The Ecology of Bacterial Individuality

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The Ecology of Bacterial Individuality

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Ph.D. Thesis
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General Introduction

It is long known that bacteria exhibit phenotypical heterogeneity within clonal populations and that heterogeneous environments have an impact on bacterial individuals that inhabit it. Nevertheless microbiologists tend to investigate bacteria as averages of populations: Be it by observing the average phenotypic properties of a population such as colony morphologies on an agar plates or the growth rate in a shaken liquid culture. A classical approach in microbiology is to determine the total number of colony forming units that can be recovered from an environment to evaluate the capability of the population of a bacterium to grow and survive in the environment. More recently, scientists turned to molecular tools to determine bacterial responses to an environment on a molecular level such as gene-expression of a population.

The concept of bacterial individuality is a recent development in microbiology. In essence it is built around the ideas that bacteria should be treated as individuals and investigated at scales that actually matter for them. In this thesis I followed these ideas and developed methods that show the value of the information that can be extracted from single-cell observations and how they increase our knowledge about the ecology of bacteria.

As the study environment, I used the leaf surface, or phyllosphere, as a natural system for bacterial colonization. By using bacterial bioreporter technology and fluorescence microscopy coupled with image cytometry I was aiming to determine colonization success and the impact of chance on colonization processes. In another bioreporter-based approach I aimed to determine local differences in fructose permeance through isolated cuticles. I then used the data derived from the previous studies to formulate a spatial-explicit, agent-based model for phyllosphere colonization.

A brief introduction to bacterial ecology

The term ecology describes the study of the distribution, abundance, and interaction of organisms with their biotic and abiotic environment (Begon, Harper et al. 1996). Bacterial ecology deals with questions on what and how Earth’s ecosystems are inhabited and influenced by bacteria. Some examples for bacterial ecosystems are: biofilms in aquatic sediments (Torsvik, Sorheim et al. 1996), oceans (Gugliandolo and Maugeri 1993), the gastro intestinal tract of animals (Brune 1998; Zoetendal, Collier et al. 2004), soil (Torsvik, Sorheim et al. 1996) and plants (Andrews and Harris 2000; Lindow and Brandl 2003; Leveau 2006). Additionally, a variety of anthropogenic types of bacterial ecosystems exist, e.g. wastewater treatment plants (Kapley, De Baere et al. 2007), vinegar or acetic acid bioreactors (Sueki, Kobayashi et al. 1991), glutamic acid bioreactors (Hermann 2003), and silage (Denoncourt, Caillet et al.
It is generally accepted that bacteria colonize and influence nearly all habitats on Earth (Whitman, Coleman et al. 1998).

**Introduction of plants as a habitat for bacteria**

As the experimental model system in this thesis involves plant-associated bacteria, a more detailed introduction into plants as a habitat for bacteria will be given in the following section. Plants are common habitats for microorganisms. Their roots and leaves are naturally colonized by a broad spectrum of bacteria, fungi and protozoa (Andrews and Harris 2000), which have a variety of interactions with the host plant ranging from mutualism to commensalism and pathogenicity (Singh, Millard et al. 2004). Especially the root system, the so-called rhizosphere, is colonized very densely. One gram of forest soil may typically contain up to $10^9$ bacterial cells (Whitman, Coleman et al. 1998), and bacterial densities in the rhizosphere may be one to two orders of magnitude greater (Liljeroth and Baath 1988; Duineveld and Van Veen 1999; Jaeger, Lindow et al. 1999). Compared to underground organs, aerial parts of plants are less densely inhabited (Holm and Jensen 1972). Leaf surfaces, for example, feature limited availability of water and nutrition, increased exposure to UV-radiation, and the waxy cuticle covering the leaf is a substrate that is very hard to penetrate (Kolattukudy 1985) and offers few niches to populate (Monier and Lindow 2004). Also, several plants possess a self-cleaning effect, which prevents the successful colonization of leaves (Barthlott and Neinhuis 1997). Therefore, only specialized microorganisms can manage to survive the conditions on leaf surfaces. Leaves are typically inhabited by $10^6$ - $10^7$ bacterial cells per cm$^2$ or up to $10^8$ bacteria cells per gram of leaves (Lindow and Brandl 2003), and the cumulative number of bacteria on plant leaves world-wide has been estimated to add up to the astonishing number of $10^{26}$ (Morris and Kinkel 2002). Most of these so-called epiphytic bacteria have developed strategies to adhere to their substrate: some synthesize and secrete extracellular polymers, whereas others possess pili to facilitate adhesion and prevent removal (Beattie and Lindow 1999). Some bacteria can also influence the substrate structure itself, e.g. by producing indole-3-acetic acid (IAA), a phytohormone. In high doses, bacterially produced IAA is thought to cause russet on pear fruits (Lindow, Desurmont et al. 1998), which result from an increased division activity of plant cells. These cells act as sinks for nutrients from which the bacteria might benefit. Also, it is known that bacteria are able to increase the permeability of leaf cuticles (Schreiber, Krimm et al. 2005), thereby increasing their access to plant-derived nutrients.

Leaves are not colonized in a uniform manner. The most frequently colonized struc-
tures are the bases of trichomes, glandular cells, leaf veins, epidermal cell wall junctions, hydathodes, wounds, and stomata (Beattie and Lindow 1999; Monier and Lindow 2004). Leaf permeance for nutrients, mostly photoassimilates such as fructose, glucose and sucrose, can differ between these features (Schlegel, Schönherr et al. 2005; Schreiber 2005). This is in part due to the heterogeneous compositions and thickness of the cuticle, which overlays the epidermis of plant leaves (Fernández, Khayet et al. 2011). The cuticle has also been shown to contain locally clustered hydrophilic pores, which allow higher rates of diffusion for hydrophilic substances through the cuticle (Schönherr and Schreiber 2004; Schlegel, Schönherr et al. 2005; Schreiber 2005). Water droplets can adhere longer at trichomes, glandular cells, leaf veins and other structures on the leaf and act as a sink for nutrients that leach out of the leaf (Linskens 1950). In addition, the nutrient flow in leaf veins is much higher than in the rest of the leaf, resulting in a steeper gradient, increasing the driving force with which nutrients permeate across the cuticle (Schreiber and Schönherr 2009). Differences in leaf geography are also thought to contribute to differential adherence and protection of bacteria (Monier and Lindow 2004). As a result, the distribution of cells on a single leaf surface can be very heterogeneous and can differ by 100 fold even between segments of 9 mm² (Kinkel, Wilson et al. 1995; Monier and Lindow 2004).

Leaves are colonized by airborne, waterborne, or biotic vector mediated (e.g. insects) events or during the emergence of the shoot through the soil (Leveau 2006). Usually, bacteria initially arrive on leaves as single cells or cell aggregates. These colonizers may encounter a variety of conditions, depending on their particular location. In laboratory experiments with bean plants, it was found that uncolonized leaf surfaces carry sufficient amounts of sugar (0.2 to 10 µg; the majority consisting of sucrose, fructose and glucose) to sustain a population of $1.7 \times 10^7$ bacteria per gram (Mercier and Lindow 2000). In another study, it was shown that most bacterial immigrants to bean leaves encounter nutrient conditions that allow them to divide several times, but after this initial phase, sugar is only available in a limited number of sites on the leaf (Leveau and Lindow 2001). Bacteria that have not already colonized these sinks likely starve and die off unless they have specialized adaptations to this environment, i.e. high tolerance to desiccation, starvation, UV-radiation, or adaptations to avoidance strategies these stresses altogether, e.g. by entering the leaf (Beattie and Lindow 1999).

**Studying the ecology of bacterial communities**

The study of bacteria in their natural environment involves experimental techniques
that typically have been developed for the study of bacteria in the laboratory, e.g. cultures grown in flasks or colonies and biofilms grown on agar plates. Many of these techniques are population-based measurements. Some of these techniques will be discussed here in more detail, using the phyllosphere as a model bacterial ecosystem.

A basic but valuable method to determine the number of bacteria on a leaf, as well as to investigate the general composition of the bacterial phyllosphere community, is plate counting. Leaves of known size or weight are washed in a known volume of a wash buffer, and samples of this buffer are spread onto a defined growth medium. After a sufficient time of incubation, colony forming units (CFUs) can be counted and used to estimate the bacterial numbers per unit of leaf surface (i.e. population size). A disadvantage of plate counting is the selectivity of the medium upon which the samples are plated, as some bacteria do not grow under artificial conditions, or they grow very slowly e.g. the pink-pigmented facultative methylotrophic bacteria (Holland 1997). Those organisms will be missed in standard plate count analyses.

Another disadvantage of plate counting is that it is a semi-quantitative method, because bacterial aggregates will form only one colony. These two problems may lead to an underestimation of the bacterial count in a sample.

Microscopic techniques can be utilized as an alternative to estimate bacterial counts. For this, leaves are washed in a known amount of buffer, and the leaf is weighed and/or the surface area is determined. Afterwards, samples are placed on a Neubauer counting chamber with a defined volume between slide and coverslip, which allows estimation of the cell count of a community. Microscopy can also be used to analyze colonized plant surfaces directly, either to examine the success of leaf washing procedures or to investigate the spatial distribution of bacteria on the leaf. To improve bacterial cell analysis, bacterial cells are commonly stained, for example with 4’,6-diamidino-2-phenylindole (DAPI) or Acridine orange, both fluorescent dyes that interact with nucleic acids and stain all bacteria including those bacteria that would be missed by plate counting due to their non-culturability.

Molecular community fingerprinting techniques can also be used to characterize the composition of an overall leaf community. For this, the total community DNA, or metagenome, is extracted and a subsequent polymerase chain reaction (PCR) with primers for conserved DNA-sequences that are connected by variable DNA-sequences is performed (Handelsman, Rondon et al. 1998). Afterwards, a step to separate PCR products of the dominant populations, such as denaturing gradient gel electrophoresis (DGGE) (Muyzer and Smalla 1998) and terminal-restriction fragment length polymorphism (T-RFLP) (Marsh 1999), is conducted. Each band
on a DGGE gel, for example, represents, in the ideal case, a population of one dominant species in the sample. These bands can be excised from the gel and sequenced to identify the species they represent. Importantly, also non-cultureable organisms will appear in these analyses. By employing DGGE, Yang et al. could show, that the phyllosphere diversity of *Citrus sinesis* is more complex than assumed based on culturable analyses (Yang, Crowley et al. 2001).

Besides improving knowledge about the qualitative composition of a leaf surface community, the quantitative composition of the community is of interest. It is possible to answer these questions by performing quantitative or real time PCR (qPCR or RT-PCR). To quantify the relative abundance of a DNA-fragment in a sample of metagenomic DNA, the produced double stranded DNA intercalates with a fluorescent dye or it is tagged with fluorescent probes, so that fluorescence intensity in a PCR becomes a measurement for the amount of fragment in the sample (Wong and Medrano 2005). By using primers for conserved DNA-sequences, it is possible to measure the content of different bacterial species or bacteria groups in a specimen (Zwielehner, Handschur et al. 2008). Furthermore, it is possible to analyze the appearance of certain genes and their abundance within a community by means of quantitative PCR methods.

To analyze the full metagenome of the phyllosphere it is possible to create a clone library by inserting large DNA-fragments of the metagenome into vectors (Jackson, Echlin et al. 2006). These vectors can then be cloned into bacteria and stored there for further analysis. Such clone libraries are able to reveal the genetic potential of a community for example which substrate the community can metabolize. The abundance of clones with certain genes, e.g. 16S RNA genes, in the library is also a proxy for the abundance of a species in a sample (Handelsman 2004).

To gain knowledge about a community it is possible to perform next generation sequencing, i.e. pyro- or Illumina-sequencing, of the metagenome. This can be done either in combination with a barcoded PCR of conserved DNA sequences to analyze the community composition (Claesson, Wang et al. 2010; Redford, Bowers et al. 2010), or by 454 shotgun pyrosequencing of the full metagenome (Kowalchuk, Speksnijder et al. 2007; Delmotte, Knief et al. 2009). The advantage of these methods is that they allow for the identification of bacterial species within a sample while providing a quantitative estimate of the relative species abundances within a sample (Fierer, Hamady et al. 2008).

A novel approach combines genomic and proteomic information of bacteria that reside in an environment, so called community proteogenomics (Delmotte, Knief et al. 2009). By generating a full metagenome database, of phyllosphere bacteria in
Figure 1 (changed from Leveau and Lindow 2002) Impact of environmental heterogeneity on individual microbial cells and the discrepancy between population and single-cell results. A) A habitat that consist of several microhabitats that feature different levels of a stimulus. B) Microbial bioreporters for the stimulus are released into the habitat. C) After a time of incubation the bioreporter will be induced to report on the stimulus in the habitat. These cells are now recovered from the environment and queried for their reporter signal either by a population-based measurement, e.g. fluorescent measurement in a cuvette, or a single-cell-based measurement, e.g. epifluorescent microscopy and image cytometry. Both methods yield different results as shown in D). The population-based measurement yields the average stimulus experienced by the bioreporter in the habitat revealing that the habitat contains average levels of the stimulus. The individual-based measurement reveals that the habitat is heterogeneous, containing sites that do not hold a stimulus, that hold average levels of the stimulus, and that hold high levels of the stimulus. This example reveals the averaging effect of population-based methods that, depending on the questions asked, can lead to wrong results. Single-cell methods yield higher information content about the habitat under study without prior knowledge of the heterogeneity of a stimulus distribution in the habitat.
combination with metaproteome information extracted from the same community, one can reveal information about the ecophysiology and possibly unique features of phyllosphere bacteria.

**Introduction of the ecology of individual bacteria**

In general, and as illustrated by the examples above, bacterial ecology has been heavily biased towards measurements of bacterial populations. Despite the fact that bacterial individuals can be observed, already since Antonie van Leeuwenhoek discovered bacteria in 1683, it has long been difficult, if not impossible, to monitor single bacterial cells and determine their identity and behavior or health status. Furthermore, the study of bacteria on or in natural substrates may be impeded by their invisibility without magnifying and staining protocols, as well as their relatively low density under natural conditions.

Given these difficulties, it is not surprising that classical bacterial ecology approaches have typically ignored single individuals. Classical bacterial ecology tends to describe the interactions of populations, not individuals, with the environment in relation to other populations (Kinkel, Andrews et al. 1987). Changes in populations are typically expressed as averages, without regard for variation that might exist at the individual level, for example in terms of generation time, response to the local microenvironment, and expression of functional traits. The heterogeneous surface of a plant leaf provides a good example to explain this. Most of the surface area is difficult to populate for bacteria, but certain spots are habitats for significant numbers of bacteria (Kinkel, Wilson et al. 1995). Figure 1 illustrates how a per leaf view, i.e. using the leaf as a unit of normalization, might give a different (and in this case incorrect) picture of how bacteria thrive in the ecosystem, compared to a single-cell perspective. On average, a plant leaf surface can be called an inhospitable environment, schematically indicated by white areas in Figure 1. If now bacteria are released into this environment (Figure 1 B) and are allowed to grow, they will do so according to the locally offered resources (Figure 1 C). As a result, the average bacterium is one that experiences the leaf surface as not very conducive to growth (Figure 1 D). However, from the perspective of single individuals, spatial variation in hospitality would result in different subpopulations on the leaf surface, some of which prosper (Figure 1 D) (Leveau and Lindow 2001). Even in environments that lack spatial heterogeneity, phenotypic variation among individual bacteria has been shown to exist. For example, it has been known for a long time that individuality occurs even in single descendants in a clonal population (Hughes 1956; Smits, Kuipers et al. 2006). An evolutionary explanation for phenotypic variation in individuals of
an isogenic culture could be the higher chance to survive hazardous conditions, so called bet-hedging (Veening, Smits et al. 2008). For instance, in case a population is subjected to environmental stress, individuals that are in a certain physiological state can be able to resist stress, and, after the stress stimulus is relieved, these individuals can regenerate a population featuring several phenotypes (Smits, Kuipers et al. 2006). Individual variation also arises from stochastic differences in cell division and bacterial growth (Kelly and Rahn 1932; Rosenfeld, Perkins et al. 2006; Strovas, Sauter et al. 2007).

Thus, there seems to be a clear need for bacterial ecologists to pay more attention to the fate of individual cells. How do single bacteria vary in their experiences of their local environment and in their reaction to it? How does the sum of all reactions shape population structure and distribution? Can we predict bacterial activities better if we get an appreciation for bacterial life at the single cell level? These are central questions to the ecology of bacterial individuality and thus central to this thesis.

**Individual based ecology and agent-based modelling**

In this context, a very useful theoretical framework to embrace is the concept of “individual based ecology” (IBE) (Grimm and Railsback 2005). IBE handles populations as a group of individuals whose growth, reproduction, food intake and ultimately death is an outcome of their adaptive behavior to their surroundings. Adapted to a bacterial system, it gives each individual bacterium its own individual environment. An individual environment results in an individual reaction, i.e. phenotype or gene expression.

The phyllosphere offers a great opportunity to investigate such relationships. It is easily accessible, highly amenable to experimentation, offers a highly heterogeneous environment at the level of individual bacteria, and it a natural and relevant habitat for bacteria.

An important tool in individual-based ecology is modeling. An individual-based bacterial model can be though of as a spatial explicit two- or three-dimensional environment inhabited by agents, representing organisms, that interact with each other and the environment. Agents are simulated individuals that feature behavior answers, e.g. resource uptake, movement, and division, which are dependent on environmental conditions, such as the local availability of resources, as well as modeler-defined thresholds, e.g. the amount resources that an agent owns before it divides. The possible reactions of agents, and the effects of these reactions on their environment, need a high amount of calculation power, which makes computers prerequisite for ecology models based on individuals (Grimm and Railsback 2005). The phyllosphere
for example can be simplified into a model by creating an environment consisting of a two-dimensional grid, which depicts the leaf, inhabited by agents that mimic bacteria.

So far, there have been several applications of individual-based ecology to explain bacterial growth patterns. Kreft et al. (1998) developed an individual-based bacterial model system BacSim. BacSim has been used to model dynamics of bacteria in biofilms, colony growth, and the migration of Salmonella enteritidis into chicken’s eggs (Kreft, Booth et al. 1998; Kreft, Picioreanu et al. 2001; Grijspeerdt, Kreft et al. 2005). Representing a second generation of biofilm models, iDynoMiCS has recently been released (Lardon, Merkey et al. 2011). An individual-based bacterial modeling systems has been developed to investigate microscale evolution of bacteria (Gregory, Saunders et al. 2006). Also, some successful efforts have been made to model individual-based chemotaxis models (Emonet, Macal et al. 2005). More individual-based bacterial model systems have recently been reviewed elsewhere (Hellweger and Bucci 2009). In an ideal case, a model should include all possible answers and behavior patterns, but the results produced by such a model are likely to be as complicated as the natural phenomena they are attempting to explain. So models typically aim for simplicity, while still trying to tackle and explain relevant biological questions.

Investigating individual bacteria

As mentioned above, bacterial ecology has traditionally been biased towards the study of populations and communities. Relatively little data and theories are available on the ecology of bacteria as individuals. However, a growing array of techniques is becoming available to interrogate individual bacteria in their natural habitats. This section will list some of these recently developed techniques and their potential in an individual-based approach to bacterial ecology.

Single cell analysis by microscopy

The most frequently used and oldest technique to investigate single bacterial cells is microscopy. Since the first bacteria have been observed while using a simple light microscope (Dobell 1933) microscopy has made many technical advances. But even to date, the basic concepts of microscopy are still the same, and light microscopy is still widely used. Microscopic images can be captured with digital cameras, and analyzed by imaging software, e.g. to determine the size of single cells or the content of a chromophore inside a cell (Brehm-Stecher and Johnson 2004). Light microscopy allows interrogation of single bacteria in their natural habitat, provided that cells are not stained prior to the analysis. To obtain a three-dimensional perspective of a habi-
tat, Confocal Laser Scanning Microscopy (CLSM) may be used (Zhang, Callaway et al. 2009). While using CLSM, several pictures are taken along the Z-axis to produce a three-dimensional image, which can be an advantage when studying thicker specimens, or a microenvironment that is characterized by an uneven surface. It is possible to create ultra-high resolution images of individual bacterial cells on surfaces by using electron microscopy (Morris, Monier et al. 1997). Prior to the microscopy, surfaces harboring communities of interest have to be fixed and sputter-coated with gold. During the observation specimen are kept under vacuum.

**Bioreporter technology in single-cell microbiology**

Reporter proteins are valuable tools in microbial ecology. In a typical application, reporter genes are fused to a constitutive- or inducible gene promoter which allows the visualization of bacteria in their natural environment (Leveau and Lindow 2002), or the quantification of gene activities (Kohler, Bubert et al. 2000). If fused to promoters that are activated by a particular stimulus, it is possible to generate bioreporters, i.e. cells that respond to the stimulus by the production of the reporter protein. Examples of such stimuli are nutrients (Leveau and Lindow 2001), heavy metals (Abd-El-Haleem, Zaki et al. 2006) or even the abundance of certain species in a certain culture volume by quorum sensing (March and Bentley 2004). The list of bioreporters is growing constantly (Hansen and Sorensen 2001; Leveau and Lindow 2002; Shimshon 2003; Nivens, McKnight et al. 2004). The following section will introduce some of the currently available reporter genes.

The green fluorescent protein (GFP) is derived from fluorescing jelly fish (Chalfie, Tu et al. 1994) and has been extensively used as a reporter protein in different types of organisms, including bacteria. Through modification, the original properties of GFP have been changed to design proteins that differ in maturation time, stability (Andersen, Sternberg et al. 1998), excitation wavelength maximum (Cormack, Valdivia et al. 1996), and emission intensity and wavelength (thereby changing the color of the resulting protein) (Cormack, Valdivia et al. 1996; Miller, Desai et al. 1999). Fluorescent proteins from other organisms, such as DsRED and its derivatives, have been added to the pool of available reporter proteins (Matz, Fradkov et al. 1999; Shaner, Campbell et al. 2004). The advantage of short-lived forms of GFP is the ability to observe temporal changes in gene activity. This can be of utility in studies that employ bacterial bioreporters for the exploration of changes in environmental conditions. For example, Leveau and Lindow (Leveau and Lindow 2001) introduced a gene for unstable GFP[AAV] (Andersen, Sternberg et al. 1998) under the control of a fructose responsive promoter into *Erwinia herbicola*. Bacteria harboring this plasmid were successfully used to quantify fructose in culture and show
the transient availability of fructose on leaf surfaces. Other reporter proteins that are used for single-cell analyses are enzymes like β-galactosidase (Chung, Conner et al. 1995). The detection is based on enzymatic cleavage of substrates into easily detectable compounds, e.g. X-Gal is cleaved by β-galactosidase to form insoluble indigo-blue homodimers, or MUGLR-substrates that result in insoluble fluorescent products after cleavage by β-galactosidase (Manafi, Kneifel et al. 1991). Due to their insolubility, cleaved MUGLR-products are not released by intact cells, so that it is possible to quantify enzyme activities and thus promoter activities in individual cells. These fluorescent proteins and substrates may be combined with a stain that only colors dead cells, e.g. propidium iodide (Monier and Lindow 2003), which allows an estimation of living and dead cells.

**Analysis of individual cells using fluorescent probes**

Beside reporter proteins, it is also possible to interrogate single cells by using fluorescence *in situ* hybridization (FISH). FISH involves fluorescently marked DNA-probes that hybridize with RNA or DNA (Amann, Fuchs et al. 2001; Zwirglmaier, Ludwig et al. 2004). FISH can be highly sensitive, which makes it an ideal tool for cell counting by microscopy or flow cytometry (see below). Furthermore, by using probes that bind to variable regions of the ribosomal RNA, it is possible to confirm the phylogenetic identity of bacteria under investigation. The use of differently colored probes with different specificities makes it possible to obtain data sets about multiple populations from a single sample. Probes that hybridize with 16SrRNA can also be used to make statements about the general activity of cells (Ruimy, Breittmayer et al. 1994; Leveau and Lindow 2001). FISH is fully compatible with other fluorescence-based methods, like GFP reporter technology (Eberl, Schulze et al. 1997). A disadvantage of FISH is that it is destructive due to the need to fix cells prior to the analysis. Several variations on the FISH theme have been developed. To enhance the detection of low copy RNA or DNA, catalyzed reporter deposition FISH (CARD-FISH) can be used: by using probes that are tagged with horseradish peroxidase instead of a fluorophore. The peroxidase activity leads to signal amplification compared to fluorophore-labeled FISH probes (Pernthaler, Pernthaler et al. 2002). The recognition of individual genes by FISH (RING-FISH) is possible by using polynucleotide probes. The method is based on the formation of a probe network that is anchored at the probe-specific site in the cell. The network is formed due to complementary regions within the polynucleotide probes. Another variation is to investigate the gene expression or genetic traits of single cells via prokaryote *in situ* polymerase chain reaction (PI-PCR) (Hodson, Dustman et al. 1995). The method
uses fixed cells that are spotted on a microscopic slide after which a PCR is executed directly on the slide. Afterwards, fluorescence probes are added onto the slide similar to FISH, which can hybridize to the amplified nucleotide strains.

