1	Conjugative IncF and IncI1 plasmids with tet(A) and class 1 integron conferring multidrug
2	resistance in F18 ⁺ porcine enterotoxigenic <i>E. coli</i> .
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28 Abstract

29 Enterotoxigenic E. coli (ETEC) bacteria are frequently causing watery diarrhea in newborn and weaned pigs. Plasmids carrying genes of different enterotoxins and fimbrial adhesins and plasmids 30 conferring antimicrobial resistance are of prime importance in the epidemiology and pathogenesis 31 32 of ETEC. Recently, the significance of the porcine ETEC plasmid pTC was revealed, carrying 33 tetracycline resistance gene tet(B) with enterotoxin genes. In contrast the role of tet(A) plasmids in transferring resistance of porcine ETEC is less understood. Objective of the present study was to 34 35 provide comparative analysis of antimicrobial resistance and virulence gene profiles of porcine post-weaning enterotoxigenic E. coli (ETEC) strains representing pork producing areas in Central-36 Europe and in the USA with special attention on plasmids carrying the *tet*(A) gene. 37

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Antimicrobial resistance phenotype and genotype of 87 porcine ETEC strains isolated from cases of 39 post-weaning diarrhea in Austria, Czech Republic, Hungary and the Midwest USA was determined 40 by disk diffusion and by PCR. Central-European Strains carrying tet(A) or tet(B) were further 41 subjected to molecular characterization of their tet plasmids. Results indicated that >90% of the 42 ETEC strains shared a common multidrug resistant (MDR) pattern of sulfamethoxazole (91%), 43 tetracycline (84%) and streptomycin (80%) resistance. Tetracyclin resistance was most frequently 44 45 determined by the tet(B) gene (38%), while tet(A) was identified in 26% of all isolates with wide ranges for both tet gene types between some countries and with class 1 integrons and resistance 46 47 genes co-transferred by conjugation. The virulence gene profiles included enterotoxin genes (lt, sta 48 and/or stb), as well as adhesin genes (k88/f4, f18). Characterization of two representative tet(A) plasmids of porcine F18⁺ ETEC from Central-Europe revealed, that the IncF plasmid (pES11732) 49 50 of the Czech strain (~120 kb) carried tet(A) in association with catA1 for chloramphenicol resistance. The IncI1 plasmid (pES2172) of the Hungarian strain (~138 kb) carried tet(A) gene and 51 52 a class 1 integron with an unusual variable region of 2,735 bp composed by two gene cassettes: estX-aadA1 encoding for streptothricin-spectinomycin/streptomycin resistance exemplifying 53 54 simultaneous recruitment, assembly and transfer of multidrug resistance genes by tet(A) plasmid of porcine ETEC.. 55

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57 By this we provided the first description of IncF and IncI1 type plasmids of F18⁺ porcine 58 enterotoxigenic *E. coli* responsible for co-transfer of the tet(A) gene with multidrug resistance. 59 Additionally the unusual determinant *estX*, encoding for streptothricin resistane was first reported 60 here in porcine enterotoxigenic *E. coli*. 62 **Keywords**: enterotoxigenic *Escherichia coli*, tetA plasmid, multiresistance, antimicrobial 63 resistance, virulence

64 Introduction

Severe watery diarrhoea of newborn- and weaned pigs as well as of newborn calves are often caused by enterotoxigenic *Escherichia coli* (ETEC) strains (Nagy and Fekete, 2005). On the other hand the ETEC infection often leads to traveler's diarrhoea among humans and may cause significant morbidity and mortality of children in the developing countries (Quadri et al., 2005). Due to the pathogenic similarities of ETEC infections in animals and humans, ETEC has been the subject of intensive studies in human and veterinary medicine over the past three decades.

In weaned pigs two main types of virulence factors are essential for development of the ETECcaused diarrhoea: bacterial attachment to the mucosal surface of the small intestine mediated by fimbrial adhesins (*i.e.* K88/F4 and/or F18ac), and the enterotoxins - heat-labile toxin (LT) and/or heat-stable toxins (STa, STb) - changing the absorptive function of the small intestine to secretion leading to diarrhoea and loss of water and electrolytes.

Genes encoding the above virulence factors are located on ETEC virulence plasmids (Gyles, 1994; Fekete et al., 2012), however several other mobile virulence elements such as pathogenicity islands (PAIs), bacteriophages and transposons may also contribute to transfer of virulence genes between *E. coli* strains (Hacker et al., 1997). As an example, heat-stable enterotoxin genes *sta* and *stb* are part of transposons: *sta* being carried by Tn*1681* (So and McCarthy, 1980), and *stb* was described as a part of Tn*4521* (Hu and Lee, 1988). The spread of the heat labile toxin gene *lt* is realized by IS-mediated transfer mechanisms (Schlör et al., 2000).

In our previous studies on the representative porcine post-weaning F18⁺ ETEC strain 2173 we 83 described and sequenced the large conjugative plasmid pTC (sta^+ , stb^+ , $tetB^+$) as responsible for the 84 enterotoxigenicity and tetracycline resistance [tet(B)] of the host strain (Nagy et al., 1990; Fekete, et 85 al., 2012). Among Enterobacteriaceae the tetracycline resistance is encoded mainly by tetracycline 86 efflux proteins, removing the tetracycline from the bacterial cytoplasm most often ecoded by tetA, 87 tetB, tetC, tetD, tetE and tetG genes.. Tetracycline resistance genes are often located on mobile 88 elements, such as plasmids, transposons and/or conjugative transposons, which can sometimes be 89 transferred between bacterial species (Michalova et al., 2004; Chopra and Roberts, 2001), Studies on 90 the tetracycline resistance mechanisms are justified by data about tetracyclines being the far most 91 92 frequently used antimicrobials in the EU (Schwarz and Chaslus-Dancla, 2001., Moulin, 2008).

