Isolation and characterization of a new candidate human inactivated rotavirus vaccine strain from hospitalized children in Yunnan, China: 2010-2013

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Author contributions: Wu JY performed the majority of experiments and analyzed the data; Zhou Y, Zhang GM, Yi S, Mu GF, Yin N, Xie YP, and Lin XC performed the molecular investigations; Li HJ and Sun MS designed and coordinated the research; Wu JY and Zhou Y wrote the paper.

Supported by the CAMS Initiative for Innovative Medicine, No. 2016-12M-1-019 and No. 2016-12M-3-026; National Natural Science Foundation of China, No. 31700154; Major Science and Technology Special Project of Yunnan Province (Biomedicine), No. 2018ZF006; Science and Technology Project of Yunnan Province—general program, No. 2016FB034; and The State Project for Essential Drug Research and Development, the national “Twelfth Five-Year” plan, No. 2014ZX09102041004.

Institutional review board statement: All experiments were approved by the Institutional Animal Care and Use Committee of Institute of Medical Biology, CAMS (Kunming, China).

Institutional animal care and use committee statement: All experiments were conducted in accordance with the ethical guidelines for animal experiments and safety guidelines.

Conflict-of-interest statement: The authors declare that they have no conflicts of interest.

ARRIVE guidelines statement: The manuscript was revised according to the ARRIVE guidelines.

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Manuscript source: Unsolicited manuscript

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Received: May 24, 2018
Peer-review started: May 24, 2018
First decision: July 3, 2018
Revised: July 16, 2018
Accepted: July 31, 2018
Article in press: August 2, 2018
Published online: October 6, 2018

Abstract

**AIM**
To determine the distribution of rotavirus VP7 gene in...
hospitalized children in Yunnan, China.

METHODOLOGY
A total of 366 stool specimens were collected from hospitalized children in hospitals in Yunnan Province from September 2010 to December 2013. The genomic RNA electropherotypes and the G genotypes of the rotaviruses were determined. A phylogenetic analysis of the VP7 gene was performed. Rotavirus isolation was performed, and characterized by plaque, minimum essential medium, and all genes sequence analysis. Quantification of antibodies for inactivated vaccine prepared with ZTR-68 was examined by enzyme-linked immunosorbent assay and microneutralization assay.

RESULTS
Group A human rotavirus was detected in 177 of 366 (48.4%) stool samples using a colloidal gold device assay. The temporal distribution of rotavirus cases showed significant correlation with the mean air temperature. Rotaviruses were isolated from 13% of the rotavirus-positive samples. The predominant genotype was G1 (43.5%), followed by G3 (21.7%), G9 (17.4%), G2 (4.3%), G4 (8.7%), and mixed (4.3%) among a total of 23 rotavirus isolates. A rotavirus strain was isolated from a rotavirus-positive stool sample of a 4-month-old child in The First People’s Hospital of Zhaotong (2010) for use as a candidate human inactivated rotavirus vaccine strain and for further research, and was designated ZTR-68. The genotype of 11 gene segments of strain ZTR-68 (RVA/Human-wt/CHN/ZTR-68/2010/G1P[8]) was characterized. The genotype constellation of strain ZTR-68 was identified as G1-P[8]-I1-R1-C1-M1-A1-N1-T1-E1-H1. The VP7 and VP4 genotypes of strain ZTR-68 were similar to Wa-like strains.

CONCLUSIONS
A high prevalence of the G1, G2, and G3 genotypes was detected from 2010 to 2012. However, a dominant prevalence of the G9 genotype was identified as the cause of gastroenteritis in children in Yunnan, China, in 2013. A candidate human inactivated rotavirus vaccine strain, designated ZTR-68 was isolated, characterized, and showed immunogenicity. Our data will be useful for the future formulation and development of a new inactivated rotavirus vaccine in China.

