Introduction

*Escherichia coli* is a bacterium that normally inhabits the intestines of humans and warm-blooded animals. Most strains of *E. coli* are not harmful for them, but some strains are pathogenic causing gastrointestinal infections (Dhamu et al., 2013; Ray & Bhunia, 2014; Shuahong et al., 2015; Awadallah et al., 2016). Nowadays, more than 700 different serotypes of *E. coli* have been identified and divided into different serotypes according to their "O" (somatic or lipopolysaccharide) and "H" (flagellar) antigens. Pathogenic strains of *E. coli* are divided into six groups (enterotoxigenic, enteropathogenic, enteroinvasive, enterohemorrhagic, enteroc aggregative and diffuse-adhering) based on their ability to produce toxins and to adhere and to invade intestinal epithelial cells (Rani et al., 2017; Vijayan et al., 2017). But the most pathogenic strains of *E. coli* of all known strains are enterohemorrhagic *E. coli* (EHEC). Serogroup O157:H7 of EHEC is the most frequently associated with serious food poisoning, which is accompanied by severe bloody diarrhea, hemorrhagic uremic syndrome (HUS) and can be lethal for humans.

Other serogroups (O26, O45, O103, O111, O121, O145) of EHEC are less common. One of the main characteristics of all of these serogroups is the production of Shiga-toxin (Stx), which is an important virulence factor, responsible for HUS. Because all enterohemorrhagic *E. coli* produce Shiga-toxin, they are also known as Shiga-toxigenic *E. coli* (STEC) or verotoxigenic *E. coli* (VTEC) because of their cytotoxic effect on vero cells (Montzkar & Jamshidi, 2013; Haugum et al., 2014; Ray & Bhunia, 2014; Sudeshan et al., 2014).

There are two types of Shiga-like toxins (Stx1 and Stx2) produced by STEC. Shiga-toxigenic *E. coli* can produce Stx1 only, Stx2 only, or both. The production of these toxins is regulated by bacteriophages which carry the stx genes and which lyosgenize producing strains of *E. coli*. Shiga-toxigenic *E. coli* or verotoxigenic *E. coli* because of their cytotoxic effect on vero cells (Montzkar & Jamshidi, 2013; Haugum et al., 2014, Ray & Bhunia, 2014; Sudeshan et al., 2014).

*Escherichia coli* is part of the normal microflora of the intestinal tract of humans and warm-blooded animals, but its presence in raw material and food of animal origin is considered as fecal contamination and can be very dangerous for consumers. The determination of the number of *E. coli* in raw material and food is important because among them can be pathogenic strains. The most dangerous strains are considered enterohemorrhagic *E. coli* as a causative agent of severe bloody diarrhea and hemorrhagic uremic syndrome in humans through the production of Shiga-toxin, which is the main virulence factor, responsible for disease. The aim of this study was to identify the prevalence of Shiga toxin-producing strains of *E. coli* (STEC) from swabs of beef and swine carcass in slaughterhouses in Ukraine and characterize their genes, which are responsible for pathogenic properties. A total of 230 samples of swabs from beef (130) and swine (100) carcasses were obtained from 5 slaughterhouses in Ukraine between 2012 and 2015. Samples of swabs from carcasses were randomly selected at the final point of the process after the final washing of the carcass from the following areas: distal hind limb, abdomen (lateral and medial) from swine carcasses, brisket, flank and flank groin areas from beef carcasses. All samples were examined by culture-dependent method, after that each positive isolate of STEC was analyzed by multiplex PCR to detect the stx1, stx2, and eae genes. Out of 230 collected samples, seven (7.2%) were contaminated with STEC. The highest prevalence of STEC was found in swabs from beef carcasses (8.1%) in comparison to swabs from swine carcasses (5.7%). The stx1 gene was the predominant gene detected in all STEC positive samples. The eae gene was found in one of the examined isolates from beef carcass. Three isolates from swabs of beef carcass carried both stx1 and stx2 genes, one isolate showed association between stx1 and eae genes, one isolate was positive for stx1 gene only. In swabs from swine carcasses (2 isolates) stx1 and stx2 genes were presented simultaneously. The results of this study suggested that fresh raw meat could be a potential vehicle for transmission of the Shiga toxin-producing strain of *E. coli* to humans. This is the first report of STEC prevalence in beef and swine carcasses in Ukraine and these data will be valuable for microbiological risk assessment and help the appropriate services to develop strategies to mitigate health risk.

