Molecular surveillance and genetic divergence of rotavirus A antigenic epitopes in Gabonese children with acute gastroenteritis

Gédéon Prince Manouana, Sandra Niendorf, Alexandru Tomazatos, Mirabeau Mbong Ngwese, Moustapha Nzamba Maloum, Paul Alvyn Nguema Moure, Gedeon Bingoulou Matsougou, Simon Ategboe, Elie Gide Rossatanga, C. Thomas Bock, Mirabeau Mbong Ngwese, Moustapha Nzamba Maloum, Paul Alvyn Nguema Moure, Gedeon Bingoulou Matsougou, Simon Ategboe, Elie Gide Rossatanga, C. Thomas Bock, Steffen Bormann, Benjamin Mordmüller, Daniel Eibach, Peter G. Kremsner, Thirumalaisamy P. Velavan, Ayola Akim Adegnik*

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**ABSTRACT**

**Background:** Rotavirus A (RVA) causes acute gastroenteritis in children <5 years of age in sub-Saharan Africa. In this study, we described the epidemiology and genetic diversity of RVA infecting Gabonese children and examined the antigenic variability of circulating strains in relation to available vaccine strains to maximize the public health benefits of introducing rotavirus vaccine through the Expanded Programme on Immunization (EPI) in Gabon.

**Methods:** Stool samples were collected consecutively between April 2018 and November 2019 from all hospitalized children <5 years with gastroenteritis and community controls without gastroenteritis. Children were tested for rotavirus A by quantitative RT-PCR and subsequently sequenced to identify circulating rotavirus A genotypes in the most vulnerable population. The VP7 and VP4 (VP8*) antigenic epitopes were mapped to homologs of vaccine strains to assess structural variability and potential impact on antigenicity.

**Findings:** Infections were mostly acquired during the dry season. Rotavirus A was detected in 98/177 (55%) hospitalized children with gastroenteritis and 14/67 (21%) of the control children. The most common RVA genotypes were G1 (18%), G3 (12%), G8 (18%), G9 (2%), G12 (25%), with G8 and G9 reported for the first time in Gabon. All were associated either with P[6] (31%) or P[8] (38%) genotypes. Several non-synonymous substitutions were observed in the antigenic epitopes of VP7 (positions 94 and 147) and VP8* (positions 89, 116, 146 and 150), which may modulate the elicited immune responses.

**Interpretation:** This study contributes to the epidemiological surveillance of rotavirus A required before the introduction of rotavirus vaccination in the EPI for Gabonese children.

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1. Introduction

Diarrheal diseases remain one of the major cause of illness among children <5 years, causing over 500,000 deaths worldwide each year, mainly in Africa and South Asia [1]. One of the most important etiological agents of gastroenteritis in infants and children is group A rotavirus (RVA). The non-enveloped, triple-layered viral particle has...
a genome of 11 double-stranded RNA (dsRNA) encoding six structural (VP1-VP4, VP6, VP7) and six non-structural (NSP1-NSP6) proteins [2]. The outer capsid layer is formed by the VP7 glycoprotein (G) and the VP4 spike protease-sensitive (P) protein, both encoded by genomic segments on which a binary taxonomical system of rotaviruses is based at intraspecific level, determining their classification into G and P genotypes [3].

The Rotavirus classification work group of the International Committee on Virus Taxonomy has currently identified 41 G and 57 P genotypes in mammal and bird species worldwide [3,4], although in humans only a few genotypes are responsible for the disease burden (G1-G4, G9, G12, P[4], P[6], P[8]) [5–8]. Additionally, genotype G5, G6 and G8 are considered relevant to human health in Africa and Asia [7]. In addition to rapid genetic drift caused by the error-prone RNA-dependent RNA polymerase, segment reassortment (genetic shift) is another major driver of rotavirus evolution, leading to new genetic combinations and enabling interspecies transmission [9,10]. Previous studies reported G1P[8] as the most frequent RVA strain globally [5,11], while G2P[4], G3P[8], G4P[8], and G9P[8] were reported to be common strains worldwide in the pre-vaccination era [11].

