Cheaters shape the evolution of phenotypic heterogeneity in *Bacillus subtilis* biofilms

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ABSTRACT

Biofilms are closely packed cells held and shielded by extracellular matrix composed of structural proteins and exopolysaccharides (EPS). As matrix components are costly to produce and shared within the population, EPS-deficient cells can act as cheaters by gaining benefits from the cooperative nature of EPS producers. Remarkably, genetically programmed EPS producers can also exhibit phenotypic heterogeneity at single cell level. Previous studies have shown that spatial structure of biofilms limits the spread of cheaters, but the long-term influence of cheating on biofilm evolution is not well understood. Here, we examine the influence of EPS nonproducers on evolution of matrix production within the populations of EPS producers in a model biofilm-forming bacterium, Bacillus subtilis. We discovered that general adaptation to biofilm lifestyle leads to an increase in phenotypical heterogeneity of eps expression. Apparently, prolonged exposure to EPS-deficient cheaters, may result in different adaptive strategy, where eps expression increases uniformly within the population. We propose a molecular mechanism behind such adaptive strategy and demonstrate how it can benefit the EPS-producers in the presence of cheaters. This study provides additional insights on how biofilms adapt and respond to stress caused by exploitation in long-term scenario.

INTRODUCTION

Cooperative interactions are prevalent for all life forms [1], even for simple microbes that often exist in communities of matrix bound surface-attached cells called biofilms [2–6]. However, when costly products such as siderophores [7, 8], extracellular polymeric substances [9, 10], digestive enzymes [11], and signaling molecules [12, 13] are secreted and shared, cooperative behavior becomes susceptible to cheating [2, 14–16], where mutants defective in cooperation can still benefit from cooperative community members [4, 5, 17]. It has been shown that spatially structured biofilms, where interactions with clone mates are common and diffusion of public goods is limited, may serve as natural defense against cheating [18–20]. However, long time scale studies have recently reported that biofilm defectors can spontaneously emerge and spread in biofilms by exploiting other matrix-proficient lineages [21–24]. In fact, a pioneering microbial evolution study on *Pseudomonas fluorescens* has already pointed towards dynamic evolutionary interplay between cooperation and exploitation in a biofilm mat [25], where emergence of cellulose overproducer (Wrinkly) allowed mat formation, but also created an opportunity for exploitation by non-producers (Smooth), eventually leading to so called 'tragedy of the commons' [4, 26, 27].

Taken together, biofilms are a suitable model to understand social interactions in an evolutionary time scale [23, 28–31]. Modelling and empirical data confirm that mutualism (beneficial to both actor and recipient) and altruism (beneficial to recipient but not to actor) play crucial roles in biofilm enhancement [32] but at the same time can lead to biofilm destabilization [25]. Can cooperators evolve tactics to evade exploitation and in turn, can cheats utilize evolution to enhance their selfish actions?

Recent studies showed that in well-mixed environment, cooperators adapt to cheats by reducing cooperation [14, 15, 33]. Such reduction could be achieved by various strategies, for instance decrease in motility [15], down regulation or minimal production in public goods [14, 15, 33], up-regulation of other alternative public goods [14], or bi-stable expression in virulence gene [2]. Interestingly, populations of cooperators often exhibit phenotypic heterogeneity at the single cell level [34, 35]. Therefore, an alternative and simple mechanism to modulate levels of cooperation in a population would be through changes in phenotypic heterogeneity pattern. However, the long-term effects of cheats on costly goods' expression at individual cell level, have never been examined. Understanding how heterogeneity of gene expression within the population is affected in the presence of cheats would provide better insight on microbial adaptation and stress response mechanisms.

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Here, we address this question using pellicle biofilm model of *Bacillus subtilis* [36, 37]. Pellicle formation in *B. subtilis* involves, amongst others, aerotaxis driven motility and subsequent matrix production [38]. Aerotaxis is important for oxygen sensing to aid cells reach the surface, while matrix formation is significant to sustain cells to adhere to the surface and to each other. Exopolysaccharide (EPS) is a costly public good in *Bacillus subtilis* biofilms [10, 18, 39] and is heterogeneously expressed during biofilm formation with approximately 40% of cells exhibiting the ON state [39, 40]. We aimed to investigate the cheat-dependent alteration related to phenotypic heterogeneity in *eps* expression by the producer.

We reveal that cheating mitigation by the EPS producers involves a shift in phenotypic heterogeneity towards stronger *eps* expression, which can be achieved by a loss-of-function mutation in a single regulatory gene. Our study uncovers an alternative anti-cheating mechanism based on changes in public goods' expression pattern and highlights meandering trajectories prior cooperation collapse.