**Keywords:** STEC; beef carcasses; swine carcasses; stx1; stx2; eae genes; multiples PCR
In addition to Shiga toxin production, another virulence factor expressed by STEC is intimin. Intimin is a membrane protein produced by all attaching enteric pathogens including STEC as an adherence factor for attachment to the intestinal epithelial cells. The eae gene codes the first toxin (stx1), or the gene that codes the second toxin (stx2), or both genes (stx1 and stx2) at the same time, or other combinations of three virulence genes (stx1, stx2 and eae) (Croxen et al., 2013; Awadallah et al., 2016; Soledad-Cadona et al., 2018).

Humans primarily become infected by Shiga toxin-producing E. coli through consumption of food of animal origin (Ju et al., 2012; Montzau & Jamshidi, 2013; Lozinak et al., 2016). Consumption of undercooked ground beef is the main source of infection as the meat can be easily contaminated with cattle feces during slaughter and butchering (Taye et al., 2013; Abdissa et al., 2017; Premarathne et al., 2017; Vijayan et al., 2017; Omoruyi et al., 2018). Although beef meat is considered as the main source of STEC for people, alimentary infections caused by STEC have also been described after consumption of swine meat (Troz-Williams et al., 2012; Tseng et al., 2014).

The disease caused by the toxin usually has very serious consequences for human health and can manifest by three different syndromes: hemorrhagic colitis (HC), hemolytic uremic syndrome (HUS), and thrombotic thrombocytopenic purpura and which can even cause death (Majowicz et al., 2014; Smith et al., 2014; ECDC, 2016). Due to the danger of STEC at the international level and to ensure food safety, it is officially recognized as a requirement to carry out research on the testing of these microorganisms in meat and meat products (Tafida et al., 2014; ECDC, 2016). It is possible to determine STEC by bacteriological methods that include the preenrichment of studied samples in broth with antibiotics, then isolation on agar typical colonies and then serotyping. To control STEC in each country, culture-dependent methods are different, but PCR – methods are the most rapid, sensitive and highly specific. Therefore, development and modifying a general technique of PCR – methods for detection of Shiga toxin-producing strain of E. coli are the focus of attention of scientists in different countries. Often, for rapid detection of the presence of several virulence factors of STEC in the samples, polymerase chain reaction (PCR) in multiplex version is used (Puttalingamma et al., 2012; Rantsiou et al., 2012; Haugum et al., 2014; Hara-Kudo et al., 2016).

It should be noted that at present in Ukraine rapid control techniques for STEC, including PCR, have not been developed yet, which is a deterrent to ensuring access of Ukrainian goods to foreign markets, especially the EU. In this regard, it is important to modify or adapt a general specific and rapid method of detection and modifying a general technique of PCR – methods for STEC, including PCR, have not been developed yet, which is a deterrent to ensuring access of Ukrainian goods to foreign markets, especially the EU. It is possible to determine STEC by bacteriological methods that include the preenrichment of studied samples in broth with antibiotics, then isolation on agar typical colonies and then serotyping. To control STEC in each country, culture-dependent methods are different, but PCR – methods are the most rapid, sensitive and highly specific. Therefore, development and modifying a general technique of PCR – methods for detection of Shiga toxin-producing strain of E. coli are the focus of attention of scientists in different countries. Often, for rapid detection of the presence of several virulence factors of STEC in the samples, polymerase chain reaction (PCR) in multiplex version is used (Puttalingamma et al., 2012; Rantsiou et al., 2012; Haugum et al., 2014; Hara-Kudo et al., 2016).

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Methods of isolation and identification of E. coli from samples. Samples were examined for 2–12 hours. Swab samples were examined by standard methods: serial dilutions of samples were plated on the surface of commercial medium “Compact Dry™ EC” for isolation of E. coli in Petri dishes (NISSUI Pharma). It is a ready-to-use chromogenic medium for performing E. coli and coliform counts. Petri dishes with isolates were incubated for 24 h at 37 °C. Interpretation of the results was performed by the following indicators: blue colonies were considered as E. coli.

