



BMJ Open is committed to open peer review. As part of this commitment we make the peer review history of every article we publish publicly available.

When an article is published we post the peer reviewers' comments and the authors' responses online. We also post the versions of the paper that were used during peer review. These are the versions that the peer review comments apply to.

The versions of the paper that follow are the versions that were submitted during the peer review process. They are not the versions of record or the final published versions. They should not be cited or distributed as the published version of this manuscript.

BMJ Open is an open access journal and the full, final, typeset and author-corrected version of record of the manuscript is available on our site with no access controls, subscription charges or pay-per-view fees (<http://bmjopen.bmj.com>).

If you have any questions on BMJ Open's open peer review process please email [info.bmjopen@bmj.com](mailto:info.bmjopen@bmj.com)

## Seroprevalence and Demographic Characteristics of SARS-CoV-2– Infected Residents of Kibera Informal Settlement, Nairobi, Kenya, during the COVID-19 Pandemic

|                               |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
|-------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Journal:                      | <i>BMJ Open</i>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| Manuscript ID                 | bmjopen-2024-094546                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| Article Type:                 | Original research                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
| Date Submitted by the Author: | 02-Oct-2024                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| Complete List of Authors:     | <p>Carter, Jane; Amref Health Africa, Regional Laboratory Programme Khamadi, Samoel; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Mwangi, Joseph; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Muhula, Samuel; Amref Health Africa, Learning and Impact</p> <p>Munene, Stephen; Amref Health Africa, Regional Laboratory Programme</p> <p>Kanyara, Lucy; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Kinyua, Joyceline; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Lagat, Nancy; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Chege, Judy; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Oira, Robert; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Maiyo, Alex; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Stewart, Roy; University Medical Centre Groningen, Department of Health Sciences, Community &amp; Occupational Medicine</p> <p>Postma, Maarten; UMCG, Department of Health Sciences; UMCG, Department of Economics, Econometrics and Finance</p> <p>Stekelenburg, Jelle; Medisch Centrum Leeuwarden, Department of Obstetrics and Gynaecology; Universitair Medisch Centrum Groningen, Department of Health Sciences, Global Health</p> <p>Osur, Joachim; Amref International University</p> <p>Hulst, Marinus; University Medical Centre Groningen, Department of Health Sciences, Community &amp; Occupational Medicine; Martini Hospital, Department of Clinical Pharmacy and Toxicology</p> |
| Keywords:                     | COVID-19, EPIDEMIOLOGY, PUBLIC HEALTH, Community-Based Participatory Research                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
|                               |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |



I, the Submitting Author has the right to grant and does grant on behalf of all authors of the Work (as defined in the below author licence), an exclusive licence and/or a non-exclusive licence for contributions from authors who are: i) UK Crown employees; ii) where BMJ has agreed a CC-BY licence shall apply, and/or iii) in accordance with the terms applicable for US Federal Government officers or employees acting as part of their official duties; on a worldwide, perpetual, irrevocable, royalty-free basis to BMJ Publishing Group Ltd ("BMJ") its licensees and where the relevant Journal is co-owned by BMJ to the co-owners of the Journal, to publish the Work in this journal and any other BMJ products and to exploit all rights, as set out in our [licence](#).

The Submitting Author accepts and understands that any supply made under these terms is made by BMJ to the Submitting Author unless you are acting as an employee on behalf of your employer or a postgraduate student of an affiliated institution which is paying any applicable article publishing charge ("APC") for Open Access articles. Where the Submitting Author wishes to make the Work available on an Open Access basis (and intends to pay the relevant APC), the terms of reuse of such Open Access shall be governed by a Creative Commons licence – details of these licences and which [Creative Commons](#) licence will apply to this Work are set out in our licence referred to above.

Other than as permitted in any relevant BMJ Author's Self Archiving Policies, I confirm this Work has not been accepted for publication elsewhere, is not being considered for publication elsewhere and does not duplicate material already published. I confirm all authors consent to publication of this Work and authorise the granting of this licence.

Protected by copyright, including for uses related to text and data mining, AI training, and similar technologies. Enseignement Supérieur (ABES).

# Seroprevalence and Demographic Characteristics of SARS-CoV-2–Infected Residents of Kibera Informal Settlement, Nairobi, Kenya, during the COVID-19 Pandemic

Jane Y Carter,<sup>1</sup> Samoel Khamadi,<sup>2</sup> Joseph Mwangi,<sup>2</sup> Samuel Muhula,<sup>1</sup> Stephen M Munene,<sup>1</sup> Lucy Kanyara,<sup>2</sup> Joyceline Kinyua,<sup>2</sup> Nancy Lagat,<sup>2</sup> Judy Chege,<sup>2</sup> Robert Oira,<sup>2</sup> Alex Maiyo,<sup>2</sup> Roy Stewart,<sup>3</sup> Maarten Postma,<sup>3,4</sup> Jelle Stekelenburg,<sup>3,5</sup> Joachim Osur,<sup>6</sup> Marinus van Hulst<sup>3,7</sup>

<sup>1</sup>Amref Health Africa Headquarters, PO Box 27691–00506, Nairobi, Kenya

<sup>2</sup>Centre for Virus Research, Kenya Medical Research Institute, PO Box 54840–00200, Nairobi, Kenya

<sup>3</sup>Department of Health Sciences, University of Groningen, University Medical Center Groningen, Groningen, Netherlands

<sup>4</sup>Department of Economics, Econometrics and Finance, University of Groningen, Faculty of Economics and Business, Groningen, Netherlands

<sup>5</sup>Department of Obstetrics and Gynaecology, Medical Centre Leeuwarden, Leeuwarden, Netherlands

<sup>6</sup>Amref International University, PO Box 27691–00506, Nairobi, Kenya

<sup>7</sup>Department of Clinical Pharmacy and Toxicology, Martini Hospital, Groningen, Netherlands

Jane Y Carter  <https://orcid.org/0000-0003-4671-3519>

Samoel Khamadi  <https://orcid.org/0000-0003-3050-5977>

Joseph Mwangi  <https://orcid.org/0000-0002-4827-185X>

Stephen Munene  <https://orcid.org/0000-0002-5834-2914>

Samuel Muhula  <https://orcid.org/0000-0001-7992-6532>

Lucy Kanyara  <https://orcid.org/0000-0002-9126-388X>

Joyceline Kinyua  <https://orcid.org/0000-0002-5659-5424>

Nancy Lagat  <https://orcid.org/0000-0002-4834-0632>

Judy Chege  <https://orcid.org/0000-0002-7929-2557>

Robert Oira  <https://orcid.org/0000-0002-2399-9801>

1 Roy Stewart  <https://orcid.org/0000-0001-9227-433X>

2 Maarten Postma  <https://orcid.org/0000-0002-6306-3653>

3 Jelle Stekelenburg  <https://orcid.org/0000-0002-2732-6620>

4 Joachim Osur  <https://orcid.org/0000-0002-6277-8753>

5 Marinus van Hulst  <https://orcid.org/0000-0003-3216-7246>

15 Correspondence should be addressed to Dr Jane Y Carter

16 Amref Health Africa Headquarters, PO Box 27691–00506, Nairobi, Kenya; email: [jane.carter@amref.org](mailto:jane.carter@amref.org)

## ABSTRACT

### Introduction

Kenya recorded relatively low numbers of COVID-19 cases during the recent pandemic. The reasons remain unclear especially in informal settlements where implementing containment measures is impractical. A study was conducted in Kibera informal settlement in Nairobi, Kenya's largest slum, to determine the seroprevalence of SARS-CoV-2 and associated demographic characteristics, before vaccination became widespread.

### Methods

Demographic data were collected from participants age  $\geq 1$  year who reported no current symptoms of COVID-19. Capillary blood (500  $\mu$ L) was collected into Microtainer® EDTA tubes and transported to the Kenya Medical Research Institute laboratories for SARS-CoV-2 antibody testing using the Standard Q COVID-19 IgM/IgG Combo rapid test and two enzyme-linked immunosorbent assays: Wantai Total Ab (IgM/IgG/IgA) and Platelia SARS-CoV-2 Total Ab (IgM/IgG/IgA).

### Results

A total of 438 participants were recruited into the study. Most (79.2%) were age 18–50 years; females (64.2%) exceeded males. More than one third (39.1%) were unemployed; only 7.4% were in formal, full-time employment. Less than one quarter (22.1%) self-reported any underlying health conditions. Nearly two-thirds (64.2%) reported symptoms compatible with COVID-19 in the previous 16 months; only one (0.23%) had been hospitalised with a reported negative COVID-19 test. 370 (84.5%) participants tested positive in any of the three tests. There was no significant difference in SARS-CoV-2 seropositivity across participants' age, sex, presence of underlying health conditions, on medication, or those ever tested for SARS-CoV-2. Multiple logistic regression analysis showed COVID-19 symptoms in the previous 16 months were the only significant independent predictor of seropositivity ( $p=0.0085$ ).

### Conclusion

This study reports high SARS-CoV-2 exposure in Kibera informal settlement with limited morbidity; only recent COVID-19 symptoms were a significant independent predictor of seropositivity. The study confirms other reports of high SARS-CoV-2 exposure with limited morbidity in slum communities. Small blood volumes are convenient for serosurveillance studies in large populations.

1     **Key questions**

2     ***What is already known on this topic***

3  
4     The COVID-19 pandemic had a limited impact on the African continent, with rates of morbidity and  
5     mortality much lower than reported elsewhere. The underlying reasons remain unclear although the high  
6     infectious disease burden and demographic age structure have been suggested. High exposure rates to  
7     SARS-CoV-2 infection in slum communities have also been reported from other resource-limited  
8     countries with less impact than expected, despite warnings of potentially major consequences.

9  
10  
11  
12     ***What this study adds***

13  
14     To our knowledge, this is the first serosurveillance study in an informal settlement community in Africa,  
15     filling an important gap in existing research. By focusing on this specific setting, the study provides  
16     valuable insights that can help collaborate and contextualise findings from previous research conducted  
17     in different geographic and socioeconomic environments, including Africa.

18  
19  
20  
21     ***How this study might affect research, practice, or policy***

22  
23     This study reports data that could inform future research on factors contributing to the unexpected low  
24     COVID-19 morbidity observed in informal settlement communities, despite high exposure rates and the  
25     inability to comply with social distancing measures. This study's findings also challenge the assumption  
26     that high exposure rates necessarily lead to high morbidity, suggesting the need to further investigate  
27     potential protective factors and immune responses. The insights gained from this study could influence  
28     policy decisions regarding the allocation of resources and development of targeted interventions for  
29     informal settlement communities during public health emergencies. The outcomes may also challenge the  
30     one-size-fits-all approach to COVID-19 mitigation measures and encourage policymakers to consider  
31     context-specific strategies that account for the unique characteristics and resilience of informal settlement  
32     communities. The successful use of capillary blood samples for seroprevalence studies could streamline  
33     and expand the reach of future surveillance efforts, making it easier to monitor the spread of COVID-19  
34     and other infectious diseases in hard-to-reach populations.



## INTRODUCTION

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the cause of coronavirus disease 2019 (COVID-19), was first reported in Wuhan, Hubei Province, China, in November 2019 and rapidly spread across China and the world [1]. On 30 January 2020, the World Health Organization (WHO) declared the outbreak a Public Health Emergency of International Concern, and on 11 March 2020 the WHO Director General declared COVID-19 a pandemic [2]. Although the WHO declared the COVID-19 global health emergency over on 5 May 2023 [3], globally over 775 million cases and more than 7 million deaths from COVID-19 had been reported by 16 June 2024, with more than 127,000 new cases and nearly 2,000 deaths in the previous 28 days [4]. In the African region, more than 9 million confirmed cases and over 175,000 deaths had been reported cumulatively by June 2024, although the numbers of cases and deaths reported in Africa were likely an underestimate due to low testing rates [5]; an assessment by the WHO Regional Office for Africa showed only one in seven (14.2%) COVID-19 infections were being detected [6]. Studies from Kenya and other countries in Africa noted low mortality from COVID-19, despite high exposure rates [7,8].

The modes of transmission of SARS-CoV-2 have been elucidated through detailed case contact studies. Respiratory transmission, with SARS-CoV-2 carried on tiny particles emitted from the respiratory tract, has been established as the clear and dominant route of spread; indirect transmission appears to be of limited importance despite initial concerns [Error! Reference source not found.]. The clinical spectrum of COVID-19 varies from asymptomatic or pre-symptomatic infection, mild to moderate illness, to severe and critical illness characterised by respiratory failure and multiple organ dysfunction; varying proportions of infected persons remain asymptomatic [10,11]. Transmission from asymptomatic individuals is estimated to account for more than half of all transmissions [12]. Diagnostic testing using reverse transcriptase polymerase chain reaction (RT-PCR)-based assays performed on respiratory specimens is the reference standard for establishing a microbiological diagnosis of COVID-19 [13] but is constrained by the presence of virus to a few days before infection and a short time after infection. Seroepidemiological studies to detect host antibodies to SARS-CoV-2 are therefore important for estimating disease burden and providing a more complete picture of exposure to SARS-CoV-2 in a population [14].

### *COVID-19 infection in Kenya*

The first confirmed case of COVID-19 was reported in Kenya on 13 March 2020 [15]. Restrictions to mitigate the spread of COVID-19 were instituted in March 2020 and were eased towards the end of 2020 until a second lockdown was instituted in five counties in March 2021; in May 2021, as the number



of cases dropped, lockdown restrictions were again lifted [16]. COVID-19 vaccination was initiated in Kenya in March 2021, with vaccination numbers rising from August 2021 with increased vaccine availability. Kenya actively promoted vaccination through the COVID-19 Vaccination Acceleration Programme [17]; as of 16 May 2023, more than 18 million vaccine doses had been administered with 30.7% of the adult population ( $\geq 18$  years) fully vaccinated. Kenya planned to vaccinate all adults and teenagers in 2022 and to provide third dose booster shots to all eligible adults [18]. On 11 March 2022, Kenya lifted all COVID-19 restrictions while urging continued personal public health measures; by June 2024, 344,000 cases and 5,700 deaths had been reported since the pandemic started, rates that remained low in comparison with global figures [4].

Available seroprevalence data from Kenya targeting different population groups painted a concerning picture of the COVID-19 pandemic’s progression within the country. Studies examining stored samples from blood donors across six regional blood transfusion centres, including the capital city Nairobi, revealed a dramatic rise in SARS-CoV-2 antibody prevalence over time. Seroprevalence increased from 4.3% (95% CI, 2.9–5.8%) in samples collected between April and June 2020 to 48.5% (95% CI, 45.2–52.1%) in samples taken just a year later from January to March 2021 [19,20]. These samples were tested using a non-commercial validated enzyme-linked immunosorbent assay (ELISA) for SARS-CoV-2 IgG against spike protein. The same assay was used to test stored blood from women attending antenatal care services at three hospitals in Kenya between August 2020 and October 2021: seroprevalence rose from 50% (95% CI, 42–58%) in August 2020 to 85% (95% CI, 78–92%) in October 2021 in Nairobi; 31% (95% CI, 25–37%) in May 2021 to 71% (95% CI, 64–77%) in October 2021 in Busia; and from 1% (95% CI, 0–3%) in September 2020 to 63% (95% CI, 56–69%) in October 2021 in Kilifi [21]. Purposive testing of venous blood from healthcare workers between July and December 2020 using the same ELISA showed significant variation in overall seropositivity (20.8%: 17.5%–24.4%). Seroprevalence varied significantly ( $p < 0.001$ ) by site: 43.8% (35.8–52.2%) in Nairobi, 12.6% (8.8–17.1%) in Busia and 11.5% (7.2–17.6%) in Kilifi; only 16 (2%) of the sampled healthcare workers reported acute respiratory symptoms at the time of sample collection [22]. Purposive testing of truck drivers and their assistants, again using the same ELISA, conducted between September and October 2020 showed an overall seropositivity of 7.4%; none reported current or previous symptoms of illness [23]. Truck drivers and their assistants continued to transport essential supplies during the COVID-19 pandemic, placing them at increased risk of being infected and of transmitting SARS-CoV-2 over a wide geographical area. In a study on samples collected from January to March 2020 from rural populations in western Kenya with and without human immunodeficiency virus (HIV) infection, 3.3% had detectable SARS-CoV-2 antibodies, with no difference between participants with and without HIV infection (3.1% vs. 4.0%,  $p = 0.68$ ); this study used

the Platelia SARS-CoV-2 Total Ab assay (Bio-Rad Laboratories, Hercules, California, USA). Participants denied symptoms except for one who reported a cough in the preceding week [24]. In our study, SARS-CoV-2 seroprevalence in an informal settlement in Nairobi was conducted to supplement these data and determine exposure in a community subjected to overcrowding and lack of reliable clean water, housing, health services and waste management facilities, where preventive measures were difficult to apply.