There are four WHO prequalified Rotavirus vaccines available thus far. Rotarix (GlaxoSmithKline Biologicals, Belgium) is a monovalent vaccine derived from a human G1P[8] strain and RotaTeq (Merck&Co., USA) is a pentavalent (G1-G4 and P[8]) containing human-bovine RVA reassortant strains [10]. Both vaccines were proven to be safe and efficacious. While Rotarix and RotaTeq are used in over 90 countries worldwide [12], WHO recently prequalified Rotavac (Bharat Biotech, India) and ROTASIIL (Serum Institute of India). Both Rotavac and ROTASIIL are in use only in India, whereas Rotavac is used in India and Palestine. Besides pronounced immune responses against the capsid protein, antibody responses against other rotavirus antigens such as VP6, VP2, NSP2 and NSP4 were also detected. However, the mechanisms by which antiviral immunity is acquired are not clearly understood [13]. Because the introduction of RVA vaccination requires a thorough documentation of disease burden, viral diversity, many countries have initiated RVA surveillance. In Africa, the first countries to introduce RVA vaccination were South Africa (2009) and Morocco (2010) [14]. Rotavirus evolution at the genotypic and sub genotypic levels helps to understand transmission dynamics, where new genotypes emerge through recombination of wild-type and vaccine strains, due to selection pressure (post-vaccine strain shift) [15], or due to gene reassortment [16,17]. Therefore, the question remains how antigenic drift can evade adaptive immunity and thus affect vaccine responses [18,19].

In Gabon, a national immunization program for RVA has not yet been implemented. Furthermore, the information on RVA diversity and burden is poorly documented. Some of the pathogenic RVA genotypes (G1, G2, G3, G12, G6, P[4], P[6], P[8]) were previously detected in four different cities of Gabon, as well as one emerging G6P[6] strain [20]. Thus, pursuing national and regional surveillance is important due to rapid evolution including zoonotic transmission with increased odds of emergence of novel strains. To this end, the aim of the present study is to describe epidemiology and genetic diversity of RVA in Gabonese children <5 years old and to investigate antigenic variability of circulating strains in relation to available vaccines.

2. Methods

2.1. Ethics statement

The study protocol was approved by the Institutional Ethical Committee of the Centre de Recherches Mériciales de Lamberéné (CERMEL) (CEI-CERMEL: 003/2017). Written informed consent was obtained from parents or legal representatives of the children.

2.2. Study population and sampling

Between April 2018 and November 2019, stool samples were collected from children under 5 years of age who were residents of semi-urban Lamberéné and its surrounding rural area. The population considered here is representative of the general population because children aged 0–5 years who presented to the outpatient clinics of the two major hospitals in Lamberéné were examined and included if they suffered from diarrhoea (defined as three or more liquid stools within 24 h during the previous three days) and lived in the study area (within a radius of approximately 20 km from Lamberéné). The sample size was calculated based on an estimated rotavirus prevalence among Gabonese children with diarrhoea (27.1%) from a previous [20]. The significance level was 0.05 (corresponding to a 95% confidence interval) with a precision of 0.07 for a sample size of 155 children. Rainy seasons (March–May, October–December) of similar length are interspersed between short (January and February) and long (June–September) dry seasons. Children with diarrhoea or history of diarrhoea within the last 24 h were recruited at outpatient department of two main hospitals (Hospital Albert Schweitzer and Centre Hospitalier Régional Georges Rawiri de Lamberéné). In addition, stool samples were randomly collected from healthy children of the same age without gastroenteritis who resided in the same compound as index cases. The collected stool samples were immediately transported to the laboratory and were stored in RNA later at –20°C for further use.
2.3. Rotavirus detection

The suspension of 200 mg stool in 1 ml of RNAlater solution was homogenized by vortexing and centrifuged. Viral RNA was then extracted from 140 μl of the supernatant with the use of QIAamp viral RNA Mini Kit (QIAGEN, Hilden Germany). All steps of the RNA isolation were performed following the manufacturer’s instructions. The genomic RNA was eluted in a total volume of 60 μl, concentrated immediately on a Qubit dsRNA XR Assay Kit and the Qubit 4.0 fluorometer (Invitrogen, Paisley, UK) and subsequently stored at –80°C.

