Prevalence of Enteropathogens and Virulence Traits in Brazilian Children With and Without Diarrhea

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The use of molecular diagnostics for pathogen detection in epidemiological studies have allowed us to get a wider view of the pathogens associated with diarrhea, but the presence of enteropathogens in asymptomatic individuals has raised several challenges in understanding the etiology of diarrhea, and the use of these platforms in clinical diagnosis as well. To characterize the presence of the most relevant bacterial enteropathogens in diarrheal episodes, we evaluated here the prevalence of diarrheagenic *E. coli* pathotypes, *Salmonella* spp., and *Yersinia enterocolitica* in stool samples of children with and without diarrhea using real-time quantitative PCR (qPCR).

We found that the presence of genetic markers associated with bacterial pathogens was significantly higher in stool samples from the diarrhea group compared to the control (*P* < 0.001). Bacterial loads in samples positive for *eae* and *aggR* markers were also determined. Compared to samples from asymptomatic children, a significantly higher number of copies of the *eae* gene were found in diarrhea samples. Also, the presence of genetic markers associated with STEC strains with clinical significance was evaluated in *eae*-positive samples by high-throughput real-time PCR. The data presented herein demonstrated that asymptomatic children of an urban area in Brazil might be enteropathogen reservoirs, especially for STEC.

**Keywords:** enteric pathogens, *Escherichia coli* pathotypes, diarrheal children, healthy children, molecular diagnostics

**INTRODUCTION**

Diarrheal disease is the second most important cause of child morbidity and mortality in the world, and is frequently due to contaminated food and water. Worldwide, 780 million individuals lack access to safe drinking water and 2.5 billion lack adequate sanitation. Diarrhea is widespread throughout developing countries. In low-income countries, children under 3 years old experience on average three episodes of diarrhea every year (World Health Organization, 2017).
Rotovirus and diarrheagenic Escherichia coli (DEC) are the two most common etiological agents of moderate-to-severe diarrhea in low-income countries (World Health Organization, 2017). Other pathogens such as Cryptosporidium, Shigella, Campylobacter, Salmonella, and Yersinia species also cause human bacterial gastroenteritis, but location-specific etiological patterns also need to be considered (Bottone, 1999; Freitas Neto et al., 2010; Marinou et al., 2012; Nunes et al., 2012; World Health Organization, 2017; Lima F. M. et al., 2019; Leli et al., 2020).

Enterotoxigenic E. coli (ETEC), enteroinvasive E. coli (EIEC), typical and atypical enteropathogenic E. coli (tEPEC and aEPEC), typical and atypical enterotoaggregative E. coli (tEAEC and aEAEC), and Shiga toxin-producing E. coli (STECE and STEC), and its subgroup enterohemorrhagic E. coli (EHEC), enteroinvasive E. coli (EIEC), and diffusely adherent E. coli (DAEC) comprise DEC pathotypes (Kaper et al., 2004; Gomes et al., 2016). These pathogens use many common features to colonize the intestinal mucosa and trigger disease, but its initiation, progression, and complications vary considerably (Croxen et al., 2013). Thus, ETEC isolates carry the enterotoxins LT and/or ST on plasmids, as well as colonization factors (CFs). tEPEC, aEPEC, and LEEl-positive STEC strains (EHEC) carry the locus of enterocyte effacement (LEE) which contains eae, the gene that encodes intimin, but only tEPEC has the bundle-forming pilus gene (bfp). STEC strains, LEE-positive and -negative, carry the Shiga toxin genes (stx1 and/or stx2). Some virulence factors for EAE are common for a certain group of isolates are found on the pAA plasmid, and the presence of the aggR gene differentiates tEAEC from aEAEC. EIEC and also Shigella spp. have the ability to invade cells mainly through the pINV plasmid and acquired additional virulence attributes mainly from chromosomal pathogenicity islands (PAIs) (Croxen et al., 2013).

