Analysis of the Clonal Relationship of Shiga Toxin-Producing Escherichia coli Serogroup O165:H25 Isolated from Cattle

Lutz Geue,* Thomas Selhorst, Christina Schnick, Birgit Mintel, and Franz J. Conraths

Friedrich-Loeffler-Institut, Federal Research Institute for Animal Health, Institute for Epidemiology, Seestrasse 55, 16868 Wusterhausen, Germany

Received 22 March 2005/Accepted 23 December 2005

Variations in time and space of a clonal group of *Escherichia coli* O165:H25 on a cattle farm were monitored. The virulence marker pattern (*stx* genes, *eae* gene, *hly*_{EHEC} gene, *katP* gene, *espP* gene, *efa* gene) suggests that *E. coli* O165:H25 of bovine origin may represent a risk for human infection.

Shiga toxin-producing Escherichia coli (STEC) is a group of zoonotic enteric pathogens (29). Human infections with some STEC serotypes, also designated enterohemorrhagic Escherichia *coli* (EHEC), result in hemorrhagic or nonhemorrhagic diarrhea, which may be complicated by hemorrhagic colitis (HC) and several renal sequelae, including the hemolytic-uremic syndrome (HUS) (20, 34, 41, 52). The mechanisms by which EHEC strains cause disease are not completely understood. The virulence factors include production of two major phageencoded toxins, Shiga toxin 1 (Stx1) and Shiga toxin 2 (Stx2), which can be produced alone or in combination. Stx1 and Stx2 are thought to cause the vascular endothelial damage observed in patients with HC and HUS (53). In addition to Stx, EHEC strains possess other virulence factors, such as the ability to cause attaching and effacing (eae) lesions in the large intestine (26). They often contain a plasmid that carries other potential virulence genes, such as an enterohemolysin gene (hly_{EHEC}), a catalase peroxidase gene (katP), and an extracellular serine protease gene (espP) (9, 10, 11, 45). The efa1 (E. coli factor for adherence) gene represents another intestinal colonization factor in an EHEC O111:H- strain (30). The efal gene of O111:H- is 99.9% homologous to the lifA gene in enteropathogenic E. coli. The efa1 locus is not physically linked to the locus for enterocyte effacement pathogenicity island (51).

Ruminants, especially cattle, are considered the primary reservoir for human EHEC infections (23). In this study, 38 bovine *E. coli* O165:H25 isolates were characterized to assess their potential to cause EHEC disease in humans. These isolates were detected over a 4-month period in eight different animals (Table 1) from a single group of beef cattle during a long-term study (19). Sporadic cases of human infections with O165:H25 and O165:H- in Europe and Canada have been described previously (6, 15, 17, 24). The four German O165:H25 and O165:H- strains of human origin (kindly supplied by H. Tschäpe, Robert Koch-Institute, Wernigerode, Germany) used in our study for comparison were associated with diarrhea in patients (54). Human HUS cases caused by O165 isolates have been reported from Denmark (15) and Germany (17).

Typing and subtyping of genes (stx_1 and/or stx_2 , eae, hly_{EHEC} , katP, and espP) associated with STEC were performed by LightCycler fluorescence PCR (40) and different Block cycler PCRs (Tables 2 and 3). A complete pattern of virulence markers was detected in most bovine isolates examined. An stx₂ gene, but not an stx_1 gene, was present in all O165 strains (Table 1). EHEC strains with stx_2 genes are significantly more frequently associated with HUS and other severe diseases than isolates with an stx_1 gene, which are more often associated with uncomplicated diarrhea or healthy individuals (6, 36). Stx2 is closely related to a family of Stx2 variants or alleles, which includes Stx2c (48), Stx2d (36), Stx2e (56), and Stx2f (47), although Stx2c and Stx2d are produced by STEC strains isolated from humans (36, 38, 43, 44, 48, 50). Additional genetic variants of the stx_2 gene have been described (5, 14, 28, 55). In contrast to STEC strains harboring stx_2 gene variants, however, STEC strains with the stx₂ genotype were significantly associated with HUS (17). An stx_2 gene with the $stx_{2-EDL933}$ genotype was found in all O165 isolates tested (Table 1). The nucleotide sequences of the A and B subunits of the stx_2 gene of the bovine O165:25 strain 02/09/010-1 (GenBank accession number AY652745) were identical to the sequences of the stx_2 gene of EHEC type strain EDL933 (35), a typical O157:H7 strain isolated from a HUS patient, with the sequences of the gene of bacteriophage 933W (37), and with the sequences of stx_2 genes of other E. coli O157:H7 strains of human origin isolated from EHEC outbreaks (25, 27). All bovine O165:H25 strains produced an Stx2 with high cytotoxicity for Vero cells as determined by an Stx enzyme-linked immunosorbent assay and by a Vero cell neutralization assay (49).

Not only factors that influence basal and inducible Stx production are important in STEC pathogenesis. In previous studies, it has been suggested that the *eae* and hly_{EHEC} genes likely contribute to STEC pathogenicity (3, 6, 42). Ritchie et al. (42) found both of these genes in all HUS-associated STEC isolates that they analyzed. The *stx*₂ genes were present in combination with *eae* genes in all O165 isolates that we obtained (Table 1). To date, 10 distinct variants of *eae* have been described (13, 21, 31, 39). Some serotypes were closely associated with a particular intimin variant (4, 12, 13, 55). Our study confirmed these associations. Like the O103 isolates, all bovine and human *E. coli* O165 strains were grouped into the *ɛ-eae* subgroup. Also, nucleotide sequencing of the bovine O165:H25 strain 02/09/010-1

^{*} Corresponding author. Mailing address: Friedrich-Loeffler-Institut, Federal Research Institute for Animal Health, Seestrasse 55, D-16868 Wusterhausen, Germany. Phone: 49 33979 80189. Fax: 49 33979 80222. E-mail: lutz.geue@fli.bund.de.

