Cohort profile: the ChicagO Multiethnic Prevention and Surveillance Study (COMPASS)

Briseis Aschebrook-Kilfoy 1,2, Muhammad G Kibria,1,2 Farzana Jasmine,1,2 Liz Stepniak,1,2 Rajan Gopalakrishnan,3 Andrew Craver,2 Paul Zakin,2 Saira Tasmin,1,2 Karen E Kim,3,4 Kathleen H Goss,3 Marcy List,3 Michelle LeBeau,3 Habibul Ahsan1,2

ABSTRACT

Purpose The ChicagO Multiethnic Prevention and Surveillance Study or COMPASS is a population-based cohort study with a goal to examine the risk and determinants of cancer and chronic disease. COMPASS aims to address factors causing and/or exacerbating health disparities using a precision health approach by recruiting diverse participants in Chicago, with an emphasis on those historically underrepresented in biomedical research.

Participants Nearly 8000 participants have been recruited from 72 of the 77 Chicago community areas. Enrolment entails the completion of a 1-hour long survey, consenting for past and future medical records from all sources, the collection of clinical and physical measurement data and the on-site collection of biological samples including blood, urine and saliva. Indoor air monitoring data and stool samples are being collected from a subset of participants. On collection, all biological samples are processed and aliquoted within 24 hours before long-term storage and subsequent analysis.

Findings to date The cohort reported an average age of 53.7 years, while 80.5% identified as African-American, 5.7% as Hispanic and 47.8% as men. Over 50% reported earning less than US$15 000 yearly, 35% were obese and 47.8% were current smokers. Moreover, 38% self-reported having had a diagnosis of hypertension, while 66.4% were measured as hypertensive at enrolment.

Future plans We plan to expand recruitment up to 100 000 participants from the Chicago metropolitan area in the next decade using a hybrid community and clinic-based recruitment framework that incorporates data collection through mobile medical units. Follow-up data collection from current cohort members will include serum samples, as well as longitudinal health, lifestyle and behavioural assessment. We will supplement self-reported data with electronic medical records, expand the collection of biometrics and biosamples to facilitate increasing digital epidemiological study designs and link to state and/or national level databases to ascertain outcomes. The results and findings will inform potential opportunities for precision disease prevention and mitigation in Chicago and other urban areas with a diverse population.

INTRODUCTION

Despite the tremendous advances in medicine and public health in recent decades, many areas of the USA have seen a growing gap in health outcomes and health equity.1,2 As one of the most racially segregated cities in the USA,3 Chicago reports alarming health disparities and an unequal burden of cancer and chronic risk and mortality, particularly on the South Side.3 The South Side of Chicago encompasses roughly 60% of the city’s land area. The South Side is one of the nation’s largest contiguous urban African-American communities, and is home to a large percentage of the city’s 81 4500 African-American residents.4 The South Side also includes Chinatown and Bridgeport, the two neighbourhoods with the highest concentrations of Asian-Americans in Illinois, making Chicago the sixth largest Asian-American city in the USA. The most recent census also
illustrated that Hispanics now account for nearly one-third of the city’s total population of 2.7 million.4

Chicago faces significant urban challenges, including concentrated poverty, violence and crime, poor housing and living standards, food deserts and environmental exposures such as toxins in the air and water due to waste sites, landfills and abandoned industrial buildings.5–17 Despite advances in cancer and chronic disease prevention, screening, diagnosis and treatment that have improved chronic disease risk and outcomes for many Americans, disparities in cancer, cardiovascular disease, diabetes, asthma and maternal child health persist in Chicago.18 Disease-related mortality rates reflect continued disparity between non-Hispanic Blacks and Chicagoans of other races.19–22 Health disparities are increasingly understood to be a substantial burden on society in terms of healthcare costs, lost productivity and general societal well-being.23 24 According to the National Institute on Minority and Health Disparities, health disparities are multidimensional, complex phenomena that result from the interaction of multiple contributing factors over time.25 These disparities create trajectories of health vulnerability for certain groups of people that vary from the majority norm.26–28 Therefore, there is an urgent need to better understand the influence of factors that drive health disparities, such as healthcare coverage and low socioeconomic status. It is also important to identify the still-unknown factors behind health disparities in order to mitigate these trends and reduce the unequal burden of disease.

