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The UCL Centre for Longitudinal Studies (CLS) is an Economic and Social Research Council (ESRC) Resource Centre based at the UCL Social Research Institute, University College London. It manages four internationally-renowned cohort studies: the 1958 National Child Development Study, the 1970 British Cohort Study, Next Steps, and the Millennium Cohort Study. For more information, visit www.cls.ucl.ac.uk.

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About the National Child Development Study

The National Child Development Study (NCDS) is a longitudinal birth cohort study, following a nationally representative sample of over 17,000 people born in Britain in a single week in March 1958.

Cohort members have been surveyed throughout their lives, since birth, creating an incredibly rich resource for a wide range of research. The study data show the very long roots of childhood, how past experiences can reverberate through the years, and the interplay between the different facets of people’s lives.

NCDS has equipped policymakers with robust evidence in areas as diverse as smoking in pregnancy, educational inequalities, adult basic skills, and social mobility. Today, with the cohort now in their sixties, the study is casting light on how people experience retirement and ageing in the 21st century.
1. Introduction

1.1 NCDS Surveys

The National Child Development Study (NCDS) started life as the Perinatal Mortality Survey, which was designed to examine the social and obstetric factors associated with stillbirth and infant mortality. In the first survey, data were collected about the births and families of just over 17,000 babies born in Great Britain during one week in March 1958.

NCDS is a continuing, multi-disciplinary longitudinal study. It began as the Perinatal Mortality Survey, which was designed to examine the social and obstetric factors associated with stillbirth and infant mortality. In this first survey, data were collected about the births and families of just over 17,000 babies born in Great Britain (England, Scotland and Wales) in one week in 1958, all of whom were eligible for comprehensive follow-up.

There have been nine surveys gathering information from respondents living in England, Scotland and Wales, in order to monitor their health, education, social and economic circumstances. These surveys were carried out in 1965 (age seven), 1969 (age eleven), 1974 (age sixteen), 1981 (age 23), 1991 (age 33), 1999/2000 (age 42), 2002-2004 (44) 2004/2005 (age 46), 2008/2009 (age 50) and 2013 (age 55). The 2002-2004 follow-up was a biomedical survey conducted by nurses. As part of the 1991 survey, information was additionally collected on the children of one in three cohort members; this included assessments of the NCDS 2013 Follow-Up – User Guide 8 behaviour and cognitive development of around 5,000 co-resident children. There have also been surveys of sub-samples of the cohort: most recently, in 1995 (age 37) a 10% representative subsample was assessed for difficulties with basic skills.

Data for NCDS have so far been collected from a number of different sources; the midwife present at birth, parents, schools, health service personnel, the cohort members themselves, their spouses, cohabitees and children. Data has been collected using a variety of methods: paper, electronic and self-completion questionnaires, clinical records, medical examinations, physical measurements,
ability tests, educational assessments and diaries. Most of the previous follow-ups have been carried out face-to-face visits to cohort members in their homes, but the age 46 survey in 2004 was conducted by telephone. The Centre for Longitudinal Studies (CLS) at the UCL Institute of Education (and formerly the Social Statistics Research Unit at City University), has been responsible for the study since 1985. CLS has been funded as an ESRC resource centre since 2004.

1.2 Introduction to the second edition of the 2002-2004 biomedical survey

The NCDS 2002-2004 biomedical survey was designed to obtain objective measures of ill-health and biomedical risk factors in order to address a wide range of specific hypotheses relating to anthropometry; cardiovascular, respiratory and allergic diseases; visual and hearing impairment; and mental ill-health.

This guide should be read in conjunction with the Technical Report produced by NatCen, which details the development of the survey, the conduct of the fieldwork and subsequent coding and processing of the biological samples.

This second version of the NCDS biomedical user guide contains information related to the NCDS Biomedical data available under End User Licence and the Special Licence. This is the result of the splitting of the original dataset, previously available under the UKDS Special Licence, into two datasets with different levels of sensitivity and disclosivity, which can therefore be made available under two different types of data access licences.

The majority of the NCDS 2002-4 biomedical data are now available under End User Licence (1079 variables), with the original data considered sensitive is still available under Special User Licence. This decision was the result of the Centre for Longitudinal Studies’ (CLS) disclosure assessment of each variable and the broad aim to make as much data available with the lowest possible barriers.