MATERIALS AND METHODS

Bacterial strains and culture conditions

Strain *B. subtilis* 168 $P_{hyperspank}$ -mKATE P_{eps} -GFP (TB869) was obtained by transforming the laboratory strain, *B. subtilis* 168 $P_{hyperspank}$ -mKATE (TB49) [10, 18], with genomic DNA from NRS2243 (*sacA*:: P_{epsA} -*gfp*(Km)*hag*::cat) and selecting for Km resistance. The $\Delta rsiX$ strain with fluorescence reporters (TB959) was obtained by transforming TB869 with genomic DNA isolated from BKE23090 (168 *trpC*2 $\Delta rsiX$::erm) [41]. Strains were maintained in LB medium (Lysogeny Broth (Lennox), Carl Roth, Germany), while 2×SG medium was used for biofilm induction [10]. The Δeps strains (TB608) was created previously [10].

Experimental evolution

Eight biological replicates of the co-cultures of 1:1 ratio of *B. subtilis* TB869 and TB608 were grown in 48-well plate containing 1ml 2×SG medium at 30°C for two days. Pellicles were harvested into Eppendorf tubes containing 500 μ l sterile 2×SG medium and 100 μ l of sterile glass sand, vortexed for 90 seconds, 10 μ l fraction was transferred into 1ml 2×SG medium of a 48 well plate and incubated at 30°C static condition for two days. Such growth cycle was continuously repeated 35 times. As a control treatment, four biological replicates of mono-cultures of *B. subtilis* TB869 were evolved using the same transfer method. Every 5th transfer (5 growth cycles), harvested cultures were mixed with 15% glycerol and stored at -80° C.

Population ratio assay

At every 5th transfer, pellicle biofilm productivities and relative frequencies of mutants and WT were qualitatively assessed (colony forming units (CFU)/ml) using LB agar containing selective antibiotics. LB agar plates were incubated at 37°C for 16 h and colonies were counted. Three

single clones of WT and of Δeps per population per timepoint were isolated from plates and stored at -80°C in the presence of 15% glycerol.

Pellicle competition assay/Fitness assay

Competition assays were performed as previously described [10]. Specifically, strains of interest were premixed at 1:1 ratio based on their OD_{600} values and the mixture was inoculated into 2×SG medium at 1%. Cultures were grown for 48 h under static conditions at 30°C and their relative frequencies were accessed using CFU counts (and selective antibiotics).

Stereomicroscopy to assess competition of WT and $\Delta rsiX$ against Δeps

Fluorescent images of pellicles were obtained with an Axio Zoom V16 stereomicroscope (Carl Zeiss, Jena, Germany) equipped with a Zeiss CL 9000 LED light source and an AxioCam MRm monochrome camera (Carl Zeiss) and HE eGFP (excitation at 470/40 nm and emission at 525/50 nm), and HE mRFP (excitation at 572/25 nm and emission at 629/62 nm) filter sets. Images were taken at 3.5× and 55× magnifications. The exposure times for green and red fluorescence were set up to maximal possible values before reaching overexposure, using range indicator function. Zeiss software was used to obtain overlaid, artificially colored images of both fluorescence channels.

Qualitative assessment of eps expression pattern via laser scanning confocal microscopy

Single isolates of evolved WT (TB869) obtained from population ratio assay were allowed to form 1-day old pellicle. Harvested pellicles were subjected to microscopic analysis using an Axio Observer 780 Laser Scanning Confocal Microscope (Carl Zeiss) equipped with a Plan-Apochromat 63×/1.4 Oil DIC M27 objective, an argon laser for stimulation of fluorescence (excitation at 488 nm for green fluorescence and 561 nm for red fluorescence, with emission at 528/26 nm and 630/32 nm respectively). Zen 2012 Software (Carl Zeiss) and FIJI Image J Software [42] were used for image recording and processing, respectively.

Flow cytometry and data analysis

Frozen stocks of evolved populations were transferred onto LB-agar plates containing kanamycin (5µg/ml) to select solely for WT colonies. The plates were incubated overnight at 37°C, followed by inoculation of 10 randomly selected single colonies into 2×SG medium. After 24h-incubation at 30°C, the pellicles were harvested, sonicated and diluted accordingly. Flow cytometry was performed using BD FACSCanto II (BD Biosciences). To separate bacterial cells from noise, mKate-fluorescence (constitutively expressed reporter) and GFP-fluorescence (P_{eps} -GFP promoter fusion) were recorded, gating was setup for mKate-positive objects and GFP signal was measured within these objects. Histograms of P_{eps} -GFP were created in OriginPro using the same binning intervals for all samples. To remove sample size differences (different amounts of measured objects) histograms were normalized to maximum count, described as Normalized Frequency. To obtained an average distribution image of *eps* expression within populations, a mean count for each histogram bin was calculated (by averaging individual counts within this bin obtained for single isolates), resulting in mean distribution of single cell level *eps* expression per

population. Such 'averaged' histograms were used solely for visual representation of data and not for statistical analysis.

