Molecular characterization of Yersinia enterocolitica strains to evaluate virulence associated genes

Eleonora Ventola¹, Sarah Lovari², Silvana Farneti³, Guido Finazzi⁴, Stefano Bilei², Slawomir Owczarek⁵ and Elisabetta Delibato¹

¹Dipartimento di Sicurezza Alimentare, Nutrizione e Sanità Pubblica Veterinaria, Istituto Superiore di Sanità, Rome, Italy ²Direzione Operativa Microbiologia degli Alimenti, Istituto Zooprofilattico Sperimentale Lazio e Toscana "M. Aleandri", Rome, Italy °Controllo Alimenti Umbria, Istituto Zooprofilattico Sperimentale dell'Umbria e delle Marche "Togo Rosati", Perugia, Italy

Early release] ⁴Dipartimento Sicurezza Alimentare, Istituto Zooprofilattico Sperimentale della Lombardia e dell'Emilia-Romagna "Bruno Ubertini", Brescia, Italy

⁵Dipartimento di Malattie Infettive, Istituto Superiore di Sanità, Rome, Italy

Abstract

Introduction. Yersinia enterocolitica (Ye) species is divided into 6 biotypes (BT), 1A, 1B, 2, 3, 4, 5 classified based on biochemical reactions and about 70 serotypes, classified based on the structure of the lipopolysaccharide O-antigen. The BT1A is considered non-pathogenic, while the BT 1B-5 are considered pathogenic.

Methods. Evaluate the distribution of eleven chromosomal and plasmid virulence genes, ail, ystA, ystB, myfA, breP, fes, fepD, ymoA, sat, virF and yadA, in 87 Ye strains isolated from food, animals and humans, using two SYBR Green real-time PCR platforms.

Results. The main results showed the presence of the *ail* and *ystA* genes in all the pathogenic bioserotypes analyzed. The ystB, on the other hand, was identified in all non-pathogenic strains biotype 1A. The target fes, fepD, sat and hreP were found in both pathogenic biotypes and in BT1A strains. The myfA gene was found in all pathogenic biotype and in some Ye BT1A strains. The virF and yadA plasmid genes were mainly detected in bioserotype 4/O:3 and 2/O:9, while ymoA was identified in all strains.

Conclusions. The two molecular platforms could be used to better define some specific molecular targets for the characterization and rapid detection of Ye in different sources which important implications for food safety and animal and human health.

INTRODUCTION

The genus Yersinia has been recently classified to Yersiniaceae family [1] and comprises of 28 species (https://lpsn.dsmz.de/genus/yersinia), three of which are pathogenic to humans Y. pestis, Y. enterocolitica and Y. pseudotuberculosis [2]. Yersinia enterocolitica (Ye) is Gram-negative and psychrophilic enteropathogen [3]. In fact, Ye has an optimal growth temperature between 28-30 °C but is able to growth from 2 and 42 °C [4]. Yersiniosis is usually a self-limiting disease and gastrointestinal symptoms pre-dominate, but extra-intestinal disorders may also appear [5]. Children under 5 years of age, immunocompromised people, and elderly are Key words

- Yersinia enterocolitica
- SYBR Green real-time PCR
- virulence genes

more likely to get sick and to have a more serious illness [4].

In Europe, yersiniosis is the third most frequently reported foodborne zoonosis in humans with a stable trend in 2015-2019 [6]. In Italy the number of reported cases is very low compared with other EU/EEA member states, because the notification of yersiniosis is voluntary, and the number of cases is probably underestimated [7].

Ye is widely distributed in the environment and pigs are the main reservoir of pathogenic strains, in particular, the bacterium colonized tonsils, lymph nodes and intestines. Ye is also isolated from other animals, such

as cattle, small ruminants, wild animals (e.g., deer) and pets (cats and dogs) [8, 9]. Humans become infected by ingesting undercooked pork contaminated with Ye, but also milk and dairy products, vegetables and untreated water [10, 11]. Although pork meat is the major source of pathogenic Ye, in recent years, fresh vegetables have been linked to several foodborne outbreaks, particularly spinach and salad [12, 13]. The Ye species is highly heterogeneous and is divided into 6 biotypes (BT), 1A, 1B, 2, 3, 4, 5, on the basis of biochemical tests and about 70 serotypes [14]. The biotypes show differing pathogenic potential, in particular: biotype 1A (BT1A) is considered non-pathogenic, biotypes 2-5 weakly pathogenic, and biotype 1B highly pathogenic [15]. Pathogenic Ye have historically been defined as carriers of a 70 kb virulence plasmid (pYV), which has genes encoding adhesin A (yadA) and a transcriptional regulator gene (virF); in addition, at the chromosomal level they harbour virulence genes as invA (invasin), ail (attachment and invasion locus). vstA (stable Yersinia toxin A), and mvfA (mucoid Yersinia factor A) [16].