In addition to our above studies on pTC plasmid of $F18^+$ ETEC carrying *tet*(B) we intended to 93 and genetic associations of tet(B) and tet(A) with emphasis on 94 study the prevalence 95 characterization and transfer of the less well explored *tet*(A) plasmids of porcine ETEC. Therefore, in this study we aimed to provide a comparative description of antimicrobial resistance and 96 97 virulence profiles of porcine post-weaning ETEC strains representing modern pig industry of different geographical regions: Hungary, Austria, the Czech Republic and the USA. Furthermore, 98 the characterization of genetic vectors for multidrug resistance and tetracycline resistance in this 99 collection was attempted, with special regards to *tet*(A). 100

101 Materials and methods

102 Bacterial strains

Porcine post-weaning enterotoxigenic *E. coli* (ETEC) strains studied here were isolated and identified between 1987-1995 from cases of post weaning diarrhoea, representing 8-12 farms/country in three neighbouring countries: Hungary (n=16), Austria (n=34) and the Czech Republic (n=17). The Central-European collection was complemented with 20 comparable ETEC strains originating from the Midwest-USA. Selected properties of some Hungarian and some US strains have been previously reported (Fekete et al. 2003, and Olasz et al., 2005) All ETEC strains were stored at -80°C in Tryptic soy broth (TSB) complemented with 10% glycerol.

110 Detection of antimicrobial resistance phenotype and identification of tetracycline resistance genes

The antimicrobial resistance phenotype was tested by disc diffusion assay against 18 111 antimicrobial compounds (Oxoid) with clinical relevance. These were as follows (lower indexes 112 113 indicates disc concentration): amoxicillin (AMX₂₅), ampicillin (AMP₁₀), cefotaxime (CTX₃₀), chloramphenicol (CHL₃₀), enrofloxacin (ENR₅), erythromycin (ERY₁₀), florfenicol (FFC₃₀), 114 115 gentamicin (GEN₁₀), kanamycin (KAN₃₀), nalidixic acid (NAL₃₀), rifampicin (RIF₅), spectinomycin $(STR_{10}),$ sulfamethoxazole (SMX₂₅), 116 $(SPE_{100}),$ streptomycin tetracycline $(TET_{30}),$ 117 trimethoprim (TMP₅) and trimethoprim-sulfamethoxazole ($SXT_{1,25/23,75}$). Interpretation of the data was performed according to Clinical and Laboratory Standard Institute (CLSI) guidelines and 118 119 interpretive standards (CLSI, 2013). ETEC strains with intermediate zone diameter values were considered susceptible. The E. coli reference strain ATCC 25922 was used as control. Multidrug 120 121 resistance (MDR) was defined as co-resistance to three or more antimicrobial classes.

ETEC strains of tetracycline resistant phenotype were subjected to PCR-based typing of the *tet* gene, using primers to detect the common *tet* genes of *Enterobacteriaceae*, as listed in Table 1.

124 *Identification and characterisation of ETEC plasmids mediating tetracycline resistance*

In order to characterize plasmids for tetracycline resistance, and to detect the possible cotransfer of tetracycline resistance genes and those encoding typical ETEC virulence factors (*sta, stb, elt, f18, k88/F4*) a total of representetive 8 *tet*(A) and 12 *tet*(B) ETEC strains from Austria, Czech Republic and Hungary were selected for conjugation experiments .

Conjugations were performed by using the plasmid-free, rifampicin resistant E. coli K12 J5-3 129 strain as recipient. Overnight LB broth cultures of the parental ETEC strains and the recipient cells 130 131 were mixed at a ratio of 1:1 and plated onto Luria-Bertani (LB) agar plates. The next day the bacterial lawn was dissolved in 5ml Phosphate buffered saline (PBS), and tenfold dilutions were 132 made up to 10⁻⁷. Selection of the transconjugants was carried out on LB agar plates complemented 133 with the combination of tetracycline (50µg/ml) and rifampicin (150µg/ml). Conjugation frequency 134 was calculated as a ratio between the number (CFU/ml) of transconjugants and that of the recipient 135 strain J5-3. The ETEC strain 2173 served as a reference for the transferability of the tet(B)-136 137 mediating plasmid pTC (Fekete et al., 2012).

138 *Plasmid profile analysis and replicon typing*

139 Parental and transconjugant strains representing successful transfer of the tet(A) and tet(B) genes were subjected to plasmid profile analysis. Plasmids were prepared according to the alkaline 140 lysis method of Kado and Liu (1981). Separation of non-digested plasmids was performed in 0.7% 141 agarose gel in a vertical system with TBE buffer at 180 V. The DNA was stained with 0.5 µg/ml of 142 ethidium bromide. Plasmid sizes were estimated in comparison with plasmid markers (2.1 - 168 kb)143 isolated from E. coli strains V517 and MD112 respectively. Due to our main focus on plasmids 144 carrying the tet(A) gene in F18⁺ ETEC strains, further analyses were done on two tet(A)-positive 145 monoplasmidic transconjugant strains (2172/11 and 11732/11), derived from a Hungarian and a 146 147 Czech isolates respectively. Plasmid incompatibility (Inc) groups were determined by PCR-based replicon typing (PBRT) using primers and conditions developed by Carattoli et al. (2005) and 148 149 García-Fernández et al (2009).