Genome re-sequencing and genome analysis

Genomic DNA of single isolates from selected evolved populations were extracted using Bacterial and Yeast Genomic DNA kit (EURx) directly from –80°C stocks grown in LB medium for 5 h at 37°C with shaking at 220 rpm. For population sequencing analysis, approx. 100 colonies belonging to the evolved populations were harvested into 2ml LB broth and incubated at 37°C shaking at 220 rpm for 2-3 h. Re-sequencing was performed on an Illumina NextSeq instrument using V2 sequencing chemistry (2x150 nt). Base-calling was carried out with "bcl2fastq" software (v.2.17.1.14, Illumina). Paired-end reads were further analyzed in CLC Genomics Workbench Tool 9.5.1. Reads were quality-trimmed using an error probability of 0.05 (Q13) as the threshold. Reads that displayed \geq 80% similarity to the reference over \geq 80% of their read lengths were used in mapping. Quality-based SNP and small In/Del variant calling was carried out requiring \geq 10× read coverage with \geq 25% variant frequency. Only variants supported by good quality bases (Q \geq 30) on both strands were considered. Gene functions (product names) in *SI Appendix* Datasets were reported based on SubtiWiki [43].

Statistical analysis

Statistical differences between two experimental time-points of the same experimental groups (e.g. changes in relative frequencies of WT and Δ*eps* during evolution) were accessed using Pair-Sample t-Test. Statistical differences between two experimental groups were calculated using Two-sample t-Test. To compare multiple samples with WT, we used One-way Repeated Measures ANOVA, and Dunnett Test. ANOVA and Tukey Test was used for multiple samples comparisons.

For analysis of P_{eps} -GFP expression in evolved populations, we used 10 randomly picked single colonies, cultivated from the frozen stocks (10th transfer) on LB-agar plate with appropriate selection marker (selecting against Δeps). All Flow Cytometry data of P_{eps} -GFP expression were transferred to histograms, and fitted to Gauss function. Differences in average *eps* expression per population compared to WT_{anc}, and differences in single-cell level distribution of *eps* expression compared to WT_{anc} were calculated using One-way Repeated Measures ANOVA, and mean comparison by Dunnett test. Deviation from WT-like distribution were assessed from changes in Adjusted R-Squared (Adj. R. Sq) values for Gauss fitting of Flow Cytometry data. All

evolved populations, where average Adj. R. Sq was significantly lower compared to WT ancestor, were suspected to have evolved different phenotypic heterogeneity pattern of *eps* expression. Corresponding histograms were visually inspected, classified as potentially bimodal and subjected to multiple peak fitting. In all such cases, fit quality was improved (Adj. R. Sq>0.98). Mean expression for *eps*-low and *eps*-high subpopulations were compared by ANOVA, Tukey test. No statistical methods were used to predetermine sample size and the experiments were not randomized. One data point was removed from the P_{eps}-GFP Flow Cytometry dataset of $\Delta rsiX$ as a significant outlier (P<4.5x10⁻⁷) confirmed by Grubbs test. All statistical tests and data fitting were performed using OriginPro 2018 software.

RESULTS

Cheaters modulate evolution of phenotypic heterogeneity of eps expression in the WT

Exopolysaccharide (EPS) is one of the major components of *B. subtilis* biofilm matrix and mutants deficient in EPS production (Δeps) are not able to form pellicle biofilms (Supplementary Fig. 1). In line with previous results [10, 18, 39], we confirmed that the Δeps can take advantage of EPS-producing wild type (WT) and incorporate into the pellicle biofilm, resulting in lower productivity of the WT (Supplementary Fig. 1b) and reduced surface complexity of the pellicle (Supplementary Fig. 1a). Interestingly, despite pellicles formed by the WT+ Δeps lacked surface complexity and were more fragile compared to the WT monoculture pellicles (as easily observed during sampling), the total numbers of viable cells (our productivity measure) in the WT and mixed pellicles, were similar (Supplementary Fig. 1b). This indicates high carrying capacity of the WT to support surface colonization by Δeps .

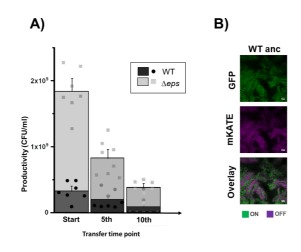


Fig. 1 Pellicle productivity and phenotypic heterogeneity. a Total colony forming unit per ml of WT and Δeps in 48 h old pellicles non-evolved (Start) (n=9), after experimental evolution at 5th (n=8 populations) and 10th transfer (n=7 populations). One population after 10th transfer was unable to form pellicle attributed to WT being totally outnumbered by Δeps . Dots represent data obtained for all individual populations, while columns represent averages. Error bars correspond to standard error. Levels of Δeps at the start, 5th and 10th transfers were 82%, 76% (p<0.29 compared to start, Pair-Sample t-Test) and 83% (p<0.42 compared to start, Pair-Sample t-Test). **b** Pellicles formed by WT^{mKATE} P_{eps}-GFP viewed under the confocal laser scanning microscope. Cells constitutively expressing mKATE are represented in magenta (OFF cells) and *eps*- expressing cells (ON cells) are represented in green. Scale bar 10µm.