BT1A strains are considered non-pathogenic because they do not have the pYV and some chromosomal virulence genes such as *ail*. However, studies have shown that some biotype 1A strains, particularly those isolated from faeces during a gastrointestinal illness, could be pathogenic [17]. Although BT1A may lack the pYV plasmid, alternative virulence factors, including the thermostable toxin *ystB* and *breP* [18] may be present. BT1A is the most heterogeneous of the six Ye biotypes and includes a wide range of serotypes of which O:5, O:6,30, O:6,31, O:7,8, O:10, as well as not-typable (NT) O strains, are most often reported [19]. The most common bioserotypes causing human yersiniosis in Europe are Ye 4/O:3 and 2/O:9 [6].

ISO 10273 (International Organization for Standardization) standard cultural method for the detection and isolation of *Ye* from food samples are laborious and time-consuming to differentiate pathogenic and nonpathogenic strains [20, 21].

To meet the requirement for faster analysis, ISO approved a standard method for rapid identification of pathogenic Ye based on detection of the chromosomelocalized *ail* gene which is present in all pathogenic bioserotype by real-time PCR [22]. However, recent studies have indicated that in some non-pathogenic 1A strains of Ye, the *ail* gene can be detected [23]. Therefore, it would be useful to consider other pathogenicity targets of the strains to have a more complete genomic characterisation. Recently, a study analyzed by realtime PCR the distribution of the yadA, virF, inv, ystA, ystB, myfA, breP and ymoA genes in Ye strains in order to select useful target genes for assessing the presence of pathogenic Ye [18].

The aim of the present study is to extend the examination of the distribution of virulence genes in nonpathogenic and pathogenic *Ye* strains, isolated from animal, food and human samples. Additional molecular targets would allow for a more complete characterization of *Ye* strains, and the evaluation of the distribution of these genes which have virulence potential and are less investigated. Thus, besides the virulence genes *ail*, *ystA*, *ystB*, *myfA*, *virF* and *yadA*, the other target genes analyzed are: *breP* (host reactive element), *sat* (streptogramin acetyltransferase), *fepD* (enterochelin transporter ABC), *fes* (enterochelin esterase) and *ymoA* (*Yersinia* modulating protein) [24], using two molecular real-time PCR SYBR Green platforms.

MATERIALS AND METHODS Identification and typing of bacterial strains

A total of 87 Ye strains, 82 isolates from 2005 to 2015 and 5 isolates from 1980 to 1985 in Italy, were identified at biochemical level and subsequently biotyped and serotyped. These strains were isolated from human (n=39), in particular stools (n=24), blood (n=3), appendix (n=1) and unknown (n=11), from animals (n=12), in particular faeces (n=7), amygdala (n=2) and unknown (n=3) and from food (n=36).

Identification of Ye species was performed using the API® 20E system (bioMérieux). Ye biotyping was carried out according to the ISO 10273 scheme, based on biochemical reactions, in particular: pyrazinamidase and lipase activity, production of indole, production of acids from xylose, trehalose and hydrolysis of esculin. Serotyping was performed using O-antisera for sero-groups O:3, O:5, O:8, O:9 and O:27 purchased from Biolife (Biolife Italiana, Milan, Italy).

Genomic DNA extraction

The strains were grown in Tryptone Soy Broth (Biolife Italiana, Milan, Italy) at 30 °C for 24-48h. Two mL of each broth culture was subjected to DNA extraction according to the protocol of Peruzy *et al.* [18]. The extracted DNA was used as a template for real-time PCR.

