1	MOLECULAR ANALYSIS OF NSP4 CODING GENE OF PORCINE ROTAVIRUS IN
2	BRAZIL
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4	(ANÁLISE MOLECULAR DO GENE CODIFICADOR DA NSP4 DE ROTAVÍRUS SUÍNOS
5	NO BRASIL)

- 6 SUMMARY
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The non-structural protein 4 (NSP4) has different roles in rotaviral replication, morphogenesis, 8 and enterotoxin-like activity causing secretory diarrhea. A total of 11 partial nucleotide 9 sequences of NSP4 coding gene were defined from group A rotavirus circulating in Brazilian 10 swine herds. On comparing the viral sequences of diarrheagenic peptide area (amino acid 114-11 135), there was a single point mutation at amino acid 135 presented by two strains with amino 12 acid alanine, and valine in the others. The NSP4 gene phylogeny showed that all strains 13 14 clustered into E1 genotype, and the nucleotide identity between Brazilian strains ranged from 92.4% and 100%, while the putative amino acid identity, between 95.8% and 100%. As a 15 conclusion, these data demonstrate the occurrence of a common NSP4 genotype described 16 elsewhere in pigs and low diversity between the samples from the surveyed areas. 17

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KEY WORDS: Genotypes. Non-structural protein. Reoviridae. 19

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RESUMO 21

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A proteína não estrutural 4 (NSP4) desempenha diferentes funções na replicação e na 23 24 morfogênese dos rotavírus, possuindo, ainda, uma atividade de enterotoxina, causando diarreia do tipo secretória. Um total de 11 seguências parcias de nucleotídeos do gene codificador da 25 NSP4 de rotavírus suínos de criações brasileiras foram definidas como pertencentes ao grupo 26 A. Comparando-se as sequências virais da área do peptídeo toxigênico, que compreende a 27 28 porção entre os aminoácidos de 114 a 135, constatou-se uma única mutação pontual no aminoácido 135, sendo que duas amostras apresentaram alanina, e as demais, valina. A análise 29 filogenética do gene demonstrou que todas as amostras pertencem ao genotipo E1, e que a 30 identidade nucleotídica das amostras brasileiras variou de 92,4% a 100%, enquanto que a 31 identidade de aminoácidos, de 95,8% a 100%. Assim, esses dados mostram a ocorrência de um 32 genotipo comum da NSP4 já descrito anteriormente em suínos, com uma baixa diversidade 33 entre as amostras encontradas. 34

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PALAVRAS-CHAVE: Genotipos. Proteína não-estrutural. Reoviridae. 36

INTRODUCTION

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Group A rotavirus (RV-A), members of the Reoviridae family, genus Rotavirus, are 40 regarded as a major cause of gastroenteritis both in humans and animals worldwide 41 (KAPIKIAN et al., 2001). The RV-A genome consists of 11 segments of double-stranded RNA, 42 encoding six structural virus proteins (VP1-VP4, VP6 and VP7) and six non-structural proteins 43 (NSP1-NSP6). The complete virus is a triple-layered particle, with VP4 and VP7 constituting 44 the outer layer whose respective encoding genes are markers for P and G genotypes, 45 respectively (KING et al., 2012). So far, 35 genotypes P and 27 G have been defined 46 (MATTHIJNSSENS et al., 2011). The inner capsid protein VP6 important on host immunity 47 and determines groups A-G, and more recently, a novel RV-H has been discovered (KING et 48 al., 2012; MATTHIJNSSENS et al., 2012). Genotypes previously described in pigs include 49 50 G1, G2, G3, G4, G5, G6, G11 and G12, usually associated with P[6], P[7], P[13], P[19], P[23], P[26], and P[27] (MATTHIJNSSENS et al., 2008b; TONIETTI et al., 2013). 51 52 The non-structural protein 4 (NSP4), encoded by gene segment 10, has multiple functions

in RVs morphogenesis and pathogenesis. It has an enterotoxin-like activity (BALL et al., 1996) 53 and has been identified as a viroporin (HYSER et al., 2012). The peptide 114-135 is considered 54 to trigger a signal transduction pathway as it increases intracellular calcium leading to chloride 55 secretion, and therefore secretory diarrhea, as it has been shown in mice (BALL et al., 1996; 56 HUANG et al., 2004; TIAN et al., 1995). Changes within this region have been associated with 57 alterations in the toxigenic activity of NSP4 and virulence of RVs (BALL et al., 1996; ZHANG 58 et al., 1998). So far 14 NSP4 genotypes have been defined from RV-A samples infecting 59 Human and animal hosts (MATTHIJNSSENS et al., 2008a). 60

The aim of this investigation was to sequence and analyze a partial fragment of NSP4 gene of RV-A from different Brazilian pig herds to define their phylogenetic relations with other animal and human isolates described elsewhere.

