Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Original Research

The 1st year of the COVID-19 epidemic in Estonia: a population-based nationwide sequential/consecutive cross-sectional study

A. Uusküla a,*, R. Kalda a, M. Solvak b, M. Jürisson a, M. Käärik c, K. Fischer c, A. Keis a, U. Raudvere d, J. Vilo d, H. Peterson d, E. Käärik c, M. Metspalu c, T. Jürgenson c,e, L. Milani e, L. Kolberg d, E.-M. Tiit c, K. Vassil f

a Institute of Family Medicine and Public Health, University of Tartu, Estonia
b Johan Skytte Institute of Political Studies, University of Tartu, Estonia
c Institute of Mathematics and Statistics, University of Tartu, Estonia
d Institute of Computer Science, University of Tartu, Estonia
e Institute of Genomics, University of Tartu, Estonia
f Vice-Rector, University of Tartu, Estonia

INTRODUCTION

During the 1st year of COVID-19 pandemic, control measures (non-pharmaceutical interventions, including business and school closures, restrictions on movement, total lockdowns, social distancing) were widely implemented to contain the spread of SARS-CoV-2 and have been effective in curbing the COVID-19 epidemic, but they do not represent desirable long-term strategies. The future trajectory of the COVID-19 pandemic hinges on the dynamics of both viral evolution and population immunity against SARS-CoV-2.

Understanding the future trajectory of this disease requires knowledge of the population-level landscape of immunity, generated by the life histories of the SARS-CoV-2 infection or vaccination among individual hosts. The drivers of future COVID-19 dynamics are complex. However, characterisation of the prevaccination
prevalence, the change of active infections and development of immunity in the population are vital data elements for adequately projecting the future course of the SARS-CoV-2 epidemic and the effect of containment measures.

In Estonia, as of January 31, 2021, 785,333 SARS-CoV-2 (RNA) tests (58,967 per 100,000 population) were undertaken, and a total of 44,208 (3326 per 100,000) COVID-19 cases were confirmed (Fig. 1). The confirmed SARS-CoV-2 case rate was the highest among people aged 15–24 years (4120/100,000), followed by the age group 45–54 years (4053/100,000), and lowest among children younger than 10 years (522 and 1279 per 100,000 among 0- to 4-year olds and 5- to 9-year olds, respectively). By January 31, 2021, of all the confirmed COVID-19 cases, 5.6% (n = 2471) had been hospitalised for treatment and 0.9% (n = 419) had died.

The first case of COVID-19 was confirmed in Estonia on February 26, 2020. A special digital referral system was developed in mid-March 2020 to simplify the referral process. Individuals who were deemed to be at a high risk for the SARS-CoV-2 infection (symptomatic patients referred by family physicians) and frontline staff members (health care, nursing home, social workers, police, border guard officers with a referral letter from their employer) were all eligible for testing. Testing eligibility was relaxed by July 2020.

On March 13, 2020, a set of lockdown rules was implemented—people were allowed to leave their homes at any time so long as they observed social distancing. By June 2020, the restrictions were gradually eased, but physical distancing requirements, that is, the 2 + 2 rule (up to two people can be in a public place together and at least a 2-m distance must be kept from others’), have remained in force. In response to the increase of new case notifications since the last week of July 2020, and attributing the new cases to visiting nightclubs and bars, the Police and Border Guard Board imposed bans on night-time alcohol sales from August 7 (in two counties), and since September 25, a nationwide restriction on the sale of alcohol has been in force. Since the beginning of November 2020, additional measures on the workplace (recommendation to work remotely and cancelling all joint events), in public places and in transport (mandatory mask wearing) were implemented. COVID-19 vaccination started in January 2021.

The evidence of the first year of the COVID-19 epidemic is frequently based on the data from symptomatic patients, seroepidemiological studies and modelling. Most studies are based on small or selected population samples (e.g., hospital admissions) providing data not representative of the community. To the best of our knowledge, large population-based studies needed

Fig. 1. The COVID-19 epidemic in Estonia: daily numbers of new confirmed cases, the number of tests, proportion of positive tests and the number of deaths, 2020–2021.
to understand risk factors and dynamics and delineate the pre-vaccination course of the COVID-19 pandemic are scarce.\(^{16,17}\)

In this study, we rely on a national survey designed to be representative of the target population to describe the course of the epidemic over the first year and risk factors for testing positive for SARS-CoV-2 in Estonia (until the end of January 2021).