150 *Detection of antimicrobial resistance and virulence genes*

Parental and transconjugant strains were tested by PCR for the presence of antimicrobial resistance genes related to mobile genetic elements and typical ETEC virulence factors, including the flanking regions of the *stb* gene specific to the pTC plasmid of F18⁺ ETEC 2173 (Fekete et al., 2012). Genes *intI1*, *qacE* $\Delta 1$ and *sul1* as parts of the 3' conserved region of the class 1 integron were also tested by PCR. Primers used for the above PCR testing are presented in Table 1 and Table 2. To reveal whether the *tet*(A) gene is located on the transposon Tn*1721*, the primer TetAR3: 5'-GGCATAGGCCTATCGTTTCCA-3' was used (Hartman et al., 2003).

158 *Characterization of class 1 integron carried by the tetA plasmid*

The variable region of the class 1 integron detected in the Hungarian mono-plasmidic *tet*(A) transconjugant strain 2172/11 was amplified with primers 5'CS-F1 and 3'CS-R (Table 1). In order to identify the gene cassette array of the variable region, the PCR product was purified with Qiagen PCR Purification Kit (Qiagen) and submitted to sequencing with primers listed in Table 1. Nucleotide sequences were analysed in comparison with the NCBI database using the BLASTN algorithm. The resulted fragment of 2,645 bp identified in the Hungarian strain 2172/11was deposited in the GenBank under accession number JQ313793.

166 **Results**

167 Antimicrobial resistance phenotype of porcine ETEC strains

168 According to the results of the antimicrobial susceptibility testing, an overwhelming majority of the 87 ETEC strains tested (94.3%) were considered as multidrug resistant (MDR), showing co-169 170 resistance to at least three antimicrobial classes. The distribution of antimicrobial resistance 171 phenotypes among ETEC strains from Hungary, Czech Republic, Austria and the USA is presented in Fig. 1. In general, the prevalence of resistance was lower in ETEC strains from the Central-172 173 European countries, as compared to those from the USA. Majority of the strains shared a common MDR backbone, most frequently being resistant to sulfamethoxazole (91%), tetracycline (84%), 174 erythromycin (84%), and streptomycin (79%). Resistance to spectinomycin (44%), rifampicin 175 (36%) and amoxicillin (32%) were also detected independently of the geographical origin of the 176 177 strains (Fig 1). Concerning resistance to kanamycin and gentamicin, ETEC strains from the Czech Republic were highly susceptible against these drugs, while kanamycin resistance was most 178 179 frequently detected among the strains from the USA and Hungary (90% and 81%). Besides, 180 Hungarian strains showed reduced resistance to gentamicin, chloramphenicol and ampicillin(13181 19%). ETEC strains from Austria showed the highest rate of chloramphenicol resistance (53%), 182 while 50% of the USA strains were resistant to ampicillin (Fig. 1).The mean prevalence of 183 resistance to trimethoprim and nalidixic-acid were relatively low (18% and 15%), while all strains 184 were susceptible to cefotaxime, enrofloxacin and florfenicol (Fig. 1).

185 Distribution of the tet gene types and their combinations among the ETEC strains

ETEC strains with tetracycline resistant phenotype have been tested by PCR for the 186 identification of the tet gene types [tet(A), tet(B), tet(C), tet(D), tet(C), tet(G)] representing the most 187 frequently described efflux pump encoding genes in Enterobacteriaceae. The tet(B) was the most 188 frequently found gene conferring tetracycline resistance in 38%, of ETEC strains, while the *tet*(A) 189 190 gene was identified in 26% (Fig. 2). Only a few strains showed the coexistence of the above two tet gene types. The combination of tet(A)/tet(B), tet(B)/tet(C) and of tet(B)/tet(D) genes averaged 3%, 191 192 1% and 1% respectively. In 31% of the strains the tetracycline resistant phenotype was not 193 confirmed by any of the tet genes tested (Fig. 2) indicating resistance mechanisms encoded by some 194 of the other less frequent tetracyclin resistance genes beside those listed in Table 1.

Distribution of the *tet* gene types varied according to geographical origin.. Hungarian and Austrian strains were characterized by prevalence of 38% and 21% of the *tet*(A) gene respectively and *tet*(B)gene (25% and 32% respectively). In contrast most strains from The Czech Republic carried the *tet*(A) gene (59%). Predominance of the *tet*(B) gene (65%) caharctezied the USA strains (Fig. 2). No ETEC strains tested carried *tet*(E) or *tet*(G) genes.

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201 Plasmid profiles and self-conjugative plasmids of tetracycline resistant ETEC strains

Out of the 8 tet(A) and 12 tet(B) ETEC strains the conjugative transfer of the tet(A) plasmids 202 was successful in 2 of the 8 tet(A) strains selected. One was an F18⁺ ETEC strain from Hungary 203 (2172), and the other one $F18^+$ ETEC strain was from the Czech Republic (11732), resulting in two 204 tet(A)-positive transconjugant strains designated as 2172/11 and 11732/71 respectively. The 205 transfer of tet(B) plasmids resulted in 6 Austrian tet(B) strains out of a total of 12 tet(B) tested. 206 207 from which 8 were of Austrian origin (Table 3). The conjugation frequency of the tet(A) plasmids ranged between 1.06×10^{-5} and 2.37×10^{-5} , while the *tet*(B)plasmids were transferred with a 208 frequency of 2.83×10^{-4} to the transconjugants of the Austrian strain AII.28. Conjugation frequency 209 of the control tet(B) plasmid pTC was 7.76×10^{-4} and the difference between the conjugative 210 transfers of these tet(A) and tet(B) plasmids was obvious but statistically not significant (Fig. 3). 211