We were further interested if such social cheating could leave a phenotypic or genetic fingerprint in the population of the wild type *B. subtilis*. Previous studies have shown that cooperators can adapt to presence of cheats for example by decreasing the amount of released public goods and therefore minimizing cheating opportunities [2, 14, 15]. As *B. subtilis* exhibits phenotypic heterogeneity in *eps* matrix gene expression [39, 40] (Fig. 1a), we investigated how such heterogeneous expression is influenced by the presence of cheats in an evolutionary perspective. To address this question, we co-cultured the EPS producers (wild type- WT) and cheaters (Δeps) for 10 biofilm growth cycles (~60 generations) starting at 1:1 ratio (see methods). Based on previous studies [21, 44], we assumed that this evolutionary timeframe will be sufficient for evolution of adaptive mechanisms in the WT in response to social cheating, at the same time preventing the diversification of the WT into a biofilm-deficient morphotype, which can be observed on longer evolutionary timescales [21].

During 10-cycle co-cultivation of WT and Δeps strain, we observed a general trend of declining pellicle productivity (Fig. 1b). The relative frequency of EPS non-producers in biofilms was maintained at the high level across all parallel populations (82% at the start, 76% after 5th transfer with p<0.29, and 83% after 10th transfer with p<0.42) (Fig. 1b), indicating that the WT strain was constantly exposed to social cheating throughout the experiment.

Using confocal laser scanning microscopy, qualitative assessment of randomly selected isolates revealed that early populations of the EPS producers (5-10 transfer) exhibited different phenotypes compared to the WT ancestral strain (WT_{anc}) (Supplementary Fig. 2). To obtain a quantitative comparison of single-cell level expression of *eps* in the WT_{anc} vs evolved WT populations, we performed flow cytometry measurements of P_{*eps*}-GFP harboring strains in pellicles formed by 10 randomly selected isolates per population (Supplementary Fig. 3).

First, we noticed that in most strains isolated from the control evolved populations exhibited an increase in phenotypic heterogeneity in *eps* expression as compared to the WT ancestor (Fig. 2A,

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Fig. 2B., Supplementary Fig. 3, Supplementary dataset 1). Specifically, while single cell level distribution of P_{eps} -GFP in the WT ancestor, ideally fitted the Gauss function (Adj. R. Sq = 0.99 ±0.01), this was no more the case for the populations of WT evolved without cheater (C1, C2 and C4; Adj. R. Sq: 0.88-0.93). Similar change was noticed for two WT populations evolved with cheaters (Pop1 and Pop8; Adj. R. Sq: 0.87-0.94). In the aforementioned populations, the *eps* expression was rather bimodal, distributed between low-*eps* and high-*eps* subpopulations (Fig. 2A, Fig. 2B., Supplementary Fig. 3, Supplementary dataset 1). Importantly, these bimodal populations evolved alone (C1, C2 and C4) or coevolved with Δeps (Pop1 and Pop8), were similar in terms of *eps* expression levels or ratios of low-*eps* and high-*eps* subpopulations (Supplementary dataset 1). In addition, an average within-population *eps* expression increased in four out of seven populations that evolved with cheater, but only in one control population evolved without cheats (Fig. 2C., Supplementary dataset 1).

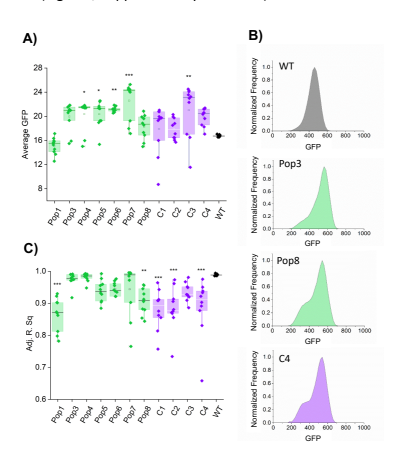


Fig. 2 Evolutionary changes in phenotypic heterogeneity pattern and expression of *eps*. a Changes in distribution of P_{eps} -GFP signal in co-evolved and evolved WT populations compared

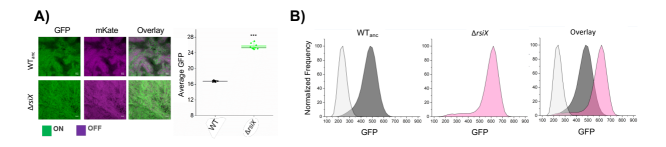
to the WT_{anc} manifested as a decline of adjusted R-square values for Gauss model fitting. **B** Flow cytometry analysis showing average distributions of P_{eps}-GFP in WT_{anc} (dark grey), WT evolved with cheaters, Pop3 and Pop8 (green) and WT evolved without cheaters, C4 (purple). **c** Average P_{eps}-GFP expression levels in co-evolved and evolved WT populations compared to the WT_{anc}, calculated from mean values of P_{eps}-GFP expression within 10 single isolates. For a and c panels *p<0.05; **p<0.01: ***p<0.001 compared to the WT_{anc} (One-way Repeated Measures ANOVA, Dunnett Test). All data and corresponding p values are provided in Supplementary data 1.

Altogether, most populations coevolved with cheater showed an increase in *eps* expression levels, retaining WT-like phenotypic heterogeneity pattern. On the contrary, in majority of control populations (evolved without cheaters) phenotypic heterogeneity level increased, without significant increase in mean *eps* expression (Fig. 2, Supplementary Fig. 3, Supplementary dataset 1).