Nevertheless, the designation of certain isolates into a pathotype is complicated due to the plasticity of the E. coli genome resulting in the emergence of hybrid strains; besides, DEC epidemiology varies in different regions of the world and may depend on the host, environmental factors (Croxen et al., 2013). It is important to note that DEC pathotypes have been largely disregarded because of the lack of laboratory capabilities (Piazza et al., 2010; Miliewsky et al., 2016). Moreover, in several case-control studies, it shows almost equal distribution, especially in EPEC and EAEC pathotypes (Bueris et al., 2007; Hernandes et al., 2009; Gomes et al., 2016; Imdad et al., 2018; Dias et al., 2020).

Besides LEE-positive pathogens and Shigella, Salmonella also uses the type III secretion system (T3SS) apparatus to deliver effector proteins into the host cell, and invA, an inner-membrane component, is critical to the functioning of Salmonella T3SS (Galán et al., 1992; Ginocchio and Galán, 1995). In addition, the most common virulence-associated gene in pathogenic Yersinia enterocolitica proved to be ystA, which can therefore be considered the target gene to be amplified to evaluate the presence of pathogenic biotypes (Peruzy et al., 2017).

The use of multiple pathogen detection platforms in epidemiological studies have allowed us to get a broader view of the pathogens associated with diarrhea, but the presence of enteropathogens in asymptomatic individuals has raised several challenges in understanding the etiology of diarrhea, as well as the use of these platforms in clinical diagnosis (Liu et al., 2012; Lima A. A. M et al., 2019). The use of real-time quantitative PCR (qPCR) to determine bacterial loads in stool samples appears as an alternative to associate a pathogen with diarrheal cases.

To determine the most prevalent bacterial pathogens of diarrhea in children, we used qPCR to detect and quantify the presence of EPEC, ETEC, EAEC, STEC, EIEC, Shigella, Salmonella, and Y. enterocolitica. Furthermore, in a subset of eae-positive samples, we analyzed by high-throughput real-time PCR the presence of genetic markers related to EHEC strains with clinical significance.

**MATERIALS AND METHODS**

**Sample Collection, Ethical, and Biosafety Procedures**

Stool samples were obtained from 110 children with diarrhea (54 boys and 56 girls) and 150 children without diarrhea (71 boys and 79 girls), aged from 1 month to 7 years old. Stool samples were collected from March 2008 through November 2010 at Hospital Municipal do Tatuapé, Hospital Infantil Cândido Fontoura, Hospital Infantil Menino Jesus and Centro Educatacional Unificado (CEU-Butantã) in Sao Paulo city, SP, Brazil. Children did not display illnesses or comorbidities other than diarrhea, and they were not under antibiotic treatment for at least 3 months prior to sample collection. Diarrhea was defined as three or more unformed stools in the 24h prior to enrollment. The Ethics Committee of the Biomedical Science Institute at the University of Sao Paulo approved this study (2006/743). All procedures followed the management of biosafety and biosecurity, since Biomedical Science Institute at São Paulo University guarantees excellent conditions for workers and population, and environmental protection from biohazards exposure.

**DNA Extraction**

Total DNA from 200 mg of stool was obtained by using a QIAamp DNA Stool Mini Kit (QIAGEN) according to the manufacturer's instructions, resuspended in 100 μL of ultrapure water and stored at −80°C until use. DNA concentrations were determined by spectrophotometer (NanoDrop 2000, Thermo Scientific, USA), and 5 μL of each DNA sample were checked for integrity on 1% agarose gel.

**Identification of the Main Enteric Pathogens by qPCR**

qPCR assays for genes eae (attaching and effacing lesions), ipaH (enteroinvasive mechanism), lt (heat-labile toxin), st (heat-stable toxin), aggR (aggregative fimbriae regulator), invA (innmembrane component of the Salmonella T3SS apparatus), and ystA (enterotoxin YstA Yersinia virulence plasmid) were carried out in duplicate and performed in a total volume of 25 μL, containing 2X TaqMan universal master mix (Applied Biosystems, USA), 10 μM of each primer, 10 μM of TaqMan probe, and 2 ng of DNA. Amplifications were performed in a
thermal cycler programmed as follows: denaturation at 95°C for 10 min, followed by 45 cycles of two steps: denaturation at 95°C for 15 s and an annealing temperature at 60°C for 1 min. The primer/probe sets used are shown in Supplementary Table 1. A standard curve was also derived using 10-fold DNA dilutions from the reference strains with their respective primer pairs. Amplifications were adjusted to R² > 0.900. A P < 0.05 was considered statistically significant. A sample was considered positive for a target gene when the detected fluorescence generated a curve above the background fluorescence, which was established by the Rotor-Gene Q6000 analytical software (Qiagen, Brazil).

