

Emergence of a new GII.17 norovirus variant in patients with acute gastroenteritis in Jiangsu, China, September 2014 to March 2015

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From September 2014 to March 2015, 23 outbreaks of norovirus (NoV) acute gastroenteritis occurred in Jiangsu, China. Partial sequencing of the NoV capsid gene suggested that 16 of the 23 outbreaks were related to a new GII.17 variant. This variant was first detected in sporadic specimens in October 2014, and became predominant in February 2015. Analysis of the RNA-dependent RNA polymerase (RdRp), and complete capsid including the protruding domain P2 sequences confirmed this GII.17 variant as distinct from previously identified GII variants.

Norovirus (NoV) is a major cause of viral gastroenteritis and is the main aetiological agent of outbreaks of acute gastroenteritis [1]. It is estimated that each year NoV cause 64,000 episodes of diarrhoea requiring hospitalisation and 900,000 clinic visits among children in industrialised countries, and up to 200,000 deaths of children <5 years of age in developing countries [2].

NoVs are classified into six genogroups, GI–GVI, of which genogroup I, II, and IV are responsible for disease in humans [3,4]. Genogroups are subdivided further into genotypes. To date, based on RNA-dependent RNA polymerase (RdRp) and capsid gene sequences, 31 and 22 genotypes of GII NoVs have been respectively determined [5]. Of these, GII.4 caused at least six epidemics of gastroenteritis worldwide over the past 20 years (1995–1996, 2002, 2004, 2006, 2009, and 2012) with the emergence and rapid global spread of viral variants [6]. In contrast, GII.17 NoV has rarely been reported as a major genotype causing diarrhoea.

In late 2014, the Emergent Public Health Event Information Management System (EPHEIM) in Jiangsu province observed an increase of NoV outbreaks compared with previous seasons. Data from these outbreaks indicated that this increase was associated with the emergence of a new variant of GII.17, which was rarely reported in China before 2014. Surveillance of NoV in both outbreak and sporadic cases was conducted from September 2014 to March 2015 to study the molecular epidemiology characteristics of GII.17-associated diarrhoea in Jiangsu province, China.

Methods

Surveillance of gastroenteritis

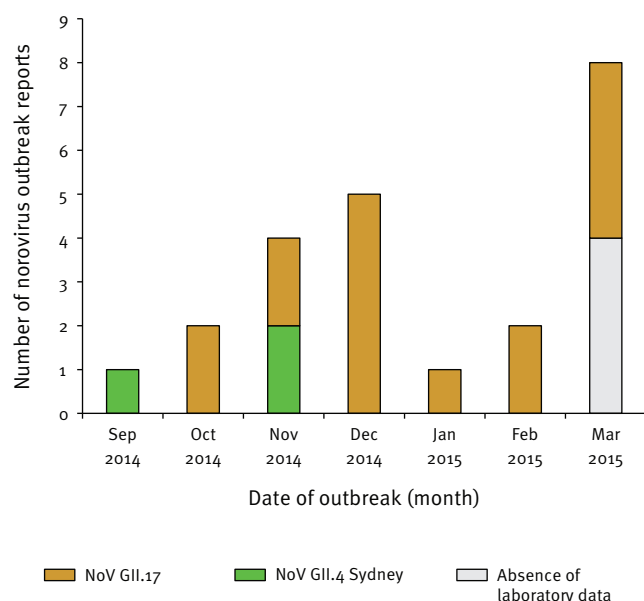
Outbreaks of gastroenteritis and the occurrence of sporadic cases were monitored through different surveillance systems.

Outbreak data were obtained from EPHEIM. An acute gastroenteritis outbreak was defined as ≥20 cases of vomiting and/or diarrhoea associated with a common exposure.

