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ABSTRACT

Introduction: *Brucella suis* is the causative agent of brucellosis in suidae and is differentiated into five biovars (bv). Biovars 1 and 3 possess zoonotic potential and can infect humans, whereas biovar 2 represents the main source of brucellosis in feral and domestic pigs in Europe. Both aspects, the zoonotic threat and the economic loss, emphasize the necessity to monitor feral and domestic pig populations. Available serological or PCR based methods lack sensitivity and specificity.

Results: Here a bioinformatics approach was used to identify a *B. suis* specific 17 bp repeat on chromosome II (BS1330_II0657 locus). This repeat is common for *B. suis* bv 1 to 4 and was used to develop a TaqMan probe assay. The average PCR efficiency was determined as 95% and the limit of detection as 12,5 fg/µl of DNA, equally to 3.7 bacterial genomes. This assay has the highest sensitivity of all previously described *B. suis* specific PCR assays, making it possible to detect 3-4 bacterial genomes per 1 µl of sample. The assay was tested 100% specific for *B. suis* and negative for other *Brucella* spp. and closely related non-*Brucella* species.

Conclusions: This novel qPCR assay could become a rapid, inexpensive and reliable screening method for large sample pools of B. suis 1 to 4. This method will be applicable for field samples after validation.

INTRODUCTION

Brucellosis is a zoonotic infectious disease caused by different species of the genus Brucella. One of the prevalent species in feral pigs and hares is Brucella suis with five different biovars. The biovars (bv) 1-3 infect mainly suidae, by 4 infects reindeers and by 5 is rarely found in rodents (Alton and others 1988, Forbes 1991). All of them possess a zoonotic potential, but only by 1 and 3 seem to cause severe diseases in humans (Munoz and others 2010). In Europe the most common cause of porcine brucellosis is due to infections with B. suis by 2, whereas by 1 and 3 are reported in Croatia only once (Garcia-Yoldi and others 2007, Cvetnic and others 2009). In most European countries feral pigs and hares seem to be the main reservoir for B. suis by 2

(Garcia-Yoldi and others 2007, Leuenberger and others 2007, Melzer and others 2007, Munoz and others 2010, Gregoire and others 2012, Wu and others 2012). Transmission to domestic pigs may occur through direct contact, sexual transmission or shared water reservoirs (Risco and others 2014). The disease then causes infertility and abortion in female pigs and orchitis in male pigs. Furthermore, lymphadenitis, arthritis and subcutaneous abscesses may occur. Studies show that interactions between feral pigs and domestic pigs occur quite often (Wu and others 2012). Regarding the growing populations of feral pigs in Europe this could explain the increase of infections in domestic pigs with brucellosis since 1990 (Godfroid and Kasbohrer 2002). Therefore, brucellosis as a zoonotic disease is of great public health importance in many countries (Melzer and others 2007, Gerst and others 2010, Von Roost and others 2010). Monitoring of Brucella spp. infections in domestic livestock and wild animals is mainly based on detection of antibodies specific for smooth lipopolysaccharide. This includes the risk of false positive reactions, especially for porcine brucellosis due to cross-reacting antibodies to the highly similar O-antigen of Yersinia enterocolitica O:9 (Hurvell 1973). Additionally, non-reactors or animals which have not yet developed an antibody level can cause false negative results. During recent years PCR-based methods became more and more important as fast and reliable methods to detect Brucella-specific DNA from colony material or directly from field samples. On one hand conventional or real-time Brucella genus-specific PCR methods are used to identify brucellae from colony material or field samples (Bricker 2002, Probert and others 2004). On the other hand species-specific multiplex PCR techniques are implemented to differentiate on species level (Lopez-Goni and others 2011). Since realtime PCR is much more sensitive compared with conventional PCR it is often used to detect specific DNA in field samples directly. Obviously, a positive result in a genus-specific



