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Cohort profile: the Healthy Life in an Urban Setting (HELIUS) study in Amsterdam, The Netherlands

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ABSTRACT

Purpose Ethnic minority groups usually have a more unfavourable disease risk profile than the host population. In Europe, ethnic inequalities in health have been observed in relatively small studies, with limited possibilities to explore underlying causes. The aim of the Healthy Life in an Urban Setting (HELIUS) study is to investigate the causes of (the unequal burden of) diseases across ethnic groups, focusing on three disease categories: cardiovascular diseases, mental health and infectious diseases.

Participants The HELIUS study is a prospective cohort study among six large ethnic groups living in Amsterdam, the Netherlands. Between 2011 and 2015, a total 24 789 participants (aged 18–70 years) were included at baseline. Similar-sized samples of individuals of Dutch, African Surinamese, South-Asian Surinamese, Ghanaian, Turkish and Moroccan origin were included. Participants filled in an extensive questionnaire and underwent a physical examination that included the collection of biological samples (biobank).

Findings to date Data on physical, behavioural, psychosocial and biological risk factors, and also ethnicity-specific characteristics (eg, culture, migration history, ethnic identity, socioeconomic factors and discrimination) were collected, as were measures of health outcomes (cardiovascular, mental health and infections). The first results have confirmed large inequalities in health between ethnic groups, such as diabetes and depressive symptoms, and also early markers of disease such as arterial wave reflection and chronic kidney disease, which can only just partially be explained by inequalities in traditional risk factors, such as obesity and socioeconomic status. In addition, the first results provided important clues for targeting prevention and healthcare.

Future plans HELIUS will be used for further research on the underlying causes of ethnic differences in health. Follow-up data will be obtained by repeated measurements and by linkages with existing registries (eg, hospital data, pharmacy data and insurance data).

INTRODUCTION

Ethnic minority groups usually have a more unfavourable disease risk profile than the host population. For example, the prevalence of viral hepatitis B and C infections, as well as the death risks from almost all infections, are generally higher among ethnic minority groups. Depressive symptoms are more common in ethnic minority groups across Europe. Also, the higher cardiovascular risk among ethnic minorities is well documented, for example, the high prevalence of coronary heart disease among South Asians and the higher risk of stroke among people originating from Africa.

In Europe, ethnic inequalities in health have been observed in relatively small studies, with limited possibilities to explore underlying causes. In addition, most large-scale population-based studies excluded ethnic minorities, possibly due to the practical challenges to include these groups and because a homogeneous study population increases the internal validity of the study. The ‘Healthy Life in an Urban Setting’ (HELIUS) study was set up to fill this gap in epidemiological health research in Europe, aiming to provide a knowledge base for

Strengths and limitations of this study

► The main strength of our cohort is the inclusion of a large number of participants from several ethnic groups living in the same city, in whom an extensive set of questionnaires, biological samples and physical measurements were collected.

► Outcomes and risk factors were measured using the same methodology across all ethnic groups, including the majority population, which allows direct comparisons between the groups.

► Another major strength is the focus on three disease categories (cardiovascular diseases, mental health and infectious diseases), allowing us to investigate potential cross-links between them.

► Response rates were relatively low, possibly resulting in selection bias. Nevertheless, large numbers of each ethnic group were included, and all socioeconomic levels are represented in the samples; moreover, our non-response analyses show that socioeconomic differences between participants and non-participants were very small.
the improvement of healthcare and the prevention of communicable and non-communicable diseases in ethnic minority groups.7

The HELIUS study is designed as a prospective cohort study, including six ethnic groups (including the Dutch as a reference) living in Amsterdam, the Netherlands. The general objective of the HELIUS study is to study the causes of (the unequal burden of) diseases across these ethnic groups, with emphasis on three disease categories: cardiovascular diseases, mental health and infectious diseases.7 These are all major causes of global disease burden and mortality,8 9 and these disease categories are characterised by large ethnic variation, shared risk factors and complex patterns of causal relations to each other.10 11 The aim of this paper is to describe the cohort (design, participation, baseline study population and measurements) and illustrate some key findings to date. In addition, the strength and limitations of the cohort are described, as well as future plans and the collaboration policy of the study.

