

International Journal of Engineering & Technology

Website: www.sciencepubco.com/index.php/IJET

Research paper



Investigating Nano particles in Bioinformatic Analysis of the Brucella Genome for Indication and Differentation by qPCR

Alfiya V. Khammadova¹, Nail I. Khammadov², Maksim A. Kosarev², Konstantin A. Osyanin² and Eduard A. Shuralev³

¹Kazan Federal University, 18 Kremlyovskaya St., Kazan, 420008, Russian Federation

²Federal Center for Toxicological, Radiation and Biological Safety, Nauchniy Gorodok-2, Kazan, 420075, Russian Federation

³Kazan State Medical Academy – Branch Campus of Russian Medical Academy of Continuous Professional Education, 36 Butlerova St., Kazan, 420012, Russian Federation

*Corresponding author E-mail: eduard.shuralev@mail.ru

Abstract

Based on the identified genes, the design of specific primers and probes for qPCR was made, which makes it possible to differentiate *B. melitensis* and *B. abortus* in a single reaction. For the differentiation of Brucella strains of the same species, the PCR method of multilocus VNTR analysis, optimized for uniform temperature conditions, is described. In addition, MLVA profiles for *B. abortus* and *B. melitensis* genomes are presented. All methods of indication and differentiation of Brucella species and strains are described in detail, which allows them to be used in the everyday work of any PCR laboratory dealing with brucellosis. Control of amplification in the indication of bacteria of the genus Brucella and determination of their belonging to *B. abortus* and *B. melitensis* species is recommended to be carried out using the plasmid DNA designed in this work.

Keywords: Brucella; genome; indication; nano particle; identification; qPCR; MLVA.

1. Introduction

There is a wide variety of species of the genus Brucella found in nature [1]: B. melitensis; B. abortus; V. suis; B. neotomae; B. ovis; B. canis; B. ceti; B. pinnipedialis; V. microti. To date, there are methods for laboratory diagnosis of brucellosis pathogens by molecular genetic methods [2]. New approaches are being introduced in the indication and identification of antigenic, vaccine and field Brucella spp. using multilocus sequence typing [3], Luminex bead-based suspension array [4], Multiple-locus variable number tandem repeat analysis (MLVA) [5]. The methodology of PCR for the detection of brucella in organs and tissues in postmortem studies is being improved [6]. In addition, there are differentiation methods for different strains of the same species, based on the Variable number tandem repeat analysis (VNTR) in the genome of brucella and other organisms [7]. Such studies supplement various databases on the genomes of infectious disease pathogens, one of which is BrucellaBase, which contains information on the genomes of more than 500 strains of Brucella [8]. In this paper we showed the available tools for gene diagnosis and differentiation of brucellosis pathogens.

2. Materials and Methods

For the genetic indication of bacteria of the genus Brucella and optimization of the methodology for their differentiation, a bioinformation analysis of the brucella genome of various species and strains was carried out, the general principles of bioinformatic analysis are similar to the development of primers for other biopathogens [9, 10]. Nucleotide sequences for genome analysis were obtained from the National Center for Biotechnology Information (NCBI), in addition, "BLAST" and "VectorNTI 9.1.0" (Invitrogen Corporation) software utilities were used.

The design of oligonucleotide primers for quantitative real-time PCR (qPCR) was performed with the expectation of the possibility of multiplex amplification of various DNA markers in a single reaction. For the control of qPCR, a positive DNA sequence was developed comprising all of the desired marker nucleotide sequences. To amplify the chromosomal DNA, nucleic acids isolated from bacterial cultures of various brucella species (*B. suis, B. abortus, B. melitensis*) were used in the work. Isolation of nucleic acids was carried out by the method of magnetic sorption with a set of reagents "Magno-Sorb" variant 100-200, according to the instructions of the manufacturer – "Amplisens" (Russia).