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Bacterial species Reference		Bacterial species	Reference
Brucella spp.		Non- <i>Brucella</i> species	
B. abortus 544	ATCC 23448	Actinobacillus pleuropneumoniae	ATCC 2708
B. abortus S99		Bacillus brevis	ATCC 8246
<i>B. abortus</i> S19		Bacillus cereus	ATCC 1087
<i>B. abortus</i> 86/8/59	ATCC 23449	Bacillus megaterium	ATCC 1458
<i>B. abortus</i> Tulya	ATCC 23450	Bacillus subtilis	ATCC 6633
B. abortus 292	ATCC 23451	Bacillus thuringiensis	ATCC 1079
B. abortus B3196	ATCC 23452	Bordetella bronchiseptica	ATCC 1939
<i>3. abortus</i> 870	ATCC 23453	Burkholderia cepacia	DSM7288
<i>3. abortus</i> C68	ATCC 23455	Burkholderia mallei	ATCC 2334
<i>3. melitensis</i> 16M	ATCC 23456	Burkholderia pseudomallei	ATCC 2334
<i>B. melitensis</i> Elberg		Escherichia coli	DSM 30083
B. melitensis 63/9	ATCC 23457	Lactobacillus ruminis	DSM 20403
<i>3. melitensis</i> Ether	ATCC 23458	Mannheimia haemolytica	ATCC 3339
<i>3. suis</i> 1330 (bv 1)	ATCC 23444	Ochrobactrum anthropi	CCUG 104
<i>3. suis</i> Thomson (bv 2)	ATCC 23445	Oligella urethralis	DSM 7531
<i>3. suis</i> 686 (bv 3)	ATCC 23446	Pasteurella multocida	DSM 5281
<i>3. suis</i> 40 (bv 4)	ATCC 23447	Pasteurella multocida subsp. multocida	ATCC 4313
<i>3. suis</i> 513 (bv 5)	NCTC 11996	Proteus mirabilis	ATCC 2990
<i>3. canis</i> RM6/66	ATCC 23365	Pseudomonas aeruginosa	ATCC 9027
3. ovis 63/290	ATCC 25840	Pseudomonas alcaligenes	ATCC 1490
3. neotomae 5K33	ATCC 23459	Pseudomonas fluorescens	ATCC 1352
<i>3. ceti</i> B1/94	NCTC 12891	Pseudomonas polymyxa	ATCC 842
3. pinnipedialis B2/94	NCTC 12890	Rhodococcus equi	DSM 2030
<i>B. microti</i> CCM4915	BCCN 07-01	Staphylococcus aureus subsp. aureus	ATCC 2517
<i>3. inopinata</i> BO1	BCCN 09-01	Stenotrophomonas maltophilia	ATCC 1363
		Streptococcus agalactiae	ATCC 2795
		Streptococcus equinus	ATCC 9812
		Streptococcus parauberis	DSM 6631
		Taylorella equigenitalis	DSM 10668
		Yersinia enterocolitica subsp. enterocolitica	ATCC 9610

ATCC, American Type Culture Collection; bv, biovar; BCCN, Brucella Culture Collection from Nouzilly; CCUG, Culture Collection, University of Göteborg; DSM, German collection of microorganisms (Deutsche Sammlung von Mikroorganismen); NCTC, National Collection of Type Culture

real-time PCR should be confirmed by a second method which ideally is a species-specific method. So far published real-time PCR methods are able to distinguish *B. abortus, B. melitensis* and *B. suis* by 1 (Al Dahouk and others 2007a). In central Europe *B. suis* by 2 is endemic in wild boar and hare populations and therefore influences monitoring of brucellosis at least in pig farms (Garcia-Yoldi and others 2007, Leuenberger and others 2007, Munoz and others 2010, Gregoire and others 2012, Wu and others 2012). The last outbreaks of brucellosis in pigs in Germany were *B. suis* by 2 infections in outdoor holdings which have been confirmed by isolation of the infectious agent (Melzer and others 2007).

The aim of this study was to develop a quantitative real-time PCR (qPCR) assay as a sensitive diagnostic tool for detection of *Brucella suis* by 1–4. These biovars are the most prevalent and are of special epidemiological importance.

MATERIALS AND METHODS

Bacterial strains and growth conditions

Bacteria used in this study are listed in Tables 1 and 2. A collection of 25 *Brucella* spp. reference strains, 75

B. suis field isolates and 30 closely related and clinically relevant non-*Brucella* species were obtained from the in house strain collection of the national reference laboratory for brucellosis (Friedrich-Loeffler-Institut, Germany). *Brucella* spp. strains were maintained on Brucella agar (Becton Dickinson) or Brucella agar, containing polymyxin B (2500 IE) bacitracin (12,500 IE), cyclohexymid (50 mg/l), nalidixic acid (2.5 mg/l), nystatin (50,000 IE) and vancomycin (10 mg/l), respectively (Farrell 1974). *Brucella* cultures were incubated with 5–10 per cent of CO₂ at 37°C for 72 hours (Alton and others 1988). All other bacterial strains were routinely cultivated on 5 per cent sheep blood agar (Becton

TABLE 2: B. suis field isolates used in this study		
Biovar	Source of isolates	No. of isolates
1	Feral pig	1
2	Hare	20
	Feral pig	32
	Domestic pig	21
5	Human	1

Dickinson) or Luria Bertani agar (Fisher Scientific) overnight at 37°C.