COHORT DESCRIPTION

Study design

Between January 2011 and December 2015, baseline HELIUS data were collected among Amsterdam residents of Dutch, Surinamese, Ghanaian, Turkish and Moroccan ethnic origin. More information on the migration backgrounds of the Dutch ethnic minority groups included in HELIUS can be found in table 1. Data were collected through a questionnaire (or interview) and a physical examination that included the collection of biological samples. All participants provided written informed consent.

Recruitment

People in the age range of 18–70 years were randomly sampled, stratified by ethnic origin, through the municipality register of Amsterdam. This register contains data on country of birth of citizens and of their parents, thus allowing for sampling based on the widely accepted Dutch standard indicator for ethnic origin. This country of birth indicator of ethnicity has the advantage of being objective and stable over time, and cross-validation studies showed a high correlation between the country of birth indicator and self-identified ethnic group indicator among Turkish, Moroccan and Surinamese people in the Netherlands.12 More specifically, a person was defined as of non-Dutch ethnic origin if he/she fulfilled one of two criteria: (1) he/she was born outside the Netherlands and has at least one parent born outside the Netherlands (first generation) or (2) he/she was born in the Netherlands but both parents were born outside the Netherlands (second generation). For the Dutch sample, we invited people who were born in the Netherlands and whose parents were born in the Netherlands. A limitation of the country of birth indicator for ethnicity is that people who are born in the same country might have a different ethnic background, which in the Dutch context is applicable to the Surinamese population (see table 1). Therefore, after data collection, participants of Surinamese ethnic origin were further classified according to self-reported ethnic origin (obtained by questionnaire) into ‘African’, ‘South-Asian’ or ‘other’.12

Selected individuals of Dutch, Surinamese, Ghanaian, Turkish and Moroccan ethnic origin received a written invitation combined with written information and a response card (in Dutch, and also in English for Ghanaian participants, in Turkish for Turkish participants and Moroccan Arabic for Moroccan participants). After a positive response, subjects received a confirmation letter of

Table 1 Information on the migration history of the ethnic minority groups included in the HELIUS study

| Ethnic group | Migration history |
|--------------|------------------|
| Surinamese   | The Surinamese migrated to the Netherlands from Suriname, a former Dutch colony in South America. Surinamese with an African background (referred to as ‘Creole’ in the Dutch context) are mainly the descendants of West Africans, and those with a South-Asian background (referred to as ‘Hindustani’ in the Dutch context) have their roots in North India. Both groups migrated to Suriname in the 19th century. Their migration from Suriname to the Netherlands was mainly due to the unstable political situation in Suriname in 1975 and 1980. Ethnic minority groups with comparable South Asian and African backgrounds can also be found in other European countries, including the UK. |
| Turks and Moroccans | Turks and Moroccans form important migrant groups in the Netherlands and in other West European countries (Belgium, France, Spain, Italy and Germany). Migration from Turkey and Morocco was encouraged in the 1960s and early 1970s to fill labour shortages in unskilled occupations. The initial period of labour migration was followed by a second period (1970–1980) in which many guest workers brought their spouses and children to the Netherlands. Since then, many young Turkish and Moroccan people have chosen partners from their region of origin. |
| Ghanaians | The migration of Ghanaians to the Netherlands occurred in two phases. The first phase (between 1974 and 1983) was due to economic reasons. The second phase (in the early 1990s) was linked to drought, political instability and the expulsion of Ghanaians from Nigeria. Ghanaians are also an important migrant group in the UK and Germany. |
Figure 1 Flow chart of recruitment for the HELIUS study. Participation rate: percentage of participants of those contacted; response rate: percentage of participants of those invited.

Figure 1 Flow chart of recruitment for the HELIUS study. Participation rate: percentage of participants of those contacted; response rate: percentage of participants of those invited.
Age and educational levels (self-reported highest level of education attained, either in the Netherlands or in the country of origin), within and across the ethnic groups. There were gender differences in educational level in some of the ethnic groups. Among Ghanaian, Turkish and Moroccan participants, the proportion of low education was higher in women than in men, whereas the proportion of medium-low education was lower (data not shown).

Differences between participants, non-participants and those not contacted

Among all ethnic groups, women were more likely to participate than men, as evidenced by the higher percentage of women among participants as compared with that of the total random samples (all invited) (table 3). Among those not contacted, the percentage of women was particularly low (except among Surinamese).