**Analysis of individual cells by flow-based techniques**

By flow cytometry, large numbers of cells can be rapidly analyzed (Davey and Kell 1996). This technique can directly or indirectly (e.g. after staining) detect and quantify single cells based on their optical properties, such as fluorescence intensity of bacterial bioreporter cells. Fluorescence-activated cell sorting (FACS) is a flow-cytometric tool to separate cells with different optical properties (Davey and Kell 1996; Brehm-Stecher and Johnson 2004). For example, it is possible to separate rod-shaped and coccus-shaped, or fluorescent from non-fluorescent bacteria. This method is a versatile tool to enrich cells with an interesting property from a given sample. Examples for FACS applications include the separation of two different populations with defined GFP contents for further analyses (Davey and Kell 1996; Shapiro 2000).

Another flow-based single-cell interrogation technique is microfluidic flow cytometry (Shapiro 2000). It has several advantages: the sensitivity is much higher than the sensitivity of a standard flow cytometer, it is possible to add and mix fluids into the sample (e.g. stains), and the flow of the sample can be stopped and reversed to investigate a cell again or investigate its behavior over time. The disadvantage of this technique is the smaller number of cells that can be analyzed compared to conventional flow cytometry.

**Raman spectrum-based analysis of individual cells**

A very interesting and advanced form of microscopy is Raman microscopy. Based on Raman spectroscopy, which uses the inelastic light scattering that is produced by molecules that were exposed to a monochromatic laser, it is used for chemical analyses in very small areas by exposing them to a monochromatic light beam and the analysis of the resulting spectra (Huang, Griffiths et al. 2004). Raman microscopy produces a spectrum that represents an enormous amount of data for the different components from a single cell. Raman microscopy has already been used to analyze single bacteria to study differences in chemical composition (Schuster, Urlaub et al. 2000; Rosch, Harz et al. 2005; Huang, Bailey et al. 2007). Sample preparation for Raman microscopy is not destructive, so whole inoculated samples can be investigated (Rosch, Harz et al. 2005). This allows the investigation of individual chemical heterogeneity in populations (Schuster, Urlaub et al. 2000). The main disadvantage
of Raman microscopy lies in the time-consuming analysis of these spectra and, as a consequence, the relatively low number of cells that can be analyzed in a given period of time.

**General Approaches**

The aim of this PhD-thesis was to develop the concept of bacterial individuality into a more quantifiable and experimentally testable science within the field of bacterial ecology. We determined the merit of a single-cell approach to explain the activity of bacteria in their natural environment. As a model bacterial ecosystem, the leaf surface, or phyllosphere, was chosen. This habitat has several experimental advantages, i.e. being readily accessible by microscopy and allowing the easy recovery of bacteria for single-cell analysis. Also, the phyllosphere can be manipulated to mimic specific environmental situations, e.g. different levels of humidity, rain, wind, and light/UV illumination levels. By using diverse methods of inoculation, such as airbrushing or the dipping of leaves into bacterial solutions, different kinds of colonization strategies can be tested.

For most of the experiments, *Phaseolus vulgaris* plants (green snap bean ‘Blue Lake Bush 274’) were used as the host system, as there are already many protocols and studies available on the colonization its leaves by bacteria (Beattie and Lindow 1999; Mercier and Lindow 2000; Leveau and Lindow 2001; Monier and Lindow 2003; Monier and Lindow 2004). Beans can also easily be cultivated in the greenhouse and thus allow independence from seasons.

In addition to intact leaves, isolated plant leaf cuticles were used. Their properties are similar to living leaf surfaces (Kirsch, Kaffarnik et al. 1997; Schreiber 2001) and allow for standardized experiments, for example to quantify diffusion of substances. The bacterial strains used in this thesis are all derived from the parental strain *Erwinia herbicola* 299R that has been used in many phyllosphere studies (Lindow, Desurmont et al. 1998; Beattie and Lindow 1999; Leveau and Lindow 2001). Erwinia herbicola (also known as Pantoea agglomerans or Enterobacter agglomerans) is a heterotrophic, flagellated, Gram-negative Gammaproteobacterium that belongs to the family Enterobacteriaceae (Rezzonico, Smits et al. 2009). It is primarily a plant epiphyte, but has been found in other ecological niches like soil, sediments, and aquatic environments (Rezzonico, Smits et al. 2009). Erwinia herbicola forms monolayer and multilayer biofilms on plant surfaces, it also forms spherical or ob-long-elliptical multicellular structures in culture and on plants which are currently under investigation (Tecon and Leveau, unpublished).
Outline of the thesis

Chapter 2
In chapter two, a proof of concept is provided demonstrating how a single-cell approach provides complementary information over a population-based approach. A bacterial bioreporter was constructed, characterized and exploited for quantification of single-cell reproductive success. Consequently, we coined the bioreporter CUSPER for reproductive success = repsuc read backwards. After establishing the proof-of-principle, basic leaf colonization experiments were performed. The bioreporter consisted of the *Erwinia herbicola* strain *Eh*299R::JBA28(pCPP39) that can be induced to produce GFP in response to Isopropyl-β-D-thiogalactopyranosid (IPTG) (Leveau and Lindow 2001). In an environment lacking IPTG, GFP production ceases and whatever GFP is present in cells is diluted with every cell doubling. In that way GFP content of any single cell is inversely proportional to the number of times its ancestor has divided since this ancestor was introduced into the IPTG-free environment. This single-cell approach revealed to what extent the total population-increase on the leaf was due to individual immigrant cells that contributed to different degrees to the final population. This cannot be answered using a population approach. Proof-of-principle characterization of the bioreporter was performed as follows: Growth of GFP-loaded CUSPER on LB without IPTG revealed a decrease in GFP concentration (as quantified by fluorescence microscopy) with a rate that is inverse to the growth rate. GFP-loaded CUSPER cells were then released on bean leaves by dipping them into a bacterial suspension. After different incubation times, the cells were retrieved, cell counts were determined by plate counts and GFP concentration of single cells were measured. Data revealed subpopulations of cells that contributed to different degrees to bacterial populations on bean leaves, thereby providing evidence for environmental heterogeneity in the phyllosphere. We determined that the phyllosphere offers a sliding scale of habitability for bacterial colonizers.

Chapter 3
In Chapter 3, I evaluated the general understanding of the term “carrying capacity” in microbiology, which in some respect differs from the term as it is used in the general ecological context. In microbiology, it is used as a synonym for maximal load, which describes the maximum number of organisms that an environment can sustain, while in ecology the term describes the equilibrium of growth and death,
which results in the maximum number of organisms that an environment can sustain. Different densities of GFP-loaded CUSPER cells were airbrushed onto bean leaves. Some of these densities were deliberately higher than the carrying capacity of the leaves under study. Bacteria were recovered from leaves at different time points, CFUs were determined and single-cell reproductive success was analyzed. Results provided evidence that the phyllosphere consists of distinct microenvironments that offered low, intermediate, or high success rates for reproduction. The study estimated the probability of successful phyllosphere colonization. We concluded from our data that the carrying capacity of the phyllosphere has to be understood as a sum of local carrying capacities that are determined by individual sites on leaves.

Chapter 4
In chapter 4, I evaluate if the success of biocontrol by the principles of preemptive colonization and preemptive exclusion are determined at the level of single cells. It is known that certain bacteria and fungi are able to attack plants, potentially causing severe damage and significantly lower crop yields (Pernezny, Raid et al. 1995; Roberts, Berger et al. 1997; Savary, Willocquet et al. 2000; Goszczynska, Botha et al. 2007). Some of these pathogens cause damage only when a certain population density is reached (Whitehead, Byers et al. 2002). There have been several attempts to control such plant pathogens by inoculating plants with non-pathogenic, but competitive, bacteria (biocontrol agents) before pathogens can reach the population density necessary for infection (Lee, Lee et al. 2006; Stockwell, Johnson et al. 2010). Among potential mechanisms of action, biocontrol agents are acknowledged to repress growth of pathogenic strains by competitive exclusion (Lindow 1987). This is the principle of preemptive colonization, and its success or failure ultimately depends on how individuals of the biocontrol agent and the pathogen interact at the micrometer level.

The CUSPER bioreporter was used in competition with preemptively inoculated red fluorescent Erwinia herbicola (Eh299R(pFru97)) cells on bean leaves. Leaves were inoculated with three densities of Eh299R(pFru97) or a water control 24 hours before GFP-filled CUSPER cells were released.

Chapter 5
In this chapter, I describe the localization of hotspots of fructose diffusion over isolated leaf cuticles of poplar (Populus canescens). Isolated leaf cuticles were measured for their bulk-permeance of fructose by using gas chromatography coupled a with flame ionization detector. Subsequently, cuticles were inoculated with the GFP-based fructose bioreporter strain Erwinia herbicola strain Eh299R(pFRU46)
(Leveau and Lindow 2001) to later be scanned by epifluorescence microscopy. Sites containing GFP-expressing cells were quantified and their size was measured. By relating the bulk-measurements with the local information obtained by fructose-exposed bacteria, it was possible to estimate the local fructose flux over the cuticles.

Chapter 6
This chapter introduces the agent-based simulation of microbial phyllosphere colonization (ASiMoPh) model. ASiMoPh was constructed around the framework of CUSPER bacteria that report reproductive success in a heterogeneous environment. A heterogeneous two-dimensional environment, mimicking a leaf surface, was generated using the agent-based modelling software NetLogo (Wilensky 1999). This environment consisted of patches that offer different amounts of “energy”. The agents released into the environment can utilize this energy. The model allow for the visualization of bacterial growth in a heterogeneous environment. It also records the number of agents that reside in the environment and the number of generations that individual agents were able to undergo since they were released into the environment. This facilitated the comparison of model results with experimental results acquired with the CUSPER reporter system.

Chapter 7
In this brief chapter, I describe the preliminary results of the de novo assembly and draft annotation of the *Erwinia herbicola* 299R genome. Genomic DNA was sequenced using the Illumina platform and de novo assembled using the bioinformatics tools Velvet and SeqMan Ngen. Draft annotation was performed using the RAST online annotation service. A finalized assembly and annotation will enable follow-up applications like expression profiling of *Erwinia herbicola* 299R in natural and artificial environments. Expression profiling and the availability of the genome can help to answer questions like, “what makes a successful phyllosphere bacterium?” and, “how does *Erwinia herbicola* 299R experience the leaf environment?”

Chapter 8
This chapter provides a summarizing discussion and examines the implications of the results presented in this thesis on phyllosphere microbiology and the ecology of bacterial individuality.
The Ecology of Bacterial Individuality
Linking environmental heterogeneity and reproductive success at single-cell resolution
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Abstract
Individual-based microbial ecology (IBME) is a developing field of study in need of experimental tools to quantify the individual experience and performance of microorganisms in their natural habitats. We describe here the conception and application of a single-cell bioreporter approach with broad utility in IBME. It is based on the dilution of stable green fluorescent protein (GFP) in dividing bacteria. In the absence of de novo synthesis, GFP fluorescence of a daughter cell approximates half of that of its mother, from which follows that the fluorescence of a progeny cell is a quantitative measure for the reproductive success of its ancestor. To test this concept, we exposed GFP-filled bacteria to different degrees of environmental heterogeneity and assessed how this affected individual cells by the analysis of GFP content in their progeny. Reporter bacteria growing in rich medium in a shaking flask showed no variation in reproductive success, confirming that life in a broth is experienced much the same from one bacterium to the next. In contrast, when reporter bacteria were released onto plant leaf surfaces, representing a microscopically heterogeneous environment, clear intrapopulation differences in reproductive success were observed. Such variation suggests that individual cells in the founding population experienced different growth-permitting conditions, resulting in unequal contributions of individual bacteria to future offspring and population sizes. Being able to assess population changes bottom-up rather than top-down, the bioreporter offers opportunities to quantify single-cell competitive and facilitative interactions, assess the role of chance events in individual survivorship and reveal causes that underlie individual-based environmental heterogeneity.
Introduction

Environmental heterogeneity, defined as spatial and temporal variation in the physical, chemical and biological environment, is a fundamental property of ecosystems (Scheiner and Willig 2008). At the scale of individual organisms, it affects the ability to survive, reproduce, co-exist and interact with other organisms. For plants and animals, environmental impact can be assessed and quantified relatively simply at the level of individual organisms (Melbourne, Cornell et al. 2007). In microbial ecology, however, the effect of environmental variability on microbial activity and diversity is commonly assessed at a scale that is several orders of magnitude greater than the dimensions of the microorganisms under study (Hellweger and Bucci 2009). This ‘coarse-grained’ (Templeton and Rothman 1978) approach to environmental heterogeneity suffers from the averaging effect that is typical of many population-based approaches (Brehm-Stecher and Johnson 2004). It is increasingly being recognized that ‘fine-grained’ environmental heterogeneity, that is the one experienced by microscopic individuals at the micrometer-scale is a key factor in explaining microbial activity, diversity, distribution and evolution (Davey and Winson 2003; Green and Bohannan 2006; Prosser, Bohannan et al. 2007; Davidson and Surette 2008). However, due to the relative lack of tools to probe environments for micrometer-scale differences in physical, chemical or biological variables, little is known about the heterogeneity that individual micro-organisms are exposed to and, more importantly, how this affects their activity and reproductive success.

Bioreporter technology (Leveau and Lindow 2001; Leveau and Lindow 2002; Harms, Wells et al. 2006; Leveau 2006; Tecon and van der Meer 2006) relies on microorganisms themselves to report on local environmental conditions. Many of these bioreporters involve the conditional expression of green fluorescent protein (GFP), a reporter that can be quantified with relative ease in individual cells by fluorescence image microscopy (Jaspers, Meier et al. 2001; Leveau and Lindow 2001) or flow cytometry (Axtell and Beattie 2002; Maksimow, Hakkila et al. 2002; Harms, Wells et al. 2006; Roostalu, Joers et al. 2008). When properly calibrated, the GFP signal becomes a measure for exposure to a particular environmental stimulus. For example, Leveau and Lindow (2001a) used a fructose-responsive promoter fused to the gene for GFP to probe the availability of this sugar to bacteria on plant leaf surfaces, also known as the phyllosphere (Leveau 2006). Temporal and spatial variation in single-cell green fluorescence indicated substantial heterogeneity in the availability of fructose to individual leaf colonizers (Leveau and Lindow 2001). Such heterogeneity has also been reported for other nutrients or stimuli that leaf bacteria are exposed to, including iron (Joyner and Lindow 2000), water (Axtell and Beattie
While bacterial bioreporters, such as the ones described above, are useful in micrometer mapping of differences in the bacterial experience of single environmental variables, they cannot communicate how each of those variables, individually or jointly, impact the fate of bacteria in the environment under study. We therefore designed a bioreporter tool that describes micrometer-scale environmental heterogeneity in general terms, that is as a sum of all variables expressed into a single, quantifiable effect on the bacterium. The bioreporter we introduce here records environmental heterogeneity in terms of past reproductive success. In concept, it is based on the observation that upon cell division, GFP in a bacterial cell is distributed in a predictable manner between its two daughter cells (Rosenfeld, Perkins et al. 2006; Roostalu, Joers et al. 2008): one division leaves cells approximately half as green fluorescent as their parent, two divisions one-fourth as fluorescent, and so on. Thus, the GFP content of an individual offspring cell becomes a quantifiable measure of reproductive success. This approach resembles the method that was used (Mailloux and Fuller 2003) to estimate in situ doubling times for bacteria released into an aquifer after staining them with carboxy-fluorescein diacetate succinimidyl ester, a fluorescent protein stain that dilutes from the bacteria with every cell division. However, whereas these authors were interested solely in population averages of in situ growth, we tested our GFP-based bioreporter by exposure to microscopic conditions of low (that is, LB broth) and high (that is, the phyllosphere) environmental heterogeneity to reveal sub-population differences in the reproduction of single bacteria. The implications of our findings extend broadly to studies on other microbial habitats dealing with the question of how individual bacteria in founder populations differ in their contribution to future population sizes.

Materials and methods

Bacterial strains and culture conditions

Erwinia herbicola 299R JBA28 (pCPP39) (Eh299R::JBA28 (pCPP39)) (Leveau and Lindow, 2001b) carries a chromosomal mini-Tn5-Km transposon insertion that expresses stable GFP from a LacIq-repressible P_{A1/O4/O3} promoter fusion to gfpmut3. The transposon confers resistance to kanamycin. The strain also harbors plasmid pCPP39, which confers tetracycline resistance and harbors a lacI^q gene for control of P_{A1/O4/O3} activity, and thus GFP production by isopropyl-β-D-thiogalactopyranoside (IPTG). Bacteria were cultivated at 28 °C on LB agar or in LB broth at 300 r.p.m. Where appropriate, IPTG, kanamycin, or tetracycline were added to final concentrations of 1mM, 50 or 15 mg ml$^{-1}$, respectively. Optical densities of bacterial cultures
were measured at 600nm (OD600) in a Unico 1100 spectrophotometer (Unico, Dayton, NJ, USA).

*GFP-loading, release and recovery of bioreporter Eh299R::JBA28 (pCPP39)*

Exponentially growing cells of Eh299R::JBA28 (pCPP39) were diluted 300-fold into fresh LB broth containing 1mM IPTG and grown to mid-exponential phase. These GFP-loaded cells were used to inoculate plant leaves (see below) or LB broth. In the latter case, 25 ml of LB was inoculated with 200 ml of GFP-loaded bacteria and incubated at 28°C and 300 r.p.m. Samples were taken every 30 min to measure OD600 and to collect bacteria for fixation (see below). For plant inoculations, GFP-loaded bacteria were diluted in Milli-Q water to a final concentration of 5 x 10⁴ colony-forming units ml⁻¹. Leaves of 12–14-day old Phaseolus vulgaris plants (green snap bean, variety Blue Lake Bush 274) were inoculated by brief submersion into this bacterial suspension, shaken to dispose of excessive liquid and transferred to a closed translucent box for high-humidity incubation at 21°C. At different time intervals, two leaves were transferred to a 50-ml Falcon tube with 20 ml 1 x PBS buffer, vortexed briefly and sonicated for 7 min. Part of the bacterial cells in the leaf washing was plated on agar for counting colony forming units, whereas the rest was collected on 0.2-µm Durapore filters (Millipore, Amsterdam, The Netherlands), recovered by vortexing for 15s in 1ml 1 x PBS, and fixed (see below).

*Fluorescence in situ hybridization, fluorescence microscopy and image cytometry*

Bacterial cells collected from LB broth or plant leaves were fixed as described previously (Leveau and Lindow 2001) and stored at -20°C in 50% 1 x PBS/50% ethanol for no longer than 2 weeks. To distinguish cells of Eh299R::JBA28 (pCPP39) from indigenous bacteria on the bean leaves, fixed leaf washings were subjected to fluorescence in situ hybridization using an Eh299R-specific, TAMRA-labeled probe (Brandl, Quinones et al. 2001) at a final concentration of 5.5 ng µl⁻¹. LB-or leaf-exposed cells were examined with an Axio Imager.M1 (Zeiss, Oberkochen, Germany) using 470/20 nm excitation for the visualization of GFP and 546/6 nm for TAMRA. Digital images were captured at 1000-fold magnification with an AxioCam MRm camera (Zeiss) in phase contrast and through a 525/25nm (GFP) or 575–640 nm (TAMRA) filter set. Using AxioVision 2.6 Software (Zeiss), single-cell GFP fluorescence was quantified as the mean-pixel intensity (Leveau and Lindow 2001), and expressed in units of Sfere (Standardized fluorescence reference), where 1 milliSfere equals one-thousandth of the average mean-pixel intensity of 1-mm Tetraspeck Fluorescent Microsphere Standards (Molecular probes, Eugene, OR, USA). Data analyses and simulations were performed in Microsoft Excel 2003 (Microsoft Cor-
poration, Redmond, WA, USA).

**Computer simulations**

For computer simulations presented in Figure 3a, reproductive success was calculated for 100 cells with a green fluorescence (GF) equal to X/2^t, in which X equals the Excel formula ‘=norminv(rand(),1000,250)’. Figure 3b shows the temporal changes in reproductive success of individual bacteria as a function of t from a population of 90 cells with GF = X and 10 * 2^t cells with GF = X/2^t. Figure 3c shows the reproductive success of 20 bacteria with GF = X, 20 * 2^0.125 * t bacteria with GF = X/2^0.125 * t, 20 * 2^0.25 * t bacteria with GF = X/2^0.25 * t, 20 * 2^0.5 * t bacteria with GF = X/2^0.5 * t and 20 * 2^t bacteria with GF = X/2^t. Figure 3d shows the reproductive success in a population of 20 bacteria with GF = X, 20 * 2^t bacteria with GF = X/2^t (t ≤ 1) or 20 * 2^t bacteria with GF = X/2^t (t > 1), 20 * 2^t bacteria with GF = X/2^t (t > 2) or 20 * 2^t bacteria with GF = X/2^t (t > 3), 20 * 2^t bacteria with GF = X/2^t (t > 3) or 20 * 2^t bacteria with GF = X/2^t (t > 3) and 20 * 2^t bacteria with GF = X/2^t.

**Results**

**GFP dilution is a quantitative measure for reproductive success**

A previously formulated mathematical model of GFP expression in bacteria (Leveau and Lindow 2001) predicts that in the absence of de novo synthesis, GFP dilutes from dividing cells at a rate equal to growth rate μ. We verified this prediction here using Eh299R::JBA28 (pCPP39), which accumulates stable GFP in the presence of the synthetic inducer IPTG. Upon transfer of IPTG-induced GFP-loaded cells to broth that lacked IPTG, GFP fluorescence declined exponentially and at a rate that was not significantly different from μ (Figure 1). No such decrease in fluorescence was observed when GFP-loaded cells were transferred to sterile Milli-Q water (data not shown).

![Figure 1](image)

**Figure 1** Optical density (■) and average single-cell green fluorescence protein (GFP) content (●) of GFP-loaded Eh299R::JBA28(pCPP39) growing in a shaken liquid LB broth. Fitted lines indicate a growth rate of 1.10 h⁻¹ and a GFP dilution rate of 1.13 h⁻¹. Also shown are representative fluorescence microscopy images of cells at t = 0, 1, 2 and 3 h (left to right).
not shown), confirming that GFP is extremely stable in this strain and that its dilution from cells depends on division. From Figure 1, it follows that the average number of cell divisions since $t = 0$ can be calculated from the GFP fluorescence at times $t$ ($GFP_t$) and $t = 0$ ($GFP_0$) as $2 \log(GFP_0/GFP_t)$. This value is essentially a measure of reproductive success; in simple terms, it means that the dimmer a cell is, the more successful its ancestor was in producing offspring. Figure 2a shows that in an exponentially growing bacterial population in LB broth, the reproductive success of individual cells (which for any individual cell was calculated as $2 \log$[average cell’s GFP fluorescence at $t = 0$ divided by single cell’s fluorescence at time $t$]) was normally distributed. The parallel lines signify that over time, all cells contributed with equal success to the population increase (Figure 2a). This observation is consistent with
the assumption that a shaking LB culture represents a homogeneous environment. A very different result was obtained when GFP-loaded cells of Eh299R::JBA28 (pCPP39) were released onto leaf surfaces of bean plants. In this environment (which is devoid of IPTG, see below), progeny bacteria showed considerably more variation in reproductive success compared with those in LB (Figure 2b), with clear deviation from the parallel lines observed in Figure 2a. This suggests that some immigrant cells to the leaves contributed more progeny to the population than others. In a control experiment, we introduced an uninduced culture of Eh299R::JBA28 (pCPP39) onto leaf surfaces. Analysis of bacteria recovered from leaves after 24h revealed that they had remained non-fluorescent, confirming that (1) the leaf surface was devoid of IPTG or other compounds that might induce de novo synthesis of GFP and (2) loss of plasmid pCPP39 (which would lead to constitutive expression of GFP) did not occur during the course of the experiment.