### Methods

#### Study design

A population-based nationwide sequential/consecutive cross-sectional study was conducted.

#### Source population

In 2020, the population of Estonia was estimated at 1,326,535 million people (equivalent to 0.02% of the total world population), with 68% of the population living in urban areas. The Estonian language is spoken by roughly 68% of the population, with approximately 28% of the population being Russian speakers.\(^1\) Historically, most of Russians-speaking population is settled in the capital, Tallinn, or the northeastern region of the country (Ida-Virumaa County)\(^{18,19}\).

**Data source: SARS-CoV-2 community prevalence studies**

The data for this work originate from sequential/consecutive nationwide cross-sectional studies. This methodology was chosen on the premise that valid inferences of change in population values can be made on the basis of repeated cross sections within the single population.\(^1\)

The listing of the Estonian Population Registry\(^2\) was used as a sampling frame, and all individuals aged 18 years and older were eligible for study participation.

Using standardised methodology (population-based, random stratified sampling), 11 cross-sectional studies were conducted with data collection during April 23–29, April 30 – May 6, May 22–31, June 11–22, August 6–25, September 21 – October 3, November 11–19, November 26 – December 6 and December 11–20 in 2020 and during January 7–18 and January 21 – February 2 in 2021. For each study, multistage stratified random sampling was used. Primary sampling strata consisted of all counties (\(n=15\)), and two most populated cities were considered separately from their respective counties. In each primary sampling stratum, stratified by gender and age (18–39, 40–64 and 65+ years), random samples (\(n=200\) in most regions, \(n=400\) in the three most populated areas) of civilian residents were recruited.

**Sample size**

The required total sample size for individual SARS-CoV-2 RNA testing studies was estimated based on the upper Clopper-Pearson confidence limit under the assumption of no positive test results. The sample size of 2000 was derived at a 5% level of significance with an upper confidence limit of 0.184%.

**Study procedures**

Participants were contacted by e-mail (original invitation and up to two reminders) or telephone (for those aged 65 years or older) for completion of a screening questionnaire regarding previous SARS-CoV-2 testing and symptoms of COVID-19. Respondents could take a phone interview in case of any problems with accessing the web questionnaire. A structured questionnaire (based on the instrument recommended by World Health Organization\(^21\)) was used to elicit respondent sociodemographic data, data on the size and age structure of the household, health status and social- and work-related contacts within two weeks before the study.

Referral and registration for SARS-CoV-2 testing at state drive-in sites or home visits by the testing station team (for those study participants unable to access drive-in stations) was undertaken by the study team.

**SARS-CoV-2 testing**

The nasopharyngeal samples collected were tested for SARS-CoV-2 RNA by quantitative reverse-transcriptase–polymerase-chain-reaction (RT-PCR) at the SYNLAB Laboratory, a private medical laboratory company (SolGent DiaPlexQT Novel Coronavirus (2019-nCoV) Detection Kit CE-IVD). Viral RNA from all samples was isolated within 24 h.

All SARS-CoV-2 test results were entered into the state E-Health service system and communicated back to participants by the authorised staff member of the testing stations. Participants who tested positive for SARS-CoV-2 were required to self-isolate for 14 days since developing symptoms. All those who tested positive were monitored by their own family doctor until recovery.

**Statistical analysis**

Descriptive statistics (i.e., proportions and means) are presented. SARS-CoV-2 prevalence (the proportion of testing positive) and 95% Clopper-Pearson confidence interval (CI) were calculated, taking into account the sample design. Prevalence rates were calculated using the Estonian population at the beginning of 2020 as a denominator.\(^1\)

A survey-adjusted logistic regression model was applied to explore associations between data collection timing (study round), age, gender, preferred language, region of residence, size and age structure of the household, pre-existing physician diagnosed chronic conditions, body mass index, number of contacts within two weeks before the study and having COVID-19—specific symptoms at the time of study with the SARS-CoV-2 RNA test positivity. Variables identified as statistically significant predictors with a significance level of \(P < 0.05\) were inserted into a multivariable logistic model.

