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# Successful direct-acting antiviral treatment of three patients with genotype 2/1 recombinant hepatitis C virus

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#### ABSTRACT

There have been a few reports on the treatment of patients infected with recombinant hepatitis C virus (HCV) genotype 2/1 strains with direct-acting antivirals (DAAs). We experienced three patients, with genotype 2/1 recombinant HCV, treated with DAAs successfully. The first, a 39-year-old man, was infected with recombinant HCV genotype 2a/1b, a rare variant. The sequence of the relapsed virus showed chimeric HCV 2a/1b with the recombinant breakpoint found at nucleotide +49 from the start of the NS3 region. Sofosbuvir plus ribavirin, a regimen recommended for HCV genotype 2, did not lead to a sustained viral response (SVR). Retreatment with grazoprevir plus elbasvir resulted in an SVR. The second case, a 70-year-old woman, was infected with recombinant HCV genotype 2b/1b. DAA therapy with sofosbuvir plus ledipasvir resulted in an SVR. The third case, a 48-year-old woman, was also infected with recombinant HCV genotype 2b/1b. DAA therapy with daclatasvir plus asunaprevir resulted in an SVR. The baseline sequences of the viruses from both the second and third cases showed chimeric HCV 2b/1b with the recombinant breakpoint found at

nucleotide +10 from the NS3 start. We report three cases with 2/1 chimeras and discuss

the prevalence and response to therapy.

Key words: Direct-acting antivirals, Recombinant hepatitis C virus

#### **INTRODUCTION**

Hepatitis C virus (HCV) infection is a global problem [1]. With the introduction of direct-acting antivirals (DAAs), approximately 95% of naïve patients with HCV can achieve a sustained virological response (SVR) [2]. Before the development of pangenotypic DAAs, the selection of an HCV-genotype-oriented DAA regimen was recommended. However, roughly 14~25% of patients in Germany and Israel with HCV genotype 2 have a recombinant (chimeric) HCV strain, which consists of two types of HCV [3]. There are few reports on the efficacy of DAA therapy for such patients [4]. In Japan, HCV typing is performed using one or two of the three available methods. The virus is serotyped as type 1 or 2 using an enzyme-linked immunosorbent assay with type-specific antibodies against the nonstructural (NS) 4 region [5]. Genotyping is determined by PCR with genotype-specific primers for the core region [6], or direct sequencing of the NS5B region [7]. The discrepancy in typing using the three methods suggests the possibility of recombinant HCV infection. It was reported that 1~2% of the patients with genotype 2 in Japan were infected with recombinant HCV [8].

We identified the nucleotide breakpoint of the recombinant HCV in three patients who achieved an SVR to interferon-free DAA regimens.

#### **Case presentation**

#### Case 1

A 39-year-old man had failed interferon-based therapy for chronic hepatitis C (**Table**). He had a hetero-type of *interleukin 28B* single-nucleotide polymorphism. He had no history of surgery, blood transfusion, illegal drug use, or tattooing. The HCV genotype was determined as 2a using PCR of the core region. Combination therapy with sofosbuvir 400 mg/day and ribavirin 800 mg/day was initiated in 2015 at previous hospital. The serum HCV RNA fell rapidly from 6.4 log<sub>10</sub> IU/mL at baseline to below the detection limit of the COBAS TaqMan assay (1.2 log<sub>10</sub> IU/mL) at 4 weeks. Although the viral load was below the detection level during treatment, HCV relapsed 4 weeks after the end of treatment.

He moved because of his job and was referred to our hospital in 2016. Our screening test for HCV-infected patients showed that his HCV serotype was 1.

Consequently, we examined the core, NS5A, and NS5B regions to determine the HCV genotype, which was determined to be 2a by PCR of the core region, but as 1b by sequencing NS5A and NS5B. After confirming that there was no substitution associated with resistance in the NS5A region, we started retreatment with grazoprevir 100 mg/day and elbasvir 50 mg/day. The viral load fell rapidly from 7.0 log<sub>10</sub> IU/mL at baseline to  $< 1.2 \log_{10}$  IU/mL at 4 weeks and an SVR at 12 weeks (SVR<sub>12</sub>) was achieved.

#### HCV 2a/1b recombinant breakpoint in Case 1

We assumed that the patient was infected with a recombinant HCV strain, which consisted of genotype 2a in the first half of the viral genome and genotype 1b in the second half. Because the recombinant sites were often identified around the NS2/NS3 junction in previous studies [3, 9, 10], we designed primer pairs to amplify a 438-bp fragment covering the NS2/NS3 junction. The HCV sequences amplified before and after sofosbuvir treatment both aligned with the HCV2a\_HC-J6 and HCV1b\_Con1 strains. The recombinant breakpoint was found at nucleotide position +49 relative to the start of the NS3 region (**Figure 1**).