All tet(B)-positive ETEC strains carried diverse plasmids ranging from ~10 to 200 kb in size, 212 and their derivative transconjugants showed different plasmid combinations as well.. In general, 213 large plasmids of ~ 120, 145 and 175 kb were the most likely transferable (Table 3). The co-transfer 214 215 of tet(B)-aadA1-catA1 or of tet(B)-catA1 genes and the corresponding phenotypes of tetracyclineaminoglycoside-chloramphenicol resistance was detected by testing the antimicrobial resistance 216 patterns of the tet(B)-positive transconjugant strains originated from Austria (Table 3). All tet(B) 217 strains carried class 1 integrons which were also transferred by conjugation to all but one of the 218 transconjugants. The transfer of certain toxin genes was observed in two transconjugant strains only 219 220 (derivatives of AII.23 and AII.27), sharing the virulence patterns sta-stb and stb-lt-astA respectively, while the adhesin genes *f18* and *k88/f4* were not transferable in the system used (Table 221 222 3). Together with the identification of antimicrobial resistance and virulence genes, parental and transconjugant strains were tested for the 5' flanking region of the stb gene, as a marker for the 223 224 toxin specific locus (TSL) of the tet(B)-mediating plasmid pTC in porcine ETEC. Results indicated the presence of this locus in four of the six tet(B) strains, and in one of the tet(A) strains without 225 226 being transferable in this system (Table 3).

227 Incl1 and IncF plasmid-mediated transfers of tet(A) gene and associated class 1 integron

The Hungarian strain 2172 and the Czech strain 11732 carrying the tet(A) plasmids were also 228 229 multi-plasmidic. Accordingly, plasmid replicon typing identified the coexistence of IncI1, IncF, IncP and $colE_{Tp}$ type plasmids in both of them. However, only the *tet*(A) plasmid was transferred 230 231 from both of these strains. The tet(A) gene was transferred by a large plasmid of IncI1 type (~138) kb) of the Hungarian strain, 2172 (designated as pES2172) and of IncF type (~120 kb) of the Czech 232 strain 11732 (designated as pES11732) respectively (Table 3). The IncI1 plasmid was responsible 233 for the co-transfer of *tet*(A)-*aadA-strA-catA1*(tetracycline-aminoglycoside-chloramphenicol 234 resistance) genes in the Hungarian strain, while the transfer of *tet*(A)-*catA1*was mediated by a large 235 IncF plasmid of the Czech strain (Table 3). PCR analysis identified the *tet*(A) gene as part of the 236 237 Tn1721 transposon in all parental and transconjugant strains. The toxin genes sta, stb and the adhesin gene *f18*, of these two ETEC strains were not transferable (Table 3). 238

239 In the Hungarian F18⁺ ETEC strain, the *aadA1* gene was part of a class 1 integron, located also on the IncI1 tet(A) plasmid (pES2172). The amplification of its variable region resulted in a 2,735 240 bp fragment composed by two gene cassettes. Adjacent to the integrase gene, the estX gene was 241 identified encoding 242 resistance to streptothricin, downstream of which the streptomycin/spectinomycin gene *aadA1* was detected. 243

244 **Discussion**

245 Antimicrobial resistance phenotypes and genotypes of ETEC carrying tet(A) or tet(B) plasmids

Most virulence and antimicrobial resistance genes of enterotoxigenic Escherichia coli (ETEC) are 246 247 located on large plasmids which makes these extra-chromosomal mobile genetic elements the essential tools of evolution through horizontal gene transfer. Plasmids carrying genes of different 248 enterotoxins and/or fimbrial adhesins and plasmids conferring antimicrobial resistance are usually 249 different, but in some cases they appear as hybrid plasmids carrying both resistance and virulence 250 genes. Recently one such hybrid plasmid of porcine post-weaning ETEC (pTC) has been 251 characterized in detail (Fekete et al, 2012). This 90kb self-conjugative plasmid proved to be 252 characteristic to F18⁺ porcine post-weaning ETEC carrying the tetracycline resistance encoding 253 tet(B) gene (Fekete et al., 2003; Olasz et al., 2005). However, the role of tet(A) plasmids in 254 255 transferring resistance and virulence of porcine ETEC seems to be much less understood. Therefore, 256 in these studies we aimed to perform a comparative analysis of antimicrobial resistance and 257 virulence gene profiles of porcine post-weaning enterotoxigenic. E. coli (ETEC) strains representing pork producing areas in Central-Europe and in the USA in order to assess the 258 259 significance of tetracycline resistance and the role of the underlying tet gene types especially tet(A), carried by conjugative plasmids. 260