We present adjusted odds ratios (ORs) together with the 95% confident estimates. Since the observed prevalence is relatively low (<3%),\(^24\) the ORs found in the logistic regression model approximate the risk ratios reasonably well.

We used the R statistical programming language for the analyses.\(^23\)

The study is registered with the ISRCTN Registry, ISRCTN10182320.

### Results

**SARS-CoV-2 community prevalence over the first year of the epidemic**

A total of 34,915 individuals, including 15,203 males and 19,712 females, participated in the series of cross-sectional studies from April 2020 to February 2021. The age of the study participants ranged from 18 to 96 years (average age = 48.1 years); 85.5% filled the survey in Estonian and 14.2% in Russian language. The average household size among the study participants was 2.7. SARS-CoV-2 prevalence declined at the beginning of the observation period (in April: 0.27%, 95% CI = 0.10%–0.59%; June 2020: 0.00%; 95% CI = 0.00%–0.12%)) and remained low until the end of September 2020.
## Table 1
Characteristics of the population-based SARS-CoV-2 prevalence studies and respective study participants, Estonia, 2020–2021.

| Study characteristics | Round 1 | Round 2 | Round 3 | Round 4 | Round 5 | Round 6 | Round 7 | Round 8 | Round 9 | Round 10 | Round 11 |
|-----------------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|----------|----------|
|                       | April 23–29, 2020 | April 30 – May 6, 2020 | May 22–31, 2020 | June 11–22, 2020 | Aug 6–25, 2020 | Sept 21–Oct 3, 2020 | Nov 11–19, 2020 | Nov 26–Dec 6, 2020 | Dec 11–20, 2020 | Jan 7–18, 2021 | Jan 21– Feb 2, 2021 |
| Total sample          | 10,209 | 12,020 | 21,830 | 28,034 | 25,998 | 22,900 | 23,187 | 20,032 | 23,921 | 21,063 | 25,135 |
| Non-contacts (n)      | 4119   | 6113   | 12,869 | 20,133 | 19,467 | 15,460 | 16,322 | 14,296 | 17,900 | 14,957 | 18,042 |
| Refusals (n)          | 2060   | 1791   | 2923   | 2414   | 3024   | 2623   | 2211   | 2294   | 2583   | 2816   | 2816   |
| Other non-response (n)| 1141   | 981    | 2538   | 1813   | 983    | 893    | 688    | 749    | 732    | 1138   | 1138   |
| Participants (n)      | 2889   | 3135   | 3500   | 3872   | 3172   | 3433   | 3349   | 2837   | 2978   | 2999   | 2999   |
| SARS-CoV-2 tested (n) | 2306   | 2666   | 2579   | 2983   | 2335   | 2532   | 2726   | 2381   | 2522   | 2370   | 2470   |
| Study characteristics |         |         |         |         |         |         |         |         |         |         |         |
| Participants characteristics |     |     |     |     |     |     |     |     |     |     |     |
| Men (n, %)             | 1254, 43.4% | 1377, 43.9% | 1627, 46.5% | 1657, 42.8% | 1504, 43.8% | 1388, 41.5% | 1345, 40.2% | 1297, 43.6% | 1202, 43.1% | 1295, 43.8% | 1295, 43.8% |
| Age (mean, SD, range)  | 47.7, 15.8, 18-94 | 46.7, 15.6, 18-94 | 49.6, 16.7, 18-93 | 47.5, 15.6, 18-92 | 48.2, 15.8, 18-96 | 47.2, 15.9, 18-99 | 48.7, 15.9, 18-95 | 49.9, 16.2, 18-94 | 47.0, 15.8, 18-91 | 48.1, 16.1, 18-93 | 47.0, 15.8, 18-91 |
| Size of the household (mean, SD) | 2.77, 1.42 | 2.79, 1.42 | 2.70, 1.41 | 2.75, 1.41 | 2.77, 1.43 | 2.68, 1.36 | 2.62, 1.36 | 2.68, 1.37 | 2.66, 1.36 | 2.67, 1.38 | 2.67, 1.38 |
| Respondent language    | 396, 13.7% | 432, 13.8% | 513, 14.7% | 554, 14.3% | 380, 12.0% | 482, 14.0% | 565, 16.9% | 411, 14.5% | 388, 13.0% | 406, 14.6% | 428, 14.5% |
| Smoking (yes; n, %)    | 685, 23.7% | 709, 22.6% | 762, 21.8% | 789, 20.4% | 706, 22.3% | 681, 19.9% | 652, 19.5% | 520, 18.3% | 600, 20.2% | 555, 19.9% | 553, 18.7% |
| Pre-existing chronic disease (yes; n, %) | 1138, 39.4% | 1217, 38.8% | 1493, 42.7% | 1352, 39.6% | 1224, 38.6% | 1375, 40.1% | 1345, 40.2% | 1176, 41.2% | 1230, 40.2% | 1152, 40.2% | 1192, 40.3% |
| Self-reported COVID-19 symptoms (yes; n, %) | 1079, 37.4% | 1132, 36.1% | 1110, 31.7% | 1142, 29.5% | 1047, 33.0% | 1234, 36.0% | 1159, 34.6% | 1005, 35.4% | 1987, 36.5% | 939, 33.6% | 945, 31.9% |
| Previous SARS-CoV-2 testing (yes; n, %) | 143, 4.95% | 159, 5.07% | 318, 9.09% | 308, 9.75% | 363, 11.4% | 810, 23.6% | 1107, 33.1% | 1061, 37.4% | 1268, 42.6% | 1341, 48.1% | 1475, 49.9% |
| Previously tested positive for SARS-CoV-2 RNA (n, %) | 12, 8.39% | 11, 6.92% | 16, 5.03% | 15, 4.87% | 6, 1.65% | 14, 1.73% | 19, 1.72% | 31, 2.92% | 43, 3.39% | 75, 5.59% | 82, 5.56% |