Data on sporadic cases of gastroenteritis in the September 2014 to March 2015 period originated from two surveillance systems in Jiangsu province. The first was the comprehensive surveillance of infectious diarrhoea, which was launched in March 2012 for epidemiological and aetiological surveillance of diarrhoea in children (≤15 years-old) and adult outpatients. This surveillance was conducted via 26 hospitals located in Nantong, Wuxi and Xuzhou, three cities chosen for their location in the southern, central and northern

FIGURE 1

Numbers of laboratory-confirmed norovirus (NoV) outbreaks per month in Jiangsu province, China, September 2014–March 2015 (n=23)



part of the province. Nantong, Wuxi and Xuzhou have a population of 7.298 million, 6.484 million and 8.591 million respectively, and totally account for 28% of the province's whole population (79.395 million). The second surveillance system for sporadic cases was that of diarrhoea of viral origin in Jiangsu province, which was launched in January 2006 for the epidemiological and aetiological surveillance of viral diarrhoea in children hospitalised in Suzhou Children's Hospital in Suzhou. This hospital is the second largest paediatric hospital in Jiangsu province.

Sporadic specimens that were laboratory-confirmed for any viral agent causing gastroenteritis were submitted to the laboratory of Jiangsu Provincial Center for Disease Control and Prevention (JSCDC) on a monthly basis for further analysis.

Testing samples for norovirus

For viral RNA extraction, a 10% (wt/vol) stool suspension in RNase-free water at a total volume of 1 ml was centrifuged for 5 min at 2,370xg. The supernatant was further processed with a MagMAX-96 Viral RNA Isolation Kit (Applied Biosystems, CA, US) on an automated MagMAX Express24 Magnetic Particle Processor (Applied Biosystems, CA, US) according to the manufacturer's instructions.

The presence of NoV RNA was tested for GII NoVs by using the Qiagen Probe RT-PCR Kit (Qiagen, Hilden, Germany) on a 7500 real-time PCR platform (Applied Biosystems, Singapore) with primers (Cog2F/Cog2R) as described previously [7].

Molecular characterisation of the norovirus

RNA from NoV positive specimens was analysed by reverse transcription-polymerase chain reaction (RT-PCR) directed at the region C of the capsid gene (open reading frame (ORF)2; 344bp), using the previously described primers G2SKF/G2SKR [8].

Region A sequences of the RdRp gene in ORF1 were obtained by using a semi-nested GII-specific primer set (NV2F/G2SKR for a first-round PCR and p289IUB/G2SKR for a second-round PCR) [8–10], which amplified a region of 1,095 bp in the ORF1/ORF2 junction of the viral genome.

Extracted viral RNA was reverse transcribed to cDNA with a VN3T20 primer by using the Superscript III cDNA synthesis kit (Invitrogen, CA, US). ORF2 gene sequences encoding the major capsid protein viral protein 1 (VP1) were obtained by using a semi-nested PCR GII-specific primer set (COG-2F/VN3T20 in the first-round PCR and G2SKF/VN3T20 for the second-round PCR) [7,8,11].

The PCR products were purified and then sent to the Sangon Biotech (Shanghai) Company for sequencing. The nucleotide sequences data of GII.17 variants were deposited in GenBank under accession numbers KR270442–KR270449.

Preliminary genotypes were assigned by using the norovirus genotyping tool (<http://www.rivm.nl/mpf/norovirus/typingtool>).

The phylogenetic analysis of aligned sequences was carried out using Molecular Evolutionary Genetics Analysis (MEGA) 5.1 [12]. The reliability of the phylogenetic tree was assessed by bootstrap sampling of 1,000 replicates, and genetic distances were calculated by Kimura's 2 parameter method [13].