On average, those who participated were slightly older than those who refused to participate (except for the Dutch) or were not contacted. Those not contacted were the youngest among all ethnic groups. Figure 2 (upper panel) shows the response rates by age groups and ethnicity. The highest response was in the age group 45–54 years among the Ghanaian, Turkish and Moroccan groups, whereas in the Dutch and Surinamese groups, the highest response was among those aged 55–64 years.

The lower response rates in the Ghanaian, Turkish and Moroccan groups of older age might be due to language difficulties experienced, in particular, in these older groups. Among those contacted, the percentage of those agreeing to participate (participation rate) also differed slightly by age group across all ethnic groups (figure 2, lower panel).

To explore to what extent the response was related to socioeconomic status (SES), we constructed two SES indicators based on registry data at the level of the six-digit postcode maintained by the Department of Research and Statistics of the Municipality of Amsterdam: (1) the average property value of dwellings and (2) the percentage of residents living on a minimum income. This six-digit postcode area is the smallest geographical unit available. On average, these units are 50 by 50 m in size and include 10–20 households. Table 3 shows that both SES indicators are more favourable (ie, higher property value and lower percentage of residents living on minimum income) among participants as compared with non-participants. However, the differences are relatively small, particularly when compared with differences in these SES indicators across ethnic groups. The differences in SES indicators between the ethnic groups are in line with the ethnic differences in individual-level measured educational level as measured among participants (table 2), showing the highest educational level among Dutch (highest

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**Table 2** Demographic characteristics of the baseline HELIUS study population by ethnicity

|            | Dutch | South-Asian Surinamese | African Surinamese | Ghanaian | Turkish | Moroccan |
|------------|-------|------------------------|--------------------|----------|---------|----------|
| N          | 4671  | 3369                   | 4458               | 2735     | 4200    | 4502     |
| Age (years)| 46.1±14.1 | 45.1±13.5            | 47.6±12.8         | 44.0±11.7| 39.9±12.5| 39.7±13.1|

Age groups

- 18–29 years: 883 (18.9), 661 (19.6), 611 (13.7), 420 (15.4), 1139 (27.1), 1328 (29.5)
- 30–39 years: 824 (17.6), 500 (14.8), 602 (13.5), 477 (17.4), 925 (22.0), 1058 (23.5)
- 40–49 years: 956 (20.5), 870 (25.8), 1075 (24.1), 920 (33.6), 1214 (28.9), 1047 (23.3)
- 50–59 years: 1114 (23.8), 902 (26.8), 1508 (33.8), 811 (29.7), 739 (17.6), 783 (17.4)
- 60–70 years: 894 (19.1), 436 (12.9), 662 (14.8), 107 (3.9), 183 (4.4), 286 (6.4)

Sex

- Female: 2525 (54.1), 1809 (53.7), 2654 (59.5), 1671 (61.1), 2281 (54.3), 2786 (61.9)

Migration generation

- 1st generation: NA, 2545 (75.5), 3689 (82.8), 2582 (94.4), 2885 (68.7), 2998 (66.6)

Educational level*

- Low: 153 (3.3), 474 (14.1), 252 (5.7), 684 (28.0), 1260 (31.1), 1303 (30.4)
- Medium-low: 660 (14.3), 1120 (33.4), 1602 (36.2), 976 (40.0), 1008 (25.0), 782 (18.2)
- Medium-high: 1018 (22.1), 1003 (29.9), 1582 (35.8), 629 (25.8), 1174 (29.1), 1468 (34.2)
- High: 2784 (60.3), 753 (22.5), 986 (22.3), 152 (6.2), 586 (14.5), 739 (17.2)

Data are presented as mean±SD or n (%). Participants of unknown/other Surinamese (n=803), or unknown/other (n=51) ethnic origin are excluded from this table.

*Low=no schooling or elementary schooling only, medium-low=lower vocational schooling or lower secondary schooling, medium-high=intermediate vocational schooling or intermediate/higher secondary schooling, high=higher vocational schooling or university.

NA, not applicable.

