Increased Pathogen Identification in Vascular Graft Infections by the Combined Use of Tissue Cultures and 16S rRNA Gene Polymerase Chain Reaction

Evelyne Ajdler-Schaeffler 1†, Alexandra U. Scherrer 1†, Peter M. Keller 2, Alexia Anagnostopoulos 1, Michael Hofmann 1, Zoran Rancic 3, Annelies S. Zinkernagel 1, Guido V. Bloemberg 2, Barbara K. Hasse 1* and the VASGRA Cohort

1 Division of Infectious Diseases and Hospital Epidemiology, University Hospital and University of Zurich, Zurich, Switzerland,
2 Institute of Medical Microbiology, University of Zurich, Zurich, Switzerland,
3 Clinic for Cardiovascular Surgery, University Hospital and University of Zurich, Zurich, Switzerland

Background: Vascular graft infections (VGI) are difficult to diagnose and treat, and despite redo surgery combined with antimicrobial treatment, outcomes are often poor. VGI diagnosis is based on a combination of clinical, radiological, laboratory and microbiological criteria. However, as many of the VGI patients are already under antimicrobial treatment at the time of redo surgery, microbiological identification is often difficult and bacterial cultures often remain negative rendering targeted treatment impossible. We aimed to assess the benefit of 16S rRNA gene polymerase chain reaction (broad-range PCR) for better microbiological identification in patients with VGI.

Methods: We prospectively analyzed the clinical, microbiological, and treatment data of patients enrolled in the observational Vascular Graft Cohort Study (VASGRA), University Hospital Zurich, Switzerland. The routine diagnostic work-up involved microbiological cultures of minced tissue samples, and the use of molecular techniques in parallel. Patient-related and microbiological data were assessed in descriptive analyses, and we calculated sensitivity, specificity, negative and positive predictive value for broad-range 16S rRNA gene PCR versus culture (considered as gold standard).

Results: We investigated 60 patients (median age 66 years (Interquartile range [IQR] 59–75)) with confirmed VGI between May 2013 and July 2017. The prevalence of antimicrobial pretreatment at the time of sampling was high [91%; median days of antibiotics 7 days (IQR 1–18)]. We investigated 226 microbiological specimens. Thereof, 176 (78%) were culture-negative and 50 (22%) were culture-positive. There was a concordance of 70% (158/226) between conventional culture and broad-range PCR (sensitivity 58% (95% CI 43–72); specificity 74% (67–80%)). Among the group of 176 culture-negative specimens, 46 specimens were broad-range PCR-positive resulting in identification of overall 69 species. Among the culture and/or broad-range PCR-positive specimens (n = 96), 74 (77%) were monomicrobial and 22 (23%) polymicrobial, whereas the rate of polymicrobial samples was higher in broad-range PCR-positive specimens (93%).
INTRODUCTION

Vascular graft infections (VGIs) are among the most serious complications in vascular surgery (1), increasing the risk of morbidity and mortality in affected patients substantially. Extravascular groin or limb graft infections occur in about 2–6%, whereas intravascular abdominal or thoracic graft infections are less common (0.5–2%). Vascular surgical site infections are combining the clinical features of biofilm-mediated and chronic wound infections due to both, the implantation of foreign material and circulatory disturbances. Therefore, therapy strategies solely relying on antimicrobial treatment oftentimes fail, and redo-surgery is needed. Traditional surgical salvage procedures include complete or partial graft removal with in-situ or extra-anatomical reconstruction and local treatment with muscle flap, omentum, or negative pressure wound therapy (NWPT) (1, 2).