Results

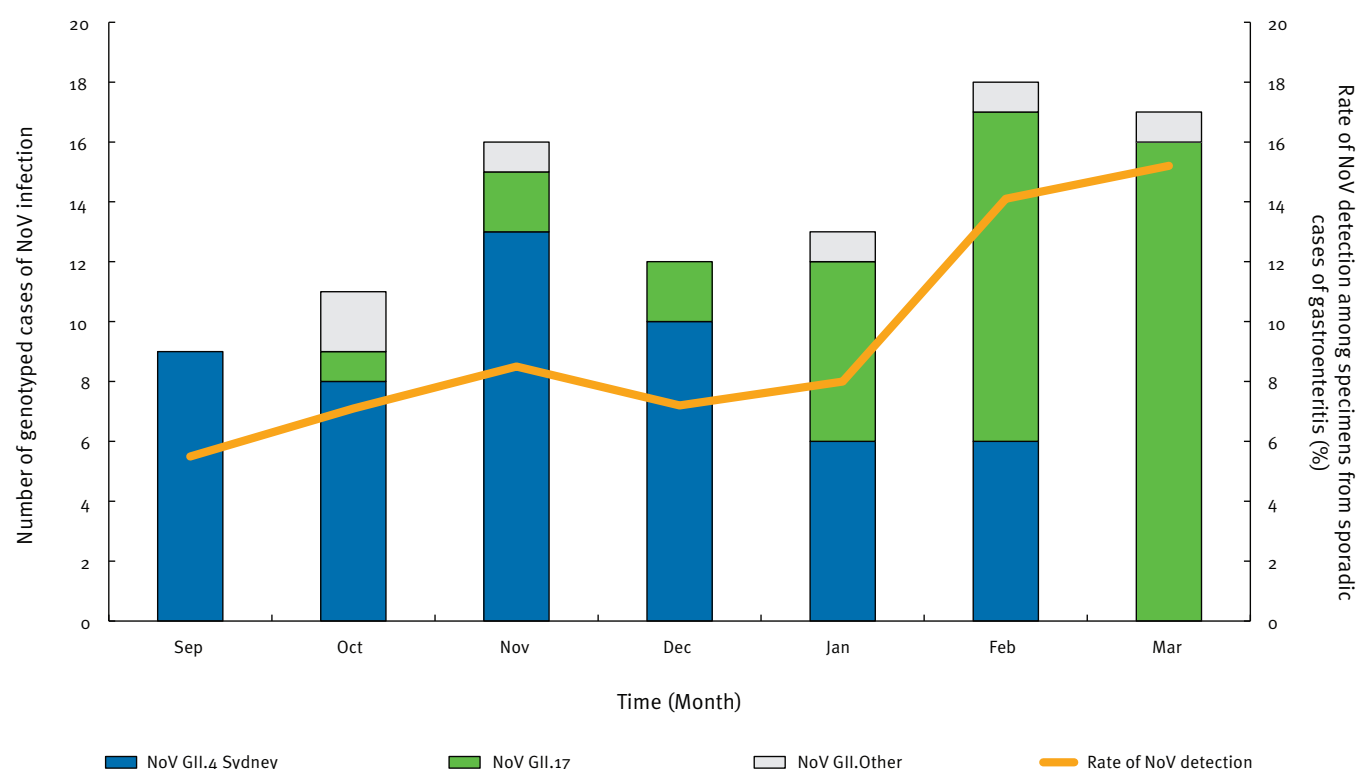
Epidemiological features and genotyping of noroviruses

From September 2014 to the end of March 2015, there were 23 norovirus laboratory-confirmed outbreaks reported to EPHEIM in Jiangsu province, substantially higher than during the same time period in the previous years 2013 to 2014. JSCDC received specimens from all outbreaks up to February 2015, however not from all the outbreaks in March 2015. This resulted in specimens being available from 19 of the 23 outbreaks. Sequence analysis of the Region C of the capsid gene from these 19 outbreaks (Figure 1) showed that three outbreaks (3/23) were caused by GII.4 Sydney, while the other 16 outbreaks (16/23) were all caused by a novel GII.17 variant. This novel variant was first identified in October 2014 and became the dominant strain from December 2014 to March 2015 (Figure 1).

During the study period, 132/1,077 sporadic specimens were positive for norovirus (79/783 outpatient,

FIGURE 2

Monthly distribution of norovirus (NoV) genotypes among sporadic NoV infections and monthly detection rate of NoV among specimens from sporadic cases of gastroenteritis, Jiangsu province, China, September 2014–March 2015



During the study period, 132/1,077 specimens obtained from sporadic cases of gastroenteritis were positive for NoV. The orange curve shows a plot of monthly detection rates of NoV among specimens.