Antimicrobial resistance phenotype and genotype of a representative collection of 87 porcine 261 262 ETEC strains isolated from cases of post-weaning diarrhoea in Austria, Czech Republic, Hungary and the Midwest USA, indicated the existence of a common resistance backbone of ETEC strains in 263 these two distant geographic regions: with an average multidrug resistance to sulfamethoxazole 264 (91%), tetracycline (84%) and streptomycin (80%). Although the USA strains have shown a 265 generally higher frequency of resistance to the clinically relevant antimicrobials tested, the 266 267 occurence of tetracycline resistance was the highest (100%) among the ETEC strains from the USA, confirming the data of Boerlin et al (2005), from Ontario, Canada in the North-American region.. 268 269 Regarding tetracycline it must be borne in mind that approx. 50-66% of antimicrobial substances 270 used in animal production in the EU was tetracycline (Schwarz and Chaslus-Dancla, 2001, Moulin 271 et al, 2008), while >80% of growing swine in the USA receive tetracylin or tylosin (Landers et al, 272 2012). No wonder that the reported prevalence of resistance genes are high in the EU and in the US. 273 Regarding the two main types of genes encoding tetracycline resistance. Hungarian and Austrian strains were characterized by somewhat similar prevalence of the tet(A) and tet(B) gene (38% vs 274 21%) and (25% vs 32%) respectively, while most strains from The Czech Republic carried the 275

tet(A) gene (59%). In contrast, the predominance of the type tet(B) gene (65%) was found in the 276 USA strains. These data about differences between countries and regions regarding types of tet 277 genes are confirmatory with earlier pu blications of Olasz et al. (2005) and of Boerlin et al.,(2005). 278 279 Their data are supported here by indicating an important role of tetracycline resistance plasmids in the epidemiology of porcine post weaning ETEC. It is acknowledged however that distribution of 280 tet(A) and tet(B) types of porcine ETEC on the same area could change over time (Maynard et al., 281 2003). Therefore, these data should form a comparative background for similar studies on recent 282 postweaning ETEC isolates from these countries. Identification of the genes encoding the three 283 284 main types of tetracylin resistance mechanisms (efflux, ribosomal protection, and enzymatic inactivation) and of the so called unknown types should also be the aim of a future molecular 285 286 analysis for >30 *tet* gene types that could come theoretically into question. (Roberts, 2005).

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An interesting difference between the two main types of *tet* plasmids was observed between the 288 frequency of their conjugative transfers. The transfer of tet(A) plasmids was >1 log₁₀ less frequent 289 290 as compared to the *tet*(B) plasmids. Although this difference was statistically not significant, it 291 could be biologically important enough to be one of the contributing factors - beside plasmid 292 incompatibility - to the wide scale dissemination of the *tet*(B) plasmids in the USA. Furthermore in contrast to tet(A) strains, relatively more tet(B) strains (2/8 vs 6/12) provided transferable plasmids 293 294 and they showed co-transfers with virulence plasmids or with virulence genes on the tet(B) plasmids. In contrast, *tet*(A) strains transferred only their *tet*(A) plasmids without virulence genes. 295 296 However they carried resistance genes against one or more other antibiotics, thereby assisting further selection and spread of multidrug resistance without specific selective pressures. 297

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Beside antimicrobial resistance and virulence genes, parental and transconjugant strains were tested for the 5' flanking region of the *stb* gene, as a marker for the toxin specific locus (TSL) of the *tet*(B)-mediating plasmid pTC. As expected, the results indicated the presence of this locus in majority of *tet*(B) strains, confirming the results of Fekete et al.(2003) and of Olasz et al., (2005) but surprisingly, in one of the *tet*(A) strains (2172) the 5' flanking region was detected as well. However, this indicator gene of TSL was present in a, non-*tet*(A) plasmid (most likely in the 174 kb plasmid) of this strain, which could be the subject of further studies about TSL associations.

308 Although tet(A) and tet(B) are the two most frequently occurring efflux pump encoding resistence 309 genes of porcine ETEC, the role of *tet*(A) plasmids in transferring resistance of porcine F18⁺ ETEC 310 is less understood as that of the tet(B) plasmids (Fekete et al., 2003., Olasz et al., 2005). One of the 311 reasons is that tet(A) seems to be less frequent than tet(B) in porcine ETEC (Boerlin et al., 2005, 312 Maynard et al. 2003). Other reasons could be the differences in the incompatility and conjugative forces of the plasmids involved...The large self conjugative *tet*(A) plasmids of IncI1 type (~138 kb) 313 314 of the Hungarian ETEC (pES2172) and of IncF type (~120kb) of the Czech ETEC (pES11732) seem to deserve attention as representative tet(A) plasmids of F18⁺ porcine ETEC in Central 315 316 Europe. Both carried tet(A) as part of Tn1721 as expected. Besides both self conjugative plasmids carried further resistance genes as well. The plasmid (pES2172) was responsible for the co-transfer 317 318 of tet(A)-aadA-strA-catA1(tetracycline-aminoglycoside-chloramphenicol resistance) genes,, while the plasmid (pES11732) mediated the transfer of *tet*(A)-*catA1*. Literature data about replicon types 319 of self conjugative multidrug resistant tet(A) plasmids of porcine F18⁺ ETEC have been missing... 320 Earlier we have determined replicon types of *f18* plasmids of porcine ETEC (*f18ab*) and VTEC 321 (f18ac) using basic replicon DNA probes and found both kinds of fibmrial plasminds possessing the 322 FIc type of replicons (Fekete et al., 2002). Recently Johnson et al (2011) provided comparative 323 324 genomic analysis of IncI plasmids of porcine ETEC and suggested the existence of a conserved IncI1 plasmid backbone with a single locus for the acquisition of accesory genes associated with 325 326 antimicrobial resistance. However, they did not study IncI or IncF plasmids of ETEC carrying 327 tetracylin resistance genes. In our present study we provided data for the first time about Incl1 and IncF replicon types of multidrug resistance tet(A) plasmids of F18⁺ porcine ETEC. 328

