (Fig. 6; unpaired two-sample t-test: *P* = 0.095, *n* = 4).
Discussion

Many cellular processes in *B. subtilis* are tightly connected through their underlying regulatory networks. Examples include motility and biofilm formation or biofilm formation and sporulation (e.g. Vlamakis et al., 2013; Marlow et al., 2014; Hölscher et al., 2015). Here, we report an additional connection between flagellar motility and competence development. We could show that mutants with impaired flagellar function have defects in competence development. Such mutants displayed a considerably lower transformation frequency and expression of late competence genes, suggesting that it is due to an altered expression of the competence master regulator gene *comK*. The rescue experiment with an inducible *comK* confirmed that indeed competence could be rescued in the Δ*hag* strain, since Δ*hag* Pₓyl*comK* exhibited a wild type transformation level.

In a recently published study, similar effects were observed even though different methods have been used: Diethmaier and colleagues found that the expression of *comK* is lower in deletion mutants of the *fla/che* operon, *hag*, and the second stator gene *motB* (Diethmaier et al., 2017). While in their study the expression of *comK* was primarily monitored using a *comK* promoter fusion, our experiments predominantly assayed transformation frequency. Both studies, however, report a negative effect of the deletion of flagellar components on competence development. The extent to which wild type and mutant differ in competence development is in the same order of magnitude between the studies: for example Diethmaier et al. observe a 10-fold reduced number of *comG* expressing-cells in Δ*hag* (Diethmaier et al., 2017), whereas our flow cytometry experiments showed a slightly lower, 5-fold reduction. Additionally, by investigating the transformation frequency in a *cheA* mutant, we could also show that the chemotactic response does not seem to have an influence on competence development.

Investigating modified variants of the response regulator DegU, we found that the transformation frequency of the *hag*-mutant could be restored to wild type level when the mutant carried a non-phosphorylatable DegU variant (*degU*146). This result suggests that a high level of DegU~P in the flagellar mutants was the reason for the decreased expression of the competence genes and *comK*, which could be counteracted by introducing a non-phosphorylatable variant of DegU. By additionally investigating a strain harboring a *degU*-yfp fusion, Diethmaier et al. also suggested an increased level of DegU~P to be present in the *hag*-mutant (Diethmaier et al., 2017), which is consistent with our conclusions. In addition, the authors detected a reduced expression of *comK* in a strain with the *degU*32 variant, which produces a form of DegU~P with higher stability (Diethmaier et al., 2017). Comparable results were obtained by Msadek et al., who found that high levels of DegU~P inhibit competence (Msadek et al., 1990). We observed a similar, although weak statistical trend towards a reduced transformation frequency in *degU*32 strain. Miras and Dubnau (2016) have recently
highlighted that differences in the DegU phosphorylation pathway among diverse \emph{B. subtilis} isolates were likely responsible for variance in DNA transformation efficiency among certain domesticated and undomesticated strains. Moreover, slight differences in competence induction levels could also be affected by strain-specific characteristics. For example, \emph{B. subtilis} 168 strains derived from different laboratories can exhibit striking variations in biofilm robustness (Gallegos-Monterrosa \textit{et al.}, 2016). As suggested by Diethmaier and colleagues, the reduced transformation frequency in \emph{degU32} might be caused by the high DegU~P levels of this strain. However, the \emph{degU32} strain exhibits a non-motile phenotype and in the undomesticated strains DegU32 is not able to interact with SwrA at the \textit{P}
\textit{A} promoter of the \textit{fla/che} operon (\textit{swrA} is inactive in domesticated strains), leading to repression of \textit{P}
\textit{A} (\textit{fla/che}) (Amati \textit{et al.}, 2004; Mordini \textit{et al.}, 2013). Due to low or no expression of the basic flagellar genes, this phenotype could mimic the situation observed in the flagellar mutants. In addition, we observed an increased transformation frequency when the \textit{hag} gene was deleted from the \emph{degU32} background. This is in contrast to the model assuming that increased levels of phosphorylated DegU in the cells lowers competence. Therefore, it is possible that yet unidentified factors are also involved in connecting motility and competence development that might be independent of DegU~P. At this point however, we cannot provide a reasonable explanation for the increased transformation frequency of \textit{Δhag degU32}.

