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Rescue of duck-origin virulent Newcastle disease virus from cloned cDNA and stable expression of the red fluorescent protein



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ABSTRACT

Ducks are generally considered as potential reservoirs for different genotypes of Newcastle disease virus (NDV) and to be resistant even to velogenic NDV strains. However, outbreaks of highly virulent genotype VII NDV lethal to ducks have been frequently reported in China in recent years. But until now, the pathogenesis and potential vaccine of duck-origin genotype VII NDV are not known. In this study, a reverse genetics system using the prevalent high virulence genotype VIId isolate SS1 was constructed. Based on this system, a red fluorescent protein (RFP)-expressing virus was generated by inserting an additional transcription cassette coding for the RFP between the noncoding region of P and M genes. The rescue of the recombinant viruses was confirmed by western blotting, fluorescence microscopy and genetic marker detection. In addition, the replication kinetics, biological characteristics and pathogenicity of the rescued viruses were indistinguishable from the parental wild-type virus. Moreover, the recombinant virus rSS1-RFP could efficiently replicate in most of the duck tissues, especially in duck immune organs. The results obtained suggest that this reverse genetics system will provide a useful tool for the analysis of duck-origin NDV pathogenesis and dissemination, as well as preparation for novel vaccine vector or genotype-matched NDV attenuated vaccines used in ducks.

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Newcastle disease virus (NDV), also known as avian paramyxovirus serotype-1 (APMV-1), is a member of the genus *Avulavirus* within the *Paramyxoviridae* family (Gogoi et al., 2015). The genome of NDV is a negative-sense, non-segmented, single-stranded RNA that contains six major genes in the order of 3'-NP-P-M-F-HN-L-5' and encodes six structural proteins as well as two non-structural proteins, V and W (Miller et al., 2010; Yusoff and Tan, 2001). Up to now, more than 250 bird species are reported to be susceptible to NDV, thus causing severe economic burden in domestic poultry industry worldwide (Ganar et al., 2014). In general, ducks are considered as the natural reservoir for all genotypes of NDV and are typically experienced asymptomatic infection or little pathology for velogenic NDV strains lethal to chickens (Liu et al., 2008). However, outbreaks of high virulence genotype

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VII NDV strain lethal to ducks have been occasionally reported in China in recent years (Dai et al., 2014; Kang et al., 2014; Kang et al., 2016), but the pathogenesis of genotype VII NDV in ducks has not been clearly clarified. Therefore, it is worth strengthening fundamental research on the genetic evolution and pathogenic mechanism of virulent NDV isolates.

Nowadays, the development of plasmid-based reverse genetics systems to rescue negative-sense RNA viruses from cloned cDNA provides a means not only to investigate the pathogenesis of virus (Kai et al., 2015; Zhao et al., 2013), but also to make it possible to generate novel vaccines and vaccine vectors (Neumann et al., 2002; Duan et al., 2015a). Since the generation of infectious NDV was firstly reported in 1999, the recovery of other NDV strains, such as Herts/33 (de Leeuw et al., 2005), ZJ1 (Liu et al., 2007), I4 (Hu et al., 2011), 9a5b-D5C1 (Yu et al., 2012) and NA-1 (Wang et al., 2015), was also successfully accomplished. It is noteworthy that all of the rescued NDV strains were isolated from chicken or goose flocks, the recovery of duck-origin virulent NDV strain has not been reported. In this study, we used the established reverse genetics system based on the full-length cDNA under the control of the T7 promoter to rescue duck-origin genotype VII NDV strain SS1. Furthermore, the recombinant virus rSS1-RFP expressing the red fluorescent protein (RFP) was produced. The stability and pathogenicity of rSS1-RFP was also evaluated.