The presence of RVA was detected using a one-step reverse transcription real time PCR (RT-qPCR) protocol for the amplification of the NSP4 gene [21]. Briefly, RNA extracts were first diluted and denatured at 95°C for 1 min, then RT-qPCR was carried out in a total volume of 12 μl using the SuperScript III/Platinum Taq OneStep kit (Invitrogen, Carlsbad, CA). The reaction mix contained PCR buffer, forward primer (RoA 25-s 5’-GCTTTAAGTT-CGTTCGCCAG), and reverse primer (RoA 26a-as 5’-ACTCAATGTGAAGGTGAGCT- probe 5’-VIC–ATCTTICGCCAGC–MGB and Platinum Enzyme mix. Samples with a cycle threshold (CT) ≤ 39 were considered positive.

2.4. Rotavirus genotyping

For the rotavirus G/P typing we used two RT-semi-nested PCR protocols, specific for each type, as described [22,23]. For the first round, pan-specific primers located close to the 3′ and 5′ end of the segments were used. In the second run, one of the pan-specific primer was used as a fluorescence labelled primer (with HEX for G types and FAM for P types). The second primer is specific for each F or G Type. This results in fluorescence labelled amplicons with different fragment length. The amplicons generated in the first and second PCRs were also independently checked using agarose gel electrophoresis. For each sample, the fluorescence labelled amplicons generated in the second PCR (nested-PCR round) were mixed with a fluorescence labelled ladder and then analyzed using capillary sequencer. Thus, specific peaks for G and P type could be determined. Additionally, the amplicons generated in the first round of the G/P typing were sequenced using the PCR primer used in the first PCR round. Those samples with a weak signal in the fragment length analysis and/or negative in the first PCR round, a P or G type specific PCR was additionally performed. Some samples showed a signal in the fragment length analysis, but no amplicons were detectable in the first PCR round. For these samples, a VP4- or VP7-long amplification was attempted to identify their genotypes. All amplicons were Sanger sequenced for further analysis. The RVA nucleotide sequences of this study are available under the following GenBank accession numbers: MZ966335 - MZ966498.

2.5. Sequence data analysis

Generated sequences were analyzed in Geneious Prime v2021.1.1 (Biomatters, Auckland, New Zealand). Genotype was assigned using NCBI BLAST and the VIPR typing tool for rotavirus A genotype determination (https://www.viprbrc.org). Nucleotide sequences of VP7 and VP4 were aligned with MAFFT implemented in Geneious and neighbor-joining phylogenetic trees were reconstructed using MEGA 7.0.26 [24] following substitution model using 1000 bootstrap iterations for evaluation of node support. Amino acid sequence similarity was calculated with the p-distance method. Phylogenetic trees were displayed with iTOL [25] and potential N-linked glycosylation sites were screened with NetNGlyc 1.0 Server (http://www.cbs.dtu.dk/services/NetNGlyc/).

2.6. Statistical analysis

Differences of RVA prevalence among children with and without diarrhea were tested using the Chi-square or Fischer exact test, with statistical significance set at two-sided p-value <0.05. Associations of RVA infection with potential risk factors were evaluated by conditional logistic regression and a multiple logistic regression model fitted using a stepwise backward procedure. The variables with a p-value ≥0.30 in the univariate analysis were removed sequentially. We adjusted for potential confounders that may influence the occurrence of RVA (e.g., gender, residence). All statistical tests were performed using R version 4.0.2 [26].

2.7. Role of funding source

The project is part of the GZ EI 1044/1-1 AOBJ 630127 grant funded by DFG. The Funders had any role in study design, data collection, data analyses, interpretation, or writing of report.

3. Results

3.1. Study population

A total of 244 children were recruited in the study. Of these 177 (73%) were symptomatic children aged between 0 and 59 months with a median age of 12 months, whereas the median age of healthy children was 24 months. Most participating children (78%; 190/244) were aged 0-24 months. A total of 152 (62%) participants lived in the semi-urban area. Male children were 57% (101/177) in the symptomatic group and 61% (41/67) in the asymptomatic group.

3.2. RVA Prevalence and seasonal distribution

RVA was detected in 98/177 (55%) of symptomatic children and 14/67 (21%) of controls. High RVA detection rate was associated with age, sex, and residence among symptomatic children (Table 1). Children between 0 and 6 months showed the highest proportion of infection (30/47, 64%), while the lowest detection rate was observed among children between 19 and 24 months (5/16, 31%). Females were more often infected (44/76, 58%) than males (54/101, 54%), with similar observations in children from rural (60%) compared to those from semi-urban settings (53%). In the control group, children between 7 and 12 months showed the highest proportion of infections (4/8, 50%). Males were more likely to be infected (10/41, 24%) than females (4/26, 15%), with similar observations in children from rural (31%) compared to those from semi-urban settings (7%).

