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Genomic analysis of diverse rubella virus genotypes

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Based on the sequence of the E1 glycoprotein gene, two clades and ten genotypes of *Rubella virus* have been distinguished; however, genomic sequences have been determined for viruses in only two of these genotypes. In this report, genomic sequences for viruses in an additional six genotypes were determined. The genome was found to be well conserved. The viruses in all eight of these genotypes had the same number of nucleotides in each of the two open reading frames (ORFs) and the untranslated regions (UTRs) at the 5' and 3' ends of the genome. Only the UTR between the ORFs (the junction region) exhibited differences in length. Of the nucleotides in the genome, 78% were invariant. The greatest observed distance between viruses in different genotypes was 8.74% and the maximum calculated genetic distance was 14.78 substitutions in 100 sites. This degree of variability was similar among regions of the genome with two exceptions, both within the P150 non-structural protein gene: the N-terminal region that encodes the methyl/guanylyltransferase domain was less variable, whereas the hypervariable domain in the middle of the gene was more divergent. Comparative phylogenetic analysis of different regions of the genome was done, using sequences from 43 viruses of the non-structural protease (near the 5' end of the genome), the junction region (the middle) and the E1 gene (the 3' end). Phylogenetic segregation of sequences from these three genomic regions was similar with the exception of genotype 1B viruses, among which a recombinational event near the junction region was identified.

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INTRODUCTION

Rubella virus is an important human pathogen that causes an acute, contagious disease known as rubella, 3-day measles or German measles, and severe birth defects (known as congenital rubella syndrome) when infection occurs during the first trimester of pregnancy (Chantler *et al.*, 2001). *Rubella virus* is the single member of the genus *Rubivirus* in the family *Togaviridae* and is an enveloped, single-stranded, positive-polarity RNA virus with a genome of approximately 10 kb. The genome contains two long open reading frames (ORFs): the 5'-proximal ORF (NSP-ORF) encodes two non-structural proteins, P150 and P90, that function in RNA replication, and the 3'-proximal ORF (SP-ORF) encodes three structural proteins: the capsid protein, C, and two envelope glycoproteins, E1 and E2. The SP-ORF is translated from a subgenomic RNA synthesized in infected cells (Frey, 1994). The genome also contains untranslated regions (UTRs) at its 5' and 3' ends and between the ORFs (known as the junction region).

Although rubella occurs worldwide, vaccination efforts with live-attenuated vaccines have been concentrated in

developed countries. Currently, approximately 50% of countries have national vaccination efforts against rubella (Robertson *et al.*, 2003). Isolation and genetic sequencing of rubella viruses has been most thorough in countries pursuing elimination (Bosma *et al.*, 1996; Frey *et al.*, 1998; Icenogle *et al.*, 2006; Katow, 2004; Katow *et al.*, 1997a, b; Reef *et al.*, 2002; Saitoh *et al.*, 2006); however, collections have recently been assembled from other regions of the world (Donadio *et al.*, 2003; Katow, 2004; Zheng *et al.*, 2003a, c). Recently, a standard taxonomy for rubella viruses was adopted based on sequences of a standard window within the E1 gene and supported by sequencing of the SP-ORF of selected viruses (WHO, 2005). The taxonomy consists of two clades [corresponding to the previous genotypes I and II (Frey *et al.*, 1998; Zheng *et al.*, 2003a)] containing a total of ten genotypes, seven in clade 1 (1a, 1B, 1C, 1D, 1E, 1F and 1g) and three in clade 2 (2A, 2B and 2c); the genotypes designated in lower case are provisional. Within the E1 gene, maximal variation among clade 1 viruses is 5.8%, that among clade 2 viruses is 8.0%, and it is 8.2% between the two clades (Zheng *et al.*, 2003a). Geographically, clade 1 viruses circulate worldwide, whilst clade 2 viruses thus far have been restricted to Eurasia.

Thus far, ten complete genomic sequences of *Rubella virus* have been reported, which represent only two of the ten

Supplementary tables are available in JGV Online.

genotypes (eight sequences of genotype 1a viruses and two sequences of genotype 2A viruses). Among these sequences, genomic genetic variability is similar to that in the E1 gene, with the exception of a 'hypervariable region' (HVR) of greater variability in the middle of the P150 non-structural protein gene (Hofmann *et al.*, 2003; Zheng *et al.*, 2003b). Given the lack of representation of the majority of the genotypes in the current genomic database, the first goal of this study was to expand the number of genomic sequences, using viruses in our collection from six additional genotypes (1B, 1C, 1D, 1E, 2B and 2C). The second goal of this study was to extend phylogenetic analysis to 5' regions of the genome, which had not previously been done. To this end, the sequence of the non-structural protease-encoding region within the P150 gene was determined and compared phylogenetically with the sequences of the junction region and the E1 gene from 43 viruses representing eight genotypes.

METHODS

Viruses, cells, RNA extraction, cDNA amplification and DNA sequencing. The viruses analysed in this study are listed in Table 1 (genomic sequences) and Supplementary Table S1, available in JGV Online (genomic regions). The Cba strain was provided by Dr Marta Zapata, University of Cordoba, Cordoba, Argentina. A monolayer of Vero cells (25 cm² T-flask or 60 mm plate) was infected with each virus. Three to five days post-infection, the culture medium was removed and total cellular RNA was extracted by using Tri-Reagent (Molecular Research Center) using the manufacturer's protocol. The RNA extracted from one 25 cm² T-flask or one 60 mm plate was resuspended in 50 µl double-distilled H₂O and stored at -80 °C until use. cDNA was synthesized in a 20 µl total reaction volume reaction containing 5 µl denatured (95 °C, 5 min)

RNA template, 4 µl 5 × Reverse Superscript buffer (Invitrogen), 4 µl 2.5 mM dNTPs, 1 µl 0.1 M dithiothreitol, 1 µl 4 µM 3'E1/808 reverse primer (5'-TTTTTTTTTCTATACAGCAAC-3'; T₉ followed by the complement of nt 9751-9762 of the rubella virus genome), 1 µl (40 units) RNasin (Promega) and 1 µl (200 units) Superscript reverse transcriptase III (Invitrogen). The reaction was incubated at 55 °C for 60 min and then stored at -20 °C prior to use in PCR. Each 50 µl PCR contained 25 µl 2 × GC buffer I&II (TaKaRa), 8 µl 2.5 mM dNTPs, 3 µl cDNA template, 1 µl 40 µM appropriate forward and reverse primers and 0.5 µl (2.5 units) LA *Taq* polymerase (TaKaRa). Cycling parameters were determined according to the manufacturer's (TaKaRa) recommendations. For genomic sequencing, 10-11 overlapping fragments encompassing the entire genome were amplified by using appropriate primer pairs. The primers used to amplify the genomic regions are listed in Supplementary Table S2, available in JGV Online. Amplified fragments were purified following agarose-gel electrophoresis by using a QIAquick gel extraction kit (Qiagen). Sequencing reactions were performed bidirectionally by using appropriate primers and cycle-sequencing kits (ABI PRISM BigDye Terminator v. 3.1; PE Applied Biosystems) and resolved by using a 3100 Genetic Analyzer (Applied Biosystems). The 5' and 3' ends of the genome were determined by using a 5'/3' FirstChoice RLM-RACE kit (Ambion Inc.).

Sequence analysis. For cataloguing and storage, sequences were input into free online sequence-alignment software (ALIGN Query, GENESTREAM SEARCH network server IGH, Montpellier, France; <http://xylian.igh.cnrs.fr/bin/align-guess.cgi>). The assembled nucleotide sequences were aligned by using the CLUSTAL_W multiple sequence-alignment program version 1.8 (Henikoff & Henikoff, 1994) and the PileUp program in the GCG software package (Genetics Computer Group, version 11.0; Accelrys Inc.). The TN93 substitution model (Tamura & Nei, 1993) with discrete gamma-distributed rate heterogeneity with eight gamma rate categories (Yang, 1994) (TN93 + Γ model) was used as a substitution model for phylogenetic reconstruction, as it was found statistically to be the best fit for our datasets. Maximum-likelihood (ML) phylogenetic analysis

Table 1. Rubella virus genomic sequences used in this study

Virus	Isolation site and year	Genotype	GenBank accession no.	Reference
Fth_USA64	Connecticut, USA, 1964	1a	M15240	Dominguez <i>et al.</i> (1990)
RA27_USA64	Pennsylvania, USA, 1964	1a	L78917	Pugachev <i>et al.</i> (1997)
M33_USA61	New Jersey, USA, 1961	1a	X05259, X72393	Clarke <i>et al.</i> (1987)
CEN_BEL63	Belgium, 1963	1a	AF188704	Lund & Chantler (2000)
TO-W_JAP67	Toyama, Japan, 1967	1a	AB047330	Kakizawa <i>et al.</i> (2001)
TO-V_JAP67	Toyama, Japan, 1967	1a	AB047329	Kakizawa <i>et al.</i> (2001)
ULR_GER84	Leipzig, Germany, 1984	1a	AF435865	Hofmann <i>et al.</i> (2003)
SUR_SVK74	Bratislava, Slovakia, 1974	1a	AF435866	Hofmann <i>et al.</i> (2003)
Cba_ARG88	Cordoba, Argentina, 1988	1B	DQ085339	This report
GUZ_GER92	Stuttgart, Germany, 1992	1B	DQ388280	This report
Anim_MEX97	Baja California, Mexico, 1997	1C	DQ085341	This report
JC2_NZL91	Auckland, New Zealand, 1991	1D	DQ388281	This report
6423_ITA97	Pavia, Italy, 1997	1E	DQ085343	This report
BRDI_CN80	Beijing, China, 1980	2A	AY258323	Zheng <i>et al.</i> (2003b)
BRI_CN79	Beijing, China, 1979	2A	AY258322	Zheng <i>et al.</i> (2003b)
AN5_KOR96	Seoul, Korea, 1996	2B	DQ085342	This report
I-11_ISR68	Tel-Aviv, Israel, 1968	2B	DQ085338	This report
C74_RUS97	Moscow, Russia, 1997	2c	DQ085340	This report
C4_RUS67	Moscow, Russia, 1967	2c	DQ388279	This report

was performed by using the TREE-PUZZLE program version 5.2 (Strimmer & von Haeseler, 1996). ML genetic distances and nucleotide-substitution statistical parameters were estimated under the selected TN93+ Γ substitution model with an initial neighbour-joining tree and then the best ML tree was reconstructed with these optimized parameters by using the quartet-puzzling method in the TREE-PUZZLE program. Fifty thousand and one hundred thousand quartet-puzzling steps were performed in constructing trees from the 19 genomic and 43 genomic region sequences, respectively. Sequence similarities and observed distances were calculated by using the Old Distance program in the GCG software package. A nucleotide sequence PLOTSIMILARITY plot across the genome (100 nt window) was generated by using the PLOTSIMILARITY program in the GCG software package. As the sequences of genotype 1a viruses were over-represented, the plot was generated by using six sequences from each clade, including members from each genotype. Nucleotide sequence substitution-rate analysis was carried out with PILEUP (GCG package), fastDNAMl (version 1.2.2) and DNArates (version 1.1.0), employing default parameters. To detect recombination, phylogenetic analysis of sequences on either side of putative break points was conducted by using TREE-PUZZLE with the same parameter settings as were used in the genomic sequence analysis. Recombination was also analysed by using the sequence recombination-detection programs TOPALi (Milne *et al.*, 2004), RIP 2.0 (Recombination Identification Program; <http://hivweb.lanl.gov/RIP/RIPsubmit.html>) and the four-cluster likelihood mapping analysis in the TREE-PUZZLE program.

