Quantitation of all Four *Gardnerella vaginalis* Clades Detects Abnormal Vaginal Microbiota Characteristic of Bacterial Vaginosis More Accurately than Putative *G. vaginalis* Sialidase A Gene Count

Elena Shipitsyna1 · Anna Krysanova1 · Guzel Khayrullina2 · Kira Shalepo1 · Alevtina Savicheva1 · Alexander Guschin2 · Magnus Unemo3

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Abstract

**Background** Bacterial vaginosis (BV) is a vaginal disorder characterized by a depletion of the normal lactobacillus-dominant microbiota and overgrowth of mainly anaerobic bacteria.

**Objectives** The study aimed to evaluate the distribution and abundance of the *Gardnerella vaginalis* clades and sialidase A gene in vaginal samples from Russian women, and investigate if the *G. vaginalis* sialidase A gene count detects an abnormal vaginal microbiota characteristic of BV more accurately than *G. vaginalis* load.

**Methods** Vaginal samples from 299 non-pregnant patients of gynecological clinics were examined using Nugent scores and *G. vaginalis* clade and sialidase A gene quantitative real-time polymerase chain reactions (PCRs). Discriminatory power for BV microbiota was evaluated with receiver operating characteristic (ROC) analysis.

**Results** The vaginal microbiota was characterized by Nugent scores as normal, intermediate, and BV microbiota in 162, 58, and 79 women, respectively. *G. vaginalis* clades 1, 2, 3, 4, and the sialidase A gene were detected in 56% (51–62%), 40% (34–45%), 20% (16–25%), 94% (91–96%), and 70% (64–75%) of vaginal samples, respectively. The frequency and abundance of clades 1, 2, 4, and the sialidase A gene as well as clade multiplicity were significantly associated with abnormal microbiota. The sialidase A gene was present in all multi-clade samples, in all single-clade samples comprising clades 1, 2, and 3, and in four of 84 (5% [2–12%]) samples comprising clade 4 only. Total *G. vaginalis* load showed significantly higher discriminatory power for abnormal microbiota than sialidase A gene count (areas under ROC curves 0.933 vs. 0.881; *p* = 0.0306).

**Conclusions** Quantifying all four *G. vaginalis* clades discriminates between BV microbiota and normal microbiota more accurately than measuring *G. vaginalis* sialidase A gene. Clade 4 is strongly associated with BV microbiota, despite most clade 4 strains lacking the sialidase A gene.

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**Key Points**

High *Gardnerella vaginalis* overall load predicts abnormal vaginal microbiota characteristic of bacterial vaginosis (BV) better than high sialidase A gene counts.

Although *G. vaginalis* clade 4 strains appear to lack the sialidase A gene, it is strongly associated with BV microbiota.
Bacterial vaginosis (BV) is a vaginal disorder characterized by a depletion of the normal lactobacillus-dominant microbiota and overgrowth of mainly anaerobic bacteria, i.e., an abnormal vaginal microbiota. Clinically, BV is manifested as abnormal malodorous vaginal discharge, but an abnormal vaginal microbiota characteristic of BV can be detected also in asymptomatic individuals [1]. BV is associated with an increased risk of many gynecological and obstetrical complications [2–5]. *Gardnerella vaginalis* is a common bacterial species in the vaginal microbiota and can be found in many healthy women. However, it is substantially more abundant and nearly ubiquitous in women with BV [6, 7]. *G. vaginalis* is considered to play a key role in the pathogenesis of BV, being an initial colonizer of the vaginal epithelium due to the higher adhesion, cytotoxicity, and greater propensity to form biofilm (compared to other BV-associated bacteria), and acting as a scaffold to which other species subsequently can attach [8–10].

*Gardnerella vaginalis* is a phenotypically and genetically heterogeneous species [11–17]. Eight metabolic biotypes of *G. vaginalis* were identified early based on the presence of β-galactosidase, lipase, and hippurate hydrolysis activities [11], and 17 biotypes based on these characteristics and additionally on utilization of xylose, arabinose, and galactose [12]. Phenotypic diversity within the *G. vaginalis* species has also been shown in relation to virulence factors, such as production of sialidase and vaginolysin, initial adhesion, cytotoxic effect, biofilm formation, susceptibility to antimicrobials, and the ability of *G. vaginalis* to displace lactobacilli adhered to HeLa cells [18–22]. Association of *G. vaginalis* biotypes with BV has been evaluated in a number of studies, but the results have been contradictory [11, 19, 20, 23–25]. Sialidase is utilized by some BV-associated bacteria, including *G. vaginalis*, for degradation and depletion of protective host mucus barriers [26, 27]. Detection of the BV virulence factor sialidase has also been used in rapid point-of-care tests for BV, e.g., BVBlue® (Sekisui Diagnostics, L.L.C., Birmingham, AL, USA) [28].

Genetic heterogeneity within the *G. vaginalis* species has been demonstrated using molecular approaches, such as amplified ribosomal DNA restriction analysis (ARDRA) [13]. Using this method, three distinct *G. vaginalis* genotypes were identified, of which two produced sialidase. Although a link with sialidase production was shown for particular *G. vaginalis* ARDRA genotypes [18, 20], no clear association between BV and any of the ARDRA genotypes has been identified [18]. Different genetic variants of *G. vaginalis* based on a comparative genomic analysis have been described, and evidence of substantial differences in metabolic and virulence potential between *G. vaginalis* clinical strains has been reported [14, 15].

