Antimicrobial Susceptibility of Microbiota in Bacterial Vaginosis Using Fluorescence In Situ Hybridization

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Abstract: Background: Testing of antibiotic resistance of intact vaginal microbiota in pure culture is not feasible. METHODS: Metronidazole, antiseptic octenisept®, antimycotic ciclopirox, bacterial probiotic Lactobacillus crispatus, yeast probiotic Saccharomyces boulardii, Gardnerella-phase-endolysin named phagolysin and phagolysin in combination with probiotics were tested for bacteriolytic activity. Included were vaginal swabs from 38 random women with Amsel-confirmed bacterial vaginosis (BV). Test aliquots were incubated by 37°C for 2 and 24 h. Gardnerella, low G+C, Atopobium, lactobacilli, Lactobacillus iners and crispatus, Prevotella-Bacteroides, and Gammaproteobacteria microbial groups were quantified using fluorescence in situ hybridization (FISH). Results: The probiotic strain Lactobacillus crispatus demonstrated the weakest bacteriolytical effects, followed by metronidazole. Both had no impact on Gardnerella species, instead lysing Prevotella-Bacteroides, Enterobacteriaceae (by L.crispatus) or LGC, Atopobium and Prevotella-Bacteroides (by metronidazole) groups of the microbiota. Cytolytic activity on Gardnerella was highly pronounced and increased from octenisept to ciclopirox, phagolysin with L.crispatus, being best in the combination of phagolysin with S.boulardii. Universally active ciclopirox and octenisept suppressed nearly all microbial groups including those which are regarded as beneficial. Phagolysin had no effect on naturally occurring Lactobacillus crispatus. Conclusions: FISH susceptibility testing allows unique efficacy evaluation of individually adjusted topical therapy without microbial isolation facilitating optimal therapy choice.

Keywords: antimicrobial resistance; microbiome susceptibility; bacterial vaginosis; dysbiosis; FISH; polymicrobiota; metronidazole; octenisept®, ciclopirox; Lactobacillus crispatus; Saccharomyces boulardii; Gardnerella-phase-endolysin; phagolysin

1. Introduction

Bacterial vaginosis (BV) is the most common cause of vaginal discharge [1]. In addition to troublesome symptoms, BV is associated with serious complications including preterm birth, infertility and higher STI risk, among others [1,2].

The disease is characterized by a marked increase in bacterial diversity and concentrations [1,2]. Although involved microorganisms are presently identified by next generation sequencing, exact details of pathogenesis are still controversial [3]. Gardnerella species are omnipresent and dominant in nearly all cases, but are also often present in healthy
women [3,4]. Occurrence of other pathogenic anaerobes, even when found exclusively in BV, is unpredictable and variable. BV is therefore generally defined as a disappearance of the normally lactobacilli-dominated vaginal microbiota overgrown and replaced by gram-labile bacteria of which Gardnerella species are most numerous, forming structured biofilms [3,4]. The recognition of such a microbial shift is already apparent on wet microscopy, but the polymicrobial nature represents a therapeutic challenge [3]. Traditionally, the central tool of infectious disease research is susceptibility testing and measuring growth-inhibition of pure cultures of individual microbial strains. However, cultured isolates respond to antimicrobials differently than microorganisms which are growing in the form of a polymicrobial community.

Current treatment principles of vaginal dysbiosis are therefore based on estimations derived from in vitro susceptibility tests of individual isolates, clinical observations and experience based upon group statistics. Theoretical models and personal preferences led to a long list of drugs and measures with general low efficacy and high recurrence rates [5]. Susceptibility tests of native polymicrobial communities, which are necessary for evidence-based therapy are lacking. Although modern gene-based detection methods allow characterization of growth/suppression dynamics of polymicrobials, their application for susceptibility testing has not emerged. Two problems interfered. First, as opposed to isolated strains, the growth of complex microbiota outside of their natural environment is unpredictable. Each utilized culture medium favors single components of the complex microbiota differently making a balanced growth of the involved polymicrobials microbiome even for the short time in resistance testing impossible. Irrelevant bystanders overgrow and suppress constitutive microorganisms in vitro with or without antimicrobial additives. Second, it is difficult to distinguish between living organisms and their DNA remnants by culture independent methods. Both difficulties can be circumvented.

Copan Liquid Amies Elution Swab (ESwab™ 493C02 COPAN) Collection and Transport System contain puffer without nutritive additives. Vaginal smears obtained from the vaginal surface include microbiota, copious slime and desquamated epithelial cells, on which the microbiota were previously growing. The transfer in eSwab maintain the pre-existing conditions exactly the same as in the individual patient from whom they came, except for the dilution effect. Test substances can be incubated with aliquots of such suspensions providing reliable reference to each other and controls.