### **Purpose of the study**

This study assessed the prevalence of SARS-CoV-2 antibodies in the residents of Kibera informal settlement in Nairobi, Kenya, one of sub-Saharan Africa's largest slums, before vaccination became widespread, and explored some of the major risk factors for infection. This study also provided an opportunity to determine pathogen exposure in large numbers of persons in a pandemic situation using small blood volumes obtained from fingerprick samples.

## **MATERIALS AND METHODS**

### **Study site**

Kibera informal settlement stands on a 2.5-kilometre square area of land located 5 kilometres southwest of Nairobi's Central Business District. The population is estimated at 170,070, although this may be an underestimate [25]. Residents comprise the major ethnic communities in Kenya [26] with reportedly 116 women for every 100 men [27]. Kibera is governed by area chiefs as the local administrative arm of the government.

### **Study design**

A cross-sectional study was performed to determine SARS-CoV-2 antibody prevalence in Kibera informal settlement, Nairobi, Kenya. The study was conducted between 2 and 13 August 2021 in 10 of the 14 Kibera villages, which provided an adequate representation of the whole settlement. The Cochran sample size formula for larger populations was used to determine sample size [28]. Since the population of Kibera is above 170,000 and approximately 500 people had tested positive for SARS-CoV-2 at the time of the study, a sample size of 389 (~400) participants was determined, a target of 40 participants in each selected village (confidence level 99%; precision 0.01). In each village every fifth household was selected; if a household declined participation, the next household was included. Entry to individual households was facilitated through community health volunteers who were formally assigned to these villages. Participants age 1 year and above who had been resident in Kibera since at least October 2019 were included.

### **Study methods**

Five data collectors, each assigned to two villages, administered a brief questionnaire to each consenting participant, or their parent or legal guardian; data were collected directly into the Open Data Kit system installed in handheld tablets. Data were collected on the following: age, sex, length of residence in Kibera, pregnancy status (for women), main occupation and nature of employment, any underlying health conditions and type of condition, any medications and name of the medication, symptoms suggestive of COVID-19 in the previous 16 months (cough, fever, difficulty in breathing, shortness of breath, new loss of taste and smell), any hospitalisation and date (if known), ever tested for SARS-CoV-2, date and result (if known), and COVID-19 vaccination(s) and date (if known). Five laboratory staff accompanying the data collectors collected 500 µL of capillary blood from each participant by fingerprick into a Microtainer® EDTA tube. Blood samples were transported to the Kenya Medical Research Institute (KEMRI) laboratories for same-day testing on whole blood using the Standard Q COVID-19 IgM/IgG Combo rapid test (SD Biosensor Inc, Suwon, Gyeonggi, Republic of Korea). At the KEMRI laboratories, blood was separated the same day and plasma samples stored at −80 °C for further testing with two enzyme-linked immunosorbent assays (ELISAs): Wantai Total Ab (IgM/IgG/IgA) ELISA for SARS-CoV-2 (Wantai Biological Pharmacy Enterprise Co. Ltd, Beijing, China), and Platelia SARS-CoV-2 Total Ab (IgM/IgG/IgA) Assay (Bio-Rad Laboratories Inc, California, USA). The SARS-CoV-2 antibody tests selected in this survey were commercial tests approved for use in Kenya at the time of the study. Test performance is described in a separate paper [Error! Reference source not found.]. Individuals with positive test results in any of the three tests were considered positive (for the ELISAs, optical density ≥1.0) and indeterminate ELISA results as negative (optical density 0.9–0.99). PCR or antigen testing was not performed in any of the study participants. Each participant was assigned a unique identification number; data files and blood sample labelling did not include participant names to ensure anonymity. Individuals who met the case definition of COVID-19 infection [30] at the time of recruitment were not included in the study and were referred to a nearby health facility for further investigation and management. The research teams followed government regulations for the prevention of COVID-19 transmission during data collection [31].

### Statistical analysis

Data were entered into Excel, then cleaned and analysed using SAS version 9.4 (SAS Institute Inc). Frequency and percentages were used to present the data; Pearson’s chi-square test was used to examine the association between seropositive results and various demographic and health-related factors, including age, sex, employment status, presence and types of underlying health conditions, on any type of medication, COVID-19 vaccination, previous symptoms compatible with COVID-19 and those ever tested for SARS-CoV-2. A simple logistic regression estimated the crude odds ratio (COR) for each variable

found to be significantly associated with seropositivity in the chi-square test ( $\leq 0.20$ ). Multiple logistic regression was used to calculate the adjusted odds ratio (AOR), controlling for the effects of multiple variables simultaneously. The COR and AOR were calculated to quantify the strength of the associations between the variables and seropositive status.

## RESULTS

### Demographic data

A total of 438 participants consented to answer the questions and have a blood sample collected by fingerprick; one participant refused to divulge her occupation. Table 1 shows the information collected from the participant questionnaire. Most participants (79.2%) were between age 18 and 50 years; only 4.8% were age  $\leq 17$  years and 16% were age  $\geq 50$  years. Females exceeded males across the 18–50 years age group by a ratio of 2:3, but males exceeded females in the  $< 18$  years and  $> 50$  years age groups. Eleven female participants were pregnant; two were uncertain of their pregnancy status. Of the 419 participants age  $\geq 16$  years (the official age for employment in Kenya [32]) more than one third (39.1%) were unemployed; by far the greatest number of unemployed participants were women in the 18–50 years age group (74.4% of unemployed participants). Less than one quarter (22.1%) of participants self-reported any type of underlying health condition; the most common was cardiovascular disease, including hypertension (9.1% of all participants); only 14 (3.2%) participants reported underlying respiratory disease (including asthma). Underlying health conditions were most common in females in the 18–50 years age group (43.8% of all participants). Less than one fifth (16%) of participants were taking one or more medications at the time of the survey; these included therapeutic drugs as well as vitamins and symptomatic treatments. Four of the 18 participants with HIV infection reported not taking antiretroviral medication. Only 1 (0.23%) of the 281 (64.2%) participants reporting symptoms suggestive of COVID-19 in the previous 16 months had been hospitalised; he reported a negative SARS-CoV-2 test at that time. Only 19 (63%) of the 30 participants who reported receiving vaccination against SARS-CoV-2 had received two doses. Most participants could not recall exact dates of vaccinations, but most received vaccinations between 1 April and 28 July 2021, within 4 months of the survey; one reported receiving vaccination in March 2021 and one in August 2021.

### Serological results in relation to demographic data

The collection of an adequate volume of blood by fingerprick required lancets of sufficient depth (at least 1.8 mm; we used 2 mm); smaller lancets used initially did not produce an adequate flow of capillary blood for collection of the full 500  $\mu$ L. Due to inadequate sample collection, there was

insufficient sample in 72 participants to conduct the Platelia SARS-CoV-2 Total Ab Assay, the last test to be conducted. Comparison of performance between the three diagnostic tests (Wantai ELISA, Platelia ELISA and the rapid diagnostic test) are described in a separate paper [Error! Reference source not found.]. In summary, the Wantai ELISA showed greater percentage positive results (82.6%) compared with the rapid test (51.8%) and the Platelia ELISA (69.7%); of the rapid test results, 23 were IgM positive, 151 were IgG positive, 53 were both IgM and IgG positive. Specificities of the rapid test and Platelia SARS-CoV-2 Total Ab Assay using the Wantai ELISA as the reference test were high (>90%). There was no significant difference in percentage positive results across participants' age, sex, presence of underlying health conditions, on any type of medication, or those ever tested for SARS-CoV-2; there was a significant difference in percentage positive results according to participants' employment status ( $p=0.0176$ ), COVID-19 symptoms in the previous 16 months ( $p=0.0099$ ), and vaccination status ( $p=0.0561$ ); this last variable was included in the multiple logistic regression analysis because of the  $p$ -value of  $\leq 0.20$ . The previously hospitalised participant tested negative with the rapid test but tested positive with the two ELISAs. The results of multiple logistic regression analysis showed that having had COVID-19 symptoms in the previous 16 months was the only significant independent predictor of seropositivity in this population ( $p=0.0085$ ) [Table 2].

DISCUSSION

This study confirmed the expected high exposure to SARS-CoV-2 (84.5%) in residents of Kibera slum at the mid-point of the COVID-19 pandemic. Although nearly two-thirds of participants (62.4%) reported COVID-like symptoms, only one had been hospitalised and recovered. There was no significant association of SARS-CoV-2 infection with age, sex, underlying health conditions, being on any type of medication, having been previously tested for SARS-CoV-2; significant associations were found with employment status, COVID-19 vaccination and previous symptoms compatible with COVID-19, but multiple regression analysis showed an association only with previous symptoms suggestive of COVID-19. The categories <18 years of age and not applicable for employment status (<16 years) have wide confidence intervals, indicating uncertainty about the true value of the odds ratio; this may also be contributed to by the low participant numbers in these categories. In addition, numbers of vaccinated participants were small. The high exposure in this population could be linked to the fact that a large proportion are casual labourers who walk up to 15 kilometres daily in search of work and mix with populations in other areas; other groupings were also at risk due to the high population density resulting in high likelihood of transmission. The overall seroprevalence in our study is similar to that of women who attended antenatal care services in Nairobi (84.5% vs 85%) at a similar time frame (August 2021 vs



October 2021) [21], which was conducted at the national referral hospital close to the Kibera slum. However, in our female population, seroprevalence was only 33.3% in the 18–30 years age group, and 34.9% in the 31–50 years age group, implying the population groups were different and suggesting the need for further investigation.

The reasons for low rates of severe disease in this infected population are unclear. Several explanations have been proposed for the overall low morbidity and mortality from COVID-19 in Africa, including the high infectious disease burden on the African continent and the demographic age structure [33,34,35,36,37]. According to the United Nations, the over 1 billion slum dwellers worldwide are mostly confined to three regions, including sub-Saharan Africa [38]. For residents of urban slums, the difficulty in implementing COVID-19 preventive measures due to severe overcrowding was recognised early [39,40] but the predicted high caseload and mortality rates from community transmission were not apparent, especially in Africa's slums [41,42,43].

This study supports seroprevalence studies in urban slums outside Africa that showed high exposure to the virus as well as low morbidity, with differing associated factors. Malani *et al* in a comparative study of slum and non-slum areas in Mumbai, India, showed markedly higher proportions of positive tests in slum areas (54.1%) than in non-slum areas (16.1%), with lower infection fatality in slums (0.076%) than in non-slums (0.263%) [44]. Nirala *et al* in a study across 10 different slums in Patna, India, found a seropositivity rate of 31.5% (95% CI: 27.9–35.1) with seropositive status significantly associated with age 18–30 years, male gender, high-risk occupations (autorickshaw drivers, rickshaw pullers, street vendors), below poverty line economic status, residing in a hut or kutcha house (makeshift dwelling) and COVID-like symptoms in the preceding one month [45]. A study by Raqib *et al* comparing slum and non-slum areas in Bangladesh showed seroprevalence was positively associated with limited years of formal education (AOR = 1.61; 95% CI = 1.43, 1.82), lower income (AOR = 1.23; 95% CI = 1.03, 1.46), overweight (AOR = 1.2835; 95% CI = 1.26, 1.97), diabetes (AOR = 1.67; 95% CI = 1.21, 2.32) and heart disease (AOR = 1.38; 95% CI = 1.03, 1.86) [46]. However, in a study by George *et al* in a large slum in South India with a reported overall COVID-19 seroprevalence of 57.9% (95% CI 53.4–62.3), age, education, occupation and presence of reported co-morbidities were not significantly associated with seroprevalence [47]. The discrepancies between high seroprevalence, low morbidity and varying underlying factors suggest the need for further research to understand the potential protective factors and immune responses that may be at play in these informal settlements. The informal community in our study may have distinct characteristics that sets it apart from other slum populations examined in previous research. Factors such as community resilience, social support networks and environmental conditions

1 may vary from place to place and could play a role in mitigating the impact of COVID-19 and influencing  
2 the observed lack of associations with reported risk factors.  
3

4 Co-morbidities associated with severe COVID-19 have also been examined in other populations.  
5 Martono Fatmawati *et al* reviewed nine studies from Asia and Europe that showed risk factors for severe  
6 COVID-19 included age, gender, chronic comorbidities, cardiovascular disease, diabetes, hypertension,  
7 kidney failure, cancer and a history of smoking [48]. Wang *et al* using self-reported data from a survey of  
8 adults in the United States reported that individuals with underlying health conditions were more likely to  
9 become infected with SARS-CoV-2 and have more severe post-infection symptoms [49]. Kompaniyets *et al*  
10 in a cross-sectional study of children age  $\leq 18$  years in the United States reported underlying health  
11 conditions, such as type 1 diabetes, cardiac and circulatory congenital anomalies and obesity, were more  
12 associated with severe COVID-19 or death [50]. Costa *et al* in a review of published data up to June 2020  
13 reported patients with metabolic disorders including obesity, diabetes, cardiovascular disease and liver  
14 disease faced a higher risk of SARS-CoV-2 infection and were associated with a significantly worse  
15 outcome [51]. Wu *et al* in a review of published papers up to April 2020 reported diabetes increased the  
16 mortality of patients with COVID-19 [52]. However, the findings from our study challenge the  
17 assumptions that COVID-19 risk factors are necessarily linked to severity of disease. Males have also  
18 been reported to be at higher risk of severe illness and increased mortality than females [53,12]; this was  
19 as well not evident in our study.  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31

32 Our study also explored the use of small blood volume samples collected through fingerprick. We  
33 found fingerprick sampling a convenient and acceptable way of conducting a serosurvey across a large  
34 population that included small children, and for evaluating new technologies. Testing kits used for  
35 serosurveys should therefore utilise whole blood or plasma separated from capillary blood. Our study used  
36 three tests concurrently to detect antibodies to SARS-CoV-2: one rapid test and two ELISA tests. Since  
37 the start of the COVID-19 pandemic, molecular testing for SARS-CoV-2 virus using RT-PCR-based  
38 assays performed on respiratory specimens has been central to disease management and control [14]. Tests  
39 for SARS-CoV-2 antigens on clinical specimens using immunoassays and rapid tests have also been used  
40 but are limited by sub-optimal sensitivity [53,55]. Molecular and antigen tests are constrained by detection  
41 of virus to a few days before infection and a short time after infection. In a systematic review of  
42 longitudinal studies of RT-PCR test results in symptomatic SARS-CoV-2 infections, the highest  
43 percentage of virus detection in nasopharyngeal swabs by PCR was 89% between 0 and 4 days post-  
44 symptom onset, dropping to 54% (95% CI 47 to 61) after 10–14 days [56]. Tests for detecting antibodies  
45 to SARS-CoV-2 applicable outside research laboratory settings include rapid tests using lateral flow  
46 immunology and ELISA tests. In a study by Guo *et al* using an ELISA based on recombinant viral  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60



nucleocapsid protein, the median time to IgM and IgA antibody detection after symptom onset was 5 (IQR 3–6) days, while IgG was detected at 14 (IQR 10–18) days [57]. Zhao *et al*, using an ELISA prepared from recombinant antigen containing the receptor binding domain of SARS-CoV-2 spike protein, reported median times from symptom onset to total antibody, IgM and IgG seroconversion were 11, 12 and 14 days, respectively [58]; the presence of antibodies was <40% among patients within 1 week of onset, and rapidly increased to 100% (total antibody – 94.3% (IgM), 79.8% (IgG)) by day 15 after onset; there were also negative antibody findings in 7% patients, possibly due to blood samples not taken at an appropriate time after symptom onset. In a report by Qu *et al* using a IgG/IgM chemiluminescent immunoassay with combined nucleocapsid protein and spike glycoprotein antigens, seroconversion time of IgG antibody was earlier than that of IgM antibody: 97.6% of patients (40/41) were positive with IgG and 87.8% (36/41) with IgM, with a median time of seroconversion for IgG at 11 days (8–16 days) and for IgM at 14 days (8–28 days) after disease onset. The level of IgG antibody reached the highest concentration on day 30, while the highest concentration of IgM antibody appeared on day 18, but then began to decline [59]. Hoffman *et al* found no statistical difference between testing samples taken from PCR-confirmed COVID-19 cases between 9 and 17 days and 18 and 29 days for either IgM or IgG seropositivity [60]. Generally, IgM is produced first with a later switch towards IgG production, but studies on SARS-associated coronaviruses suggest that IgM and IgG often develop at around the same time [61,62]. In general, detection of antibodies is better in samples taken >14 days after onset of disease, due to the time taken for antibody development [63,64]. Reported duration of antibody after natural confirmed infection is variable, including >6 months [65], up to 20 months in unvaccinated adults in the United States after confirmed COVID-19 infection [66], and more than 2 years in patients in Wuhan, China [67]. It is therefore probable that most natural infections occurring within the previous 16 months in our study population were detected, although the relevance of the IgM and IgG antibody findings is unclear.