More recently, detailed genomic analyses [16] have identified four main clades within the *G. vaginalis* species. There were substantial differences in genome size, GC ratio, and core gene content between the clades, suggesting a possibility of considering them as separate bacterial species. Four subgroups of *G. vaginalis* (subgroups A–D), based on sequencing of a region of the chaperonin-60 (cpn60) gene, have also been proposed to represent different species [17]. Subsequently, *G. vaginalis* clades 1–4 were reconciled with *cpn60* subgroups A–D [29]: *cpn60* subgroup A corresponded to clade 4, subgroup B to clade 2, subgroup C to clade 1, and subgroup D to clade 3. In the same study [29], presence and activity of the putative sialidase A gene were investigated in *G. vaginalis* isolates. All isolates of subgroups B, C, and D, but only one (3%) of the subgroup A isolates, possessed the putative sialidase A gene, while sialidase activity was displayed by all isolates of subgroup B, three (9%) subgroup C isolates, and none of the isolates of subgroups A and D, suggesting that this gene is not expressed (or not constitutively expressed) in many isolates. An association of the *G. vaginalis* sialidase A gene, as a proxy for sialidase production, with biofilm on the vaginal epithelium and with BV was subsequently examined [30]. The sialidase A gene was detected in 75% of *G. vaginalis*-positive vaginal samples, and there was a strong association between high counts of the *G. vaginalis* sialidase A gene, the presence of *G. vaginalis* biofilm, and BV [30]. These observations indicate distinct clinical and diagnostic relevance of *G. vaginalis* subgroups/clades for BV.

The objectives of this study were to evaluate the distribution and abundance of the *G. vaginalis* clades, corresponding to the *cpn60* subgroups, and putative sialidase A gene in vaginal samples from Russian women with normal and BV microbiota, and to investigate if *G. vaginalis* sialidase A gene counts can detect an abnormal microbiota characteristic of BV more accurately than *G. vaginalis* bacterial load.

## 2 Materials and Methods

### 2.1 Patients, Specimens and Bacterial Vaginosis Diagnosis

The study subjects were women of reproductive age (from 18 to 54 years) who attended gynecological clinics in St Petersburg, Russia with vaginal symptoms (mainly abnormal vaginal discharge) from October to December 2015. The exclusion criteria were pregnancy and taking oral or intravaginal antimicrobials within the last month.
Vaginal samples were collected using two Dacron® swabs. One vaginal swab was used for preparing a microscopic slide, which was Gram stained and analyzed microscopically for bacterial morphotypes according to the Nugent scores [31]. The other vaginal swab was placed into 0.5 mL of buffered saline solution supplemented with mucolytic, preservative, and stabilizing agents (Transport Medium with Mucolytic Agent, InterLabService, Moscow, Russia) and used for polymerase chain reaction (PCR) detection of the *G. vaginalis* clades and the putative *G. vaginalis* sialidase A gene.

### 2.2 Quantitative Polymerase Chain Reaction (PCR) Detection of the *Gardnerella vaginalis* Clades and the Putative *G. vaginalis* Sialidase A Gene

The detection of the four clades of *G. vaginalis* [16] was performed using a multiplex real-time PCR assay with previously described primers and probes [32]. Real-time PCR analysis of the *G. vaginalis* sialidase A gene was performed using the forward primer GVSI forward (5′-GAC GAC GGC GAA TGG CAC GA-3′), designed by Santiago et al. [18], and the newly designed reverse primer GvSia1R2 (5′-GCT GTA AGT TTA TTA TTA CTA CAA-3′) and TaqMan® probe GvSia1Z2r (5′-R6G-CTC CGC GAT TTG CGC GAA TAATC-BHQ1-3′). The design of the new oligonucleotides was based on the alignment of the sequence of the sialidase A gene from *G. vaginalis* American Type Culture Collection (ATCC) 14019 (National Center for Biotechnology Information [NCBI] reference sequence: NC_014644.1 [https://www.ncbi.nlm.nih.gov/nucleotide/NC_014644.1]), used by Santiago et al. [18], and corresponding sequences of other *G. vaginalis* strains (belonging to different clades), described in the study by Ahmed et al. [16]. In this sequence alignment, the previously used reverse primer GVSI [18] had two mismatches at the 3′-end in some strains, which could reduce the sensitivity of detecting some *G. vaginalis* strains. A new TaqMan® hybridization probe specific for the conservative part of the chosen fragment was also designed to enable accurate, sensitive, and specific detection. Specificity of the newly designed primer and probe was verified using Standard Nucleotide BLAST® (Basic Local Alignment Search Tool) software (NCBI; https://blast.ncbi.nlm.nih.gov/Blast.cgi).

DNA for both PCR assays was isolated from 100 μL of sample using the silica-based manual extraction kit DNA-Sorb-AM (InterLabService, Moscow, Russia), according to the manufacturer’s instructions, with an elution volume of 100 μL. The DNA preparations were stored at 4 °C prior to amplification, which was performed within 3 days.