| Study characteristics |         |         |         |         |         |         |         |         |         |         |         |
| SARS-CoV-2 positivity and estimated prevalence |     |     |     |     |     |     |     |     |     |     |     |
| No of test positives | 4       | 8       | 2       | 0       | 1       | 5       | 10      | 30      | 55      | 42      |         |
| Prevalence (%)        | 0.27% (0.10% 0.65%) | 0.17% (0.05% 0.65%) | 0.04% (0.00% 0.12%) | 0.00% (0.00% 0.12%) | 0.01% (0.00% 0.08%) | 0.22% (0.08% 0.49%) | 0.37% (0.18% 0.68%) | 1.34% (0.92% 1.89%) | 1.27% (0.87% 1.79%) | 2.69% (2.08% 2.96%) | 2.05% (1.53% 2.69%) |

CI, confidence interval; SD, standard deviation.
Table 2
Risk factors for testing positive for SARS-CoV-2, population-based SARS-CoV-2 prevalence studies, Estonia, 2020–2021.

| Variables | Odds ratio (OR) | Lower confidence limit (2.5%) | Upper confidence limit (97.5%) | P-value |
|-----------|----------------|------------------------------|-----------------------------|---------|
| Data collection timing | | | | |
| April 23–29, 2020 | 1 | | | |
| April 30–May 6, 2020 | 0.63 | 0.13 | 3.10 | |
| May 22–31, 2020 | 0.67 | 0.08 | 5.39 | *** |
| June 11–22, 2020 | 0.00 | 0.00 | 0.00 | |
| Aug 6–25, 2020 | 0.02 | 0.00 | 0.22 | ** |
| Sept 21–Oct 3, 2020 | 0.84 | 0.16 | 4.43 | |
| Nov 11–19, 2020 | 1.44 | 0.30 | 6.84 | |
| Nov 26–Dec 6, 2020 | 5.35 | 1.25 | 22.93 | * |
| Dec 11–20, 2020 | 5.12 | 1.21 | 21.73 | * |
| Jan 7–18, 2021 | 11.07 | 2.65 | 46.32 | *** |
| Jan 21–Feb 2, 2021 | 8.48 | 2.03 | 35.43 | ** |
| Participant Language | | | | |
| Estonian (base) | 1.00 | | | |
| Russian | 1.85 | 1.15 | 2.99 | * |
| Size of the household (number of individuals) | | | | |
| Reporting symptoms* at the time of study | | | | |
| No | 1.00 | | | |
| Yes | 2.21 | 1.59 | 3.08 | *** |
| Region of the country | | | | |
| Harju County w/o Tallinn (base) | 1.00 | | | |
| Hiiumaa County | 0.91 | 0.32 | 2.60 | |
| Ida-Virumaa County | 3.06 | 1.67 | 5.59 | *** |
| Jõgeva County | 0.14 | 0.02 | 1.03 | |
| Järva County | 0.54 | 0.14 | 2.05 | |
| Lääne-Virumaa | 1.08 | 0.38 | 3.04 | |
| County | | | | |
| Lääne County | 0.24 | 0.05 | 1.06 | |
| Põlva County | 0.13 | 0.02 | 1.00 | |
| Pärnumaa County | 0.86 | 0.37 | 1.99 | |
| Raplamaa County | 0.38 | 0.11 | 1.32 | |
| Saare County | 0.93 | 0.38 | 2.27 | |
| Tartumaa County | 1.32 | 0.74 | 2.34 | |
| Tartu County | 0.87 | 0.50 | 1.52 | |
| Tartu County w/o city | 0.50 | 0.17 | 1.52 | |
| Valga County | 0.83 | 0.19 | 3.60 | |
| Viljandi County | 2.20 | 0.95 | 5.12 | |
| Võrumaa County | 1.71 | 0.74 | 3.91 | |

* Participants reporting at least one of the three major symptoms (cough, fever, dyspnoea) or at least two of minor symptoms (fatigue, sputum production, muscle or joint aches, headache, new loss of taste or smell, sore throat, congestion or runny nose, nausea or vomiting, diarrhoea, irritability or confusion).

Discussion

The nationwide study documents substantial changes in the population prevalence of SARS-CoV-2 RNA in Estonia during the 1st year of the COVID-19 epidemic, with an initial decrease between April and June, 2020. The findings of the post-1st wave of COVID-19 prevalence and decline are in perfect agreement with a community-based SARS-CoV-2 study from England for the period of April to June 2020. In their study, SARS-CoV-2 community prevalence of 0.32% (95% credible interval 0.19–0.52%) in April 2020 declined to a very low level by the end of June 2020 (0.08%, 95% credible interval 0.05–0.12%). In Estonia, the short period of very low SARS-CoV-2 prevalence over the summer of 2020 was followed by an initially slow (in September and October) and then escalating increase since November 2020.

This study documents a clear decline in the prevalence of SARS-CoV-2 following the implementation of the nationwide non-pharmaceutical intervention (NPI) at the beginning of the epidemic. SARS-CoV-2 prevalence remained extremely low for a short period after lifting NPI measures. In the face of mitigation (slowing down transmission) rather than suppression (stopping SARS-CoV-2 community spread) of containment, an exponential increase of new COVID-19 cases occurred at the verge of the 2nd year of the epidemic.

These findings allow us to speculate that, until now, this is a very unforgiving virus. While rigorous and comprehensive NPI measures are clearly effective in stopping transmission, lifting the measures or less stringent implementation will lead to new and sizable outbreaks.

Second, findings from Estonia should be interpreted in the context of the high SARS-CoV-2 testing rate (80,630/100,000), a very low COVID-19 case fatality rate of 0.8% (both, as of March 18, 2021) and no significant excess (all cause) deaths over the first year of the epidemic.

