

# **HICDEP**

**Subject: Format Specification**

**Version: 1.90**

**Date: 2015-12-2**

## Table of Contents

<b>About HICDEP</b>	<b>5</b>
<b>Change log</b>	<b>6</b>
Draft Version 1.100	6
Version 1.90	6
Version 1.80	7
Version 1.70	8
Version 1.60	9
Version 1.50	9
Version 1.30	9
Version 1.25	9
Version 1.21	10
Version 1.2	10
Version 1.1	10
Version 1.00	10
Version 0.50 and 0.90	10
Version 0.38	10
<b>HICDEP 1.90</b>	<b>12</b>
General data format	12
Overview of data tables	12
Diagram	12
Structure of data	14
From flat files towards a normalized structure	14
Technical considerations	14
Coding Conventions	15
Date codes	15
Day unknown	15
Month and day unknown	15
Year unknown	15
Specification of precision	15
?ICD-10 codes	15
?ATC codes	15
Other codes	16
Quality Assurance	16
<b>tbIART - Antiretroviral treatment</b>	<b>17</b>
Core fields	17
Additional fields	17
QA Checks	17
<b>ART_ID field</b>	<b>19</b>
Coding Table	19
<b>ART_RS field</b>	<b>21</b>
Coding Table	21
<b>tbIBAS - Basic clinical, background and demographic information</b>	<b>23</b>
Core fields	23
Additional fields	23
QA Checks	24
<b>EDU_LVL field</b>	<b>25</b>
Coding Table	25
<b>ETHNIC field</b>	<b>26</b>
Coding Table	26
Limitations	26
<b>MODE field</b>	<b>27</b>
Coding Table	27
<b>ORIGIN field</b>	<b>28</b>
Coding Table	28
Region Codes	28
Country Codes	28
<b>tbICENTER - Center information</b>	<b>34</b>
Core fields	34
<b>tbICEP - Clinical Events and Procedures</b>	<b>35</b>
Core fields	35
Additional fields	35
QA Checks	35
<b>CEP_ID field</b>	<b>37</b>
Coding Table	37
Coding Table: Pregnancy-related adverse events	38
Coding Table: Early childhood-related adverse events	38
Case Definitions	38
<b>Case Definitions for AE_ID</b>	<b>39</b>
<b>CEP_SPEC field</b>	<b>41</b>
Coding Table	41
<b>tbIDELIVERY_CHILD - Delivery information related to the child(ren)</b>	<b>43</b>
Core fields	43

QA Checks	43
<b>tbIDELIVERY_MUM - Delivery information related to the mother</b>	<b>45</b>
Core fields	45
QA Checks	45
<b>tbIDIS - CDC-C diseases</b>	<b>47</b>
Core fields	47
Additional fields	47
QA Checks	47
<b>DIS_ID field</b>	<b>49</b>
Coding Table	49
Coding Table: Mother-to-child and paediatric specific	49
Case definitions	50
<b>DIS_WD field</b>	<b>53</b>
Coding Table	53
<b>tbILAB - Laboratory values</b>	<b>54</b>
Core fields	54
Additional fields	54
QA Checks	54
<b>LAB_ID field</b>	<b>56</b>
Coding Table	56
<b>LAB_U field</b>	<b>57</b>
Coding Table	57
<b>tbILAB_BP - Laboratory values - Blood pressure</b>	<b>58</b>
Core fields	58
QA Checks	58
<b>BP_U field</b>	<b>59</b>
Coding Table	59
<b>tbILAB_CD4 - Laboratory values</b>	<b>60</b>
Core fields	60
Additional fields	60
QA Checks	60
<b>tbILAB_RES - Resistance testing</b>	<b>61</b>
Core fields	61
Additional fields	61
QA Checks	61
<b>tbILAB_RES_LVL_1 - Nucleotide sequences (PRO, RT, GP41, GP120)</b>	<b>63</b>
Core fields	63
Additional fields	63
QA Checks	63
<b>tbILAB_RES_LVL_2 - Mutations</b>	<b>64</b>
Core fields	64
QA Checks	64
<b>tbILAB_RES_LVL_3 - Resistance test result</b>	<b>66</b>
Core fields	66
Additional fields	66
QA Checks	66
<b>tbILAB_RNA - Laboratory values</b>	<b>67</b>
Core fields	67
Additional fields	67
QA Checks	67
<b>RNA_T field</b>	<b>68</b>
Coding Table	68
<b>tbILAB_VIRO - Laboratory values - viro-/serology</b>	<b>69</b>
Core fields	69
Additional fields	69
QA Checks	69
<b>VS_ID field</b>	<b>71</b>
Coding Table	71
<b>VS_T field</b>	<b>72</b>
Coding Table	72
<b>VS_U field</b>	<b>73</b>
Coding Table	73
<b>tbILTFU - Death and drop-out</b>	<b>74</b>
Core fields	74
Additional fields	75
QA Checks	75
<b>DEATH_R1 field</b>	<b>77</b>
Coding Table	77
<b>DROP_RS field</b>	<b>81</b>
Coding Table	81
<b>tbIMED - Other medication</b>	<b>82</b>
Core fields	82
Additional fields	82
QA Checks	82

<b>MED_ID field</b>	<b>83</b>
Coding Table	83
<b>MED_RS field</b>	<b>86</b>
Coding Table	86
<b>tbINNEWBORN - Newborn</b>	<b>88</b>
Core fields	88
QA Checks	88
<b>tbINNEWBORN_ABNORM - Abnormalities</b>	<b>90</b>
Core fields	90
QA Checks	90
<b>tbIOverlap - Cross-cohort identification</b>	<b>91</b>
Core fields	91
QA Checks	91
<b>tbIPREG - Pregnancy</b>	<b>92</b>
Core fields	92
QA Checks	93
<b>Notes on the Pregnancy-Related Tables</b>	<b>94</b>
Example	94
Important relations	94
<b>tbIPREG_OBS - Obstetrical problems during pregnancy</b>	<b>95</b>
Core fields	95
QA Checks	95
<b>tbIPREG_OUT - Pregnancy outcome</b>	<b>96</b>
Core fields	96
QA Checks	96
<b>tbIPROGRAM - Program information</b>	<b>98</b>
Core fields	98
<b>tbIREFILL - Prescription refill data</b>	<b>99</b>
Core fields	99
QA Checks	99
<b>DRUG_ID field</b>	<b>100</b>
Coding Table	100
<b>tbISAMPLES - Blood Samples</b>	<b>101</b>
Core fields	101
Additional fields	101
QA Checks	101
<b>tbIVIS - Basic follow-up/visit related data</b>	<b>103</b>
Core fields	103
Additional fields	103
QA Checks	104
<b>CDC_STAGE field</b>	<b>106</b>
Coding Table	106
<b>CONTRACT field</b>	<b>107</b>
Coding Table	107
<b>EMPLOY field</b>	<b>108</b>
Coding Table	108
<b>Considerations for data management</b>	<b>109</b>
Uniqueness of PATIENT ID	109
Duplicate records	109
<b>Considerations for using the format to create a database</b>	<b>110</b>
Administrative fields	110
Further normalisation	110
Lookup tables	110
Performance	111
<b>QA checks</b>	<b>112</b>
Error Codes	112
QA checks listed nowhere else	112
All QA checks	113

## About HICDEP

HIV cohort collaborations have made substantial contributions to the knowledge of HIV epidemiology and management over the last years. So far, most collaborations have incorporated slightly different protocols for data exchange causing unnecessary workload for the people in charge of data extraction.

We were therefore asked to put together this draft consensus protocol for discussion at the 7th International Workshop on HIV Observational Databases, March 29th-30th 2003, Fiuggi, Italy. It is based on our experience with data-exchange protocols for [?D:A:D](#), the [?ART Cohort-Collaboration](#), the PLATO Collaboration and several previous studies on the safety of stopping OI prophylaxis.

This protocol is based on a relational structure (with some very minor deviations) and currently incorporates 26 data tables and numerous lookup-tables for the codes. It is evident that - depending on the questions addressed - only subsets of tables and fields will have to be extracted for data exchange.

We have not elaborated on database systems (e.g. SQL-Server, Oracle, Access) and their respective file formats as there are excellent tools for transferring data between most of the popular packages (e.g. [?StatTransfer](#)). The suggested data structure should work with most formats and software packages.

Please keep in mind that the primary purpose of this document is to provide you with formats for data-exchange but not for an operational database used for data-management on a day-to-day basis. Some considerations with that respect can be found in the appendices.

We plan to update this document on a regular basis and the most recent versions will be made available on the [HICDEP website](#). HICDEP is a format under constant improvement (currently under the EuroCoord Project, Grant Agreement No. 260694) and additions are made almost every year. Please refer to the [ChangeLog](#) for the most current updates and always use the tables available on the HICDEP website for most current coding lists for ART and MED drugs.

Members of EuroCoord WP4, September 2011

## Change log

### Draft Version 1.100

### Version 1.90

#### [tblART](#)

- added new optional field [GENERIC](#)
- added new optional field [ART\\_RS2](#)
- added new optional field [ART\\_RS3](#)
- added new optional field [ART\\_RS4](#)

#### [ART\\_ID](#)

- Added new single tablet formulations:
  - J05AR15 (Atazanavir and cobicistat)
  - J05AR16 (Lamivudine and raltegravir)
  - J05AR17 (Emtricitabine and tenofovir alafenamide)
  - J05AR18 (Emtricitabine, tenofovir alafenamide, elvitegravir and cobicistat)
  - J05AR19 (Emtricitabine, tenofovir alafenamide and rilpivirine)
- Deleted (moved to tblMED):
  - J05AF08;"Adefovir (PREVEON)"
  - J05AF10;"Entecavir"
  - J05AF11;"Telbivudine"

#### [ART\\_RS](#)

- added new (sub-)codes for stopping medication
  - 11 (Bone toxicity)
  - 15 (Social contra-indication)
  - 16 (Contra-indication unspecified)
  - 16.8 (Contra-indication expired)
  - 16.9 (Contra-indication - other)
  - 17 (MTCT regimen completed)
  - 70 (Pregnancy - toxicity concerns (during pregnancy))
  - 75 (Pregnancy - switch to a more appropriate regimen for PMTCT)
  - 92.5 (Regular treatment termination (e.g. DAA's for HCV, antibiotics))

#### [tblBAS](#)

- Added field for Center [CENTER](#)

#### [tblCENTER](#)

- Added table

#### [tblLAB](#)

##### [LAB\\_U](#)

- Deleted 18 (µkat/L, was same as 11)

#### [tblTFU](#)

##### [DEATH\\_R1](#)

- added new subcodes:
  - 03.1.3 (HCV with liver cancer)
  - 03.2.3 (HBV with liver cancer)

#### [tblMED](#)

##### [MED\\_ID](#)

- Added new drug codes:
  - A11CC (vitamin D)
  - G02CA (Tocolysis)
  - H02 (Corticosteroids)
  - J01 (Antibiotics)
  - J01GA01 (streptomycin)
  - J02AC05 (Isavuconazole)
  - J02AX04 (caspofungin)
  - J04AB05 (Rifapentine (Priftin))
  - J05AF12 (Clevudine)
  - J05AX GRAZ-ELB (Grazoprevir/Elbasvir)
  - J05AR-DAAS (Daclatasvir/Asunaprevir)
  - J07BM0 (HPV Vaccine)
  - J07BM01 (HPV Vaccine (types 6, 11, 16, 18))
  - J07BM02 (HPV Vaccine (types 16, 18))
  - J07BM03 (HPV Vaccine (types 6, 11, 16, 18, 31, 33, 45, 52, 58))
  - M05BA (bisphosphonate)
  - N05A (Antipsychotics)
  - N05CD (Benzodiazepine derivatives)
  - N05CF (Benzodiazepine related drugs)
  - N06A (Antidepressant)
  - N07BC (Other drugs used in opioid dependence)
  - N07BC01 (Buprenorphine)
  - N07BC02 (Methadone)
  - N07BC03 (Levacetylmethadol)

- N07BC04 (Lofexidine)
- N07BC51 (Buprenorphine, combinations)
- V03AB15 (Naloxone)
- Added new drug codes (moved from tblART):
  - J05AF08;"Adefovir (PREVEON)"
  - J05AF10;"Entecavir"
  - J05AF11;"Telbivudine"
- Changed temporary drug names to ATC codes:
  - J05AR-A450OM (ABT-450/r/Ombitasvir) --> J05AX67 (Ombitasvir, paritaprevir(ABT-450) and ritonavir)
  - J05AR-LESO (Ledipasvir/Sofosbuvir) --> J05AX65 (Ledipasvir/Sofosbuvir)
  - J05AX-DBV (Dasabuvir) --> J05AX16 (Dasabuvir)

#### MED\_RS

- added new (sub-)codes for stopping medication (to keep identical to ART\_RS)
  - 11 (Bone toxicity)
  - 15 (Social contra-indication)
  - 16 (Contra-indication unspecified)
  - 16.8 (Contra-indication expired)
  - 16.9 (Contra-indication - other)
  - 17 (MTCT regimen completed)
  - 70 (Pregnancy - toxicity concerns (during pregnancy))
  - 75 (Pregnancy - switch to a more appropriate regimen for PMTCT)
  - 92.5 (Regular treatment termination (e.g. DAA's for HCV, antibiotics))

#### tblPROGRAM

- Added table

#### tblIVIS

- Added field for Center CENTER
- Added field for family history of CVD/Stroke FAM\_Y
- Added field for type of clinic/service CLIN\_TYPE
- Added field for speciality of physician SPEC\_TYPE
- Added field for stage of transition from adolescence to adulthood TRANS\_STAGE

### Version 1.80

#### tblART

##### ART\_ID

- Added:
  - J05AR11 (Lamivudine, tenofovir disoproxil and efavirenz)
  - J05AR12 (Lamivudine and tenofovir disoproxil)
  - J05AR13 (Lamivudine, abacavir and dolutegravir)
  - J05AR14 (Darunavir and cobicistat)
  - J05AX-CAB (Cabotegravir (GSK-744))

#### tblBAS

##### ETHNIC

- Revised and added codes for different ethnicities
- Changed limitations
- Changed description

##### ORIGIN

- Changed description to be more specific

#### tblCANC

- Added table

#### tblITFU

##### DEATH\_R1

- Added sub-types for 22 and 23
- Added additional codes table

#### tblMED

##### MED\_ID

- Added:
  - J05AE11 (Telaprevir (INCIVEK, INCIVO))
  - J05AE13 (Faldaprevir)
  - J05AE14 (Simeprevir)
  - J05AE15 (Asunaprevir)
  - J05AX14 (Daclatasvir)
  - J05AX15 (Sofosbuvir)
  - J05AR-A450OM (ABT-450/r/Ombitasvir)
  - J05AR-DAAS (Daclatasvir/Asunaprevir)
  - J05AR-LESO (Ledipasvir/Sofosbuvir)
  - J05AX-DBV (Dasabuvir)

#### tblLAB

##### LAB\_U

- Added:
  - 14 (mg/24h)
  - 15 (mg/mmol)
  - 16 (fl)

- 17 (µg/mL = mg/L)
- 18 (µkat/L)

#### LAB\_ID

- Added:
  - ACRA (Albumin Creatinin Ratio)
  - PCRA (Protein Creatinin Ratio)
  - PROT (Protein)
  - PSA (Prostate-specific antigen)
  - PTH (Parathyroid Hormone)

## Version 1.70

- tblAE has been renamed to tblCEP

#### tblCEP

- Added field CEP\_V
- CEP\_ID
  - Added: HOSP, ICU, JAUN, LIVD, LIVT, USAB, ASP, BART, CHAG, NOCA, PCE, PMAR, REQU.
  - Added: BACT, ENDO, MENI, OSTI, PERI, PNEU, PYEL, LEIS, MCDI (Previously in tblDIS).
  - Removed: ANG, BYP, END as they are already in the field CEP\_SPEC.
  - Removed COR as now specified with SUD\_DEATH\_Y and EXP\_DEATH\_Y in tblLTFU.
- CEP\_SPEC
  - Changed description of NADM - CERV to contain "grade 2 or higher"
  - Added values: LIVB - XX, HEP - XX, NADM - HENE, NADM - BRAIN

#### tblBAS

- Added field RECART\_D (Date ART started)
- Added field LTART\_D (Date last assessed for ART)
- ETHNIC: Added Limitations section in description
- Added field EDU\_LVL (Last completed education Level)
- Added field HIV\_POS\_D (Date of first positive HIV test)

#### tblDIS

- DIS\_ID: Removed HG, BACT, ENDO, MENI, OSTI, PERI, PNEU, PYEL, LEIS, MCDI (now in tblCEP).

#### tblVIS

- Added optional field EMPLOY (What is the patient's current situation regarding labour?)
- Added optional field CONTRACT (If the patient is an employee, what is the type of the patient's employment contract?)
- Added optional field SMOKING\_Y (Is the patient currently a smoker?)
- Added optional field PREG\_Y (Is the patient currently pregnant?)
- Added optional field CDC\_STAGE
- Added optional field WHO\_STAGE

#### tblLTFU

- Added optional fields for ICD9 cause of death coding (ICD9\_#)
- Added optional field DEATH\_SOURCE
- Added optional fields SUD\_DEATH\_Y and EXP\_DEATH\_Y. Replacing COR
- CoDe
  - Typos in the coding table (HBF -> HBV)
  - New codes for cancer (04.XX)
  - New codes for AMI (08.XX)
  - New code: 92.1 - Unknown, competing risks

#### tblART

- ART\_ID: Coding extended/updated to match current ATC codes.
  - Lopinavir / Ritonavir changed code from "J05AE06" to "J05AR10".
  - Elvitegravir changed code from "J05AX-EVG" to "J05AX11"
  - J05AR07 - Triomune
  - J05AR08 - Eviplera / Complera
  - J05AR09 - Stribild
  - J05AR10 - Kaletra / Aluvia
  - J05AX12 - Dolutegravir
  - V03AX03 - Cobicistat
- ART\_FR: Added 0.33 and 0.5. Added special value -1 for "frequency unknown".
- ART\_RS
  - Added new codes: 6.X, 92.31, 92.32, 92.33, 92.4, 92.9, 94.2, 96.1, 96.2, 97.1, 97.2, 97.6

#### tblLAB

##### LAB\_ID

- Removed redundant "APT" code. Use code "ALP"
- Removed "GLYCE" (synonymous to Glucose in blood)
- Added "AFP" (Alfa Fetoprotein) and "DIPP" (Dipstick result for protein in Urine).

##### LAB\_U

- Removed WBC codes (WBC was already removed from LAB\_ID)
- Added generic "13 - µg/L" and "99 - No units" code

##### LAB\_ST

- Added codes U and U24.

#### tblLAB\_RES

- SUBTYPE Modified description to also include HCV

#### tblLAB\_RNA



RNA\_T

- New code: 42 - Abbott RealTime HIV-1 m2000
- New code: 59 - Monitor unspecified

tbIMEDMED\_ID

- Changed description from "Other HIV-related drugs" to "Other medication"
- Added codes for Boceprevir and Telaprevir

MED\_RS

- Same changes as in tbiART ART\_RS

tbILAB\_VIRO

- VS\_T New codes 6,7,8.

**Version 1.60**tbIBAS:

- The ORIGIN field is now coded using UN region and country codes.
- Added optional SEROHOW field indicating how the seroconversion date was determined.
- Added optional CENS\_D field holding the last date the database was updated for a patient.
- tbIREFILL created to hold prescription refill data.
- tbILTFU: The DEATH\_R# fields now use CoDe codes for coding cause of death.

tbILAB:

- Added codes for Gamma-glutamyltransferase and Prothrombin rate.

tbILAB\_VIRO:

- Changed code for "Other" of field VS\_T to value 9.
- Added "HBVACIGM", "HBVACIGG", "HCVBD" and "HDVA" codes for VS\_ID field.

tbILAB\_RES, tbILAB\_RES\_LVL\_1, tbILAB\_RES\_LVL\_2 and tbILAB\_RES\_LVL\_3:

- Renamed SAMP\_ID to TEST\_ID, as it gives a better description of what the values should encode.

tbILAB\_RES:

- Added a VIRUSTYPE variable allowing to distinguish HIV- and HCV-resistance tests.
- Added recommended HIV-1 subtype codes and HCV geno- and subtype codes.

tbILAB\_RES\_LVL\_3:

- ATC\_CODE replaces ART\_ID as HCV medication in tbIMED may be encoded.

tbIDIS:

- Added optional DIS\_ED field for the end date of the disease.

tbIMED:

- Added MED\_RS describing the reason for stopping the treatment; needed for HCV.

tbIAE:

- Added a number of codes for AE\_ID: ANG, ASCI, AVN, BYP, CERC, END, FIBS, FRA, HEP, HESY, LAC, LIVB, OESO, PAN and PERI.
- QA checks have been extended and updated where necessary.

**Version 1.50**

- Added the pediatric tables: tbIPREG, tbIPREG\_OBS, tbIPREG\_OUT, tbIDELIVERY\_MUM, tbIDELIVERY\_CHILD, tbINEWBORN and tbINEWBORN\_ABNORM.
- Added new table tbISAMPLES which holds information regarding sample storage.
- tbIART: Updated list of drugs

**Version 1.30**

- tbIART: Updated list of drugs
- tbIAE:
  - added EVENT\_ID as unique identifier and link to detailed tables for each event (see <sup>1</sup>) ? this replaces the optional AE\_NO field.
  - added AE\_SPEC to further specify an event by coding
  - a series of basic verification fields have been added to allow for tracking of event status for source documentation availability, verification of documentation (through monitoring) and final approval of the event.
  - in AE\_R\_Y - Relation to treatment: added more detailed codes.
- tbILAB: added several codes for various biomarker tests.
- tbILAB\_CD4: added CD4\_U as optional fields to discriminate between CD4% and CD4 cell count, so that the tbILAB\_CD4 table can hold both types of measurements.
- tbILAB\_VIRO: added several codes for various virology and serology tests.
- CaseDefinitions updated with end stage renal disease, chronic liver disease and non-AIDS defining malignancies

<sup>1</sup>:Detailed table definitions for the D:A:D events are available at <http://www.cphiv.dk/HICDEP/Documents/tabid/159/Default.aspx>

**Version 1.25**

- tbIART: Updated list of drugs
- tbIMED: Updated list of drugs
- tbIDIS:
  - Changed wording for CANO to ?Candidiasis, oesophageal, bronchi, trachea, or lungs?
  - Added COCC - Coccidioidomycosis, disseminated or extrapulmonary
- tbILAB: Added LAB\_ST as additional field to code for type of specimen used for the measurement
- tbILAB\_CD4: Added CD4\_U as additional field so the table can hold both percentage and absolute CD4 measurements
- tbILAB\_RNA: Added RNA\_UL (upper limit of detection) to the list of additional fields.