Gabon, a tropical country has both wet and dry seasons. RVA infection among children with diarrhea was not significantly associated with either dry or wet season ($\chi^2 = 0.31$, p=0.5). However, a significant difference in RVA positive cases was observed between the cumulative long dry and wet seasons ($\chi^2 = 5.06$, p=0.02), with peak

| Table 1 | Rotavirus burden in the study population. |
|---------|----------------------------------------|
|          | RVA in Children with acute gastroenteritis cases n (%) | RVA in Children without acute gastroenteritis cases n (%) | p-value |
| Total    | 98/177 (55) | 14/67 (21) | <0.0001 |
| Age (months) |          |                    |        |
| 0-6      | 30/47 (64) | 0/7 (0) | 0.002 |
| 7-12     | 24/47 (51) | 4/8 (50) | 1 |
| 13-18    | 27/45 (60) | 0/4 (0) | 0.0345 |
| 19-24    | 5/16 (31) | 3/16 (19) | 0.6851 |
| 25-59    | 12/22 (55) | 7/32 (22) | 0.0205 |
| Gender    |          |                    |        |
| Female   | 44/76 (58) | 4/26 (15) | 0.0002 |
| Male     | 54/101 (54) | 10/41 (24) | 0.0016 |
| Residential area |          |                    |        |
| Rural    | 32/53 (60) | 12/39 (31) | 0.0062 |
| Semi-urban | 66/124 (53) | 2/28 (7) | <0.0001 |
RVA infection during September and October 2018 (turn of the seasons) and August-September 2019 (dry season) (Fig. 1a, 1b).

3.3. Genotype distribution and risk factors

In the 112 RT-qPCR positives, we identified two P types and five G types, either in single or mixed infections (Table 2). The P[8] (38.4%) was predominant, followed by P[6] (31%). The most prevalent G type identified was G12 (25%) followed by G1 (18%), G8 (18%), G3 (12%) and G9 (2%). The G type could not be assigned in 31 cases (26%) cases, while the P type identification failed for 34 positive samples (30%) (Supplementary Table S1). The frequent G/P combinations were G12P[6] (21%), G1P[8] (17%) and G8P[8] (15%) (Table 2).

Crude analysis indicated that nutrition, source of drinking water and diarrhea were associated with RVA. Children nourished with formula milk (OR=1.5; 95% CI: 0.55–4.12, p-value= 0.013) and drinking water from wells (OR=1.29; 95%CI: 0.61–2.80; p-value= 0.009) were at higher risk of acquiring a RVA infection. The RVA detection rate was significantly associated with diarrhea (OR=4.63; 95%CI: 2.45–9.29, p-value = <0.001). After adjusting for potential confounders, RVA infection remained significantly associated with diarrhea (OR=4.63; 95%CI: 1.5–8.58, p-value=0.004), and equally for semi-urban population (OR=0.46; 95%CI: 0.21-0.96, p-value= 0.04) (Supplementary Table S2).

3.4. Phylogenetic analysis with vaccine strains

RVA strains belonging to G8, G9, G12, P[6] and P[8] clustered within a single lineage per genotype (Figs. 2, 3). The Gabonese G1 clustered as lineage I and II, showing genetic homogeneity to African and Asian strains, respectively. The Gabonese G1 (lineage II) revealed a higher amino acid pairwise identity with the Rotarix G1 (98%) than the homologous component of RotaTeq. A low pairwise identity was observed between the protein sequence of Gabonese G3 strains and homologous component of RotaTeq (94-95%), as indicated also by their phylogenetic branching. Gabonese G3 strains grouped in lineage III with European, Indian and African strains. Additionally, we have detected the VP7 nucleotide sequence of a G3 strain (GAB/449) closely related to a bat-borne G3P[3] earlier detected in Gabon. The Gabonese G8 clustered in lineage I, showing a close relationship with Eastern Asian RVA (Japan), similarly to the G9 sequences of lineage VI (China and Japan). The G12 sequences formed a relatively homogeneous cluster within lineage III, along African, Asian and European strains.