TABLE 1. Strains examined

Sampling day or year ^a	Source	Strain	Serotype	Virulence profile											
				<i>stx</i> ₁ gene	<i>stx</i> ₂ gene	Stx1 (toxin)	Stx2 (toxin)	<i>stx</i> ₂₋ Subtype	<i>eae</i> Subtype	<i>efa1/lifA</i> gene	hly _{EHEC} gene	<i>katP</i> gene	espP gene	Plasmid(s) (bp)	Genetic subcluster
Day 1	Cattle 17	02/17/009-9	O165:H25	_	+	_	+	stx2=EDI 933	ε	+	+	+	+	70, 55	7
	Cattle 24	02/24/007-1	O165:H25	_	+	_	+	Stx2 EDL033	3	+	+	+	+	70, 55	1
		02/24/007-2	O165:H25	_	+	_	+	Stx2-EDL933	ē.	+	+	+	+	70, 55	1
		02/24/007-3	O165·H25	_	+	_	+	stra EDL933	£	+	+	+	+	70 55	1
		02/24/007-4	O165·H25	_	+	_	+	stra EDL933	e	+	+	+	+	70,55	1
		02/24/007-5	O165:H25	_	+	_	+	stra EDL933	e	+	+	+	+	70,55	1
		02/24/007-6	O165:H25	_	+	_	+	stra EDL933	e	+	+	+	+	70,55	3
		02/24/007-7	O165:H25	_	+	_	+	str_=	e	+	+	+	+	70, 55	1
		02/24/007-9	O165:H25	_	+	_	+	str	e	+	+	+	+	70, 55	1
		02/24/007-10	O165:H25	_	+	_	+	str2-EDL933	c	+	+	+	+	70, 55	1
	Cattle 26	02/24/00/-10	0165.1125	_	+	_	+	str2-EDL933	c		- -	-		70, 55	1
	Cattle 20	02/26/006-1	0105.1125	_	- T		- T	strange Strang	ε			- -	- -	70, 55	1
		02/26/006-2	0105.H25	_	+	_	-	SIX _{2-EDL933}	ε	- -	- -	+		70, 55	1
		02/20/000-3	O105.H25	_	-	_	-	SIX _{2-EDL933}	ε	+	+	-	- T	70, 55	1
		02/26/006-4	O165:H25	_	+	_	+	SIX _{2-EDL933}	3	+	+	+	+	70, 55	9
		02/26/006-5	O165:H25	_	+	_	+	SIX _{2-EDL933}	3	+	+	+	+	70, 55	9
		02/26/006-6	O165:H25	_	+	-	+	stx _{2-EDL933}	8	+	+	_	+	67, 55	9
		02/26/006-7	O165:H25	_	+	-	+	$stx_{2-EDL933}$	з	+	_	_	_	_ 55	4
		02/26/006-8	O165:H25	_	+	_	+	$stx_{2-EDL933}$	в	+	+	+	+	70, 55	5
		02/26/006-9	O165:H25	_	+	-	+	$stx_{2-EDL933}$	З	+	-	_	_	55	6
		02/26/006-10	O165:H25	-	+	—	+	stx _{2-EDL933}	в	+	+	+	+	70, 55	1
Day 28	Cattle 25	02/25/007-1	O165:H25	-	+	_	+	$stx_{2-EDL933}$	ε	+	+	+	+	70, 55	1
		02/25/007-4	O165:H25	-	+	-	+	$stx_{2-EDL933}$	ε	+	+	+	+	70, 55	1
		02/25/007-5	O165:H25	_	+	-	+	$stx_{2-EDL933}$	ε	+	-	_	-	55	1
		02/25/007-6	O165:H25	_	+	-	+	$stx_{2-EDL933}$	ε	+	+	+	+	70, 55	1
		02/25/007-7	O165:H25	-	+	—	+	stx _{2-EDL933}	ε	+	+	+	+	70, 55	1
		02/25/007-8	O165:H25	_	+	-	+	stx2-EDL933	З	+	+	+	+	70, 55	1
		02/25/007-9	O165:H25	_	+	-	+	stx _{2-EDL933}	З	+	+	+	+	70, 55	1
		02/25/007-10	O165:H25	_	+	_	+	stx2-EDI 933	в	+	+	+	+	70, 55	1
Day 56	Cattle 9	02/09/010-1	O165:H25	_	+	_	+	stx2-EDI 933	ε	+	+	+	+	70, 55	8
5		02/09/010-2	O165:H25	_	+	_	+	stx2-EDL933	ε	+	+	+	+	70, 55	8
	Cattle 18	02/18/011-1	O165:H25	_	+	_	+	Stx2 EDL033	ε	+	+	+	+	70	10
		02/18/011-5	O165:H25	_	+	_	+	Stx2 EDL022	3	+	+	+	+	70, 55	10
		02/18/011-6	O165:H25	_	+	_	+	Stx2-EDL933	ē.	+	+	+	+	70, 55	10
	Cattle 25	02/25/008-1	O165:H25	_	+	_	+	Stx2-EDL933	ē.	+	+	+	+	70, 55	1
		02/25/007-3	O165·H25	_	+	_	+	stra EDL933	e	+	+	+	+	70,55	2
Day 119	Cattle 19	02/19/013-7	O165:H25	_	+	_	+	stra EDL933	e	_	+	+	+	70,55	11
	Cuttle 15	02/19/013-8	O165:H25	_	+	_	+	stra EDL933	e	+	+	+	+	70,55	8
		02/10/013-10	O165:H25	_	+	_	+	stv2-EDL933	6	+	+	+	+	70,55	8
1008	Uuman Cormony	02/19/013-10	O165.H	_	+	_	+	str2-EDL933	c c		-			75	0
1998	diarrhea	90-4902	0105.11-		Ŧ		T	<i>stx</i> _{2/2c}	ε	т	т	т	T	15	
1998	Human, Germany, diarrhea	98-8419-1 ^b	U165:H-	-	+	_	+	stx _{2/2c}	ε	+	+	+	+	95, 75	
1999	Human, Germany, diarrhea	99-2258 ^b	O165:H25	-	+	-	+	$stx_{2/2c}$	ε	+	+	+	+	95, 75	
2002	Human, Germany, diarrhea	02-11228 ^b	O165:H25	-	+	-	+	$stx_{2/2c}$	ε	+	+	+	+	95, 75	