Interestingly, induction of competence state has negative impact on motility in \emph{B. subtilis}. ComK negatively controls \textit{hag} gene expression by stimulating the transcription of \textit{comFA-C} operon and the downstream located anti-sigmaD factor coding gene, \textit{flgM} (Liu and Zuber, 1998). This feedback loop presents another intriguing connection between these two cellular processes.

Diethmaier \textit{et al.} proposed that increased DegU~P and lower \textit{comK} expression in the flagellar mutants and in a strain with straight flagella was caused by a lower viscous load (Diethmaier \textit{et al.}, 2017). In line with this report, we also observed that higher viscosity in the medium resulted in an increased transformation frequency. Nevertheless, a possible role of the DegSU two-component system in sensing incomplete assembly of flagella and dysfunction as suggested previously (Hsueh \textit{et al.}, 2011; Cairns \textit{et al.}, 2013) could also explain the increased DegU~P levels in the flagellar mutants.

Together, our results identify a connection between two major physiological processes, providing another example of the complexity of intracellular regulatory networks and the vast amount of tasks a single regulator can cover.
Experimental Procedures

Strains and cultivating conditions

The strains used in this study and their mutant derivatives are listed in Table S1. Mutants constructed in this study were obtained by natural transformation of a *B. subtilis* receptor strain with genomic DNA from a donor strain. Strain TB831 was created by transformation of strain 168 P<sub>xyl</sub>-comK with genomic DNA of strain GP902 (J. Stülke lab collection). To obtain strains TB926 and TB925, genomic DNA of strain 168 P<sub>comG</sub>-gfp was used to transform strain TB710 and TB689, respectively. Strain TB928 was obtained by transforming strain 168 P<sub>xyl</sub>-comK with genomic DNA of GP901 (J. Stülke lab collection). To create strain TB935 and TB936, strain 168 was transformed with genomic DNA obtained from strain QB4371 (Kunst *et al.*, 1994) and QB4458 (Dahl *et al.*, 1991), respectively. Their derivatives harbouring also a mutation of hag (TB923 and TB924) were created by transformation with genomic DNA, which was obtained from GP901. In-frame deletions of motA, flgE, and cheA were created using plasmids pEC1, pDP306, and pDP338, respectively, as previously described (Courtney *et al.*, 2012; Chan *et al.*, 2014; Calvo and Kearns, 2015). Strains were verified by fluorescence microscopy (P<sub>comG</sub>-gfp reporter), PCR (hag mutants), or sequencing (degU variants), using the oligonucleotides listed in Table S2. For experiments with strains harboring the inducible construct P<sub>xyl</sub>-comK, 1% of xylose (final concentration) was added for induction (see van den Esker *et al.*, 2017). To increase medium viscosity, 10% Ficoll400 (Carl Roth) was added to the medium before culture inoculation and the mix was vortexed vigorously for ca 20 s.

Transformation frequency assay

To assess the transformation frequency of different strains, a modified version of the transformation protocol from Konkol *et al.* (2013) was used. 1 ml of each culture grown in 3 ml Lysogeny broth (LB) medium (LB-Lennox, Carl Roth; 10 g L<sup>-1</sup> tryptone, 5 g L<sup>-1</sup> yeast extract, and 5 g L<sup>-1</sup> NaCl) for 16 h was centrifuged for 2 min at 11,000 x g. The pellet was washed twice in de-ionized water and was re-suspended in 100 µl de-ionized water. The re-suspended culture was diluted (1:80) in complete competence medium (MC: 1.8 ml de-ionized water, 6.7 µl 1M MgSO<sub>4</sub>, 50 µl 0.2% L-Tryptophan, 200 µl 10xMC per 100 ml 10xMC: 14,036 g K<sub>2</sub>HPO<sub>4</sub>[x3H<sub>2</sub>O], 5,239 g KH<sub>2</sub>PO<sub>4</sub>, 20 g glucose, 10 ml 300 mM tri-sodium citrate, 1 ml 83.97 mM ammonium iron (III) citrate, 1 g casein hydrolysate, 2 g potassium glutamate [H<sub>2</sub>O]) and incubated at 37°C, 225 rpm. For experiments with strains harboring P<sub>xyl</sub>-comK, 10x MC with fructose instead of glucose was used. After 6 h incubation time, 5 µl DNA with an antibiotic marker (PY79 safA::Tet gDNA, 60 ng/µl) was added to 500 µl culture. Any alteration in incubation time before addition of the DNA is indicated in the results section. Each culture was incubated for 30 min, then 500 µl fresh LB medium was added and the culture was incubated for
another 1 h under the conditions mentioned above. Serial dilutions of cultures supplemented with DNA were prepared and plated on LB medium supplemented with 1.5 % agar to determine the number of colony forming units (cfu). Additionally, 50 µl and 100 µl undiluted cultures supplemented with gDNA as well as controls were plated on tetracycline (Tet) containing LB-agar plates (10 µg ml⁻¹ Tet) to determine the number of transformant colonies. The transformation frequency was calculated by dividing the number of transformants per ml by cfu per ml.