Ribosomal-gene-based in situ hybridization (FISH) targets intact microorganisms at species, group and domain level. Presence of microbial DNA in samples is irrelevant, since FISH signals must correlate with definite morphology to be distinguished from biases. Although FISH cannot discriminate between living and dead microorganisms (microbes are fixed at the time of FISH), the visual disappearance of the previously documented intact microbial cells indicates, that they were lysed by test substances. In the present study, we quantified the bacteriolytic effects of six common antimicrobials reported to have clinical efficacy and representing different modes of action.

2. Patients, Materials and Methods

Samples from 40 randomly selected 21–48-year-old (mean 29 years) white symptomatic women with BV diagnosed according to the Amsel criteria. Presence of sexually transmitted disease was excluded. None of the women had received an antibiotic in the previous 6 months.

ESwab™ 493C02 COPAN Diagnostics were used for collection of vaginal smears and stored at 4 °C Celsius for not longer than 24 h prior to resistance testing.

2.1. Antimicrobial Susceptibility Test

To avoid uneven distribution on the microscopic slide of heterogeneous suspensions of vaginal smears, quantifications of bacteria need a reliable reference. The best reference provides a definite number of epithelial cells with surrounding and attached bacteria, since
this is independent of dispersion biases. The sufficient number and maintenance of the epithelial cells over time of testing is therefore crucial for measurement validity.

In total, 10 µL of suspensions from vaginal smears taken in Copan E-swabs contained upon microscopy at least 50 epithelial cells within a 10 mm × 10 mm area at magnification of 400. In most cases however, more than 1000 epithelial cells were observed, comfortably allowing reliable quantification of bacteria related to epithelial cells and minimizing possible dilution and manipulation biases. Dilution by addition of antimicrobial test substances, incubation, Carnoy-fixation and resuspension steps led to about 30% loss of the initial number of epithelial cells, which needed to be compensated for by the reduced volume in which the samples were dissolved after all procedural steps. Based on this algorithm, 1000 µL of Amies medium in Copan eSwab provided sufficient material for 9 different antimicrobial tests (100 µL of the suspension + 50 µL of the test substance) for 2 h and 24 h incubation (75 µL of the mix for each) still leaving material (left directly on the swab) for other cultural, PCR and sequencing based techniques when necessary.

The incubated 100 µL of the vaginal suspension with 50 µL of test substance, terminated by Carnoy, centrifuged and resuspended in volume of 70 µL were in turn sufficient for about 10 multicolor hybridizations for each set.

2.2. Susceptibility Test

The filled eSwab-vials were vortexed gently, resulting in a suspension that contained epithelial cells and vaginal bacteria. The suspensions were divided in 10 portions of 100 µL each. All reactions and the following steps were performed in Eppendorf 1.5 mL vessels.

In total, 50 µL of the following test substances or their combinations were added dissolved in required puffer/medium to achieve the following concentrations of additives: none for untreated controls, metronidazole (1 mg/mL), ciclopirox (0.5 mg/mL), octenisept®vaginal (dilution 1:20), *Lactobacillus crispatus* (10⁸ cfu/mL suspension), *Saccharomyces boulardii* (10⁷ cfu/mL suspension), phagolysin (20 µg/mL), phagolysin + *Lactobacillus crispatus*, phagolysin + *Saccharomyces boulardii*. 50 µL Amies medium were added to untreated controls or to tests of antimicrobials to equalize the dilution.

Metronidazole (B. Braun Melsungen AG, Germany), ciclopirox (Almirall Hermal GmbH, Reinbek, Germany), octenisept®(Schülke, Norderstedt, Germany) and *Saccharomyces boulardii* CNCM I-745 (Biocodex, Gentilly, France) were purchased from the pharmacy. *Lactobacillus crispatus* was obtained from Leibnitz Institute DSMZ German Collection of Microorganisms and Cell Cultures GmbH, Phagolysin was provided by PhagoMed Biopharma GmbH. For this study engineered *Gardnerella* phage endolysin PG100 was used. For convenience reasons we refer to this preparation as “phagolysin” throughout the manuscript. In comparative investigations of lytic activity, phagolysin proved to be 100-fold more effective (data not shown) than the previously extensively tested phage endolysin pH477. We could therefore use lower concentrations than in the previous study [6].

The 150 µL of test aliquots were incubated aerobically at 37 °C and divided. After 2 and also 24 h, the action was terminated by adding 1000 µL Carnoy solution (alcohol/chloroform/acetic acid 6/3/1 by volume) [7] for 10 min. The samples were then centrifuged at 6000 RPM for 8 min, and taken and stored in 50 µL of Carnoy solution.

2.3. FISH

All identifications and quantifications of microorganisms in this study were performed exclusively based on FISH methodology.

Fields of 10 mm × 10 mm were marked on SuperFrost slides (Langenbrinck, Emmendingen, Germany) with a PAP pen (Kisker-Biotech, Steinfurt, Germany). Five µL aliquots of vortexed fixed antimicrobial resistance tests were dropped onto the marked field. The slides were dried for 60 min at 50 °C before FISH analysis.