RESULTS

Genomic sequences and comparisons

A representative rubella virus phylogenetic tree based on the standard E1 gene window recommended by the WHO (nt 8291–9469) and containing the reference viruses for each genotype and the ten viruses for which the genomic sequence has been determined is shown in Fig. 1. As the genomic sequences were from genotype 1a and 2A viruses, the genomic sequences of nine representative viruses from six additional genotypes were determined (Table 1). Among these 19 viruses, with three exceptions, the genomes were 9762 nt in length and consisted (5'–3') of a 40 nt 5' UTR, a 6351 nt NSP-ORF, a 120 nt junction region, a 3192 nt SP-ORF and a 59 nt 3' UTR. All three exceptions were in the junction region: the genome of one of the genotype 1B viruses (GUZ_GER92) was 9760 nt in length because it had a deletion of 2 nt at positions 6480–6481 (between the end of the NSP-ORF and the SG RNA start site) and the genomes of both genotype 2B viruses were 9761 nt in length because they had a deletion of 1 nt at position 6422 (between the SG RNA start site and the start of the SP-ORF). Fig. 2 shows a similarity plot across 12 genomic sequences proportionally representing all eight genotypes. Overall variability averaged approximately 7% and was roughly comparable across the genome, with the exceptions of the 5'-terminal approximately 400 nt, a region encoding the methyl/guanlyltransferase (MT) domain within the P150 gene that exhibited variability of approximately 4%, and a region of the P150 gene encompassing nt 2100–2400, the HVR, in which local variability peaked at up to 18%. Pairwise observed genomic distances between viruses in different genotypes (see

Supplementary Table S3, available in JGV Online) ranged from 2.0 to 8.7%. The range of pairwise observed distances for genomic regions (the five genes and domains within P150 and P90) and the 3' *cis*-acting element (3'CAE) is given in Table 2. Maximal observed distances of these regions were shown to range from 8.29 to 11.44%, with the exception of the MT domain (5.24%) and the HVR (21.18%). Given this limited degree of variability across most of the genome, it was not surprising that 78% of the nucleotides in the genome were invariant across the 19 sequences (Table 2).

The additional sequences contributed by this report greatly expand information on the genomic diversity of rubella viruses and, therefore, we took the opportunity to calculate a number of evolutionary parameters (Table 2). The maximum pairwise genetic distance (see Supplementary Table S3, available in JGV Online) was 14.78 substitutions in 100 sites, greater than the largest observed distance (8.74 observed substitutions in 100 sites). The maximal genetic distances for the genomic regions (the five genes, domains within P150 and P90 and the 3'CAE) ranged from 13.97 to 23.0 substitutions in 100 sites, with the exception of the MT domain (6.77 substitutions in 100 sites) and the HVR (35.85 substitutions in 100 sites). The transition/transversion site parameter (K) was 7.04 for the entire genome and ranged among genomic regions from 4.5 (HVR) to 13.35 (3'CAE). The pyrimidine and purine (Y/R) transition parameter of entire genome was 2.7 and varied among the genomic regions from 0.93 (X domain of the P150 gene) to 3.7 (C gene). To test whether the observed distance simply underestimates the genetic distance or whether substitution saturation has been reached, the pairwise number of transitions and transversions was plotted as a function of the calculated genetic distance (by using DAMBE; Xia & Xie, 2001), with the result that both transitions and transversions increased linearly with genetic distance, with the number of transitions being higher than transversions (data not shown). Neither reached a plateau, indicating that substitution saturation had not occurred.

Among the 19 genomic sequences, 78% of the nucleotides were invariant. Not surprisingly, the parameter of rate heterogeneity, α , was 0.22 for the entire genome and varied between 0.19 and 0.33 across the genomic regions, with the exception of the HVR, within which $\alpha = 1.35$. These small α values indicated a strong substitution-rate heterogeneity among nucleotide sites across most of the genome (i.e. more than three-quarters of the nucleotides remained constant, whilst fewer than one-quarter exhibited variability). Within the HVR, 46% of the nucleotides were variable.

Phylogenetic analysis

ML phylogenetic trees constructed from the complete genomic sequences, as well as from the NSP- and SP-ORFs, are displayed in Fig. 3. As in the E1-based tree, the six clade 2 sequences formed a clear, consistent branching pattern with high support values in all three trees, indicating that

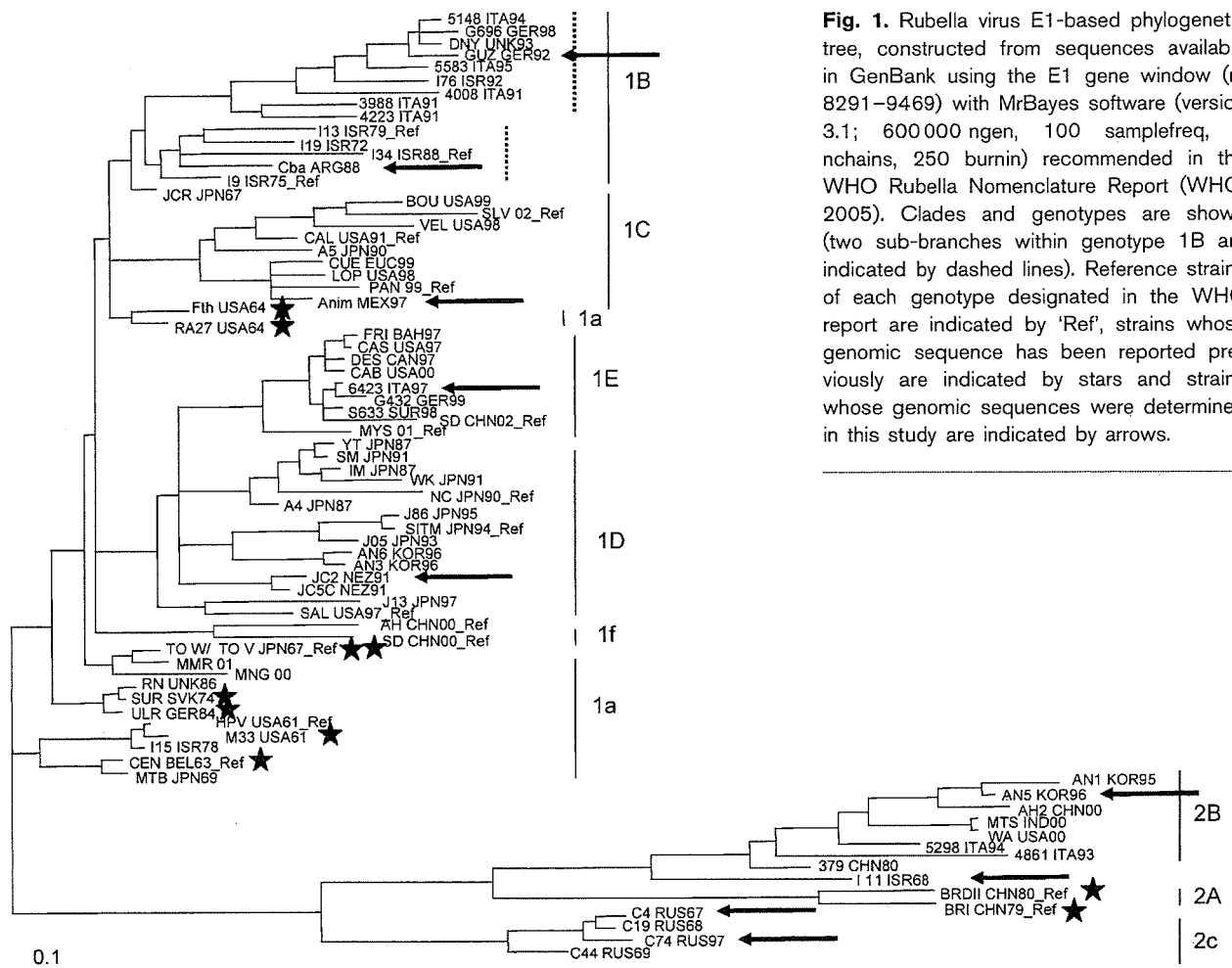


Fig. 1. Rubella virus E1-based phylogenetic tree, constructed from sequences available in GenBank using the E1 gene window (nt 8291–9469) with MrBayes software (version 3.1; 600 000 ngen, 100 samplefreq, 4 nchains, 250 burnin) recommended in the WHO Rubella Nomenclature Report (WHO, 2005). Clades and genotypes are shown (two sub-branches within genotype 1B are indicated by dashed lines). Reference strains of each genotype designated in the WHO report are indicated by 'Ref', strains whose genomic sequence has been reported previously are indicated by stars and strains whose genomic sequences were determined in this study are indicated by arrows.

genotypes 2A and 2B are related more closely to each other than to genotype 2c. In clade 1, the groupings of genotypes 1B, 1C, 1D and 1E on the three trees were consistent: the two genotype 1B sequences formed a branch, as did the individual genotype 1D and 1E sequences, whilst the individual genotype 1C sequence extended from the baseline with no relative relationship to other genotypes. On the genomic and SP-ORF trees, the eight genotype 1a sequences grouped into four pairs, indicated in Fig. 3 as a1 (TO-w and TO-v; a wild-type parent and the attenuated vaccine derived from it), a2 (Fth and RA27/3; both isolated in the north-eastern USA in 1964), a3 (CEN and M33; isolated from Europe and the USA in 1961–1962) and a4 (SUR and ULR; both isolated from eastern Europe in 1974 and 1984). Interestingly, on the NSP-ORF tree, M33 separated from CEN (a3) and clustered with the a4 grouping. We also constructed trees by using the sequences of the genes and regions within the NSP-ORF and SP-ORF (data not shown), with the result that they had the same general topology as the ORF-generated trees. The exception was the HVR-generated tree, on which each of the clade 1 viruses formed an individual branch, apart from the a1 and M33-SUR groupings, which were preserved.