The broadly used definition of VGI is based on clinical, radiological, and laboratory criteria (3–5). Even though histopathologic work-up is routinely recommended, no standardized cut-off levels for polymorphonuclear neutrophil counts per high power field have been defined. Microbiological evidence of pathogens remains the cornerstone of diagnosis, and the detection of microorganisms in blood cultures and/or the extraction of bacteria from the periprosthetic tissue biofilm by tissue homogenization are essential for diagnosis. However, blood cultures have a low sensitivity in detecting pathogens causing a VGI, since biofilm formation on the implant material oftentimes precludes planktonic seeding of bacteria and therefore their detection in blood (4). Cultures from the prosthetic sonicate fluid have only rarely been validated for VGIs (6), especially due to the lack of standardization caused by high rates of polymicrobial infections and different involved anatomical regions and graft materials. Broad-range 16S rRNA gene polymerase chain reaction (broad-range PCR) has not been established in the routine diagnostic work-up of VGIs yet. These molecular genetic techniques (7) are providing sensitive methods for more accurate pathogen identification, especially in case of infections with antimicrobial pretreatment and fastidious to grow or non-cultivable pathogens like Coxiella burnetii, Tropheryma whippelii, and Mycobacteria (8–10). The method bears the risk of sample contamination and is allegedly less effective in case of polymicrobial infections. It is estimated that up to one third of all VGI are polymicrobial, which is potentially affecting the sensitivity of broad-range PCR and therefore its effectiveness in the reliable identification of causative pathogens. We aimed to assess whether broad-range PCR in addition to conventional microbiological cultures would improve diagnostics of VGIs of patients included in the prospective, observational Vascular Graft Cohort Study (VASGRA).

MATERIALS AND METHODS

Study Design and Study Population

All participants of VASGRA with proven VGIs between May 2013 and July 2017 were eligible for this analysis. In the VASGRA cohort clinical, laboratory and microbiological information is collected at three-monthly follow-up visits. The prospective, observational cohort is located at the University hospital of Zurich (Switzerland), a 880-bed tertiary-care referral center. We obtained written informed consent from all participants. The Independent Ethics Committee of Zurich approved the study (KEK_2012-0583).

Definitions and Classifications

We used the Management of Aortic Graft Infection Collaboration (MAGIC) criteria (5) for the diagnosis of VGI. A multidisciplinary team of infectious disease specialists, microbiologists, cardiovascular surgeons and radiologists adjudicated each criterion. In addition, each VGI was additionally classified according to the Samson classification (11). Extravascular VGIs mainly included the groin and/or the lower extremities, whereas intravascular VGIs involved the abdomen or less commonly the thorax (1).

Microbiological Analyses

Clinical specimens from suspected VGIs were sent to the Institute for Medical Microbiology, Zurich, Switzerland for conventional microbiological culture as well as bacterial broad-range PCR. For strict pathogens, we considered the microbiologic criterion as fulfilled when there was one positive culture and/or a positive broad-range PCR from an intraoperative specimen. For potential “contaminant” pathogens (e.g., coagulase-negative Staphylococci, Corynebacterium spp., Propionibacterium acnes), at least two positive intraoperative specimens were required.

Microscopy

We performed Gram stains of clinical specimens according to standard procedures. Ten 1,000-fold visual fields were used to

Conclusions: Combined cultures and broad-range 16S rRNA gene PCR from periprosthetic tissue and/or explanted vascular grafts increased the diagnostic accuracy in VGI, particularly in patients already under antimicrobial treatment at the time of redo surgery. Ideally, antimicrobial treatment should be withheld until surgical sampling in order to optimize microbiological diagnostics.

Clinical trials.gov identifier: NCT01821664

Keywords: 16S rRNA gene polymerase chain reaction, vascular graft infection, pathogen identification, microbiological diagnosis, tissue culture
determine the mean bacterial count (no bacteria; detection of bacteria after enrichment; detection of some scattered bacteria; detection of bacteria); and ten 100-fold visual fields were used to determine the mean count for leucocytes (no leucocytes, some scattered amounts of leucocytes, few leucocytes, leucocytes in lumps).

**Cultures**

Tissue specimens were placed in sterile 0.9% sodium chloride solution in the operating theater and immediately transferred to the laboratory. Upon receipt, the specimens were minced using an IKA Ultra-Turrax tissue homogenizer (Sigma-Aldrich, Buchs, Switzerland) and culture media were inoculated and incubated under aerobic and anaerobic conditions. Sheep agar plates without antibiotics (COS; bioMérieux, Marcy l’Etoile, France), and with colistin-nalidixic acid (CNA) were used for the detection of Gram-positive bacteria (bioMérieux), while MacConkey (MCK) agar plates (bioMérieux) were used for Gram-negative bacteria and Sabouraud agar plates for fungi. In addition, thioglycolate broth, aerobic selective chocolate agar plates (HAE2; bioMérieux), anaerobic sheep blood agar plates with hemin and vitamin K1 (Brucella agar; BD), laked sheep blood Brucella agar plates with kanamycin and vancomycin (BD), and phenylethylalcohol agar plates with vitamin K1 (BD) were used. Growth on agar plates was examined at 24, 48, and 72 h as well as after one week; the thioglycolate broth was examined up to day 10 and the Sabouraud agar plates up to day 20 (12). Antibiotic susceptibility testing was performed according to EUCAST recommendations (13).