**Modeling of GFP dilution in individual cells**

To facilitate interpretation of the experimental data from the leaf surface, we ran four simple simulations (see Materials and methods) to examine the effects of different scenarios of environmental heterogeneity on the shape of reproductive success distributions. In the first scenario, all cells experienced the same conditions for maximal growth, much like the experiment in LB broth. This resulted in normal distributions of reproductive success (Figure 3a), as expected and as observed experimentally.

**Figure 3 Simulations of reproductive success in bacterial populations with different intrapopulation growth rates.** See text for details on panels (a-d). Data points with the same shape correspond to the same single simulated sampling time. Time is expressed as generations, not in hours.
In a second scenario, the starter population was split into two sub-populations, one of which, representing an arbitrary 10% of the cells, had a maximum reproductive success rate, whereas 90% were unsuccessful at producing offspring (Figure 3b). In the other two simulations, the starter population was divided into five equal sub-populations, each of which produced progeny at different rates (Figure 3c) or produced progeny at the same maximum rate but ceased doing so at different times during the course of the simulation (Figure 3d).

Comparison of the simulated distribution curves to the experimental ones suggests that it is unlikely that the leaf surface consists of only two types of locales: one that fully supports bacterial growth and another that does not. Based on Figure 3b, this would have resulted in the clear separation of two sub-populations in the distribution curves. Instead, it seems more likely that leaf locales represent a sliding scale in their ability to support growth of initial colonizers. Figures 3c and d show that patterns of increased heterogeneity can be simulated by assuming sub-populations that differ in their ability to reproduce, either through being offered less than favorable growth conditions or by being offered less time or resources to reproduce. Which one of these scenarios applies to the leaf surface, or whether it is a combination of the two, cannot be easily resolved by comparison of experimental to simulated data. However, both simulations prove the point that heterogeneity in reproductive success is indicative of a starter population in which cells are exposed to different growth-permitting conditions, resulting in unequal contributions to future offspring and population sizes.

Bacterial immigrants to the phyllosphere contribute differentially to leaf population sizes. Based on these simulations, we interpreted the experimental leaf data (Figure 2b) as follows. During the first 3 h, GFP content did not differ significantly from \( t = 0 \) across the population, suggesting that the bacteria did not reproduce during that time. Six hours after inoculation, > 90% cells appeared in a straight line more or less parallel to the \( t = 0 \) distribution, but with an average reproductive success of 2.9 divisions. This suggests that during the early period of colonization, most cells encountered similar conditions, allowing them to contribute equally to an approximately \( 2^{2.9} \), that is 7.5-fold population increase. About 5% of the cells appeared brighter than expected. These might represent cells with ancestors that settled in spots unfavorable for growth. With time, the shape of the distribution curve changed (Figure 2b). At \( t = 9 \) h, approximately 3% of the cells had divided 0 times, 2% 1 time, 8% 2 times, 36% 3 times and 51% 4 times or more. Extrapolated to \( t = 0 \), this means that 22%, 7%, 15%, 33% and 23% of the starter population contributed 3%, 2%, 8%, 36% and 51%, respectively, of the population at \( t = 9 \) (Figure 4). In other words, while nearly one-
Individual-based reproductive success of bacteria

Figure 4 Unequal contribution of bacterial founder cells to leaf population sizes. Plotted on the y axis is the cumulative percentage of cells with 0, 1, 2, 3 or ≥4 divisions 9h after inoculation onto the leaf surface, as determined by their green fluorescence protein (GFP) content, versus on the x axis the cumulative percentage that these cells represented in the founder population at t = 0. The latter was calculated by extrapolation to t = 0; for example, the 8% of bacteria that appeared to have divided twice by t = 9 constitute $8^2$ divided by the sum of $3/2^0$, $2/2^1$, $8/2^2$, $36/2^3$ and $51/2^4$ equals 15% at t = 0. The convex shape of the line indicates uneven contribution of individual founder cells to the total population size. For example, the subpopulations at the bottom-left and the top-right of the graph represent similarly sized fractions (29% and 23%, respectively) of the founder population, but differ 10-fold (5% and 51%, respectively) in their contribution to the population at t = 9.

third of the starter cells contributed only 5% to the population size at t = 9, less than one-fourth eventually contributed more than half. These data confirm that bacterial immigrants to the leaf surface contributed differently to population sizes, which is consistent with the hypothesis that the fate of individual immigrants is determined in large part by the environmental heterogeneity at the microscopic leaf level. We were unable to make estimates of relative contribution for the t = 24 population, as 79% of the cells had divided beyond the limit of GFP detection.

Estimating population changes from single-cell data

Figure 5a shows the changes in bacterial population sizes on the leaves, as determined by plate counting. The growth curve follows a pattern that is typical for this type of plant inoculation experiment, including a short lag, a phase of rapid growth and a level-off to apparent carrying capacity. It is interesting to note that this pattern can be reproduced quite well using only the single-cell data. From $N = N_0 \times e^{\mu t}$ (in which N is the number of cells at time t, and $N_0$ is the number of cells at $t = 0$) and $\text{GFP} = \text{GFP}_0 \times e^{\mu t}$ (in which $\text{GFP}$ is the average single-cell GFP content at time t, and $\text{GFP}_0$ is the average single-cell GFP content at $t = 0$), $\mu$ and $t$ can be eliminated to reveal that $N/N_0$ equals $\text{GFP}_0/\text{GFP}$. In other words, a plot of $\text{GFP}_0/\text{GFP}$ as a function of time produces in essence a growth curve, which is indeed confirmed for our data.
in Figure 5b. The underestimation of growth by the single-cell data at \( t = 9 \) and \( t = 24 \) is likely because of the fact that the reproductive success of a portion of the cells was undervalued because their GFP content was below the limit of fluorescence detection. Combined, Figures 4 and 5 demonstrate that our GFP bioreporter allows the assessment of population growth at the individual as well as population level.

**Discussion**

The experimental data we present here demonstrate the utility of the ‘reproductive success’ concept by offering new insight into bacterial phyllosphere colonization. The heterogeneity that we observed in the ability of individual immigrants to produce offspring on the leaf surface is a novel observation. It corroborates findings of others who have documented leaf-based heterogeneity in environmental stimuli (Joyner and Lindow 2000; Leveau and Lindow 2001; Axtell and Beattie 2002; Gunasekera and Sundin 2006), each of which is likely to affect reproduction. Together, these data support the hypothesis that local conditions are the key determinants of the abundance and dynamics of microbes on plant leaf surfaces (Woody, Ives et al. 2007).

We noted striking similarities in the interpretation of leaf colonization patterns based on our bioreporter data and those from a previously described bioreporter that is also *E. herbicola*-based but measures fructose availability (Leveau and Lindow 2001). In both cases, bacterial cells appeared to experience a period of adaptation immediately after immigration to the leaf surface. In the experiment presented here, this period of adaptation was accompanied by a reduction in cell size (data not shown), which concentrated the GFP fluorescence signal in the bacteria, resulting in brighter green fluorescent cells and lower apparent values for reproductive success (Figure 2b).

![Figure 5 Bacterial growth on leaf surfaces as determined by plate counts of colony forming units (a) and by estimation from green fluorescence protein (GFP) content of individual cells (b). The construction of panel b is described in more detail in the text. Owing to the unavailability of comparable colony counts for leaves at \( t = 0 \), both graphs compare population sizes relative to time point \( t = 1 \) h. The stippled line in panel b serves as a reference representing the shape of the curve in panel a.](image-url)
This initial period of adaptation was followed by a period of reproduction for nearly all immigrants. The fructose bioreporter revealed that bacteria differ substantially in their subsequent access to fructose, causing them to deplete their resources and cease dividing at different times during colonization. This parallels our observation here of unequal contributions to the population size (Figure 2b). These lines of evidence suggest that heterogeneity in fructose availability at the micrometer scale has an important role in the reproductive success of individual bacterial immigrants to the leaf surface.

Our observations of intra-population variability in the reproduction of bacteria on leaf surfaces are compatible with current theories of aggregative behavior of bacteria in the phyllosphere. Various studies (Morris, Monier et al. 1998; Monier and Lindow 2004) have shown that many or most bacteria on naturally or experimentally inoculated leaves occur not as isolated cells but in aggregates. Aggregation has been explained to result from the differential survival and growth of solitary and aggregated cells (Monier and Lindow 2003). In the case of *Pseudomonas syringae* (Monier and Lindow 2004), the frequency distribution of the number of cells per aggregate was found to be right-hand skewed, representing a sliding scale from many aggregates with few bacteria to few aggregates with many bacteria. Assuming that each aggregate arose from a single founder cell, a right-hand skewed distribution of aggregate sizes would indeed translate into a curved distribution of reproductive success, much like we observed for *E. herbicola* cells on leaves after prolonged exposure to the leaf surface (Figures 2b and 5b).

A limitation of our bioreporter is the inability to interpret reproductive success for cells in which GFP is diluted beyond the limit of detection. In our most optimal setup, this corresponded to six doublings, or 64 progeny cells from a single ancestor. This is sufficient for studies that are relatively short-term, involve bacteria with low rates of reproduction, or habitats with low or intermediate degrees of environmental heterogeneity. The need for fluorescence in situ hybridization to distinguish bioreporters from indigenous cells made it impossible to follow bacterial reproduction beyond four divisions. In future versions of the bioreporter, this may be solved by complementation of the bioreporter with a GFP-compatible, constitutively expressed fluorescent protein, for example red fluorescent mCherry (Shaner, Campbell et al. 2004). An additional advantage is the prospect of in situ observation of the bioreporter independent of its GFP fluorescence. Thus, one can start to interpret ancestral success of individual bacteria in the context of their location in the micrometer landscape.

Despite this room for improvement, the bioreporter in its current form offers several unique opportunities and advantages. One of its strengths is that reproductive success
is recorded in the GFP content of each cell, which is a major advantage for studies that allow only intermittent observation or that necessitate destructive sampling of the environment, as most experiments in microbial ecology do. Another plus of the reproductive success bioreporter is that it offers microbial ecologists low-ambiguity output. Most GFP bioreporters are promoter-based, and although promoters can be quite specific in response to the environmental variable under investigation, their activity can be modulated in unpredictable ways by other input from the environment (Leveau and Lindow 2001). Such ambiguity makes promoter-based bioreporters susceptible to misinterpretation, particularly in the absence of proper controls (Leveau and Lindow 2001). The reproductive-success bioreporter is promoter-independent in that it is solely based on dilution of previously synthesized GFP from the cell by division, hence with the minimal likelihood of misinterpretation of GFP output. Such types of bioreporters are expected to have the broadest and most reliable utility in microbial ecology. Another major advantage of our bioreporter is its compatibility with many other single-cell interrogation techniques (Davey and Kell 1996; Brehm-Stecher and Johnson 2004) with the ultimate goal to link reproductive success at single-cell resolution to specific bacterial behaviors or environmental experiences, and to identify the sources of heterogeneity and their impacts on bacterial individuals and on population structure and activity.

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Individual-based reproductive success of bacteria
Variation in local carrying capacity and the individual fate of bacterial colonizers in the phyllosphere

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Abstract

Using a phyllosphere model system, we demonstrate that the term “carrying capacity”, as it is commonly used in microbial ecology, needs to be understood as the sum of many “local carrying capacities” in order to better explain and predict the course and outcome of bacterial colonization of an environment. Using a green fluorescent protein-based bioreporter system for the quantification of reproductive success in individual \textit{Erwinia herbicola} cells, we were able to reconstruct the contribution of individual immigrants to the observed changes in population size. This analysis revealed that to bacteria the leaf represents an environment where individual fate is determined by the local carrying capacity of the site where an immigrant cell lands. With increasing inoculation densities, the reproductive success of most immigrants declined, suggesting that local carrying capacity under the tested conditions was linked to local nutrient availability. Fitting the observed experimental data to an adapted model of phyllosphere colonization suggested that there might exist three types of sites on bean leaves, which differ in their frequency of occurrence and local carrying capacity. Specifically, our data were consistent with a leaf environment that is characterized by few sites where individual immigrants can produce high numbers of offspring, while the remainder of the leaf offered roughly an equal number of sites with low and medium reproductive success. Our findings contribute to a bottom-up understanding of leaf surface colonization, which includes a quantifiable role of chance in the experience at the individual level and in the outcome at the population level.
Introduction
In ecology, the “carrying capacity” of an environment is the maximum number of individuals that the environment can support (McArthur 2006). In phyllosphere microbiology, i.e. the study of microorganisms that colonize leaf surfaces, this term is invariably referred to as the number of microbial cells that a leaf is able to sustain (Wilson and Lindow 1994; Wilson, Savka et al. 1995; Woody, Ives et al. 2007; Nix, Burpee et al. 2009). Carrying capacity on leaf surfaces is typically determined through counts of colony-forming units (CFUs) on agar plates. It is a conditional value that depends on several biotic and abiotic factors (Leveau 2006). For example, different plant species are known to sustain different microbial population sizes (Kinkel, Wilson et al. 2000; Knief, Ramette et al. 2010). Furthermore, bacterial species may vary in the densities they can achieve on leaf surfaces depending on their ability to deal with the environmental conditions on the leaf surface (Jacques, Kinkel et al. 1995; Jacobs, Carroll et al. 2005). Water availability on the leaf has frequently been highlighted as yet another driver for microbial abundance in the phyllosphere (Leben 1965; Elad and Kirshner 1993; Wilson and Lindow 1994).

One key factor in determining foliar carrying capacity is the availability of nutrients. Carbon is a limiting resource during phyllosphere colonization (Mercier and Lindow 2000; Leveau and Lindow 2001). The process of leaching (Tukey 1966) deposits photosynthates such as the sugars fructose, sucrose and glucose on the leaf surface where they become readily available for consumption by microbes (Mercier and Lindow 2000; Leveau and Lindow 2001; Van Der Wal and Leveau 2010). Spraying of a carbon source onto plant leaves (Wilson and Lindow 1995) or making an additional carbon source available through genetic modification of the plant (Wilson, Savka et al. 1995) can also increase the carrying capacity of leaves for bacterial populations.

The use of bacterial bioreporters has confirmed that sugars such as fructose are not distributed evenly across the leaf surface (Leveau and Lindow 2001). It has also been reported that unused sugars could still be detected on leaves that are colonized by bacteria at the apparent carrying capacity (Mercier and Lindow 2000). These two observations are in line with the hypothesized existence of discrete sites on the leaf surface which contain disparate amounts of resources (e.g. sugars) and which may or may not be colonized by bacteria (Kinkel, Newton et al. 2002; Lindow and Brandl 2003). At high inoculation densities, all sites on a leaf can be expected to be colonized by bacteria, and in those cases, carrying capacity will be determined by the amount of nutrients available in all sites combined. At low inoculation densities, some sites will remain uncolonized, including some that contain nutrients. In
the latter case, the carrying capacity is not reached and instead becomes a function of how many, and which; sites are actually occupied by bacterial colonizers. Part of this model of leaf colonization was verified in experiments with the epiphytic bacterium Pseudomonas syringae (Wilson and Lindow 1994), revealing that populations derived from low inoculum concentrations never were able to reach the carrying capacity that was realized at high inoculum densities. However, what is still lacking is an experimental, quantitative demonstration of the variation that exists in nutrient availability, or local carrying capacity, between sites on a leaf surface. How big is this variation, and how does it contribute to the course and outcome of leaf colonization? Population-based measurements are not suited to address these questions, as they only examine averages across the population as a whole. In order to tackle these issues properly, an individual-based approach seems necessary.

Here, we used the bacterial bioreporter CUSPER (reverse of REPSUC, for REPro ductive SUcces) (Remus-Emsermann and Leveau 2010) to describe how increasing inoculation densities are experienced by individual bacterial colonizers of bean leaf surfaces. From this information, we were able to derive estimates for the variation in local carrying capacity and assess the impact of this variation on achieving final foliar population sizes. The CUSPER bioreporter is based on dilution of green fluorescent proteins (GFP) from dividing cells in such a way that GFP fluorescence of a single cell is inversely proportional to the reproductive success of the original immigrant from which the observed cell was derived. The CUSPER bioreporter was recently used to reveal that in a heterogeneous environment, such as the phyllosphere, individual bacterial immigrants contribute unequally to future population sizes (Remus-Emsermann and Leveau 2010). In this study, we utilized CUSPER for an individual-based examination of leaf colonization at different inoculation densities in order to gain a quantitative understanding of variation in local carrying capacities in the phyllosphere. Such knowledge is potentially important to general ecological understanding as well as downstream applications such as plant pathogen control.

Materials and Methods
Bacterial strain, culture conditions, and plant experiments
CUSPER is a bacterial bioreporter of reproductive success (Remus-Emsermann and Leveau 2010). It is based on Erwinia herbicola 299R::JBA28 (pCPP39) (Leveau and Lindow 2001; Remus-Emsermann and Leveau 2010), which constitutively expresses LacI\(^q\) from plasmid pCPP39 and carries a chromosomal mini-\(Tn5\)-Km insertion that expresses stable GFP from a LacI\(^q\)-repressible \(P_{A1/04/03}\) promoter fusion to \(gf\)-
The transposon confers resistance to kanamycin and the plasmid to tetracycline. The concept of CUSPER lies in the loading of bacterial cells with GFP through derepression of the PA104/03 promoter with isopropyl β-D-1-thiogalactopyranoside (IPTG), followed by release of these GFP-loaded cells into an IPTG-free environment. Without de novo GFP production, GFP is diluted every time a cell divides so that GFP fluorescence in each cell becomes an inverse measure for reproductive success. CUSPER cells were cultured to mid-exponential phase at 300 rpm and 28°C in LB broth supplemented with 50 μg kanamycin and 15 μg tetracycline per mL in addition to 1 mM IPTG. Bacteria were harvested by centrifugation at 3900×g for 10 minutes, and resuspended in 1x PBS to approximate densities of $10^5$, $10^6$, $10^7$, and $10^8$ colony-forming units (CFU) per mL. Two fully expanded cotyledon leaves of 14-day-old *Phaseolus vulgaris* plants (green snap bean, variety Blue Lake Bush 274; Burpee, Warminster, PA) were inoculated by evenly spraying the adaxial surface of each leaf with approximately 0.5 mL of one of the four bacterial dilutions using an airbrush paint gun (Pilot I Spray gun, Walther Pilot, Wuppertal-Vohwinkel, Germany). Inoculated plants were transferred to closed translucent boxes and incubated at high relative humidity and 21°C.

*Recovery of bacteria and analysis by microscopy*

After 0, 3, 5, 7, and 24 hours, three leaves were sampled from the plants that had been sprayed with one of the four bacterial suspensions, for a subtotal of 12 leaves per time point and a grand total of 60 leaves. Each leaf was transferred to a 50 ml Falcon tube with 20 ml 1x PBS buffer, vortexed briefly, sonicated for 7 min, and vortexed briefly again. Fifty-μL aliquots from each leaf wash were plated using an Eddy Jet spiral-plater (IUL, Barcelona, Spain) onto duplicate LB agar-plates supplemented with 50 μg kanamycin per mL in order to determine CFUs. For two of the three leaves per time point and treatment (i.e. 40 leaves total), the remainder of the leaf wash was filtered over a 0.2-μm Isopore filter (Millipore, Amsterdam, The Netherlands). CUSPER cells were recovered from the filter by vortexing for 15 sec in 1 ml 1x PBS, fixed as previously described (Leveau and Lindow 2001) and stored in 1x PBS at 4°C for no longer than 3 days. Cells were analyzed with an Axio Imager. M1 epifluorescent microscope (Zeiss, Oberkochen, Germany) using standard phase contrast and Zeiss Filter 38 (BP 470/ 40, FT 495, BP 525/ 50) for the visualization of GFP. Digital images were captured using an EC Plan-Neofluar 100x/1.30 oil objective (Zeiss) with an AxioCam MRm camera (Zeiss). Single-cell fluorescence was quantified as the mean-pixel intensity of individual cells (Leveau and Lindow 2001) using the AxioVision 2.8 software package (Zeiss, Jena, Germany).
Downstream data analysis was performed in Microsoft Excel 2010 (Microsoft Corporation, Redmond, WA, USA). To reconstruct the relative contribution of original leaf immigrants to the final population based upon single-cell green fluorescence measurements, we divided the green fluorescence of every one of the N cells in each sample at $t=24$ by the average cell’s green fluorescence in the inoculum at time $t=0$. For each cell, this relative GFP content was divided by the sum of all GFP contents within the same sample ($\Sigma\text{GFP}$) to give a GFP/$\Sigma\text{GFP}$ value for each cell. For each cell, we also calculated the reproductive success (RS) of its original progenitor as $2\log(1/\text{GFP})$. Cells were then binned into one of the following RS classes: $\text{RS}<0.5$, $0.5\leq\text{RS}<1.5$, $1.5\leq\text{RS}<2.5$, $2.5\leq\text{RS}<3.5$, $3.5\leq\text{RS}<4.5$, $4.5\leq\text{RS}<5.5$, and $\text{RS}\geq5.5$. For each bin, the GFP/$\Sigma\text{GFP}$ values of all cells were summed to give $\Sigma(\text{GFP}/\Sigma\text{GFP})$. Values of $\Sigma(\text{GFP}/\Sigma\text{GFP})$ were then plotted as a function of RS in a histogram (see Figures 2B, 3, and 5).

Simulation of phyllosphere colonization using a logistic growth model

We adapted a previously developed logistic-growth model for bacteria in a heterogeneous environment (Kinkel, Newton et al. 2002). The model simulates a leaf with $S=10^4$ sites that vary in their carrying capacity $J$, i.e. the maximal number of bacteria that a single site can sustain. Bacteria and their offspring were not allowed to move from one site to another. At the beginning of the simulation, bacteria were distributed over the S sites following a Poisson distribution assuming an initial population of $10^3$, $10^4$, or $10^5$ bacteria per leaf. With each iteration $n$, the number of bacteria $N$ in a site doubled, up to a maximum value $N=J$. Heterogeneity in $J$ was introduced by assuming a normal or log-normal distribution that was created using a custom script for NetLogo 4.2.2 (Wilensky 1999). Other distributions were created with a custom Visual Basic Script for MS Excel (http://azzalini.stat.unipd.it/SN/stephen_gersuk.excel). The trimodal distribution that provided the best fit to the observed experimental data assumed that 48.5% of the sites had a value $J$ of 2, another 48.5% had a value $J$ of 48, while the remaining 3% of sites had a value $J$ of 1152.

Results

Determining the carrying capacity of bean leaves

To determine the bacterial carrying capacity of bean leaves under laboratory conditions, we inoculated four sets of cotyledon leaves with a suspension of *Erwinia herbicola* strain 299R (Eh299R) CUSPER bacteria to reach four different densities, i.e. $4.5\times10^3$, $6.9\times10^4$, $5.7\times10^5$ or $1.2\times10^7$ CFUs per leaf (Figure 1A). Plants were
incubated at conditions of high relative humidity, and we monitored changes in the bacterial population sizes by determining CFU counts per leaf at 3, 5, 7, and 24 hours after inoculation (Figure 1A). With the two lowest inoculation densities, population sizes increased during the first 7 hours of the experiment, albeit at different rates. At the two highest inoculation densities, population sizes either remained constant or decreased during this time period. Carrying capacity was estimated from Figure 1B, which shows a plot of the fold-increase in population size between $t=0$ and $t=24$, i.e. $N_{24}/N_0$, as a function of the population size at time $t=0$. The trendline crosses the x-axis at 3.4x10^6 CFUs per leaf, which represents the inoculation density where no net growth is expected to occur, i.e. the total carrying capacity under these conditions.