Preparation of DNA templates

DNA was extracted from single colonies and resuspended in 200 µl of sterile distilled water after heat inactivation for two hours at 80°C. DNA isolation was carried out using the HighPure PCR Template Preparation Kit (Roche Diagnostics, Mannheim, Germany) according to the manufacturer's instructions for bacteria. DNA quantity and purity was determined using a NanoDropTM 1000 spectrophotometer (Thermo Fisher Scientific, Wilmington, USA). Serial 1:10 dilutions ranging from 100 ng to 1 fg/µl of DNA were prepared for qPCR efficiency and specificity testing. DNA was also directly isolated from one *Brucella* culture-positive spleen sample from a hare with the same extraction kit according to the manufacturer's instructions for mammalian tissue. This was done as an example to show suitability of this assay for field samples.

Bioinformatic analyses

For identification of a *B. suis*-specific target whole genome alignments were performed using the Geneious software (Geneious 6.0.3, Biomatters Ltd., Auckland, New Zealand). Genome sequences from various *Brucella* spp. (Table 3) were obtained from the NCBI GenBank database and analysed for *B. suis* by 1 and 2 conserved regions. Identified targets were then tested for specificity using BLAST search with a Discontiguous Megablast against the NCBI GenBank database.

Quantitative real time PCR

Oligonucleotides *B. suis* for 5'-GCC AAA TAT CCA TGC GGG AAG-3' and *B. suis* rev 5'-TGG GCA TTC TCT ACG GTG TG-3' targeting a 106 bp fragment of the

Brucella spp.	Accession no.*	
B. abortus	NC 016777	
D. abonus	NC 006933	
	NC 007624	
B. melitensis	NC 003318	
	NC 012442	
	NC 017247	
	NC_017245	
	NC_017283	
B. ovis	NC_009504	
B. canis	NC_010104	
B. microti	NC_013118	
B. pinnipedialis	NC_015858	
B. suis bv 1	NC_017250	
B. suis bv 2	NC_010167	
B. suis bv 3	NZ_DS999731	
B. suis bv 4	NZ_GG703794	
B. suis bv 5	NZ_DS999712	

BS1330 II0657 locus encoded on chromosome 2 of B. suis by 1 with the FAM-labelled hydrolysis probe B. suis probe 6-carboxyfluorescein (FAM)-TTG CGC TTT TGT GAT CTT TGC GCT TTA TGG-BHQ1 were used for the qPCR detection assay (Jena Bioscience, Germany). Primer and probe were designed with Geneious software (Geneious 6.0.3, Biomatters Ltd., Auckland, New Zealand). For qPCR analysis the Stratagene Mx3000P **OPCR** System (Agilent Technologies, Santa Clara, USA) was used. The qPCR was performed in a 25 µl format using the 2× TaqMan Universal PCR Mastermix (Applied Biosystems, Foster City, California, USA), 0.3µM of each primer, 0.1µM probe and 2µl of template DNA. The qPCR was carried out on an Mx3000P QPCR Instrument (Agilent Technologies, Santa Clara, California, USA) using the following cycling conditions: 50°C for 2 minutes, 95°C for 10 minutes and 50 cycles of 95°C for 15 seconds and 60°C for 1 minute. Data were analysed with the Mx3000Pro software. Raw data were normalised using an adaptive baseline correction and a smoothing with a moving average of three amplification points. Initially cycle threshold (C_t) was set with backgroundbased threshold algorithm. For better comparison of the results a threshold of 0.05 (dRn) was applied to all qPCR signals, as the highest of all computed thresholds from all experiments. Fluorescence signals above this value were considered as positive, below as negative results.