Mutations in rsiX lead to high-eps phenotype

To unravel the genetic basis of the high-*eps* phenotype that evolved in presence of cheats, several single isolates from the evolved populations were subjected to genome resequencing (for details see methods). The comparative analysis of sequencing data revealed that Population 3 and Population 7, co-evolved with Δeps , shared mutations in *rsiX* gene (Supplementary dataset 2). The *rsiX* gene encodes for an anti-sigma factor that controls the activity of extracellular cytoplasmic function (ECF) sigma factor X which is involved in cationic antimicrobial peptide resistance important for cell envelope stress response [45]. Detected mutations resulted either in substitution of Valine 106 to Alanine or frameshift mutations in Serine 104 or Isoleucine 347 that could lead to change or loss of anti-SigX function. Indeed, we were able to recreate the evolved high-*eps* phenotype in the pellicle solely by deleting the *rsiX* gene in the WT ancestor (Fig. 3a,b). Interestingly, a different type of frameshift mutation in Lysine 200 was found in one population of evolved WT alone but this population demonstrated a bimodal phenotypic heterogeneity pattern (Fig. 2, Supplementary Fig. 3, Supplementary dataset 1), suggesting that

only certain types of mutations in *rsiX* lead to the uniform shift in *eps* expression or additional



mutations have antagonistic effects in this isolate.

Fig. 3 Effect of *rsiX* **deletion on** *eps* **expression in pellicles. a** Qualitative assessment of *eps* gene expression based on confocal laser scanning microscopy of pellicles formed by $\Delta rsiX$ showing high-*eps* compared to WT_{anc}. Cells constitutively expressing mKATE (OFF) are shown in magenta and *eps*- expressing cells (ON) are represented in green. Scale bar 10µm. Right panel: Average P_{eps}-GFP expression levels in WT_{anc} and $\Delta rsiX$ the WT_{anc} calculated from mean values of P_{eps}-GFP expression within 10 single isolates. ***p<0.001 (Two-Sample t-Test). All data and corresponding P values are provided in Supplementary data 1. b Flow cytometry results showing average distribution of fluorescence intensities of WT_{anc} cells (dark grey), $\Delta rsiX$ cells (pink) and overlay of the 2 in comparison to WT non-labelled (light grey).

Mutation in rsiX contributes to competitive advantage of producer strains against cheats

As mutation in *rsiX* resulted in high-*eps* phenotype that may be linked to elevated secretion of EPS, we hypothesized that $\Delta rsiX$ producers could support the spread of cheats. To better understand how ancestor WT and $\Delta rsiX$ interact with Δeps , we cultivated the Δeps in presence of EPS-containing supernatants obtained from the WT and $\Delta rsiX$ (Supplementary Fig. 4). Both supernatants could partially restore pellicle formation by Δeps resulting in similar productivities of Δeps , thereby not supporting our hypothesis on improved performance of the mutant in presence of high-*eps* $\Delta rsiX$ strain.

In order to determine the effect of *rsiX* deletion on fitness of the WT in presence of cheats, we performed a series of competition assays. Apparently, the $\Delta rsiX$ showed two-fold increase in relative frequency (40%) (Fig. 4a, Supplementary Fig. 5) when competed against the Δeps , as compared to the WT ancestor (20%). Additionally, even higher fitness improvement was observed for the WT co-evolved with cheats 5th transfer and 10th transfer, mutually with

occurrence of high-*eps* phenotype in those populations. This was not the case for the WT evolved alone at 5th transfer (20%) (Fig. 4a, Supplementary Fig. 5). These results suggest that *rsiX* mutation, which is associated with high-*eps* phenotype, does not fully explain, but contributes to the early improvement of WT competitive strategies against cheats.

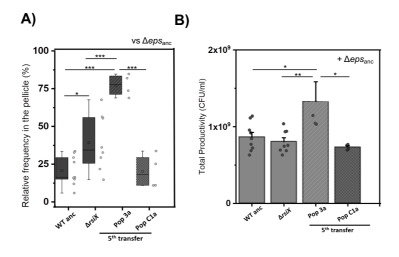
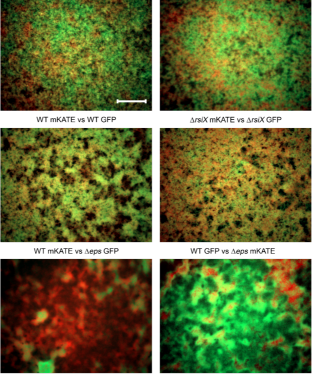


Fig. 4 Performance of evolved WT and $\Delta rsiX$ **in co-cultures with** Δeps_{anc} **. a** Relative frequencies of single isolates belonging to producer populations (WT_{anc} (n=9), $\Delta rsiX$ (n=8), WT evolved with (n=4) and without cheaters (n=4)) in mixed pellicles with Δeps_{anc} . **b** Productivity assay based on total CFU/mI of pellicles of co-cultures of Δeps_{anc} and single isolates belonging to producer populations (WT_{anc} (n=9), $\Delta rsiX$ (n=8), WT evolved with (n=4) and without cheaters (n=4)). Mean is represented in square within the box plots; median is denoted by horizontal line inside the boxes; whiskers represent the min and max; Error bars in bar graph are based on Standard error; single dots represent the individual data points. For a and b panels: *p<0.05; **p<0.01: ***p<0.001 (ANOVA, Tukey Test).