The University of Chicago is home to the Institute for Population and Precision Health (IPPH), as well as an National Cancer Institute (NCI)-designated comprehensive cancer centre. Given its location in the largest contiguous African-American community in the USA, its extensive community engagement infrastructure and cohort building experience and the availability of innovative and well-established population science resources, we are in a unique position to focus efforts on understudied populations and to study health disparities on Chicago’s South Side. To meet the need for a rigorous study of cancer and chronic disease and disparities in the Chicago metropolitan area, we established the ChicagO Multi-ethnic Prevention and Surveillance Study (COMPASS) in 2013. COMPASS is a population-based longitudinal study of cancer and chronic disease risk and progression with the overall goal of recruiting a diverse study population in Chicago. As those with lowest socioeconomic status and possibly the greatest vulnerability to poor health outcomes have historically been the least likely to enrol in biomedical and biobanking research,29 30 we have implemented a study design in which recruitment occurs in both the community and the clinic, and involves extensive engagement activities. Because of the growing recognition of health disparities in Chicago, COMPASS data collection efforts have focussed on potentially important factors for minority health and have emphasised recruitment from minority populations. COHORT DESCRIPTION

Purpose

The aims and primary scientific goals of COMPASS are to focus on the discovery of cancer and chronic disease in general, with the perspective that social, economic, racial and political marginalisation are often primary contributing factors associated with health disparities.31–41 The COMPASS scientific focus areas are illustrated in figure 1. The primary COMPASS aims driving the design are as follows:

1. To discover novel genetic and epigenetic factors related to the risk of cancers among the urban population through comprehensive yet cost-efficient genome-wide screening of germline DNA for single-nucleotide polymorphism and methylation variants. This will be accomplished by efficient typing of case-cohort samples using custom gene-chips based on all novel genetic and epigenetic variants identified through next-generation whole genome deep sequencing and methylation profiling of cohort samples supplemented with previously identified and validated variants from 1000 Genome Project. Similar screening of tumour tissues of cancer cases will also be done to distinguish between inherent and acquired genomic alterations.

2. To elucidate and characterise cancer risk defining germline genetic/epigenetic discoveries, we will prospectively screen baseline serum/plasma and RNA samples from representative samples of matched subcohorts of all four risk-groups (carriers with cancer, non-carriers with cancer, carriers without cancer and non-carriers without cancer) for proteins/hormones/enzymes and messenger RNAs/non-coding RNAs, respectively. Similar screening of tumour tissues of the two cancer groups will also be done to distinguish between inherent and acquired alterations to further elucidate the biology and identification of novel preventive and therapeutic approaches.

3. To integrate the complex interactions of the genomic determinants with non-genomic environmental, nutritional, behavioural and psychosocial determinants to develop a more complete understanding of cancer risk and causation. This includes consideration of the microbiome. This will be accomplished through tailored statistical and causal modelling approaches developed by statisticians and epidemiological methodologists at the University of Chicago.

4. To discover novel environmental (including air, water and household/workplace pollutants) and dietary (including food additives/toxicants and deleterious food items/nutrients) carcinogens that predispose urban populations to a higher risk for various cancers and chronic disease. These discoveries will be made through interdisciplinary efforts involving environmental sensors using nanotechnology and molecular dosimetry for high-throughput screening for biomarkers of exposure to carcinogens, coupled with sophisticated environmental, statistical and systems modelling.
5. To identify previously unrecognised factors related to urban lifestyle and behaviour (including stress, energy-balance/obesity, sleep and ageing) that increase the risk of various cancers and chronic diseases. These investigations will be conducted through integrated application of behavioural and psychosocial assessment tools, biomarkers of stress/ageing/obesity and novel multilevel and agent-based statistical modelling.

6. To consider new approaches to prevention that are personalised to one’s specific risk profile based on their host, environmental/behavioural and genomic profiles by exploiting new knowledge about the risk-driving inherent genomic variants, their interactions with environmental, host, behavioural and psychosocial dynamics, and the downstream molecular alterations in serum/plasma as well as tumour tissues that potentially mediate disease risk. COMPASS will function as a platform for focussed molecular, mechanistic and clinical research.

7. To specifically integrate the diverse data sources named here to investigate opportunities for prevention and early diagnosis, with an emphasis on the mitigation of health disparities.

The recruitment of this diverse cohort presents important opportunities for scientific discovery, disease mitigation, and the alleviation of the dramatic disparities observed in Chicago and other urban centres across the USA. According to the National Cancer Institute, overcoming cancer health disparities is one of the best opportunities for lessening the burden of cancer. COMPASS data and biosample collection leverages the advanced population science infrastructure at the University of Chicago and preserves the opportunity to use advanced analytical methods and technologies in the future. In the short-term, much will be learnt about the health of Chicago communities. In the long-term, the investigation of the aims above will offer insight into the relative contribution of key health drivers to decrease health inequalities, understand disease risk more generally and elucidate opportunities for prevention and mitigation.