Determination of qPCR efficiency

Standard curves were done with serial dilutions 1:10 from 10^6 to 10^0 fg/µl of bacterial DNA from *B. suis* by 2. The efficiency of the PCR was calculated from the slope of the logarithmic regression of C_t values plotted against DNA concentrations by $E=e^{(-1/slope)-1}$. For statistical analysis and assessment of reproducibility, standard curve experiments were done in quadruplicate and repeated five times on five different days. Additionally, qPCR efficiencies for *B. suis* by 1–4 were compared by analysing serial dilutions 1:10 of genomic DNA from 10^6 to 10^2 fg/µl. Dilutions were chosen within the linear dynamic range of the assay and expected concentrations of DNA within possible specimens. Samples were analysed in duplicate.

Determination of the qPCR limit of detection

The limit of detection (LOD) was defined as the DNA concentration with 95 per cent positive qPCR results. Therefore we used 100, 50, 25, 12.5 and 6.25 fg/µl of *B. suis* by 2 DNA spiked into DNA preparations from anti-*Brucella* antibody and *Brucella* DNA-free swine sera. To proof *Brucella* DNA free status of the sera a published real-time PCR (Probert and others 2004) was used as described. LOD samples were analysed in octuplicate for each concentration and repeated three times on three different days.

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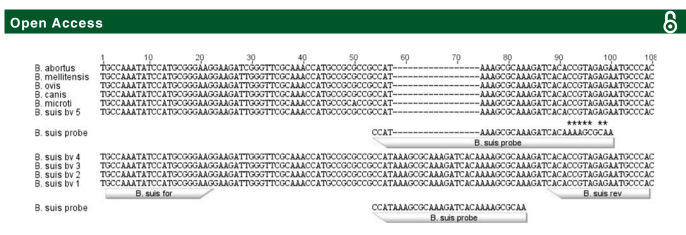


FIG 1: Alignment of the targeted 3' end of BS1330_II0657 locus from different *Brucella* spp. strains and *B. suis* bv 1–5. The alignment shows the 17 bp repeat specific for *B. suis* bv 1–4 and the resulting 7 bp mismatch of the probe (indicated by asterisks) for all other *Brucella* spp

Capillary gel electrophoresis

Real time PCR products were analysed by capillary gel electrophoresis using the QIAxcel DNA high resolution kit (1200) with an analytical range of 15 bp to 10 kb on the QIAxcel Advanced system (QIAGEN, Hilden, Germany). Briefly, 10 μ l of each PCR product were added to the template tray and a standardised volume of 0.1 μ l was injected automatically into the QIAxcel cartridge for analysis. For accurate fragment size determination the QX alignment marker 15 bp/3 kb and the QX DNA size marker FX174/HaeIII with a range of 50– 1350 bp was used. Electrophoresic separation of amplicons was then carried out by applying the OM500 program. For data analysis and generation of a virtual gel image the Biocalculator QIAxcel software was used.

RESULTS

Identification of a *B. suis* by 1–4 specific qPCR target by whole genome comparison.

Annotated genome sequences from 12 *Brucella* spp. and *B. suis* by 1 and 2 (Table 3) were analysed by whole

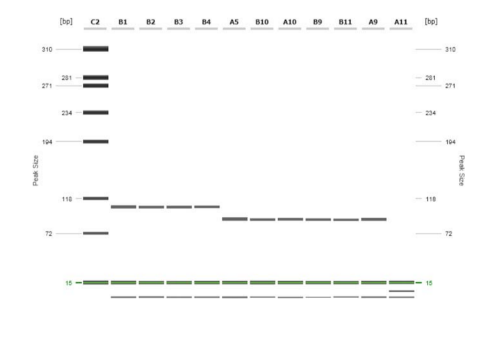
FIG 2: High-resolution capillary gel electrophoresis of the B. suis-targeted quantitative real-time PCR amplicons. Primer B. suis for and B. suis rev were used for amplification of the targeted region within BS1330_II0657 locus. Lack of the 17 bp repeat in Brucella spp. or B. suis by 5 are visualised and resulted in a fragment size of 106 bp for *B. suis* bv 1-4 and 89 bp for all other strains. Lane C2: Size marker; lane B1: B. suis by 1; lane B2: B. suis by 2; lane B3: B. suis bv 3; lane B4: B. suis by 4; lane A5: B. suis by 5; lane B10: B. abortus 544; lane A10: B. melitensis 16M: lane B9: B. ovis: lane B11: B. canis; lane A9: B. microti; lane A11: no template control

