Swedish Covid-19 Investigation for Future Insights – A Population Epidemiology Approach Using Register Linkage (SCIFI-PEARL)

Fredrik Nyberg, Stefan Franzén, Magnus Lindh, Lowie Vanfleteren, Niklas Hammar, Björn Wettermark, Johan Sundström, Ailiana Santosa, Staffan Björck, Magnus Gisslén

School of Public Health and Community Medicine, Institute of Medicine, Sahlgrenska Academy, University of Gothenburg, Gothenburg, Sweden; National Diabetes Register, Centre of Registers Västra Götaland, Gothenburg, Sweden; Department of Infectious Diseases, Institute of Biomedicine, Sahlgrenska Academy, University of Gothenburg, Gothenburg, Sweden; Department of Clinical Microbiology, Sahlgrenska University Hospital, Gothenburg, Sweden; Department of Internal Medicine and Clinical Nutrition, Institute of Medicine, Sahlgrenska Academy, University of Gothenburg, Gothenburg, Sweden; Unit of Epidemiology, Institute of Environmental Medicine, Karolinska Institute, Stockholm, Sweden; Pharmacoepidemiology & Social Pharmacy, Department of Pharmacy, Uppsala University, Uppsala, Sweden; Clinical Epidemiology Unit, Department of Medical Sciences, Uppsala University, Uppsala, Sweden; The George Institute for Global Health, University of New South Wales, Sydney, Australia; Centre of Registers Västra Götaland, Gothenburg, Sweden; Region Västra Götaland, Department of Infectious Diseases, Sahlgrenska University Hospital, Gothenburg, Sweden

Background: In response to the Covid-19 pandemic, we designed and initiated a nationwide linked multi-register, regularly updated, observational study for timely response to urgent scientific questions.

Aim: To describe the SCIFI-PEARL (Swedish Covid-19 Investigation for Future Insights – A Population Epidemiology Approach using Register Linkage) linked database encompassing essentially all known diagnosed Swedish Covid-19 patients plus a large general population comparison cohort and outline its utility in the current and future phases of the pandemic.

Methods: Individuals with Covid-19 from the entire country are identified on a regularly updated basis, from different sources: all individuals from SmiNet, the national database of notifiable diseases, with positive SARS-CoV-2 polymerase chain reaction (PCR) test results; patients identified in the healthcare system by condition (ICD-10) or procedure codes in the National Patient Register or Cause-of-Death Register; patients identified through several disease-specific national quality registers (NQRs); and in two regions additionally patients identified in primary care. A comparison population was obtained by stratified random sampling from Swedish national population registers. Data from all these registers plus the National Prescribed Drug Register, the Cancer Register, national sociodemographic registers, some additional NQRs, the National Vaccination Register, and further data sources, are then linked to all study subjects (Covid-19 cases and population cohort). New cases in the study population and all data for all subjects are updated every few months, as required.

Conclusion and Utility: The SCIFI-PEARL study cohort captures Swedish residents with Covid-19 on an ongoing basis, includes a representative general population comparison cohort, and links to a broad range of national and regional healthcare data for a comprehensive longitudinal view of the Covid-19 pandemic. By combining high-quality national registers with short time delay and continuous repeated linkage and updating, the project brings timely and internationally relevant data for epidemiological research on SARS-CoV-2. Our efforts provide an example and important learnings for similar efforts internationally in the future.

Keywords: SARS-CoV-2, population cohort, data updates, observational research, longitudinal

Introduction

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) first emerged in Wuhan in December 2019, and was soon found to cause the disease now named coronavirus disease 2019 (Covid-19). During 2020, the disease developed into a large-scale pandemic, spreading to most countries in the world, including enterin
a second or third wave in many places, with cumulative numbers of over 83 million cases and 1.8 million deaths reported globally during the first year until end of 2020.\textsuperscript{2} By the end of 2020, vaccines had been developed and vaccination campaigns were initiated in multiple countries, including Sweden. The first Covid-19 case in Sweden was reported on 31 January 2020, and in parallel with Europe becoming the epicenter for the epidemic during the Spring of 2020, the disease rapidly expanded in Sweden, causing severe strain on society and the healthcare system.\textsuperscript{3}