Extensive phylogenetic analysis has not been done previously using sequences from 5' regions of the genome. To do so, trees were constructed from the sequences of the non-structural protease (NP) region (nt 3035–3973; Fig. 2), the junction region and adjacent sequences (JR; nt 6351–6829, which includes the 3' end of the P90 gene, the UTR between the ORFs and the 5' end of the C gene) and the E1 gene [nt 8731–9469, the recommended window for routine genotyping (WHO, 2005)] of 43 viruses representing eight genotypes. ML phylogenetic trees constructed from these sequences are shown in Fig. 4(a). Clustering of viruses on the three trees was similar, with the exception of a group of seven genotype 1B viruses that formed a single branch on the NP tree and two branches (one of five and one of two viruses) on the JR tree, but did not form a cluster on the E1 tree. The single-nucleotide deletion at nt 6422 in the junction region detected in the genome sequence of the two genotype 2B viruses was confirmed in the JR sequences determined from three additional genotype 2B viruses, and the 2 nt deletion at nt 6480–6481 detected in the genome sequence of one of the two genotype 1B viruses was discovered in the JR sequences determined from four of the five additional genotype 1B viruses. Interestingly, this 2 nt

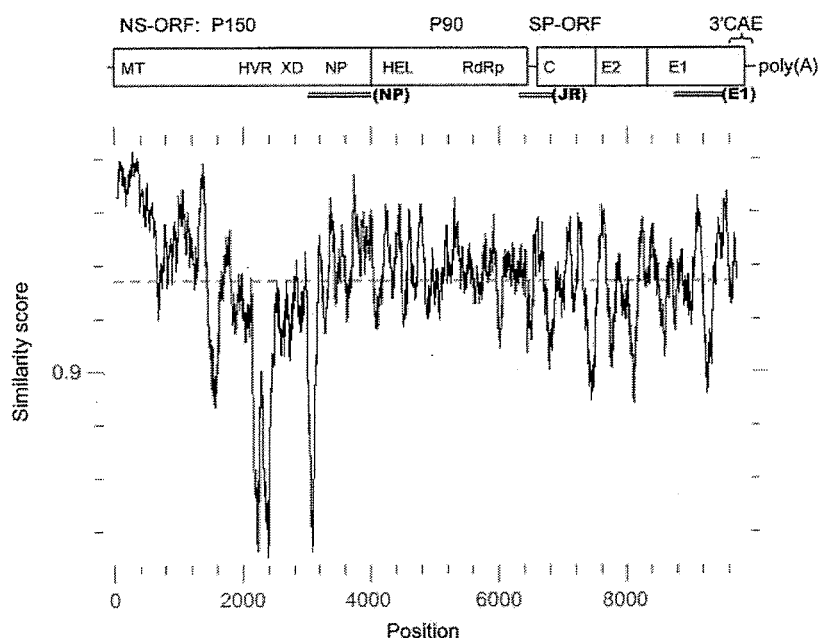


Fig. 2. Rubella virus genome nucleotide-similarity plot. The plot was generated from 12 genome sequences proportionally representing the two clades and eight genotypes by using the PLOTSIMILARITY program in the GCG software package with a 100 nt window. A genomic map is overlaid above the plot that shows the two ORFs, five genes, regions/domains within the NS-ORF (MT, methyl/guanylyltransferase; HVR, hypervariable region; XD, X domain; NP, non-structural protease; HEL, helicase; RdRp, RNA-dependent RNA polymerase or replicase) and the 3' *cis*-acting elements (3'CAE). The regions from which sequences were determined from 43 viruses for comprehensive phylogenetic analysis (see Fig. 5) are also denoted (NP, non-structural protease; JR, junction region; E1, WHO standard E1 window).

Table 2. Maximum-likelihood parameters calculated for regions of the rubella virus genome

Region	G+C content (mol%)	No. sites	No. invariant sites (%)	Observed distance (range)*	Genetic distance (range)*	K*	Y/R*	α^*
Genome†	69.5	9762	7611 (78.0)	0.22–8.74	0.22–14.78	7.04	2.7	0.22
P150†	71.8	3903	3033 (77.7)	0.26–9.3	0.26–15.89	5.97	2.25	0.23
P150: MT†	67.4	210	189 (90.0)	0–5.24	0–6.77	9.9	3.05	0.23
P150: HVR†	81.1	321	174 (54.2)	2.49–21.18	0.31–35.85	4.5	1.78	1.35
P150: XD†	74.2	507	377 (74.4)	0–11.44	0–23	7.2	0.93	0.24
P150: NP†	72.7	900	700 (77.8)	0.33–10	0.34–17.04	6.31	2.4	0.24
P90†	67.2	2448	1952 (79.7)	0.12–8.29	0.12–13.97	10.67	3.08	0.21
P90: HEL†	66.8	756	596 (78.8)	0.4–8.73	0.4–16.9	9.95	3.39	0.26
P90: RdRp†	66.9	1563	1250 (80.0)	0–8.83	0–15.3	11.06	2.74	0.19
C†	72.8	900	697 (77.4)	0.11–9.44	0.11–17.59	7.26	3.7	0.26
E2†	71	846	625 (73.9)	0–10.05	0–15.86	6	3.13	0.33
E1†	66.2	1446	1153 (79.7)	0.14–9.2	0.14–15.49	7.6	2.82	0.19
3'CAE†	64.9	307	244 (79.5)	0–8.79	0–14.67	13.35	2.98	0.2
NP‡	73	939	670 (71.4)	0–10.44	0–16.69	5.98	2.35	0.28
JR‡ ^c	69.9	479	334 (69.7)	0–10.67	0–16.66	5.43	2.69	0.32
E1‡	67.5	739	546 (73.9)	0–10.42	0–21.06	7.74	3.42	0.21

*Parameters were calculated by using the TREE-PUZZLE program with setting of the TN93 substitution model with gamma discrete distribution-rate heterogeneity. K, Transitions/transversions; Y/R, pyrimidine transitions (T↔C)/purine transitions (A↔G); α , rate-heterogeneity shape parameter.

†Parameters were calculated from the indicated regions by using the 19 genomic sequences. P150: nt 41–3943; MT: methyltransferase, nt 230–439; HVR: hypervariable region, nt 2120–2440; XD: X domain, nt 2492–2998; NP: non-structural protease, nt 3041–3940; P90: nt 3944–6391; HEL: helicase region, nt 4043–4798; RdRp: RNA-dependent RNA polymerase region, nt 4826–6388; C, nt 6512–7411; E2: nt 7412–8257; E1: nt 8258–9703; 3'CAE: 3' *cis*-acting elements, nt 9456–9762.

‡Parameters were calculated from the sequences of the NP (nt 3035–3973), JR (nt 6351–6829) and E1 (nt 8731–9469) regions determined for 43 viruses.

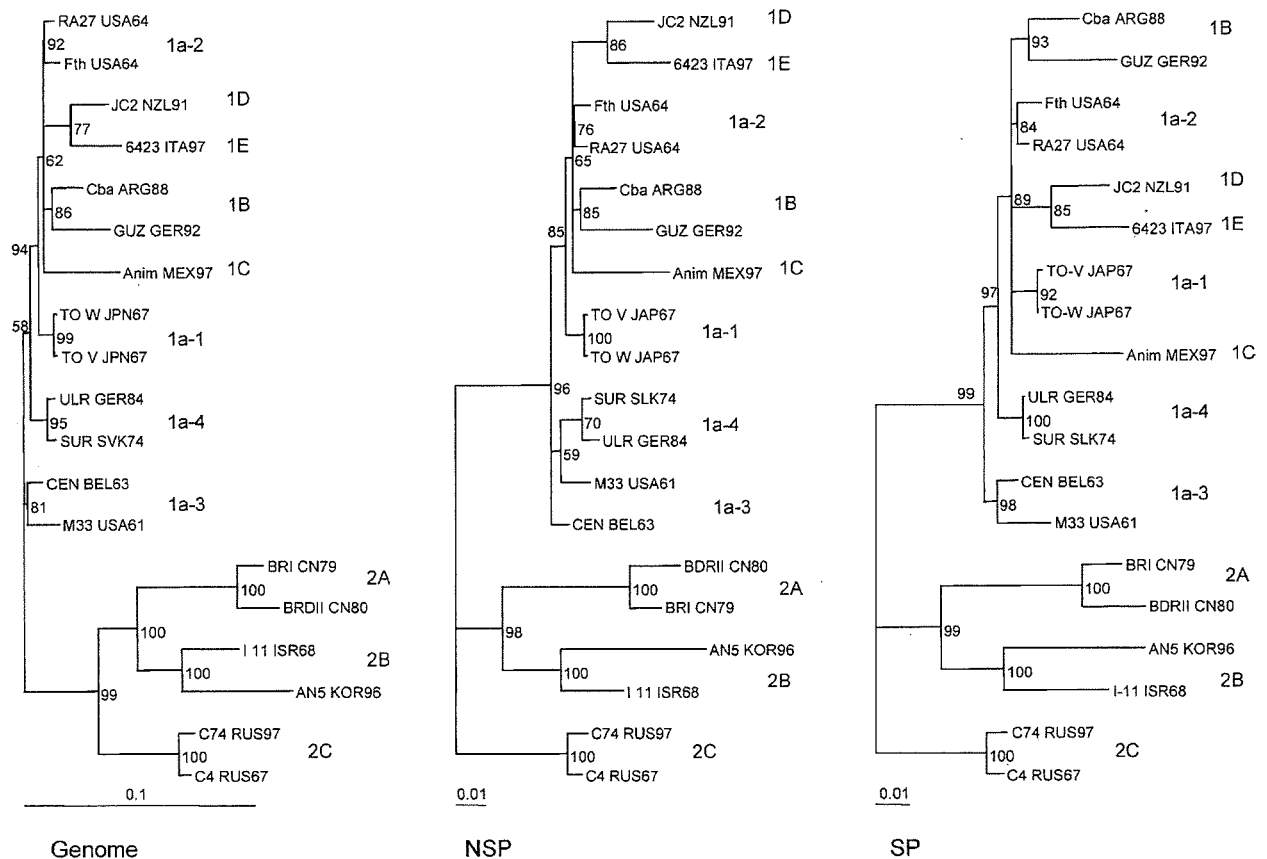


Fig. 3. Phylogenetic trees of genomic, NS-ORF and SP-ORF sequences. Trees were constructed by using TREE-PUZZLE (version 5.2) with 50 000 puzzle steps; reliability values are indicated on each node. Genotypes are denoted, including four subclusters of genotype 1a (1a-1 to 1a-4). Genetic distance (substitutions in 100 nt) calculated by using the TN93+ Γ substitution model is indicated by the bar below each tree.