 a^{a} On days -120 to 0 O165:H25 was not detected on eight occassions, on day 91 O165:H25 was not detected, and on days 147 to 511 O165:H25 was not detected by 17 investigations.

^b Kindly provided by H. Tschäpe.

(GenBank accession number AF479581) revealed a high level of sequence homology (99.7%) to the eae gene of an O103:H2 strain (31). E. coli O103:H2 strains have frequently been associated with human HUS cases in Europe. Like the eae gene, the hly_{EHEC} gene was found in association with severe disease in humans (45, 46). In our study, the $hly_{\rm EHEC}$ gene was detected in the O165 strains in which a 70-kb plasmid was also found (Table 1). The presence of a 70-kb plasmid was associated with the occurrence of additional virulence markers, such as the espP gene, and all but one isolate contained the katP gene (10, 11). The reason for the slightly smaller size (67 kb) of the plasmid in this isolate may be linked to fact that the katP gene was not present in this isolate (Table 1). The efa1 genes were detected in 37 of 38 bovine O165:H25 isolates with two DNA probes by colony hybridization, and the results were confirmed by Southern hybridization after pulsed-field gel electrophoresis (PFGE); in this analysis the DNA probes were

labeled with digoxigenin (DIG), primers lifA1 and lifA2 and primers lifA3 and lifA4 (Table 3) were used with a PCR DIG probe synthesis kit (Roche Diagnostics, Mannheim, Germany), and DIG Easy Hyb solution (Roche) was used for prehybridization and hybridization. The efa-1 gene was located on an approximately 240-kb XbaI fragment or on an approximately 440-kb NotI fragment. These fragments were missing in the isolate that was negative as determined by colony hybridization (Fig. 1). To determine this, slices of the plugs were digested for 4 h with XbaI, NotI (New England Biolabs GmbH, Frankfurt am Main, Germany), BlnI (AvrII), or SpeI (Amersham Biosciences Inc., Buckinghamshire, United Kingdom). The resulting fragments were separated in a 1.0% agarose gel (SeaKem Gold agarose; Cambrex) in $0.5 \times$ Tris-borate-EDTA at 10°C with a CHEF Mapper XA system. The pulse times for XbaI and NotI digests were increased from 5 to 50 s (gradient, 6 V/cm) during 25 h at a constant angel of 120°. The switch time

Target(s)	Primer	Sequence $(5'-3')^a$	Reference
stx_1 and stx_2	STEC-1	GA(AG) C(AG)A AAT AAT TTA TAT GTG	40
. 2	STEC-2	TGÀ TĠA TG(AG) CAA TTC AGT AT	33
stx_1	STEC-I HP-1	TTT ACG TTT TCG GCA AAT ACA GAG GGG AT-(FL)	40
1	STEC-I HP-2	(Red 640)-TCG TAC AAC ACT GGA TGA TCT CAG TGG G-Ph	
stx_2	STEC-II HP-1	TCA GGC ACT GTC TGA AAC TGC TCC TGT GTA-(FL)	40
2	STEC-II HP-2	(Red 705)-ACC ATG ACG CCG GGA GAC GTG GAC CT-Ph	
eae	eaeAF	GAC CCG GCA CAA GCA TAA GC	32
	eaeAR	CC ACCT GCA GCA ACA AGA GG	
eae	eae HP1	ACA GTT CTG AAA GCG AAA TGA TGA AGG c-(FL)	40
	eae HP2	(Red 640)-CCT GGT CAG CAG ATC ATT TTG CCA CT-Ph	
hly _{EHEC}	hlyAF	GCA TCA TCA AGC GTA CGT TCC	32
) Lille	hlyAR	AAT GAG CCA AGC TGG TTA AGC T	
hlyEHEC	hlyA HP1	GCA TGG CTC TTG ATG AAT TGC T-(FL)	40
7 LILC	hlyA HP2	(Red 705)-CAA CGG GAA GGA GAG GAT ATA AGT CAG-Ph	

TABLE 2. Oligonucleotide primers and LightCycler hybridization probes used for LightCycler PCR

^a FL, fluorescein; Red 640, LC Red 640-N-hydroxy-succinimide ester; Red 705, LC Red 705-phosphoramidite; Ph, 3-phosphate.

values for BlnI and SpeI were set using the Auto Algorithm function of the CHEF Mapper XA to separate fragments in the range from 50 to 450 kb (BlnI) or from 30 to 350 kb (SpeI). All fragments larger than 45 kb (up to 27 fragments with XbaI, up to 24 fragments with NotI, up to 25 fragments with BlnI, and up to 29 fragments with SpeI) were included in the clonal analysis of the isolates.