It was clear at the time that research both in the short and long term would be required, and the research communities in Sweden and internationally quickly geared up to respond to the challenge. The study described here based on Swedish data was rapidly designed as a large-scale effort to assemble a rich database for the study of Covid-19 epidemiologically in real time, ie, with the requirement that data would be regularly and sufficiently frequently updated. The importance of not only focusing on Covid-19 patients was also recognized, making the inclusion of a large comparison cohort reflecting the general population another key asset.

Even against the background of fast-track processes for ethical approval and data access for Covid-19 research developing at the time,\textsuperscript{4} this has been quite a complex task without clear precedent in the Swedish registry data ecosystem. This paper describes the database now established under the name of SCIFI-PEARL (Swedish Covid-19 Investigation for Future Insights – a Population Epidemiology Approach using Register Linkage), and outlines its research potential and utility in the ongoing and future phases of the pandemic.

### Materials and Methods

Sweden, similar to other Nordic countries, has high quality national health and population registers that can be linked with a high degree of accuracy, providing an extraordinary suitable environment for epidemiologic research. The Swedish Personal Identity Number (PIN; personnummer) uniquely linked to each individual enables compilation and linkage of data from diverse register data to the same individuals, with relative ease and with extremely high accuracy.\textsuperscript{5} This SCIFI-PEARL study exploits the possibility to assemble a unique large database of individuals with Covid-19 plus a population cohort. After linkage, the database is pseudonymised, according to usual procedures.

### Study Population

Newly identified individuals with Covid-19 from the entire country are included in the linked database on a regularly updated basis, from different sources (Figure 1). This includes all individuals with a positive SARS-CoV-2 polymerase chain reactions (PCR) test results identified from SmiNet,\textsuperscript{6} the national register of notifiable communicable diseases managed by the Public Health Agency of Sweden. In addition, patients who may not have a positive test registered but have been identified by the healthcare system are included, eg, by Covid-19 ICD codes (ie, ICD-10 U07.1 = Covid-19, virus demonstrated; U07.2 = Covid-19, virus not shown) or relevant procedure codes on a national basis from the National Patient Register (NPR) for outpatient and hospitalized inpatient specialist care.\textsuperscript{7} Similarly, individuals recorded in the National Cause-of-Death Register (NCDR)\textsuperscript{8} as deaths due to Covid-19 are included. In addition, registration of test result and ICD diagnoses made by several disease-specific national quality registers (NQRs) are scrutinized to identify additional potential Covid-19 patients with healthcare contacts related to their particular disease and therefore captured in that register. Further, since there is no national coverage for primary care data, healthcare contacts for Covid-19 in primary care are captured on a regional basis, for Western Sweden (Västra Götaland Region) and the Stockholm Region.

In addition, a general population comparison group was obtained using the national Register of the Total Population (RTB) at Statistics Sweden,\textsuperscript{9} consisting of approximately 10% of the Swedish population (~1,000,000 individuals), based on individuals of all ages resident in Sweden on 1 January 2020, through stratified random sampling in 20 age (5-year age categories) by 2 sex strata. This population can be used either in analyses as it is, weighted to the Swedish population through sampling weights, or subsampled, as required for different types of statistical analyses and research questions.

### Data Acquisition

To this combined study population of individuals with Covid-19 from the whole Swedish population plus the 10% general population comparison group, we linked diverse data from a range of registers with relevant information, to get a comprehensive and detailed view of patient history (Figure 1, Table 1). For practical and ethical reasons the actual linkage is performed by the National
Board of Health and Welfare (NBHW), and the PIN for each subject is replaced by an individual study subject identifier, to pseudonymise the data.