For qPCR amplification, the following composition of the reaction mixture was used, per one sample: $1.5 \ \mu$ l of a 25 mM MgCl₂ solution; 2 μ l of a 2.5 mM dNTP solution; $1.5 \ \mu$ l of 10x PCR buffer; 0.5 μ l of 10 pM probe solution for PCR; 10 pM solution of forward and reverse primers of 0.5 μ l; 0.5 μ l Taq polymerase; 5 μ l DNA and 3.5 deionized water. All the reagents were produced by Syntol (Russia). The final volume of the reaction mixture was 15 μ l. The probe for PCR, forward and reverse primers were applied individually (specifically) for each type of PCR (Table 1) and in the multiplex variant (indication of generic and specific DNA markers in one reaction mixture). The reaction mixture for each PCR in MLVA was identical, except for the probe for PCR, not used in this reaction, and 10x PCR buffer with EvaGreen stain (instead of the 10x PCR buffer described above) was used to monitor PCR amplification.

PCR was performed on a C1000 amplifier with an optical unit CFX96 (BioRad). The amplification program was as follows: (i) DNA denaturation at 95° C for 2 minutes; (II) 5 cycles consisting



Copyright © 2018 Authors. This is an open access article distributed under the <u>Creative Commons Attribution License</u>, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

of: 10 seconds at 95°C, 30 seconds at 60°C, 10 seconds at 72°C; (III) 40 cycles consisting of: 10 seconds at 95°C, 30 seconds at 60°C, 10 seconds at 72°C. Detection of the result of PCR (fluorescence) occurs each 40th cycle of the third stage of the PCR, at 60°C via the FAM, Rox and R6G channels.

Electrophoresis was performed in a chamber for vertical electrophoresis (glass size 100×100 mm) manufactured by Helicon, with supply of 120 V into the chamber for 180-300 minutes. The buffer used for electrophoresis was from the "EF" kit (Central Research Institute of Epidemiology, Russia).

3. Results and Discussion

3.1 Analysis of Brucella Genes

The study on selecting the most informative genes for gene indication involved analysis of genomes of various Brucella strains of species B. abortus, B. melitensis. The alignment of the nucleotide sequence of different strains and isolates of each gene and the search for the genus-specific and species-specific sequences revealed optimal nucleotide sequences that can be successfully used for gene identification of bacteria of the genus Brucella and differentiation of Brucella species of interest to us (B. abortus and B. melitensis). Thus, for the indication of the genus Brucella, the gene BSCP31 (the sequence cipher in the gene bank - KX529834) was chosen, with a molecular mass of 1034 bp localized in the first chromosome. The locus BAW_20982 with a molecular mass of 1220 bp was chosen as the optimal display for B. abortus, the locus is located on the second chromosome at 1003355 to 1004575 bp (strain Wisconsin). For the indication of *B. melitensis*, the locus COR52_12390 with a molecular mass of 482 bp, localized in the second chromosome at 906042 to 906524 bp (strain B3), showed to be the most specific, and was used for the further design of the primers and a probe.

3.2 Design of Primers and Probes

Based on the nucleotide sequence of the above marker loci, the design of primers and probes for PCR was performed (Table 1); the main criteria were: to achieve the most identical annealing temperature of the primers; minimum content of dimers and secondary structures in primers and probes; a minimum of guanine and cytosine at the 3' end of the primers; the absence of guanine at the 5' end of the probe.

Table 1:. Nucleotide sequence of primers and probes for PCR indication of brucella

Detectable pathogen	Primer / probe en- coding	5 '-> 3` nucleotide sequence
0000110	Bru F	tcgaatggctcggttgcca
genus	Bru R	cgggtaaagcgtcgccagaa
Бисена	Bru P	(Fam)cgatcaagtcgggcgctctgga(BHQ1)
	Bru abortus F	gcaatcgtcgtattgccactataatcat
B. abortus	Bru abortus R	gacggcgcagttctcgaacaa
	Bru abortus P	(Rox)ccgaaaggatcagcgtgccagaa(BHQ2)
D	Bru meli F	tgettggcaceteggaaacae
в. melitensis	Bru meli R	cattcccgaaagccgatagagtttga
	Bru meli P	(R6G)tgaagcgctgcagacaaattttgacttcc(BHO2)

For the convenience of interpreting the results of PCR with the above primers and probes, the latter were labeled with various fluorescent stains, which are also indicated in Table 1, in parentheses. The characteristics of the primers and probes resulting from the design are indicated in Table 2.

