Survey of Minas frescal cheese from Southwest Minas Gerais for virulence factors and antimicrobial resistance in *Escherichia coli* isolates

Avaliação em queijos Minas frescal provenientes do Sudoeste de Minas Gerais da presença de fatores de virulência e resistência antimicrobiana em isolados de *Escherichia coli*

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**ABSTRACT**

The soft cheese Minas frescal is one of the most popular cheese in Brazil, which is typically manufactured in small dairy farms under unsatisfactory hygiene conditions. To assess the risk involved in consumption of this cheese, virulence markers were investigated in 330 *Escherichia coli* strains isolated from 30 Minas frescal cheeses inspected by official government agency (SIF - serviço de inspeção federal), from 50 cheeses not inspected by SIF and 31 cheeses not inspected by SIF with spice added, all of them collected in the southwest of Minas Gerais State. The *E. coli* isolates were screened for the presence of Shiga toxin-encoding (stx 1 and stx 2), intimin (eae) genes and for the presence of (pap, sfa, afa) genes related to adhesion in epithelial cells. The only gene detected by PCR was the sfa gene at one isolate. The strains were also screened for resistance to 9 antimicrobial drugs. Predominant resistance was to cephalothin, tetracycline and streptomycin. Multidrug resistance was found among isolates from cheese with SIF (16.6%), cheese without SIF (8.0%) and cheese without SIF with spice added (30.0%) what is a reason for concern due to the high consumption of raw milk cheese by the Brazilian population.

**Key words:** *Escherichia coli*, cheese, STEC, ExPEC, antimicrobial resistance, Minas frescal.

**INTRODUCTION**

Minas frescal cheese is one of the most popular cheeses produced in Brazil, this soft white cheese is made of pasteurized or raw milk. It is characterized by a high pH and moisture content (>55%) and low percentage of salt (1.4-1.6%) (MORAES et al., 2009). *Escherichia coli* is used as an indicator of direct or indirect fecal contamination of foods, and therefore the possible presence of enteric pathogens. In cheese, *E. coli* is used as an indicator to assess the contamination and its presence may indicate poor hygiene conditions during processing.
or post-processing contamination (O'BRIEN et al., 2009), in Brazil there are specific rules for the production and commercialization of dairy products (BRASIL, 1996, 2001), however the production and retail sale of no inspected dairy products is common in Brazil (MORAES et al., 2009).

Since the occurrence of an outbreak of foodborne disease caused by enteropathogenic E. coli associated with consumption of soft ripened cheese (MARIER et al., 1973), the presence of these microorganisms in cheese acquired additional significance. Verocytotoxin (VT)-producing E. coli (VTEC) also known as Shiga toxin-producing E. coli (STEC) are among the most important causes of foodborne diseases. They are responsible for overall human gastrointestinal diseases, including watery or blood diarrhea (PATON & PATON, 1998). The STEC strains can gain access to milk by fecal contamination or by direct excretion (mastitis) from the udder into the milk (LIRA et al., 2004). Outbreaks and sporadic cases of illnesses have been traced to consumption of STEC-contaminated cheese (BAYLIS, 2009).

Some pathotypes of E. coli are capable of causing intestinal disease, while others referred to as extraintestinal pathogenic E. coli (ExPEC), are responsible for extraintestinal infections. Usually, ExPEC isolates have specialized virulence factors enabling them to colonize host surfaces, injure host tissues, and avoid or subvert host defense systems (JOHNSON et al., 2005). Among them the P fimbriae (encoded by the pap gene), Sfa fimbriae (encoded by sfa genes) and Afa afimbral (encoded by afa gene) are important virulence factors related to adhesion and usually found among E. coli strains causing urinary tract infection (UTI) in humans. Some studies have suggested that food may give raise to human acquired ExPEC strains (JOHNSON et al., 2005; VINCENT et al., 2010).

During the last decade there has been an increasing awareness of potential problems to human health caused by the selection of antimicrobial resistance among food producing animals (AARESTRUP, 2005). Since pathogenic or commensal bacterial species from animals can be transmitted to humans through milk or milk products, it seems prudent to explore not only resistance patterns to antimicrobials used in veterinary medicine, but also to antimicrobials used in human medicine. In Brazil only a limited number of studies concerning the antibiotic resistance of commensal E. coli isolated from cheese have been performed (PANETO et al., 2007).

Cattle are an important reservoir of toxigenic E. coli, and have been implicated as a source of E. coli that infect and cause disease in human beings (BAYLIS, 2009). The aims of the present study were to determine the presence of STEC and ExPEC strains in Minas frescal cheese and also to verify the resistance of E. coli strains isolated from cheese to antimicrobial drugs.