The bulk of the data are from different national registers. Date and cause of death are obtained from RTB and the NCDR. In-patient and specialist outpatient healthcare data from 2015 onward, comprising a complete specialist care medical history for all individuals, are obtained from the National Patient Register (NPR). Cancer incidence data from 2015 are extracted from the National Cancer Register (NCR). A complete drug history for prescription drugs from 2018 is obtained from the National Prescribed Drug Register (NPDR).

Sociodemographic data including education, family situation, income and occupational data from 2018 onward are obtained from the Statistics Sweden Longitudinal Integrated Database for Health Insurance and Labour Market Studies (LISA; Longitudinell Integrationsdatabas för Sjukförsäkrings- och Arbetsmarknadsstudier). Finally, data from a range of national quality registers are included. These registers are national in scope and provide detailed information on Covid-19 management, as well as relevant comorbidities and risk factors of importance for Covid-19 patients, with high granularity and detail, on all recorded individuals with the respective underlying disease or condition covered by that particular NQR (Table 1). They include registers for acute emergency care and intensive care (particularly important for Covid-19 patient data) and the Swedish Register of Cardiopulmonary Resuscitation (SRCR) with data extracted from 1 Jan 2020, as well as registers for infectious diseases, several cardiovascular disease registers, 2 respiratory disease registers, and the National Diabetes Register (NDR) with data from 2015. The database is also linked to the National Vaccination Register (NVR), which following urgent pandemic legislative changes in Dec 2020 captures information on vaccinations against Covid-19 in Sweden.

In addition, some data not available on a consolidated national basis are included in the database for a more detailed regional view. For Western Sweden (Västra Götaland Region) and the Stockholm Region, with a population of 4,902,162 out of 10,327,589 nationally (39.7%), healthcare contacts and diagnoses recorded in primary care are captured from 2015. For Western Sweden, a data source with data on SARS-CoV-2 testing from the start of the pandemic (both positive and negative tests), is also included in the database, to provide data on test penetration over time in the population.

All data are updated with the most recent data from all included registers on a regular basis, every few months, as required by the study objectives and the evolution of the

Figure 1 Structure of SCIFI-PEARL database. Included registers; identification of study population of COVID-19 patients (blue arrows) and general population sample (purple arrow); followed by linkage to all registers for relevant data from each register.
pandemic, in an agreed process triggered each time by a request from the study team to all registers. The complex update process from extraction of data at each data source to linkage and delivery of a new complete dataset is designed to take around 1 month.

Data Management
The study population selection, data linkages and pseudonymisation of the data are managed in secure environments at the NBHW and Statistics Sweden. The pseudonymized research database thus produced and delivered is then housed on a dedicated server at the Centre of Registers Västra Götaland, which has extensive experience with large databases, as it houses and manages a number of NQRs. Data transfers use state-of-the-art encryption and the project-specific hardware is maintained with high protection levels relevant for sensitive personal data, with regular backups. Access protection includes disc encryption, network segmentation and two-level firewall protection with differential access rights based on project role.

Ethics
The study, including regular updates until at least end of 2025, has received ethical approval from the Swedish Ethical Review Authority.

Research Objectives
The core initial research objectives outlined in the project plan are relatively broad and cover descriptive, analytical and predictive modeling analyses.

Primary Aims
- To descriptively characterize individuals with Covid-19 (overall, hospitalized, intensive care unit (ICU) treated, deaths, and in important subgroups) in terms of sociodemographics, various risk factors (eg, comorbidity, medication), geographic distribution, timing, etc. Further, describe how the epidemic and the characteristics of the affected patient group change over time, ie, “natural history” of the epidemic in relation to the population.
- To quantify the impact and importance of comorbidities (in particular respiratory and cardiovascular, diabetes, obesity), severity of and co-medications for concomitant disease, sociodemographics, and other risk factors, as well as protective measures and vaccinations, on risk of various endpoints. Some anticipated endpoints include developing Covid-19, being hospitalized for Covid-19, emergency/ICU care, treatments, specific care, hospitalization duration, prognosis, death and in survivors study long-term outcomes up to 3 years. Various study designs and methods as appropriate will be employed, including case-cohort, case-control and interrupted time series study designs, and machine learning methods.
- To develop, and periodically update prediction models for Covid-19 occurrence and prognosis, and evaluate how this impacts healthcare resource use, including the need for ICU treatment. Use of machine learning methods will be important here.