INTRODUCTION
Rotavirus is a major cause of diarrhea in children aged below five years, and causes approximately 600,000 deaths in both developed and developing countries each year[1,2]. It is estimated that approximately 35000 children die per year in China from rotavirus, which is the second largest number of rotavirus deaths in the world[3,4]. Rotavirus is a non-enveloped RNA virus, and the viral genome contains 11 segments of dsRNA. The surface structural proteins VP7 and VP4 define the virus’ G and P genotypes, respectively[5]. Globally, human infections have been mainly caused by five G types, which are G1-G4 and G9[6,7]. Together, the genome encodes for six structural proteins and five nonstructural proteins. Rotaviruses are now classified into G-genotypes based on the relatedness of the genes encoding VP7[8,9]. Molecular sequencing of rotaviruses has also led to the development of a classification system. In this system, each internal gene is assigned a particular genotype based on established nucleotide identity cutoff percentages. Now, the acronym Gx-P[x]-I[x]-Rx-Cx-Mx-Ax-Nx-Tx-Ex-Hx is used to classify the VP7-VP4-VP6-VP1-VP2-NSP1-NSP2-NSP3-NSP4-NSP5/6-encoding segments[10].

In China, 70% of children’s deaths caused by rotavirus occur in rural areas each year, mostly among children aged below five years[11]. Many changes in China, such as economic development, improved access to medical care, improved sanitation, and the one child family rule, may all contribute to this decline in overall diarrheal deaths and in the number of rotavirus deaths as well. In western China, urbanization is ongoing, and the economy is still poor and developing. City Zhaotong, County Xiangyun and XishuangBanna are relatively impoverished districts in Yunnan province, located in southwestern China. In this study, stool samples from hospitalized children admitted for diarrhea were collected to determine the distribution of rotavirus. All hospitalized children were under five years old. The prevalence of rotavirus infection peaks during the winter season, when the temperature is low. We wanted to isolate and characterize a human virus for use as a
candidate human inactivated rotavirus vaccine strain and for further research. A rotavirus strain was isolated from a rotavirus-positive stool sample of a 4-month-old child in The First People’s Hospital of Zhaotong (2010), which was designated ZTR-68.

**MATERIALS AND METHODS**

**Ethics statement**

Collection and use of human stool specimens was approved by the Ethics Committee of the Institute of Medical Biology (YISHENGLUNZI [2016] 3), and was provided following children stool samples with written informed consent.

**Collection of stool specimens**

Stool samples were collected from hospitalized children in Yunnan, China. Group A rotavirus antigens were detected using an enzyme immunoassay (colloidal gold device assay, Rotavirus Group A Diagnostic assay, Colloidal gold device, Beijing Wantai Biological Pharmacy Enterprise Co., Ltd.) according to the manufacturer’s instructions. These samples were collected from September 2010 through December 2013.

**Rotavirus isolation**

An extract of child stool sample was agitated in 20% (wt/vol) PBS, pH 7.0 ± 0.2. The suspension sample was centrifuged for clarification at 8000 g for 20 min. The extract was treated with trypsin (10 μg/mL) for 60 min and inoculated onto a monolayer of MA104 cells for 90 min. After washing, the monolayer was maintained in serum-free minimum essential medium (MEM, Institute of Medical Biology, IMBCAMS, Kunming, China) supplemented with trypsin (1.5 μg/mL) and streptomycin (50 μg/mL) for 4 d. A viral lysate was treated with the freeze-thawing procedure three times and centrifuged at 7700 g for 30 min before infected in MA104 cells. The rotavirus strain ZTR-68 was isolated from a 4-month-old child hospitalized in The First People’s Hospital of Zhaotong, Yunnan Province, China. This patient had acute diarrhea associated with a positive result of fecal rotavirus antigen detected by Rotavirus Group A diagnostic assay (Colloidal gold device, Beijing Wantai Biological Pharmacy Enterprise Co., Ltd.). The virus isolation procedure was performed by previously described methods[12,13].

**Extraction and electropherotyping of viral RNA**

Rotavirus dsRNA was extracted from stools and infected cell cultures by using the MiniBEST Viral RNA Extraction Kit (TaKaRa Biotechnology, Dalian, China) in accordance with the manufacturer’s instructions. Rotavirus dsRNA was analyzed by 10% polyacrylamide gel electrophoresis (PAGE) analysis. For the analysis, 15 μL of RNA was electrophoresed, and the gels were stained with silver nitrate and photographed.