Here, we present this prospective multiethnic cohort study, including a description of the cohort enrolled to date. We also outline the research resources offered by COMPASS and implications for concurrent ancillary studies, as well as future observational and intervention opportunities within the cohort.

**Study population**

As of 2019, there were an estimated 2.7 million people and 1.1 million households within the city limits of Chicago. According to a 2018 estimate, the Chicago metropolitan area is the third largest in the USA, with 9.9 million living in city and suburbs combined. More than half the population of the state of Illinois lives in the Chicago metropolitan area. The 2010 US Census shows the population density of the city itself at 11,842 people per square mile, making it one of the nation’s most densely populated areas.
cities. The median income for a household in the city as of 2018 was US$55,198, with 19.5% of persons in poverty. The goal of COMPASS is to recruit 100,000 persons, oversampling the majority minority population (ie, more than 45% African-Americans and 20% Hispanics).\(^4\)\(^3\)

COMPASS is a longitudinal cohort study including persons recruited using a population-based approach, a community-based recruitment approach and a hospital/clinic-based recruitment approach. The multiple recruitment modalities were considered in the early phase of this project in order to capitalise on the strengths and challenges of our Chicago context. As such, here we describe participant recruitment methods using a targeted or community partner approach, as well as at University of Chicago Medicine and on mobile medical units. Parallel data are collected from both community and clinic source populations. These data consist of survey responses, biospecimens including blood, urine, saliva and stool, clinical measures, electronic health records and environmental samples from the home.

To be eligible for participation in COMPASS, participants had to meet the following criteria:

1. Residence in the designated geographical area (census tract).
2. Age 35 and above at the time of contact (to enhance efficiency to obtain cancer outcomes).
3. Ability to complete the consent and interview in English or Spanish.
4. Willingness to provide blood, urine and saliva samples.
5. Provision of a social security number (for outcomes follow-up).

Targeted community-based recruitment

To explore the feasibility of a population-representative sample for COMPASS, we implemented a recruitment plan to target potentially eligible participants using a two-stage cluster sampling approach, first randomising at the census tract level and then identifying potentially eligible persons from within the tract using commercially available address lists. Field staff would then knock on doors and invite the targeted individuals to participate. Specifically, ‘cluster sampling’ involves dividing the specific population of interest into geographically distinct groups or clusters. Census tract was the primary sampling unit for COMPASS, as census tracts are designed to be homogeneous with respect to population characteristics. In the City of Chicago, there are 798 census tracts; 169 are identified as majority Hispanic, 239 as majority White, 290 as majority African-American and 6 as majority Asian. We identified 120 (15% of Chicago) study areas of interest using census tracts that would confer the racial/ethnic distribution desired for recruitment. We then used Stata module gsample\(^4\) to randomly select approximately 120 tracts (40 original census tracts to which 80 tracts have been added) and addresses within blocks. The number of tracts selected was based on an assumption of a 15% response rate and the assumption that 46% of residents would be 35 years or older. By sampling at the census tract level, study interviewers would recruit from limited geographical areas where many households could be contacted efficiently, minimising travel time and cost. In particular, we used a probability proportional to size sampling design as this design reduces sampling errors if characters of interest are related to population size. In the first stage, census tracts were randomly selected with probability proportional to the population of census blocks. In the second stage, we randomly chose households within the tract. As two locations within the same census block do not each contribute completely independent information (this is known as the ‘intra-cluster correlation’), we selected more clusters rather than more points within any cluster in order to improve precision while maximising study efficiency and minimising cost.

The commercially available address lists used for sampling are available from the US Postal Service’s Address Management System and demographic data is available for addresses from commercial sources. The address lists used for COMPASS contained all residential addresses along a mail delivery route, and excluded commercial addresses as well as PO Boxes. The commercial list of addresses provided data for households in the targeted census tracts on age and race and a telephone number (if available). This allowed us to weight the household based on the probability that they would meet eligibility criteria and would reflect the desired racial/ethnic distribution. We excluded all households that did not appear to have a resident 35 or older.

Engagement for targeted community-based recruitment

We used the list of addresses in the chosen census blocks to mail a postcard introducing the study and outlining the inclusion/exclusion criteria. The postcard explained that the resident has been selected to participate in COMPASS, what participation would involve, the time commitment, study eligibility criteria, compensation for participation and that multiple eligible household members are welcome to participate. The postcard also provided a time frame in which an interviewer would be requesting participation.

After the postcards were mailed to targeted households, the field interviewers went door-to-door on the targeted study census blocks based on a predetermined schedule to request participation. If a subject did not respond, the interviewer revisited the house within the following 2 weeks. If the interviewer received no response at the third visit, the household would be documented as a non-responder. By ‘dropping in’ on all targeted participant homes, a goal was to recruit as least some portion of those individuals at greatest risk of adverse health outcomes.