The Special Licence dataset contains 123 variables including questions on child adversity that were collected between 2002 and 2004, but not released in the previous NCDS42-4 Biomedical dataset. The Special Licence dataset also contains specific sensitive topics including: child adversity, CIS-R specific questions on mental health and questions which contains categories with small frequencies.
related to personal details such as skin colour, pregnancy, a surgical operation, specific height and unusual high number of children.

Information about the medication taken by the cohort members of the study is also deposit under End User Licence for the first time. These data were collected in 2002-4, but it was never released via the UKDS.

The number of cases in these datasets has changed from 9377 to 9293, as participants who did not provide consent to deposit their responses at the UK Data Service were excluded from the final dataset.
2. Development of the Biomedical Survey and Fieldwork

The survey content was proposed in detail in the MRC grant application. Practical development began in October 2001. Between then and the first pilot in March 2002 the project management team and the collaborators met on a regular basis. The content and order of the interview was determined, including the time allocated to each module, the exact measurements and questions to be included, the equipment that would be used, nurse protocols, and the way information would be recorded in CAPI (computer-assisted personal interviewing) and on paper. The issued sample was defined, and sample management procedures outlined. The CAPI was programmed and the documents were designed. Associated procedures were agreed, such as sample management, the packing and transport of equipment, the mailing of blood and saliva samples, and the exchange of information between the nurses, NatCen’s Operations Department and the participating laboratories. Three field pilots were carried out, two using convenience samples recruited from the general public, and the third using a sample of cohort members. These are described in full in the Technical Report.

2.1 Target Sample

As in previous waves of the NCDS, the target sample for the biomedical survey was all cohort members currently living in England, Scotland or Wales, excluding permanent refusals – 14,737 cohort members in August 2002. The sample definition was subsequently refined, and some cohort members excluded for various reasons, so that the sample issued to field (i.e. cohort members invited to take part in the study) comprised 12,037 cohort members.

2.2 Nurse briefings

Briefings for nurses began on 30th July 2002. Each briefing lasted three days and was attended by an average of seven nurses.

Day 1 was led by researchers from NatCen and CLS, and covered the background to the project; contact procedures; survey documents, including the ARF, consent
booklets and self-completion questionnaires; and the CAPI programme, including the CASI and the CIS-R interview.

Days 2 and 3 covered the measurement protocols in detail, including practical sessions, and were led by one of the principal investigators and the survey doctor, supported by NatCen nurse supervisors, and, on occasions, other survey collaborators.

Before leaving the briefing, each nurse carried out a dummy interview, including measurements, observed by members of the survey team and nurse supervisors. If there was any doubt about a nurse’s ability to carry out the interview satisfactorily, additional training and supervision was arranged. All nurses who were new to NatCen were supervised on their first interviews.

Nineteen briefings were held in London and six regional centres. The first briefing, of nurse supervisors, took place at the end of July 2002. Six briefings were held in August, five in September and two in October 2002, to enable fieldwork to begin promptly. Subsequent briefings were held by demand in order to ensure coverage throughout the fieldwork period, in November and December of 2002, and February, May and July of 2003. In total, 122 nurses were briefed to work on the project during its lifetime.

2.3 Issue of work

Fieldwork took place between 9th September 2002 and 26th March 2004. The sample was issued in fifteen monthly waves, starting in September 2002 and finishing in December 2003.20 It was divided in to 667 assignments or ‘points’, each a month’s workload for a nurse, clustered into geographically convenient groupings, with an average size of 18 addresses (see Chapter 2 of the Technical Report for further details).
3. Response

9378 cohort members were contacted and consented to the Biomedical Survey. One participant subsequently withdrew their consent and 9 interviews were lost. The dataset contains 9377 cases (the 9339 productive from the main survey and an additional 29 cases from the Dress Rehearsal.(Pilot 3) and the 9 lost interviews, for which blood was obtained).

84 participants of those 9377 cases did not provide consent for sharing the data and were excluded from the current data deposit, bringing the total number of cases to a total of 9293.

The full response is shown in the table below.