The SS1 strain that caused about 30% mortality was isolated from Sansui Sheldrake duck flocks in Guizhou Province, in 2014 (Duan et al., 2015b). The complete genomic sequence of the isolate has been determined and deposited in the GenBank database (accession no. KP742770). To construct the three helper plasmids, the open reading frames (ORFs) of the NP, P and L genes were amplified from the SS1 cDNA using specific primers (Supplemental Table S1), and then cloned into pCI-neo vector (Promega, USA) to generate pCI-NP, pCI-P, and pCI-L, respectively. To assemble the full-length SS1 cDNA into the TVT7R(0.0) vector, seven overlapping PCR fragments spanning the full-length cDNA of SS1 were generated by RT-PCR with specific primers (Table 1). To conveniently insert foreign genes into SS1 genome as previously reported (Hu et al., 2007; Zhao et al., 2015), the F3 fragment containing the Apal restriction site at the 4118–4123 nt in the M ORF region was eliminated by introducing one synonymous mutation at position 4120 (G to A), and served as the genetic marker, while another Apal restriction site at the 3139-3144 nt in the P-M noncoding region was reserved as an insertion locus for the foreign genes (Fig. 1A). The obtained seven fragments were subsequently ligated into the TVT7R(0.0) vector to generate the full-length cDNA cloning plasmid pTVT/SS1 by utilizing unique restriction enzyme sites (Fig. 1A). For the construction of pTVT/SS1-RFP, the "gene end" (GE), intergenic (IG), "gene start" (GS) and the Apal restriction sites sequences (Hu et al., 2007) were added to the RFP ORF by PCR amplification using primers RFP-F and RFP-R (Table 1), according to the "rule of six" (Fig. 1B). The amplicon was inserted into the F3 fragment, which was then digested with AgeI and BstZ17I to replace the corresponding region in the full-length cDNA clone pTVT/SS1 (Fig. 1B). The resulting plasmid was named pTVT/SS1-RFP.

To rescue infectious virus from the plasmids, BSR-T7/5 cells stably expressing the T7 phage RNA polymerase were grown to 80% confluence in 35-mm-diameter dishes and then cotransfected with a total of 3 µg DNA consisting of a mixture of pTVT/SS1 or pTVT/SS1-RFP together with three helper plasmids at a ratio of 2:2:1:1 using the FuGENE HD transfection reagent (Roche, Germany). At 60 h post-transfection, the cell monolayers and culture supernatants were harvested and inoculated into the allantoic cavities of 10-day-old embryonated SPF eggs. The hemagglutination (HA) test revealed that HA-positive allantoic

Table 1

Primers used for the construction of full-length cDNA and helper plasmids.

Primer name	Position	Nucleotide sequence $(5' \rightarrow 3')$	Length (bp)
P1F	1-22	CGTCTCGTATAGGGACCAAACAGAGAATCTGTGAGG (BsmBI)	1803
P1R	1779-1803	TTCTAATACCTTGGGCTCTGGGCGG	
P2F	1738-1763	CCACCCGGGACAACACAGGCACAGC	1204
P2R	2911-2941	ACGACGCGTTCAGGCCCGCTTGCAGTGGCAGAC (MluI)	
P3F	2859-2884	ACGCTCAATAAACTCTCACAACCGG	1876
P3R	4710-4734	ATGATTGACCCAGTCTGAGACGAGG	
P4F	4661-4689	GCAGCTGCAGGAATTGTAGTAACAGGAG	2713
P4R	7348-7373	CCTCCGTAAACTGGGAACCATACACG	
P5F	7318-7346	GGAGTGGGAGGAGGGTCTTTTATTGACG	4704
P5R	11,997-12,021	GACATGTGGGCTATTTTCGCGAGCG	
P6F	11,969–11,994	TCGAAGACTCAAGAGAGAGAGAGCCG	1126
P6R	13,070-13,094	ACAGCAGATCGTACTTATTCCCTGC	
P7F	13,028-13,053	TTGTTAGTCTCTTGCACTCGACGCG	2165
P7R	15,168-15,192	CGTCTCTACCCACCAAACAGAGATTTGGTGAATGAC (BsmBI)	
NP-F	78–99	CG GAATTC GAGCGCGAGGCCGAAGCTTGAAC (EcoRI)	1544
NP-R	1585-1606	AT GTCGAC CTGGGTGTTGTCGATCAGTAC (Sall)	
P-F	1877-1901	CCG CTCGAG GTGAATTAGGGTGAAGATGGCCACT (XhoI)	1256
P-L	3092-3144	GCC GTCGAC AGTGACGGGAGCCTGTTGTGAGT (Sall)	
L-F1	8385-8412	ACGGTCGACACATGGCGGGCTCCGGTCCAGAAAGGGC (Sall)	4696
L-R1	13,041-13,071	GCATATAAACCTGAGACCACGCGTCGAGTGC (MluI)	
L-F2	13,034-13,059	GTCTCTTGCACTCGACGCGTGGTCTC (MluI)	1980
L-R2	14,983-15,001	ATATGCGGCCGCTTAAGAGTCATTATTACTG (Notl)	
RFP-F		CTGGGGCCCTCttagaaaaaaaTacgggtagaaGTACCATGGCCTCCTCCG AGAACGTC (Apal)	
RFP-R		TTGGGCCCTCTTACAGGAACAGGTGGTGGCGGCCCTCGGTG CGCTCG (Apal)	

The restriction enzyme sites used in the primers are marked in italic and bold type. The gene end sequence is lowercased and italicized. The gene start sequence is lowercased.

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