Secondary Aims
Secondary aims will include rapidly upcoming research questions based on primary prevention and healthcare needs, and include:

- Treatment effects of medicines and other healthcare interventions, including novel indications for established treatments and vaccinations, and possible side-effects related to Covid-19.
- Resource use in the healthcare system, at the individual level, and at societal level (hospital resources, measures, medicines, costs, etc.)
- Study the effectiveness of vaccination programs and possible adverse reactions in the population.
- Consequences of the covid-19 pandemic including future vaccination programs on drug use, healthcare and other societal functions in the population.
- Other upcoming questions and information gaps from society, healthcare and the scientific community (if these are outside the above specified issues, may need additional supplemental ethical application).

Current and Future Activities
A first data delivery of all data was compiled in October 2020. Data merging and data management, as well as analysis variable and analysis database construction, are performed on an ongoing basis. We have received three complete data updates by Spring 2021. The updating process has thus been tested and found to be functional, and regular updates of the study population and data may now be considered routine. A number of research analyses are now in various stages of completion.

Based on the first data delivery, the number of study subjects and their age and sex distribution are described in Table 2 and Figure 2. As the data are updated, the population comparison cohort (as sampled on 31 Dec 2019) will remain initially static, with a view to update at suitable
### Table 1  Source Registers and Types of Data Currently Available in the SCIFI-PEARL Study Database

| Register                                      | Register Holder + Supporting Register Centre / Personal Data Holding Authority* | Type of Information                                      | Example Variables                                                                 |
|-----------------------------------------------|---------------------------------------------------------------------------------|-----------------------------------------------------------|-----------------------------------------------------------------------------------|
| RTB - Register of the Total Population        | SCB** (Statistics Sweden) / same                                                 | Demographic data                                          | Birth/death/immigration/emigration dates.                                          |
| LISA - (Statistics Sweden Longitudinal Integrated Database for Health Insurance and Labour Market Studies) | Same as above                                                                   | Socio-demographic data                                    | Sex, date of birth, education, family situation, income, occupation, country of birth data |
| NPR - National Patient Register               | Swedish National Board of Health and Welfare (NBHW; Socialstyrelsen) / same      | In-patient and specialist outpatient healthcare data      | Dates of hospitalization (start/end) and specialist hospital physician visits Primary and secondary diagnoses as ICD-10 codes Date of diagnosis |
| NCDR - National Cause-of-Death Register       | Same as above                                                                   | Cause of death data                                       | Date of death, underlying and contributing causes of death as ICD-10 codes         |
| NCR - National Cancer Register                | Same as above                                                                   | Cancer incidence data                                     | Site of tumor, stage, type of tumor, Date of diagnosis                             |
| NPDR - National Prescribed Drug Register      | Same as above                                                                   | Prescription drugs data                                   | Date of prescription fill, ATC code for drug, dose                                 |
| SmiNet – National surveillance system for reportable infectious diseases | Swedish Public Health Agency / same                                              | Communicable diseases data                                | Date of positive SARS-CoV-2 PCR test                                               |
| NVR - National Vaccination Register           | Same as above                                                                   | Vaccination data                                          | Type, dose, place and date of vaccination                                          |
| SIR - Swedish Intensive Care Registry         | SIR Steering Group + QRC Stockholm / Region Värmland                              | Intensive care data                                        | Diagnoses, interventions, inpatient care, Patient Reported Outcome Measures (PROM) or other patient-reported health effects |
| ANSWER, A National SWedish Emergency Registry (SVAR**) | ANSWER Steering Group + UCR / Karolinska University Hospital                   | Emergency department data                                 | Patient diagnoses, assessments, interventions and care Date, time and type of admission Discharge details or date/cause of death Process details, dates and times |