PCR amplifications were performed in reactions (25 μL) containing 10 μL of Mix-1 (0.7 pmol/μL of each primer, 0.2 pmol/μL of each probe, 0.44 mM deoxyribonucleotide triphosphates (dNTPs), 0.1 mM Tris, and 0.1 mM EDTA), 5 μL of Mix-2 (20% glycerol, 0.17 M Tris–HCl (pH 8.3), 42 mM (NH₄)₂SO₄, 0.025% Tween-20, 7.5 mM MgSO₄, 0.25 mg/mL of bovine serum albumin (BSA), 0.02% xilenethanol, 0.125% sodium azide, 0.5 μL of Taq-polymerase), and 10 μL of DNA template. The PCR program parameters were as follows: 95 °C for 15 min followed by 40 cycles of 95 °C for 15 s, 60 °C for 20 s, and 72 °C for 15 s, with fluorescence read at 60 °C. All PCR amplifications were performed on a Rotor-Gene Q instrument (QIAGEN, Hilden, Germany). Negative controls (sterile nucleic acid-free water) were included in each run.

For quantification of PCR-amplified fragments, quantitative standard samples were constructed by cloning PCR fragments of the targeted genes into pGEM®-T Vector Systems (Promega, Madison, USA). DNA concentration of plasmid preparations was tested using digital PCR with the same specific primers and probes on a QX100™ Droplet Digital™ PCR System (BioRad, Hercules, CA, USA). Standard curves were generated by testing quantitative standard samples (in duplicate) in concentrations of 10³, 10⁵, 10⁷, and 10⁹ genome equivalents (geqs)/mL for all targets. Concentrations in the clinical samples were expressed as geqs/mL of DNA extract.

### 2.3 Statistical Analyses

Based on the Nugent scores, all women were divided into three groups, i.e., women with normal vaginal microbiota (a score of 0–3), intermediate microbiota (4–6), and BV microbiota (7–10) [31]. Categorical variables (the presence of the *G. vaginalis* clades and sialidase A gene and the presence of multiple *G. vaginalis* clades) were evaluated using Pearson’s Chi squared test for all three groups of patients, with Cramer’s V coefficient measured to evaluate the strength of association, which was followed by pairwise comparisons using Fisher’s exact test. Continuous data (the loads of the *G. vaginalis* clades and sialidase A gene), after being characterized as not normally distributed in the Shapiro–Wilk test, were analyzed using the Kruskal–Wallis test for all three groups followed by post hoc pairwise comparisons using Dunn’s multiple comparisons test. All percentages were reported with 95% confidence intervals (CIs), computed with the use of Wilson’s method with Brown’s modification. To assess the ability to differentiate between samples with normal and abnormal vaginal microbiota using total *G. vaginalis* DNA load, *G. vaginalis* clades and sialidase A gene counts, receiver operating characteristic (ROC) curves were plotted, and classification performance characteristics (areas under the curves [AUCs], as well as true positive proportion [sensitivity], true negative proportion [specificity], and Youden’s index in optimum points) were computed. Differences between AUC values were assessed using the DeLong method. Statistical analyses were performed with △ Adis
the use of the statistical packages GraphPad Prism version 7.04 for Windows (GraphPad Software, San Diego, CA, USA) and Analyse-It for Microsoft Excel 5.11 (Analyse-it Software, Leeds, UK). All tests for significance were two-sided, and statistically significant differences were assumed when \( p < 0.05 \).

### 3 Results

In total, 299 women aged 18–54 years (median age 30 years; mean age 32 years) were enrolled in the study. The vaginal microbiota was characterized as normal in 162 women (median age 29 years), intermediate in 58 (median age 31 years), and BV microbiota in 79 (median age 30 years). *G. vaginalis* DNA was detected in 288 women: 151 women with normal microbiota and in all women with intermediate microbiota and abnormal microbiota (Table 1). The most prevalent *G. vaginalis* clade was clade 4 (94% [91–96%]), followed by clades 1 (56% [51–62%]), 2 (40% [34–45%]), and 3 (20% [16–25%]). The *G. vaginalis* sialidase A gene was detected in 208 women (70% [64–75%]). Clades 1, 2, and 4 and the sialidase A gene were significantly associated with BV microbiota (Table 1).

Loads of *G. vaginalis* clades 1, 2, and 4 as well as total *G. vaginalis* load and sialidase A gene counts were significantly higher in intermediate microbiota than in normal microbiota, in abnormal versus intermediate, and in abnormal versus normal samples. For clade 3, no difference was observed between loads in abnormal versus normal and in abnormal versus intermediate samples (Table 2).

Of the 288 *G. vaginalis*-positive women, 89 (31% [26–36%]) harbored a single clade and 199 (69% [64–74%]) multiple clades (two or more). Multiple clades were detected significantly more often in abnormal than in intermediate microbiota (odds ratio [OR] 0.234; \( p = 0.0019 \)) and in abnormal than in normal microbiota (OR 0.163; \( p < 0.0001 \)). No significant difference between normal and intermediate microbiota was observed (OR 0.701; \( p = 0.3416 \) (Fig. 1).

The large majority (84 of 89; 94% [88–98%]) of the single-clade samples contained clade 4, with 57 samples obtained from women with normal microbiota, 19 samples from women with intermediate microbiota, and eight samples from women with BV microbiota. The remaining five samples contained clades 1 (\( n = 3 \)), 2 (\( n = 1 \)), and 3 (\( n = 1 \)) with all of them obtained from women with normal microbiota. The sialidase A gene was present in all single-clade samples comprising clades 1, 2, and 3 and in 4 (5% [2–12%]) single-clade samples comprising clade 4: two samples with normal microbiota and two samples with intermediate microbiota. In all multi-clade samples, the sialidase A gene was present.

