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> To the Editor of Microbes and Infection

#### Département de pathologie et de médecine de laboratoire

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Lausanne, 29 January 2014

Dear Editor,

Please find attached a manuscript entitled " **ESCMID postgraduate technical workshop on intracellular bacteria: from biology to clinic**", that we submit as an article in Microbes and Infection.

In this meeting report, we present the main events that took place during this meeting and describes main scientific presentations.

This manuscript is not submitted or accepted for publication elsewhere. The current version of the manuscript has been seen and accepted by both authors. We hope that this original work will fall within the scope of your Journal.

Sincerely yours,

Gilbert Greub



1	ESCMID postgraduate technical workshop on intracellular
2	bacteria: from biology to clinic
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# 32 Abstract

Infection by intracellular bacteria can lead to several diseases in both veterinary and human medicine. Unfortunately, the biology of these intracellular bacteria is highly complex due to their interactions with their host cells. Thus, it is very important to develop several tools in order to better understand the complex intracellular life of these pathogens, so allowing to improve the diagnosis options and the treatments of infectious diseases that they are causing. The workshop organised in Villars-sur-Ollon (Switzerland) by the ESCMID Study group on intracellular bacteria was a good opportunity to enhance our knowledge on these fastidious pathogens. During 5 days, 15 speakers gave 41 talks, covering all fields, from biology to clinic of different intracellular bacteria such as Bartonella, Chlamydia, Coxiella, Ehrlichia, Listeria, *Parachlamydia, Risckettsia,* and *Waddlia*. The format of this postgraduate course, which took place in the Swiss mountains, allowed interactive sessions and living discussions between the participants coming from all around the world. One of the major strength was to gather epidemiologists, clinical microbiologists, infectious diseases specialists, entomologists, veterinarians as well as bioinformaticians, biochemists and biologists to deliver a unique "one-health science" on intracellular bacteria. Here, we summarize the main take-home messages delivered during this meeting.

- 0-

- 64 Main text
- 65

#### 66 1) Introduction

67 Obligate intracellular bacteria, such as Chlamydia, Rickettsia and Coxiella, have to infect their eukaryotic host in order to survive. Some of them are able to infect a wide range of hosts 68 69 and some may even cause an infectious disease. These infections can be asymptomatic but in 70 most cases, they result in significant deleterious effects for the host, often associated with 71 significant mortality and morbidity. Due to the difficulty to detect obligate intracellular 72 bacteria, they have been often described for the first time during outbreaks. Thus, some 73 intracellular bacteria are likely yet unknown whereas others are still only considered as 74 emerging pathogens since it is difficult to confirm their pathogenic role. Moreover, despite 75 significant pathogenicity of several intracellular bacteria, they are still poorly studied. Indeed, 76 due to the historical lack of genetic tools, and due to the need of cell culture, it is very difficult 77 to study their strict intracellular lifestyle and to precise the virulence factors involved in their 78 pathogenesis. Fortunately, several people are working in this field and try to better understand 79 the biology of intracellular bacteria and the pathogenic mechanisms that are at play.

The ESCMID study group for intracellular bacteria (ESCAR) is currently composed of approximately 300 members, including physicians, microbiologists, veterinarians, and other specialists, interested in the biology, epidemiology, and pathogenicity of *Coxiella, Ehrlichia, Anaplasma, Rickettsia, Bartonella, Chlamydia* and *Chlamydia*-related bacteria. Since the aim of ESCAR is to encourage basic and applied research in the field of intracellular bacteria and related diseases, ESCAR regularly organise postgraduate courses.

86 Thus, an ESCMID postgraduate workshop was organised by Pr. Gilbert GREUB in Villars-sur-87 Ollon, in Switzerland, from 26 to 30 August 2013, and it was entitled "Intracellular Bacteria: 88 From Biology to Clinic". This workshop gathered 55 participants and included interactive 89 sessions, fostering living discussions and a high level of interaction and cooperation about our 90 shared fascinating interest, intracellular bacteria. During this workshop, the sessions 91 alternated between i) biology, with talks about the pathogenesis, molecular biology, genomics 92 and cell biology of intracellular bacteria, and ii) medicine, with talks about clinical 93 presentation, diagnostic approaches, epidemiology and treatment of infections caused by 94 obligate intracellular bacteria.