Serologic studies that assess prior infection and immunity to infectious diseases are essential for epidemiologic studies, ongoing surveillance, vaccine studies and potentially for risk assessments of healthcare workers, and provide an indication of previous as well as recent exposure in a population. Surveillance therefore remains fundamental to understanding the evolution of SARS-CoV-2 infection, the risk factors for severe disease, and the impact of vaccination and public health and social measures [68]. Although hospital admissions and severe outcomes from COVID-19 have substantially decreased since the start of the pandemic, COVID-19-related deaths remain substantial: in the United States, COVID-19 still ranks as the 10th most common cause of death. The percentage of positive tests for SARS-CoV-2, a key indicator of community spread, reached peak levels of 12.9% in January 2024 [69]. Since disease mitigation measures implemented in many countries severely affected the global economy and financial

markets [70], and substantially impacted the inadequately prepared health systems in Africa [71], health systems need to be vigilant of emerging SARS-CoV-2 variants and the possibility of new surges in cases and deaths. As WHO recommends countries transition from emergency mode to managing COVID-19 alongside other infectious diseases, providing a more complete picture of the total number of people infected with SARS-CoV-2 remains an important measure for guiding public health responses [72]. The recent establishment of a global network to detect and monitor novel coronaviruses of public health importance will facilitate early detection, risk assessment, and response to coronavirus-related health challenges [73]. Well-designed surveillance systems remain core to monitoring acute viral respiratory infections to inform public health measures, health system capacities, impact of vaccination programmes and other control measures [74].

**Conclusions**

This study reports high SARS-CoV-2 exposure in a slum community in Nairobi, Kenya, with limited morbidity; only recent COVID-19 symptoms were a significant independent predictor of seropositivity. This study confirms other reports of high SARS-CoV-2 exposure with limited morbidity in slum communities. The study also supports the convenient use of small blood volumes for conducting population serosurveys. Use of serological assays to conduct seroprevalence studies will continue to remain important as more robust tests are developed. Public health systems would greatly benefit from serosurveillance to supplement and strengthen existing COVID-19 case-based infectious disease surveillance strategies [75].

**Study limitations**

This study was conducted during the working week during working hours, which may explain the low number of male participants of working age and low number of those formally employed. The tests used were limited by availability of approved tests at the time.

**Acknowledgements**

Approval for the study was obtained from the Amref Health Africa Ethics and Scientific Review Committee, from the National Commission for Science, Technology and Innovation, and from the Nairobi Metropolitan Services, Ministry of Health. Verbal permission to conduct the study was obtained from five area chiefs administering the 10 Kibera villages. The study was funded by American Tower Corporation.

**Author contributions**

Conceptualisation: JYC, SK, SM, JM, JO; Methodology: JYC, SK, SM, SMM, LK, JK, NL, JC, RO, AM; Analysis: JYC, RS, MH; writing—original draft preparation: JYC, RS, MH; writing—review and editing: JYC, SK, JM, SM, SMM, RS, MH, MP, JS, JO. All authors have read and agreed to the published version of the manuscript.

## Competing interests

All authors declare they have no competing interests.

## REFERENCES

1. Zhu H, Wei L, Niu P. The novel coronavirus outbreak in Wuhan, China. *Glob Health Res and Policy* 2020;**5**(6). Available <https://doi.org/10.1186/s41256-020-00135-6>
2. Cucinotta D, Vanelli M. WHO declares COVID-19 a pandemic *Acta Biomed* 2020;**91**(1)157–160. doi: <https://doi.org/10.23750/abm.v91i1.9397>
3. World Health Organization. WHO Director-General's opening remarks at the media briefing. 5 May 2023. Available <https://www.who.int/news-room/speeches/item/who-director-general-s-opening-remarks-at-the-media-briefing---5-may-2023>
4. World Health Organization. *Coronavirus disease (COVID-19) Epidemiological Updates and Monthly Operational Updates*. Available <https://www.who.int/emergencies/diseases/novel-coronavirus-2019/situation-reports>
5. Mwananyanda L, Gill CJ, MacLeod W, *et al.* Covid-19 deaths in Africa: prospective systematic postmortem surveillance study. *BMJ* 2021;**372**:n334. doi: <http://dx.doi.org/10.1136/bmj.n334>
6. World Health Organization Regional Office for Africa. Six in seven COVID-19 infections go undetected in Africa. October 2021. Available <https://www.afro.who.int/news/six-seven-covid-19-infections-go-undetected-africa>
7. Ngere I, Dawa J, Hunsperger E, *et al.* High seroprevalence of SARS-CoV-2 but low infection fatality ratio eight months after introduction in Nairobi, Kenya. *Int J Infect Dis* 2021;**112**:25–34. doi: [10.1016/j.ijid.2021.08.062](https://doi.org/10.1016/j.ijid.2021.08.062). Epub 2021 Sep 2. PMID: 34481966; PMCID: PMC8411609
8. McKay T, Robinson RS, Musungu S, *et al.* The missing millions: uncovering the burden of Covid-19 cases and deaths in the African region. *Pop Dev Review* 2024;**50**:7–58. doi: <https://doi.org/10.1111/padr.12608>
9. Meyerowitz EA, Richterman A, Gandhi RT, *et al.* Transmission of SARS-CoV-2: a review of viral host, and environmental factors. *Ann Intern Med* 2020;**174**(1):69–79. doi: [10.7326/M20-5008](https://doi.org/10.7326/M20-5008). PMCID: PMC7505025. PMID: [32941052](https://pubmed.ncbi.nlm.nih.gov/32941052/)
10. Nishiura H, Kobayashi T, Miyama T, *et al.* Estimation of the asymptomatic ratio of novel coronavirus infections (COVID-19). *Int J Infect Dis* 2020;**94**:154–155. doi: <https://doi.org/10.1016/j.ijid.2020.03.020>

11. Ogalo EA, Gudu E, Andale T, Korir D, Ndege S, Simiyu T, Olekuyo R, Mwangi H, Kimaiyo S, Aruasa W. Clinical spectrum of COVID-19 at a national referral hospital in western Kenya during the period 2020–2021. *Front Virol* 2023;**3**:1202742. doi: 10.3389/fviro.2023.1202742

12. Johansson MA, Quandelacy TM, Kada S, *et al.* SARS-CoV-2 transmission from people without COVID-19 symptoms. *JAMA Network Open* 2021;**4**(1):e2035057. doi:10.1001/jamanetworkopen.2020.35057

13. Oliveira MC, Scharan KO, Thomés BI, *et al.* Diagnostic accuracy of a set of clinical and radiological criteria for screening of COVID-19 using RT-PCR as the reference standard. *BMC Pulm Med* 2023;**23**:81 <https://doi.org/10.1186/s12890-023-02369-9>

14. Cheng MP, Yansouni CP, Basta NE, *et al.* Serodiagnostics for severe acute respiratory syndrome-related coronavirus-2: a narrative review. *Annals of Internal Medicine* 2020;**173**:6. <https://doi.org/10.7326/M20-2854>

15. Aluga, MA. Coronavirus disease 2019 (COVID-19) in Kenya: preparedness, response and transmissibility. *J Microbiol Immunol Infect* 2020;**53**(5):671–673. doi: <https://doi.org/10.1016/J.JMII.2020.04.011>

16. World Bank. Socioeconomic impacts of COVID-19 in Kenya. 2021. Available <https://documents1.worldbank.org/curated/en/949721626096781344/pdf/Socioeconomic-Impacts-of-COVID-19-in-Kenya.pdf>

17. Ministry of Health, Republic of Kenya. National COVID-19 Vaccine Deployment Plan. August 2021.

18. Ministry of Health, Republic of Kenya. Kenya COVID-19 Vaccination Program: Daily Situation Report. 5 February 2022.

19. Uyoga S, Adetifa IMO, Otiende M, *et al.* Seroprevalence of anti–SARS-CoV-2 IgG antibodies in Kenyan blood donors. *Science* 2021;**371**(6524):79–82. doi:10.1126/science.abe1916. Epub 2020 Nov 11.

20. Uyoga S, Adetifa IMO, Otiende M, *et al.* Prevalence of SARS-CoV-2 antibodies from a national serosurveillance of Kenyan blood donors, January–March 2021. *JAMA* 2021;**326**(14):1436–1438. doi: 10.1001/jama.2021.15265

21. Lucinde R, Mugo D, Bottomley C, *et al.* Serosurveillance for IgG to SARS-CoV-2 at antenatal care clinics in three Kenyan referral hospitals: repeated cross-sectional surveys 2020–21. *PLoS ONE* 2022;**17**(10):e0265478. doi: <https://doi.org/10.1371/journal.pone.0265478>

22. Etyang AO, Lucinde R, Karanja H, *et al.* Seroprevalence of antibodies to severe acute respiratory syndrome coronavirus 2 among healthcare workers in Kenya. *Clin Infect Dis* 2021;**74**(2):288–293. <https://doi.org/10.1093/cid/ciab346>
23. Kagucia EW, Gitonga JN, Kalu C, *et al.* Seroprevalence of anti-SARS-CoV-2 IgG antibodies among truck drivers and assistants in Kenya. Book Review. *Open Forum Infect Dis* 2021;**8**(7):1–4. doi: <https://doi.org/10.1101/2021.02.12.21251294>
24. Crowell TA, Daud II, Maswai J, *et al.* 2021. SARS-CoV-2 antibody prevalence in people with and without HIV in rural western Kenya, January to March 2020. *AIDS* 35(14):2401–2404. doi:10.1097/QAD.0000000000003054
25. Kenya National Bureau of Statistics. The 2009 Kenya Population and Housing Census. August 2010.
26. Mutisya E, Yarime M. Understanding the grassroots dynamics of slums in Nairobi: the dilemma of Kibera informal settlements [Internet]. Vol. 2, *Int Transact J Eng Manage Appl Sci Technol*. 2011 [Accessed 2020 May 18]. Available: <http://tuengr.com/V02/197-213.pdf>
27. UN Habitat. COVID-19 exposes the harsh realities of gender inequality in slums. 2020. <https://unhabitat.org/news/03-jun-2020/covid-19-exposes-the-harsh-realities-of-gender-inequality-in-slums#:~:text=In%20Kibera%2C%20Kenya%2C%20the%20world's,women%20for%20every%20100%20men>
28. Statistics How To. Sample size in Statistics (how to find it): Excel, Cochran's formula, general tips. <https://www.statisticshowto.com/probability-and-statistics/find-sample-size/>
29. Carter JY, Khamadi S, Mwangi J, *et al.* 2023. Performance of one rapid test and two ELISAs for a SARS-CoV-2 seroprevalence survey in Kibera informal settlement, Nairobi, Kenya. Manuscript submitted for publication.
30. World Health Organization. COVID-19: Case definitions. 16 December 2020.
31. Ministry of Health, Kenya. Interim guidelines on management of COVID-19 in Kenya. COVID-19, Infection Prevention and Control (IPC) and Case Management. June 2020.
32. Laws of Kenya. Employment Act. Chapter 226. Revised Edition 2012 [2007].
33. Wamai RG, Hirsch JL, Van Damme W, *et al.* What could explain the lower COVID-19 burden in Africa despite considerable circulation of the SARS-CoV-2 virus? *Int J Environ Res Public Health* 2021;**18**(16):8638. doi: <https://doi.org/10.3390/ijerph18168638>
34. Musa HH, Musa TH, Musa IH, *et al.* Addressing Africa's pandemic puzzle: perspectives on COVID-19 transmission and mortality in sub-Saharan Africa. *Int J Inf Diseases* 2021;**102**:483–488. doi: [10.1016/j.ijid.2020.09.1456](https://doi.org/10.1016/j.ijid.2020.09.1456). PMID: 33010461. PMCID: [PMC7526606](https://pubmed.ncbi.nlm.nih.gov/PMC7526606/)



35. Adams J, MacKenzie MJ, Amegah AK, *et al.* The conundrum of low COVID-19 mortality burden in sub-Saharan Africa: myth or reality? *Glob Health Sci Pract* 2021;**9**(3):433–443. doi: <https://doi.org/10.9745/GHSP-D-21-00172>

36. Tessema SK, Nkengasong JN. Understanding COVID-19 in Africa. *Nat Rev Immunol* 2021;**21**:469–470. <https://doi.org/10.1038/s41577-021-00579-y>

37. Kusi KA, Frimpong A, Partey FD, *et al.* High infectious disease burden as a basis for the observed high frequency of asymptomatic SARS-CoV-2 infections in sub-Saharan Africa [version 3; peer review: 2 approved] *AAS Open Research* 2021;**4**:2. doi: <https://doi.org/10.12688/aasopenres.13196.3>

38. United Nations. New York: United Nations Publication; 2016. [2020-04-23]. Urbanization and development: emerging futures. World Cities report. <https://tinyurl.com/y4ekncmh>.

39. Gibson L, Rush D. Novel coronavirus in Cape Town informal settlements: feasibility of using informal dwelling outlines to identify high risk areas for COVID-19 transmission from a social distancing perspective. *JMIR Public Health Surveill* 2020;**6**(2):e18844. Available: <http://publichealth.jmir.org/2020/2/e18844/> doi: 10.2196/18844. PMID: 32250283

40. Wasdania KP, Prasada A. The impossibility of social distancing among the urban poor: the case of an Indian slum in the times of COVID-19. *Local Environment* 25(5):414–418. <https://doi.org/10.1080/13549839.2020.1754375>

41. UN Habitat. Coronavirus will travel 'incredibly fast' in Africa's slums. April 2020. Available: <https://unhabitat.org/coronavirus-will-travel-incredibly-fast-in-africas-slums-un-habitat-chief-warns>

42. Arora RK, Joseph A, Van Wyk J, *et al.* SeroTracker: a global SARS-CoV-2 seroprevalence dashboard. *Lancet Infect Dis* 2021;**4**:E75–76. doi: 10.1016/S1473-3099(20)30858-6

43. Ilesanmi O, Oderinde T, Afolabi A. The urban slums: potential source of COVID-19 spikes in Africa. *Public Health Pract (Oxf)* 2020; Nov:1:100052. doi: 1:10.1016/j.puhip.2020.100052.

