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Cohort profile: The Jundiaí Zika cohort (JZC), a pregnancy and birth cohort in Sào Paulo state, Brazil

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ABSTRACT

Purpose The Jundiaí Zika Cohort (JZC) is a prospective pregnancy and birth cohort setup in the State of São Paulo, Brazil, to investigate the epidemic of cases of microcephaly and other neurological disorders, presumed to be associated with Zika virus (ZIKV) infection.

Participants A total of 748 women with high-risk pregnancies were recruited in the period of March 2016 to August 2017.

Findings to date Baseline sociodemographic and medical data were collected at recruitment from 737 pregnant women. Biological samples (ie, blood, saliva and urine) were collected from 695 of the pregnant women (94.3%), of whom 53 (7.6%) were ZIKV-positive on subsequent testing by reverse transcription polymerase chain reaction (RT-PCR) in urine. Biological sample (ie, blood, saliva, urine and cerebrospinal fluid) were collected within 10 days of birth from 409 (57.4%) of the liveborn infants, of whom 19 (4.6%) were ZIKV-positive on subsequent testing by RT-PCR in urine. All remaining biological specimens, as well as colostomy, umbilical cord and placental samples, have been stored in a secure biorepository. Antenatal and postnatal imaging studies and neonatal anthropometry were carried out.

Future plans The JZC provides a unique data set which will continue to be explored to study the effects of pregnancy comorbidities on Zika virus infection during pregnancy, the long-term outcomes of children with congenital Zika infection and how physiotherapy and group interventions can improve outcomes for congenitally-infected children. All women in the cohort have reached the end of their pregnancy and currently the oldest children are 2 years old. The study will continue until all the children reach their third birthday (April 2021).

INTRODUCTION

The Jundiaí Zika Cohort (JZC) is an ongoing multidisciplinary longitudinal study which is following a cohort of pregnant women and their children to study the effects of prenatal Zika virus (ZIKV) exposure. The cohort was set-up in response to the clusters of cases of severe microcephaly and associated neurological disorders that were reported in areas affected by ZIKV in the Northeast of Brazil in October 20151 and that provoked the Public Health Emergency of International Concern declared by the WHO on 1st February 2016.2 At this point in time, the São Paulo Research Foundation (FAPESP) encouraged all researchers who had been part of the Brazilian Genome Project to submit thematic research project proposals addressing the potential causes of the cluster of cases of microcephaly.

Laboratory confirmation of autochthonous ZIKV transmission in Brazil was first established in the north-eastern states of Pernambuco, Rio Grande do Norte and Bahia, and later in other states of the central-west and south-eastern regions of Brazil. When the JZC was set-up, there was no published data about the ZIKV epidemic in São Paulo state. It was not known how the differences in climate and socioeconomic status between this south-eastern state and the poorer and more tropical north-eastern regions, where the ZIKV epicentre was focused, would influence the

Strengths and limitations of this study

► The Jundiaí Zika Cohort (JZC) is one of the few prospective Zika cohort studies that has recruited both asymptomatic and symptomatic pregnant women and therefore benefits from having a large control group.

► The high-risk profile of the pregnant women provides a unique opportunity to study comorbidities that may contribute to, or potentially be protective for, the development of negative sequelae associated with ZIKA virus (ZIKV) exposure.

► The prevalence of ZIKV reverse transcription polymerase chain reaction positivity was relatively low in this study population resulting in small numbers for estimating absolute and relative risks.

► Because high-risk pregnant women are at higher risk of developing some of the outcomes of interest, this may have reduced the power to detect differences between Zika-exposed and unexposed dyads.
epidemic. The study site of Jundiaí in the south-eastern state of São Paulo was therefore chosen in order to explore these variations.

This cohort profile aims to describe the Jundiaí Zika Cohort including: The context behind its creation, materials and methodology, recruitment and follow-up of pregnant women and children as well as some of its preliminary results. The oldest children of the cohort, as of November 2018, are 2 years old; all the children will be followed up until their third birthday.