(Continued)
| Register | Register Holder + Supporting Register Centre / Personal Data Holding Authority* | Type of Information | Example Variables |
|----------|---------------------------------------------------------------------------------|---------------------|-------------------|
| SRCR - Swedish Registry of Cardiopulmonary Resuscitation (HLR***) | SRCR Steering Group + Register Centre (RC) Västra Götaland (VG) / Region VG through RC VG | Medical cardiac arrest and cardiopulmonary resuscitation data | Cardiopulmonary resuscitation (CPR) before arrival of the emergency medical services (EMS) staff Information on treatment provided during CPR Outcome (ie 30-day survival) |
| InfectionReg - National Quality Register for Infectious Diseases | InfectionReg Steering Group + RC South / Region Örebro | Infectious diseases data | Diagnoses, interventions, treatment and Patient Reported Outcome Measures (PROM) for some types of infection (incl. Pneumonia) Severity of disease Patients’ risk profile, medical and medical device treatment Outcomes, and complications from the time of intervention for all procedures and surgical interventions |
| SWEDHEART - National Quality Registry for Enhancement and Development of Evidence-Based Care in Heart Disease | Swedeheart Steering Group + Uppsala Clinical Research (UCR) / Region Uppsala | Inpatient and outpatient cardiac care data | |
| RiksSwikt - National Quality Registry for Heart Failure | NQR Steering Group + UCR / Region Östergötland | Inpatient and outpatient heart failure data | Hospital discharge Outpatient clinic visit Diagnostics Laboratory values Medical treatment Follow up |
| RiksStroke - National Quality Registry for Stroke | RiksStroke Steering Group + RC North / Region Västerbotten | Stroke patient’s data | Admission to stroke unit Diagnostic procedures Medical complications during hospital stay Discharge status Management and outcomes in routine clinical settings. |
| SNAR - Swedish National Airway Register (LVR***) | SNAR Steering Group + RC VG / Region VG through RC VG | Asthma patient data (both children and adults) and COPD outpatient and inpatient’s data from primary, secondary and tertiary care | Diagnostic investigations (spirometry, blood sample, skin prick test), symptom-scores, comorbidities and prescribed treatments |
intervals to the current Swedish population, whereas the number of Covid-19 patients will grow successively with each data delivery.

Currently, a number of research projects have been started at various stages. The core project team has initiated collaborations with a number of external researchers, and collaboration proposals from other external partners nationally and internationally are encouraged. The study database is a valuable resource that should be used as broadly as feasible for important research to provide epidemiological and clinical evidence to benefit patients, the healthcare system and society.

We are also exploring the potential to link the database to some existing large-scale well-characterized population-based epidemiological studies in Sweden that have more detailed clinical and questionnaire data on specific populations, such as the West Sweden Asthma Study (WSAS) and SCAPIS (the Swedish CArdioPulmonary bioImage Study). These potential spin-off projects will likely require separate setup and ethics review, but may provide interesting insights, also on a more mechanistic level.

An international outlook is also important. We have established collaboration with the Nordic COHERENCE project, aiming to facilitate large-scale register-based observational analyses across several Nordic Covid-19 national databases, especially for research questions where larger sample sizes are desirable, or where comparisons across different populations, societies or healthcare systems may be relevant. The analysis strategy will build on a Nordic common data model (CDM) and distributed analyses, retaining data at each country site to protect individual data. Other international collaborations are also encouraged.

**Discussion**

The Covid-19 pandemic is a significant concern globally as well as in Sweden, given the pace of spread and breadth of health impacts on the population level, posing a massive challenge to global health. A major obstacle to understanding, managing and preventing Covid-19 continues to be inadequate, non-transparent, and out-of-date Covid-19 data as the pandemic progresses. Many scientists continue to search for innovative sources and uses of real-world data for comprehensive and current Covid-19 research. Since knowledge on Covid-19 is constantly evolving, population-based research at a national scale is crucial, using the latest, point-of-care data that can be leveraged to support research and development. Carefully assembled and used, the high-quality national register data available in Sweden will help research
### Table 2: Study Sample for SCIFI-PEARL Based on Initial October 2020 Data Delivery, with Age and Sex Distributions

