Single-Cell Microfluidics to Study the Effects of Genome Deletion on Bacterial Growth Behavior

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ABSTRACT: By directly monitoring single cell growth in a microfluidic platform, we interrogated genome-deletion effects in Escherichia coli strains. We compared the growth dynamics of a wild type strain with a clean genome strain, and their derived mutants at the single-cell level. A decreased average growth rate and extended average lag time were found for the clean genome strain, compared to those of the wild type strain. Direct correlation between the growth rate and lag time of individual cells showed that the clean genome population was more heterogeneous. Cell culturability (the ratio of growing cells to the sum of growing and nongrowing cells) of the clean genome population was also lower. Interestingly, after the random mutations induced by a glucose starvation treatment, for the clean genome population mutants that had survived the competition of chemostat growing and nongrowing cells) of the clean genome population was also lower. Interestingly, after the random mutations induced by a glucose starvation treatment, for the clean genome population mutants that had survived the competition of chemostat culture, each parameter markedly improved (i.e., the average growth rate and cell culturability increased, and the lag time and heterogeneity decreased). However, this effect was not seen in the wild type strain; the wild type mutants cultured in a chemostat retained a high diversity of growth phenotypes. These results suggest that quasi-essential genes that were deleted in the clean genome might be required to retain a diversity of growth characteristics at the individual cell level under environmental stress. These observations highlight that single-cell microfluidics can reveal subtle individual cellular responses, enabling in-depth understanding of the population.

KEYWORDS: single-cell, microfluidics, cell growth, genome reduction, mutation

Synthetic biology has adopted the idea of chassis organisms with reduced genomes that are engineered to efficiently express essential endogenous proteins. The reduced genomes of these chassis present a stable and predictable platform upon which synthetic genes and circuits can be rationally designed and then embedded.1-4 The development of a chassis genome involves identifying a set of essential genes, usually via sequence alignment with strains of the same species, and then deleting anything deemed nonessential.5 This should theoretically have the additional advantage of reducing the associated energy cost (energy per gene) of replicating a larger genome and expressing genes that are not essential for the desired environment (protein burden).5 However, rather paradoxically, genome reductions appear to slow growth rates.6,7 This has been taken as evidence for a reduction in fitness and has led to the suggestion that in addition to essential genes, genomes might also contain quasi-essential genes that function to promote fast growth.2

We know that even in supposedly clonal populations of bacteria there can be unexplained “stochastic” variability in the phenotypical behavior of individual cells.7-9 Indeed, there is evidence from natural microbial communities that a substantial number of cells do not contribute to genetic diversity of subsequent generations, which suggests that not all cells contribute to growth events. This result has been inferred by estimating the effective population size.10 It is not known whether this stochastic variability in phenotype is increased or diminished in minimal genomes, because we typically observe growth at the population level rather than in individuals. Furthermore, synthetic organisms are usually tested in highly controlled laboratory conditions, so it is unclear if the heterogeneity in phenotype is exacerbated by environmental

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The chassis is fundamental to any synthetic biotechnology, therefore not only is it essential to work out the complexities of chassis gene components, it is also important to be able to reliably characterize growth and, where variability occurs, potentially select fractions of the population that offer the highest growth and the least variability.

Microfluidics provides an ideal platform to culture and monitor individual bacterial cells over many generations. It allows multiparameter, near-continuous measurements of dividing cells in high throughput devices with a bewildering range of spatial architectures and with low sample consumption. Recently, we have developed a simple gradient microfluidic system for a rapid bacterial growth inhibition test at the single-cell level. With this approach, quantitative and dynamic growth information about individual cells within a population can be obtained in parallel under a wide range of conditions. Importantly, it provided a powerful platform to mimic naturally occurring habitats of environmental microbes, enabling a close-to-reality analysis of bacterial responses to stresses and consequently the discovery of new phenomena.