**Polymerase chain reaction amplification and sequence analysis of VP7**

The rotavirus dsRNA was used as a template for reverse transcription polymerase chain reaction (RT-PCR) using the Prime Script® One Step RT-PCR Kit (TaKaRa Biotechnology, Dalian, China) in accordance with the manufacturer’s instructions. The rotavirus VP7 general primers GeneralF/GeneralR designed in this study and previously described primers, Beg9/End9[14,15] (Table 1) were used. RT-PCR amplification was carried out in a 50 μL reaction volume containing PrimeScript 1 Step Enzyme Mix, 2 × 1 Step Buffer, 10 μmol/L primers, and template RNA (less than 1 μg). Thermocycling was performed for 30 min at 50 °C, 2 min denaturation step at 94 °C, and 30 cycles of 30 s denaturation step at 94 °C, 30 s annealing step at 55 °C-60 °C, and a 1 min extension step at 72 °C for each assay. The RT-PCR products were analyzed by electrophoresis on a 1% agarose gel (Invitrogen, Spain) in Tris-borate buffer containing ethidium bromide and visualized under ultraviolet light. The 2000 bp DNA ladder marker (Fermentase) was used as a size marker to estimate the lengths of the products. The PCR amplicons were purified using a column-based purification kit (OMEGA, the United States) and sequenced with an automated DNA sequencer (ABI 3730XL, the United States).

**Phylogenetic analysis of nucleotide and amino acid sequences**

The VP7 nucleotide and amino acid sequences were analyzed. The selected sequences were aligned with ClustalX[16]. The phylogenetic tree (neighbor-joining, maximum parsimony, maximum likelihood) was constructed using the MEGA 4 from dissimilar distances and pairwise comparisons with the Kimura 2-parameter model[17].

**Plaque assay**

The virus plaque was analyzed by previously described methods with modifications[12]. Virus stocks were activated with trypsin (10 μg/mL) for 60 min at 37 °C. The virus was diluted in MEM and 300 μL/well were plated on 6-well plates (Corning, the United States) with a monolayer of MA104. Then the inoculums were covered with 4 mL per well of MEM with 3.5% agarose. The

### Table 1 Primer sequences used in reverse transcription-polymerase chain reaction

| Primer     | Sequence                                      | References |
|------------|-----------------------------------------------|------------|
| GeneralF   | 5'- AATGATATGGTATTGAAATACCAC -3'              | This study |
| GeneralR   | 5'- TAATGATCTTGGACCTTTGGACAGA -3'             | This study |
| Beg9       | 5'- GAGAGAATTTCCGTTTGG -3'                   | [14,15]    |
| End9       | 5'-GGTCACATCATACAAATTCTAACCTAAG -3'          | [14,15]    |

Wu JY et al. New candidate human inactivated rotavirus vaccine strain

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WJCC | www.wjgnet.com | 428 | October 6, 2018 | Volume 6 | Issue 11 |
aggar was natural solidification, and then the plates were incubated at 37 ℃ for 4 d. Visualized plaques could be read after adding 2% neutral red in 1 mL MEM.

**Structural genes and nonstructural genes RT-PCR and nucleotide sequencing**

RT-PCR was performed for amplification of all structural and nonstructural genes using the primers listed in the supplementary data. Briefly, the extracted RNA genome of rotavirus was denatured at 96 ℃ for 5 min, and RT-PCR was carried out by using a Prime Script® One Step RT-PCR Kit (TAKARA, China). This involved an initial reverse transcription step of 30 min at 50 ℃, 2 min denaturation step at 94 ℃, and 30 cycles of 30 s denaturation step at 94 ℃, 30 s annealing step at 55 ℃-60 ℃ and 1 min extension step at 72 ℃ for each assay. PCR products were electrophoresed in 2% agarose gels containing ethidium bromide and visualized under UV. PCR amplicons were purified by a column-based purification kit (OMEGA, the United States) and sequenced with an automated DNA sequencer (ABI 3730XL, Uthe United States). Phylogenetic tree (neighbor-joining, maximum parsimony, maximum likelihood) was constructed using the MEGA 4 from dissimilar distances and pairwise comparisons with 1000 bootstrap replicates and the Kimura 2-parameter model.

The genotype of 11 gene segments of strain ZTR-68 (RVA/Human-wt/CHN/ZTR-68/2010/G1P[8]) was characterized. The genotype constellation of strain ZTR-68 was identified as G1-P[8]-I1-M1-A1-N1-T1-E1-H1.

**Electron microscopy**

The morphology of the rotavirus strain ZTR-68 stained by phosphotungstic acid negative staining was checked by electron microscopy (HTACHI H-600, Japan) to ensure that the virus contained triple-layered particles (TLP).