COHORT DESCRIPTION

Study site

The JZC is housed in the paediatric department of the Jundiaí Medical School in the city of Jundiaí, São Paulo state. Jundiaí is 50 km northwest of the city of São Paulo (figure 1) and has a population of 405,740 inhabitants. It has a relatively high Human Development Index ranking 11th out of the total 5565 municipalities in the country. The climate in the area is humid subtropical, according to the Köppen classification, with a mean annual temperature of 20.9°C. The majority (64%) of the land in the municipality of Jundiaí is considered rural and 31% of this is made up of the Japi Mountain, a Biosphere Reserve of Atlantic Forest recognised by UNESCO since 1994. Jundiaí University Hospital is the only public maternity facility in the municipality of Jundiaí and is the local referral centre for high-risk pregnancies. In 2017, 37.6% of babies born in the municipality were born in this hospital; the remaining 62.4% were born in one of the three private hospitals in Jundiaí.

Given the severity of the phenotype assumed to be associated with prenatal exposure to ZIKV and the urgent need to establish a cause, the JZC, like many other Zika Cohorts in Brazil, commenced without any formal funding on 1st March 2016. It later received seed funding from The London School of Hygiene and Tropical Medicine prior to obtaining formal FAPESP funding for a thematic research project over a year after the initiation of the project on 1st April 2017 (figure 2).
Following ethical approval for the study from the research ethics committee of Jundiaí Medical School (protocol number 1446577), written informed consent was obtained from participating women for themselves and for future follow-up of their child.

This cohort profile describes the recruitment and follow-up of the pregnant women and children in the JZC. The oldest children of the cohort, as of November 2018, are 2 years old; all the children will be followed up until their third birthday.

**Study participant characteristics**

During the recruitment period, 752 women were enrolled in the study (figure 3). Of these, 15 (2%) were excluded as the baseline questionnaire was not fully administered. The mean age of women was 27.5 years (13 to 46). Three hundred and sixty-eight (53.7%) of the women were of white ethnicity, 238 (34.7%) mixed race or brown (known as ‘para’ in Portuguese), 66 (9.6%) black, 11 (1.6%) Asian and 2 (0.3%) indigenous. Five hundred and sixty-nine (77.2%) of the women reported being married or living with their partner (table 1). During pregnancy, 231 (33.9%) of the women had diabetes and 129 (18.9%) had hypertension. The presence of other risk factors, or comorbidities, is detailed in table 2.

Of the 737 pregnant women with full baseline information collected, 695 (94.3%) provided biological samples (ie, blood, saliva and urine from each) for ZIKV testing. Of the 695 women whose urine was tested by RT-PCR, 53 (7.6%) were ZIKV-positive. Due to financial constraints during the outbreak, blood and saliva collected from both women and neonates have been stored in a biorepository for future testing. Twenty-six (3.4%) women were lost to follow-up during pregnancy (ie, before the birth outcome), and there was one first trimester maternal death. Of the 710 women who were followed until the birth outcome, 23 (3.2%) had foetal losses and 687 (96.9%) had live births, of which 25 (3.6%) were twin pregnancies.

Of the 712 live births, 376 (52.7%) were female (table 3). Of these, 655 (92.0%) had anthropometry at birth (a minimum of weight and head circumference measured at birth); the mean birth weight among live births was 3001 g (590 g to 4525 g), mean length at birth was 47.5 cm (28.5 cm to 58.5 cm) and mean head circumference was 33.7 cm (22 cm to 38.5 cm). The number of liveborn infants who had urine tested for ZIKV RT-PCR within 10 days of birth was 409 (57.4%). Of these, 19 (4.6%) were positive. Notably, no neonates presented with symptoms consistent with postnatal ZIKV infection in the first 10 days. Of the 712 live births, 271 (38.1%) babies were seen in the JZC paediatric clinic between 0 to 2 months of age, 186 (26.1%) between 3 to 4 months of age, 173 (24.3%) between 5 to 6 months of age, 218 (30.6%) between 7 to 12 months of age and 74 (10.4%) between 13 to 24 months of age. So far, no infants have been followed up beyond 2 years of age.

**External validity and possible participation biases**

The recruitment of only high-risk pregnant women brought advantages, as discussed earlier, in both logistics and maximisation of follow-up rates, however, these advantages also introduced limitations in external validity when generalising JZC findings to the general pregnant population of Brazil. Moreover, as recruitment was carried out in a specific population of pregnant women that were users of a particular health service, it is possible that there are systematic differences between those recruited and those not recruited, which we were
not able to measure, and these may have introduced bias. However, when comparing the sociodemographic profile of the pregnant women in our cohort and the profile of pregnant women living and using public maternity facilities in the State of São Paulo at the time of the study,9 10 we can see that they are quite similar. For example, around half of the women were white, around half had vaginal deliveries and the majority had finished
high school and were co-habiting and/or married to their partner (table 1).