**Bacterial and Fungal Identification**

16S broad-range PCR was performed as a conventional, semi nested PCR using three amplification primers as described before (14, 15). PCR amplification was assessed optically on silver-stained polyacrylamide gel and compared to the buffer background as described by Rampini et al. (15). Reagents and buffers were routinely controlled to be low bacterial DNA-containing prior to use. Broad-range fungal inter spacer region (ITS) polymerase chain reaction (PCR) for the detection and identification of fungi in clinical specimens was performed as described before (16).

We used matrix-assisted laser desorption ionization–time of flight (MALDI-TOF) and 16S rRNA gene or ITS sequencing for identification (7, 14, 16–18). 16S rRNA sequence homology analyses were done using GenBank (NCBI) and the SmartGene IDNS database and software (SmartGene GmbH, Zug, Switzerland).

**Statistical Analysis**

To analyze the patient characteristics and the characteristics of the VGI, we used all available diagnostic samples (performed in internal or external laboratories). For the further analysis, we restricted the analysis to patients where i.a. deep or superficial wound tissue specimens, NPWT foams or explanted vascular grafts were investigated in parallel with broad-range PCR and conventional microbiological cultures. Samples with only broad-range PCR or microbiological cultures were excluded. We present results as medians with interquartile range (IQR), crude numbers, percentages and 95% confidence intervals (CI) of specimens/species. We compared sensitivity, specificity, positive predictive value and negative predictive value of 16S rRNA gene broad-range PCR and conventional culture (considered as gold standard) (19). We performed statistical analyses with Stata (version 15.1/SE; Stata Corporation, College Station, Texas, USA).

**RESULTS**

**Patient Characteristics**

The VASGRA dataset included information on 508 participants. We excluded 387 control patients because they underwent routine vascular surgery without infection. Of the remaining 121 patients with suspected VGI, 61 patients were excluded due to unconfirmed VGIs (n = 30) or due to missing parallel sampling of intraoperative tissue (n = 31) (Figure 1). Considering also patients with positive blood cultures (n = 17) we finally included 60 prospective patients with proven VGIs (82% men) in the analysis (Table 1). The median age, body mass index (BMI) and Charlson Comorbidity Index of participants were 66 years (IQR 58.5–75), 28 kg/m² (25–30), and 2 (1–3), respectively. VGIs occurred after a median of 3 months (IQR 1–22) after the index surgery.

**Microbiological Specimens**

We finally investigated 226 intraoperative tissue samples in which both, tissue cultures and broad-range PCR were performed in parallel (Table 1, Figure 2). The tissue samples originated from extracavitary (22%), intracavitary abdominal (55%), or thoracic (20%) VGIs. The remaining tissue samples (3%) were isolated from secondary VGIs due to endovascular treatment of mycotic aneurysms. Microbiological samples were obtained after a median of 25 days (9–91) after diagnosis of VGI. At the time of sampling, 91% of patients were under antimicrobial treatment for a median of 7.5 days (IQR 1–18.5).