- Added more viral assays to the list of [RNA\\_T codes](#)
- [tblLAB\\_VIRO](#): Added unit field to [tblLAB\\_VIRO](#) into the general format and [VS\\_LL](#) (lower limit of detection), [VS\\_UL](#) (upper limit of detection) and [VS\\_T](#) (type of test) and list of tests to the list of additional fields.
- [tblLTFU](#): Added [DEATH\\_RC#](#) to code for causal relation of the [DEATH\\_R#](#) code to the death in order to comply with [CoDe](#) and still maintain a format to be used for cohorts not using CoDe. [ICD10\\_#](#) fields have been moved to the list of additional fields.

### Version 1.21

- Added reasons for stopping treatment to table [tblART\\_CODE\\_RS](#):

Code	Coding for Reason of Stopping Treatment
1.1	Virological failure
1.2	Partial virological failure
1.3	Immunological failure ? CD4 drop
1.4	Clinical progression
90	Side effects ? any of the above but unspecified
90.1	Comorbidity
92.1	Simplified treatment available
92.2	Treatment to complex
92.3	Drug interaction
93.1	Structured Treatment Interruption (STI) ? at high CD4
94.1	Non-compliance
96	Pregnancy
97	Study treatment

### Version 1.2

- added [CaseDefinitions](#)

### Version 1.1

#### [tblBAS](#):

- The table was split into [tblBAS](#) and [tblLTFU](#). [tblLTFU](#) holds data on death and drop-out
- Renamed [LOS\\_Y](#) to [LOSS\\_Y](#)
- Renamed [GAL\\_Y](#) to [GAIN\\_Y](#)

#### [tblLAB\\_BLP](#):

- Renamed table to [tblLAB\\_BP](#)
- Renamed [BLP\\_D](#) to [BP\\_D](#)
- Renamed [BLP\\_SYS](#) to [BP\\_SYS](#)
- Renamed [BLP\\_DIA](#) to [BP\\_DIA](#)
- Renamed [BLP\\_U](#) to [BP\\_U](#)

### Version 1.00

#### [tblBas](#):

- Renamed [BIRTHDAY](#) to [BIRTH\\_D](#)
- Renamed [FIRSTVIS](#) to [FRSVIS\\_D](#)
- Renamed [REC\\_ART](#) to [RECART\\_Y](#)

#### [tblLAB](#):

- [LAB\\_U](#): has been dropped ? please use the ?unit codes/strings? as that is a safer way to code/represent the units ? prefixing all ?unit codes/strings? with a numeric value should however make analysis easier.

#### [tblLAB\\_VIRO](#):

- New table added to capture mainly hepatitis measurements/tests

#### [tblLAB\\_RES](#):

- [SEQ\\_DT](#) was added to capture the time of sequencing in order to facilitate quality assurance of the data for contamination that might have happened during the sequencing.

#### [tblLAB\\_RES\\_LVL1](#):

- Renamed [SEQ\\_ST](#) to [SEQ\\_STAR](#)
- [SEQ\\_STOP](#): Added to the table to specify at which position in the sequence the sequencing was terminated

#### [tblLAB\\_RES\\_LVL2](#):

- The table has been optimised for ease of analysis so that the mutation codes have been split into their components of amino acid position, sub position for insertions and 4 our more fields for mixtures of amino acids found in the sample

### Version 0.50 and 0.90

First public versions that incorporated comments and corrections received from attendees at the 7th International Workshop on HIV Observational Databases, March 29th-30th 2003, Fiuggi, Italy and Stephen Hart.

### Version 0.38

Version presented at 7th International Workshop on HIV Observational Databases, March 29th-30th 2003, Fiuggi, Italy

## HICDEP 1.90

This article describes the current draft version HICDEP 1.90. For a more detailed version history, please refer to the [ChangeLog](#).

### General data format

The table pages referenced [in the overview](#) describe the specific tables' structure in detail and present a list of suggested codes, both standard and human readable.

All codes apart from trivial no, yes or unknown codes are presented as lookup tables, the usage of these are described in the the article [Considerations for using the format to create a database](#).

Along with the basic structure described in each ?Core fields? section, additional fields containing additional or more specific data are described in the ?Additional fields? sections. These fields were taken from several cohort collaborations but with the required changes that were needed for the specific data structures. This is presented to the reader to show that the core structure is not a fixed proposal but rather a basic structure, which can be altered by adding fields.

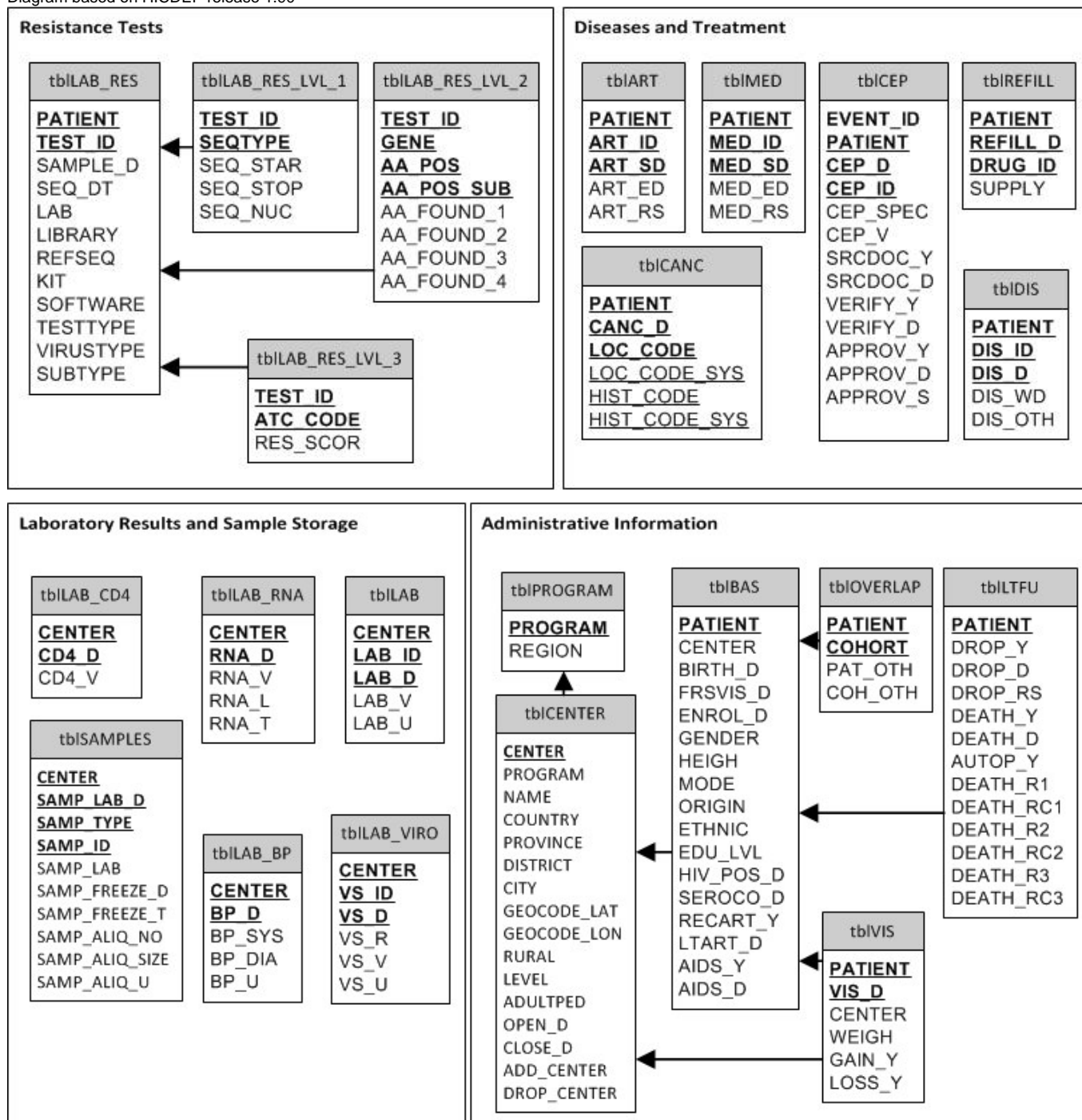
Issues regarding duplicates are discussed in [Considerations For Data Management](#).

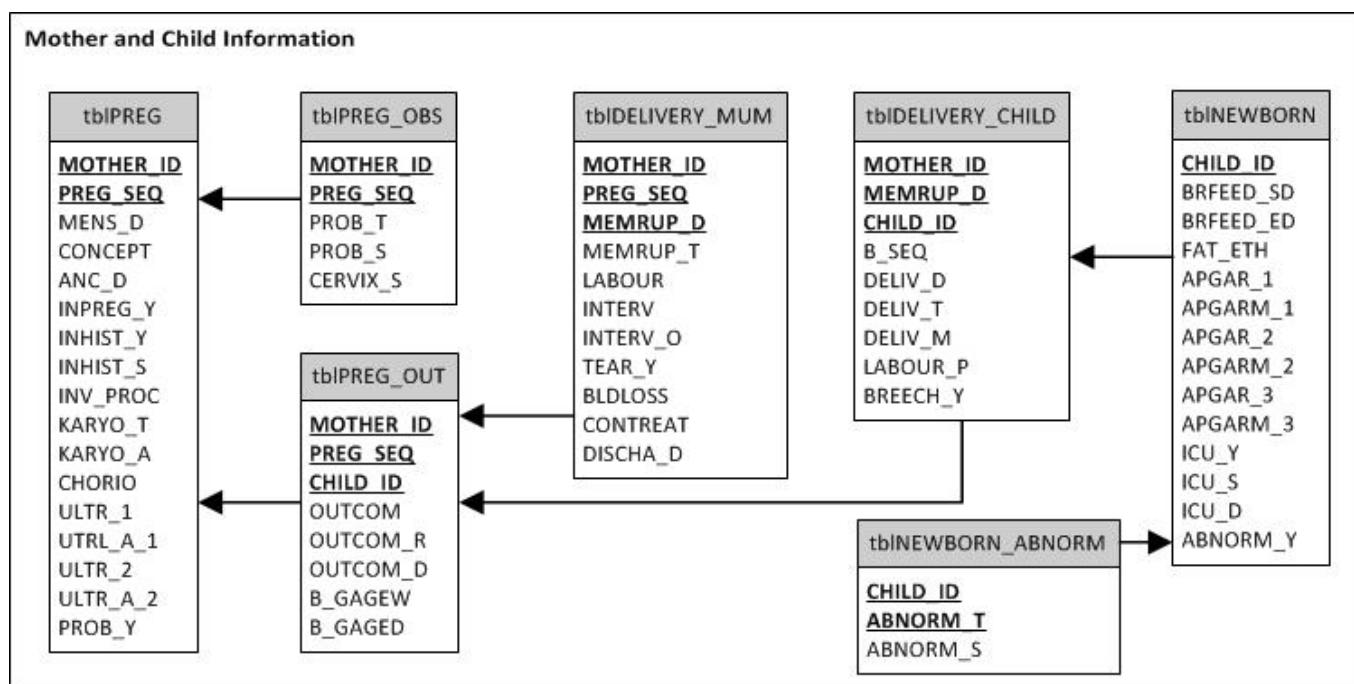
### Overview of data tables

Table	Content
<a href="#">tblART</a>	holds type of <b>antiretroviral drug</b> , start and stop dates and reason for stopping
<a href="#">tblBAS</a>	holds <b>basic</b> information such as demographics, basic clinical information and date of AIDS diagnosis
<a href="#">tblCANC</a>	holds type and date of <b>diagnosis of cancer</b>
<a href="#">tblCENTER</a>	holds information about the <b>Center</b> (e.g. geographical localisation, type of clinic) where the patient is receiving HIV care
<a href="#">tblCEP</a>	holds type and date of <b>clinical events and procedures</b> including serious non-AIDS conditions. Former known as tblAE (adverse event).
<a href="#">tblDELIVERY_CHILD</a>	holds <b>delivery</b> information related to the child
<a href="#">tblDELIVERY_MUM</a>	holds <b>delivery</b> information related to the mother
<a href="#">tblDIS</a>	holds type and date of CDC-C <b>diseases</b> and malignancies.
<a href="#">tblLAB</a>	holds type, date, value and unit of <b>laboratory tests</b> .
<a href="#">tblLAB_BP</a>	holds date, diastolic and systolic values and unit of <b>blood pressure</b> measurements.
<a href="#">tblLAB_CD4</a>	holds date and value of <b>CD4 measurements</b> .
<a href="#">tblLAB_RNA</a>	holds date, value, detection limit and type of <b>viral assay</b> .
<a href="#">tblLAB_RES</a>	holds <b>background information</b> on the resistance test, <b>laboratory</b> , library, kit, software and type of test
<a href="#">tblLAB_RES_LVL_1</a>	holds nucleoside sequence for the PRO and RT sequences
<a href="#">tblLAB_RES_LVL_2</a>	holds mutations and positions of these.
<a href="#">tblLAB_RES_LVL_3</a>	holds resistance result in relation to antiretroviral drug.
<a href="#">tblLAB_VIRO</a>	holds test results for <b>viro-/serological tests</b> (hepatitis etc.)
<a href="#">tblLTFU</a>	holds data on <b>death and drop-out</b>
<a href="#">tblMED</a>	holds type, start and stop dates for <b>other medication/treatments</b> .
<a href="#">tblNEWBORN</a>	holds information related to <b>newborns</b>
<a href="#">tblNEWBORN_ABNORM</a>	holds information related to <b>abnormalities</b> of newborns
<a href="#">tblOVERLAP</a>	holds information on the patient's <b>participation in other cohorts</b>
<a href="#">tblPREG</a>	holds general <b>pregnancy</b> -related information
<a href="#">tblPREG_OBS</a>	holds information on <b>obstetrical problems during pregnancy</b>
<a href="#">tblPREG_OUT</a>	describes the <b>pregnancy outcome</b>
<a href="#">tblPROGRAM</a>	holds information on the <b>program with which the center is associated</b>
<a href="#">tblREFILL</a>	holds information on <b>prescription refills</b>
<a href="#">tblSAMPLES</a>	holds information on the storage of blood, urine and other <b>biological samples</b>
<a href="#">tblVIS</a>	holds <b>visit related information</b> such as weight, wasting, smoking, occupational status etc.

### Diagram

Diagram based on HICDEP release 1.90





## Structure of data

### From flat files towards a normalized structure

The data collected in HIV collaborations is presented on the following pages in a set of data files/tables. Typically data would be put into one data file that would hold one line/record per patient where each field is represented as a separate column in that dataset. Often a dataset could contain more than 3000 columns of data.

The implication of going from thousands of fields to fewer fields means that data is in fact transposed from the flat format into the normalised format.

#### Example of a flat file structure:

PATIENT	ALAT_D	ALAT_V	ALAT_U	ASAT_D	ASAT_V	ASAT_U
999999	01-01-2000	15	U/l	01-01-2000	12	U/l

#### The normalised structure would then be like this:

PATIENT	TYPE_ID	LAB_DATE	LAB_VAL	LAB_UNIT
999999	1	01-01-2000	15	U/l
999999	2	01-01-2000	12	U/l

The type of measurement is identified through the TYPE\_ID field. Here 1 codes for ALAT and 2 codes for ASAT:

Code	Description
1	ALAT - Alanin-Aminotransferase
2	ASAT - Aspartat aminotransferase

## Technical considerations

To enable a normalised structure that minimises the number of columns dramatically, the one file solution must be broken into several minor tables. These breakdowns are driven by the different data characteristics.

Each table has a basic structure that includes the patient identifier, a code that represents e.g. drug, adverse event or laboratory test performed. Along with this combination values like date, result, unit etc are present for each record.

A record for a laboratory measurement would include:

- Patient identifier
- Measurement type identifier
- Measured value
- Unit of value
- Date of determination

A record for usage of an antiretroviral drug would include:

- Patient identifier
- Drug identifier
- Start date for usage
- End date for usage
- Reason for discontinuation

These issues imply that a set of distinct tables must be generated based on the ?nature? of the data. Since laboratory, medication and event data both cannot and should not be mixed at least 3 tables must be designed. Additionally there are other types of information that need their own domains: background information on the patient (height, birth date etc.), visit related data (weight, blood pressure, wasting etc.), and resistance testing (the latter requires more consideration due to the diversity of data present).

In this protocol further separation of data into different tables are presented. These separations are not only based on the rules for the relational model and normalisation, but they are ?culturally? related.

For example: antiretroviral treatment medication is kept in one table and other medication in another table; CD4 cell measurements and HIV-RNA measurements are put into separate tables, that are also different from the general laboratory table. These separations are done simply because data in these tables are of distinct importance in analysis and often are gathered more frequently and with more attention than other variables.

## Coding Conventions

### Date codes

Although it is best to have precise dates in the format of YEAR-MONTH-DAY [?ISO standard](#), it might be that some cohorts are limited to representing date data at the level of the month only, or information kept on the patient in the charts only defines dates to the month and in some cases only to the year. To solve this a set of date codes are presented here.

#### Day unknown

In this case the date should be coded as the 15th of the month ? so that 1999-12-?? becomes 1999-12-15. This enables the date to be no more than 15 days away from the actual date.

#### Month and day unknown

Best approach to this is to apply something similar, as with unknown dates, this would then mean that 1999-??-?? becomes 1999-07-01.

#### Year unknown

If the year is unknown but the presence of the date value is needed as in case of opportunistic infections or adverse events (see later in this document) a fictive date should be used that couldn't be mistaken with an actual date. An unknown year should be coded as 1911-11-11.

### Specification of precision

An alternative to the above is to apply an additional field to each date field for which it is known that there might be issues regarding the precision of the dates. The field is then used to specify at which degree of the day, month or year the date is precise:

Code	Precision of date
<	Before this date
D	Exact to the date
M	Exact to the month
Y	Exact to the year
>	After this date
U	Unknown

### [?ICD-10 codes](#)

The coding system is the official standard for coding of diseases, however there is a wide set of ?homebrew? codes used within the HIV field in data coding in general, often it's a 3 or 4 letter code which is an abbreviation for the AIDS defining disease. ICD-10 doesn't have single codes that represent all single CDC-C events and as a consequence of this a list of 3 to 4 letter codes is the recommended way of coding for all CDC stage C events

ICD-10 codes are however the recommended for codes AE?s since it would become impossible for this protocol to maintain a complete list of all possible AE?s. ICD-10 is also recommended for causes of death.

### [?ATC codes](#)

ATC is a hierarchical structure for coding medication. The structure and hierarchy are best explained with an example of how a drug code is defined. Here it is on Indinavir:

```
J
  ANTIINFECTIVES FOR SYSTEMIC USE (1st level, anatomical main group)
J05
  ANTIVIRALS FOR SYSTEMIC USE (2nd level, therapeutic subgroup)
J05A
  DIRECT ACTING ANTIVIRALS (3rd level, pharmacological subgroup)
J05AE
  Protease inhibitors (4th level, chemical subgroup)
J05AE02
  Indinavir (5th level, chemical substance)
```

This hierarchy has some benefits as will be explained later, but one of its limitations is that it's impossible to read the code compared to the widely used 3 letter mnemonic codes for antiretroviral drugs.

Example:

Drug	Code	ATC code
Indinavir	IDV	J05AE02

The difference is that the IDV code is easily readable, where the ATC code is not; going from a flat file structure to a normalised structure the human readable aspect becomes increasingly important. In the flat file format the column names and the possibility of labels makes data more or less readable; in the normalised format only the coding can help. Because of this the 3 letter codes are being presented in this document. However it must be stressed that usage of the ATC coding should be used to diminish the risk of several homebrew and non-compatible coding schemes.

**Currently however, the ATC scheme does not provide sufficient detail on the specific drugs, there is e.g. no official way to code Saquinavir as hard or soft gel. Thus a slight alteration to the set of codes will be presented in the sections of the ART and MED tables. The alterations are designed to extend the existing structure of ATC.**

One of the benefits is that the structure of ATC allows easier statistics on e.g. drug classes

- J05AE Protease inhibitors
  - J05AE01 Saquinavir
  - J05AE02 Indinavir
  - J05AE03 Ritonavir
  - J05AE04 Nelfinavir
  - J05AE05 Amprenavir
  - J05AE06 Lopinavir
- J05AF Nucleoside and nucleotide reverse transcriptase inhibitors
  - J05AF01 Zidovudine
  - J05AF02 Didanosine
  - J05AF03 Zalcitabine
  - J05AF04 Stavudine
  - J05AF05 Lamivudine
  - J05AF06 Abacavir
  - J05AF07 Tenofovir disoproxil
  - J05AF30 Combinations1
- J05AG Non-nucleoside reverse transcriptase inhibitors
  - J05AG01 Nevirapine
  - J05AG02 Delavirdine
  - J05AG03 Efavirenz

Although the codes might be harder to read they provide grouping mechanisms in the way they are coded. Interested readers should go to the [ATC Website](#) to learn about the structure of ATC. A fully updated database of ATC codes and DDD (Defined Daily Dosage) is available for querying.

### Other codes

It is often necessary to code for values like ?Yes?, ?No? and ?Unknown?, this document suggests that the following codes should be used:

Code	Description
0	No
1	Yes
9	Unknown

Unknown should be used to identify the difference between a value that has not yet been collected (Empty) and a value that cannot be collected (Unknown). Empty values should be required where Unknown values make little sense to keep querying for a value.