Table 2:. Characteristics of the PCR-intended primers and probes

Detectable pathogen	Primer / probe encoding	Melting point	Number / length of dimers	Number / length of secondary structures
genus Bru-	Bru F	59.6°C	5 / 3 and 4	$\frac{2}{3}$ and
cena				4
	Bru R	59.7°C	2/3	0
	Bru P	65.1°C	4 / 3 and 4	1/3
B. abortus	Bru abortus F	59.5°C	7 / 3, 4 and	3 / 3 and
			6	6
	Bru abortus R	59.8°C	4/4	0
	Bru abortus P	64.0°C	1/4	0
B. melitensis	Bru meli F	59.6°C	2/3	1/3
	Bru meli R	60.2°C	4/3	1/3
	Bru meli P	66.6°C	7 / 4 and 6	1/4

As can be seen from Table 2, the developed primers are characterized by the same melting point (59.73±0.23°C), which allows for amplification under unified reaction conditions (one amplification program), from one to seven dimers and secondary structures in primers and probes, without the formation of critical polyneomas for PCR. This in aggregate will allow the use of these primers and probes for the indication of the brucella DNA. Complementarity of the above primers and probes (Table 1) to the nucleotide sequences of the desired bacteria is shown in Figure 1.

	Part of the second seco
418103	THEOREM DEPOSITORIA DESCRIPTION CHARACTER DEPOSITION SUBJECTION SUBJECTS DEPOSITOR DEPOSITORIES DEPOSITORIE
0.0001	HARD FILM MORE TRUE AND A TRUE TRUE AND A TR
A	аноссыли наласти наласти соститор сарости ресолиции аносто постоять начиные атальное токаност посотися сопстание намалися основных начитие таковать саростир таковать основных токанов такование <u>ресолиции</u>
	DEACCEPTOR STUDIAL TO ANTISSICS SCATSSICS CATSSICS AND TO AN ADDRESS TO AND THE ADDRESS AND ADDRESS AD
SECOND.	AND
В	CHORENEL TRACKARY OLIVERNAL REPORT CONTROL CONFORM TRALETA CONSULT ANALYTIK CONSULT ANALYTIK ANALYTIK ANALYTIKA ENDERING ANALYTIK CONTROL SOCCESSION CONTROL CONTROL CONTROLS ANALYTIKA ENDERING CONTROL CONTROL OF ANALYTIK CONTROL OF ANALYTIKA CONTROL OF ANALYTIKA ENDERING CONTROL OF ANALYTIK CONTROL OF ANALYTIKA CONTROL OF ANALYTIKA ENDERING CONTROL OF ANALYTIK CONTROL OF ANALYTIKA CONTROL OF ANALYTIKA ENDERING CONTROL OF ANALYTIK CONTROL OF ANALYTIKA ENDERING CONTROL OF ANALYTIKA CONTROL OF ANALYTIKA ENDERING CONTROL OF ANALYTIKA ENDERIN
DEPER	NAMES AND
C. seren	тольныция технольные мотоление мотоление такиности сонитолы Алионалети тостольсти: Слоковсто: Алеонетись ини ини ини ини ини ини ини ини ини ини

Fig. 1:. Complementarity of primers and probes for the indication of bacteria of the genus Brucella and species *B. abortus* and *B. melitensis*: A - nucleotide sequence of the gene BSCP31 *B. canis* strain YH-C16; B - nucleotide sequence of the locus BAW_20982 *B. abortus* strain Wisconsin; C - nucleotide sequence of the locus COR52_12390 *B. melitensis* strain B3.

Primers and a probe for the indication of bacteria of the genus Brucella (in Figure 1 denoted as brucella F, R, P) amplify a gene characteristic of all genus species, including *B. abortus* and *B. melitensis*. Primers and a probe for the indication of bacteria belonging to species *B. abortus* (in Figure 1 denoted as brucella abortus F, R, P) are complementary only for different strains and isolates of this species. Primers and a probe for indicating bacteria belonging to species *B. melitensis* (in Figure 1 denoted as brucella melitensis F, R, P) are also characterized by specific amplification of bacteria of this species only.