**Oligonucleotide primers.** Alignment of nucleotide sequences and their homology analysis was performed by the module Clustalx software "Vector NTI" v.10.0.1 (Invitrogen) and Blast-analysis resource www.ncbi.nlm.nih.gov (National Center for Biotechnology Information, USA). Primers for PCR multiplex variant were calculated by the software "Vector NTI" v.10.0.1 and synthesized in the "Lytech" (Russia). Lympholized primers were diluted to a concentration of 100 pmol/µl "Ultra Pure Distilled Water" (Invitrogen, Cat. # 10977-023, USA) and stored at −20 °C until use. Specific oligonucleotide primers that were used for the detection of STEC are shown in Figure 1.

![Fig. 1. Specific oligonucleotide primers specific to the toxin gene stx1 (a), stx2 (b) and intimin (eae) (c) that were used for the detection of Escherichia coli (STEC)](image-url)

DNA isolation. DNA isolation was performed by three different methods: 1st – a colony of E. coli cultures was taken in a test tube with 0.5 microliters of sterile deionized water and was heated for 3 minutes at 100 °C, after that the tube was immediately transferred to ice; 2nd – selection using a commercial kit "DNA Sorb-B" (Amphimsens, Russia); 3rd – using silica-modified (~ 15 nm) magnetic particles (synthesized and provided by N. Volkova, Institute of Physics and Biophysics NAS) with a concentration of 10 mg ml⁻¹ and saturation magnetization of 37 emu (Am²/kg), purification of bacterial DNA for spectrophotometric analysis was performed using a set of "Ultra Clean DNA Purification Kit" (Cat. # 12500-100; Mobio, USA). The concentration and purity of drugs obtained DNA was measured on a spectrophotometer "NanoDrop 2000c" (USA).
Polymerase chain reaction (PCR). PCR was performed in thermocycles “Tertsyk” (DNA technology, Russia) and “T1” (Biometra, Germany). The reaction was carried out by “hot” start in a volume of 0.025 cm³. In order to minimize the formation of nonspecific dimers primer matrix and its amplification the method of preparation of the reaction mixture with the physical separation of PCR components was used.

To prepare the “lower” reaction mixture, nucleotidyrophosphatase (2 mM) was mixed with appropriate primers in one tube at the rate of 0.025 cm³ each (final concentration from each primer 10–15 pmol/ sample). After mixing in a vortex, the mixture was dropped in prepared for PCR microtubes in volume 0.005 cm³ in each and on the top of it molten wax in volume 0.015 cm³ was added. After solidification of wax in the tube, the “upper” reaction mixture in volume 0.017 cm³ and 2 drops of mineral oil were added. The “upper” reaction mixture (1 sample calculation) consisted of 0.005 cm³ (x5) PCR buffer; 0.0025 cm³ 50 mM MgSO4; 0.009 cm³ 1 H2O MilliQ and 0.0005 cm³ Taq-polymerase (5 units/ml). Samples of bacterial DNA were placed under oil in the volume 0.003 cm³.

Thermal cycling parameters were as follows: 95 °C for 3 min, followed by 35 cycles of 94 °C for 30 s, 56 °C for 30 s and 72 °C for 30 s; and a terminal extension step of 72 °C for 4 min. Negative control was a non-pathogenic E. coli strain.

Electrophoretic analysis of PCR products. Analysis of amplification products was performed by separation of DNA fragments in a 1.5% (w/v) agarose gel with containing 0.5 µg/cm³ ethidium bromination products was performed by separation of DNA fragments in a microorganisms under oil in the volume 0.003 cm³.

Key isolation of Shiga toxin-producing E. coli by PCR. Multiplex PCR was used for detection of three target genes stx1, stx2 and eae in 97 E. coli isolates. Only 7 E. coli isolates (7.2%) had at least one of 3 genes. Most of the isolates (685.7%, 571.4%) carried the stx1 or/stx2 genes and only one isolate had the eae gene. Fragments of positive result of detection of PCR products of fragment gene of stx2 and gene eae are shown on Figure 2.