deletion did not co-segregate with the JR sequence of these viruses, as it was present in three members of the five-virus genotype 1B branch and both members of the two-virus genotype 1B branch [marked by an asterisk in Fig. 4(a)]. Evolutionary parameters calculated from these larger and more genotypically representative sequence sets, shown in Table 2, were similar to those calculated by using the smaller genome sequence set.

Detection of genomic recombination among genotype 1B viruses

The lack of co-segregation of the genotype 1B JR sequences and the 2 nt deletion led us to hypothesize that a recombinational event had occurred in this region of the genome of these viruses, at or downstream of the deletion. To test this hypothesis, we expanded the sequence determined upstream into the 3' end of the P90 gene (the RdRp domain) and employed several software programs designed to detect recombination events, as well as secondary phylogenetic analysis, to detect putative break points. The RIP program predicted a break point at nt 6555,

within the 5' end of the C gene (which begins at nt 6512). As shown in Fig. 4(b), this prediction is supported by trees of sequences up- and downstream of this break point. Similar to the NP tree in Fig. 4(a), all seven genotype 1B sequences form a branch on the tree constructed by using sequences upstream of this break point (nt 5720–6554). On the tree constructed by using sequences downstream of the break point (nt 6555–6814), two sequences (TOM_UNK86 and 0754_GER92) were on a branch distinct from the other genotype 1B sequences, similar to the JR tree in Fig. 4(a). We thus conclude that a recombinational event occurred at or near this site during the evolution of these viruses.

DISCUSSION

The goal of this study was to expand the rubella virus genomic sequence database to include viruses in the majority of the currently defined genotypes. Whilst ten genomic sequences had been reported previously, only two of the ten currently defined genotypes were represented. This study added nine genomic sequences representing an additional six genotypes, encompassing the most widely

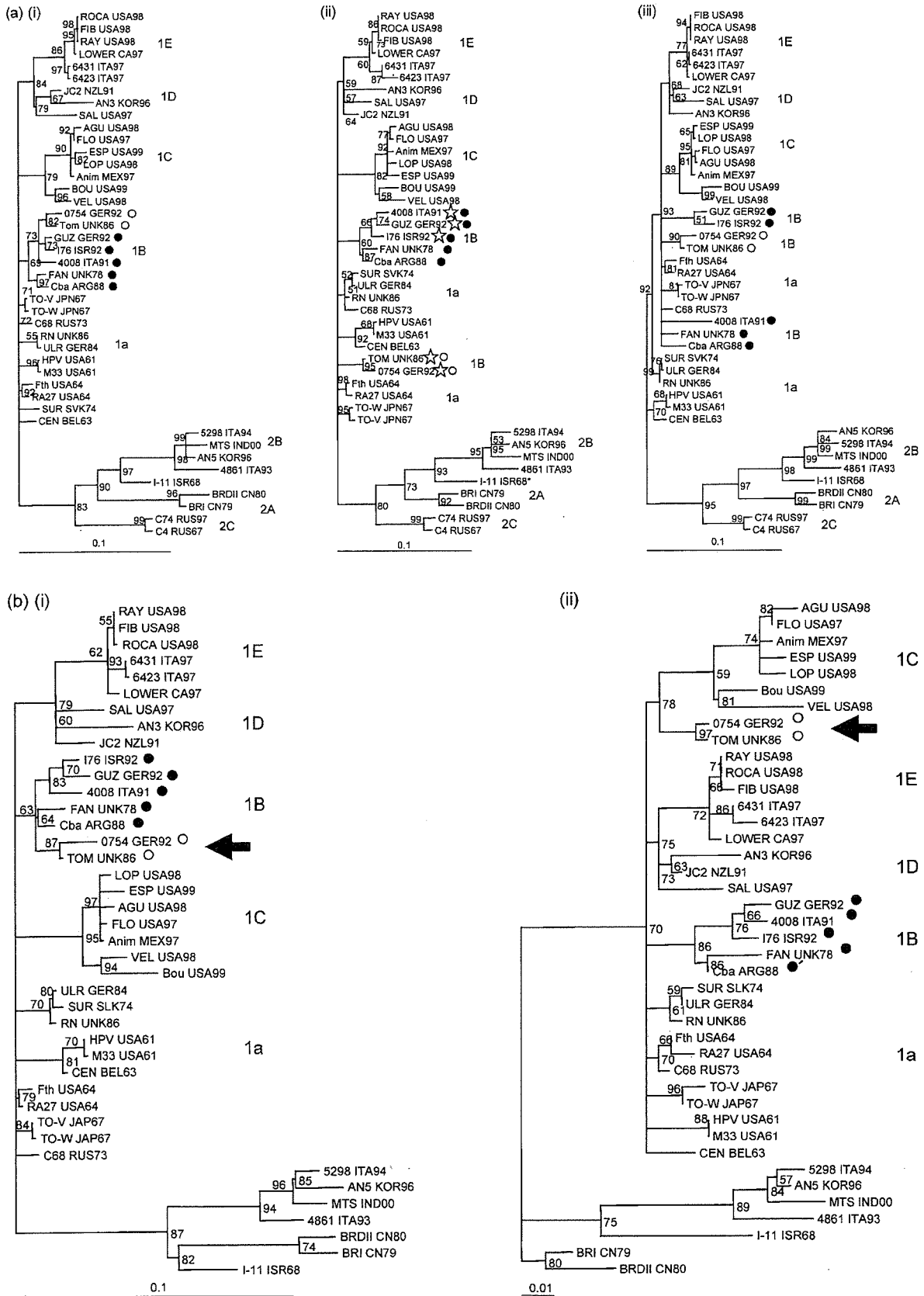


Fig. 4. Phylogenetic trees based on sequence of genomic regions. (a) Trees were constructed using the sequence of the (i) non-structural protease (NP, nt 3035–3973), (ii) junction region (JR, nt 6351–6829, including the 3' end of the NS-ORF, the UTR between the ORFs and the 5' end of the C gene) and (iii) WHO standard E1 window (nt 8731–9469). On all three trees, genotype 1B viruses are indicated by dots (filled or empty for the clusters of five or two viruses, respectively, on the JR-based tree), and in (ii), genotype 1B viruses with a 2 nt deletion at nt 6480–6481 are indicated by stars. (b) Trees were constructed by using nt 5720–6554 (i) or 6555–6814 (ii), on either side of the putative recombination break point. Differential segregation of two of the genotype 1B viruses is indicated by arrows. All trees were constructed with TREE-PUZZLE (version 5.2; 100 000 puzzle steps). Reliability values are indicated on each node and genetic distance (substitutions in 100 nt) calculated by using the TN93+ Γ substitution model is indicated by the bar below each tree.

divergent genotypes. The most striking finding was the genomic uniformity of rubella viruses, as it was discovered that 78 % of the nucleotides in the genomes of the viruses from the eight genotypes were invariant and these viruses preserved identical genomic dimensions across the two ORFs and two of the three UTRs. Only in the junction region (the UTR between the ORFs) of genotype 2B viruses, which had a 1 nt deletion, and a subset of genotype 1B viruses, which had a 2 nt deletion, was any plasticity observed. Such strict uniformity of genomic topology is highly unusual among RNA viruses (Huang *et al.*, 2004; Kang *et al.*, 2004; Kinney *et al.*, 1998; Saleh *et al.*, 2003; Takahashi *et al.*, 2003; Tarbatt *et al.*, 1997; van Cuyck *et al.*, 2003; Yang *et al.*, 2004). Sequence diversity was also low; across the eight genotypes, the maximum observed distance was <9 % and the maximum calculated genetic distance was 14.8 substitutions in 100 sites. Regardless of this difference, substitution saturation had not occurred, indicating that, despite the limited sequence diversity among rubella viruses, sufficient phylogenetic signal was retained to support the groupings observed (Salemi & Vandamme, 2003; Xia, 2000; Xia & Xie, 2001).

A sequence-similarity profile revealed, with two exceptions, a comparable pattern across the genome with local windows of similarity and dissimilarity varying about a relatively uniform mean, indicating that most genomic regions, including both virion protein and replicase protein genes, were equally divergent. Both observed and genetic distances between these genomic regions were comparable. The two exceptions were both within the P150 gene, with the N-terminal MT domain exhibiting less variability and the internal HVR exhibiting greater variability. Although the MT domain was predicted to encode both methyl- and guanylyltransferase activities (Rozanov *et al.*, 1992; neither activity has been demonstrated experimentally), the fact that 90 % of the nucleotide residues within this region are conserved raises the possibility that this region serves as a CAE in addition to encoding protein sequence. Consistent with this possibility, the phenotype of a cell culture-potentiating mutation discovered at nt 164 of the RA27/3 genome was found to be due to the nucleotide itself rather than to the encoded amino acid (Pugachev *et al.*, 2000). Conservation of the MT domain sequence has also been observed in other alpha-like family viruses (Gouvea *et al.*, 1998). The HVR encodes a proline- and arginine-rich domain of P150 termed the 'proline hinge' (Koonin *et al.*,

1992), although this domain contains several adaptor motifs that could serve to facilitate the association of P150 with other proteins. If this domain serves as a structural hinge between functional domains within the P150 protein, this could explain the lower constraint on sequence conservation within the HVR in comparison with the rest of the genome. On the other hand, Hofmann *et al.* (2003) reported data suggesting that the HVR among clade 1 viruses was under positive selection at the amino acid level. It should be pointed out that 'hypervariable region' is a relative term, in that HVRs in the genomes of other viruses are often more variable than the rubella virus HVR. For example, in the hepatitis E virus HVR, variability is > 50 % (Arankalle *et al.*, 1999; Gouvea *et al.*, 1998; Nishizawa *et al.*, 2003; van Cuyck *et al.*, 2003).