genome. A region within the 3' end of BS1330_II0657 locus was identified to be highly specific for *B. suis* with a 17 bp repeat exclusively found in *B. suis* bv 1–4 (Fig 1). All other *Brucella* spp. and *B. suis* bv 5 lack this repeat. Designed real-time primer were located up- and downstream of this repeat generating a 106 bp fragment for *B. suis* bv 1–4 and a 89 bp fragment for all other *Brucella* spp. and *B. suis* bv 5 (Fig 2). BLAST search performed for the 106 bp amplicon revealed 100 per cent complementarity for *B. suis* bv 1–4. The probe spanning the 17 bp repeat in *B. suis* bv 1–4 shows a 7 bp mismatch within the 5'-end in all other *Brucella* spp., lacking this repeat. However, this mismatch lowers the T_m to 57°C and prevents binding of the probe at the assay-specific annealing temperature of 60°C.

genome comparison with B. suis by 2 as reference

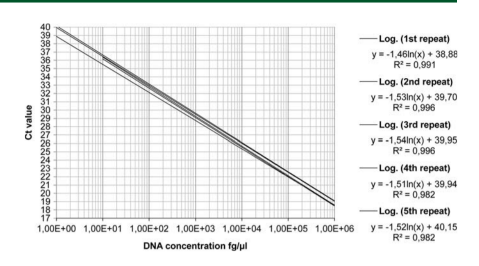
PCR efficiency

For PCR efficiency determination genomic DNA standards from 10^6 to 10^0 fg/µl of *B. suis* by 2 DNA were analysed. Efficiency was calculated from the slope and



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FIG 3: Logarithmic regression of quantitative real-time PCR efficiency testing experiments. Formulas show the slope values used for the efficiency calculation (slope In). The values at the end of each formula show the y interceptions, which represent the theoretical maximal C_t values



ranged between 90.8 and 97.6 per cent with a yinterception between 38.88 and 40.15 and a R^2 value between 0.9821 and 0.9965 (Fig 3). Analysis of *B. suis* by 1, 3 and 4 DNA standards revealed similar efficiencies of 94.3–95.5 per cent with y-intercepts between Ct- value 44.24 and 45.24 and an R^2 value between 0.9567 and 0.9965 (Fig 4). In general we observed a decreasing accuracy, assessed as an increasing standard error (se) for the resulting C_t values with decreasing DNA concentrations. The se increased from 1 ng/µl to 10 fg/µl from 0.20 to 0.83 during the PCR efficiency testing and from 100 fg/µl to 12.5 fg/µl from 0.36 to 1 for the LOD determination (Fig 5).

Limit of detection

The developed qPCR assay reproducibly detected 12.5 fg/µl of *B. suis* by 2 DNA (3.7 bacterial genomes). At this DNA concentration all qPCR assays were 100 per cent positive. However, at a DNA concentration of 6.25 fg/µl (1.8 bacterial genomes), still 83 per cent of performed reactions were positive (Fig 5).

PCR specificity

To evaluate the specificity of the *B. suis* qPCR, DNA preparations from 30 non-*Brucella* species (Table 1), 25

FIG 4: Logarithmic regression of the *B. suis* by 1–4 genomic DNA standards. Formulas show the slope values used for the efficiency calculation (slope In). The values at the end of each formula show the y-interceptions, which represent the theoretical maximal C_t values

Brucella spp. reference strains including five B. suis reference strains (Table 1), representing all five biovars were tested at a concentration of 100 pg/µl. For all 30 non-Brucella species negative results were obtained with Ct values not exceeding the calculated y-intercept from qPCR efficiency testing of 40 for any DNA concentration used (data not shown). Similar results were obtained for all Brucella spp. analysed. Only DNA from B. suis by 1-4 strains showed positive results with Ct values of approximately 22. There were no suspicious or undefined signals for non-B. suis by 1-4 samples (Fig 6). Furthermore we analysed 75 field isolates (Table 2) from confirmed B. suis cases from various animal species and geographical regions which were all qPCR positive for B. suis by 1-4. One sample, which was a confirmed human case of B. suis by 5 was tested negative.

