Development and Characterization of *recA* Mutants of *Campylobacter jejuni* for Inclusion in Attenuated Vaccines

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Isogenic recA mutants of Campylobacter jejuni have been constructed for evaluation of their usefulness in attenuated vaccines against this major worldwide cause of diarrhea. The recA⁺ gene of C. jejuni 81-176 was cloned by using degenerate primers to conserved regions of other RecA proteins in a PCR. The C. jejuni recA⁺ gene encodes a predicted protein with an M_r of 37,012 with high sequence similarity to other RecA proteins. The termination codon of the recA⁺ gene overlaps with the initiation codon of another open reading frame which encodes a predicted protein which has >50% identity with the N terminus of the Escherichia coli enolase protein. A kanamycin resistance gene was inserted into the cloned recA⁺ gene in E. coli and returned to C. jejuni VC83 by natural transformation, resulting in allelic replacement of the wild-type recA gene. The resulting VC83 recA mutant displayed increased sensitivity to UV light and a defect in generalized recombination as determined by natural transformation frequencies. The mutated recA gene was amplified from VC83 recA by PCR, and the product was used to transfer the mutation by natural transformation into C. jejuni 81-176 and 81-116, resulting in isogenic recA mutants with phenotypes similar to VC83 recA. After oral feeding, strain 81-176 recA colonized rabbits at levels comparable to wild-type 81-176 and was capable of eliciting the same degree of protection as wild-type 81-176 against subsequent homologous challenge in the RITARD (removable intestinal tie adult rabbit diarrhea) model.

Campylobacter spp., particularly Campylobacter jejuni, are a major cause of enteric disease worldwide. Acute symptoms most often include dysentery, fever, and abdominal pain (5). Sequelae can include colitis, reactive arthritis, and Guillain-Barré syndrome (5, 36). An estimated 400 million cases occur annually, with 2.4 million occurring in the United States (48). Campylobacters have recently been recognized as a significant cause of travelers diarrhea, with attack rates approaching 25 to 37% per year among individuals traveling to highly endemic areas (15, 34, 50). Outbreaks and sporadic cases of Campylobacter gastroenteritis are also well documented among U.S. military personnel (6, 7, 14, 50, 52). While the need for a vaccine against Campylobacter spp. for the general population in developed nations is arguable, the emergence of these organisms as a major cause of diarrhea has led to interest in development of a vaccine for selected high-risk populations.

Vaccine strategies against *Campylobacter* spp. to date have focused largely on identification of potential protective antigens for inclusion into carrier vaccine strains (42). One of the most promising of such antigens is flagellin, the subunit of the flagella filament. Although development of antibodies against flagellin seems to correlate with acquisition of immunity to infection (32), it remains to be shown that flagellin is protective. Moreover, since a nonflagellated mutant strain (22) was able to fully protect against subsequent challenge in a rabbit model (40), other antigens are capable of eliciting protective antigen, there exist other problems in its inclusion in a carrier vaccine, including antigenic diversity among serotypes (30), phase and antigenic variability (11, 25), and the presence of posttranslational modifications on the surface (2, 31). Similar uncertainties exist for other individual candidate antigens (42).

An alternative approach to enteric vaccines is the development of living attenuated strains expressing the full complement of native surface antigens, such as *Salmonella typhi* Ty21a (17) and several living candidate vaccine strains of *Vibrio cholerae* (29, 51). Some of the *V. cholerae* candidates have included a *recA* mutation for two reasons. First, amplification of cholera toxin genes, a process which enhances virulence, has been shown to be *recA*-dependent (18), and second, the presence of a *recA* allele would reduce concerns about reversion of virulence defects in vaccine strains once introduced into endemic areas (26, 51). Furthermore, it has recently been shown that *recA* mutants of *Salmonella typhimurium* are avirulent (8).

Campylobacter spp. are unique among enteric pathogens in that they are naturally transformable (53). This ability to take up and incorporate exogenous DNA may increase the possibility of reversion of attenuated strains of C. jejuni in the environment. One way of precluding this problem, and at the same time achieving some degree of attenuation, would be the incorporation of recA mutations into vaccine strains. In this study, we report the cloning, sequencing, and site-specific mutagenesis of the recA⁺ gene of C. jejuni and demonstrate that a C. jejuni recA mutant is still capable of colonizing and eliciting protection in a rabbit model. In addition, we have also demonstrated that natural transformation can be exploited to generate primary mutations in C. jejuni without the need for conjugative transfer of suicide vectors from Escherichia coli and used a combination of PCR and natural transformation to transfer the isogenic recA mutation among other strains of C. jejuni.

