1 Anti-Coccidiosis Potential of Autoclaveable Antimicrobial Peptides from Xenorhabdus budapestensis Resistant to Proteolytic (Pepsin, 2 Trypsin) Digestion based on in vitro studies 3 4 5 András Fodor^{1*}, László Makrai², László Fodor², István Venekei³, Ferenc Husvéth⁴, László Pál⁴, Andor Molnár⁴, Károly Dublecz⁴, Csaba Pintér^{4^} Sándor Józsa^{4^} and Michael G. Klein⁵ 6 7 8 9 ¹1Department of Genetics, Faculty of Science and Informatics, University of Szeged, 10 University of Szeged, Közép fasor 52, Szeged, H-6726, Hungary, 11 12 ²Department of Microbiology and Infectious Diseases, University of Veterinary Medicine 13 Budapest, Hungary ³Department of Biochemistry, Eötvös Loránd University, Budapest H-1117, Hungary 14 15 ⁴Department of Animal Sciences and Animal Husbandry, Georgikon Faculty, University of Pannonia, Széchenyi Street, 11 Keszthely, H-8360 Hungary). 16 17 ⁴Adjunct Emeritus Professor, Georgikon Faculty, University of Pannonia, Széchenyi Street, 11 Keszthely, Hungary (The place of the in vivo experiment) 18 19 ⁵Adjunct Professor, Department of Entomology, The Ohio State University, 1680 Madison Ave., Wooster, OH-44691, USA 20 21 22

24 ABSTRACT

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Aims: To elucidate the anticoccidial potential of antimicrobial peptides from *Xenorhabdus budapestensis* on both causative pathogens (prokaryotic *Clostridium perfringens* and eukaryotic *Eimeria tenella*). **Objectives:** (1) To establish if the antimicrobial compounds of the cell-free culture media (CFCM) of the entomopathogenic symbiotic bacterium species, *X. budapestensis* DSM 16342 (EMA) and *X. szentirmaii* DSM 16338 (EMC) were active against 13 independent pathogenic isolates of *Clostridium perfringens in vitro*; (2) To create a sterile, autoclaved, bio-preparation called "XENOFOOD", for future *in vivo* feeding studies, aimed at determining the efficacy, and side-effects, of EMA and EMC on *C. perfringens* in chickens.

Study design: Clostridium perfringens samples (LH-1-LH24) were collected from chickens and turkeys, and were deposited in the frozen stock collection of Department of Microbiology and Infectious Diseases, Faculty of Veterinary Science, Szent István University, Budapest, Hungary, where the *in vitro* assays were carried out on 13 of these isolates.

Place and Duration of Study: Department of Microbiology and Infectious Diseases, Faculty of Veterinary Science, Szent István University, Budapest, Hungary between September 2013 and February 2014.

Methodology: Adaptation of our previously published *in vitro* bioassays for aerobic tests for the anaerobic bacteria *Clostridium perfringens*. When preparing "XENOFOOD" we benefitted from our experimental data about the heat tolerance and endurance to proteolytic enzymatic digestion of the studied antimicrobial peptides.

Results: The studied antimicrobial peptides were heat-stable, trypsin and pepsin resistant. All but one of 13 *C. perfringens* isolates was sensitive to EMA-CFCM. XENOFOOD (made here) is not toxic for chicken, (unpublished).

Conclusion: Since these cell-free cultures killed *E. tenella* cells, but were toxic to permanent chicken liver (LMH) cells, we need to run *in vivo* feeding tests to determine the gastrointestinal (ileac), anti-*Clostridium* and anti-*Eimeria* biological effects of the these heat, - and proteolysis tolerant antimicrobial peptides.

26 27

Keywords: Clostridium perfringens, Xenorhabdus Antimicrobial Peptides; in-vitro Bioassay, Xenofood

28 29 30

31 **1. INTRODUCTION**

32

33 Multi-drug resistance (MDR) has gradually been increasing in both Gram-positive [1] and 34 Gram-negative [2] pathogenic bacterium species. MDR has always been a phenotypic 35 consequence of sequential accumulation of simultaneously appearing mutations, or the up-36 take of resistance plasmids harboring mobile genetic elements or genomic islands with 37 resistance genes. These encode for either enzymes capable of destructing the antibiotics, or 38 catalyzing biochemical reactions resulting in inhibition of either binding to, or permeation 39 through, the cellular membrane (CM). The poultry gastro-intestinal (GI) flora is a seed-bed 40 of MDR, as shown by the spectacular on-going evolution in Enterococcus [3] [4] [5], in 41 Clostridium [6], as well as in Salmonella genera [7]. The explanation is that the poultry GI is 42 an ideal "market place" for exchange and horizontally transferring resistance gene -carrying

43 plasmids, and mobile genetic elements, between coexisting bacteria. Enterococcus

44 cecorum, for instance, once a simple commensal member of the intestinal microbiota, has

45 become the causative pathogen of arthritis and osteomyelitis worldwide in chickens, such as

in Hungary [8] and Poland [9]. Evidences of multidrug-resistant plasmid transfer from Gram

47 positive [10] and Gram negative [11] [12] chicken pathogens via consumed chicken meat to

48 human pathogens, has been accumulating. Apart from the veterinary aspects, this horizontal

49 gene transfer is of critical clinical importance.