We saw that those with a larger household size were at a higher risk of SARS-CoV-2 infection with no attributable risk either from the age of the individual or from the age structure of the household (very similar to the results of the study from the UK). Ongoing household transmission with occasional spill over to other households could act as an important driver for ongoing transmission and is estimated to be responsible for roughly 70% of SARS-CoV-2 transmission when widespread community control measures are in place.

Our findings of higher SARS-CoV-2 risk among those reporting symptoms characteristic to COVID-19 are clearly not new. Yet, it highlights the need to focus on symptomatic cases rather than mass-testing in the face of resource constraints or competing resource needs (i.e., vaccination). Focus on symptomatic COVID-19 cases has a solid evidence base—the majority of COVID-19 cases are asymptomatic (~60–80%) and are significantly more likely to infect their close contacts than their asymptomatic counterparts.

Last but not least, we saw regional and ethnic (main language spoken) differences in SARS-CoV-2 positivity. Disproportionately affected racial and ethnic minority groups have been reported elsewhere (United States, UK). In Estonia, ethnic disparities are not unique to COVID-19 outcomes. The reasons for ethnic disparities in COVID-19 outcomes are multilayered and underline...
the regional differences in Estonia. Ida-Viru County is in the northeastern part of Estonia bordering the Russian Federation. The overwhelming majority (82%) of residents are Russian speaking. It is important to note that nearly 75% of Russian speakers in Estonia regularly follow TV channels and online media originating from the Russian Federation\(^{34}\) and are more likely to trust Russian than domestic (Estonian) or EU media.\(^{35}\) Whether the Russian Federation’s pandemic-related disinformation campaign\(^{36}\) has had some effect on the beliefs and behaviours of the Russian-speaking population in Estonia (and other neighbouring countries with sizable Russian speaking minorities) is unknown at this stage. There are anecdotal reports from Ida-Viru County on residents of declining state-provided COVID-19 vaccines and demands to be vaccinated with the Russian Sputnik vaccine.\(^{37}\) There is a risk that COVID-19 vaccine uptake will be lower among minority ethnic groups in Estonia, thereby widening the health gap further. COVID-19 risk communication and community engagement is a priority for information provision and to counter misinformation.

In conclusion, a rather limited number of studies have assessed the prevalence of SARS-CoV-2 infection in the general population (seroprevalence,\(^{38,39}\) SARS-CoV-2 RNA\(^{10,40,41}\) ). Population-based studies assessing temporal changes in SARS-CoV-2 prevalence, either via repeated cross-sectional studies\(^{42}\) or following subjects longitudinally,\(^{3}\) are, to our best knowledge, exceedingly rare. It is critically important to create a knowledge base to inform future strategies, and a range of real-life COVID-19 epidemic scenarios over extended periods needs to be documented to assist in understanding the infection risk factors at the individual and population levels. Analyses based on patients in need of hospital treatment, and/or with comorbidities reported during the early phases of the COVID-19 epidemic, were unable to disentangle infection from virulence risks. Yet, primary prevention operates through the control of (the true) infection risk factors.

Our study has several limitations. The degree to which the study is representative of the larger population is influenced by the low response rate and potential selective factors associated with responses. To minimise non-response bias, the prevalence estimates were weighted (age, gender and region) to ensure representativeness of the source population. Yet, there could be other factors for which we did not have detailed information about population distributions which are also associated with testing positive for SARS-CoV-2. The number of people testing SARS-CoV-2 RNA positive in the cross-sectional studies is low, leading to relatively large uncertainty around estimates.

We see the long period of observation and population-based nationwide study design as strengths of our work. Interpretation of changes in SARS-CoV-2 incidence and positivity rates originating from case notification or clinical cases is likely to be confounded by substantial changes in testing practice over time. Our study is based on a series of cross-sectional studies with a standardised methodology and is thereby very unlikely to be influenced by the testing practice. As this evaluation is based upon observing a single population over time, we speculate that selection bias or unmeasured confounders would operate rather uniformly over the period of observation, though presenting a less-threatening trend of SARS-CoV-2 prevalence and analysis of factors associated with SARS-CoV-2 positivity.