We also analyzed the spatial and temporal behavior of the clonal group of O165:H25 strains in the herd by genomic typing with PFGE (Fig. 1). During a 3-year monitoring program on four cattle farms (19), the O165:H25 clone was detected on only one farm for 4 months. This serotype was not

detected before or after this period, although many other potential EHEC strains belonging to other serotypes were found in this herd (19). Twenty O165:H25 isolates were found in four different cattle on the first date of detection. Different subclusters were already present and were even isolated from a single animal (Table 1). These results suggest that the O165 clone either had been introduced into the farm shortly before the first detection or was the result of recombination due to horizontal gene transfer (8, 18). The occurrence of different subclusters at the same time could have been the result of interactions between different bacteria or between bacteria and the host in the bovine intestine. At later sampling dates the num-

TABLE 3. PCR primers used for detection and characterization of STEC by conventional PCR

TargetPrimerSequence $(5'-3')$ DenaturationAnnealingExtensionNo. of cycles $stxB_{2/2c}$ GK3ATG AAG AAG ATG TTT ATG GK494305260724035 $stxB_{2d}$ VT2-cmAAG AAG ATG TTT GTA GCG G VT2-f94305260724035 $stxB_{2c}$ FK1CCG GAT CCA AGA AGA TG TTT ATA GCG G VT2-f94305560724035 $stxB_{2e}$ FK1CCG GAT CCA AGA AGA TG TTT ATA GCG G VT2-f94305560724035 $stxB_{2e}$ FK1CCG GAT CCA AGA AGA TG TTTA TAG FK294305560724035 $stxB_{2f}$ 128-1AGA TTG GGC GTC ATT CAC TGG TTG 128-294305760726035 $aceae$ SK1CCC GAA TTC GGC ACA AGC ATA AGC LP2943055607212030 $LP2$ CCC GAA TTC GGC ACA AGC ATA AGC LP2STC GCAA TTC GGC ACA AGC ATA AGC943055607212030		Reference
$\frac{1}{\text{Temp}} \xrightarrow{\text{Time}}{\text{Temp}} \xrightarrow{\text{Time}}{\text{Temp}} \xrightarrow{\text{Time}}{\text{Temp}} \xrightarrow{\text{Time}}{\text{CC}} \stackrel{\text{of}}{(s)} \stackrel{\text{of}}{(cc)} \stackrel{\text{of}}{(s)} \stackrel{\text{of}}{(cc)} \stackrel{\text{of}}{(cc)} \stackrel{\text{of}}{(s)} \stackrel{\text{of}}{(cc)} \text{o$	Product length	
$stxB_{2/2c}$ GK3ATG AAG AAG ATG TTT ATG94305260724035 $gK4$ TCA GTC ATT ATT AAA CTG94305260724035 $stxB_{2d}$ VT2-cmAAG AAG ATA TTT GTA GCG G94305260726035 $vT2-f$ TAA ACT GCA CTT CAG CAA AT94305260724035 $stxB_{2c}$ FK1CCG GAT CCA AGA AGA TGT TTA TAG94305560724035 $FK2$ CCC GAA TTC TCA GTT AAA CTT CAC CStxB_{2f}128-1AGA TTG GGC GTC ATT CAC TGG TTG94305760726035 $128-2$ TAC TTT AAT GGC CGC CCT GTC TCC943055607212030 μ -eaeSK1CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 μ -2CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 μ -2CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 μ -2CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 μ -2CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 μ -2CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 μ -2CCC GAA TTC GGC ACA AGC ATA AGC943055	(bp)	
GK4TCA GTC ATT ATT AAA CTG $stxB_{2d}$ VT2-cmAAG AAG ATA TTT GTA GCG G94305260726035 $VT2$ -fTAA ACT GCA CTT CAG CAA AT94305560724035 $stxB_{2e}$ FK1CCG GAT CCA AGA AGA TGT TTA TAG94305560724035 $stxB_{2f}$ 128-1AGA TTG GGC GTC ATT CAC TGG TTG94305760726035 $128-2$ TAC TTT AAT GGC CGC CCT GTC TCC943055607212030 α -eaeSK1CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 $LP2$ CCC GAA TTC GGC ACA AGC ATA AGC943055607212030	260	22
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
VT2-fTAA ACT GCA CTT CAG CAA AT $stxB_{2e}$ FK1CCG GAT CCA AGA AGA TGT TTA TAG FK294305560724035 $stxB_{2f}$ 128-1AGA TTG GGC GTC ATT CAC TGG TTG 128-294305760726035 α -eaeSK1CCC GAA TTC GGC ACA AGC ATA AGC LP2943055607212030 μ <th< td=""><td>256</td><td>36</td></th<>	256	36
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
FK2CCC GAA TTC TCA GTT AAA CTT CAC C $stxB_{2f}$ 128-1AGA TTG GGC GTC ATT CAC TGG TTG94305760726035128-2TAC TTT AAT GGC CGC CCT GTC TCC943055607212030 α -eaeSK1CCC GAA TTC GGC ACA AGC ATA AGC943055607212030LP2CCC GAA TTC GGC ACA AGC ATA AGC943055607212030 α -eaeSK1CCC CAA TTC GGC ACA AGC ATA AGC943055607212030	280	16
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	428	47
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		
LP2 CCC GAA TTC GGC ACA AGC ATA AGC	2,807	31
0_{100} SV1 CCC C A A TTC C C C A C A A C C A T A A C C 04 20 55 60 72 120 20		
p-eae SKI CCC GAA ITC GOC ACA AGC ATA AGC 94 50 55 00 72 120 50	2,287	31
LP4 CCC GTG AT ACCA GTA CCA ATT ACG GTC		
γ -eae SK1 CCC GAA TTC GGC ACA AGC ATA AGC 94 30 45 60 72 120 3	2,792	31
LP3 CCC GAA TTC TTA TTC TAC ACA AAC CGC 94 30 52 60 72 120 28		
δ-eae Int-d TAC GGA TTT TGG GGC AT 95 20 45 60 72 60 30	544	1
Int-Ru TTT ATT TGC AGC CCC CCA T		
ε -eae SK1 CCC GAA TTC GGC ACA AGC ATA AGC 94 30 55 60 72 120 30	2,608	31
LP5 AGC TCA CTC GTA GAT GAC GGC AAG CG		
<i>katP</i> wkat-B CTT CCT GTT CTG ATT CTT CTG G 94 30 56 60 72 150 30	2,125	10
wkat-F AAC TTA TTT CTC GCA TCA TCC		
<i>espP</i> EspA AAA CAG CAG GCA CTT GAA CG 94 30 56 60 72 150 30	1,830	11
EspB GGA GTC GTC AGT CAG TAG AT		
<i>lifA/efa1</i> lifA1 AGC CAT TCC ATC AAT CCG AT 95 30 50 60 72 60 30	532	7
lifA2 TCC CTG CCA AAC TAC CGA CAC		
<i>lifA/efa1</i> lifA3 CAG CTA CAG GAG ACC GTT TTT 95 30 50 60 72 60 30	560	7
lifA4 CAA TAT CAG GCC AAT CAA		