If the eligible person/s agreed to participate, they enrolled in the study, completed the interview and provided biospecimens and physical assessment data onsite in their home.

The results of the door-to-door recruitment response are presented by Press et al.\(^4\)\(^5\) Taken together, the team
achieved a much higher response rate in African-American communities than anticipated.

Untargeted community-based recruitment using mobile medical units
Community-based COMPASS recruitment is supported by two large mobile medical units (figure 2) and the community engagement infrastructure including partner community-based organisations. The mobile medical units have two fully equipped exam rooms with phlebotomy capabilities, a bathroom to facilitate urine collection, are WiFi-enabled and include CLIA (Clinical Laboratory Improvement Amendments) certified labs with processing/storage capabilities. All study procedures can be accommodated on-board the mobile medical units.

Untargeted community-based engagement
Untargeted community-based recruitment typically occurs both in conjunction with community partners, as well as without partners in communities targeted for participation. Examples of community partners include faith-based organisations, non-profit organisations, retail stores, civic groups, Chicago Park District facilities and Chicago Public Schools. These partners publicise COMPASS enrolment events and invite participation on days scheduled for mobile unit presence.

Hospital/clinic-based recruitment
Study population
In conjunction with ongoing epidemiologic efforts at University of Chicago Medicine (UCM) administered by the UChicago IPPH (University of Chicago IPPH), COMPASS participants are recruited in a UCM research clinic. All UCM patients and their guests aged 35 and older are eligible for participation in COMPASS. In the clinic context, eligibility is not restricted based on place of residence. As such, the eligible study population includes ~60,000 UCM patients per year, over 60% of whom are African-American. However, this population differs from the community-based sampling in some key ways, and will reflect a lower-risk population from the perspective of cancer health disparities.

Hospital/clinic-based engagement
In-person outreach in the medical centre is conducted by trained research assistants who approach persons in waiting areas and inform them about COMPASS. Those who meet the eligibility criteria are invited to participate. Digital outreach will be expanded in the near future to use email and text introduction. Participants will be able to self-schedule an enrolment and data collection visit at our IPPH research clinic by following a link in the email or text and completing a REDCap (ResearchElectronic Data Capture) scheduling form.

DATA COLLECTION
Overview
Daily enrolment of COMPASS participants was initiated in May of 2013.

Enrolment of cohort members
The COMPASS interview has been revised and is currently divided into three online modules. These modules are administered via the Qualtrics (Qualtrics, Provo, Utah) survey platform and have been optimised for mobile devices. During their visit to the UChicago research clinic, one of the mobile data collection units, or during community recruitment, participants complete the consent process and survey module #1 using their own mobile device or an IPPH-provided Apple iPad. Survey modules #2 and #3 can be completed onsite during the same visit, or the participant can elect to receive the survey links via email or SMS to complete offsite. COMPASS staff are available onsite to help participants navigate the online consent and survey modules, and participants who choose to complete survey modules #2 and #3 offsite are encouraged to reach out to staff via phone or email with any questions or to receive technical assistance.

Once the consent and 60 to 80 min survey is completed, the following clinical measurements are ascertained: anthropometry, blood pressure and other haemodynamic measures. Study staff measure weight and body fat percentage using a scale and height, hip and waist circumference using a measuring device, and they then measure blood pressure and other haemodynamic parameters using a device that monitors pulse (ie, Dyna-Pulse). An overview of the data collection is provided in table 1. In table 2, the key topics and measurements in the COMPASS questionnaire are outlined. We conducted an extensive review of questionnaires used in other large cancer studies, including the American Cancer Societies Cancer Prevention Study and National Cancer Institute cohorts (including PLCO (Prostate, Lung, Colorectal, and Ovarian Cancer Screening Trial), NIH-AARP and the Agricultural Health Study) in addition to the NHANES (National Health and Nutrition Examination Survey) questionnaires, such that we would be able to harmonise our data in cohort consortium collaboration and maximise our ability to enhance study power for key diseases...
and populations. Our data dictionary can be found at https://compass.uchicago.edu/.

