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10	Onya Opota, René Brouillet, Gilbert Greub, Katia Jaton
11	
12	Institute of Microbiology, University Hospital Center and University of Lausanne, Switzerland.
13	
14	
15	
16	Corresponding author:
17	Prof. Gilbert GREUB
18	Phone: +41 (0)21 314 49 79
19	Fax: +41 (0)21 314 40 60
20	gilbert.greub@chuv.ch
21	
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24 Summary

25

26 The advances in molecular biology of the last decades have dramatically improved the field of 27 diagnostic bacteriology. In particular PCR-based technologies have impacted the diagnosis of 28 infections caused by obligate intracellular bacteria such as pathogens from the Chlamydiacae 29 family. Here, we describe a real-time PCR-based method using the Taqman technology for the 30 diagnosis of Chlamydia pneumoniae, Chlamydia psittaci and Chlamydia abortus infection. The 31 method presented here can be applied to various clinical samples and can be adapted on opened 32 molecular diagnostic platforms. 33 34 Key words: Chlamydia pneumoniae, Chlamydia psittaci, Chlamydia abortus, molecular

35 diagnostic, real-time PCR, DNA extraction, Taqman

36 **1. Introduction**

37 *Chlamydiacae* are obligate intracellular bacteria, among which several species are pathogenic 38 towards humans and can cause a broad range of diseases. Chlamydia trachomatis is involved in 39 urogenital infection as well as ocular, joint and oropharyngeal infections. Chlamydia 40 pneumoniae and Chlamydia psittaci are primarily associated with community-acquired 41 pneumonia (CAP). C. pneumoniae have been essentially associated with infections in humans [1] 42 but some studies suggest an association with other mammals such as koalas [2-8]. C. 43 pneumoniae might also be a causative agent of an asthma-like syndrome in children [9]. C. 44 *psittaci* is the etiologic agent of a respiratory zoonosis transmitted by birds, in particular parrots 45 and parakeets, but also chicken from the food industries as well as feral pigeons [2]. C. abortus is 46 genetically closely related to C. psittaci but displays a distinct animal and tissue tropism as it can 47 colonize the placenta of cows, goats, cattles, pigs and horses, which can lead to abortion. C. 48 *abortus* can be transmitted to humans, for instance through the exposure to infected animal 49 abortive tissues, with the same outcome for pregnant women [2].

50 The diagnosis of infections due to intracellular bacteria has been dramatically improved by PCR-51 based methods developed in the last decades. This is particularly true for the diagnosis of C. 52 trachomatis, for which several commercial systems have been developed. The laboratory of 53 molecular diagnostic of the Institute of Microbiology of the Lausanne University Hospital 54 (Lausanne, Switzerland), developed a molecular diagnostic platform recently described by 55 Greub and colleagues, which allows one to perform multiple PCRs simultaneously targeting 56 different pathogens, thanks to the standardisation of the parameters (i.e. amplicon length, and 57 primers and probes Tm) of the reactions [10]. This molecular platform based on the Tagmanprobes technology (Applied Biosystems) allows for performance of up to 91 different PCR 58 59 reactions corresponding to 69 pathogens and/or resistance genes in a single microplate [10]. Among them are a real-time PCR for the detection of *C. pneumoniae* [11] and a real-time duplex 60 61 PCR for the detection of C. psittaci and C. abortus infection from various clinical samples 62 (Table 1), both based on the Taqman probe technology [12]. The C. pneumoniae PCR is based 63 on the pst1 gene and the C. psittaci-C. abortus duplex PCR was designed as follows: 1) PCR1 64 targets a DNA sequence of the 16S–23S rRNA operon allowing the detection of both C. psittaci and C. abortus and 2) PCR 2 targets the coding DNA sequence CPSIT 0607 so far unique to C. 65 66 psittaci [12]. In this chapter, the methods to achieve the C. pneumoniae PCR and the C. psittaci*C. abortus* duplex PCR on an opened molecular diagnostic platform are detailed. For this
specific chapter, part of the samples' processing methods and infrastructure descriptions have
been extracted from the accredited documentation of the Laboratory of Diagnostic of the
Institute of Microbiology of the University of Lausanne.

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75 **2. Materials**

76 **2.1 Laboratory organization**

All the procedures should be carried-out according to molecular diagnostic principles aimed to
 avoid contaminations with microorganisms or nucleic acids (See Note 1).