It is worth to mention that we could not detect any significant fitness costs or benefits linked to *rsiX* deletion in pairwise competition between $\Delta rsiX$ and WT in the liquid medium (Supplementary Fig. 6; relative fitness of $\Delta rsiX = 1.00 \pm 0.02$ S.D.). Furthermore, we did not observe significant differences in productivities of WT and the $\Delta rsiX$ mutant, when grown in monoculture pellicles (Fig. 4b), indicating that positive effect of *rsiX* mutation only manifests in presence of cheats. Similarly, different relative frequencies of Δeps in pellicles formed by the ancestor or evolved matrix producers, did not result in different productivities of mixed pellicles (Supplementary Fig. 7). These results suggest that high-*eps* phenotypes are vested on the

increase in *eps*-expressing cells or limiting the spread of cheats but do not result in an increase in total yield.

It was previously demonstrated that increased matrix production can allow favorable positioning of a bacterial strain in the biofilm, thereby providing fitness advantage [46]. To test whether high*eps* phenotype can allow better positioning of the $\Delta rsiX$ in presence of Δeps , we visualized 48 h grown pellicles formed by $\Delta rsiX$: Δeps and WT: Δeps mixtures inoculated at 1:1 initial frequencies (Fig. 5, Supplementary Fig. 8). While WT and Δeps were 'well-mixed' with both strains present on the oxygen-rich surface of the pellicle, the $\Delta rsiX$ strain clearly dominated over the Δeps occupying majority of the biofilm surface and marginalizing the Δeps into small clusters (Fig. 5, Supplementary Fig. 8). Therefore, deletion of *rsiX* and an associated high-*eps* phenotype provides fitness advantage in the presence of Δeps most likely by allowing the EPS producers to occupy upper, oxygen-rich layers of the pellicle. Therefore, *rsiX* frameshift mutation found in certain coevolved WT populations could be an adaptive mechanism to resist cheating by EPS-deficient strain.



∆*rsiX* mKATE vs ∆eps GFP

 $\Delta rsiX \text{ GFP vs } \Delta eps \text{ mKATE}$

Fig. 5 Effect of *rsiX* **deletion on positioning of EPS producers in the pellicle.** Competition assay between WT+ Δeps and $\Delta rsiX+\Delta eps$. Strains labelled with constitutively expressed GFP and mKate proteins, were inoculated in 1:1 initial frequency, pellicles were cultivated for 48 h at 30°C and visualized using stereomicroscope. Upper panels represent controls (two isogenic WT or $\Delta rsiX$ strains labelled with different fluorescent markers), middle panel represents pellicles formed by WT+ Δeps and bottom panels represent pellicles formed by $\Delta rsiX+\Delta eps$, each in two alternative combinations of fluorescent markers. Scale bar corresponds to 500 µm.

High-eps phenotype serves as a transient adaptive response to cheating

As the evolved WT isolates carrying point mutation in *rsiX* as well as the recreated $\Delta rsiX$ mutant (in ancestral genetic background) performed better in competition with Δeps as compared to the WT_{anc}, we reasoned that loss-of-function mutation in *rsiX* together with an associated high-*eps* phenotype, might be an efficient evolutionary strategy against social cheating. Surprisingly however, prolonged evolution experiment eventually led to so called 'tragedy of the commons' as the Δeps mutant took over in 6 out of 8 populations, completely abolishing the pellicle formation (Supplementary Fig. 9).

To investigate the genetics behind this phenomenon all evolved WT populations from the last evolutionary time point (or the last time point prior the collapse) were re-sequenced. Curiously, in contrast to the WT populations that were outcompeted by Δeps , both WT populations which resisted the invasion (Pop5 and Pop8), carried mutations in *yvrG* gene (Supplementary Fig. 9, Supplementary Dataset 2) encoding for two-component histidine kinase involved in cell wall process. Finally, the *rsiX* mutation was not detected neither in the last populations before the collapse, with an exemption of population 7, nor at the last transfer point for the non-collapse populations (Supplementary dataset 2), implying that this mutation was lost in the late populations.