50 The anaerobic, Gram-positive, C. perfringens was first published as a globally threatening

51 danger by Van Immerseel and his associates, [13] as the causative pathogen of necrotic

52 enteritis. Since then it has become alarming from both veterinary and human clinical

aspects. The incidence of *C. perfringens*-associated necrotic enteritis in poultry has also

54 increased in countries that stopped using antibiotic growth promoters. Both the disease and

its subclinical forms are caused by *C. perfringens* type A strains, which produce either the

alpha toxin, (to a lesser extent type C), or both alpha and beta toxins [14]. A few C.

57 *perfringens* type A isolates produce an enterotoxin at sporulation as well, causing disease in

58 humans, [14].

As for the pathogenesis of necrotic enteritis in chickens [15], it is a result of a "joint venture"

60 the eukaryotic *Eimeria* species and *C. perfringens*, [16] [17]. The lesions and damages in the

61 gut wall tissues (mainly in the lamina muscularis mucosae and in the lamina mucosa)

62 provide anaerobic conditions needed for propagation of the toxin-producing *Clostridium*,

63 especially in the ileum. The *Eimeria* (most frequently) *tenella* infection is usually preceded by

64 previous unfavorable changes in the GI biota. The latter might be an indirect consequence of

65 non-appropriate diets which increases the viscosity of the intestinal contents and makes it

66 predisposed to necrotic enteritis [15]. This important discovery provides an opportunity for

67 nutrient scientists to help reduce *Clostridium* infections. In other words, the discovery that

68 the gastrointestinal microbiota could significantly be restructured by nutritional factors,

69 provides additional opportunities for nutrition scientists working on the problem of coccidiosis

70 [18] [19] or similar problems such as *Campylobacter jejuni*, [20]

71 *Clostridium perfringens* type A cells release different toxins that causing diseases not only in 72 chickens, but also in humans. One of them, the necrotic enteritis B-like toxin (NetB), is a β -

barrel pore-forming one, which used to be a candidate vaccine [21]. Another one, called

74 perfringolysin O (PFO, also called θ toxin), is a pore-forming cholesterol-dependent cytolysin

75 (CDC) [22]. PFO is secreted as a water-soluble monomer that recognizes and binds

76 membranes via cholesterol. Membrane-bound monomer molecules undergo chemical

577 structural changes that culminate in the formation of an oligomerized pre-pore complex on

the membrane surface [22]. The pre-pore then undergoes conversion into the bilayer-

spanning pore, playing an important role in so-called gas gangrene progression and necro-

80 hemorrhagic enteritis in some mammals [22].

81 Clostridium perfringens strains which were isolated from epidemic outbreaks of necrotic

82 enteritis, and were capable of secreting factors that inhibit growth of other (competitor) C.

83 *perfringens* strains, including those isolated from the guts of healthy chickens [23]. This

84 feature lends a selective virtue to respective NetB-toxin producing virulent strains, the

85 causative factor of gut lesions. The factor providing this selective virtue to the virulent strains

- 86 is a novel, chromosomally encoded, heat-labile, trypsin and proteinase-K sensitive protein
- 87 with bacteriocin activity called perfrin [23]. The gene, which can only be found in *C*.
- 88 perfringens NetB strains and nowhere else, (despite the fact that the NetB is a plasmid
- 89 encoded toxin), could be transferred to and expressed in *E. coli*. Theoretically, it may also
- 90 happen in the chicken GI at any time, and the recombinant gene product is antibacterial

91 active at a large pH range [23].

- 92 Several data from the literature seem to support our opinion that although vaccination is an
- 93 effective, but probably not an omnipotent, veterinary tool for controlling Gram-positive MDR
- 94 pathogens such as *Clostridia*. The vaccination projects involving *Enterococcus* seem to be in
- a promising, but only very experimental stage [24]. (None of the seven respective

96 publications have recently been available in PubMed include anything on poultry).

- 97 As for *Clostridia*, the vaccination of chickens against the fatal human pathogen type C
- 98 (causing botulism), have fortunately been successful [25]. The vaccination against C.
- 99 *perfringens* however, although seeming to be not too far from realization, but maybe not in
- 100 the near future. The immunization with NetB genetic, or formaldehyde toxoids, seemed to be
- 101 the most plausible approach [26], but only the double vaccination (on 3 and 12 days, with
- 102 crude supernatant), were effective. Immunization with a single toxin molecule did not give
- 103 satisfactory protection to chickens against necrotic enteritis lesions, which probably is not a
- 104 realistic option for practical application [27].
- 105 This observation led Professor Dr. Van Immersee (Universiteit Gent, Belgium) and his
- associates to the conclusion that "immunization with single proteins is not protective against
- 107 severe challenge. Therefore combinations of different antigens are needed as alternative. In
- 108 most published studies multiple dosage vaccination regimens were used. It is not a relevant
- 109 way for practical use in the broiler industry", [28]. Some other less pessimistic reports, such
- as suggesting the use of C. *perfringens* recombinant proteins in combination with
- 111 Montanide[™] ISA 71 VG adjuvant as a vaccine [29] or anticoccidial live vaccine [30] have
- been noted. Nevertheless, we think that we'd better to accept the opinion of the Expert #1 in

that research field: the vaccination against avian *C. perfringens* type A strains in broiler