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MATERIALS AND METHODS

Bacterial strains and growth conditions. C. jejuni 81-176 is a clinical isolate which has been described previously (4). C. jejuni 81-116 was the gracious gift of Diane Newell and has been described previously (38). C. jejuni VC83 is a clinical isolate from Canada (1). E. coli DH5 α was used as the host for molecular cloning experiments. C. jejuni was routinely cultivated on Mueller-Hinton (MH) agar supplemented when indicated with kanamycin (50 µg/ml) at 37°C in an atmosphere of 10% CO₂-85% N₂-5% O₂. E. coli was grown on Luria agar (45) supplemented as indicated with ampicillin (50 µg/ml) or kanamycin (50 µg/ml).

Recombinant DNA techniques and plasmids. Restriction enzymes and T4 DNA ligase were purchased from Boehringer Mannheim (Indianapolis, Ind.) and used as recommended by the supplier. pUC18 (GIBCO, Gaithersburg, Md.) was used for cloning.

PCR conditions. PCRs were run in a Perkin-Elmer Cetus thermal cycler with reagents supplied by Perkin-Elmer Cetus (Norwalk, Conn.) at the concentrations recommended by the supplier. The conditions for primers PMP1 and REV3 were 40 cycles of denaturation at 94°C for 1 min, annealling at 22°C for 1 min, and extension at 72°C for 1 min. Conditions for primers Rec1 and Rec2 were 30 cycles of denaturation at 94°C for 1 min, and extension at 72°C for 2 min, annealling at 42°C for 1 min, and extension at 72°C for 2 min.

DNA hybridizations. DNA restriction fragments separated in agarose gels were transferred to nitrocellulose sheets as described by Sambrook et al. (45) and hybridized under conditions previously described (23). Probes were labelled with $[\alpha$ -³²P]dCTP by nick translation performed with a commercial kit (DuPont-New England Nuclear, Wilmington, Del.).

DNA sequencing. DNA sequencing reactions were performed on double-stranded plasmid templates with commercially available dideoxy-terminator cycle sequencing kits from Applied Biosystems, Inc. (Foster City, Calif.). Sequencing reactions were run on an Applied Biosystems model 373 automated DNA sequencer. Custom oligonucleotide primers were synthesized on an Applied Biosystems model 392 DNA synthesizer. Sequences were analyzed by using MacVector and Assemblylign software packages (IBI, New Haven, Conn.).

Natural transformation. A slight modification of the biphasic method of natural transformation of *C. jejuni* in which the biphasic cultures are incubated in 5% CO₂–95% air instead of air was used (1, 21, 53). *C. jejuni* cells were harvested from overnight MH plates and resuspended in MH broth to an approximate optical density at 600 nm of 0.5. Two-hundredfifty-microliter aliquots were dispensed into polypropylene tubes (12 by 75 mm) containing 1.5 ml of MH agar. The tubes were incubated in an atmosphere of 5% CO₂–95% air for 2 h at 37°C, at which time 1 to 2 µg of DNA was added. After 4 to 5 h of incubation at 37°C in 5% CO₂–95% air, the cells were plated to selective media and grown in an atmosphere of 10% CO₂–85% N₂–5% O₂.

UV sensitivity measurements. Cells were harvested after 18 h of growth on MH agar and resuspended in 0.1 M MgSO₄ to an optical density at 600 nm of approximately 0.5. The cells were transferred to a petri dish and exposed to UV light in a UV cross-linker (Hoeffer Scientific Instruments, San Francisco, Calif.). After increasing doses of UV exposure, aliquots were removed, serially diluted in MH broth, and plated to MH agar.

Rabbit experiments. Female New Zealand White rabbits (Hazelton Research Products, Denver, Pa.), weighing 1.0 to 2.0 kg, were fed either strain 81-176, strain 81-176 *recA* mutant, or

sterile broth via a nasogastric tube after neutralization of gastric acidity as described previously (10, 40). One month later, all three groups were challenged by the removable intestinal tie adult rabbit diarrhea (RITARD) procedure (10, 40) with strain 81-176.

Frozen stocks of 81-176 and 81-176 *recA* were thawed and inoculated on MH or MH supplemented with kanamycin, respectively, and incubated at 37°C microaerobically. After 18 h of incubation, cells were suspended in brucella broth to a concentration giving an optical density at 625 nm of approximately 0.1. Eight milliliters of the suspension was overlaid in 25-cm² tissue culture flasks (Becton Dickinson, Oxnard, Calif.) containing 5 ml of brucella blood agar, and these biphasic cultures were incubated ungassed at 37°C (10, 40). After 10 h of incubation, the broth phase of the biphasic culture system was collected and used for animal challenge. All challenge doses were monitored by plate counts. Oral feeding doses were approximately 10^{10} per animal. For the RITARD challenge, the pooled culture was diluted 10^4 -fold in brucella broth, resulting in a challenge dose of approximately 10^6 per animal.