**Growth curve experiments**

To examine growth properties, cultures were inoculated in LB medium from frozen glycerol stocks and incubated for ca. 16 h at 37°C shaking at 225 rpm. Cultures were diluted 1:100 in 200 µl fresh completed MC medium (see above) and the OD₅₉₀ nm was recorded for 16 h using a TECAN Infinite F200 PRO microplate reader. The cultures were incubated with orbital shaking with a duration of 800 s and an amplitude of 3 mm at 37°C and the OD₅₉₀ was measured every 15 min.

**Fluorescence microscopy**

Strains were investigated using a confocal laser scanning microscope (LSM 780, Carl Zeiss) equipped with an argon laser and a Plan-Apochromat/1.4 Oil DIC M27 63× objective. Cultures were grown prior microscopy for 5 h (if not indicated otherwise) in competence medium under the same conditions as described above (see section transformation frequency assay). Excitation of the fluorescent reporter (GFP) was performed at 488nm and the emitted fluorescence was recorded at 493-598nm. For image visualization, Zen 2012 software (Carl Zeiss) was used, brightness and contrast were adjusted equally in all images.

**Flow cytometry**

Flow cytometric measurements were performed using a Partec CyFlow® Space (Sysmex Partec GmbH, Germany), which was equipped with a solid-state laser for excitation of green/yellow fluorescent proteins at 488 nm. Single cells were detected in forward and sideward scatter channels as well as in one fluorescent channel. A minimum of 40,000 cells were analyzed for the experiments. To define the background fluorescence signal, non-labelled *B. subtilis* cultures were analyzed as control. Cultures used for measurements were grown for 5 h (if not indicated otherwise) in competence medium under the same conditions as described above (see section transformation frequency assay). For evaluation of the data, the FlowJo® software (FlowJo LLC, Ashland, USA) was used and a gate was set at 3 fluorescence units for all samples to isolate the fluorescent population and determine the percentage of fluorescent cells.
Statistical analyses were performed using OriginPro 2016 (V93E, OriginLab Northampton, USA). Unpaired two-sample t-test with Welch Correction or a Kruskal-Wallis test was used to test for significant differences.
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Conflict of interest. None declared
References


Figure legends

**Fig. 1.** Transformation frequency is reduced in a mutant lacking flagellin protein. (A) Transformation frequency of *B. subtilis* wild type and hag-mutant after 6 h incubation in competence medium (unpaired two-sample t-test with Welch Correction: $P = 3.1 \times 10^{-5}$, $n = 9$). The inset shows a zoom-in of the hag-mutant data. (B) Growth dynamics of wild type and hag-mutant during 12 h incubation in competence medium. Standard deviations for the measurements are depicted in light grey (unpaired two-sample t-test with Welch Correction: $P = 0.001$, $n = 5$). Arrows indicate the time points of DNA addition to investigate the transformation frequency over time, which is shown as box-and-whisker plot in (C). The line in the boxes represents the median, the box indicates the 25th-75th percentile. Asterisks indicate statistically significant differences between wild type and hag-mutant (unpaired two-sample t-test with Welch Correction for WT - Δhag: $P = 0.125$ for 3 h, $P < 0.01$ for 4 h, 5 h, 6 h; $n = 6$).