### TABLE 1 | Characteristics of patients and samples.

| Characteristic                                    | Patient characteristics |
|--------------------------------------------------|-------------------------|
| Male, n (%)                                      | 49 (100)                |
| Age in years at infection, median (IQR)          | 66 (58–75)              |
| Body mass index (kg/m²), median (IQR)            | 28 (25–30)              |
| Charlson Comorbidity Index, median (IQR)         | 2 (1–3)                 |
| Months since index surgery, median (IQR)         | 3 (1–22)                |
| Days since graft infection, median (IQR)         | 6 (0–22)                |
| Localization of vascular graft                   |                         |
| Abdominal intracavitary, n (%)                   | 33 (55)                 |
| Thoracic intracavitary, n (%)                    | 11 (20)                 |
| Extracavitary, n (%)                             | 13 (22)                 |
| Other, n (%)                                     | 3 (3)                   |
| Samson Classification                            |                         |
| 3, n (%)                                         | 29 (49)                 |
| 4, n (%)                                         | 15 (25)                 |
| 5, n (%)                                         | 16 (28)                 |
| Specimens, n (%)                                 | 226 (100)               |
| Number of specimens per patient, median (IQR)    | 3 (1–4)                 |
| Material, n (%)                                  |                         |
| Deep wound, n (%)                                | 156 (69)                |
| Superficial wound, n (%)                         | 30 (13)                 |
| Vascular graft, n (%)                            | 6 (3)                   |
| Other (e.g., biopsy, blood culture), n (%)       | 20 (9)                  |
| NPWT foam, n (%)                                 | 14 (6)                  |
| Antibiotic treatment at sampling, n (%)          |                         |
| Yes, n (%)                                       | 206 (91)                |
| Culture-positive on antimicrobial treatment, n (%)| 40/206 (19)             |
| PCR-positive on antimicrobial treatment, n (%)    | 71/206 (34)             |
| Days on continuous antimicrobial treatment, Median (IQR) | 7 (1–18) |

IQR, Interquartile range; NWPT, Negative pressure wound therapy.

### Performance of Broad-Range PCR as Compared to Intraoperative Tissue Cultures

Among the 226 intraoperative specimens, 176 (78%) were culture-negative and 50 (22%) were culture-positive (Figures 2, 3). Among the 176 culture-negative samples, 46 were broad-range PCR-positive (≥2 organisms 20%). Among the 50 culture-positive samples (≥2 organisms 26%), 21 were broad-range PCR-negative and 29 broad-range PCR-positive (≥2 organisms 38%). Among the latter, 29 specimens, 11/29 (38%) yielded a non-identical species. Taken together, there was a concordance of 70% (159/226) between conventional culture and broad-range PCR. Compared to microbiological cultures as the gold standard, broad-range PCR had a sensitivity of 58% (95% CI 43–72), a specificity of 74% (67–80), a positive predictive value (PPV) of 39% (28–51), and a negative predictive value (NPV) of 86% (80–91). Combining culture and molecular diagnostics, the number of positive microbiological results almost doubled rising from 50/226 (22%) to 96/226 (42%). We did a sub-analysis were we ignored the presence of Candida sp, because panfungal PCR was not done for all samples. The diagnostic accuracy did not change markedly [sensitivity: 58% (39–75), specificity 76% (70–82), PPV 29% (19–42), NPV 91% (86–95)].

Among the culture and/or broad-range PCR-positive specimens (n = 96), 74 (77%) were monomicrobial and 22 (23%) polymicrobial. We detected in 14% two species per sample, and >3 species in 9%. The polymicrobial specimens were culture- and broad-range PCR-positive in 11 cases, culture-positive and broad-range PCR-negative in two cases and culture-negative, broad-range PCR-positive-positive in nine cases.

### Detection of Intraoperative Species by Broad-Range PCR as Compared to Conventional Cultures

The detected species in the culture and/or broad-range PCR-positive specimens were not always identical. Details on the diagnostics of the detected species are presented in Table 2 and Supplementary Table 1. Of all bacterial species, Gram-positive bacteria clearly predominated (44%) compared to Gram-negative bacteria (28%), fungi (6%), anaerobes (8%), and miscellaneous other bacteria (14%). Among all detected species, 22 (17%) were culture- and broad-range PCR-positive, 40 (30%) were culture-positive/broad-range PCR-negative and 69 (53%) were culture-negative/broad-range PCR-positive.

Culture- and broad-range PCR-positive species: Among the 22 species with positive results in both methods, *Candida* sp. (45%) and *S. epidermidis* (23%) predominated. At the time of sampling, 21/22 patients were on antimicrobial treatment for a median of 6 days (IQR 3–6).

Culture-positive/broad-range PCR-negative species: The 40 species with only positive cultures, mainly included *Candida* sp. (20%), *S. epidermidis* (13%) and *Enterococcus* spp. (15%). As the bacterial load was relatively low, no bacteria were detected (or only bacteria after enrichment) in 85% (Figure 4A). Leucocytes were detected in some scattered amounts, in a few amounts or in lumps in 35, 12.5, and 12.5%, respectively (Figure 4B).