Principally, any of the microbial groups composing the biofilms can be quantified. However, the number of hybridizations performable with a single sample are limited. The main focus of the present investigation was on populations of *Gardnerella species*, *Atopo-
bium, low GC = LGC (guanine + cytosine) bacteria (Mycoplasmatales, Firmicutes, Bacillales, Lactobacillales), lactobacilli, Lactobacillus iners, Lactobacillus crispatus, Prevotella-Bacteroides, Gammaproteobacteria and Candida species. We do not know which of these groups if any are crucial for dysbiosis. However, Gardnerella and LGC bacteria were especially suitable to quantify the antimicrobial effects since they were most numerous and ubiquitous within vaginal dysbiotic community and their FISH signals do not overlap, which was especially valuable for the quantitative assessment.

The following probes were applied: Gard662 (Gardnerella), Lab158 (lactobacilli) Liner23-2 (L.iners), Lcrips16-1 (L.crispatus), Ato291 (Atopobium), Caal (Candida) [8], low G+C probe LGC35 [9], Bact1080 [10] (Prevotella-Bacteroides), GAM42a (Gammaproteobacteria) and in special cases also Eub338 (all bacteria) [11] and Ebac1790 (Enterobacteriaceae) [12].

For multicolor analysis each the applied oligonucleotide probes were synthesized with a carbocyanine Cy3 (orange) and Cy5 (dark red) fluorescent dye. The hybridizations were performed at 50 °C as previously described [7]. DAPI stain was used to visualize DNA rich structures of bacteria and eukaryotic cells. In some cases, we applied dobe-FITC stained Eub338 probes to see all bacteria in the same set in addition to DAPI.

Hybridizations were performed with at least two differently stained probes. The Cy3 stained probe served the enumeration of the targeted bacterial cluster, Cy5 stained probe served as reference to other microbiome groups. We used Gard662-C3/LGC-C5 initially and combinations of C3 stained other microbial probes with Gard662-C5 probe as a universal reference. In the presence of high density or local microbial accumulations, the differentiation of single microorganism became sometimes difficult using the universal for all bacteria Eub338 probe as reference, because it hybridizes with exactly the same microorganisms as the FISH probe of interest.

A Nikon e600 fluorescence microscope was used. The images were photo-documented with a Nikon DXM 1200F color camera and software (Nikon, Tokyo, Japan).

The exact protocols with single steps and solutions are also presented at: http://www.swidsinski.de/zusatzdateien/fishmethode/fishmethode.htm (accessed on 1 April 2022).

2.4. Bacterial Quantification

Bacteria were enumerated in a representative area surrounding 10 epithelial cells. When low numbers of bacteria were present (for example lactobacilli, Prevotella, Gammaproteobacteria, etc.) larger areas, including at least 10 microscopic fields were evaluated and the mean number referred to 10 epithelial cells was used. When moderate numbers with spaces between bacteria were present, all countable bacteria were enumerated. In case of too high local concentrations within biofilms with no spaces between individual bacteria the counting was impossible. In such cases, the 10 × 10 µm surface of such agglomerates was assigned to 500 bacteria. The total surface covered with such biofilm was measured in area around 10 representative epithelial cells, expressed in numbers of full quadrats, multiplied with 500 and rounded to 100. Individually countable numbers of bacteria within areas of 10 epithelial cells were added to biofilm derived counts when exceeding 100.

Such mode of enumeration makes a direct comparison between moderate and high-density samples questionable. To reduce the possibility of quantification bias, we subdivided the patients in two subgroups with moderate (<500 bacteria, N = 22) and high density (>500 to 5000 bacteria, N = 18) dysbiosis and evaluated them separately. This graduation in high and moderate density was performed based on enumeration of bacteria seen in the DAPI stain.

2.5. Exceptions and Anomalies

An excessive growth of Candida completely destroyed epithelial cells after 24 h of incubation, making the comparable quantification impossible in two of the patients with moderate density dysbiosis. These two samples/patients were removed from analysis.

The incubation with phagolysin for 2 h at 37 °C led to a decrease in Gardnerella numbers (mean ± SD 840 ± 920), while LGC bacteria remained unchanged (mean ± SD 161 ± 139).
In the 2h sample from the control group the decreased values were (mean ± SD 1380 ± 1371 and 171 ± 168). No relevant changes in the mean numbers of *Gardnerella* or LGC bacteria were observed with all other antimicrobials as compared to the untreated controls after 2 h incubation (data of other test groups are not shown for irrelevance). No further hybridization or analysis of samples incubated for 2 h were performed.

The differentiated response of bacteria to other than phagolysin antimicrobials became obvious after 24 h at 37 °C of incubation. These samples were hybridized with FISH probes for all chosen microbial groups.