For both P[6] and P[8] genotypes, their VP4 sequences clustered within lineage I and lineage III, respectively. However, the analysis of all symptomatic children revealed a pattern of P type segregation in Gabonese strains, indicative of their association with different G types (Figs. 2, 3). The Gabonese P[8] strains had 93-96% similarity with RotaTeq strain and 90% with the Rotarix P[8]. Furthermore, the P[8] associated with G1 were more similar with RotaTeq and Rotarix (96% and 90%, respectively) than G8P[8] (93-94% and 89-90%, respectively).

3.5. Analysis of VP7 and VP4 neutralizing epitopes

Vaccine efficacy can be undermined if structural differences accumulate in the antigenic epitopes of circulating RVA strains. We used comparative protein analysis to highlight potential antigenic differences between Gabonese RVAs and strains of vaccines that may be introduced. Mutation sites with potential for neutralization escape by monoclonal antibodies reside in the VP7 trimer on two structurally defined antigenic epitopes: 7-1 and 7-2 [27] (Fig. 4). The immunodominant 7-1 epitope is further divided into 7-1a and 7-1b. Overall, the highest number of amino acid differences was observed in the 7-1b subunit of VP7, followed by the 7-2 subunit and the 7-1a subunit. We observed only 5 of 29 amino acid residues completely conserved (W98, Q104, Q201, G264, K291). Most of these sites (W98, Q104, K291) were located on 7-1a. Due to the low viral load and high ct values, we could only obtain only sequence information from one asymptomatic child.

The VP7 antigenic epitopes of Gabonese G1 strains revealed four residue differences relative to Rotarix G1 and five changes compared to the RotaTeq G1. These sites were found on the epitopes 7-1a and
The analysis of G3 showed four to five amino acids different from the RotaTeq G3, distributed on 7-1b and 7-2 epitopes and up to 13 differences relative to Rotarix G1. Several exhibited features of escape mutants [28]. The change in the N94S residue was only observed in strain GAB/675, while the entire antigenic region 7-1b of the G3 strains matches the RotaTeq. All G3 strains carried a mutation at position 238 (epitope 7-1), with most having a substitution (K238N) that modulates N-glycosylation, with one exception in the GAB/449 strain (K238D), which was replaced by aspartic acid. In the 7-2 subunit, mutations at positions 147-148 can modulate responsiveness to specific antibodies; however, only position 147 had amino acid substitutions for both G1 and G3 strains. Most G1 strains had dissimilar residues (N/S147D), while GAB/449 (G3), phylogenetically related to the bat RVA, had a T147A substitution previously associated with immune escape [28,29].

Gabonese G9 strains had only two residues changes relative to all RotaTeq strains (sites 94 and 242), each located on the two subunits of the 7-1 epitope, but up to 12 amino acid difference in relation to the closest RotaTeq strain (G3). When compared to the Rotarix strain, the Gabonese G9 showed 14 amino acid changes. G12 strains were the most divergent, having a minimum of 9 different amino acids when compared to VP7 epitopes of vaccine strains. Relative to the closest RotaTeq strain (G3), G12 contained 15 residue changes. The most extensive epitope divergence was observed in relation to Rotarix G1, with 17 residues (Fig. 4).

Following the virion's trypsin activation in the intestine, the VP4 spike protein is proteolytically cleaved in two components: the globular head (VP8*) placed on top of the stalk (VP5*). We analyzed the region coding for the VP8* globular head, the main determinant of RVA P type containing four surface-exposed antigenic epitopes: 8-1, 8-2, 8-3 and 8-4 [31]. Among the Gabonese P[8], P[6] and all vaccine strains we found only two completely conserved sites in epitopes 8-2 (E180) and 8-3 (N132), of the 25 present in VP8* (Fig. 5). The Gabonese P[8] contained up to 24 identical amino acids compared to the P[8] of RotaTeq and 19 identical amino acids relative to P[8] of Rotarix. Six amino acids differences located mostly on epitopes 8-1, 8-3 and 8-4 were shared with both P[8] vaccine strains. The highest number of differences (10 residues) was noted between Gabonese G1P[8] strain GAB/675 and Rotarix P[8].

The epitope composition of the Gabonese P[6] strains relative to vaccine VP8* showed extensive dissimilarity, consistent with the levels of sequences divergence. P[6] strains associated with G3 contained 18-20 residue differences, whilst the G12P[6] contained 17-19 different amino acids.