After oral and RITARD challenge, the colonization of rabbits was monitored by daily rectal swabs. The swabs were cultured for *C. jejuni* by direct plating on *Campylobacter*-selective agar (Remel, Lenexa, Kans.). *Campylobacter* spp. were identified by a positive oxidase test (Oxidase Reagent Droppers; Marion Scientific, Kansas City, Kans.) and by microscopic examination. Presumed 81-176 *recA* colonies were confirmed by testing for kanamycin resistance (Km^r). An animal was considered to be no longer colonized by *Campylobacter* spp. after 3 consecutive days of negative cultures.

Experiments reported herein were conducted according to the principles set forth in the *Guide for the Care and Use of Laboratory Animals* (Institute of Laboratory Animal Resources, National Research Council, U.S. Department of Health and Human Services).

Nucleotide sequence accession number. The DNA sequence shown in Fig. 2 has been deposited with GenBank under accession number UO3121.

RESULTS

Cloning and sequence analysis of the $recA^+$ gene of C. jejuni. Two degenerate PCR primers were designed on the basis of the known G+C bias of Campylobacter spp. which would encode two highly conserved regions of RecA proteins from several species (3, 16, 27, 33, 37, 39, 43, 46, 47). These primers were PMP1 (GAAAT[AT]TATGGTCCTGA) and REV3 (TT CACC[AT]GTATC[AT]GGTT). PMP1 encodes amino acids EIYGP (Fig. 3, residues 62 to 66) which are highly conserved in RecA proteins. The inverse of REV3 encodes amino acids PDTGE corresponding to residues 118 to 122, another highly conserved region. When these primers were used in a PCR with strain 81-176 DNA, they produced a product of the predicted 180-bp size. This product was cloned into pUC18, and the insert was sequenced by using forward and reverse primers. The DNA sequence encoded a predicted protein with 70% sequence identity with the RecA protein of E. coli within this region, as can be seen in Fig. 3. This PCR product was labelled with ³²P by nick translation and used as a probe to clone a 2.7-kb HindIII fragment into pUC18. This plasmid was termed pPMP100. Sequence analysis of the insert in pPMP100 indicated that it lacked the final 153 bp of the RecA open reading frame (ORF1; see below). pPMP100 was used as a probe to clone an overlapping 1.8-kb XbaI-PstI fragment, and this plasmid was termed pPMP101. The plasmids are shown diagrammatically in Fig. 1.



FIG. 1. Restriction map of the $recA^+$ region. The top line represents the chromosome of strain 81-176. The position of the inserts in plasmids pPMP100 and pPMP101 are indicated. Restriction site abbreviations: B, *Bgl*II; C, *Cla*I; H, *Hind*III; R1, *Eco*RI; RV, *Eco*RV; P, *PsI*; X, *Xba*I. The kanamycin resistance cassette was inserted into the *Cla*I site on pPMP100 to generate the plasmid pPMP102 which was used to generate the *recA* mutation.

The DNA sequence of the $recA^+$ information on pPMP100 and pPMP101 is shown in Fig. 2. There is an ORF (ORF1) of 1,029 nucleotides encoding a predicted protein of 343 amino acids with a predicted M_r of 37,012. The G+C content within the coding region is 37%. There is a putative ribosome binding site (shown by overlining in Fig. 2) 5 bp upstream from the translational start of the ORF. The protein encoded by ORF1 shows significant sequence identity with other RecA-like proteins, as shown in Fig. 3, with the best match (66% identity overall) being to the $recA^+$ gene of *Neisseria gonorrheae* (16).

A second ORF begins in the sequence at the second A of the TAA termination codon of the $recA^+$ gene, as shown in Fig. 2. The predicted N terminus of this ORF shows sequence similarity with the N-terminal sequences of enolases from several sources. The predicted protein is 53% identical to the N terminus of enolase of *E. coli* (MSKIVKIIGREIIDSRGNP TVEAEVHLEGGFVG [55]) and 47% identical to rat enolase (MSIQKIWAREILDSRGNPTVEVDLHTAKG) [44]). A putative ribosome binding site is found 8 bp upstream from the translational initiation site of ORF2 and within the $recA^+$ gene.