### Summary of cohort, issued sample and response to biomedical survey

<table>
<thead>
<tr>
<th>Valid</th>
<th>Frequency</th>
<th>Percent</th>
<th>Valid Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>101 Productive</td>
<td>9339</td>
<td>50.3</td>
<td>50.3</td>
</tr>
<tr>
<td>102 Productive - dress rehearsal</td>
<td>29</td>
<td>.2</td>
<td>.2</td>
</tr>
<tr>
<td>103 Lost productive - some data, no CAPI</td>
<td>9</td>
<td>.0</td>
<td>.0</td>
</tr>
<tr>
<td>201 Issued but ineligible - dead</td>
<td>28</td>
<td>.2</td>
<td>.2</td>
</tr>
<tr>
<td>202 Issued but ineligible - living outside GB</td>
<td>65</td>
<td>.4</td>
<td>.4</td>
</tr>
<tr>
<td>301 Refused</td>
<td>1803</td>
<td>9.7</td>
<td>9.7</td>
</tr>
<tr>
<td>302 Data withdrawn after interview</td>
<td>1</td>
<td>.0</td>
<td>.0</td>
</tr>
<tr>
<td>401 Invalid address</td>
<td>12</td>
<td>.1</td>
<td>.1</td>
</tr>
<tr>
<td>402 No contact made at address</td>
<td>191</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>403 Moved to unknown address</td>
<td>494</td>
<td>2.7</td>
<td>2.7</td>
</tr>
<tr>
<td>404 Unavailable for interview</td>
<td>86</td>
<td>.5</td>
<td>.5</td>
</tr>
<tr>
<td>405 Other reason for no interview</td>
<td>12</td>
<td>.1</td>
<td>.1</td>
</tr>
<tr>
<td>501 Ineligible: dead</td>
<td>1196</td>
<td>6.4</td>
<td>6.4</td>
</tr>
<tr>
<td>502 Ineligible - living outside GB</td>
<td>1236</td>
<td>6.7</td>
<td>6.7</td>
</tr>
<tr>
<td>601 Permanent refusals</td>
<td>1041</td>
<td>5.6</td>
<td>5.6</td>
</tr>
<tr>
<td>Valid</td>
<td>Frequency</td>
<td>Percent</td>
<td>Valid Percent</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-----------</td>
<td>---------</td>
<td>---------------</td>
</tr>
<tr>
<td>901 NCDS6 by proxy</td>
<td>31</td>
<td>.2</td>
<td>.2</td>
</tr>
<tr>
<td>902 Not issued for another reason</td>
<td>2506</td>
<td>13.5</td>
<td>13.5</td>
</tr>
<tr>
<td>903 Not in sample data base</td>
<td>479</td>
<td>2.6</td>
<td>2.6</td>
</tr>
<tr>
<td>Total</td>
<td>18558</td>
<td>100.0</td>
<td>100.0</td>
</tr>
</tbody>
</table>
4. The interview

4.1 CAPI main interview

The CAPI interview used in the main survey included the following elements.

- **Vision**: measures of near vision in right and left eyes (using appropriate visual correction), with and without pinhole viewer; stereo vision; distance vision (using appropriate visual correction).

- **Blood pressure and pulse**: three measures of systolic and diastolic blood pressure and resting pulse.

- **Prescription drugs**: all prescribed drugs taken, by name and BNF code.

- **Hearing**: thresholds of hearing in right and left ears at 1kHz and 4kHz.

- **Anthropometry**: standing height, sitting height, weight, waist circumference, hip circumference.

- **Lung function**: three measures (from up to five attempts) of forced vital capacity (FVC), forced expiratory volume (FEV1) and peak flow (PF).

- **Eye measurements using autorefractor**: sphere, cylinder and axis of right and left eyes.

- **Non-fasting blood sample**: four tubes filled and sent by nurses to laboratories in London, Newcastle and Bristol.

- **Computer Administered Self Completion (CASI)**:
  - AUDIT and supplementary questions about drinking alcohol
  - Adverse childhood experiences
CIS-R\(^1\) interview: modules covering appetite, fatigue, concentration and forgetfulness, sleep problems, irritability, depression, depressive ideas, anxiety, phobias, and panic.

4.2 Paper self-completion questionnaires

Two paper self-completion questionnaires covered the following topics.

**Questionnaire 1 (SC1) (‘yellow questionnaire’, completed in advance):**

- sun exposure
- physical activity connected with work
- hearing
- eyesight
- pain
- working conditions
- household circumstances
- social support.

**Questionnaire 2 (SC2) (‘lilac questionnaire’, completed during the interview):**

- general health and diet
- leisure exercise
- employment
- partnership status and children
- life events and
- (women only) contraception and HRT

Appendix A of the Technical Report includes the CAPI and self-completion questionnaires

---

4.3 Consents

Written consents were collected for:

- all measurements;
- questions about psychological health;
- collection, testing, storage and future use of blood;
- feedback of blood test results to the cohort member;
- feedback of measurements and results to the cohort member’s GP;
- analysis, storage and future use of saliva;
- deposit of data in the ESRC data archive;
- use of NHS administrative data.