44. Malani A, Shah D, Kang G, *et al.* Seroprevalence of SARS-CoV-2 in slums versus non-slums in Mumbai, India. *The Lancet Global Health* 2021;**9**(2)February E110-111. doi: [https://doi.org/10.1016/S2214-109X\(20\)30467-8](https://doi.org/10.1016/S2214-109X(20)30467-8)

45. Nirala SK, Naik BN, Chaudhary N, *et al.* Sero-epidemiological survey of SARS-CoV2 in urban slums of a capital city: a cross- sectional study. *J Family Med Prim Care* 2022;**11**(6):2709–16. doi: 10.4103/jfmpe.jfmpe\_2127\_21

46. Raqib R, Sarker P, Akhtar E, *et al.* Seroprevalence of SARS-CoV-2 infection and associated factors among Bangladeshi slum and non-slum dwellers in pre-COVID-19 vaccination era: October 2020 to February 2021. *PLoS ONE* 2022;**17**(5):e0268093. doi: <https://doi.org/10.1371/journal.pone.0268093>



47. George CE, Inbaraj LR, Chandrasingh S, *et al.* High seroprevalence of COVID-19 infection in a large slum in South India; what does it tell us about managing a pandemic and beyond? *Epid Inf* 2021;149:e39,1–6. doi: <https://doi.org/10.1017/S0950268821000273>
48. Martono, Fatmawati F, Mulyanti S. Risk factors associated with the severity of COVID-19. *Malays J Med Sci* 2023;30(3):84–92. doi:10.21315/mjms2023.30.3.7. Epub 2023 Jun 27. PMID: 37425387; PMCID: PMC10325129.
49. Wang B, Yuan S, Ruan S, *et al.* Associations between underlying diseases with COVID-19 and its symptoms among adults: a cross-sectional study. *Front Public Health* 2023;11:1210800. doi: 10.3389/fpubh.2023.1210800
50. Kompaniyets L, Agathis NT, Nelson JM, *et al.* Underlying medical conditions associated with severe COVID-19 illness among children. *JAMA Netw Open* 2021;4(6):e2111182. doi:10.1001/jamanetworkopen.2021.11182. PMID: 34097050. PMCID: PMC8185607.
51. Costa FF, Rosário WR, Farias ACR, *et al.* Metabolic syndrome and COVID-19: an update on the associated comorbidities and proposed therapies. *Diabetes Metab Syndr: Clinical Research & Reviews* 2020;14(5):809–814. doi: <https://doi.org/10.1016/j.dsx.2020.06.016>
52. Wu ZH, Tang Y, Cheng Q. Diabetes increases the mortality of patients with COVID-19: a meta-analysis. *Acta Diabetol* 2021;58:139–144. doi: <https://doi.org/10.1007/s00592-020-01546-0>. PMID: 32583078; PMCID: PMC7311595.
53. Jin JM, Bai P, He W, *et al.* Gender differences in patients with COVID-19: focus on severity and mortality. *Front Public Health* 2020;8:152. doi: 10.3389/fpubh.2020.00152. eCollection 2020.
54. Adnan N, Khandker SS, Haq A, *et al.* Detection of SARS-CoV-2 by antigen ELISA test is highly swayed by viral load and sample storage condition. *Expert Rev Anti Infect Ther* 2022;20(3):473–481. doi: 10.1080/14787210.2021.1976144. Epub 2021 Sep 11. PMID: 34477019; PMCID: PMC8442762.
55. Høeg TB, Prasad V. Rapid antigen testing for COVID-19: decreasing diagnostic reliability, potential detrimental effects and a lack of evidence to support continued public funding of community-based testing. *Public Health Pract (Oxf)* 2023;6:100451. doi: 10.1016/j.puhip.2023.100451. PMID: 38075468; PMCID: PMC10701347.
56. Mallett S, Allen AJ, Graziadio S, *et al.* At what times during infection is SARS-CoV-2 detectable and no longer detectable using RT-PCR-based tests? A systematic review of individual participant data. *BMC Medicine* 2020;18:346. doi: <https://doi.org/10.1186/s12916-020-01810-8>
57. Guo L, Ren L, Yang S, *et al.* Profiling early humoral response to diagnose novel coronavirus disease (COVID-19). *Clin Infect Dis* 2020;71(15):778–785. <https://doi.org/10.1093/cid/ciaa310>

58. Zhao J, Yuan Q, Wang H, *et al.* Antibody responses to SARS-CoV-2 in patients with novel coronavirus disease 2019. *Clin Infect Dis* 2020;**71**(16):2027–34. doi: 10.1093/cid/ciaa344.

59. Qu J, Wu C, Li X, *et al.* Profile of Immunoglobulin G and IgM antibodies against severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). *Clin Infect Dis* 2020;**71**(16):2255–58. doi: 10.1093/cid/ciaa489.

60. Hoffman T, Nissen K, Krambrich J, *et al.* Evaluation of a COVID-19 IgM and IgG rapid test: an efficient tool for assessment of past exposure to SARS-CoV-2. *Infect Ecol Epidemiol* 2020;**10**(1):1754538. doi:10.1080/20008686.2020.1754538. PMID: 32363011. PMCID: PMC7178815.

61. Hsueh PR, Huang LM, Chen PJ, *et al.* Chronological evolution of IgM, IgA, IgG and neutralisation antibodies after infection with SARS-associated coronavirus. *Clin Microbiol Infect* 2004;**10**(12):1062–6. doi:10.1111/j.1469-0691.2004.01009.x. PMID: 15606632. PMCID: PMC7129952.

62. Li G, Chen X, Xu A. Letter to the Editor. Profile of specific antibodies to the SARS-associated coronavirus. *N Engl J Med* 2003;**349**(5):508–509. doi: [10.1056/NEJM200307313490520](https://doi.org/10.1056/NEJM200307313490520)

63. Hanssen DAT, Slaats M, Mulder M, *et al.* Evaluation of 18 commercial serological assays for the detection of antibodies against SARS-CoV-2 in paired serum samples. *Eur J Clin Microbiol Infect Dis* 2021;**40**(8):1695-1703. doi:10.1007/s10096-021-04220-7. Epub 2021 Mar 17. PMID: 33733395; PMCID: PMC7968571.

64. Gong F, Wei H, Li Q, *et al.* Evaluation and comparison of serological methods of COVID-19 diagnosis. *Front Mol Biosci* 2021;**8** Sec. Structural Biology. doi: <https://doi.org/10.3389/fmolb.2021.682405>

65. Dan JM, Mateus J, Kato Y, *et al.* Immunological memory to SARS-CoV-2 assessed for up to 8 months after infection. *Science* 2021;**371**(6529):eabf4063. doi: 10.1126/science.abf4063

66. Alejo JL, Mitchell J, Chang A, *et al.* Prevalence and durability of SARS-CoV-2 antibodies among unvaccinated US adults by history of COVID-19. *Research Letter JAMA* 2022;**327**(11):1085–1087. doi:10.1001/jama.2022.1393

67. Guo L, Zhang Q, Gu X, *et al.* Durability and cross-reactive immune memory to SARS-CoV-2 in individuals 2 years after recovery from COVID-19: a longitudinal cohort study. *Lancet Microbe* 2024;**5**(1):E24–E33. doi: [https://doi.org/10.1016/S2666-5247\(23\)00255-0](https://doi.org/10.1016/S2666-5247(23)00255-0)

68. World Health Organization. 2023. Update to requirements for reporting COVID-19 surveillance data under the International Health Regulations (IHR 2005): Addendum to Public health surveillance for

- COVID-19 interim guidance, 25 August 2023. <https://www.who.int/publications/i/item/WHO-2019-nCoV-Surveillance-Guidance-Addendum-2023-1>
69. National Centre for Immunization and Respiratory Diseases. 2024. The changing threat of COVID-19. 23 February 2024. <https://www.cdc.gov/ncird/whats-new/changing-threat-covid-19.html>
70. Pak A, Adegboye OA, Adekunle AI, *et al*. Economic consequences of the COVID-19 outbreak: the need for epidemic preparedness. *Front Public Health* 2020;**8**:241. doi: 10.3389/fpubh.2020.00241
71. Tessema GA, Kinfu Y, Dachew BA, *et al*. The COVID-19 pandemic and healthcare systems in Africa: a scoping review of preparedness, impact and response. *BMJ Global Health* 2021;**6**(12):e007179. doi:10.1136/bmjgh-2021-007179
72. World Health Organization. Media Briefing. 5 May 2023. Available: <https://www.who.int/director-general/speeches/detail/who-director-general-s-opening-remarks-at-the-media-briefing---5-may-2023>
73. World Health Organization. 2024. WHO launches CoViNet: a global network for coronaviruses. <https://www.who.int/news/item/27-03-2024-who-launches-covinet--a-global-network-for-coronaviruses>
74. European Centre for Disease Prevention and Control. <https://www.ecdc.europa.eu/en/infectious-disease-topics/z-disease-list/covid-19/facts/surveillance-covid-19>
75. Haselbeck AH, Im J, Prifti K, *et al*. Serology as a tool to assess infectious disease landscapes and guide public health policy. *Pathogens* 2022;**11**(7):732. doi: <https://doi.org/10.3390/pathogens11070732>

Table 1. Seroprevalence of antibodies to SARS-CoV-2, by sociodemographic and other participant characteristics

| Overall                                                                           | n                       | positive (%)           |                     |
|-----------------------------------------------------------------------------------|-------------------------|------------------------|---------------------|
|                                                                                   | 438                     | 370 (84.5)             |                     |
| Variables                                                                         | n (%)                   | positive [% (95% CI)]  | p (chi-square test) |
| Age                                                                               |                         |                        |                     |
| ≤17                                                                               | 21 (4.8)                | 14 [3.2 (1.54–4.9)]    | 0.0670              |
| 18–30                                                                             | 173 (39.5)              | 146 [33.3 (28.9–37.8)] |                     |
| 31–50                                                                             | 174 (39.7)              | 153 [34.9 (30.5–39.4)] |                     |
| >50                                                                               | 70 (16)                 | 57 [13.0 (9.9–16.2)]   |                     |
| Sex                                                                               |                         |                        |                     |
| Female                                                                            | 281 (64.2)              | 241 [55.0 (50.3–59.7)] | 0.3185              |
| Male                                                                              | 157 (35.8)              | 129 [29.5 (25.2–33.7)] |                     |
| Employment status*undisclosed was not included in the calculation of significance |                         |                        |                     |
| Unemployed                                                                        | 164 (39.1)              | 137 [31.4 (27.0–35.7)] | 0.0176              |
| Self-employed                                                                     | 102 (24.3)              | 82 [18.8 (15.1–22.4)]  |                     |
| Casual/part-time                                                                  | 88 (21.0)               | 83 [19.0 (15.3–22.7)]  |                     |
| Student                                                                           | 34 (8.1)                | 27 [6.2 (3.9–8.4)]     |                     |
| Formal/regular                                                                    | 31 (7.4)                | 28 [6.4 (4.1–8.7)]     |                     |
| Not applicable (<16 years)                                                        | 18                      | 12 [2.7 (1.2–4.3)]     |                     |
| Undisclosed                                                                       | 1                       | 1 [0.23]               |                     |
| Underlying health conditions                                                      |                         |                        |                     |
| No                                                                                | 341 (77.9)              | 287 [65.5 (61.1–70.0)] | 0.7364              |
| Yes                                                                               | 97 (22.1)               | 83 [19.0 (15.3–22.6)]  |                     |
| Types of underlying health conditions                                             | (% of all participants) |                        |                     |
| Cardiovascular disease                                                            | 40 (9.1)                | 32 [7.3 (4.9–9.8)]     | 0.4123              |
| Stomach ulcer                                                                     | 25 (5.7)                | 20 [4.6 (2.6–6.5)]     | 0.5246              |
| HIV                                                                               | 18 (4.1)                | 17 [3.9 (2.1–5.7)]     | 0.2330              |
| Respiratory disease                                                               | 14 (3.2)                | 12 [2.7 (1.2–4.3)]     | 0.8964              |
| Diabetes                                                                          | 11 (2.5)                | 10 [2.3 (0.9–3.7)]     | 0.5506              |
| On medication                                                                     |                         |                        |                     |
| No                                                                                | 343 (78.3)              | 291 [66.4 (62.0–70.9)] | 0.6888              |
| Yes                                                                               | 95 (21.7)               | 79 [18.0 (14.4–21.7)]  |                     |
| COVID-19 symptoms (previous 16 months)                                            |                         |                        |                     |
| No                                                                                | 157 (35.8)              | 142 [32.4 (28.0–36.8)] | 0.0099              |
| Yes                                                                               | 281 (64.2)              | 228 [52.1 (47.4–56.8)] |                     |
| Ever tested for SARS-CoV-2                                                        |                         |                        |                     |
| No                                                                                | 371 (84.7)              | 311 [71.0 (66.7–75.3)] | 0.3787              |
| Yes                                                                               | 67 (15.3)               | 59 [10.3–16.7]         |                     |
| Previous SARS-CoV-2 test                                                          |                         |                        |                     |
| Tested with negative result                                                       | 54 (80.6)               | 48 [71.6 (60.6–82.7)]  | *                   |
| Tested with positive result                                                       | 4 (6.0)                 | 4 [6.0 (0.1–11.8)]     |                     |
| Tested with unknown result                                                        | 9 (13.4)                | 7 [10.4 (2.0–18.0)]    |                     |
| COVID-19 vaccination                                                              |                         |                        |                     |
| No                                                                                | 408 (93.2)              | 341 [77.9 (74.0–81.8)] | 0.0561              |
| Yes                                                                               | 30 (6.8)                | 29 [6.6 (4.3–9.0)]     |                     |

\* = no chi square possible due to a negative value in the tested negative category

**Table 2. Predictors of positivity to SARS-CoV-2 antibodies (n=437)**

| Variables                                     | COR* (95% CI) | p      | AOR† (95% CI)  | p      |
|-----------------------------------------------|---------------|--------|----------------|--------|
| <i>Age</i>                                    |               |        |                |        |
| <18                                           | 0.4 (0.1–1.0) | 0.0853 | 0.4 (0.0–5.0)  | 0.5574 |
| 18–30                                         | 1             |        | 1              |        |
| 31–50                                         | 1.3 (0.7–2.5) |        | 1.2 (0.6–2.3)  |        |
| >50                                           | 0.8 (0.4–1.7) |        | 0.7 (0.3–1.6)  |        |
| <i>Employment status</i>                      |               |        |                |        |
| Unemployed                                    | 1             | 0.0320 | 1              | 0.1739 |
| Self-employed                                 | 0.8 (0.4–1.5) |        | 0.9 (0.4–1.7)  |        |
| Casual/part-time                              | 3.3 (1.2–8.8) |        | 3.2 (1.2–8.7)  |        |
| Student                                       | 0.8 (0.4–1.5) |        | 0.8 (0.3–2.1)  |        |
| Formal/regular                                | 1.8 (0.5–6.5) |        | 1.8 (0.5–6.6)  |        |
| Not applicable (<16 years)                    | 0.8 (0.3–1.9) |        | 1.2 (0.1–19.9) |        |
| <i>COVID-19 symptoms (previous 16 months)</i> |               |        |                |        |
| No                                            | 1             | 0.0121 | 1              | 0.0085 |
| Yes                                           | 0.5 (0.3–0.8) |        | 0.4 (0.2–0.8)  |        |
| <i>COVID-19 vaccination</i>                   |               |        |                |        |
| No                                            | 0.2 (0.0–1.3) | 0.0893 | 0.2 (0.0–1.3)  | 0.0822 |
| Yes                                           | 1             |        | 1              |        |

\*Crude odds ratio; †Adjusted odds ratio



# BMJ Open

## Seroprevalence and demographic characteristics of SARS-CoV-2- infected residents of Kibera Informal Settlement during the COVID-19 pandemic in Nairobi, Kenya. A cross-sectional study

|                                 |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
|---------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Journal:                        | BMJ Open                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         |
| Manuscript ID                   | bmjopen-2024-094546.R1                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| Article Type:                   | Original research                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| Date Submitted by the Author:   | 20-Feb-2025                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |
| Complete List of Authors:       | <p>Carter, Jane; Amref Health Africa, Regional Laboratory Programme Khamadi, Samoel; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Mwangi, Joseph; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Muhula, Samuel; Amref Health Africa, Learning and Impact</p> <p>Munene, Stephen; Amref Health Africa, Regional Laboratory Programme</p> <p>Kanyara, Lucy; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Kinyua, Joyceline; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Lagat, Nancy; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Chege, Judy; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Oira, Robert; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Maiyo, Alex; Kenya Medical Research Institute, Centre for Virus Research</p> <p>Stewart, Roy; University Medical Centre Groningen, Department of Health Sciences, Community &amp; Occupational Medicine</p> <p>Postma, Maarten; UMCG, Department of Health Sciences; UMCG, Department of Economics, Econometrics and Finance</p> <p>Stekelenburg, Jelle; Medisch Centrum Leeuwarden, Department of Obstetrics and Gynaecology; Universitair Medisch Centrum Groningen, Department of Health Sciences, Global Health</p> <p>Osuri, Joachim; Amref International University, Office of the Vice Chancellor</p> <p>van Hulst, Marinus; Martini Hospital, Clinical Pharmacy; UMCG, Department of Health Sciences</p> |
| <b>Primary Subject Heading</b>: | Epidemiology                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| Secondary Subject Heading:      | Public health                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| Keywords:                       | COVID-19, EPIDEMIOLOGY, PUBLIC HEALTH, Community-Based Participatory Research                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
|                                 |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |





I, the Submitting Author has the right to grant and does grant on behalf of all authors of the Work (as defined in the below author licence), an exclusive licence and/or a non-exclusive licence for contributions from authors who are: i) UK Crown employees; ii) where BMJ has agreed a CC-BY licence shall apply, and/or iii) in accordance with the terms applicable for US Federal Government officers or employees acting as part of their official duties; on a worldwide, perpetual, irrevocable, royalty-free basis to BMJ Publishing Group Ltd ("BMJ") its licensees and where the relevant Journal is co-owned by BMJ to the co-owners of the Journal, to publish the Work in this journal and any other BMJ products and to exploit all rights, as set out in our [licence](#).