The purpose of COMPASS anthropometry is to collect body measurements using standardised examination procedures and calibrated equipment. To ensure the collection of high-quality data, COMPASS staff members are trained to follow standardised examination protocols, calibrate equipment according to a prescribed schedule

Table 1  An overview of data collection

| Construct                     | Variable                                                                 |
|--------------------------------|--------------------------------------------------------------------------|
| Socioeconomic status          | Households, neighborhoods, education, employment, immigration, marriage, housing, incarceration, income, insurance, legal status, marital status, migration, occupation, poverty, race, residence, retirement, sex, socioeconomic status, status, wealth, work status, age, birth date, education level, marital status, occupation, place of birth, race/ethnicity, sex/gender, work status, age, birth date, education level, marital status, occupation, place of birth, race/ethnicity, sex/gender, work status, age, birth date, education level, marital status, occupation, place of birth, race/ethnicity, sex/gender, work status |
| Medical history               | Age at menarche; breast feeding; fertility interventions; hormonal birth control use; hormone replacement therapy; menopausal status; pregnancy history |
| Psychosocial                  | Alcohol use; cannabis use; e-cigarettes/vaping; heroin use; lifetime passive smoke exposure; physical activity; self-reported health; smoking cessation; tobacco use history |
| Lifestyle behaviours          | Alcohol use; cannabis use; e-cigarettes/vaping; heroin use; lifetime passive smoke exposure; physical activity; self-reported health; smoking cessation; tobacco use history |
| Follow-up data                | Address; email address; phone number; contact information of friends or family members; driver’s license number; social security number |
| Immune status                 | Allergies; blood transfusions; influenza immunisation history |
| Medication use                | Antihistamine use; anti-hypertensive use; H2 receptor blocker use; insulin use; non-steroidal anti-inflammatory drug use; opioid use; pain status; proton pump inhibitor use; sleep medication use; statin/cholesterol/lipid lowering drug use |
| Psychological status          | Anxiety; depression; stress |
| Recent medical care utilisation | Barriers to healthcare access; emergency room/urgent care utilisation; healthcare coverage type and duration; healthcare utilisation; screening activities; usual healthcare provider/location |
| Significant medical history   | Arthritis; asthma; autoimmune disorders; blood cholesterol; dental history; diabetes; family history of cancer; heart attack/myocardial infarction; hepatitis B; hepatitis C; HIV/AIDS; hypertension; number of children; number of siblings; other chronic diseases; personal history of cancer; sleep apnoea; sexually transmitted diseases; surgeries |
| Sociodemographic status       | Age; birth date; household income; household size; education level; marital status; occupation; place of birth; race/ethnicity; sex/gender; work status |
| Social environment            | Caregiver status; community social cohesion; night shift work; religion; self-reported patient experience/satisfaction; sleep habits; stress at home; stress at work |
| Blood pressure                | Resting blood pressure; resting heart rate |
| Body composition              | Height; hip circumference; waist circumference; weight |
| Environmental exposures       | Drinking water sources; lifetime proximity to point sources of pollution; residence during adolescence; self-report crime and violence concerns; self-report environmental concerns; work commute (current and historical) |
and method and precisely measure and record the survey data. Portable digital scales are used in the field on the mobile units by interviewers to measure weight, as well as percentage body fat and water. Body length measurements are made with waist circumference tape and retractable steel measurement tape.

The DynaPulse Pathway System measures central systolic, diastolic and mean arterial blood pressures, and the data are also transmitted to the DynaPulse Analysis Center online for an array of haemodynamic profiles/values. Briefly, DynaPulse, is a cuff-sphygmanometry-based method, which applies Pulse Dynamics waveform analysis principle and has demonstrated and validated its capability of deriving cardiac output simultaneously with blood pressure, mean arterial pressure, systemic vascular resistance, systemic vascular compliance, brachial artery compliance, distensibility and resistance (brachialartery compliance, brachialartery distensibility and brachialartery resistance), LV dP/dtmax (maximalleft ventricular pressure rise) and other haemodynamic parameters. These haemodynamic values, collected for a subset participants, obtained simultaneously allow physicians to correlate the dynamic changes of each parameter to evaluate the physiological conditions of a patient’s circulatory system.

Following survey data collection and clinical measurements, COMPASS staff members draws a blood sample. For each participant, the blood sample is collected in cryogenic barcode-labelled tubes as follows: (a) Lavender top EDTA vacutainer – 10mL; (b) Red and black top SST vacutainer – 10mL; (c) Green top Lithium Heparin vacutainer – 10mL; (d) Blue top Trace Element K2 EDTA vacutainer – 10mL; (e) Gold top SST vacutainer – 3.5mL; (f) Red top serum vacutainer – 5mL; and (g) cfDNA collection tube – 10mL. The vacutainers are inverted per protocol for proper mixing, ensuring a source of blood material for haematological, biochemical, hormonal, serological and other special tests that require high-quality DNA or RNA.