**Differentiation of Enteric Pathogens by PCR**

Samples positive for eae (n = 57) and ipaH (n = 15) by real-time PCR were further screened for the eae, bfpA (bundle-forming pilus for typical EPEC), ipaH, lacY (lactose permease), and stx1 and stx2 (Shiga toxins) genes by multiplex PCR according to the protocol of Aranda et al. (2004) and Pavlovic et al. (2011) for differentiation between EPEC/STEC and EIEC/Shigella samples.

**Characterization of the eae-positive Samples by High-Throughput Real-Time PCR**

A LightCycler® 1536 (Roche, Meylan, France) was used to perform high-throughput real-time PCR amplifications as described previously (Delannoy et al., 2012b), except that 1 µL of sample DNA was used in each reaction for a final reaction volume of 2 µL. The thermal profile was modified as follows: 95°C for 1 min, followed by 45 cycles of 95°C for 0 s, and 58°C for 30 s. All ramp rates were set to 2°C/s. The primers and probes used targeted genes encoding intimin (eae, eae-alpha, eae-beta, eae-gamma, eae-epsilon, and eae-theta), O-group-associated genes for the top 7 serogroups (wzxO26, rfbE O157, wzy O145, wxx O93, wbdd O11, wzx O121, wzx O45), flagellar antigens H11, H19, H2, H28, H7, and H8 (fliCH11, fliCH19, fliCH2, fliCH28, fliCH7, and fliCH8), urease (ureD), effector proteins translocated by T3SS (EspL2 [ent or espL2]), NleB [nleB], NleE [nleE], NleA [nleA], NleF [nleF], Efa1 [efa1], EspN [espN], EspK [espK], and EspM1 [espM1]), and other genetic markers including Z6065, Z2098, Z2099, terE, hlyA, ehxA, and pagC and markers related to the CRISPR loci SP_O157_A, SP_O157_B, SP_O157_C, SP_O26_C, SP_O26_D, SP_O26_E (Perelle et al., 2004; Fratamico et al., 2009, 2010; Bugarel et al., 2010; Delannoy et al., 2012a, 2013, 2015, 2016; Piazza et al., 2013). The wecA gene was used as a reference genetic marker for E. coli (Bugarel et al., 2011). An inhibition control (IC) was included in each sample to check for potential inhibition of the PCR reaction due to intrinsic characteristics of the sample. IC is a recombinant pBluescriptIIISK+ plasmid containing the dsb gene from Ehrlichia canis (Michelet et al., 2014). The plasmid was added to each sample at a concentration of ~0.3 pg/µL. Primers and probe specific for the Ehrlichia canis dsb gene were used to detect IC (Michelet et al., 2014).

**Statistical Analyses**

We compared the presence of pathogen between groups using the Fisher exact test. A comparison of bacterial loads in eae- and aggR-positive samples between samples from children with and without diarrhea was performed with a Mann-Whitney U-test. A P < 0.05 was considered statistically significant. All statistical analyses were performed in Stata v.12.1.

**RESULTS**

**Prevalence of Enteropathogens in Stool Samples Obtained From Children With and Without Diarrhea**

A total of 260 stool samples (110 from children with diarrhea and 150 children without diarrhea) were collected from public hospitals and schools, respectively, in Sao Paulo city (the largest city in South America) from March 2008 through November 2010. The groups studied consisted of children between 1 month and 7 years old, representing much of childhood. Each group was further divided into 3 age groups: 0–11 months, 12–23 months, and older than 24 months. The pathogens detected through their representative genetic markers in the stool samples from children with and without diarrhea of different age are reported in Table 1.