#### Case 2

The patient was a 70-year-old woman with chronic hepatitis C (**Table**). She had undergone an appendectomy at the age of 48. She had no history of blood transfusion or anti-HCV therapy. The HCV serotype was determined as 1 and the genotype as 2b by PCR of the core region. Therefore, we checked the genotype by sequencing NS5A and NS5B. This showed that her HCV genotype was 2b by PCR of the core region, but as 1b by sequencing NS5A and NS5B. Combination therapy with sofosbuvir 400 mg/day and ledipasvir 90 mg/day was initiated in 2016. The viral load fell rapidly from 6.2  $log_{10}$  IU/mL at baseline to < 1.2  $log_{10}$  IU/mL at 4 weeks and SVR<sub>12</sub> was achieved.

#### Case 3

The patient was a 48-year-old woman with HCV-related cirrhosis (**Table**). She had no history of surgery, blood transfusion, illegal drug use, tattooing, or anti-HCV therapy. HCV serotype was determined to be 1 and genotype as 2b by PCR of the core region. Therefore, we checked the genotype by sequencing NS5A and NS5B. This

determined that her HCV genotype was 2b by PCR of the core region, but as 1b by sequencing NS5A and NS5B. Combination therapy with daclatasvir 60 mg/day and asunaprevir 200 mg/day was initiated in 2014, after confirming that there was no substitution associated with resistance in the NS5A region. The viral load fell from 6.5  $log_{10}$  IU/mL at baseline to < 1.2  $log_{10}$  IU/mL at 12 weeks and SVR<sub>12</sub> was achieved. Of interest, her daughter was also infected with HCV and the serotype was determined to be 1 and the genotype as 2b by PCR of the core region.

#### HCV 2b/1b recombinant breakpoint in Cases 2 and 3

We assumed that the patients were infected with recombinant HCV strains in which the first half was genotype 2b and the second half was genotype 1b. Both HCV strains were aligned with the HCV2b\_HC-J8 and HCV1b\_Con1 strains. In both cases, the recombinant breakpoint was found to be at nucleotide position +10 relative to the NS3 start (**Figure 2**).

#### DISCUSSION

We treated three patients with genotype 2 and 1 recombinant HCV strains using interferon-free DAAs. One patient (Case 1), who was initially treated with the DAA regimen recommended for genotype 2, did not achieve an SVR, but the DAA regimen recommended for genotype 1 led to an SVR in all three patients: as the initial DAA treatment in Cases 2 and 3 and retreatment in Case 1.

In our hospital, the HCV type is determined using NS4 serotyping and PCR with genotype-specific primers for the core region. Of the 887 patients treated with DAA therapy between September 2014 and July 2017, three (0.3%) patients had typing results that differed with the two methods. Further examination by direct sequencing of NS5B showed that the serotyping results matched the genotypes. To our knowledge, the HCV recombinant viruses in previous studies consisted of the structural region of genotype 2 and the NS4 to NS5 region of genotype 1 [3, 10]. We identified the recombinant breakpoint of the HCV genotype 2/1b in all cases. The breakpoint in Case 1 was located +49 nucleotides from the NS3 start, which was the same position reported by Susser *et al.* (**Figure 1**) [3]. The breakpoint in Cases 2 and 3 was at the same position at +10 nucleotides (**Figure 2**). In previous reports, the breakpoint of 2/1 chimera HCV was located within 80 acids of the NS2/NS3 junction [10]. The prevalence and characteristics of chimeric HCV differ regionally. The first report was on an HCV genotype 2k/1b chimera derived from a Russian patient in 2001 [11]. The original 2k/1b chimera is prevalent (14~25%) in Russia, Georgia, Germany, and Israel [3, 9, 10]. In comparison, the original 2b/1a chimeric HCV was predominant in Western Europe and North America [10, 12], while the original 2b/1b chimera was identified in Japan and the Philippines [13, 14]. The first 2a/1b chimeric HCV variant originated from Russia [3], and our case was the second report of 2a/1b recombinant HCV.

There are several issues that need to be solved regarding chimeric HCV, including the mechanism of recombination, the cause of the limited G2/G1 recombination pattern, and how HCV is spread. Our cases did not share clinical characteristics, including the infection route. However, mother-to-child transmission of chimeric HCV was suspected in Case 3.