**Vaccine preparation and vaccination of mice**

Rotavirus strain ZTR-68 was cultivated in MA104 cells and the virus was purified by isopycnic CsCl gradient centrifugation as described previously[18]. Triple-layered particles and double-layered particles were collected separately. Both purified viruses were dialyzed in 10 mmol/L PBS pH 7.0 to remove CsCl. TLPs were inactivated with formalin at 400 μg/mL concentration at 37 ℃ for 120 h. The inactivated vaccine was stored at 4 ℃ before use. The vaccine was quantified for proteins by Lowry method [19] and rotavirus antigen was quantified by enzyme-linked immunosorbent assay (ELISA). Balb/c mice (18-22 g) were purchased from Institute of Medical Biology, CAMS (Kunming, China). Mice were separated in groups of 20, and vaccinated with 0 μg, 10 μg or 20 μg inactivated rotavirus vaccine (IRV) formulated with alum by injection into the abdominal cavity. Pre-bleed was taken before injection. Serums were collected 2 wk after injection. All experiments were approved by the Institutional Animal Care and Use Committee of Institute of Medical Biology, CAMS (Kunming, China) and conducted in accordance with the ethical guidelines for animal experiments and safety guidelines.

**Fluorescent cross-rotavirus antibody test**

MA104 cell line was maintained in MEM with 10% fetal calf serum (GIBCO, the United States) in 6-well microtiter plates (Coming, the United States). The cell monolayers were incubated at 37 ℃ in an incubator containing 50 mL/L CO2 under humid conditions, and infected with different viruses (strain Wa, SA11, S2, ZTR-68) with virus titer in 104-105 PFU/mL, and incubated for 16 h. The plates were fixed in chilled acetone (-20 ℃) for 15 min and air dried, then washed with 0.01 mol/L PBS buffer and blocked with 3% BSA for 1 h at 37 ℃. Mouse anti rotavirus strain ZTR-68 20 μg inactivated vaccine antibody (1:500 dilution, 0.5 mL volume) was added to microtitre plate wells and incubated at 37 ℃ for 1 h. After washing with 0.01 mol/L PBS, goat anti mouse FITC conjugate (MILLIPORE, 1:1000) was added and incubated for 1 h. The slides were examined under fluorescent microscope (Nikon, Japan) indicative of rotavirus.

**ELISA and microneutralization assay on sera**

RV-specific IgG antibodies were detected in sera on days 0, 14, and 28. Goat anti rotavirus antibodies (Millipore, the United States) coated 96-well ELISA plates overnight at 4 ℃. The plates were washed with PBS pH 7.0 and blocked with 5% BSA in PBS. The plates were washed and incubated with supernatant of rotavirus Wa infected MA104 cells (~105 PFU/well) for 1 h at 37 ℃. Serially diluted mouse serum was added in each well and incubated for 1 h at 37 ℃, then washed. HRP-conjugated rabbit anti mouse antibody (Millipore, the United States, 1:1000) was added. TMB (Tiangen, China) substrate was added for detection and was stopped with 2mol/L H2SO4. OD value was determined with an EIA reader (BioTek, the United States). Cutoff value was twofold of negative well value, and the antibody titer was defined as the highest dilution OD greater than the cutoff value. Geometric mean titer was given. Microneutralization assay was performed to measure rotavirus neutralizing activity as described previously [20]. Serial diluted mouse serum was added to 96-well plates and incubated with 1000 PFU of Wa per well for 1 h at 37 ℃. MA104 cells were prepared in 96-well plates as a monolayer and cultivated in MEM with 5 μg/mL trypsin (GIBCO, the United States) at 37 ℃ for 3 d. Then all 96-well plates of neutralization were transferred into 96-well plates MA104 cells grown in a monolayer and incubated at 37 ℃ for another 7 d. The plates were lysed by freeze/thaw three times. RV antigen was detected by ELISA method with HRP-labeled goat anti rotavirus antibody (Institute of Medical Biology, CAMS, Kunming, China). Cutoff value was twofold of cells well without virus value, neutralizing antibody titer was defined as the highest dilution that gave a lower than antigen cutoff value. Virus-only controls were performed to confirm the effectiveness of ELISA system. Graph analysis was performed using GraphPad Prism 5.02, and statistical
analysis was performed using Excel 2016 with the T test (two tailed). P values lower than 0.05 were considered statistically significant.

