Development of a Flow-Trough Microarray based Reverse Transcriptase Multiplex Ligation-Dependent Probe Amplification Assay for the Detection of European Bunyaviruses

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Abstract It is suspected that apart from tick-borne encephalitis virus several additional European Arboviruses such as the sandfly borne Toscana virus, sandfly fever Sicilian virus and sandfly fever Naples virus, mosquito-borne Tahyna virus, Inkoo virus, Batai virus and tick-borne Uukuniemi virus cause aseptic meningo-encephalitis or febrile disease in Europe. Currently, the microarray technology is developing rapidly and there are many efforts to apply it to infectious diseases diagnostics. In order to arrive at an assay system useful for high throughput analysis of samples from aseptic meningo-encephalitis cases the authors developed a combined multiplex ligation-dependent probe amplification and flow-through microarray assay for the detection of European Bunyaviruses. These results show that this combined assay indeed is highly sensitive, and specific for the accurate detection of multiple viruses.

Keywords European Bunyaviruses · MLPA · Flow-through chip · Aseptic meningo-encephalitis

Introduction

While climate warming in Europe and global social changes have led to focussing attention to novel emerging infectious diseases [1, 2], there are many hardly noticed European arboviral infections causing neurological disorders like encephalitis or meningo-encephalitis [3, 4] and fevers [5, 6]. The aetiology of aseptic meningo-encephalitis in 50% of the cases in Europe for example remains unclear [3, 7]. In this regard seven Bunyaviruses, particularly, sandfly borne Toscana virus (TOSV), sandfly fever Sicilian virus (SFSV) and sandfly fever Naples-Virus (SFNV), mosquito-borne Tahyna virus (TAHV), Inkoo virus (INKV), Batai virus (BATV) and tick-borne Uukuniemi virus (UUKV), may be of greater importance than hitherto realized. The laboratory diagnosis of viral infections is generally performed by various serological assays e.g., haemagglutination-inhibition (HAI), enzyme-linked immunoassay (EIA or ELISA), viral cell culture in combination with immuno-fluorescence (IF) or electron microscope [8, 9]. These conventional methods are well established but often time-consuming, and in addition the sensitivity is often low [8]. Molecular methods have become more and more important in the diagnosis of viral infections and diseases [10, 11]. These methods are based on the direct detection of viral-specific DNA or RNA (via PCR or reverse transcription (RT)-PCR) without the need of virus cultivation. One of the most successful technologies in recent years is real-time PCR, which allows amplification of nucleic acids and simultaneous confirmation of the amplified product by specific probes [12, 13]. Although these approaches are rapid, accurate, and offer high specificity and sensitivity, they also introduce new disadvantages. As successful identification depends on appropriately chosen primer sets and probes used in separate parallel amplifications. In order to solve this disadvantage for virus detection, researchers have developed multiplex real-time PCR assays to detect many viruses in one tube [14, 15]. The disadvantage of multiplex PCR and real-time PCR is that only a limited number of targets can
effectively be identified simultaneously due to restrictions caused by the spectral overlap of the fluorescent dyes used [16].

Currently, DNA microarray technology appears attractive in this regard [17, 18] for its ability of simultaneous screening for several pathogens in a single step reaction-array [19, 20]. However, the microarray technology is not yet widely used in clinical virology laboratories, because of problems with sensitivity and the requirement of large starting amounts of nucleic acids, and expense [21, 22].

After thorough analysis of the literature the authors decided to try an amplification and hybridisation approach, which amplifies a synthetic signal molecule, which is dependent on sequence-specific hybridisation rather than an amplification technique amplifying the target molecule itself. The authors have developed an approach that combines Multiplex ligation-dependent probe amplification (MLPA) with a flow-through chip hybridisation system (PamGene, Netherlands). The principle of this assay is presented and described on Fig. 1.

This set up allows an easy design of several MLPA systems that can be hybridised simultaneously to the microarray without the need for optimisation of hybridisation sequences and specific microarray probe printing. In addition, this technology reduces chip costs and allows rapid hybridisation and reduced input amounts of nucleic acid samples in the comparison to classic DNA arrays. The flow-through microarray offered full process automation, including incubation, washing, hybridisation, altogether reducing the potential for cross contamination.