3.3 Design of Primers for Multilocus VNTR Analysis

Marker loci for MLVA of bacteria of the genus Brucella were determined based on the analysis of VNTR loci of brucella presented on the website: http://mlva.upsud.fr/mlvav4/genotyping/query.php. The nucleotide sequence of VNTR loci is determined from the data published by S.O. Sadikalieva and co-authors [7]. For the convenience of the reaction, the primers were modified, the nucleotide sequences are shown in Table 3. This modification was performed to optimize the molecular weights of the amplification products obtained (for convenient analysis of the amplicon after Polyacrylamide Gel

Electrophoresis (PAGE)) and unify the PCR conditions for ampli- fication of any locus.

Table 3:. Nucleotide sequence of primers and characteristics of amplicons for	VNTR ana	lysis of brucella
---	----------	-------------------

Item	5` -> 3`sequence	Variable region	The length of the amplicon excluding the variable repeats
VBru4F	ctgacgaagggaaggcaa	ataaaaaa	147 ba
VBru4R	aggcgatctggagattatcg	gtagggca	147 bp
VBru6F	gggatgtggtagggtaatcg	ggatgggcagcagacgttacccgcccgcagccgcgaagcgagcg	126 bp
VBru6R	gcgtgacaatcgactttttg	caa agg caa a ag cgg g cgttt g cct g g g g g g caa t g a t g g ca cg c g g g c g catt t a t ct a c g a ca c c c c c g g c g c g c g c	120 bp
VBru7F	agctgacggggaagaacat	ttecceta	125 bn
VBru7R	cctttttcagtcaaggcaaat	liccceta	125 00
VBru8F	attattcgcaggctcgtga	tccagectgtcattetgt	279 bp
VBru8R	caaacagaaggttttccagct	totagootgtomtotgt	
VBru9F	catggcggattcgttctt	actorect	103 hn
VBru9R	gggagtatgttttggttgtacatag	acigette	105 bp
VBru11F	gctgttgatctgaccttgca	actaccattaaccagaatcatacctcgattaataccaataccaagaacattaccgacaacaag	195 bp
VBru11R	ccagacaacaacctacgtcct	acisecultzzecuzantezactiezatuztaecultzecultzecultzzezaciaeaaz	195 Op
VBru16F	ggagtttttgttgctcaatgtt	taennee	132 hn
VBru16R	gccatgtttccgttgatttat	aagggagt	152 Up
VBru18F	atgttagggcaatagggcag	nneetnen	137 hn
VBru18R	gatggttgagagcattgtgaag	gagtaagg	137 00
VBru19F	cgacccggaccatgtct	cacgat	161 bn
VBru19R	tcaccgtaacgtcgtggat	cacga	101.00
VBru21F	gctcatgcgcaaccaaa	ctattacc	120 bp
VBru21R	atctcgtggtcgataatctcatt	changee	120 0p
VBru30F	tgaccgcaaaaccatatcc	tacanae	113 hn
VBru30R	aatatgtgcagagcttcatgttc	agggtagt	115 bp
VBru42F	tcgcctcaactataccgtca	aagcgccatcatttttcgtgggtgacgatgacggggatcaagagatcaccccagtt-	162 hn
VBru42R	accgcaaaatttacgcatc	gccgtcgccgttcccgtgttgccgaaggcacacaaagagaagaaacatggcggcgacggcaattgttga	102 op
VBru43F	tcaagcccgatatggagaat	ccatttacccac	152 hn
VBru43R	tccgcctgcccataaac	ecguaceoge	132 op
VBru45F	tcatccttgcctctccctac	caacaacaacacttcaaa	116 bn
VBru45R	gggtaaatatcaatggcttgg	~55~55~5gcgciicggg	110 bp
VBru55F	aggctgtttcgtcatgtcttt	acageacteeacgeetettagttatateeetgeegtgeet	183 hn
VBru55R	tctggcgttcgagttgttc	geggeacheacgeennggnalaneergegegeen	105 00

The nucleotide sequence may contain point replacements. The molecular weight of the amplified fragment consists of the length of the amplicon without taking into account the variable repeats and the value of a particular variable region, with the multiplicity represented in the genome of a particular strain. The analysis of the variable amount of such repeats (determination of the number of these repeats), respectively, was carried out by calculating the difference in the molecular weight of the amplified fragment excluding the variable repeats (Table 3), the resulting digit was divided by the number of nucleotides of the variable region (the nucleotide sequence is represented in Table 3); an example of the calculation of the number of variable repeats is presented below. The typical profile of MLVA for some *B. abortus* and *B. melitensis* strains and isolates is presented at: http://vnivi.ru/images/mlva.xls.