MATERIALS AND METHODS

Sample collection, microbiological analysis

A hundred-eleven samples of Minas frescal cheese were randomly purchased in the Triangulo Mineiro region in the southwest of Minas Gerais State from supermarkets or small food stores at different cities. Thirty cheese samples have been inspected by official government agencies (SIF-servico de inspecção federal), fifty were not inspected (without SIF) and thirty-one were not inspected and have spice added (Origanum vulgare and/or Pretoselium sativum). All samples were kept under refrigeration in plastic bags; information about dates of production and of assigned shelf lives sometimes were not presented. Cheese samples were collected over a period of 10 months between March and December 2007, and were analyzed on the day of acquisition. Samples were transported under refrigeration (4-6°C) in thermal boxes containing ice packs and were tested immediately after arrive at the laboratory. A 25g portion of each cheese was blended with 225ml of nutrient broth (Difco, Detroit, USA) for two min at normal speed using a Stomacher lab blender and incubated at 37°C for 24h. An 1ml sample of the nutrient broth culture was mixed with 9ml of MacConkey broth (Oxoid, Hampshire, UK) and further incubated at 37°C for 24h. One loop of each tube was streaked on MacConkey agar. At least ten colonies from each plate with typical E. coli morphology were selected and examined by biochemical tests, including hydrogen sulphide, citrate, urease and indole (KONEMAN et al., 1997). The technique used to determine the most probalbe number (MPN) of coliforms was developed according to the AMERICAN HEALTH ASSOCIATION (2001) and U.S. FOOD AND DRUG ADMINISTRATION (2001). Microbiological standards for soft cheese are determined by the official Brazilian parameters of the Ministries of Agriculture and Health (BRASIL 1996, 2001).

Polymerase chain reaction

Bacterial strains (E. coli isolates) grown overnight in nutrient broth (Sigma Chemical Co, St Louis, USA) at 37°C were tested for the presence of stx (stx 1 and stx 2) and eae genes using the polymerase chain reaction (PCR) protocol of CHINA et al. (1996)
and the protocol of LE BOUGUENEC et al. (1992) for the presence of pap, sfa, afa genes. DNA templates were prepared by boiling in sterile distilled water. The DNA was subjected to PCR performed in an Eppendorf Mastercycler (Eppendorf AG, Hamburg, Germany). Stx1, Stx2, eae, pap, sfa and afa genes were detected using primers and PCR conditions in the above-mentioned protocols. The amplified DNA products were separated by electrophoresis on 1.5% agarose gel, stained with ethidium bromide and detected under ultraviolet light. Reference E. coli strains used as controls were EDL 933 (O157:H7, stx1, stx2, eae), FV L2 (sfa, pap, iuc, hly, cnf-1) and DH5α (negative control).

Susceptibility testing

Antimicrobial susceptibility tests were performed using the disk diffusion method recommended by the Clinical and Laboratory Standards Institute (CLSI, 2008). Drug-impregnated disks (CEFAR, São Paulo, BR) were placed on agar surfaces using a disk dispenser. The following nine antimicrobial agents were tested: tetracycline (TET, 30 μg); sulfamethoxazole/trimethoprim (SUT, 25 μg); amikacin (AMK, 30 μg); cephalothin (CFL, 30 μg); ceftriaxone (CEF, 30 μg); gentamicin (GEN, 5 μg); streptomycin (STR, 10 μg); nalidixic acid (NAL, 30 μg); ciprofloxacin (CIP, 5 μg).

RESULTS AND DISCUSSION

A total of one thousand-two hundred-and-forty-three E. coli isolates were obtained from cheese samples, 216 isolates from cheese SIF + (with SIF inspection), 30 cheeses) and 449 isolates SIF - / spice (without SIF inspection). The following nine antimicrobial agents were tested: tetracycline (TET, 30 μg); sulfa- methoxazole/trimethoprim (SUT, 25 μg); amikacin (AMK, 30 μg); cephalothin (CFL, 30 μg); ceftriaxone (CEF, 30 μg); gentamicin (GEN, 5 μg); streptomycin (STR, 10 μg); nalidixic acid (NAL, 30 μg); ciprofloxacin (CIP, 5 μg).