4.4 Saliva sample

Cohort members were asked to collect and return two samples of saliva, accompanied by a short self-completion questionnaire.

Appendix B of the Technical Report includes information about the equipment used and nurse protocols.

Consent booklets and saliva forms can be found with other survey documents in Appendix C of the Technical Report.
5. Biochemical analyses of blood samples

Glycosylated haemoglobin
Glycosylated haemoglobin (hba1c) was measured on whole citrated blood by ion exchange high performance liquid chromatography, using the Tosoh A1c 2.2 Glycohemoglobin Analyser HLC-723GHb.

Triglycerides, total and HDL cholesterol
Triglycerides, total and HDL cholesterol variables (trig, chol, hdl, ldl) were measured on non-fasting serum by Olympus model AU640 autoanalyser.

LDL cholesterol was derived by the following formula:

\[ \text{ldl} = \text{Total cholesterol (chol)} - (\text{hdl} + (\text{trig}/2.2)) \]

Insulin-like growth factor
Insulin-like growth factor (igf1) was measured on serum by chemiluminescence-immunoassay.

Immunoglobulin E, House Dust Mite Allergen, Cat Allergen, Grass Pollen Allergen
Immunoglobulin E (ige) was measured on serum by the HYTEC enzyme immunoassay, with positive and negative controls. Total IgE was assayed on all specimens

Allergen-specific IgE to house dust mite (hdm), mixed grasses (grass), and cat fur (cat) were measured on specimens with a total IgE concentration above the median (30kU/L).

Fibrinogen
Fibrinogen (fib) was measured on citrated plasma by the Clauss method using a MDA 180 coagulometer

Tissue plasminogen activator antigen and von Willebrand factor antigen
Tissue plasminogen activator antigen (t-pa) and von Willebrand factor antigen (vWF) were measured on citrated plasma by enzyme-linked immunosorbent assays (ELISA) employing a double sandwich technique.
C-reactive protein

C-reactive protein (crp) contains data on C-reactive protein measured on citrated plasma by high-sensitivity nephelometric analysis of latex particles coated with CRP-monoclonal antibodies.

D-Dimer

Data on fibrin D-dimer was received from Dr A Rumley and Professor GDO Lowe (University Department of Medicine, Royal Infirmary) on 1st September 2006. The D-dimer assays were performed at the end of the field study using two different manufacturer kits. Within in manufacture kit quality control checks of the D-dimer assays were within set limits but there was a systematic difference between the two different kits (see Appendix 1).
6. Research data

6.1 Structure of the data

The files available under the UKDS are

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ncds_biomedical_2002_eul.sav</td>
<td>End User Licence data</td>
</tr>
<tr>
<td>ncds_biomedical_2002_sl.sav</td>
<td>Special Licence data</td>
</tr>
<tr>
<td>NCDS_biomedical_2002_coded_manikins.pdf</td>
<td>Diagram relating to Pain coding</td>
</tr>
<tr>
<td>ncds_biomedical_2002_capi_questionnaire.pdf</td>
<td>Questionnaire</td>
</tr>
<tr>
<td>NCDS_biomedical_2002_data_dictionary.xlsx</td>
<td>List of variables, topics and data collection instruments (EUL)</td>
</tr>
</tbody>
</table>

Variables labels are prefixed to indicate which part of the survey excluding the main CAPI interview they were collected in, as follows:

- CASI – Computer Aided Self Completion
- CIS-R – CASI – CIS-R Malaise questions
- SC1 – Self Completion Booklet 1 (variable label indicates the question number)
- SC2 – Self Completion Booklet 2 (variable label indicates the question number)
- LAB – Data relating to blood sampling, biochemical measures.

6.2 End User Licence Dataset

The End User Licence (EUL) dataset contains 1079 variables. It includes the majority of the data previously available under Special Licence, as well as CIS-R summary scores and data on taken medications that were previously collected in the Biomedical Survey 2002-4 but never made available via the UKDS before.
This EUL dataset contains modified variables generated as a de-identified version of variables previously available under Special Licence. The modified variables can be identify with the “_M” at the end of their name. The original variables have been kept in a dataset available under Special Licence.

A complete list of the variables is available at the UKDS website and can be downloaded with the dataset.