Conclusions

The population-based effect of the novel vaccines against SARS-CoV-2 is highly contingent on the infection-blocking (or transmission-blocking) action of the vaccine and population uptake.\(^{8}\) SARS-CoV-2 population prevalence needs to be carefully monitored to inform containment decisions as vaccine programmes are rolled out.

Author statements

Acknowledgements

We thank our study partners Kantar-Emor AS, OÜ Medicum Eliarsiabi, and SYNLAB Eesti OÜ.

Ethical approval

Ethical approval for the study was obtained from the Research Ethics Committee of the University of Tartu.

Funding

This study was funded by grants SMVPT20243, SMVPT20599, IUT34-4 from the Estonian Ministry of Education and Research;
Estonian Research Council Grants (PRG1095, PRG59, PRG1197), and Project No 2014-2020.4.01.16-0271, ELIXIR.

Competing interests

The authors report no competing of interest.

Author contributions

Anneli Uusküla - Conceptualisation, Methodology, Writing - Original Draft; Ruth Kaldla, Miikkel Solvak, Mikk Jurisson, Krista Fischer, Aime Keis, Kristjan Vassil, Jaak Vilo, Hedi Peterson, Lili Milani, Mait Metspalu - Conceptualisation, Writing - Review & Editing; Meelis Kääriik, Krista Fischer, Uku Raudvere, Ene Kääriik, Liis Kolberg, Tuuli Jürgenson, Ene-Margit Tiit - Investigation, Data analysis, Writing - Review & Editing; Meelis Kääriik, Krista Fischer - Visualisation. All listed authors reviewed and edited the manuscript and approved the final, submitted version. All authors confirm that they had full access to the data in the study and accept responsibility to submit for publication.