Real-time PCR-based protocol

For the molecular characterization of Ye virulence genes, two SYBR Green real-time PCR platforms were developed with two different thermal profile. The first platform, a PCR reaction in singleplex, containing: 1X SsoAdvanced SYBR Green PCR Master Mix (Bio-Rad), one of the primer (Table 1) at different concentrations, of the virulence genes ail (250 nM), vstA (100 nM), ystB (150 nM), myfA (300 nM) and 3 µl of DNA in a final volume of 25 µl. The thermal profile was: 95 °C for 5 min, 35 cycles at 95 °C for 10 s and 60 °C for 30 s, followed by a thermal cycle (65-95 °C) necessary for the analysis of the melting curve. The second platform, a PCR reaction in singleplex, containing: 1X SsoAdvanced SYBR Green PCR Master Mix (Bio-Rad, USA), one of the primer (Table 1), at different concentrations, of the virulence genes *breP* (300 nM), and 250 nM of the other virulence genes virF, yadA, ymoA, fes, fepD, and sat and 3 μ l of DNA in a final volume of 25 μ l. The thermal profile of the reaction was: 95 °C for 5 min, 35 cycles at 95 °C for 60 s, 60 °C for 60 s, 72 °C for 60 s, followed by a thermal cycle (65-95 °C) necessary for the analysis of the melting curve. The specificity of the reaction is given by the detection of the melting temperature (T_m) of the amplification products after the last reaction cycle. The melting curve was visualized with the software MxPro (Mx3005P v 4.00 - Agilent).

Table 1

Primers used for SYBR Green real-time PCR

Gene	Primer sequence (5'->3')	Amplicon size (bp)	Reference
ail	ACTCGATGATAACTGGGGAG	170	[25]
	CCCCCAGTAATCCATAAAGG		
ystA	ATCGACACCAATAACCGCTGAG	79	[26]
	CCAATCACTACTGACTTCGGCT		
ystB	GTACATTAGGCCAAGAGACG	146	[26]
	GCAACATACCTCACAACACC		
myfA	CAGATACACCTGCCTTCCATCT	272	[27]
	CTCGACATATTCCTCAACACGC		
hreP	GCCGCTATGGTGCCTCTGGTGTG	757	[24]
	CCCGCATTGACTCGCCCGTATC		
virF	GGCAGAACAGCAGTCAGACATA	591	[25]
	GGTGAGCATAGAGAATACGTCG		
yadA	TAAGATCAGTGTCTCTGCGGC	747	[28]
	TAGTTATTTGCGATCCCTAGCAC		
утоА	GACTTTTCTCAGGGGAATAC	330	[29]
	GCTCAACGTTGTGTGTGTCT		
fes	GCCGGCAGGCACAGCGTAAT	561	[30]
	GGCCAACCCACCCAAAACTT		
fepD	GTGTGATTGCCTTACTATTG	381	[30]
	CGGTCATCCTTTTATTACGG		
sat	CCGATGGTGGGGTTTTCTCAAG	456	[24]
	GGGATTACCGCCGACCACACTA		

RESULTS

The results obtained of serotyping and biotyping of the 87 Ye strains have identified four biotypes and several serotype: 1A/O:5 (n=15), 1A/O:8 (n=9), 1A/NT (n=25), 2/O:9 (n=5); 3/O:5,27 (n=1); 4/O:3 (n=32). The results obtained analysing these strains through SYBR Green real-time PCR showed a close association between the *ail* gene, considered an exclusive indicator of pathogenicity, and the *ystA* gene as they were found in all strains belonging to the pathogenic biotypes (*Table 2*). In contrast, *ystB* was found exclusively in all

Table 2

The distribution of genes in Ye strains grouped by biotype/serotype

	· · · · · ·										
Biotype/serotype (n)	ail	ystA	ystB	myfA	hreP	virF	yadA	fepD	fes	утоА	sat
4/O:3 (32)	100%	100%	-	100%	100%	40.1%	40.1%	43.8%	43.8%	100%	96.8%
2/O:9 (5)	100%	100%	-	100%	100%	60.0%	60.0%	100%	100%	100%	100%
3/0:5.27 (1)	100%	100%	-	100%	100%	-	-	100%	100%	100%	100%
1A/NT (25)	-	-	100%	12.0%	72.0%	-	-	96.0%	96.0%	100%	84.0%
1A/O:5 (15)	-	-	100%	-	93.3%	-	-	100%	100%	100%	100%
1A/O:8 (9)	-	-	100%	-	100%	-	-	100%	100%	100%	100%

non-pathogenic biotype 1A strains. The presence of the *virF* and *yadA* plasmid genes appears to be exclusive to pathogenic biotypes, found in 42.1% (n=16) of the total of the pathogenic strains (*Table 2*).