- 114 chicken is not yet available [28].
- 115 Consequently, there is a room for working on novel antimicrobials, especially on novel
- antimicrobial peptides which might be used to control *C. perfringens* A and also MDR
- 117 pathogens in the GI system of broiler chickens. This approach needs a comprehensive
- 118 strategy, based on Quantitative Structure Activity Relation (QSAR) analysis and in silico
- modeling [31]. Chemical synthesis of modified analogs leading to new antimicrobial agents
- 120 with novel modes of action should follow the molecular design to get new antimicrobial
- 121 peptides, [31]. The structural design of AMP candidate molecules has aimed at improving
- 122 endurance to proteolytic degradation, binding to, and the penetration through cellular
- membranes and other biological barriers [32]. This can be achieved by adding modules for
- 124 passive or active transport [32]. Another approach is searching for efficient synergisms [33].
- 125 Another (ever-green) alternative research line is to search for new antimicrobials of
- 126 completely novel modes of action in nature. Our research team has been searching for novel
- 127 antimicrobials, which are not used in human medicine, are toxic only for chicken pathogens,
- 128 but not toxic for the organisms to be protected. We expect to find the best candidates among

129 the natural antimicrobial peptides (AMPs), synthesized by the obligate bacterial symbionts 130 (EPB) of entomopathogenic nematodes (EPN) [34]. These EPB-released AMPs are 131 evolutionary products developed under severe selective pressure, and comprise a powerful 132 chemical arsenal against a large scale of prokaryotic and eukaryotic organisms. They 133 provide monoxenic conditions for a given respective EPN / EPB symbiotic complex in 134 polyxenic (insect gut, soil) conditions. Many EPN-EPB complexes exist, and many AMP 135 profiles could be determined. Considering that all but one [35] of the known AMPs can be 136 produced by the bacterium in vitro, the EPN/EPB complexes provide a gold mine for 137 researchers interested in new antimicrobials. The majority of EPB-produced AMPs were 138 identified in the last 15 years [36] [37] [38] [39]. Each of these evolutionarily designed 139 antibiotic arsenals has effectively overcome intruders representing a full scale of antibiotic 140 resistance repertoire in their respective niche. Each EPB-AMP discovered so far is a non-141 ribosomal peptide (NRP), synthesized by multi-enzyme thiotemplate mechanisms, using 142 non-ribosomal peptide synthetases (NRPS), fatty acid synthases (FAS), and / or related 143 polyketide synthases (PKS), or a hybrid biosynthesis thereof [40]. The biosynthetic enzymes 144 are encoded by gene clusters [41], determining the biosynthetic pathways.

145 Cabanilasin, from *X. cabanillasii*, exerts of a strong antifungal activity [42]. In our

146 experiments, the cell-free culture media (CFCM) of *X. cabanillasii* was also extremely toxic

147 to Staphylococcus aureus, Escherichia coli and Klebsiella pneumoniae, isolated from cows

with mastitis syndromes [43]. In that experiment, the antibacterial activities of the CFCM of
 several *Xenorhabdus* species were compared.

150 We found that and those of X. budapestensis DSM 16342 (EMA), and X. szentirmaii DSM 151 16338 (EMC) [44] proved far the best. The CFCM of EMA and EMC were also effective 152 against S. aureus MRSA, (Fodor, McGwire and Kulkarni, unpublished). Furthermore, the 153 CFCM from EMA and EMC also was effective against plant pathogens, including both 154 prokaryotic Erwinia amylovora, E. carotavora, Clavibacter michigenense and several 155 Xanthomonas species [45] [46] [47] and all tested eukaryotic Oomycetes (Phytophthora) 156 species [42] (Muvevi et al., <u>unpublished</u>). Gualtieri confirmed our data, declaring that X. 157 szentirmaii DSM16338 (EMC) was really a source of antimicrobial compounds of great 158 potential, and he sequenced this strain [48]. One of the products (szentiamide) has been 159 chemically synthesized [46].

160 We suppose that these antimicrobial peptides act in concert. The idea of a preparing a bio-161 product for oral administration to via chicken food, ("XENOFOOD"), is based on the intention 162 to benefit from the joint action of cooperating AMP molecules produced by EMA and EMC 163 cells, not only on a single molecule. We know that the strongest, predominant antibacterial 164 peptide produced by both EMA and EMC species is fabclavine [51], but there are also others 165 acting on eukaryotic pathogens as well, especially in EMC [48] [49]. (This is the explanation 166 why we did not use only EMA CFCM alone, but a mixture of EMA and EMC CFCM instead in 167 the experiments reported here).