Generation of a primary recA mutation in C. jejuni. Initial strategies to generate a site-specific mutation in the $recA^+$ gene utilized standard methods of shuttle mutagenesis (20, 22, 28). Several plasmids were constructed in which a Km^r cassette (28) was inserted into either the ClaI site or the unique BglII site in pPMP100 (Fig. 1). The inserts were transferred into the suicide vector pGK2003 (22) and mobilized from DH5 (RK212.2) hosts into strain 81-176 selecting for Km^r. Transconjugants were screened by hybridization to pUC18 DNA to determine whether the suicide vector had integrated into the chromosome by a single crossover event. All transconjugants examined from numerous crosses contained vector DNA. In an effort to circumvent this unexpected result, we introduced the mutated recA allele into C. jejuni by natural transformation. Plasmid pPMP102 is the construction in which the Km^r cassette was inserted into the ClaI site in pPMP100 (Fig. 1). To increase the likelihood of a double crossover event, this plasmid was linearized with BamHI and used to transform 81-176 and VC83 by the biphasic method described by Wang and Taylor (53). VC83 is a strain of C. jejuni which has been shown previously to be highly transformable (1, 21). No transformants were obtained from 81-176, but a single VC83 transformant was obtained, and colony blot hybridization with the pUC18 probe indicated that no vector sequences were present in this transformant (data not shown). DNA was

GGAGAATGTG CAATAGTAAT TTTTTTCATC TATGCTTCCT TGTAAATCTT TGCTTTAATT 60 CTAGCAAAAAT AAATCATTAT TTAATTITAT TITGTTAGAA TTCTAGCTAA AAATTTAGAA 120 TTTTTAAGGA AAGTT ATG GAT GAT AAT AAA AGA AAA TCT CTA GAC GCT GCC D D N K R K S L D A A 170 12 215 28 CTA AAA AGT TTA GAT AAA ACC TTT GGA AAA GGC ACT ATT TTA AGA CTA L K S L D K T F G K G T I L R L 265 44 GGG GAT AAA GAA GTT GAG CAA ATC GAT AGC ATA GGC ACA GGT G D K E V E Q I D S I G T G 315 60 GGG CTT GAT CTT GCT TTA GGT ATA GGC GGT GTT CCA AAA GGA AGA ATT G I D L A L G I G G V P K G R I 360 76 GGA CCT GAA AGT TCA GGT AAA ACC ACT CTA ACT TTA 410 92 455 108 GAT GCA GAA CAT GCG CTT GAT GTA AAA TAT GCT AAA D A F H A I D V K Y A K 505 124 AAT ACA GAT GAT TTG TAT GTT TCT CAG CCT GAT N T D D L Y V S Q P D 555 140 TTA GAA ATT GTA GAA ACT ATA GCA AGA AGT GGC GCA GTA GAT CTT ATC L E I V E T I A R S G A V D L I 600 156 GTA GTA GAT AGC GTT P 650 172 CAA CAT 0 H GTA GGA CTT CAA GTT CAT AAA 695 188 ATT TTT ATC AAC CAA ATT CGT ATG AAA ATC GGT GCT ATG GGT TAT GGC I F I N Q I R M K I G A M G Y G 745 204 ACT CCT GAA ACT ACA ACA GGT GGA AAT GCA TTA AAA ATC TAT GCT TCT T P E T T T G G N A L K I Y A S 795 220 GTG CGT TTA GAT GTT AGA AAA GTA GCA ACC TTA AAA CAA AAC GAA GAA V R L D V R K V A T L K O N E E 840 236 890 252 CCT ATA GGA AAC CGC GTT AAA GTA AAA GTA GTA GTT AAA AAT AAA GTT GCT P I G N R V K V K V V K N K V A 935 268 GCT GAA TTT GAT E F D GTG ATG TTT V M F AGC CGT GAA GGT GAA TTG ATC GAT TAT GGT GTA AAA S R E G E L I D Y G V K 985 284 1035 300 GAT AAA AGT GGT GCG D K S G A TGG TTT TCT TAT AAA W F S Y K 1080 316 GGT AGA GAA AAT TCA AAA GCT TTC TTA AAA G R F N S K A F L K 1128 332 AGC GAA GAT GAC GAA GGA GAA GAA TA A TG TTA GTA ATT S E D D E G E E> M L V T 1210 343/4 GAA GAT GTT AG<u>A GCC TAT GAA GTT CTT GAT AGT AGA GGA</u> E D V R A Y E V L D S R G 1258 20 AAT CCA ACC 1300 GTA AAA GCC GAA GTT ACG CTA AGC GAT GGA AGT GTA GGT GCG V K A E V T L S D G S V G A