Example ? weight:

Depending on the unit in which weight is measured, a different value for Unknown should be applied. In the case of kg the ?Unknown? code should be 999 and not just 9 or 99, the last two could be actual values.

Blank values, for SAS users also known as " ." and for database programmers known as NULL, should be used wherever specified in this protocol. However, sometimes it might be more correct just to omit the record if no value has been recorded, test has not been performed etc.

### Quality Assurance

In order to verify the consistency and correctness of the data, QA checks are made before the data is used. The QA checks applying to a given table are listed at the bottom of its article. Additionally, a list of all QA checks, including checks which do not directly apply to the HICDEP tables themselves, is available [here](#).



## tbIART - Antiretroviral treatment

holds type of antiretroviral drug, start and stop dates and reason for stopping

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	identifies patient
<a href="#">ART_ID</a>	character. see <a href="#">coding table</a> for valid codings.	represents the antiretroviral treatment
<a href="#">ART_SD</a>	yyyy-mm-dd	date of initiation of treatment
<a href="#">ART_ED</a>	yyyy-mm-dd	date of stopping treatment
<a href="#">ART_RS</a>	character. see <a href="#">coding table</a> for valid codings.	reason for stopping treatment

### Additional fields

Depending on the aim of the study it might be needed to gather both the dosage and the frequency of the dosage taken. However many cohorts do not collect this date and thus these fields are optional.

Field name	Format	Description
<a href="#">ART_RS2</a>	character. see <a href="#">coding table</a> for valid codings.	Second reason for stopping treatment
<a href="#">ART_RS3</a>	character. see <a href="#">coding table</a> for valid codings.	Third reason for stopping treatment
<a href="#">ART_RS4</a>	character. see <a href="#">coding table</a> for valid codings.	Fourth reason for stopping treatment
<a href="#">ART_DO</a>	numeric	Dosage (mg or mL) per intake unless ART_FR=-1
<a href="#">ART_FR</a>	numeric: <ul style="list-style-type: none"> <li>-1 = Frequency not known. ART_DO contains dosage per day</li> <li>0.33 = 1 dose every third day</li> <li>0.5 = 1 dose every second day</li> <li>1 = 1 daily dose/qd</li> <li>2 = 2 daily doses/bid</li> <li>3 = 3 daily doses/tid</li> <li>4... = code gives number of daily doses</li> </ul>	Frequency
<a href="#">GENERIC</a>	numeric: <ul style="list-style-type: none"> <li>1 = Branded</li> <li>2 = Generic</li> <li>9 = Unknown</li> </ul>	Was this a branded or generic drug?

It may also be necessary to record the start and end time:

Field name	Format	Description
<a href="#">ART_ST</a>	hh:mm	Start hour and minute of the day
<a href="#">ART_ET</a>	hh:mm	Stop hour and minute of the day

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES

AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblART	WithinTable	AW001	ART_RS not null, but end date NULL		YES
tblART	WithinTable	AW002	ART_RS null but end-date non NULL		YES
tblART	WithinTable	AW003	ART_RS=98 yet ART_OTH is null	EPPICC	NO
tblART	WithinTable	AW004	Duplicate records for same cohort, patient, art_id and art_sd		YES
tblART	WithinTable	AW005	Dose out of range for those dose units and frequency	PaediatricOnly	YES
tblART	WithinTable	AW006	Missing art_fr	PaediatricOnly	YES
tblART	WithinTable	AW007	Missing art_do	PaediatricOnly	YES
tblART	WithinTable	AW008	Missing patient		YES
tblART	WithinTable	AW009	Missing art_id		YES
tblART	WithinTable	AW010	Missing art_sd		YES
tblART	WithinTable	AW011	Overlapping periods of same drug		YES
tblART	WithinTable	AW012	Double reporting - records reported for both combination drugs and their components		YES
tblART	WithinTable	AW013	Periods of overlap of contra-indicated drugs		YES
tblART	WithinTable	AW014	Restart of same drug without a stop		YES
tblART	WithinTable	AW015	ART_SD greater than or equal to ART_ED		YES
tblART	CrossTable	AC001	Patient has no record in table BAS		YES
tblART	CrossTable	AC002	Records exist in tblART yet RECART_Y=0 in tblBAS		YES

## ART\_ID field

containing table

[tblART](#)

explanation of variable

Code representing the antiretroviral treatment

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

A set of extended ATC codes are being presented here in order to code both more specific on subtypes of the drugs, e.g. saquinavir hard and soft gel, but also to enable coding of drugs that are at their trial stage and have not yet been assigned an ATC code. To do this the drug will be assigned the code elements as far down the levels as possible. Given two examples to illustrate this:

Saquinavir - Hard Gel

J05AE01-SQH

Saquinavir - Soft Gel

J05AE01-SQS

Saquinavir - not specified

J05AE01

This will ensure the fidelity needed to distinguish between hard and soft gel and not specified, but also for analysis easily include all records which coding starts with J05AE01, regardless if the drug is hard or soft gel.

See the [?ATC Index](#) for the individual codes. The extended ATC codes are listed below along with a subset of ATC codes relevant to ART.

[Download this table as CSV file](#) [Download this table as XML file](#)

Code (Extended ATC Codes)	Anti-Retroviral Drugs
J05A	ART unspecified
J05A-BEV	Bevirimat
J05A-PBT	Participant in Blinded Trial
J05AE	PI unspecified
J05AE-MOZ	Mozenavir (DMP-450)
J05AE01	Saquinavir (gel, not specified)
J05AE01-SQH	Saquinavir hard gel (INVIRASE)
J05AE01-SQS	Saquinavir soft gel (FORTOVASE)
J05AE02	Indinavir (CRIXIVAN)
J05AE03	Ritonavir (NORVIR)
J05AE03-H	Ritonavir high dose (NORVIR)
J05AE03-L	Ritonavir low dose (NORVIR)
J05AE04	Nelfinavir (VIRACEPT)
J05AE05	Amprenavir (AGENERASE)
J05AR10	Lopinavir/Ritonavir (Kaletra). Former code: J05AE06
J05AE07	Fos-amprenavir (Telzir, Lexiva)
J05AE08	Atazanavir (Reyataz)
J05AE09	Tipranavir (Aptivus)
J05AE10	Darunavir (TMC-114, Prezista)
J05AF	NRTI unspecified
J05AF-ALO	Alovudine
J05AF-AMD	Amdoxovir (DADP)
J05AF-FOZ	Fozivudine tidoxi
J05AF-LDN	Lodenoine (trialdrug)
J05AF-RVT	Reverset
J05AF01	Zidovudine (AZT, RETROVIR)
J05AF02	Didanosine (ddl) (VIDEX)
J05AF03	Zalcitabine (ddC) (HIVID)
J05AF04	Stavudine (d4T) (ZERIT)
J05AF05	Lamivudine (3TC, EPIVIR)

J05AF06	Abacavir (1592U89) (ZIAGEN)
J05AF07	Tenofovir (VIREAD)
J05AF09	Emtricitabine
J05AG	NNRTI unspecified
J05AG04	Etravirine (TMC 125)
J05AG05	Rilpivirine (TMC-278)
J05AG-CPV	Capravirine
J05AG-DPC083	DPC 083
J05AG-DPC961	DPC 961
J05AG-EMV	Emivirine (MKC442)
J05AG-LOV	Loviride
J05AG01	Nevirapine (VIRAMUN)
J05AG02	Delavirdine (U-90152) (RESCRIPTOR)
J05AG03	Efavirenz (DMP-266) (STOCRIN, SUSTIVA)
J05AR01	Combivir (Zidovudine/Lamivudine)
J05AR02	Kivexa (Lamivudine/Abacavir)
J05AR03	Truvada (Tenofovir/Emtricitabine)
J05AR04	Trizivir (Zidovudine/Lamivudine/Abacavir)
J05AR05	Douvir-N (Zidovudine/Lamivudine/Nevirapine)
J05AR06	Atripla (Emtricitabine/Tenofovir/Efavirenz)
J05AR07	Triomune (Stavudine/Lamivudine/Nevirapine)
J05AR08	Eviplera/Complera (Emtricitabine/Tenofovir/Rilpivirine)
J05AR09	Stribild (Emtricitabine/Tenofovir/Elvitegravir/Cobicistat)
J05AR10	Kaletra/Aluvia (Lopinavir/Ritonavir)
J05AR11	Lamivudine, tenofovir disoproxil and efavirenz
J05AR12	Lamivudine and tenofovir disoproxil
J05AR13	Triumeq (Lamivudine, abacavir and dolutegravir)
J05AR14	Darunavir and cobicistat
J05AR15	Atazanavir and cobicistat
J05AR16	Lamivudine and raltegravir
J05AR17	Emtricitabine and tenofovir alafenamide
J05AR18	Emtricitabine, tenofovir alafenamide, elvitegravir and cobicistat
J05AR19	Emtricitabine, tenofovir alafenamide and rilpivirine
J05AX-VIC	Vicriviroc (Schering)
J05AX07	Enfuvirtide (Fuzeon, T-20)
J05AX08	Raltegravir (Merck)
J05AX09	Maraviroc (Pfizer)
J05AX11	Elvitegravir
J05AX12	Dolutegravir
J05AX-CAB	Cabotegravir (GSK-744)
L01XX05	Hydroxyurea/Hydroxycarbamid (Litalir)
V03AX03	Cobicistat

## ART\_RS field

containing table

[tblART](#)

explanation of variable

Reason for stopping treatment

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Reason for Stopping Treatment
1	Treatment failure (i.e. virological, immunological, and/or clinical failure)
1.1	Virological failure
1.2	Partial virological failure
1.3	Immunological failure - CD4 drop
1.4	Clinical progression
2	Abnormal fat redistribution
3	Concern of cardiovascular disease
3.1	Dyslipidaemia
3.2	Cardiovascular disease
4	Hypersensitivity reaction (skin eruption etc.)
5	Toxicity, predominantly from abdomen/G-I tract
5.1	Toxicity - GI tract
5.2	Toxicity - Liver
5.3	Toxicity - Pancreas
6	Toxicity, predominantly from nervous system
6.1	Toxicity - peripheral neuropathy
6.2	Toxicity - neuropsychiatric
6.3	Toxicity - headache
7	Toxicity, predominantly from kidneys
8	Toxicity, predominantly from endocrine system
8.1	Diabetes
9	Haematological toxicity (anemia etc.)
10	Hyperlactataemia/lactic acidosis
11	Bone toxicity
15	Social contra-indication
16	Contra-indication unspecified
16.8	Contra-indication expired
16.9	Contra-indication ? other
17	MTCT regimen completed
70	Pregnancy - toxicity concerns (during pregnancy)
75	Pregnancy - switch to a more appropriate regimen for PMTCT
88	Death
90	Side effects - any of the above but unspecified
90.1	Comorbidity
91	Toxicity, not mentioned above
91.1	Toxicity, unspecified
92	Availability of more effective treatment (not specifically failure or side effect related)
92.1	Simplified treatment available
92.2	Treatment too complex

92.3	Drug interaction
92.31	Drug interaction - commencing TB/BCG treatment
92.32	Drug interaction - ended TB/BCG treatment
92.33	Change in eligibility criteria (e.g. child old enough for tablets; refrigerator no longer available)
92.4	Protocol change
92.5	Regular treatment termination (used in tbIMED e.g. for DAA's against HCV, antibiotics)
92.9	Change in treatment not due to side-effects, failure, poor adherence or contra-indication
93	Structured Treatment Interruption (STI)
93.1	Structured Treatment Interruption (STI) - at high CD4
94	Patient's wish/decision, not specified above
94.1	Non-compliance
94.2	Defaulter
95	Physician's decision, not specified above
96	Pregnancy (see also more specific codes 70 and 75)
96.1	Pregnancy intended
96.2	Pregnancy ended
97	Study treatment
97.1	Study treatment commenced
97.2	Study treatment completed
97.6	Drug not available
98	Other causes, not specified above
99	Unknown

Although most reasons mentioned above could be coded in ICD-10 codes, it is still certain that many reasons will never be defined in terms of ICD-10 (STI, physicians and patients decision), so to avoid a mixture of two different coding systems it is advised to go with the above list and build upon that for now. This coding list is identical to MED\_RS used for non-ART medication recorded in tbIMED

## tbIBAS - Basic clinical, background and demographic information

holds basic information such as demographics, basic clinical information, date of AIDS diagnosis, death and drop-out information

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">CENTER</a>	character	Code for Clinic/Centre/Hospital where patient is seen. Links to <a href="#">tbICENTER</a> .
<a href="#">BIRTH_D</a>	yyyy-mm-dd	Birth date
<a href="#">FRSVIS_D</a>	yyyy-mm-dd	First seen at clinic
<a href="#">ENROL_D</a>	yyyy-mm-dd	Date of enrolment into the cohort
<a href="#">GENDER</a>	numeric: <ul style="list-style-type: none"> <li>1 = Male</li> <li>2 = Female</li> <li>9 = Unknown</li> </ul>	Gender/sex
<a href="#">HEIGH</a>	numeric (metric): 999 = Unknown	Height of patient at visit/most current
<a href="#">MODE</a>	numeric. see <a href="#">coding table</a> for valid codings.	Mode of infection
<a href="#">ORIGIN</a>	character (1-3 letter/numeric codes). see <a href="#">coding table</a> for valid codings.	Country or region of birth
<a href="#">ETHNIC</a>	numeric. see <a href="#">coding table</a> for valid codings.	Ethnicity of patient. Please take the <a href="#">additional notes</a> into consideration when using this field.
<a href="#">EDU_LVL</a>	numeric. see <a href="#">coding table</a> for valid codings.	Last completed education Level. ISCED97 refers to the <a href="#">1997 International Standard Classification of Education</a>
<a href="#">HIV_POS_D</a>	yyyy-mm-dd	Date of first positive HIV test
<a href="#">SEROCO_D</a>	yyyy-mm-dd	Date of seroconversion
<a href="#">RECART_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Has the patient received antiretroviral treatment?
<a href="#">RECART_D</a>	yyyy-mm-dd	Date ART started
<a href="#">LTART_D</a>	yyyy-mm-dd	Date last assessed for ART. If started ART, last date known to be on ART, or if not on ART, last date ART free.
<a href="#">AIDS_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Has patient been given an AIDS diagnosis?
<a href="#">AIDS_D</a>	yyyy-mm-dd	IF YES, date of AIDS diagnosis

### Additional fields

For mode of infection and origin a set of other fields are often used to capture what cannot be coded. These fields are represented here as optional fields as it is the intention that the suggested codes applied to the MODE and ORIGIN should be able to cover all possible values.

Field name	Format	Description
<a href="#">MODE_OTH</a>	character	Mode of infection OTHER
<a href="#">ORI_OTH</a>	character	Origin of patient OTHER
<a href="#">CENS_D</a>	yyyy-mm-dd	The last date the database has been updated for this patient
<a href="#">SEROHOW</a>	numeric: <ul style="list-style-type: none"> <li>1 = Midpoint between last neg/first pos test</li> <li>2 = Lab evidence of seroconversion</li> <li>3 = Seroconversion illness</li> <li>4 = Other</li> <li>9 = Unknown</li> </ul>	For Seroconverters only: How was the seroconversion date determined?

<a href="#">NAIVE_Y</a>	numeric: <ul style="list-style-type: none"> <li>• 0 = No</li> <li>• 1 = Yes</li> <li>• 9 = Unknown</li> </ul>	Is the patient ART-naïve upon enrollment?
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### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblBAS	WithinTable	BW001	AIDS date < SEROCO_D		YES
tblBAS	WithinTable	BW002	Duplicate patients		YES
tblBAS	WithinTable	BW003	First 3 chars of PATIENT don't form valid cohort code	CascadeOnly	YES
tblBAS	WithinTable	BW004	Missing PATIENT		YES
tblBAS	WithinTable	BW005	Missing CENTER		YES
tblBAS	WithinTable	BW006	Missing BIRTH_D		YES
tblBAS	WithinTable	BW007	Missing FRVIS_D		YES
tblBAS	WithinTable	BW008	Missing ENROL_D		YES
tblBAS	WithinTable	BW009	Missing GENDER		YES
tblBAS	WithinTable	BW010	Missing HEIGH		YES
tblBAS	WithinTable	BW011	Missing MODE		YES
tblBAS	WithinTable	BW012	Missing MODE_OTH if MODE=90		YES
tblBAS	WithinTable	BW013	Missing ORIGIN		YES
tblBAS	WithinTable	BW014	Missing ETHNIC		YES
tblBAS	WithinTable	BW015	Missing SEROCO_D		YES
tblBAS	WithinTable	BW016	Missing RECAT_Y		YES
tblBAS	WithinTable	BW017	Missing AIDS_Y		YES
tblBAS	WithinTable	BW018	Missing AIDS_D if AIDS_Y=1		YES
tblBAS	WithinTable	BW019	BIRTH_D out of range (15-85 yrs)	CascadeOnly	YES
tblBAS	WithinTable	BW020	BIRTH_D out of range (<18)	PENTA	YES
tblBAS	CrossTable	BC001	RECAT_Y=1 but no records in tblART		YES
tblBAS	CrossTable	BC002	AIDS_Y=0, but AIDS-defining records in tblDIS		YES
tblBAS	CrossTable	BC003	AIDS_Y=1 but no AIDS-defining records in tblDIS table		YES



## EDU\_LVL field

containing table

[tblBAS](#)

explanation of variable

Last completed education Level. ISCED97 refers to the [1997 International Standard Classification of Education](#)

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.70](#)

## Coding Table

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Description
1	primary education (ISCED97-1)
2	lower secondary (ISCED97-2) OR end of basic education
3	upper secondary or post-secondary non-tertiary (ISCED97 3 and 4)
4	university or post-graduate (ISCED97 5A and 5B)
8	other, only if none of the codes 0 to 4 applies
9	Unknown

## ETHNIC field

containing table

[tblBAS](#)

explanation of variable

Ethnicity of patient

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

Codes are hierarchically structured, therefore please indicate most detailed code as possible.

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Ethnicity of patient
10	White/Caucasian
20	Black
21	> Sub-Saharan African
22	> Caribbean
23	> African-American
24	> Other Black
30	Hispanic/Latino/Latin American
40	Asian
41	> Chinese
42	> Southeas Asia (e.g. Thai, Vietnamese, Philippino)
43	> Indian Subcontinent (Indian, Pakistani, Bangladeshi)
44	> Japanese
45	> Other asian
50	Indigenous people from Americas or Alaska Native
60	Indigenous people from other continents/locations
70	Other ethnic groups
71	> Maghrebian
72	> Middle East/Arabic
73	> Turkish
74	> Roma people/Gypsy (whichever is term is acceptable)
xyyy(zz)	Mixed race/ethnicity. Combine 2-digits numbers from above (e.g. 1020 for white+black)
98	Prohibited (there are countries with legal restrictions to collect information on ethnicity)
99	Unknown

## Limitations

The definition of ethnicity is complex and there is no ideal definition for all countries and for all times. Likewise is the definition of race and though conceptually different from ethnicity, they are often used interchangeably. As described by many authors, ethnicity is a fluid and imprecise concept heavily influenced by societal views. If definition of ethnicity is complex, inevitably its categorization will be complex too. The definition and categorization used in HICDEP acknowledges these limitations and aims by no means to solve the intense international debate of this issue but to provide a homogeneous and practical approach for HIV research. We have partially used existing administrative classifications as they provide the advantage to have, in some instances, census population denominators but are invariably too detailed for practical use in the context of HICDEP.

We suggest users to ask themselves "why is this variable necessary to answer my research question?" to avoid some of the common mistakes highlighted in the publications below which have attributed to exclusively biological and/or genetic traits differences heavily influenced by the profound social, cultural and political differences inherent to those categories. We aim to provide a standardized definition that, in addition to the information on country or region of birth already collected within HICDEP, can be used by cohort studies of HIV infected people from different countries. Therefore, in order to encompass these different scenarios, some terms may have little meaning for some settings. Finally, this classification allows for multiple options and whenever possible, should be based on the patients' self-identification.

1. Ahdieh L, Hahn RA. Use of the terms "race", "ethnicity", and "national origins": a review of articles in the American Journal of Public Health, 1980-1989. *Ethnicity and Health* 1996; 1:95-8
2. Bhopal R. Glossary of terms relating to ethnicity and race: for reflection and debate. *J. Epidemiol. Community Health* 2004; 58:441-445
3. Cooper RS, Kaufman JS, Ward R. Race and Genomics. *N Engl J Med* 2003; 348; 12: 1166-1170
4. European Centre for Disease Prevention and Control. Improving HIV data comparability in migrant populations and ethnic minorities in EU/EEA/EFTA countries: findings from a literature review and expert panel. Stockholm: ECDC; 2011. [www.ecdc.europa.eu](http://www.ecdc.europa.eu)

## MODE field

containing table

[tbIBAS](#)

explanation of variable

Mode of infection

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Mode of infection
1	homo/bisexual
2	injecting drug user
3	(1+2)
4	haemophiliac
5	transfusion, non-haemophilia related
6	heterosexual contact
7	(6+2)
8	Perinatal
90	other, (specify)
99	unknown

## ORIGIN field

containing table

[tblBAS](#)

explanation of variable

Country or region of birth

format of data

character (1-3 letter/numeric codes). see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Region codes](#) & [country codes](#)

### Region Codes

[UN Region Codes](#) - Composition of macro geographical (continental) regions, geographical sub-regions, and selected economic and other groupings

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Region
001	World
002	Africa
014	- Eastern Africa
017	- Middle Africa
015	- Northern Africa
018	- Southern Africa
011	- Western Africa
019	Americas
419	- Latin America and the Caribbean
029	- - Caribbean
013	- - Central America
005	- - South America
021	- Northern America
142	Asia
143	- Central Asia
030	- Eastern Asia
034	- Southern Asia
035	- South-Eastern Asia
145	- Western Asia
150	Europe
151	- Eastern Europe
154	- Northern Europe
039	- Southern Europe
155	- Western Europe
009	Oceania
053	- Australia and New Zealand
054	- Melanesia
057	- Micronesia
061	- Polynesia

### Country Codes

[UN Country Codes](#) - Countries or areas, codes and abbreviations, including translation from ISO ALPHA-3 codes.