79

80 2.2 Sample processing

- N-acetyl cysteine solution: in a 50 ml conical tube, containing 1.0 g of N-acetyl cysteine,
 add 50 ml of Tris-Sodium-Citrate-di-hydrate buffer. The solution should be used the
 same day.
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 2. Molecular biology grade PBS: DNase-Free, RNase-Free, does not contain detectable
 85 amounts of nucleic acid or any nucleic-acid extractions' compatible molecular biology
 86 grade solution.
- 87

88 2.3 Material for DNA extraction

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 1. Several automated instruments exist for DNA extraction, in the Institute of microbiology,
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 2. Molecular biology grade PBS: DNase-Free, RNase-Free, does not contain detectable
 98 amounts of nucleic acid or any nucleic-acid extractions' compatible molecular biology
 99 grade solution.
- 100

101 **2.4 Assembly of the PCR plate and amplification**

- 102
- 103 1. TaqMan Universal Master Mix (Applied Biosystems).
- 104 2. Primers and probes (Table 1).

105	3.	Molecular biology grade water: DNase- and RNase-free water, which does not contain
106		detectable amounts of nucleic acid.

- 4. Positive control. The positive controls consist of synthetic plasmids containing the exact
 PCR amplicon [12,11].
- 109 5. PCR instrument (see Notes 3 and 4).
- 110
- 111 **3. Methods**
- 112

113 **3.1 Sample processing**

In clinical practice, the detection of *C. pneumoniae*, *C. psittaci* and *C. abortus* can be achieved from a broad range of samples (Table 2). Depending on the nature of the sample (liquid, viscous or solid) a specific processing will be required either to concentrate (liquid samples) or to homogenize (viscous or solid samples) the specimen.

- 118
- Introduce the lab request into the laboratory information system (LIS) to generate the
 barcode labels necessary to manage tubes during the analytic process (See Note 5).
- 121 2. For each sample, label three screw cap tubes (2ml) with the barcode as follows: one
 122 "native" tube, one "native-aliquot" tube and one "DNA tube".
- 123 3. Under the laminar flow dedicated for molecular diagnostics in the "specimen-receiving 124 laboratory," distribute the sample into the native-aliquot tube and the native tube. The 125 native tube will be frozen at -80°C and kept as backup in case of need. If necessary 126 several native tubes can be stored; a suggestion for biopsies would be to store both the 127 native samples and the processed samples in distinct tubes. The aliquot-native tube 128 together with the DNA tube will be transported to the molecular diagnostic laboratory 129 and used for nucleic acids extraction. Viscous samples such as respiratory secretions 130 should be homogenized by liquefaction where as solid samples such as biopsies and 131 fragments should be crushed as described below.
- 4. Liquefaction of viscous secretions: respiratory secretions (sputum, bronchial aspirate, stomach tube) for the detection of *C. pneumoniae* and *C. psittaci* can be too viscous to be homogenized by simple vortexing. Liquefaction can be achieved using N-acetyl-L-cysteine, a reducing substance having free thiol groups (-SH), to reduce specimen

136 viscosity by breaking the disulfide bonds of glycoproteins that constitute the bronchial 137 mucus. Any other viscous fluids shall also be liquefied. To do so, visually control the 138 viscosity of the sample; if liquefaction is necessary, transfer a suitable amount of the 139 sample in a 15 or 50 ml conical tube using a single use plastic Pasteur pipette, under the 140 hood with laminar flow. Add an equal amount of N-acetyl-L-cysteine solution. A larger 141 amount of N-acetyl-L-cysteine solution can be needed for very sticky samples. Vortex 142 and visually check the liquefaction of the sample and if necessary, leave the sample on a 143 rotor for up to 30 minutes. At the end of the incubation, visually check the liquefaction of 144 the sample and centrifuge the sample 30 minutes at 3000 g then remove the supernatant 145 using a pipette with filter tips, leaving 1 to 2 mL of liquid in which the pellet will be 146 resuspended. Transfer the amount necessary for nucleic acids extraction (more than 200 147 μ l) into the tube labelled as aliquot-native" (See Note 6), and keep the rest in the tube 148 labelled as "native". Store the native tube at -80°C in the core specimen receiving 149 laboratory and transport the aliquot-native tube and the DNA tube to the molecular 150 diagnostic laboratory.