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# 98 2) Participants and venue

99 The workshop on intracellular bacteria, with 55 participants coming from 21 countries, 100 was organised in Villars-sur-Ollon, in the beautiful scenery offered by the Swiss Alps. The 101 parity was appropriate during this postgraduate course, with 30 men and 25 women, and with 102 people coming from all around the world. Majority of the participants were from European 103 countries, but some people made a long trip, such as 2 investigators from Korea, 3 from 104 Australia, 1 from India, 1 from Sri Lanka, 2 from Tunisia and 1 from USA. The atmosphere was 105 very relaxed and friendly between all participants, allowing fruitful discussions and 106 interactions, particularly during the lunch and diner breaks around delicious specialities like 107 the famous Swiss Raclette (Fig.1). This week of scientific presentations on intracellular 108 bacteria was brighten up by an afternoon of hiking in the pastures, allowing also to know each 109 other better (Fig.1). For the most motivated people, a jogging session was organised all 110 morning and it was also very nice to discuss and to have the opportunity to create some 111 collaborations in parallel to non-scientific activities (Fig.1). The last day, it was time to come 112 down again to the lemanic area, since this day was dedicated to the practical session organised 113 in Lausanne at the University Hospital and at the Institute of Microbiology of the University. 114 The practicals represented a good opportunity for all participants, distributed in small groups 115 of 5 persons, to discover some techniques used to study intracellular bacteria, and/or to 116 improve their practical skills (see below).

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### 118 3) Epidemiology

The workshop was opened by the chairman of ESCAR, Pr. Amel Letaief, and by the organizer, Pr. Gibert Greub. During the first afternoon, talks focused on the epidemiology of infections caused by three different obligate intracellular bacteria: *Rickettsia, Coxiella* and *Chlamydia*.

123 P.E. Fournier (Marseille, France) started this session with an overview of Rickettsia present 124 around the world. Rickettsia are intracellular bacteria widely distributed and vectorized by 125 arthropods such as ticks, fleas and body lice. These bacteria are human pathogens, which may 126 cause either a spotted fever rickettsial infection or the typhus. As underlined by P.E. Fournier, 127 the dogma associating a given rickettsiosis to a given arthropod vector is false, since exceptions 128 may commonly occur and geographic repartition of rickettsiosis has been recently redefined 129 [1]. Arthropods are key vectors for *Rickettsia* but could also be vectors of others intracellular 130 bacteria. This hypothesis was formulated by A. Croxatto (Lausanne, Switzerland) that reported 131 the results of its project investigating the transmission of *Chlamydia* and *Chlamydia*-related 132 bacteria by arthropods. Thanks to a new pan-*Chlamydiales* PCR, Croxatto et *al.* showed the high

133 prevalence (30 to 45 %) of *Chlamydiles* present in ticks collected in Switzerland and in Algeria 134 (Croxatto *et al.* in press). Thus, arthropods and more specifically the ticks, already known as 135 vector for *Rickettsiales*, seem to also act as a reservoir and possibly as a vector for some 136 *Chlamydia*-related bacteria. This is important since human and animals are commonly exposed 137 to ticks. Small ruminants are particularly susceptible to chlamydial and rickettsial infections 138 [2]. D. Longbottom (Edinburgh, UK) focused his talk on five strict intracellular bacteria: (i) 139 Ehrlichia ruminatum responsible for Heartwater, (ii) Anaplasma ovis and (iii) Anaplasma 140 phagocytophilum responsible for Anaplasmosis and Tick-borne fever respectively, (iv) Coxiella burnetii responsible for Q fever, and finally (v) Chlamydia abortus responsible for ovine 141 142 chlamydiosis. These five ovine infectious diseases were described in details including clinical 143 pictures, and approaches used for their diagnosis and treatment options in the veterinarian 144 field. The zoonotic risk was also addressed, being thus an excellent introduction for the second 145 part of the afternoon sessions dedicated to three Coxiella outbreaks.