*Estimating reproductive success of CUSPER immigrants from GFP content in offspring*

Prior to introduction into the bean phyllosphere, the CUSPER cells were loaded with stable GFP. This initial amount of GFP gets diluted as each cell starts to divide (Remus-Emsermann and Leveau, 2010). A hypothetical example is given in Figure 2A to illustrate how GFP content of single CUSPER cells was used to reconstruct the reproductive success (RS) of individual immigrants to the leaf surface. In this example, we consider an initial population of $N_0=8$ cells. Four of these do not divide, which equals an RS of 0 for each of those cells. One cell divides once (RS=1), two divide twice (RS=2) to produce four cells each, and one divides three
Quantifying local carrying capacity in the phyllosphere

GFP GFP/ΣGFP $^2\log(1/GFP)$

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$N_0=8$

$N_t=22$

Figure 2. Illustrative example of how GFP content in CUSPER offspring at time $t$ can be retropolated to reveal the reproductive success of bacterial cells at time $t=0$. Details are provided in the text.

times (RS=3) for an offspring of eight cells, resulting in a final population at time $t$ of $N_t=22$ (Figure 2A). Thus, at time $t$ this yields 4 cells with a relative GFP content of 1 (i.e. these cells did not divide and GFP was not diluted), 2 cells with a relative GFP content of 0.5 (each divided once, thus diluting the original GFP into two daughter cells), 8 cells with a relative GFP content of 0.25, and 8 cells with a relative GFP content of 0.125. For each cell from this population at time $t$, we can calculate two values: 1) GFP/ΣGFP, which is its GFP content divided by the sum of all GFP contents in the same population, and 2) $^2\log(1/GFP)$, which is the reproductive success (RS) of its progenitor. Cells can then be binned into one of the following RS classes: 0, 1, 2, or 3, and for each bin, the GFP/ΣGFP values of all cells are summed to give Σ(GFP/ΣGFP). Figure 2B shows Σ(GFP/ΣGFP) charted as a function of RS in a bar graph. In this particular representation, the width of each bar equals $2^{RS}$.

It is important to realize that this histogram is constructed solely on the basis of data from time $t$ and on the average GFP content at $t=0$. Yet, it tells us many things about the fate of individual cells and the increase in population size since $t=0$. Firstly, the total area covered by all bars in the histogram, i.e. $Σ(Σ(GFP/ΣGFP) x2^{RS})=2.75$, equals the increase in population size between $t=0$ and time $t$, or $N_t/N_0 (22/8=2.75)$. This value can also be calculated directly from $N_t/ΣGFP$, i.e. the number of cells at time $t$ divided by the sum of all GFP contents. Secondly, the
Figure 3. Variation in reproductive success of Eh299R CUSPER immigrants to the bean leaf surface, as inferred from GFP content in bacteria that were retrieved from leaves after 24 hours of incubation. Cells were retrieved from the same experimental setup that was used to construct Figure 1. For each of the four inoculation densities, cells from two independent leaf samples were analyzed: panels A and B, 4.5x10³ CFU per leaf; panels C and D, 6.9x10⁴ CFU per leaf; panels E and F, 5.7x10⁵ CFU per leaf; panel G and H, 1.2x10⁷ CFU per leaf. Histograms were fashioned in the same way as was done for the fictional data in Figure 2.
Graph reveals how successful different fractions of the original population were in producing offspring. For example, it shows that half (i.e. \( \Sigma \text{GFP}/\Sigma \text{GFP} = 0.5 \)) of the cells in the population at \( t=0 \) did not divide (RS=0) and that a quarter of the cells (\( \Sigma \text{GFP}/\Sigma \text{GFP} = 0.25 \)) at \( t=0 \) divided 2 times to produce 4 cells at time \( t \) (RS=2). Thirdly, the area of any single bar in the graph divided by the total area covered by all bars in the histogram gives the proportional contribution of cells from the corresponding RS bin to the population size at time \( t \). For example, the area of the third bar (RS=2) is \( \Sigma \text{GFP}/\Sigma \text{GFP} \times 2^{\text{RS}} = 0.25 \times 2^2 = 1 \), which, divided by 2.75 (=total area, see above), equals 0.36, meaning that 25% of the cells in the original population divided twice to produce enough offspring to account for 36% of the population at time \( t \).

**Single-cell experiences of leaf colonization after inoculation at different densities**

Using the analysis illustrated in Figure 2, we retropolated GFP content of leaf-recovered bacteria to the reproductive success of individual bacteria in the \( t=0 \) population. Specifically, we analyzed bacterial cells retrieved at the \( t=24 \) time point from eight leaves (4 inoculation densities, 2 leaf washings each) from the experiment for which population data are shown in Figures 1. The results are summarized in Figure 3, which shows eight histograms (A-H), i.e. two leaves from different plants for each density. These graphs indicate great variation in the fate of individual bacteria after they were introduced onto the leaf surface at different starting densi-
On the two leaves that received the lowest inoculation density (Figure 3A and B), 41 and 70 percent, respectively, of the initial colonizers divided once after arrival on the bean leaf surface. We interpret this to mean that a substantial portion of the immigrant population landed in a spot that did not support much growth. On the other hand, 59 and 21 percent, respectively, of the cells at \( t=0 \) was very successful, with an RS value of 5 or greater, i.e. they produced \( \geq 32 \) offspring. These very successful immigrants reproduced to become 98 and 80 percent, respectively, of the final population size, compared to only 2 and 13 percent, respectively, for the cells that underwent only one division upon arrival.

With increasing inoculation density, the relative fraction of very successful cells (RS\( \geq 5 \)) diminished (Figure 3), down to 2.4 and 0.7 percent, respectively, on the two leaves that were sprayed with the highest inoculation density (Figure 3G and H). At this density, 91 and 93 percent, respectively, of the bacterial immigrants each produced 4 offspring at most, with a majority not dividing at all or only once. Despite their low initial abundance of 2.4 and 0.7 percent, the very successful cells contributed a considerable fraction, 31 and 11 percent, respectively, of the final population size.

It is evident from Figure 3 that with increasing inoculation density, the sum of the area underneath all of the bars (representing \( \frac{N_{24}}{N_0} \), as explained in Figure 2) decreases. This decline in fold-increase in population size is consistent with our findings in Figure 1B, and can be explained by assuming that at higher inoculation densities, similar amounts of resources need to be shared by more bacteria, resulting in fewer offspring per cell. The notion that there were still few very successful immigrants at the highest inoculation density suggests that a small portion of the leaf surface (2.4/0.7\% in this case) consists of sites that allow considerably more reproduction than most other sites. Another trend that is apparent in Figure 3 is the increase in resemblance of the two histograms from each inoculation density, as inoculation density increases. This suggests that the course and outcome of leaf colonization become more predictable at higher inoculation densities.

For each of the eight histograms shown in Figure 3, we plotted in Figure 4 the sum of the area underneath all of the bars as a function of the \( \frac{N_{24}}{N_0} \)-fold increase that was obtained from CFU-based data. The fact that data points do not fall on a line with slope 1 through the origin indicates that GFP- and CFU-based results do not fully agree with each other. At the lowest inoculation density, this discrepancy can be explained by the known limitation of CUSPER (Remus-Emsermann and Leveau, 2010) that after 5 cell divisions GFP gets diluted to the point where we...
can no longer reliably detect and quantify it. For the two leaves inoculated at this density, we calculated that in order to get the GFP-based estimates for increase in population size up to the one derived from CFU data, one would have to assume average RS values of 7.4 and 9.0, respectively, for cells that were binned in the RS>5 category in Figure 3. This corresponds to 169 and 512 offspring, respectively, for each of these cells. At the highest inoculation density, the discrepancy between GFP- and CFU-based results may be explained by assuming that a proportion of the cells on the leaf were not able to form a colony on the plate. For the two leaves that were inoculated at this density, we calculated that in order to get the CFU-based estimates for increase in population size to match those derived from GFP data, one would have to assume that 90 and 87 percent, respectively of the cells lost their ability to form a colony between $t=0$ and $t=24$. Another explanation for the discrepancy between CFU- and GFP-based data is that aggregates that were washed from the leaf surface did not break up into single cells during the wash procedure, so that one CFU might represent more than one cell. However, we can exclude this possibility, based on the fact that such aggregates were never observed under the microscope.

**Modeling single-cell experiences in the phyllosphere**

We used our experimental data to test a modified version of a previously formulated logistic growth model for bacterial colonization of a heterogeneous leaf environment (Kinkel, Newton et al. 2002). Our model consisted of a virtual leaf that contains $10^4$ independent sites. Leaves were inoculated with $10^3$, $10^4$, or $10^5$ virtual bacteria following a Poisson distribution. For the lowest inoculation density, this means that approximately 90% of all sites were devoid of bacteria, whereas at the highest inoculation density, each site contained 10 cells on average. In each site, bacteria were allowed to reproduce up to the site’s local carrying capacity $J$. We tested different distributions of $J$ and recorded the reproductive success of individual bacteria. We found that of all tested scenarios, a trimodal (low, medium, high) distribution of $J$ was the one that best matched the general patterns we observed for the experimental data (Figure 5), i.e. a co-occurrence of very successful and nearly unsuccessful immigrants at low inoculation densities and the existence of a small portion of very successful colonizers among a majority of less successful immigrants at high inoculation densities. The best-fitting trimodal distribution assumed roughly similar numbers of sites with low or medium local carrying capacity, while the remaining, very small portion of sites was characterized by high local carrying capacity. In the model, a high inoculation density meant that medium-$J$ sites
received immigrants in numbers that for the most part approximated or exceeded \( J \), such that these cells did not differ much in reproductive success from cells that landed in low-\( J \) sites. In contrast, the few high-\( J \) sites were occupied by cells in numbers well below \( J \), allowing for a substantial number of offspring per cell. At low inoculation density, the low frequency of high-\( J \) sites prevented most, if not all, cells from landing in one of these sites, but since medium-\( J \) sites received cells in numbers well below \( J \), there was still an opportunity for these cells to proliferate much more than cells that landed in low-\( J \) sites, where no or few doublings occurred.

**Discussion**

We have determined the bacterial carrying capacity of bean leaves under controlled conditions in the laboratory. By bioreporter analysis, we were able to demonstrate that under these conditions, realized carrying capacity at different inoculation densities is a function of variation in the local carrying capacity of different sites on the leaf surface. Our findings are consistent with a model of phyllosphere colonization that features different fates for bacterial immigrants, depending on where and with how many others they land on the leaf surface. Our study offers a quantitative estimate for these fates, which is a significant step towards a more predictive understanding of bacterial colonization of leaf surfaces.

It is evident from our study that individual- and population-based data sets each offer an incomplete understanding of how bacteria colonize a leaf. CUSPER-
based measurements have the limitation that information is lost beyond a reproductive success of 5 (Remus-Emsermann and Leveau, 2010), so that the contribution of some immigrants to the population will be underestimated, while CFU-based measurements provide only the average contribution of cells to the population, while ignoring the variability that clearly exists among immigrants. In combination, however, these measurements can offer significant novel insights. For example, the substantial number of cells that divide at most once upon arrival to the leaf surface at low inoculation density, and the corresponding high increase in population size as derived from CFU counts, suggests that the reproductive success of cells that actually did divide was underestimated and that some cells can produce offspring in numbers that exceed 500 cells. Bacterial aggregates of this size and much larger have been found on bean leaves under appropriate conditions (Monier and Lindow 2004), although it was not established whether they were derived from single cells. On the other hand, to explain high RS values for individual bacterial immigrants at high inoculation densities in the context of a constant or decreasing CFU count under the same conditions, one needs to assume that a substantial number of cells lose their ability to form a colony on agar plates. Indeed, such nonculturability has been demonstrated experimentally for Pseudomonas syringae, another model colonizer of the phyllosphere, even under favorable conditions of relative high humidity (Wilson and Lindow, 1992).

In our current model of bacterial colonization of the phyllosphere, each leaf colonization event can be thought of as a bottleneck occurrence, where some percentage of the incoming population is prevented from reproducing, or even surviving, while other immigrants in that same population are numerically enriched because they settle at sites conducive to growth. Based on our derived estimates on the quantity and quality of sites, landing in a high-\( J \) site is unlikely for any individual bacterium, and at low inoculation densities, chance will determine whether such sites will be filled or not. Thus, the outcome of an inoculation event would be less predictable for low inoculation densities than for high ones. Ecological theory holds that if a nutrient resource is limited and the number of organisms competing for this resource is high, the average success of the population goes down (Le Galliard, Fitze et al. 2005; Rankin and Kokko 2006). This is precisely what we observed in our experiments and simulations: high inoculation densities forced more bacterial individuals to share a site and resulted in overall reduced reproductive success. This seems to support the notion that local carrying capacity on leaves is determined by local nutrient availability. Being perfectly isogenic, cells of \( Eh299R \) can be considered each other’s ultimate competitors,
with a niche overlap index (NOI) of 1 (Wilson and Lindow 1994). Thus, unless a site offered very high levels of nutrients, bacterial immigrants would be relatively unable to reproduce, representing a convincing case of the “tragedy of the commons” (Hardin 1968).

However, a one-time immigration event of up to $10^7$ bacteria per leaf is not a likely event in nature (Lindemann and Upper 1985). Instead, immigration is a gradual process, with cells arriving to the leaf from the surrounding air at different times and typically at low rates (Upper and Hirano 2002). Another aspect of our experimental setup that can be considered artefactual is that all immigrants were of clonal descent. In reality, the leaf surface is colonized by very diverse bacteria and other microorganisms with different NOIs, which allows for various degrees of co-existence (Knief, Frances et al. 2010). However, it is important to note that the ability to set up such artificial colonization events allowed us to extract data on the quality of the leaf surface as a habitable environment. This high level of experimental amenability is quite unique to microbial ecology and is one of the advantages microbial ecologists have over ‘macro’-bial ecologists (Fierer 2007). Future applications of CUSPER will address more realistic scenarios, by assessing variation in reproductive success of CUSPER cells in competition with bacteria other than $Eh299R$ and after immigration onto pre-colonized leaves.

Our observation that a small fraction (0.7-2.4%) of the cells was very successful in reproducing even at the highest inoculation density suggests the existence of sites on the leaf surface that offer high levels of nutrients, allowing these cells to contribute disproportionally (11-31%) to the final population size. Interestingly, this finding agrees with reports showing that relatively rare features on the leaf surface tend to harbor large numbers of bacteria on colonized leaves. Specifically, it is worth noting that while only 0.6% of the bean leaf surface is covered by hooked and glandular trichomes (Monier and Lindow 2005), yet 44.1% of $Eh299R$ bacteria in aggregates were found to be associated with these structures after incubation under moist conditions for 5 days (Monier and Lindow 2005). In at least one study (Yadav, Karamanoli et al. 2005), the population size of epiphytic bacteria was positively correlated with the densities of glandular and non-glandular trichomes. This link between bacterial fate, population size and leaf topography calls for further investigation into the predictive value of trichome density for bacterial colonization of leaf surfaces.

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Single-cell experience of bacterial immigrants to pre-colonized leaf surfaces

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Abstract

In this study, we assessed how preemptive inoculation of plant leaves with bacteria affected the establishment of secondary colonizers. We quantified the latter in two ways: 1) at the population level, i.e. as changes in the number of colony-forming units and 2) at the level of single cells, i.e. as reproductive success of individual bacteria. Both analyses showed that the ability of secondary immigrants to establish on the leaf was negatively correlated with the level of pre-population by primary colonizers. This effect was best described by an inverse dose-response curve with an apparent half-point inhibition efficacy of approximately $10^6$ cells of primary colonizer per gram leaf. Calculated efficacy was the same whether we interpreted population data or average single-cell data. Interestingly, under conditions of heavy pre-population with primary colonizers, we still observed that a small fraction of secondary immigrants still produced offspring despite the fact that the corresponding population measurement showed no net increase in population size. This difference in population-level and single-cell level data on the colonization success of secondary inoculants has direct relevance for biocontrol strategies that are based on the principle of preemptive exclusion of foliar bacterial pathogens. For example, even at saturating levels of primary inoculum, some individual cells of a pathogen may still be able to divide, providing them with the opportunity to reach a quorum and trigger behaviors that favor survival or contribute to virulence.

Keywords: Phyllosphere, biological control, BCA, \textit{Erwinia herbicola}, \textit{Pantoea agglomerans}, \textit{Phaseolus vulgaris}, preemptive colonization, biocontrol
Introduction
The plant leaf surface, or phyllosphere, is an extreme and uninviting microbial environment due to abiotic stress conditions such as drought, rain, heat, cold, and exposure to UV-radiation (Beattie and Lindow 1999; Andrews and Harris 2000; Leveau and Lindow 2001; Lindow and Brandl 2003; Leveau 2006). Nevertheless, a variety of microbes, including archaea, yeasts, filamentous fungi, and bacteria, is well adapted to life in the phyllosphere. Phyllosphere colonizers can have a variety of interactions with their hosts, ranging from mutualism to commensalism and pathogenicity (Leveau 2006). Since outbreaks of aboveground plant pathogens can cause a significant decrease in crop yield (Savary, Willocquet et al. 2000; Anderson, Cunningham et al. 2004), management of such pathogens is crucial in modern, often intense, agricultural systems. Additionally, it has been shown that human pathogenic bacteria are able to survive on fresh produce and cause outbreaks in human populations (Beuchat 1996; Brandl 2008; Whipps, Hand et al. 2008; Frank, Faber et al. 2011), suggesting that such targets should also be taken into consideration in crop management applications.

Many strategies have been developed to prevent or meditate the establishment of plant pathogens, including the application of chemicals such as copper hydroxide (Wimalajeewa, Cahill et al. 1991), fungicides such as azoxystrobin (Bertelsen, De Neergaard et al. 2001), antibiotics such as streptomycin and oxytetracycline (Lindow and Brandl 2003), as well as biocontrol agents (BCAs) (Andrews 1992). BCAs are nonpathogenic microorganisms that help reduce pathogen load by one or more different mechanisms. One important mechanisms is preemptive exclusion (PE) by which BCAs effectively compete for space and/or resources (Lindow 1987; Wilson and Lindow 1994; Wilson and Lindow 1995; Wilson, Savka et al. 1995; Monier and Lindow 2005; Stockwell, Johnson et al. 2011). An example of such a strategy is the preemptive inoculation of \textit{Pantoea vagans} (Smits, Rezzonico et al. 2010; Stockwell, Johnson et al. 2010) in high densities on blossoms and leaves to minimize the chance of secondary colonization by the plant pathogens \textit{Erwinia amylovora}, the causal agent of fireblight (Venisse, Barny et al. 2003).

Many studies have dealt with biocontrol by preemptive exclusion (PE), mostly via top-down population-based approaches, i.e. by looking at the effect of pre-inoculation of a BCA strain and a plant pathogenic strain on the colony forming units of both strains that can be recovered and/or on the development of disease symptoms on plants (Wilson and Lindow 1994; Braun-Kiewnick, Jacobsen et al. 2000; Nix, Burpee et al. 2009; Stockwell, Johnson et al. 2010; Xu, Salama et al. 2010). However, BCA applications sometimes fail (Andrews 1992; Kinkel, Newton et al. 2002),
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and the reasons for this failure are not always clear. For some cases, it has been suggested (Kinkel, Newton et al. 2002) that the applied density of a BCA might not have been sufficient to reach all growth-conducive patches on a leaf or that the time since after BCA application might not have been long enough for the BCA to fill up these patches and to prevent all secondary plant pathogens to find and colonize unfilled or partially filled patches. To address this question from a single-cell, bottom-up perspective, we utilized a recently established single-cell bioreporter for bacterial reproductive success, \textit{Eh299RCUSPER} (Remus-Emsermann and Leveau 2010); this thesis chapter 3). This bioreporter is based on the model phyllosphere colonizer \textit{Erwinia herbicola} 299R (also known as \textit{Pantoea agglomerans} 299R), which can be loaded with green fluorescent protein. Bacterial reproduction can then be followed at the single-cell level by determining the dilution of the fluorescent signal, which occurs at each cell division. Prior to the release of CUSPER cells, the bean plants were pre-inoculated with different densities of the red fluorescent \textit{E. herbicola} bioreporter strain, \textit{Eh299RdsRed}. Because \textit{Eh299RCUSPER} and \textit{Eh299RdsRed} are nearly isogenic (they only differ in the color of fluorescent protein that they produce), they can be considered each others’ most effective competitors, in that they have a nutrient overlap index of 1, which is considered optimal for biocontrol through preemptive exclusion (Wilson and Lindow 1994).

\textbf{Materials and Methods:}

\textit{Bacterial strains and culture conditions}

\textit{Eh299RCUSPER} is based on \textit{Erwinia herbicola} 299R::JBA28 (pCPP39) (Leveau and Lindow 2001). Plasmid pCPP39 constitutively expresses the repressor LacI\textsuperscript{q} and confers resistance to tetracycline. The chromosomal mini-Tn5-Km transposon insertion JBA28 carries a LacI\textsuperscript{q} repressable GFPmut3 gene and confers resistance to kanamycin. \textit{Erwinia herbicola} 299R (pFRU97) (\textit{Eh299RdsRed}) carries plasmid pFRU97 (Leveau, unpublished) which contains a constitutively expressed red-fluorescent dsRed gene and confers resistance to kanamycin.

\textit{Eh299RCUSPER} cells were cultured in LB broth supplemented with 50 µg kanamycin per mL, 15 µg tetracycline per mL and 1 mM of the LacI\textsuperscript{q} derepressor IPTG, which activates the expression of GFPmut3, resulting in cells loaded with green fluorescent protein. \textit{Eh299RdsRed} was cultured in LB broth supplemented with 50 µg kanamycin per mL. After growth to mid-exponential phase at 300 rpm and 28°C, bacteria were harvested by centrifugation at 3900 \( \times g \) for 10 minutes, and resuspended in 1× PBS (8 g NaCl, 0.2 g KCl, 1.44 g Na\textsubscript{2}HPO\textsubscript{4}, and 0.24 g KH\textsubscript{2}PO\textsubscript{4} per liter H\textsubscript{2}O, adjusted to pH 7.4).
Inoculation of bean plants with \( Eh299R_{\text{dsRed}} \)

Cells of \( Eh299R_{\text{dsRed}} \) were grown and harvested as described above and resuspended in PBS to concentrations of \( 10^6 \), \( 10^7 \), and \( 10^8 \) CFU per mL, as approximated by optical density. Fully expanded cotyledon leaves of two-week-old \( Phaseolus vulgaris \) plants (green snap bean, variety Blue Lake Bush 274) were each airbrushed as described previously (this thesis chapter 3) with 500 µL of one of the bacterial suspensions, or with 500 µL sterile PBS. Plants were then transferred to translucent plastic boxes that were closed to retain high relative humidity and incubated for 24 hours under constant illumination at room temperature. Subsequently, each pre-inoculated leaf was inoculated by airbrushing with 500 µL of a freshly prepared suspension of \( 10^6 \) colony-forming units (CFUs) of \( Eh299R_{\text{CUSPER}} \) per mL. Three leaves per treatment were sampled 0, 3, 6, 8, and 24 hours after inoculation with \( Eh299R_{\text{CUSPER}} \).