Taken together the author developed an assay based on the combination of RT-MLPA and flow-through 3D universal microarray for the detection of seven European Bunyaviruses, particularly (TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV) for an expanded accurate diagnosis of aseptic meningo-encephalitis and fevers of unknown origin in Europe.

Materials and Methods

Viral Cultures

The virus-isolates (TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV) were grown in VeroB4 cells or VeroE6 cells in Minimum essential or Dulbecco’s modified Eagle medium, respectively, supplemented with 2% foetal calf serum and penicillin/streptomycin, at 37°C in 5% CO₂
atmosphere. Each strain was passaged for at least four times. The supernatants were harvested when the cytopathic effect (CPE) of the infected cells was more than 90%. Subsequently, the supernatants were used for viral RNA extraction.

Enrichment of the Viral Particles and Viral RNA Extraction

The viral particles were enriched from supernatants using 30% Polyethylene glycol (PEG). In brief, 20 ml cell culture supernatant was centrifuged at 50 ml Falcon tubes at 4°C, 2,000 rpm for 10 min to eliminate cell debris, transferred into new 50 ml Falcon tubes, and supplemented with 1.48 ml of 5 M NaCl and 10.8 ml of PEG8000. Afterwards the supernatant was shaken carefully on a spinning-wheel at 4°C for 30 min and centrifuged at 4°C, 6,000 rpm for 1 h. The enriched virus particle pellets were processed for RNA isolation. The viral RNA genome of all investigated viruses was extracted using PeqGold TriFast™ (PeqLab, Germany) according to the manufacturer’s instructions.

Generation of DNA and RNA Standards

Isolated RNAs were reverse transcribed using the Transcriptor First Strand cDNA Synthesis Kit (Roche, Germany) and the corresponding specific primers for the S-segment according to the manufacturer’s instructions. The resulting cDNAs were amplified using AccuPrime Pfx Polymerase (Invitrogen, Germany), dNTPs and specific primers designed for the UUKV S-segment forward primer: 5' ACACAAGACCTCAACTTAGCTC-3', reverse primer: 5'-ACACAAAGACCC TCAAACATTAAGC-3' and the S-segment of SFNV, forward primer: 5'-TGCCCCACCCCCCTACAT-3', reverse primer: 5'-ACACAAAAAGAT CCGTTATTTAAATTC-3', and specific primers for TOSV, SFSV, TAHV, INKV and BATV published previously by us [23, 24]. A total of 14 MLPA probes were designed using the DNA primer design software program (PamGene, The Netherlands) and selected using the following criteria: The MLPA probes never overlap and locate directly adjacent to each other, total probe length (LPO + RPO) 100–132 bp, PCR primer sequence 42 nucleotides (nt), unique Cpam 20 nt, max length of LHS and RHS 70 nt at a melting temperature (Tm) of 67 and 70°C. In addition, Tm was checked using Exiqon Tm prediction (http://lna-tm.com/) and the Raw program from MRC-Holland and secondary structure was assessed using the mfold server (http://www.bioinfo.rpi.edu/applications/mfold/cgi-bin/rna-form1.cgi). The specificity of the designed probes was checked using a Blastn-analysis against the sequences in the NCBI database and against the 124 zip-codes spotted on the PamChip® 4U. The designed MLPA probes were synthesized by Biolegio (Nijmegen, The Netherlands) and are listed in Table 1, showing the size (100–113 bp) of MLPA/RT-MLPA PCR amplicons.
MLPA and RT-MLPA Reaction