No genetic markers were detected in 39 of 110 (35%) children with diarrhea and 92 out of 150 (61%) without diarrhea. The presence of genetic markers associated with bacterial pathogens was significantly higher in stool samples from the diarrhea group (71/110) compared to the control (58/150) (P < 0.001). A significant increase in the number of samples positive for one pathogen was found in the diarrhea samples (46/110) compared to asymptomatic group (39/150) (P = 0.0036). Although we observed statistical differences between groups in samples with 3 or 4 pathogens, in both groups, the number of samples is low, not allowing further conclusions (Table 2).

**Bacterial Loads in eae- and aggR-Positive Samples**

Quantification of bacterial loads was shown to be a promising tool to identify the etiological agent of diarrheal diseases. Of all the genes evaluated, aggR and eae were the markers most found in diarrheal and asymptomatic children (71 and 56 samples, respectively). We compared the copy number of eae and aggR genes in all the samples positive for these markers in both diarrhea and asymptomatic groups (Figure 1). A significantly higher copy number of eae was found in diarrhea samples (P < 0.039); no differences in aggR copy number were found between groups (P = 0.43).

**Prevalence of EHEC-Associated Virulence Markers in eae-positive Stool Samples From Children With and Without Diarrhea**

In the control group, we detected 13% eae-positive samples and 11% stx-positive samples, while in the diarrhea group, 34% eae-positive samples, and 27%-stx positive samples were present,
TABLE 1 | Distribution of virulence genes from enteric pathogens in fecal samples from children with and without diarrhea.

| Pathogens detected | Genetic markers | Asymptomatic group (n = 150) | Diarrhea group (n = 110) | P-value |
|--------------------|-----------------|-----------------------------|-------------------------|---------|
|                    |                 | 0–11 months (n = 54) | 12–23 months (n = 59) | 24 months (n = 37) | Total (n = 55) | 12–23 months (n = 50) | 24 months (n = 5) | Total (n = 5) |         |
| STEC               | eae, stx1/stx2  | 2 | 4 | 0 | 6 | 2 | 9 | 0 | 11 | 0.19 |
| EAECE              | aggR            | 15 | 9 | 1 | 25 | 10 | 8 | 0 | 18 | 0.016 |
| EPEC Typ           | eae, bfp       | 0 | 1 | 0 | 1 | 1 | 0 | 0 | 1 | 0.44 |
| EPEC atyp          | eae            | 0 | 0 | 0 | 0 | 3 | 1 | 0 | 4 | 0.003 |
| ETEC               | tt/st          | 3 | 1 | 0 | 4 | 1 | 3 | 0 | 4 | 0.38 |
| EIEC               | ipaH, LacY     | 0 | 1 | 2 | 3 | 4 | 3 | 1 | 8 | 0.088 |
| STEC EAECE         | eae, aggR, stx1/stx2 | 3 | 1 | 0 | 4 | 1 | 2 | 0 | 3 | 0.25 |
| STEC EPECtyp       | eae, bfp, stx1/stx2 | 1 | 1 | 0 | 2 | 3 | 5 | 0 | 8 | 0.049 |
| STEC ETEC          | aggR, tt/st    | 3 | 3 | 0 | 6 | 1 | 0 | 0 | 1 | 0.013 |
| EAEC EPECtyp       | aggR, eae, bfp | 2 | 1 | 0 | 3 | 0 | 0 | 0 | 0 | 0.02 |
| EAEC EIEC          | aggR, ipaH, LacY | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0.182 |
| EAEC EPEC atyp     | aggR, eae      | 2 | 1 | 0 | 3 | 0 | 0 | 0 | 0 | 0.098 |
| EAEC Salmonella    | aggR, invA     | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0.18 |
| STEC EAEC ETEC     | eae, stx1/stx2, aggR, tt/st | 1 | 1 | 0 | 2 | 0 | 0 | 0 | 0 | 0.057 |
| STEC EAECE EPECtyp | eae, aggR, stx1/stx2, bfp | 0 | 1 | 0 | 1 | 2 | 0 | 1 | 3 | 0.2 |
| STEC EPECtyp EIEC  | eae, stx1/stx2, bfp, ipaH, LacY | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0.18 |
| STEC EPEC ETEC     | eae, stx1/stx2, bfp, tt/st | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0.18 |
| STEC EPEC EIEC     | eae, stx1/stx2, tt/st, invA, LacY | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0.18 |
| STEC EPEC ETEC EIEC| eae, stx1/stx2, bfp, tt/st, ipaH, LacY | 0 | 0 | 0 | 0 | 2 | 0 | 2 | 0 | 0.98 |