The *myfA* gene was detected in all pathogenic strains (n=38), but also in 3 strains of BT1A strains (6.1%). The *fes* and *fepD* genes, which encode for factors capable of capturing and utilizing host iron, were found in synergy with each other in both pathogenic and non-pathogenic biotypes. The *breP* and sat genes appear present in high percentages in all biotypes analyzed, as is the *ymoA* gene, which is present in 100% of strains (*Table 2*). The distribution of bioserotypes showed the prevalence of 4/O:3 and 2/O:9 mainly in human samples, and 4 of 32 strains of *Ye* 4/O:3 were found in pigs, whereas BT1A was found mainly in a wide variety of food samples, but also, less frequently, in animals and humans (*Table 3*).

DISCUSSION

The results showed a non-uniform distribution of the different target genes in the various *Ye* strains. Chromosomal virulence genes are very important elements that determine the pathogenic capabilities of *Ye*. These include the attachment invasion locus (*ail*), which encodes the outer membrane proteins responsible for adhesion, and is considered the target gene for detection of pathogenic *Ye* according to ISO/TS 18867:2015.

Furthermore, Ye has the ability to produce three types of YstI toxins (YstA, YstB and YstC) encoded by ystA, ystB and ystC genes, respectively, which play a crucial role in the origin of diarrhea [31]. As expected, all strains belonging to pathogenic biotypes, analyzed in this work, contain the *ail* gene together with the ystA gene. Ye BT1A mainly produce the enterotoxin YstB and rarely YstC. The ability of some BT1A strains to cause illness gives indirect evidence that YstB plays an important role in yersiniosis, as suggested in some studies [31, 32]. Indeed, in the present study the ystB was detected in 100% of BT1A strains, results in agreement with literature data [24, 33].

The pYV plasmid undoubtedly plays an important role in pathogenicity and carry virulence factors, such as *virF*, a regulatory gene, which encodes the transcriptional activator of several genes, including the plasmid virulence gene *yadA* involved in the *Yersinia* invasion process. However, plasmids are unstable structures, and *Ye* BT1A generally do not harbour plasmids [31, 34].

Table 3

The distribution of different sources (food, animals and human) in Yersinia enterocolitica biotypes (BT)

Source (n)	Specific source	BT1A	BT2	BT3	BT4
Animals (12)	Bear	1			
	Bovine	1			
	Sheep	1		1	
	Swine	4			4
Food (36)	Beef	1			
	Chicken meat	4			
	Fish	2			
	Fresh vegetables	2			
	Milk	1			
	Minced meat of beef and pork	3			
	Pork meat	8			
	Ready to eat pork meat	2			
	Ready to eat vegetables	12			
	Sheep meat	1			
Humans (39)	Humans	6	5		28

The results reported in our study confirm that plasmid virulence genes are not present in any strain belonging to the non-pathogenic biotype (BT1A), whereas they are found in about 42.1% of the total pathogenic biotypes, with a different distribution between bioserotype 4/O:3 where plasmid genes are present in 40.1% and bioserotype 2/O:9 where *virF* and *yadA* are present in 60%.

MyfA, plays an important role at the beginning of infection by promoting adhesion to enterocytes. The *myfA* gene has been found in *Ye* strains of bioserotype 4/O:3 isolated from human and pigs [33] and has also been detected in some *Ye* strains of BT1A [24].

These results are confirmed in this study, and we also found *myfA* in all strains of bioserotype 2/O:9 isolated from human and in the only strain belonging to bioserotype 3/O:5,27 isolated from animal.

The ymoA gene was present in all Ye isolates studied, as reported in several studies [18, 24, 33]. According to recent studies, the ymoA gene is the main regulator of yst gene expression and other virulence genes [34].

The *breP* gene, which encodes the bacterial subtilisin/ Kexin-like protease, and the *sat* gene, which encodes the acetyltransferase streptogramin, are both present in high percentage in all Ye strains. These results are in agreement with data from some other Authors [18, 24], but seem to be in contrast with data reported by Morka *et al.* [33]. Few studies have described the *breP* gene, encoding for a bacterial protease, relevant for full virulence of Ye [35]. Finally, the *fes* and *fepD* genes, involved in iron capture and utilization, contribute to the growth of microorganism [30].