In this work, we have exploited single-cell microfluidics to investigate the influence of genome reduction on microbial growth, aiming to reveal the diversity of individual behaviors to gain an in-depth understanding of the population. For this purpose, the lab reference strain Escherichia coli, K12 MG1655 (denoted as the wild type, WT), and the derived genome-reduced strain, MDS42 (denoted as the clean genome, CG), were used. This CG strain is formed by deleting 14.3% of the WT’s genome, with the deleted sections including nonessential genes, mobile DNA elements and cryptic prophages. The CG strain has been reported to be a good chassis for synthetic biology applications because of plasmid stability, low mutation rate, and protein production. However, this low mutation rate appears to hold only when the E. coli are in optimal laboratory condition; Couto et al. showed that under glucose starvation stress, as described in Ferenci et al., mutation rates indexed in the rpoB gene (RifR) or cycA gene (cycR), were orders of magnitude higher in both the WT and CG organisms. Most mutations correlated with a decrease in growth rate and ultimately failed to gain a foothold in the population. A few, however, appeared to remain fit enough to be retained in the population over time. However, a comparison of the diversity in growth characteristics of the WT and CG mutants that can survive these harsh conditions has not been made. Thus, here, we cultured mutant subpopulations in oligotrophic conditions, minimal media supplemented with 1 mM glucose in a chemostat with a dilution rate of 0.1 h⁻¹. This dilution rate meant that only mutants capable of growing at a rate that was greater than or equal to 0.1 h⁻¹ would be retained; the slower growers would be washed out. As described below, using microfluidics allows us to rapidly characterize the diversity in growth characteristics of the mutants that could survive.

To enhance throughput and flexibility of cell handing, a new generation of microfluidic device was developed that consisted of an array of microchambers to allow monolayer cell culture and real-time monitoring of cell growth (Figure 1). Single-cell growth characteristics, such as specific growth rate (μ, h⁻¹), lag time (λ, h), and culturability, were quantified. This enables a direct correlation of growth rate and lag time of individual cells that reveals the population heterogeneity—information that is difficult to obtain using conventional well-plate approaches. For the first time, the effects of genome reduction and random mutation on population heterogeneity were measured, which indicate that the deleted genes are not as redundant as previously believed, but play an important role to resist stress and homogenize the population.
RESULTS

Effects of Genome Reduction on Cell Growth. Growth rates of both WT (i.e., MG1655) and CG (i.e., MDS42) strains have been previously shown to be unequal. We confirmed these findings via batch growth, where the specific growth rate $\mu$ is $1.04 \text{ h}^{-1}$ (or doubling time of 40 min) for the WT strain and $0.78 \text{ h}^{-1}$ (or doubling time of 53 min) for the CG strain (Supporting Information, Figure S1). However, since these measures represented average responses of the population, it is unclear what the main contributor to this discrepancy is. For instance, does the lower growth rate in the CG strain result from a reduced growth of all the cells or only from the slowest growing subset? In this context, microfluidics devices (Figure 1) were employed to characterize the growth kinetics of individual cells, revealing quantitative information on diverse behaviors within the populations.

Figure 2 illustrates typical time-lapse images of single cells as they gradually grew and divided to form microcolonies in the microchambers. From these images, growth curves for randomly selected single cells were obtained (Supporting Information, Figure S2). On-chip culture was under a continuous flow condition, which meant that after a short lag phase, exponential growth was observed and sustained for the duration of the experiment (because the substrates were continually replenished, Supporting Information, Figure S2). This enabled an accurate derivation of colony specific (individual) growth rate ($\mu$) and lag time ($\lambda$) parameters for a given substrate concentration as described in the Materials and Methods.