Previously, it was reported that the RF1-2k/1b recombinant HCV was responsive to PEG-interferon therapy in a chimeric mouse model [15]. However, Hoshino et al. showed that patients with recombinant HCV 2b/1b did not achieve an SVR with PEG-interferon and ribavirin combination therapy [8]. Excluding recent pangenotypic DAA regimens, the guidelines recommend an HCV-genotype-specific DAA regimen. It was not clear which DAA regimen was effective for patients with HCV genotype 2/1 chimera. In our patient with HCV 2a/1b, sofosbuvir plus ribavirin, which is recommended for genotype 2, did not result in an SVR, while three DAA regimens recommended for genotype 1 (daclatasvir + asunaprevir, sofosbuvir + ledipasvir, and elbasvir + grazoprevir) all led to an SVR. We speculate that the HCV genotype 2/1b chimera has NS3/4, NS5A, and NS5B derived from genotype 1b. These HCV regions are targets for DAAs. Our data suggest that the exact determination of HCV genotype is important for choosing the optimal treatment for patients with chronic hepatitis C.

In conclusion, we experienced three patients with chimeric HCV that consisted of genotypes 2 and 1b who achieved an SVR to DAA regimens recommended for genotype 1. It is necessary to evaluate the efficacy of pangenotypic DAA regimens for chimeric HCV, including rare types of HCV.

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	Case 1	Case 2	Case 3
Age (years)	39	70	48
Gender	male	female	female
Cirrhosis/non-cirrhosis	non-cirrhosis	non-cirrhosis	cirrhosis
Liver stiffness (kPa)	4.7	4.1	20.2
Past IFN based therapy	no responder	-	-
WBC (/µL)	4300	3900	2800
Hb (g/dL)	16.8	12.2	9.8
Plt (×104/µL)	17.7	15.2	9.7
T-Bil (mg/dL)	1.4	0.7	0.8
ALT (IU/L)	20	19	50
FIB-4 index	0.94	3.11	4.89
AFP (ng/mL)	3.2	2.3	375.7
HCV RNA viral load (log10 IU/mL)	7	6.2	6.5
Serotype	1	1	1
Genotype; core/NS5B	2a/1b	2b/1b	2b/1b
Interleukin 28B rs8099917 SNP	TG	TT	TT
DAA regimen	SOF+RBV	SOF+LDV	DCV+ASV
Outcome of DAA therapy	relapse	SVR	SVR
	SVR		
Outcome retreatment	by GZR+EBR	-	-

Table. Clinical characteristics of the three patients

IFN, interferon; WBC, white blood cell; Hb, hemoglobin; Plt, platelet; T-Bil, total

bilirubin; ALT, alanine aminotransferase; FIB-4 index, Fibrosis-4 index; AFP,

α-fetoprotein; HCV, hepatitis C virus; NS5B, non-structural 5B; DAA, direct-acting

antiviral; SOF, sofosbuvir; RBV, ribavirin; GZR, grazoprevir; EBR, elbasvir; LDV,

ledipasvir; DCV, daclatasvir; ASV, asunaprevir; SVR, sustained viral response.

#### FIGURE LEGENDS

**Figure 1**. HCV 2a/1b recombination breakpoint in Case 1. Alignment of the nucleotide sequences from the recombinant strains before and after sofosbuvir treatment with reference strains HCV2a\_HC-J6 and HCV1b\_Con1. The breakpoint is at +49, referring to the nucleotide position relative to the NS3 start.