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MATERIAL AND METHODS

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A total of 11 stool samples from pigs with diarrhea from three cities in São Paulo's State,
Brazil, were collected in 2008 and screened with polyacrylamide gel electrophoresis (PAGE),
ELISA, and characterized in P and G genotypes as previously described (GOUVEA et al.,
1994a,b).

Feces suspensions (v/v; 50%) were prepared with phosphate-buffered saline 0.01M, pH
72 7.2, clarified at 5000g/15 min at 4°C, and the supernatants used in the assays.

Extraction of total RNA from the reference RVs strain (NCDV) and the supernatants of
the field samples were carried out with TRIzol ReagentTM (Invitrogen, Carlsbad, CA, USA)
according to the manufacturer's instructions.

For RT-PCR, 5.6µL of RNA solution was mixed with 1.4µL of DMSO and denatured at 95°C for 5 minutes and kept in ice. Then it was added to a solution of 1x First Strand Buffer (InvitrogenTM), 1 mM of each dNTP, 10 mM DTT, and 1 µM of each primer targeting NSP4 coding gene (10BEG16 and 10END722) as described by Lee et al. (2000) and 200U of reverse transcriptase (InvitrogenTM), to a 13µL final reaction volume. This mixture was then heated at 42°C for 1 hour and 70°C for 15 min at thermal cycler.

PCR amplification was carried out by adding 5μ L of cDNA of the RT reaction in a mix containing 1x PCR Buffer (InvitrogenTM), 0.2 mM of each dNTP, 0.5 μ M of each primer (10BEG16 and 10END722), as described by Lee et al. (2000), 2 mM of MgCl₂, and 2.5U of Taq DNA Polymerase (InvitrogenTM) and ultra-pure water for a final reaction volume of 50 μ L. This mixture was heated at 94°C for 2 min, followed by 30 cycles each at 95°C for 45 s, 49°C for 30 s, 72°C for 1.5 min, and one cycle at 72°C for 10 min. The products of the PCR were resolved on a 1.5% agarose gel stained with 0.5µg/mL ethidium bromide.

Amplicons of 725 bp in length were purified with Illustra GFXTM PCR DNA and Gel
Band Purification Kit, according to the manufacturer's instructions (GE Healthcare) and
submitted to bi-directional sequencing with BigDye 3.1TM (Applied Biosystems, Carlsbad, CA,
USA) according to the manufacturer's instructions. The reaction products of the sequencing
reactions were resolved in the automatic sequencer ABI-377TM (Applied Biosystems, Carlsbad,
CA, USA).

95 Nucleotide sequences obtained in this study (nt 66 to nt 566, using as reference Gottfried Strain accession number GU199490) (Table 1) were aligned among them and with 96 representative strains belonging to different NSP4 genotypes according to Matthijnssens et al. 97 (2008a) using Bioedit 7.0.5.3 software (HALL, 1999) and Clustal W 1.83 (THOMPSON et al., 98 1994) downloaded from the NCBI GenBank database. The strains used were 99 100 (genotype/accession number/host/strain): a) E1/ GU199490/ Swine/ Gottfried; b) E1/ DQ494398/ Bovine/ KJ75; c) E1/ AF144799/ Swine/ A411; d) E1/ D88831/ Swine/ OSU; e) 101 E1/X69485/Swine/YM; f) E1/U59109/Human/M37; g) E2/AF144805/Bovine/B223; h)E3/ 102 AF144806/ Canine/ CU 1; i) E4/ AB065285/ Avian/ Ty 1; j) E5/ AF533535/ Lapine/ 160 01; 103 k) E6/ DQ490560/ Human/ RV176 06; 1) E7/ U96337/ Murine/ EC; m) E8/ EF442742/ Canine/ 104 RV52 96; n) E9/ DQ534017/ Swine/ CMP034; o) E10/ FJ169862/ Avian/ 02V0002G3 and p) 105 E12/ FJ347120/ Bovine/ Arg B383. 106

The nucleotide and amino acid similarities were calculated using Bioedit v. 7.0.5.3
software (HALL, 2009). The phylogenetic tree from nucleotide sequences was built using
MEGA software version 4 (TAMURA et al., 2007) based on Neighbor-joining method using
Maximum Composite Likelihood (1,000 bootstrap trials).