2.6. Use of Term Lysis

Within the frame of “classical antibiotic resistance concepts” the lytic action of antibiotics is used to describe direct degrading effects of the antimicrobial substances on the microorganisms. The direct impact of antimicrobials on bacterial cells was not und could not be demonstrated in our study, which monitors the numbers of physically present bacterial cells at initial, 2 h, and 24 h timepoints. We cannot say what exactly happens within these periods of time with the bacteria. Are bacteria directly disintegrated upon impact of antibiotics, are they weakened, dying or being decayed by other factors? What we actually measure with FISH are numeric dynamics of specific microbial groups within a polymicrobial population-rise of some and disappearance of others.

However, the scientific literature uses the word “lysis” not only as in the above-described content but also in context of dissolution, disappearance, solving, termination, etc. One should bear in mind, that speaking about bacteriolytic effects in our manuscript we do not mean direct disintegration of bacteria upon antimicrobial impact, but their disappearance following exposure to the test antimicrobials.

3. Results

General dynamics in vaginal microbiome upon incubation without therapeutic additives.

*Gardnerella* and LGC bacteria were present in all baseline samples. The baseline microbial concentrations of *Gardnerella* were mean ± SD 1494 ± 1474 for all, 2576 ± 1609 for high density dysbiosis and 156 ± 110 for moderate density dysbiosis. Other microbial groups were detected only in some samples, with lactobacilli (Lab 158), *Atopobium* (Ato291), *Prevotella* (Bacto1080) being most often (Figure 1, Tables 1 and 2).

The mean numbers of *Gardnerella* species were nearly a Log$_{10}$ higher than that of all other studied microorganism. Gram-positive *Gardnerella* composed 87%, *Atopobium* 9%, LGC 5%, and Gram-negative *Prevotella + Gammaproteobacteria* together 2% of all investigated vaginal bacteria in high density microbiome. In moderate density dysbiosis the ratio of *Gardnerella* numbers was lower with 61% (Gard), 8% (Ato), 22% (LGC) 9% (Gram-negative). The low GC bacteria are a large group including *Mycoplasmaeae, Firmicutes, Bacillales, Lactobacillales*. The increase of its ratio in moderate density dysbiosis is probably due to the higher occurrence and concentrations of *Lactobacillus iners*.

All investigated microbial groups tended to fall upon incubation at 37 °C in similar rates leaving in each of the individual vaginal smears enough bacteria for comparison of antimicrobial effects. The proportion of bacterial groups to each other remained approximately the same in 28 of 38 samples with 24h incubation (exact relation for individual data not shown). The only changes observed were due to excessive increase (one to two < Log$_{10}$) of *Candida* (4 of 8 were positive for *Candida samples*) or *Gammaproteobacteria* (6 of 18) with resulting unproportioned suppression of most other bacterial groups. This constancy of the individual microbiome profiles beyond the overgrowth exceptions was astonishing, since the vaginal fluid suspensions were incubated at aerobic conditions and the vaginal microbiome is composed of aerobe and oxygen sensitive anaerobes. Obviously, the polymicrobial environment in native vaginal slime can sufficiently sustain its components. The group specific overgrowth in some samples indicates that deprived of actively present vaginal immunity, some strains within vaginal microbiome escape the restraints of the
previously protective community environment. The additional hybridization of samples with *Gammaproteobacteria* overgrowth demonstrated, that the increment is solely due to *Enterobacteriaceae* (Ebac1790 probe) probably representing some recent incomer. In two samples *Candida* overgrowth was thus massive that all epithelial cells were destroyed and had to be removed from the resistance evaluation. In four, the growth did not visually affect the epithelial cell numbers. These four *Candida* positive and six samples with *Enterobacteriaceae* overgrowth were not removed from the antibiotic resistance testing, since despite chaotic changes in ratios of the initially observed microbial group, each sample sustained enough microbial representatives for a comparison of antimicrobial effects.

![Figure 1](image-url)

**Figure 1.** Mean ± SD concentrations of omnipresent *Gardnerella* and LGC bacteria surrounding 10 epithelial cells in the antimicrobial tests. The abbreviations are the same for all figures and tables in this manuscript: N-number of patients, c0—baseline controls, c2/c24—controls after 2 h and 24 h incubation, M—metronidazole, Ci—ciclopirox, O octenisept®, L—*Lactobacillus crispatus*, S—*Sacharomyces boulardii*, P—phagolysin, PL—phagolysine + *Lactobacillus crispatus*, PS—phagolysin + *Sacharomyces boulardii*. HDD—high density dysbiosis, MDD moderate density dysbiosis. "*" indicates statistical significance with \( p < 0.05 \) as assessed with one-way ANOVA with multiple comparisons against the c24 control.
**Table 1.** Mean concentrations of omnipresent *Gardnerella* and LGC bacteria surrounding 10 epithelial cells in the antimicrobial tests.