151 5. Crushing of solid specimens. Solid samples such as biopsies and fragments that cannot be 152 homogenized either using the vortex or by liquefaction should be crushed as follows. 153 Under the hood with laminar flow put the little pieces in the crushing device with part of 154 the solution of the native sample (if any) or add a solution adapted to the DNA extractor 155 (PBS) using a pipette with filtered tips (see Note 7). If the specimen is too big, take a 156 piece of it with sterile forceps and place it in a sterile Petri dish to cut it into several small 157 pieces. Crush a piece of the specimen and keep the rest in reserve. Start the crushing 158 device. Generally, the fragments are successfully crushed, if not, this procedure is still 159 sufficient to release microorganisms from the specimen by compression. In this case, 160 avoiding some remaining large fragments, transfer the crushed sample (volume) in the 161 aliquot-native tube, and the rest in the native tube for storage at -20° C. If necessary add 162 some molecular biology grade water into the aliquot-native tube to reach the minimum 163 volume suitable for nucleic acids extraction. Vortex the tube. Transport the aliquot-164 native tube and the DNA tube to the molecular diagnostic laboratory.

165

166 **3.2 DNA extraction**

Using the STARlet^R liquid handling instrument, transfer 200 ul of the sample tube in the
 extraction plate of the MagNA Pure 96^R instrument.

- Extraction control (EC). The EC corresponds to a tube submitted to the same extraction
 protocol that the clinical specimen but in which the volume of the clinical sample is
 replaced by an equal volume of a solution adapted to the DNA extractor (PBS). An
 extraction control is needed for each run of extraction (see Note 8). The extraction
 control must be negative when the PCR specific for the pathogen (s) tested is (are) made.
- Transfer the 96-well microplate into the MagNAPure 96^R instrument and start the nucleic acids extraction program according to the user manual. Nucleic acids can be eluted either in 50µl or 100µl of elution buffer depending on the chosen program and are maintained at 4°C (See Note 9).
- 178

179 **3.3 Preparation of the PCR controls**

- Negative control of PCR. Each run of amplification should contain a PCR negative control that consists of the extraction control used as a template. The negative control is used to test the reactivity of the component of the reaction mixture. It should not be contaminated with target DNA and should not allow non-specific amplification.
- 2. Positive control. Positive controls consist of synthetic plasmid DNA containing the target sequence of the PCRs [12,11] (See Note 10). For each run of PCR, three reactions with three dilutions of the positive controls containing 10, 10² and 10³ DNA copies per reaction should be added. They will serve both to generate the standard curve that will be used for the quantification of the positive control and to determine the sensitivity of the reaction based on the positive amplification, the reaction containing 10 copies of DNA.
- Inhibition control. The presence of PCR inhibitors should be tested for each sample. To
 do so the inhibition control reaction consists of a reaction in which 200 copies of the
 positive control is added to the reaction mixture containing the DNA specimen to be
 tested [10].
- 194

3.4 Preparation of the PCR mix, design and assembly of the PCR microplate and
 amplification

197 1. Design of the Tagman PCR plate. This can be achieved using dedicated software such as 198 the SDS 2.4.1 software (Applied Biosystems) allowing the design of either 96 or 384 well 199 PCR plates [10]. The same final volume (20 µl) is convenient for both type of plate. It is 200 strongly recommended to do each analysis in duplicate (or even in triplicate). When an 201 internal control is not used, an additional well for the inhibition control is needed, if not 202 using internal inhibition control. Moreover, there is a need for at least one well for the 203 inhibition control, one well for the negative control and 3 wells for the standard curve. A 204 standard curve is required for each PCR. One regression curve is required for the PCR1 205 and another for the PCR2 [12].

206
2. In the DNA-free laboratory, prepare the PCR mix. A single mix is needed for the detection of *C. psittaci* and *C. abortus*. In a final volume of 20 µl, add 5 µl of the extracted DNA, the forward and reverse primers and the probes at the final concentration indicated in table 2. The inhibition control reaction consists of the same reaction in which, 200 copies of the control plasmids are added. As an extraction negative control the DNA is replaced by the same volume of the extraction control. As a negative control of the PCR, the DNA is replaced by molecular grade water.