## Figure 1

		-60 I	-40	-20
HCV2a_HC-J6	CCCGTGTCCGCCCGA	CTTGGCCGGGAG	GTCCTCCTTGGCCCAG	CTGATGGCTATACCTCC
Case 1 pre-SOF	<b>T</b>	T	C	CG
Case 1 post-SOF			<b>C</b>	
HCV1b_Con1	C C	AGG. GA	A . A . AT G A G	A CA CT . GAAGGG
		<sup>1</sup> Start NS3	20	40
HCV2a_HC-J6	AAGGGGTGGAGTCTT	CTCGCCCCCATC	ACTGCTTATGCCCAGC	AGACACGCGGCCTTTTG
Case 1 pre-SOF	G		C	
Case 1 post-SOF			C	
HCV1b_Con1		GTT	GCCTA	<mark>G</mark> AAC.T
	+49	60	80	100
HCV2a_HC-J6	GGCACCATAGTGGTG	AGCATGACGGGG	CGCGACAAGACAGAAC	AGGCCGGGGGAGATTCAG
HCV2a_HC-J6 Case 1 pre-SOF	New York Contractor Contractor			AGGCCGGGGGAGATTCAG
_		C.C.AC	GGC.ACC.GG GG.ACC.GG	ГТ. А.G G. G А ГТ. А.G G. G А
Case 1 pre-SOF		C.C.AC	GGC.ACC.GG GG.ACC.GG	TT.AGG.G.A
Case 1 pre-SOF Case 1 post-SOF		C.C.AC	GGC.ACC.GG GG.ACC.GG	ГТ. А.G G. G А ГТ. А.G G. G А
Case 1 pre-SOF Case 1 post-SOF HCV1b_Con1		C.C.AC C.C.AC C.C.AC 120	GGC.ACC.GGT GG.ACC.GGT GG.ACC.GGT 140 1	ГТ. А.G G. G А ГТ. А.G G. G А ГС. А.G G. С А
Case 1 pre-SOF Case 1 post-SOF HCV1b_Con1	GTCCTGTCCACGGTC	C.C.AC C.C.AC C.C.AC 120 1 ACTCAGTCCTTC	GGC.ACC.GGT GG.ACC.GGT GG.ACC.GGT 140 1	TT.AGG.G.A TT.AGG.G.A TC.AGG.C.A <sup>160</sup> L CGGGGGGTCTTATGGACT
Case 1 pre-SOF Case 1 post-SOF HCV1b_Con1 HCV2a_HC-J6	GTCCTGTCCACGGTC GG.TCA	C.C.AC C.C.AC C.C.AC 120 ACTCAGTCCTTC G.AT	GGC.ACC.GGT GG.ACC.GGT GG.ACC.GGT 140 I CTCGGAACAACCATCT	TT.AGG.G.A TT.AG.G.G.A TC.AG.G.C.A 160 1 CGGGGGGTCTTATGGACT AT.C.NNNNG.ACTN AT.C.G.GT

Figure legend: HCV 2a/1b recombination breakpoint. Alignment of nucleotide sequences from the recombinant strains pre- and post-SOF treatment with reference strains HCV2a\_HC-J6 and HCV1b\_Con1 was shown. The breakpoint is at +49, referring to nucleotide position relative to NS3 start..

**Figure 2**. HCV 2b/1b recombination breakpoint in Cases 2 and 3. Alignment of the nucleotide sequences from the recombinant strains in Case 2 (daclatasvir + asunaprevir) and Case 3 (ledipasvir + sofosbuvir) with the reference strains HCV2b\_HC-J8 and HCV1b\_Con1. The breakpoint is at +10, referring to the nucleotide position relative to the NS3 start.

### Figure 2

	-1	20	-100	-80
HCV2b_HC-J8	TCATTGTGTGGG	GGGCTGAGAC	AGTGGCGTGTGGAGA	CATCCTGCATGGCCTCCCGGTCT
Case 2			<b>A</b>	
				ΤΤ
HCV1b_Con1	. T CACC	<mark>AC</mark>	C.CG	A.CTTGGC
	-6	0	-40	-20 I
HCV2b_HC-J8	CCGCGAGGCTAG	GTAGGGAGGT	TCTGCTCGGCCCTGC	CGACGGCTACACCTCCAAGGGGT
Case 2	G .			TAGAC.
Case 3				ΤΤ
HCV1b_Con1	CC.CAGG.	. G A .	A.AT. G. A. G.	AACTTGAAGGGC
	1	Start NS3 +1	0 20	40
HCV2b_HC-J8	GGAAGCTCCTAG	CTCCCATTAC	TGCTTACACTCAGCA	AACTCGTGGTCTCCTGGGTGCTA
Case 2	1	0		and the second
		. G	G C I . C A	G A A C G T TGC.
Case 3			GCT.CA	G A A C G T TGC. G G G C A T TGC.
			G C T. C A	G A A C G T TGC. G G G C A T TGC. G G A C A T CTGC.
			G C T. C A	G G G C A T TGC.
HCV1b_Con1			G C T . C A G C T . C A <sup>80</sup> I	G G G C A T TGC.
HCV1b_Con1 HCV2b_HC-J8		G.T.ACGGGCCG	GCT.CA GCT.CA <sup>80</sup> I CGACAAAAATGAGCA	GGGCATTGC. GGACATCTGC. 100 1
HCV1b_Con1 HCV2b_HC-J8 Case 2			G C T. C A G C T. C A 80 1 CGACAAAAATGAGCA G G C GT	GGGCATTGC. GGACATCTGC. 100 I GGCTGGGCAGGTCCAGGTTCTGT

Figure legend: HCV 2b/1b recombination breakpoint. Alignment of nucleotide sequences from the recombinant strains Case 2 and Case 3 with reference strains HCV2b\_HC-J8 and HCV1b\_Con1 was shown. The breakpoint is at +10, referring to nucleotide position relative to NS3 start.