Of the 132 specimens testing positive for norovirus, genotype was determined for a total of 95 samples. The chart vertical bars represent the total monthly number of specimens of each genotype.

and 53/294 hospitalised children), and partial capsid sequences (Region C) of 95 strains were sequenced, including 57/79 outpatient, and 38/53 hospitalised children. The sequencing of the 95 sporadic specimens resulted in the identification of five capsid genotypes (GII.2, GII.3, GII.13, GII.4 Sydney, and GII.17). A novel GII.17 variant was first identified in October 2014 with only one strain, compared with eight strains of GII.4 Sydney. GII.4 Sydney variant remained the dominant strain from September to December 2014 (9/9 in September, 8/11 in October, 13/16 in November, and 10/12 in December). In January 2015, the proportion of GII.4 Sydney decreased to 6/13, but that of the GII.17 variant increased to 6/13. In February 2015, the proportion of GII.17 variant further increased to 11/17 making it the predominant variant. This predominance continued in March 2015 when GII.4 Sydney was no longer detected. Along with the increased number of confirmed NoV GII.17 specimens, a higher detection rate of NoV among samples from sporadic cases of gastroenteritis was observed in February and March of 2015 (Figure 2).

Phylogenetic Analysis

The Region A and complete VP1 region of eight GII.17 variants (3 specimens from 3 respective outbreaks and 5 sporadic specimens) were further compared with other GII.17 strains by phylogenetic analysis.

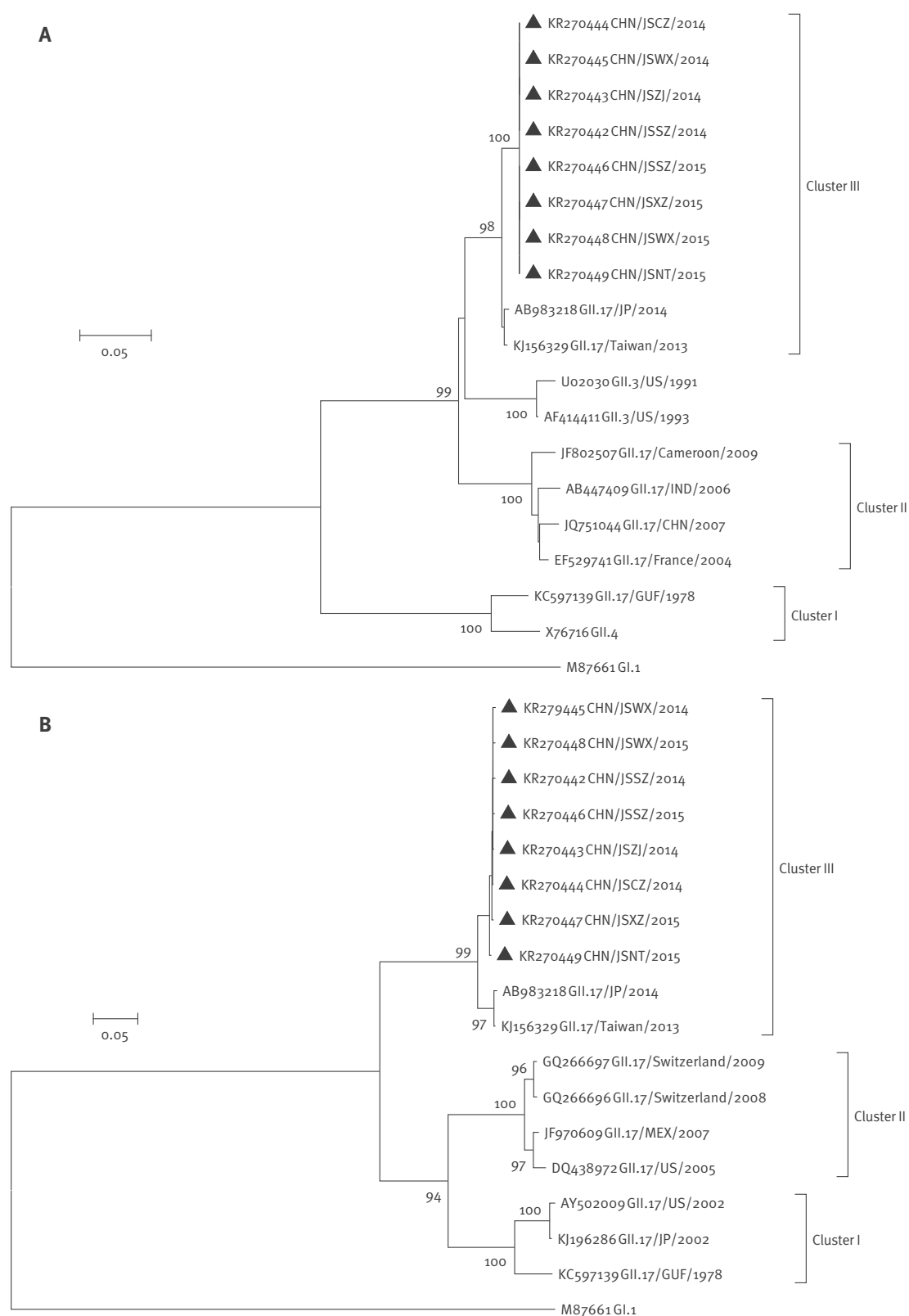
The GII.17 strains were segregated into three distinct genetic groups both in Region A (Figure 3a) and VP1 region (Figure 3b).