Furthermore, as there is no known association between high-risk pregnancies and congenital Zika syndrome (CZS), we do not anticipate that the restricted selection criteria of the cohort would have a significant impact on our interpretation of the underlying biology of congenital ZIKV infections.

**FOLLOW-UP**

**Pregnant women**

Women who at enrolment reported not having had any symptoms consistent with ZIKV infection during their pregnancy (and who were currently asymptomatic), according to the WHO ZIKV clinical case definition,\(^\text{11}\) were followed up as per Group 1 and women who were symptomatic at any point during pregnancy were followed up as per Group 2 (figure 3). Women were asked to contact the research team and attend the hospital if they experienced any symptoms consistent with ZIKV infection at any point in their pregnancy so that biological samples could be collected as detailed below. In addition, trained volunteers carried out weekly telephone follow-up consultations at pre-arranged times that were convenient for the women until the time of birth to ask specifically about the occurrence of any ZIKV symptoms and any women who had experienced symptoms were advised to go to the hospital.

Regardless of symptom occurrence, women in both groups were seen 14 to 21 days after enrolment for biological sample collection and then in 2 to 3 monthly intervals thereafter (sample collection details and laboratory procedures are described below). Antenatal ultrasound scanning was carried out in months 3, 5, 7 and 8 in asymptomatic (Group 1) women and monthly in symptomatic (Group 2) women at the São Paulo foetal medicine centre (CPMF). Additionally, where malformations, signs of congenital ZIKV infection or intra-uterine growth restriction were found, ultrasound scanning was carried out weekly at Jundiaí University Hospital or CPMF (figure 4).

**Neonates, infants, children**

Women whose birth outcomes resulted in a live birth were invited to come to the JZC paediatric follow-up clinic.

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At each stage of follow-up there was a significant non-response rate (see figure 3). Pregnant women and mothers who did not attend a scheduled follow-up appointment were contacted by phone, mobile or Facebook. In addition, JZC teams visited and spoke to staff at the basic health units in Jundiaí and requested them to refer any children belonging to the cohort. Despite these efforts, for many families, a complex and precarious social situation precluded them from being able to attend the follow-up appointments.

DATA COLLECTION

The JZC is a multidisciplinary study containing a rich range of information regarding the follow-up and outcomes of the JZC mothers and children, both exposed and unexposed to ZIKV during pregnancy. The main health, medical and laboratory data collected to date are listed in table 4.

The initial baseline questionnaire administered to the pregnant women was designed by the JZC researchers before any internationally standardised collection tool had been developed. In August 2016, the WHO created a standardised questionnaire in order to streamline data collected by all the Zika cohorts; new questions which were not already in the initial JZC tool (mainly related to environmental exposures, for example mosquito repellent and bednet use, type of housing and water sources) were added.

Four specialist foetal medicine doctors carried out the antenatal ultrasound scans at the CPMF with Voluson S10, Voluson E6 e Voluson E8 GE Healthcare equipment.

Table 3 Description of liveborn infants in the Jundiaí Zika cohort

<table>
<thead>
<tr>
<th>Liveborn infants (n=712)</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Twin pairs</td>
<td>3.5%</td>
<td>(n=25)</td>
</tr>
<tr>
<td>Sex (female)</td>
<td>52.8%</td>
<td>(n=376)</td>
</tr>
<tr>
<td>Delivery method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vaginal</td>
<td>47.1%</td>
<td>(n=330)</td>
</tr>
<tr>
<td>Caesarean section</td>
<td>50.3%</td>
<td>(n=352)</td>
</tr>
<tr>
<td>Forceps</td>
<td>2.6%</td>
<td>(n=18)</td>
</tr>
<tr>
<td>Unknown</td>
<td>1.7%</td>
<td>(n=13)</td>
</tr>
<tr>
<td>Weight at birth in grams (mean and range)</td>
<td>3001 (590 – 4525)</td>
<td></td>
</tr>
<tr>
<td>Length at birth in cm (mean and range)</td>
<td>47.5 (28.5–58.5)</td>
<td></td>
</tr>
<tr>
<td>Head circumference at birth (mean and range)</td>
<td>33.7 (22–38.5)</td>
<td></td>
</tr>
<tr>
<td>Biological sample tested for ZIKV RT-PCR at&lt;10 days</td>
<td>57.4% (n=409)</td>
<td></td>
</tr>
<tr>
<td>Positive urine ZIKV RT-PCR in first 10 days of life</td>
<td>4.6% (n=19)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Percentages for all categories were calculated with exclusion of those with missing data from the denominator.