The Submitting Author accepts and understands that any supply made under these terms is made by BMJ to the Submitting Author unless you are acting as an employee on behalf of your employer or a postgraduate student of an affiliated institution which is paying any applicable article publishing charge ("APC") for Open Access articles. Where the Submitting Author wishes to make the Work available on an Open Access basis (and intends to pay the relevant APC), the terms of reuse of such Open Access shall be governed by a Creative Commons licence – details of these licences and which [Creative Commons](#) licence will apply to this Work are set out in our licence referred to above.

Other than as permitted in any relevant BMJ Author's Self Archiving Policies, I confirm this Work has not been accepted for publication elsewhere, is not being considered for publication elsewhere and does not duplicate material already published. I confirm all authors consent to publication of this Work and authorise the granting of this licence.

1

2

3

4

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

# Seroprevalence and demographic characteristics of SARS-CoV-2-infected residents of Kibera Informal Settlement during the COVID-19 pandemic in Nairobi, Kenya. A cross-sectional study

5

6

7

8

9

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

Jane Y Carter,<sup>1</sup> Samoel Khamadi,<sup>2</sup> Joseph Mwangi,<sup>2</sup> Samuel Muhula,<sup>1</sup> Stephen M Munene,<sup>1</sup> Lucy Kanyara,<sup>2</sup> Joyceline Kinyua,<sup>2</sup> Nancy Lagat,<sup>2</sup> Judy Chege,<sup>2</sup> Robert Oira,<sup>2</sup> Alex Maiyo,<sup>2</sup> Roy Stewart,<sup>3</sup> Maarten Postma,<sup>3,4</sup> Jelle Stekelenburg,<sup>3,5</sup> Joachim Osur,<sup>6</sup> Marinus van Hulst<sup>3,7</sup>

<sup>1</sup> Amref Health Africa Headquarters, PO Box 27691–00506, Nairobi, Kenya

<sup>2</sup>Centre for Virus Research, Kenya Medical Research Institute, PO Box 54840–00200, Nairobi, Kenya

<sup>3</sup>Department of Health Sciences, University of Groningen, University Medical Center Groningen, Groningen, Netherlands

<sup>4</sup>Department of Economics, Econometrics and Finance, University of Groningen, Faculty of Economics and Business, Groningen, Netherlands

<sup>5</sup>Department of Obstetrics and Gynaecology, Medical Centre Leeuwarden, Leeuwarden, Netherlands

<sup>6</sup>Amref International University, PO Box 27691–00506, Nairobi, Kenya

<sup>7</sup>Department of Clinical Pharmacy and Toxicology, Martini Hospital, Groningen, Netherlands

24

25

26

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

58

59

60

Correspondence should be addressed to Dr Jane Y Carter

Amref Health Africa Headquarters, PO Box 27691–00506, Nairobi, Kenya; email: [jane.carter@amref.org](mailto:jane.carter@amref.org)

## ABSTRACT

**Objectives** To assess the prevalence of SARS-CoV-2 antibodies in the residents of Kibera informal settlement in Nairobi, Kenya, before vaccination became widespread, and explore demographic and health-related risk factors for infection.

**Design** A cross-sectional study.

**Setting** Kibera informal settlement, Nairobi, Kenya.

**Participants** Residents of Kibera informal settlement between October 2019 and August 2021, age 1 year and above who reported no current symptoms of COVID-19.

**Main outcome measures** Associations were determined between SARS-CoV-2 positive tests measured with one rapid test and two enzyme-linked immunosorbent assays and demographic and health-related factors, using Pearson's chi-square test. Crude odds ratio and adjusted odds ratio were calculated to quantify the strength of associations between variables and seropositive status.

**Results** A total of 438 participants were recruited. Most (79.2%) were age 18–50 years; females (64.2%) exceeded males. More than one third (39.1%) were unemployed; only 7.4% were in formal, full-time employment. Less than one quarter (22.1%) self-reported any underlying health conditions. Nearly two-thirds (64.2%) reported symptoms compatible with COVID-19 in the previous 16 months; only one (0.23%) had been hospitalised with a reported negative COVID-19 test. 370 (84.5%) participants tested positive in any of the three tests. There was no significant difference in SARS-CoV-2 seropositivity across age, sex, presence of underlying health conditions, on medication, or those ever tested for SARS-CoV-2. Multiple logistic regression analysis showed COVID-19 symptoms in the previous 16 months were the only significant independent predictor of seropositivity ( $p=0.0085$ ).

**Conclusion** High SARS-CoV-2 exposure with limited morbidity was found in residents of Kibera informal settlement. The study confirms other reports of high SARS-CoV-2 exposure with limited morbidity in slum communities. Reason cited include the high infectious disease burden on the African continent, demographic age structure, and underreporting due to limited testing and lack of access to healthcare services; genetic factors may also play a role. These factors require further investigation.





## INTRODUCTION

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the cause of coronavirus disease 2019 (COVID-19), was first reported in Wuhan, Hubei Province, China, in November 2019 and rapidly spread across China and the world [1]. On 30 January 2020, the World Health Organization (WHO) declared the outbreak a Public Health Emergency of International Concern, and on 11 March 2020 the WHO Director General declared COVID-19 a pandemic [2]. Although the WHO declared the COVID-19 global health emergency over on 5 May 2023 [3], globally more than 777 million cases and more than 7 million deaths from COVID-19 had been reported by 5 January 2025, with more than 161,000 new cases and nearly 3,000 deaths in the previous 28 days [4]. In the African region, more than 9.5 million confirmed cases and more than 175,000 deaths had been reported cumulatively by 5 January 2025, although the numbers of cases and deaths reported in Africa are likely an underestimate due to low testing rates [5]; an assessment by the WHO Regional Office for Africa showed only one in seven (14.2%) COVID-19 infections were being detected [6]. Studies from Kenya and other countries in Africa noted low mortality from COVID-19, despite high exposure rates [7,8].

The modes of transmission of SARS-CoV-2 have been elucidated through detailed case contact studies. Respiratory transmission, with SARS-CoV-2 carried on tiny particles emitted from the respiratory tract, has been established as the clear and dominant route of spread; indirect transmission appears to be of limited importance despite initial concerns [Error! Reference source not found.]. The clinical spectrum of COVID-19 varies from asymptomatic or pre-symptomatic infection, mild to moderate illness, to severe and critical illness characterised by respiratory failure and multiple organ dysfunction; varying proportions of infected persons remain asymptomatic [10,11]. Transmission from asymptomatic individuals is estimated to account for more than half of all transmissions [12]. Diagnostic testing using reverse transcriptase polymerase chain reaction (RT-PCR)-based assays performed on respiratory specimens is the reference standard for establishing a microbiological diagnosis of COVID-19 [13] but is constrained by the presence of virus to a few days before infection and a short time after infection. Seroepidemiological studies to detect host antibodies to SARS-CoV-2 are therefore important for estimating disease burden and providing a more complete picture of exposure to SARS-CoV-2 in a population [14].

### COVID-19 infection in Kenya

The first confirmed case of COVID-19 was reported in Kenya on 13 March 2020 [15]. Restrictions to mitigate the spread of COVID-19 were instituted in March 2020 and were eased towards the end of 2020

1 until a second lockdown was instituted in five counties in March 2021; in May 2021, as the number of  
2 cases dropped, lockdown restrictions were again lifted [16]. COVID-19 vaccination was initiated in Kenya  
3 in March 2021, with vaccination numbers rising from August 2021 with increased vaccine availability.  
4 Kenya actively promoted vaccination through the COVID-19 Vaccination Acceleration Programme [17];  
5 as of 16 May 2023, more than 18 million vaccine doses had been administered with 30.7% of the adult  
6 population (≥18 years) fully vaccinated. Kenya planned to vaccinate all adults and teenagers in 2022 and  
7 to provide third dose booster shots to all eligible adults [18]. On 11 March 2022, Kenya lifted all COVID-  
8 19 restrictions while urging continued personal public health measures; by June 2024, 344,000 cases and  
9 5,700 deaths had been reported since the pandemic started, rates that remained low in comparison with  
10 global figures [4].

11 Seroprevalence data from Kenya targeting different population groups painted a concerning picture of  
12 the pandemic’s progression across the country. Studies examining stored samples from blood donors in  
13 six regional blood transfusion centres, including the capital city Nairobi, revealed a dramatic rise in SARS-  
14 CoV-2 antibody prevalence over time. Seroprevalence increased from 4.3% (95% CI, 2.9–5.8%) in  
15 samples collected between April and June 2020 to 48.5% (95% CI, 45.2–52.1%) in samples taken just one  
16 year later from January to March 2021 [19,20]. These samples were tested using a non-commercial  
17 validated enzyme-linked immunosorbent assay (ELISA) for SARS-CoV-2 IgG against spike protein. The  
18 same assay was used to test stored blood from women attending antenatal care services at three  
19 hospitals in Kenya between August 2020 and October 2021: seroprevalence rose from 50% (95% CI, 42–  
20 58%) in August 2020 to 85% (95% CI, 78–92%) in October 2021 in Nairobi; 31% (95% CI, 25–37%) in May  
21 2021 to 71% (95% CI, 64–77%) in October 2021 in Busia; and from 1% (95% CI, 0–3%) in September 2020  
22 to 63% (95% CI, 56–69%) in October 2021 in Kilifi [21]. Purposive testing of venous blood from healthcare  
23 workers between July and December 2020 using the same ELISA showed significant variation in overall  
24 seropositivity (20.8%: 17.5%–24.4%). Seroprevalence varied significantly ( $p<0.001$ ) by site: 43.8% (35.8–  
25 52.2%) in Nairobi, 12.6% (8.8–17.1%) in Busia and 11.5% (7.2–17.6%) in Kilifi; only 16 (2%) of the sampled  
26 healthcare workers reported acute respiratory symptoms at the time of sample collection [22]. Purposive  
27 testing of truck drivers and their assistants, using the same ELISA, conducted between September and  
28 October 2020 showed an overall seropositivity of 7.4%; none reported current or previous symptoms of  
29 illness [23]. Truck drivers and their assistants continued to transport essential supplies during the COVID-  
30 19 pandemic, placing them at increased risk of being infected and of transmitting SARS-CoV-2 over a  
31 wide geographical area. In a study on samples collected from January to March 2020 from rural

populations in western Kenya with and without human immunodeficiency virus (HIV) infection, 3.3% had detectable SARS-CoV-2 antibodies, with no difference between participants with and without HIV infection (3.1% vs. 4.0%,  $p=0.68$ ) [24]; this study used the Platelia SARS-CoV-2 Total Ab assay (Bio-Rad Laboratories, Hercules, California, USA). Participants denied symptoms except for one who reported a cough in the preceding week. In our study, SARS-CoV-2 seroprevalence in an informal settlement in Nairobi was conducted to supplement these data and determine exposure in a community subjected to overcrowding and lack of reliable clean water, housing, health services and waste management facilities, where preventive measures were difficult to apply. At the time of the study, Kenya was experiencing wave 4 of the pandemic predominated by the *Delta* variant of concern (VoC) [25].

### Purpose of the study

This study assessed the prevalence of SARS-CoV-2 antibodies in the residents of Kibera informal settlement in Nairobi, Kenya, one of sub-Saharan Africa's largest slums, before vaccination became widespread, and explored some of the major risk factors for infection. This study also provided an opportunity to determine pathogen exposure in large numbers of persons in a pandemic situation using small blood volumes obtained from fingerprick samples.

## MATERIALS AND METHODS

### Study site

Kibera informal settlement stands on a 2.5-kilometre square area of land located 5 kilometres southwest of Nairobi's Central Business District. The population is estimated at 170,070, although this may be an underestimate [26]. Residents comprise the major ethnic communities in Kenya [27] with reportedly 116 women for every 100 men [28]. Kibera is governed by area chiefs as the local administrative arm of the government.

### Study design

A cross-sectional study was performed to determine SARS-CoV-2 antibody prevalence in Kibera informal settlement, Nairobi, Kenya. The study was conducted between 2 and 13 August 2021 in 10 of the 14 Kibera villages, which provided a representation of the whole settlement. The Cochran sample size formula for larger populations was used to determine sample size [29]. Since the population of Kibera is above 170,000 and approximately 500 people had tested positive for SARS-CoV-2 at the time of the study, a sample size of 389 (~400) participants was determined, a target of 40 participants in each selected village (confidence level 99%; precision 0.01). In each village every fifth household was selected;

1 if a household declined participation, the next household was included. Entry to individual households  
2 was facilitated through community health volunteers who were formally assigned to these villages.  
3 Participants age 1 year and above who had been resident in Kibera since at least October 2019 were  
4 included.

5 **Study methods**

6 Five data collectors, each assigned to two villages, administered a brief questionnaire to each consenting  
7 participant, or their parent or legal guardian; data were collected directly into the Open Data Kit system  
8 installed in handheld tablets. Data were collected on the following: age, sex, length of residence in  
9 Kibera, pregnancy status (for women), main occupation and nature of employment, any underlying  
10 health conditions and type of condition, any medications and name of the medication, symptoms  
11 suggestive of COVID-19 in the previous 16 months (cough, fever, difficulty in breathing, shortness of  
12 breath, new loss of taste and smell), any hospitalisation and date (if known), ever tested for SARS-CoV-  
13 2, date and result (if known), and COVID-19 vaccination(s) and date (if known). Five laboratory staff  
14 accompanying the data collectors collected 500 µL of capillary blood from each participant by fingerprick  
15 into a Microtainer® EDTA tube. Blood samples were transported to the Kenya Medical Research Institute  
16 (KEMRI) laboratories for same-day testing on whole blood using the Standard Q COVID-19 IgM/IgG  
17 Combo rapid test (SD Biosensor Inc, Suwon, Gyeonggi, Republic of Korea). At the KEMRI laboratories,  
18 blood was separated the same day and plasma samples stored at -80 °C for further testing with two  
19 enzyme-linked immunosorbent assays (ELISAs): Wantai Total Ab (IgM/IgG/IgA) ELISA for SARS-CoV-2  
20 (Wantai Biological Pharmacy Enterprise Co. Ltd, Beijing, China), and Platelia SARS-CoV-2 Total Ab  
21 (IgM/IgG/IgA) Assay (Bio-Rad Laboratories Inc, California, USA). The SARS-CoV-2 antibody tests selected  
22 in this survey were commercial tests approved for use in Kenya at the time of the study. Test  
23 performance is described in a separate paper [30]. Individuals with positive test results in any of the  
24 three tests were considered positive (for the ELISAs, optical density ≥1.0) and indeterminate ELISA results  
25 as negative (optical density 0.9–0.99). PCR or antigen testing was not performed for any of the study  
26 participants. Each participant was assigned a unique identification number; data files and blood sample  
27 labelling did not include participant names to ensure anonymity. Individuals who met the case definition  
28 of COVID-19 infection [31] at the time of recruitment were not included in the study and were referred  
29 to a nearby health facility for further investigation and management. The research teams followed  
30 government regulations for the prevention of COVID-19 transmission during data collection [32].