FIG. 2. DNA sequence of the $recA^+$ gene and flanking DNA. The predicted amino acid residues are shown in the single-letter code. The numbers on the right refer to the nucleotide numbers (top) and predicted amino acid residues (bottom). ORF1 marks the beginning of the $recA^+$ gene, and ORF2 marks the beginning of the putative enolase gene. Putative ribosome binding sites are indicated by overlining. The positions of the 2 PCR primers, Rec1 and Rec2, are shown by underlining.

purified from this transformant and characterized by Southern blot hybridization to pPMP101. The results, shown in Fig. 4, indicate that when digested with XbaI and PstI, the mutant recA gene has increased in molecular mass by 1.4 kb (the size of the Km^r cassette; lane B) to 3.2 kb compared with the corresponding 1.8-kb fragment from wild-type VC83 (lane A). When the Km^r cassette was used as the probe, it hybridized to the same 3.2-kb fragment in the mutant (data not shown) but did not hybridize to the wild type. UV kill curves on VC83 and VC83 recA confirmed that the mutant was more sensitive to UV damage than the parent (Fig. 5), and subsequent DNA

		10		20		30		40		50		60	
Ci	MDDNK	RKSLD	AALKS	LDKTF	GKGTI	LRLGD	KEVEO	IDSIG	TGSVG	LDLAL	GIGGV	PKGRI	
Ng	MsDdK	sKaLa	AALaq	ieKsF	GKGaI	mkmGs	qqeEn	levIs	TGSLG	LDLAL	GvGGl	rrGRI	
Mf	MDeNr	sKaLa	AALsq	ieKqF	GKGsI	mRmGD	tdVad	Iqavs	TGS1G	LDIAL	GIGGI	PrGRI	
Pa Tf	MDear	skale	AALGO	iDKaF	GKGav	mR/mGD	nerqa	Ipais	TGSIG	LDLAL	GIGGI	PKGRI	
Ec	iDeNK	aKaLa	AALaa	ieKaF	GKGsI	mRLGe	drsmn	vetIs	TGS1s	LDVAL	GaGGI	PrGRT	
				•••••									
		70		80		90		100		110		120	
				*				*		*		*	
ĊÌ	IEIYO	PESSO	S KTTLI	LHII	ECOK/	GGVC	FIDAE	HALD	KYAK	LGVNT	DDLYN	SQPDF	•
Ng	VEIFG	PESSG	KTTLC	LeavA	qCQKn	GGVCA	FVDAE	HATDp	VYArk	LGVkv	eeLYL	SOPDE	
Pa	VEING	PESSG	KTTLT	LSVIA	FaOKa	GatCA	FUDAE	HALDO	dYAck			SOPO+	
Tf	VEIYG	PESSG	KTTLT	LHaIA	sCQaA	GGtaA	FIDAE	HALDP	gYAhk	LGVdl	enLli	SOPDE	
Ec	VEIYG	PESSG	KTTLT	LqvIA	aaQre	GktCA	FIDAE	HALDP	iYArk	LGVdi	DnLlc	SQPDt	
		130		140		150		160		170		180	
C -1		*	TADC C	*	NAUDCH	*		*	-	*		*	
Na	GEOAL	ELVET	LARSG 1VRSG	aiDev	VVDSV	AALIP	KAELE	GDMGD	CHVGL		SQALK	KLIGI	
Mf	GEOAL	Eladm	LVRSG	sVDvv	VVDSV	AALTP	KAEIE	GeMGD	shingL	OARLM	SOALR	KLTan	
Pa	GEQAL	EItdm	lvRSn	AVDvI	iVDSV	AALVP	KAEIE	GeMGD	aHVGL	QARLM	SQALR	KiTGn	
Tf	GEQAL	Eladm	1vRSG	AVDLI	ViDSV	AALTP	KAEIE	GeMGD	sHVGk	QARLM	SQALR	nLTan	
EC	GEQAL	FICGO	LARSG	AVDV1	VVDSV	AALTP	KAEIE	Ge1GD	shmGL	aARmM	SQAmR	KLaGn	
		190		200		210		220		230		240	
Cj	VHKMN	TTVIF	INQIR	MKIGA	MGYGT	PETTT	GGNAL	KIYAS	VRLDV	RKVAT	LKONE	EPIGN	
Ng	ikKtN	TlVvF	INQIR	MKIGV	M-fGs	PETTT	GGNAL	KfYsS	VRLDi	Rrtgs	iKkgE	E∨lGN	
Mf	ikrtN	TIVIF	INQIR	MKIGL	M-fGn	PETT	GGNAL	KEYAS	VRLDi	Rrtga	iKkgd	EvtGs	
Tf	isrsN	TIVIE	TNOTR	MKTGV	M-YGs	PETTT	GGNAL	KEYAS	VRIDI	Rriga	iKked	EVVUS	
Ec	lkqsN	TILIF	INQIR	MKIGV	M-fGn	PETTT	GGNAL	KFYAS	VRLDi	Rriga	vKegE	nvvGs	
		250		260		270		280		290		300	
. .		*		*						•		•	
	RVKVK	VYKNK	VAPPE	RQAEF	DVMFG	EGLSR	EGELI	DYGVK	LDIVD	KSGAW	FSYKD	KKLGQ	
Mf	etrVK	VVKNn	VAPPE	klAFF	DilvG	FGiSR	FGFiT	elQVn	lklie	KAGAW	VSYKA	aKiGO	
Pa	etrVK	VVKNK	VsPPF	ROAEF	qilyG	kGiyR	tGEiI	DLGVa	LalVe	KSGAW	vSYaa	sKiGO	
Tf	dtrVK	VVKNK	VAPPF	ReAEF	aiyyG	EGISR	lsELv	D ເຊນ ເ	fĎIVe	KSGAW	ySYqg	hriGQ	
Ec	etrVK	VVKNK	iAaPF	kQAEF	qilyG	EGinf	yGELv	DIGVK	eklie	KaGAW	ySYKg	eKiGQ	
		314		374		330		340					
		*		\$		\$		340 *					
Cj	GRENS	KAFLK	ENPEI	ADEIT	KAIQN	SMGIE	GMISG	SEDDE	GEE				
Ng		rvwLK	ENPEI	SDEId	akIra	InGvE	mhIte	gtqDE	td				
Pa	GkaNa	akvie	ANDET		K+Trd	alaks	Gnyka	doeev	SOL				
TF	GkdNa	ravLK	VHPEL	AanIe	arIra	aaagh	plafa	eEves	pg				