With the use of ROC analysis, we evaluated the usefulness of DNA counts of the *G. vaginalis* clades and putative sialidase A gene for distinguishing BV microbiota from normal microbiota (Fig. 2). Total *G. vaginalis* load possessed a superior discriminatory power (AUC 0.933), followed by the counts of the sialidase A gene (AUC 0.881), clade 4 (AUC 0.857), clade 1 (AUC 0.827), and clade 2 (AUC 0.818).

### Table 1 Frequency of the Gardnerella vaginalis clades and sialidase A gene in women with normal microbiota, intermediate microbiota, and abnormal vaginal microbiota characteristic of bacterial vaginosis

| Clade/gene | Frequency (% [95% CI]) | Pearson’s Chi squared test: Chi square statistic; Cramer’s V coefficient (\( p \) value) | Fisher’s exact test: \( OR ( p \) value) |
|------------|------------------------|-----------------------------------------------|----------------------------------|
| All women  | (\( n = 299 \))         | Normal vs. intermediate                        | Normal vs. abnormal |
| Clade 1    | 169 (56 [51–62])        | 31.3; 0.324 (< 0.0001)                         | 0.502 (0.0318) 0.360 (0.0116) 0.180 (0.0001) |
| Clade 2    | 119 (40 [34–45])        | 34.0; 0.337 (< 0.0001)                         | 0.754 (0.4066) 0.261 (0.0002) 0.196 (0.0001) |
| Clade 3    | 59 (20 [16–25])         | 0.73; 0.049 (0.6941)                           | 1.018 (1.0000) 1.322 (0.6550) 1.347 (0.4895) |
| Clade 4    | 282 (94 [91–96])        | 15.2; 0.226 (0.0005)                           | 0.000 ND 0.000 (0.0010) |
| Any clade  | 288 (96 [94–98])        | 9.7; 0.180 (0.0080)                            | 0.000 ND 0.000 (0.0179) |
| Sialidase A| 208 (70 [64–75])        | 23.6; 0.281 (< 0.0001)                         | 0.604 (0.1554) 0.274 (0.0065) 0.165 (0.0001) |

CI confidence interval. ND not determined (because the frequency rates in the groups were the same). \( OR \) odds ratio

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Quantitation of *G. vaginalis* Clades and Sialidase A Gene to Detect Abnormal Vaginal Microbiota

The clade 3 count, as expected, showed no discriminatory ability for BV microbiota (AUC 0.517). The AUC for sialidase A gene count significantly exceeded those for clade 1 and 2, showed no difference with that for clade 4, and was significantly lower than that for total *G. vaginalis* load (Table 3).

### 4 Discussion

Since the four clades of *G. vaginalis* were described, several studies have investigated their association with BV. Balashov et al. [32] developed a real-time PCR assay for detection of the four clades in clinical vaginal samples and found positive correlations between BV and clades 1 and 3, but not clades 2 and 4. The presence of multiple clades, which was detected in 70% of samples, was associated with BV. More recently, it was shown that clades 1 and 2 were significantly more common in BV samples (Nugent score 7–10), whereas clades 3 and 4 had no association with BV, and multi-clade *G. vaginalis* communities were associated with Nugent score ≥ 4 [33]. In a recent cohort study [34], it was demonstrated that young women who developed BV were more likely to have clade 4 than those reporting no sex or who practiced non-coital activities. Furthermore, vaginal samples were more likely to contain multiple *G. vaginalis* clades rather than a single clade if women engaged in penile–vaginal sex or were diagnosed with BV. Hilbert et al. [35], studying *G. vaginalis* population dynamics in BV, revealed a predominance
of clades 1 and 4 in vaginal specimens, and the abundance of clades varied with different Nugent score, Amsel criteria, and success of treatment.

In a previous study by our group [36], which evaluated a PCR test for diagnosis of BV, significant difference between BV and normal vaginal samples was observed for *G. vaginalis* clades 1/2 (the PCR assay did not differentiate between clades 1 and 2), 3, and 4 in both detection rates and concentrations, and the presence of multiple clades was strongly associated with higher *G. vaginalis* loads. In our present study, for detection of the *G. vaginalis* clades we used the PCR assay developed by Balashov et al. [32], for comparability with the previous studies by other groups [32–35].

The most common *G. vaginalis* clade among the Russian women examined in our study was clade 4 (94%), followed by clade 1 (56%), clade 2 (40%), and clade 3 (20%). In concordance with previous studies, although the prevalence figures slightly differed, clades 4 and 1 were the most prevalent [32, 33, 35]. In the present study, *G. vaginalis* clades 1, 2, and 4 were detected significantly more often and in significantly higher loads in women with BV microbiota than in those with normal vaginal microbiota. Clade 3 showed no association with abnormal microbiota.