It is of further interest that in the IncI1 tet(A) plasmid (pES2172) the aadA1 gene was part of a 329 class 1 integron, with a 2,735 bp fragment composed by two gene cassettes. Adjacent to the 330 integrase gene, the unusual *estX* gene was identified encoding resistance to streptothricin,. The 331 aminoglycosid antibiotic streptothricin has not been licenced for use in veterinary medicine but it 332 was known as a growth promoter in the former German Democratic Republic (Witte, 1997: Roberts, 333 2005). Genes encoding resistance to streptothrichin acetiltransferase, (sat1 and sat2), as well as the 334 putative esterase (estX) have been described to occur on class 2 integrons of avian pathogenic and 335 commensal E. coli of turkeys in Italy (Piccirillo, 2014) of commensal E. coli from healthy chicks in 336 Korea (Dessie et al, 2013), and of urinary pathogenic E. coli (UPEC) of swine and dog in Germany 337 (Kadlecz and Schwarz, 2008), and in commensal porcine E. coli as well as on class 2 integrons of 338

Aeromonas and *E. coli* from a slaughterhouse wastewater plant without specification of slaughtered animals in Portugal (Moura et al., 2007). The *estX* gene has been reported as part of class 1 integron in two *E. coli* strains isolated from diarrhoeal swine without definition of their pathotype (Cocchi et al., 2007), teherfore its occurence in a porcine ETEC is a novel observation.

343 In summary, our comparative analysis of antimicrobial resistance and virulence gene profiles of porcine post-weaning enterotoxigenic E. coli (ETEC) isolated within less than a decade 344 representing Central-Europe and the USA revealed that almost all ETEC were mutidrug resistant, 345 346 sharing a common pattern of sulfamethoxazole tetracycline and streptomycin resistance. By PCR and sequencing on tetracycline resistance genes and on associated integrons as well as on further 347 antimicrobial resistance and virulence genes we provided the first description of IncF and IncI1 type 348 plasmids of $F18^+$ porcine enterotoxigenic *E. coli* carrying *tet*(A) and multidrug resistance. 349 Additionaly the unusal resistance determinant *estX*, encoding for streptothricin resistance was first 350 described here in a porcine enterotoxigenic E. coli. 351

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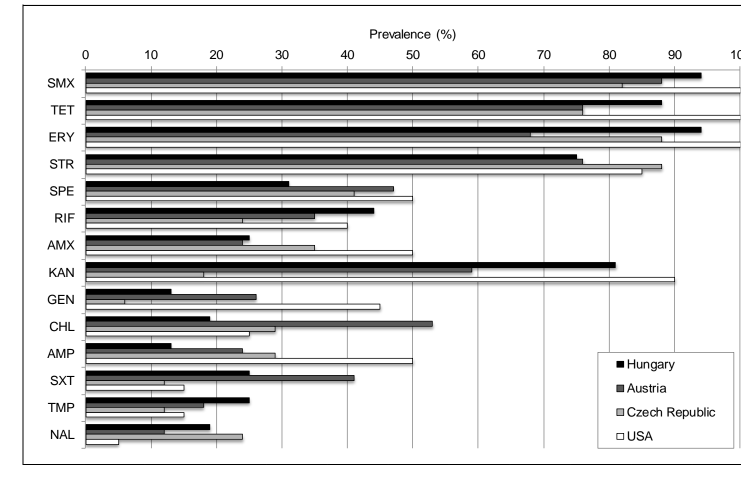
492 **Figure legends**

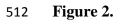
Fig. 1. Distribution (%) of antimicrobial resistance phenotypes among ETEC strains from Hungary,
Austria, the Czech Republic and the USA. Antimicrobial compounds are abbreviated as follows:
SMX, sulfamethoxazole; TET, tetracycline; ERY, erythromycin; STR, streptomycin; SPE,
spectinomycin; RIF, rifampicin; AMX, amoxicillin; KAN, kanamycin; GEN, gentamicin; CHL,
chloramphenicol; AMP, ampicillin; SXT, trimethoprim-sulfamethoxazole; TMP, trimethoprim;
NAL, nalidixic acid.

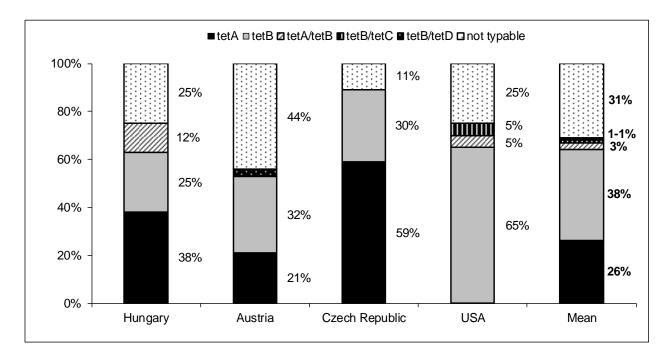
Fig. 2. Distribution (%) of the *tet* gene types and their combinations among ETEC strains from
Hungary, Austria, the Czech Republic and the USA.

Fig. 3. Conjugation frequency of tetracycline resistance plasmids *tet*(A) *andtet*(B) in ETEC strains. Conjugation frequency was calculated as the ratio between the number of transconjugant CFUs (grown on tetracycline and rifampicin) and the number of recipient J5-3 CFUs (which were resistant only to rifampicine). The strain 2173 served as a pTC conjugative transfer control. The CFUs were calculated in two independent experiments with three parallel samples each. Distribution of conjugation frequencies were compared to the 2173 using two-tailed Student's t-test.

Figure 1.