References

1. WHO. WHO Director-General’s opening remarks at the media briefing on COVID-19 – 11. https://www.who.int/dg/speeches/detail, March 2020. [Accessed 21 August 2020].
2. COVID-19 situation update worldwide, as of week 49. Updated 16 December 2021. https://www.ecdc.europa.eu/en/geographical-distribution-2019-covn-cases, 2021. [Accessed 21 December 2021].
3. Saad-Roy CM, Wagner CE, Baker RE, Morris SE, Farrar J, Graham AL, et al. Immune response to SARS-CoV-2 in the Icelandic population. Lancet 2020;395(10233):1395–8. https://doi.org/10.1016/S0140-6736(20)30201-3.
4. Saad-Roy CM, Wagner CE, Baker RE, Morris SE, Farrar J, Graham AL, et al. Spread of SARS-CoV-2 in the Icelandic population. SARS-CoV-2 IgG antibodies in an area of northeastern Italy with a high incidence of COVID-19 incidence by age, sex, and period and period countries worldwide as of 13 April 2021 (per million population). https://www.statista.com/statistics/1104645/covid19-testing-rate-select-countries-worldwide/, 2021. [Accessed 18 March 2021].
5. European. Graphs and maps (18.03.2021). https://www.europa.eu/en-graphs-and-maps/excess-mortality, 2021. [Accessed 18 March 2021].
6. Martin CA, Jenkins DR, Minhas JS, Gray LJ, Tang J, Williams C, et al. Socio-demographic heterogeneity in the prevalence of COVID-19 during lockdown is associated with ethnicity and household size: results from an observational cohort study. EClinicalMedicine 2020;25:100466. https://doi.org/10.1016/j.eclinm.2020.100466.
7. Nande A, Adlam B, Sheen J, Levy MZ, Hill AL. Dynamics of COVID-19 under social distancing measures are driven by transmission network structure. PLoS Comput Biol 2021;17(2):e1009684. https://doi.org/10.1371/journal.pcbi.1009684.
8. Ronchina F, Peng Z, Guo Y, Song R, Li Y, Xiao Y, et al. Assessing the effects of metropolitan-wide quarantine on the spread of COVID-19 in public space and households. Int J Infect Dis 2020;96:503–5. https://doi.org/10.1016/j.ijid.2020.05.019.
9. Byamugashira O, Cardon M, Bell C, Clark J, MvLaws ML, Glasziou P. Estimating the extent of asymptomatic COVID-19 and its potential for community transmission: systematic review and meta-analysis. JAMMI 2020;5(4):223–34. https://doi.org/10.3186/jammi-2020-0030.
10. Sayanpanathan AA, Heng CS, Pin PH, Fang J, Leong TY, Lee VJ. Infectivity of asymptomatic versus symptomatic COVID-19. Lancet 2021;397(10269):93–4. https://doi.org/10.1016/S0140-6736(20)32651-9.