Our results confirm that clade multiplicity is strongly associated with abnormal microbiota characteristic of BV. The presence of multiple clades in the vagina might be associated with having multiple sexual partners and engaging in unprotected penile–vaginal sex, which has also been previously suggested [32, 33, 35]. It should also be noted that the *G. vaginalis* clade multiplicity is consistent with the overall increased microbial diversity associated with abnormal vaginal microbiota. Another reason can be the biofilm character of BV. It is well-recognized that bacteria in BV are frequently present in biofilms, including densely populated microbial communities, which at least partly explains the significantly higher loads of BV-associated bacteria in the vagina of women with BV than in healthy women. BV biofilm might incorporate different bacteria, including different *G. vaginalis* strains, over time, so it can be hypothesized that biofilm forms of *G. vaginalis* in BV are commonly multi-clade, whereas dispersed forms of the organism in healthy women are single clade. Furthermore, these multi-clade *G. vaginalis* biofilms can be transmitted between sexual partners, as shown by Swidsinski et al. [37], who found cohesive *G. vaginalis* forms in all patients with proven BV and their partners using fluorescence in situ hybridization (FISH) analysis. Studies into *G. vaginalis* forms in normal vaginal and BV samples using clade-specific FISH analysis might help to elucidate this issue.

Sialidase production is considered an important virulence factor in BV and aerobic vaginitis [26, 38], and is associated with adverse pregnancy outcomes [39]. Sialidase is a

| Clade/gene | AUC [95% CI] | DeLong test for difference from AUC = 0.5 (no discrimination line): Z statistic (p value) | Optimal threshold (geqs/mL) | True positive proportion (sensitivity) at the optimal threshold | True negative proportion (specificity) at the optimal threshold | Youden’s index at the optimal threshold | DeLong test for difference from sialidase A gene AUC | Difference [95% CI] | Z statistic (p value) |
|------------|--------------|--------------------------------------------------------------------------------|-----------------------------|-------------------------------------------------|-------------------------------------------------|--------------------------------|-----------------------------|----------------------|----------------------|
| Clade 1    | 0.827 [0.764–0.891] | 10.12 (<0.0001) | 3.4 × 10⁴ | 0.646 | 0.975 | 0.621 | −0.054 [−0.100 to −0.008] | −2.30 (0.0216) |
| Clade 2    | 0.767 [0.700–0.833] | 7.81 (<0.0001) | 6.7 × 10⁴ | 0.481 | 0.981 | 0.462 | −0.115 [−0.172 to −0.058] | −3.94 (<0.0001) |
| Clade 3    | 0.517 [0.463–0.570] | 0.62 (0.2865) | ND | ND | ND | ND | ND | ND |
| Clade 4    | 0.857 [0.797–0.917] | 11.63 (<0.0001) | 7.2 × 10⁴ | 0.759 | 0.744 | 0.704 | −0.024 [−0.094 to 0.046] | −0.67 (0.5031) |
| All clades | 0.933 [0.890–0.976] | 19.77 (<0.0001) | 3.4 × 10⁶ | 0.848 | 0.975 | 0.823 | 0.051 [0.005 to 0.098] | 2.16 (0.0306) |
| Sialidase  | 0.881 [0.824–0.938] | 13.05 (<0.0001) | 2.1 × 10⁶ | 0.734 | 0.981 | 0.716 | | |

*AUC* area under the curve, *CI* confidence interval, *ND* not determined (because of the marker lacking discriminatory ability for abnormal vaginal microbiota indicating bacterial vaginosis)
hydrolytic enzyme that degrades protective mucus barriers cleaving sialic acid from terminal glycans of sialoglycoproteins, which are major constituents of mucus [40]. It has been previously demonstrated that not all G. vaginalis strains, including those possessing the sialidase gene, produce sialidase [18, 20, 27, 29]. In the present study, the G. vaginalis sialidase A gene was detected in 208 (70%) women, and its distribution and abundance were strongly associated with abnormal microbiota. Due to the presence of multiple clades in the majority of samples, it was not possible to evaluate the exact frequency of the sialidase A gene in the strains of each G. vaginalis clade. Nevertheless, our results suggest that most clade 4 strains lack the sialidase A gene. Previous analysis of cultured G. vaginalis strains demonstrated the presence of the G. vaginalis sialidase A gene in all clade 1, 2, and 3 strains and in only 3% of clade 4 strains [29]. Furthermore, in the study by Janulaitiene et al. [33], the G. vaginalis sialidase A gene was detected in all clade 1 and clade 2 isolates, but in none of the clade 4 strains.