GenBank Accession Numbers
The sequences were deposited in GenBank under the following accession numbers: KM247264-KM247286. ZTR-68 genomic sequences: JX509930-JX509940.

RESULTS

Epidemiologic features of rotavirus in hospitalized children
Group A human rotavirus was detected in 177 of 366 (48.4%) stool samples using the colloidal gold device assay (Table 2) and in 133 of 177 (75.14%) by dsRNA-PAGE between September 2010 and December 2013 (Figure 1A). On an average, the prevalence of rotavirus during each year was 48.4% and 51.2% during the peak season. The age of the 177 patients ranged from one and a half months to four and a half years old (Figure 2A), although the majority were less than 2 years old (n = 157, 88.69%). The temporal distribution of rotavirus cases showed a significant correlation with the mean air temperature. The number of rotavirus cases peaked during the fall and winter seasons when the temperature began to decrease (Figure 2B and 2C).

All 11 segments were visible in 133 samples, corresponding to 75.14% of the rotavirus-positive samples and 36.33% of the total 366 stool samples. In total, 97 samples presented a long electropherotype, 36 presented a short electropherotype and none of the samples had a mixed pattern. The electropherotypes of the samples obtained from September 2011 to August 2012 presented a relatively higher level of diversity than the samples obtained from September 2010 to August 2011 and from September 2012 to December 2013. There was relative concordance between the electropherotypes and the distribution of G genotypes (Figure 1A and 3A).

Partial sequence analysis of VP7 gene
We determined the VP7 nucleotide sequences for G1, G2, G3, G4, and G9 for the G genotype isolates obtained during the study period (Figure 1B, 3A). It was determined that phylogenetically, the G3 isolates clustered in one lineage, the G1 isolates clustered in two lineages, and the G2 and G4 isolates clustered in one lineage. Several of the G1 isolates maintained a similarity of 97.4% to 99.9% for 2010, 2011, and 2013 in cluster 1-2, and the others maintained a similarity of 98.5% to 99.2% for 2011 in cluster 1-3. The G3 isolates analyzed during this study showed 99.8% to 99.9% similarity for 2010 and 2011, and they composed one cluster with the strains RV3, YO, and AU-1. In the case of G9, one sequence in 2012 (12N-T107) was related to the rotavirus strains BJ-CR7440, BJ-Q794, and MRC-DPRU1102, which originated in China and Zimbabwe. Three sequences in 2013 showed high similarity (99.9%) with each other and were related to strains that originated in Spain (98.2%), Ecuador (97.9%-98.1%), Saudi Arabia (97.5%-97.9%), Belgium (98.6%-98.9%) and Thailand (98.3%-98.9%). One G2 isolate 11O-T65 was related to strains RMC/G66, TB-Chen, and DS-1, which originated in Ecuador. One mixed genotype isolate (12N-T66) was found, which contained a mixed G2 and G9 genotype and was phylogenetically located between the G2 cluster and the G9 cluster. In the case of G4, two 2011 sequences (11D-T100, 11D-T101) were related to the Hochi (85.5%) and Gottfried (86.5%-86.6%) strains. In the case of G1, seven isolates analyzed in one cluster, closed to Wa and KU cluster. One G1 isolate 10O-T68R was related to strains Wa and KU but not in one cluster. This isolate then cultured in MA104 cells and purified by plaque (Figure S3), and was designated as strain ZTR-68.

Rotavirus isolation and characterization
Positive rotavirus antigen was diagnosed in the diarrheal stool of a 4-month-old child who was infected naturally with rotavirus and developed severe, acute diarrhea sustained for four days and was hospitalized in The First People’s Hospital of Zhaotong, Yunnan Province, China. PAGE analysis of the rotavirus dsRNA segments from stool specimen revealed a typical 4-2-3-2 pattern of rotavirus, specific to the group A rotaviruses. (Figure 3B) The presence of virus-like particles, almost 70 nm in diameter was confirmed in this specimen by electron microscopy. (Figure 3C, Figure S2) The rotaviruses from isolate extract 10O-T68R were adapted to infect MA104 cells and purified by plague technique. ZTR-68 showed a typical CPE of rotavirus and virus titers grew to over 10^{12} PFU/mL.