A sample of the calculation of the number of variable repeats upon amplification of the "Bru4" locus, if: the molecular weight of the amplicon is 187 bp; the magnitude of the amplified fragment without taking into account the variable repeats 147 bp; nucleotide sequence of the 8 bp variable region is "gtagggca". The number of variable repeats is calculated as follows: 187-147=40, 40/8=5. The number of variable repetitions of the locus "Bru4" is 5.

3.4 Design and Construction of Positive Control

To verify the correctness of the design of the primers and the probe, the correct composition of the reaction mixtures in PCR and the amplification program, a positive control was created, presented as a plasmid DNA with a specific insert for amplification of bacteria of the genus Brucella and determining their belonging to *B. abortus* and *B. melitensis* (complementary developed primers and probes). The nucleotide sequence of the specific insert is an alternation of loci complementary to direct primers for indicating each of the three DNA markers, then alternation of the loci complementary to the reverse primers to

indicate each of the three DNA markers. Thus, the original nucleotide sequence of all three marker regions is imitated. The specific insert created had the following nucleotide sequence: "5'tcgaatggctcggttgccagcaatcgtcgtattgccactataatcattgcttggcacctcgg aaacaccgatcaagtcgggcgctctggaccgaaaggatcagcgtgccagaatgaagcgctgcagacaaattttgacttccttctggcgacgctttacccgttgttcga-

gaactgcgccgtctcaaactctatcggctttcgggaatg3^{**}. The complementarity of the primers developed and the positive control is illustrated in Figure 2.





Synthesis and an insert into the plasmid vector "pAL2-T" were provided by Evrogen, Russia. As a result, PCR with this positive control accumulates amplification products of molecular weight from 162 bp to 164 bp, which allows to control PCR (with any complex of oligonucleotide primers indicated in Table 1) both in electrophoretic format and in real time.

3.5 Polymerase Chain Reaction with Developed Primers and Probes

The performance of all primers and probes is confirmed by amplification. Primers and probes for indicating bacteria of the genus Brucella and determining their belonging to *B. melitensis and B. abortus* were amplified from positive control DNA and from the DNA of vaccine strains of *B. abortus and B. melitensis*; the amplification result is shown in Figure 3.



Nomenclature: PCR with DNA matrix of *B. suis, B. abortus* and *B. melitensis* (A); PCR with a positive control DNA matrix at a concentration of 1 μ g/ μ l (B); the PCR result for the Fam channel (2, 3, 5, 7); the PCR result for the Rox channel (1, 6); the PCR result for the R6G channel (4, 8); amplification of *B. suis* DNA (3); amplification of *B. abortus* DNA (1, 2); amplification of *B. melitensis* DNA (4, 5).

The PCR used a mixture of primers and probes to simultaneously show (in one tube) the bacteria of the genus Brucella and determine their belonging to *B. abortus* and *B. melitensis*. As a result of amplification with *B. suis* DNA, PCR was performed only with generic primers and a probe (via the FAM channel). Amplification with the *B. abortus* and *B. melitensis* DNA was successful not only for generic primers, but also for specific primers and probes (via Rox and R6G, respectively). Specific amplification occurred only with the desired DNA. With positive control DNA, amplification was successful in all three channels, which confirms their correct design.

The use of primers for MLVA was performed using a DNA of *B. melitensis* strain Rev.1 as DNA matrix; the amplification results (indicating the molecular weight of amplicons and the number of variable repeats) are shown in Table 4.

Locus	Molecular mass of the am-	Number of re-
	plicon	peats
VBru4	155	1
VBru6	390	2
VBru7	150	3
VBru8	330	3
VBru9	150	6
VBru11	260	1
VBru16	150	2
VBru18	170	4
VBru19	160	0
VBru21	150	4
VBru30	150	5
VBru42	410	3
VBru43	180	2
VBru45	150	2
VBru55	260	2

Table 4:. Amplification results for MLVA

Comparison of the MLVA profiles of different isolates and strains of brucellas allows us to establish in a relatively short time (2-3 days) the strain belonging to the analyzed species of brucella, and to find identical, similar or different isolates.