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Country	ISO ALPHA-3 code
4	Afghanistan	AFG
248	Åland Islands	ALA

8	Albania	ALB
12	Algeria	DZA
16	American Samoa	ASM
20	Andorra	AND
24	Angola	AGO
660	Anguilla	AIA
28	Antigua and Barbuda	ATG
32	Argentina	ARG
51	Armenia	ARM
533	Aruba	ABW
36	Australia	AUS
40	Austria	AUT
31	Azerbaijan	AZE
44	Bahamas	BHS
48	Bahrain	BHR
50	Bangladesh	BGD
52	Barbados	BRB
112	Belarus	BLR
56	Belgium	BEL
84	Belize	BLZ
204	Benin	BEN
60	Bermuda	BMU
64	Bhutan	BTN
68	Bolivia (Plurinational State of)	BOL
535	Bonaire, Saint Eustatius and Saba	BES
70	Bosnia and Herzegovina	BIH
72	Botswana	BWA
76	Brazil	BRA
92	British Virgin Islands	VGB
96	Brunei Darussalam	BRN
100	Bulgaria	BGR
854	Burkina Faso	BFA
108	Burundi	BDI
116	Cambodia	KHM
120	Cameroon	CMR
124	Canada	CAN
132	Cape Verde	CPV
136	Cayman Islands	CYM
140	Central African Republic	CAF
148	Chad	TCD
830	Channel Islands	
152	Chile	CHL
156	China	CHN
344	China, Hong Kong Special Administrative Region	HKG
446	China, Macao Special Administrative Region	MAC
170	Colombia	COL
174	Comoros	COM
178	Congo	COG
184	Cook Islands	COK
188	Costa Rica	CRI
384	Côte d'Ivoire	CIV

191	Croatia	HRV
192	Cuba	CUB
531	Curaçao	CUW
196	Cyprus	CYP
203	Czech Republic	CZE
408	Democratic People's Republic of Korea	PRK
180	Democratic Republic of the Congo	COD
208	Denmark	DNK
262	Djibouti	DJI
212	Dominica	DMA
214	Dominican Republic	DOM
218	Ecuador	ECU
818	Egypt	EGY
222	El Salvador	SLV
226	Equatorial Guinea	GNQ
232	Eritrea	ERI
233	Estonia	EST
231	Ethiopia	ETH
234	Faeroe Islands	FRO
238	Falkland Islands (Malvinas)	FLK
242	Fiji	FJI
246	Finland	FIN
250	France	FRA
254	French Guiana	GUF
258	French Polynesia	PYF
266	Gabon	GAB
270	Gambia	GMB
268	Georgia	GEO
276	Germany	DEU
288	Ghana	GHA
292	Gibraltar	GIB
300	Greece	GRC
304	Greenland	GRL
308	Grenada	GRD
312	Guadeloupe	GLP
316	Guam	GUM
320	Guatemala	GTM
831	Guernsey	GGY
324	Guinea	GIN
624	Guinea-Bissau	GNB
328	Guyana	GUY
332	Haiti	HTI
336	Holy See	VAT
340	Honduras	HND
348	Hungary	HUN
352	Iceland	ISL
356	India	IND
360	Indonesia	IDN
364	Iran (Islamic Republic of)	IRN
368	Iraq	IRQ
372	Ireland	IRL

833	Isle of Man	IMN
376	Israel	ISR
380	Italy	ITA
388	Jamaica	JAM
392	Japan	JPN
832	Jersey	JEY
400	Jordan	JOR
398	Kazakhstan	KAZ
404	Kenya	KEN
296	Kiribati	KIR
414	Kuwait	KWT
417	Kyrgyzstan	KGZ
418	Lao People's Democratic Republic	LAO
428	Latvia	LVA
422	Lebanon	LBN
426	Lesotho	LSO
430	Liberia	LBR
434	Libya	LBY
438	Liechtenstein	LIE
440	Lithuania	LTU
442	Luxembourg	LUX
450	Madagascar	MDG
454	Malawi	MWI
458	Malaysia	MYS
462	Maldives	MDV
466	Mali	MLI
470	Malta	MLT
584	Marshall Islands	MHL
474	Martinique	MTQ
478	Mauritania	MRT
480	Mauritius	MUS
175	Mayotte	MYT
484	Mexico	MEX
583	Micronesia (Federated States of)	FSM
492	Monaco	MCO
496	Mongolia	MNG
499	Montenegro	MNE
500	Montserrat	MSR
504	Morocco	MAR
508	Mozambique	MOZ
104	Myanmar	MMR
516	Namibia	NAM
520	Nauru	NRU
524	Nepal	NPL
528	Netherlands	NLD
540	New Caledonia	NCL
554	New Zealand	NZL
558	Nicaragua	NIC
562	Niger	NER
566	Nigeria	NGA
570	Niue	NIU

574	Norfolk Island	NFK
580	Northern Mariana Islands	MNP
578	Norway	NOR
275	Occupied Palestinian Territory	PSE
512	Oman	OMN
586	Pakistan	PAK
585	Palau	PLW
591	Panama	PAN
598	Papua New Guinea	PNG
600	Paraguay	PRY
604	Peru	PER
608	Philippines	PHL
612	Pitcairn	PCN
616	Poland	POL
620	Portugal	PRT
630	Puerto Rico	PRI
634	Qatar	QAT
410	Republic of Korea	KOR
	Republic of Kosovo	
498	Republic of Moldova	MDA
638	Réunion	REU
642	Romania	ROU
643	Russian Federation	RUS
646	Rwanda	RWA
652	Saint-Barthélemy	BLM
654	Saint Helena	SHN
659	Saint Kitts and Nevis	KNA
662	Saint Lucia	LCA
663	Saint-Martin (French part)	MAF
666	Saint Pierre and Miquelon	SPM
670	Saint Vincent and the Grenadines	VCT
882	Samoa	WSM
674	San Marino	SMR
678	Sao Tome and Principe	STP
680	Sark	
682	Saudi Arabia	SAU
686	Senegal	SEN
688	Serbia	SRB
690	Seychelles	SYC
694	Sierra Leone	SLE
702	Singapore	SGP
534	Sint Maarten (Dutch part)	SXM
703	Slovakia	SVK
705	Slovenia	SVN
90	Solomon Islands	SLB
706	Somalia	SOM
710	South Africa	ZAF
728	South Sudan	SSD
724	Spain	ESP
144	Sri Lanka	LKA
729	Sudan	SDN



740	Suriname	SUR
744	Svalbard and Jan Mayen Islands	SJM
748	Swaziland	SWZ
752	Sweden	SWE
756	Switzerland	CHE
760	Syrian Arab Republic	SYR
762	Tajikistan	TJK
764	Thailand	THA
807	The former Yugoslav Republic of Macedonia	MKD
626	Timor-Leste	TLS
768	Togo	TGO
772	Tokelau	TKL
776	Tonga	TON
780	Trinidad and Tobago	TTO
788	Tunisia	TUN
792	Turkey	TUR
795	Turkmenistan	TKM
796	Turks and Caicos Islands	TCA
798	Tuvalu	TUV
800	Uganda	UGA
804	Ukraine	UKR
784	United Arab Emirates	ARE
826	United Kingdom of Great Britain and Northern Ireland	GBR
834	United Republic of Tanzania	TZA
840	United States of America	USA
850	United States Virgin Islands	VIR
858	Uruguay	URY
860	Uzbekistan	UZB
548	Vanuatu	VUT
862	Venezuela (Bolivarian Republic of)	VEN
704	Viet Nam	VNM
876	Wallis and Futuna Islands	WLF
732	Western Sahara	ESH
887	Yemen	YEM
894	Zambia	ZMB
716	Zimbabwe	ZWE

## tbICENTER - Center information

Holds information about the Center in general, such as geographical location, start date of data collection, type of care provider etc.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b><u>CENTER</u></b>	character	Code for Clinic/Center/Hospital where patient is seen. Needs to be unique.
<u>PROGRAM</u>	character	Program or region with which the center is associated. Links to <a href="#">tbIPROGRAM</a> .
<u>NAME</u>	character	Proper name to identify center
<u>COUNTRY</u>	character	3-letter ISO code
<u>PROVINCE</u>	character	(Optional) Proper name to identify province
<u>DISTRICT</u>	character	(Optional) Proper name to identify district
<u>CITY</u>	character	(Optional) Proper name to identify city
<u>GEOCODE_LAT</u>	numeric	Latitude (e.g. 47.376739 for University Hospital Zurich)
<u>GEOCODE_LON</u>	numeric	Longitude (e.g. 8.549156 for University Hospital Zurich)
<u>RURAL</u>	numeric: <ul style="list-style-type: none"> <li>• 1 = Urban</li> <li>• 2 = Mostly urban</li> <li>• 3 = Mostly rural</li> <li>• 4 = Rural</li> <li>• 9 = Unknown</li> </ul>	Code for the site situation (facility location)
<u>LEVEL</u>	numeric: <ul style="list-style-type: none"> <li>• 1 = Health centre</li> <li>• 2 = District hospital</li> <li>• 3 = Regional, provincial or university hospital</li> <li>• 9 = Unknown</li> </ul>	Code for level of care
<u>ADULTPED</u>	character: "PED", "ADULT", or "BOTH"	Population the center serves
<u>OPEN_D</u>	yyyy-mm-dd	(Optional) Date of opening of dataset: earliest date for which data were included from this site
<u>CLOSE_D</u>	yyyy-mm-dd	(Optional) Date of closing of dataset
<u>ADD_CENTER</u>	yyyy-mm-dd	Inclusion date: date that the site was added to the cohort
<u>DROP_CENTER</u>	yyyy-mm-dd	(Optional) Exclusion date: date that the site was dropped from the cohort

## tbICEP - Clinical Events and Procedures

holds type and date of adverse events including serious non-AIDS conditions

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">EVENT_ID</a>	numeric	Unique Event Identifier (foreign key to the different event tables)
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">CEP_D</a>	yyyy-mm-dd	date of event
<a href="#">CEP_ID</a>	character. see <a href="#">coding table</a> for valid codings.	identifies type of event
<a href="#">CEP_SPEC</a>	character. see <a href="#">coding table</a> for valid codings.	further specification
<a href="#">CEP_V</a>	numeric. See <a href="#">coding table</a> for interpretation.	Depending on CEP_ID and CEP_SPEC: value of given event
<a href="#">SRCDOC_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> </ul>	whether the source documentation is available
<a href="#">SRCDOC_D</a>	yyyy-mm-dd	date for source documentation verification
<a href="#">VERIFY_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> </ul>	Has the monitor verified the source documentation?
<a href="#">VERIFY_D</a>	yyyy-mm-dd	date for monitor verification
<a href="#">APPROV_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> </ul>	final verification/approval
<a href="#">APPROV_D</a>	yyyy-mm-dd	final verification date
<a href="#">APPROV_S</a>	character	signature for final verification

### Additional fields

Field name	Format	Description
<a href="#">CEP_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	has the patient had an event?
<a href="#">CEP_NAME</a>	character	full name of the event
<a href="#">CEP_DESCRIP</a>	character	full description of the event
<a href="#">CEP_R_Y</a>	numeric: <ul style="list-style-type: none"> <li>0 = not related</li> <li>1 = definitive</li> <li>2 = remote/unlikely</li> <li>3 = possible</li> <li>4 = probable</li> </ul>	relation to treatment

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES

AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES

## CEP\_ID field

containing table

[tblCEP](#)

explanation of variable

Code to identify event

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Event
AMI	Acute myocardial infarction
ASCI	Ascites
ASP	Invasive aspergillosis
AVN	Avascular necrosis in the femoral head
BACT	Bacteremia
BART	Bartonellosis
CERC	Cervical cerclage
CHAG	Chagas disease (American trypanosomiasis) of the CNS
CLD	Chronic liver disease
DIA	Diabetes mellitus
ENDO	Endocarditis
ESRD	End stage renal disease
FAT	Fatal case with insufficient data
FIBS	Fibroscan stiffness (add elasticity value in CEP_V)
FRA	Bone fracture (irrespective of location)
HEP	Hepatic encephalopathy
HESY	Hepatorenal syndrome
HOSP	Hospitalisation
ICP	Invasive Cardiovascular Procedures
ICU	Admission for the ICU
JAUN	Jaundice
LAC	Lactic acidosis
LEIS	Leishmaniasis, visceral
LIVB	Liver biopsy
LIVD	Liver decompensation
LIVT	Liver transplantation
MENI	Meningitis
MCDI	Microsporidiosis diarrhoes (dur. > 1 month)
NADM	Non-AIDS defining malignancies
NOCA	Nocardiosis
OESO	Oesophageal variceal bleeding
OSTI	Ostitis
PAN	Pancreatitis
PCE	Pneumocystis carinii extrapulmonary
PMAR	Penicillium marneffeii, disseminated
PERI	Spontaneous bacterial peritonitis
PNEU	Pneumonia
PYEL	Pyelonephritis
REQU	Rhodococcus equi disease
STR	Stroke (infarction or haemorrhagia)

USAB	Ultrasound imaging of the abdomen
------	-----------------------------------

### Coding Table: Pregnancy-related adverse events

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Event
ANEM	Anemia (<10g/l)
FEV	Fever (> 38 0C and > 1 day)
URITINF	Urinary tract infection
HEMATOMA	Hematoma
ENDOM	Endometritis
WOUINF	Wound infection
PERIT	Peritonitis
PNEU	Pneumonia
SEPSIS	Sepsis (fever and pos blood culture)
THROMB	Thromboembolism
DIC	Disseminated intravascular coagulation (DIC)
SUBI	Subileus/ ileus
HEMOR	Hemorrhage
PSY	Psychosis
PREECL	Preeclampsia/eclampsia
ANEMBL	Severe anemia requiring blood transfusion

### Coding Table: Early childhood-related adverse events

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Event
DEV	Developmental delay
SEIZ	Seizures
ONEU	Other neurological symptoms (use with AE_DESCRIP)
OABN	Other abnormal findings (use with AE_DESCRIP)

### Case Definitions

see [separate article](#).

Case Definitions for [AE\\_ID](#)

HICDEP Code	ICD-10 codes	Adverse Event	Definition
AMI	I21.9	Acute myocardial infarction	Definitive myocardial infarction (MI) i) definitive electrocardiogram (ECG), ii) symptoms together with probable ECG and abnormal enzymes, iii) typical symptoms, abnormal enzymes and ischaemic/non-codable/not available ECG, or iv) fatal cases with naked-eye appearance of fresh MI and/or recent coronary occlusion found at necropsy. Please see the <a href="#">?MONICA manual</a> for further criteria.
STR	I64.9	Stroke, not specified as haemorrhage or infarction	Rapidly developed clinical signs of focal or global disturbance of cerebral function lasting more than 24 hours (unless interrupted by surgery or death), with no apparent cause other than a cardiovascular origin. Secondary stroke caused by trauma should be excluded. The differentiation between infarction and haemorrhage should be based on results of cerebral scanning or necropsy. In case of uncertainty (results not interpretable, or test not performed), please indicate so on the event form. Please see the <a href="#">?MONICA manual</a> and the <a href="#">?DAD MOOP (Manual of Operations)</a> for further criteria
DIA	E14 (also E10 ? insulin dependent and E11 non-insulin-dependent)	Unspecified diabetes mellitus	The diagnostic criteria is: fasting blood glucose > 7 mmol/l Please see the ADA (the American Diabetes Association) criteria for classification.
ICP - BYP	n/a	Coronary artery by-pass grafting	Procedure
ICP - END	n/a	Carotid endarterectomy	Procedure
ICP - ANG	n/a	Coronary angioplasty/stenting	Procedure
LAC		Lactate acidosis	Elevated S-lactate > 2.5 mM (>2.3 mg/dL) AND plasma pH < 7.35 (alternatively: Bicarbonate/HCO <sub>3</sub> ≤ 20 mM (≤ 20 meq/L)) AND otherwise unexplained recent onset of at least one of the following: Abdominal distension, anorexia, abdominal pain, nausea, vomiting, diarrhea, increased liver function enzymes, jaundice, dyspnea, fever, neuropathy, generalized weakness, ascending neuromuscular weakness, myalgias, paresthesias, weight loss or hepatomegaly.
PAN		Pancreatitis	Typical clinical history (i.e. severe abdominal pain), plus one or more of the following: elevated serum amylase > 1.5x ULN, elevated serum lipase, radiological findings.
ESRD	N18.0 (N18.8/9, N25.9, N26, N0.5, N04, N08)	End stage renal disease	A. Hemodialysis or peritoneal dialysis expected to last at least three months, documented in a clinical note B. A kidney transplant, documented in a clinical note Confirmed: A or B Probable: Not applicable
AVN		Avascular necrosis in the femoral head	Diagnosed by the combination of clinical symptoms (pain, walking difficulties) and imaging findings (MRI, bone scintigraphy)
FRA	Several depending on location	Bone fracture	Diagnosed by X-ray
HEP		Severe hepatic encephalopathy (stage III or IV)	Stage III: marked confusion, incoherent speech, asterixis, sleeping but arousable - Stage IV: coma

CLD		Chronic liver disease ?severe clinical manifestations	<p>A. 1. Clinical symptoms of end-stage liver failure in patients with chronic liver disease, based on the diagnosis documented in a clinical note of either</p> <p>(i) bleeding from gastric or esophageal varices</p> <p>(ii) hepatic encephalopathy stage III or IV</p> <p>(iii) hepatorenal syndrome</p> <p>A. 2 liver transplantation documented in a clinical note</p> <p>B. Pathology report or fibro-scan report documenting severe liver fibrosis or cirrhosis (Metavir F3 or F4 or fibroscan liver stiffness <math>\geq</math> 8 kPa)</p> <p>Confirmed: A1 and B; or A2</p> <p>Probable: A1</p>
NADM		Non AIDS defining cancers	<p>A. Diagnosis of cancer (other than: AIDS defining (non-Hodgkin?s lymphoma, Kaposi's sarcoma ), or invasive cervical cancer); and basal and squamous cell skin cancers) in a pathology report that established the diagnosis</p> <p>B. Diagnosis of cancer (other than: AIDS defining (non-Hodgkin?s lymphoma, Kaposi's sarcoma, or invasive cervical cancer); and basal and squamous cell skin cancers) in a hospital discharge summary or consultation note from the hospitalization or clinic visit during which the diagnosis was established</p> <p>C. In the absence of A or B: Strong suspicion of cancer supported by (i) evidence from radiological or other imaging technique, (ii) or biochemical assay</p> <p>D. In the absence of A, B or C: Strong suspicion of cancer by visual inspection (e.g. skin metastasis, suspected malignant melanoma, tissue growth resembling cancer visualized during endoscopy/anoscopy) not explained by other known conditions.</p> <p>Confirmed: A or B</p> <p>Probable: C</p> <p>Possible: D</p> <p>* The date of diagnosis is the month, day and year the tumor was first diagnosed by a recognized medical practitioner, whether clinically or microscopically confirmed.</p>



## CEP\_SPEC field

containing table

[tblCEP](#)

explanation of variable

Code to further specify the event identified by [CEP\\_ID](#).

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

Note: The codes for NADM are tentative and subject to change for the next release.

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code (CEP_ID)	Code (CEP_SPEC)	Description
AMI	DAMI	Definitive Myocardial infarction
AMI	PAMI	Possible Myocardial infarction
LIVB	F0	no fibrosis
LIVB	F1	portal fibrosis without septa
LIVB	F2	portal fibrosis with few septa
LIVB	F3	numerous septa without cirrhosis
LIVB	F4	cirrhosis
HEP	I	Hepatic encephalopathy stage I
HEP	II	Hepatic encephalopathy stage II
HEP	III	Hepatic encephalopathy stage III
HEP	IV	Hepatic encephalopathy stage IV
HEP	III+IV	Hepatic encephalopathy stage III or IV
ICP	ANG	Invasive Cardiovascular Procedures: Coronary angioplasty/stenting
ICP	BYP	Invasive Cardiovascular Procedures: Coronary artery by-pass grafting
ICP	END	Invasive Cardiovascular Procedures: Carotid endarterectomy
NADM	ALL	Leukemia: Acute lymphoid
NADM	AML	Leukemia: Acute myeloid
NADM	ANAL	Anal dysplasia, grade 2 or higher
NADM	ANUS	Anal cancer
NADM	BLAD	Bladder cancer
NADM	BRCA	Breast cancer
NADM	BRAIN	Brain cancer
NADM	CERV	Cervical dysplasia/carcinoma in situ, grade 2 or higher
NADM	CLL	Leukemia: Chronic lymphoid
NADM	CML	Leukemia: Chronic myeloid
NADM	COLO	Colon cancer
NADM	COTC	Connective tissue cancer
NADM	HDL	Hodgkin lymphoma
NADM	HENE	Head and neck cancer
NADM	KIDN	Kidney cancer
NADM	LEUK	Leukemia: unspecified
NADM	LIPC	Lip cancer
NADM	LIVR	Liver cancer
NADM	LUNG	Lung cancer
NADM	MALM	Malignant melanoma
NADM	MEAC	Metastasis: of adenocarcinoma
NADM	MEOC	Metastasis: of other cancertype

NADM	MESC	Metastasis: of squamous cell carcinoma
NADM	META	Metastasis: unspecified
NADM	MULM	Multiple myeloma
NADM	PENC	Penile cancer
NADM	PROS	Prostate cancer
NADM	RECT	Rectum cancer
NADM	STOM	Stomach cancer
NADM	TESE	Testicular seminoma
NADM	UTER	Uterus cancer
STR	SHAE	Stroke: Haemorrhagia
STR	SINF	Stroke: Infarction
STR	SUNK	Stroke: Unknown

## tbDELIVERY\_CHILD - Delivery information related to the child(ren)

This table holds information about a delivery **specific to a child**. Mother-specific data is kept in [tbDELIVERY\\_MUM](#).