Recovery of bacteria from leaves and measurement of CFU counts and single-cell GFP content

Sampled leaves were individually transferred to a 50-ml Falcon tube with 20 ml 1× PBS buffer, vortexed briefly, sonicated for 7 min, and vortexed briefly again. Two 50-µL aliquots of each leaf wash were plated onto duplicate LB agar-plates supplemented with 50 µg kanamycin per mL or 20 µg tetracycline per mL using an Eddy Jet spiral-plater (IuL, Barcelona, Spain) in order to determine the number of CFUs of \( Eh299R_{\text{CUSPER}} \) (i.e. the number of colonies on the tetracycline plates) and \( Eh299R_{\text{dsRed}} \) (i.e. the number of colonies on the kanamycin plate minus the number of colonies on the tetracycline plates). The remainder of the leaf wash was filtered through a 0.2-µm Isopore filter (Millipore, Amsterdam, The Netherlands). Cells were recovered from the filter by vortexing for 15 sec in 1 ml 1x PBS, fixed as described before (Leveau and Lindow 2001) and stored in 1x PBS at 4°C for no longer than 3 days. Cells were analyzed using an Axio Imager.M1 epifluorescent microscope (Zeiss, Oberkochen, Germany) using phase contrast and Zeiss Filter 38 (BP 470/ 40, FT 495, BP 525/ 50) for the visualization of GFP and Filter 20 (BP 546/ 12, FT 560, BP 575-640) for the visualization of dsRed. Digital images were captured at 1000-fold magnification with an AxioCam MRm CCD-camera (Zeiss, Jena, Germany). Single-cell green fluorescence intensity of non-red fluorescent cells was quantified using the AxioVision 2.8.2 software package (Zeiss, Jena, Germany) as the mean-pixel intensity of individual cells (Leveau and Lindow 2001). Single-cell green fluorescence measurements were used to reconstruct the relative contribution of original leaf immigrants to the final population as explained elsewhere (this thesis chapter 3). Data analysis
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Results

Changes in population size of \( \text{Eh299R}_{\text{CUSPER}} \) cells on bean leaves pre-colonized with \( \text{Eh299}_{\text{dsRed}} \)

Bean leaves were inoculated with different amounts of \( \text{Eh299R}_{\text{dsRed}} \) (primary colonizers) to reach population sizes of \( 2.09 \times 10^6 \), \( 6.31 \times 10^6 \), and \( 6.76 \times 10^7 \) CFU per gram leaf after 24 hours of incubation. Subsequent inoculation of these pre-colonized leaves with approximately \( 5 \times 10^5 \) cells of \( \text{Eh299R}_{\text{CUSPER}} \) revealed that the increase in population size of these secondary colonizers over the next 24 hours was inversely correlated with the level of pre-colonization by \( \text{Eh299}_{\text{dsRed}} \) (Figure 1A). The data fitted a downward-sloping response-curve (Figure 1B), with an approximate threshold of \( 1 \times 10^5 \) CFUs of \( \text{Eh299}_{\text{dsRed}} \) per gram of leaf tissue, i.e. below this population density the secondary colonizers would do as well as when the leaves were not first sprayed with \( \text{Eh299}_{\text{dsRed}} \). The effective range of \( \text{Eh299R}_{\text{dsRed}} \) reached up to approximately \( 1 \times 10^8 \) CFUs per gram of leaf, with an apparent IC50 value of 6.116, or \( 1.31 \times 10^6 \) CFUs of \( \text{Eh299R}_{\text{dsRed}} \) per gram leaf tissue.
The level of pre-colonization by $Eh299R_{dsRed}$ had a strong impact on the reproductive success of individual secondary immigrants of $Eh299R_{CUSPER}$ (Figure 2). The higher the population of $Eh299R_{dsRed}$ bacteria, the lower the average reproductive success of $Eh299R_{CUSPER}$ cells, resulting in a shift to the left in the corresponding histograms. At the highest levels of pre-colonization, we observed almost no reproduction of $Eh299R_{CUSPER}$, although a small number of immigrants (about 1%) underwent two cell divisions after 24 hours (Figure 2, bottom row).

By reconstructing the increase in population size from single-cell measurements (as was done in chapter 3), we were able to show the increases in secondarily inoculated populations as a function of the CFU of $Eh299R_{dsRed}$ at time $t_0$ (Figure 3). The resulting graph had a similar shape as the one in Figure 1B, i.e. the data fit a downward-sloping response curve (Figure 3). The IC50 for this curve was calculated to be 6.204, or $1.59 \times 10^6$ CFUs of $Eh299R_{dsRed}$ per gram leaf.

**Discussion**

In situations where primary and secondary colonizers use the exact same resources,
Secondary colonizers are expected to be less successful in the colonization of a pre-populated environment (Wilson and Lindow 1994). We found that the ability to increase in population size of the secondary colonizer, *Eh*$_{299R}$$_{CUSPER}$, decreased with increasing levels of preemptively inoculated *Eh*$_{299R}$$_{dsRed}$ cells. This phenomenon was observed both at the population level and the single-cell level (Figure 1B and 3). These findings are in line with a previous study, where it was found that increased densities of *Eh*$_{299R}$$_{CUSPER}$ inoculants decreased the reproductive success of the overall population (this thesis chapter 3). Interestingly, despite this agreement between the two levels of analysis, there were instances where we found a small proportion of the cells to reproduce and form offspring in the absence of a total increase in population size. For example, 3% of *Eh*$_{299R}$$_{CUSPER}$ cells were able to reproduce three times to create 8 offspring each on leaves that were heavily pre-populated at 6.31x10$^6$ CFUs per gram leaf (Figure 2, column 4, row 3). This is a relevant observation if one considers that for foliar pathogens such as *Pseudomonas syringae* the quorum size, i.e. the number of cells in a local population to trigger a common behavior by signaling molecules, can be as low as 13 cells or even fewer (Dulla and Lindow 2008). Thus, a microcolony of 8 cells might be sufficiently large to trigger quorum

Figure 3. The increase in population size of *Eh*$_{299R}$$_{CUSPER}$ as derived from single-cell measurements at t=24 was plotted against the population size of *Eh*$_{299R}$$_{dsRed}$ at $t_0$. The curve shows the fitted dose response (value = best-fit ± std. error, log top-response = 1.38 ± 0.022, log bottom-response = 0.1835 ± 0.022, log IC50 = 6.204 ± 0.041, r$^2$ = 0.99, formula: $y = bottom$-response + (top-response − bottom-response) / (1 + 10$^{(x−LogIC50)}$)). The slope of the fitted curve was fixed to a standard slope, or Hill-slope, of 1. X and Y Error-bars represent standard deviations of the mean.
sensing behavior, which has been shown for *P. syringae* to be important for pathogenicity (Quiñones, Dulla et al. 2005). This observation supports the notion that bacterial reproduction or the expression of pathogenicity factors, on leaf surfaces may not always be predicted accurately from population-based measurements. This pertains not only to plant pathogens, but possibly also to other unwanted contaminants of leaf surfaces, such as the human pathogen *Escherichia coli* O157:H7. Several studies have documented the rapid decline of O157:H7 or proxy isolates on leafy greens (Delaquis, Bach et al. 2007; Moyne, Sudarshana et al. 2011), and typically this decline is measured at the population level as CFUs that can be washed off leaf surfaces. Our study suggests that, masked by this population decline, there might be some cells that actually do quite well and are able to survive, or even reproduce into microcolonies that offer advantages for survival (Monier and Lindow 2005). The ability of some bacterial immigrants to be highly successful on leaf surfaces while most others are not can be explained by the existence of “oases” of relative abundant nutrients (Lindow and Brandl, 2003). It would appear that the challenge for biocontrol strategies that depend on the principal of preemptive exclusion lies in 1) the occupation of all of nutritional oases and 2) the complete exhaustion of available resources in each of these oases.

Secondary colonizers might face different situations after reaching the leaf: they arrive either in sites that are un-occupied or already occupied by primary colonizers. In sites that are un-occupied, colonization success will be a function of the local carrying capacity (this thesis chapter 3). In occupied sites on the other hand they will have to compete with the previous colonizer, and colonization success will then be determined by the amount of resources left in the sites. This can include those resources that the primary colonizers is incapable of utilizing, or might depend on the amount of resources that are leaching locally from the leaf to the phyllosphere (Remus-Emsermann, de Oliveira et al. 2011). Interestingly, it has been suggested that if secondary colonizers land in already established aggregates, their survival is higher compared to cells that land in un-occupied sites (Monier and Lindow 2005). Thus, it is not necessarily all precolonized sites that do not allow growth of a secondary colonizer. These sites could allow growth of secondary colonizers, if they still carry enough resources at the time of colonization or the replenishment of resources is high enough.

The presented results show that increasing numbers of precolonizers effectively reduces the reproductive success of secondary colonizers. These results also suggest that to prevent foliar growth of a secondary strain or pathogen by the principle of PE, even a BCA that shares a high resource overlap with the secondary colonizer and
that is inoculated at high densities might still not be able to prevent the reproduction of certain immigrant pathogens. In a field situation, a preemptive colonizer is likely to perform even less well than under the ideal conditions in the lab, for example due to incomplete or uneven application onto leaves. To avoid the presence of sites that are free for the colonization by secondary colonizers, unoccupied sites would need to be re-occupied by the biocontrol agent by migration processes or have to be reinoculated.

Our observation that a small fraction (0.7-2.4%) of the cells was very successful in reproducing even at the highest inoculation density suggests the existence of sites on the leaf surface that offer high levels of nutrients, allowing these cells to contribute disproportionally (11-31%) to the final population size. Interestingly, this finding agrees with reports showing that relatively rare features on the leaf surface tend to harbor large numbers of bacteria on colonized leaves. Specifically, it is worth noting that while only 0.6% of the bean leaf surface is covered by hooked and glandular trichomes (Monier and Lindow 2005), yet 44.1% of Eh299R bacteria in aggregates were found to be associated with these structures after incubation under moist conditions for 5 days (Monier and Lindow 2005). In at least one study (Yadav, Karamanoli et al. 2005), the population size of epiphytic bacteria was positively correlated with the densities of glandular and non-glandular trichomes. This link between bacterial fate, population size and leaf topography calls for further investigation into the predictive value of trichome density for bacterial colonization of leaf surfaces.

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Quantification of lateral heterogeneity in carbohydrate permeability of isolated plant leaf cuticles
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Abstract
In phyllosphere microbiology, the distribution of resources available to bacterial colonizers of leaf surfaces is generally understood to be very heterogeneous. However, there is little quantitative understanding of the mechanisms that underlie this heterogeneity. Here, we tested the hypothesis that different parts of the cuticle vary in the degree to which they allow diffusion of the leaf sugar fructose to the surface. To this end, individual, isolated cuticles of poplar leaves were each analyzed for two properties: 1) the permeability for fructose, which involved measurement of diffused fructose by gas chromatography and flame ionization detection (GC-FID), and 2) the number and size of fructose-permeable sites on the cuticle, which was achieved using a green fluorescent protein (GFP)-based bacterial bioreporter for fructose. Bulk flux measurements revealed an average permeance $P$ of $3.39 \times 10^{-9}$ m s$^{-1}$, while the bioreporter showed that most of the leaching fructose was clustered to sites around the base of shed trichomes, which accounted for only 0.37% of the surface of the cuticles under study. Combined, the GC-FID and GFP measurements allowed us to calculate an apparent rate of fructose diffusion at these preferential leaching sites of $9.2 \times 10^{-7}$ m s$^{-1}$. To the best of our knowledge, this study represents the first successful attempt to quantify cuticle permeability at a resolution that is most relevant to bacterial colonizers of plant leaves. The estimates for $P$ at different spatial scales will be useful for future models that aim to explain and predict temporal and spatial patterns of bacterial colonization of plant foliage based on lateral heterogeneity in sugar permeability of the leaf cuticle.
Introduction
For terrestrial plants, the cuticle represents the direct interface between aerial plant parts and the atmosphere. It predominantly consists of cutin, an esterified aliphatic polymer (Kolattukudy 1980) that is impregnated and overlaid by intra- and epicuticular waxes, respectively (Jetter and Schäffer 2001). The thickness of plant cuticles ranges from 30 nm for Arabidopsis thaliana leaves to 30 µm for Malus domestica fruit (Schreiber and Schönherr 2009). From a plant perspective, the cuticular membrane plays a pivotal role in preventing the loss of water, minerals, and nutrients, and in protecting the plant against pathogenic attacks (Riederer and Müller 2006). The hydrophobic nature of plant cuticles correlates with a limited permeability for polar substances such as water and ions (Schönherr and Schreiber 2004). By comparison, plant cuticles are less permeable for water than synthetic polymers such as polyethylene or polypropylene of equal thickness (Riederer and Schreiber 2001). Another difference between plant cuticles and synthetic membranes is that the former are characterized by a pronounced lateral heterogeneity. Plant leaves are oftentimes decorated with features such as trichomes, gland cells, and stomata, (Schlegel, Schönherr et al. 2005; Schreiber 2005), and these structures may differ substantially in the thickness and wax composition of the cuticle that covers them, as shown recently for trichomes of a pubescent peach fruit variety (Fernández, Khayet et al. 2011). For example, the leaf cuticle tends to be thicker over anticlinal cell walls between epidermal cells than on top of epidermal cells (Jeffree 2007).

Despite the design of the leaf cuticle to prevent the loss of water and nutrients, it has long been recognized that plant leaves may lose measurable amounts of carbohydrates and other substances to the leaf surface through a process called leaching (Tukey 1966). These “leachates” are used by microorganisms that inhabit the leaf surface, or phyllosphere (Andrews and Harris 2000). For many of these bacteria, yeasts, and fungi, the phyllosphere is considered a carbon-limited environment (Wilson and Lindow 1995; Wilson, Savka et al. 1995), which means that these microbes rely on a constant flux of photosynthates such as glucose, fructose and sucrose from the leaf interior. A recent study (van der Wal and Leveau 2011) estimated that the diffusion of fructose through the isolated cuticular membrane (CM) of Juglans regia (walnut) leaves is sufficient to sustain microbial populations as they are typically found in the phyllosphere. Others have come to the same conclusion using isolated CMs of Prunus laurocerasus, or cherry laurel (Krimm 2005).

So far, the quantification of sugar diffusion across plant leaf cuticles has been restricted mainly to bulk measurements integrating sugar flux over entire leaves (Tukey 1966; Mercier and Lindow 2000) or over isolated CM discs (Schönherr
Estimations of sugar permeance at smaller scales, for example the scale of individual bacteria, are not available, although the use of fluorescent bioreporters has previously revealed the existence of ‘hotspots’ of fructose availability to bacteria on the plant leaf surface (Leveau and Lindow 2001; Krimm 2005). Possibly, such sites represent areas where the cuticle is more permeable than in other locales on the leaf surface. A quantitative understanding of this lateral heterogeneity in carbohydrate permeance at the micrometer scale will improve our ability to explain and predict foliar growth of microorganisms, including those that are plant pathogenic.

To achieve such understanding, we employed an integrated approach that combined bulk flux measurements of fructose across discs of isolated cuticles with the analysis of single-cell bioreporter information on local fructose availability on one and the same set of cuticles. The bioreporter was based on *Erwinia herbicola* 299 (*Eh*299), a natural colonizer of the phyllosphere (Brandl and Lindow 1996) which has been used in many phyllosphere studies (Brandl, Quinones et al. 2001; Leveau and Lindow 2001; Krimm 2005; Remus-Emsermann and Leveau 2010; van der Wal and Leveau 2011). *Eh*299 is equipped with a construct allowing expression of the green fluorescent protein (GFP) in response to fructose exposure (Leveau and Lindow 2001). On the plant side, we chose to work with CMs isolated from the adaxial, astomatous, mature leaves of poplar (*Populus x canescens*). Being a model system for the molecular biology of trees (Bradshaw, Ceulemans et al. 2000), the industrial production of wood, and phytoremediation applications (Stanton, Eaton et al. 2002), poplar also has played an important role in research on the permeability of plant leaf cuticles for ions (Schönherr and Schreiber 2004; Schreiber, Elshatshat et al. 2006). Experiments with isolated leaf cuticles of *Populus x canescens* suggested the existence of aqueous pores in the cuticle, of sufficient hydrophilicity to allow the passage of ions such as Ca$^{2+}$, Cl$^-$, and Ag$^+$ (Schönherr and Schreiber 2004; Schreiber, Elshatshat et al. 2006). As leaves of *Populus x canescens* mature, trichomes are shed, and ion passage (more specifically, the formation of microscopically visible AgCl crystals) was most abundantly found to occur at the bases of these shed trichomes. It was ruled out that shedding resulted in the formation of holes; instead, the authors argued for the existence of polar pathways, also referred to as aqueous pores (Schönherr and Schreiber 2004) at these sites. This finding challenged the model of slow diffusion of polar substances through a laterally homogeneous cuticle in favor of a model featuring much more lateral heterogeneity (Franke 1967; Schreiber and Schönherr 2009). For a fundamental understanding of microbial phyllosphere colonization, this lateral heterogeneity is of great importance, because it could be used to explain the
nonrandom patterns of leaf colonization by bacteria (Monier and Lindow 2004) In
the study presented here, we confirm that shed trichome sites are more permeable for
sugars and provide quantitative estimates for the permeance and frequency of sugar-
permeable sites on the poplar leaf surface.

**Material and Methods**

*Isolation of cuticular membranes*
Isolation of CMs was conducted as described elsewhere (Schreiber and Schön-
herr 2009). In short, it involved the digestion of 20-mm diameter discs of
*Populus x canescens* (Aiton) Sm. leaves harvested in Spring 2002 near Sarstedt,
Germany, with 2% cellulase and pectin-
ase in 0.01 M citric acid buffer adjusted
to pH 3-4 with KOH. To prevent growth
of microorganisms, NaN$_3$ was added to
the solution at a final concentration of 1
mM. After complete digestion, isolated
CMs were flattened with compressed air
on Teflon discs and stored at room tem-
perature and low ambient humidity.

*Measuring of fructose bulk flux by gas chromatography and flame ionization detection (GC-FID)*
To measure the bulk flux of fructose over
cuticles, cylindrical stainless steel cham-
bers were used as described in Schreiber
and Schönher (2009). The chambers
(Figure 1) consisted of a donor-chamber,
filled with 1 mL of a 180 g per L solution
of fructose, and a receiver-chamber filled
with deionized water. Both donor and receiver had an inner diameter of 1.2 cm and
were equipped with lockable sampling holes to allow sampling and exchange of so-
lutions with a syringe. With their physiological outside facing the receiver-chamber,
isolated CMs were mounted on top of the donor chamber ring and sealed with high

![Figure 1 Setup of the cuticle diffusion chamber. See text section Measuring of fructose bulk flux by gas chromatography and flame ionization detection (GC-FID) for details.](#)
vacuum silicon grease to the receiver chamber. The chambers were placed on a rolling bench at 28 °C and samples were taken from the receiver chamber after 5 and 24 hours. Before analysis of these samples by GC/FID (see below), we tested 100 µl of each for fructose using 20 µl Benedict’s reagent (Benedict 1909) to verify the integrity of the CMs and the integrity of the chambers with mounted CMs. Those CMs or chambers that were found to be leaky because they had similar fructose concentrations in donor and receiver compartments after 5 hours were discarded and not used for further analysis. Receiver samples harvested from intact CMs and chambers were supplemented with 3 µg xylitol to serve as an internal standard, dried under constant N2-flow at 70 °C, derivatized to trimethylsilyl-ethers by adding 30 µl of the catalyst Pyridine and 30 µl of Bis(trimethylsilyl)-trifluoracetamid and incubated for 40 min at 70 °C. Fructose concentration was determined by GC-FID (5890 Series II Plus, HP, Agilent Technologies, Böblingen, Germany), equipped with a dimethylpolysiloxane column (DB1 GC column, 30 m x 0.32 mm, 0.1 µm, J&W Scientific, Folsom, USA). One µl of each sample was injected. H2 flow was set to 37 kPa. The initial temperature was 65 °C for 3 minutes, after which the temperature was raised at a rate of 8 °C per min to a temperature of 240 °C, after which the rate was increased by 12 °C per min to a final temperature of 310 °C for 35 minutes. Data analysis was performed with GC ChemStation (Rev. B.03.02 (341), Agilent Technologies, Böblingen, Germany). Permeances were calculated using the following equation (Schreiber, Elshatshat et al. 2006): P=F/(AxAΔc). Fructose flow F (g s⁻¹) was obtained from the regression line fitted to the points that were obtained by plotting the amount of fructose (in grams) in the receiver chamber as a function of time, while A (m²) represents the area of exposed cuticle (1.13x10⁻⁴ m²) and Δc (g m⁻³) the difference in fructose concentration between the receiver and donor compartment (i.e. 180 g per L or 1.8 *10⁵ g per m³).

Inoculation of isolated CMs with bacteria
To measure localized diffusion of fructose, the receiver chambers were unmounted, the donor chambers were emptied, and the outside and inside of the CMs were washed with deionized water. The donor chamber was refilled with one mL of an 18 g L⁻¹ fructose solution. Cells of Eh299R carrying (pPfruB-gfp[AAV]) (Leveau and Lindow 2001) were grown overnight on LB agar plates supplemented with 50 ng kanamycin per mL at 30 °C, spun down for 10 min at 3000x g, washed twice in sterile water, and subsequently diluted with sterile water to an optical density of 1 at 600 nm, which corresponds to approximately 4x10⁸ bacterial colony forming units per mL. Plasmid (pPfruB-gfp[AAV]) confers resistance to kanamycin and codes for the expression of a short-lived version of the green fluorescent protein GFP[AAV] under
the control of the fructose-responsive *fruB* promoter (Leveau and Lindow 2001). CMs were inoculated by pipetting 400 µl of this bacterial suspension on the physiological outer side of the cuticle as described by Knoll and Schreiber (Knoll and Schreiber 2004). Diffusion chambers were incubated with the inoculated side facing up for 6 hours at high relative humidity and 28 °C to facilitate adhesion of bacteria to the CM. Subsequently the bacterial suspension was carefully removed by pipetting and drying at room humidity for not more than 5 minutes. Next, the setup was transferred into a high humidity chamber and incubated at 28 °C for 16 hours.

**Fluorescence microscopy and image analysis**

For analysis of inoculated cuticles by fluorescence microscopy and to reveal the location of reporter bacteria responding to the local availability of fructose, the donor solution was removed and a 10 µl droplet of 75% glycerol supplemented with DAPI, to counterstain all bacteria (whether they were reporting fructose or not), was applied to the center of the inoculated side of the cuticle and covered by a cover slip. Microscopy was performed using a Zeiss AxioPlan stereomicroscope at 400-fold magnification and two filter sets: BP365/FT395/LP397 (Zeiss filter set 1) for visualization of DAPI-stained cells and leaf autofluorescence, and 450-490/FT510/515-565 (Zeiss filter set 10) for visualization of bacteria expressing GFP. Images were taken in grey scale mode with a DMX1200 charged coupled device (Nikon Corporation, Japan) and the Software Act-1 version 2.70 (Nikon Corporation, Japan). Images were analyzed with the software package ImageJ 1.45b (Abramoff, Magelhaes et al.

Figure 2 Diffusion of fructose across isolated Populus x canescens leaf cuticles. A) Accumulation of fructose (in grams) as a function of time at a donor concentration of 0.18 g per mL. Shown is the linear regression ($r^2 = 0.99$) with 95% confidence intervals (broken lines) of the mean (n=10) diffusion of fructose over Populus x canescens leaf cuticles. Error-bars represent the standard error of measurement. B) Probability plot of the log-transformed fructose permeability ($P$, in m s$^{-1}$) of individual Populus x canescens CMs. The dataset passed the D’Agostino & Pearson omnibus test for normality ($P = 0.28$)
Figure 3 GFP-based bioreporting of fructose availability on isolated *Populus x canescens* cuticles. A-E: Epifluorescent images of *Populus x canescens* CMs inoculated with *Eh299*(pPfruB-gfp[AAV]). White scale bars represent 20 µm. Images are pseudo-colored merges of DAPI (blue) and GFP (green) channels; when fluorescence occurred in both channels the visible color is yellow. The blue channel mainly shows autofluorescence of the CM and DAPI counterstained, non-fructose reporting bacteria. A) Area with many bacterial bioreporter cells but none that fluoresce to indicate exposure to fructose; the right-hand panel is a magnification of the photograph shown on the left-hand site, to show individual DAPI-stained bacteria. B-D) Typical arrangements of bioreporting bacteria on isolated *Populus x canescens* CMs. F) Light microscopy image of a *Populus x canescens* CM. Arrows point to sites with shed-off trichomes.
2004) by creating pseudo-RGB images from photographs taken from the same area of the cuticle using the different filter sets. The area covered by green-fluorescent bacteria was determined using imageJ’s measurement tools.

Results

*Estimating fructose diffusion across isolated* *Populus x canescens* *cuticles*

The bulk permeance of ten individual CMs from poplar leaves was analyzed using a two-compartment experimental setup (Figure 1) that allowed the measurement of fructose diffusion from a donor compartment with a constant high fructose concentration across an isolated cuticle into a receiver compartment. Fructose abundances in the receiver compartment were quantified by gas chromatography and plotted as a function of time (Figure 2 A). From the slope of the best-fit line through these points, bulk permeances of *Populus x canescens* CMs for fructose were calculated as ranging from $2.54 \times 10^{-10}$ to $2.78 \times 10^{-8}$ m s$^{-1}$. Values were log-normally distributed around an average log $P$ of $-8.46 \pm 0.67$, which back-transformed to an average $P$ of $3.39 \times 10^{-9}$ m s$^{-1}$ (Figure 2 B).