FIG. 1. (a) Dendrogram of DNA divergence (distance) generated by using the Dice coefficient for pairwise comparisons of banding patterns for XbaI, NotI, BlnI, and SpeI restriction of potential EHEC O165:H25 and O165:H- isolates. (b to e) Schematic representations of restriction patterns of bovine and human *E. coli* O165:H25 and O165:H- strains after digestion with XbaI (b), NotI (c), BlnI (d), and SpeI (e). Ellipses indicate the missing XbaI and NotI fragments in strain 02/19/013-7. The *efa1* gene is located at this position in the other bovine strains.

ber of O165:H25 isolates was reduced. The dominant subcluster, subcluster 1, could still be found, but many other subclusters were detected at the same time. The genetic distance between the O165:H25 isolates increased, and in some isolates plasmid-encoded virulence markers or complete virulence plasmids were missing. Variations in the O165:H25 clonal group might have been caused by increasing competition between the bacterial populations of various subtypes in the bovine intestine or by potential interactions between the O165:H25 EHEC and the host. The O165:H25 clonal group finally disappeared from the herd after it had persisted for 4 months. Perhaps the loss of the efa-1 gene in one isolate obtained on the last date when O165:H25 was isolated from the herd can help us understand why the clone disappeared. Efa1 is considered an E. coli factor for colonization of the bovine intestine by non-O157 STEC (51). Also, this O165:H25 strain exhibited the greatest genetic distance compared to the remaining strains in the clonal group (Fig. 1). The distances were calculated from the fragmentation patterns produced by each of the four PFGE enzymes by using the RAPDistance program, version 1.04 (2). The Euclidean distance in three dimensions was not calculated because all O165:H25 strains were isolated from the same farm (i.e., the geographic distance was zero for all isolates). A cluster analysis, using the unweighted pair group method with arithmetic averages, was performed for PFGE by using Statistica 6.1 for Windows (StatSoft Inc., Tulsa, OK). In any case, our results suggest that the persistence of a distinct clone of EHEC may be limited in time and space (i.e., in a cattle herd). This is apparently not a unique property of O165:H25, since we obtained similar results for clonal groups of other EHEC serotypes (O26:H11 and O157:H7) in cattle. The four human O165H25/H- isolates belonged to different other clonal groups (Fig. 1).

In conclusion, bovine O165:H25 isolates can carry virulence factors of EHEC that are associated with EHEC-related disease in humans, particularly HC and HUS. Therefore, strains of bovine origin may represent a considerable potential risk for human infection.

We thank J. Bockemühl, H. Tschäpe, T. Kuzius, and A. Fruth for serotyping the E. coli strains and H. Tschäpe for providing E. coli O165:H25 and O165:H- strains of human origin.

This study was funded by the German Federal Ministry of Consumer Protection, Food and Agriculture.