11. Van Dyke ME, Mendoza MCB, Li W, Parker EM, Belay D, Davis EM, et al. Racial and ethnic disparities in COVID-19 incidence by age, sex, and period among US counties June 25 Years - 16 U.S. Jurisdictions, January 1-December 31, 2020. Mortal Mortal Wkly Rep 2021;70(11):382–8. https://doi.org/10.15585/mmwr.mm7011e1.
12. Razai MS, Kankam HKN, Majeed A, Esmaill Williams DR. Mitigating ethnic disparities in covid-19 and beyond. BMJ 2021;372:m4921. https://doi.org/10.1136/bmj.n9231.
13. Rahu V, Kiviöpp F, Villand K, Pehme L, Kahu M. Respiratory tuberculosis incidence and mortality in Estonia: 30-year trends and sociodemographic determinants. Int J Tubercul Lung Dis 2021;25(1):112–8. https://doi.org/10.5588/ijtlkd.20.0388.
14. Loit U, Harro-Loit H. Media Pluralism Monitor 2016. In: Role of Russian Media in the Baltics and Moldova. https://www.usagwp.wp-content/uploads/2012/08/RBRC-Gazette-Russian Media-pg-2/02-04-16.pdf, 2021. [Accessed 18 March 2021].
15. Weits R. Assessing the Russian disinformation campaign during COVID-19. International Center for Defence and Security. 2020; 11.5.2021. https://icsd.ee/en/assessing-the-russian-disinformation-campaign-during-covid-19/, 2021. [Accessed 18 March 2021].
16. Eesmaa M. Sotsiaalmisinspektiion hakukoti. Sputniku vaktsinili veenma. 10.02.2021. https://www.delfi.ee/news paevaandaduit/estvideo/sotsiaalmisinspektiion-hakukoti-sputniku-vaktsinili-veenma-veenma–id–9253421, 2021. [Accessed 18 March 2021].
17. Stefaneli P, Bella A, Fedele G, Pancheri S, Leone P, Vacca P, et al. Prevalence of SARS-CoV-2 in a veterinary population in an area of northeastern Italy with a high incidence of COVID-19 cases: a population-based study. Clin Microbiol Infect 2020;36(2):2138.e1-2138.e10. https://doi.org/10.1016/j.clinmicinfe.2020.01.014.
18. Pollán M, Pérez-Gómez B, Pastor-Barriuso R, Óteo J, Hernández MA, Pérez-Díaz M, et al. Prevalence of SARS-CoV-2 in Spain (ENCE-COVID): a nationwide, population-based soropospectrological study. Lancet 2020;396(10250):535–44. https://doi.org/10.1016/S0140-6736(20)31483-5.
19. Menachemi N, Yiannoutsos CT, Dixon BE, Duszynski TJ, Fadel WF, Wools-K卡拉orom J, et al. Population point prevalence of SARS-CoV-2 infection based on a statewide random sample - Indiana, April 25-29, 2020. Morb Mortal Wkly Rep 2020;69(29):960–4. https://doi.org/10.15585/mmwr. mm6929e1.
20. Riley S, Ainslie KE, Eales O, et al. High prevalence of SARS-CoV-2 swab positivity in England during November 2020: interim report of round 5 of REACT-1 study. medRxiv. 2020. https://doi.org/10.1101/2020.10.09.20204727.
21. Lavezzi E, Franchin E, Cavallero C, et al. Suppression of a SARS-CoV-2 outbreak in the Italian municipality of Vo’. Nature 2020;584:425–9.