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Table 3  Sex, age, and postal code-based socioeconomic status (SES) indicators among participants, non-participants and those not contacted by ethnicity

|                | Dutch | Surinamese* | Ghanaian | Turkish | Moroccan |
|----------------|-------|-------------|----------|---------|----------|
| **Sex (% women)** |       |             |          |         |          |
| All invited (random samples) | 50.0  | 54.9        | 52.9     | 47.7    | 49.3     |
| Participants | 54.1  | 57.3        | 61.1     | 54.3    | 61.9     |
| Non-participants | 55.5  | 52.1        | 48.6     | 49.9    | 59.7     |
| Not contacted | 44.3  | 55.1        | 48.3     | 42.8    | 38.9     |
| **Age (years)** |       |             |          |         |          |
| All invited (random samples) | 43.2±14.6 | 43.5±13.6 | 42.8±12.3 | 37.6±13.3 | 37.9±13.5 |
| Participants | 46.0±14.1 | 46.2±13.1 | 43.7±11.8 | 39.6±12.5 | 39.5±13.1 |
| Non-participants | 47.9±14.7 | 43.6±13.9 | 42.2±13.4 | 37.8±13.5 | 38.4±14.1 |
| Not contacted | 38.8±13.6 | 41.3±13.4 | 42.4±12.1 | 36.5±13.3 | 37.1±13.3 |
| **SES indicator (average property value, kEuro)** |       |             |          |         |          |
| All invited (random samples) | 264.9±139.0 | 192.3±70.7 | 154.3±47.0 | 194.9±61.6 | 192.8±60.3 |
| Participants | 273.6±139.6 | 196.2±72.0 | 151.9±43.9 | 194.2±58.1 | 195.2±56.9 |
| Non-participants | 258.8±136.7 | 191.9±69.0 | 150.9±43.0 | 193.0±61.4 | 194.5±61.9 |
| Not contacted | 261.3±139.4 | 189.5±70.8 | 158.1±50.9 | 196.4±63.4 | 191.0±60.8 |
| **SES indicator (% on minimum income)** |       |             |          |         |          |
| All invited (random samples) | 10.0 (2.7–20.7) | 22.0 (9.5–34.4) | 30.0 (18.5–39.4) | 25.0 (12.5–36.4) | 28.0 (17.1–38.5) |
| Participants | 9.1 (1.7–19.0) | 20.9 (7.8–33.3) | 30.8 (19.2–40.0) | 26.1 (13.3–37.0) | 28.6 (18.2–38.7) |
| Non-participants | 11.1 (2.9–22.7) | 21.8 (9.1–34.6) | 31.0 (18.9–40.9) | 26.3 (14.3–26.3) | 29.0 (17.6–39.1) |
| Not contacted | 10.2 (3.0–20.8) | 22.9 (10.9–35.0) | 28.6 (17.1–37.9) | 23.8 (11.1–35.3) | 27.3 (16.3–37.9) |

Data are presented as percentages, mean±SD or median (IQR).
*Non-response data were only available for the Surinamese sample as a whole, because municipality registers do not distinguish between Surinamese subgroups.

percentage of medium-high plus high education) and the lowest among Ghanaians (highest percentage of low plus medium-low education).

**Measurements of risk factors and health outcomes**

Table 4 provides an overview of the variables that were measured either by questionnaire or during the physical examination (more details on measurements are available from the authors on request). Risk factors include general factors such as physical, behavioural, psychosocial and biological factors, as well as ethnicity-specific characteristics such as culture, migration history, ethnic identity, socioeconomic factors and discrimination. In addition, more extensive measures were measured not in the total study population but in subsamples only. For example, dietary intake was estimated by an additional extensive food frequency questionnaire in a subsample of about 5200 participants. Table 5 lists the biological samples that were collected (and, for those who gave permission, also stored) and the laboratory measurements that are already available. Faeces microbiome data will be available from a subsample of about 6000 participants; this will provide a unique cohort to extend research on the role of gut microbiota composition in the development of diabetes, cardiovascular diseases and other health outcomes.

**Cross-cultural validity of measurements**

Whenever possible, we used standard validated questionnaires. For example, the Patient Health Questionnaire-9 (PHQ-9) was used to measure depressive symptoms, the 12-Item short form survey (SF-12) was used to measure quality of life and the short questionnaire to assess health-enhancing physical activity (SQUASH) was used to assess physical activity. However, because validity studies for these instruments were performed in the Dutch general population or in only a few ethnic groups in the Netherlands only, it was unknown whether they had similar validity and reliability for all the ethnic groups included in HELIUS. Validity studies within the HELIUS study indicated that the PHQ-9 and SF-12 indeed measure the same concepts in all ethnic groups and that there are no systematic differences in reporting between the groups. Also, the validity of self-reported physical activity with the SQUASH was similar across ethnic groups. However, consistent with findings in the literature, we observed low agreement between self-reported physical activity and objectively measured physical activity by accelerometer and heart rate monitor. In addition, low test–retest reliability was found for the SQUASH in all groups, implying no valid basis for the comparison of physical activity between the different ethnic groups.
FINDINGS TO DATE