MLPA and RT-MLPA reagents were obtained from MRC-Holland (Amsterdam, The Netherlands). The reactions were performed according to the manufacturer’s protocol. The purified S-segment amplicons of TOSV, SFSV, SFNV, TAHV, INKV BATV and UUKV (100–150 ng) or water as a negative control were diluted in TE buffer (pH 8) to 5 µl and heated at 98 °C for 5 min. After cooling at 25 °C, 1.5 µl of the corresponding MLPA probe mix and 1.5 µl of MLPA buffer were added to each sample, followed by denaturation at 95 °C for 1 min and hybridization at 60 °C for 12–16 h. Ligation was performed by adding 30 µl of ligation mix at 54 °C for 15 min to each sample. After the inactivation of ligase at 98 °C for 5 min, the MLPA PCR was carried out in a final volume of 50 µl using the SALSA forward primer 5’-GGGTTCCCTAAAGGGTGGA-3’ and reverse primer 5’-GTGCCAGACAGATCCAATCTAGA-3’ PCR primers complementary to the primer binding sites of the LPO (5’ FAM label) and RPO probes (MRC-Holland, The Netherlands) using the following temperature profile: 35 cycles of 95 °C/30 s, 60 °C/30 s and 72 °C/60 s, and a final elongation step at 72 °C for 20 min.

For the RT-MLPA reactions RNA standards or water as negative control were transcribed to cDNA using MMLV reverse transcriptase enzyme (Promega, Germany) and specific or hexamer primers according to Promega protocol instructions. In order to increase the sensitivity of MLPA probe sets 20% formamide buffer was added to the hybridization step of MLPA probes to the target cDNAs and assayed at a temperature range of 56–60 °C. In order to check the specificity of MLPA probes, MLPA/RT-MLPA negative control reactions were performed with water instead of MLPA probes.

Gel Electrophoresis by Agilent Bioanalyzer

Prior the microarray hybridization, the amplified MLPA and RT-MLPA products were analysed by gel electrophoresis using the Agilent Bioanalyzer (Agilent Technology, Germany). The short-length amplicons were analysed using the DNA 1000 LabChip® kit.

3D Microarray Hybridization and Detection System

The MLPA, RT-MLPA PCR and negative control products were subsequently analysed by flow-through microarray using a universal customized PamChip® 4U which consisted of a 4-arrays of a porous aluminium-oxide solid matrix. 124 zip-code addresses spotted as duplicate onto each array (PamGene, The Netherlands). The technology is available for DNA and RNA applications (Fig. 1). Hybridization, washing and detection was performed on the PamStation®12 which is an automated microarray processing system described elsewhere [26, 27], that also allows automated data analysis. 11 µl of each MLPA, RT-MLPA or negative control PCR reaction was denaturated at 95 °C for 5 min, 5.5 µl hybridization buffer containing 5× SSPE and 0.5 N-lauroylsarcosine and 11.5 µl of water was added. The array was subjected to an initial threefold wash with 1× hybridisation buffer containing 1× SSPE and 0.1% N-lauroylsarcosine (pH 7.0). The samples (25 µl)

### Table 1 Specific viral hybridization sequences of MLPA probes

| Virus Specific hybridization sequence of the left probe (LHS) | Specific hybridization sequence of the right probe (RHS) | Size of RT or MLPA amplicons |
|-------------------------------------------------------------|-------------------------------------------------------------|-----------------------------|
| PAM16 Toscan-1 GCATCCATAGTGTCGACCAC | ACGGGCAGGCATTCTGACAGG | 104 bp |
| PAM20 Toscan-2 CAGGGATTCTGGCAAGCACAT | TCAGCCGACAGATCCACCAGGAAC | 107 bp |
| PAM13 SFS-1 CCCGGCAATTAACATTGATGTGAGG | TGTCATAGACTTCTCCAG | 107 bp |
| PAM15 SFS-2 TGCAACGGCTCTAATCTCCT | TTGCTCCCTCTGGAGACATTCTC | 105 bp |
| PAM44 SFN S-1 GAACCCCCACCAAATCAATGACC | AATTATTCACCTTTACCTCCCTTC | 113 bp |
| PAM45 SFN S-2 ATTCCACCTACCTCCTCCTC | CCCACATTGGGACTGCTACCTGG | 104 bp |
| PAM84 Tahyna-1 GAGCTTAAGGAGGTAGGACGAC | TTAATGGGGTAGGGGCGCC | 106 bp |
| PAM85 Tahyna-2 GGTGTAGACGAGTTCAAGGAAATG | AAGACCCAGCTAAAGGGAGATG | 111 bp |
| PAM78 Inkoo-1 TTTGGAAGTGGCAGGTGGAGATTG | TCAATAATCTTTTTCTGAAA + C+A +G+G* | 113 bp |
| PAM75 Inkoo-2 GAGATGTGCGAGGAGATTGTC | AATAATCATTTTCTGGAACAGG | 106 bp |
| PAM100 Batai-1 CAGATGGGAGAGGTTTACC | CTTCTATTTTTCCAGG + C+T + G | 105 bp |
| PAM102 Batai-2 CAGATGGGAGAGGTTTACC | TCTATTTTCTCCAGGCTG | 104 bp |
| PAM42 Uuk S-1 ACCATCAAGTGAAGCAGG | GGAATGGGGCAAGGAAGATC | 104 bp |
| PAM64 Uuk S-2 CACTCTACTACCTAAGGCAAAC | CCACTGAGGCGATATCCACACC | 106 bp |