The genes *fepD* and *fes* were found in 100% or percentage near 100% in all isolates of all bioserotypes except for BT4/O:3 where the prevalence of these genes was about 44%. BT1A strains possessing the genes *ystB*, *breP*, *sat*, *fes*, *fepD* and *myfA* genes may have a virulence potential with respect to causing infections in humans and animals [24, 36]. Although in the study of Campioni and Falcão [36], *myfA* appears to have a higher prevalence in the BT1A strains (55%) than in our study.

Furthermore, the lack of identification of BT1B strains in this work, appears to be in line with what has been reported in the literature regarding the limited presence or absence of this biotype in Europe [4, 6].

Overall, the study seems to confirm what other authors have already pointed out regarding the distribution of these virulence genes in *Ye* strains.

CONCLUSIONS

The isolation and biotyping of Ye are currently difficult and time-consuming, but biotyping remains important as a basis for assessing the pathogenicity of isolated strains. The detection of this microorganism by means of molecular biology tools as a real-time PCR allows the quick detection of pathogenic Ye in food, animal and human and can be a valid support to classical microbiology techniques. Although, the ail gene remains the main virulence marker as reported by ISO/ TS 18867:2015, other virulence genes are important in evaluating the pathogenicity of Ye, such as the *ystB* gene to identify strains of biotype 1A. Therefore, the use of two SYBR Green real-time PCR platforms in this study, allowed the rapid detection of the eleven virulence genes, in pathogenic biotypes and in BT1A of Ye. It also allowed to highlight a diversity in the distribution of virulence genes in Ye strains isolated from different sources, which has valuable implications in terms of food safety and animal and human health from a One Health perspective.

Authors' contributions

Conceptualization: ED and EV; methodology: ED; investigation: ED, EV, SF, SL, GF, SB and SO; re-

sources: ED, SF, SL, GF, SB and SO; writing-original draft preparation: EV and ED; writing-review and editing: ED, SF, SL, GF, SB; supervision: ED. All authors have read and agreed to the published version of the manuscript.

REFERENCE

- 1. Adeolu M, Alnajar S, Naushad S, Gupta RS. Genomebased phylogeny and taxonomy of the 'Enterobacteriales': proposal for Enterobacterales ord. nov. divided into the families Enterobacteriaceae, Erwiniaceae fam. nov., Pectobacteriaceae fam. nov., Yersiniaceae fam. nov., Hafniaceae fam. nov., Morganellaceae fam. nov., and Budviciaceae fam. nov. Int J Syst Evol Microbiol. 2016;66(12):5575-99. doi: 10.1099/ijsem.0.001485
- Hammerl J, Barac A, Erben P, Fuhrmann J, Gadicherla A, Kumsteller F, Lauckner A, Müller F, Hertwig S. Properties of two broad host range phages of *Yersinia enterocolitica* isolated from wild animals. Int J Mol Sci. 2021;22:11381. doi: 10.3390/ijms222111381
- Gill CO, Reichel MP. Growth of the cold-tolerant pathogens Yersinia enterocolitica, Aeromonas hydrophila and Listeria monocytogenes on high-pH beef packaged under vacuum or carbon dioxide. Food Microbiol. 1989;6:223-30 doi:10.1016/S0740-0020(89)80003-6
- 4. Bari ML, Hossain MA, Isshiki K, Ukuku D. Behavior of Yersinia enterocolitica in Foods. J Pathog. 2011;2011:420732. doi: 10.4061/2011/420732
- Rodio DM, Bressan A, Ambrosi C, Scribano D, Tolli R, Mansour W, Speziale F, Antonelli G, Trancassini M, Pietropaolo V. Yersinia enterocolitica in Italy: A case of septicemia and abdominal aortic aneurysm infection. Front Med (Lausanne). 2018;5:156. doi: 10.3389/ fmed.2018.00156
- EFSA. The European Union One Health 2020 Zoonoses Report. EFSA J. 2021;19(12):6971. doi: 10.2903/j.efsa.2021.6971
- European Centre for Disease Prevention and Control (ECDC). Yersiniosis. In: ECDC. Annual epidemiological report for 2019. Stockholm: ECDC; 2021.
- Rakin A, Garzetti D, Bouabe H, Sprague LD. Yersinia enterocolitica. In: Tang Y-W, Sussman M, Liu D, Poxton I, Schwartzman J (Eds). Molecular medical microbiology, 2nd ed. London: Academic Press; 2015. p. 1319-44. doi: 10.1016/B978-0-12-397169-2.00073-1
- 9. Ye Q, Wu Q, Hu H, Zhang J, Huang H. Prevalence and characterization of *Yersinia enterocolitica* isolated from retail foods in China. Food Control. 2016;61:20-7 doi: 10.1016/j.foodcont.2015.09.016
- Chlebicz A, Śliżewska K. Campylobacteriosis, salmonellosis, yersiniosis, and listeriosis as zoonotic foodborne diseases: A review. Int J Environ Res Public Health. 2018;15(5):863. doi: 10.3390/ijerph15050863
- Shoaib M, Shehzad A, Raza H, Niazi S, Khan I.M, Akhtar W, Safdar W, Wang Z. A comprehensive review on the prevalence, pathogenesis and detection of *Yersinia enterocolitica*. RSC Adv. 2019;70:41010-21 doi: 10.1039/ C9RA06988G
- Espenhain L, Riess M, Müller L, Colombe S, Ethelberg S, Litrup E, Jernberg C, Kühlmann-Berenzon S, Lindblad M, Hove NK, Torpdahl M, Mörk MJ. Cross-border outbreak of *Yersinia enterocolitica* O3 associated with imported fresh spinach, Sweden and Denmark, March 2019. Euro Surveill. 2019;24:1900368 doi: 10.2807/1560-7917. ES.2019.24.24.1900368