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>MOTHER_ID</b>	Character (or numeric if possible)	Patient ID of mother of the child
<b>MEMRUP_D</b>	yyyy-mm-dd	Date of rupture of membranes
<b>CHILD_ID</b>	Character (or numeric if possible)	Patient ID of the child
<b>B_SEQ</b>	numeric	If multiple births, indicate number (1=first born)
<b>DELIV_D</b>	yyyy-mm-dd	Date of delivery
<b>DELIV_T</b>	hh:mm	Time of delivery
<b>DELIV_M</b>	character: <ul style="list-style-type: none"> <li>1=Vaginally, spontaneous</li> <li>2=Vaginally, forceps</li> <li>3=Vaginally, vacuum</li> <li>10= Cesarean section, primary/elective (before onset of labour and rupture of membrane)</li> <li>11=Cesarean section, Secondary</li> </ul>	Mode of delivery
<b>LABOUR_P</b>	numeric	Duration of labour in hours (from cervical dilatation (>=3 cm) until delivery)
<b>BREECH_Y</b>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Was the child born from a breech presentation?

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbDELIVERY_CHILD	CrossTable	DCC001	MOTHER_ID doesn't exist in tbIPREG		YES
tbDELIVERY_CHILD	CrossTable	DCC002	CHILD_ID doesn't exist in tbIPREG_OUT		YES
tbDELIVERY_CHILD	WithinTable	DCW001	Duplicate records for same Mother_ID, CHILD_ID		YES
tbDELIVERY_CHILD	WithinTable	DCW002	B_SEQ >4		YES
tbDELIVERY_CHILD	WithinTable	DCW003	B_SEQ>1, but no record exists for same Mother, Child with B_SEQ=1		YES

tblDELIVERY_CHILD	WithinTable	DCW005	DELIV_D < MEMRUP_D		YES
tblDELIVERY_CHILD	WithinTable	DCW004	LABOUR_P >72		YES
tblDELIVERY_CHILD	WithinTable	DCW006	2 children, same mother, same MEMRUP_D, DELIV_D >DELIV_D , yet B_SEQ < B_SEQ		YES

## tbDELIVERY\_MUM - Delivery information related to the mother

This table contains information about the delivery **specific to the mother**. Child-specific information is recorded in [tbDELIVERY\\_CHILD](#) instead.

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>MOTHER_ID</b>	Character (or numeric if possible)	Patient ID of mother of the child
<b>PREG_SEQ</b>	numeric	Sequence number of the pregnancy for the specified mother
<b>MEMRUP_D</b>	yyyy-mm-dd	Date of rupture of membranes
<b>MEMRUP_T</b>	hh:mm	Time of rupture of membranes
<b>LABOUR</b>	character: <ul style="list-style-type: none"> <li>1=Spontaneous</li> <li>2=Induced</li> <li>3=No labour (elective C-section)</li> </ul>	Onset of labor
<b>INTERV</b>	character: <ul style="list-style-type: none"> <li>0=No</li> <li>1=Fetal blood sampling FBS</li> <li>2=Internal electrodes</li> <li>3=1+2</li> <li>90=Other</li> <li>91=1+Other</li> <li>92=2+Other</li> <li>93=1+2+Other</li> </ul>	Interventions during delivery
<b>INTERV_O</b>	character	Interventions during delivery - other
<b>TEAR_Y</b>	numeric: <ul style="list-style-type: none"> <li>1=Yes</li> <li>0=No</li> <li>9=Unknown</li> </ul>	Episiotomy/tear
<b>BLDLOSS</b>	numeric (mL)	Estimated blood loss during delivery
<b>CONTREAT</b>	character: <ul style="list-style-type: none"> <li>0=No, treatment has been interrupted</li> <li>1=Yes, at the foreseen intervals</li> <li>2=Yes, but not at foreseen time points</li> <li>9=Unknown</li> </ul>	Did the patient continue the usual antiretroviral therapy?
<b>DISCHA_D</b>	yyyy-mm-dd	Date of discharge from hospital

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES

tblDELIVERY_MUM	CrossTable	DMC001	MOTHER_ID doesn't exist in tblPREG		YES
tblDELIVERY_MUM	CrossTable	DMC002	MOTHER_ID doesn't exist in tblDELIVERY_CHILD with same MEMRUP_D		YES
tblDELIVERY_MUM	WithinTable	DMW001	PREG_SEQ invalid (PREG_SEQ>1 and PREG_SEQ=x, but PREG_SEQ=x-1 doesn't exist)		YES
tblDELIVERY_MUM	WithinTable	DMW002	DISCHA_D < MEMRUP_D		YES
tblDELIVERY_MUM	WithinTable	DMW004	INTERV=0-3 but INTERV_O non null		YES
tblDELIVERY_MUM	WithinTable	DMW003	INTERV=90,91, 92 or 93 but INTERV_O null		YES

## tbIDIS - CDC-C diseases

holds type and date of CDC-C diseases and malignancies.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">DIS_ID</a>	character. see <a href="#">coding table</a> for valid codings.	Code to identify event
<a href="#">DIS_D</a>	yyyy-mm-dd	Date of event
<a href="#">DIS_WD</a>	numeric. see <a href="#">coding table</a> for valid codings.	Means of diagnosis
<a href="#">DIS_OTH</a> <sup>1</sup>	character	Other location, only to be filled out if code alone is not sufficient

<sup>1</sup> DIS\_OTH might be part of the record's unique identification

### Additional fields

Please see [tbICEP](#) for specification on optional fields.

Field name	Format	Description
<a href="#">DIS_ED</a>	yyyy-mm-dd	End of opportunistic infection

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbIDIS	WithinTable	DW001	Duplicate records for same DIS_ID and same DIS_D		YES
tbIDIS	WithinTable	DW002	Miscoded DIS_WD as codes on table definition		YES
tbIDIS	WithinTable	DW003	Miscoded DIS_ID - as in code list attached to table definition		YES
tbIDIS	WithinTable	DW004	DIS_D missing		YES
tbIDIS	WithinTable	DW005	DIS_ID missing		YES
tbIDIS	WithinTable	DW006	Same event recorded twice - 2 records, same DIS_ID, DIS_D within 6 months		YES
tbIDIS	WithinTable	DW007	DIS_ED present but before DIS_D		YES
tbIDIS	CrossTable	DC001	Patient has no record in BAS		YES
tbIDIS	CrossTable	DC002	AIDS-defining records, yet AIDS=0 in tbIBAS		YES

tblDIS	CrossTable	DC003	First AIDS-defining DIS_D not equal to AIDS_D in tblBAS		YES
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## DIS\_ID field

containing table

[tblDIS](#)

explanation of variable

Code to identify event

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Severe Opportunistic Infection/Malignancies
DEM	AIDS dementia complex
BCNE	Bacterial pneumonia, recurrent (>2 episodes within 1 year)
CANO	Candidiasis, oesophageal, bronchi, trachea, or lungs
COCC	Coccidioidomycosis, disseminated or extrapulmonary
CRCO	Cryptococcosis, extrapulm.
CRSP	Cryptosporidiosis (duration > 1 month)
CMVR	Cytomegalovirus (CMV) chorioretinitis
CMVO	CMV ? other location
HERP	Herpes simplex virus ulcers (duration > 1 month) or pneumonitis/esophagitis
HIST	Histoplasmosis, extrapulm.
WAST	HIV Wasting Syndrome
ISDI	Isosporiasis diarrhoea (duration > 1 month)
MC	Mycobact. avium complex (MAC) or Kanasii, extrapulm.
MCP	Mycobact. tuberculosis pulm.
MCX	Mycobact. tuberculosis extrapulm
MCPO	Mycobact. pulm., other
MCXO	Mycobact. extrapulm., other
PCP	Pneumocystis carinii pneumonia (PCP)
LEU	Progressive multifocal leucoencephalopathy
SAM	Salmonella bacteraemia (non-typhoid) (recurrent)
TOX	Toxoplasmosis, brain
FBLS	Focal Brain lesion
KS	Kaposi Sarcoma
NHG	Non-Hodgkin Lymphoma -not specified
NHGB	Non-Hodgkin Lymphoma ? Burkitt (Classical or Atypical)
NHGI	Non-Hodgkin Lymphoma ? Diffuse large B-cell lymphoma (Immunoblastic or Centroblastic)
NHGU	Non-Hodgkin Lymphoma - Unknown/other histology
NHGP	Non-Hodgkin Lymphoma - Primary Brain Lymphoma
CRVC	Cervical Cancer

## Coding Table: Mother-to-child and paediatric specific

[Download this table as CSV file](#), [Download this table as XML file](#)

Stage	Code	CDC disease description
A	CA-LYM	Lymphadenopathy
A	CA-HEY	Hepatomegaly
A	CA-SPL	Splenomegaly
A	CA-DER	Dermatitis
A	CA-PAR	Parotitis
A	CA-URI	Recurrent or persist. UR infection, sinusitis, or otitis media

B	CB-ANE	Anemia
B	CB-BMP	Bacterial meningitis, pneumonia, or sepsis
B	CB-CAN	Candidiasis oropharyngeal for >2 months (age>6)
B	CB-CMY	Cardiomyopathy
B	CB-CMN	CMV onset before 1 month
B	CB-DIA	Diarrhea (recurrent or chronic)
B	CB-FEV	Fever (lasting >1 month)
B	CB-HEP	Hepatitis
B	CB-HSS	Herpes simplex stomatitis (>2 episodes in 1 year)
B	CB-HCV	HSV bronchitis, pneumonitis, esophagitis (<1 month)
B	CB-HZO	Herpes zoster, multidermatomal or relapse
B	CB-LEI	Leiomyosarcoma
B	CB-LYM	Lymphoid interstitial pneumonia or pulmonary lymphoid hyperplasia complex
B	CB-NEP	Nephropathy
B	CB-NOC	Nocardiosis
B	CB-TON	Toxoplasmosis (start before 1 month)
B	CB-VAR	Varicella, disseminated
C	CC-EPD	Pneumocystis disease, extrapulmonary
C	CC-TOD	Toxoplasmosis disseminated
C	CC-COM	Cryptococcal meningitis
C	CC-GEN	M. genavense disease
C	CC-LOB	Lymphoma, primary, cerebral
C	CC-ICC	Carcinoma, cervical, invasive
C	CC-ILE	Intracerebral lesions, indetermined

### Case definitions

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Severe Opportunistic Infection/Malignancies	Definitive/Autopsy or presumptive?	Definition
DEM	AIDS dementia complex	D	Disabling cognitive and/or motor dysfunction, or milestone loss in a child, and no other causes by CSF exam and brain imaging or by autopsy
DEM	AIDS dementia complex	P	Same as above but no CSF and brain imaging performed
BCNE	Bacterial pneumonia, recurrent (>2 episodes within 1 year)	D	New X-ray evidence not present earlier and culture of pathogen that typically causes pneumonia (other than P .carinii or M. tuberculosis)
BCNE	Bacterial pneumonia, recurrent (>2 episodes within 1 year)	P	Acute radiological findings (new X-ray evidence not present earlier) and acute clinical findings
CANO	Candidiasis, oesophageal, bronchi, trachea, or lungs	D/A	Gross inspection by endoscopy/autopsy or by microscopy (histology)
CANO	Candidiasis, oesophageal, bronchi, trachea, or lungs	P	Recent onset retrosternal pain on swallowing and confirmed oral or pharyngeal candidiasis
CRCO	Cryptococcosis, extrapulm.	D/A	Microscopy, culture of, or antigen detection in affected tissue
CRSP	Cryptosporidiosis (duration > 1 month)	D/A	Microscopy. Duration of diarrhoea for more then 1 month
CMVR	Cytomegalovirus (CMV) chorioretinitis	P	Loss of vision and characteristic appearance on serial ophthalmoscopy, progressing over serial months
CMVO	CMV ? other location	D/A	Microscopy (histology or cytology)

HERP	Herpes simplex virus ulcers (duration > 1 month) or pneumonitis/esophagitis	D	Microscopy, culture of, or antigen detection in affected tissue
HIST	Histoplasmosis, extrapulm.	D/A	Microscopy, culture of, or antigen detection in affected tissue
WAST	HIV Wasting Syndrome	D	Weight loss (over 10% of baseline) with no other cause, and 30 days or more of either diarrhoea or weakness with fever
ISDI	Isosporiasis diarrhoea (duration > 1 month)	D/A	Microscopy (histology or cytology). Duration of diarrhoea for more than 1 month
LEIS	Leishmaniasis, visceral	D/A	Histology or culture of Leishmania amastigotes in bone marrow or detection of amastigotes in tissue/fluid from affected organ in a patient with symptoms and signs consistent with disseminated Leishmaniasis
MCDI	Microsporidiosis diarrhoea (dur. > 1 month)	D/A	Stool microscopy or rectal biopsy in patient with persistent diarrhoea
MC	Mycobact. avium complex (MAC) or Kanasil, extrapulm.	D	Culture
MCP	Mycobact. tuberculosis pulm.	D	Culture
MCX	Mycobact. tuberculosis extrapulm	D	Culture
MCPO	Mycobact. pulm., other	D	Culture (indicate type)
MCPO	Mycobact. pulm., other	P	Acid fast bacteria (species not identified by culture) in sputum
MCXO	Mycobact. extrapulm., other	D	Culture (indicate type)
MCXO	Mycobact. extrapulm., other	P	Acid fast bacteria (species not identified by culture) on microscopy of normally sterile body fluid/tissue
PCP	Pneumocystis carinii pneumonia (PCP)	D	Microscopy (histology or cytology)
PCP	Pneumocystis carinii pneumonia (PCP)	P	Recent onset of dyspnoea on exertion or dry cough, and diffuse bilateral infiltrates on chest X-ray and pO <sub>2</sub> <70 mmHg and no evidence of bacterial pneumonia
LEU	Progressive multifocal leucoencephalopathy	D/A	Microscopy (histology or cytology)
LEU	Progressive multifocal leucoencephalopathy	P	Progressive deterioration in neurological function and CT/MR scan evidence
SAM	Salmonella bacteraemia (non-typhoid) (recurrent)	D	Culture
TOX	Toxoplasmosis, brain	D	Microscopy (histology/cytology)
TOX	Toxoplasmosis, brain	P	Recent onset focal neurological abnormalities or reduced level of consciousness, and mass effect lesion on scan, and specific therapy response
FBLS	Focal Brain lesion	?	TODO:: To be updated ASAP
KS	Kaposi Sarcoma	D/A	Histology
KS	Kaposi Sarcoma	P	Characteristic erythematous/violaceous plaque-like lesion on skin or mucous membranes
NHG	Non-Hodgkin Lymphoma -not specified	?	TODO:: To be updated ASAP
NHGB	Non-Hodgkin Lymphoma ? Burkitt (Classical or Atypical)	D	Histology
NHGI	Non-Hodgkin Lymphoma ? Diffuse large B-cell lymphoma (Immunoblastic or Centroblastic)	D	Histology
NHGU	Non-Hodgkin Lymphoma - Unknown/other histology	?	TODO:: To be updated ASAP
NHGP	Non-Hodgkin Lymphoma - Primary Brain Lymphoma	D	TODO:: To be updated ASAP

NHGP	Non-Hodgkin Lymphoma - Primary Brain Lymphoma	P	Recent onset of focal neurological symptoms and signs or reduced level of consciousness, CT/MR scan evidence of a lesion or lesions having mass effect, no response to toxo therapy, no evidence of lymphoma outside the brain
CRVC	Cervical Cancer	D/A	Histology

## DIS\_WD field

containing table

[tblDIS](#)

explanation of variable

Means of diagnosis

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Means of diagnosis
1	Definitive diagnosis
2	Presumptive diagnosis
3	Diagnosis from autopsy
4	Diagnosis from registry

## tblLAB - Laboratory values

holds type, date, value and unit of laboratory tests.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">LAB_ID</a>	character. see <a href="#">coding table</a> for valid codings.	Code representing the measurement
<a href="#">LAB_D</a>	yyyy-mm-dd	Date of measurement/sample
<a href="#">LAB_V</a>	numeric: -1 = undetectable or detection limit as negative value  for dipstick results:  <ul style="list-style-type: none"> <li>• 0 = Negative</li> <li>• 9 = Trace</li> <li>• 1 = 1+</li> <li>• 2 = 2+</li> <li>• 3 = 3+</li> <li>• 4 = 4+</li> </ul>	Value of measurement
<a href="#">LAB_U</a>	numeric. see <a href="#">coding table</a> for valid codings.	Unit of measurement

### Additional fields

Other detailed information regarding the patient and the measurement would be relevant, like the proposed fasting information shown below.

Field name	Format	Description
<a href="#">LAB_FA</a>	numeric:  <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	Was the blood sample taken while fasting?
<a href="#">LAB_ST</a>	character:  <ul style="list-style-type: none"> <li>• WB = Whole Blood</li> <li>• P = Plasma</li> <li>• S = Serum</li> <li>• U24 = 24h Urine</li> <li>• U = Urine</li> </ul>	Specimen type

Depending on the set of measurements collected and the mandatory fields applicable to these individual measurements, it might be useful to separate the LAB table into several sub tables. This is already shown for the CD4 and RNA measurements: the level of detail needed for CD4 is less than for the LAB variables in general (no unit since it's always the same), while for RNA the data required is more detailed (assay and detection limit).

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblLAB	WithinTable	LW001	Duplicate records		YES

tblLAB	WithinTable	LW002	Missing LAB_D		YES
tblLAB	WithinTable	LW003	Missing LAB_V		YES
tblLAB	WithinTable	LW004	Missing LAB_U		YES
tblLAB	WithinTable	LW005	Missing LAB_ID		YES
tblLAB	WithinTable	LW008	Missing LAB_FA		YES
tblLAB	WithinTable	LW009	Missing LAB_ST		YES
tblLAB	WithinTable	LW010	LAB_V out of range for unit LAB_U		YES
tblLAB	CrossTable	LC001	Patient has no record in BAS table		YES

## LAB\_ID field

containing table

[tblLAB](#)

explanation of variable

Code representing the measurement

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Measurement
ACRA	Albumin Creatinin Ratio
ALB	Albumine
AFP	Alfa Fetoprotein
ALP	Alkaline Phosphatase
ALT	Alanin-Aminotransferase
AMY	Amylase
AST	Aspartat aminotransferase
BIL	Total Bilirubin
CD3	CD3
CD3P	% CD3 of leukocytes
CD8	CD8
CD8P	% CD8 of leukocytes
CHOL	Total Cholesterol
CL-	Cl-
CRE	Creatinine
DIPP	Dipstick result for protein in Urine
GGT	Gamma-glutamyltransferase
GLUC	Glucose
HAEM	Haemoglobin
HDL	Serum HDL
HEMA	Hematocrit
INR	Quick/INR
LACT	Lactate
LEUK	Leukocytes
LYM	Lymphocytes
LYMP	% Lymphocytes of leukocytes
MCV	MCV
NA+	Na+
NEU	Neutrophils
PCRA	Protein Creatinin Ratio
PHA	PH arterial
PHV	PH venous
PP	PP factor (II, VII, X)
PROT	Protein
PSA	Prostate-specific antigen
PTH	Parathyroid Hormone
PTR	Prothrombin rate
THR	Thrombocytes
TRIG	Serum Triglyceride
URA	Uric acid



## LAB\_U field

containing table

[tblLAB](#)

explanation of variable

Unit of measurement

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Unit Code	Unit String
1	mmol/L
2	g/L
3	g/dL
4	mg/dL
5	IU/L (u/L)
6	µmol/L
7	INR
8	1E+9/L
9	1E+6/L
10	cells/µL
11	µkat/L
12	%
13	µg/L = ng/mL
14	mg/24h
15	mg/mmol
16	fl (Femtoliter)
17	µg/mL = mg/L
99	no units (e.g. for Dipstick results)

It is recommended to use the string codes from the above table since this makes the data human readable.

## tbILAB\_BP - Laboratory values - Blood pressure

holds date, diastolic and systolic values and unit of blood pressure measurements.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<u><a href="#">PATIENT</a></u>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<u><a href="#">BP_D</a></u>	yyyy-mm-dd	Date of Measurement/Sample
<u><a href="#">BP_SYS</a></u>	numeric	Systolic Blood Pressure
<u><a href="#">BP_DIA</a></u>	numeric	Diastolic Blood Pressure
<u><a href="#">BP_U</a></u>	numeric. see <a href="#">coding table</a> for valid codings.	Unit of measurement

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES

## BP\_U field

containing table

[tblLAB\\_BP](#)

explanation of variable

Unit of measurement

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Unit for blood pressure
1	mmHg
2	cmHg
3	Kpa

## tbILAB\_CD4 - Laboratory values

holds date and value of CD4 measurements.

Note: If needed, a CD8 table (tbILAB\_CD8) could be formed from the same structure.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">CD4_D</a>	yyyy-mm-dd	Date of measurement
<a href="#">CD4_V</a>	numeric (per microliter): -1 = undetectable or detection limit as negative value	Value of CD4 measurement

### Additional fields

[CD4\\_V](#) is assumed to contain absolute CD4 cell counts per mL as standard. In case CD4 % (with respect to CD45+ lymphocytes as denominator) should be collected as well, please append the following field to the table:

Field name	Format	Description
<a href="#">CD4_U</a>	numeric with codes (or full string): <ul style="list-style-type: none"> <li>1 = cells/<math>\mu</math>l</li> <li>2 = %</li> </ul>	Unit of measurement

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbILAB_CD4	WithinTable	CW001	CD4 value out of range		YES
tbILAB_CD4	WithinTable	CW002	Duplicate records for same date		YES
tbILAB_CD4	WithinTable	CW003	Missing CD4_D		YES
tbILAB_CD4	WithinTable	CW004	Missing CD4_V		YES
tbILAB_CD4	WithinTable	CW005	Missing CD4_U		YES
tbILAB_CD4	WithinTable	CW006	Miscoded CD4_U as coding list on table definition		YES
tbILAB_CD4	WithinTable	CW007	CD4_U=2 (percentage) and CD4_V>100		YES
tbILAB_CD4	WithinTable	CW008	CD4_U=(1 or 3) and CD4_V>3000		YES
tbILAB_CD4	WithinTable	CW009	CD4 counts spike up or down suddenly (large change in less than a year)		YES
tbILAB_CD4	CrossTable	CC001	Patient has no record in BAS		YES

## tbLAB\_RES - Resistance testing

holds background information on the resistance test, laboratory, library, kit, software and type of test

Note: This table is tightly linked to [tbLAB\\_RES\\_LVL\\_1](#), [tbLAB\\_RES\\_LVL\\_2](#) and [tbLAB\\_RES\\_LVL\\_3](#).