All sequences are given in 5’ to 3’ orientation, * nucleotides marked with + are locked nucleotides
were loaded onto the arrays and pumped back and forth through the porous membrane by air pressure during hybridisation at 50, 53, 56, 59 and 62 °C for 20 min. Positive signals of the hybridization were detected by the integrated CCD camera of the PamStation®12 at the following exposure times (ET), 100, 300, 500, 800 and 1000 ms, using the Cy3 filter set to detect the FAM labelled hybridized RT-MLPA products.

Microarray Data Analysis

The recorded images of each array in the corresponding data files were transferred to the BioNavigator software (PamGene, The Netherlands) and were converted into spot intensity values. Median signal intensities and the local background were calculated for each spot on the hybridized array. The signal of local background was subtracted and the fluorescence saturation was limited to <1% of the spots. These calculated median signals were transported as text files, and the mean of signal intensity and standard deviation (SD) were calculated by Microsoft Exel and presented on melting curve diagrams.

Results

MLPA Reaction

An outline of the MLPA reaction is shown in Fig. 1. This reaction allows the detection of up to 45 different targets in one simple reaction, using a single PCR primer pair [28, 29]. Since the MLPA reaction was originally developed for genomic DNA applications [28], the authors started MLPA reactions using 100–150 ng highly purified PCR amplicons of the S-segments of the European Bunyaviruses TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV as target template for the MLPA hybridisation.

The target DNA was denatured and corresponding MLPA probes were added and hybridized for 12–16 h at 60°C in a thermocycler with heated lid. Ligation and inactivation of ligase was followed by PCR amplification of the ligated MLPA probe system using a FAM labelled forward primer. The analysis of the MLPA amplicons using the Agilent Bioanalyzer confirmed the corresponding amplicon of all investigated MLPA systems. The amplicon bands of one virus migrated separately from each other but banded a little higher than expected. This was attributed to a combination of secondary DNA structure of the amplicons and the sizing error of the Bioanalyzer which is given as 10%. Furthermore the SFSV, INKV and TOSV showed additionally bands presenting an excess of primer dimers and unligated MLPA probe in the reaction (Fig. 2). Finally, the detection of these MLPA amplicons on the PamChip®4U was performed using a continuous hybridisation temperature profile protocol at 50, 53, 56, 59 and 62 °C to identify the optimal hybridisation temperature of each MLPA probe set for each virus. The results showed that the authors could identify all seven European Bunyaviruses on the DNA level, however, the signal intensity of MLPA PCR amplicons of TOSV, BATV, INKV and UUKV were stronger than those of SFNV, SFSV and TAHV. The specific signals were accompanied by some background/cross-hybridisation at a hybridisation temperature of 50°C but disappeared at increased temperatures. The signals at optimal microarray hybridisation temperature are presented on Fig. 3. TOSV, SFSV, SFNV, INKV and UUKV showed optimal hybridisation temperature at 53°C, while TAHV and BATV hybridised optimal at 56°C.