The contents of the EUL dataset appear in the following order:

**CAPI**
- Interview information
- Vision Tests
- Blood Pressure Tests
- Audiology
- Anthropometry: height, sitting height, weight, waist, hip
- Lung Function
- Eye Measurement of Refractive Error
- Blood Sampling
- Alcohol Audit (CASI)
- Mental Health Interview: CIS-R summary scores
- Medications
- Administrative and data consent variables

**Self completion 1**
A. Sun exposure
B. Physical Activity connected with work
C. Hearing
D. Eyesight
E. Pain
F. Work
G. Household circumstances
H. Social Life

**Self completion 2**
A. General Health and Diet
B. Leisure Activities
C. Work and Home Circumstances
D. Life Events
E. Women Only (contraception and HRT)

**Blood samples /Biomedical Assays**

- Blood results
- Saliva collection

**Derived variables**

### 6.3 Special Licence Dataset

The Special License dataset contains 122 biomedical variables, which can be divided into three different sections:

**Childhood Adversity (CASI)**

This battery of questions regarding adverse childhood conditions and upbringing (CHAD) was previously available via the UKDS (25 variables).

The summary variable adversity has been derived by Charlotte Clark at QMUL - see Appendix 2.

**Mental Health Interview: CIS-R questions**

The Clinical Interview Schedule-Revised (CIS-R) questions on appetite, fatigue, concentration and forgetfulness, sleeping problems and patterns, irritability, depression, depressive ideas, anxiety, phobias and panic (87 variables).

**Detailed personal characteristics**

A number of variables have been kept under Special Licence as they may contain potentially disclosive information such as height, skin colour, pregnancy, a surgical operation and number of children (10 variables).
### 6.4 Recoding of disclosive information for availability under End User Licence

A less disclosive version of the 10 variables potentially disclosive kept under Special Licence has been created in order to make this information available under End User Licence. The original variables names have been kept in the Special Licence dataset. The modified variables available under End User Licence have been renamed to have the “_m” suffix at the end of the name.

<table>
<thead>
<tr>
<th>Original Variable (Special Licence)</th>
<th>Recoded Variable (EUL)</th>
<th>Variable label</th>
<th>Amendment</th>
</tr>
</thead>
<tbody>
<tr>
<td>bheight</td>
<td>bheight_m</td>
<td>Height</td>
<td>Cap to 195 if values &gt;= 195&lt;br&gt;Cap to 145 if values &lt;= 145</td>
</tr>
<tr>
<td>estht</td>
<td>estht_m</td>
<td>Final measured or estimated height</td>
<td>Cap to 195 if values &gt;= 195&lt;br&gt;Cap to 145 if values &lt;= 145</td>
</tr>
<tr>
<td>skincol</td>
<td>skincol_m</td>
<td>Self-reported natural skin colour</td>
<td>Medium and dark skin combined</td>
</tr>
<tr>
<td>childnow</td>
<td>childnow_m</td>
<td>Number of children living with you, aged 18 or less</td>
<td>Cap to 6 for values &gt;= 6</td>
</tr>
<tr>
<td>childnum</td>
<td>childnum_m</td>
<td>Number of natural (biological) children</td>
<td>Cap to 7 if values &gt;= 7</td>
</tr>
<tr>
<td>hadop01a</td>
<td>hadop01a_m</td>
<td>Age at the time of removal of uterus and both ovaries</td>
<td>Cap to 20 if values &lt;= 20</td>
</tr>
<tr>
<td>dvhght44</td>
<td>dvhght44_m</td>
<td>Height at interview (age 44)</td>
<td>Cap to 194 if values &gt;= 194&lt;br&gt;Cap to 148 if values &lt;= 148</td>
</tr>
</tbody>
</table>
6.5 Data Coding and Editing

This section offers additional information to the coding and editing referred to in Section 7 of the Technical Report.

Missing values

Where no data entry in CAPI (e.g. because of routing) cells have been left empty. Not on route is coded –1 in data from the self-completion questionnaires. Codes -8, -9 were used to label cases that did not provide an answer or cases without applicable answers.

For variables where ‘refused’ is a listed option this has been retained and has not been recoded to merge with –9 entries. Where values are measures, no response recorded as 99, 999 etc (Note: some value ranges have negative values).

Refused consent/not attempted

Participants who did not provide consent to deposit their responses at the UK Data Service have been excluded, thus lowering the number of cases from 9377 to 9294.

These data have not been edited from the original CAPI, although the outcomes for particular measures may be inconsistent in cases where consent was withdrawn during measurement (e.g. blood pressure).