A list of all publications that are based on data from the HELIUS study is available at this website: http://www.heliusstudy.nl/nl/researchers/publications. The first results have confirmed large inequalities in health between ethnic groups, such as diabetes and depressive symptoms, which can only partially be explained by inequalities in traditional risk factors, such as obesity and SES. In addition, the first results provided important clues for targeting prevention and healthcare. Below, we highlight some of these key findings emerging from the HELIUS study.

Ethnic inequalities in health

The prevalence of important cardiovascular disease risk factors varies largely between the ethnic groups. Figure 3A (upper panel) shows that, while the prevalence of diabetes in the Dutch group remains below 5%, the prevalence of diabetes ranges from 10% to 12% in participants of African Surinamese, Ghanaian, Turkish and Moroccan origin and is particularly high among the South-Asian Surinamese (20%). Ethnic minority groups also have a 1.3 (Moroccans) to 3.6 (Ghanaians) times higher prevalence of hypertension as compared with the Dutch groups, and hypertension prevalence is particularly high in the two groups of African origin (Ghanaians and African Surinamese). The higher prevalence of hypertension among the Turkish and Moroccan groups compared with the Dutch group suggests that risk patterns may be changing unfavourably over time because, just over a decade ago, these groups had a lower prevalence of hypertension compared with the Dutch. A large variation is also seen in depressive symptoms, with particularly high prevalence rates of depressed mood among the South-Asian Surinamese (19%), Turkish (23%) and Moroccan (21%) participants (figure 3B) (middle panel).

Overweight is one of the most important causes of increased cardiovascular risk and is also associated with depression. As expected, large differences in overweight and obesity were found between the ethnic groups (figure 3C) (lower panel). While 60% of the Dutch population is considered to have a normal weight based on their body mass index, this percentage is only about 30% in the ethnic minority groups. Particularly in the Ghanaian and Turkish groups, the prevalence of normal weight is very low, and the prevalence of obesity (extreme overweight) is extremely high, that is, up to 35% as compared with 10% in the Dutch. However, the ethnic inequalities in both hypertension and diabetes are not explained by differences in overweight, or other ‘traditional’ risk factors such as fat distribution, educational level or health behaviours. Ethnic inequalities were not only observed for cardiovascular risk factors, but also for early markers of cardiovascular disease or markers of end-organ damage, such as arterial wave reflection and chronic kidney disease; again, this could not be attributed to traditional cardiovascular disease risk factors. Our results suggest ethnic inequalities in the aetiology of
| Theme | Variables questionnaire | Explanatory factors | Outcomes | Physical examination |
|-------|-------------------------|---------------------|----------|----------------------|
| **General** | Demographic and ethnicity-specific factors: sex, age, marital status, household composition, country of birth of participant and his/her (grand)parents, self-reported ethnicity, migration history, educational level, occupational status, occupational level, religion, cultural distance (acculturation), ethnic identity, perceived ethnic discrimination (Everyday Discrimination Scale) Health-related behaviours: smoking, alcohol intake, cannabis use, physical activity (SQUASH questionnaire), weight perception, fruit intake, vegetarian diet, dietary pattern (breakfast, lunch and evening meal), coffee/tea intake, sugary drinks intake Healthcare use and related factors: subjectively measured health literacy (SBS-Q), compliance with medication, perceived quality of GP, healthcare use (GP, specialists, psychological care and alternative healthcare), healthcare use in country of origin Subsample*: dietary intake by extensive food frequency questionnaire (n~5200) | Perceived general health, quality of life (SF-12), list of 20 chronic conditions, functional limitations (in those aged >55 years) | Anthropometry (weight, height and circumferences of waist, hip, thigh, arm and calf) Body fat percentage (using bioelectrical impedance) Hand grip strength Current medication use Subsamples*: ► physical activity by Actiheart accelerometer and heart rate monitor, 5 days (n~500) ► objectively measured health literacy (REALM-D test) (n~9700) |
| **Cardiovascular health** | History of high blood pressure/hypercholesterolaemia/diabetes (including family history), family history of cardiovascular disease/sudden death, fainting history, age of menarche, age of menopause | Angina pectoris, possible myocardial infarction and intermittent claudication (by Rose questionnaire), self-reported and suspected myocardial infarction, self-reported and suspected cerebrovascular events | Blood pressure (sitting position, 5 min of rest, WatchBP Home, Microlife) Electrocardiogram (supine position, MAC 1600 System, GE Healthcare) Subsamples*: ► ankle-brachial blood pressure index (supine position, 7 min of rest, WatchBP Office ABI, Microlife) (n~14600) ► arterial stiffness by oscillometrically measured pulse wave velocity (supine, 10 min of rest, Arteriograph) (n~15000) ► non-invasive haemodynamics such as stroke volume, cardiac output, systemic vascular resistance (supine position, Nexfin) (n~14500) ► glycoalyx measurement (under tongue, Glycocheck) (n~6800) |
| **Mental health** | Perceived social support (DES subscale of SSQT with SSQS), childhood trauma, parental psychiatric history, mastery (Pearlin-Schooler Mastery Scale), neuroticism and extraversion (NEO Five Factor Inventory), stressful life events | Depressive symptoms (PHQ-9), nicotine use-related disorder (Fagerstrom), alcohol use-related disorder (AUDIT), cannabis use-related disorder (CUDIT) | |
| **Infectious diseases** | History and presence of allergy/asthma/rhinitis, family history of allergy/asthma, food allergy, urogenital infections, travel behaviour, use of self-tests, history of blood transfusions, history of surgery in other countries, injecting drug use, sexual behaviour, use of contraceptives (women), vaccination against human papilloma virus (women), circumcision (men) | Self-reported current respiratory symptoms Self-reported vaginal hygiene (women) | |