168 Many of our experiments with EMA were repeated in the laboratory of Professor Helge B

169 Bode (Goethe-Universität, Frankfurt – am – Main, Germany). They confirmed that EMA

170 CFCM exhibited broad-spectrum bioactivity against Bacillus subtilis, E. coli, Micrococcus

171 Iuteus, Plasmodium falciparum, Saccharomyces cerevisiae, Trypanosoma brucei, and T.

- 172 cruzi [51] as well. They subjected the CFCM from X. budapestensis to MALDI-MS analysis
- 173 and found altogether 4 isomers of fabclavine, one of which was then purified, and its
- 174 structure was determined. The details of biosynthesis were impressively reconstructed by
- the authors, but no data about the mode of action has been published so far [51].
- 176 Fabclavines are considered a novel class of biosynthesized hybrid peptide-polyketide-
- 177 polyamino natural compounds with extremely high antimicrobial potential in both prokaryotic
- and eukaryotic pathogen targets, but also with unwanted eukaryotic cell-toxicity. They are
- 179 unambiguously the most effective antimicrobial *Xenorhabdus* peptide-products that have
- 180 ever been discovered, and they are released by *X. budapestensis* and *X. szentirmaii* [44].
- 181 (This is a spectacular example of present-day science, when on group of scientists are
- 182 "sowing" while the other ones are "harvesting").
- We tested CFCM of EMA and EMC were in 2009 in the McGwire laboratory (Ohio State
 University, Columbus, OH, USA) against different targets, and found that, similarly to several
- 185 other antimicrobial peptides [52] [53] they exerted apoptotic effects on eukaryotic cells of
- 186 Leichmania donovanii. They were also active against Candida sp., and Phytophthora
- 187 *infestans* (A. Fodor et al., unpublished).
- 188 Considering that not only prokaryotic, but eukaryotic pathogens also exist, we decided to 189 continue the "EMA-EMC" project. Coccidiosis is the best example of when a prokaryotic and 190 a eukaryotic pathogen act together. Dr. Petra Ganas tested both CFCMs on a permanent 191 chicken liver cell line at the Vet Med University of Vienna, Austria, and found them toxic to 192 the tissue cultures (Ganas, personal communication, for details, see Discussion), even if the 193 toxic cell concentration was 1 order of magnitude higher than the bactericide concentration. 194 These data, and the identification of the most active component (fabclavine), might seem 195 discouraging for the continuation of the project.
- 196 However, considering the presence of multidrug resistance, and even pan-resistance, 197 problems in the GI system of broiler chicken, which may also threaten human health, and the 198 limitation of vaccinations, we reconsidered it as a potential tool, on the prospects that orally 199 applied compounds would not be absorbed into the meat of broiler chickens. Prior to in vivo 200 feeding tests we carried out the *in vitro* bioassays presented here, and formulated a chicken 201 food, Xenofood, to test in the in vivo tests. From this aspect, we believe that the results of 202 this in vivo experiment are worthwhile, and our conclusions will be taken into consideration 203 by coccidiosis specialists.
- 204

205 2. MATERIAL AND METHODS

206

207 **2.1 Bacterium Strains**

- 208
- 209 Clostridium perfringens NCAIM 1417 strain was obtained from the National Collection of
- 210 Agricultural and Industrial Microorganisms –WIPO (of Hungary, Faculty of Food Sciences,
- 211 Szent István University Somlói út 14-16 1118 Budapest, Hungary). Clostridium perfringens
- 212 LH1-LH8; LH11-LH16; LH19, and LH20 are of chicken origin, and LH24 came from a pig;
- 213 each has been deposited in the (frozen) stock collection of Department of Microbiology and
- 214 Infectious Diseases, University of Veterinary Medicine Budapest, Hungary.

- 215 Xenorhabdus strains, X. budapestensis DSM 16342 (EMA), X. szentirmaii DSM 16338
- 216 (EMC) [44] and X. bovienii NYH which had been isolated from the entomopathogenic
- 217 nematodes Steinernema bicornutum [Tallósi] [54], S. rarum and S. feltiae HU1 [55], are
- 218 originated from the Fodor laboratory, Eötvös University, Budapest, Hungary. EMA and EMC
- 219 had also been deposited by us in the DSMZ, (Leinbniz Institute Deutsche Sammlung von
- 220 Mikroorganismen und Zellkulturen, Braunchweig, Germany) as DSM 16342 and DSM
- 221 16338, respectively. Xenorhabdus nematophila ATTC 19061, was from Forst Laboratory at
- the University of Wisconsin Milwaukee, USA) and X. nematophila DSM 3370 DSMZ,
- Braunschweig, Germany). *Steinernema cabanillasii* BP was isolated by us from infective
 dauer juveniles from the EPN *S. riobrave*.
- 225 **2.2 Overlay Bioassays for Comparing the Antibacterial Potential of Different**
- 225 **2.2 Overlay Bioassays for Comparing the Antibacterial Potential of Differer** 226 **Xenorhabdus Strains**
- Overlay bioassays for comparing the antibacterial potential of different *Xenorhabdus* strains
 (each representing a species), were carried out as previously described [43]. To make sure
 that we use the proper bacterium, an earlier experiment was repeated in which we compared
 the antibacterial activities of 5 different *Xenorhabdus* strains on *K. pneumoniae*.
- 231 To determine if the antimicrobial compounds from EMA were effective against *C*.
- 232 *perfringens*, an overlay experiment was carried out [43]. To be sure that the intestinal
- 233 proteolytic activities would not inactivate our compounds, samples of EMA CFCM were
- 234 digested with pepsin, following the professional guidance of our coauthor Professor Ferenc
- 235 Husvéth (University of Pannonia, Keszthely, Hungary), while another sample was digested
- 236 with trypsin by István Venekei (Eötvös University, Budapest, Hungary)

237 2.3 Agar-Diffusion Assay of EMA CFCM against *Clostridium perfringens* NCAIM 1417 238 Laboratory Strain

- Agar Diffusion Tests were similarly carried out, as described by [46], but we converted the method for the anaerobic specimen, *C. perfringens*. An agar diffusion test was conducted as follows: In a hole at the center of the agar plate, 100 ul of EMA CFCM were pipetted and overlaid with 3 ml of a log phase *C. perfringens* suspension diluted to 1:250 with soft (0.6
- 243 V/V%) agar. They were incubated for 24h under anaerobic conditions at 40 °C.