FIG. 3. Amino acid alignments of the *C. jejuni* RecA protein with other bacterial RecA proteins. Abbreviations: Cj, *C. jejuni*; Ng, *N. gonorrhoeae* (16); Mf, *Methylobacillus flagellatum* (19); Pa, *Pseudomonas aeruginosa* (47); Tf, *Thiobacillus ferrooxidans* (43); Ec, *E. coli* (46). The *C. jejuni* sequence is shown in capital letters; residues in the other RecA proteins which are identical to the *C. jejuni* RecA are capitalized. A dash indicates a gap introduced to maximize alignment. The numbers refer to the amino acid residue of the *C. jejuni* protein.

transformation experiments revealed a defect in generalized recombination (see below).

Transfer of the recA mutant allele among C. jejuni strains by natural transformation of a PCR product. Since all attempts to introduce a mutation into 81-176 by using plasmid pPMP102 by conjugation or natural transformation were unsuccessful, an alternate mutational strategy was developed. Two PCR primers were designed, Rec1 and Rec2, which were capable of amplifying the entire recA gene. The Rec2 primer also includes the beginning of the putative enolase gene. The position of these primers is indicated in Fig. 2. These primers amplified a 1.2-kb fragment from VC83 and a 2.6-kb fragment from VC83 recA, further indication of the presence of the Km^r cassette in the mutant gene. The PCR product from VC83 recA was purified from agarose and used to transform 81-176 to Km^r. Approximately 50 transformants were obtained, and one was further characterized. Southern blot analysis of strain 81-176 (Fig. 4, lane C) and this Km^r transformant (lane D) indicated that the patterns were identical to those of VC83 wild type (lane A) and VC83 recA (lane B). UV sensitivity experiments indicated that the 81-176 mutant showed an increase in UV sensitivity similar to that seen in Fig. 5 for the VC83 mutant



FIG. 4. Southern blot analysis of *Campylobacter* DNAs probed with pPMP101. DNAs were digested with *XbaI* and *PstI*, electrophoresed on a 0.7% agarose gel, and transferred to nitrocellulose. Lanes: A, VC83; B, VC83 *recA*; C, 81-176; D, 81-176 *recA*; E, 81-116; F, 81-116 *recA*. The bands in lanes A and C correspond to the 1.8-kb insert in pPMP101, and those in lanes B and D are 3.2 kb, corresponding to an increase in 1.4 kb due to addition of the Km⁻ cassette.

(data not shown). The *recA* allele was similarly transformed by using the same PCR product in *C. jejuni* 81-116. Southern blot analysis of one transformant from this experiment is also shown in Fig. 4 (lane F) in comparison with 81-116 (lane E). The *recA*⁺ gene of 81-116 shows restriction polymorphisms from the *recA*⁺ genes of VC83 and 81-116, but the pattern of the transformant (Fig. 4, lane D) includes the 3.2-kb band of VC83 *recA* and 81-176 *recA* (lanes B and D, respectively). The UV sensitivity of the 81-116 mutant was also confirmed (data not shown).