|                      | COVID-19 Cases in Sweden (Included in the Study) | General Population Based Comparison Group |
|----------------------|-------------------------------------------------|------------------------------------------|
|                      | Total Registered in the Study Healthcare Data at Some Time Point * | Test-Positive | Hospitalized | Weighted to the Age-Sex Distribution of the Swedish Population** | Stratified Random Sample (ie Actual number of Study Participants) |
| Study sample, n      | 126,953                                         | 86,798         | 17,026       | 10,327,589                                                 | 972,723 |
| Age, mean (SD)       | 49.36 (21.70)                                   | 48.51 (21.42) | 64.11 (18.37) | 40.80 (24.01)                                               | 48.11 (28.05) |
| Age categories, n (%) |                                                |                |              |                                                          |                    |
| 0–9 years            | 1605 (1.3%)                                     | 698 (0.8%)     | 84 (0.5%)    | 1,225,802.0 (11.9)                                         | 100,000 (10.3)     |
| 10–19 years          | 6444 (5.1%)                                     | 4841 (5.6%)    | 106 (0.6%)   | 1,177,928.0 (11.4)                                         | 100,000 (10.3)     |
| 20–39 years          | 19,337 (15.2%)                                  | 14,234 (16.4%) | 514 (3.0%)   | 1,327,395.0 (12.9)                                         | 100,000 (10.3)     |
| 40–49 years          | 19,396 (15.3%)                                  | 13,211 (15.2%) | 1027 (6.0%)  | 1,366,489.0 (13.2)                                         | 100,000 (10.3)     |
| 50–59 years          | 20,027 (15.8%)                                  | 14,300 (16.5%) | 1077 (6.0%)  | 1,298,355.0 (12.6)                                         | 100,000 (10.3)     |
| 60–69 years          | 20,901 (16.5%)                                  | 15,046 (17.3%) | 3064 (18.1%) | 1,297,863.0 (12.6)                                         | 100,000 (10.3)     |
| 70–79 years          | 13,123 (10.3%)                                  | 8370 (9.6%)    | 3024 (17.8%) | 1,108,438.0 (10.7)                                         | 100,000 (10.3)     |
| 80–89 years          | 10,921 (8.6%)                                   | 6040 (7.0%)    | 3279 (19.3%) | 989,013.0 (9.6)                                            | 100,000 (10.3)     |
| ≥90 years            | 10,538 (8.3%)                                   | 6680 (7.7%)    | 3000 (17.6%) | 436,679.0 (4.2)                                            | 100,000 (10.3)     |
| Sex, N (%)           |                                                |                |              |                                                          |                    |
| Men                  | 54,703 (43.1%)                                  | 37,264 (42.9%) | 9907 (58.2%) | 5,195,814.0 (50.3%)                                        | 480,320 (49.4%)    |
| Women                | 72,250 (56.9%)                                  | 49,534 (57.1%) | 7119 (41.8%) | 5,131,775.0 (49.7%)                                        | 492,403 (50.6%)    |

**Notes:** Test-Positive and hospitalized COVID-19 patient subgroups are subsets of the total identified COVID-19 patient pool. *Test-Positive in SmiNet; or diagnosis in the National Patient Register, the National Cause-of-death Register, or in primary care data from Västra Götaland; or registered as COVID-19 in National Quality Registers included in the study, based on initial October 2020 data delivery. **The comparison group in the study, weighted using sampling weights based on 20 strata (sex and 10 age categories). N in this column is thus estimated and reflects the number of persons in the Swedish population represented by each category of sampled study subjects. Proportions are estimated proportions in the full Swedish population.
immensely, for example, to characterize risk patients, identify and quantify risk factors, develop machine learning tools to predict COVID-19 severity and mortality, and follow vaccination and pandemic developments, critical insights for driving care to highest-risk patients and better managing the outbreak, with broad global applicability. We have therefore assembled a continuously updated database of Covid-19 patients and general population comparator individuals.