244 2.4 Comparison of the Sensitivities (MID Values) of 13 *C. perfringens* Strains, Isolated 245 from Poultry, to Cell-Free Culture Media (CFCM) of *X. budapestensis* (EMA) in Liquid 246 Cultures

247 2.4.1 Determination of MID Values

248 To quantify the sensitivity of the strains, the maximum inhibiting dilution (MID) values [43] 249 [56] [46] [47] were determined as below. These studies were carried out in sterile 24-hole 250 tissue culture plates, with 4 (A-D) rows and 6 (1-6) Columns, in 1 ml final volumes. Each 251 Clostridium strain was used in a different tissue culture plate. Each hole contained 0.5 ml of 252 2XRCM Reinforced Clostridium Media [57] liquid medium, and 0.5 ml of sterile, diluted EMA 253 CFCM, with the following distribution: 100, 80, 60, 40, 20 and 0 volume / volume (V/V) % in 254 column 1, 2, 3, 4, 5, and 6, respectively. There were 50, 40, 30, 20, 10 and 0% V/V final 255 concentration of EMA CFCM in columns 1, 2, 3, 4, 5, and 6. Each culture in rows A, B and C 256 were inoculated with loopful of the respective bacteria obtained from three separate colonies 257 grown on sheep blood agar plates. The holes in row D were not inoculated, and served as 258 sterile (negative) controls. Columns 6 served did not contain EMA CFCM and served as

259 positive controls. Each 1-ml culture was overlaid by 0.5 ml sterile (freshly autoclaved),

260 paraffin oil to provide anaerobic conditions. Plates were then incubated at 37°C for 24h and

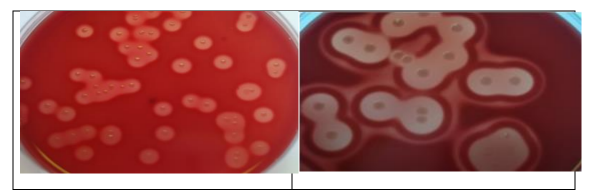
then scored visually. After 24h culturing, the growing and inhibited cultures could

unambiguously be identified. We considered the concentration as MID where none of the 3

263 replicates contained visible growth.

264 2.4.2 Enumeration of Clostridium perfringens colony forming units (CFU)

265 Samples were taken from the first hole in which bacterial proliferation was not visually 266 detected. 0.5 ml of culture were sucked out cautiously from below the paraffin oil and serial dilutions were prepared up to 10⁻⁵, and 100 µl volumes were simultaneously spread onto the 267 surface of sheep blood agar (by D. László Makrai, see Fig. 1) and Tryptose-Sulfite-268 269 Cycloserine (TSC) agar [59] plates. The latter was designed as a highly selective solid 270 medium for growing and enumerating C. perfringens colony forming units. The TSC allows 271 virtually complete recovery C. perfringens, while it inhibits practically all facultative 272 anaerobes tested, and is known as being more selective than SFP Agar. Three replicates 273 were used for each dilution. In preliminary experiments, carried out by András Fodor and 274 Andor Molnár, both then at the Department of Animal Sciences and Animal Husbandry 275 (Georgikon Faculty, University of Pannonia, Keszthely, Hungary), TSC plates were 276 incubated under anaerobic conditions at 40°C, and found the best readability between 48 -277 72h. The C. perfringens colonies were recognized by colony color and the black reduced 278 sulfides granules around them, but the color of the agar also gave a kind of qualitative 279 information (Fig. 1). The colonies used in these preliminary experiments were obtained from 280 chicken ileal digests, and from the stock collection of Dr. L. Makrai, were reproducibly 281 counted.



282

Figure 1 shows the *Clostridium* colonies to be counted on a blood agar plate (Photo: Dr.
László Makrai, (Department of Microbiology and Infectious Diseases, University of
Veterinary Science, Szent István University, Budapest, Hungary).

285 Veterinary Science, Szent Istvan University, Budapest, Hungary).

286 2.5 Study of the Endurance of the Antimicrobial Compounds in the Cell-Free Culture 287 Media (CFCM) of *X. budapestensis* and *X. szentirmaii* to Proteolytic Degradation

288 <u>2.5.1. Trypsin-digested samples</u> were tested on Gram-positive (*Staph. aureus*) and Gram
 289 negative (*E.coli*) targets in agar diffusion assay, and compared with untreated CFCM
 290 samples. No differences were demonstrated.

291 <u>2.5.2. Pepsin resistance</u> was studied as follows: in the center of a Luria Broth plate, a
 292 Millipore filter of 0.22 um pore size was laid and infiltrated with HCl and pepsin. Then EMA
 293 CFCM was pipetted onto it. The pepsin preparations were prepared by Professor Ferenc
 294 Husvéth. After that the plate was overlaid with a *Pseudomonas aeruginosa* suspension
 295 diluted with soft agar as described [46] [47]. After 24 h incubation at 40 °C, the growth of the
 296 test bacterium lawn was checked.

297 2.6 Preparation of XENOFOOD

298 299 XENOFOOD: XENOFOOD contained 5% soy-meal, which had been suspended with equal 300 amount (w/w) of EMA, and another 5% suspended in equal amount (w/w) of EMC cells 301 obtained from 5 days-old shaken (2000 rpm) liquid cultures, followed by high-speed 302 (Sorwall; for 30 minute) centrifugation. The liquid cultures were in 2XLB (DIFCO), 303 supplemented with meat extract equivalent to the yeast extract. Five days was optimal for 304 antibiotic production at 25°C under these conditions [43] [45]. It had previously been 305 discovered that both EMA and EMC grow and produce antibiotics in autoclaved soy-meal containing some water and yeast extract, or in autoclaved 0.5% w/w yeast (Fodor, 306 307 unpublished). Therefore the original chicken food served as a semi-solid culture media for 308 the Xenorhabdus cells. Both the separate EMA and EMC culturing semi-solid chicken food that we (Dr. László Pál) prepared daily were incubated under sterile conditions for another 309 310 five days. Then the EMA and EMC culture media were combined, autoclaved (20 min, 121 311 °C), and then dried by heat (70°C) overnight. The Xenorhabdus cells were killed in such a 312 way, while the heat stabile [43] antimicrobial compounds remained active.