P < 0.05 was considered statistically significant.

TABLE 2 | Association of enteric pathogens in fecal samples from children with and without diarrhea determined by virulence genes presence.

| Number of pathogens detected | Asymptomatic group (n = 150) | Diarrhea group (n = 110) | P-value |
|-----------------------------|------------------------------|-------------------------|---------|
|                             | Total | Total |              |         |
| None                        | 92    | 39    | >0.001       |         |
| Positive for 1 or more      | 58    | 71    | >0.001       |         |
| pathogens                   |       |       |              |         |
| 1 pathogen                  | 39    | 46    | 0.0036       |         |
| 2 pathogens                 | 16    | 16    | 0.1734       |         |
| 3 pathogens                 | 3     | 7     | 0.0354       |         |
| 4 pathogens                 | 0     | 2     | 0.0486       |         |

DISCUSSION

A better knowledge of the distribution of pathogens linked to stool contamination is important for evaluating the pathogen loads in the environment as well as the associated health risks (Zhang et al., 2016). Over the last few decades, new molecular methods have become essential in allowing the rapid detection and characterization of pathogens (Miliwebsky et al., 2016).

The present study was performed to determine the molecular epidemiology of enteropathogens among children under seven years old with and without acute gastroenteritis in Sao Paulo, SP, Brazil. Virulence markers for DEC pathotypes were largely detected confirming some other surveys in Latin America and China (Ochoa et al., 2009; Moreno et al., 2010; Zhang et al., 2016; Lima A. A. M et al., 2019; Lima F. M. et al., 2019). Different results were reported in Western Kenya among bacterial pathogens, where two species of Shigella were the most prevalent (Swierzewski, 2013).

Our results showed that the genes encoding virulence factors for Salmonella and Yersinia were detected in very low frequencies, in agreement with another survey in counties near Sao Paulo city (Lima F. M. et al., 2019).

Herein, DEC-associated genetic markers were detected in 71 out of 110 (65%) children with diarrhea and 58 out of 150 (39%) without diarrhea and significant differences were observed between the two groups. These results are in agreement with other studies in South American countries. In Northeast Brazil, the presence of EPEC and EAEC were consistently associated...
with diarrhea (Maranhão et al., 2008; Moreno et al., 2010). Furthermore, EAEC and STEC showed by multivariate logistic regression analysis significant odds of being associated with a risk for diarrheal diseases as well as norovirus, adenovirus, rotavirus, and Giardia infection. Furthermore, EAEC enteric infections are associated with a high clinical severity score plus moderate-to-severe dehydration (Lima A. A. M et al., 2019).

The shifting in DEC epidemiology has been described in different studies and geographical regions (Pérez et al., 2010; Torres, 2017; Imdad et al., 2018; Pérez-Corrales and Leandro-Sandi, 2019) including Brazil, where until the 1990s, EPEC, mainly the typical EPEC serotypes producing EPEC adherence factor (EAF), was the main cause of infantile diarrhea (Gomes et al., 1991; Rosa et al., 1998; Regua-Mangia et al., 2004), and also Salmonella species (8%), ETEC (7%), and Shigella species (5%) were associated with diarrhea (Gomes et al., 1991), but it seems that they are becoming more and more rare, as presented herein and described elsewhere (Franzolín et al., 2005; Bueris et al., 2007; Moreno et al., 2010; Lima A. A. M et al., 2019; Lima F. M. et al., 2019; Ori et al., 2018).