REFERENCES

- 1. Adu-Bobie, J., G. Frankel, C. Bain, A. G. Goncalves, L. R. Trabulsi, G. Douce, S. Knutton, and G. Dougan. 1998. Detection of intimins alpha, beta, gamma, and delta, four intimin derivatives expressed by attaching and effacing microbial pathogens. J. Clin. Microbiol. 36:662-668.
- 2. Armstrong, J. S., Gibbs, A. J., Peakall, R., and Weiller, G. 1994. The RAP-Distance package, version 1.04. [Online.] http://life.anu.edu.au/molecular /software/rapd.html.
- 3. Barrett, T. J., J. B. Kaper, A. E. Jerse, and I. K. Wachsmuth. 1992. Virulence factors in Shiga-like toxin-producing Escherichia coli isolated from humans and cattle. J. Infect. Dis. 165:979-980.
- 4. Bertin, Y., K. Boukhors, V. Livrelli, and C. Martin. 2004. Localization of the insertion site and pathotype determination of the locus of enterocyte effacement of Shiga toxin-producing Escherichia coli strains. Appl. Environ. Microbiol. 70:61-68.
- 5. Bertin, Y., K. Boukhors, N. Pradel, V. Livrelli, and C. Martin. 2001. Stx2 subtyping of Shiga toxin-producing Escherichia coli isolated from cattle in France: detection of a new Stx2 subtype and correlation with additional virulence factors. J. Clin. Microbiol. 39:3060-3065.
- 6. Boerlin, P., S. A. McEwen, F. Boerlin-Petzold, J. B. Wilson, R. P. Johnson, and C. L. Gyles. 1999. Associations between virulence factors of Shiga

toxin-producing Escherichia coli and disease in humans. J. Clin. Microbiol. 37:497-503

- 7. Böhnke, U. 2002. Untersuchung zur Lokalisation und Regulation des als "lymphocyte inhibitory factor" (lifA) oder "EHEC factor for adherence" (efa1) bezeichneten Gens des bovinen enterohämorrhagischen E. coli-Stammes RW1374 (O103:H2). Diplomarbeit. Freie Universität Berlin, Berlin, Germany.
- 8. Brunder, W., and H. Karch. 2000. Genome plasticity in Enterobacteriaceae. Int. J. Med. Microbiol. 290:153-165.
- 9. Brunder, W., H. Schmidt, M. Frosch, and H. Karch. 1999. The large plasmids of Shiga-toxin-producing Escherichia coli (STEC) are highly variable genetic elements. Microbiology 145:1005-1014.
- 10. Brunder, W., H. Schmidt, and H. Karch. 1996. KatP, a novel catalaseperoxidase encoded by the large plasmid of enterohaemorrhagic Escherichia coli O157:H7. Microbiology 142:3305-3315.
- 11. Brunder, W., H. Schmidt, and H. Karch. 1997. EspP, a novel extracellular serine protease of enterohaemorrhagic Escherichia coli O157:H7, cleaves human coagulation factor V. Mol. Microbiol. 24:767-778.
- 12. China, B., F. Goffaux, V. Pirson, and J. Mainil. 1999. Comparison of eae, tir, espA and espB genes of bovine and human attaching and effacing Escherichia coli by multiplex polymerase chain reaction. FEMS Microbiol. Lett. 178:177-182.
- 13. China, B., E. Jacquemin, A. C. Devrin, V. Pirson, and J. Mainil. 1999. Heterogeneity of the eae genes in attaching/effacing Escherichia coli from cattle: comparison with human strains. Res. Microbiol. 150:323-332.
- 14. De Baets, L., D. T. Van, I., M. De Filette, D. Pierard, L. Allison, H. De Greve, J. P. Hernalsteens, and H. Imberechts. 2004. Genetic typing of Shiga toxin 2 variants of Escherichia coli by PCR-restriction fragment length polymorphism analysis. Appl. Environ. Microbiol. 70:6309-6314.
- 15. Ethelberg, S., K. E. Olsen, F. Scheutz, C. Jensen, P. Schiellerup, J. Enberg, A. M. Petersen, B. Olesen, P. Gerner-Smidt, and K. Molbak. 2004. Virulence factors for hemolytic uremic syndrome, Denmark. Emerg. Infect. Dis. 10: 842-847
- 16. Franke, S., D. Harmsen, A. Caprioli, D. Pierard, L. H. Wieler, and H. Karch. 1995. Clonal relatedness of Shiga-like toxin-producing Escherichia coli O101 strains of human and porcine origin. J. Clin. Microbiol. 33:3174-3178.
- 17. Friedrich, A. W., M. Bielaszewska, W. L. Zhang, M. Pulz, T. Kuczius, A. Ammon, and H. Karch. 2002. Escherichia coli harboring Shiga toxin 2 gene variants: frequency and association with clinical symptoms. J. Infect. Dis. 185:74-84.
- 18. Fruth, A., R. Prager, A. Friedrich, T. Kuczius, P. Roggentin, H. Karch, A. Ammon, J. Bockemühl, and H. Tschäpe. 2002. Infektionen des Menschen durch enterohämorrhagische Escherichia coli (EHEC) in der Bundesrepublik Deutschland von 1998 bis 2001-Prävalenz und Typenspektrum. Bundesgesundheitsblatt Gesundheitsforsch. Gesundheitsschutz 45:715-721.
- 19. Geue, L., M. Segura-Alvarez, F. J. Conraths, T. Kuczius, J. Bockemuhl, H. Karch, and P. Gallien. 2002. A long-term study on the prevalence of shiga toxin-producing Escherichia coli (STEC) on four German cattle farms. Epidemiol. Infect. 129:173-185.
- 20. Griffin, P. M., and R. V. Tauxe. 1991. The epidemiology of infections caused by Escherichia coli O157:H7, other enterohemorrhagic E. coli, and the associated hemolytic uremic syndrome. Epidemiol. Rev. 13:60-98.
- 21. Jores, J., K. Zehmke, J. Eichberg, L. Rumer, and L. H. Wieler. 2003. Description of a novel intimin variant (type zeta) in the bovine O84:NM verotoxin-producing Escherichia coli strain 537/89 and the diagnostic value of intimin typing. Exp. Biol. Med. 228:370-376.
- 22. Karch, H., H. I. Huppertz, J. Bockemühl, H. Schmidt, A. Schwarzkopf, and R. Lissner. 1997. Shiga toxin-producing Escherichia coli infections in Germany. J. Food Prot. 11:1454-1457.
- 23. Karmali, M. A. 1989. Infection by verocytotoxin-producing Escherichia coli. Clin. Microbiol. Rev. 2:15-38.
- 24. Keskimaki, M., M. Saari, T. Heiskanen, and A. Siitonen. 1998. Shiga toxinproducing Escherichia coli in Finland from 1990 through 1997: prevalence and characteristics of isolates. J. Clin. Microbiol. 36:3641-3646.
- 25. Makino, K., K. Yokoyama, Y. Kubota, C. H. Yutsudo, S. Kimura, K. Kurokawa, K. Ishii, M. Hattori, I. Tatsuno, H. Abe, T. Iida, K. Yamamoto, M. Onishi, T. Hayashi, T. Yasunaga, T. Honda, C. Sasakawa, and H. Shinagawa. 1999. Complete nucleotide sequence of the prophage VT2-Sakai carrying the verotoxin 2 genes of the enterohemorrhagic Escherichia coli O157:H7 derived from the Sakai outbreak. Genes Genet. Syst. 74:227-239.
- 26. McKee, M. L., A. R. Melton-Celsa, R. A. Moxley, D. H. Francis, and A. D. O'Brien. 1995. Enterohemorrhagic Escherichia coli O157:H7 requires intimin to colonize the gnotobiotic pig intestine and to adhere to HEp-2 cells. Infect. Immun. 63:3739-3744.
- 27. Muniesa, M., M. de Simon, G. Prats, D. Ferrer, H. Panella, and J. Jofre. 2003. Shiga toxin 2-converting bacteriophages associated with clonal variability in Escherichia coli O157:H7 strains of human origin isolated from a single outbreak. Infect. Immun. 71:4554-4562.
- Nakao, H., K. Kimura, H. Murakami, T. Maruyama, and T. Takeda. 2002. Subtyping of Shiga toxin 2 variants in human-derived Shiga toxin-producing Escherichia coli strains isolated in Japan. FEMS Immunol. Med. Microbiol. **34:**289–297.