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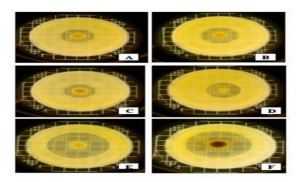
314 2.7. Statistical Analysis

315 ANOVA procedures were used following the procedures of the SAS 9.4 Software, mostly 316 due to the unbalanced data set. The significant differences ($\alpha = 0.05$) between treatment 317 means were assessed using the Least Significant Difference (LSD).

318 3. RESULTS

319 320 3.1. Results of Experiments, Aimed at Helping to Choose the Best Xenorhabdus 321 Strains for This Study

Results shown in Fig. 2, and a qualitative evaluation of the inactivation zones, indicated the appropriate bacteria to use. As expected, *X. budapestensis* (EMA) and *X. szentirmaii* were the best. Results of the overlay bioassay experiment with different *Xenorhabdus* strains on *K. pneumoniae* helped to make the right decision when choosing antimicrobial producing strains.



A: X. nematophila DSM 3370

B: X. cabanillasii BP

C: X. nematophila ATTC19061

- D: X. bovienii NYH
- E: X. budapestensis DSM16342^T
- F: X. szentirmaii DSM16338^T

327

- 328 Figure 2 Comparison of the antimicrobial potential of different Xenorhabdus strains
- 329 (representing species) in overlay bioassays [43]. (Photo: Andrea Máthé Fodor. The Ohio
- 330 State University, Wooster, OH, USA)

331 3.2. Endurance of the antimicrobial peptides of *X. budapestensis* to pepsin, - and 332 trypsin digestion

- 333
- As demonstrated by Fig. 3, the overnight pepsin-digested EMA CFCM remained active
- 335 against Pseudomonas aeruginosa. The trypsin-digested samples also preserved their anti-
- 336 Gram-positive (on S. aureus) and anti-Gram-negative (Es. coli) activities, (not shown).

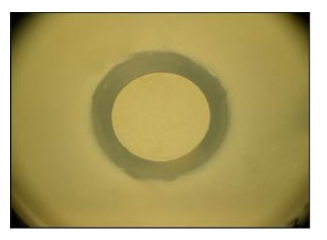


Fig. 3. Experimental evidence that the antimicrobial compounds of *X. budapestensis* cell-free media are resistant to the proteolytic activity of pepsin. After 24 h incubation at 37 °C a large inactivation zone could be seen, demonstrating a significant antimicrobial activity of the pepsin-treated EMA CFCM.

337 **3.3. Efficacy of EMA CFCM on** *C. perfringens* Laboratory Strain NCAIM 1471

The cell-free EMA CFCM exerted strong antimicrobial activity on *C. perfringens* laboratory strain NCAIM 1471 in an agar diffusion test. The large inactivation zone of 3.7 cm diameter shows the anti – *Clostridium* activity (Fig 3). The question arises as to whether the pathogenic poultry isolates were also sensitive.

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- 343



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Figure 4. Anti- Clostridium activity of cell-free culture medium of Xenorhabdus budapestensis 347 on Clostridium perfringens NCAIM 1417 strain in agar diffusion test [46] [47]. (Photo: Dr. 348 349 Csaba Pintér, University of Pannonia, Keszthely, Hungary)

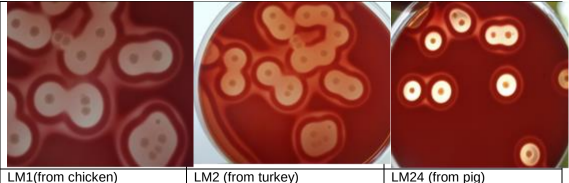
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351 3.4 Results of the Comparison of the Sensitivities (MID values) of 13 Clostridium 352 perfringens strains isolated from Poultry to Cell-Free Culture Media (CFCM) of 353 Xenorhabdus budapestensis (EMA) in Liquid Cultures

354 Table 1 lists the MID values as a qualitative parameter of the sensitivity of each of the 355 poultry isolates to the antibacterial compounds of X. budapestensis. A majority of the 356 examined strains are sensitive but one of the 13 was resistant (LM24). No direct interrelation 357 between the degree of EMA sensitivity and other behavior could be demonstrated. The 358 results provide a good message: The majority of C. perfringens isolates are sensitive. 359 However, they also provide a bad message: There are EMA-resistant resistant C. 360 perfringens isolates, even if they are rare.

361 None of the samples taken from cultures with no visible proliferation contained any CFU, 362 indicating that the toxicity was complete. Whether the differences in the sensitivities could 363 relate to the cellular phenotype was not revealed by this experiment, although the C. perfringens isolates were rather different concerning colony morphology and hemolytic 364 365 behavior (Fig 5).

366



LM1(from chicken)

LM2 (from turkey)

- 367 368
- 369 Figure 5 *Clostridium perfringens* isolates LM1, LM2 and LM24 differing in colony
- 370 morphology, sporulation, and hemolytic behavior. (Photo: Dr. László Makrai, (Department of
- 371 Microbiology and Infectious Diseases, University of Veterinary Science, Hungary).