The distribution of genes encoding either for one single pathogen in the diarrhea group (66%) and in control (67%) or in multiple coinfections, 34 and 33%, respectively, was similar in the present study. However, except for Salmonella plus E. coli and Yersinia plus E. coli, we cannot assure that we detected different strains, since they were not isolated from stools. Nevertheless, coinfections have been described, suggesting associations among potential enteropathogens in the etiology of diarrhea (Orlandi et al., 2006; Lima F. M. et al., 2019). Indeed, some hybrid strains possessing typical genes from different pathotypes (heteropathogens) have recently emerged (Santos et al., 2020). For example, the O104:H4 strain associated with the severe German outbreak in 2011 was a hybrid of both EAE and EHEC (Brzuszkiewicz et al., 2011). More recently, enterohemorrhagic E. coli hybrid pathotype O80:H2 has emerged in Europe as an important serotype responsible for hemolytic uremic syndrome (HUS) associated with extraintestinal infections in children (Cointe et al., 2018). Also, coinfections have been reported, detecting in stool samples the presence of EPEC and EAEC and S. enterica houtenae and EAEC (Lima F. M. et al., 2019).

The incidence of intestinal infection caused by STEC is low in the Brazilian population (de Souza et al., 2011; Ori et al., 2018; Lima F. M. et al., 2019). Thus, it was surprising to find more EHEC-associated genetic markers in the asymptomatic group than in the diarrhea group, since these genetic markers are usually more associated with E. coli strains of greater clinical significance (Delannoy et al., 2012a,b, 2016; Piazza et al., 2013). This suggests that these potential EHEC strains are not the etiological agent of diarrhea in this sampled group, which remains to be investigated.

Current methods for either screening or detecting of enteropathogens by PCR, multiplex PCR, and qPCR have been used worldwide, given the high sensitivity and specificity, which are satisfactory parameters to ensure an adequate diagnosis (Panchalingam et al., 2012). However, the presence of several pathogens in diarrhea samples, as we observed, and the high frequency of enteropathogens in asymptomatic children has made it difficult to use these methods as a tool to identify the etiological agent of diarrhea.

Here, we found a significantly higher copy number of eae in diarrhea samples compared to samples from asymptomatic children, suggesting that LEE-containing pathogens could be the etiological agents responsible for diarrhea in children. Even though we did not find differences between groups in the number of copies for aggR, our data must be complemented with a larger number of samples and/or selection of other EAEC markers to support the use of qPCR for molecular diagnosis for this pathogen.

Our study had the following limitations. The number of samples was limited and we did not search for other...
relevant pathogens, such as rotavirus, norovirus, protozoa, and helminthes. For bacterial loads, we only evaluated two markers, which might be insufficient, considering that DEC harbors several virulence determinants that can be used as a target for diagnosis. Another limitation of the study was that we did not isolate the pathogenic strains. This would be helpful to identify if asymptomatic children might shed the bacteria and possibly be contagious.

In conclusion, we found that the presence of genetic markers associated with bacterial pathogens was significantly higher in stool samples from the diarrhea group compared to the control. Also, compared to samples from asymptomatic children, there was a significantly higher copy number of the eae gene in diarrhea samples. The data presented herein demonstrated that asymptomatic children of an urban area in Brazil might be enteropathogen reservoirs, especially for STEC, thus favoring their widespread.

**DATA AVAILABILITY STATEMENT**

All datasets generated for this study are included in the article/Supplementary Material.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Ethics Committee of the Biomedical Science Institute at the University of Sao Paulo. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

**AUTHOR CONTRIBUTIONS**

VM, VN, SD, PF, RP, and MA-C participated in the design of the study. VM, VN, SD, PF, MF, RP, and MA-C participated in data analysis. MF, RP, SD, and PF participated in writing the manuscript. VM and VN carried out the qPCR experiments. GA carried out the multiplex PCR experiments. SD and PF carried out the high-throughput real-time PCR experiments. All authors read and approved the final manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb.2020.549919/full#supplementary-material

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