**Resistance should be reported at lowest level of interpretation possible ? so if the nucleotide sequence is available this should be reported rather than the list of mutations or resistance scores. However, the resistance test results should be captured if they have been part of the physician's treatment decisions for the patient.**

**Non-amplifiable resistance tests should not be reported.**

These four tables are designed to capture several possible formats the clinics and cohorts might have recorded resistance test data in. Once this data is gathered it should like all other tables be quality assessed.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">TEST_ID</a>	character (or numeric if possible)	An arbitrary value identifying a resistance test result
<a href="#">SAMPLE_D</a>	YYYY-mm-dd	Date of the actual sample taken (NOT the test date)
<a href="#">SEQ_DT</a>	YYYY-mm-dd hh:mm	Date and time when the sequencing was performed
<a href="#">LAB</a>	character	Name of laboratory where the test was performed
<a href="#">LIBRARY</a>	character	Library/algorithm used to identify resistance mutations
<a href="#">REFSEQ</a>	character	Name/identifier of reference strain used to find mutations
<a href="#">KIT</a>	character	Vendor and version/name of the kit used for the test
<a href="#">SOFTWARE</a>	character	Software and version used to determine resistance
<a href="#">TESTTYPE</a>	numeric: <ul style="list-style-type: none"> <li>• 1 = Genotype</li> <li>• 2 = Phenotype</li> <li>• 9 = Other</li> </ul>	Type of test
<a href="#">VIRUSTYPE</a>	numeric: <ul style="list-style-type: none"> <li>• 1 = HIV</li> <li>• 2 = HCV</li> </ul>	Type of Virus
<a href="#">SUBTYPE</a>	character	Subtype of HIV- or HCV-RNA

### Additional fields

As shown with the core fields, the *TEST\_ID* is the link between the 3 levels of data and the test background information table.

Some prior assessment of the assigned test identifiers has to be done in order to avoid duplicates.

In a running database the duplicate issues are easily resolved by adding a unique auto-generated key as the identifier between 3 levels of data and the test background information table.

Along with the *TEST\_ID* it might be necessary to store the ID assigned to the sample at both the testing laboratory but also the centres laboratory in order to track the sample.

Field name	Format	Description
<a href="#">SAMP_LAB</a>	character (or numeric if possible)	The assigned sample ID at the lab where the resistance test is performed.
<a href="#">SAMP_INT</a>	character (or numeric if possible)	The assigned sample ID from the centre.

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
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AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblLAB_RES	WithinTable	LRW001	Duplicate records for same patient on same date		YES
tblLAB_RES	WithinTable	LRW002	Missing PATIENT		YES
tblLAB_RES	WithinTable	LRW003	Missing TEST_ID		YES
tblLAB_RES	WithinTable	LRW004	Missing SAMPLE_D		YES
tblLAB_RES	WithinTable	LRW005	Missing SEQ_DT		YES
tblLAB_RES	WithinTable	LRW006	Missing LAB		YES
tblLAB_RES	WithinTable	LRW007	Missing LIBRARY		YES
tblLAB_RES	WithinTable	LRW008	Missing REFSEQ		YES
tblLAB_RES	WithinTable	LRW009	Missing KIT		YES
tblLAB_RES	WithinTable	LRW010	Missing SOFTWARE		YES
tblLAB_RES	WithinTable	LRW011	Missing TESTTYPE		YES
tblLAB_RES	WithinTable	LRW012	Missing SUBTYPE		YES
tblLAB_RES	WithinTable	LRW013	SEQ_DT has no time part		YES
tblLAB_RES	WithinTable	LRW014	Missing VIRUSTYPE		YES
tblLAB_RES	CrossTable	LRC001	PATIENT has no record in tblBAS		YES
tblLAB_RES	CrossTable	LRC002	This TEST_ID has both LVL_1 and LVL_2 records		YES

## tbILAB\_RES\_LVL\_1 - Nucleotide sequences (PRO, RT, GP41, GP120)

holds nucleoside sequence for the PRO and RT sequences. No entry is made if the test was a phenotype test.

Note: This table is tightly linked to [tbILAB\\_RES](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">TEST_ID</a>	character (or numeric if possible)	Identifier linking this record to <a href="#">tbILAB_RES</a>
<a href="#">SEQTYPE</a>	character: <ul style="list-style-type: none"> <li>PRO = PRO sequence</li> <li>RT = RT sequence</li> <li>GP41 = GP41 sequence</li> <li>GP120 = GP120 sequence</li> </ul>	Type of nucleotide sequence if available
<a href="#">SEQ_START</a>	numeric	Start position for the sequence
<a href="#">SEQ_STOP</a>	numeric	Stop position for the sequence
<a href="#">SEQ_NUC</a>	character	Nucleotide sequence if available

### Additional fields

In cases where the amino acid sequence is collected rather than the nucleotide sequence, the field *SEQ\_NUC* might be replaced with *SEQ\_AA*, which is the nucleotide sequence, expressed in an amino acid sequence:

Field name	Format	Description
<a href="#">SEQ_AA</a>	character	Amino acid sequence if available (empty if test was phenotype)

However using the amino acid sequence does not give the same detail of data as the nucleoside sequence: wobbles in the nucleoside sequence can either complicate the reading and alignment of the amino acid sequence or the wobbles can be lost and silent mutations are lost.

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbILAB_RES_LVL_1	WithinTable	L1W001	Duplicate records per TEST_ID and SEQTYPE		YES
tbILAB_RES_LVL_1	WithinTable	L1W002	SEQ_START > SEQ_STOP		YES
tbILAB_RES_LVL_1	WithinTable	L1W003	SEQ_NUC contains invalid IUPAC character		YES
tbILAB_RES_LVL_1	WithinTable	L1W004	Missing TEST_ID		YES
tbILAB_RES_LVL_1	WithinTable	L1W005	Missing SEQ_START		YES
tbILAB_RES_LVL_1	WithinTable	L1W006	Missing SEQ_STOP		YES
tbILAB_RES_LVL_1	WithinTable	L1W007	Missing SEQ_NUC		YES
tbILAB_RES_LVL_1	CrossTable	L1C001	TEST_ID not in tbILAB_RES.TEST_ID		YES

## tbILAB\_RES\_LVL\_2 - Mutations

holds mutations and positions of PRO and RT sequences.

Note: This table is tightly linked to [tbILAB\\_RES](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">TEST_ID</a>	character (or numeric if possible)	Identifier linking this record to <a href="#">tbILAB_RES</a>
<a href="#">GENE</a>	character: <ul style="list-style-type: none"> <li>PRO = PRO sequence</li> <li>RT = RT sequence</li> <li>GP41 = GP41 sequence</li> <li>GP120 = GP120 sequence</li> </ul>	Type of sequence/gene (PRO, RT, GP41, GP120)
<a href="#">AA_POS</a>	numeric	Position of the mutation in the sequence
<a href="#">AA_POS_SUB</a>	character: <ul style="list-style-type: none"> <li>a = first</li> <li>b = second</li> <li>etc.</li> </ul>	Subposition used to code insertions
<a href="#">AA_FOUNDED_1</a>	character. empty = Amino acid has been deleted.	Mutation (Amino acid) found in the sequence
<a href="#">AA_FOUNDED_2</a>	character. empty = Amino acid has been deleted.	Mutation (Amino acid) found in the sequence (if more than 1)
<a href="#">AA_FOUNDED_3</a>	character. empty = Amino acid has been deleted.	Mutation (Amino acid) found in the sequence (if more than 2)
<a href="#">AA_FOUNDED_4</a>	character. empty = Amino acid has been deleted.	Mutation (Amino acid) found in the sequence (if more than 3)

[AA\\_FOUNDED\\_#](#) could be extended if mixtures with more than 4 amino acids are found.

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbILAB_RES_LVL_2	WithinTable	L2W001	AA_FOUNDED_x but nothing in AA_FOUNDED(x-1)		YES
tbILAB_RES_LVL_2	WithinTable	L2W002	Duplicate records per TEST_ID, GENE, AA_POS and AA_POS_SUB		YES
tbILAB_RES_LVL_2	WithinTable	L2W003	Missing TEST_ID		YES
tbILAB_RES_LVL_2	WithinTable	L2W004	Missing GENE		YES
tbILAB_RES_LVL_2	WithinTable	L2W005	Missing AA_POS		YES
tbILAB_RES_LVL_2	WithinTable	L2W006	Missing AA_POS_SUB		YES
tbILAB_RES_LVL_2	WithinTable	L2W007	Missing AA_FOUNDED_1		YES



tblLAB_RES_LVL_2	CrossTable	L2C001	TEST_ID not in tblLAB_RES.TEST_ID		YES
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## tbILAB\_RES\_LVL\_3 - Resistance test result

holds resistance result in relation to antiretroviral drug.

Note: This table is tightly linked to [tbILAB\\_RES](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">TEST_ID</a>	character (or numeric if possible)	Identifier linking this record to <a href="#">tbILAB_RES</a>
<a href="#">ATC_CODE</a>	character	?ATC code of the medication
<a href="#">RES_SCOR</a>	character	Score of resistance or recommendation given from the test.

### Additional fields

For phenotype test results it will be necessary to extend the table with a field to store the cut-off value:

Field name	Format	Description
<a href="#">RES_CUT</a>	character	Cut-off value for phenotype test result

However using the amino acid sequence does not give the same detail of data as the nucleoside sequence: wobbles in the nucleoside sequence can either complicate the reading and alignment of the amino acid sequence or the wobbles can be lost and silent mutations are lost.

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbILAB_RES_LVL_3	WithinTable	L3W001	Duplicate records for same TEST_ID and ATC_CODE		YES
tbILAB_RES_LVL_3	WithinTable	L3W002	Missing TEST_ID		YES
tbILAB_RES_LVL_3	WithinTable	L3W003	Missing ATC_CODE		YES
tbILAB_RES_LVL_3	WithinTable	L3W004	Missing RES_SCOR		YES
tbILAB_RES_LVL_3	CrossTable	L3C001	TEST_ID not in tbILAB_RES.TEST_ID		YES

## tbILAB\_RNA - Laboratory values

holds date, value, detection limit and type of viral assay.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">RNA_D</a>	yyyy-mm-dd	Date of Measurement/Sample
<a href="#">RNA_V</a>	numeric: -1 = undetectable or detection limit as negative value	HIV-RNA measurement value
<a href="#">RNA_L</a>	numeric	Lower Limit of HIV-RNA Assay
<a href="#">RNA_T</a>	numeric. see <a href="#">coding table</a> for valid codings.	IF AVAILABLE, What type of VIRAL ASSAY was used for this measurement?

### Additional fields

Field name	Format	Description
<a href="#">RNA_UL</a>	numeric	IF AVAILABLE, Upper Limit of assay

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbILAB_RNA	WithinTable	RW001	Duplicate records for same RNA_D		YES
tbILAB_RNA	WithinTable	RW002	RNA_V < 0 and RNA_L missing		YES
tbILAB_RNA	WithinTable	RW003	RNA_V > 10 000 and (RNA_V modulo 1000) = 1 and RNA_UL missing		YES
tbILAB_RNA	WithinTable	RW004	Missing RNA_D		YES
tbILAB_RNA	WithinTable	RW005	Missing RNA_V		YES
tbILAB_RNA	WithinTable	RW006	RNA_V > 10 million		YES
tbILAB_RNA	WithinTable	RW007	RNA_V < RNA_L and RNA_V >= 0		YES
tbILAB_RNA	WithinTable	RW008	RNA_V > RNA_UL		YES
tbILAB_RNA	CrossTable	RC001	Patient has no record in BAS		YES

**RNA\_T field**

containing table

[tblLAB\\_RNA](#)

explanation of variable

IF AVAILABLE, What type of VIRAL ASSAY was used for this measurement?

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

**Coding Table**

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Viral assay used
5	Roche TaqMan
10	Roche 1.0
15	Roche 1.5 ultra-sensitive
19	Any Roche (unspecified)
20	NASBA
21	NASBA ultra-sensitive
29	Any NASBA (unspecified)
31	Chiron b-DNA 1.0
32	Chiron b-DNA 2.0
33	Chiron b-DNA 3.0
39	Any Chiron (unspecified)
40	Abbott ultra-sensitive
41	Abbott LCx
42	Abbott RealTime HIV-1 m2000
50	Monitor 1.0
51	Monitor 1.0 ultra-sensitive
55	Monitor 1.5
56	Monitor 1.5 ultra-sensitive
59	Monitor unspecified
65	Cobas 1.5
66	Cobas 1.5 ultra-sensitive
90	Other
99	Unknown

## tbILAB\_VIRO - Laboratory values - viro-/serology

holds test results for viro-/serological tests (hepatitis etc.)

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b><u>PATIENT</u></b>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<b><u>VS_ID</u></b>	character. see <a href="#">coding table</a>	Viral test
<b><u>VS_D</u></b>	yyyy-mm-dd	Measurement date
<b><u>VS_R</u></b>	numeric: <ul style="list-style-type: none"> <li>• 1 = Positive</li> <li>• 0 = Negative</li> <li>• 9 = Unknown/borderline</li> </ul>	Measurement result
<b><u>VS_V</u></b>	numeric	Measurement value (HCV-RNA & HBV-DNA only) (copies/ml)
<b><u>VS_U</u></b>	character. see <a href="#">coding table</a> for valid codings.	Measurement unit

### Additional fields

Field name	Format	Description
<b><u>VS_LL</u></b>	numeric	IF AVAILABLE, Lower limit of assay
<b><u>VS_UL</u></b>	numeric	IF AVAILABLE, Upper limit of assay
<b><u>VS_T</u></b>	character. see <a href="#">coding table</a> for valid codings.	IF AVAILABLE, type of ASSAY used for this measurement

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbILAB_VIRO	CrossTable	LVC001	Patient doesn't have a record in BAS		YES
tbILAB_VIRO	WithinTable	LVW002	Missing patient		YES
tbILAB_VIRO	WithinTable	LVW003	Missing VS_ID		YES
tbILAB_VIRO	WithinTable	LVW004	Missing VS_D		YES
tbILAB_VIRO	WithinTable	LVW005	Missing VS_R		YES
tbILAB_VIRO	WithinTable	LVW006	Missing VS_V		YES
tbILAB_VIRO	WithinTable	LVW007	Missing VS_U		YES
tbILAB_VIRO	WithinTable	LVW008	More or less than exactly 1 positive HIV test per patient		YES
tbILAB_VIRO	WithinTable	LVW009	More than 1 negative HIV test for a patient		YES
tbILAB_VIRO	WithinTable	LVW010	Date of negative test after date of positive test		YES

tblLAB_VIRO	WithinTable	LVW011	Duplicate records		YES
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## VS\_ID field

containing table

[tblLAB\\_VIRO](#)

explanation of variable

Viral test

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Viral Test
BVA	Bacterial vaginosis unspecified method
BVAC	Bacterial vaginosis - clinical
BVAG	Bacterial vaginosis - gram stain
CHLA	Chlamydia
CMVA	CMV anitbodies
GONO	Gonorrhoe
HBV	Marker for hepatitis B infection (=HBVAC) - test unknown
HBVAC	HBV antibody (core)
HBVACIGM	HBV antibody (core IgM)
HBVACIGG	HBG antibody (core IgG)
HBVAE	HBV antibody (envelope)
HBVAS	HBV antibody (surface)
HBVD	HBV-dna
HBVGE	HBV antigen (envelope)
HBVGS	HBV antigen (surface)
HCV	Marker for hepatitis C infection - test unknown
HCVA	HCV antibody
HCVG	HCV antigen
HCVBD	HCV b-dna
HCVR	HCV-rna
HDVA	Hepatitis delta antibody
HIV-1	HIV-1 test
HIV-2	HIV-2 test
HIVAE	HIV antibodies ELISA
HIVAWB	HIV antibodies Western blot
MYCO	Mycoplasma
P24AG	P24 Ag
RUB	Rubella
STR	Streptococcus, group B
TOXA	Toxo antibodies
UREP	Ureaplasma

**VS\_T field**

containing table

[tblLAB\\_VIRO](#)

explanation of variable

IF AVAILABLE, the type of ASSAY used for this measurement

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

**Coding Table**

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Viral test used
1	Roche qualitative (Amplicor) [HCV and HBV]
2	Roche quantitative test for HBV (Cobas Amplicor HBV monitor)
3	Bayer Bdna quantitative [HCV]
4	Bayer Bdna quantitative [HBV]
5	Roche Taqman
6	Abbott Real Time [HCV and HBV]
7	Siemens VERSANT [HCV and HBV] DNA (bNA)
8	Quiagen artus [HCV and HBV] PCR kit
9	Other



**VS\_U field**

containing table

[tblLAB\\_VIRO](#)

explanation of variable

Measurement unit

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

**Coding Table**

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Test measurement unit
1	copies/mL
2	IU/mL
3	Geq (millions of genome equivalent)

## tbILTFU - Death and drop-out

holds data in death and drop-out

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">DROP_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> </ul>	Has the patient DROPPED OUT?
<a href="#">DROP_D</a>	yyyy-mm-dd	IF YES, Date of Last Visit
<a href="#">DROP_RS</a>	numeric. see <a href="#">coding table</a> for valid codings.	IF YES, Reason for DROP
<a href="#">DEATH_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> </ul>	Has the patient died?
<a href="#">DEATH_D</a>	yyyy-mm-dd	Date of Death
<a href="#">SUD_DEATH_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Sudden Death?
<a href="#">EXP_DEATH_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Expected Death?
<a href="#">AUTOP_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Was an autopsy Performed?
<a href="#">DEATH_R1</a>	character. see <a href="#">coding table</a> for valid codings.	Cause of death
<a href="#">DEATH_RC1</a>	character with codes: <ul style="list-style-type: none"> <li>I = Immediate cause</li> <li>U = Underlying cause/condition</li> <li>C = Contributing cause</li> <li>N = Not available</li> </ul>	Coding of causal relation of the code given in DEATH_R1 to the death
<a href="#">DEATH_R2</a>	character. see <a href="#">coding table</a> for valid codings.	Cause of death
<a href="#">DEATH_RC2</a>	character with codes: <ul style="list-style-type: none"> <li>I = Immediate cause</li> <li>U = Underlying cause/condition</li> <li>C = Contributing cause</li> <li>N = Not available</li> </ul>	Coding of causal relation of the code given in DEATH_R2 to the death
<a href="#">DEATH_R3</a>	character. see <a href="#">coding table</a> for valid codings.	Cause of death
<a href="#">DEATH_RC3</a>	character with codes: <ul style="list-style-type: none"> <li>I = Immediate cause</li> <li>U = Underlying cause/condition</li> <li>C = Contributing cause</li> <li>N = Not available</li> </ul>	Coding of causal relation of the code given in DEATH_R3 to the death
<a href="#">DEATH_SOURCE</a>	character	Source of information for coding of death (e.g. CoDe within own cohort, CoDe from D:A:D, CoDe from ART-CC, etc.)

List of *DEATH\_R#* and *DEATH\_RC#* should be continued for as many reasons that are recorded.

The *DEATH\_RC#* fields should enable cohorts to transfer data in accordance with the [?Coding of Death project \(CoDe\)](#). You are welcome to contact the CoDe group for electronic sample forms for detailed collection of data used for the CoDe review process.

CoDe defines 1 immediate, 2 contributing and 1 underlying cause of death.

### Additional fields

Field name	Format	Description
<a href="#">ICD10_1</a>	character	Cause of death as ICD-10 if available
<a href="#">ICD10_2</a>	character	Cause of death as ICD-10 if available
<a href="#">ICD10_3<sup>1</sup></a>	character	Cause of death as ICD-10 if available
<a href="#">ICD9_1?</a>	character	Cause of death as ICD-9 if available
<a href="#">ICD9_2</a>	character	Cause of death as ICD-9 if available
<a href="#">ICD9_3<sup>1</sup></a>	character	Cause of death as ICD-9 if available
<a href="#">DEATH_OT</a>	character	Reason for death ? other - description
<a href="#">L_ALIVE</a>	yyyy-mm-dd	Last date known to be alive

<sup>1</sup>: List of [ICD10\\_#](#) and [ICD9\\_#](#) in place of or together with [DEATH\\_R#](#) and together [DEATH\\_RC#](#) and should be continued for as many reasons that are recorded.

CoDe defines 1 immediate, 2 contributing and 1 underlying cause of death.

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblLTFU	WithinTable	LFW001	DROP_Y and DEATH_Y both non-null		YES
tblLTFU	WithinTable	LFW002	DEATH_Y and DROP_RS both non-null		YES
tblLTFU	WithinTable	LFW003	Any of DEATH_Rx or DEATH_RCx non-null but DEATH_Y=0		YES
tblLTFU	WithinTable	LFW004	R2/RC2 non-null but R1/RC1 null		YES
tblLTFU	WithinTable	LFW005	R3/RC3 non-null but R2/RC2 null		YES
tblLTFU	WithinTable	LFW006	Duplicate patients		YES
tblLTFU	WithinTable	LFW007	Missing DROP_Y		YES
tblLTFU	WithinTable	LFW008	Missing DROP_D if DROP_Y=1		YES
tblLTFU	WithinTable	LFW009	Missing DROP_RS if DROP_Y=1		YES
tblLTFU	WithinTable	LFW010	Missing DEATH_Y		YES
tblLTFU	WithinTable	LFW011	Missing DEATH_D if DEATH_Y=1		YES
tblLTFU	WithinTable	LFW012	DEATH_D non null but DEATH_Y=0		YES
tblLTFU	CrossTable	LFC001	Patient not found in tblBAS		YES

tbLTFU	CrossTable	LFC002	patient in tbIBAS hasn't got a record in tbLTFU		YES
tbLTFU	CrossTable	LFC003	tbIBAS has AIDS=0 but DEATHRx =8.1		YES

## DEATH\_R1 field

containing table

[tblLTFU](#)

explanation of variable

Cause of death

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

The following table represents the Coding of Death ([?CoDe](#)) standard. The newest version is available in the Review Form [?2.2](#).