DISCUSSION

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Studies on evolution of cooperative behavior is important to understand how social behaviors are shaped in longer time scale. Moreover, exploring long term consequences of exposure to cheating allows to better understand how cooperation prevails in nature where environmental stress and exploitation exist inherently. Here, we took a reductionist approach, focusing on evolution of a single cooperative trait - the expression of *eps*, which plays a crucial role in biofilm lifestyle of *B. subtilis* and other bacteria. As we focused on the single cell level expression of *eps* in multiple single strain, isolated from the ancestral or evolved populations, we could obtain a multi-level insight into evolutionary changes in *eps* expression. Our study revealed previously observed population-level diversification of matrix genes expression, indicating the strainindependence and reproducibility of adaptation in biofilms [21, 24]. Strikingly, next to population-level diversification, we also observe an increase in phenotypic heterogeneity of *eps* expression within single isolates (Fig. 6). Based on co-culture studies performed for WT and Δeps (this work) as well as for WT and spontaneously evolved biofilm-deficient lineage [21], we believe that low-eps subpopulations may be acting as conditional cheater, supported by 'hypercooperative' subpopulations of high-eps. It remains to be determined whether increased levels of eps expression translate into higher amount of released EPS, but based on previous studies it is likely to be the case [44].

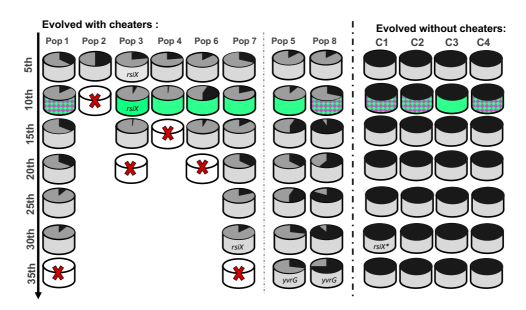


Fig. 6 Changes in relative frequencies and *eps* expression pattern during evolution with and without cheaters. Summary figure shows the population dynamics based on producer and cheater frequency per population from 5th transfer to 35th transfers for populations evolved with cheaters with collapse (Pop 1, 2, 3, 4, 6 and 7) and without collapse (Pop 5 and 8) and populations evolved without cheaters (C1, C2, C3, C4) with indications of phenotypes based on high-*eps* or heterogenous *eps*-expression. Key mutations on single clone level of evolved WT and evolved Δeps or population level are specified. The **rsiX* mutation differs from mutation observed in strains evolved with cheaters.

Previous evolution studies on cheater-cooperator interactions in spatially structured environment showed cheater mitigation via minimization of the cooperative trait [2, 14, 15]. On the contrary, here we show that cooperators respond to cheating by intensifying the cooperative behaviors through uniform shift towards higher *eps* expression. Further molecular analysis of the high-*eps* isolates strongly suggests that this phenotype is triggered by loss-of-function mutation in *rsiX* gene. The product of *rsiX* represses the activity of ECF sigma factor, SigX that is involved in cell envelope stress response against cationic antimicrobial peptides [47]. Importantly, SigX has been previously shown to induce expression of *epsA-O* in *B. subtilis* via a complex regulatory pathway involving Abh and SIrR [48], explaining the observed enhanced in *eps* gene expression in *rsiX* mutant. Another example of matrix overproduction via ECF adaptation was also reported in Gram-negative bacterium *Pseudomonas aeruginosa* where mutations in another ECF called AlgT led to alginate overproduction and increased resistance to antimicrobials [49]. Therefore, adaptive boosts in matrix production through modulation of ECF is not exclusive for *B. subtilis* but seems to occur also in medically relevant Gram-negative pathogens like *P. aeruginosa*.

In contrast to previous studies that addressed long term cheating on diffusible siderophores [50– 53], we explored evolutionary interplay between biofilm producers and non-producers in structured environment. Our results support previous observations on evolution of specific cheating-resisting mechanisms in co-operators, pointing towards ubiquity of this phenomenon. In addition, our work brings up two major findings 1) matrix producers can adapt to matrix nonproducers by shifting phenotypic heterogeneity towards increased levels of matrix-expression,

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2) high-*eps* phenotype is associated with favorable positioning of the matrix producers in the biofilm in presence of cheats, thereby limiting their numbers, 3) high-*eps* anti-cheating strategy is a short-term solution followed by tragedy of the commons. As EPS-deficient strain took over in all but two mixed populations (including populations, without *rsiX* mutation and homogenous shift towards higher *eps* expression), we do not interpret the collapse as a direct consequence of mutation in *rsiX* gene. However, we argue that an emergence of several matrix overproducing lineages, may facilitate the spread of cheats [21], especially if a substantial number of cells within the high-*eps* lineage serves as facultative (phenotypic) cheaters. As recently demonstrated, EPS-deficiency is not a dead-end strategy for *B. subtilis* population, because alternative EPS-independent biofilm formation strategies can emerge by single amino acid change is TasA [44]. It remains to be discovered whether shifts in phenotypic heterogeneity in response to long term cheating is general phenomenon that applies to different types of public goods.

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Competing Interests

The authors declare that there are no competing financial interests in relation to the work described.

Authors contributions

Á.T.K. conceived the project, M.M., A.D., S.B. and D.S. performed the experiments. G.M contributed with methods. M.M., A.D. and Á.T.K. wrote the manuscript, with all authors contributing to the final version.