RT-MLPA Reaction

After establishing the 3D microarray based MLPA assay at the DNA level, the authors introduced a RT step into the MLPA reaction for the transcription of target RNA into cDNA to adapt the method for the detection of European Bunyaviruses. Initially, the RT step was performed with specific primers but later the procedure was simplified using hexamer primers. The RT step preceded MLPA probe hybridization, MLPA probe ligation and probe amplification. Again the RT-MLPA amplicon size was
examined and confirmed at the correct size by Agilent Bioanalyzer and then the amplicons were detected on the PamChip® 4U using the optimised hybridization protocol. The observed signals after the RT-MLPA amplicon microarray hybridization of all investigated viruses were similar to those of the MLPA amplicon microarray hybridisation (Figs. 3 and 4).

Melting Curve Analysis

The authors generated the melting curves for each MLPA set from the continuous microarray hybridization fluorescence read out data at 50, 53, 56, 59 and 62 °C. The mean of the signal intensities of each MLPA probe set was calculated and plotted in Fig. 5, showing the signal intensity of each MLPA probe set at different hybridization temperatures. It was observed that in general the signal strength of each MLPA probe set decreased with increasing hybridisation temperature and with decreasing RNA molecule numbers as expected. The signal intensities of different MLPA probe sets for the detection of one particular virus showed distinct sensitivities. This type of analysis helped to identify highly sensitive probe sets. Remarkably, the optimal hybridisation temperature for all RT-MLPA amplicons was now observed at 53°C—ideal for a multiplex MLPA assay.

Sensitivity Analysis

In order to determine the sensitivity of the 3D microarray hybridization of RT-MLPA PCRs products on the PamChip® 4U, RNA standards for TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV

![Fig. 3 MLPA hybridisation pattern on the flow-through chip. Representative images of MLPA amplicon hybridisation on flow-trough microarray for the detection of seven the European arboviruses, TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV](image1)

![Fig. 4 Sensitivity of RT-MLPA hybridisation patterns on the flow-through chip. Representative microarray images of RT-MLPA PCR products for the detection of seven European Arboviruses, showing detection limits for TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV](image2)
TAHV, INKV, BATV and UUKV were generated (see Materials and Methods). Serial dilutions of RNA standards were prepared and subjected to real-time PCR. Subsequently these serial dilutions of RNA standards were used for RT-MLPA reactions. The RT-MLPA hybridization step was performed with and without 20% formamide buffer to improve specific and reduce unspecific hybridisation. The RT-MLPA amplicons were detected on PamChip® 4U using the optimized microarray hybridization protocol described above.

It was observed an increased probe sensitivity using 20% formamide during the MLPA probe target hybridization reaction step in comparison to the standard procedure. Starting the procedure from the transcribed RNA
standards the sensitivity of detection of the combined RT-MLPA and hybridisation on the PamChip® 4U for
TOSV, BATV, UUKV and INKV was 10 copies, while SFSV, SFNV and TAHV was 1000 copies (Fig. 4).

Probe Specificity

In order to evaluate the specificity of the developed assay, all of the designed MLPA probes were tested in a mono-
plex and in a multiplex RT-MLPA reaction, containing the generated RNA standards of TOSV, SFSV, SFNV, TAHV,
INKV, BATV and UUKV. The generated RT-MLPA PCR amplicons were purified to remove the unligated MLPA
probes, analysed by gel electrophoresis and subsequently hybridized on the PamChip® 4U. The results clearly show
that even in the multiplex mixture containing all investigated probe sets only specific MLPA hybridization, liga-
tion and hybridisation to the microarray occurred. Cross detection was not observed.

In order to test for cross reactions of the microarray-based MLPA assays on the Pamgene system, all total
RNAs of TOSV, SFSV, SFNV, TAHV, INKV, BATV and UUKV were extracted from the supernatants of the corre-
sponding viral cell cultures and used as template for RT-MLPA reaction. Afterwards the RT-MLPA amplicons
were examined by gel electrophoresis on their amplicon size and analysed by PamGene microarray hybridization on
PamChip® 4U. The data analysis showed similar results to those with standards RNAs for all investigated European
Bunyaviruses. The pattern of the specific signal for each virus was observed as expected. Again cross detection was
not observed.