Conflict of interest statement

The Authors declare no conflict of interest.

Received on 2 May 2023. Accepted on 26 September 2023.

- MacDonald E, Einöder-Moreno M, Borgen K, Thorstensen Brandal L, Diab L, Fossli Ø, Guzman Herrador B, Hassan AA, Johannessen GS, Johansen EJ, Jørgensen Kimo R, Lier T, Paulsen BL, Popescu R, Tokle Schytte C, Sæbø Pettersen K, Vold L, Ørmen Ø, Wester AL, Wiklund M, Nygård K. National outbreak of Yersinia enterocolitica infections in military and civilian populations associated with consumption of mixed salad, Norway, 2014. Euro Surveill. 2016;21:30321. doi: 10.2807/1560-7917.es.2016.21.34.30321
- Fredriksson-Ahomaa M. Yersinia enterocolitica. In: Foodborne diseases, 3rd ed. London: Elsevier, Academic Press; 2017. p. 223-33 doi: 10.1016/B978-0-12-385007-2.00009-7
- 15. Reuter S, Connor TR, Barquist L, Walker D, Feltwell T, Harris SR, Fookes M, Hall ME, Petty NK, Fuchs TM, Corander J, Dufour M, Ringwood T, Savin C, Bouchier C, Martin L, Miettinen M, Shubin M, Riehm JM, Laukkanen-Ninios R, Sihvonen LM, Siitonen A, Skurnik M, Falcão JP, Fukushima H, Scholz HC, Prentice MB, Wren BW, Parkhill J, Carniel E, Achtman M, McNally A, Thomson NR. Parallel independent evolution of pathogenicity within the genus *Yersinia*. Proc Natl Acad Sci USA. 2014;18:6768-73 doi: 10.1073/pnas.1317161111
- Rivas L, Strydom H, Paine S, Wang J, Wright J. Yersiniosis in New Zealand. Pathogens. 2021;10:191. doi:10.3390/ pathogens10020191
- Bhagat N, Virdi JS. The enigma of Yersinia enterocolitica biovar 1A. Crit Rev Microbiol. 2011;37(1):25-39 doi: 10.3109/1040841X.2010.506429
- Peruzy MF, Murru N, Perugini AG, Capuano F, Delibato E, Mercogliano R, Korkeala H, Proroga YTR. Evaluation of virulence genes in *Yersinia enterocolitica* strains using SYBR Green real-time PCR. Food Microbiol. 2017;65:231-5. doi: 10.1016/j.fm.2017.03.004
- Tennant SM, Grant TH, Robins-Browne RM. Pathogenicity of *Yersinia enterocolitica* biotype 1A. FEMS Immunol Med Microbiol. 2003;38:127-37. doi: 10.1016/ S0928-8244(03)00180-9
- European Committee for Standardization. Microbiology of the food chain - Horizontal method for the detection of pathogenic *Yersinia enterocolitica*. Geneva: International Organization for Standardization; 2017. (EN ISO 10273:2017).
- 21. Fredriksson-Ahomaa M, Joutsen S, Laukkanen-Ninios R. Identification of *Yersinia* at the species and subspecies levels is challenging. Curr Clin Microbiol Rep. 2018;5:135-42. doi: 10.1007/s40588-018-0088-8
- 22. International Organization for Standardization. Microbiology of the food chain - Polymerase chain reaction (PCR) for the detection of food-borne pathogens - Detection of pathogenic *Yersinia enterocolitica and Yersinia pseudotuberculosis*. Geneva: International Organization for Standardization; 2015. (ISO/TS 18867:2015).
- Joutsen S, Johansson P, Laukkanen-Ninios R, Björkroth J, Fredriksson-Ahomaa M. Two copies of the *ail* gene found in *Yersinia enterocolitica* and *Yersinia kristensenii*. Vet Microbiol. 2020;247:108798. doi: 10.1016/j.vetmic.2020.108798