The blood collection approach was conceived to confer the ability to implement a variety of future analyses relevant for cancer, environmental and population and precision health research. Specifically, the lavender top collection can accommodate assays of whole blood DNA, PBMC (peripheral blood mononuclear cell) - RNA, lymphocyte markers, cytokines, cardiac/cardiovascular disease markers, metabolic panels, cancer panels and investigation of viral and bacterial load (PCR). The green top collection can accommodate assays including thyroid parameters, drug screening and other biochemical tests. The gold top collection accommodates a variety of hormone assays, measurement of lithium, iron, vitamin B12, folate, immunoglobulins, autoantibodies and antibiotic assays. The blue top tubes can accommodate a variety of environmental measurements, including the measurement of some metals.

Study participants are asked to provide a buccal cell sample, using an Oragene DNA self-collection kit, as well as a urine sample.

Participants are eligible to receive three US$25 Visa gift cards in recognition of their time. The first gift card is distributed when the participant completes the consent process, survey module #1, and contributes biospecimens and physical measurement data. The second gift card is given to the participant on completion of survey modules #2 and #3. Those who participate in optional home air quality monitoring receive the third gift card when IPPH staff retrieve the air quality devices from their home and the participant completes a short survey on characteristics of their home and environmental exposures.

Study participants are also asked to provide an optional at-home stool sample using a stool sample collection kit. This kit includes instructions to transfer a small swab from toilet paper to a vial. The participant then places the sample vial into a prepaid and addressed envelope and mails it to IPPH.

Biospecimens are stored in medical-grade refrigerators at the IPPH research clinic and aboard the mobile medical units, and are transferred to our laboratory on the University of Chicago campus within 4 to 6 hours of collection.

**Biorepository**

At the UChicago IPPH laboratories, the EDTA-mixed blood sample is centrifuged to separate out the plasma sample and aliquoted in multiple 0.5 mL screw-capped plastic tubes for preservation (to avoid repeated thaw and freeze cycles in future). All components of blood are stored at −80°C and urine samples are stored at −20°C with proper inventory. Freezers are connected to emergency power supply line (red line) and a central alarm system. The saliva samples collected using the Oragene DNA self-collection kit are processed by warming in a hot water bath and stored at −20°C in the IPPH laboratory. Stool samples are snap frozen and stored at −20°C.

**Environmental sample collection**

Study participants may be asked to allow IPPH staff to collect air quality data from their home using one or more air quality sensors. The sensors provide real-time measures of the presence of particulate matter (PM2.5 and PM10) in addition to other contaminants such as carbon monoxide, volatile organic compounds and radon. Air quality data are collected using the COMPASS best practices protocol. Study participants are provided directions to not disturb the sensors and contact the study team with issues. The IPPH staff member schedules a follow-up visit to retrieve the sensors.

Indoor air quality data will be integrated with ambient levels from the Array of Things (AoT) initiative (https://arrayofthings.github.io/), a joint effort by UChicago and Argonne National Laboratories and supported by the City of Chicago. AoT is currently collecting environmental information on air pollution in addition to light intensity, precipitation, air quality, heat, sound volume and foot traffic through decorative sensors on traffic poles throughout the City of Chicago. The result is a system that collects data about the city at the micro level, including fine-grained pollution levels in different
neighbourhoods. The diversity of pollutants measured, the number of monitors that will be installed across the city (>500) and the availability of a complimentary sensor for indoor measures, provides an unprecedented opportunity to more accurately assess exposure to air pollution in Chicago. AotT is placing devices in communities participating in COMPASS. This presents a significant opportunity to understand the role of air pollution in disease aetiology, as well as the potential that findings could inform interventions targeting at-risk Chicago communities.

Details on air quality in-home measurements, ambient concentration modelling and total exposure estimation can be found in accompanying manuscripts.47

Interested study participants may be referred to the Chicago Water Quality Study, an initiative of the City of Chicago Department of Water Management. Under this initiative, a water quality kit consisting of sample collection bottles and instructions is mailed to the participant. The participant collects and documents samples per instructions and contacts the Department of Water Management for sample pick up. During the consent process, participants are asked to allow COMPASS to retrieve water quality analysis results from the Department of Water Management.

Data management
A secure HIPAA-compliant database was created by the Information Technology Group at Dartmouth for the first 4 years of the project. Going forward, all study data is being collected and stored in REDCap and/or Qualtrics (Qualtrics, Provo, Utah). In addition, the University of Chicago Medicine Comprehensive Cancer Center (UCCCC) has created an address database to track the response rate as well as details of non-responders and those who opt not to participate. This database is maintained by the COMPASS informatics team.

Tracking health outcomes
All participants have provided their social security numbers so that their cancer and vital status can be tracked long-term through the Illinois State Cancer Registry, which is maintained by the Illinois Department of Public Health.