RT-PCR, reverse transcription polymerase chain reaction; ZIKV, Zika virus.

Figure 4 Full follow-up protocol for Jundiaí Zika cohort study (01 March 2016 to 23 August 2017), Jundiaí, São Paulo, Brazil. JZC, Jundiaí Zika Cohort; ZIKV, Zika virus.
Foetal anthropometry was carried out and gestational age estimated.

Anthropometry was carried out in neonates in the first hour of life unless their condition was unstable. Head circumference was measured using a non-elastic tape measure placed between forehead and occiput, weight was measured using digital scales and length using a recumbent baby length scale. Z-scores for these measurements were then calculated using the INTER-GROWTH-21st curves which take into account the sex and gestational age of the neonate. Gestational age was calculated using first-trimester ultrasound when available and last menstrual period when not available.

Clinical samples and laboratory procedures

Blood, saliva and urine were collected from the women and children by nursing staff, healthcare assistants and auxiliary nurses (all registered at the nursing professional registration body of the State of São Paulo (COREN-SP)). They completed a baseline questionnaire detailing sociodemographic details, past medical history, family history, past obstetric history (parity, miscarriages, mode of delivery, malformations), current obstetric history (if pregnancy was planned, use of tobacco, alcohol, drug and medications and vaccinations received), presence of symptoms/signs consistent with ZIKV infection (fever, rash, non-purulent conjunctivitis, arthritis/arthralgia, lymphadenopathy, myalgia, headache) at any point throughout pregnancy, the woman’s environment (type of housing, number of rooms, number of people per household), preventative measures (use of repellent, protective clothing, window or bed nets, barrier contraception) and their knowledge of ZIKV and its forms of transmission as well as what their sources of information were.

Table 4 The Jundiaí Zika cohort, summary of health, medical and laboratory data collected from women and their children

<table>
<thead>
<tr>
<th>Phase</th>
<th>Measurements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pregnant women at enrolment</td>
<td>Baseline questionnaire - sociodemographic details, past medical history, family history, past obstetric history (parity, miscarriages, mode of delivery, malformations), current obstetric history (if pregnancy was planned, use of tobacco, alcohol, drug and medications and vaccinations received), presence of symptoms/signs consistent with ZIKV infection (fever, rash, non-purulent conjunctivitis, arthritis/arthralgia, lymphadenopathy, myalgia, headache) at any point throughout pregnancy, the woman’s environment (type of housing, number of rooms, number of people per household), preventative measures (use of repellent, protective clothing, window or bed nets, barrier contraception) and their knowledge of ZIKV and its forms of transmission as well as what their sources of information were.</td>
</tr>
<tr>
<td>Pregnant women follow-up 14–21 days after enrolment</td>
<td>Sample collection (blood, saliva, urine) for ZIKV RT-PCR and IgG/IgM. Symptoms questionnaire - presence of symptoms/signs consistent with ZIKV infection (timing, duration, intensity, action taken)</td>
</tr>
<tr>
<td>Pregnant women subsequent 2–3 monthly follow-ups</td>
<td>Sample collection (blood, saliva, urine) for ZIKV RT-PCR and IgG/IgM. Symptoms questionnaire - presence of symptoms/signs consistent with ZIKV infection (timing, duration, intensity, action taken)</td>
</tr>
<tr>
<td>Pregnant women weekly phone follow-up</td>
<td>Symptoms questionnaire - presence of symptoms/signs consistent with ZIKV infection (timing, duration, intensity, action taken)</td>
</tr>
<tr>
<td>Birth (mother and neonate)</td>
<td>Sample collection (blood, saliva, urine) for ZIKV RT-PCR and IgG/IgM. Colostrum (mother) and cerebrospinal fluid (neonate exposed to ZIKV and/or with microcephaly) for ZIKV PCR. Anthropometry – weight, length, head circumference (neonate) Placenta and umbilical collection - pathology</td>
</tr>
<tr>
<td>Neonatal, infant and child follow-ups</td>
<td>Sample collection (blood, saliva, urine) for ZIKV RT-PCR and IgG/IgM in months 1, 3, 6 and 15 for neonates. (Women found to be ZIKV RT-PCR positive during pregnancy also had blood, saliva and urine collected during paediatric follow-up appointments). Paediatric follow-up questionnaire – problems, significant events, feeding, vaccinations, developmental milestones reached, review of lab test results (including heel-prick test) Anthropometry – weight, length, head circumference Paediatric physical examination – general, cardiovascular, respiratory, gastrointestinal, neurological, developmental Physiotherapy assessment Speech and language assessment</td>
</tr>
<tr>
<td>ZIKV exposed infants and/or with microcephaly/other neurological abnormalities</td>
<td>Ophthalmology assessment (with Teller-CAT Cambridge Colour Test, funduscopy and extrinsic ocular motility tests) Specialist neurodevelopmental assessment (using Bayley-III developmental scales) Specialist audiology assessment Gastrointestinal assessment Imaging – Cranial ultrasound and CT brain at birth and 12 months</td>
</tr>
</tbody>
</table>