Inconsistencies, improbabilities

The edit has taken note of soft checks within CAPI. Within the self-completion booklets, the approach has been inclusive: that is, inconsistencies and improbabilities have not always been corrected: the policy on this was decided in consultation with the scientific collaborators.

Sex of cohort member as key variable

This is the “gold standard” variable for the sex of the cohort member, other variables which relate to the sex of the cohort member are left, to explain any consequent erroneous routing.

Lung function values
Values have not been cleaned, and there are no ‘not obtained’ or missing value codes (for FVC these have been left as 9.96, for FEV 9.95, for Peak Flow 995).

**Nurse number**

The nurse number (nurseno) has been recoded for reasons of anonymity.

**Retinomax codes**

Where the quality score has been recorded as 0 or 99, all measures for that eye have been recoded as not obtained. Measures with unsatisfactory quality scores of 1 to 6 have been left in the data.

**CIS-R summary scores**

The EUL dataset includes the summary scores for CIS-R modules:

- `totsum` CIS-R Total summary variable
- `overalla` CIS-R Overall stopped from doing things used to do in past 7 days
- `overallb` CIS-R Overall stopped from doing things more than once
- `overallc` CIS-R Overall things more difficult
- `ideasum` Depressive ideas summary score
- `depsum` Depresion summary score
- `pansum` Panic summary score
- `phobsum` Phobias summary score
- `anxsum` Anxiety summary score
- `fatsum` Fatigue summary score
- `irritsum` Irritability summary score
- `slpsum` Sleep summary score
- `forgsum` Concentration and forgetfulness summary score
Pain Coding

Variables [e3_acr1 to e3_acr10] and [e3_man1 to e3_man29] should be used in conjunction with the accompanying document coded_mannikins.pdf.

Lab Data Batch numbers

There are four different laboratory numbers, to be used as follows:

Newcastle batch number (nclbatch): for use with IgE and IGF-1 measurements

vWF batch number (vwfbatch): for use with vWF measurement

t-PA batch number (tpabatch): for use with t-PA measurement

Glasgow batch number (glabatch): for use with fibrinogen and CRP measurements

Blood Collection

For variables clorec and bldrec the values are (-3) "Old/Insufficient" (-2) "No blood received at lab" (-1.0) "Not recorded" (0) "No" (1) "Yes".

All variables with dates for blood reception at the lab: (bldday, bldmonth, bldyear, dayrnl, monthrnl and yearrnl) are set to missing if recorded as (-2) "No blood received at lab" or (-1) "Not recorded".

Consent given for storing blood samples

Variables consn, plstor, sestor refer to consent given for storing blood, plasma and serum. (-2.0) "No blood received at lab" (-1.0) "Not recorded" (0.0) "No" (1.0) "Yes".

Recoding of samples processed at Newcastle Laboratory

The result was re-coded to missing if the original lab value was coded as NSR (no sample received), NOSEP (no separation), INSUFF (insufficient), UNOLD (old sample - more than 5 days old), UNUS (atypical chromatographic pattern relating to HbA1c), RNCALB (LDL result not calculable), UNSUIT (for HDL calculation as triglyceride>13), LDLUS (for LDL calculation as HDL>4.5), HAEM (?), UNAVH, UNAVL, UNCALC or coded NR (for not required, but only if IgE >30).
<table>
<thead>
<tr>
<th>Variable</th>
<th>Recode</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>ld1</td>
<td>&quot;&lt;0.1&quot; re-coded to MISSING</td>
<td>ld1 &lt;0.1 - According to Newcastle this is the default (nonsense) value reported by the computer system for a calculated result when the calculation components are non-numeric.</td>
</tr>
<tr>
<td>ld1</td>
<td>If hdl measured and hdl&gt;4.5 then ld1 re-coded to MISSING</td>
<td>According to Newcastle LDL calculation invalid if HDL&gt;4.5</td>
</tr>
<tr>
<td>igf1</td>
<td>&quot;&lt;2&quot; re-coded to 0</td>
<td></td>
</tr>
<tr>
<td>ige</td>
<td>ige &quot;&gt;2000&quot; re-coded to 2000</td>
<td></td>
</tr>
<tr>
<td>hdm</td>
<td>&quot;&lt;0.35&quot; re-coded to 0</td>
<td></td>
</tr>
<tr>
<td>hdm</td>
<td>if Ige measured and Ige&lt;=30 re-coded to 0</td>
<td>According to Newcastle if IgE&lt;=30, house dust mite allergen, cat and grass allergen results are not required.</td>
</tr>
<tr>
<td>cat</td>
<td>&quot;&lt;0.35&quot; re-coded to 0</td>
<td></td>
</tr>
<tr>
<td>grass</td>
<td>&quot;&lt;0.35&quot; re-coded to 0</td>
<td></td>
</tr>
<tr>
<td>cat</td>
<td>if Ige measured and Ige&lt;=30 re-coded to 0</td>
<td>According to Newcastle if IgE&lt;=30, house dust mite allergen, cat and grass allergen results are not required.</td>
</tr>
<tr>
<td>grass</td>
<td>if Ige measured and Ige&lt;=30 re-coded to 0</td>
<td>According to Newcastle if IgE&lt;=30, house dust mite allergen, cat and grass allergen results are not required.</td>
</tr>
<tr>
<td>hdl</td>
<td>If triglyceride measured and triglyceride&gt;13 then hdl re-coded to MISSING</td>
<td>According to Newcastle blood unsuitable for HDL analysis if triglyceride&gt;13.</td>
</tr>
</tbody>
</table>
If there was a delay between sampling date and date received > 5 days (or dates missing) then all results set to MISSING. Analysis was considered not valid if a sample was more than 5 days old. (NB Delay could not be calculated for 9 study subjects as information on time blood taken missing).