Genotype classification of ZTR-68
The ORF nucleotide sequences for the 11 genome
Figure 1  Polyacrylamide gel electrophoresis profiles and VP7 gene sequences of rotavirus strains isolated from diarrhea stools collected in Yunnan, China. A: Electrophoretic migration pattern of RNA from 24 rotavirus-positive stool samples during September 2010 through December 2013. Rotavirus strains Wa and SA11 were used as the markers. The viral RNAs were analyzed by electrophoresis in a 10% polyacrylamide gel and visualized by staining with silver nitrate. L: long electropherotype; S: short electropherotype. Genes 10 and 11 of rotavirus RNA of some samples from September 2012 to December 2013 were not clear in this pattern; B: The partial sequences determined in this study are in bold. The most closely related sequences found in the GenBank database are also included. References for the sequences used in VP7 gene comparisons marked with “G genotype/isolate/country/collected year”. The scale bar represents 5% nucleotide sequence difference. Bootstrap values of > 50% (for 1000 iterations) are shown.

Wu JY et al. New candidate human inactivated rotavirus vaccine strain
segments of cell culture-adapted ZTR-68 strain were determined. The sequences deduced in this study are either identical or show a few changes from those that are already in GenBank for ZTR-68. Genotypes were assigned for each genome segment based on the nucleotide percent identity cut-off values defined by the RCWG and by submission to RotaC. Our analysis shows that ZTR-68 can be classified as a G1P[8] strain. Specifically, the gene segment 1 (VP1) of ZTR-68 showed 90.7%-99.2% nucleotide identity with the R1 genotype cluster. The sequence identity was found to be 98.6%-99.2% with strains Dhaka16-03, Dhaka6, and

Figure 2 Age distribution and trends in number and proportion of gastroenteritis of rotavirus diarrhea cases from 2010 to 2013 in Yunnan, China. A: Age distribution of rotavirus diarrhea cases from 2010 to 2013; B: Trends in number and proportion of gastroenteritis by rotavirus antigen over the study period. The scale bar on the left represents the number of rotavirus cases and total diarrhea cases; C: The mean daily temperature of Yunnan from January 2011 to December 2013 compared to the number of rotavirus cases.
Dhaka12 of genotype R1 lineage. Percent nucleotide sequence identities with other genotypes were in the range of 77.6%-85.8%. Sequence analysis of the VP2 gene of rotavirus strain ZTR-68 showed genetic relatedness with the strains of the C1 genotype with 86.7%-95.2% nucleotide identity. The strain clustered with strains Wa, Dhaka6, YO, KU, and Gottfried. Percent nucleotide sequence identities with other genotypes (C2-C3) were in the range of 75.8%-77.5% (< the cut off value of 84%). Nucleotide sequence analysis of the VP3 gene of rotavirus strain ZTR-68 showed genetic relatedness with the strains of M1 genotype with 86.0%-95.2% nucleotide identity. The sequence identity was found to be 95.2%, 91.9%, and 86% with strains ST3, Wa, and Dhaka6, respectively. Percent nucleotide sequence identities with other genotypes (M2-M5) were in the range of 69.6%-71.6% (< the cut off value 81%). Sequence analysis of the VP4 gene of rotavirus strain ZTR-68 showed genetic relatedness with the strains of the P[8] genotype with 88.1%-97.3% nucleotide identity. The sequence identity was found to be 91.9%-97.1% with strains DRC88, B3458, MO, YO, and CH32 and 88.1%-89.1% with Wa and Hochi. Phylogenetically the strain ZTR-68 clustered with the strains of I1 genotype of the VP6 gene indicating 91.8%-97.7% nucleotide sequence identity and identified to be closer to strain Hosokawa. The nucleotide identity with other genotypes (I2-I7) ranged from 80.2% to 85.2%. Sequence analysis of the VP7 gene of rotavirus strain ZTR-68 showed genetic relatedness with the strains of G1 genotype with 91%-91.5% nucleotide identity. The sequence identity was found to be 91% and 91.5% with strains Wa and KU of genotype G1 lineage, respectively. Percent nucleotide sequence identities with other genotypes (G2-G10) were in the range of 53.7%-66.7% (< the cut off value 80%) (Figure 4).