With the exception of the HVR, nucleotide residues or sites across the genome showed a strong heterogeneity in rate of divergence, as indicated by the low value of the rate-heterogeneity parameter α . Sequence collections with low α values exhibit an L-shaped distribution on a graph of number of sites versus rate of divergence, rather than the bell-shaped curve generated when α is 1 (or > 1). The low α value reflects the fact that roughly 80 % of the residues in the rubella virus genome were invariant in this collection of genomic sequences. The percentage of invariant residues at first and second codon positions was 93 %, compared with 48 % at third codon positions (Y. Zhou, unpublished data), and thus maintenance of amino acid sequence is a substantial component of the conservation of nucleotide sequence. Among third codon positions, the G + C content was 81 mol%, compared with 63 mol% among first and second codon positions (Y. Zhou, unpublished data), and thus there was selection for G and C residues. This selection was also evident in the HVR, within which the G + C content was 81 mol%, compared with 70 mol% for the genome.

Among the nucleotide substitutions at the 20 % of genomic sites that exhibited variability, transitions were strikingly more abundant than transversions; across the genome, the transition to transversion ratio, K, was 7.0 and varied among genomic regions from 4.5 to 13.4. Thus, the rubella virus genome exhibited the transition over transversion preference that has been well documented in DNA genomes (Meyer *et al.*, 1999; Salemi & Vandamme, 2003). This preference has been attributed to the facts that transitions are more likely to lead to silent mutations in amino acid

sequence and that, during replication, it is more likely that a mutation to a nucleotide of equal size (transition) will occur than to a nucleotide of different size (transversion). In RNA genomes, the possibility of both G–C and G–U pairing would also favour transitions in the replication process. Interestingly, pyrimidine transitions were favoured over purine transitions by a ratio (Y/R) of 2.7 across the entire genome; Y/R varied from 0.9 to 3.7 in genomic regions. Within the HVR, the most variable region of the genome, both K and Y/R were lower than for the entire genome and most of the other genomic regions, indicating that the variability in this region was generated in part by relaxing of the genomic preference for pyrimidine transitions over transversions and purine transitions.

Phylogenetic analysis of rubella viruses has traditionally been done on the basis of E1 gene or subE1 gene sequences and a standard taxonomy was proposed recently, based on a window within the E1 gene, that was substantiated by using complete SP-ORF sequences (WHO, 2005). The second goal of this study was to extend phylogenetic analysis to the 5' region of the genome and we found that generally comparable trees, in terms of both overall variability and phylogenetic clustering, were generated with sequence windows in the NSP-ORF. The exception was a group of seven genotype 1B viruses that formed a branch in a tree based on NP sequence, but formed two branches on the basis of JR sequence. Intriguingly, a deletion in the junction region of five of these seven viruses did not segregate with the two phylogenetic branches on the JR tree. Analysis revealed a recombinational event, putatively near the 5' end of the C gene, that led to the generation of the two branches on the JR tree. There was one previous report of a natural recombination event in *Rubella virus* (in the E1 gene; Zheng *et al.*, 2003a), but the origin of the recombinant strain was in doubt because one of the parents was related closely to a commonly used laboratory strain. Thus, this was the first conclusive evidence of rubella virus recombination in nature.

Interestingly, the E1 sequences of the seven genotype 1B viruses did not cluster on the E1-based tree and, in comparison with the NP- and JR-based trees, this could be due to divergence or additional recombinational events. As can be seen in the tree in Fig. 1, genotype 1B consists of two sub-branches that would not necessarily appear to be related if fewer sequences were employed (e.g. the E1-based tree in Fig. 4). It is also to be noted that all of the WHO reference strains are on one of these sub-branches. Thus, for this genotype, phylogenetic analysis using sequences from the NSP-ORF region of the genome could be useful in assessing relatedness.

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Editor-Communicated Paper

Tandem Repeats of Lactoferrin-Derived Anti-Hepatitis C Virus Peptide Enhance Antiviral Activity in Cultured Human Hepatocytes

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Abstract: Previously, we found that bovine and human lactoferrin (LF) specifically inhibited hepatitis C virus (HCV) infection in cultured non-neoplastic human hepatocyte-derived PH5CH8 cells, and we identified 33 amino acid residues (termed C-s3-33; amino acid 600–632) from human LF that were primarily responsible for the binding activity to the HCV E2 envelope protein and for the inhibiting activity against HCV infection. Since the anti-HCV activity of C-s3-33 was weaker than that of human LF, we speculated that an increase of E2 protein-binding activity might contribute to the enhancement of anti-HCV activity. To test this possibility, we made two repeats [(C-s3-33)₂] and three repeats [(C-s3-33)₃] of C-s3-33 and characterized them. Far-Western blot analysis revealed that the E2 protein-binding activities of (C-s3-33)₂ and (C-s3-33)₃ became stronger than that of the C-s3-33, and that the binding activity of (C-s3-33)₂ was stronger than that of (C-s3-33)₃. Using an HCV infection system in PH5CH8 cells, we demonstrated that the anti-HCV activities of (C-s3-33)₂ and (C-s3-33)₃ became stronger than that of the C-s3-33. Furthermore, using a recently developed infection system with a VSV pseudotype harboring the green fluorescent protein gene and the native E1 and E2 genes, we demonstrated that the antiviral activities of (C-s3-33)₂ and (C-s3-33)₃ were stronger than that of C-s3-33. These results suggest that tandem repeats of LF-derived anti-HCV peptide are useful as anti-HCV reagents.

Key words: Hepatitis C virus, Lactoferrin, Anti-HCV peptide, E2 protein-binding activity

Hepatitis C virus (HCV) infection frequently causes chronic hepatitis, which progresses to liver cirrhosis and hepatocellular carcinoma (28). HCV is an enveloped positive single-stranded RNA (9.6 kb) virus belonging to the *Flaviviridae*. The HCV genome encodes a large polyprotein precursor of about 3,000 amino acids (aa), which is cleaved by the host and viral proteases into at least 10 proteins: the core, envelope 1 (E1), E2, p7, and non-structural 2 (NS2), NS3, NS4A, NS4B, NS5A, and NS5B (7, 8, 18). These HCV proteins function not only in virus replication but may also

affect a variety of cellular functions, including gene expression, signal transduction, and apoptosis (1, 17).

Approximately 170 million people worldwide are infected with HCV (32). The combination of a pegylated interferon with ribavirin is the current standard therapy for chronic hepatitis C and yields a sustained virological response rate of about 55% (6). This means that about 45% of patients with chronic hepatitis C are still threatened by the progress of the disease to cirrhosis and hepatocellular carcinoma.

Although the entry mechanism of HCV remains unclear, to date, several candidates for HCV receptors

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Abbreviations: aa, amino acids; DMEM, Dulbecco's modified Eagle's medium; E2, envelope 2; GFP, green fluorescent protein; HCV, hepatitis C virus; LF, lactoferrin; MBP, maltose-binding protein; NS2, non-structural 2; TF, transferrin; VSV, vesicular stomatitis virus.

have been reported: CD81, the scavenger receptor class B type I, the mannose-binding lectins DC-SIGN and L-SIGN, low-density lipoprotein receptors, etc. (4). Most of them have been identified as interacting materials with a soluble and truncated form of the HCV E2 protein, because of the lack of efficient HCV proliferation in cell cultures, although several culture systems using PCR for detection of HCV infection have been reported (20). However, a major advance in investigating HCV entry has been achieved by the development of pseudotype viruses bearing HCV E1 and E2 proteins assembled onto retrovirus particles (2, 9) or vesicular stomatitis virus (VSV) particles (3, 23, 30). Extensive characterization of the pseudotype viruses has shown that these mimic the early steps of the HCV life cycle. This system has allowed the study of the role of candidate receptors in the early steps of HCV infection (4).

We previously found that bovine and human lactoferrins (LFs) specifically prevented HCV infection in cultured human non-neoplastic hepatocyte PH5CH8 cells using the PCR method for detection of HCV infection (10, 12). Regarding these findings, some clinical studies have demonstrated that monotherapy with bovine LF improves the serum HCV RNA and/or alanine aminotransferase levels in patients with chronic hepatitis C (15, 16, 27, 31).

LF is an 80-kDa mammalian iron-binding glycoprotein and consists of two homologous globular lobes (an N-lobe and a C-lobe), each with a single iron (Fe^{3+}) binding site. It is structurally related to the plasma iron-transport protein transferrin (TF). LF's biological roles include activities in the host defense mechanism as well as in iron metabolism (21, 22). Unlike TF, LF is a primary defense protein against microbial infection. LF possesses strong bacteriostatic and bactericidal activities against pathogenic bacteria, as well as inhibitory activity against pathogenic viruses (5, 21, 22, 33).

LF's preventive mechanism against HCV infection has been thought to be the direct interaction between LF and HCV; indeed, by Far-Western blot analysis using thioredoxin-fused LF fragments expressed in *Escherichia coli* (*E. coli*) and the soluble E2 protein expressed in Chinese hamster ovary cells, we demonstrated that the 93 carboxyl aa of LF (human, bovine, and horse), termed C-s3, specifically bound to the E2 protein (25). On the other hand, Yi et al. (34) independently reported that the E1 and E2 proteins could bind to human and bovine LFs, although the binding region of LF was not identified. Furthermore, we identified the 33 aa of human LF (termed C-s3-33; aa 600–632), which was primarily responsible for the E2 protein-binding activity, and demonstrated that maltose-binding protein (MBP)-fused C-s3-33 prevented HCV infection

in PH5CH8 hepatocyte cells (25). However, the E2 protein-binding activity and the anti-HCV activity of C-s3-33 were obviously weaker than those of human LF. Therefore, we presumed that the increase of the E2 protein-binding activity would lead to the enhancement of anti-HCV activity.