In the tree based on Region A, the GII.17 strain characterised in 1978 grouped with a GII.4 Rpd, forming Cluster I. This cluster appeared to be ancestral to GII.17 sequences reported after 2003. The GII.17 variants identified in the period from 2004 to 2011 formed Cluster II and the variants originating from 2013 to 2015 formed Cluster III. Sequences from Cluster II and Cluster III shared a high nucleotide identity with GII.3 reference strains, especially those in Cluster III (Figure 3a).

In the tree based on the VP1 gene, the GII.17 variants from 1978 to 2002 formed Cluster I, those from 2005

FIGURE 3

Phylogenetic trees based on (a) Region A including the RNA-dependent RNA polymerase in open reading frame (ORF) 1 and (b) the complete major capsid protein viral protein 1 (VP1) in ORF 2



The scale bars indicate the number of nucleotide substitutions per site for the phylogenetic trees.

The numbers on the branches represent the percentage bootstrap support for the nodes after 1,000 iterations. Genbank accession numbers of all sequences used for the phylogenetic analysis figure in the respective trees.

In the two trees, the GII.17 respective VP1 and Region A sequences obtained from the eight strains characterised in the present study (Jiangsu province, China, October 2014–March 2015) are indicated by black triangles.

FIGURE 4

Alignment of sequences of viral protein 1 protruding P2 domain derived from norovirus (NoV) GII.17 strains

		RGD/K	Epitope A	Site I	Epitope A	Site II
		285	295	350	375	385
	
		↓	↓	↓	↓	↓
	
KC597139	GUF/1978	SGICAFRGKL	TADVHQSHDD	NPNTTRAHEA	INFGSTSDD-	FQLQQPTKFT
AY502009	US/2002	SGICAFRXXL	TADV D GSH--	NPNTTRAHEA	VNFGSTSTD-	FQLQQPTKFT
KJ196286	JP/2002	SGICAFRGRL	TADV D GSHDD	NPNTTRAHEA	VNFGSTSTD-	FQLQQPTKFT
DQ438972	US/2005	TGICAFRGKI	SADVQNSHQD	GNNTTRAHEV	INFGSESED-	FQVGPPTKFT
JF970609	MEX/2007	TGICAFRGKI	SADAQNSHQD	GNNTTRAHEV	INFGSESDD-	FQVGPPTKFT
GQ266696	SUI/2008	TGICAFRGKI	SADVQSSHQD	NXNTTRAHEV	INFGSESED-	FQIGPPTKFT
GQ266697	SUI/2009	TGICAFRGKI	SADVHSSHQD	NXNTTRAHEV	INFGSESDD-	FQISPPTKFT
KJ156329	TW/2013	SGICAFRGRV	TA--ETDHRD	APGSTRAHEA	VNFRSNDND-	FQXXQPTKFT
AB983218	JP/2014	SGICAFRGRV	TA--ETDHRD	APGSTRAHEA	VNFRSNDND-	FQXXQPTKFT
KR270442	CHN	SGICAFRGRV	TA--QINQRD	APGSTRAQQA	VNLRISDND	FQXXQPTKFT
KR270445	CHN	SGICAFRGRV	TA--QINQRD	APGSTRAQQA	VNLRISDND	FQXXQPTKFT
KR270447	CHN	SGICAFRGRV	TA--QINQRD	APGSTRAQQA	VNLRISDND	FQXXQPTKFT