Comparison of RT-MLPA Microarray assays
with Real-time RT-PCR assays

The capacity of both assays to detect each targeted virus was investigated by using RNA standards and RNAs from
viral cell cultures. As mentioned above serial dilution of standard RNAs and RNA from cell culture for TOSV,
SFSV, SFNV, TAHV, INKV, BATV and UUKV were performed and used for real-time RT-PCR, MLPA, as well
as RT-MLPA flow-through microarray assays. The results are shown in Table 2, indicating that the sensitivity of each
monoplex real-time PCR assay ranged from 10 to 1000 copies of target RNA per reaction, depending on the virus
targeted. The sensitivity of MLPA or RT-MLPA flow-through microarray assay ranged from 10 copies for TOSV,
INKV, BATV and UUKV to 1000 copies for SFSV, SFNV and TAHV. In most cases, the sensitivity of both methods
was comparable.

Discussion

The authors have developed a combined MLPA and uni-
versal flow-through microarray detection assay for Euro-
pean Bunyaviruses, particularly, TOSV, SFSV, SFNV,
TAHV, INKV, BATV and UUKV. The combined assay
design used the MLPA technique the amplicons of which
were hybridized to the 3D universal flow-through micro-
array (PamChip® 4U). MLPA probes (LPO/RPO) were
designed for two conserved regions of the Bunyavirus
S-segments. After MLPA or RT-MLPA amplification using
a FAM labelled driving primer the amplicon samples were
actively pumped back and forth through the porous struc-
ture of the array membrane of the PamChip® 4U by air
pressure, and hybridised to the zip-code addresses which
are immobilized on the array membrane.

The aluminium-oxide matrix significantly improves the
signal capture, as compared with spotted arrays or in situ
synthesised arrays on glass surfaces [30]. The signal
intensities observed for the hybridisation results were
indeed very strong and specific for all analyzed European
Bunyaviruses, indicating high efficiency and fast hybridisation on the array during their identification.

The procedure of the hybridisation on the chip at the
temperature range used (50, 53, 56, 59 and 62 °C) took
only about 2 h. This is an advantage over longer (over-
night) hybridisation periods used in many glass slide based
microarray systems [27, 31]. The MLPA probe hybridisa-
tion, however, is still very long and at 12–18 h almost
negates the advantages of the rapid microarray
hybridisation.

Although the Cpam sequences conveying the hybrid-
isation onto the microarray zip- code (Pam4U sequence)
probes, have all been designed to meet a narrow hybridis-
atation reaction window, the authors observed differing hybridisation characteristics of the various RT-MLPA-PCR
amplics to the respective microarray probes (Fig. 5).
This may be explained by the variant sequences flanking
the Cpam hybridisation sequence (especially the virus-
specific MLPA hybridisation sequences RHS/LHS) and the
influence they elicit on the T.M of the Cpam sequence and

Table 2  Analytical sensitivities of various assay types during the
MLPA-microarray development given as molecules detected

| Virus assay | RT-PCR | MLPA | MLPA-microarray |
|------------|--------|------|-----------------|
| TOSV       | 10^1   | 10^1 | 10^1            |
| SFSV       | 10^2   | 10^2 | 10^3            |
| SFNV       | 10^2   | 10^2 | 10^3            |
| TAHV       | 10^2   | 10^2 | 10^3            |
| INKV       | 10^3   | 10^1 | 10^1            |
| BATV       | 10^1   | 10^1 | 10^1            |
| UUKV       | 10^1   | 10^1 | 10^1            |
on the multiple hybridisation states that can occur between hybridisation partners as described by Santa Lucia [32, 33].

Quan et al. [21] reported that randomly primed amplification of DNA microarray assays allow the detection of a large number of pathogens. The sensitivity of such a generic approach is, however, much reduced in comparison to pathogen-specific priming [34, 35] and is the main reason why microarray diagnostics have not yet arrived in the clinical diagnostic laboratory [36].