Figure 3 shows the distributions of such individual $\mu$ and $\lambda$ parameters from 80 randomly picked dividing cells. The average growth rates calculated using the on-chip single cell approach were consistent with the results from batch culture (Supporting Information, Figure S1), with the WT strain ($\mu = 1.0 \text{ h}^{-1}$) growing significantly faster than the CG strain ($\mu = 0.8 \text{ h}^{-1}$) ($p < 0.001$) (Figure 3A). A histogram of the growth rate, $\mu$, of single cells showed that this slower average growth in the CG strain was a result of a majority of CG cells growing slower than those of the WT strain (Figure 3B). Approximately 80% of the WT cells had $\mu > 1.0 \text{ h}^{-1}$ in comparison to only 40% of the CG cells having $\mu > 1.0 \text{ h}^{-1}$ (Figure 3B). Furthermore, it was found >90% of the WT cells showed no lag time (Figure 3C), whereas approximately 80% of the CG cells had a lag time between 0.5 and 3 h. As a result, the mean lag time of the CG strain was significantly longer ($p < 0.00001$, Figure 3A). It should be noted an extended lag time is indicative of the weak ability of a cell to acclimatize to a new environment (e.g., from broth suspension to the surface of a glass substrate), therefore indicating a negative effect of genome reduction on cell growth.

Effects of Starvation Induced Random Mutations. To investigate the impact of emerging genetic variation on the growth kinetics of these strains, we developed two populations of mutants, one from each strain. Each community was created by harvesting mutants from a glucose-limited chemostat that maintained a population at a dilution rate of 0.1 h$^{-1}$ for 21 days. Glucose-limitation has been shown to promote random mutations in strains of E. coli that can be screened via acquired resistance to Rifampicin or Cycloserine, two commonly used antibiotics in lab evolution experiments. We exploited this system to produce organisms with mutations in either the rpoB gene (rifR) or cyaA gene (cycR) in both the WT and CG strain backgrounds. Once harvested from the...
chemostats, mutants from each strain background were designated WT_mut (mutants in the WT background) and CG_mut (mutants in the CG background). We started two new glucose-limited chemostat runs to grow up and maintain each of these communities of mutants, in the absence of any nonmutants. The chemostats were seeded with a subsample from each of the exponentially growing mutant communities, and operated with a dilution rate 0.1 h$^{-1}$ for a period of 20 h, by which time the density of organisms had reached a steady state (OD$_{600}$ = 0.1). Then samples of such cultures were taken for loading into the microfluidic devices.

As shown in Figure 4, the starvation induced random mutations generated contrasting growth effects for the WT and CG strains. The population of WT_mut cells showed a slightly

Figure 4. (A) Average values of the growth rates for the WT and CG strains, and the WT_mut and CG_mut populations together with (B, C) histograms of their distribution. (D) Average values of the lag times for the four populations together with (E, F) histograms of their distribution. Arrows in (B) indicate the subsets of slow growing cells. $p < 0.0001$ (****); $p < 0.0005$ (**); ns denotes not significant.

Figure 5. Scatter diagrams of the lag time $\lambda$ versus the specific growth rate $\mu$ of individual cells for the four populations. Each spot denotes a single cell.
decreased average growth rate (0.93 ± 0.14 h⁻¹) but larger standard deviation in comparison to its parent WT strain (1.07 ± 0.11 h⁻¹) (p < 0.0001) (Figure 4A). The diversity of growth rates in the mutants and the presence of the slower growing mutants (Figure 4B, indicated by the arrows) is perhaps surprising, given that they were sampled from a fixed growth-rate chemostat. In contrast, the average growth rate for the CG mut population (0.99 ± 0.11 h⁻¹) increased and the standard deviation dropped in comparison to the CG strain (0.87 ± 0.19 h⁻¹) (p < 0.0001) (Figure 4A), indicating a narrowing in the distribution of individual growth rates in the CG-mut population (Figure 4C). No significant difference was found in the lag time between the WT-mut population and the WT strain (Figure 4D,E). However, the mean lag time for the CG-mut cells was substantially shorter than that of the CG cells (p < 0.00001) (Figure 4D), since not only is there a high percentage of the CG-mut cells with a lag time ≤1.0 h but also no CG-mut cells showed a lag time >1.5 h (Figure 4F). Clearly, the diversity of mutants that could persist in a chemostat is, in part, dependent on the genetic background of that parent strain.