**Removal of redundant blood pressure variables**

Variables sysres1, sysres2 and sysres3 were excluded in this latest deposit because they contain exactly the same data available in the variables sys, sys2 and sys3. These variables contain information about systolic readings from blood pressure.

Similarly the variable nobp was deleted because the same information related to no BP measure due to pregnancy was already included in the variable pregmes. This later variable contained a more detailed label and is available in the Special Licence dataset.

The variable bresps2 only contained value 1 for all cases with the label ‘some data’. This variable was considered redundant and removed from the new deposit.

**6.6 Coding of medication variables**

Variables drc1_m to drc19_m contain data of prescribed medication taken by the cohort member. The data was self-reported by the cohort members in open-end question text format and subsequently coded using 6-digits BNF codes.

There are up to 19 drugs per person in the EUL dataset. This detailed information is sensitive and could potentially lead to data disclosure. The structure of the BNF codes is shown below as an example:

<table>
<thead>
<tr>
<th>BNF Chapter</th>
<th>BNF Section</th>
<th>BNF Paragraph</th>
</tr>
</thead>
<tbody>
<tr>
<td>05</td>
<td>03</td>
<td>05</td>
</tr>
<tr>
<td>Infections</td>
<td>Antiviral Drugs</td>
<td>Macrolides</td>
</tr>
</tbody>
</table>
To reduce the data disclosure risk and deposit the variables under the End User Licence the BNF codes have been truncated to 4 digits. The first two digits refer to the BNF Chapter and the second two digits refers the BNF Section. In addition, categories with less than 10 cases were grouped together in a residual category ‘other’ for each of the chapter.

The set of variables [ytake11 - ytake50] denote the reason for taking the drug. A taken medication can have up to three reasons.

6.7 Derived Variables

The derived variables included in the EUL dataset are:

<table>
<thead>
<tr>
<th>Name</th>
<th>Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>smok42_1</td>
<td>Smoking at 42 codes 0 to 2</td>
</tr>
<tr>
<td>smok42_2</td>
<td>Smoking at 42 codes 0 to 6</td>
</tr>
<tr>
<td>sc0_1</td>
<td>Social Class at birth</td>
</tr>
<tr>
<td>sc42_1</td>
<td>Social Class at 42</td>
</tr>
<tr>
<td>reg0_58</td>
<td>Region at birth (based on pre-1974 regions)</td>
</tr>
<tr>
<td>reg46_58</td>
<td>Region at age 46 (based on pre-1974 regions)</td>
</tr>
<tr>
<td>reg46_04</td>
<td>Region at age 46 (based on post-1974 regions)</td>
</tr>
<tr>
<td>bresps0</td>
<td>Final outcome code</td>
</tr>
<tr>
<td>bresps1</td>
<td>Summary of cohort, issued sample and response</td>
</tr>
</tbody>
</table>
Appendix 1: D-Dimer Edits

Author: Alicja R Rudnicka, St George’s, University of London

Data on fibrin D-dimer was received from Dr A Rumley and Professor GDO Lowe (University Department of Medicine, Royal Infirmary) on 1 September 2006. The D-dimer assays were performed at the end of the field study using two different manufacturer kits. Within in manufacture kit quality control checks of the D-dimer assays were within set limits but there was a systematic difference between the two different kits.