In order to compare the sensitivity of Taqman probe realtime RT-PCR assays and RT-MLPA assays, the respective amplicons and MLPA probes were designed in the same target region. The comparative analysis showed that the analytical sensitivity of the real-time PCR and MLPA assays was almost identical (Table 2 columns 2 and 3) and only a little loss of sensitivity was observed for the detection of SFSV, SFNV and TAHV viruses when the RT-MLPA amplicons were hybridised to the microarray (Table 2 columns 2 and 4). Remarkably, the sensitivity of INKV virus MLPA amplicon hybridisation on the chip was higher than real-time PCR. These results confirm that an amplification step is necessary to achieve a high sensitivity in microarray hybridisation assays. It should be noted that in the case of the MLPA the detection signal and not the target sequence is amplified, which significantly reduces the problem of target molecule carry over contamination in a diagnostic setting.

The data analysis was easy to perform thus simplifying the microarray data management. The obtained data were highly reproducible, as the standard deviation and the variance of the calculated mean for the individual spot signal intensity was negligible (data not shown). Altogether the results support other reports on MLPA assays combined with flow through microarrays describing high sensitivity, specificity, speed, and reproducibility [37, 38].

Although real-time RT-PCR is fast and sensitive, it is typically designed for detecting one to three targets per reaction. For the detection of seven Bunyaviruses, the samples would need to be divided into multiple reactions. This may be an issue when the amount of sample (e.g., cerebrospinal fluid) is small. In contrast, the MLPA method combined with the narrow reaction window hybridisation (zip-codes) on a 3D flow-through array allows the highly complex analysis of up to 45 targets and the authors believe that the assay format is easily extensible to include more viruses.

For the evaluation of the specificity of the monoplex and the multiplex RT-MLPA for the mentioned viruses, the authors used the corresponding standard RNAs or viral RNAs extracted from cell cultures as target molecules for RT-MLPA reaction. The results showed that MLPA probes were specific and did not cross-hybridize to any of the other tested European Bunyaviruses or to the cellular background. The MLPA products showed specific hybridisation to the microarray.

In order to optimize the multiplex assay it was therefore intend to extend the MLPA probe design to M- and L-segments of these viruses to create a higher specificity since amplification probes of any kind suffer from sequence variation mismatch.

Currently, there is no report on the use of a microarray for the detection of Bunyaviruses, but similar assays have been reported for the detection of respiratory viruses [39, 40], West Nile virus and Dengue virus in clinical materials [38, 41].

A multiplex MLPA assay using an RT-PCR pre-amplification step developed for the simultaneous detection of six DNA viruses causing infections of the central nervous system termed MeningoFinder [42] was evaluated using extracts from cerebrospinal fluid (CSF) samples. Similarly the RespiFinder RT-MLPA system was used for the simultaneous detection of fifteen respiratory RNA viruses in clinical samples. It included a pre-amplification step by RT-PCR protocol to reduce the hybridisation time of the MLPA reaction from 12 to 1 h. The improved hybridisation time of the RespiFinder, however, is achieved at the cost of another contamination risk during pre-amplification [39].

In another approach for the detection of respiratory viruses a multiplex PCR followed by primer extension and microarray hybridization was also used and showed good agreement with real-time RT-PCR when using clinical samples [40]. The most comparable study is the development of a multiplex reverse transcriptase PCR-ligase detection assay (RT-LDR) which hybridised to a universal microarray for the detection of several West Nile virus genes [38] and a similar setup for the four Dengue viruses which was successfully tested using clinical samples [41]. It therefore seems highly likely that the RT-MLPA combined with flow-through universal microarray assay presented here, which also uses a ligation step without pre-amplification of the target sequence could be as sensitive in clinical samples as it was shown using synthetic RNA standards and viral cell cultures.

In conclusion, the authors have shown that combined multiplex MLPA and 3D universal microarray assay is sufficiently sensitive and specific for the detection of European Bunyaviruses and show good correlation to monoplex real-RT-PCR, therefore the authors expect that this method can be developed into a diagnostic tool for the rapid analysis of infections due to European Arboviruses.

Acknowledgments The authors thank gratefully Monique Mommersteeg and Faris Naji (PamGene company, Netherlands) for the technical assistance by the establishment of this method and data analysis, respectively. The study was supported by the Federal Ministry of Education and Research (BMBF), grant number 01Kl0710.
“Research on Zoonotic Infectious Diseases” programme, Emerging arthropode-borne viral infections in Germany: Pathogenesis, diagnostics and surveillance.

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