- 373 Table 1 MID values of *Clostridium perfringens* isolates from chicken differing in colony
- 374 morphology and hemolytic behavior

C. perfringens isolates from	Minimum Inhibiting Dilutions (MID)	Conclusion
poultry (L. Makrai, unpublished)	Values (V/V%) of the cell-free	
	culture medium (CFCM) of	
	Xenorhabdus budapestensis	
	(EMA) Inhibiting Bacterial	
	Proliferation	
LM 1	< 10	Extremely sensitive
LM 2	< 30	Sensitive
LM 3	< 10	Extremely sensitive
LM 4	< 10	Extremely sensitive
LM 5	< 10	Extremely sensitive
LM 8	< 30	Sensitive
LM 11	< 10	Extremely sensitive
LM 14	< 10	Extremely sensitive
LM 15	< 10	Extremely sensitive
LM 16	< 10	Extremely sensitive
LM19	< 10	Extremely sensitive
LM20	< 30	Sensitive
LM 24	> 50	Resistant

375 376

377 4. Discussion

378 The *in vitro* experiments demonstrated that antimicrobial peptides of X. budapestensis 379 (EMA) were highly toxic for all but one (LM 24) C. perfringens isolates. Dr. Klaus Teichmann 380 (Biomin, Tulln, Austria), as a courtesy, tested EMA and EMC CFCM preparations, obtained 381 from us. He declared that the CFCM of EMA exerted an extremely strong anticoccidial 382 activity on both Clostridium and Eimeria cells. He declared that he had not ever worked with 383 such an efficient anticoccidial preparation before as EMA CFCM. Dr. Teichmann found a 384 lower concentration range within which E. tenella cells died, while the cells of the chicken 385 tissue culture were not affected, (Klaus Teichmann, personal communication). These facts 386 are arguments for taking the potential use of EMA and EMC antimicrobial peptides, as 387 potential anticoccidial agents administered per os, into consideration.

388 But there are arguments against using XENOFOOD as well, and they are those data which 389 showed in vitro cytotoxicity on the permanent chicken liver cell line LMH [60]. Dr. Ganas and 390 her associates (Aziza Amin, Irina Profjeva, and Micheal Hess) tested the cytopathogenic 391 effect of different dilutions of the same samples of sterile cell-free media (CFCM) of EMA 392 and EMC on permanent chicken liver LMH cells, as Dr. Teichmann. They demonstrated that 393 EMA CFCM at a dose of < 5% V/V concentration was harmless, but at >5% V/V 394 concentrations they seriously damaged the cell layer. Doses >10% V/V caused total 395 destruction of the cell layer, while that of 5 - 10% V/V resulted in about a 50% damage 396 within the first 24h, and this damage was not repaired in the next 72 hrs. As for EMC, only 397 the dose of 32% resulted in complete cell layer destruction, but the lower doses of 1-20% 398 V/V also resulted in ~ 50% permanent damage, calculated on the base of the score scale of 399 Amin et al. [60] (2012); (Petra Ganes et al., personal communication).

Fabclavines are the predominant antimicrobial compound produced by both EMA and EMC and were isolated and purified [51], and was not suggested as a future drug because of its extremely large target size and toxicity to eukaryotic targets. This kind of "certification" is usually quite enough to place a candidate drug molecule into the wastebasket, despite its super strong antimicrobial effects. However, an exception with fabclavine may be considered because of the following arguments:

First, there are not only prokaryotic, but eukaryotic pathogens also exist. Coccidiosis is the best example where a prokaryotic *C. perfringens* and a eukaryotic *E. tenella* cooperate in causing the disease, and both should be controlled.

Second, there is practically no vaccination technique against *C. perfringens* [28]. So the
introduction of new antimicrobial compounds should be taken into consideration.

We are not the only team walking on this road. Recently, there have been several research
directions attempting to solve the coccidiosis problem. A project includes a search for novel
antibiotic-delivery systems, such as using ovotransferrin as a targeting molecule [61].

Another approach is to improve the usefulness of commonly used anticoccidials and

antibiotics, which have recently been tested on a subclinical necrotic enteritis model [62].

416 Recently proline-rich antimicrobial peptides are considered as potential therapeutics against

417 antibiotic-resistant bacteria [63]. The designer proline-rich antibacterial peptide A3-APO

prevents the Gram-positive *Bacillus anthracis* mortality by deactivating bacterial toxins [64].

- Even more recently two (NZ2114 and MP1102) novel plectasin-derived peptides have been designed for targeting Gram-positive bacteria, and the tests on gas gangrene-associated *C*.
- 421 *perfringens* provided encouraging results [65].

422 The hopes of applying probiotics have been also emerging [66] [67] [68]. The use of

423 vegetative *Bacillus amyloliquefaciens* cells did not justify the hopes: they did not confer

424 protection against necrotic enteritis in broilers, despite the high antibacterial activity of its

425 supernatant against *C. perfringens* in vitro [69].

426 **5. Conclusions**

427 There are two alternative approaches to control coccidiosis in broiler chicken: the

428 vaccination and the "chemotherapy", (that is, a search for gastro-intestinally active,

429 autoclaveable antimicrobial peptides active against both C. perfringens and E. tenella).

Considering that there are publications about antibiotic resistant and multiresistant pathogen *C. perfringens* [70] [71], and that the coccidiosis problem has not yet seem to be solved by
using vaccination, the search for new efficient antimicrobials to control coccidiosis have
probably been justified.