To evaluate this idea, we made tandem repeats of C-s3-33, and compared their E2 protein-binding activities and anti-HCV activities with those of the C-s3-33. Here, we report our findings that the anti-HCV activity of the tandem repeats were stronger than that of the monomer when accompanied by the enhancement of the E2 protein-binding activity, by analyses using not only the HCV infection system but also the infection system of a VSV pseudotype bearing the native forms of HCV E1 and E2 proteins.

Materials and Methods

Cell cultures. Simian virus 40 large T antigen-immortalized non-neoplastic human PH5CH8 hepatocytes were maintained as described previously (11, 24). Human hepatoblastoma HepG2 cells were maintained in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum.

Construction of expression plasmids for *E. coli*. The pMAL-c2X (hLF600–632) (25) expression plasmid for the MBP-fused C-s3-33 LF fragment, was used as a template for the PCR using a primer set of hLFB6 5'-TGATAGGATCCGTTGGTGTCTCGGATGATAAGG-3' containing the *Bam*HI recognition site (underlined) (25) and 632R6A 5'-ATCCATCCGAGACACCA-CAAACTTGTCCGGGCAGTCAGATCC-3' containing an extra 18 nts (underlined) encoding the amino-terminal 6 aa of the C-s3-33 LF fragment. After PCR (20 cycles) using KOD-plus DNA polymerase (Toyobo, Osaka, Japan), the amplified PCR product was used as a template for a second PCR using the primer set of hLFB6 and 632R 5'-TAATAAAGCTTT-TAAAACCTTGTCCGGGCAGTCAGATCC-3' containing the *Hind*III recognition site (underlined) (25). After PCR (35 cycles) using KOD-plus DNA polymerase, amplified PCR products (approximately 200 bp for the two-repeat form and approximately 300 bp for the three-repeat form) were subcloned into the *Bam*HI and *Hind*III sites of pMAL-c2X, and were used as expression plasmids for the production of the MBP-fused (C-s3-33)₂ and (C-s3-33)₃.

To prepare an expression plasmid for the production of the MBP-fused C-s3-33-relevant fragment (aa 587–619) of human TF, pCXbsr/huTF (29) encoding full-length human TF was used as a template for the PCR using a primer set of hTF587F 5'-TGATAG-

GATCCGTGGTCACACGG-3' containing the *Bam*HI recognition site (underlined) and hTF619R 5'-TAATAAAGCITTTTAAAAGTTGCCCG-3' containing the *Hind*III recognition site (underlined). After PCR (35 cycles) using KOD-plus DNA polymerase, the amplified PCR product was subcloned into the *Bam*HI and *Hind*III sites of pMAL-c2X, and was used as the expression plasmid.

Expression and purification of the MBP-fused protein. Expression and purification of the MBP-fused LF fragment [C-s3-33, (C-s3-33)₂, or (C-s3-33)₃] or the MBP-fused C-s3-33-relevant fragment of human TF were carried out as described previously (25). Briefly, the expression plasmid for MBP-fused protein was transformed into the *E. coli* strain JM109. The transformants were cultured at 37 °C for several hours, and the harvested cells were sonicated. After removal of insoluble cellular debris by centrifugation, the supernatant obtained as the soluble fraction was applied onto an amylose resin affinity column (New England Biolabs) to obtain the MBP-fused protein. The purity of the obtained MBP-fused protein was evaluated to be more than 95% by electrophoresis on 10% SDS-PAGE gels. The concentration of the purified MBP-fused protein was determined by using Coomassie protein assay reagent (Pierce). The MBP2 (43 kDa) produced from the pMAL-c2X with a stop codon inserted into the *Xmn*I site was used as a control protein.

Far-Western blot analysis. Far-Western blot analysis was carried out as described previously (25). Briefly, 0.5 µg of human LF, MBP2, and MBP-fused LF fragments were resolved by 10% SDS-PAGE and transferred to polyvinylidene difluoride membranes. After blocking with N-buffer (25), a binding reaction was carried out using the secreted form of the E2 protein (E2-681) consisting of aa 384–681 expressed in Chinese hamster ovary cells as a probe (14), and then rat monoclonal antibody, MO-12 (13), against E2 protein was used for the detection of E2 protein-bound MBP-fused LF fragments.

Assay for anti-HCV activity of MBP-fused protein. An assay for anti-HCV activity of the MBP-fused LF fragment was carried out by the method described previously (25). Briefly, 2 µl (2 × 10⁴ HCV) of the HCV-positive serum HCV-O (previously described as 1B-2 (19)) (genotype 1b) and the MBP-fused LF fragment (final concentration, 0.5, 1.0, and 2.0 mg/ml) were pre-incubated for 60 min at 4 °C and then inoculated onto the PH5CH8 cells (1.5 × 10⁴ cells were cultured for 2 days before viral inoculation on a 96-well plate). After incubation of the cells for 90 min at 37 °C, the cells were washed three times with PBS and further cultured for 1 day at 32 °C. Cellular RNA (0.5 µg) prepared by

ISOGEN extraction kit (Nippon Gene Co., Toyama, Japan) was used for the quantitative analysis of HCV RNA using LightCycler PCR as described previously (26). As the positive and negative controls for anti-HCV activity, human LF and MBP2, respectively, were used.

Assay for anti-VSV pseudotype activity of MBP-fused protein. For this assay, the VSV pseudotype VSVΔG*(HCV), bearing the native forms of HCV E1 and E2 proteins from the O strain (19), was used. VSVΔG*(HCV) was prepared by introducing the native form of E1 and E2 proteins into recombinant VSV, VSVΔG*, which harbors the green fluorescent protein (GFP) gene instead of the VSV G envelope protein gene (30). An assay for the anti-VSV pseudotype activity of the MBP-fused LF fragment was carried out by a method described previously (30). Briefly, VSVΔG*(HCV) (Approximately 100 IU/assay) was pre-incubated with the MBP-fused LF fragment (final concentration, 0.1–1.0 mg/ml) at 37 °C for 60 min and inoculated onto PH5CH8 or HepG2 cells (1.5 × 10⁴ cells were cultured for 2 days before viral inoculation on a 96-well plate). After incubation of the cells for 90 min at 37 °C, the cells were washed with DMEM three times and incubated with fresh culture medium. VSVΔG*G was used as a control in this assay. After 24 hr of incubation, each infectious titer was determined by counting the number of GFP-expressing cells under a fluorescence microscope. As the positive and negative controls for the assay, human LF and MBP2 were used, respectively. Human TF and an MBP-fused C-s3-33-relevant fragment of human TF were also used for the assay.

Results

Two and Three Repeats of the Human LF Fragment (C-s3-33) Strengthened the E2 Protein-Binding Activity

Previously we found that bovine and human LFs prevented HCV infection in PH5CH8 cells via direct interaction between LF and HCV (10, 12), and we further identified 33 aa residues (C-s3-33; aa 600–632 of human LF) as an essential and minimum domain possessing binding activity for the HCV E2 protein (secreted form consisting of aa 384–681) and inhibiting activity against HCV infection (25). This result suggested that the E2 protein-binding activity contributes to the anti-HCV activity. However, the E2 protein-binding activity of C-s3-33 was somewhat weaker than that of human LF (25), and the anti-HCV activity of C-s3-33 (IC₅₀ = 20 µM) in the infection system using PH5CH8 cells was also weaker than that of human LF (IC₅₀ = 5 µM) (25). To improve these points, we first tried to

enhance the E2 protein-binding activity of C-s3-33 by the multiplication of C-s3-33. Initially, we made pMAL-c2X-based expression vectors encoding two, three, and four repeats of C-s3-33 as MBP-fused proteins, and then expressed them in *E. coli*. We successfully purified two repeats (C-s3-33)₂ and three repeats (C-s3-33)₃ of C-s3-33 as soluble forms of the MBP-fused protein; the purification of the four repeats of C-s3-33 failed due to problems with solubility. Using the

MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃, we performed Far-Western blot analysis to compare their E2 protein-binding activities. The result revealed that the E2 protein-binding activities of (C-s3-33)₂ and (C-s3-33)₃ became stronger than that of the C-s3-33, and the binding activity of (C-s3-33)₃ was stronger than that of (C-s3-33)₂ (Fig. 1). Although the E2 protein-binding activity of C-s3-33 was weaker than that of human LF, the binding activities of (C-s3-33)₂ and (C-s3-33)₃