146 The prevalence of the Q fever is very low in Switzerland but one of the largest Coxiella 147 outbreaks ever described worldwide occurred in 1983 in the "Val de Bagnes", a few dozen of kilometres away from the meeting venue. Up to 415 persons were infected, as underlined by 0. 148 149 Peter (Sion, Switzerland). The source of the outbreak was sheep flocks coming from alpine 150 pastures [3]. More recently, another small outbreak was reported in Lavaux (Switzerland), and 151 G. Greub (Lausanne, Switzerland), who presented this topic, stressed the importance of the fast 152 implementation of different public health measures in order to impact favourably the outcome 153 of outbreaks. Indeed, rapid implementation of public health measures may explain why the 154 outbreak remained limited to only 14 human cases (Bellini et al. New microbes and new 155 infections, in press). Unfortunately, outbreaks do not always have such a favourable issue and 156 may last for years. This is the case of the huge Q fever epidemic that started in Netherlands in 2007. As highlighted by C. Bleekers-Rovers (Nijmegen, Netherlands), due to a delay to 157 158 implement measures and to environmental conditions favourable to bacterial dissemination, 159 the epidemic expanded and more than 4000 cases have been identified to date [4]. The 160 epidemic is now under control but clinicians currently face a subsequent huge outbreak of 161 patients suffering from chronic Q fever. This long lasting Q fever epidemic has been a major 162 opportunity to obtain precise information concerning the control of Q fever outbreaks, the 163 diagnostic, the treatment of the disease and of course, the transmission of *Coxiella*, which may 164 hopefully help public health actors and infectious diseases specialists to avoid another such 165 large epidemic in the future.

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168 4) Clinical presentations, diagnostic approaches and treatments

169 During all morning sessions, speakers detailed the clinical presentation of infections due 170 to different intracellular bacteria and provided state of the art information on diagnosis 171 approaches and treatment options of these infections. Thanks to the huge Dutch Q fever 172 epidemic, a lot of information about *Coxiella* infections has been gathered [4]. As reported by C. 173 Bleekers-Rovers (Nijmegen, Netherlands), the infection is symptomatic in only about 40% of 174 infected people, that present generally with an acute Q fever. This infection is generally 175 diagnosed by using PCR and/or serology, as underlined by O. Péter (Sion, Switzerland) and is 176 resolved spontaneously within 2 or 3 weeks. Despite that, 1-5% of the patients develop a 177 chronic Q fever, so it is necessary to treat acute Q fever for the patients at risk, with 178 doxycycline 200 mg daily for 21 days. In case of chronic 0 fever, the challenge is to early 179 diagnose the infection. Indeed after an asymptomatic phase, severe complications occur with endocarditis as the most common manifestation, leading to high morbidity and mortality 180 181 varying from 13% with a good treatment to 60% in absence of treatment. The recommended 182 treatment is a long-term association of doxycycline and hydroxychloroquine, and the resection 183 of the infected tissues. Two others intracellular bacteria play also a major role in the 184 development of infectious endocarditis, *Bartonella* sp., and *T. whipplei*. Indeed, as explained by 185 G. Greub (Lausanne, Switzerland), 14 species of Bartonella may infect humans, being 186 responsible for several diseases such as Oroya fever (B. bacilliformis), cat scratch disease (B. 187 *henselae*), trench fever (*B. quitana*) [5]. This infection is difficult to diagnose, so most of time it is better to use PCR, serology and/or histology. For uncomplicated cases there is no need of 188 189 antibiotic, but for patients at risk or for specific clinical presentations, personalised antibiotic 190 treatment might be used. T. whipplei are also responsible for endocarditis in case of localized 191 chronic infection, but in the majority (80%) of the cases, the clinical manifestations are non 192 specific, different organs can be targeted and when the gut is involved with associated weight 193 loss, malabsorption syndrome and arthritis, we name this syndrome "classical Whipple's 194 disease". (F. Fenollar, Marseille, France). The diagnosis is mainly based on PCR, and the 195 treatment is empiric but long-term (for lifetime) association of doxycycline and 196 hydroxychloroquine seems to be the ideal treatment.

The diagnosis of blood culture negative endocarditis caused by *T. whipplei* and intracellular bacteria such as *Bartonella* and *Coxiella* is often difficult. Indeed, as reported by P-E Fournier (Marseille, France) (Fig. 2) blood culture being negative, it is necessary to use other diagnostic approaches such as PCR on blood and valves, serology and immunohistochemistry (or autoimmunohistochemistry) on valves [6] [7]. Another interesting clinical characteristic of intracellular bacteria is the ability to induce adverse pregnancy outcomes [8]. D. Baud 203 (Lausanne, Switzerland) (Fig. 2) summarized the role of some intracellular bacteria in humans' 204 and/or animals' miscarriage, stillbirths, and preterm labour. *Listeria monocytogenes* is one of 205 these bacteria. As described by M. Lecuit (Paris, France), this bacterium is very well known as 206 an entheropathogen but it can also be responsible for septicemia, central nervous system 207 infections, and maternal-foetal infections [9]. The diagnosis can be done by culture (blood, 208 cerebro-spinal fluid, placenta,...), PCR, serology, and immunohistochemistry. The 209 recommended treatment is amoxicillin combined to gentamicin for severe infections. Among 210 the other intracellular bacteria implicated in adverse pregnancy outcomes, the importance of 211 *Chlamydia* and *Chlamydia*-related bacteria is increasingly recognised. D. Baud (Lausanne, 212 Switzerland) exposed the characteristics of Chlamydia trachomatis infections that include 213 trachoma and urogenital infections. These infections may be diagnosed by PCR or serology, and 214 are best treated with doxycycline or azithromycin. D. Baud presented also some data about 215 Waddlia chondrophila, a Chlamydia-related bacterium considered as an abortigenic agent in 216 bovine, and associated with human adverse pregnancy outcomes [10] [11].