31 **Statistical analysis**

Data were entered into Excel, cleaned and analysed using SAS version 9.4 (SAS Institute Inc). Frequency and percentages were used to present the data; Pearson's chi-square test was used to examine the association between seropositive results and various demographic and health-related factors, including age, sex, employment status, presence and types of underlying health conditions, on any type of medication, COVID-19 vaccination, previous symptoms compatible with COVID-19 and those ever tested for SARS-CoV-2. Simple logistic regression estimated the crude odds ratio (COR) for each variable found to be significantly associated with seropositivity in the chi-square test ( $\leq 0.20$ ). Multiple logistic regression was used to calculate the adjusted odds ratio (AOR), controlling for the effects of multiple variables simultaneously. The COR and AOR were calculated to quantify the strength of associations between the variables and seropositive status. Vaccinated individuals were included in the overall analysis as numbers were small (6.8%).

### Patients and public involvement

This study was planned, designed and conducted as a shared activity with Kenya's national Virus Research Centre. This included development of the research protocol and study tools, training of data collectors and field laboratory staff, and sample collection. The Virus Research Centre staff conducted all the laboratory tests. The study was discussed with the five area administrative chiefs, and entry to the households was facilitated through the community health volunteers assigned to the study villages. The study results will be presented to the Kibera chiefs, community health volunteers and the participants through a community meeting.

## RESULTS

### Demographic data

A total of 438 participants consented to answer the questions and to a blood sample collection by fingerprick; one participant refused to divulge her occupation. Table 1 shows the information collected from the participant questionnaire.

**Table 1. Seroprevalence of antibodies to SARS-CoV-2 by sociodemographic and other participant characteristics**

| Overall    | <i>n</i>     | positive (%)           |                            |
|------------|--------------|------------------------|----------------------------|
|            | 438          | 370 (84.5)             |                            |
| Variables  | <i>n</i> (%) | positive [% (95% CI)]  | <i>p</i> (chi-square test) |
| <i>Age</i> |              |                        |                            |
| $\leq 17$  | 21 (4.8)     | 14 [3.2 (1.54–4.9)]    | 0.0670                     |
| 18–30      | 173 (39.5)   | 146 [33.3 (28.9–37.8)] |                            |
| 31–50      | 174 (39.7)   | 153 [34.9 (30.5–39.4)] |                            |
| >50        | 70 (16)      | 57 [13.0 (9.9–16.2)]   |                            |



|    |                                                                                            |            |                        |        |
|----|--------------------------------------------------------------------------------------------|------------|------------------------|--------|
| 1  | Sex                                                                                        |            |                        |        |
| 2  | Female                                                                                     | 281 (64.2) | 241 [55.0 (50.3–59.7)] | 0.3185 |
| 3  | Male                                                                                       | 157 (35.8) | 129 [29.5 (25.2–33.7)] |        |
| 4  | <i>Employment status (undisclosed was not included in the calculation of significance)</i> |            |                        |        |
| 5  | Unemployed                                                                                 | 164 (39.1) | 137 [31.4 (27.0–35.7)] | 0.0176 |
| 6  | Self-employed                                                                              | 102 (24.3) | 82 [18.8 (15.1–22.4)]  |        |
| 7  | Casual/part-time                                                                           | 88 (21.0)  | 83 [19.0 (15.3–22.7)]  |        |
| 8  | Student                                                                                    | 34 (8.1)   | 27 [6.2 (3.9–8.4)]     |        |
| 9  | Formal/regular                                                                             | 31 (7.4)   | 28 [6.4 (4.1–8.7)]     |        |
| 10 | Not applicable (<16 years)                                                                 | 18         | 12 [2.7 (1.2–4.3)]     |        |
| 11 | Undisclosed                                                                                | 1          | 1 [0.23]               |        |
| 12 | <i>Underlying health conditions</i>                                                        |            |                        |        |
| 13 | No                                                                                         | 341 (77.9) | 287 [65.5 (61.1–70.0)] | 0.7364 |
| 14 | Yes                                                                                        | 97 (22.1)  | 83 [19.0 (15.3–22.6)]  |        |
| 15 | <i>Types of underlying health conditions (% of all participants)</i>                       |            |                        |        |
| 16 | Cardiovascular disease                                                                     | 40 (9.1)   | 32 [7.3 (4.9–9.8)]     | 0.4123 |
| 17 | Stomach ulcer                                                                              | 25 (5.7)   | 20 [4.6 (2.6–6.5)]     | 0.5246 |
| 18 | HIV                                                                                        | 18 (4.1)   | 17 [3.9 (2.1–5.7)]     | 0.2330 |
| 19 | Respiratory disease                                                                        | 14 (3.2)   | 12 [2.7 (1.2–4.3)]     | 0.8964 |
| 20 | Diabetes                                                                                   | 11 (2.5)   | 10 [2.3 (0.9–3.7)]     | 0.5506 |
| 21 | <i>On medication</i>                                                                       |            |                        |        |
| 22 | No                                                                                         | 343 (78.3) | 291 [66.4 (62.0–70.9)] | 0.6888 |
| 23 | Yes                                                                                        | 95 (21.7)  | 79 [18.0 (14.4–21.7)]  |        |
| 24 | <i>COVID-19 symptoms (previous 16 months)</i>                                              |            |                        |        |
| 25 | No                                                                                         | 157 (35.8) | 142 [32.4 (28.0–36.8)] | 0.0099 |
| 26 | Yes                                                                                        | 281 (64.2) | 228 [52.1 (47.4–56.8)] |        |
| 27 | <i>Ever tested for SARS-CoV-2</i>                                                          |            |                        |        |
| 28 | No                                                                                         | 371 (84.7) | 311 [71.0 (66.7–75.3)] | 0.3787 |
| 29 | Yes                                                                                        | 67 (15.3)  | 59 [10.3–16.7)]        |        |
| 30 | <i>Previous SARS-CoV-2 test</i>                                                            |            |                        |        |
| 31 | Tested with negative result                                                                | 54 (80.6)  | 48 [71.6 (60.6–82.7)]  | *      |
| 32 | Tested with positive result                                                                | 4 (6.0)    | 4 [6.0 (0.1–11.8)]     |        |
| 33 | Tested with unknown result                                                                 | 9 (13.4)   | 7 [10.4 (2.0–18.0)]    |        |
| 34 | <i>COVID-19 vaccination</i>                                                                |            |                        |        |
| 35 | No                                                                                         | 408 (93.2) | 341 [77.9 (74.0–81.8)] | 0.0561 |
| 36 | Yes                                                                                        | 30 (6.8)   | 29 [6.6 (4.3–9.0)]     |        |

37 \* = no chi square possible due to a negative value in the tested negative category

38

39 Most participants (79.2%) were between age 18 and 50 years; only 4.8% were age ≤17 years and 16%

40

41 were age ≥50 years. Females exceeded males across the 18–50 years age group by a ratio of 2:3, but

42

43 males exceeded females in the <18 years and >50 years age groups. Eleven female participants were

44

45 pregnant; two were uncertain of their pregnancy status. Of the 419 participants age ≥16 years (the

46

47 official age for employment in Kenya [33]) more than one third (39.1%) were unemployed; the greatest

48

49 number of unemployed participants were women in the 18–50 years age group (74.4% of unemployed

50

51

52

53

54

55

56

57

58

59

60

9

For peer review only - <http://bmjopen.bmj.com/site/about/guidelines.xhtml>

participants). Less than one quarter (22.1%) of participants self-reported any type of underlying health condition; the most common was cardiovascular disease, including hypertension (9.1% of all participants); only 14 (3.2%) participants reported underlying respiratory disease (including asthma). Underlying health conditions were most common in females in the 18–50 years age group (43.8% of all participants). Less than one fifth (16%) of participants were taking one or more medications at the time of the survey; these included therapeutic drugs as well as vitamins and symptomatic treatment. Four of the 18 participants with HIV infection reported not taking antiretroviral medication. Only 1 (0.23%) of the 281 (64.2%) participants reporting symptoms suggestive of COVID-19 in the previous 16 months had been hospitalised; he reported a negative SARS-CoV-2 test at that time. Only 19 (63%) of the 30 participants who reported receiving vaccination against SARS-CoV-2 had received two doses. Most participants could not recall exact dates of vaccinations, but most received vaccinations between 1 April and 28 July 2021, within 4 months of the survey; one reported receiving vaccination in March 2021 and one in August 2021.

#### **Serological results in relation to demographic data**

The collection of an adequate volume of blood by fingerprick required lancets of sufficient depth (at least 1.8 mm; we used 2 mm); smaller lancets used initially did not produce an adequate flow of capillary blood for collection of the full 500 µL. Due to inadequate sample collection, there was insufficient sample in 72 participants to conduct the Platelia SARS-CoV-2 Total Ab Assay, the last test to be conducted. Comparison of performance between the three diagnostic tests (Wantai ELISA, Platelia ELISA and the rapid diagnostic test) are described in a separate paper [30]. In summary, the Wantai ELISA showed greater percentage positive results (82.6%) compared with the rapid test (51.8%) and the Platelia ELISA (69.7%); of the rapid test results, 23 were IgM positive, 151 were IgG positive, 53 were both IgM and IgG positive. Specificities of the rapid test and Platelia SARS-CoV-2 Total Ab Assay using the Wantai ELISA as the reference test were high (>90%). There was no significant difference in percentage positive results across participants' age, sex, presence of underlying health conditions, on any type of medication, or those ever tested for SARS-CoV-2; there was a significant difference in percentage positive results according to participants' employment status ( $p=0.0176$ ), COVID-19 symptoms in the previous 16 months ( $p=0.0099$ ), and vaccination status ( $p=0.0561$ ); this last variable was included in the multiple logistic regression analysis because of the  $p$ -value of  $\leq 0.20$ . The previously hospitalised participant tested negative with the rapid test but positive with the two ELISAs. Simple and multiple logistic regression

1 analysis showed that having had COVID-19 symptoms in the previous 16 months was the only significant  
2 independent predictor of seropositivity in this population ( $p=0.0085$ ) (Table 2).

3 **Table 2. Predictors of positivity to SARS-CoV-2 antibodies ( $n=437^*$ )**

| Variables                                     | Simple logistic regression analysis<br>COR <sup>†</sup> (95% CI) | <i>p</i> | Multiple logistic regression analysis<br>AOR <sup>‡</sup> (95% CI) | <i>p</i> |
|-----------------------------------------------|------------------------------------------------------------------|----------|--------------------------------------------------------------------|----------|
| <i>Age</i>                                    |                                                                  |          |                                                                    |          |
| <18                                           | 0.4 (0.1–1.0)                                                    | 0.0853   | 0.4 (0.0–5.0)                                                      | 0.5574   |
| 18–30                                         | 1                                                                |          | 1                                                                  |          |
| 31–50                                         | 1.3 (0.7–2.5)                                                    |          | 1.2 (0.6–2.3)                                                      |          |
| >50                                           | 0.8 (0.4–1.7)                                                    |          | 0.7 (0.3–1.6)                                                      |          |
| <i>Employment status</i>                      |                                                                  |          |                                                                    |          |
| Unemployed                                    | 1                                                                | 0.0320   | 1                                                                  | 0.1739   |
| Self-employed                                 | 0.8 (0.4–1.5)                                                    |          | 0.9 (0.4–1.7)                                                      |          |
| Casual/part-time                              | 3.3 (1.2–8.8)                                                    |          | 3.2 (1.2–8.7)                                                      |          |
| Student                                       | 0.8 (0.4–1.5)                                                    |          | 0.8 (0.3–2.1)                                                      |          |
| Formal/regular                                | 1.8 (0.5–6.5)                                                    |          | 1.8 (0.5–6.6)                                                      |          |
| Not applicable (<16 years)                    | 0.8 (0.3–1.9)                                                    |          | 1.2 (0.1–19.9)                                                     |          |
| <i>COVID-19 symptoms (previous 16 months)</i> |                                                                  |          |                                                                    |          |
| No                                            | 1                                                                | 0.0121   | 1                                                                  | 0.0085   |
| Yes                                           | 0.5 (0.3–0.8)                                                    |          | 0.4 (0.2–0.8)                                                      |          |
| <i>COVID-19 vaccination</i>                   |                                                                  |          |                                                                    |          |
| No                                            | 0.2 (0.0–1.3)                                                    | 0.0893   | 0.2 (0.0–1.3)                                                      | 0.0822   |
| Yes                                           | 1                                                                |          | 1                                                                  |          |

4 \*The one participant with undisclosed employment status was omitted

5 <sup>†</sup>Crude odds ratio; <sup>‡</sup>Adjusted odds ratio

7 **DISCUSSION**

8 This study confirmed the expected high exposure to SARS-CoV-2 (84.5%) in residents of Kibera slum at  
9 the mid-point of the COVID-19 pandemic. Although nearly two-thirds of participants (62.4%) reported  
10 previous COVID-like symptoms, only one had been hospitalised and recovered. There was no significant  
11 association of SARS-CoV-2 infection with age, sex, underlying health conditions, being on any type of  
12 medication, having been previously tested for SARS-CoV-2; significant associations were found with  
13 employment status, COVID-19 vaccination and previous symptoms compatible with COVID-19, but  
14 multiple regression analysis showed an association only with previous symptoms suggestive of COVID-  
15 19. The categories <18 years of age and not applicable for employment status (<16 years) have wide  
16 confidence intervals, indicating uncertainty about the true value of the odds ratio; this may also be  
17 contributed to by the low participant numbers in these categories. In addition, numbers of vaccinated

participants were small. The high exposure in this population could be linked to the fact that a large proportion are casual labourers who walk up to 15 kilometres daily in search of work and mix with populations in other areas; other groupings were also at risk due to the high population density in the Kibera slum resulting in high likelihood of transmission. The overall seroprevalence in our study is similar to that of women who attended antenatal care services in Nairobi (84.5% vs 85%) at a similar time frame (August 2021 vs October 2021) [21], which was conducted at the national referral hospital close to the Kibera slum. However, in our female population, seroprevalence was only 33.3% in the 18–30 years age group, and 34.9% in the 31–50 years age group, implying the population groups were different and suggesting the need for further investigation. The reasons for low rates of severe disease in this infected population are unknown. Analysis of data from an ongoing population-based infectious disease surveillance platform showed that in Kibera during the COVID-19 period observed (March 2020–December 2021) all-cause mortality was slightly lower with no significant change in mortality due to leading specific causes of death [34].

Several explanations have been proposed for the overall low reported morbidity and mortality from COVID-19 in Africa despite high infection rates, including the high infectious disease burden on the African continent, demographic age structure [35,36,37,38,39] and underreporting due to limited testing and lack of access to healthcare services [8]. An integrative review concluded that low COVID-19 mortality and morbidity in Africa was largely a result of the combined effect of the younger African population and underreporting of COVID-19 cases [40], although the protective effect of a younger population has been questioned [8]. In our study, the number of participants age >50 years was low (16%) which may also have contributed to the lower rate of severe disease in this population. Genetic factors may also explain the diversity observed in severity of COVID-19 in African populations due to single nucleotide polymorphisms (SNPs) within the SARS-CoV-2 receptor genes; these have been demonstrated to have both detrimental and protective effects across ethnic groups [41]. Indeed, the population of Kibera represents a variety of the ethnic groups found in the eastern Africa region.

According to the United Nations, over 1 billion slum dwellers worldwide are mostly confined to three regions, one being sub-Saharan Africa [42]. For residents of urban slums, the difficulty in implementing COVID-19 preventive measures due to severe overcrowding was recognised early [43,44] but the predicted high caseload and mortality rates from community transmission were not apparent, especially in Africa's slums [45,46,47]. This study supports seroprevalence studies in urban slums outside Africa that showed high exposure to the virus as well as low morbidity, with differing associated factors.