|                  | *Gardnerella* | LGC Group |                  |
|------------------|---------------|-----------|------------------|
|                  | All HDD MDD   | All HDD MDD |
| N                | 38 20 18      | 38 20 18  |
| c0               | 1494 2576     | 156 202  |
| c2               | 1380          | 161      |
| c24              | 1097 1917     | 179 224  |
| M                | 1032 1803     | 67 70    |
| Ci               | 93 153        | 28 40    |
| O                | 331 587       | 29 39    |
| L                | 997 1756      | 175 229  |
| S                | 816 1430      | 194 251  |
| P                | 97 161        | 154 188  |
| PL               | 88 150        | 156 202  |
| PS               | 63 120        | 1 242    |

k24/c,o,p,pl,ps  p < 0.001

24/c,o,p,pl,ps  p < 0.001

k24/c,o,p,pl,ps  p < 0.05

k24/p,ps  p < 0.01

k24/ps  p = 0.03

k24/m, c,o p < 0.001

k24/p/ps  p = 0.012

k24/ps  p = 0.03

k24/m p < 0.02

p/ps  p < 0.01

p/ps  p = 0.01

p/ps  p = 0.03

SD values are not demonstrated since irrelevant for presentation. The abbreviations are the same for all tables in this manuscript: N—number of patients, c0—baseline controls, c2/c24—controls after 2 h and 24 h incubation, M—metronidazole, Ci—ciclopirox, O octenisept®, L—*Lactobacillus crispatus*, S—*Sacharomyces boulardii*, P—phagolysin, PL—phagolysine + *Lactobacillus crispatus*, PS—phagolysin + *Sacharomyces boulardii*. HDD—high density dysbiosis, MDD moderate density dysbiosis. Bold numbers within the table are the numbers significantly reduced as compared to 24 h controls.
Table 2. Mean concentrations of bacteria detected only in some vaginal samples in the antimicrobial tests.

|                | Atopobium | Lactobacilli (Lab158) | L.iners | L.crisp | Prevotella | Gammaproteobacteria | Candida |
|----------------|-----------|-----------------------|---------|---------|------------|--------------------|---------|
|                | All       | HDD | MDD | All       | HDD | MDD | All       | HDD | MDD | All       | HDD | MDD | All       | HDD | MDD | All       | HDD | MDD | All       |
| N=             | 24 (64%)  | 16  | 8   | 25 (66%)  | 15  | 10  | 14 (36%)  | 5   | 9   | 5 (30%)  | 24  | 13  | 11 (25%)  | 15  | 5(2) | 10 (48%)  | 5(2) | 13  | 15 (63%) |
| c0             | 92 (32%)  | 125 | 36  | 43 (28%)  | 44  | 22  | 12 (22%)  | 49  | 172 | 49 (20%) | 17  | 13  | 24 (18%)  | 23  | 22  | 15 (63%) |
| c24            | 62 (76%)  | 86  | 38  | 38 (54%)  | 27  | 14  | 16 (22%)  | 40  | 204 | 40 (28%) | 46  | 37  | 54 (11%)  | 470 | 412 | 631 (24%)|
| M              | 48 (75%)  | 73  | 7   | 37 (16%)  | 26  | 18  | 12 (20%)  | 40  | 192 | 37 (6%)  | 0.7 | 0.7 | 0.5 (90%) | 458 | 377 | 597 (20%)|
| Cl             | 10 (20%)  | 17  | 3.3 | 12 (20%)  | 2.8 | 22  | 8.4 (10%) | 1.2 | 13  | 2.8 (10%) | 16  | 19  | 12 (15%)  | 148 | 193 | 111 (20%)|
| O              | 2.6 (70%) | 3.6 | 1.4 | 5.3 (60%) | 3.7 | 6.9 | 4.9 (60%) | 2.8 | 6.8 | 1.8 (90%) | 1   | 1.5 | 0.8 (40%) | 9.6 | 9.2 | 9.9 (30%)|
| L              | 61 (90%)  | 84  | 31  | 36 (60%)  | 30  | 41  | 170 (90%) | 45  | 214 | 38 (60%) | 13  | 16  | 11 (10%)  | 118 | 84  | 157 (20%)|
| S              | 46 (75%)  | 78  | 12  | 23 (60%)  | 19  | 26  | 56 (60%)  | 22  | 87  | 40 (60%) | 48  | 49  | 46 (10%)  | 556 | 460 | 652 (20%)|
| P              | 33 (55%)  | 55  | 5.8 | 27 (55%)  | 18  | 36  | 132 (60%) | 41  | 150 | 39 (60%) | 60  | 39  | 80 (10%)  | 510 | 425 | 595 (20%)|
| PL             | 32 (50%)  | 52  | 5.5 | 29 (50%)  | 25  | 33  | 131 (60%) | 50  | 176 | 37 (60%) | 12  | 13  | 12 (10%)  | 115 | 115 | 116 (20%)|
| PS             | 22 (35%)  | 40  | 3.2 | 25 (35%)  | 18  | 33  | 23 (10%)  | 12  | 28  | 44 (60%) | 103 | 88  | 116 (90%) | 858 | 569 | 1168 (20%)|

k24/ps

p = 0.001

k24/c,o

p < 0.001

k24/m

p < 0.05

k24/o

p < 0.05
4. Measurement of Antibiotic Resistance

All tested antimicrobials demonstrated significant lytic effects on at least one of the microbial groups, and the overall lytic patterns were agent specific and differed strongly from each other.