Results

Modifying of PCR for identification of stx1, stx2 and intimin (eae) genes in isolated STEC. For the modifying of PCR for detection and identification of Shiga toxin-producing E. coli in beef and swine carcasses, first we performed research on the products received in PCR with oligonucleotide primers specific to the toxin gene stx1, stx2 and intimin (eae), which are pathogenicity markers for STEC. To design specific oligonucleotide primers, gene sequences in the databases GenBank, EMBL (European Molecular Biology Library), DDBJ (Japanese database nucleotide sequences) and PDB sequences were analyzed. As a basis, the following sequence had been selected: AB647443, AF022236, BA000007, ECOSTLII, AB647559, AB647430, AB647374, AB334567, AJ308552.1, AB647553.1, DQ523611.1, AB647374.1, H19BLSTA, KF771380.1, AB647493.1, AB647432.1, AB647437.1, AB647449.1, DQ523603.1, AB647365.1, EU700490.1, EF441589.1, EF441588.1.

As a basis, we have chosen a pair of oligonucleotide primers for the detection of E. coli O157:H7, which had been developed by Puttagangamma et al. (2012). We found that aforementioned primers contain palindromes and loops, and they influence the formation of numerous homo- and heterodimers that generally affect the specificity of the reaction. Simultaneous use of these primers in PCR multiplex option is not effective. The correct choice of oligonucleotide primers is very important because it determines the effectiveness and reproducibility of PCR.

The first task in the development of PCR was to improve the current method of detecting DNA of Shiga toxin-producing E. coli (STEC) by changing the composition of nucleotide sequences in primers for amplification of specific nucleic acid fragments. The developed primers could be used also in multiplex variant for detection of specific DNA fragments (stx1, stx2 and eae) of Shiga toxin-producing E. coli (STEC) in the samples.

Primer specificity was confirmed in test strains of heterologous microorganisms Salmonella enterica, Listeria monocytogenes, Bacillus anthracis, Campylobacter jejuni, Pasterella multocida and Yersinia enterocolitica. To evaluate specificity and sensitivity of the reaction, optimal annealing temperature of the primers was determined (Fig. 2). To determine the sensitivity of the developed primers, 10-fold serial dilutions of purified bacterial DNA were prepared. Concentration of purified DNA was determined with a spectrophotometer. Sensitivity for eae gene of O145 strain was 0.220 and 0.021 ng for genes stx1 and stx2 of O157 strain.

Results of isolation and identification of E. coli from samples. Among the 230 swabs, a total of 97 E. coli isolates (42.2%) with typical cultural properties (blue colonies) was isolated by using commercial medium “Compact Dry” for isolation of E. coli in Petri dishes. The study was conducted on 62 isolates from the surfaces of beef and 35 isolates from swine carcasses (Table 1). More positive samples were detected from surfaces of beef (47.7%) than from swine carcasses (35.0%).

<table>
<thead>
<tr>
<th>Sample type</th>
<th>No. investigated samples</th>
<th>No. positive samples</th>
<th>Percentages, %</th>
</tr>
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<tbody>
<tr>
<td>Swabs from beef</td>
<td>130</td>
<td>62</td>
<td>47.7</td>
</tr>
<tr>
<td>Swabs from swine</td>
<td>100</td>
<td>35</td>
<td>35.0</td>
</tr>
<tr>
<td>Total</td>
<td>230</td>
<td>97</td>
<td>42.2</td>
</tr>
</tbody>
</table>