Effect of recA mutations on natural transformation in C. *jejuni*. To determine the effect of recA mutations on natural transformation, we initially used DNA from a streptomycin-



FIG. 5. UV kill curves of strain 81-176 (open circles) and 81-176 *recA* (closed circles). Cultures were exposed to increasing doses of UV light and serially diluted as described in Materials and Methods.

	No. of Str ^r transformants/100 μ l ^b					
Recipient	C. coli VC167 Str ^{rc}	<i>C. jejuni</i> VC83 Str ^r	<i>C. jejuni</i> 81-176 Str ^r			
VC83	1.1×10^{4}	1.25×10^{4}	1.5×10^{3}			
VC83 recA	0	0	0			
81-176	1.4×10^{2}	22	2.3×10^{2}			
81-176 recA	0	0	0			
81-116	2.1×10^{2}	$5 imes 10^2$	2.2×10^{3}			
81-116 recA	0	0	0			

TABLE 1. Transformation of wild-type and *recA* mutants with Str^r marker from various strains"

^{*a*} Results represent the average of two experiments. ^{*b*} Viable counts were approximately 5×10^6 to $2 \times 10^7/100$ µl.

^c Source of DNA.

resistant (Str^r) strain of *Campylobacter coli*, VC167 (25). Cross-species transformations between *C. coli* and *C. jejuni* have been reported at different frequencies (1, 49). The results, shown in Table 1, indicate that the Str^r marker from *C. coli* VC167 can be transformed into all three strains of *C. jejuni*, although VC83 shows the highest frequency. *recA* mutants in all three strains showed no detectable transformation with VC167 DNA.

The differences in transformation frequencies among the three strains could be due to differences in inherent transformability or to differences in restriction of incoming DNA. To distinguish these possibilities, and to try to maximize the transformation frequencies in the wild-type strains, we purified DNA from cells of VC83 and 81-176 which had been transformed to Str^r with VC167 DNA and used this DNA to transform the C. jejuni strains to Strr. VC83 Strr DNA transformed VC83 at approximately the same frequency as VC167 Str^r, indicating that VC83 does not appreciably restrict incoming C. coli DNA. 81-176 Str^r DNA, however, transforms VC83 at a lower frequency than either VC83 Str^r or VC167 Str^r DNA. Transformation of VC167, VC83, and 81-176 Str^r DNAs into strain 81-116 occurs at approximately the same frequency. Transformation into 81-176 was consistently the lowest with all DNAs used but was particularly low with VC83 Str^r DNA. This reduction in transformation frequency was not due to restriction of VC83 DNA, however, since 6.9×10^2 transformants were obtained when VC83 recA DNA was used to transform 81-176 to Km^r.

Evaluation of ability of 81-176 recA to immunize rabbits against intestinal colonization. Similar to previously studied strains of C. jejuni (10, 40), rabbits fed either 81-176 or 81-176 recA for the first time were colonized without signs of diarrhea for 1 to 3 weeks. The mean duration of primary colonization for 81-176 and 81-176 recA mutants were 19.5 \pm 8.9 and 15.0 \pm 7.4 days, respectively (Table 2). One month after oral immunization, both immunized and control rabbits were challenged by the RITARD procedure with 10⁶ CFU of strain 81-176 per animal. This dose is approximately 4 logs higher than the 100% infectious dose (9). The mean number of days that rabbits were colonized after RITARD challenge is shown in Table 2. For both 81-176 and 81-176 recA, the duration of colonization after challenge was significantly shorter than in control animals fed only sterile broth. In both immunized groups, the majority of rabbits had negative fecal swabs by 2 days postchallenge. By day 1 postchallenge, fecal cultures were negative in two of four of the 81-176 recA-immunized rabbits and in two of five of the 81-176-immunized animals.

TABLE 2. Colonization	n of rabbits after	oral immunization with
10 ¹⁰ CFU of strain 81-17	6 or 81-176 recA	and RITARD challenge
with 10	⁶ CFU of strain	81-176

Immunization strain	No. of days colonized post- feeding (mean ± SD) ^a	No. of days colonized post-RITARD challenge (mean \pm SD) ^b
81-176	19.5 ± 8.9	1.6 ± 1.8
81-176 recA	15.0 ± 7.4	1.5 ± 1.9
None		7.0 ± 0.8

" The number of animals per group was six.