Quantification of the loads of the individual G. vaginalis clades and sialidase A gene in the vaginal samples enabled us not only to estimate a difference in those between women with normal microbiota and abnormal microbiota characteristic of BV, but also, using ROC analysis, to investigate if measuring the levels of the G. vaginalis sialidase A gene can differentiate between normal and BV microbiota than measuring the loads of G. vaginalis clades, individually and combined. The AUC value for the sialidase A gene (0.881) significantly exceeded that for clade 1 (0.827) and 2 (0.767), but showed no main difference compared with that for clade 4 (0.857) (clade 3 showed no discriminatory capacity for abnormal microbiota). However, the superior AUC value was obtained for the total G. vaginalis load (0.933), which was significantly higher than that for the G. vaginalis sialidase A gene. Accordingly, the sialidase A gene PCR mostly detects G. vaginalis clades belonging to clades 1, 2, and 3, which are known to contain this gene. However, quantitative detection of all G. vaginalis clades, including clade 4, which appears mostly to lack the sialidase A gene, significantly increased the ability to discriminate between normal and BV microbiota. Thus, in the present study G. vaginalis clade 4 was significantly associated with abnormal vaginal microbiota characteristic of BV. This is in disagreement with the studies by Balashov et al. [32] and Janulaitiene et al. [33], where clade 4 did not relate to BV. This discrepancy can be explained by a higher rate of detection of clade 4 in normal vaginal samples in those studies. Furthermore, in contrast to those studies, we compared not only the distribution but also the abundance of the individual G. vaginalis clades between groups of patients and demonstrated that in the vagina of women with normal microbiota clade 4 was present in significantly lower loads than in women with abnormal microbiota, which was confirmed by the results of our ROC analysis. It is noteworthy that about 10% (eight of 79) of all G. vaginalis-positive samples from women with BV microbiota in our study comprised solely clade 4, with all samples lacking the sialidase A gene. It is not clear whether G. vaginalis clade 4 strains can produce other enzymes with mucinase activity, or if other coexisting BV-associated bacteria possessing sialidase activity, e.g., Prevotella spp., Porphyromonas spp., and Bacteroides spp. [27, 41], can degrade mucin. In the present study, the specimens were not analyzed for phenotypic sialidase activity, which limits the interpretation of our results. Harwich et al. [14] suggested that there is no direct link of sialidase production in G. vaginalis with biofilm formation or BV, because both of two investigated G. vaginalis BV isolates, one sialidase negative and one sialidase positive, had a significantly increased biofilm-forming capacity, aggregation, and epithelial cell adherence compared with the non-BV sialidase-negative isolate. This suggestion was recently confirmed by Janulaitiene et al. [42], who found significant difference between G. vaginalis clades 1, 2, and 4 in sialidase activity (clade 2 had the highest, clade 1 intermediate, and clade 4 the lowest sialidase activity) but no significant difference between the clades in biofilm-forming ability.

Another limitation of our study is that no data on clinical manifestations specific for BV were available. It should be emphasized that the Nugent scores applied in this study, as well as other laboratory methods for BV diagnosis, reveal abnormal vaginal microbiota, which can be asymptomatic or manifest as the clinical syndrome BV. It cannot be excluded that there were no main symptoms of BV in the women comprising solely sialidase-free G. vaginalis strains. Future studies are warranted to verify associations between the presence of the G. vaginalis sialidase A gene, sialidase production, biofilm formation, clinical signs of BV, and BV recurrence.

It should also be noted that we used the combined loads of the four G. vaginalis clades as a proxy for total G. vaginalis load. This assumption was based on the data of Balashov et al. [32], who validated their clade-specific PCR using 39 cultured G. vaginalis strains and 60 vaginal samples in comparison with three G. vaginalis species-specific PCRs, which targeted the tuf gene [32], the 16S ribosomal RNA (rRNA) gene [43], and the cpn60 gene [44]. All G. vaginalis strains were positive by the PCRs targeting the tuf and 16S rRNA genes, and all but one of the G. vaginalis strains were successfully typed by the clade-specific PCR. The cpn60 PCR missed all strains (n = 14) typed as clades 2 or 4 by the clade-specific PCR. Of the 60 clinical vaginal samples, 59 were positive for G. vaginalis by both the tuf PCR and clade-specific PCR, 58 samples were positive by the 16S rRNA PCR, and 44 samples were positive by the cpn60 PCR. These results suggest that the clade-specific PCR detects the large majority of G. vaginalis-positive samples. Nevertheless, there are data on a limited sensitivity of this
clade-specific PCR for the detection of clade 2 and 3 (cpn60 subgroup B and D, respectively) [29]. It is possible that there are molecular variants within the subgroups or additional G. vaginalis subgroups to the four that have already been described. Ideally, only thoroughly validated G. vaginalis PCRs proven to detect all G. vaginalis clades/subgroups/variants should be used for diagnostic purposes.

5 Conclusions

The findings of the present study indicate that the detection of all four G. vaginalis clades discriminates between normal vaginal microbiota and abnormal vaginal microbiota characteristic of BV more accurately than measuring G. vaginalis sialidase A gene count, and there is a strong association of G. vaginalis clade 4 with BV microbiota, although most of the clade 4 strains appear to lack the sialidase A gene. Our results support previous observations that G. vaginalis clades 4 and 1 are the most prevalent in vaginal samples, and clade multiplicity is strongly associated with BV microbiota.

Compliance with Ethical Standards

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Conflict of interest Elena Shipitsyna, Anna Krysanova, Guzel Khayrullina, Kira Shalepo, Alekstina Savicheva, Alexander Guschin, and Magnus Unemo declare that they have no conflicts of interest that are directly relevant to the content of this study.

Ethical approval The study was approved by the Ethical Committee at the D.O. Ott Research Institute of Obstetrics, Gynaecology and Reproductology, St Petersburg (approval number 73/2015).

Informed consent Informed consent was obtained from all individual participants included in the study.