		Epitope D	Epitope E	Site III
		395	405	415
	
		↓		↓
	
KC597139	GUF/1978	PVGIXXXESG	HDFDQWALPR	YSGHLTLNMN
AY502009	US/2002	PVGIXXIESG	HEFDQWALPR	YSGHLTLNMN
KJ196286	JP/2002	PVGIXXIESG	HEFDQWALPR	YSGHLTLNMN
DQ438972	US/2005	PVGIXXXETG	HSFRQWDPPN	YSGALTLNMN
JF970609	MEX/2007	PVGIXXXETG	HPFRQWDLPN	YSGALTLNMN
GQ266696	SUI/2008	PVGIXXXETG	HPFNQWDLPN	YSGALTLNMN
GQ266697	SUI/2009	PVGIXXXETG	HPCNQWDLPN	YSGALTLNMN
KJ156329	TW/2013	PVGINXXDGD	HPFRQWELPD	YSGLLTLNMN
AB983218	JP/2014	PVGINXXDGD	HPFRQWELPD	YSGLLTLNMN
KR270442	CHN	PVGVNDDDDG	HPFRQWELPN	YSGELTLNMN
KR270445	CHN	PVGVNDDDDG	HPFRQWELPN	YSGELTLNMN
KR270447	CHN	PVGVNDDDDG	HPFRQWELPN	YSGELTLNMN

The VP1 amino acid numbering is based on the GII.17 prototype strain KC597139/GUF/1978. Dots indicate sequence identity among sequences presented in the alignment.

Amino acid positions corresponding to GII.4 predicted antibody binding regions epitope A, D, E [14] are marked by symbols: black circles, epitope A; black diamonds, epitope D; black square, epitope E.

Grey regions indicate the RGD motif [15] and site I, II, and III, which are putative histo-blood group antigen (HBGA)-binding sequences of GII NoV genotypes [16].

to 2009 formed Cluster II and those from 2013 to 2015 formed Cluster III.

All of the eight GII.17 variants reported here from Jiangsu province were in Cluster III and grouped with GII.17 strains from Taiwan (KJ156329, 2013) and Japan (AB983218, 2014) in both the polymerase- and VP1 region-based trees.

Twelve GII.17 capsid protein VP1 sequences from 1978 to 2014 were aligned, including nine GII.17 sequences released previously in the GenBank database and three sequences from representative strains obtained in our study. Sequence data showed that amino acid (aa) differences occurred mostly in the protruding P2 domain (Figure 4), particularly at aa 295–297, 376, 398–400 and 414 which form the predicted antibody binding regions in variant GII.4; epitope A (aa295–297, 376), epitope D (aa398–400) and epitope E (aa 414) [14]. In addition, some of the GII.17 strains, including the new GII.17 strains reported here, had a K289R mutation in the alert RGD/K-like motif, located at positions 287–289 [15]. In the three histo-blood group antigen (HBGA) binding sites, a single aa change (H353Q) at site I and a single aa (D380) insertion peripheral to site II occurred in the outbreak representative strains [16].

Discussion

Through the web-based surveillance system EPHEIM, increased levels of NoV activity were detected in late 2014 compared with previous season in Jiangsu province, China. Our findings suggest that this coincided with the emergence of a novel GII.17 variant, which caused most (16/23) of the NoV outbreaks reported between September 2014 and March 2015 in the province. Due to unavailability of laboratory data from four outbreaks in March, the number of GII.17-associated outbreaks could have been underestimated.