Some learnings from this initiative are broadly relevant in an international setting. Assembling and accessing data in real time from many data sources can be challenging and very time consuming due to process, legal and other aspects – even in a country such as Sweden - and therefore good planning is essential. Plans for such efforts should be made in any country that aims for better future pandemic preparedness, tailored to the particular situation in that area. It is particularly important to pay close attention to local legal frameworks when establishing processes for data sharing. This is unfortunately not the last time there will be urgent need of scientifically robust data on a new health concern, and it will be essential to plan in advance to have rich data available for research use when the need arises.

This initiative has several strengths. The unique PIN in Sweden allows for high-quality linkage of many registers, assembling rich data for each person. Swedish national healthcare registers and national quality registers are generally of very high quality, with reliable data. The complete and accurate national population register (RTB) that is frequently updated and available for sampling enables us to sample essentially all identifiable individuals with Covid-19 in Sweden as well as a very large stratified random sample of general population comparison subjects that can provide context and comparison for planned analyses. This cannot be easily done outside the Nordic countries. The study has a large sample size and we have also built in regular updating of the database with new Covid-19 subjects and current data over time during the pandemic, to ensure optimal power and data availability.

The study builds on real life clinical data from a health care system with essentially equal access to all residents in Sweden. Thus, unequal or biased testing/diagnosis/hospital care for eg,
cost reasons is a limited problem. However, other factors such as resource limitations and prioritization, especially during the most intensive pandemic phases, may still affect access to care, and should be adequately considered to avoid bias in any research performed.

Potential weaknesses include those common to most observational data. Data quality is dependent on the completeness and accuracy of the information registered. There are inherent weaknesses related to eg, ICD10 coding and the way different conditions are classified, recorded and grouped. Similar considerations apply to procedure and drug coding. Some data are just not available, eg, despite a high-quality prescription drug register being available, in-hospital drug treatment is not readily available in Sweden. Register research is by its very nature retrospective in the sense that it relies on data collected primarily for another purpose, which may affect content, accuracy, and completeness of data. This can also be a strength, eg, for inference and potential relevance, and avoidance of some types of bias. Finally, register data most often is used for observational research, which, irrespective of study designs – descriptive, case-cohort, case control etc. – may suffer from residual bias even after use of standardization, weighting or adjustment to eliminate confounding, and therefore as high quality as possible of the underlying register data is important.

A potential weakness that needs to be considered in studies of COVID-19 is the uncertainty regarding who is tested for Covid-19 and how this affects estimates of disease occurrence or severity in different groups. Rich data to help address this is present in our study. It is also clear that the probability of being infected varies across groups depending on eg, housing conditions and social and behavioral factors, although to the extent that these can be captured in the database this will be taken into account or specifically studied. However, these weaknesses are present everywhere in epidemiologic research on Covid-19 and may in many cases be less prominent in Sweden than in some other countries with greater socioeconomic differences and more restricted access to health care. This is also not unique for Covid-19. For almost any disease there is uneven diagnosis with both over- and underdiagnosis, which varies by other factors. Similarly, most diseases lack stable gold standard criteria that can be applied in clinical practice over time, and observational research must rely on the actual clinical diagnostic efforts and clinical reality.

**Conclusions**

The SCIFI-PEARL (Swedish Covid-19 Investigation for Future Insights – a Population Epidemiology Approach using Register Linkage) study cohort captures essentially all Swedish residents with Covid-19 on an ongoing basis, includes a representative general population comparison cohort, and links to a broad range of national and regional healthcare data for a comprehensive longitudinal view of the Covid-19 pandemic. By combining high quality national registers with short time delay and continuous repeated linkage updating, the project will bring timely and internationally unique data for epidemiological research on SARS-CoV-2. Broad research efforts are ongoing and further collaborative efforts are invited. Finally, our experience in establishing the project and database carries important lessons for building data repositories elsewhere to address urgent health problems.

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**Disclosure**

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