RESULTS

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For the 11 samples, a common fragment of 501 nt (nt 66 to nt 566, using as reference 113 Gottfried Strain accession number GU199490) from RVs NSP4-coding gene was investigated. 114 The strain identification, P and G genotypes, and respective NSP4 gene accession numbers are 115 shown in Table 1. The Genbank accession numbers for the NSP4 partial gene sequences of 116 porcine rotaviruses determined in this study are: HQ840943, HQ840944, HQ840945, 117 HQ840946, HQ840947, HQ840948, HQ840949, HQ840950, HQ840952, HQ840953, 118 HQ840954. 119 120 Nucleotide identity ranged from 92.4% to 100% while amino acid ranged from 95.8% to 100%. The comparison of NSP4 genes sequenced in this study with other strains classified as 121 genotype E1 from GenBank revealed a nucleotide identity ranging from 94.4% (strain PORV6 122 with porcine strains Gottfried and OSU) to 84.1% (strain PORV9 with human strain EF672589) 123 and amino acid identity ranging from 98.8% (strains PORV1; PORV2; PORV3; PORV4; 124 PORV6; PORV7 and PORV11 with Venezuelan porcine strain AF165219) to 89.4% (strain 125 PORV5 with human strain EF672589). 126

Deduced amino acids of the sequences generated herein revealed a moderate variation among the strains (Fig. 1). Moreover, considering the toxigenic peptide (amino acid 114-135) it was shown that there was a single point mutation on aa 135 presented as alanine in two RV strains and as valine in the other strains. In addition, six other amino acid changes at residues 136 (valine, alanine and serine), 137 (arginine and glycine), 139 (isoleucine and valine), 154 (arginine and lysine), 161 (serine and asparagine) and 174 (serine and proline) were found.

The phylogenetic tree (Fig. 2) depicts that the strains of the present study clustered with E1 genotype representatives, while the others segregated in separate clusters with a resolved genealogy, according to its genotypes.

DISCUSSION

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Among non-murine NSP4 amino acid sequences, most of the divergence was observed in the VP4-binding domain (aa 112-148) and in the double-layered particle-binding region (aa 161-175) (IOSEF et al., 2002). Strains of the present study showed low degree of polymorphism in both regions, with four and two mutations respectively, as shown in Fig. 1.

Strains PORV6 and PORV10 presented amino acid residue alanine at position 135, while 142 the others presented amino acid valine. By comparisons of NSP4 sequences, Zhang et al. (1998) 143 suggested that changes between amino acids 131 and 140 are important for viral pathogenesis, 144 145 showing that a change from amino acid valine to alanine in the NSP4 protein at this position was important in OSU attenuated strains, as it was associated with loss of the ability of inducing 146 diarrhea in mice, which was also observed in a piglet model with virulent and tissue culture-147 attenuated human RVs Wa strains (WARD et al., 1996). On the other hand, Kirkwood et al. 148 (1996) found isoleucine at position 135 in symptomatic children, as well as did Mascarenhas et 149 150 al. (2007).

Tyrosine residue at position 131 of NSP4 coding gene has been postulated to be critical for the diarrheagenic activity of the toxic peptide (BALL et al. 1996), but histidine was also found in diarrheic young children (CUNLIFFE et al., 1997; MASCARENHAS et al., 2007). Sequence analysis from porcine strains revealed amino acids serine, alanine and histidine at residue 131 (CIARLET et al., 2000; MATTHIJNSSENS et al., 2010; STEYER et al., 2007). In the present study, all strains showed histidine, as shown in Fig. 1. Therefore, the enterotoxin domain (aa 114-135) is conserved among them, except for one mutation at aa 135.