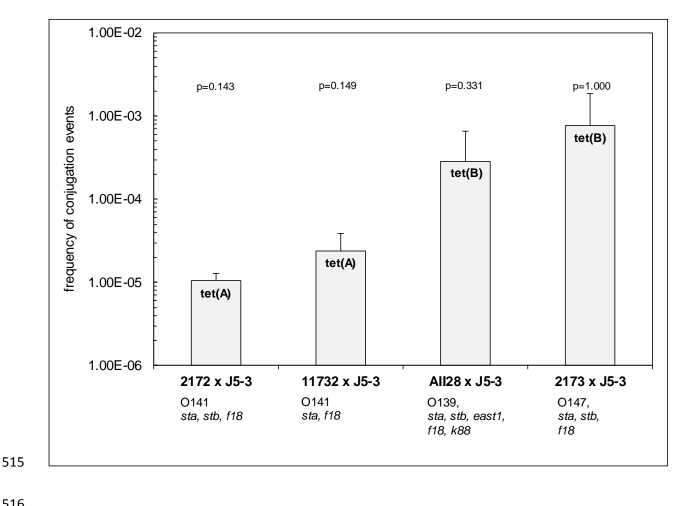


Table 1. Primers used for the detection of antimicrobial resistance genes and for the 520 characterization of class 1 integrons 521

Antimicrobial family and genes	Primer	Sequence $(5' \rightarrow 3')$	Amplicon (bp)	Method	Reference
Tetracyclines					
tet (A)	tetA f	GGCCTCAATTTCCTGACG	372	PCR	Guillaume et al., 2000
	tetA r	AAGCAGGATGTAGCCTGTGC			
tet (B)	tetB f	GAGACGCAATCGAATTCGG	228	PCR	Guillaume et al., 2000
()	tetB r	TTTAGTGGCTATTCTTCCTGCC			
tet(C)	tetC f	TCCTTGCATGCACCATTCC	635	PCR	Guillaume et al., 2000
	tetC r	AACCCGTTCCATGTGCTCG		_	,,
tet (D)	tetD f	GGATATCTCACCGCATCTGC	436	PCR	Guillaume et al., 2000
	tetD r	CATCCATCCGGAAGTGATAGC			
tet(E)	tetE f	TCCATACGCGAGATGATCTCC	442	PCR	Guillaume et al., 2000
	tetE r	CGATTACAGCTGTCAGGTGGG	-1-12	TOR	Califadirie et al., 2000
tet (G)	tetG f	GCTCGGTGGTATCTCTGCTC	468	PCR	Frech and Schwarz, 200
	tetG r	AGCAACAGAATCGGGAACAC	400	TOR	Treen and Ochwarz, 200
Aminoglyappidoo	leigi	AGCAACAGAATCGGGGAACAC			
Aminoglycosides	aacC2 f		600		Francist al. 2001
aacC2		GGCAATAACGGAGGCAATTCGA	698	PCR	Frana et al., 2001
	aacC2 r	CTCGATGGCGACCGAGCTTCA	047	DOD	
aacA4	aac(6')lb f	GTTACTGGCGAATGCATCACA	217	PCR	Frana et al., 2001
	aac(6')Ib r	TGTTTGAACCATGTACACGGC			
aadB	aadB1 fw	GTTGGACTATGGATTCTTAGC	248	PCR	This study
	aadB1 rv	GCCTGTAGGACTCTATGTG			
aadA	aadA fw	GTACGGCTCCGCAGTGGATGG	193	PCR	This study
	aadA rv	GATGATGTCGTCATGCACG		PCR/SQ	
strA	strA fw	CCTGGTGATAACGGCAATTC	546	PCR	Rosengren et al., 2009
	strA rev	CCAATCGCAGATAGAAGGC			
strB	strB fw	ATCGTCAAGGGATTGAAACC	509	PCR	Rosengren et al., 2009
	strB rev	GGATCGTAGAACATATTGGC			
β-lactams					
bla _{CTX-M}	CTX-M f	CGATGTGCAGTACCAGTAA	585	PCR	Batchelor et al., 2003
	CTX-M r	TTAGTGACCAGAATCAGCGG			
bla _{TEM}	TEM f	CATTTTCGTGTCGCCCTTAT	793	PCR	Hopkins et al., 2007
	TEM r	TCCATAGTTGCCTGACTCCC			•
bla _{SHV}	SHV f	ATTTGTCGCTTCTTTACTCGC	1018	PCR	Yagi et al., 2000
	SHV r	TTTATGGCGTTACCTTTGACC		_	3 ,
Phenicols	0				
catA1	catl f	AGTTGCTCAATGTACCTATAACC	680	PCR	Rosengren et al., 2009
00011	catl r	TIGTAATTCATTAAGCATTCTGCC	000	1 OIX	
floR	floR f	CGCCGTCATTCCTCACCTTC	888	PCR	Rosengren et al., 2009
nor	floR r	GATCACGGGCCACGCTGTGTC	000	TOR	Rosengren et al., 2009
am/A	cmlA f	TTGCAACAGTACGTGACAT	293	PCR	Becongrep et al. 2000
cmlA			293	PCR	Rosengren et al., 2009
	cmlA r	ACACAACGTGTACAACCAG			
Class 1 integron-related		000701100170700177700	100	505	
intl1	intl1 f	GGGTCAAGGATCTGGATTTCG	483	PCR	Mazel et al., 2000
	intl1 r	ACATGGGTGTAAATCATCGTC			
qacE∆1	qac F	GGCTGGCTTTTTCTTGTTATCG	273	PCR	Mazel et al., 2000
	qac R	TGAGCCCCATACCTACAAAGC		PCR/SQ	
sul1	sul1 f	TGGTGACGGTGTTCGGCATTC	789		Sáenz et al., 2004
	sul1 r	GCGAGGGTTTCCGAGAAGGTG			
Variable region	5CS-F1	ATGTTACGCAGCAGGGC	variable	PCR/SQ	Libisch et al., 2004
	3CS-R	GGAATTCGACCTGATAGTTTGGCTGTG		PCR	
	sqpr 1 fw	CCTTGCCCTCCCGCACGATG		SQ	This study
	 sqpr 2 rv	CACCACACCGCAGACGACATT		SQ	This study
	sqpr 3 fw	TGGCGAATCAACTCAGGTACTG		SQ	This study
	sqpr 4 fw	CAGAGGTAGTTGGCGTCATC		SQ	This study