**Fig. 2.** Mutants impaired in flagellar function exhibit lower transformation frequencies. Deletion of the gene encoding a flagellar stator (*motA*; A) or the gene encoding the hook protein (*flgE*; B) results in significantly lower transformation frequency of the respective strain compared to the wild type after incubation in competence medium for 6 h. The line in the boxes represents the median, the box indicates 25th-75th percentile. Asterisks indicate statistically significant differences (unpaired two-sample t-test with Welch Correction: $P = 0.01$ for ΔmotA; $P = 0.039$ for ΔflgE; $n = 9$ for all).

**Fig. 3.** Fewer cells of the hag-mutant express competence genes compared to the wild type. (A) Representative microscopy images of strains harboring the $P_{comG}$-gfp reporter in wild type, Δhag or ΔcomK genetic background. Images were recorded after incubation in competence medium for 5 h. The scale bar represents 50 µm. (B) Histograms of flow cytometric measurements showing the cell count and the fluorescence in arbitrary units for wild type, Δhag, and ΔcomK including background fluorescence. Representative images are shown for each strain. (C) Percentage of fluorescent cells determined from the data in (B) for wild type and hag-mutant by isolating the fluorescent population with fluorescence intensities above 3 A.U. Asterisks indicate significant differences (unpaired two-sample t-test with Welch Correction: $P = 0.036$, $n = 3$).

**Fig. 4.** Competence gene expression is reduced in mutants lacking a functional flagellum. (A) Representative microscopy images of strains harboring the $P_{comG}$-gfp reporter in wild type, ΔmotA, or ΔflgE genetic background. Images were recorded after 5 h incubation in competence medium. The scale bar represents 50 µm. (B) Histograms of flow cytometric measurements showing the cell count.
and the fluorescence in arbitrary units for wild type, ΔmotA or ΔflgE. Representative images are shown for each strain. (C) Percentage of fluorescent cells determined from the data in (B) by isolating the fluorescent population with fluorescence intensities above 3 A.U. showing a significant difference (asterisks) between wild type and ΔmotA (P = 0.017) as well as wild type and ΔflgE (P < 0.001) with n = 3 for both (unpaired two-sample t-test with Welch Correction).

Fig. 5. Synthetically induced comK and degU146 increase competence of Δhag. (A) Transformation frequencies of wild type compared to strains harboring a xylose-inducible copy of comK (P<sub>xyl</sub>-comK) with wild type or Δhag genetic background (Kruskal-Wallis test for WT - WT P<sub>xyl</sub>-comK: P = 3.4·10^{-4}; for WT P<sub>xyl</sub>-comK - Δhag P<sub>xyl</sub>-comK: P = 3.4·10^{-4}, n = 9 for both). (B) Transformation frequencies of WT compared to strains harboring either a phosphorylated DegU variant (degU32) or a non-phosphorylatable DegU variant (degU146) in wild type or Δhag background. Strain Δhag degU32 is significantly different from all other strains (Kruskal-Wallis test: P < 0.05 for all, n=6). The line in the boxes represents the median, the box indicates 25<sup>th</sup>-75<sup>th</sup> percentile. Asterisks indicate statistically significant differences (Kruskal-Wallis test: P = 0.007, n=6).

Fig. 6. Competence is improved in viscous medium. Transformation frequency of the wild type strain grown in normal competence medium and in medium with increased viscosity. The line in the boxes represents the median, the box indicates 25<sup>th</sup>-75<sup>th</sup> percentile, # indicates marginally significant differences (unpaired two-sample t-test: P = 0.095, n = 4).
Figure 1

302x275mm (300 x 300 DPI)
Figure 2

Transformation frequency (×10^1)

A

B

WT  ΔmotA

WT  ΔflgE

332x130mm (300 x 300 DPI)
Figure 3
410x216mm (300 x 300 DPI)
Figure 4

487x217mm (300 x 300 DPI)
Figure 5

169x263mm (300 x 300 DPI)
Figure 6

Transformation frequency ($10^{-5}$)

WT

WT + Ficoll400

177x190mm (300 x 300 DPI)