IRV ZTR-68 induces increase of IgG and neutralizing antibodies levels

Immunogenicity of inactivated rotavirus vaccine prepared with ZTR-68 was examined by IgG ELISA and neutralization assay (Figure 5A and 5B). RV-specific
Wu JY et al. New candidate human inactivated rotavirus vaccine strain
Wu JY et al. New candidate human inactivated rotavirus vaccine strain
IgG and neutralizing antibodies were detected after one dose vaccination, and its levels increased after two vaccinations. Ten micrograms of IRV induced lower levels of total IgG antibodies and neutralizing antibodies compared to 20 μg IRV. Both 10 μg and 20 μg of IRV induced increased IgG and neutralizing antibody levels. After the first vaccination (at 0 d), differences between 10 μg and 20 μg IRV were not statistically significant both in the IgG titer and neutralizing antibodies titer. In addition, a significant difference (P = 0.0043) was found in the IgG titer between 10 μg and 20 μg at 14 d and 28 d, but there was no significant difference (P = 0.0565) in the neutralizing antibodies titer. IRV ZTR-68 also could induce increased cross-typing anti-rotavirus antibodies anti-rotavirus strain Wa (G1P[8]), S2(G2P[4]), and SA11(G3P[2]) by the fluorescent antibody test (Figure 5C).

**DISCUSSION**

Rotavirus is recognized as a major public health problem for children in China[3,21-24]. Diarrhea remains the major common cause of Chinese children hospitalizations[3,25-28]. In Chinese rural areas, rotavirus caused about 33% of hospitalizations for severe pediatric gastroenteritis. The numbers of outpatient visits and all diarrhea episodes in the community are 28% and 7%, respectively[3,12,13,27,29,30]. The percentage of diarrhea patients hospitalized for rotavirus is consistent with findings in other Asian countries. In a 2004 study of rotavirus surveillance throughout Asia, 44%-53% of children hospitalized for gastroenteritis in Myanmar, Thailand, Indonesia, and Vietnam were detected as rotavirus-positive[31,32]. Our study demonstrated that the G1, G2, and G3 genotypes had a high prevalence from 2010 to 2011. However, the G9 genotype was the predominant cause of gastroenteritis in children in Yunnan, China, in 2012 and 2013. The most prevalent rotavirus G genotype shifted from G1-G4 and G9 to G1 + G9. Although the reason for this trend is unclear, the emergence of the G9 rotavirus has been documented in many studies over the last decade[33-35]. Previous studies have reported G3 as the most common strain in China[3,28], however, in this study, we found that G1 was predominant (43.3%), followed by G3 (21.7%), and G9 (17.4%). Notably, from 2010 to 2011, G1 + G3 were detected, while there was no G9 detected. However, from 2012 to 2013, G1 + G9 were predominant, and no G3 was detected. In this study, we observed an increase in the incidence of G9 rotavirus. In addition, G9P[4] rotavirus strains have been described as emerging in several Latin American countries[33,35]. The results of the phylogenetic sequence analyses demonstrating that some of the G9 rotavirus sequences detected in this study were closely related to sequences that originated in Ecuador may support this view. Some G9 rotavirus isolates were obtained during the surveillance, and full genomic sequencing and characterization will be performed in future studies.

Rotavirus diarrhea presents a serious health burden in China. A previous study suggested that every child in
China experiences at least one episode of diarrhea due to rotavirus. The use of rotavirus vaccines should be part of a comprehensive strategy to control diarrheal disease. WHO/UNICEF recommend that all children receive solutions of low-osmolality oral rehydration salts to prevent and treat dehydration due to diarrhea. Rotarix and RotaTeq are two approved vaccines used worldwide. The Lanzhou Lamb Rotavirus Vaccine has already been licensed and used in China for several years. However, the immunization program for rotavirus is unpopular in the area of Yunnan. Our study may be useful for decisions regarding the need for rotavirus vaccines for use in rural China, such as Zhaotong, Xiangyun, and Xishuangbanna. It is assumed that G1 type rotavirus may be the most common strain in these areas. However, some issues should be considered, such as the reason for the G3 and G9 shift over time and the impact of the changes in the genetic diversity of rotavirus on the use of a rotavirus vaccine in China.