The novel variant was first noted in October of 2014, and spread rapidly throughout the province, causing an increasing number of outbreaks. During the course of the winter and early spring it became the predominant cause of NoV outbreaks (Figure 1), replacing the GII.4 Sydney variant starting from December 2014. Newly identified emerging variant that become predominant have been previously reported, in particular GII.4 variants, and these can also be associated with atypical increases in the incidence of acute gastroenteritis [17,18]. In Jiangsu province for example, an earlier dominant variant, GII.4-2006b, had been replaced by GII.4 Sydney in 2012/13 [19].

Based on the surveillance of NoV sporadic cases in this study, while the number of NoV GII.4 Sydney cases decreased in January 2015, sharp increases in the number of GII.17 cases were observed in February and March (Figure 2). In these two months the detection rate of NoV-positive specimens among specimens of sporadic cases of gastroenteritis also increased.

Phylogenetic analysis of GII variants, including GII.17 strains obtained in this and other studies, suggests that RNA recombination, a significant driving force in viral evolution [20,21] led to some characteristics of the novel GII.17 variant reported here. In the RdRp phylogenetic tree obtained in this study, the GII.17 strains formed three Clusters. Cluster I comprised the only GII.17 strain reported before the year 2000 as well as a GII.4 variant. Cluster II and Cluster III contained RdRp sequences of GII.17 strains, all found after 2003, and which additionally shared a high nucleotide identity with GII.3 reference strains characterised in the 1990s, especially those in Cluster III (Figure 3a). The fact that the only one GII.17 strain reported before 2000 had a GII.4 RdRp genotype, while most GII.17 NoVs detected in the 2000s possessed a GII.3-like RdRp genotype leads to hypothesise that the new GII.17 variant may be a recombinant strains with a GII.3-like RdRp gene and a GII.17 capsid gene. Interestingly, most GII.3 strains detected in the 2000s were recombinant strains, possessing a non-GII.3 RdRp genotype [22].

The VP1 protein P2 domain is the most exposed region of the viral particle and is well positioned to interact with potential neutralising antibodies and HBGA ligands. Mutations in this domain may have a significant effect on virus receptor binding and the host immune response to viral infection [23,24] and mutations in the P2 domain were observed in the new GII.17 variant. We speculate that, through the accumulation of mutations at several sites in the P2 domain, a new antigenic variant of the GII.17 lineage which gains the potential to escape herd host immunity could occur eventually. However, more studies, such as studies including virus-like particles (VLP)-HBGA binding assays, are needed to provide insights into the complex interaction between NoV GII.17 and their ligands.

The limitation of our study was that our results were not from nationwide surveillance but from Jiangsu province accounting for only 5.4% of the total China population. However, a similar situation to the one reported here was observed in 2012/13 with the emergence of the GII.4 Sydney variant, which was first detected in a NoV outbreak in late 2012, and soon afterwards led to large increases in NoV activity nationwide [19].

In conclusion, the new GII.17 variant which emerged in October 2014 appears to have subsequently increasingly caused NoV outbreaks in Jiangsu province, China. This study reveals that the variant presents a number of mutations in the P2 domain of VP1. Simultaneous dominance by GII.17 in outbreaks and sporadic infections indicates that this genotype might be established in Jiangsu Province. Nationwide surveillance for NoV outbreaks will be needed to understand epidemiological or outbreak trends related to the emergence of relatively rare GII.17 variants. Furthermore research into the mechanisms driving the evolution of NoV strains is also important for the development of effective prevention and control strategies.

Acknowledgments

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Conflict of interest

None declared.

Authors' contributions

Yefei Zhu, Miao Jin, Changjun Bao, and Fenyang Tang designed the study. Jianguang Fu, Jing Ai, Jun Zhang, Chao Shi, Qin Lin, and Zhaohu Yuan collected, analysed, and interpreted data. Jianguang Fu and Xian Qi characterised the specimens. Jianguang Fu, Jing Ai, and Yefei Zhu drafted the article. All authors reviewed and revised the first and final drafts of this manuscript.

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