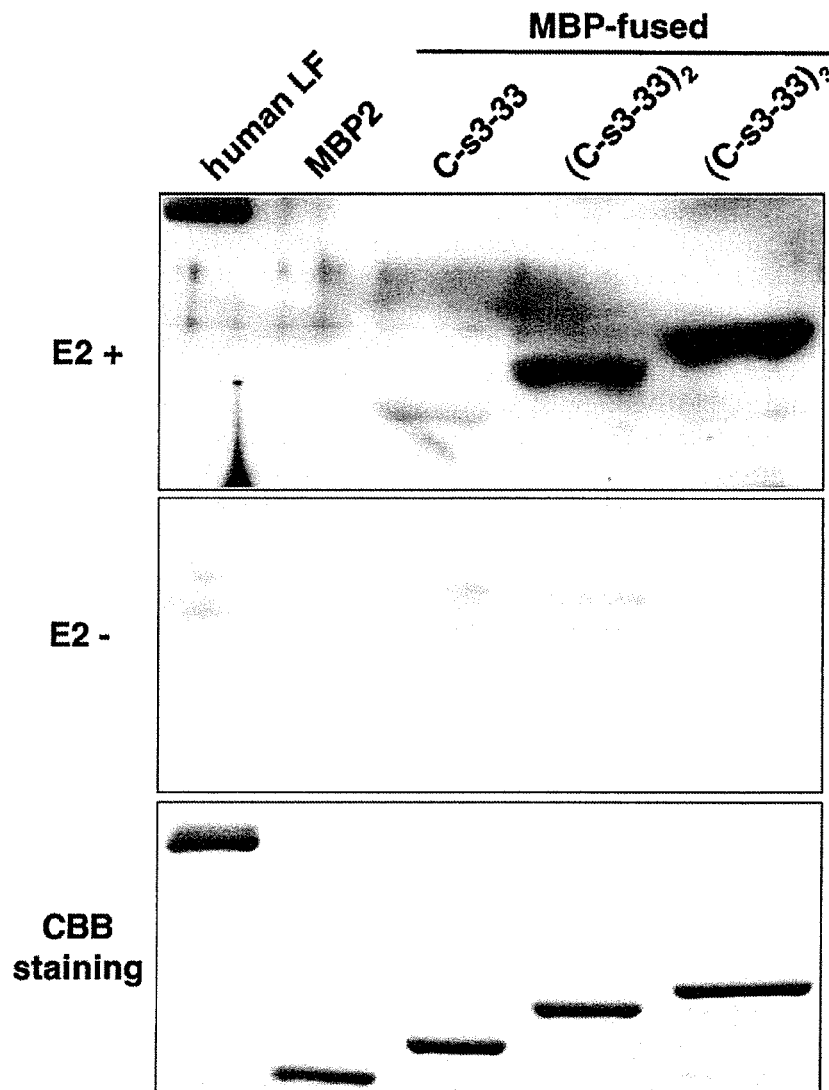


Fig. 1. Comparison of the E2 protein-binding activities of MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃. MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ (0.5 μ g each) were resolved by 10% SDS-PAGE. Human LF and MBP2 (0.5 μ g each) were also used for the assay as control materials. Far-Western blot analysis using the E2 protein expressed in Chinese hamster ovary cells (14) as a probe was performed as described under "Materials and Methods." Rat monoclonal antibody MO-12 (13) against the E2 protein was used for the detection of the E2 protein bound to MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃, as well as human LF. Far-Western blot analysis in the absence of the E2 protein was also performed. The bottom panel shows the results for human LF, MBP2, and MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ detected by staining with Coomassie Brilliant Blue.

became comparable with that of human LF (Fig. 1). To exclude the possibility of cross-reactions between C-s3-33 and the anti-E2 antibody, we performed a Far-Western blot analysis in the absence of the E2 protein. No significant bands were obtained in this control experiment (Fig. 1). The Far-Western blot analysis using normal rat serum instead of anti-E2 antibody also detected no significant bands (data not shown). These results suggest that the specific E2 protein-binding activities of (C-s3-33)₂ and (C-s3-33)₃ increase with the degree of multiplication of C-s3-33.

(C-s3-33)₂ and (C-s3-33)₃ Efficiently Prevented HCV Infection in PH5CH8 Cells

Since we obtained the expected results that the E2-binding activities of (C-s3-33)₂ and (C-s3-33)₃ were stronger than that of C-s3-33, we next compared their anti-HCV activities in our HCV infection system using PH5CH8 cells (10, 25). The obtained result (Fig. 2)

revealed that the anti-HCV activities of (C-s3-33)₂ and (C-s3-33)₃ (IC₅₀ = 10 μM in both) became stronger than that of the C-s3-33 (IC₅₀ = 23 μM), although their activities were somewhat weaker than that of human LF (IC₅₀ = 5 μM). These results support the previous suggestion that the E2 protein-binding activity of C-s3-33 contributes to the inhibition of HCV infection (inoculum HCV-O) in human hepatocyte cells (25). However, in our HCV infection system, we failed to clearly show a difference in inhibiting activities between (C-s3-33)₂ and (C-s3-33)₃, because each standard deviation became somewhat large value due to the low level of cell culture-based HCV infection (20, 25, 31). In order to improve this point, we developed an infection system with VSVAG*(HCV), a VSV pseudotype bearing the native E1 and E2 proteins derived from HCV-O (30), and this VSV pseudotype was used for further analysis as described below.

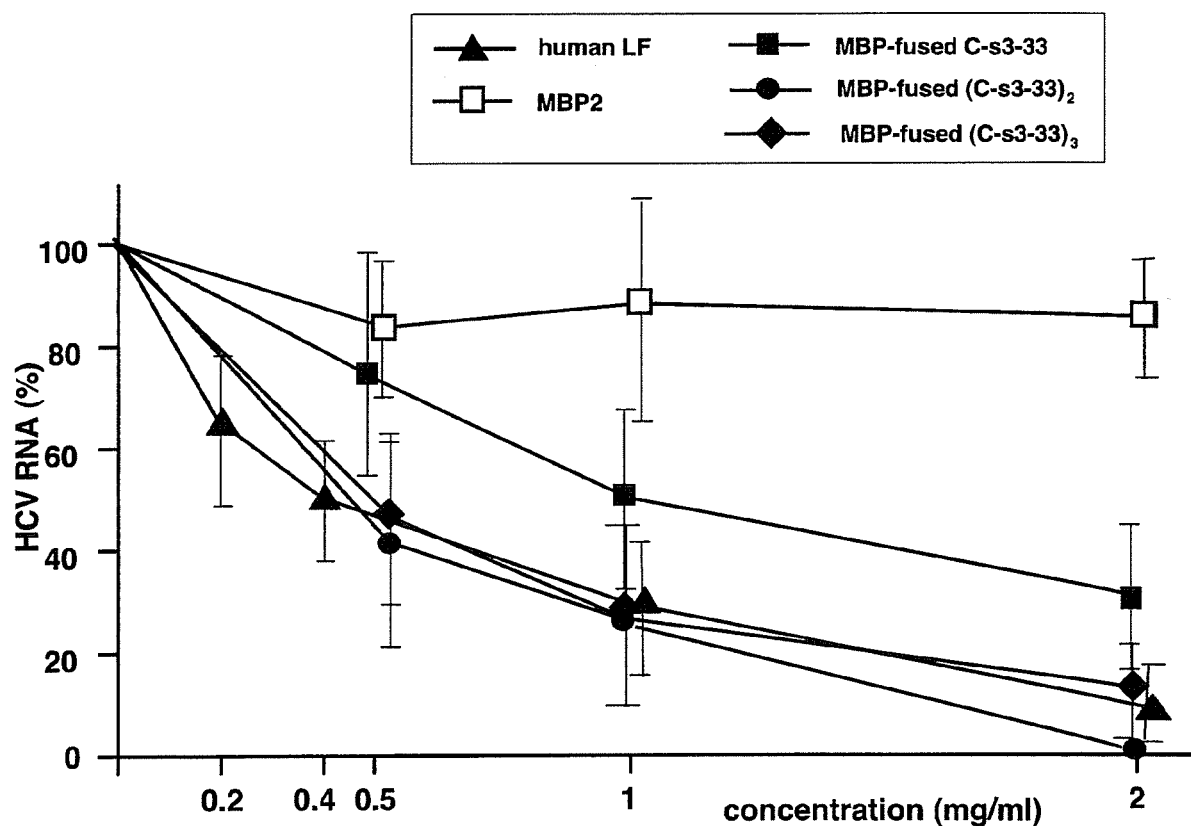


Fig. 2. Anti-HCV activities of MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ in an HCV infection system using PH5CH8 cells. PH5CH8 cells and the inoculum HCV-O were used for the HCV-inhibiting assay, as described under "Materials and Methods." The number in the ordinate axis indicates the percent of HCV RNA determined by real-time LightCycler PCR (26). Approximately 2,000 copies of HCV RNA per μg of cellular RNA were reproducibly obtained using this HCV infection system (10, 26). In addition to the MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃, human LF and MBP2 were also used for the assay as control materials. The data are means ± SD of triplicates from three independent experiments.

Antiviral Effects of (C-s3-33)₂ and (C-s3-33)₃ against VSVΔG(HCV) Infection in PH5CH8 Cells*

Since PH5CH8 cells showed good susceptibility to our developed VSV pseudotype, VSVΔG*(HCV) (30), we examined the antiviral effects of (C-s3-33)₂ and (C-s3-33)₃ against VSVΔG*(HCV) infection in PH5CH8 cells, and compared them with those of the C-s3-33 and human LF. In this experiment, the antiviral effects of human TF and a C-s3-33-relevant fragment of human TF were also examined. The results (Fig. 3) clearly showed that human LF (IC₅₀=0.6 μM) strongly inhibited VSVΔG*(HCV) infection, but that human TF and the C-s3-33-relevant fragment of human TF did not, nor did MBP2, suggesting that inhibition against VSVΔG*(HCV) infection also occurred in an LF-specific manner as observed previously in the HCV infection system (25, 31). These results support previous findings (23, 30) using the VSV pseudotype infection

system. Furthermore, we obtained clear results that C-s3-33 showed inhibiting activity against VSVΔG*(HCV) infection, and that its inhibiting activity was increased with multiplication of C-s3-33. The IC₅₀ doses of C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ were 17 μM, 5.0 μM, and 3.0 μM, respectively. This result indicates that antiviral activity of C-s3-33 is improved by the duplication and triplication of C-s3-33, although the antiviral activity of (C-s3-33)₃ is still weaker than that of human LF. We confirmed that these LF fragments did not inhibit VSVΔG*G (control virus without E1 and E2 proteins) infection in PH5CH8 cells (data not shown). In summary, our results suggest that direct interaction of the C-s3-33 fragment with the E2 protein in VSVΔG*(HCV) prevents the virus infection in PH5CH8 cells.