PCR, polymerase chain reaction; RT-PCR, reverse transcription polymerase chain reaction; ZIKV, Zika virus.
tubes which were mixed and labelled appropriately and placed in the fridge.

**ELISA for identification of anti-ZIKV immunoglobulins (IgM and IgG)**

Detection of anti-ZIKV antibodies (IgM and IgG) was performed by ELISA using commercial Zika IgG and Zika IgM ELISA kits Euroimmun (Euroimmun BR 2015), approved by the National Sanitary Surveillance Agency (ANVISA) for the diagnosis of ZIKV.

Serum, blood or plasma were diluted 1:101 in sample buffer. For IgM detection, the buffer contains an IgG/RF absorbent (preparation of anti-goat IgG). After, 100µL of this dilution was applied to each well of the plates. The plates were covered with protective film and incubated for 60 min at +37°C. After the first incubation, the plates were washed with 400µL of wash buffer (1X). One hundred microliters of the conjugate (peroxidase-labelled human IgG and IgM) were applied to each well, followed by incubation for 30 min at room temperature. Another wash was performed, under the same conditions described previously. In each well 100µL of the substrate/chromogen was added, followed by incubation for 30 min at room temperature, protected from light. After washing, 100µL of Stop solution was added and the plates were read in Automatic Biochemical Analyzer, model preiest TOUCH (2009 ROBONIK India), in 450 nm absorbance.

For the detection of IgG, three calibraters were used plus the negative and positive controls contained in the kit. For IgM detection a calibrator plus the positive and negative controls were used. The cut-off was calculated by the ratio between the absorbance of the controls and that of the calibraters. Samples with a cut-off <0.8 and positive samples with a cut-off of ≥1.1 were considered negative. The samples with a cut-off between ≥0.8 and 1.1 were considered equivocal.

**Real time PCR for ZIKV detection**

ZIKV RNA detection was performed by real time PCR (RT-qPCR), as recommended by the WHO, according to the protocol developed by Lanciotti et al.\(^6\) on maternal and neonatal urine samples.