*In subsamples, we strived for equal numbers in each ethnic group.

ABI, ankle-brachial index; AUDIT, Alcohol Use Disorders Identification Test; BP, blood pressure; CUDIT, Cannabis Use Disorders Identification Test; DES, Daily Emotional Support subscale; GP, General practitioner; NEO, Neuroticism, Extraversion and Openness; PHQ-9, Patient Health Questionnaire-9; REALM-D, Rapid Estimate of Adult Literacy in Medicine in Dutch; SBS-Q, Chew’s Set of Brief Screening Questions; SF-12, 12-Item Short Form Survey; SQUASH, Short Questionnaire to Assess Health-enhancing physical activity; SSQS, Social Support Questionnaire for Satisfaction; SSQT, Social Support Questionnaire for Transactions;
Table 5 Overview of available laboratory measures and stored biological samples

| Type of sample                      | Laboratory measurements available                                                                 | Biological samples stored in biobank                                      |
|------------------------------------|-----------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------|
| Fasting blood (including DNA)      | ► glucose, HbA1c, haemoglobin, triglycerides, total cholesterol, HDL, LDL (calculated), creatinine  | ► citrate plasma (−80°C)                                                |
|                                    | ► D-dimer, fibrinogen, Lpa, ApoB, CRP (subsample n=6000)                                           | ► serum (−80°C)                                                         |
|                                    | ► cholesteryl ester fatty acids and carotenoids (subsample n=1000)                                 | ► heparin plasma (−80°C)                                                |
|                                    | ► metabolites (subsample n=500)                                                                   | ► EDTA-plasma (−80°C)                                                   |
|                                    | ► acylcarnitines, amino acids, sphingolipids (subsample n=700)                                     | ► whole blood (−80°C)                                                   |
|                                    | ► antibodies against human papillomavirus, human T-lymphotropic virus-1, Helicobacter pylori, Herpesvirus and Chlamidia trachomatis (subsample n=4680) | ► isolated DNA from pellets (4°C)                                      |
|                                    | ► angioplasma, CRP (subsample n=1200)                                                              |                                                                          |
|                                    | ► hepatitis B infection (anti-Hbc, anti-HBs, HBeAg, anti-HBe, HBV-DNA) and hepatitis C infection (anti-HCV, HCV RNA) (subsample n=2990) |                                                                          |
|                                    | ► whole genome SNP genotypes (GSA Illumina) (subsample n=12000)*                                   |                                                                          |
|                                    | ► citrate plasma (−80°C)                                                                          |                                                                          |
|                                    | ► serum (−80°C)                                                                                    |                                                                          |
|                                    | ► heparin plasma (−80°C)                                                                          |                                                                          |
|                                    | ► EDTA-plasma (−80°C)                                                                             |                                                                          |
|                                    | ► whole blood (−80°C)                                                                             |                                                                          |
|                                    | ► isolated DNA from pellets (4°C)                                                                  |                                                                          |
| Morning urine                      | ► microalbumin, creatinine                                                                        |                                                                          |
|                                    | ► urine dipstick: pH, glucose, ketones, leucocytes, nitrite, protein, erythrocytes                  |► urine (−80°C)                                                         |
| Faeces samples                     | ► faecal microbiome*                                                                               |                                                                          |
| (subsample, n=6000)                |                                                                                                   |                                                                          |
| Vaginal swabs                      | ► vaginal human papillomavirus (subsample n=600)                                                  |► vaginal swabs (−20°C)                                                 |
| (subsample, n=6000)                |                                                                                                   |                                                                          |
| Vaginal microbiome (subsample n=600)|                                                                                                   |                                                                          |
| Nasal and throat swabs             | ► respiratory viruses (subsample n=600)                                                            |► material (cells, mucus) in medium (−80°C)                              |
| (subsample, n=6600)                |                                                                                                   |                                                                          |