**Correlated Individual Growth Characteristics Revealed Changes in Heterogeneity.** To further understand how genome reduction affects growth kinetics of the overall population, the individual μ and λ values from these four populations were plotted in a scatter diagram (Figure 5). The diagrams reveal the heterogeneity of the populations, information that cannot be obtained from batch culture. If individual cells performed similarly, i.e., they were homogeneous, then all data spots would cluster due to similarity in both the μ and λ values. In contrast, cellular heterogeneity is marked by an increased amount of scatter. These plots also provided clues as to which parameter drives the diversity of the population and whether there is dependency between these two parameters. Accordingly, individual growth characteristics for the CG strain were the most heterogeneous, with a broad scatter from both the μ and λ parameters (Figure 5A). Interestingly, the CG-mut population were much more clustered compared with their parent strain, indicating that a far more homogeneous population of mutants survived in the chemostat environment (Figure 5B). The WT strain showed diversity in the μ parameter only; almost all the cells of that strain had no detectable lag time λ (Figure 5C). The WT-mut population showed a slightly increased diversity in λ and similar diversity in the μ parameter, thus the range of phenotypes that survived the chemostat was similar for the mutants (Figure 5D).

**Cell Culturability of the Populations.** A large number of nondividing cells were observed in all the populations under investigation and so a measure of culturability was investigated. Long-term observations indicated that these cells did not divide and were thus either quiescent or need a very long lag time (>24 h) to proliferate under the current nutrient condition (Supporting Information, Figure S3). Considering the purpose of this study, such nondividing cells were excluded from the above-mentioned growth kinetics assay.

We defined the culturability as the fraction of cells that grow, which was quantified by enumerating both dividing and nondividing cells. The culturability of the WT strain (62 ± 11%) was similar to that of the derived WT-mut population (63 ± 11%; p > 0.05), both of which are three times higher than that of the CG strain (20 ± 6%; p < 0.005). The culturability of the CG-mut population (45 ± 2%) was twice as high as that of its parent CG strain (p < 0.05). It should be remembered that the chemostat selects for the fittest mutants and so this is to be expected. It is interesting that the WT-mut populations culturability is equal to that of the parent population and so the WT-mut strains’ culturability has not been enhanced by the selection imposed by the chemostat.

**DISCUSSION**

The main motivations for producing genome-reduced strains of bacteria for synthetic biotechnologies are to alleviate protein burden and retain genome integrity. However, some reduced-genome strains have been reported to grow more slowly than their parental strains in batch culture, calling into question their efficiency. Genome integrity, or its effect on growth kinetics in a reduced genome strain has only recently been investigated, and it was found that in the context of metabolic stress, mutations accumulated quickly in these strains. The majority of these mutations ultimately led to less competitive phenotypes in chemostat populations. Therefore, it would be unlikely that most mutants would survive in the long chemostat run. Indeed, in a chemostat that is limited by a single resource, we would expect the single fittest, fastest growing, phenotype, mutant or otherwise, to “win out” and ultimately become monodominant. Yet even in clonal populations there appears to be variation in growth rates and hence phenotypes. What is not known is if that growth variability persists in the populations that remain in the harsh conditions in a chemostat. Thus, here, we have grown supposedly clonal populations of the WT strain (i.e., MG1655) and the derived reduced-genome CG strain (i.e., MDS42CG) in oligotrophic conditions in chemostats. We also grew mixed communities comprising solely of organisms that had acquired mutations in their rpoB gene (rif⁶) or cyaA gene (cyc⁵) for both strains. Each chemostat was sampled and the variability in the growth rates were characterized at the individual cell level using a facile microfluidic platform.