Use the most specific coding available whenever possible.

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Cause of Death
01	AIDS (ongoing active disease)
01.1	Infection
01.2	Malignancy
02	Infection (other than 01.1)
02.1	Bacterial
02.1.1	Bacterial with sepsis
02.2	Others
02.2.1	Others with sepsis
02.3	Unknown aetiology
02.3.1	Unknown with sepsis
03	Chronic viral hepatitis (progression of/complication to)
03.1	HCV
03.1.1	HCV with cirrhosis
03.1.2	HCV with liver failure
03.1.3	HCV with liver cancer
03.2	HBV
03.2.1	HBV with cirrhosis
03.2.2	HBV with liver failure
03.2.3	HBV with liver cancer
04	Malignancy (other than 01.2 and 03, 03.1, 03.2)
04.03	ANUS - Anal cancer
04.04	BLAD - Bladder cancer
04.05	BONE - Bone cancer
04.06	BRAC - Brain cancer
04.07	BRCA - Breast cancer
04.10.1	ALL - Leukaemia: Acute lymphoid
04.10.2	AML - Leukaemia: Acute myeloid
04.10.3	CLL - Leukaemia: Chronic lymphoid
04.10.4	CML - Leukaemia: Chronic myeloid
04.10.9	LEUK - Leukaemia: unspecified
04.18	COLO - Colon cancer
04.11	COTC - Connective tissue cancer
04.12	ESOP - Esophagus cancer
04.13	GALL - Gallbladder cancer
04.14	GYCA - Gynaecologic cancer
04.15	HDL - Hodgkin lymphoma
04.16	HENE - Head and neck (incl. face) cancers
04.17	KIDN - Kidney cancer

04.19	LIPC - Lip cancer
04.20	LIVR - Liver cancer
04.21	LUNG - Lung cancer
04.22	MALM - Malignant melanoma
04.27	MULM - Multiple myeloma
04.29	PANC - Pancreas cancer
04.31	PENC - Penile cancer
04.32	PROS - Prostate cancer
04.33	RECT - Rectum cancer
04.34	STOM - Stomach cancer
04.35	TESE - Testicular seminoma
04.36	UTER - Uterus cancer
04.40.1	MEAC - Metastasis: of adenocarcinoma
04.40.2	MEOC - Metastasis: of other cancer type
04.40.3	MESC - Metastasis: of squamous cell carcinoma
04.40.9	META - Metastasis: unspecified
04.90	OTH - Other Malignancy Type
04.99	UNKP - Unknown Malignancy Type
05	Diabetes Mellitus (complication to)
06	Pancreatitis
07	Lactic acidosis
08	MI or other ischemic heart disease
08.1	AMI
08.1.1	Definitive AMI (Dundee 1)
08.1.2	Possible AMI (Dundee 2/9)
08.2	Other ischemic heart disease
09	Stroke
10	Gastro-intestinal haemorrhage (if chosen, specify underlying cause)
11	Primary pulmonary hypertension
12	Lung embolus
13	Chronic obstructive lung disease
14	Liver failure (other than 03, 03.1, 03.2)
15	Renal failure
16	Accident or other violent death (not suicide)
17	Suicide
18	Euthenasia
19	Substance abuse (active)
19.1	Chronic Alcohol abuse
19.2	Chronic intravenous drug-use
19.3	Acute intoxication
20	Haematological disease (other causes)
21	Endocrine disease (other causes)
22	Psychiatric disease (other causes)
22.1	Mental and behavioural disorders due to use of psychoactive substances (other than alcohol and intravenous opioids)
22.2	Schizophrenia, schizotypal and delusional Disorders
22.3	Mood /Affective disorders (Major depressive disorder, Bipolar disorder and other mood disorders)
22.4	Neurotic, stress-related and somatoform disorders (including anxiety disorders, phobias, OCD, stress reaction, dissociative disorders, somatoform disorders)
22.5	Behavioral syndromes associated with physiological disturbances and physical factors (including eating disorders, sleep disorders, sexual disorders)

22.90	Other psychiatric disorders
23	CNS disease (other causes)
23.1	Movement disorders (Parkinson's disease; dystonias and Parkinson-like syndromes)
23.2	Degenerative disorders of the central nervous system (Alzheimer's disease; Creutzfeldt-Jakob disease and other degenerative diseases of nervous system)
23.3	Demyelinating diseases of the central nervous system (Multiple sclerosis, other demyelinating diseases)
23.4	Epilepsy (including localised and generalized epilepsy and epileptic syndromes)
23.5	Polyneuropathies (Guillain-Barré syndrome and other polyneuropathies/disorders of the peripheral nervous system)
23.6	Diseases of myoneural junction and muscle (Myasthenia gravis and other myoneural disorders)
23.90	Other disorders of the nervous system
24	Heart or vascular (other causes)
25	Respiratory disease (other causes)
26	Digestive system disease (other causes)
27	Skin and motor system disease (other causes)
28	Urogenital disease (other causes)
29	Obstetric complications
30	Congenital disorders
31	Symptoms caused by mitochondrial toxicity (other than 06, 07)
32	Bleeding (haemophilia)
33	Sudden infant death
33.1	Child abuse
90	Other causes
91	Unclassifiable causes
92	Unknown
92.1	Unknown, Competing risks

The following additional codes are used in the [?TB-HIV](#) study, but not yet officially included in [?CoDe](#).

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Cause of Death
01.1.1	Tuberculosis
01.1.1.1	Pulmonary tuberculosis
01.1.1.2	Extrapulmonary TB
01.1.1.2.01	Pleura (isolated, without lungs involvement)
01.1.1.2.02	Lymphatic intrathoracic (isolated, without lungs involvement)
01.1.1.2.03	Lymphatic extrathoracic
01.1.1.2.04	Pericardia
01.1.1.2.05	Spine
01.1.1.2.06	Bone/joints other than spine
01.1.1.2.07	Meningitis
01.1.1.2.08	CNS other than meningitis
01.1.1.2.09	Genito-urinary tract
01.1.1.2.10	Peritoneal/digestive
01.1.1.2.11	TB with poly-organ failure/ TB sepsis
01.1.1.3	Disseminated tuberculosis
01.1.1.3.01	Lungs
01.1.1.3.02	Miliary
01.1.1.3.03	Pleura
01.1.1.3.04	Lymphatic intrathoracic
01.1.1.3.05	Lymphatic extrathoracic

01.1.1.3.06	Pericardia
01.1.1.3.07	Spine
01.1.1.3.08	Bone/joints other than spine
01.1.1.3.09	Meningitis
01.1.1.3.10	CNS other than meningitis
01.1.1.3.11	Genito-urinary tract
01.1.1.3.12	Peritoneal/digestive
01.1.1.3.13	TB with poly-organ failure/ TB sepsis
01.1.2	other AIDS-defining infection
01.1.2.1	with lungs involvement
01.1.2.2	with CNS involvement
01.2.1	AIDS Malignancy localised to lungs
01.3	IRIS after cART initiation (can only be used as contributing cause of death)
02.1.2	Bacterial pneumonia 02.1.3 Bacterial meningitis
02.2.2	other pneumonia 02.2.3 other with CNS involvement
19.3.1	Toxicity to anti-TB drugs



## DROP\_RS field

containing table

[tblLTFU](#)

explanation of variable

IF [DROP\\_Y](#) = 1, Reason for DROP

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Reason for Drop Out
0	Patient was not infected (mainly for children)
1	Patient lost to follow-up / not known to be dead
2	Patient has not had visit within required amount of time
2.1	Patient did not respond to several invitations
3	Patient moved away
3.1	Patient moved to another country
4	Patient moved and is followed by another centre
4.1	Paediatric patient transferred to adult care
5	Patients decision
5.1	Patients caretaker wanted to discontinue (for children)
6	Consent withdrawn
7	Incarceration/jail
8	Institutionalisation (drug treatment, psychological ?etc.)
9	Other

*Note:* If consent is withdrawn, all patient data except for the patient id and reason for drop out *may* have to be deleted.

## tbIMED - Other medication

holds type, start and stop dates for other medication/treatments.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b><u>PATIENT</u></b>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<b><u>MED_ID</u></b>	character. see <a href="#">coding table</a> for valid codings.	Code representing the treatment
<b><u>MED_SD</u></b>	yyyy-mm-dd	Date of Initiation of Treatment
<b><u>MED_ED</u></b>	yyyy-mm-dd	Date of stopping treatment
<b><u>MED_RS</u></b>	character. see <a href="#">coding table</a> for valid codings.	reason for stopping treatment

### Additional fields

Please see [tbIART - Antiretroviral treatment](#).

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbIMED	WithinTable	MW001	Duplicate records		YES
tbIMED	WithinTable	MW002	MED_ONG=0 and MED_ED null	EPPICC	NO
tbIMED	WithinTable	MW003	MED_ONG=1 and MED_ED non-null	EPPICC	NO
tbIMED	WithinTable	MW004	Missing MED_ID		YES
tbIMED	WithinTable	MW005	Missing MED_SD		YES
tbIMED	WithinTable	MW006	Mising MED_ED		YES
tbIMED	WithinTable	MW007	Overlapping periods of same drug		YES
tbIMED	WithinTable	MW008	MED_ED < MED_SD		YES
tbIMED	WithinTable	MW009	MED_RS not null, but end date NULL		YES
tbIMED	CrossTable	MC001	Patient has no records in tbIBAS		YES

## MED\_ID field

containing table

[tblMED](#)

explanation of variable

Code representing the treatment

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.30](#)

## Coding Table

See also the [notes on the extended ATC-Codes](#) and the [?ATC Index](#).

Similar to the drugs listed below you can report any other non-ART medication with it's ATC-Code.

[Download this table as CSV file](#), [Download this table as XML file](#)

Codes (Extended ATC-Codes)	Other medication
A10A	Insulin or derivatives hereof
A10B	Oral antidiabetic agents
A11CC	vitamin D
A14A	Anabolic steroids/appetite stimulants
B01AC	Anti-platelets
C-HYP	Other anti-hypertensive agents [C02, C03, C04, C07, C08]
C09	ACE inhibitors
C10	Lipid lowering agents
G02CA	Tocolysis
H02	Corticosteroids
J01	Antibiotics
J01AA08	Minocycline (MINOCIN)
J01EA01	Trimethoprim (MONOTRIM, NOFIL)
J01EC02	Sulfadiazine
J01EE	Cotrimoxazole - Comb. of sulfonamides and trimethoprim (BACTRIM, EUSAPRIM, NOFIL)
J01EE01	Sulfamethoxazole and trimethoprim (Bactrim)
J01EE03	Sulfametrole and trimethoprim - Cosoltrime (MADERAN)
J01FA09	Clarithromycine (KLACID)
J01FA10	Azithomycine (ZITHROMAX)
J01FF01	Clindamycine (DALACIN)
J01GA01	streptomycin
J01GB06	Amikacine (AMIKINE)
J01MA02	Ciprofloxacin (CIPROXINE, CILOXAN)
J01MA12	Levofloxacin (TAVANIC)
J01MA14	Moxifloxacin
J01RA02	Cosoltrime (MADERAN)
J02AA01	Amphotericin B (FUNGIZON)
J02AB	Imidazoles (DAKTARIN, NIZORAL, PEVARYL ?)
J02AB02	Ketoconazole
J02AC01	Fluconazole (DIFLUCAN)
J02AC02	Itraconazole (SPORANOX)
J02AC03	Voriconazole
J02AC04	Posaconazole
J02AC05	Isavuconazole
J02AX01	Flucytosine
J02AX04	caspofungin
J04AB02	Rifampin (RIMATICIN)

J04AB04	Rifabutin (MYCOBUTIN)
J04AB05	Rifapentine (Priftin)
J04AC01	Isoniazide (RIMIFON)
J04AK01	Pyrazinamide (PYRAZINAMID)
J04AK02	Ethambutol (EMB, MYAMBUTOL)
J04AM05	RIFATER
J04BA01	Clofazimine (LAMPREN)
J04BA02	Dapsone
J05AB01	Aciclovir (ZIVORAX)
J05AB04	Ribavirin
J05AB06	Ganciclovir (CYMEVENE)
J05AB09	Famciclovir
J05AB11	Valaciclovir (VALTEX)
J05AB12	Cidofovir (VISTIDE)
J05AB15	Valganciclovir
J05AD01	Foscarnet (FOSCAVIR)
J05AE11	Telaprevir (INCIVEK, INCIVO)
J05AE12	Boceprevir (VICTRELIS)
J05AE13	Faldaprevir
J05AE14	Simeprevir
J05AE15	Asunaprevir
J05AF08	Adefovir (PREVEON)
J05AF10	Entecavir
J05AF11	Telbivudine
J05AF12	Clevudine
J05AR-DAAS	Daclatasvir/Asunaprevir
J05AX GRAZ-ELB	Grazoprevir/Elbasvir
J05AX14	Daclatasvir
J05AX15	Sofosbuvir
J05AX16	Dasabuvir
J05AX65	Ledipasvir/Sofosbuvir
J05AX67	Ombitasvir, paritaprevir (ABT-450) and ritonavir
J07BM0	HPV Vaccine
J07BM01	HPV Vaccine (types 6, 11, 16, 18)
J07BM02	HPV Vaccine (types 16, 18)
J07BM03	HPV Vaccine (types 6, 11, 16, 18, 31, 33, 45, 52, 58)
L01AA01	Cyclophosphamide (ENDOXAN)
L01AD02	CCNU (LOMUSTINE)
L01AX04	Dacabazine (DTIC - Dome)
L01BA01	Methotrexate
L01CA01	Vinblastin (VELBE)
L01CA02	Oncovin (VINCRISTINE)
L01CB01	Etoposide (VEPESIDE, EXITOP 100)
L01DB01	Doxorubicine, Adriamycine (DOXIL, CAELYX, ADRIBLASTIN)
L01DC01	Bleomycine
L01XB01	Procarbazine (NATULAN)
L03AA02	G-CSF/Filgrastim (NEUPOGEN)
L03AB	Interferons
L03AB-AL2	Peginterferon alfa-2a/alfa-2b (PEGINTRON, PEGASYS)
L03AB10	Peginterferon alfa-2b (PEGINTRON)
L03AB11	Peginterferon alfa-2a (PEGASYS)

L03AC-IL2	Interleukin 2 (PROLEUKIN)
M05BA	bisphosphonate
N05A	Antipsychotics
N05CD	Benzodiazepine derivatives
N05CF	Benzodiazepine related drugs
N06A	Antidepressant
N07BC	Other drugs used in opioid dependence
N07BC01	Buprenorphine
N07BC02	Methadone
N07BC03	Levacetylmethadol
N07BC04	Lofexidine
N07BC51	Buprenorphine, combinations
P01AX06	Atovaquone (WELLVONE, MEPRONE)
P01BA03	Primaquine
P01BD01	Pyrimethamine (DARAPRIM)
P01BD51	Pyrimethamine/Sulfadoxine (FANSIDAR)
P01CX01	Pentamidine aerosol (PENTACARNET)
V03AB15	Naloxone
V03AF03	Folate of calcium (LEUCOVORINE)

## MED\_RS field

containing table

[tblMED](#)

explanation of variable

Reason for stopping treatment

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.70](#)

## Coding Table

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Reason for Stopping Treatment
1	Treatment failure (i.e. virological, immunological, and/or clinical failure)
1.1	Virological failure
1.2	Partial virological failure
1.3	Immunological failure - CD4 drop
1.4	Clinical progression
2	Abnormal fat redistribution
3	Concern of cardiovascular disease
3.1	Dyslipidaemia
3.2	Cardiovascular disease
4	Hypersensitivity reaction (skin eruption etc.)
5	Toxicity, predominantly from abdomen/G-I tract
5.1	Toxicity - GI tract
5.2	Toxicity - Liver
5.3	Toxicity - Pancreas
6	Toxicity, predominantly from nervous system
6.1	Toxicity - peripheral neuropathy
6.2	Toxicity - neuropsychiatric
6.3	Toxicity - headache
7	Toxicity, predominantly from kidneys
8	Toxicity, predominantly from endocrine system
8.1	Diabetes
9	Haematological toxicity (anemia etc.)
10	Hyperlactataemie/lactic acidosis
11	Bone toxicity
15	Social contra-indication
16	Contra-indication unspecified
16.8	Contra-indication expired
16.9	Contra-indication ? other
17	MTCT regimen completed
70	Pregnancy - toxicity concerns (during pregnancy)
75	Pregnancy - switch to a more appropriate regimen for PMTCT
88	Death
90	Side effects - any of the above but unspecified
90.1	Comorbidity
91	Toxicity, not mentioned above
91.1	Toxicity, unspecified
92	Availability of more effective treatment (not specifically failure or side effect related)
92.1	Simplified treatment available
92.2	Treatment too complex

92.3	Drug interaction
92.31	Drug interaction - commencing TB/BCG treatment
92.32	Drug interaction - ended TB/BCG treatment
92.33	Change in eligibility criteria (e.g. child old enough for tablets; refrigerator no longer available)
92.4	Protocol change
92.5	Regular treatment termination (e.g. DAA's for HCV, antibiotics)
92.9	Change in treatment not due to side-effects, failure, poor adherence or contra-indication
93	Structured Treatment Interruption (STI)
93.1	Structured Treatment Interruption (STI) - at high CD4
94	Patient's wish/decision, not specified above
94.1	Non-compliance
94.2	Defaulter
95	Physician's decision, not specified above
96	Pregnancy
96.1	Pregnancy intended
96.2	Pregnancy ended
97	Study treatment
97.1	Study treatment commenced
97.2	Study treatment completed
97.6	Drug not available
98	Other causes, not specified above
99	Unknown

Although most reasons mentioned above could be coded in ICD-10 codes, it is still certain that many reasons will never be defined in terms of ICD-10 (STI, physicians and patients decision), so to avoid a mixture of two different coding systems it is advised to go with the above list and build upon that for now. This list is identical to the stopping reasons for ART (ART\_RS)

## tbNEWBORN - Newborn

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>CHILD_ID</b>	Character (or numeric if possible)	Patient ID of the child
BRFEED_SD	yyyy-mm-dd	Breastfeeding, start date.
BRFEED_ED	yyyy-mm-dd	Breastfeeding, end date.
FAT_ETH	See <a href="#">Coding Table</a> .	Ethnicity of father
APGAR_1	numeric	1 <sup>st</sup> APGAR score
APGARM_1	numeric	minute at which the 1 <sup>st</sup> APGAR test was performed
APGAR_2	numeric	2 <sup>nd</sup> APGAR score
APGARM_2	numeric	minute at which the 2 <sup>nd</sup> APGAR test was performed
APGAR_3	numeric	3 <sup>rd</sup> APGAR score
APGARM_3	numeric	minute at which the 3 <sup>rd</sup> APGAR test was performed
ICU_Y	numeric: <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	Referral to intensive/intermediate care unit?
ICU_S	character	if yes, specify reason
ICU_D	yyyy-mm-dd	Date of entry to intensive/intermediate care unit
ABNORM_Y	numeric: <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	did any abnormalities occur? (if yes, recorded in <a href="#">tbNEWBORN_ABNORM</a> )

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbNEWBORN	CrossTable	NC001	CHILD_ID doesn't exist in tbIDELIVERY_CHILD		YES
tbNEWBORN	CrossTable	NC002	ABNORM_Y = 1, yet no records in tbNEWBORN_ABNORM		YES
tbNEWBORN	CrossTable	NC003	ABNORM_Y = 0 or 9, yet records in tbNEWBORN_ABNORM		YES
tbNEWBORN	WithinTable	NW001	BRFEED_SD>BRFEED_ED		YES



tblNEWBORN	WithinTable	NW002	APGARM_x's out of order (e.g APGARM_3 < APGARM_2)		YES
tblNEWBORN	WithinTable	NW003	ICU_Y=1, but ICU_S or ICS_D null		YES
tblNEWBORN	WithinTable	NW004	ICU_Y=0 or 9, but ICU_S or ICU_D non null		YES

## tbNEWBORN\_ABNORM - Abnormalities

Abnormalities in newborns are recorded here, one abnormality per row. The absence of a record is to be interpreted as "unknown whether the abnormality existed" since most cohorts only record positive events.

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>CHILD_ID</b>	Character (or numeric if possible)	Patient ID of the child
<b>ABNORM_T</b>	character: <ul style="list-style-type: none"> <li>• 1 = Birth defect(s) (detectable in physical examination including skin abnormalities)</li> <li>• 2 = Congenital infection(s)</li> <li>• 3 = Drug withdrawal syndrome</li> <li>• 4 . 1 = Neurological disorder(s): abnormal reflexes</li> <li>• 4 . 2 = Neurological disorder(s): abnormal motility</li> <li>• 4 . 3 = Neurological disorder(s): abnormal tonus</li> <li>• 90 = Other health problems</li> </ul>	type of abnormality
<b>ABNORM_S</b>	character	further specification of the abnormality

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbNEWBORN_ABNORM	CrossTable	NAC001	CHILD_ID doesn't exist in NEWBORN		YES
tbNEWBORN_ABNORM	WithinTable	NAW001	ABNORM_T=90 but ABNORM_S null		YES

## tbOVERLAP - Cross-cohort identification

holds information on the patient's participation in other cohorts

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<u>PATIENT</u>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<u>COHORT</u>	character	Code/name of the cohort
<u>PAT_OTH</u>	character	Unique patient identifier in other cohorts
<u>COH_OTH</u>	character	Name of the cohort

Patients of an "original"-cohort who also participate in a "super"-cohort should be analysed within the "original"-cohort only. To suppress these patients from the datasets of the "super"-cohorts the identifier used in the "super"-cohort is needed. It is suggested that "original"-cohorts report id's from the "super"-cohorts, since the "super"-cohorts might not even know the other ID's. Often this information is only available at centre level.

A record should be present for each cohort that the patient is participating in (apart from it's own "original"-cohort).