- Nataro, J. P., and J. B. Kaper. 1998. Diarrheagenic *Escherichia coli*. Clin. Microbiol. Rev. 11:142–201.
- Nicholls, L., T. H. Grant, and R. M. Robins-Browne. 2000. Identification of a novel genetic locus that is required for in vitro adhesion of a clinical isolate of enterohaemorrhagic *Escherichia coli* to epithelial cells. Mol. Microbiol. 35:275–288.
- Oswald, E., H. Schmidt, S. Morabito, H. Karch, O. Marches, and A. Caprioli. 2000. Typing of intimin genes in human and animal enterohemorrhagic and enteropathogenic *Escherichia coli*: characterization of a new intimin variant. Infect. Immun. 68:64–71.
- Paton, A. W., and J. C. Paton. 1998. Detection and characterization of Shiga toxigenic *Escherichia coli* by using multiplex PCR assays for *stx*₁, *stx*₂, *eaeA*, enterohemorrhagic *E. coli hlyA*, *rfb*_{O111}, and *rfb*_{O157}. J. Clin. Microbiol. 36:598–602.
- 33. Paton, A. W., J. C. Paton, P. N. Goldwater, and P. A. Manning. 1993. Direct detection of *Escherichia coli* Shiga-like toxin genes in primary fecal cultures by polymerase chain reaction. J. Clin. Microbiol. 31:3063–3067.
- Paton, J. C., and A. W. Paton. 1998. Pathogenesis and diagnosis of Shiga toxin-producing *Escherichia coli* infections. Clin. Microbiol. Rev. 11:450–479.
- 35. Perna, N. T., G. Plunkett III, V. Burland, B. Mau, J. D. Glasner, D. J. Rose, G. F. Mayhew, P. S. Evans, J. Gregor, H. A. Kirkpatrick, G. Posfai, J. Hackett, S. Klink, A. Boutin, Y. Shao, L. Miller, E. J. Grotbeck, N. W. Davis, A. Lim, E. T. Dimalanta, K. D. Potamousis, J. Apodaca, T. S. Anantharaman, J. Lin, G. Yen, D. C. Schwartz, R. A. Welch, and F. R. Blattner. 2001. Genome sequence of enterohaemorrhagic *Escherichia coli* O157:H7. Nature 409:529–533.
- Pierard, D., G. Muyldermans, L. Moriau, D. Stevens, and S. Lauwers. 1998. Identification of new verocytotoxin type 2 variant B-subunit genes in human and animal *Escherichia coli* isolates. J. Clin. Microbiol. 36:3317–3322.
- Plunkett, G., III, D. J. Rose, T. J. Durfee, and F. R. Blattner. 1999. Sequence of Shiga toxin 2 phage 933W from *Escherichia coli* O157:H7: Shiga toxin as a phage late-gene product. J. Bacteriol. 181:1767–1778.
- Ramachandran, V., M. A. Hornitzky, K. A. Bettelheim, M. J. Walker, and S. P. Djordjevic. 2001. The common ovine Shiga toxin 2-containing *Escherichia coli* serotypes and human isolates of the same serotypes possess a Stx2d toxin type. J. Clin. Microbiol. 39:1932–1937.
- Reid, S. D., C. J. Herbelin, A. C. Bumbaugh, R. K. Selander, and T. S. Whittam. 2000. Parallel evolution of virulence in pathogenic *Escherichia coli*. Nature 406:64–67.
- Reischl, U., M. T. Youssef, J. Kilwinski, N. Lehn, W. L. Zhang, H. Karch, and N. A. Strockbine. 2002. Real-time fluorescence PCR assays for detection and characterization of Shiga toxin, intimin, and enterohemolysin genes from Shiga toxin-producing *Escherichia coli*. J. Clin. Microbiol. 40:2555– 2565.
- Remuzzi, G., and P. Ruggenenti. 1998. The hemolytic uremic syndrome. Kidney Int. Suppl. 66:S54–S57.
- Ritchie, J. M., P. L. Wagner, D. W. Acheson, and M. K. Waldor. 2003. Comparison of Shiga toxin production by hemolytic-uremic syndrome-asso-