**The Single-Cell Approach Enables Novel Measures of Population Heterogeneity.** The facile microfluidic platform developed allowed us to observe cell growth in real time and quantify properties such as specific growth rate μ and lag time λ of a population at the single-cell level. As illustrated in Figure S, the direct correlation of μ and λ of individual cells depicted key constitutional features of a population. Hidden information, such as overall heterogeneity of the growing cells and characteristics of each subset, are revealed at a glance. In combination with the culturability of a population, a clear relationship was shown where the culturability increased with the decrease of heterogeneity. For instance, the least cultivable CG strain has the greatest degree of heterogeneity. It is apparent that these two parameters can serve as novel measures of robust growth. Notably, this crucial information, especially
the individual $\mu - \lambda$ distributions, are inherently overlooked by
conventional methods, therefore our method could contribute
to more accurate understanding, modeling, and predicting of
the bacterial responses to environmental changes.

The Effects of Genome Reduction on Cell Growth. We
showed that average specific growth rates in the exponential
growth phase obtained from on-chip culture are in excellent
agreement with that from batch culture (Supporting
Information, Figure S1 and Table 1), both of which show
that the CG strain grow slower than the WT strain as reported
previously.2 Importantly, with the ability of obtaining growth
characteristics at single-cell resolution, our study provides the
first measures of heterogeneity and culturability in these
populations. We found that the genome-reduced CG strain not
only had lower culturability than the WT strain but also had
more heterogeneous growth characteristics than the parent WT
strain. This observed increase in heterogeneity was present in
both the growth rate and the lag time of individual cells. Given
that one of the purposes of developing reduced genome chassis
is to increase stability then this variability in growth might be
unexpected. Furthermore, the microfluidics devices were loaded
with cells sampled directly from the chemostats, where the
organisms in the population were in a steady-state of growth.
However, the variability might reflect the ability of individual
microorganisms to respond to changes in local environment via
modulations in gene expression,13,25–31 with more of the CG
organisms being affected by the temporary stress of being
transferred into the microfluidics device. Indeed, functional
analyses of genome reductions in the CG strains show that they
impacted global gene expression in these organisms by altering
transcription of nondeleted regions of these genomes.6 This
could be a crucial contributing factor to the growth
heterogeneity observed in the CG strain.

Interestingly, our discoveries echo a recent study of the
minimal genome organisms derived from Mycoplasma mycoides,
which showed that as a greater number of genes are deleted
from a genome, growth rates slow down.2 This trade-off is the
result of genomes containing quasi-essential genes, which are
thought to be coexpressed with the essential genes for robust
growth.2 These genes could also be essential for increasing
culturability of a strain, features that have not yet been
investigated in any other reduced or minimal genomes strains.
It is noteworthy that quasi-essential genes can be elusive,
especially when comparative genomics are used as the main
strategy for defining the minimal set of required genes.7 The
704 genes that were deleted to create the E. coli (MDS42) CG
strains were targeted largely based on an in silico comparative
genomics strategy to eliminate nonessential gene clusters.5
However, when strains bearing the systematic deletions were
analyzed in depth for function, specifically growth properties in
different environments, it was found that their impact was far
more global than previously anticipated.6 Defects could not be
attributed to single genes, instead clusters or genomic segments
appeared to be involved. These segments impacted growth in
specific environmental niches where they played a role in
building new cellular material and conveying stress tolerance.
In addition, these segments would have been acquired via
horizontal gene transfer, thus comparative genomics, although
a convenient strategy to define deep branching core genes, can
miss recently acquired advantageous genomic changes that appear
to be essential for good growth.6

Mutations at Different Genetic Backgrounds. In a
chemostat, one would expect the fastest growing organisms to
be the most abundant, and ultimately, if all the organisms in the
chemostat community are competing for the same limiting
resource, the fittest organism should theoretically become
monodominant. We grew up populations in chemostats that
had acquired mutations through starvation and maintained the
density at steady state. We know from a previous study24 that
most starvation induced mutations have a deleterious effect on
growth and thus we might expect a significant reduction in the
diversity of growth characteristics in the mutants that survive
and grow to dominate the chemostat. That is exactly what we
found with the CG_mut strains. Indeed, our microfluidic
analysis showed that the average growth rate of cells sampled
from the CG_mut population that grew to prominence in the
chemostat exceeded that of the nonmutants (Table 1).