Even though nucleotide and amino acid polymorphism were found both at the toxigenic peptide and VP6-binding domain (aa 112-175) observed in Fig. 1, it was not possible to speculate on the significance of these changes for the virulence of the RV strains since all the animals studied had diarrhea. In other studies, this correlation between virulent and attenuated
strains was not observed (ANGEL et al., 1998; WARD et al., 1997), showing the possibility
that virus attenuation can occur by several mechanisms, including mutations in other viral
proteins. Moreover, the extreme C terminus, including aa methionine at position 175 was shown
to be important for double-layered particle (DLP)-binding activity (TAYLOR et al., 1992). As
shown in Fig. 1, all the porcine strains presented methionine at this site.

This study revealed the occurrence of genotypes G10 and G11 in association with P[6] or P[7] in the swine population. G10 genotype has been widely detected in bovine rotaviruses in Brazil (ALFIERI et al., 2004) and other countries (FALCONE et al., 1999; GARAICOECHEA et al., 2006; HOWE et al., 2008), and also in humans (RAMANI et al., 2009; URASAWA et al., 1993). A study in Thailand also revealed this genotype in pigs (PONGSUWANNA et al., 172 1996).

G11 rotaviruses were first detected in pigs in Mexico and Venezuela (CIARLET et al., 174 1994; ROSEN et al., 1994; RUIZ et al., 1988) and are believed to be circulating in this 175 population, although in low numbers. In subsequent years, no additional G11 strains were 176 detected in the same or nearby pig farms, but in the last decade, several reports have described 177 the isolation of G11 RVs strains from humans (MATTHIJNSSENS et al., 2010). These authors 178 also showed that multiple reassortment events have occurred between porcine or human G11 179 rotaviruses and co-circulating human Wa-like RVs strains.

The phylogenetic tree (Fig. 2) showed that the circulating Brazilian RVs strains belong to E1 genotype, also reported elsewhere in humans, swine, equine, and bovine (MATTHIJNSSENS et al., 2008a), reinforcing the association between E1 genotype and pig RVs previously described. Although evidences for independent segregation of the VP6- and NSP4-encoding genes have been described in porcine RV-A (GHOSH et al., 2006; ITURRIZA-GÓMARA, 2002), considering the limited number of surveyed samples and occurrence of undefined P and G genotypes, it was not possible to observe this pattern among Braziliansamples.

Interspecies transmission of rotaviruses may occur in natural and experimental conditions 188 (MARTELLA et al., 2010). The introduction of a new human-animal reassortant RVs strain 189 into the human population could have an impact on the spread of rotavirus disease and also on 190 prevention measures (STEYER et al., 2008). This study also revealed (data not shown) that 191 192 strain PORV6 had 96,4% amino acid identity with Brazilian strain NB-150, a human strain previously isolated by Mascarenhas et al. (2007) from a newborn with diarrhea who lived in 193 the outskirts of Belém do Pará, Brazil, that reinforce the hypothesis that interspecies 194 195 transmission may occur naturally, without loss of virulence (VARGHESE et al., 2004).

There are numerous examples of RVs interspecies transmission, but there are few 196 documented evidences in which whether the transmission event has involved the whole genome 197 (PALOMBO, 2002). In fact, pigs may serve as a reservoir of RVs for humans, as described by 198 several authors in different countries, such as India, Ecuador and Hungary (BANYAI et al., 199 2004; BANYAI et al., 2009; VARGHESE et al., 2004). It has been proposed that human RVs 200 Wa-like strains and swine strains have a common origin (MATTHIJNSSENS et al., 2008b), 201 and, recently, a new virus isolated from pigs was closely related to a novel group of human 202 203 rotaviruses (WAKUDA et al., 2011).

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CONCLUSIONS

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As a conclusion, NSP4 genes of porcine RVs isolated in Brazil during 2008 had only a moderate polymorphism and belonged all to E1, in an extent previously unknown in this country.

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TABLE 1

Strain	Genotype P	Genotype G	NSP4 Accession number
PORV1	P[6]	G[11]	HQ840943
PORV2	P[6]	G[11]	HQ840944
PORV3	P[6]	G[11]	HQ840945
PORV4	P[6]	G[11]	HQ840946
PORV5	-	-	HQ840947
PORV6	-	G[10]	HQ840948
PORV7	P[7]	-	HQ840949
PORV8	P[6]	-	HQ840952
PORV9	P[7]	-	HQ840950
PORV10	P[7]	G[10]	HQ840953
PORV11	-	G[10]	HQ840954

Table 1: P and G genotypes and accession numbers of partial NSP4 sequences RVs from
piglets samples in São Paulo State, Brazil. Gaps indicate genotypes that were not defined.