434 On the basis of *in vitro* studies, fabclavine alone (and / or as a component of interacting 435 antimicrobial active peptide complexes present in the CFCM of EMA and EMC) fulfil the 436 criteria of a promising chemotherapeutic agent *in vitro,* that is, acting as strong antibacterial 437 on *C. perfringens* and as strong apoptotic cytotoxic compounds on the unicellular eukaryotic 438 pathogen, *E. tenella*.

However, the cytotoxicity may pose a serious problem of practical use. Indeed, we found
that the CFCM of both EMA and EMC were cytotoxic *in vitro* in permanent chicken liver cells.
But the *in vitro* and the *in vivo* situation are completely different.

443

If it happened that the orally administered fabclavine (and/or the whole AMP complex), due to their proteolytic endurance), might act *in vivo* as strong anti-Clostridia and anti-Eimeria agents in the GI, without causing any harm of the organism to be protected, it would have a chance to be register and use Xenofood as an anticoccidial bio-preparation. This option cannot be ruled out if the adsorption from the gut, were similarly low as that of the orally administred vancomycin [72].

450

We believe that an *in vivo* XENOFOOD feeding experiment would be necessary to learn whether the orally administrated antimicrobial peptides produced by *X. budapestensis* (EMA) and *X. szentirmaii* (EMC), *in vitro* against both the prokaryotic (*C. perfringens*) and the eukaryotic (E. tenella) pathogens causing coccidiosis in chicken, could be used in broiler cockerels.

- 456 We are ready for *in vivo* bioassay and looking for cooperative partners.
- 457

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459

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483 **COMPETING INTERESTS**

484

The authors declare that the research was conducted in the absence of any commercial or
financial relationships that could be construed as a potential conflict of interest.
Authors have declared that no competing interests exist.

488

489 **AUTHORS' CONTRIBUTIONS**

490

491 Dr. Habil. András Fodor was the project initiator and put the MS together. Furthermore, he
 492 provided the bacterium cultures and inoculated and fermented Xenofood with *X*.
 493 budapestensis and *X. szentirmaii*.

494 Professors László Makrai and László Fodor (University of Veterinary Sciences, Budapest,
495 Hungary) were the intellectual leader and governing the *Clostridium* tests. We worked on the
496 *Clostridium perfringens* isolates (LM1 - LM24), previously collected, identified and deposited
497 by them in the Department's stock collection. All but a few experiments were carrying on in
498 their Laboratory at Department of Microbiology and Infectious Diseases, Faculty of
499 Veterinary Science, Szent István University, Budapest, Hungary.

Associate Professor Dr. István Venekei carried out the trypsin digestions and bio-assayed
 the antimicrobial activities of the digested preparations with A. Fodor at the Department of
 Biochemistry at Eötvös University in Budapest, Hungary.

503 **Professor Emeritus Ferenc Husvéth**, (University of Pannonia, Keszthely, Hungary 504 biochemist, independently from the others, guided our experiments on testing the pepsin 505 sensitivity of the antimicrobial active compounds EMA and EMC CFCM in Keszthely, many 506 thanks for it. This work was essential and Dr. Husvéth should definitely be our coauthor.

507 **Associate Professor Dr. László Pál** (University of Pannonia, Keszthely, Hungary) prepared 508 the media by mixing up the food components before A. Fodor inoculated it with X. 509 budapestensis and X. szentirmaii. He weighted the animals daily and calculated the food 510 conversion. His work was essential and Dr. Pál should definitely be one of the coauthors.

511 **Research Associate Dr. Andor Molnár** (University of Pannonia, Keszthely, Hungary) and

A. Fodor has applied previously published overlay,- and agar diffusion technique to C.

- 513 perfringens and he was the pioneer of the *Clostridium* experiments at the Department of
- 514 Animal Sciences and Animal Husbandry, Georgikon Faculty, University of Pannonia,
- 515 (Keszthely, H-8360 Hungary) working with chicken ileal digestions.
- 516 The statistical analyses were carried out or guided by **Adjunct Emeritus Professor Dr.**
- 517 Sándor Józsa (University of Pannonia, Keszthely, Hungary).
- 518 Adjunct Emeritus Professor Dr. Csaba Pintér (University of Pannonia, Keszthely,
- 519 Hungary) made all but one (made by Andrea *Máthé Fodor*, Ohio State University,
- 520 Wooster, OH, USA) photos published here

521 Adjunct Professor Michael G. *Klein* (The Ohio State University) made the final shaping

522 and proofreading of the MS and was the spiritual rector of putting the data and ideas in the 523 present form, which has been an invaluable contribution.

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834		

ABBREVIATIONS

- 836 List of Abbreviations
- **AMP** = Antimicrobial Peptides
- **CFCM** = cell-free culture media
- **EMA** = Xenorhabdus budapestensis, (obligate bacterium symbiont of the nematode
- *Steinernema bicornutum* but can easily be grown in vitro, even in supplemented chicken 841 food)
- **EMC** = Xenorhabdus szentirmaii, (obligate bacterium symbiont of the nematode
- 843 Steinernema rarum but can easily be grown in vitro, even in supplemented chicken food)
- **EPB** = entomopathogenic (nematode-symbiotic) bacterium
- **EPN** = entomopathogenic nematode
- **GI** = gastro-intestinal system
- **MDR** = multi drug resistance (multiple antibiotic resistance)