ciated and bovine-associated Shiga toxin-producing *Escherichia coli* isolates. Appl. Environ. Microbiol. **69**:1059–1066.

- Russmann, H., E. Kothe, H. Schmidt, S. Franke, D. Harmsen, A. Caprioli, and H. Karch. 1995. Genotyping of Shiga-like toxin genes in non-O157 *Escherichia coli* strains associated with haemolytic uraemic syndrome. J. Med. Microbiol. 42:404–410.
- Russmann, H., H. Schmidt, A. Caprioli, and H. Karch. 1994. Highly conserved B-subunit genes of Shiga-like toxin II variants found in *Escherichia coli* O157 strains. FEMS Microbiol. Lett. 118:335–340.
- Schmidt, H., L. Beutin, and H. Karch. 1995. Molecular analysis of the plasmid-encoded hemolysin of *Escherichia coli* O157:H7 strain EDL 933. Infect. Immun. 63:1055–1061.
- 46. Schmidt, H., C. Kernbach, and H. Karch. 1996. Analysis of the EHEC *hly* operon and its location in the physical map of the large plasmid of enterohaemorrhagic *Escherichia coli* O157:H7. Microbiology 142:907–914.
- Schmidt, H., J. Scheef, S. Morabito, A. Caprioli, L. H. Wieler, and H. Karch. 2000. A new Shiga toxin 2 variant (Stx2f) from *Escherichia coli* isolated from pigeons. Appl. Environ. Microbiol. 66:1205–1208.
- Schmitt, C. K., M. L. McKee, and A. D. O'Brien. 1991. Two copies of Shiga-like toxin II-related genes common in enterohemorrhagic *Escherichia coli* strains are responsible for the antigenic heterogeneity of the O157:Hstrain E32511. Infect. Immun. 59:1065–1073.
- Segura-Alvarez, M., H. Richter, F. J. Conraths, and L. Geue. 2003. Evaluation of enzyme-linked immunosorbent assays and a PCR test for detection of Shiga toxins for Shiga toxin-producing *Escherichia coli* in cattle herds. J. Clin. Microbiol. 41:5760–5763.
- Stephan, R., and L. E. Hoelzle. 2000. Characterization of Shiga toxin type 2 variant B-subunit in *Escherichia coli* strains from asymptomatic human carriers by PCR-RFLP. Lett. Appl. Microbiol. 31:139–142.
- Stevens, M. P., P. M. van Diemen, G. Frankel, A. D. Phillips, and T. S. Wallis. 2002. Efa1 influences colonization of the bovine intestine by Shiga toxin-producing *Escherichia coli* serotypes O5 and O111. Infect. Immun. 70:5158–5166.
- Su, C., and L. J. Brandt. 1995. Escherichia coli O157:H7 infection in humans. Ann. Intern. Med. 123:698–714.
- Tesh, V. L., and A. D. O'Brien. 1991. The pathogenic mechanisms of Shiga toxin and the Shiga-like toxins. Mol. Microbiol. 5:1817–1822.
- 54. Tschäpe, H. 2003. Personal communication.
- 55. Vernozy-Rozand, C., M. P. Montet, Y. Bertin, F. Trably, J. P. Girardeau, C. Martin, V. Livrelli, and L. Beutin. 2004. Scrotyping, stx₂ subtyping, and characterization of the locus of enterocyte effacement island of Shiga toxin-producing *Escherichia coli* and *E. coli* O157:H7 strains isolated from the environment in France. Appl. Environ. Microbiol. **70**:2556–2559.
- 56. Weinstein, D. L., M. P. Jackson, J. E. Samuel, R. K. Holmes, and A. D. O'Brien. 1988. Cloning and sequencing of a Shiga-like toxin type II variant from *Escherichia coli* strain responsible for edema disease of swine. J. Bacteriol. 170:4223–4230.