Electronic medical records
UChicago is a member of the Chicago Area Patient Centered Outcomes Research Network (CAPriCORN)—a platform that integrates EMR across the nine largest academic medical centres in Chicago. COMPASS participants provide consent for access to medical records, which will allow for the efficient collection of comprehensive health history data from EMRs of participants who receive care both at the University of Chicago Medical Centers and outside of our system. The collection of medical records will be completed continuously, when available.

Follow-up data and biosample collection
Previously enrolled participants will be re-contacted to schedule in-person visits to our IPPH research clinic and our mobile research units. They will be asked to sign an updated consent form and to complete an online follow-up survey in order to understand changes in health status, health behaviours and lifestyle. This visit will also include collection of follow-up biospecimens and physical measurements. Follow-up survey data collection will be requested annually and frequency of in-person biospecimen collection will be funding dependent. In addition, following COMPASS notification that a participant has had a cancer diagnosis, either from the participants themselves or the Illinois State Cancer Registry, we submit a request to the diagnosing and treatment institutions for medical records such that we may ascertain more detailed tumour characteristic and treatment information. We note in the consent that we plan to follow study participants for at least 10 years. In a recent effort to collect follow-up data, we were able to reach 25% of participants by phone. In the future, we will reach out via text and phone to enhance response rates.

Patient and public involvement
None of the participants was involved in the questionnaire design, biological measurements or outcome measures; they were likewise not involved in the design, recruitment and implementation of the study. Furthermore, all participants were informed of the use of the data for research in this study. There were no plans to disseminate the study results to participants. When genetic results become available, participants will be contacted and asked if they would like to receive them per the protocol/consent documents.

All study participants signed a consent form prior to enrolment.

Data analyses
Baseline data from the cohort were analysed for tabular and descriptive statistics (table 3). Means and SD were calculated for variables with a continuous distribution and proportions were calculated for variables with categorical distributions. Body mass index was computed based on weight and height measured by the interviewers. All analyses were performed using Stata 15 (StataCorp, College Station, Texas) and GeoDa for Windows.48

CHARACTERISTICS OF STUDY PARTICIPANTS RECRUITED TO DATE
Among the 7728 participants enrolled in the study (table 3), 80.5% identified as African-American, 5.7% as Hispanic and 47.8% as men. In addition, over 50% reported earning less than US$15 000 yearly, 15.2% reported working full-time, 35% were obese, 47.8% were current smokers and 38% reported having hypertension. High school graduates comprised 19.8% of the cohort, and 17% of the participants were married. Consistent
with findings by the Illinois Department of Public Health, the prevalence of diabetes was higher among women than men, and non-Whites compared with Whites. The overall prevalence of type 2 diabetes in this study was 10.1%.

The maps in figure 3 show the prevalence of the specified variable in the 77 community areas in Chicago. Only community areas with at least 20 observations were included, resulting in 45 communities being analysed. There is an alarming prevalence of high blood pressure, with the majority of the community areas sampled being classified as having stage 1 or 2 hypertension. In addition, the majority of community areas sample have a smoking prevalence of at least 50%.

**STRENGTHS AND LIMITATIONS**

COMPASS efficiently recruits individuals previously under-represented in biomedical research. The pilot analysis of the first 8000+ individuals demonstrates our capacity to recruit such a diverse, urban population. Our UCM research space coupled with our mobile research units allow our team to collect and process baseline and follow-up data on campus and in Chicago communities.

Another strength of the study is the prospective cohort design, which reduces possible recall bias and selection bias. Furthermore, the geographically defined cohort design offers a number of advantages for the study of health disparities. The principal reason that we are recruiting a geographical cohort (rather than an occupational cohort, for example) is because it allows for the exploration of an array of exposures potentially affecting health outcomes. In many other cohort designs, subjects are recruited given their unique exposure, such as nurses, farmers, miners and so on. However, the factors that disproportionately affect African-Americans’ health status and outcomes in Chicago are largely unknown or under-characterised. Similarly, the relevant risk factors in Hispanic and Asian populations are likely unique and remain under-characterised.