FIGURE 1

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	10	20	30	40	50	60	70	80	90
PORV1	~~~~~NYTLSVIT	LMNDTLHSII	ODPGMAYFPYIA	SVLTVLFTL	HKASIPTMKIA	LRTSKCSYKV	IKYCMVTIIN	TLLKLAGYKE(TVC
PORV2	~~~~ ADL								
PORV3	~~~ T ADT								
POPV4	~~~~ ADT .								
POPVE	T	•••••			т	v	37	•••••	•••
PORVS							•••••••••••		•••
PORVE	~~~~~	•••••	••••••••••	••••••	••••••	• • • • • • • • • • •	•••••	•••••	•••
PORV /	~~~~~	• • • • • • • • • • •	••••••		•••••	• • • • • • • • • •	• • • • • • • • • • •	•••••	•••
PORV8	MDKLADL	• • • • • • • • • •	••••••••••	• • • • • • • • • •		• • • • • • • • • •	•••••	•••••	•••
PORV9	~~~~L					.K	• • • • • • • • • •	.I	•••
PORV10	MDKLADL	V				.K			
PORV11	MDKLADL								
	100	110	120	130	140	150	160	170	
	100	110	120	130	140	150	160	170	
PORV1	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 LTTREIEQVELI	130 	140 RPIDAIDMSKE	150 . FNQKNIRTLD	160 . EWESGKNPYE	170 •• ••• PSE VTASM	
PORV1 PORV2	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 . LTTREIEQ V ELI	130 	140 . RPIDAIDMSKE	150 . FNQKNIRTLD	160 . EWESGKNPYE	170 PSEVTASM	
PORV1 PORV2 PORV3	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 LTTREIEQVELI	130 	140 RPIDAIDMSKE	150 . FNQKNIRTLD	160 	170 	
PORV1 PORV2 PORV3 PORV4	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 LTTREIEQVELI	130 	140 RPIDAIDMSKE	150 	160 	170 	
PORV1 PORV2 PORV3 PORV4 PORV5	100 TKDEIEQQMDRIVKE	110 III MRRQLEMIDK	120 	130 	140 . RPIDAIDMSKE	150 	160 . EWESGKNPYE	170 	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV5	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 . LTTREIEQVELI	130 	140 . RPIDAIDMSKE	150 	160 . EWESGKNPYE	170 	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV6 PORV6	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 . LTTREIEQVELI	130 	140 	150 	160 	170 PSE VTASM	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV6 PORV7 PORV7	100 TKDEIEQQMDRIVKE	110 III MRRQLEMIDK	120 	130 	140 	150 	160 	170 PSE VTASM	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV6 PORV7 PORV8 PORV9	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 . LTTREIEQVELI	130 	140 	150 	160 	170 PSEVTASM	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV5 PORV5 PORV5 PORV5 PORV5 PORV5 PORV5	100 TKDEIEQQMDRIVKE	110 I MRRQLEMIDK	120 . LTTREIEQVELI	130 	140 	150 	160 	170 PSE VTASM P.	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV5 PORV7 PORV7 PORV8 PORV9 PORV10	100 TKDEIEQQMDRIVKE	110 MRRQLEMIDK	120 	130 	140 	150 	160 	170 	
PORV1 PORV2 PORV3 PORV4 PORV5 PORV5 PORV5 PORV7 PORV8 PORV9 PORV10 PORV11	100 TKDEIEQQMDRIVKE	110 I MRRQLEMIDK	120 	130 	140 	150 	160 	170 PSEVTASM	

Fig. 1. Section of the alignment of the deduced 175 amino acids (aa 9-175) of the NSP4-coding

391 gene from rotavirus detected in porcine stool samples from Brazilian herds. The marked area

refers to the toxigenic peptide (residues 114 to 135).

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FIGURE 2



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Fig. 2. Unrooted neighbor-joining tree for a stretch of 501 nucleotides (nt 25-525) of the NSP4coding gene, showing the proposed E genotypes. Taxa designated as "PORV (1 to 11)" are related to the Brazilian field strains from the present study; numbers at each node are the bootstrap values greater than 50% obtained with 1,000 replicates.

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