*Frequency and size of fructose permeable sites on* *Populus x canescens* *cuticular membranes*

We performed fluorescence microscopy analysis on three CMs that immediately after the bulk flux measurements (see above) were exposed to the bacterial bioreporter *Eh299R(pP$_{fruB}$-gfp[AAV])) which responds to the presence of fructose by producing green fluorescence. The inoculation resulted in a homogeneous and random distribution of bacteria on the CMs as shown by DAPI counterstaining (Figure 3 A). In contrast, the distribution of DAPI-stained cells that also were green fluorescent was
not random. Many occurred as part of circle- or ring-shaped clusters of different diameters (Figure 3B, C, D). The sizes of these clusters from three CMs were measured and found to be lognormally distributed (Figure 4), with many small clusters and few large clusters. The largest site covered by green fluorescent bacteria was $4.7 \times 10^4 \mu m^2$. Values were distributed around a log average of $3.42 \pm 0.74$ which back-transformed to an average size of $2.62 \times 10^3 \mu m^2$.

The majority (79%) of clusters of green fluorescent bacteria were associated with sites of shed trichomes (e.g. Figures 3B and 3C). Thus, these structures appeared to be the main sites of fructose diffusion. We determined that 1 cm$^2$ of the *Populus x canescens* leaf cuticles we used contained on average 138±67 shed trichome sites. Assuming a sugar-permeable area of $2.62 \times 10^3 \mu m^2$ for each of these (see above), we can calculate that only 0.37% of the leaf cuticle surface allowed diffusion of fructose high enough to induce GFP expression. Correspondingly, the permeance of the *Populus x canescens* CM at these sites would be $100/0.37 = 270$ times higher than our bulk estimate, i.e. $270 \times 3.39 \times 10^{-9} = 9.15 \times 10^{-7}$ m s$^{-1}$.

Discussion

The GC-FID-based detection method used here allowed us to determine that the average bulk permeance of *Populus x canescens* CMs for fructose was one orders of magnitude higher than that reported for isolated CMs of *Juglans regia*, i.e. $2.79 \times 10^{-10}$ m s$^{-1}$ (van der Wal and Leveau 2011), but two orders of magnitude higher than *Prunus laurocerasus*, i.e. $6.64 \times 10^{-11}$ m s$^{-1}$ (Krimm 2005). This is consistent with earlier findings (Kirsch, Kaffarnik et al. 1997) that cuticles of evergreens such as *P. laurocerasus* are less permeable than cuticles of deciduous trees such as *Populus x canescens* and *J. regia*.

Aqueous pores are thought to be made up of hydrated polar functional groups in the cuticle (Chamel, Pineri et al. 1991). Values reported for aqueous pore sizes of plant leaf cuticles were determined for a series of isolated cuticles and showed average radii of 5 Å estimated from trans-cuticular Ag$^+$- and Cl$^-$-diffusion (Schreiber, Elshatshat et al. 2006). The hydrodynamic radius of a fructose molecule has been calculated to be 3.61 Å (Ribeiro, Santos et al. 2006), thus permeance of fructose through these pores should be possible. Interestingly, the estimates derived from our study can be used to get a better appreciation for the density of aqueous pores. The permeance $P$ of an individual pore can be calculated from diffusion coefficient $D$ for fructose in water, i.e. $7.44 \times 10^{-10}$ m$^2$ s$^{-1}$ at 30°C (Ribeiro, Santos et al. 2006) and the length $L$ of the path along which diffusion takes place. If we assume that the latter is approximated by the thickness of the cuticle, $2.8 \times 10^{-6}$ m (Todeschini, Lingua et al. 2011), $P$ can be calculated as $D/L$ or $2.66 \times 10^{-4}$ m s$^{-1}$. This value is $7.84 \times 10^4$ times
higher than the bulk permeance we measured for the poplar cuticle. If diffusion of fructose only takes place through aqueous pores we can now estimate the relative area of the cuticle that is effectively permeable to fructose to be \(100\% / 7.84 \times 10^4 = 1.28 \times 10^{-3}\%\). Assuming a pore radius of 5 Å, i.e. a pore area of \(2.5 \times 10^{-19}\) m\(^2\), the total number of pores per m\(^2\) can be calculated as \(5.1 \times 10^{13}\), which is close to the value of \(3.5 \times 10^{14}\) to \(2 \times 10^{15}\) pores per m\(^2\) reported for isolated Citrus aurantium cuticles (Schreiber and Schönherr 2009). Assuming that pores only occur at permeable sites (i.e. the basis of trichomes), the density of aqueous pores at these sites may be calculated as 270 times higher, i.e. \(1.38 \times 10^{18}\) aqueous pores per m\(^2\). However, these calculated values have to be used with great caution, and will need to be verified in future studies.

Our results with the GFP-based fructose bioreporter indicate that the poplar leaf cuticle features sites that are more permeable to fructose than most other sites on the cuticle. These preferential sites seemed to be clustered around shed trichomes. It has been shown that the base of plant leaf trichomes can be heavily cutinized (Fernández, Khayet et al. 2011), and so our findings and those of others (Schreiber, Elshatshat et al. 2006) can be explained by assuming that the observed increased permeability is specific to shed trichomes, representing sites that are structurally less integer than non-modified cuticular areas, as has been suggested for lenticels on fruit (Glenn and Poovaiah 1985) In the absence of measurement of cutin and waxes, however, this remains speculative and needs to be confirmed in future experiments. The predominant presence of sugar-diffusible sites around trichomes on the Populus x canescens cuticle is in good agreement with general observations that bacterial leaf colonizers often establish large populations around the base of trichomes (Monier and Lindow 2004). Trichomes have also been identified as sites where sugars are commonly and abundantly available to leaf-colonizing bacteria (Leveau and Lindow 2001). Heterogeneity in the availability of nutrients on the leaf surface is consistent with the recent observation that bacterial immigrants to the bean phyllosphere experience very different fates (Remus-Emsermann and Leveau, 2010): while most cells divided a few times upon arrival, a few divided many more times. The latter was explained by assuming that those cells landed in sites of high nutrients access. Plants other than Populus x canescens, for example Vicia vaba and P. vulgaris posses aqueous pores not only around trichomes but also associated with other leaf features, such as guard cells of stomates, anticlinal cell walls and glandular trichomes (Schlegel, Schönherr et al. 2005; Schreiber and Schönherr 2009). Based on this difference, our prediction would be that patterns of bacterial colonization on leaves from these plants would be different from those observed for poplar.
In conclusion, the poplar leaf cuticle features sites that are more permeable to fructose than most other sites on the cuticle. These preferential sites seemed to be clustered around shed trichomes. The estimated values for diffusion rates at these sites provide the first numerical estimation of micrometer scale carbohydrate availability in the phyllosphere. They will help to explain microbial growth patterns in the phyllosphere and contribute to modeling of microbial growth in the phyllosphere, which so far has been hampered by experimentally derived estimates of carbohydrate diffusion and availability at scales that matter most to potential colonizers of the phyllosphere.

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Agent-based Simulation of Phyllosphere Colonization

ASiMoPh – Agent-based Simulation of Microbial Phyllosphere Colonization

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Abstract

ASiMoPh is an agent-based model of phyllosphere colonization, which was developed to unravel the underlying causes of heterogeneity that influence bacterial growth in the phyllosphere. ASiMoPh describes the phyllosphere as a two-dimensional domain of patches that contain different amounts of energy. Agents, representing bacteria, are released in this environment and reproduce based on the energy in their patch. The number of divisions that agents undergo is recorded and expressed as reproductive success. The total number of agents in the phyllosphere environment and in individual patches is also tracked. ASiMoPh reported a direct dependence of agent inoculation density on single-agent reproductive success in that higher initial densities lowered reproductive success. Furthermore, ASiMoPh reported that preemptively inoculated agents had a similar effect on secondary agents, lowering the reproductive success of secondary colonizers dramatically at high inoculation densities. Both results are in good agreement with \textit{in vitro} experiments performed with a bacterial bioreporter of reproductive success based on the bacterium \textit{Erwinia herbicola} on \textit{Phaseolus vulgaris} cotyledon-leaves (data obtained from chapter 3 and 4 in this thesis). In future studies ASiMoPh can serve as a foundation for phyllosphere
modeling to answer more complex questions and incorporate more agent-agent and agent-environment interactions.

**Introduction**

Microorganisms have traditionally been investigated at the level of populations and studies at the level of individual organisms and their interactions are still under-represented in microbiology. However, there are many good reasons for shifting the microbiological perspective to the micrometer scale at which microorganisms experience their habitat, e.g. since microscopical techniques and fluorescent reporter and probe technologies made revolutionary advances in the last decades, the observation of individual bacteria is well possible (Pernthaler, Glöckner et al. 2001; Leveau and Lindow 2002; Musat, Halm et al. 2008; Remus-Emsermann, de Oliveira et al. 2011). Even microorganisms from a clonal population exhibit individuality in a homogeneous environment (Davidson and Surette 2008; Roostalu, Joers et al. 2008). In a microbiological context, individuality means that microorganisms occupy a unique place in time and space that they do not share with others and have a unique cellular composition (Hughes 1956; Levin, Morton-Firth et al. 1998; Strovas, Sauter et al. 2007) which can create different stable phenotypes even within a clonal (i.e. genotypically identical) population (Davidson and Surette 2008).

Two types of tools are used in the study of single bacterial cells. One is based on experimental techniques and allows us to manipulate and interrogate individual bacteria (Brehm-Stecher and Johnson 2004; Müller and Nebe-von-Caron 2010), the other tool involves modelling approaches to derive testable hypotheses, facilitate experimental planning, and analyze ecological processes. The number of applications of individual-based models (IBMs) in microbial ecology is rapidly growing (Hellweger and Bucci 2009). IBMs often follow a so-called pattern-oriented modeling (POM) approach (Grimm, Revilla et al. 2005) using patterns observed in real systems to compare them to the output of simplified models to justify the simplifications. IBMs in microbiology have been used to investigate phenomena such as biofilm and colony formation (Kreft, Booth et al. 1998; Lardon, Merkey et al. 2011), phytoplankton dynamics in aquatic systems (Woods and Onken 1982; Woods 2005; Hellweger 2008), the lag phase of bacterial growth (Dens, Bernaerts et al. 2005), and evolution of microorganisms (Lenski, Ofria et al. 2003; Chow, Wilke et al. 2004). More examples of microbial IBMs were recently summarized in Hellweger and Bucci (2009).

A microbial environment for which a spatially explicit IBM approach is still lacking is the plant leaf surface, or phyllosphere (Ruinen 1961). The phyllosphere is an extreme environment: e.g. widely fluctuating temperatures (Lindow and Brandl...
2003), highly localized and variable amounts of nutrition, e.g. fructose, glucose, and sucrose (Leveau and Lindow 2001; Remus-Emsermann and Leveau 2010; Remus-Emsermann, de Oliveira et al. 2011), and erratic periods of water stress (Beattie 2011). Leaf surfaces also feature a rich variation in topography full of grooves, veins, trichomes, stomata and glandular cells at the microscopic level. Despite the appearance of a seemingly stressful environment, the phyllosphere is a significant habitat for microbes of all kinds. For example, leaves worldwide have been estimated to cumulatively carry up to $10^{26}$ bacterial cells (Morris and Kinkel 2002).

In a previous experimental phyllosphere study (Remus-Emsermann and Leveau 2010) we created a bioreporter that tracks individual reproductive success of bacteria based on dilution of green fluorescent protein (GFP) during cell division and growth. Using this bioreporter, we have generated a database of patterns that show individual variation in bacterial reproduction on bean leaves under different conditions (this thesis chapter 3 and 4). So far, we have interpreted these patterns using a logistic-growth model that assumed a maximum number of bacteria that the leaf is able to sustain, which we called carrying capacity. However, this logistic model did not take into account death of bacterial individuals or resource replenishment in habitable sites in the phyllosphere. There is ample evidence that both these processes occur: living plant leaves leak sufficient amounts of substances that sustain microbial life in the phyllosphere (Tukey and Mecklenburg 1964; Leveau and Lindow 2001; Remus-Emsermann, de Oliveira et al. 2011), and bacteria do die (Monier and Lindow 2005; Monier and Lindow 2005). Here, we expanded the logistic growth model into a model that includes a replenishment component and bacterial death. By also including cell-to-cell heterogeneity and stochasticity of cell division, resource uptake, and movement we essentially translated the logistic model into one that is individual based and made it spatially explicit using the modelling program NetLogo (Wilensky 1999). Named “Agent-based Simulation of Microbial Phyllosphere colonization”, or short ASiMoPh, it is intended to facilitate the evaluation and interpretation of bacterial growth on the leaf surface to better understand the relationships between population increase, individual reproductive success, and environmental heterogeneity.

**Methods**

*Study organism and pattern-generating experiment*

The bacterium *Erwinia herbicola* 299R (a.k.a. *Pantoea agglomerans*) is a model phyllosphere colonizer (Brandl and Lindow 1998; Leveau and Lindow 2001; Remus-Emsermann and Leveau 2010; van der Wal and Leveau 2011), originally isolated from a pear tree leaf (Brandl and Lindow 1996). Patterns of reproductive suc-
cess were obtained from experiments described in chapters 3 and 4 of this thesis, and were generated to document the individual experience of bacteria under various conditions of competition with other bacteria.

**Model description**

We used the Design concepts and Details (ODD) format proposed by Grimm et al. for the description of the model (Grimm, Berger et al. 2010). A working code example of ASiMoPh for NetLogo is available in the Supporting information.

**Purpose**

ASiMoPh is intended to facilitate the evaluation and interpretation of bacterial growth on the leaf surface to better understand the relationships between population increase, individual reproductive success, and environmental heterogeneity.

**State variables and Scales**

Individual bacteria are represented as agents in NetLogo. Another type of agents are patches, representing parts of the leaf environment. Each bacterial agent possesses

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<td>Abbreviation/abbreviation initial value</td>
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<td>Energy storage</td>
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<th>Patch properties</th>
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<td>Occupied space in simulation</td>
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<td>Energy storage</td>
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<td>Energy replenishment</td>
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Abbreviation/abbreviation initial value

Mean value

Unit
the following attributes: an internal energy pool $E_a$; an initial pool of energy $E_i$; an ability to take up energy from its environment with rate $R_u$ per model time step; a threshold of energy $E_d$, at which reproduction of the agent is triggered, followed by a redistribution of the two newly created agents with a dispersal distance $D$; bacteria metabolize a portion of their energy during every model time step to maintain their metabolism with the respiration-rate $R_m$; agents record their generation $G$. It is possible to release two sets of agents side by side or with a time interval into the simulation.

Patch agents have the following attributes: initial energy $E_{pi}$, current energy $E_p$, and energy replenishment rate $R_r$. The size of an individual patch corresponds to 2500 $\mu$m$^2$, which is about the average size of a fructose-conductive site in the poplar phyllosphere (Remus-Emsermann, de Oliveira et al. 2011). We used this area estimate since it is the only one available for fructose-conductive sites in the phyllosphere and the only estimate for patch size. All agent properties are summarized in table 1.

Due to computational constraints the simulation did not cover a whole leaf, but was restricted to a representative sample of the leaf environment measuring 1.05 mm x 1.05 mm or 1.1025 mm$^2$, which represents approximately 0.02% of an average bean cotyledon. One model time step corresponds to the shortest average generation time

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**Algorithm 1:**

**Model initialization step:**
Randomized distribution of patches and agents; $E_a = E_i$; $E_p = E_{pi}$

**Repetitive process list:**
1) if $E_p^{\text{(current)}} > 0$, bacterial agents take up energy from the patch they are in, so that $E_p^{\text{(new)}} = E_p^{\text{(current)}} - R_u$ and $E_a^{\text{(new)}} = E_a^{\text{(current)}} + R_u$, if $E_p^{\text{(current)}} < R_u$. $R_u$ is reduced so that $R_u = E_p^{\text{(current)}}$

2) bacterial agents divide and daughter cells are relocated with distance $D$ if $E_a > E_d$; the generation of daughter agents increases by one; $G = G + 1$

3) bacterial agents metabolize $R_m$ energy for maintenance of metabolism; $E_a = E_a - R_m$; bacterial agents die if $E_a \leq 0$

4) patches that contain less than their initial-energy replenish energy until $E_p = E_{pi}$; if $E_p < E_{pi}$; $E_p = E_p + E_r$
of the bacterial strain *Erwinia herbicola* 299R on bean leaves, which is approximately 1 hour 44 min (Leveau and Lindow 2001). Most experimental data derived from experiments that lasted 24 hours, which correspond to 14 time steps.

**Process overview and scheduling**

In each time step, bacterial agents take up energy, divide, redistribute, consume energy, and survive or die (see algorithm 1). For every process step bacterial agents are randomized in a list that is processed concurrently. Patch agents replenish energy, simulating the natural process of leaching.

**Design concepts**

**Emergence:**
Bacterial agents form micro-colonies within patches. The frequency distribution of bacterial numbers in patches is recorded and the current generation of all individual agents is recorded.

**Prediction:**
ASiMoPh predicts bacterial population sizes from simulated individual cell reproduction which is a function of the variation in available energy and number of agents in patches. By separating immigration events of two strains in time, the impact of primary colonizers on the subsequent growth of secondary colonizers can be investigated.

**Interaction:**
Interaction between bacterial agents is indirect and based on competition for energy that is heterogeneously available in patches. Individual patches are not connected, e.g., no lateral diffusion of energy is possible. Due to the redistribution of daughter cells after division, bacterial agents can reach adjacent patches.

**Stochasticity:**
Stochastic processes in ASiMoPh include the following: (I) the initial positions of bacterial agents are randomized within the model grid. (II) \( E_i \) and \( E_{pi} \) are chosen randomly from a normal distribution with a standard deviation of 25% of the given average. \( E_i \) is on average 1. \( E_{pi} \) averages are 2, 48, or 1500 (see initialization) (III) The bacterial agent attributes \( R_u, E_d, D \), are chosen randomly around an average of 1, 2, and 0, respectively, with standard deviations of 0.01, 0.25, 0.015 every time step. (IV) For every process step that queries agents, agents are queried in a randomized
The random number generator used in the simulation was based on Mersenne twister (Matsumoto and Nishimura 1998; Wilensky 1999). For the comparison of simulations the random number seed can be preset.

**Collectives:**
Up to two different breeds of agents can be released into the world representing different bacterial strains with different properties in terms of energy uptake, division threshold, energy consumption and movement after division.

**Observation:**
The recorded model output consists of the distribution of reproductive success within agent groups, frequency distribution of agent counts per patch, total growth and death of each breed in the simulation, and the distribution of the available energy in patches.

**Initialization:**
Initialization of ASiMoPh is user-controlled. As an exemplary study we setup a leaf-portion that contained three differential types of patches and a total of 441 patches. Patch kind one contained $E_p = 2$ and covered 48.5 % of the leaf, patch kind two contained $E_p = 48$ and covered 48.5 %, and patch kind three carried $E_p = 1500$ and covered 3 % of the leaf. These values were extracted from a logistic model presented in a previous study (Chapter 3). In this study the values represented local carrying

![Figure 1: Computational domain, two-dimensional grid representation of a heterogeneous surface consisting of 21 x 21 patches, approximately 1.05 mm x 1.05 mm, including 48.5% patches with low $E_p$, 48.5% of patches with intermediate $E_p$, and 3% patches with high $E_p$.](image)
capacities without considering replenishment so that one unit of energy sustained 
one cell. In the here presented study one unit of energy relates to the amount of energy needed for one cell division. Patch kind two and three replenished energy during each time step, patch kind two 5 percent of the initial value and patch kind three 1 percent of the initial value. These choices were made to obtain a total amount of energy in the system sufficient to sustain 3300 agents, but since movement in the model is limited, there will be unreached or unreachable patches if patches are not completely colonized. Related to the average size of a bean leaf the total number of agents would be 1.5×10^7, which is in good agreement with CFUs often reported in the phyllosphere (Lindow and Brandl 2003). Agents enter the simulation with an average $E_i$ of 1, an average reproductive threshold of $E_d$ 1, an average energy uptake $E_u$ of 1, an energy consumption $E_m$ of 20 percent of the energy uptake, and an average dispersal distance of $D = 0$ with a standard deviation of 0.015.

**Simulation experiments:**

In study 1, one set of agents was released in different densities into the simulation, the average densities ranged from 2^2 to 2^5 per patch or 0.25 to 32 agents per patch. In study 2, two sets of agents were released subsequently with a difference of 14 time steps, or 24 hours, into the simulation, densities of both sets ranged from 2^4 to 2^5, or 0.0625 to 32 average agents per patch. The reproductive success of the individual agents was recorded and compared to laboratory experiments.

![Figure 2: A – D Representative results of ASiMoPh after 24 simulated hours. Simulations were initiated with different numbers of agents. Average numbers of agents per patch were 0.25 (A), 2 (B), 4 (C), and 16 (D), respectively. Error bars represent the standard deviation of three replicate simulations. E - H) Data obtained in the lab from CUSPER cells that were inoculated on Phaseolus vulgaris cotyledons that were recovered after 24 hours. CUSPER-cells were inoculated in rising CFU densities per mL: 10^5, 10^6, 10^7, 10^8 (this thesis chapter 3). Error bars represent the standard deviation of two leaf replicates.](image-url)
Submodels:

The computational domain: The computational domain is an evenly spaced rectangle grid of 21 x 21 patches (Figure 1).

Results

Study 1

Simulations conducted with different densities of one breed yielded reproducible patterns of reproductive success (RS) after 14 model time steps. Low average densities of agents per patch, e.g. less than 2 initial agents per patch (figure 2A), resulted in a relative contribution of the initial population of agents that contributed to the observed population that were split in to subpopulation of approximately equal size, one with low final RS of 1-2 and population that was very successful with an RS >5. Intermediate densities of agents per patch, i.e. 2 and 4 initial agents per patch, produced initial populations that were approximately equal distributed over the whole range of RS (figure 2 B and C). High initial densities of agents per patch, i.e. 16 agents per patch, produced a final population in which only 3% of the initial popula-
tion were very successful with a RS of >5. The rest of the initial population created a pattern resembling a right-tailed or lognormal distribution (figure 2D). The first peak was at a RS of 1 while the second peak was at a RS of >5. More than 50% of all initial agents experienced less than 3 doublings while the second peak encompassed only 2 – 4% of all initial agents doubled more than 5 times.

Study 2
The second experiment was conducted by releasing a set of agents 14 time steps prior to a second set of agents. The RS of the secondarily released set of agents was recorded and analyzed after an additional 14 time steps. Releasing low amounts of primary colonizing agents, i.e. ≤ 0.5 agents on average per patch, did not have large impacts on pattern observed at the single agent level (figure 3A). The initial population was separated into two subpopulations, depending on the site of immigration. One successful population that achieved RS of 5 and more and one less successful population that encompassed about 70% of the total initial population and divided once on average. With rising numbers of preinoculated agents, secondary colonizers became less successful. Especially colonizers that were exceptionally successful decrease until they vanish (figure 3B, C and D). With the highest amount of preinoculated agents only about 0.7% of the initial agents were able to divide twice (figure 3D).

Discussion:
We successfully translated a logistic growth model for bacterial phyllosphere colonization into a spatial-explicit, agent-based model. In study one the impact of inoculation density on the RS of the colonizers in silico was investigated. Using parameters extracted from a previously established model, the results that were obtained in silico are in good agreement with patterns found in vitro (figure 2E-H). The spatially explicit approach used in this study here could recreate results that were obtained with a logistic growth model approach (this thesis chapter 3). This emphasizes the importance of space in this system and supports the notion that the Phaseolus vulgaris phyllosphere offers generally at least three different qualities of habitats, in different quantities, to bacterial colonizers. One habitat that does not permit many doublings of the colonizers even if they reach a resource source on their own and do not have to share resources, one habitat that permits more than one doubling for individual colonizers but not if several colonizers have to share the resources, and finally one or more kinds of habitats that allow growth even if large numbers of colonizers have to share resources. In vitro studies suggested that patch kinds one and two represent approximately equal and large portions of the
phyllosphere while patch kind three covers only small portions (this thesis chapter 3). Due to limitations of the in vitro dataset to an observation of 5 doublings it is uncertain if these latter patches have to be distinguished into even more patches that offer differential amounts of resources to colonizers on average.