- 3. Assembly of the PCR plate. The assembly of the PCR plate can be done manually for 96
 well plates (See Note 11) or using automated instruments for 384 well plates [10]. The
 reactions are achieved in a final volume of 20 µl with 5 µl of DNA sample [12,11].
- 4. Amplification. Run the ABI 7900 instrument or similar thermocycler using the following
 cycling conditions: 2 min at 50°C, 10 min at 95°C followed by 45 cycles of 15s at 95°C
 and 1min sec at 60°C.
- 219

220 **3.5 Interpretation of the results**

- The SDS 2.4.1 software is used for analysis and interpretation of the results.
- 222 The results (qualitative and quantitative) are then checked and introduced into the LIS system.
- 223 At the end of the process the final results are validated by a clinical microbiologist.
- 224
- 1. Analyze the results of the positive controls with adequate software [10].
- 226
 2. Internal quality control. For an analysis to be valid: a) the run should pass the internal
 quality control and b) the positive controls 10 copies must be detected.
 - 9

3. Negative controls. Control that there is no amplification in the negative control reaction.

- 4. Inhibition control. Control that there is amplification in the inhibition control tubecontaining 200 copies of the positive control plasmid (at least of 50 copies).
- 5. Positive result. A result is positive if the fluorescence reaches the threshold automatically
 set by the software or manually set by the user according to the instrument.

233 6. Interpretation of the *C. psittaci-C. abortus* PCR (<u>See Note 12</u>).

- 7. If you suspect a contamination, when for instance only 1 reaction out of 3 reactions is
 positive or when amplification occurs in the extraction control or in the PCR negative
 control, you can follow the procedure described by Greub and colleagues for these
 situations [10].
- 238

4. Notes

240 Note 1

241 As described by Greub and colleagues it is important to organize the laboratory of molecular 242 diagnostic in different rooms/spaces corresponding to pre-amplification and post-amplification 243 area [10]. Moreover, the processing of clinical samples in the laboratory where the samples are 244 received should be achieved under a laminar flow dedicated for molecular diagnostics that is 245 distinct from laminar flows dedicated for conventional culture based microbiology diagnostics. 246 Sterile samples should also be processed in dedicated laminar flows. The use of disposable lab 247 coats, gloves and pipettes with filter tips is recommended. Thus, we recommend the following 248 infrastructures and instruments: A) the sample reception laboratory should be equipped with a 249 hood with laminar flow and UV, a vortex, a crushing instrument, a centrifuge, disposable lab 250 coats and gloves, pipettes and filter tips, 2ml screw cap micro-tubes, sterile forceps and sterile 251 Petri dishes; **B**) the nucleic acids' extraction laboratory should also be equipped with a hood with 252 laminar flow and UV, a vortex, pipettes and filter tips, disposable lab coats and gloves without 253 mineral powders such as talc to prevent any deposition in the extraction tubes; C) the PCR 254 master-mix should be prepared in a DNA free laboratory equipped with a hood with UV but 255 without laminar flow; for all the post extraction area, hood with laminar flow should be 256 prohibited in order to avoid the deposition of nucleic acids (NA) in opened tested tubes. In 257 addition, the following instruments should be available in this laboratory: a vortex, a micro-258 centrifuge, micro-tubes (0.2, 0.7 and 1.5 ml), 2ml screwed tube, pipettes and filter tips (0-10, 2-

259 20, 20-200 et 200-1000 μ); **D**) the positive control should be prepared and stored in a "DNA" 260 laboratory" equipped with a hood with UV but without laminar flow, a vortex, a micro-261 centrifuge, micro-tubes, pipettes and filter tips; E) It is recommended to assemble the PCR plate 262 a dedicated pre-amplification laboratory different from the positive control laboratory. 263 264 Note 2: 265 If processing a large number of samples, it is recommended to select automated distribution 266 systems that can recognize bar-coded tubes. 267 Note 3: 268 269 The assembly of the PCR plate can be achieved manually for 96 well plates or alternatively can 270 be done by automated instruments to increase the precision and to avoid mistakes, especially 271 when preparing 384 well plates [10]. 272 273 Note 4: 274 The PCR conditions described herein for the detection of C. pneumoniae, C. psittaci and C. 275 abortus have been optimized for the PCR instrument ABI 7900 HT (Applied Biosystems) [12]. 276 The reagent concentrations and the amplification program should be adapted if using other 277 instruments [10]. 278 279 Note 5: 280 Tubes management is crucial all along the analytic process to avoid major errors such as 281 inversions or contaminations. We strongly recommend the use of barcoded tubes. 282 283 *Note 6:* If using an automated system such as the STARlet^R instrument (Hamilton^R), do not forget to 284 285 provide the dead volume of the device. 286 287 Note 7: 288 To avoid the dilution of the specimen, which could negatively impact the sensitivity of the PCR, 289 do not add too much solution for the crushing of the sample.

290

291 Note 8:

We recommend placing the extraction control at the end of the series where contamination are more expected to occur rather than at the first position of a series.