However, the effect of acquired mutations was less obvious in
the WT_mut population; it had a similar average growth rate and
retained the heterogeneity and culturability of the parent
WT strain. This diversity is perhaps surprising since both
mutant populations are subject to the same stressful chemostat
conditions. Thus, one might expect the chemostat to exert the
same selection pressure on the WT_mut populations as for the
CG_mut population and thus reduce the heterogeneity and
increase cultureability; which does not seem to be the case. The
fact that a wider range of growth characteristics are maintained
and that no one phenotype wins out in the WT_mut population
might suggest that growth is not entirely limited by
the carbon substrate and that the complexity of the genome
allows individuals to find or create niches in a way that the
minimal genomes of the CG_muts cannot. Bearing in mind
that CG strain was derived by deleting “nonessential” parts,
including IS-elements, from the WT strain, and functional
analyses of these deleted regions now show that they impact
important phenotypes such as improved growth, stress
tolerance and metabolic flexibility.6 It stands to reason that
removing these sections could lead to organisms that are in
general less phenotypically robust to mutations and thus a
population where a few CG_muts have a clear advantage of
others in way that does not occur with the WT_muts. It has
previously been shown that if mutations do occur in these two
strain backgrounds, IS elements change their impact on the
overall observed phenotype.22

The reduced genome chassis was conceived based on the
hypothesis that the emergence of new traits, even if favorable,
would be minimal in these systems. This is mainly to keep
control of the process and product of the underlying
biotechnology.33 However, spontaneous mutations do occur in
populations over time, and so, their effect should be
interrogated in these reduced genome strains. Our work here
shows that in clonal populations of the reduced genome strains,
the emerging mutations that are not deleterious and can survive
in a selective chemostat environment are far more homoge-
nous than those that emerge in the wild type populations.

CONCLUSION

In this study, we developed a microfluidic system that enables
fast quantification of growth rates and lag time of E. coli strains
at the single-cell level. With this capability, we demonstrated
the capacity to measure heterogeneity (from the correlated
individual $\mu - \lambda$ distributions) and culturability of a population.
We showed the increased culturability was associated with the
decreased heterogeneity of a population for a reduced genome
strain of E. coli grown in continuous culture in a chemostat. For
a wild type strain and mutants that were derived through
starvation stress, a surprising diversity of growth characteristics was retained in continuous culture populations. These discoveries illustrate that single-cell microfluidics can reveal subtle cellular responses that are obscure when observations are made at the population level.

**MATERIALS AND METHODS**

**Fabrication of Microfluidic Platforms.** Figure 1a shows the structure of the microfluidic platform designed for this study. There are two parallel 50(w) × 10(h) μm channels, between which are eight identical 200(l) × 80(w) × 0.74(h) μm microchambers. The submicron height of the chamber guaranteed cell capture and monolayer cell culture. Several circular pillars (diameter = 6 μm) are added to maintain the chamber height. To prevent cells from passing straight through the chambers during cell loading process (see below), a region with an even lower “head” height (ca. 0.4 μm) was designed at the opposite side of the chamber (i.e., between the medium channel and main chambers).

Common soft lithography techniques were used to fabricate this structure. Briefly, SU-8-2002 and SU-8-3010 photoresist were diluted 1:4 and 1:1 respectively with cyclopentanone and then sequentially used to make features with differing heights, i.e., the low head-height region, microchambers, and two channels. After each photolithography/development step, the SU8 mold was hard baked at 180 °C for 15 min to avoid distortion in subsequent fabrication steps. The structure of the SU8 mould was measured using Bruker Dektak XT Height Profiler. When necessary, the fabrication process was optimized to meet the designed dimensions for each feature. After the silanization of the mold, polydimethylsiloxane (PDMS)/curing agent (Sylgard 184; Dow Corning Co., UK) (10/1) mixture was poured onto it and cured at 80 °C for overnight. Punching holes for both inlet and outlet, then the PDMS replica and clean coverglass were O2 plasma treated before bonding them together to make the microfluidic platform. The ca. 0.4 μm height in parts of the device was verified using microscopic optical absorption measurements of a rhodamine solution, showing that these low height structures did not collapse (Supporting Information, Figure S4).