^b Number of animals per group: 81-176, five; 81-176 recA, four; control, four.

DISCUSSION

The predicted RecA protein of C. jejuni is highly conserved compared with other RecA proteins, showing the highest similarity to that of N. gonorrhoeae (16). The genomic organization is interesting in that the $recA^+$ gene apparently overlaps the enolase gene. Overlapping, unrelated genes have been described in C. jejuni before. The ATG start of the glyA gene overlaps the TGA stop codon of lysS (12, 13), and the transcriptional start of the cloned glyA gene has been mapped in E. coli and shown to be within the lysS gene (12, 13), suggesting that the two genes are transcribed independently in C. jejuni also. Transcriptional units remain to be determined for C. jejuni recA and enolase genes. It also remains to be determined what, if any, effect the insertional inactivation of recA has on the expression of the downstream enolase gene. The rather unusual occurrence of overlapping genes may offer a mechanism by which C. jejuni selects against deleterious mutations. Thus, mutations affecting the carboxy terminus of the RecA protein would also affect either the promoter or N terminus of enolase. Such a mechanism may be particularly important in a naturally transformable organism like C. jejuni.

It is not clear why conventional methods of conjugative suicide vector mutagenesis were unsuccessful in generating a recA mutation. The technique has been used in one strain of C. coli (20, 22) and several strains of C. jejuni (28, 54), including the generation of an lcrD mutant in 81-176 (35). The original recA mutant in VC83 was ultimately generated by introducing linearized pPMP102 into VC83 by natural transformation, a method which presumably forced a double crossover event. Subsequent analysis has indicated that circular plasmid DNA is also capable of generating the mutation (data not shown). This method of using natural transformation to introduce mutated alleles into Campylobacter spp. greatly simplifies mutant construction and should have general application. A similar method has been reported in the related organism, Helicobacter pylori (24), and we have constructed mutations in other genes by using this approach in both C. jejuni and C. coli (21, 56). The recA mutation was moved into other strains by using a PCR product from VC83 recA as the source of DNA. The use of VC83 recA DNA as the template allowed for synthesis of a full-length mutated recA allele without any adjacent DNA sequences. This is particularly significant in terms of transferring isogenic mutations among different strains without affecting adjacent markers. While the frequency of transformants using the PCR product was generally low (≤ 50 transformants), it was reproducible and a suitable method for introducing isogenic mutations into other strains. In Haemophilus influenzae and N. gonorrhoeae, there exist specific DNA uptake sequences which are involved in the early steps of natural transformation. It has been suggested that similar uptake sites exist for Campylobacter spp. (53). If this is the case, such an uptake site must be present on the DNA between the Rec1 and

Rec2 primers used to generate the PCR product. Comparison of this sequence with other clones of *C. jejuni* DNA capable of natural transformation should help to elucidate this uptake sequence.

Natural transformation of C. jejuni is clearly dependent on a functional RecA gene product. No transformation was detectable when a standard assay system was used in any of the three recA mutant strains. Both Wang and Taylor (53) and Alm et al. (1) reported considerable variation in the ability of different strains to be transformed, and similar observations were made in this study. VC83 transformed consistently better than strain 81-176 or 81-116, even with C. coli donor DNA. The highest numbers of transformants were seen when VC83 was transformed with DNA from Str^r VC83, and this level of transformation is approximately 10-fold higher than the highest reported by Wang and Taylor (53). Our transformation data do not suggest any significant degree of restriction of DNA, including C. coli DNA. The very low level of transformation seen with 81-176 when Str VC83 DNA was used was surprising but was not due to restriction since the kanamycin marker from VC83 recA transformed 81-176 at a high frequency. The genomic map may vary between 81-176 and VC83 in such a way that the crossover of the Str^r marker is markedly reduced in 81-176.

The introduction of a recA allele into the live oral cholera vaccine strain CVD103 diminished colonization ability and immunogenicity in human feeding studies (26). However, the introduction of a recA allele into another V. cholerae strain resulted in minimal reduction in colonization ability (41). In this study, 81-176 recA colonized rabbits almost as well as the wild type and was capable of protecting against subsequent homologous wild-type and was capable of protecting against subsequent homologous wild-type challenge. The absence of detectable natural transformability without loss of immunogenicity would indicate that recA mutations would be useful for inclusion into vaccine strains. Further characterization will be needed to determine what, if any, effect the introduction of a recA allele has on virulence in in vivo and in vitro models. Ultimately, it could be envisioned that antibiotic-sensitive recA mutations could be combined with mutations in virulence genes to generate living attenuated vaccine strains of Campylobacter spp. expressing the full complement of surface antigens.

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