The logistic model did not take individual agents into account instead reproductive success of all “bacteria” per site was averaged. ASiMoPh showed that it is possible to achieve the same results with stochastic differences between individual agents. ASiMoPh additionally provides an excellent framework for other various factors that are potentially of importance during phyllosphere colonization, e.g. movement within a leaf, changing environmental surroundings, like desiccation and environmental engineering of bacterial agents, antibiosis and many more. Although ASiMoPh is not restricted to leaf colonization, it can be used as a basis for simulations of many given heterogeneous environments, like soil, if simplified to a 2-dimensional domain.

**Acknowledgements**

Funding was provided by the Netherlands Organisation of Scientific Research (NWO) in the form of a personal VIDI grant to JHJL.
Draft sequence and partial genome annotation of the phyllosphere model bacterium Erwinia herbicola strain 299R

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Introduction
Erwinia herbicola 299R (Eh229R; syn. Pantoea agglomerans, Enterobacter agglomerans) is a spontaneous rifampicin-resistant derivative (Brandl and Lindow 1996) of Erwinia herbicola 299, a plant epiphytic bacterium that was isolated from pear leaves (‘Bartlett’ Pyrus communis) in April 1985 (Lindow, Desurmont et al. 1998). Since, it has been employed as a model strain for the study of commensal, bacterial colonizers of the phyllosphere. For example, Eh299R was used to assess how bacterial production of indole-3-acetic acid (IAA), contributes to phyllosphere fitness and to the formation of russets on fruit (Brandl, Clark et al. 1996; Brandl and Lindow 1996; Brandl and Lindow 1997; Brandl and Lindow 1998; Lindow, Desurmont et al. 1998; Manulis, Haviv-Chesner et al. 1998; Brandl, Quinones et al. 2001). Eh299R has also been used extensively in conjunction with fluorescent-protein based bioreporter technology to unravel and quantify the bacterial colonization and experience of plant surfaces, which revealed among other things, the uneven distribution of bacterial aggregates and nutrient resources such as sugars on leaves (Jaeger, Lindow et al. 1999; Leveau and Lindow 2001; Miller, Brandl et al. 2001; Monier and Lindow 2005; Monier and Lindow 2005; Remus-Emsermann and Leveau 2010; Remus-Emsermann, de Oliveira et al. 2011).

Rationale
The availability of the full genome sequence of Eh299R will allow deeper understanding of its ability to successfully colonize leaf surfaces as well as omic-type investigations such as transcriptional profiling during leaf colonization and comparative genomics with other plant-associated relatives for which genome sequences are available, i.e. Erwinia amylovora, Erwinia pyrifoliae, Pantoea ananatis, and Pantoea vagans (De Maayer, Chan et al. 2010; Smits, Jaenicke et al. 2010; Smits, Rezzonico et al. 2010; Smits, Rezzonico et al. 2010). Some of these are plant pathogens, others are not, or are in fact beneficial as biocontrol agents. The inclusion of a commensal in this group is likely to shed light on what genes or gene clusters define the pathogen or beneficial phenotype. Furthermore, given that the classification of the
species *Erwinia herbicola* is still ambiguous (Rezzonico, Smits et al. 2009), a full genome sequence may further enlighten this issue and help distinguish and reclassify bacterial strains that are currently grouped under the name *Erwinia herbicola*.

**Results**

Eh299R genomic DNA of an overnight culture, isolated using the Qiagen midi kit genomic DNA isolation (Venlo, The Netherlands), was sequenced using the Illumina platform technology by BaseClear (Leiden, The Netherlands) resulting in ~12.9 million paired-end reads with an average length of 48 bp and an average insert size of 336 bp. Quality filtered reads were assembled de novo with different tools: Velvet 1.1.02 (Zerbino and Birney 2008; Zerbino, McEwen et al. 2009) using the full sequence set, and SeqMan NGen 3.04 build 10 (DNASTAR, Madison WI, USA) using a subset of 5 million reads. A comparison of both assembly results is given in Table 1. Annotation was performed on all contigs greater than 100 bp using the RAST prokaryotic genome annotation server (Aziz, Bartels et al. 2008). Depending on the assembly tool that was used, the genome was predicted to posses between 4303 and 4436 coding sequences of which 77 – 81 % were predicted to be non-hypothetical. RAST predicted 68 to 80 RNA coding sequences to be present in the genome. The

<table>
<thead>
<tr>
<th></th>
<th>Velvet 1.1.02</th>
<th>Lasergene NGen 3.06 build 10</th>
<th>SeqMan NGen 3.06 build 10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contigs &gt; 100 (bp)</td>
<td>314</td>
<td>159</td>
<td>159</td>
</tr>
<tr>
<td>N50 (kb)</td>
<td>35</td>
<td>59</td>
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<tr>
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<td>92</td>
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<tr>
<td>Scaffold &gt; 100 (bp)</td>
<td>55</td>
<td>92</td>
<td>92</td>
</tr>
<tr>
<td>Scaffold average length</td>
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<td>29</td>
<td>29</td>
</tr>
<tr>
<td>(kb)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Scaffold max length (kb)</td>
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<td>265</td>
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</tr>
<tr>
<td>Total sequence length</td>
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<td>4.77</td>
<td>4.77</td>
</tr>
<tr>
<td>(Mb)</td>
<td></td>
<td></td>
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<tr>
<td>Coding sequences predicted by RAST</td>
<td>4239</td>
<td>4504</td>
<td></td>
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<tr>
<td>Non hypothetical genes predicted by RAST</td>
<td>3275</td>
<td>3456</td>
<td></td>
</tr>
<tr>
<td>RNA CDS predicted by RAST</td>
<td>64</td>
<td>68</td>
<td></td>
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</tbody>
</table>
The software r2cat, which allows comparison of genomes that are somewhat related (Husemann and Stoye 2010) was used as a tool to orient Eh299R contigs, by using the recently published genome of the related bacterial strain Pantoea vagans C9-1 as a reference (Smits, Rezzonico et al. 2010). r2cat was able to align 61 out of 112 scaffolds assembled by Velvet (Figure 1 A). The other 51 contigs which accounted for 44 Kb of genetic information could not be aligned to the C9-1 genome. r2cat was able to align 145 out of 159 contigs assembled by SeqMan NGen (Figure 1 B). 14 contigs which account for 157 kb genetic information could not be aligned to the C9-1 sequence. Sequences that r2cat was not able to align against Pantoea vagans C9-1 may represent genetic information that is unique to Eh299R.

Future work on the Eh299R genome will include the filling-in of sequence gaps with either traditional Sanger sequencing or next generation sequencing using larger insert sizes to allow better resolution of scaffolds and more in-depth annotation.

Figure 1 r2cat dot-plot of two Eh299R genome assemblies (Velvet, panel A; SeqMan NGen, panel B) versus the reference genome of Pantoea vagans C9-1. Queried contigs and scaffolds were ordered against the reference using the automatic order function of r2cat. Grey lines represent scaffold and contig termini.

Acknowledgements
Funding was provided by the Netherlands Organisation of Scientific Research (NWO) in the form of a personal VIDI grant to JHJL.
General discussion and synthesis

The notion of bacterial individuality, although around for several decades (Hughes 1956; Spudich and Koshland Jr 1976), was only recently revived in the scientific literature (Avery 2006; Davidson and Surette 2008; Zengler 2009; Müller and Nebe-von-Caron 2010). In studying bacterial individuality, one works under the premise that bacteria are individuals, and should be treated as such, as opposed to only as populations, as is usually the case in classical microbiology. This thesis has focused on the development of a framework that allows bacterial individuality to be considered as a more quantifiable concept in ecology. To date, most studies examining bacterial individuality do so from the perspective of cellular processes e.g. differences in gene regulation, mRNA, or protein stochasticity (Shapiro 2000; Avery 2006; Smits, Kuipers et al. 2006; Musat, Halm et al. 2008). In contrast to these studies, my aim in this thesis has been to focus on bacterial individuality by regarding bacteria as spatial individuals: individuals that occupy a specific place in time and space and make use of, and influence, their local environment. By using the experimentally amendable system of the leaf surface, or phyllosphere, I seek to offer a case study in a natural, yet relatively low-complexity (compared to other habitats, e.g. soil) environment. I used colonization experiments in the phyllosphere to develop a proof of concept for bacterial individuality in ecology.

The phyllosphere is not only interesting as an ecological case study, but many questions concerning the colonization of microbes in the phyllosphere are still unanswered. This thesis has addressed the following specific issues (1) the development and characterization of a bacterial bioreporter for reproductive success (RS) called CUSPER, (2) a (re)-definition of the term carrying capacity in the phyllosphere to a sum of local carrying capacities, (3) the impact of pre-emptive competition on secondary colonizers at the single-cell level, (4) the analysis of spatially variable local flux of fructose across leaf cuticles, (5) the creation of an individual-based model that aims to explain differences in individual reproductive success as a function of environmental heterogeneity and local competition, (6) and the draft assembly and preliminary genome annotation of the phyllosphere model strain *Erwinia herbicola* 299R.

Reproductive success as a measurement of colonization and environmental heterogeneity

The main tool that was established for this thesis is a bacterial bioreporter that is able to report on the RS of single-cells. Called CUSPER, this bioreporter is based on the dilution of stable green fluorescent protein (GFP) from growing cells of *Erwinia*
herbicola 299R that no longer produce GFP de novo. To achieve this functionality, cells were filled with GFP by induction with Isopropyl β-D-1-thiogalactopyranoside (IPTG) prior to the experiment and introduced into an IPTG-free environment. CUSPER thereby enabled me to determine the reproduction of single cells in artificial environments such as a shaken liquid culture (see chapter 2) or agar plates (data not shown) by analyzing GFP content of individual bacterial cells and follow their growth for up to six doublings, at which point the GFP signal was too low to be differentiated from background fluorescence. CUSPER also showed differences in individual growth success of Erwinia herbicola 299R cells (Eh299R) on Phaseolus vulgaris leaves. The results in chapter 2 show that cells arriving in the phyllosphere experienced various degrees of local ‘habitability’. The results yielded a picture of a population where inoculated bacterial cells contributed unequally to the observed final population, e.g. we found that after 9 hours 29% of the initial cells contributed to only 5% of the observed population, whereas another 23% of the initial population contributed to 51% in the final population. We could show that growth curves based on colony forming unit (CFU) measurements and CUSPER single-cell data yielded similar results.

Describing the ecological paradigm of carrying capacity at scales that matter for bacteria

While many ecological theories have been well developed for higher organisms, proper translations of these theories to microbiological contexts are often lacking. For example, the term carrying capacity, as typically used in microbial ecology literature, refers to environmental scales that can hardly be perceived by bacterial individuals, e.g. the total surface of a plant leaf (Wilson and Lindow 1995; Nix, Burpee et al. 2009). For bacteria residing on a leaf it is most likely that their activity and fate is based on perception of and response to their immediate local environment (Leveau 2006). My goal has been to record variation in this environment, which encompasses many factors such as nutrient availability and exposure to stress and competitors, not through a description of the dimensions and quality of that environment, but by quantifying the bacterial response to this environment.

In chapter 2, I described that the phyllosphere exhibits a heterogeneity of patches with different levels of habitability, while in chapter 3, I showed that saturating the leaf with different densities of bacteria yielded information about the relative occurrence and quality of these habitable sites. More specifically, low inoculation densities revealed that 45-50% of evenly spread bacteria arriving on a leaf experienced an environment that did not allow them to divide more than once within 24 hours.
The remaining population experienced more than 5 doublings. Higher inoculation densities, on the other hand, revealed that 45-50% of the tested bean phyllosphere consists of sites of intermediate lognormal distributed carrying capacity, i.e. the sum of resources available that allow growth and stress factors that inhibit growth. Interestingly, even at very high inoculation densities, 2-3% of the population experienced more than 5 doublings within 24 hours, providing evidence for the existence of distinct sites in the phyllosphere that are highly conducive to growth. The more bacteria inoculated, the more predictable the resulting growth pattern, which is in line with the assumption that we saturated a heterogeneous environment to different degrees. These results were used to formulate a logistic growth model that we adapted from Kinkel et al. (Kinkel, Newton et al. 2002), and that was able to reproduce the experimental results. Thus, phyllosphere carrying capacity that can be observed by CFU measurements is a sum of local carrying capacities. These findings greatly improve our current picture of the ecology of the phyllosphere by adding quantitative knowledge of the distribution of site qualities and an estimation of the probability for successful colonization of the phyllosphere by bacteria.

Impact of secondary colonization in the phyllosphere on reproductive success

Chapter 4 sought to investigate the mechanism of preemptive exclusion by combining a preemptively inoculated red fluorescent strain, *Erwinia herbicola* 299R (pFRU97), with near-isogenic cells of CUSPER, i.e. *Erwinia herbicola* 299R::JBA28 (pCPP39).

One strategy that is available and sometimes used to prevent the onset of disease in agriculture is to preemptively inoculate crops with biocontrol agents (BCAs) prior to outbreaks of plant pathogenic organisms such as *Erwinia amylovora* (Stockwell, Johnson et al. 2010). The mechanisms by which BCAs control pathogens have typically been observed via top-down population-based measurements (Wilson and Lindow 1994; Braun-Kiewnick, Jacobsen et al. 2000; Nix, Burpee et al. 2009; Stockwell, Johnson et al. 2010; Xu, Salama et al. 2010). By making use of CUSPER, it was possible to evaluate the impact of preemptively inoculated bacteria on secondary colonizers using a bottom-up approach.

Colony forming unit data and single-cell data both showed that the success of secondary colonizers depended on the density of primary colonizers. By applying the previously formulated model for phyllosphere colonization (chapter 3), I showed how different timing and densities of preinoculated colonizers inhibited the growth of secondary colonizers to different degrees. These results provide insights into how biocontrol agents (BCAs) might impede pathogen establishment, thereby providing
crop protection. In cases of preemptive exclusion, biocontrol functionality is facilitated by an even distribution, in high numbers, and with a sufficient time advantage.

**Determining the local permeability of carbohydrates through a plant cuticle**

On plant leaf surfaces, carbohydrates more specifically photoassimilates such as fructose, glucose and sucrose, are the major source of energy and carbon for most phyllosphere colonizers (Mercier and Lindow 2000; Leveau and Lindow 2001). Being produced on the inside of the leaf during photosynthesis, they reach the leaf surface in a process called leaching (Tukey 1966).

One of the major gaps in our understanding of the phyllosphere environment, and modeling of this environment, involves the rate by which carbohydrates are locally released into the phyllosphere. Quantitative measurements of local sugar permeation are difficult (Leveau and Lindow 2001; Miller, Brandl et al. 2001) and permeation studies for carbohydrates have typically been restricted to bulk measurements of larger leaf areas (Tukey and Mecklenburg 1964; Mercier and Lindow 2000; van der Wal and Leveau 2011). By combining local semi-quantitative observations with area-dependent bulk measurements, I was able overcome this limitation and address in quantitative terms the lateral heterogeneity of carbohydrate diffusion across leaf cuticles.

For this, isolated, trichome- and stomata-free, leaf cuticles from poplar, *Populus x canescens*, were used. By combining gas chromatography, coupled with a flame ionization detector (GC-FID) and a GFP-based bacterial bioreporter, I was able to pinpoint the lateral differences in permeance of fructose and to estimate the permeability of these sites. GC-FID results revealed an average bulk permeance for fructose of $3.39 \times 10^{-9} \text{ m s}^{-1}$. Following this analysis, bioreporter cells were introduced on the same isolated cuticles. The analysis via epifluorescence microscopy revealed a patchy, non-random pattern of fructose-sensing bioreporters, corresponding to sites with higher fructose permeance. These sites were predominantly co-occurring with shed trichome sites. By relating the total area that these sites cover on the cuticle, the permeance of these sites for fructose appeared 270 times higher than the bulk permeance of the isolated cuticle.

Together with the studies described in chapters 2-4 that dealt with the heterogeneity of colonization success, the heterogeneity of available nutrients demonstrates, at least in part, that the differences in growth conduciveness between sites can be linked to differences in nutrient availability.
Modeling bacterial colonization in the phyllosphere

The data collected for chapter 2-5 were used as a foundation to create an agent-based model, called agent-based simulation of microbial phyllosphere colonization or simply “ASiMoPh”. The aim of ASiMoPh is to aid our understanding of the underlying mechanism of heterogeneity and how such habitat heterogeneity determines variability in the success of bacterial colonizers of the leaf environment. Another reason was that, even though the logistic growth model formulated in chapter 3 and 4 was able to recreate the patterns that were observed experimentally, it completely dismissed the possibility of death within the bacterial population on the leaf. It also did not take into account the possibility of resource replenishment. Both factors are crucial to capture the essence of carrying capacity, an equilibrium around which a population oscillates. Another reason to shift to a model that is rule-based and uses agents was to explicitly follow the concept of bacterial individuality in modeling. Furthermore, ASiMoPh is spatially explicit Other models for growth and BCA plant pathogen interaction are available, but they too function under the premise that the leaf environment is not spatially explicit (Kinkel, Newton et al. 2002; Xu, Salama et al. 2010).

The model was created in NetLogo (Wilensky 1999) and features agents representing bacteria and patches on which bacterial agents can feed. The model parameters include the uptake of nutrients, a threshold of energy that agents need to reach before they divide, a metabolism uptake that agents have to pay every time step and death of individual agents if they have consumed all energy. Patches possess energy and can replenish it during the simulation while agents are feeding on it. By adding both processes, it is possible for agents to reach an equilibrium between growth and death rate, or carrying capacity. Based on the observations of the average size of fructose permeable sites in a cuticle, the patch size in the model was determined to represent 2500 µm² (chapter 5). To keep computational demands feasible, 441 patches per simulation were assumed, which corresponds to about 0.02% of an average bean leaf. Thus far, ASiMoPh was able to recreate population patterns as were observed in chapter 3 and 4 and explain the results in an individual-based modeling setup.

Sequencing of the phyllosphere model bacterium Erwinia herbicola strain 299R

I successfully performed the first steps to sequence the genome of the phyllosphere model bacterium Erwinia herbicola strain 299R. A draft genome sequence was obtained and annotated. The availability of the full genome sequence will enable further research of mechanisms used by this strain to survive the harsh conditions in the phyllosphere. The species Erwinia herbi-
Cola is rather heterogeneous and contains strains that are opportunistic pathogens as well as plant commensals that are used as certified BCAs (Rezzonico, Smits et al. 2009; Smits, Rezzonico et al. 2010). With the availability of a genome that derives from a non-human pathogenic-, plant commensal strain, and together with other recently published sequences of related Pantoea and Erwinia strains (De Maayer, Chan et al. 2010; Smits, Jaenicke et al. 2010; Smits, Rezzonico et al. 2010), I hope to shed light into this issue and maybe help to reclassify strains collected under the current name.

**Perspectives for future lines of research**

One of the major limitations of the CUSPER bioreporter used during the larger part of this PhD project lies in the fact that the limit of detection is reached after 6 doubling even under the best possible experimental circumstances. By using the same principle of dilution but other means of detection and reporter molecules, it might be possible to lower this limit of detection significantly. One might consider using isotope-labeled molecules to incorporated them into cells and apply Nano secondary ion mass spectrometry (NanoSIMS) to detect the amount of label left after incubation (Li, Wu et al. 2008). This would greatly increase the time that a reporter working with a dilution principle is able to track population development and eliminates the necessity to modify the bacteria used for experimentation with plasmids and/or transposons.

To enhance the possibilities of multi-strain experiments, it will be useful to create differentially colored variants of the CUSPER bioreporter and to equip additional bacterial strains with CUSPER. Experiments with CUSPER-equipped pathogenic bacteria as secondary and co-colonizers together with biocontrol strains will yield interesting results that could be combined with enhanced versions of ASiMoPh (see below). Due to analytical problems, i.e. uneven background due to autofluorescence of the leaf, the analysis of CUSPER directly on the leaf surface has been difficult but promising advances have recently been made and will hopefully soon provide insightful results (Tecon, unpublished).

Bottom-up approaches using CUSPER bioreporter-equipped bacteria is expected to be of use in describing the experience of bacterial colonizers in other environments, too, especially in complex heterogeneous environments like soil or other, non-water saturated habitats.

The determination of the size of carbohydrate permeable sites in the phyllosphere was crucial for phyllosphere modeling. Since we now have the first estimates of the magnitude of permeance, future models, including future versions of ASiMoPh, can
employ these estimates to increase our knowledge of growth patterns in the phyllosphere.
Up until now, ASiMoPh has been able to predict the growth of bacteria in a heterogeneous environment that is only based on resource competition. In future versions, ASiMoPh could be enhanced with features that include interactions based on agent counts on patches to trigger behavior like production of antibiotic production and leaf bioengineering. This will help to predict the interaction with antibiotic-producing BCAs and leaf-bioengineering pathogens.
In the future, bacterial individuality will prove to be an important framework for microbial ecology. It will be possible to address fundamental ecological and evolutionary questions in bacterial systems, which will allow, in comparison to macroecological systems, rapid validation of hypotheses with virtually unlimited number of individuals in replicate experiments. The development of new techniques that allow or facilitate the analysis of bacterial individuals has made tremendous advances in the areas of fluorescence in situ hybridization (FISH), multicolored fluorescence bioreporters, as well as of flow cytometry and microscopical techniques. As these techniques further ease the analysis and interpretation of single-cells, they will become extremely useful for testing of general ecological and economical theories and models pertaining to individuality.
The Ecology of Bacterial Individuality