**Bacterial Strains.** All bacteria strains used in this study are derivatives of E. coli K12. The E. coli wide type strain, MG1655 (F- lambda- ilvG- rfb-50, rph-1)34 was purchased from the DSMZ culture collection (Braunschweig, Germany). The multiple deletion series (MDS) MDS42 strain has been described and characterized in detail previously1 and was purchased from Scarab Genomics (Wisconsin, USA).

**Chemostat Culture.** For each growth experiment, precultures were always started from bacteria that had been cryopreserved at −80 °C as glycerol stocks. Cells were first streaked onto LB agar plates and after overnight incubation at 37 °C a colony was transferred into 50 mL of M9 minimal medium that was supplemented with 10 mM glucose. When this preculture had reached an OD600 of 0.1, 8 mL of the culture was transferred to a computer-controlled glass bioreactor (modified miniBio 500, Applikon Biotechnology, Delft, NL myControl, Applikon Biotechnology Inc., CA) for continuous culture, with a working volume of 0.35 L. The chemostat was fed M9 minimal medium supplemented with 1 mM of glucose. Air passed into the reactor was via a 0.20 μm filter. The impeller speed was set at 80 rpm, the temperature was 37 °C, and pH within the vessel was maintained at around 7.0. The medium-feed speed was set to maintain a dilution rate of 0.1 h−1. A steady state OD600 value of about 0.1 was reached approximately 20 h after this manipulation, at which point the cell cultures were ready for loading (after dilution, see below) into microchambers.

**Generating Mutant Populations.** To prepare mutant populations of the MG1655 and MDS42 strains, triplicate populations of each strain were maintained at steady state in glucose limited (M9 medium supplemented with 1 mM glucose at a dilution rate of 0.1 h−1) chemostats for a duration of 21 days. During this time the cultures were assayed for spontaneously occurring mutations at either the rpoB23 or the cyclA20 loci by plating on selective solid media. A 1 mL of subsample from each chemostat was plated and monitored for the number of colonies resistant to the antibiotics Rifampicin or D-Cycloserine. Mutation accumulation data from the chemostat experiments were surveyed to find a population from each strain with a similar mutant fraction. For the MG1655 strain, mutants were sampled from a population with an estimated total of 34 660 mutants/mL (2.8 × 10−2 %). Similarly, for the MDS42 strain, mutants were chosen from a population with an estimate of 38,108 mutants/mL (2.2 × 10−2 %). Both populations had been in continuous, glucose-limited culture for 14 days, and had thus gone through an equivalent number of generations (N.B.: generation time is fixed in a chemostat). At the time of sampling, up to 140 resistant colonies were picked and stored as glycerol stocks at −80 °C. These were recovered in LB medium and pooled to make the WT_mut (mutants derived from a MG1655 chemostat population) and the CG_mut (mutants derived from a MDS42 chemostat population). Before inoculating the microfluidics device, a 1 mL subculture of each population was used to start up a chemostat culture, which grew using the conditions described below.