5. Gardnerella and LGC

While some of the antimicrobials led to dramatic changes in *Gardnerella* and LGC concentrations, distributions and form were hardly or not distinguishable from the controls (Figures 1 and 2; Tables 1 and 2).

![Figure 2](image_url). Multicolor FISH demonstrates the effects of antimicrobials on *Gardnerella* (Gard662 C3 orange fluorescence) and Low GC bacteria (LGC 35-Cy5 dark red fluorescence) at magnification ×400.
in two BV patients representing high and moderate density dysbiosis. The changes in corpuscular integrity of bacteria in vaginal microbiome can be well seen. Susceptible bacteria either disappear or they lose their structure according to the grade/range of disintegration after the applications of specific antimicrobial drugs. Due to selective effects of antimicrobials on different species, the proportions of bacteria in the samples change dramatically. Low concentrated species previously outnumbered by dominant representatives and imperceptible become apparent. Resistant to antimicrobial species start to grow and become predominant groups within the in vitro cultured microbiome. This leads in multicolor FISH to dramatic changes in color of the predominant population. Gardnerella species (orange fluorescence) dominate the microbiome population in controls after 24 h incubation without antimicrobials. Because of the high numbers of Gardnerella, Low-GC bacteria cannot be seen well (red fluorescence). Gardnerella still is still dominant in metronidazole, L. crispatus and Saccharomyces boulardii (high density dysbiosis) exposed samples. Gardnerella is decimated in other test sets dismantling LGC bacteria, when those were not likewise lysed with octenisept or ciclopirox.

Generally, the 24 h incubation led to a reduction in Gardnerella species and an increase in LGC bacteria, however the changes compared to baseline remained statistically insignificant $p = 0.2$ (Table 1). Statistically insignificant were the bacteriolytic impact of metronidazole, Lactobacillus crispatus or Saccharomyces boulardii although the mean concentrations of Gardnerella were 6%, 9%, and 26% lower in these groups than in the non-treated controls.

Most profound lytic effects ($p < 0.001$) were observed with ciclopirox, octenisept® and phagolysin, phagolysin combined with Lactobacillus crispatus, and Saccharomyces boulardii. Ciclopirox and octenisept® decimated both Gardnerella and LGC bacteria. No effect on LGC bacteria was observed with phagolysin, and an increase of LGC bacteria in phagolysin + Saccharomyces boulardii groups ($p < 0.05$).

Dividing the patients into those with moderate and high density dysbiosis (according to enumeration procedure described in the method section) did not change the observed effects of antimicrobials on Gardnerella and LGC bacteria for all but phagolysin + Saccharomyces boulardii group, where the previously observed tendency to lower Gardnerella numbers after exposing to probiotics became clearly statistically significant ($p < 0.001$), nearly completely eradicating Gardnerella in more than 75% of the samples.

5.1. 24 h Susceptibility of Other Microbial Groups

Measurable quantities of the Atopobium, lactobacilli, Lactobacillus iners, Lactobacillus crispatus, Prevotella-Bacteroides, Gammaproteobacteria and Candida groups were detected in 13% to 64% of the baseline vaginal samples and were still detectable in all corresponding control samples incubated for 24 h. Despite the low number of patients within single groups many of the antimicrobial effects reached a level of statistical significance (Table 2).

5.2. Atopobium

Octenisept® and ciclopirox significantly reduced Atopobium in all subgroups ($p < 0.05-0.001$). The effects of other antimicrobials were less impressive and often not clear in context of direct or mediated effects. Atopobium numbers fell already with incubation of controls and mean numbers in all other test groups were lower than in controls reaching statistical significance ($p < 0.05$) only for phagolysin and phagolysin in combination with L. crispatus and Saccharomyces boulardii groups in moderate density dysbiosis. However, the parallelism of Atopobium and Gardnerella changes together with general instability of this microbial group indicates that the suppressive effects may be not due to direct antimicrobial action but be partially mediated by concomitant changes in Gardnerella availability.
5.3. Lactobacillus Species

Bacteria positive for Lab158 probe were sensitive to octenisept® and ciclopirox especially in the subgroup with high density dysbiosis. All other antimicrobials had no visible effects on them. *Lactobacillus iners* was sensitive towards *Saccharomyces boulardii* alone (ns) and in combination with phagolysin (*p* < 0.05).

*Lactobacillus crispatus* was detected in samples from three patients with high and two patients with moderate density dysbiosis. Only octenisept® and ciclopirox demonstrated direct effects on *L. iners* and *crispatus*.

5.4. Prevotella

Bacteria of the *Prevotella-Bacteroides* group were sensitive against metronidazole, octenisept®, intermediate to ciclopirox and *Lactobacillus crispatus* in combination with phagolysin.