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References

1. Koumans EH, Sternberg M, Bruce C, McQuillan G, Kendrick J, Sutton M, et al. The prevalence of bacterial vaginosis in the United States, 2001–2004; associations with symptoms, sexual behaviors, and reproductive health. Sex Transm Dis. 2007;34:864–9.
2. Taylor BD, Darville T, Haggerty CL. Does bacterial vaginosis cause pelvic inflammatory disease? Sex Transm Dis. 2013;40:117–22.
3. Allsworth JE, Peipert JF. Severity of bacterial vaginosis and the risk of sexually transmitted infection. Am J Obstet Gynecol. 2011;205(113):e1–6.
4. Donati L, Di Vico A, Nucci M, Quaggiozzi L, Spagnuolo T, Labianca A, et al. Vaginal microbial flora and outcome of pregnancy. Arch Gynecol Obstet. 2010;281:589–600.
5. Denney JM, Culhane JF. Bacterial vaginosis: a problematic infection from both a perinatal and neonatal perspective. Semin Fetal Neonatal Med. 2009;14:200–3.
6. Shipitsyna E, Roos A, Datcu R, Hallén A, Fredlund H, Jensen JS, et al. Composition of the vaginal microbiota in women of reproductive age-sensitive and specific molecular diagnosis of bacterial vaginosis is possible? PLoS One. 2013;8:e60670.
7. Fredricks DN, Fiedler TL, Thomas KK, Oakley BB, Marrazzo JM. Targeted PCR for detection of vaginal bacteria associated with bacterial vaginosis. J Clin Microbiol. 2007;45:3270–6.
8. Alves P, Castro J, Sousa C, Cerejeira TB, Cerca N. Gardnerella vaginalis outcompetes 29 other bacterial species isolated from BV patients in an in vitro biofilm formation model. J Infect Dis. 2014;210:593–6.
9. Schwabe JR, Muzny C, Josey WE. Role of Gardnerella vaginalis in the pathogenesis of bacterial vaginosis: a conceptual model. J Infect Dis. 2014;210:338–43.
10. Patterson JL, Stull-Lane A, Girerd PH, Jefferson KK. Analysis of adherence, biofilm formation and cytotoxicity suggests a greater virulence potential of Gardnerella vaginalis relative to other bacterial-vaginosis-associated anaerobes. Microbiology. 2010;156:392–9.
11. Piot P, Van Dyck E, Peeters M, Hale J, Totten PA, Holmes KK. Biotypes of Gardnerella vaginalis. J Clin Microbiol. 1984;20:677–9.
12. Benito R, Vazquez JA, Berron S, Fenoll A, Saez-Neito JA. A modified scheme for biotyping Gardnerella vaginalis. J Med Microbiol. 1986;21:357–9.
13. Ingianti A, Petruzelli S, Morandotti G, Pompei R. Genotypic differentiation of Gardnerella vaginalis by amplified ribosomal DNA restriction analysis (ARDRA). FEMS Immunol Med Microbiol. 1997;18:61–6.
14. Harwich MD, Alves JM, Buck GA, Strauss JF, Patterson JL, Oki AT, et al. Drawing the line between commensal and pathogenic Gardnerella vaginalis through genome analysis and virulence studies. BMC Genom. 2010;11:375.
15. Yeoman CJ, Yildirim S, Thomas SM, Durkin AS, Torralba M, Sutton G, et al. Comparative genomics of Gardnerella vaginalis strains reveals substantial differences in metabolic and virulence potential. PLoS One. 2010;5:e12411.
16. Ahmed A, Earl J, Retchless A, Hillier SL, Rabe LK, Cherpes TL, et al. Comparative genomic analyses of 17 clinical isolates of Gardnerella vaginalis provide evidence of multiple genetically isolated clades consistent with subspeciation into genovars. J Bacteriol. 2012;194:3922–37.
17. Paramel Jayaprakash T, Schellenberg JJ, Hill JE. Resolution and characterization of distinct cpn60-based subgroups of Gardnerella vaginalis in the vaginal microbiota. PLoS One. 2012;7:e43009.
18. Santiago GL, Deschaght P, El Aila N, Kiama TN, Verstraelen H, Jefferson KK, et al. Gardnerella vaginalis comprises three distinct genotypes of which only two produce sialidase. Am J Obstet Gynecol. 2011;204(450):e451–7.
19. Udayalaxmi J, Bhat GK, Kotigadde S. Biotypes and virulence factors of Gardnerella vaginalis isolated from cases of bacterial vaginosis. Indian J Med Microbiol. 2011;29:165–8.
Quantitation of G. vaginalis Clades and Sialidase A Gene to Detect Abnormal Vaginal Microbiota

20. Pleckaityte M, Janulaitiene M, Lasickiene R, Zvyrbiene A. Genetic and biochemical diversity of Gardnerella vaginalis strains isolated from women with bacterial vaginosis. FEMS Immunol Med Microbiol. 2012;65:69–77.

21. Castro J, Alves P, Sousa C, Cereia T, França Â, Jefferson KK, et al. Using an in vitro biofilm model to assess the virulence potential of bacterial vaginosis or nonbacterial vaginosis Gardnerella vaginalis isolates. Sci Rep. 2015;5:11640.

22. Schuyler JA, Mordechai E, Adelson ME, Sobel JD, Gygax SE, Hillert DW. Identification of intrinsically metronidazole-resistant clades of Gardnerella vaginalis. Diagn Microbiol Infect Dis. 2016;84:1–3.

23. Briselden AM, Hillier SL. Longitudinal study of the biotypes of Gardnerella vaginalis. J Clin Microbiol. 1990;28:2761–4.