**On-Chip Bacteria Culture.** The microfluidic platform was initially filled with M9 minimal medium containing 10 μM glucose, which was delivered through the two parallel channels at 10 μL/min with a syringe pump (NE-1000, New Era Pump Systems Inc.) (Figure 1). Prior to introducing to the microfluidic chip, bacterial culture from the chemostat was diluted 10- or 20-fold with glucose-containing fresh medium. After twice centrifuging at 3000 rpm × 5 min and resuspending in the same fresh medium, the sample was loaded into the device through the “loading” channel. The time was ca. 30 min between subsampling from the chemostat and introducing to the microfluidic chip. After loading the cells into the device, the same fresh medium was allowed to flow through either side of the chambers at a suitable flow rate to remove nontrapped cells in both channels. Meanwhile cell density of the microchambers could be adjusted by changing the flow rate. Thereafter, the flow rate was changed to 0.1 μL/min to start on-chip cell culture. This continuous flow culture provided constant nutrient supply to the cells and simultaneously eliminated metabolic waste. The whole microfluidic platform was mounted into a commercial temperature-controlled chamber (Heating Insert P Lab-Tek 2000, PeCon GmbH, Germany) (Figure 1B). It was preheated to 37 °C and mounted on motorized microscope stage. Simple calculations show that at the flow rates used, the medium reaches 37 °C a few seconds after being driven into the platform. Unless denoted, the on-chip cell culture usually lasted for ∼6 h before the chamber was fully filled with bacteria, which affects clear discrimination of the boundaries between the colonies.

**Optical Image Acquisition and Analysis.** Time-lapse images were recorded for five or 6 h with an IX71 epi-
fluorescent inverted microscope (Olympus Inc., UK), equipped with a long working distance Olympus LCP FL 100 × /0.85 objective lens, an Andor iXon Ultra 897 CCD camera, and a motorized stage (SCAN IM 120 × 80−2 mm, Marzhauser Wetalar GmbH&Co.KG, Germany). The whole process was automatically controlled through the interface of the open source software Micro-Manager. The high magnification of the objective lens enabled the bacteria images clear enough for quantitative image analysis. To calculate the total area of a single colony starting from a single cell, whereby to obtain its growth rate (μ) and lag time (λ) parameters (see below), all the time-lapse images were processed with ImageJ (Supporting Information, Figure S2), as detailed previously. With the chosen microscope focus setting, the bacteria typically appeared as bright objects surrounded by a dark rim (Figure 2), which made it easy to identify and calculate colony area.

**Calculation of μ and λ Parameters.** For traditional batch culture, cell growth rates are defined as those measured during the exponential growth phase. The specific growth rate (μ) can be derived from optical density measurements using eq 1.

\[ \mu = \frac{\ln(OD_t/OD_0)}{T} \]  

where OD₀ and ODₜ are the optical density (OD) of suspended cells (i.e., an indirect measure of cell density as mass per unit volume), T is the length of time for this change.

In our microfluidic platforms, the bacteria grow as monolayer microcolonies starting from a single cell, so the cell mass can be considered to be proportional to the colony’s area (A). Therefore, the growth rate of bacteria within the colony was analyzed via the following equation:

\[ \mu = \frac{\ln(S_t/S_0)}{(t - \lambda)} \]  

where S₀ and Sₜ are the areas of a single colony at the initial time (t = 0) and at time t, and λ is the lag time, which is considered as the time taken for the initial cell division to occur. This is close to the point of intersection of the steepest slope of the growth curve with the time-axis (Supporting Information, Figure S2).

**Statistics.** A total of 80 randomly selected single cells colonies from three independent experiments were analyzed for each population so as to allow the derivation of statistically meaningful characteristics. Unless noted, average values and standard deviation from the triplicate are given. All statistical analysis was performed using Prism software. A difference was regarded as significant when p < 0.05 (*), p < 0.01 (**), p < 0.001 (***), and p < 0.0001 (****) following a two-way ANOVA with post hoc testing unless otherwise denoted.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acssynbio.7b00177.

Figure S1, Growth rates of MG1655 and MDS42 strains in batch culture; Figure S2, Image analysis and calculation of specific growth rate (μ) and lag time (λ) of single cells; Figure S3, Long-term culture of cells in the microfluidic device; Figure S4, Determination of PDMS chamber height using microscopic absorbance–concentration measurements (PDF)

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**Notes**

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