Culture-negative/broad-range PCR-positive species: Among the 69 broad-range PCR-negative/ culture-negative species, *S. epidermidis and Streptococcus* spp. were detected in 33%, whereas Gram-negative bacteria, anaerobes and mycobacteria were detected in 38, 4, and 7% of the remaining isolates, respectively. No bacteria per visual field were detected in 88% of these instances (Figure 4A), and no or only scattered amounts of leucocyte count were detected by visual field in 62% (Figure 4B).

### DISCUSSION

Combining culture-based and molecular diagnostics increased the microbiological yield in a substantial proportion of patients with VGs. Our study is the first prospective cohort study to explore the diagnostic performance of broad-range PCR for a large number of patients. Our study showed that intraoperative tissue-/explanted vascular graft sequencing combined with culture showed more sensitivity to identify the causative
pathogen as compared to intraoperative tissue cultures alone. As many of the patients were under antimicrobial treatment at the time of sampling, the rate of culture-negative tissue samples was as high as 78%. In addition, potential contaminant bacterial pathogens were taken into account. The specificity (74%) of broad-range PCR was better than its sensitivity (58%).

To date, microbiological results of pre- or intraoperatively obtained cultures from different tissue samples have only rarely been assessed in VGIs (20–22). We previously analyzed the diagnostic value of bacteria cultured from NPWT-foams as compared to bacteria obtained from deep tissue perivascular samples and found that sensitivity (58%) as well as specificity (86%) from NPWT-foams were poor (20). Another study assessed the diagnostic accuracy of blood cultures in identifying the causative pathogen among 17 patients with VGIs (22). Bisharat et al. found that microorganisms in blood cultures do not necessarily indicate a causal relationship as only 17.6% of these patients had the same microorganisms in blood cultures, excised vascular grafts, or perigraft tissue cultures, respectively.

The use of molecular techniques for the diagnosis of VGI has been suggested in the past (3, 23). However, only one retrospective study has assessed the diagnostic accuracy of broad-range PCR and sonicate-fluid cultures as compared to perivascular deep tissue and/or blood cultures among 22 patients (24 cases) with VGIs (6). The study found high rates of sonicate-fluid cultures (79%) and broad-range 16S rRNA PCR-positivity (67%). However, blood cultures, intraoperative tissue specimens and parallel PCR/deep tissue cultures were not taken routinely, therefore preventing adequate statistical analyses and exclusion of potential contaminant pathogens. In detailed analyses, we found that broad-range PCR results contributed to the diagnosis of VGI in 21/34 (62%) patients with culture-negative infections.

Comparing perivascular tissue cultures (current gold standard) and broad-range PCR, we observed a lack of sensitivity of the molecular diagnosis. Indeed, broad-range PCR was not confirmatory in molecular species identification in 40 out of 61 instances with a positive culture result. Additionally, cultures were negative in 176 cases with 46 broad-range PCR-positive samplings. We rated 14 of 69 (16%) species found in broad-range PCR-positive and culture-negative samplings as clinically irrelevant (single detection of a “contaminant” bacterial
pathogen such as \textit{S. epidermidis}, \textit{P. acnes}, \textit{Corynebacterium} spp.). Low bacterial inoculum and small amounts of bacterial DNA substantiated our decision. The mismatch of 30\% comparing broad-range PCR and culture is consistent with recent studies (6, 15).

We found remarkably high rates (176 specimens) of negative microbiological results among samples, obtained after preoperative therapeutic administration of antimicrobial agents. Many patients with VGIs receive antimicrobial treatment prior to hospital admission thus complicating the microbiological diagnosis. Broad-range PCR from specimens with (n = 206) and without antimicrobial pretreatment (n = 20) contributed to a microbiological diagnosis in 19 and 50\% of culture-negative VGI, respectively (Supplementary Table 2). The rate of culture-negative findings was higher than in previously reported studies (6) and from studies on orthopedic infections (24, 25) that combined diagnostics with PCR and wound culture was more sensitive than deep wound cultures alone.

The high prevalence of Gram-positive cocci (\textit{S. epidermidis}, \textit{Streptococcus} spp., \textit{Enterococcus} spp. and \textit{Corynebacterium} spp.) has been reported before (3, 4, 21, 26). In contrast to our expectations (6, 26, 27), we found only limited numbers of \textit{S. aureus}, whereas Gram-negative pathogens, anaerobes and fungi were predominantly found in intracavitary abdominal or extracavitary graft infections consistent with previous reports (1).