*Available in 2018.

ApoB, Apolipoprotein B; CRP, C reactive protein; GSA, global screening array; HbA1c, glycosylated hemoglobin A1C; HDL, high density lipoprotein; LDL, low density lipoprotein; Lpa, lipoprotein a; pH, potential of hydrogen; SNP, single nucleotide polymorphism.

these diseases and emphasise the importance of further research on the underlying causes of ethnic differences in cardiovascular health.

Within HELIUS, we also examined some risk factors that are less often investigated. For example, hand grip strength (a marker of muscle function) was found to be strongly related with diabetes, and large differences in hand grip strength were found between the ethnic groups (highest in Dutch and lowest in South-Asian Surinamese). In addition, short sleep duration (<7 hours per night) was more prevalent in the ethnic minority groups as compared with the Dutch, and short sleep duration was related to overweight, diabetes and hypertension. However, despite these strong relations, both short sleep duration and hand grip strength only marginally contributed to the large ethnic inequalities in cardiovascular risk factors. Chronic stress is also suggested to contribute to cardiovascular risk, and an important chronic stress factor specific for ethnic minority groups is the level of perceived ethnic discrimination (PED). In HELIUS, a positive association was found between PED and the clustering of traditional cardiovascular risk factors (metabolic syndrome) in some ethnic groups, with PED contributing about 5%–7% to the metabolic syndrome in Surinamese and Moroccans. PED was also related to depressive symptoms. In ethnic minority groups, PED is reported to account for about 25% of depressive symptoms.

Human papillomavirus (HPV) is an easily transmittable sexually transmitted infection, and persistent infections with high-risk HPV (hrHPV) types cause nearly all cases of cervical cancer. Higher incidences of cervical cancer (the fourth most common female cancer globally) have been observed among ethnic minority women as compared with women of Dutch origin. In HELIUS, the seroprevalence of hrHPV seroprevalence in blood, and the prevalence of vaginal high-risk hrHPV infection, differed between the ethnic groups. South-Asian Surinamese, Moroccan and Turkish women had an overall lower seroprevalence of hrHPV (22%, 14% and 15%, respectively) as compared with Dutch women (30%). Vaginal hrHPV prevalence was highest in the Dutch (42%) followed by the African Surinamese (32%), Turkish (29%), Ghanaian (26%), Moroccan (26%) and South Asian Surinamese women (18%). When adjusting
for sexual risk behaviour, the odds to be hrHPV positive were similar for all ethnic groups. However, because of the higher incidence of cervical cancer in ethnic minority women than in Dutch women. Future studies should explore other factors potentially responsible for the higher incidence of cervical cancer in these ethnic minorities (eg, differences in clearance/persistence of hrHPV infection or differences in participation in cervical cancer screening programmes).

**Prevention and healthcare**

Results of the earlier Surinamese in The Netherlands: Study on Ethnicity and Health (SUNSET) study in 2001–2003 showed that, at that time, Surinamese participants were less often aware of their high blood pressure and, therefore, were also less often treated, as compared with Dutch participants. This suggested a reduced access or quality of healthcare among Surinamese as compared with the Dutch, which could contribute to ethnic disparities in health. Our recent HELIUS results show that, as compared with the Dutch, the Surinamese groups (and the other ethnic minority groups) are currently even more often aware of their high blood pressure and are more often treated for hypertension.