4. Discussion

In preparation for introducing rotavirus vaccine by the EPI for Gabonese children, we investigated the prevalence and genetic characteristics of RVA among children with and without diarrhea over a 20-month period. The RVA detection rate in diarrheal cases is twice as high as previously reported in Gabon [20]. RVA was present to a greater extent in children with diarrhea than in asymptomatic ones, with similar or higher detection rate than in West/East Africa and South Asia [32,33]. As observed in various studies, high infection was observed in most children aged 0–24 months [33,34]. The significant association of RVA infection with the semi-urban environment could be related to population density and contact rates, but also to living conditions.
The AOR analysis showed no significant association with pit toilet use and RVA infections, this toilet type was present in about 90% of RVA positive households, in contrast to flush toilets (9%). The use of public taps as a source of drinking water was reported for more than half (55%) of the infected children, as opposed to water from the household tap (13%). Therefore, such factors may discretely contribute to the acquisition of RVA infections, especially in semi-urban areas with high population density. Over the sampling period, the proportion of

Fig. 2. Phylogenetic analysis of representative RVA strains based on a VP7 gene fragment (856 nucleotides). Gabonese strains analyzed are marked by black dots and vaccine strains are marked by red dots. Guineafowl RVA strain NIE13A1146 was used as an outgroup and bootstrap support >80% is indicated by asterisk. Tree scale bar represents number of substitutions per site.
infections differed statistically only between rainy seasons and cumulative long dry seasons. RVA infections peaked in September-October (2018) or August-September (2019), a pattern consistent with previous observations in Gabon [20], West Africa [35,36], Central Africa [37], Southeast Asia [38] and generally in the tropics [39].

The frequencies of G1 and G3 genotypes in the present study population are higher than previously reported in the country [20] and in
Fig. 4. Comparison of VP7 antigenic epitopes of Gabonese and vaccine strains. Amino acids of vaccine strains are indicated in bold.

Fig. 5. Comparison of VP4 (VP8*) antigenic epitopes of Gabonese and vaccine strains. Amino acids of vaccine strains are indicated in bold.
neighboring Cameroon [40], while G8 and G9 were detected for the first time in Gabon. In contrast to other Central African countries where G1 is dominant, this genotype was the second most common in our study (18%), while the dominant G12 (25%) was found in similar proportions [34,38]. Although G12 was found at a lower frequency in this study than in Cameroon (67.4%) [40], previous work by Lekana-Douki et al. [20] in Gabon also showed that G12 was less common (11.8%). In contrast to previous work by Lekana-Douki et al. [20], we observed P[8] to be the most frequent P genotype (38%). The other P genotype in our study is P[6] (31%), had a lower detection rate than those previously reported in the country (71.4%). This high incidence of P[8] has been observed not only in Central African countries, but also in Central and South-eastern Europe [41,42]. We observed a high proportion of G1P[8] (17%), although it was significantly lower than in neighboring country (Republic of Congo, 44%) [37]. G12P[8] and G12P[6] were found at lower frequencies compared to Cameroon [40]. In addition, we found G9P[8] and G8P[8], genotypes frequently detected in Southeast Asia [38]. These genotypes considered rare or uncommon, were also reported with low frequencies in other African countries [36]. 28 % of non-typeable types (Gx-Px) were observed and there were many non-typeable types among the P-types. All qPCR-positive samples that were negative in semi-nested RT-PCR were designated as non-typeable. There are two likely reasons that have been raised as limitations of this study. One probable reason is that qPCR has a high sensitivity compared to RT-PCR; this could well be observed in non-typeable samples that gave high ct values in the independently performed tests. The other reason could be due to mutations in the primer binding sites, which impairs annealing. However, the latter explanation is unlikely.

RVA zoonotic genotypes such as G8 (bovine), P[11] (bovine) and P[6] (porcine) are widespread in developing countries and cause infections in humans [43–45]. High diversity of RVA (including untypeable strains) in human populations, livestock or wildlife [46–48] are common. In our study, a G3 strain (GAB/449) obtained from a symptomatic child was phylogenetically closer to a Gabonese strain earlier discovered in a giant round bat (H. gigas, GenBank no. MN528121). Poor sanitary conditions and high contact rates with livestock and wildlife may contribute to spill-over-spillback transmission patterns. Consequently, the rate of infection is also higher (~20%) compared to middle- and high-income countries (~5%) [43].