5.5. Gammaproteobacteria

Different to all other investigated bacterial groups, the numbers of *Gammaproteobacteria* tended to be significantly (*p* < 0.05) increased upon incubation even in the controls. The increase was by Log$_{10}$2 in two patients with high density dysbiosis and four patients with moderate density dysbiosis. The increase was clearly inhibited by octenisept®, ciclopirox and *Lactobacillus crispatus* alone and in combination with phagolysin, however the effects were not statistically significant for most because of the low number of patients (*p* = 0.07/0.08). Octenisept® appeared to be the only substance which significantly reduced *Gammaproteobacteria* (*p* < 0.05).

5.6. Candida

Similar to *Enterobacteriaceae, Candida* increased by more than one Log$_{10}$ in four out of six included and by more than Log$_{10}$2 in two patients, which had to be excluded from the investigation for excessive overgrowth. Only ciclopirox, octenisept® and *Sachharomyces boulardi* had a marked cytolytic, or growth suppressive effect on *Candida* (*p* < 0.01).

6. Discussion

Our data demonstrate that the differential lytic effects of antibiotics on complex microbial dysbiosis can be clearly quantified by FISH for each of the agents tested.

Interestingly, all of the chosen agents had a statistically significant lytic effect on at least one of the studied microbial groups. As expected, the single components of microbiota responded disparately to different antimicrobials with some of the groups being decimated, while other promoted. The effect varied in patients with high and moderate density dysbiosis. However, all observed effects in this study differed from those which we had anticipate based on our previous clinical practice and experience.

The mean values of multiple subgroups are difficult to compare at glance. To get a better overview using typical for clinicians susceptibility grading, in Table 3, (additional information only) we have sorted the effects presented in Figure 1 in the following steps: ±25% changed (R—resistant), decreased between 25%≤ and ≤Log$_{10}$ (I—intermediately susceptible), decreased between Log$_{10}$1 and ≤Log$_{10}$2 (S—sensitive), decreased by more than ≥Log$_{10}$2 (SS—highly sensitive), or increased accordingly 25%≥ <Log$_{10}$1 (P—promoting).

Absolutely surprising was the complete indifference of *Gardnerella* species to metronidazole, the most often used antibiotic. The strong effects of metronidazole on Gram negative *Prevotella-Bacteroides* group were expected. However, both occurrence and numerical ratio of these bacteria on the whole dysbiosis was marginal. Unexpectedly, metronidazole displayed also intermediate effects on LGC bacteria in all and *Atopobium* in moderate density dysbiosis subgroup. However, the LGC group is very broad including *Mycoplasmales, Firmicutes, Bacillales, Lactobacillales* [9] and the anaerobe *Firmicutes* species are sensitive to metronidazole.
Table 3. Susceptibility of vaginal microbiota to lysis as related to untreated controls.

|               | Gardnerella | LGC | Atopobium | Lactobacilli | Liners | L.crisp. | Prevotella-Bact. | Gammaproteobac. | Candida |
|---------------|-------------|-----|-----------|--------------|--------|----------|------------------|-----------------|---------|
|               | All         | HDD | MDD       | All          | HDD    | MDD      | All              | HDD            | MDD     |
| M             | R           | R   | R         | I            | I      | I        | R                | R              | R       |
| C             | S           | S   | I         | I            | I      | I        | S                | S              | I       |
| O             | I           | I   | I         | I            | I      | I        | I                | I              | I       |
| L             | R           | R   | R         | R            | R      | R        | R                | R              | R       |
| S             | I           | I   | I         | R            | R      | R        | I                | I              | I       |
| P             | S           | S   | I         | R            | R      | R        | I                | I              | I       |
| PL            | S           | S   | S         | R            | R      | R        | I                | I              | I       |

R-resistant; S-sensitive; SS-highly sensitive; I-intermediate; P-promoting.
As expected, metronidazole was the only of the tested substances, which promoted the growth of Candida ($p < 0.05$).

Likewise, highly impressive was the pronounced effect of the antifungal ciclopirox on all investigated participants of the vaginal microbiome regardless of bacterial density. The effect of ciclopirox on Gardnerella clearly exceeded that of octenisept® and was achieved only by phagolysin and exceeded that of the phagolysin + probiotics combination. Although ciclopirox is known to be suppressive for Gram-positive bacteria, it was not considered appropriate for antibacterial therapy. Ciclopirox was, as expected, best in the lysis of Candida cells.

The antimicrobial effects of octenisept® were in turn predictable and generally in accordance with all what we presently know. Compared with phagolysin and ciclopirox, octenisept® was less effective against Gardnerella (especially in high density dysbiosis) and Candida but exceeded effects of ciclopirox on Atopobium, lactobacilli (including Lactobacillus iners and crispatus), Prevotella-Bacteroides and Gammaproteobacteria groups.