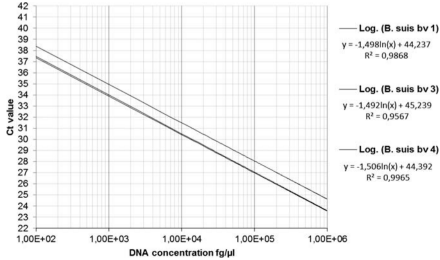
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Isolated DNA of the true positive spleen sample from the field was tested positive by *B. suis* qPCR also.

DISCUSSION

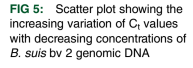
importance for direct detection of pathogens in clinical samples. For the diagnosis of brucellosis it has been already established and shown its usefulness. But till now

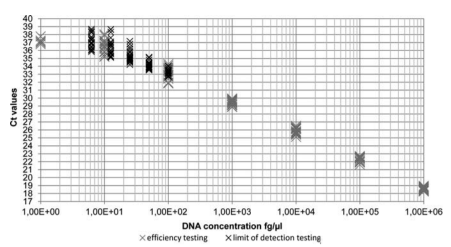
The qPCR is a fast and reliable tool with still growing



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and to the knowledge of the authors there is no description of a single probe qPCR that is able to detect all practically relevant *B. suis* by 1–4. A test panel of overall 100 *Brucella* and 30 non-*Brucella* strains showed 100 per cent specificity of the new assay for *B. suis* by 1–4. Our *B. suis* qPCR has an average efficiency of 95 per cent. This value shows its suitability for quantification of clinical samples even at very low concentrations of isolated DNA. A comparison of *B. suis* by 1–4 showed no significant differences in C_t-values for a serial dilution 1:10 from 1 ng/µl to 1 pg/µl of DNA. For the efficiencies calculated from standard curves only *B. suis* by 4 showed a decreased value of 88 per cent whereas *B. suis* by 1–3 had efficiencies of 93–94.5 per cent.

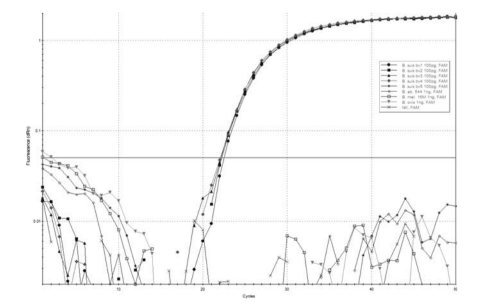
The limit of detecting was set at a concentration of 12.5 fg/µl of template DNA. This means the assay is able to detect three to four bacterial genomes in 1 µl of DNA isolated from clinical samples. To our knowledge this is the highest sensitivity reached with a *B. suis*-specific qPCR assay (Redkar and others 2001, Bogdanovich and others 2004, Probert and others 2004, Al Dahouk and others 2007b, Fretin and others 2008). Nevertheless we observed an increased variance of C_t

values for DNA concentrations of $100 \text{ fg/}\mu\text{l}$ down to the LOD with 12.5 fg/ μ l. We think the only reason for this is the growing impact of dilution errors and the resulting variance of used DNA for a single qPCR reaction. By spiking DNA into eluates from negative swine sera we also showed that the assay could be useful for diagnosis of clinical samples. DNA directly isolated from tissue of a spleen sample of one, by isolation proofed *Brucella* infected hare, was tested positive in the *B. suis* qPCR. To use the method for clinical samples like sera or biopsy material a true validation procedure has to be done.

A real-time PCR based single nucleotide polymorphism analysis (Gopaul and others 2008) differentiates *B. abortus*, *B. melitensis*, *B. ovis*, *B. suis*, *B. canis*, *B. neotomae* and marine *Brucella* spp. but with lower sensitivity than what seems to be necessary to examine field samples.

Regarding the problems with specificity and sensitivity in serology and the insufficient limits of detecting in most qPCR assays, this new assay could play a key role in improving the performance of large-scale brucellosis screenings of wild boar or hare populations and surveillance in domestic pigs.

FIG 6: Amplification curves of *B.* suis bv 1–4 (100 pg/µl DNA) with C_t values from 22.08 to 22.52 and of *B.* suis bv 5, *B.* abortus, *B.* melitensis and *B.* ovis (1 ng/µl DNA). Raw data were processed with an adaptive baseline correction and a moving average of three amplification points. The threshold was set to 0.05 (dRn)



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Data sharing statement No additional data are available.

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