4. Conclusion

In the course of the research work the following **conclusions** were drawn:

1. As a result of the bioinformational analysis, DNA marker loci for indicating bacteria of the genus Brucella and determining their belonging to the species *B. abortus* and *B. melitensis* have been determined. The design of primers and probes for qPCR is made, which allows amplifying these loci.

2. Nucleotide sequences for MLVA are adapted for use in the conventional PCR laboratory settings, under the same conditions and modes of research.

3. A unique positive control, containing the nucleotide sequences of the indicated loci, has been created.

4. The described MLVA-technique allows for effective differentiation of different isolates and strains of brucella and their possible belonging to certain strains of brucella from genomic databases.

Acknowledgements

The work is performed according to the Russian Government Program of Competitive Growth of Kazan Federal University.

References

- Sankarasubramanian J., Vishnu U.S., Gunasekaran P., Rajendhran J. Identification of genetic variants of Brucella spp. through genomewide association studies. Infect Genet Evol., 2017, 56, 92-98, doi: 10.1016/j.meegid.2017.11.016
- [2] Bariev Y.A., Musiev D.G. Comparative sensitivity of PCR and serological reactions to brucellosis. Proceedings of Gorsky State Agrarian University, 2015, 52(1): 107-110
- [3] Shome R., Krithiga N., Shankaranarayana P.B., Jegadesan S., Udayakumar S.V., Shome B.R., Saikia G.K., Sharma N.K., Chauhan H., Chandel B.S., Jeyaprakash R., Rahman H. Genotyping of Indian antigenic, vaccine, and field Brucella spp. using multilocus sequence typing. J Infect Dev Ctries, 2016, 10(3): 237-244, doi: 10.3855/jidc.6617
- [4] Lusk Pfefer T.S., Timme R., Kase J.A. Identification of Brucella genus and eight Brucella species by Luminex bead-based suspension array. Food Microbiol., 2018, 70: 113-119, doi: 10.1016/j.fm.2017.09.015
- [5] Daugaliyeva A., Sultanov A., Usserbayev B., Baramova S., Modesto P., Adambayeva A., Acutis P.L., Peletto S. Genotyping of Brucella melitensis and Brucella abortus strains in Kazakhstan using MLVA-15. Infect Genet Evol., 2018, 58: 135-144, doi: 10.1016/j.meegid.2017.12.022
- [6] Çiftci A., İça T., Savaşan S., Sareyyüpoğlu B., Akan M., Diker K.S. Evaluation of PCR methods for detection of Brucella strains from culture and tissues. Trop Anim Health Prod., 2017, 49(4): 755-763, doi: 10.1007/s11250-017-1256-1
- [7] Sadikalieva S.O., Strochkov V.M., Orynbaev M.B., Shorayeva K.A., Espembetov B.A., Sansyzbay A.R., Sultankulova K.T. Molecular-genetic typing of bacteria of genus Brucella circulated in the Republic of Kazakhstan. Vestnik Bashkirskogo Universiteta, 2017, 22(2): 403-408
- [8] Sankarasubramanian J., Vishnu U.S., Khader L.K., Sridhar J., Gunasekaran P., Rajendhran J. BrucellaBase: Genome information resource. Infect Genet Evol., 2016, 43: 38-42, doi: 10.1016/j.meegid.2016.05.006
- [9] Ndayishimiye E.W., Khammadov N.I., Osyanin K.A., Faizov T.Kh., Shuralev E.A., Mukminov M.N. Bioinformatic analysis of oligonucleotides for molecular genetic indication of honey-bee aspergillosis and ascospherosis pathogens. Veterinarny Vrach, 2015, 2: 3-9
- [10] Khammadova A.V., Shuralev E.A., Khammadov N.I., Oumarou B.M., Faizov T.K., Mukminov M.N. Design of primers for identification of honey bee viruses in multiplex-PCR. Astra Salvensis, 2017, 5(S1): 481-489