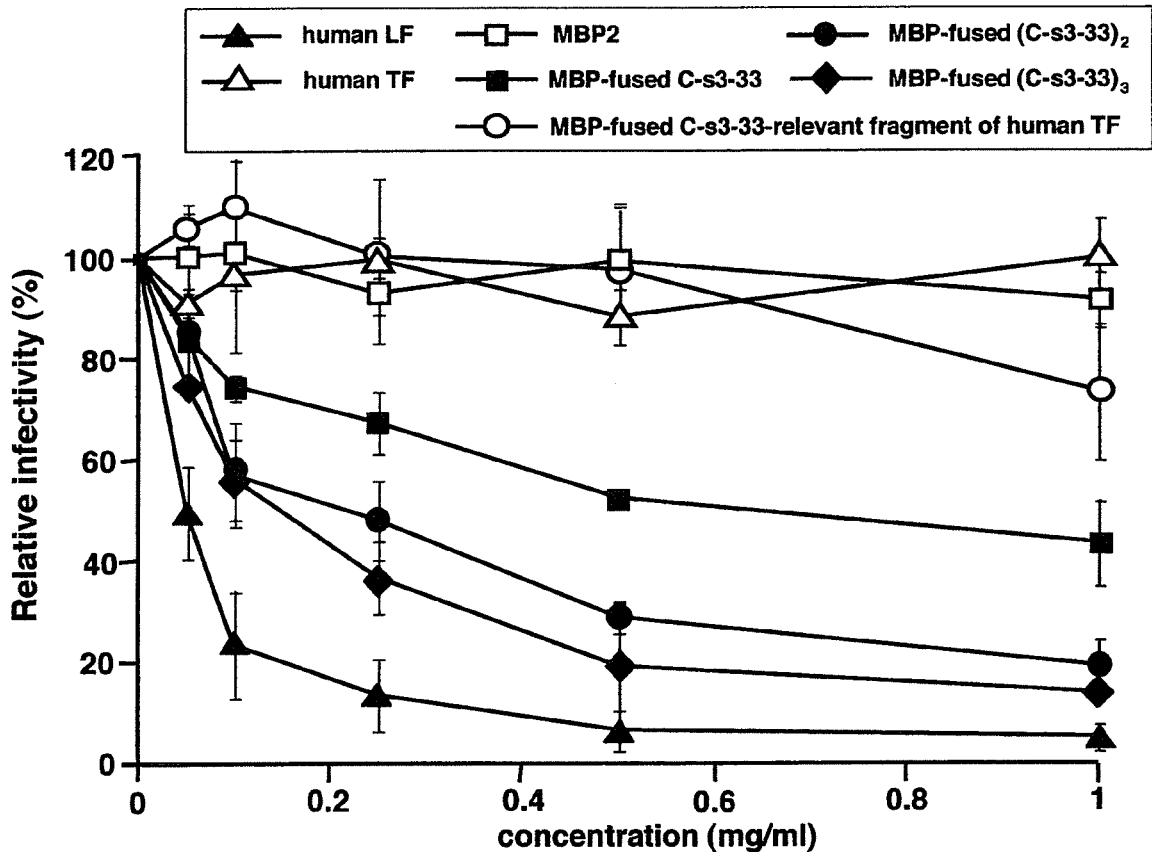


Fig. 3. Antiviral activity of the MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ in the infection system of pseudotype virus using PH5CH8 cells. PH5CH8 cells and the VSV pseudotype, VSVΔG*(HCV), were used for the HCV-inhibiting assay, as described under "Materials and Methods." The number in the ordinate axis indicates the relative infectivity (%) calculated by counting GFP-positive cells. Approximately 100 GFP-positive cells per one assay were reproducibly obtained using this pseudotype infection system (30). In addition to the MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃, human LF, human TF, MBP2, and an MBP-fused C-s3-33-relevant fragment of human TF were also used for the assay as controls. The data are means ± SD of three independent experiments.

Antiviral Effects of (C-s3-33)₂ and (C-s3-33)₃ against VSVΔG(HCV) Infection in HepG2 Cells*

We have shown the inhibiting activities of LF fragments against HCV infection or VSV pseudotype infection in PH5CH8 cells; however, it is not clear whether or not the LF fragments used in this study show inhibiting activities against virus infection in cells other than PH5CH8 cells. To clarify this point, HepG2 cells were used for the analysis, because HepG2 cells showed the highest susceptibility to VSVΔG*(HCV) among 25 cell lines examined (30). As a consequence, we obtained similar results (Fig. 4) with those obtained in the infection system using PH5CH8 cells. The IC₅₀ doses of C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ were >12 μM, 7.6 μM, and 3.9 μM, respectively, indicating that, again, the inhibiting activity was increased with multiplication of C-s3-33, although antiviral activity of (C-s3-33)₃ was still weaker than that of human LF (IC₅₀=1.2 μM). In conclusion, our results indicated that tandem repeats of

C-s3-33 enhanced the inhibiting activity in cell culture-based HCV infection.

Discussion

In our previous (30) and present studies, we showed that pretreatment of VSV pseudotypes with bovine and human LFs reduced the infectivity of VSVΔG*(HCV) and VSVΔG*(E2) bearing only the E2 protein in a dose-dependent manner, whereas pretreatment with TF did not. In contrast, LFs partially inhibited the infectivity of VSVΔG*(E1) bearing only the E1 protein (30). These results suggested that the interaction of LF and the E2 protein is the main contributing factor to the prevention of HCV infection. This idea has been strongly supported by the results obtained in this study. We demonstrated that tandem repeats of C-s3-33, an anti-HCV peptide derived from human LF, enhanced the E2 protein-binding activity and the inhibiting activity

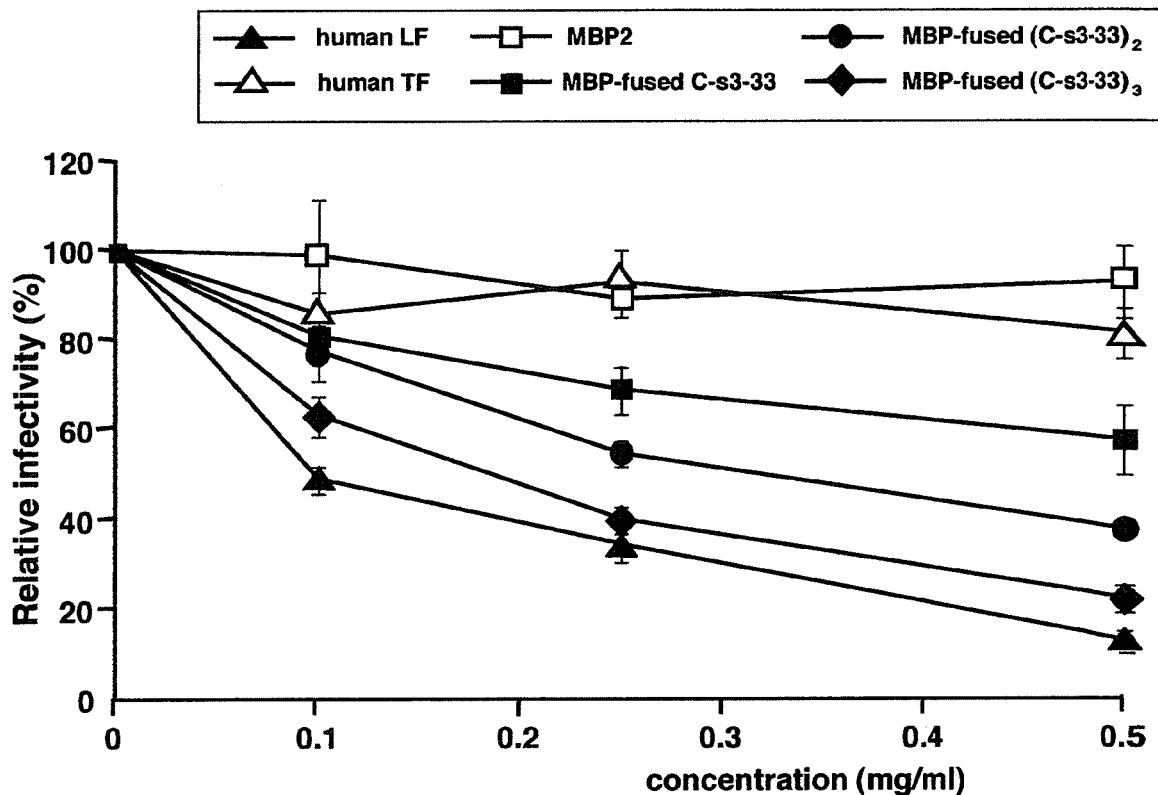


Fig. 4. Antiviral activity of the MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃ in the infection system of the pseudotype virus using HepG2 cells. HepG2 cells and the VSV pseudotype, VSVΔG*(HCV), were used for the HCV-inhibiting assay, as described in "Materials and Methods." The number in the ordinate axis indicates the relative infectivity (%) calculated by counting GFP-positive cells. Approximately 100 GFP-positive cells per one assay were reproducibly obtained using this pseudotype infection system (30). In addition to the MBP-fused C-s3-33, (C-s3-33)₂, and (C-s3-33)₃, human LF, human TF, and MBP2 were used for the assay as controls. The data are means ± SD of three independent experiments.

against infection by HCV or the VSV pseudotype, VSVΔG*(HCV), in human hepatic cell lines. These results strongly suggest that the direct interaction between C-s3-33 and the E2 protein plays a central role in the inhibition of HCV infection by LF.

Since C-s3-33 or repeated forms of C-s3-33 could prevent HCV and VSVΔG*(HCV) infection, C-s3-33 must bind to a region other than the region (aa 441–500 of E2 protein) required for heteromeric complex formation between E1 and E2 proteins. Our preliminary results suggested that the C-s3-33 bound to aa 411–500 and aa 600–661 of the E2 protein, indicating that the target sites of C-s3-33 may be plural. This result suggests a rather complex interaction between C-s3-33 and the E2 protein. To clarify this point, further comprehensive analysis will be needed.

Although tandem repeats of C-s3-33 enhanced the anti-HCV activity compared with that of the C-s3-33, the fact that their antiviral activities were still several-fold weaker than that of original human LF remains a subject to be resolved. As one approach to increase anti-HCV activity, tandem repeats of C-s3-33-relevant fragment of bovine LF may be useful, because we previously observed that the anti-HCV activity of bovine LF ($IC_{50}=1.5\ \mu M$) was stronger than that of human LF ($IC_{50}=5.0\ \mu M$) (26), and that the E2 protein-binding activity of the C-s3 (93 aa)-relevant fragment of bovine LF was stronger than that of C-s3 (25). Since 10 aa out of 33 aa differ between C-s3-33 and its relevant fragment of bovine LF, some aa substitutions between both fragments may help to further increase the anti-HCV activity of LF-derived peptides. Alternatively, some spacer between the C-s3-33 repeats may be needed. Therefore, further trials will be needed to achieve the maximum anti-HCV activity of C-s3-33.

In conclusion, the results of the present study demonstrated that tandem repeats of human LF-derived 33 aa prevented HCV infection more strongly than the 33 aa, and suggest that this repeated form will be useful as a novel anti-HCV reagent.

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