

Identification of the STEC serogroups mainly associated with human infections by Real-Time PCR amplification of O-associated genes

1. Aim and field of application

The present method concerns the identification by Real-Time PCR amplification of the genes associated with the O-antigens of the STEC serogroups mainly associated to severe human disease. The method is intended for the identification of the serogroup of *E. coli* strains isolated in pure culture. The serogroups in the field of application of the present method are: O26, O45, O55, O80, O91, O103, O104, O111, O113, O121, O128, O145, O146 and O157. All these serogroups have been frequently reported in human infections.

2. Definitions

STEC: strains possessing the genes encoding the Shiga-toxins. **O-antigen:** serogroups or “O” antigens are identified by numbers, counting from 1 to 187, and the serogroups list is evolving constantly.

Primers: oligonucleotides used to prime the amplification of a template by DNA polymerase.

Taqman probes: oligonucleotides labelled with a fluorophore covalently attached to the 5'-end and a quencher at the 3'-end, used to increase the specificity of Real-Time PCR.

3. Procedure

3.1 DNA extraction and purification

An appropriate nucleic acid extraction procedure for Gram-negative bacteria should be used to prepare DNA, from a pure culture from liquid or from solid media, according to the selected procedure.

3.2 Real-Time PCR amplification

The protocol is based on the 5' nuclease PCR assay. Considering that Real-Time PCR may use different probes labelling chemistry and run on different instruments, the amplification conditions to be applied may vary depending on the system used. Refer to the instructions supplied with the instrument and kit of choice.

The primers and probes to be used are listed in Table 1. The chemistry of the reporter and quencher fluorophores is not indicated being largely dependent on the Real-Time PCR systems available in each laboratory. The bibliographic references for the primers and probes sequences are indicated in the table. The Real-Time PCR procedures for the detection of the genes associated to the top-5 serogroups (O157, O26, O103, O111 and O145) and to serogroup O104 correspond to those illustrated in the two EURL-VTEC methods, available online in the website, and respectively named as follow:

- "Identification and characterization of Verocytotoxin-producing *Escherichia coli* (VTEC) by Real Time PCR amplification of the main virulence genes and the genes associated with the serogroups mainly associated with severe human infections" (EURL-VTEC_Method_02_Rev_n);
- "Detection and identification of Verocytotoxin-producing *Escherichia coli* (VTEC) O104:H4 in food by Real Time PCR" (EURL-VTEC_Method_04_Rev_n).

As stated above, amplification conditions will depend on the system used and need to be fine-tuned in each laboratory. However, a standard two-step thermal profile used at EURL-VTEC is the following:

95 °C X 10'
35 cycles of
95 °C X 15''
60 °C X 1'

The following thermal profile applies to the detection of O80 and O103 serogroup-associated genes only:

95 °C X 10'
35 cycles of
95 °C X 15''
55 °C X 1'

The reaction should be assembled applying the following instructions:

Buffer 10X	to 1X (MgCl ₂ 3mM)
Primer Fwd	500 nM
Primer Rev	500 nM
Probe	200nM
DNA	2 µl of DNA purified from 1 ml of culture and diluted 1:10 can be sufficient
Water	to final volume

Table 1.

Primers and probes used for Real Time PCR assays.

Target gene (Ref)	Forward primer, reverse primer and probe sequences (5'-3')	Amplicon size (bp)	Location within sequence	GenBank accession number
wzxO26 (Perelle <i>et al.</i> , 2004)	FWD: CGCGACGGCAGAGAAAATT	135	5648–5666	AF529080
	REV: AGCAGGCTTTTATATTCTCCAACCTT		5757–5782	
	Probe: CCCCCTTAAATCAATACTATTTTCACGAGGTTGA		5692–5724	
wzxO45 (USDA, 2012)	FWD: CGTTGTGCATGGTGGCAT	72	7472-7489	AY771223
	REV: TGGCCAAACCAACTATGAACTG		7543-7522	
	Probe: ATTTTTTGCTGCAAGTGGGCTGTCCA		7494-7519	
wzxO55 (EURL-VTEC)	FWD: AATTAACGAACATAACACCCAACC	101	11516-11493	AF461121
	REV: ATATCTCTTCGTTACTGTGTGTATTTTC		11416-11442	
	Probe: ACCTCCCGCTAAAACCCCAACTCTAGTAG		11489-11461	
wzyO80 (EURL-VTEC)	FWD: TGAGAGCCAAGATCCAAGCA	158	15098-15079	AB812032
	REV: TGGGCCATATTCGAAGTTTGAA		14941-14962	
	Probe: TCCAAGATTCCACGTTGAT		15013-14994	