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NetLogo code:

```netlogo
breed [agents_set1]
breed [deadset1]
breed [agents_set2]
breed [deadset2]

agents_set1-own      [agents_set1_generation
agents_set1_internal_energy_storage]
agents_set2-own      [agents_set2_generation
agents_set2_internal_energy_storage]
patches-own          [patch-current-energy
patch-energy-T0]
deadset1-own         [agents_set1_generationdead
agents_set1_internal_energy_storage_dead]
deadset2-own         [agents_set2_generationdead
agents_set1_internal_energy_storage_dead]

to setup
  random-seed random-seed-number
  setup-world
  distribute-agents
  plots
  reset-ticks
end

to release2
  release-agents-set2
end

to setup-world
  ca

  let desert count patches
  ask n-of desert patches [set pcolor yellow]

; Add green oasis
  let n_oasis_green (G_oasis_green_cover / (patchsize_increase_green + 1)) * count patches / 100
  ask n-of n_oasis_green patches [set pcolor green]
  ask-concurrent patches
    [if pcolor = green [ask-concurrent n-of patchsize_increase_green neighbors4 [set pcolor green]]]

; Add blue oasis
  let n_oasis_blue (G_oasis_blue_cover / (patchsize_increase_blue + 1)) * count patches / 100
  ask n-of n_oasis_blue patches [set pcolor blue]
  ask-concurrent patches
```
[if pcolor = blue [ask-concurrent n-of patchsize_increase_blue_neighbors4 [set pcolor blue]]]

ask patches
[if pcolor = yellow [  
  let standarddeviation2 ln (1 + (desert_stdev_energy * desert_stdev_energy) / (desert_mean_energy * desert_mean_energy))  
  let desert-standarddeviation sqrt standarddeviation2  
  let desert-mean (ln desert_mean_energy) - (standarddeviation2 / 2)  
  set patch-energy-T0 exp random-normal desert-mean desert-standarddeviation]  
if pcolor = green [  
  let standarddeviation2 ln (1 + (oasis_green_energy_stdev * oasis_green_energy_stdev) / (oasis_green_mean_energy * oasis_green_mean_energy))  
  let green-standarddeviation sqrt standarddeviation2  
  let green-mean (ln oasis_green_mean_energy) - (standarddeviation2 / 2)  
  set patch-energy-T0 exp random-normal green-mean green-standarddeviation]  
if pcolor = blue [  
  set patch-energy-T0 random-normal oasis_blue_mean_energy oasis_blue_energy_stdev]  
set patch-current-energy patch-energy-T0]  
end
to distribute-agents  
  let n_agents_set1 441 * 2 ^ (#-agentset1)  
  create-agents_set1 n_agents_set1  
  [set size 0.35  
   set shape "circle"  
   setxy random-xcor random-ycor  
   set agents_set1_generation 0  
   set agents_set1_internal_energy_storage random-normal  
   breed1-start-energy ((breed1-start-energy / 100) * 25)  
   set color black]  
end
to release-agents-set2  
  let n_agents_set2 10 ^ (#-agentset2)  
  create-agents_set2 n_agents_set2  
  [set size 0.35  
   set shape "circle"  
   setxy random-xcor random-ycor  
   set agents_set2_generation 0  
   set agents_set2_internal_energy_storage random-normal  
   breed2-start-energy ((breed2-start-energy / 100) * 25)  
   set color black]  
end
set color blue]
end

to go
repeat total-cycles
[ agents-eat
agents-divide
agents-energy-consumption
replenish-patch-energy
plots
tick]
end
to agents-eat
ask turtles-on patches with [patch-current-energy > 0]
[ let energy_uptake 0
ifelse breed = agents_set1
[ set energy_uptake random-normal agents_set1_energy_uptake
((agents_set1_energy_uptake / 100))
if (patch-current-energy - energy_uptake) < 0 [set en-
ergy_uptake patch-current-energy]
set agents_set1_internal_energy_storage agents_set1_in-
ternal_energy_storage + energy_uptake
set patch-current-energy patch-current-energy - energy_uptake]
[ if breed = agents_set2
[ set energy_uptake random-normal agents_set2_energy_up-
take ((agents_set2_energy_uptake / 100))
if (patch-current-energy - energy_uptake) < 0 [set en-
ergy_uptake patch-current-energy]
set agents_set2_internal_energy_storage agents_set2_in-
ternal_energy_storage + energy_uptake
set patch-current-energy patch-current-energy - energy_uptake]]]
end
to agents-divide
ask-concurrent agents_set1 with [agents_set1_internal_en-
ergy_storage >= random-normal breed1-reproduction-threshold
((breed1-reproduction-threshold / 100) * 25)]
[hatch 2 [;set color color - 1
setxy xcor + random-normal 0 breed1-dispersal ycor +
random-normal 0 breed1-dispersal
set agents_set1_internal_energy_storage agents_set1_in-
ternal_energy_storage / 2
set agents_set1_generation agents_set1_generation + 1]
die]
ask-concurrent agents_set2 with [agents_set2_internal_en-
ergy_storage >= random-normal breed2-reproduction-threshold
((breed2-reproduction-threshold / 100) * 25)]
    [hatch 2 [
        setxy xcor + random-normal 0 ycor + random-normal 0 breed2-dispersal
        set agents_set2_internal_energy_storage agents_set2_internal_energy_storage / 2
        set agents_set2_generation agents_set2_generation + 1]
    die]
end
to agents-energy-consumption
  let protection_from_antibiotics 0
  let energy-consumption1 (( agents_set1_energy_uptake / 100) * agents_set1_energy_consumption )
  let energy-consumption2 (( agents_set1_energy_uptake / 100) * agents_set1_energy_consumption )
  ask agents_set1
    [set protection_from_antibiotics (count agents_set1-here + count deadset1-here) / 10 * protection_factor
      let antibiotic_factor ( count agents_set2-here ) * Antibiotic_strength - protection_from_antibiotics
      if-else antibiotic_factor > 0
        [set agents_set1_internal_energy_storage agents_set1_internal_energy_storage - energy-consumption1 + ( (agents_set1_energy_uptake / 100) * (agents_set1_energy_consumption + antibiotic_factor ))]
      [set agents_set1_internal_energy_storage agents_set1_internal_energy_storage - energy-consumption1]
    ]
  ask agents_set1 with [agents_set1_internal_energy_storage < 0]
    [let current_agents_set1_generation agents_set1_generation
      hatch-deadset1 1
      [set color red
       set size 0.35
       set shape "circle"
       set agents_set1_generationdead current_agents_set1_generation
       ht
     ]
    die]
  ask agents_set2
    [set agents_set2_internal_energy_storage (agents_set2_internal_energy_storage - energy-consumption2 )]
  ask agents_set2 with [agents_set2_internal_energy_storage <
[let current_agents_set2_generation agents_set2_generation hatch-deadset2 1

[ set color red
  set size 0.35
  set shape "circle"
  set agents_set2-generationdead current_agents_set2_generation
  ht
]
die]
end

to replenish-patch-energy
  let rep-energy 0
  let engineer1 0
  let engineer2 0
  ask-concurrent patches with [count turtles > 0]

  [ if count agents_set1-here >= Bioengineer1-threshold
    [set engineer1 Bioengineer1-ability]
    if count agents_set2-here >= Bioengineer2-threshold
      [set engineer2 Bioengineer2-ability]
      if pcolor = green
        [set rep-energy ((patch-energy-T0 / 100) * replenishing_rate_oasis_green) + ((patch-energy-T0 / 100) * engineer1) + ((patch-energy-T0 / 100) * engineer2)]
      if pcolor = blue
        [set rep-energy ((patch-energy-T0 / 100) * replenishing_rate_oasis_blue) + ((patch-energy-T0 / 100) * engineer1) + ((patch-energy-T0 / 100) * engineer2)]
      if pcolor = yellow
        [set rep-energy 0]
      if patch-current-energy < patch-energy-T0
        [ifelse patch-current-energy + rep-energy > patch-energy-T0
          [set patch-current-energy patch-energy-T0]
          [set patch-current-energy patch-current-energy + rep-energy]
        ]
      end
    ]
  ]
end

to plots
  set-current-plot "population1-histogram-current-tick" ; plot current ticks histogram breed 1

  set-current-plot-pen "alive"
  let listalive [agents_set1_generation] of agents_set1
  set-plot-pen-mode 0
histogram listalive

set-current-plot-pen "dead"
let listdead [agents_set1_generationdead] of deadset1
  set-plot-pen-mode 0
histogram listdead

set-current-plot-pen "all"
let listall sentence listalive listdead
  set-plot-pen-mode 0
histogram listall

set-current-plot "population2-histogram-current-tick"
set-current-plot-pen "alive"
let listalive2 [agents_set2_generation] of agents_set2
  set-plot-pen-mode 0
histogram listalive2

set-current-plot-pen "dead"
let listdead2 [agents_set2-generationdead] of deadset2
  set-plot-pen-mode 0
histogram listdead2

set-current-plot-pen "all"
let listall2 sentence listalive2 listdead2
  set-plot-pen-mode 0
histogram listall2

set-current-plot "population-breed1"
let cpp_name word "tick " (ticks + 1)
create-temporary-plot-pen cpp_name
  set-plot-pen-mode 0
  set-plot-pen-color ticks + 5
set listall sort listall
let n_b count agents_set1 + count deadset1
let counter-current-population-breed1 0
let step-current-population-breed1 100 / (n_b)
foreach listall
  [ plotxy counter-current-population-breed1 ?
    set counter-current-population-breed1 counter-current-population-breed1 + step-current-population-breed1]

set-current-plot "population-breed2"
let cpp_name2 word "tick " (ticks + 1)
create-temporary-plot-pen cpp_name2
  set-plot-pen-mode 0
  set-plot-pen-color ticks + 5
set listall2 sort listall2
let n_b2 count agents_set2 + count deadset2
let step-current-population-breed2 100 / (n_b2 + 1)
let counter-current-population-breed2 0
foreach listall2
  [ plotxy counter-current-population-breed2 ?
    set counter-current-population-breed2 counter-current-population-breed2 + step-current-population-breed2]

set-current-plot "current-population-pattern"
clear-plot
set-current-plot-pen "breed1"
set-plot-pen-mode 0
set-plot-pen-color black
let counter-current-population-pattern-breed1 0
foreach listall1
  [ plotxy counter-current-population-pattern-breed1 ?
    set counter-current-population-pattern-breed1 counter-current-population-pattern-breed1 + step-current-population-breed1]
set-current-plot-pen "breed2"
set-plot-pen-color red
let counter-current-population-pattern-breed2 0
foreach listall2
  [ plotxy counter-current-population-pattern-breed2 ?
    set counter-current-population-pattern-breed2 counter-current-population-pattern-breed2 + step-current-population-breed2]

set-current-plot "patch-energy"
clear-plot
set-plot-pen-interval 10
set-histogram-num-bars 50
histogram [patch-current-energy] of patches

set-current-plot "Agent count"
let tick-counter word "tick " (ticks + 1)
set-current-plot-pen "Br1 Alive"
set-plot-pen-mode 0
let count-alive count agents_set1
plot count-alive
set-current-plot-pen "Br1 Dead"
set-plot-pen-mode 0
let count-dead count deadset1
plot count-dead
set-current-plot-pen "Br1 Total"
set-plot-pen-mode 0
let count-total count-alive + count-dead
plot count-total
; breed 2 plots
set-current-plot-pen "Br2 Alive"
set-plot-pen-mode 0
let count-alive2 count agents_set2
plot count-alive2
set-current-plot-pen "Br2 Dead"
set-plot-pen-mode 0
let count-dead2 count deadset2
plot count-dead2
set-current-plot-pen "Br2 Total"
set-plot-pen-mode 0
let count-total2 count-alive2 + count-dead2
plot count-total2

; agents on patches
set-current-plot "agent-counts on patches"
clear-plot
set-current-plot-pen "agentset1"
let agents-on-patches [count agents_set1-here + count dead-set1-here] of patches with [count agents_set1 + count deadset1 > 0]
  histogram agents-on-patches
set-current-plot-pen "agentset2"
let agents-on-patches2 [count agents_set2-here + count dead-set2-here] of patches with [count agents_set2 + count deadset2 > 1]
  histogram agents-on-patches2
end
ASiMoPh example interface

Run Controls
- Setup World and distribute Breed 1
- Distribute Breed 2

Simulation - Output
- Exports all plots to the Model's root folder
- Energy in World

World setup
- Desert: mean energy
- Oasis: mean energy

Agent setup
- Breed one
  - # of individuals at T0 (1D-p)
  - Amount of energy at startup
  - Dispersal from master-agent position
  - Energy uptake per cycle
- Breed two
  - # of individuals at T0 (1D-p)
  - Amount of energy at startup
  - Dispersal from master-agent position
  - Energy uptake per cycle
Mitja Nandi Paul Remus-Emsermann was born on the 7th of December 1980 in Siegburg, Germany. After his high school degree in 2000 he spend one year of civil service working in the IT-support department of the Johanniter hospital in Bonn, Germany. In 2001 he started the study of biology at the Rheinische-Friedrich-Wilhelms University in Bonn. Early 2005 he started to work as a student assistant in the department of ecophysiology of plants under supervision of Prof. Lukas Schreiber and Dr. habil. Rochus Franke mainly working with Arabidopsis thaliana surface polymers such as Suberin and Cutin. From January 2006 till March 2006 Mitja took the opportunity to conduct an internship at Syngenta, Stein, Switzerland with the topic “movement of insecticides in soil” under the supervision of Dr. Anke Buchholz. In March 2007 he earned his diploma degree in biology (major subject ecophysiology of plants, minor subjects chemistry and microbiology) and successfully finished his thesis “Characterization of the Beta-Acyl-CoA-Synthase Gene At5g43760 in Arabidopsis thaliana” under supervision of Prof. Lukas Schreiber and Dr. habil. Rochus Franke.

Also during 2007, Mitja started his PhD-project “The Ecology of Bacterial Individuality” in the department microbial ecology at the Netherlands institute of ecology, NIOO-KNAW, in Wageningen, the Netherlands, initiated and supervised by Prof. Johan Leveau and co-supervision by Prof. George Kowalchuk. Since then he performed projects as guest researcher in the department of plant pathology at the university of California, Davis CA, USA and in the department of ecophysiology of plants at the Rheinische-Friedrich-Wilhelms University, Bonn, Germany. Mitja presented his research at many seminars and several national and international meetings and conferences. The results of his PhD-project are the presented in this booklet. In November 2011, he joined the group of Julia Vorholt at the ETH Zurich to focus further on plant-microbe and microbe-microbe interactions at the single-cell level.
The Ecology of Bacterial Individuality
First of all, I want to thank my supervisor and promoter Johan. Johan, you gave me the opportunity to start this project and gave me a great head start during the first year of the project. We certainly managed this project well enough even though we had a long distance relationship with nine hours of time difference for the last three years. You taught me many things, and formed me to the scientist I am today.

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All the others, and there is so many of you since the recent merge of our institutes more than I can remember, with most of you I had beautiful and entertaining lunch and coffee breaks. It was rare that I didn’t enjoy my days at the NIOO thanks to all you guys.

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thesis and of course for giving me the confidence to start a PhD-project in the first place. Also thanks to Sheron with whom I had important discussions and who gave me more than valuable advises during the time.

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Daniela, Frau, beste Freundin, Kleinkind, Kritikerin in einem. Ohne dich hätte ich nicht durchgehalten
Summary
Classical microbial ecology generally describes and quantifies bacteria at a population or community level, be it for example in a liquid culture, as a colony on an agar plate, or as found in environmental samples. Such an approach necessarily yields results that reflect population or community averages, e.g. an average enzyme activity, an average doubling rate, or an average experience of the environment. The latter is at odds with the growing realization that considerable environmental heterogeneity exists in physicochemical and biological properties at the micrometer-scale. By averaging the bacterial experience of such heterogeneous environments, potentially valuable information is lost about the behavior of the organisms under study and about the variation in that behavior. Recent technical advances have introduced new possibilities to interrogate individual bacteria for their perception of and response to their local environment.

In this thesis, we used the phyllosphere, or leaf surface, as a model bacterial environment to address the question if and how individual cells that colonize this habitat are impacted by micro-scale environmental heterogeneity and how these impacts help to shape bacterial populations. To this end, a novel bioreporter was introduced that enabled analysis of single-cell reproductive success of individual bacteria. This new bioreporter was termed CUSPER (reproductive success = repsuc read backward) and is based on the dilution of green fluorescent protein (GFP) from growing cells of the natural leaf-colonizing bacterial strain *Erwinia herbicola* 299R.

By applying CUSPER cells to bean leaf surfaces, it was revealed that individual immigrants differed in their ability to produce offspring and contributed differently to the final observed populations. This was interpreted as an imposed effect of the environmental heterogeneity on the immigrating population, suggesting that the phyllosphere represents a patchy environment that offers various degrees of habitability to members of an immigrant population.

Subsequently, CUSPER was used to quantify the impact of different leaf inoculation densities on the reproductive success of cells within a population of bacterial immigrants. Some of the chosen densities were deliberately in excess of the carrying capacity of the bean leaves under study. This approach made it possible to show that even under circumstances in which immigrants arrive at numbers higher than the leaf can sustain, some cells were still able to reproduce. This finding suggested that carrying capacity has to be understood as the sum of many local carrying capacities.

A complementary modeling approach revealed that experimental observations can be explained by the existence of three types of microenvironments. Two of these environments represented approximately equal parts of the bean leaf surfaces under
study and offered low or intermediate reproductive success. A third class of micro-environment made up only 2% of the environment and offered high reproductive success.

A subsequent study dealt with the effect of pre-colonization of leaves on individual secondary colonizers. This is highly relevant in the context of preemptive colonization, a biocontrol approach that is often used to prevent the growth of plant-pathogenic bacteria. The study revealed that, even at high densities of precolonization, individual secondary colonizers were able to divide often enough to create as many as eight offspring, while the corresponding total population showed no net increase. Much of the heterogeneity of the leaf surface is due to local differences in nutrients that leak from the leaf interior to the surface. To obtain estimates for the rate at which this process takes place at the micrometer scale, we used an integrated approach of fructose-bioreporting bacteria and gas-chromatography, coupled to flame ionization detection, to determine the local permeability of isolated poplar leaf cuticles for carbohydrates. The average permeability of the poplar cuticles for fructose was estimated to be $3.39 \times 10^9$ m s$^{-1}$. Microscopic analysis of cuticles that were inoculated with bioreporting bacteria showed a non-random distribution of fructose-reporting bacteria. By relating the area covered by fructose-reporting bacteria with the measured average permeability, it could be shown that the permeability of poplar cuticles is locally 270 times higher than the rest of the cuticle. These findings gave an indication that resource partitioning might play an important role in the explanation of different reproductive success rates of bacterial immigrants to the phyllosphere.

In an attempt to combine the findings and models established in the previous parts of the thesis, a spatially explicit, agent-based modeling approach for bacterial colonization of the phyllosphere was formulated. The model, coined ASiMoPh for Agent-based Simulation of Microbial Phyllosphere colonization, was able to simulate the patterns observed in the previous studies using a simple set of rules for reproduction and nutrient uptake of bacterial agents. The model consisted of a two dimensional world that offered immigrant agents three different classes of microenvironments featuring different qualities in initial resource availability and resource replenishment.

The final study in this thesis presents the preliminary results of the sequencing of the strain *Erwinia herbicola* 299R. With the availability of the genomic information of this model phyllosphere organism, several new approaches will be possible, for example expression studies and comparative analysis of the genomes and functions between other phyllosphere strains to address the question: “What makes a successful phyllosphere bacterium?”
Taken together, the studies presented in this thesis demonstrated that single-cell observations yielded important information of the heterogeneity that shapes bacterial populations in the phyllosphere that were thus far overlooked. The CUSPER bioreporter offers an excellent tool for future studies investigating bacterial individuality not only in the phyllosphere, but also for many other environments. Important first steps were taken in this thesis to build a conceptual framework for bacterial individuality in natural environments that may serve as a foundation for future research in individual-based microbial ecology.
Samenvatting

In de microbiologie en microbiële ecologie wordt er over het algemeen aan bacteriën gemeten als populaties of gemeenschappen, bijvoorbeeld in een vloeibare cultuur, als een kolonie op een agarplaat of in hun natuurlijke leefomgeving. Dit heeft er toe geleid dat onze kennis van bacteriën in deze populaties en gemeenschappen veelal gebaseerd is op gemiddelden, bijvoorbeeld een gemiddelde enzymactiviteit of een gemiddelde groeisnelheid. Een groot nadeel van deze benadering is dat informatie verloren gaat over de eventuele variatie in het gedrag van individuele bacteriën binnen een populatie of gemeenschap. Er is een groeiend besef dat deze informatie belangrijk is voor een integraal inzicht in hoe bacteriële populaties en gemeenschappen tot stand komen en functioneren. Recente technische ontwikkelingen hebben het mogelijk gemaakt om onderzoek te doen aan hoe individuele bacteriën de fysisch-chemische en biologische kenmerken waarnemen van hun lokale leefomgeving. Het wordt steeds duidelijker dat op micrometerschaal variaties in deze kenmerken eerder regel dan uitzondering zijn en een belangrijke rol spelen in hoe individuele bacteriën reageren, overleven en vorm geven aan een populatie of gemeenschap.

In dit proefschrift werd gebruikt gemaakt van de fyllosfeer, oftewel het planten-bladoppervlak, als een natuurlijke, microscopisch heterogene modelleefomgeving. Binnen deze fyllosfeer werd onderzocht of en hoe het lot van individuele bacteriën verschilt van dat van de gemiddelde bacterie. Daartoe werd een nieuwe zogenaamde bacteriële bioreporter ontwikkeld die het mogelijk maakt het reproductieve succes van individuele bacteriën te kwantificeren. Deze bioreporter werd CUSPER genoemd en is gebaseerd op het uitverdunnen van groen fluorescerend eiwit (GFP) in reproducerende cellen van de bacterie Erwinia herbicola 299R, een natuurlijke kolonist van het bladoppervlak.

Toepassing van CUSPER op het oppervlak van een bonenblad liet duidelijke verschillen zien in hoeveel nageslacht individuele bacteriën produceerden en dus welke bijdrage zij leverden aan de totale bacteriële populatie op het blad. We leidden hieruit af dat de fyllosfeer een fragmentarische omgeving is die bacteriën verschillende mates van bewoonbaarheid en realisatie van nageslacht biedt.

Met CUSPER werd ook getest hoe oplopende bacteriële inoculatiedichtheden invloed hebben op het reproductieve succes van individuele cellen binnen een populatie van immigranten op het bladoppervlak. Deze aanpak gaf duidelijk aan dat zelfs wanneer het aantal bacteriële immigranten groter was dan wat het blad aankon (het zogenaamde bacteriële draagvermogen van het blad), sommige cellen nog steeds in staat waren nageslacht te produceren. Dit resultaat duidt er op dat het bacteriële draagvermogen van een blad moet worden gezien als de som van vele lokale...
draagvermogens. De experimentele waarnemingen konden worden nagebootst met de hulp van een mathematisch model, maar alleen indien aangenomen werd dat er drie soorten lokale leefomgeving op het blad voorkwamen, dat wil zeggen plekken met een relatief lage, medium of hoge waarschijnlijkheid dat bacteriën succesvol zijn in het produceren van nageslacht. Een bijkomende aannames was dat er veel meer plekken met lage en middelbare waarschijnlijkheid op het blad voorkwamen dan plekken met hoge waarschijnlijkheid.

Ook werd onderzocht wat het effect was van pre-kolonisatie van bladeren op het invasiesucces van secundaire immigranten. Dit is relevant in het kader van de zogenaamde biocontrole, waarbij de groei van plant-pathogene bacteriën, dat wil zeggen bacteriële ziekteverwekkers op planten, wordt voorkomen door het bladoppervlak te verzadigen met voor de plant onschuldige bacteriën waardoor het bladoppervlak minder mogelijkheden biedt voor het produceren van nageslacht door een pathogene bacterie. De studie toonde echter aan dat zelfs bij hoge dichtheden van pre-kolonisatie door zulke onschuldige bacteriën, een aantal individuele secundaire immigranten wel degelijk in staat was te reproduceren.

Een deel van de waargenomen verschillen in het reproductieve succes van bacteriën op het bladoppervlak is waarschijnlijk te wijten aan lokale verschillen in voedingsstoffen, zoals plantensuikers die vanuit het blad naar het oppervlak lekken. Om inzicht te krijgen in de variatie tussen verschillende delen van het bladoppervlak werd gekozen voor een geïntegreerde aanpak, een combinatie van bacteriële bio-reporterdetectie voor de plantsuiker fructose, en gaschromatografie gekoppeld aan vlamionisatiedetectie. Zo kon de lokale doorlaatbaarheid van cuticula van populierbladeren worden bepaald, op 3.39x10^9 m s^-1 voor fructose. Microscopische analyse van deze cuticula liet echter een nonrandom distributie zien van bacteriën die aan fructose blootgesteld waren. Door de twee bevindingen te combineren kon worden aangetoond dat de doorlaatbaarheid van de populierbladcuticula op sommige plekken wel 270 keer zo hoog als de gemiddelde doorlaatbaarheid. Deze resultaten geven aan dat de verdeling van suikerlekkage op het bladoppervlak niet uniform is, wat een verklaring kan zijn voor de waargenomen verschillen in het reproductieve succes van bacteriële immigranten in de fyllosfeer.

In een poging om de verschillende onderzoeksresultaten te combineren werd een ruimtelijk expliciet, ‘agent-based’ model voor bacteriële kolonisatie van de fyllosfeer geformuleerd. Met dit model, genaamd ASiMoPh voor ‘Agent-based Simulation of Microbial Phyllosphere colonisation’, werd het mogelijk te simuleren hoe bacteriën het bladoppervlak koloniseren aan de hand van eenvoudige regels voor de voortplanting van bacteriële cellen en hun opname van voedingsstoffen.
Het laatste hoofdstuk in dit proefschrift geeft een samenvatting van de stappen die tot dusver genomen zijn om de genoomsequentie van Erwinia herbicola 299R te bepalen. Met dit genoom in handen zullen genexpressiestudies en vergelijkende genoomanalyses mogelijk worden en antwoord geven op de vraag wat voor genen en genfuncties nodig zijn om als bacterie te overleven in de fylosfeer.

De experimenten en analyses in dit proefschrift bieden nieuw inzicht in de manier waarop individuele cellen van elkaar kunnen verschillen in hun lot tijdens de kolonisaatie van een nieuwe leefomgeving als gevolg van heterogeniteit in de micrometer-schaalbeleving van die leefomgeving. De CUSPER bioreporter belooft een geschikt hulpmiddel te zijn voor toekomstige studies in het kader van bacteriële individualiteit, niet alleen in de fylosfeer, maar ook in andere bacteriële habitats. Met dit proefschrift zijn een aantal belangrijke stappen gezet in de richting van een betere waardering voor de rol van bacteriële individualiteit in de microbiële ecologie.