Preliminary results of the difference between two manufacture kits used to measure D-dimer

D-dimer results from Kit (1)  D-dimer results from Kit (2)
<table>
<thead>
<tr>
<th></th>
<th>D- Dimer kit (1)</th>
<th>D-dimer kit(2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N=4609 (ng/mL)</td>
<td></td>
<td>N=3182 (ng/mL)</td>
</tr>
<tr>
<td>Median values</td>
<td>160</td>
<td>93</td>
</tr>
<tr>
<td>5th Centile</td>
<td>71</td>
<td>40</td>
</tr>
<tr>
<td>90th centile</td>
<td>325</td>
<td>197</td>
</tr>
</tbody>
</table>

There was a systematic difference between the two kits, with the second kit producing considerably lower readings.

Repeatability of repeat measures of D-dimer on same 1958 samples using two different manufacturer’s kits was assessed in 124 samples. The association between the two different kits is displayed graphically below. Figure 1a shows that on the arithmetic scale the difference between the two kits decreases as values of D-dimer increase. Figure 1b shows that the difference between the two kits is constant with log(d-dimer). There is therefore a constant proportional difference between kit (1) and kit(2). All analyses and adjustments were therefore performed using the log transformation of D-dimer.
Figure 1: Difference in d-dimer values of kit(2) minus kit(1) on the same samples plotted against average of D-dimer values from kit(1) and kit(2) on the arithmetic scale in (a) and logarithmic scale in (b).

On average D-dimer values from kit (2) were 40% lower than with kit (1). After discussions with Dr A Rumley and Professor GDO Lowe a decision was made to adjust the values from kit(2) to be in line with that on kit (1). All values from kit (2) were increased by 40%.
Appendix 2: Deriving Childhood Adversity

**Author:** Charlotte Clark, Barts School of Medicine & Dentistry, University of London

The scale used in the biomedical survey was taken from the Australian Path Through Life Study (Rosenman and Rodgers, 2004, 2006). This consists of 17 items which are retrospectively rated as having occurred or not (coded 0=no, 1=yes) by 16 years of age.

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Father or father figure was not affectionate towards me</td>
</tr>
<tr>
<td>2</td>
<td>Father or father figure suffered nervous or emotional trouble or depression</td>
</tr>
<tr>
<td>3</td>
<td>Father or father figure had trouble with drinking or other drug use</td>
</tr>
<tr>
<td>4</td>
<td>Mother or mother figure was not affectionate towards me</td>
</tr>
<tr>
<td>5</td>
<td>Mother or mother figure suffered nervous or emotional trouble or depression</td>
</tr>
<tr>
<td>6</td>
<td>Mother or mother figure had trouble with drinking or other drug use</td>
</tr>
<tr>
<td>7</td>
<td>There was much conflict and tension in the household whilst I was growing up</td>
</tr>
<tr>
<td>8</td>
<td>I was neglected</td>
</tr>
<tr>
<td>9</td>
<td>I had a strict, authoritarian or regimented childhood</td>
</tr>
<tr>
<td>10</td>
<td>I grew up in poverty or financial hardship</td>
</tr>
<tr>
<td>11</td>
<td>I was verbally abused by a parent</td>
</tr>
<tr>
<td>12</td>
<td>I suffered humiliation, ridicule, bullying or mental cruelty from a parent</td>
</tr>
<tr>
<td>13</td>
<td>I witnessed physical or sexual abuse of others in the family</td>
</tr>
<tr>
<td>14</td>
<td>I was physically abused by a parent – punched, kicked, hit or beaten with an object or needed medical treatment</td>
</tr>
<tr>
<td>15</td>
<td>I received too much physical punishment – hitting, smacking etc</td>
</tr>
<tr>
<td>16</td>
<td>I was sexually abused by a parent</td>
</tr>
</tbody>
</table>
Did your parents divorce or permanently separate when you were a child – not asked in Biomedical Survey

**NOTE**: in the survey, the question about parental divorce by age 16 was NOT asked, but details of divorce can be obtained from childhood measures, as well as data about 33y.

The derived variable adds up the total number of adversities reported – so the scale ranges from 0 to 16, as divorce is not included.

A total score has not been provided for those who were missing one or more item on the scale (coded as -8 in the variable).