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Fig S1. Pellicle biofilm formation and total productivity assessment.

a) Pellicle biofilms formed by mono-cultures of WT, Δeps , and co-culture of WT+ Δeps in 2xSG medium incubated for 48 hours at 30°C recorded using Samsung Galaxy S6 Phone Camera. Scale bar, 1cm. **b)** Productivity assessment based on CFU/ml were performed on pellicle biofilms of WT, Δeps , WT+ Δeps co-culture and control co-culture of two WT strains (WT^{mKate} used in the evolution experiment, and non-labelled WT). Productivity of Δeps dramatically increased (p<3x10⁻⁵, Pair-Sample t-Test) when co-cultured with WT, while productivity of WT decreased (p<0.002, Pair-Sample t-Test) in the presence of Δeps , indicating the ability of the mutants to act as cheaters (n=5, error bar based on standard error).

Fig S2. Qualitative assessment of *eps* gene expression based on confocal laser scanning microscopy. Pellicles formed by randomly selected WT strains 168 mKATE P_{eps} -GFP evolved in the presence of cheaters (Pop1-8) and in the absence of cheaters (C1-C4) were visualized using confocal laser scanning microscope. Cells constitutively expressing mKATE are represented in magenta (OFF cells) and *eps*- expressing cells (ON cells) are represented in green. Scale bar 10µm.

Fig S3. Single cell level distribution of *eps* **expression in 10 randomly selected single isolates from WT**_{anc} **and evolved WT populations.** Flow low cytometry data (BD Facscanto II, BD biosciences) showing single cell level distributions of fluorescence intensity of 24-hour old pellicles established by 10 randomly selected single isolates from populations of WT evolved in the absence (C1-C4) or presence of cheaters (Pop1-8) as well as autofluorescence distribution in non-labelled WT control. **Fig S4. Complementation assay of** Δ*eps* **with supernatant from** Δ*rsiX* **or WT.** Productivity data of pellicles produced by the complementation showed that hyper ON Δ*rsiX* mutant does not contribute to improved performance of Δ*eps*. Mean is represented in square inside the box plots; median is denoted by horizontal line within the boxes; whiskers represent the min and max (n=3 for Δ*eps*; n=6 for Δ*eps*+SMs). ***p<0.001 compared to Δ*eps*; Δ*eps*+ WT SM is not significantly different from Δ*eps*+ Δ*rsiX* SM (p=0.58) (ANOVA, Tukey Test).

Fig S5. Performance of $\Delta rsiX$ and evolved and co-evolved WT in mixed pellicles with Δeps_{anc} . Pellicle competition assay of single clones belonging to producer populations (WT_{anc} (n=9), $\Delta rsiX$ (n=8), WT evolved with (n=4) and without cheaters (n=4)) against Δeps_{anc} . Mean is represented in square inside the box plots; median is denoted by horizontal line within the boxes; whiskers represent the min and max; single dots represent the individual data points (n). ***p<0.001 compared to the WT_{anc} (One-way Repeated Measures ANOVA, Dunnett Test).

Fig S6. Fitness effects of *rsiX* deletion. Selection rate based on fitness assay in pairwise competition of $\Delta rsiX$ and WT ancestor showing no significant fitness cost brought about by *rsiX* mutation. Relative fitness of $\Delta rsiX$ is 1.00 ± 0.024 SD. Mean is represented in square within the boxplots; median is denoted by horizontal line inside the boxes; whiskers represent the min and max.

Fig S7. Pellicle productivity of monocultures. Total CFU/ml of pellicles produced by monocultures of WT_{anc} (n=7), $\Delta rsiX$ (n=3) and evolved with cheaters (n=8) from population 3 (5th, 10th) and 7 (30th) and single clones of WT evolved without cheaters (n=4). Mean is represented in square; median is denoted by horizontal line inside the box; whiskers represent the min and max; single dots represent the individual datapoints (n). All p values >0.05 (ANOVA, Tukey Test).

Fig S8. Effect of *rsiX* **mutation on positioning in the pellicle.** Competition assay between fluorescently WT+Δ*eps* and Δ*rsiX*+Δ*eps*. Strains labelled with constitutively expressed GFP and mKate proteins, were inoculated in 1:1 initial frequencies, pellicles were cultivated for 48h at 30°C and visualized using stereomicroscope. Upper panels represent controls (two isogenic WT or Δ*r*siX strains labelled with different fluorescent markers), middle panel represents pellicles formed by WT+Δ*eps* and bottom panels represent pellicles formed by Δ*rsiX*+Δ*eps*, each in two alternative combinations of fluorescent markers. Well size = 1.5cm.

Figure S9. Productivity changes during short and long-term co-evolution of WT+ Δeps . Pellicle Total colony forming unit per ml of WT and Δeps in 48-hour old pellicles **A**) non-evolved (start) (n=9), after experimental evolution at 5th (n=8 populations) and 10th transfer (n=7 populations) One population after 10th transfer was incapable to form pellicle attributed to WT being outnumbered by Δeps . **B**) Line graph showing the fate of populations. Data was obtained from CFU assay using selective antibiotic marker, Kanamycin selecting for WT and Tetracycline for selecting Δeps .

Supplementary dataset 1. Statistical comparisons of *eps* expression and single cell level distribution within ancestral and evolved populations.

Supplementary dataset 2. List of detected SNPs of evolved strains and their functional analysis.

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