Whole-genome sequencing has shown that human RVs with the genotype constellation of G1/G3/G4/G9/G12-P[8]-I1-C1-M1-A1-N1-T1-E1-H1 or G2-P[4]-I2-R2-C2-M2-A2-N2-T2-E2-H2 are globally dominant. Based on all 11 rotavirus gene segments and the for-
mation of RCWG classification system, rotavirus ZTR-68 has been designated G1-P[8]-I1-R1-C1-M1-A1-N1-T1-E1-H1 and was known as Wa-like rotavirus group A (RVA). Human Wa-like RVA strains are believed to have a common ancestor with porcine RVA strains. The ZTR-68 and its plaque clone 3C were fully adapted to grow in MA-104 and Vero cell cultures (data not shown). ZTR-68 was identified as a Group A rotavirus, which was a Wa-like strain. The characterization of ZTR-68 G and P types are similar to Wa, but different in RNA electropherotype. It was interesting that our data demonstrated several rotavirus isolates with high similarity to ZTR-68 during the period of this study. It is assumed that G1P[8] type rotavirus may be the most common strain in the Zhaotong area. But some issues should be considered, such as the reason that the G3 and G9 type shifted as time went on, and the impact of the variation of genetic diversity of rotavirus for the use of rotavirus vaccine in China. This newly isolated human Group A rotavirus, ZTR-68 is being used to establish a new candidate rotavirus vaccine for use in studying the immune effectiveness.

Unfortunately, we were not able to acquire purified G3 and G9 strains after several passages in cell culture. Determining the circulation of various rotavirus strains in this area is of interest. However, no P types were identified during the genotype detection, which is another limitation of our study. The usefulness of next-generation DNA sequencing has previously been evaluated for the direct detection of bovine rotavirus from fecal samples. More meticulous rotavirus surveillance should be conducted in Yunnan, China. In this study, a high prevalence of the G1, G2, and G3 genotypes was detected from 2010 to 2012. However, the G9 genotype was the predominant cause of gastroenteritis in children in Yunnan in 2013. Our data will be useful for future efforts to formulate and develop a vaccine in China.

ARTICLE HIGHLIGHTS

Research background
Human rotavirus genotypes G1-G4 and G9 are major causes of acute gastroenteritis in children worldwide. Approximately 35000 children die per year in China due to rotavirus, which is the second highest rate of rotavirus deaths in the world.

Research motivation
In China, the elimination and control of the two kinds of diseases have experienced the process of alternating use of two kinds of vaccines, which proved to be feasible and effective. These successful experiences provide a theoretical basis and an approach to the development of an inactivated rotavirus vaccine. This newly isolated human Group A rotavirus, ZTR-68 is being used to establish a new candidate inactivated rotavirus vaccine strain and for further studies, which was designated ZTR-68. The genotype of 11 gene segments of strain ZTR-68 was identified as G1P[8]-I1-R1-C1-M1 A1-N1-T1-E1-H1. The VP7 and VP4 genes were sequenced and characterized, and showed immunogenicity.

Research results
Group A human rotavirus was detected in 177 of 366 stool samples using a colloid gold device assay. The temporal distribution of rotavirus cases showed significant correlation with the mean air temperature. Rotaviruses were isolated from 13% of the rotavirus-positive samples. The predominant genotype was G1 (43.5%), G2 (21.7%), G9 (17.4%), G4 (4.3%), G4 (8.7%), and mixed (4.3%). A rotavirus strain was isolated from a rotavirus-positive stool sample of a 4-month-old child in The First People’s Hospital of Zhaotong (2010) for use as a candidate human inactivated rotavirus vaccine strain and for further studies, which was designated ZTR-68. The genotype of 11 gene segments of strain ZTR-68 (RVA/Human-wt/CHN/ZTR-68/2010/G1P[8]) was characterized. The genotype constellation of strain ZTR-68 was identified as G1-P[8]-I1-R1-C1-M1-A1-N1-T1-E1-H1. The VP7 and VP4 genes of strain ZTR-68 was similar to Wa-like strains.

Research conclusions
Group A human rotavirus was detected in 48.4% stool samples. A high prevalence of the G1, G2, and G3 genotypes was detected from 2010 to 2012. However, a dominant prevalence of the G9 genotype was identified as the cause of gastroenteritis in children in Yunnan, China, in 2013. A candidate human inactivated rotavirus vaccine strain, designated ZTR-68 was isolated, characterized, and showed immunogenicity.

Research perspectives
Our data will be useful for the future formulation and development of a vaccine in China.

ACKNOWLEDGEMENTS
We are grateful to the members of the Pediatrics Department of the First People’s Hospital of Zhaotong City for introducing the patients and the collection of clinical samples.

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P- Reviewer: Islek A, Krishnan T S- Editor: Dou Y L- Editor: Filipodia E- Editor: Song H