This study has several strengths. To our knowledge, it represents the most comprehensive prospective assessment of the use of molecular microbiological methods in VGIs to date. Additionally, molecular and culture methods were obtained in parallel in order to preclude false-positive results by contaminant pathogens (23). Moreover, a multi-disciplinary team of ID specialists, cardiovascular surgeons, microbiologists, and nuclear medicine specialists adjudicated each VGI's diagnosis. Some limitations should also be noted. Molecular approaches surveying bacterial diversity provide a less biased depiction of skin microbiota than culture-based assays, but molecular approaches currently on use are unable to distinguish between 16S rRNA genes that are derived from living versus dead bacteria. Furthermore sequencing of 16S rRNA genes does not provide information regarding the gene content of plasmids. However, DNA extraction of complex microbial communities is highly influenced by the quality of the recovered DNA. Molecular techniques are cost-intensive, bear the risk of sample contamination and are allegedly less effective in case of polymicrobial infections. Ideally, samples from suspected VGIs

### Table 2

Overview on the detected species obtained during surgical revisions among 60 patients with confirmed vascular graft infections.

| Species in culture-positive, PCR-positive species \(^a\) | n  | (%) | Species in culture-positive, PCR-negative species \(^b\) | n  | (%) | Species in culture-negative, PCR-positive species \(^b\) | n  | (%) |
|-----------------------------------------------|----|-----|-------------------------------------------------|----|-----|-------------------------------------------------|----|-----|
| Candida sp.                                   | 10 | (45)| Candida sp.                                     | 8  | (20)| Staphylococcus epidemidis                      | 21 | (30)|
| \textit{Staphylococcus} epidemidis           | 5  | (23)| Enterococcus spp.                               | 6  | (15)| \textit{Streptococcus} spp.                    | 9  | (13)|
| \textit{Streptococcus} spp.                   | 2  | (9 )| \textit{Propionibacterium} acnes                | 5  | (12)| \textit{Enterobacter} spp.                     | 4  | (6 )|
| \textit{Enterobacter} spp.                    | 1  | (5 )| \textit{Staphylococcus} epidemidis              | 5  | (12)| \textit{Corynebacterium} spp.                  | 4  | (6 )|
| \textit{Propionibacterium} acnes              | 1  | (5 )| \textit{Enterobacter} spp.                      | 2  | (5 )| \textit{Klebsiella} spp.                       | 4  | (6 )|
| \textit{Pseudomonas aeruginosa}               | 1  | (5 )| \textit{Eikenella condronis}                    | 2  | (5 )| \textit{Citrobacter freundii}                  | 3  | (4 )|
| \textit{Streptococcus} spp.                   | 1  | (5 )| \textit{Escherichia coli}                       | 2  | (5 )| \textit{Pseudomonas aeruginosa}                | 3  | (4 )|
| \textit{Mycoplasma hominis}                   | 1  | (5 )| \textit{Klebsiella} spp.                        | 2  | (5 )| \textit{Salmonella enterica}                   | 3  | (4 )|
| Aspergillus fumigatus                        | 1  | (3 )| \textit{Bacteroides fragilis}                   | 1  | (3 )| \textit{Proteus mirabilis}                     | 2  | (3 )|
| \textit{Bacteroides} fragilis                | 1  | (3 )| \textit{Bacteroides} xylosiisovolens            | 1  | (1 )| \textit{Raoultella} spp.                      | 2  | (3 )|
| \textit{Corynebacterium} spp.                 | 1  | (3 )| \textit{Finegoldia magna}                      | 1  | (3 )| \textit{Dialister inviscus}                    | 1  | (1 )|
| \textit{Mycobacterium} chimaera              | 1  | (3 )| \textit{Escherichia coli}                      | 1  | (1 )| \textit{Escherichia fergusonii}                | 1  | (1 )|
| \textit{Pseudomonas aeruginosa}               | 1  | (3 )| \textit{Fusobacterium nucleatum}                | 1  | (1 )| \textit{Listeria innocua}                     | 1  | (1 )|
| \textit{Streptococcus} spp.                   | 1  | (3 )| \textit{Mycobacterium} intracellular            | 1  | (1 )| \textit{Pasteurella multocida}                 | 1  | (1 )|
| \textit{Aspergillus} fumigatus                | 1  | (3 )| \textit{Pluralibacter pyrinus}                  | 1  | (1 )| \textit{Propionibacterium} acnes              | 1  | (1 )|
| \textit{Bacteroides} fragilis                | 1  | (3 )| \textit{Shigella} spp.                         | 1  | (1 )| \textit{Yokenella} spp.                       | 1  | (1 )|
| \textit{Corynebacterium} spp.                 | 1  | (3 )| \textit{Pasteurella multocida}                 | 1  | (1 )| \textit{Polymicrobial}                        | 2  | (3 )|