Initially, the total RNA was extracted from 140µL of urine using the QIAamp Viral RNA Mini Kit (QIAGEN), following manufacturer's instructions. The final RNA was eluted in 60µl of ultrapure H2O Nuclease-Free Water (2018 Merck KGaA, Darmstadt, Germany) and RT-qPCR was performed on the same day. The remaining RNA was stored in a freezer at ~80°C. RT-qPCR was performed with GoTaq Probe qPCR and RT-qPCR Systems (2018 Promega Corporation Brasil, Ltda). For the final volume of 20µl reaction, 8µl of RNA template was used. The Mix was created with 10µl of GoTaq Probe qPCR Master Mix with dUTP (1x), 0,4µl of GoScript RT Mix for 1-Step RT-qPCR (1x), 1µl of Forward primer (10pmol/µL), 1µl of Reverse primer (10pmol/µL), 1µl of probe (10pmol/µL) (table 5) and of Nuclease-Free Water to complete the final volume. Two sets of primers and probe were used on the RT-qPCR reaction. The reaction occurred in ABI Prism 7500 SDS Real-Time cycler (Applied Biosystems), where the amplification cycles consisted of: One cycle of 15 min at 45°C for reverse transcription, one cycle of 2 min at 95°C for reverse transcriptase inactivation and for polymerase activation, 40 cycles of 15s at 95°C for denaturation and 1 min at 60°C for annealing and extension. The primers and probes used for this quantification are complementary to the gene encoding the NS1 protein of ZIKV (table 5). The probe contains a fluorescent 6-carboxyfluorescein (FAM) reporter dye at the 5’ end and the fluorescent dye 6-carboxyethylmethyldihydramine (TAMRA) at the 3’ end. All reactions followed positive and negative controls previously quantified.

The placenta and umbilical cord were collected and stored in formaldehyde and sent to the pathology laboratory where they were examined by specialist placental pathologists.

**Patient and public involvement**

As can be seen in the timeline of the creation of the JZC (figure 2), the wider community were invited to participate in the design of the JZC study and to join the research team to be trained as volunteers. Some of these members of the public were then involved with recruitment to the study and responsible for the dissemination of information about the study to their local communities. As the JZC involves a long follow-up period of pregnant women and their children, personal clinical results have been

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**Table 5** Primer and probe sets used for RT-qPCR ZIKV detection

<table>
<thead>
<tr>
<th>Primer/probe set</th>
<th>Primer and/or probe</th>
<th>Genome position</th>
<th>Sequence (5’ – 3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set 1</td>
<td>ZIKV 835</td>
<td>835–857</td>
<td>TTGGTGTACTGATCTGCTGATTGC</td>
</tr>
<tr>
<td></td>
<td>ZIKV 860-FAM</td>
<td>860–886</td>
<td>CGGCATACAGCATCGATCGAGGAG</td>
</tr>
<tr>
<td></td>
<td>ZIKV 911 c</td>
<td>911–890</td>
<td>CCTCCCAAAATGCTCATTGCC</td>
</tr>
<tr>
<td></td>
<td>ZIKV 1086</td>
<td>1086–1102</td>
<td>CGCAGCCCGCAACACAGAG</td>
</tr>
<tr>
<td></td>
<td>ZIKV 1107-FAM</td>
<td>1162–1139</td>
<td>AGCCTACCTTGGACAGCAGTCAGCACACTCA</td>
</tr>
<tr>
<td></td>
<td>ZIKV 1162 c</td>
<td>1107–1137</td>
<td>CCACTAACGTTTTGCAGACAT</td>
</tr>
</tbody>
</table>

Primers designed by Lanciotti et al, 2008 (Based on ZIKV MR 766 GenBank accession no. AY632535).

FAM, fluorescent 6-carboxyfluorescein; PCR, polymerase chain reaction; RT-qPCR, real time quantitative PCR; ZIKV, Zika virus.
regularly communicated to patients and their families. In addition, any relevant study findings have also been carefully and appropriately explained.

Findings to date

ZIKV clinical features during pregnancy
The clinical features of ZIKV infection among pregnant women in the JZC have been described. They have been used to assess the sensitivity of the current standard clinical case definitions, and to investigate whether adverse foetal outcomes are more likely to occur among pregnant women with symptomatic ZIKV infection during pregnancy compared with asymptomatic infection.

Foetal outcomes after congenital ZIKV exposure
This includes studies comparing the incidence of negative foetal outcomes, namely low birth weight, small-for-gestational age, prematurity and foetal death among ZIKV-exposed and unexposed women. Studies have also looked at the prevalence of Chikungunya IgG among mothers who had foetal losses.

Congenital Zika syndrome
The spectrum of congenital Zika virus syndrome in the JZC children has been explored as well as visual acuity alterations among ZIKV exposed babies, including among dizygotic twins.

The placenta
Placental histological findings among ZIKV exposed infants with microcephaly have been reported as well as placental histological findings among mothers with Chikungunya and Dengue infection.