The phylogenetic analysis of Gabonese G types revealed well supported terminal clades indicating close relatedness with strains of Asian origin. Although P[8] is widespread in Africa, we cannot rule out the possibility of cross-border transmission in Gabon due to the constant influx of migrants from Asia and other countries. The phylogenetic sub-lineage segregation of Gabonese VP4 in discrete clusters following a pattern of G type association (e.g., P[6] with G3 and G12, P[8] with G1 and G8) might be a result of reassortment.

Strains of P[8] contained amino acid disparities in their VP8* epitopes in accordance with their G1, G8 and G9 combinations, although with notable exceptions. G1P[8] was one of the frequent type combinations and most divergent from vaccine cognates, similarly to the less frequent G9P[8]. These genetic and antigenic differences should be considered when planning immunization, since the chosen vaccines will shape the antigenic landscape for circulating RVA. The antigenic G1-VP7 epitopes were mostly conserved and homogeneous with the vaccine strains. Of all three VP7 epitopes, the RVA genome was apparently conserved with vaccine strains, with an exception in G1P[8] (GAB/675), which revealed the N94S substitution associated with immune escape [28,49]. The VP7 epitopes of the Gabonese G3 strains had a relatively low number of different residues compared to RotaTeq G3. Despite this observation, the presence of K238N (epitope 7-1b) may be of concern as it affects glycosylation and has been shown to neutralize RVA in mammals by monoclonal antibodies and hyperimmune sera [46,50]. K238N has also been reported previously in European and North African G3 strains [10,51], with additional sites (70–72) contributing to N-linked glycosylation.

In the case of RVA immunization, a vaccine’s failure to eradicate the virus can result in selective pressures that enhance the pathogen’s ability to evade host immunity [47]. Viral evolution is often linked to travel and domesticated livestock populations in the community that facilitate the virus shedding and recombination of RVA strains in antigenically naive hosts [48,52]. Although many studies have examined RVA burden in symptomatic patients, this study investigated children without gastroenteritis. The comparison of the RVA genotype distribution between symptomatic and asymptomatic children would have been useful. However, due to the low viral load, we could only obtain a sequence from the control group (asymptomatic children). Another limitation of our study is the small size of the asymptomatic group. This is because sampling began in the second year of the study and therefore precludes observation of the seasonal distribution of RVA cases among asymptomatic children.

The presence of the main pathogenic strains in Gabon, namely G1 and G3 in combination with P[6], the emerging G12 and the less common G9 with antigenic divergence, makes the implementation of vaccination necessary and urgent. This study provides the genetic data needed to assess the epidemiological situation prior to the introduction of vaccination in the Gabonese population, where a wide variety of RVA are present, including strains that are likely to be of zoonotic origin. Additional longitudinal studies at temporal and spatial scales are needed to determine the functional significance of these genetic divergences observed in Gabonese RVA strains and to establish a link with available vaccines, proven to be efficacious.

Contributors

AAA designed, supervised, and coordinated the study procedures. GPM performed all experimental procedures and drafted the manuscript. AT supported experimental procedures, analyzed genetic data, and drafted the manuscript. GPM, MMN, MNM, PANM, GBM, SA and EGR were involved in sampling procedures and collating the baseline data. SN and CTB performed RVA genotyping for independent verification in another laboratory and supported with data analysis. SB, BM, DE, and PGK contributed to the study design and patient recruitment. TPV and AAA contributed to the materials. TPV supervised experimental procedures on RVA screening and genotyping, data analysis, and revised the manuscript from the first draft. All authors read and approved the final version of the manuscript. TPV, AAA, GPM, SN, AT verified the underlying data in this manuscript.
STROBE Statement—Checklist of items that should be included in reports of **case-control studies**