- 294
- 295 Note 9:
- 296 It is recommended to store the DNA sample at $-80 \degree C$.
- 297
- 298 Note 10:

299 The concentration (DNA copies per ml) of the positive control should be precisely determined.

300 The stock solution of the positive control should be stored in a separate, dedicated room.

301

302 Note 11:

303 The assembly of the PCR plate can be achieved manually especially for 96 well plates. In the 304 DNA free laboratory prepare the PCR mix without the DNA. Place a 96-well micro-plate on a 305 chilled metal rack and distribute the PCR master mix in the 96-well plate according to the 306 Taqman plate set-up. Add the molecular grade water to the corresponding wells (*i.e* in the 307 negative controls) and transfer the plate with the chilled metal plate in the assembly laboratory. 308 In the assembly laboratory, add the DNA samples starting with the extraction control to avoid 309 any contamination at this stage. Close the tubes when the DNAs of a patient are pipette in order 310 to avoid contamination by aerosols. Repeat this for all the patients' DNAs. When all the patients' 311 DNAs are added, cover the corresponding wells and carefully add the positive control in the 312 wells corresponding to the "inhibition control". It is important to avoid any contamination of the 313 patients' test-tube with the positive controls which would lead to false positive results. Exit the 314 room with the plate in a chilled metal rack and transfer it to the amplification room where the 315 thermocycler is located.

316

317 Note 12:

As described in Opota et al 2015, if both PCR are positive (PCR1 and PCR2) in a respiratory sample, this indicates the presence of *C. psittaci* DNA [12]. Samples positive for PCR1 and

- 320 negative for PCR2 can be considered as positive for *C. psittaci* for respiratory specimens and
- 321 positive for *C. abortus* for genital specimens.
- 322

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- 326

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368

370 Tables

371

Table 1: Primers and probes

Pathogen	Oligonucleotide name	Target gene	Sequence (5'-3')	Modification, fluorochrome	Amplicon lengh	Final Concentration (µM)
	CPTM1	pst1 gene	CATGGTGTCATTCGCCAAGT	-		0.2
C. pneumoniae	CPTM2	pst1 gene	CGTGTCGTCCAGCCATTT TA	-		0.2
	СР	pst1 gene	TCTACGTTGCCTCTAAGAGAAAACTTCAAGTTGGA	3'-VIC, 5'-TAMRA	82	0.1
	CPSI_F	16S-23S operon	AAGGAGAGAGGCGCCCAA	-		0.35
C. psittaci and C. abortus	CPSI_R_LNA	16S-23S operon	CAA[C]CTAGTCAAACCGTCCTAA	LNA		0.35
	CPSI_P_MGB	16S-23S operon	ACTGGGATGAAGTCGTAAC	FAM, DQ	133	0.2
	CPSI_00F	CDS CPSI_0607	AGCATTAGCCAGCGCTTTAGA	-		0.35
C. psittaci	CPSI_00R_147C/G	3CDS CPSI_0607	TCTCTGAGCAAAAAC/GACTGCGT	-		0.35
	CPSI_00P_MGB	CDS CPSI_0607	ACAAAGACCTGGCGAGTA	VIC, DQ	118	0.2

LNA = Locked nucleic acid, MGB = Minor groove binder, FAM = 6-carboxy-fluorescein, VIC = TaqMan VIC reporter dye, DQ = Dark quencher, BHQ = Black hole quencher

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Table 2. Clinical specimens and	nathogons general	ly used for the diagnos	sis of Chlamydia inf	octions
raple 2. Chinical specimens and	pathogens general	ly used for the diagnos	sis of <i>Chianiyula</i> init	actions

	Sample type	C. pneumoniae	C. psittaci	C. trachomatis	C. abortus
Oral	Mouth swab			х	
Respiratory specimen	Nasal swab	х	x		
Respiratory specimen	Nasopharyngeal seretions	х	x		
Respiratory specimen	Sputum	х	x		
Respiratory specimen	Bronchial secretion	х	x		
Respiratory specimen	Bronchoalveolar lavage	х	x		
Urogenital specimen	Uterus and urethral smear			х	х
Urogenital specimen	Urethral swab			х	
Urogenital specimen	Prostate biopsies			х	
Urogenital specimen	Fragment of placenta			х	x
Anal	Anal swab			х	
Osteo-articular	Joint fluid			х	
Osteo-articular	Prosthetic fragment			х	
Vascular	Drain fluid from aortic valve				
Animal specimen	Bird spleen		x		
Animal specimen	Birds choanal or cloacal swabs		x		
Animal specimen	Fragment of sheep placenta				x

Non-comprehensive list