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<tr>
<th>Table 1 - Distribution of Most Probable Number (MPN) of fecal coliforms at 45°C among the Minas frescal cheese from Southwest Minas Gerais.</th>
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<tr>
<td>MPN g⁻¹ of fecal coliforms</td>
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<tr>
<td>Samples (%)</td>
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<tr>
<td>3.0 - 5.0 x 10¹</td>
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<td>5.0 x 10¹ - 5.0 x 10²</td>
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<td>5.0 x 10² - 5.0 x 10³</td>
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* Cheese with SIF inspection; ** cheese without SIF inspection; ***cheese without SIF inspection with spice added; **** number of samples cheese. Anvisa permitted limits =5.0 x 10² MPN g⁻¹ for Minas frescal cheese; 10⁻² for Minas frescal cheese with spices.
(3.0%) gene and two positive strains for stx 2 (6.0%) among the examined samples. ZWEIFEL et al. (2010) in a survey of a 3 year monitoring program in Swiss examined 1,502 cheeses, among them 80 samples of soft cheese and reported the detection of 8 soft cheese (10%) contaminated by non-O157 STEC strains, all of them carrying the stx 2 gene. GONZALES et al. (2000) examined 44 soft white cheeses (Minas frescal) in Rio de Janeiro, Brazil and isolated 386 E. coli strains none of them with stx genes detected. In Brazil, at the best of our knowledge, only PANETO et al. (2007) reported the analysis of 50 samples of Minas frescal cheese with the recovered of 3 E. coli isolates carrying the vt 2 (stx 2) gene (2 isolates) and vt 1 (stx 1) gene (1 isolate).

In the present work also the adhesions encoding genes pap, sfa and afa were almost absent. Only one isolate (number 301) from cheese SIF- presented the sfa gene (0.6%) but it can not be characterized as an ExPEC strain, what means there is no ExPEC strains among the cheese isolates tested. ALTALHI & HASSAN (2009) in Saudi Arabia did not found pap and sfa genes among the examined raw milk samples but they reported a great percentage of other genes such as trat T (serum resistance, 51.5%), iut A (aerobactin, 33.3%) and fyu A (yersiniabactin, 24.2%) among the E. coli isolates, all of them considered virulence-associated traits related with ExPEC.

The emergence of antimicrobial resistance among pathogens that impact animal health has been a growing concern in veterinary medicine (VAN DEN BOGAARD & STOBBERING, 2000). Some E. coli strains which contaminate dairy products represent a direct hazard to human health due to their virulence factors (BAYLIS, 2009), others represent a possible source of genes of transmissible resistance to antimicrobial drugs used for the treatment and prophylaxis of infections diseases in animals (VAN DEN BOGAARD & STOBBERING, 2000; AARESTRUP, 2005).

The susceptibility to 9 antimicrobial agents for the three hundred-thirty E. coli isolates is shown in table 3. Among the isolates from cheese the highest resistance was observed against cephalothin, streptomycin and tetracycline (Table 3). PANETO et al. (2007) examined 48 E. coli strains isolated from Minas frescal cheese and they reported resistance to cephalothin (60.0%), nalidixic acid (40.0%), tetracycline (31.0%), ampicilin (29.0%) among the isolates examined. However it is necessary to mention that the author above mentioned, combine intermediate sensibility value with the resistance value to create the resistance percentage. Looking in that way the percentage of resistance reported in the present study (Table 3) were quite similar to those above mentioned, but higher than the resistance reported by CAMPOS et al. (2006) for E. coli strains isolated from 24 samples of Minas frescal cheese from Goias, Brazil (4.0% for ampicilin and tetracycline).

UNLU et al. (2011) examined 120 white cheese samples in Turkey and 56 of those (46.6%) were found to be contaminated with E. coli. The authors reported a high level of antimicrobial resistance among the E. coli isolates to gentamycin (60.0%), tetracycline (52.0%) and enrofloxacine (30.0%), all of them higher than the percentages of antimicrobial resistance reported in the present study. E. coli multidrug resistant (MDR) was found in the three types of Minas frescal cheese examined in the present work. The most commonly found MDR were to cephalothin + tetracycline + streptomycin and sulfamethoxazole/trimethoprim + tetracycline + streptomycin. The results were 16.6% for cheese with SIF inspection, 8.0% for cheese without SIF inspection and 30.0% for cheese without SIF inspection with spice added. AMADOR et al. (2009) examined 20 cheese samples in Portugal and isolated 172 bacterial samples, 31.4% of those displayed a MDR phenotype, a value higher than the
results described in the present study. Resistance of \textit{E. coli} MDR isolated from foods to antimicrobial drugs used in human medicine has been a reason for concern due to the risk of dissemination of these genes among the human microflora.

To conclude, the present study did not find STEC or ExPEC strains among 330 \textit{E. coli} isolates from Minas frescal cheese. \textit{E. coli} commensal strains isolated from cheeses showed a high level of antimicrobial resistance and multidrug resistance, what it is a reason for concern due to the high consumption of raw milk cheese by the Brazilian population.

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