| Item No | Recommendation |
|---------|----------------|
| **Title and abstract** | 1. (a) Indicate the study’s design with a commonly used term in the title or the abstract<br>Page 2 – Lines 40 – 66<br>(b) Provide in the abstract an informative and balanced summary of what was done and what was found<br>Page 2 – Lines 40 – 66 |
| **Introduction** | **Background/rationale** 2. Explain the scientific background and rationale for the investigation being reported<br>Pages 4 and 5 Lines 112 – 161 |
| **Objectives** | 3. State specific objectives, including any prespecified hypotheses<br>Pages 4 and 5 Lines 134 – 161 |
| **Methods** | **Study design** 4. Present key elements of study design early in the paper<br>Page 5, 169 – 186 |
| **Setting** | 5. Describe the setting, locations, and relevant dates, including periods of recruitment, exposure, follow-up, and data collection<br>Page 5, 169 – 186 |
| **Participants** | 6. (a) Give the eligibility criteria, and the sources and methods of case ascertainment and control selection. Give the rationale for the choice of cases and controls<br>Page 5, 169 – 186<br>(b) For matched studies, give matching criteria and the number of controls per case<br>Page 5, 169 – 186 |
| **Variables** | 7. Clearly define all outcomes, exposures, predictors, potential confounders, and effect modifiers. Give diagnostic criteria, if applicable<br>Supplementary data Table S2 |
| **Data sources/ measurement** | 8. For each variable of interest, give sources of data and details of methods of assessment (measurement). Describe comparability of assessment methods if there is more than one group<br>Page 5, 169 – 186, (cases and controls) Supplementary data Table S2 (cases) |
| **Bias** | 9. Describe any efforts to address potential sources of bias<br>Not applicable |
| **Study size** | 10. Explain how the study size was arrived at<br>Page 5, 169 – 186 |
| **Quantitative variables** | 11. Explain how quantitative variables were handled in the analyses. If applicable, describe which groupings were chosen and why<br>Page 7, 235 – 242 |
| **Statistical methods** | 12. (a) Describe all statistical methods, including those used to control for confounding<br>Page 7, 235 – 242<br>(b) Describe any methods used to examine subgroups and interactions<br>Page 7, 224 – 233<br>(c) Explain how missing data were addressed<br>Not applicable<br>(d) If applicable, explain how matching of cases and controls was addressed<br>(e) Describe any sensitivity analyses |
| **Results** | **Participants** 13. (a) Report numbers of individuals at each stage of study—eg numbers potentially eligible, examined for eligibility, confirmed eligible, included in the study, completing follow-up, and analysed<br>Page 5, 169 – 186, Page 9, 10, 256 – 289<br>(b) Give reasons for non-participation at each stage<br>Not applicable<br>(c) Consider use of a flow diagram |
| **Descriptive data** | 14. (a) Give characteristics of study participants (eg demographic, clinical, social) and information on exposures and potential confounders<br>Page 5, 169 – 186, (controls and cases) and Supplementary Table S2, (Cases)<br>(b) Indicate number of participants with missing data for each variable of interest<br>Supplementary Table S2 |
| **Outcome data** | 15. Report numbers in each exposure category, or summary measures of exposure<br>Page 9, 10, 256 – 289 (controls) and Supplementary Table S2 (cases) |
| **Main results** | 16. (a) Give unadjusted estimates and, if applicable, confounder-adjusted estimates and their precision (eg, 95% confidence interval). Make clear which confounders were adjusted for and why they were included<br>Page 9, 10, 256 – 289 and Supplementary Table S2<br>(b) Report category boundaries when continuous variables were categorized<br>Supplementary Table S2<br>(c) If relevant, consider translating estimates of relative risk into absolute risk for a meaningful time period<br>Page 9, 10 256 – 289 |
| **Other analyses** | 17. Report other analyses done—eg analyses of subgroups and interactions, and sensitivity analyses<br>Page 8 – 10, 248 – 371 |
| **Discussion** | **Key results** 18. Summarise key results with reference to study objectives<br>Page 10 – 371 – 379 |
| **Limitations** | 19. Discuss limitations of the study, taking into account sources of potential bias or imprecision. Discuss both direction and magnitude of any potential bias<br>Page 13, 450 – 469 |
| **Interpretation** | 20. Give a cautious overall interpretation of results considering objectives, limitations, multiplicity of analyses, results from similar studies, and other relevant evidence<br>Pages 11 – 13, 373 – 469 |

(continued)
Data sharing statement

All data relevant to the study are included in the article or uploaded as supplementary information.

Declaration of Competing Interest

The authors declare no conflict of interest.

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Supplementary materials

Supplementary material associated with this article can be found in the online version at doi: 10.1016/j.ebiom.2021.103648.

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