\(^a\) Sample origin: vascular prosthesis (5\%), deep wound cultures (80\%), superficial wound culture (5\%), negative pressure wound therapy foams (10\%); \(^b\) Sample origin: vascular prosthesis (12\%), deep wound cultures (78\%), superficial wound cultures (10\%), biopsy (5\%); \(^c\) Sample origin: vascular prostheses (5\%), biopsy (5\%), negative pressure wound therapy foams (5\%), deep wound cultures (78\%), superficial wound cultures (6\%).
would have been investigated as well. However, as re-operative surgery is performed only in case of confirmed infection this was not feasible.

In conclusion, in about one third of cases with culture-negative specimens, the broad-range PCR allowed a microbiological diagnosis. Culture-positive, PCR-negative isolates were present among tissue samples with either low bacterial load, fungi or in 28% of cases contaminants were detected. With regard to enhanced microbiological diagnosis of VGI, we suggest to apply deep wound cultures as well as broad-range PCR in order to increase microbiological diagnosis especially in patients already undergoing antimicrobial treatment.

**AUTHOR CONTRIBUTIONS**

BH and GB designed the study. AS analyzed the data. EA-S, AS, GB, and BH wrote the first draft, and EA-S, AS, PK, AA, MH, AZ, GB, and BH wrote the final version of the manuscript. All investigators contributed to data collection and interpretation of the data, reviewed drafts of the manuscript, and approved the final manuscript.

**FUNDING**

This study was financed within the framework of the Vascular Graft Cohort Study (VASGRA), supported by the Swiss National Science Foundation (SNF) grant #32473B_163132/1, a SNF protected-research-time for clinicians grant #32473B_163132/2; the Vontobel Foundation and the Rozalia Foundation (to BH). AA is supported by the academic career program Filling the gap of the Medical Faculty of the University of Zurich. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
VASGRA COHORT STUDY

The members of the VASGRA Cohort Study are (in alphabetical order): A. Anagnostopoulo, B. Hasse (PI), M. Hofmann, L. Husmann, B. Ledergerber, M. Lachat, Z. Rancic, A. Scherrer, A. Weber, R. Weber, A. Zinkernagel.

ACKNOWLEDGMENTS

We are grateful to our patients for their commitment and thank B. Ruehe, C. Rüegg, A. Wolfensberger, U. Matt, M. Kaelin, M. Greiner and R. Pop for their excellent patient care. We also thank C. Mueller and S. Buergin (study nurses) and C. Voegtl (administrative assistance). We thank Reinhard Zbinden, head diagnostics and the technicians of the Institute of Medical Microbiology of the University of Zurich for their expert help and assistance.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmed.2018.00169/full#supplementary-material
prosthetic joint infection: a prospective multicenter cross-sectional study. J Clin Microbiol. (2014) 52:3583–9. doi: 10.1128/JCM.01459-14
25. Trampuz A, Piper KE, Jacobson MJ, Hanssen AD, Unni KK, Osmon DR, et al. Sonication of removed hip and knee prostheses for diagnosis of infection. N Engl J Med. (2007) 357:654–63. doi: 10.1056/NEJMoa061588
26. Hasse B, Husmann L, Zinkernagel A, Weber R, Lachat M, Mayer D. Vascular graft infections. Swiss Med Wkly. (2013) 143:w13754. doi: 10.4414/smw.2013.13754
27. Rawson TM, Lee MJ, Khanna P, Gopal Rao G, Renton S, Buckley J. Microbiological characterisation of prosthetic vascular graft infection. J Infect. (2015) 71:400–2. doi: 10.1016/j.jinf.2015.04.008

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2018 Ajdler-Schaeffler, Scherrer, Keller, Anagnostopoulos, Hofmann, Rancic, Zinkernagel, Bloemberg, Hasse and the VASGRA Cohort. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.