COMPASS is also involved in community engagement efforts to address the needs of the South Side community, through collaboration with the UCCCC Office of Community Engagement and Cancer Health Equity (OCECHE). The overall goals of the OCECHE are to: (1) establish strong partnerships with underserved communities and organisations; (2) support community and local capacity building; and (3) establish sustainable educational programmes. To ensure both academic and community relevance and to help shape future programming and direction, the OCECHE has both a Community and Internal Advisory Board comprising key community and institutional stakeholders.

| Demographics | Mean | % | N |
|--------------|------|---|---|
| Average age  | 53.7 |   | 7728 |
| % Male       | 47.8 |   | 3693 |
| % Married    | 17.4 |   | 1343 |
| % African-American | 80.5 |   | 6224 |
| % Hispanic  | 5.7  |   | 439  |

| Lifestyle factors | % | N |
|------------------|---|---|
| % Working full-time | 15.2 | 1174 |
| % High school graduate/GED | 26.3 | 2030 |
| % Bachelor's degree or higher | 8.0 | 615 |
| % Yearly income less than US$15,000 | 51.4 | 3975 |
| % Currently smoke marijuana | 20.2 | 1559 |
| % Currently smoke cigarettes | 47.8 | 3694 |

| Medical history (self-report) | % | N |
|-----------------------------|---|---|
| % History of cancer | 5.7 | 441 |
| % High cholesterol | 24.2 | 1874 |
| % Type 2 diabetes | 10.1 | 781 |
| % Hypertension | 38.0 | 2835 |
| % Health insurance | 76.6 | 5918 |
| % Heart attack | 4.7 | 367 |

| Observed values | % | N |
|-----------------|---|---|
| Average systolic BP (mm Hg) | 136.0 | 6646 |
| Average diastolic BP (mm Hg) | 84.0 | 6216 |

| Obesity | % | N |
|--------|---|---|
| Obese | 35.0 | 2704 |
In addition, the University of Chicago is a member of the CAPriCORN. CAPriCORN makes it possible for investigators at its partner institutions in Chicago to meet the growing need for the integration of EMRs in epidemiological research, including comprehensive multi-institution tracking/follow-up, such that comprehensive EMRs are available for participants who receive care at multiple institutions. The newly recruited participants will be enrolled through integrated healthcare systems, such as UChicago, which will allow for the efficient collection of serial biospecimens and tumour tissue as well as comprehensive health history data from EMRs. They will be followed longitudinally to study a broad range of risk factors, early markers and outcomes. We will be able to capitalise on the latest advances in the understanding of cancer aetiology, technology for data capture and analytical tools.

The collection of environmental data in COMPASS to understand the role of the environment in health disparities is supported by a new NIEHS (National Institute of Environmental Health Sciences)-sponsored environmental health centre in the region—ChicAgO Center for Health and Environment (CACHET). CACHET was established to address EHS (Environmental Health Sciences)-related health disparities as the first centre of its kind in the greater Chicago tri-state area (ie, Illinois, Indiana and Wisconsin). We have developed an environmental monitoring team (used for many NIEHS P30 investigations), infrastructure and expertise to explore the impact of urban environmental exposures that are embedded in COMPASS data collection. The CACHET mission is to elucidate the biological and social underpinnings between relevant urban environmental exposures and human disease and translate the findings to reduce health inequities within our communities. COMPASS is a key cohort synergised by the CACHET Integrated Health Sciences Facility Core at UChicago.

There are some limitations in our study. Thus far, there is limited variation in socioeconomic strata within race and ethnicity categories. Also, the longitudinal nature of the cohort makes it unlikely that we will be able to study cancer-related hypotheses for at least a decade. For rarer diseases, more people and time will be needed. In addition, the oversampling of African-Americans and individuals with low socioeconomic status calls into question the representativeness of the study population. However, given the paucity of cohort studies focused on this population, COMPASS provides a unique window into the factors impacting health in Chicago. There is also an oversampling of individuals out of work; however, this is possibly also a strength of the study, as some of the highest risk populations not represented in previous works and studies included.

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Contributors The study protocol was designed by HA, ML, BA-K, MLe, KHG and MGK and BA-K drafted the manuscript. BA-K and LS oversaw field operations and MGK and FJ oversaw the biosample processing and biobanking. HA, BA-K and KEK oversaw engagement. PZ, RG, AC and ST performed the data quality control and the statistical analysis. All authors participated in the study design, revised the article and approved the final version.

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Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, conduct, or reporting, or dissemination plans of this research.

Patient consent for publication Not required.

Ethics approval All study procedures and materials were reviewed and approved by the University of Chicago Biological Sciences Division Institutional Review Board Committee A (approval RB12-1660).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. COMPASS will continue to collect a rich set of data on multiple exposure domains and health outcomes. For more information, refer to the website: compass.uchicago.edu. Researchers interested in collaboration are invited to propose research questions based on the data available within COMPASS or to submit a request for additional data collection. Requests can be submitted electronically on the COMPASS website and will be reviewed by the COMPASS scientific board. The COMPASS study team is particularly interested in collaborations that will enhance research methods for this type of work, assess the impact of environmental exposure, highlight exposures of key significance in urban communities and address health issues of concern in Chicago and other urban centres.

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ORCID iD Brites Aschebrook-Kilfoy http://orcid.org/0000-0003-1918-7816

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