1 Malani *et al* in a comparative study of slum and non-slum areas in Mumbai, India, showed markedly  
2 higher proportions of positive tests in slum areas (54.1%) than in non-slum areas (16.1%), with lower  
3 infection fatality in slums (0.076%) than in non-slums (0.263%) [48]. Nirala *et al* in a study across 10  
4 different slums in Patna, India, found a seropositivity rate of 31.5% (95% CI: 27.9–35.1) with seropositive  
5 status significantly associated with age 18–30 years, male gender, high-risk occupations (autorickshaw  
6 drivers, rickshaw pullers, street vendors), below poverty line economic status, residing in a hut or kutcha  
7 house (makeshift dwelling) and COVID-like symptoms in the preceding one month [49]. A study by  
8 Raqib *et al* comparing slum and non-slum areas in Bangladesh showed seroprevalence was positively  
9 associated with limited years of formal education (AOR = 1.61; 95% CI = 1.43, 1.82), lower income (AOR  
10 = 1.23; 95% CI = 1.03, 1.46), overweight (AOR = 1.2835; 95% CI = 1.26, 1.97), diabetes (AOR = 1.67; 95%  
11 CI = 1.21, 2.32) and heart disease (AOR = 1.38; 95% CI = 1.03, 1.86) [50]. However, in a study by George  
12 *et al* in a large slum in South India with a reported overall COVID-19 seroprevalence of 57.9% (95% CI  
13 53.4–62.3), age, education, occupation and presence of reported co-morbidities were not significantly  
14 associated with seroprevalence [51]. The discrepancies between high seroprevalence, low morbidity and  
15 varying underlying factors suggest the need for further research to understand the potential protective  
16 factors and immune responses that may be at play in these informal settlements. The informal  
17 community in our study may have distinct characteristics that sets it apart from other slum populations  
18 examined in previous research including genetic variability. Factors such as community resilience, social  
19 support networks and environmental conditions may vary from place to place and could play a role in  
20 mitigating the impact of COVID-19 and influencing the observed lack of associations with reported risk  
21 factors.

22 Co-morbidities associated with severe COVID-19 have also been examined in other populations.  
23 Martono Fatmawati *et al* reviewed nine studies from Asia and Europe that showed risk factors for severe  
24 COVID-19 included age, gender, chronic comorbidities, cardiovascular disease, diabetes, hypertension,  
25 kidney failure, cancer and a history of smoking [52]. Wang *et al* using self-reported data from a survey  
26 of adults in the United States reported that individuals with underlying health conditions were more  
27 likely to become infected with SARS-CoV-2 and have more severe post-infection symptoms [53].  
28 Kompaniyets *et al* in a cross-sectional study of children age ≤18 years in the United States reported  
29 underlying health conditions, such as type 1 diabetes, cardiac and circulatory congenital anomalies and  
30 obesity, were more associated with severe COVID-19 or death [54]. Costa *et al* in a review of published  
31 data up to June 2020 reported patients with metabolic disorders including obesity, diabetes,



cardiovascular disease and liver disease faced a higher risk of SARS-CoV-2 infection and were associated with a significantly worse outcome [55]. Wu *et al* in a review of published papers up to April 2020 reported diabetes increased the mortality of patients with COVID-19 [56]. However, the findings from our study challenge the assumptions that COVID-19 risk factors are necessarily linked to severity of disease. Males have also been reported to be at higher risk of severe illness and increased mortality than females [57,12]; this was also not evident in our study.

Our study also explored the use of small blood volume samples collected through fingerprick. We found fingerprick sampling a convenient and acceptable way of conducting a serosurvey across a large population that included small children, and for evaluating new technologies. Testing kits used for serosurveys should therefore ideally use whole blood or plasma separated from capillary blood. Our study used three tests concurrently to detect antibodies to SARS-CoV-2: one rapid test and two ELISA tests. Since the start of the COVID-19 pandemic, molecular testing for SARS-CoV-2 virus using RT-PCR-based assays performed on respiratory specimens has been central to disease management and control [14]. Tests for SARS-CoV-2 antigens on clinical specimens using immunoassays and rapid tests have also been used but are limited by sub-optimal sensitivity [58,59]. Molecular and antigen tests are constrained by detection of virus to a few days before infection and a short time after infection. In a systematic review of longitudinal studies of RT-PCR test results in symptomatic SARS-CoV-2 infections, the highest percentage of virus detection in nasopharyngeal swabs by PCR was 89% between 0 and 4 days post-symptom onset, dropping to 54% (95% CI 47 to 61) after 10–14 days [60]. Tests for detecting antibodies to SARS-CoV-2 applicable outside research laboratory settings include rapid tests using lateral flow immunology and ELISA tests. In a study by Guo *et al* using an ELISA based on recombinant viral nucleocapsid protein, the median time to IgM and IgA antibody detection after symptom onset was 5 (IQR 3–6) days, while IgG was detected at 14 (IQR 10–18) days [61]. Zhao *et al*, using an ELISA prepared from recombinant antigen containing the receptor binding domain of SARS-CoV-2 spike protein, reported median times from symptom onset to total antibody, IgM and IgG seroconversion were 11, 12 and 14 days, respectively [62]; the presence of antibodies was <40% among patients within 1 week of onset, and rapidly increased to 100% (total antibody – 94.3% (IgM), 79.8% (IgG)) by day 15 after onset; there were also negative antibody findings in 7% patients, possibly due to blood samples not taken at an appropriate time after symptom onset. In a report by Qu *et al* using a IgG/IgM chemiluminescent immunoassay with combined nucleocapsid protein and spike glycoprotein antigens, seroconversion time of IgG antibody was earlier than that of IgM antibody: 97.6% of patients (40/41) were positive with IgG



1 and 87.8% (36/41) with IgM, with a median time of seroconversion for IgG at 11 days (8–16 days) and  
2 for IgM at 14 days (8–28 days) after disease onset. The level of IgG antibody reached the highest  
3 concentration on day 30, while the highest concentration of IgM antibody appeared on day 18 but then  
4 began to decline [63]. Hoffman *et al* found no statistical difference between testing samples taken from  
5 PCR-confirmed COVID-19 cases between 9 and 17 days and 18 and 29 days for either IgM or IgG  
6 seropositivity [64]. Generally, IgM is produced first with a later switch towards IgG production, but  
7 studies on SARS-associated coronaviruses suggest that IgM and IgG often develop at around the same  
8 time [65,66]. In general, detection of antibodies is better in samples taken >14 days after onset of  
9 disease, due to the time taken for antibody development [67,68]. Reported duration of antibody after  
10 natural confirmed infection is variable, including >6 months [69], up to 20 months in unvaccinated adults  
11 in the United States after confirmed COVID-19 infection [70], and more than 2 years in patients in  
12 Wuhan, China [71]. It is therefore probable that most natural infections occurring within the previous 16  
13 months in our study population were detected, although the relevance of the IgM and IgG antibody  
14 findings is unclear.

15 Serologic studies that assess prior infection and immunity to infectious diseases are essential for  
16 epidemiologic studies, ongoing surveillance, vaccine studies and potentially for risk assessments of  
17 healthcare workers, and provide an indication of previous as well as recent exposure in a population.  
18 Surveillance therefore remains fundamental to understanding the evolution of SARS-CoV-2 infection, the  
19 risk factors for severe disease, and the impact of vaccination and public health and social measures [72].  
20 Although hospital admissions and severe outcomes from COVID-19 have substantially decreased since  
21 the start of the pandemic, COVID-19-related deaths remain substantial: in the United States, COVID-19  
22 still ranks as the 10th most common cause of death. The percentage of positive tests for SARS-CoV-2, a  
23 key indicator of community spread, reached peak levels of 12.9% in January 2024 [73]. Since disease  
24 mitigation measures implemented in many countries severely affected the global economy and financial  
25 markets [74], and substantially impacted the inadequately prepared health systems in Africa [75], health  
26 systems need to be vigilant of emerging SARS-CoV-2 variants and the possibility of new surges in cases  
27 and deaths. As WHO recommends countries transition from emergency mode to managing COVID-19  
28 alongside other infectious diseases, providing a more complete picture of the total number of people  
29 infected with SARS-CoV-2 remains an important measure for guiding public health responses [76]. The  
30 recent establishment of a global network to detect and monitor novel coronaviruses of public health  
31 importance will facilitate early detection, risk assessment, and response to coronavirus-related health

challenges [77]. Well-designed surveillance systems remain core to monitoring acute viral respiratory infections to inform public health measures, health system capacities, impact of vaccination programmes and other control measures [78].

## Conclusions

This study reports high SARS-CoV-2 exposure in a slum community in Nairobi, Kenya, with limited morbidity; only recent COVID-19 symptoms were a significant independent predictor of seropositivity. This study confirms other reports of high SARS-CoV-2 exposure with limited morbidity in slum communities. The study also supports the convenient use of small blood volumes for conducting population serosurveys. Use of serological assays to conduct seroprevalence studies will continue to remain important as more robust tests are developed, especially point of care tests. Public health systems would greatly benefit from serosurveillance to supplement and strengthen existing COVID-19 and other case-based infectious disease surveillance strategies [79].

## Study limitations

This study was conducted during the working week during working hours, which may explain the low number of male participants of working age and low number of those formally employed. The tests used were limited by availability of approved tests at the time.

## Author contributions

Conceptualisation: JYC, SK, SM, JM, JO; Methodology: JYC, SK, SM, SMM, LK, JK, NL, JC, RO, AM; Analysis: JYC, RS, MH; writing—original draft preparation: JYC, RS, MH; writing—review and editing: JYC, SK, JM, SM, SMM, RS, MH, MP, JS, JO. All authors have read and agreed to the published version of the manuscript. JYC is the guarantor.

## Patient and public involvement

Patients or the public were not involved in the design, reporting or dissemination plans of our research. Entry to individual households was facilitated through community health volunteers who were formally assigned to these villages.

## Ethics statement

Approval for the study was obtained from the Amref Health Africa Ethics and Scientific Review Committee (ESRC), the National Commission for Science, Technology and Innovation (NACOSTI), and the Nairobi Metropolitan Services, Ministry of Health. Verbal permission to conduct the study was obtained from the five area chiefs administering the study villages.

## Competing interests

1 All authors declare they have no competing interests.

2 **Funding**

3 The study was funded by American Tower Corporation, USA. The funder had no role in the study design,  
4 data collection and analysis, decision to publish, or preparation of the manuscript.

5 **Data availability**

6 Data are available on reasonable request.

7 **REFERENCES**

8 1. Zhu H, Wei L, Niu P. The novel coronavirus outbreak in Wuhan, China. *Glob Health Res and Policy*  
9 2020;**5**(6). Available <https://doi.org/10.1186/s41256-020-00135-6>

10 2. Cucinotta D, Vanelli M. WHO declares COVID-19 a pandemic *Acta Biomed* 2020;**91**(1)157–160. doi:  
11 <https://doi.org/10.23750/abm.v91i1.9397>

12 3. World Health Organization. WHO Director-General's opening remarks at the media briefing. 5 May  
13 2023. Available <https://www.who.int/news-room/speeches/item/who-director-general-s-opening-remarks-at-the-media-briefing---5-may-2023>

14 4. World Health Organization. Health Emergencies Programme. WHO COVID-19 dashboard. Available  
15 <https://data.who.int/dashboards/covid19/cases?n=c>

16 5. Mwananyanda L, Gill CJ, MacLeod W, *et al.* Covid-19 deaths in Africa: prospective systematic  
17 postmortem surveillance study. *BMJ* 2021;**372**:n334. doi: <http://dx.doi.org/10.1136/bmj.n334>

18 6. World Health Organization Regional Office for Africa. Six in seven COVID-19 infections go  
19 undetected in Africa. October 2021. Available <https://www.afro.who.int/news/six-seven-covid-19-infections-go-undetected-africa>

20 7. Ngere I, Dawa J, Hunsperger E, *et al.* High seroprevalence of SARS-CoV-2 but low infection fatality  
21 ratio eight months after introduction in Nairobi, Kenya. *Int J Infect Dis* 2021;**112**:25–34. doi:  
22 [10.1016/j.ijid.2021.08.062](https://doi.org/10.1016/j.ijid.2021.08.062). Epub 2021 Sep 2. PMID: 34481966; PMCID: PMC8411609

23 8. McKay T, Robinson RS, Musungu S, *et al.* The missing millions: uncovering the burden of Covid-19  
24 cases and deaths in the African region. *Pop Dev Review* 2024;**50**:7–58.  
25 doi: <https://doi.org/10.1111/padr.12608>

26 9. Meyerowitz EA, Richterman A, Gandhi RT, *et al.* Transmission of SARS-CoV-2: a review of viral host,  
27 and environmental factors. *Ann Intern Med* 2020;**174**(1):69–79. doi: [10.7326/M20-5008](https://doi.org/10.7326/M20-5008).  
28 PMCID: PMC7505025. PMID: [32941052](https://pubmed.ncbi.nlm.nih.gov/32941052/)

10. Nishiura H, Kobayashi T, Miyama T, *et al.* Estimation of the asymptomatic ratio of novel coronavirus infections (COVID-19). *Int J Infect Dis* 2020;**94**:154–155. doi: <https://doi.org/10.1016/j.ijid.2020.03.020>
11. Ogalo EA, Gudu E, Andale T, Korir D, Ndege S, Simiyu T, Olekuyo R, Mwangi H, Kimaiyo S, Aruasa W. Clinical spectrum of COVID-19 at a national referral hospital in western Kenya during the period 2020–2021. *Front Virol* 2023;**3**:1202742. doi: 10.3389/fviro.2023.1202742
12. Johansson MA, Quandelacy TM, Kada S, *et al.* SARS-CoV-2 transmission from people without COVID-19 symptoms. *JAMA Network Open* 2021;**4**(1):e2035057. doi:10.1001/jamanetworkopen.2020.35057
13. Oliveira MC, Scharan KO, Thomés BI, *et al.* Diagnostic accuracy of a set of clinical and radiological criteria for screening of COVID-19 using RT-PCR as the reference standard. *BMC Pulm Med* 2023;**23**:81 <https://doi.org/10.1186/s12890-023-02369-9>
14. Cheng MP, Yansouni CP, Basta NE, *et al.* Serodiagnostics for severe acute respiratory syndrome-related coronavirus-2: a narrative review. *Annals of Internal Medicine* 2020;**173**:6. <https://doi.org/10.7326/M20-2854>
15. Aluga, MA. Coronavirus disease 2019 (COVID-19) in Kenya: preparedness, response and transmissibility. *J Microbiol Immunol Infect* 2020;**53**(5):671–673. doi: <https://doi.org/10.1016/J.JMII.2020.04.011>
16. World Bank. Socioeconomic impacts of COVID-19 in Kenya. 2021. Available <https://documents1.worldbank.org/curated/en/949721626096781344/pdf/Socioeconomic-Impacts-of-COVID-19-in-Kenya.pdf>
17. Ministry of Health, Republic of Kenya. National COVID-19 Vaccine Deployment Plan. August 2021.
18. Ministry of Health, Republic of Kenya. Kenya COVID-19 Vaccination Program: Daily Situation Report. 5 February 2022.
19. Uyoga S, Adetifa IMO, Otiende M, *et al.* Seroprevalence of anti–SARS-CoV-2 IgG antibodies in Kenyan blood donors. *Science* 2021;**371**(6524):79–82. doi:10.1126/science.abe1916. Epub 2020 Nov 11.
20. Uyoga S, Adetifa IMO, Otiende M, *et al.* Prevalence of SARS-CoV-2 antibodies from a national serosurveillance of Kenyan blood donors, January–March 2021. *JAMA* 2021;**326**(14):1436–1438. doi: 10.1001/jama.2021.15265