Target gene (Ref)	Forward primer, reverse primer and probe sequences (5'-3')	Amplicon size (bp)	Location within sequence	GenBank accession number
wzyO91 (Perelle <i>et al.</i> , 2004)	FWD: CGATTTTCTGGAATGCTTGATG	105	9433–9454	AY035396
	REV: CAATACATAGTTTGTATTTGTGTTTAAAGTTTAAT		9504–9537	
	Probe: CCTGGGTTGTTAGGAACAATTTTCAGCACTTC		9457–9487	
wzxO103 (Perelle <i>et al.</i> , 2005)	FWD: CAAGGTGATTACGAAAATGCATGT	99	4299–4323	AY532664
	REV: GAAAAAAGCACCCCGTACTTAT		4397–4375	
	Probe: CATAGCCTGTTGTTTTAT		4356–4373	
wzxO104 (Bugarel <i>et al.</i> , 2010)	FWD: TGTCGCGCAAAGAATTTCAAC	100	2,333,750–2,333,730	CU928145
	REV: AAAATCCTTTAACTATACGCC		2,333,673–2,333,651	
	Probe: TTGGTTTTTTTTGTATTAGCAATAAGTGGTGTC		2,333,724–2,333,693	
wbdI011 (Perelle <i>et al.</i> , 2004)	FWD: CGAGGCAACACATTATATAGTGCTTT	146	3464–3489	AF078736
	REV: TTTTGAATAGTTATGAACATCTTGTTTAGC		3579–3609	
	Probe: TTGAATCTCCCAGATGATCAACATCGTGAA		3519–3548	
wzyO113 (Perelle <i>et al.</i> , 2004)	FWD: GAGCGTTTCTGACATATGGAGTGA	107	3689–3712	AF172324
	REV: TTGCTATAAATGGAAGCCATTCTTT		3771–3795	
	Probe: TGCATGAAATGTTTAAATGCAGCGGGT		3738–3764	



Target gene (Ref)	Forward primer, reverse primer and probe sequences (5'-3')	Amplicon size (bp)	Location within sequence	GenBank accession number
wzxO121 (USDA, 2012)	FWD: AGGCGCTGTTTGGTCTCTTAGA	189	6839-6860	AY208937
	REV: GAACCGAAATGATGGGTGCT		7027-7008	
	Probe: CGCTATCATGGCGGGACAATGACAGTGC		6898-6925	
wzxO128 (Lin <i>et al.</i> , 2011)	FWD: tcgatcgtctgttcaggtt	196	8857-8876	AY217096
	REV: gaatgcaatgggcaattaac		9052-9033	
	Probe: ggggtgcacaattggcctcc		8918-8937	
wzxO145 (USDA, 2012)	FWD: AAA CTG GGA TTG GAC GTG G	135	2600222–2600240	AP019708
	REV: CCC AAA ACT TCT AGG CCC G		2600106–2600124	
	Probe: TGC TAA TTG CAG CCC TTG CAC TAC GAG GC		2600190–2600162	
wzyO146 (EURL-VTEC)	FWD: acattcggcggttttatctcgt	106	9144-9165	DQ465249
	REV: ggtcaaatctcgtgccataga		9228-9249	
	Probe: aattcaaggtgccactttca		9205-9227	
rfbEO157 (Perelle <i>et al.</i> , 2004)	FWD: TTTCACACTTATTGGATGGTCTCAA	88	348–372	AF163329
	REV: CGATGAGTTTATCTGCAAGGTGAT		412–435	
	Probe: AGGACCGCAGAGGAAAGAGAGGAATTAAGG		381–410	

3.3 Controls

DNA extracted from cultures of *E. coli* strains belonging to serogroups O26, O45, O55, O80, O91, O103, O104, O111, O113, O121, O128, O145, O146 and O157 should be used as positive control. The isolates provided by EURL-VTEC in the framework of the proficiency testing program can be used as reference strains. Moreover, the Real-Time PCR procedure requires an inhibition control.

3.4 Safety and protection devices

STEC strains can infect human beings at a very low infectious dose and can cause severe disease. Laboratory acquired infections have been reported. Therefore, handling STEC strains requires compliance to safety procedures in place (including the use of protection devices) and good laboratory practices. STEC are class 3 pathogens and in some countries their handling is allowed in CL 3 laboratory only.

4. References

- Bugarel M, Beutin L, Martin A, Gill A, Fach P. Micro-array for the identification of Shiga toxin-producing *Escherichia coli* (STEC) seropathotypes associated with Hemorrhagic Colitis and Hemolytic Uremic Syndrome in humans. *Int J Food Microbiol.* 2010 Sep 1;142(3):318-29.
- Lin A, Sultan O, Lau HK, Wong E, Hartman G, Lauzon CR. O serogroup specific real time PCR assays for the detection and identification of nine clinically relevant non-O157 STECs. *Food Microbiol.* 2011 May;28(3):478-83.
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- United States Department of Agriculture. Primer and Probe Sequences and Reagent Concentrations for non-O157 Shiga Toxin-Producing *Escherichia coli* (STEC) Real-Time PCR Assay. MLG 5B Appendix 1.01, https://www.fsis.usda.gov/wps/wcm/connect/0330211c-81ab-4e97-a9f3-d425f5759ee1/MLG_5B_Appendix_1_01.pdf?MOD=AJPERES