Environmental risk factors, prevention, educational and vector control activities
These studies have included investigations into the environmental risk factors for ZIKV infection, assessment of the peri-domicile environment of women in the JZC and identification of favourable conditions for replication evaluation of educational activities for children to help combat the proliferation of Aedes aegypti and assessment of the knowledge around the modes of transmission and prevention of ZIKV infection as well as the practice of preventative measures among pregnant women in the cohort.

Susceptibility of ZIKV infection of neural progenitor cells among dizygotic twins
Analysis of neural progenitor cells (NPCs) of dizygotic twins discordant for CZS have shown that the development of CZS depends on the intrinsic susceptibility of the NPCs.17

Seroepidemiological arbovirus studies
Studies quantifying the seroprevalence of ZIKV, Chikungunya and Dengue IgG antibodies among pregnant women in the JZC have been carried out.

Of note, unless referenced, these manuscripts are awaiting final publication.

Strengths and limitations
The JZC is one of the few prospective Zika cohort studies that has recruited both asymptomatic and symptomatic women and therefore benefits from having a large control group. It also provides the necessary study population to carry out analyses on ZIKV symptomatology. Women were recruited over more than a 1 year period of time (March 2016 to August 2017) and therefore seasonality can be explored. The diversity and frequency of biological samples collected from the women during pregnancy (and after), as well as their children, mean that JZC now has a rich and invaluable biorepository of clinical material. The high-risk profile of the pregnant women also provides an additional unique opportunity to study other factors that may contribute to, or potentially be protective for, the development of negative sequelae associated with ZIKV exposure. The JZC implemented the use of standardised WHO research method tools, as soon as they were available, and therefore has placed itself in an optimal position to collaborate in Brazilian and international consortia that will ultimately be aiming to perform meta-analyses on all Zika cohort study data.

The limitations of the JZC in part relate to the pressing nature of the ZIKV and microcephaly epidemic and the urgency to start the investigation. The JZC, like many other Zika studies, commenced without any formal funding. Recruitment and data collection commenced in paper form, before formal data management systems could be put in place. In addition, as WHO standardised research protocols were produced after the start of the investigation, some variables contained in the WHO protocol were not in our original questionnaire and therefore some of this data is missing for the earliest recruits in our cohort. As this cohort was built in the midst of the ZIKV epidemic with limited financial resources, the scientific leadership team opted to prioritise the testing of urine samples by RT-PCR due to its wider window of detection (eg, up to 2 weeks in urine vs 1 week in serum).16 18 Although important tests (eg, IgG and IgM assays and plaque reduction neutralization test (PRNT)) and the infrastructure required to perform them have continued to be too costly for testing using available resources, the relevant serum samples have been stored in a secure biorepository for future evaluation on procurement of additional funding. An additional limitation related to testing is the possibility of misclassification among a minority of the ZIKV-positive newborns who may have been infected postnatally during the first 10 days of life. The choice of study population (high-risk pregnant women) had several advantages as stated above. However, there are also a few drawbacks that should be highlighted. First, because women were not recruited based on a suspicion of having been exposed to ZIKV, the prevalence of ZIKV RT-PCR positivity was relatively low, and this equated to small numbers for estimating absolute and relative risks. Furthermore,
because high-risk pregnant women are at higher risk of developing some of the outcomes of interest, this may have reduced the power for us to detect differences in frequency of outcomes between Zika-exposed and unexposed dyads. The differential follow-up of women who were symptomatic for ZIKV infection in terms of a more intensive antenatal scanning schedule prioritised women with clinical ZIKV infection (consistent with the WHO definition)\textsuperscript{11} to try to improve our understanding of the sequence of events that occur in utero after ZIKV infection. As we were unable to offer monthly ultrasounds to all the women in the cohort, this differential follow-up may have also had consequences in terms of the detection of problems in the antenatal period. With regards to the neonatal outcomes, the loss to follow-up that we inevitably experienced may also reduce the power for us to detect differences in the frequency of outcomes between Zika-exposed and unexposed neonates.

**COLLABORATION**
The JZC headquarters is at the Jundiaí Medical School under the direction of Professor Saulo Duarte Passos. The cohort has a Facebook page: https://www.facebook.com/zikacoorjedi/ which contains information about the functioning and organisation of the JZC, fundraising events, support for Zika affected families and media coverage. Any researcher wanting to use JZC data must apply to the Jundiaí Zika Cohort Group via Professor Passos (sauloduarte@uol.com.br).

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