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#### 5) Cell and molecular biology

219 Several intracellular bacteria are human and animal pathogens. In order to understand 220 mechanisms involved in their pathogenicity, it is essential to study the cellular and molecular 221 biology of these pathogens. To enter, survive and grow within their host cells, intracellular 222 bacteria need to evade the defence mechanisms of these host cells and to generate a replicative 223 niche. As reported by A. Croxatto (Lausanne, Switzerland), secretion systems are one of the 224 tools used by intracellular bacteria to interact with the host cell. Nowadays, seven secretion 225 systems have been identified but Croxatto's talk was focusing on the Type Three Secretion 226 System of the *Chlamydiales* and on the difficulty to identify its effectors. However, it is essential 227 to identify these secreted effectors. Indeed, as explained by J. S. Dumler (Baltimore, USA), the 228 pathogenesis of Anaplasmataceae infections is better understood since the identification of 229 type II and type IV secretion systems effectors, which (i) interfere with host membrane traffic, 230 (ii) interact with MAP kinase signalling pathway, and (iii) bind DNA in the nucleus leading to 231 reduced the transcription of important genes [12]. Chlamydia pneumoniae are other 232 intracellular bacteria using type III secretion system to modulate and interact with host cell 233 signalling pathways. Within their inclusions, these bacteria are able to evade host defence 234 mechanisms and to survive in a persistent stage resulting in a chronic infection. M. Puolakainen 235 (Helsinki, Finland) (Fig. 2) described the particular way of life of these bacteria and the large 236 panel of pathologies associated with chronic chlamydial infections. Always associated to 237 Chlamydia pathogenesis, an important group of proteins specific to Chlamydia was described 238 by D. Longbottom (Edinburgh, United Kingdom). The polymorphic membrane proteins (Pmps), 239 are highly immunogenic and play an important role in bacterial virulence by acting as 240 autotransporter proteins of the type V secretion system [13] [14]. Catalases are another key 241 virulence factor for intracellular bacteria enabling their survival within phagocytic cells, but no 242 catalase have been described so far in classical *Chlamydia*. B. Rusconi (Lausanne, Switzerland) 243 showed the presence of genes encoding for catalases in *Chlamydia*-related bacteria and based 244 on a phylogenetic analysis, she highlighted the important role of these catalases and their 245 evolutionary history in the chlamydial order [15].

246 Virulence is a key feature of intracellular bacteria, and the panel of strategies deployed by 247 these bacteria is huge. The Bartonella genus is composed of 14 pathogenic species, able to 248 infect and multiply within endothelial cells and erythrocytes [16]. Then, as reported by P-E. 249 Fournier(Marseille, France), these bacteria are able to promote angiogenesis allowing 250 dissemination in the host organism, and are vectorized within erythrocytes, by insect vectors 251 such as blood-sucking arthropods [16]. *Rickettsiae* are also transmitted by these arthropods, 252 and then disseminated throughout the body via the bloodstream. These strict intracellular 253 bacteria are able to infect a large panel of cells, but the main targets are endothelial cells. G. 254 Greub (Lausanne, Switzerland) described their capability (i) to corrupt the host cells after 255 escaping to the phagocytosis vacuole, (ii) to induce secretion of proinflammatory cytokines, 256 and (iii) to inhibit apoptosis. Some intracellular bacteria are involved in several serious human 257 pathogenesis such as *Listeria*. M. Lecuit (Paris, France) reported the dual roles of ActA protein 258 involved in the virulence and persistence of Listeria during intestinal infections. Moreover, he 259 detailed the strategy used by *Listeria*, with the interaction between internalin and E-cadherin, 260 to target and cross both intestinal and placental barriers [17]. Finally, pathogenesis induced by 261 intracellular bacteria are sometimes very insidious. As explained by F. Fenollar (Marseille, 262 France), it is the case for *Tropheryma whipplei* pathogenesis. Virtually nothing is known about 263 these bacteria that seems to be opportunistic and might cause chronic infections among 264 genetically predisposed patients.