The antimicrobial efficacy of probiotic strains was relatively low, when used alone. Lactobacillus crispatus had intermediate effects only on Prevotella-Bacteroides and Gammaproteobacteria (statistically not significant). Saccharomyces boulardii demonstrated intermediate effect on Gardnerella, on Atopobium (in patients with moderate dysbiosis density), on lactobacilli (in high density dysbiosis) and specifically on Lactobacillus iners in both dysbiosis groups, but only on Candida did the effects of Saccharomyces boulardii reach the level of statistical significance despite a very low number of the patients in this group. We are aware that in clinical practice, the probiotic Lactobacillus species were often used as combination of strains and were judged solely on evaluation of symptoms. However, Lactobacillus crispatus is regarded as the most potent, and none of the other components of such combinations were previously tested, while our data clearly indicate the necessity of such tests. On the other hand, probiotic microorganisms applied together with the biologic phagolysin potentiated the effect of phagolysin (or vice versa) on Gardnerella, especially in patients with moderate density dysbiosis leading to Log$_{10}$2 reductions of Gardnerella after addition of Saccharomyces boulardii. Phagolysin combined with Lactobacillus crispatus had also intermediate effect on Atopobium in high density dysbiosis group ($p < 0.05$), on Prevotella-Bacteroides ($p < 0.05$), on Gammaproteobacteria (ns). Phagolysin in combination with Saccharomyces boulardii suppressed Gardnerella more intensively, lysed Atopobium in moderate density dysbiosis (ns), L iners ($p < 0.05$) and Candida in all dysbiosis groups (ns), displaying effects which were not observed by application of phagolysin alone.

The clear potentiating effects of combined applied biological agents demonstrate that in treating microbial communities scientific data driven combination therapy may be preferable to achieve optimal clinical effects. Depending on initial composition of the microbiome and the individual aims, these combinations may include different antimicrobials, biologics, immune modulating substance and more.

7. Concluding Comments and Limitations

Although single forms of bacterial vaginosis are presumably sustained by a limited number of essentially responsible species, multiple accidently associated and transient microorganisms including bacteria, fungi, and phages make true relations difficult to untangle. The pathways by which polymicrobials prevail are poorly understood [3]. The present therapy of bacterial vaginosis is therefore highly problematic, purely empiric and employs a broad range of antibiotics, antiseptics, probiotics, and recently phage-derivatives [5,6].

In spite of the unclear etiopathogenesis, testing of antimicrobial susceptibility of the entire dysbiotic population could be crucial for effective therapy. This was impossible previously with methods based on isolation of single microorganisms. The development of FISH enabled direct monitoring of changes within complex microbial populations. We are aware of the fact, that chosen for this study marker-microorganisms, FISH probes and test conditions are far from optimal for the adequate assessment of clinically relevant antibiotic
resistance. However, the classic culture based antibiotic susceptibility testing of isolated strains similarly needed years and many studies before it took its place in clinical practice. Antibiotic susceptibility of polymicrobials may similarly take many years.

Critically seen, the lytic activity measured may not be the same as killing activity, “not lysed” bacteria may be still affected by antimicrobials and non-vital. We are also aware that the ranges of lytic effects observed in vitro may strongly differ from those in clinical application. However, the same is true for any antimicrobial testing, and the in vitro determined minimal inhibitory concentration of classic tests have to be adjusted by corresponding clinical efficacy for each individual antimicrobial substance, regardless of effects observed in vitro. However, this new possibility of testing the efficacy of single agents alone and in combination for an individual dysbiotic microbiome opens compelling horizons in the treatment of challenging polymicrobial diseases such as BV and may prove to be a springboard for managing of all other polymicrobial diseases.

We acknowledge that this report does not answer all questions involved in this new methodology. However, this is the first effort made to approach susceptibility of a composite complex community of bacterial rather than each individual species involved. Additional studies are needed. This report serves as the first introductory study addressing the problem and supports several clinical observations.

Author Contributions: Each author named in the byline participated actively and sufficiently in the study. The author contributions: A.S. and Y.D. designed the study, A.S., L.C., L.P.T. and S.S., conducted the study, J.D.S. and V.L.-B. critically revised the manuscript, and A.S., A.G. and Y.D. performed FISH, A.S., L.C., L.P.T. and S.S. analyzed the data. All authors have read and agreed to the published version of the manuscript.

Funding: The study was supported by the Charité University research promotion grant (2016-033) and The German Federation of Industrial Research Associations ZIM Project ZF4143701AJ5. Both funding sources were not involved in study design, collection, analysis and interpretation of data, in writing a report or the decision to submit the article for publication.

Institutional Review Board Statement: The ethical committee approval number is EA1/088110, the IRB form is attached.

Informed Consent Statement: Not applicable.

Data Availability Statement: The data of this publication were not presented previously. The work has not been published previously and is not under consideration for publication elsewhere. The publication is approved by all authors and tacitly or explicitly by the responsible authorities where the work was carried out. If accepted, it will not be published elsewhere in the same form, in English or in any other language, including electronically without the written consent of the copyright-holder.

Conflicts of Interest: The authors have no conflict of interest.

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