These results suggest an improvement in hypertension management among ethnic minorities in Amsterdam in the last decade. However, despite medical treatment, ethnic minority groups suffering from hypertension still have lower rates of adequately controlled blood pressure levels. A similar pattern is observed for diabetes: although similar or even higher awareness and treatment levels are observed in ethnic minority groups compared with the Dutch, poor glycaemic control is still highly prevalent. As poor blood pressure control and poor glycaemic control are serious risk factors for cardiovascular diseases, it is important to investigate the causes of this poor control among the ethnic minority groups. One important factor to consider is the use of healthcare outside the Netherlands. In HELIUS, we observed that some ethnic minority groups use healthcare in their country of origin more often than other ethnic minority groups, ranging from 4% among Surinamese to over 20% among Turkish ethnic groups. We need to evaluate to what extent this influences their healthcare use in the Netherlands and their health.

Of all participants included in HELIUS, although 5% reported a history of cardiovascular disease (myocardial infarction, stroke and/or revascularisation), many of them did not have the risk factors under control; that is, 33% was still smoking, 76% was overweight, 55% did not achieve the norm for physical activity, 31% had uncontrolled blood pressure and 58% still had increased low-density lipoprotein (LDL)-cholesterol levels. These results show that risk factor control in secondary prevention is poor in the majority of individuals with cardiovascular diseases. The results suggest that secondary prevention should perhaps focus on different risk factors, depending on the ethnic origin of the individual patient.
STRENGTHS AND LIMITATIONS

The main strengths of our cohort are the inclusion of a large number of participants from several ethnic groups living in the same city and the collection of an extensive set of questionnaires, biological samples and physical measurements. Outcomes and risk factors are measured based on the same methodology across all ethnic groups, including the majority population.

Several supportive measures were taken to enhance the enrolment of ethnic minority groups. For example, we used ethnic-specific communication strategies, for example, working with faith communities (churches and mosques) and endorsement from local key figures. In addition, after invitation by mail, we visited participants at home, used translated questionnaires and had ethnically matched interviewers and research assistants to provide help during data collection. Despite these labour-intensive measures, response rates were relatively low, and this may have resulted in selection bias. However, we were able to include large numbers of each ethnic group in which all social-economic levels are represented; moreover, our non-response analyses show that socioeconomic differences between participants and non-participants were very small.

In HELIUS, the unique ethnic differentiation can be used to investigate which factors might explain ethnic inequalities in health, such as socioeconomic factors, culture, migration history, ethnic identity, discrimination and genetic factors. Furthermore, we are able to validate existing questionnaires (such as the SF-12 and PHQ-9) across these ethnic groups. The addition of more complex or specific measures performed among subsamples of the cohort (such as gut microbiome data) will provide additional unique data bases. Finally, a major strength is the focus on three disease categories that each are a major cause of the global burden of disease, thereby enabling studies on the potential crosslinks between these disease categories.

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Collaborators The HELIUS study has an open policy with regard to collaboration with other research groups and welcomes collaborations from a wide variety of disciplines. Information on procedures to acquire the available data and/or samples are described in the HELIUS Collaboration Policy, which can be found at www.heliusstudynl/nl/researchers/collaboration. In brief, to make use of the available data for research, we request a publication proposal describing background, aim, research questions, methods (analysis plan), and time table. All proposals should be submitted to HELIUS via the corresponding author of this article, or via info@heliusstudy.nl. The proposals are discussed in the HELIUS Executive Board regarding the study aims (compatibility with the general objectives of the HELIUS study/informed consent), the quality of the research proposal, and potential overlap with on going studies. After approval, the requested data will be provided after a Data Transfer Agreement has been signed. For more information, please visit our website or contact the corresponding author.

Contributors MBS, MP, EMD, RUPG, AHZ and KS participated in the conceptual design of the study, MS drafted the manuscript and performed the analyses. All authors critically revised the manuscript and contributed to interpretation of the data. All authors read and approved the final version of the manuscript.

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Competing interests None declared.

Patient consent Obtained.

Ethics approval The HELIUS study has been approved by the Ethical Review Board of the Academic Medical Center Amsterdam.

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