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#### 6) Genomics

The Wednesday afternoon wad dedicated to talks about genomics of intracellular bacteria. Twenty years ago, it was the beginning of genome sequencing, and this tool is now used almost in routine. Nowadays, we are in a new genomic era focusing the advent of genomics of medical importance [18]. We have access to an ever increasing number of genomes and this provides significant information about intracellular bacteria. Genomics provides an insight in bacterial evolution and bacterial metabolism. Moreover, genomics is a very useful tool for research of new drugs or new drug targets, and provides information on the presence of a variety of virulence factors, including secretion systems, autotransporters, adhesins, and catalases. These virulence factors may be identified by various functional genomics approaches, as outlined by Marie de Barsy [19]. Lessons gathered from genomics of Listeria, Risckettsia, Chlamydia, and Chlamydia-related bacteria have been presented during this session.

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### 280 7) Practicals

The last day of the workshop was dedicated to 9 different practicals (Fig. 3). They were organised at the Institute of Microbiology (IMUL) of the University Hospital Center (CHUV) and at the School of Medicine of the University of Lausanne.

The main techniques for bacterial identification, isolation, staining and observation have been taught with a special focus on specific phenotypes and tools applied to intracellular bacteria, such as *Legionella* and *Listeria*. Participants had the opportunity to have some explanations about bacterial identification by MALDI-TOF

288 Amoebal co-culture and amoebal enrichment have also been presented to the 289 participants, to get them familiarized with approaches used to isolate amoeba-resisting 290 microorganisms (ARM) and to recover free-living amoebae from clinical and environmental 291 samples [20] [21]. To show how to obtain cells from different organisms, allowing further 292 functional assays, isolation of mouse Bone Marrow-Derived Macrophages (BMDM) was shown. Ticks dissection was also explained in details. The importance of techniques allowing 293 294 visualisation of infected cells or bacteria alone was highlighted by a detailed description of the 295 immunofluorescence technique, as well as Gram and Diff-Quick stainings. Finally, bioinformatic 296 tools useful for genome assembly and annotation were presented, emphasizing the importance 297 of an interdisciplinary approach for biological data analysis.

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# 300 8) Conclusions

This Workshop, organised by Pr. Gilbert Greub on behalf of ESCAR, took place in the heart of the Swiss mountains, in a convivial environment, where biologists and clinicians had the possibility to meet, exchange ideas and learn about epidemiology, genomics, diagnosis and treatment of intracellular bacteria. The daily scientific program was brightened up by highly didactic and interesting clinical quizzes. Every day, participants were able to interact thanks to evening extra-activities such as ping-pong, swimming and running. Not less worthy was the Thursday afternoon, spent in mountains surrounding Villars-sur-Ollon. The next postgraduate technical workshop organised by ESCAR will be on the practical diagnosis of arthropod-borne
infections, and will take place in Marseille (France) from 17<sup>th</sup> to 19<sup>th</sup> of March 2014. For sure,
this will be another unique occasion to familiarize with intracellular bacteria, their intriguing
biology and the important diseases they cause.

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9) Acknowledgements

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375		Figure 1: Participants and venue. The top part is on overview of the hiking in the Swiss	
376	pastures. On the bottom left, the small but motivated group of early risers joggers, and on the		
377	bottom right, the friendly atmosphere that was present during lunch and diner.		
378			
379	Figure 2: Talks and speakers. On the top left, the beautiful presentation room with		
380	attentive participants, and three of the speakers, (i) Dr. M. Puolakainen (Helsinki, Finland), (ii)		
381	Dr. P.E. Fournier (Marseille, France), and (iii) Dr. D. Baud (Lausanne, Switzerland).		
382			
383		Figure 3: Practicals. An overview of the nine practicals organized in the Institute of	
384	microbiology of the University Hospital Center of Lausanne (IMUL-CHUV). From top to the		
385	bottom and left to the right: (1) dissection of ticks, (2) Listeria, phenotypes and phenotypic		
386	identification in the diagnostic laboratory, (3) amoebal enrichment, (4) immunofluorescence,		
387	(5) amoebal co-culture, (6) cell culture: isolation of bone marrow-derived macrophages, (7)		

doing a Gram and Diff-Quick staining, (8) bioinformatics for dummies: how to assemble a
genome and how to annotate a genome, and (9) MALDI-TOF identification of strict intracellular
bacteria: the *Chlamydiales* example.