- Bhagat N, Virdi JS. Distribution of virulence-associated genes in *Yersinia enterocolitica* biovar 1A correlates with clonal groups and not the source of isolation. FEMS Microbiol Lett. 2007;266:177-83 doi: 10.1111/j.1574-6968.2006.00524.x
- Bhaduri S, Cottrell B, Pickard AR. Use of a single procedure for selective enrichment, isolation, and identification of plasmid-bearing virulent *Yersinia enterocolitica* of various serotypes from pork samples. Appl Environ Microbiol. 1997;63:1657-60 doi: 10.1128/aem.63.5.1657-1660.1997
- Thoerner P, Bin Kingombe CI, Bögli-Stuber K, Bissig-Choisat B, Wassenaar TM, Frey J, Jemmi T. PCR detection of virulence genes in *Yersinia enterocolitica* and *Yersinia pseudotuberculosis* and investigation of virulence gene distribution. Appl Environ Microbiol. 2003;69:1810-6 doi: 10.1128/AEM.69.3.1810-1816.2003
- 27. Kot B, Trafny EA. The application of PCR to the identification of selected virulence markers of *Yersinia* genus. Pol J Vet Sci. 2004;7:27-31.
- Estrada CS, Velázquez LC, Favier GI, Genaro MS, Escudero ME. Detection of *Yersinia* spp. in meat products by enrichment culture, immunomagnetic separation and nested PCR. Food Microbiol. 2012;30:157-63. doi: 10.1016/j.fm.2011.10.014
- Grant T, Bennett-Wood V, Robins-Browne RM. Identification of virulence-associated characteristics in clinical isolates of *Yersinia enterocolitica* lacking classical virulence markers. Infect Immun. 1998;66:1113-20. doi: 10.1128/ IAI.66.3.1113-1120.1998

- Schubert S, Fischer D, Heesemann J. Ferric enterochelin transport in *Yersinia enterocolitica*: molecular and evolutionary aspects. J Bacteriol. 1999;181:6387-95. doi: 10.1128/JB.181.20.6387-6395.1999
- Platt-Samoraj A. Toxigenic properties of *Yersinia entero-colitica* biotype 1A. Toxins (Basel). 2022;14:118. doi: 10.3390/toxins14020118
- 32. Singh I, Virdi JS. Production of *Yersinia* stable toxin (YST) and distribution of *yst* genes in biotype 1A strains of *Yersinia enterocolitica*. J Med Microbiol. 2004;53:1065-8. doi: 10.1099/jmm.0.45527-0
- 33. Morka K, Wałecka-Zacharska E, Schubert J, Dudek B, Woźniak-Biel A, Kuczkowski M, Wieliczko A, Bystroń J, Bania J, Bugla-Płoskońska G. Genetic diversity and distribution of virulence-associated genes in *Y. enterocolitica* and *Y. enterocolitica*-like isolates from humans and animals in Poland. Pathogens. 2021;10:65. doi: 10.3390/ pathogens10010065
- Bancerz-Kisiel A, Pieczywek M, Łada P, Szweda W. The most important virulence markers of *Yersinia enterocolitica* and their role during infection. Genes (Basel). 2018;9:235. doi: 10.3390/genes9050235
- Young GM, Miller VL. Identification of novel chromosomal loci affecting *Yersinia enterocolitica* pathogenesis. Mol Microbiol. 1997;25:319-28. doi: 10.1046/j.1365-2958.1997.4661829.x
- Campioni F, Falcão JP. Genotypic diversity and virulence markers of *Yersinia enterocolitica* biotype 1A strains isolated from clinical and non-clinical origins. APMIS. 2014;122:215-22 doi: 10.1111/apm.12126