These 97 isolates were further subjected to polymerase chain reaction (PCR). Multiplex PCR was used for detection of three target genes stx1, stx2 and eae in 97 E. coli isolates. Only 7 E. coli isolates (7.2%) had at least one of 3 genes. Most of the isolates (685.7%, 571.4%) carried the stx1 or/stx2 genes and only one isolate had the eae gene. Fragments of positive result of detection of PCR products of fragment gene of stx2 and gene eae are shown on Figure 2.
but the detection of STEC from different products of animal origin is an achievable goal. STEC can be a source of STEC for humans through contaminated raw meat products. Therefore, within the framework of trade agreements between Ukraine and European countries, food and raw food of animal origin must be tested for safety before being shipped for export. One of the important criteria of current microbiological safety of raw meat and meat products (especially beef and pork) is the control of Shiga-toxin-producing _E. coli_ (STEC), because they are food-borne pathogens that are very serious threats to public health. STEC infection is associated with sporadic outbreaks of clinical diseases in humans, including severe hemorrhagic colitis, hemorrhagic uremic syndrome, thrombotic thrombocytopenic purpura, which can even cause death (Majowicz et al., 2014; Smith et al., 2017; Bhunia, 2014; Bonardi et al., 2015; ECDC, 2016).

The combinations of virulence genes of STEC (stx1, stx2, eae) in isolated strains (n = 7) are shown in Table 3. Three combinations of virulence genes were identified in isolated strains (n = 7): stx1 + stx2 + eae, stx1 + stx2, and stx1 + eae. The total number of positive samples with stx1, stx2, and eae genes was 5, 1, and 0, respectively. The highest percentage of positive samples was found in stx1 + stx2 combinations (36.4%). The second most common combination was stx1 + eae (14.3%). The presence of the eae gene was detected in 100% of positive samples. However, the combination of all three genes (stx1, stx2, and eae) was not found in any of the samples.

Discussion

According to international epidemiological data, in all countries of the world, cases of foodborne illness among people are increasing. Therefore, within the framework of trade agreements between Ukraine and European countries, food and raw food of animal origin must be tested for safety before being shipped for export. One of the important criteria of current microbiological safety of raw meat and meat products (especially beef and pork) is the control of Shiga-toxin-producing _E. coli_ (STEC), because they are food-borne pathogens that are very serious threats to public health. STEC infection is associated with sporadic outbreaks of clinical diseases in humans, including severe hemorrhagic colitis, hemorrhagic uremic syndrome, thrombotic thrombocytopenic purpura, which can even cause death (Majowicz et al., 2014; Smith et al., 2017; Bhunia, 2014; Bonardi et al., 2015; ECDC, 2016).

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findings (Troz-Williams et al., 2012; Colello et al., 2016). But prevalence of STEC isolates carrying virulence genes stx1, stx2, intimin (eae) has been reported in lactating cows and in contact workers in dairy farms at Sharkia Province, Egypt (Merwad et al., 2014). Three isolates from swabs of beef carcasses carried both stx1 and stx2 genes, one isolate showed association between stx1 and eae genes, one isolate had positive result for stx1 genes only (Table 3). In contrast to this study, some results have shown that E. coli O157:H7 was absent in raw minced beef samples in Tripoli, Lebanon by using real-time PCR-based method (Omari et al., 2018).

Although a small number of research articles which implicate pork as a source of human infection have been reported, the results of these investigations consider that meat other than beef also can be potential vehicles of STEC transmission (Troz-Williams et al., 2012). It is important to note that in swabs from swine carcasses (2 isolates stx1 and stx2 genes were presented). So, the results of our study indicate that swine can be a potential reservoir of STEC strains. The same researchers isolated from 465 not ready to eat pork samples 65 (14.0%) stx-positive E. coli: the stx2 gene was detected more frequently (13.3%) than the stx1 gene (1.3%) and associations of genes in pork samples were next: stx1+eae (0.4%), stx2+eae (8.0%) and stx1+stx2+eae (0.7%) (Bardasi et al., 2017). At slaughter houses in Argentina 4.1% of carcasses of swine were stx positive: 50% of isolates positive for stx2 and 16.0% for stx1/stx2. (Colello et al., 2016).

Conclusion

This study is the first report on the presence of Shiga toxin-producing strains of E. coli (STEC) in beef and swine carcasses in Ukraine. The results indicate that fresh raw meat (beef and pork) could be potential vehicles for transmission of enterohaemorrhagic E. coli infections to humans since the highest prevalence these pathogenic microorganisms was found in swabs from carcasses and swine carcasses. These data will be valuable for microbiological risk assessment and helps authorities to develop strategies to mitigate health risk.

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References