24. Numanovic F, Hukić M, Nurkić M, Grgić M, Delibegović Z, Imamović A, et al. Importance of isolation and biotyping of Gardnerella vaginalis in diagnosis of bacterial vaginosis. Bosn J Basic Med Sci. 2008;8:270–6.

25. Aroutcheva AA, Simoes JA, Behbakh K, Faro S. Gardnerella vaginalis isolated from patients with bacterial vaginosis and from patients with healthy vaginal ecosystems. Clin Infect Dis. 2001;33:1022–7.

26. Lewis WG, Robinson LS, Gilbert NM, Perry JC, Lewis AL. Degradation, foraging, and depletion of mucus sialoglycans by the vagina-adapted Actinobacterium Gardnerella vaginalis. J Biol Chem. 2013;288:12067–79.

27. Briselden AM, Moncla BJ, Stevens CE, Hillier SL. Sialidases (neuraminidases) in bacterial vaginosis and bacterial vaginosis-associated microflora. J Clin Microbiol. 1992;30:663–6.

28. Bradshaw CS, Morton AN, Garland SM, Horvath LB, Kuzevska I, Fairley CK. Evaluation of a point-of-care test, BVBlue, and clinical and laboratory criteria for diagnosis of bacterial vaginosis. J Clin Microbiol. 2005;43:1304–8.

29. Schellenberg JJ, Paramel Jayaparakash T, Withana Gamage N, Patterson MH, Vaneechouette M, Hill JE. Gardnerella vaginalis subgroups defined by cpn60 sequencing and sialidase activity in isolates from Canada, Belgium and Kenya. PLoS One. 2016;11:e0146510.

30. Hardy L, Jespers V, Van den Bulck M, Buyse J, Mwamba-rangwe L, Musengamana V, et al. The presence of the putative Gardnerella vaginalis sialidase A gene in vaginal specimens is associated with bacterial vaginosis biofilm. PLoS One. 2017;12:e0172522.

31. Nugent RP, Krohn MA, Hillier SL. Reliability of diagnosing bacterial vaginosis is improved by a standardized method of gram stain interpretation. J Clin Microbiol. 1991;29:297–301.

32. Balashov SV, Mordechai E, Adelson ME, Gygax SE. Identification, quantification and subtyping of Gardnerella vaginalis in noncultured clinical vaginal samples by quantitative PCR. J Med Microbiol. 2014;63(Pt 2):162–75.

33. Janulaitiene M, Paliiyte V, Grinceviciene S, Zakareviciene J, Vladisauskienė A, Marcinkute A, et al. Prevalence and distribution of Gardnerella vaginalis subgroups in women with and without bacterial vaginosis. BMC Infect Dis. 2017;17:394.

34. Vodstrcil LA, Twin J, Garland SM, Fairley CK, Hocking JS, Law MG, et al. The influence of sexual activity on the vaginal microbiota and Gardnerella vaginalis clade diversity in young women. PLoS One. 2017;12:e0171856.

35. Hilbert DW, Schuyler JA, Adelson ME, Mordechai E, Sobel JD, Gygax SE. Gardnerella vaginalis population dynamics in bacterial vaginosis. Eur J Clin Microbiol Infect Dis. 2017;36:1269–78.

36. Rumyantseva T, Shiptsyna E, Guschin A, Unemo M. Evaluation and subsequent optimizations of the quantitative AmpliSens flo-roacenosis/bacterial vaginosis-FRT multiplex real-time PCR assay for diagnosis of bacterial vaginosis. APIMIS. 2016;124:1099–108.

37. Swidsinski A, Doerfler Y, Loening-Baucke V, Swidsinski S, Vrastraelen H, Vaneechouette M, et al. Gardnerella biofilm involves females and males and is transmitted sexually. Gynecol Obstet Investig. 2010;70:256–63.

38. Marconi C, Donders GG, Bellen G, Brown DR, Parada CM, Silva MG. Sialidase activity in aerobic vaginitis is equal to levels during bacterial vaginosis. Eur J Obstet Gynecol Reprod Biol. 2013;167:205–9.

39. Caucci S, Cullane JF. High sialidase levels increase preterm birth risk among women who are bacterial vaginosis-positive in early gestation. Am J Obstet Gynecol. 2011;204:e1–9.

40. Lewis AL, Lewis WG. Host sialoglycans and bacterial sialidases. A mucosal perspective. Cell Microbiol. 2012;14:1174–82.

41. Moncla BJ, Braham P, Hillier SL. Sialidase (neuraminidase) activity among gram-negative anaerobic and capnophilic bacteria. J Clin Microbiol. 1990;28:422–5.

42. Janulaitiene M, Paliulyte V, Baranauskiene L, Bulavaite A, Simanavicius M, Pleckaityte M. Phenotypic characterization of Gardnerella vaginalis subgroups suggests differences in their virulence potential. PLoS One. 2018;13:e0200625.

43. Fredricks DN, Fiedler TL, Thomas KK, Mitchell CM, Marrazzo JM. Changes in vaginal bacterial concentrations with intravaginal metronidazole therapy for bacterial vaginosis as assessed by quantitative PCR. J Clin Microbiol. 2009;47:721–6.

44. Menard JP, Fenollar F, Henry M, Bretelle F, Raoult D. Molecular quantification of Gardnerella vaginalis and Atopobium vaginae loads to predict bacterial vaginosis. Clin Infect Dis. 2008;47:33–43.

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