522 523 SQ: primers used for sequencing

Gene/Region	Primer	Sequence $(5' \rightarrow 3')$	Amplicon (bp)	Reference	
sta	sta fw	TTTCTGTATTATCTTTCCCC	167	Alexa et al., 1997	
	sta rev	ATTACAACAAAGTTCACAGC			
stb	stb fw	TCTTCTTGCATCTATGTTCG	138	Alexa et al., 1997	
	stb rev	TCTCTAACCCCTAAAAAACC			
stbfl 5'	is1 rev	ACAGCGACTTCCGTCCCAGCC	987	Alexa et al., 1997	
	stb rev	TCTCTAACCCCTAAAAAACC	907		
lt	lt fw	TTACGGCGTTACTATCCTCTCTA	274	Alexa et al., 1997	
	lt rev	GGTCTCGGTCAGATATGTGATTC			
f18	f18 fw	GTGAAAAGACTAGTGTTTATTTC	511	Imberechts et al., 199	
	f18 rev	CTTGTAAGTAACCGCGTAAGC			
k88	k88 fw	GGTGATTTCAATGGTTCGGTC	764	Alexa et al., 1997	
	k88 rev	AATGCTACGTTCAGCGGAGCG			
fedA	f18 fw	GTGAAAAGACTAGTGTTTATTTC	511	Imberechts et al., 199	
	f18 rev	CTTGTAAGTAACCGCGTAAGC			
fanA	fanA fw	AATACTTGTTCAGGGAGAAA	230	Boerlin et al., 2005	
	fanA rev	AACTTTGTGGTTAACTTCCT	230		
fasA	fasA fw	GTAACTCCACCGTTTGTATC	409	Boerlin et al., 2005	
	fasA rev	AAGTTACTGCCAGTCTATGC	409		
east1	astA fw	TCGGATGCCATCAACACAGT	125	Boerlin et al., 2005	
	astA rev	GTCGCGAGTGACGGCTTTGTAAG			
paa	paa fw	GGCCCGCATACAGGCCTTG	282	Boerlin et al., 2005	
	paa rev	TCTGGTCAGGTCGTCAATACTC			
aidA-I	AIDA fw	ACAGTATCATATGGAGCCA	585	Boerlin et al., 2005	
	AIDA rev	TGTGCGCCAGAACTATTA			
sepA	sepA fw	TAAAACCCGCCGCCTGAGTA	611	Boerlin et al., 2005	
	sepA rev	TGCCGGTGAACAGGAGGTTT	611		

524	Table 2. Primers used for the detection of virulence genes and corresponding flanking regions
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Strain	O-type	Country	tet type	Resistance gene patterns	Integron type	Virulence genes	stb 5' flanking*	Plasmid sizes (~kb)
2172	O141	Hungary	tetA	aadA1, strA	intl1	sta, stb, f18	+	174, 138, 38
2172/11 tc			tetA	aadA1, strA	intl1		-	138 (Incl1)
11732	O141	The Czech Republic	tetA	aadA1, strA, catA1		sta, f18	-	138, 106, 60, 5, 4
11732/71 tc			tetA	catA1			-	106 (IncF)
AII.23	O138	Austria	tetB	aadA1, aadB, strA, catA1	intl1	sta, stb, east1, f18, k88	-	174, 145, 120, 106, 60, 47, 15,
AII.23/2 tc			tetB	catA1		sta, stb	-	174, 120
All.23/3 tc			tetB	aadA1, catA1	intl1		-	174, 145, 120
AII.25	O138	Austria	tetB	aadA1, catA1	intl1	sta, stb, f18	+	174, 145, 120, 97, 50, 15
AII.25/1 tc			tetB	catA1	intl1		-	174, 145, 120
AII.27	nt	Austria	tetB	aacC2, aadA1, catA1	intl1, intl2	stb, lt, east1, f18	-	200, 135, 52, 49
AII.27/2 tc			tetB	aadA1, catA1	intl1	stb, lt, east1	-	200, 120
AII.28	O139	Austria	tetB	aadA1, catA1	intl1	sta, stb, east1, f18, k88	+	174, 145, 120, 15
AII.28/2 tc			tetB	aadA1, catA1	intl1		-	174, 145
AII.29	O138	Austria	tetB	aadA1, aadB, catA1	intl1	sta, stb, f18	+	174, 145, 120, 15
AII.29/1 tc			tetB	aadA1, catA1	intl1		-	174, 145, 120
AII.34	O138	Austria	tetB	aacC2, aadA1, catA1	intl1	sta, stb, f18	+	174, 145, 120, 15
All.34/5 tc			tetB	aadA1, catA1	intl1		-	174, 145, 120

Table 3. Plasmid profiles of selected tetracycline resistant parental ETEC strains and of their transconjugants 526

*sstb 5' flanking positive PCR result indicate the presence of toxin specific locus (TSL) characteristic to pTC-like plasmids