1 1 21. Lucinde R, Mugo D, Bottomley C, *et al.* Serosurveillance for IgG to SARS-CoV-2 at antenatal care  
2 2 clinics in three Kenyan referral hospitals: repeated cross-sectional surveys 2020–21. *PLoS ONE*  
3 3 2022;17(10):e0265478. doi: <https://doi.org/10.1371/journal.pone.0265478>  
4 4 22. Etyang AO, Lucinde R, Karanja H, *et al.* Seroprevalence of antibodies to severe acute respiratory  
5 5 syndrome coronavirus 2 among healthcare workers in Kenya. *Clin Infect Dis* 2021;74(2):288–293.  
6 6 <https://doi.org/10.1093/cid/ciab346>  
7 7 23. Kagucia EW, Gitonga JN, Kalu C, *et al.* Seroprevalence of anti-SARS-CoV-2 IgG antibodies among  
8 8 truck drivers and assistants in Kenya. Book Review. *Open Forum Infect Dis* 2021;8(7):1–4. doi:  
9 9 <https://doi.org/10.1101/2021.02.12.21251294>  
10 10 24. Crowell TA, Daud II, Maswai J, *et al.* 2021. SARS-CoV-2 antibody prevalence in people with and  
11 11 without HIV in rural western Kenya, January to March 2020. *AIDS* 35(14):2401–2404.  
12 12 doi:10.1097/QAD.0000000000003054  
13 13 25. Nasimiya C, Matoke-Muhia D, Rono GK, *et al.* 2022. Imported SARS-CoV-2 Variants of Concern  
14 14 Drove Spread of Infections across Kenya during the Second Year of the Pandemic. *COVID* 2022, 2,  
15 15 586–598. <https://doi.org/10.3390/covid2050044>  
16 16 26. Kenya National Bureau of Statistics. The 2009 Kenya Population and Housing Census. August 2010.  
17 17 27. Mutisya E, Yarime M. Understanding the grassroots dynamics of slums in Nairobi: the dilemma of  
18 18 Kibera informal settlements [Internet]. Vol. 2, *Int Transact J Eng Manage Appl Sci Technol*. 2011  
19 19 [Accessed 2020 May 18]. Available: <http://tuengr.com/V02/197-213.pdf>  
20 20 28. UN Habitat. COVID-19 exposes the harsh realities of gender inequality in slums. 2020.  
21 21 <https://unhabitat.org/news/03-jun-2020/covid-19-exposes-the-harsh-realities-of-gender-inequality-in-slums#:~:text=In%20Kibera%2C%20Kenya%2C%20the%20world's,women%20for%20every%20100%20men>  
22 22 29. Statistics How To. Sample size in Statistics (how to find it): Excel, Cochran’s formula, general tips.  
23 23 <https://www.statisticshowto.com/probability-and-statistics/find-sample-size/>  
24 24 30. Carter JY, Khamadi S, Mwangi J, *et al.* 2023. Performance of one rapid test and two ELISAs for a  
25 25 SARS-CoV-2 seroprevalence survey in Kibera informal settlement, Nairobi, Kenya. Manuscript  
26 26 submitted for publication.  
27 27 31. World Health Organization. COVID-19: Case definitions. 16 December 2020.  
28 28 32. Ministry of Health, Kenya. Interim guidelines on management of COVID-19 in Kenya. COVID-19,  
29 29 Infection Prevention and Control (IPC) and Case Management. June 2020.  
30 30  
31 31



33. Laws of Kenya. Employment Act. Chapter 226. Revised Edition 2012 [2007].
34. Oduor C, Audi A, Kiplangat S, *et al.* Estimating excess mortality during the COVID-19 pandemic from a population-based infectious disease surveillance in two diverse populations in Kenya, March 2020–December 2021. *PLOS Glob Public Health* 2023;**3**(8):e0002141.  
<https://doi.org/10.1371/journal.pgph.0002141>
35. Wamai RG, Hirsch JL, Van Damme W, *et al.* What could explain the lower COVID-19 burden in Africa despite considerable circulation of the SARS-CoV-2 virus? *Int J Environ Res Public Health* 2021;**18**(16):8638. doi: <https://doi.org/10.3390/ijerph18168638>
36. Musa HH, Musa TH, Musa IH, *et al.* Addressing Africa's pandemic puzzle: perspectives on COVID-19 transmission and mortality in sub-Saharan Africa. *Int J Inf Diseases* 2021;**102**:483–488. doi: [10.1016/j.ijid.2020.09.1456](https://doi.org/10.1016/j.ijid.2020.09.1456). PMID: 33010461. PMCID: [PMC7526606](https://doi.org/10.1016/j.ijid.2020.09.1456)
37. Adams J, MacKenzie MJ, Amegah AK, *et al.* The conundrum of low COVID-19 mortality burden in sub-Saharan Africa: myth or reality? *Glob Health Sci Pract* 2021;**9**(3):433–443. doi: <https://doi.org/10.9745/GHSP-D-21-00172>
38. Tessema SK, Nkengasong JN. Understanding COVID-19 in Africa. *Nat Rev Immunol* 2021;**21**:469–470. <https://doi.org/10.1038/s41577-021-00579-y>
39. Kusi KA, Frimpong A, Partey FD, *et al.* High infectious disease burden as a basis for the observed high frequency of asymptomatic SARS-CoV-2 infections in sub-Saharan Africa [version 3; peer review: 2 approved] *AAS Open Research* 2021;**4**:2. doi: <https://doi.org/10.12688/aasopenres.13196.3>
40. Manna OK, Costa Clemens SA, Clemens R. Investigating the Possible Reasons for the Low Reported Morbidity and Mortality of COVID-19 in African Countries: An Integrative Review. *The Pediatric Infectious Disease Journal* 42(7):p e222–e228, July 2023. | DOI: 10.1097/INF.0000000000003916
41. Adimulam, T.; Arumugam, T.; Gokul, A.; Ramsuran, V. Genetic Variants within SARS-CoV-2 Human Receptor Genes May Contribute to Variable Disease Outcomes in Different Ethnicities. *Int. J. Mol. Sci.* 2023, **24**, 8711. <https://doi.org/10.3390/ijms24108711>
42. United Nations. New York: United Nations Publication; 2016. [2020-04-23]. Urbanization and development: emerging futures. World Cities report. <https://tinyurl.com/y4ekncmh>.
43. Gibson L, Rush D. Novel coronavirus in Cape Town informal settlements: feasibility of using informal dwelling outlines to identify high risk areas for COVID-19 transmission from a social



1 distancing perspective. *JMIR Public Health Surveill* 2020;**6**(2):e18844. Available:  
2 <http://publichealth.jmir.org/2020/2/e18844/> doi: 10.2196/18844. PMID: 32250283  
3  
4 44. Wasdania KP, Prasada A. The impossibility of social distancing among the urban poor: the case of  
5 an Indian slum in the times of COVID-19. *Local Environment* 25(5):414–418.  
6  
7 <https://doi.org/10.1080/13549839.2020.1754375>  
8  
9 45. UN Habitat. Coronavirus will travel 'incredibly fast' in Africa's slums. April 2020. Available:  
10 [https://unhabitat.org/coronavirus-will-travel-incredibly-fast-in-africas-slums-un-habitat-chief-](https://unhabitat.org/coronavirus-will-travel-incredibly-fast-in-africas-slums-un-habitat-chief-warns)  
11 [warns](https://unhabitat.org/coronavirus-will-travel-incredibly-fast-in-africas-slums-un-habitat-chief-warns)  
12  
13 46. Arora RK, Joseph A, Van Wyk J, *et al.* SeroTracker: a global SARS-CoV-2 seroprevalence dashboard.  
14 *Lancet Infect Dis* 2021;**(4)**:E75–76. doi: 10.1016/S1473-3099(20)30858-6  
15  
16 47. Ilesanmi O, Oderinde T, Afolabi A. The urban slums: potential source of COVID-19 spikes in Africa.  
17 *Public Health Pract (Oxf)* 2020; Nov:1:100052. doi: 1:10.1016/j.puhip.2020.100052.  
18  
19 48. Malani A, Shah D, Kang G, *et al.* Seroprevalence of SARS-CoV-2 in slums versus non-slums in  
20 Mumbai, India. *The Lancet Global Health* 2021;**9**(2)February E110-111. doi:  
21 [https://doi.org/10.1016/S2214-109X\(20\)30467-8](https://doi.org/10.1016/S2214-109X(20)30467-8)  
22  
23 49. Nirala SK, Naik BN, Chaudhary N, *et al.* Sero-epidemiological survey of SARS-CoV2 in urban slums of  
24 a capital city: a cross- sectional study. *J Family Med Prim Care* 2022;**11**(6):2709–16. doi:  
25 10.4103/jfmpc.jfmpc\_2127\_21  
26  
27 50. Raqib R, Sarker P, Akhtar E, *et al.* Seroprevalence of SARS-CoV-2 infection and associated factors  
28 among Bangladeshi slum and non-slum dwellers in pre-COVID-19 vaccination era: October 2020 to  
29 February 2021. *PLoS ONE* 2022;**17**(5):e0268093. doi:  
30 <https://doi.org/10.1371/journal.pone.0268093>  
31  
32 51. George CE, Inbaraj LR, Chandrasingh S, *et al.* High seroprevalence of COVID-19 infection in a large  
33 slum in South India; what does it tell us about managing a pandemic and beyond? *Epid Inf*  
34 2021;**149**:e39,1–6. doi: <https://doi.org/10.1017/S0950268821000273>  
35  
36 52. Martono, Fatmawati F, Mulyanti S. Risk factors associated with the severity of COVID-19. *Malays J*  
37 *Med Sci* 2023;**30**(3):84–92. doi:10.21315/mjms2023.30.3.7. Epub 2023 Jun 27. PMID: 37425387;  
38 PMCID: PMC10325129.  
39  
40 53. Wang B, Yuan S, Ruan S, *et al.* Associations between underlying diseases with COVID-19 and its  
41 symptoms among adults: a cross-sectional study. *Front Public Health* 2023;**11**:1210800. doi:  
42 10.3389/fpubh.2023.1210800  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

54. Kompaniyets L, Agathis NT, Nelson JM, *et al.* Underlying medical conditions associated with severe COVID-19 illness among children. *JAMA Netw Open* 2021;**4**(6):e2111182. doi:10.1001/jamanetworkopen.2021.11182. PMID: 34097050. PMCID: PMC8185607.
55. Costa FF, Rosário WR, Farias ACR, *et al.* Metabolic syndrome and COVID-19: an update on the associated comorbidities and proposed therapies. *Diabetes Metab Syndr: Clinical Research & Reviews* 2020;**14**(5):809–814. doi: <https://doi.org/10.1016/j.dsx.2020.06.016>
56. Wu ZH, Tang Y, Cheng Q. Diabetes increases the mortality of patients with COVID-19: a meta-analysis. *Acta Diabetol* 2021;**58**:139–144. doi: <https://doi.org/10.1007/s00592-020-01546-0>. PMID: 32583078; PMCID: PMC7311595.
57. Jin JM, Bai P, He W, *et al.* Gender differences in patients with COVID-19: focus on severity and mortality. *Front Public Health* 2020;**8**:152. doi: 10.3389/fpubh.2020.00152. eCollection 2020.
58. Adnan N, Khandker SS, Haq A, *et al.* Detection of SARS-CoV-2 by antigen ELISA test is highly swayed by viral load and sample storage condition. *Expert Rev Anti Infect Ther* 2022;**20**(3):473–481. doi: 10.1080/14787210.2021.1976144. Epub 2021 Sep 11. PMID: 34477019; PMCID: PMC8442762.
59. Høeg TB, Prasad V. Rapid antigen testing for COVID-19: decreasing diagnostic reliability, potential detrimental effects and a lack of evidence to support continued public funding of community-based testing. *Public Health Pract (Oxf)* 2023;**6**:100451. doi: 10.1016/j.puhip.2023.100451. PMID: 38075468; PMCID: PMC10701347.
60. Mallett S, Allen AJ, Graziadio S, *et al.* At what times during infection is SARS-CoV-2 detectable and no longer detectable using RT-PCR-based tests? A systematic review of individual participant data. *BMC Medicine* 2020;**18**:346. doi: <https://doi.org/10.1186/s12916-020-01810-8>
61. Guo L, Ren L, Yang S, *et al.* Profiling early humoral response to diagnose novel coronavirus disease (COVID-19). *Clin Infect Dis* 2020;**71**(15):778–785. <https://doi.org/10.1093/cid/ciaa310>
62. Zhao J, Yuan Q, Wang H, *et al.* Antibody responses to SARS-CoV-2 in patients with novel coronavirus disease 2019. *Clin Infect Dis* 2020;**71**(16):2027–34. doi: 10.1093/cid/ciaa344.
63. Qu J, Wu C, Li X, *et al.* Profile of Immunoglobulin G and IgM antibodies against severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). *Clin Infect Dis* 2020;**71**(16):2255–58. doi: 10.1093/cid/ciaa489.
64. Hoffman T, Nissen K, Krambrich J, *et al.* Evaluation of a COVID-19 IgM and IgG rapid test: an efficient tool for assessment of past exposure to SARS-CoV-2. *Infect Ecol Epidemiol*

2020;**10**(1):1754538. doi:10.1080/20008686.2020.1754538. PMID: 32363011. PMCID: PMC7178815.

65. Hsueh PR, Huang LM, Chen PJ, *et al.* Chronological evolution of IgM, IgA, IgG and neutralisation antibodies after infection with SARS-associated coronavirus. *Clin Microbiol Infect* 2004;**10**(12):1062–6. doi:10.1111/j.1469-0691.2004.01009.x. PMID: 15606632. PMCID: PMC7129952.

66. Li G, Chen X, Xu A. Letter to the Editor. Profile of specific antibodies to the SARS-associated coronavirus. *N Engl J Med* 2003;349(5):508–509. doi: [10.1056/NEJM200307313490520](https://doi.org/10.1056/NEJM200307313490520)

67. Hanssen DAT, Slaats M, Mulder M, *et al.* Evaluation of 18 commercial serological assays for the detection of antibodies against SARS-CoV-2 in paired serum samples. *Eur J Clin Microbiol Infect Dis* 2021;**40**(8):1695-1703. doi:10.1007/s10096-021-04220-7. Epub 2021 Mar 17. PMID: 33733395; PMCID: PMC7968571.

68. Gong F, Wei H, Li Q, *et al.* Evaluation and comparison of serological methods of COVID-19 diagnosis. *Front Mol Biosci* 2021;**8** Sec. Structural Biology. doi: <https://doi.org/10.3389/fmolb.2021.682405>

69. Dan JM, Mateus J, Kato Y, *et al.* Immunological memory to SARS-CoV-2 assessed for up to 8 months after infection. *Science* 2021;371(6529):eabf4063. doi: 10.1126/science.abf4063

70. Alejo JL, Mitchell J, Chang A, *et al.* Prevalence and durability of SARS-CoV-2 antibodies among unvaccinated US adults by history of COVID-19. *Research Letter JAMA* 2022;327(11):1085–1087. doi:10.1001/jama.2022.1393

71. Guo L, Zhang Q, Gu X, *et al.* Durability and cross-reactive immune memory to SARS-CoV-2 in individuals 2 years after recovery from COVID-19: a longitudinal cohort study. *Lancet Microbe* 2024;**5**(1):E24–E33. doi: [https://doi.org/10.1016/S2666-5247\(23\)00255-0](https://doi.org/10.1016/S2666-5247(23)00255-0)

72. World Health Organization. 2023. Update to requirements for reporting COVID-19 surveillance data under the International Health Regulations (IHR 2005): Addendum to Public health surveillance for COVID-19 interim guidance, 25 August 2023. <https://www.who.int/publications/i/item/WHO-2019-nCoV-Surveillance-Guidance-Addendum-2023-1>

73. National Centre for Immunization and Respiratory Diseases. 2024. The changing threat of COVID-19. 23 February 2024. <https://www.cdc.gov/ncird/whats-new/changing-threat-covid-19.html>

74. Pak A, Adegboye OA, Adekunle AI, *et al.* Economic consequences of the COVID-19 outbreak: the need for epidemic preparedness. *Front Public Health* 2020;**8**:241. doi: 10.3389/fpubh.2020.00241

75. Tessema GA, Kinfu Y, Dachew BA, *et al.* The COVID-19 pandemic and healthcare systems in Africa: a scoping review of preparedness, impact and response. *BMJ Global Health* 2021;**6**(12):e007179. doi:10.1136/bmjgh-2021-007179
76. World Health Organization. Media Briefing. 5 May 2023. Available: <https://www.who.int/director-general/speeches/detail/who-director-general-s-opening-remarks-at-the-media-briefing---5-may-2023>
77. World Health Organization. 2024. WHO launches CoViNet: a global network for coronaviruses. <https://www.who.int/news/item/27-03-2024-who-launches-covinet--a-global-network-for-coronaviruses>
78. European Centre for Disease Prevention and Control. <https://www.ecdc.europa.eu/en/infectious-disease-topics/z-disease-list/covid-19/facts/surveillance-covid-19>
79. Haselbeck AH, Im J, Prifti K, *et al.* Serology as a tool to assess infectious disease landscapes and guide public health policy. *Pathogens* 2022;**11**(7):732. doi: <https://doi.org/10.3390/pathogens11070732>