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbOVERLAP	WithinTable	OW001	Invalid other cohort		YES
tbOVERLAP	CrossTable	OC001	PATIENT not found in tblBAS for that cohort		YES
tbOVERLAP	CrossTable	OC002	PAT_OTH not found in tblBAS for that overlapping cohort		YES

## tbIPREG - Pregnancy

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>MOTHER_ID</b>	Character (or numeric if possible)	Patient ID of mother of the child
<b>PREG_SEQ</b>	numeric	Sequence number of the pregnancy for the specified mother
<b>MENS_D</b>	yyyy-mm-dd	Date of last menstrual period (If date not known exactly please give approximated date)
<b>CONCEPT</b>	character: <ul style="list-style-type: none"> <li>• 1=Natural</li> <li>• 2=Infertility treatment unspecified</li> <li>• 2. 1=IVF (In Vitro Fertilisation)</li> <li>• 2. 2=ICSI (IntraCytoplasmic Sperm Injection)</li> <li>• 2. 3=Ovulation induction</li> <li>• 3=Artificial insemination</li> <li>• 4=Self insemination</li> <li>• 9=Unknown</li> </ul>	Conception
<b>ANC_D</b>	yyyy-mm-dd	Date of first antenatal care contact
<b>INPREG_Y</b>	character: <ul style="list-style-type: none"> <li>• 1=Yes</li> <li>• 2=No: ectopic</li> <li>• 3=No: missed abortion</li> <li>• 4=No: death in utero (IUFT)</li> </ul>	At first gynaecological visit: intact intrauterine pregnancy?
<b>INHIST_Y</b>	numeric: <ul style="list-style-type: none"> <li>• 0=No</li> <li>• 1=Yes</li> <li>• 9=Unknown</li> </ul>	If no in INPREG_Y was a histological investigation of reason made.
<b>INHIST_S</b>	character	If yes, please specify reason
<b>INV_PROC</b>	<ul style="list-style-type: none"> <li>• 0=No</li> <li>• 1=Yes, chorionic villus sampling (CVS)</li> <li>• 2=Cordocentesis</li> <li>• 3=Amniocentesis</li> <li>• 9=Unknown</li> </ul>	Invasive procedure
<b>KARYO_T</b>	<ul style="list-style-type: none"> <li>• 0=Not done</li> <li>• 1=Normal</li> <li>• 2=Abnormal</li> <li>• 9=Unknown</li> </ul>	Karyotype
<b>KARYO_A</b>	character	If abnormal, please specify
<b>CHORIO</b>	character: <ul style="list-style-type: none"> <li>• 1=Monochorionic</li> <li>• 2=Dichorionic</li> <li>• 3=Trichorionic</li> <li>• etc.</li> <li>• 9=Unknown</li> </ul>	For multiple pregnancies
<b>ULTR_1</b>	character: <ul style="list-style-type: none"> <li>• 0=No</li> <li>• 1=Yes, normal</li> <li>• 2=Yes, abnormal</li> <li>• 9=Unknown</li> </ul>	Ultrasound 1. trimester
<b>ULTR_A_1</b>	character	If abnormal, please specify

<i>ULTR_2</i>	character: <ul style="list-style-type: none"> <li>• 0=No</li> <li>• 1=Yes, normal</li> <li>• 2=Yes, abnormal</li> <li>• 9=Unknown</li> </ul>	Ultrasound 2. trimester
<i>ULTR_A_2</i>	character	If abnormal, please specify
<i>PROB_Y</i>	numeric: <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	did any obstetrical problems occur? (if yes, recorded in <a href="#">tblPREG_OBS</a> )

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblPREG	WithinTable	PW001	ANC_D		YES
tblPREG	WithinTable	PW002	INPREG_Y=2,3,4 but INHIST_Y null		YES
tblPREG	WithinTable	PW003	INPREG_Y=1 but INHIST_Y non null		YES
tblPREG	WithinTable	PW004	INHIST_Y=1 but INHIST_S null		YES
tblPREG	WithinTable	PW004	INHIST_Y=0 or 9 but INHIST_S non null		YES
tblPREG	WithinTable	PW005	KARYO_T=2 but KARYO_A null		YES
tblPREG	WithinTable	PW006	KARYO_T<>2 but KARYO_A non null		YES
tblPREG	WithinTable	PW007	ULTRA_x =2, but ULTR_A_x null		YES
tblPREG	WithinTable	PW008	ULTRA_x <>2, but ULTR_A_x non null		YES
tblPREG	CrossTable	PC001	PROB_Y = 1 but no records in tblPREG_OBS		YES
tblPREG	CrossTable	PC002	PROB_Y = 0 or 9 but records in tblPREG_OBS		YES

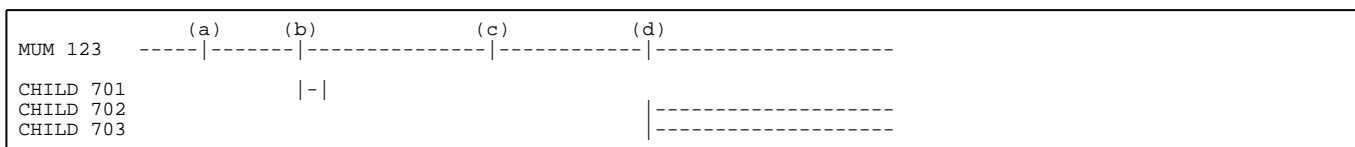
## Notes on the Pregnancy-Related Tables

The correct and accurate modelling of a pregnancy and its outcomes is rather involved, and some assumptions had to be made. This page illustrates the model used in the pregnancy tables to which these assumptions lead.

### Example

Consider the following example:

1. A female patient (123) gets pregnant at time (a).
2. At time (b) she has an abortion which causes the dead fetus of CHILD 701.
3. At time (c), she gets pregnant again.
4. The delivery of this pregnancy is at time (d), when the twins (CHILD 702 and CHILD 703) are born.



This story is described in the different pregnancy tables as follows:

There are two records in [tblPREG](#), one for each pregnancy:

MOTHER_ID	PREG_SEQ	MENS_D	...
123	1	(a)	...
123	2	(c)	...

There are three records in [tblPREG\\_OUT](#) describing the pregnancy outcome for each fetus:

MOTHER_ID	PREG_SEQ	CHILD_ID	OUTCOM	...
123	1	701	21	...
123	2	702	1	...
123	2	703	1	...

There is one record in [tblDELIVERY\\_MUM](#):

MOTHER_ID	PREG_SEQ	MEMRUP_D	...
123	2	(d)	...

There are two records in [tblDELIVERY\\_CHILD](#):

MOTHER_ID	MEMRUP_D	CHILD_ID	...
123	(d)	702	...
123	(d)	703	...

There are two records in [tblNEWBORN](#):

CHILD_ID	...
702	...
703	...

### Important relations

The following statements are true for a well-encoded data set:

1. Every record in [tblPREG\\_OUT](#) refers to some existing record in [tblPREG](#).
2. For every record in [tblPREG\\_OUT](#) that describes a delivery, there is exactly one record in [tblDELIVERY\\_CHILD](#) referring to it.
3. For every record in [tblDELIVERY\\_MUM](#), there are 1 or more records in [tblDELIVERY\\_CHILD](#) referring to it.
4. For every record in [tblNEWBORN](#) there is exactly one record in [tblDELIVERY\\_CHILD](#) referring to the same child.
5. For every record in [tblDELIVERY\\_MUM](#), there is some record in [tblPREG](#) to which it refers.

## tbIPREG\_OBS - Obstetrical problems during pregnancy

This table describes problems **during a pregnancy**. Abnormalities in newborns are recorded in [tblNEWBORN\\_ABNORM](#) instead.

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>MOTHER_ID</b>	Character (or numeric if possible)	patient id of mother
<b>PREG_SEQ</b>	numeric	Sequence number of the pregnancy for the specified mother
<b>PROB_T</b>	character: <ul style="list-style-type: none"> <li>• 1 = Preterm contractions</li> <li>• 2 = Shortened cervix</li> <li>• 3 = Preterm rupture of membranes</li> <li>• 4 = Antepartum bleeding</li> <li>• 5 = Intrauterine growth retardation (IUGR)</li> <li>• 6 = Preeclampsia/HELLP</li> <li>• 7 = Hypertension</li> <li>• 8 = Gestational diabetes (unspecified type)</li> <li>• 8.1 = Gestational diabetes (Diet)</li> <li>• 8.2 = Gestational diabetes (Insulin)</li> <li>• 9 = Placental abruption</li> <li>• 10 = Placenta praevia</li> <li>• 99 = Other, specify in <b>PROB_S</b></li> </ul>	type of obstetrical problem
<b>PROB_S</b>	character.	description of other (99) obstetrical problem
<b>CERVIX_S</b>	numeric (mm)	In case of shortened cervix, the length of the cervix in millimeters.

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbIPREG_OBS	CrossTable	POC001	MOTHER_ID+PREG_SEQ doesn't exist in tbIPREG		YES
tbIPREG_OBS	WithinTable	POW001	PROB_T=99 but PROB_S null		YES
tbIPREG_OBS	WithinTable	POW002	PROB_T<>99 but PROB_S non null		YES
tbIPREG_OBS	WithinTable	POW003	PROB_T=2 but CERVIX_S null		YES
tbIPREG_OBS	WithinTable	POW003	PROB_T<>2 but CERVIX_S non null		YES

## tbIPREG\_OUT - Pregnancy outcome

This table describes outcomes of pregnancies described in [tbIPREG](#).

Please also read the [notes on pregnancy tables](#).

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>MOTHER_ID</b>	Character (or numeric if possible)	Patient ID of mother of the child
<b>PREG_SEQ</b>	numeric	Sequence number of the pregnancy for the specified mother
<b>CHILD_ID</b>	Character (or numeric if possible)	Patient ID of the child
<b>OUTCOM</b>	character: <ul style="list-style-type: none"> <li>1=Born alive, HIV negative</li> <li>2=Born alive, HIV positive</li> <li>3=Born alive, unknown HIV status</li> <li>10=Stillborn</li> <li>11=Spontaneous miscarriage</li> <li>20=Termination: surgical</li> <li>21=Termination: medication</li> </ul>	Pregnancy outcome
<b>OUTCOM_R</b>	<ul style="list-style-type: none"> <li>1=Fetus with malformation</li> <li>2=Dead fetus</li> <li>3=Unwanted pregnancy</li> <li>8=Other</li> <li>9=Unknown</li> </ul>	Reason for termination
<b>OUTCOM_D</b>	yyyy-mm-dd	Date of birth or termination of pregnancy
<b>B_GAGEW</b>	numeric	Gestational age in complete weeks at birth or termination
<b>B_GAGED</b>	numeric	Gestational age in days in addition to weeks at birth or termination

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbIPREG_OUT	CrossTable	PTC001	MOTHER_ID+PREG_SEQ doesn't exist in tbIPREG		YES
tbIPREG_OUT	CrossTable	PTC002	OUTCOM=1,2 or 3 and CHILD_ID doesn't exist in tbINEWBORN or tbIDELIVERYCHILD		YES
tbIPREG_OUT	CrossTable	PTC003	OUTCOM=10,11,20 or 21 and CHILD_ID exists in tbINEWBORN or tbIDELIVERYCHILD		YES
tbIPREG_OUT	WithinTable	PTW001	OUTCOME=20 or 21 and OUTCOM_R null		YES



tblPREG_OUT	WithinTable	PTW002	OUTCOME not 20 or 21 and OUTCOM_R non null		YES
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## tbIPROGRAM - Program information

Holds information about the program with which the center is associated

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b><u>PROGRAM</u></b>	character	Program name
<b><u>REGION</u></b>	character: <ul style="list-style-type: none"> <li>• AP = Asia-Pacific</li> <li>• CA = Central Africa</li> <li>• CN = Caribbean, Central and South America</li> <li>• EA = East Africa</li> <li>• EU = Europe</li> <li>• NA = North America</li> <li>• SA = Southern Africa</li> <li>• WA = West Africa</li> </ul>	Region of operation

## tbIREFILL - Prescription refill data

holds drug, refill date and supply of prescriptions

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<u><a href="#">PATIENT</a></u>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<u><a href="#">REFILL_D</a></u>	yyyy-mm-dd	Date of the prescription refill
<u><a href="#">DRUG_ID</a></u>	character. see <a href="#">coding table</a> for valid codings.	Drug which was prescribed
<u><a href="#">SUPPLY</a></u>	numeric	How many days supply of the drug was supplied

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tbIREFILL	CrossTable	PRC001	Patient doesn't have a record in BAS		YES
tbIREFILL	WithinTable	PRW002	Missing PATIENT		YES
tbIREFILL	WithinTable	PRW003	Missing REFILL_D		YES
tbIREFILL	WithinTable	PRW004	Missing DRUG_ID		YES
tbIREFILL	WithinTable	PRW005	Missing SUPPLY		YES
tbIREFILL	WithinTable	PRW006	SUPPLY < 1		YES

## DRUG\_ID field

containing table

[tbIREFILL](#)

explanation of variable

Drug which was prescribed

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.70](#)

## Coding Table

ATC-Code of drug. See the [MED\\_ID coding table](#) and [ART\\_ID coding table](#), and the ART\_ID notes on extended ATC-Codes.

## tbISAMPLES - Blood Samples

This table contains information on the storage of blood, urine and other samples stored in a laboratory.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<b>PATIENT</b>	character (or numeric if possible)	patient cohort identifier
<b>SAMP_LAB_D</b>	YYYY-mm-dd	date when the sample was taken
<b>SAMP_TYPE</b>	character: <ul style="list-style-type: none"> <li>BS = blood serum</li> <li>BP = blood plasma</li> <li>C = viable cells</li> <li>D = cell pellet (DNA)</li> <li>S = semen</li> <li>OTH:x = other sample type x (none of the above)</li> </ul>	type of the sample
<b>SAMP_ID</b>	character	identification symbol allowing the localization of the sample in the laboratory
<b>SAMP_LAB</b>	character	laboratory where the samples are stored
<b>SAMP_FREEZE_D</b>	YYYY-mm-dd	date when the sample was frozen
<b>SAMP_FREEZE_T</b>	hh:mm	time when the sample was frozen
<b>SAMP_ALIQ_NO</b>	numeric	number of aliquots available
<b>SAMP_ALIQ_SIZE</b>	numeric	size of the aliquot: <ul style="list-style-type: none"> <li>in ml for serum, plasma and cell pellet aliquots</li> <li>in millions of cells for viable cell aliquots</li> </ul>
<b>SAMP_ALIQ_U</b>	character: <ul style="list-style-type: none"> <li>0 = millions of cells</li> <li>1 = ml</li> </ul>	unit of measurement for the SAMP_ALIQ_SIZE value

### Additional fields

Field name	Format	Description
<b>SAMP_LAB_T</b>	hh:mm	time when the sample was taken
<b>SAMP_TEMP</b>	numeric	temperature of the storage unit containing the samples (in °C)
<b>SAMP_DEFROST</b>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	have the samples already been defrosted?

### QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tbILTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tbILTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tbIBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES

AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
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## tbIVIS - Basic follow-up/visit related data

holds visit related information such as weight, wasting, smoking, occupational status etc.

### Core fields

Note: Fields marked **bold** form the unique identifier for a record of the table.

Field name	Format	Description
<a href="#">PATIENT</a>	character (or numeric if possible)	Code to identify patient (Cohort Patient ID)
<a href="#">VIS_D</a>	yyyy-mm-dd	Date of patient visit
<a href="#">CENTER</a>	character	Center the patient visits. Links to <a href="#">tbICENTER</a> .
<a href="#">WEIGH</a>	numeric (metric: kg): 999 = Unknown	Weight of patient at visit
<a href="#">GAIN_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Is the patient gaining fat in the abdomen, neck, breast or other defined locations?
<a href="#">LOSS_Y</a>	numeric: <ul style="list-style-type: none"> <li>1 = Yes</li> <li>0 = No</li> <li>9 = Unknown</li> </ul>	Is the patient experiencing loss of fat from extremities, buttocks or face?

Depending on the collaboration this data might be collected in intervals of a year, e.g. from July last to July this year. In that case all visit dates or a fixed number of visit dates for that period should be gathered, if the patient did not have a visit in the defined period, a record with the PATIENT id and empty fields for VIS\_D etc. should be included.

### Additional fields

Field name	Format	Description
<a href="#">CDC_STAGE</a>	character. see <a href="#">coding table</a> for valid codings.	Clinical CDC stage at time of visit?
<a href="#">WHO_STAGE</a>	numeric. <ul style="list-style-type: none"> <li>1</li> <li>2</li> <li>3</li> <li>4</li> <li>9 = Unknown</li> </ul>	Clinical WHO stage at time of visit?
<a href="#">FAM_Y</a>	numeric. <ul style="list-style-type: none"> <li>0 = No</li> <li>1 = Yes</li> <li>9 = Unknown</li> </ul>	Family history of CVD: Have any first degree relatives experienced myocardial infarction or stroke before the age of 50 years?

The following optional fields are meant to be used to **document the transition process from adolescent to adult**.

Field name	Format	Description
<a href="#">CLINIC_TYPE</a>	numeric <ul style="list-style-type: none"> <li>1 = paediatric</li> <li>2 = adolescent within paediatric care</li> <li>3 = adolescent within adult care</li> <li>4 = adolescent stand alone</li> <li>5 = adult</li> <li>9 = missing</li> </ul>	Type of clinic/service the patient is currently attending
<a href="#">SPEC_TYPE</a>	numeric <ul style="list-style-type: none"> <li>1 = Physician providing paediatric care</li> <li>2 = Physician providing adolescent care</li> <li>3 = Physician providing adult care</li> <li>4 = Physician providing paediatric and adult care</li> <li>5 = other healthcare provider (e.g. nurse)</li> <li>9 = missing</li> </ul>	Type of specialist providing care. Combinations if multiple specialists are involved (e.g. 23, 45).

<a href="#">TRANS_STAGE</a>	numeric <ul style="list-style-type: none"> <li>• 0 = transition not started</li> <li>• 1 = transition in progress</li> <li>• 2 = transition completed</li> <li>• 9 = not applicable/missing</li> </ul>	Stage of transition from pediatric to adult care at current visit. Transition has not yet started when the patient only sees paediatricians. Transition is complete when the patient only sees adult physicians.
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The following fields are meant to be used **for adolescents and adults**.

Field name	Format	Description
<a href="#">EMPLOY</a>	numeric. see <a href="#">coding table</a> for valid codings.	What is the patient's current situation regarding labour?
<a href="#">CONTRACT</a>	numeric. see <a href="#">coding table</a> for valid codings.	If the patient is an employee, what is the type of the patient's employment contract?
<a href="#">SMOKING_Y</a>	numeric: <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	Is the patient currently a smoker?
<a href="#">PREG_Y</a>	numeric: <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	Is the patient currently pregnant? If possible, provide additional details in <a href="#">tblPREG</a>

The following fields are meant to be used **for children and infants**.

The following fields are meant to be used **for infants**:

Field name	Format	Description
<a href="#">HEIGH_P</a>	numeric	Height/length of patient at visit in percentiles
<a href="#">WEIGH_P</a>	numeric	Weight of patient at visit in percentiles
<a href="#">HEADC</a>	numeric	Head circumference measured in millimeters (mm)
<a href="#">HEADC_P</a>	numeric	Head circumference in percentiles
<a href="#">BREASTF_Y</a>	numeric: <ul style="list-style-type: none"> <li>• 1 = Yes</li> <li>• 0 = No</li> <li>• 9 = Unknown</li> </ul>	Currently Breastfeeding?

## QA Checks

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
tblVIS	WithinTable	VW001	Duplicate records, same VIS_D		YES
tblVIS	WithinTable	VW002	Height decreasing over time		YES
tblVIS	WithinTable	VW003	Height out of acceptable range		YES



tblVIS	WithinTable	VW004	Weight out of acceptable range		YES
tblVIS	CrossTable	VC001	patient has no record in BAS table		YES
tblVIS	CrossTable	VC002	No weights within 3 mths of starting FPV/DRV	EPPICC	YES

## CDC\_STAGE field

containing table

[tbVIS](#)

explanation of variable

Clinical CDC stage at time of visit?

format of data

character. see [coding table](#) for valid codings.

exists since HICDEP version

[1.70](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Description
A	A
A1	A1
A2	A2
A3	A3
B	B
B1	B1
B2	B2
B3	B3
C	C
C1	C1
C2	C2
C3	C3
9	Unknown

## CONTRACT field

containing table

[tbVIS](#)

explanation of variable

If the patient is an employee, what is the type of the patient's employment contract?

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.70](#)

## Coding Table

[Download this table as CSV file](#), [Download this table as XML file](#)

Code	Description
1	Indefinite duration
2	Fixed-term
3	Other
9	Unknown

## EMPLOY field

containing table

[tbIVIS](#)

explanation of variable

What is the patient's current situation regarding labour?

format of data

numeric. see [coding table](#) for valid codings.

exists since HICDEP version

[1.70](#)

## Coding Table

[Download this table as CSV file](#) [Download this table as XML file](#)

Code	Description
1	Employed as an employee
2	Self-employed / Family worker
3	Apprentice / trainee
4	Unemployed and actively seeking work
5	Student
6	Retired / Pre-retired
7	Engaged in family duties
8	Unable to work due to ill health
9	Other situation
99	Unknown

## Considerations for data management

### Uniqueness of *PATIENT ID*

Each table in this document is shown with the *PATIENT* field as part of the unique identifier for each record, in many cases it might be necessary to specify both *PATIENT* and *COHORT* as identifier since the *PATIENT* id might not be unique across cohorts, this should however be implemented only if needed.

Another solution, which might be worth considering when building the final dataset for the analysis, is to concatenate the patient id and the cohort id into a single unique new patient id.

### Duplicate records

Transposition of the flat format data, where there is one record per patient, into a normalised structure that has multiple rows per patient requires that the combination of *PATIENT*, *TYPE\_ID* and *LAB\_DATE* is unique for each row in the table.

The normalised structure and the relational model does not allow for duplicate records like in the following table to exist:

<i>PATIENT</i>	<i>TYPE_ID</i>	<i>LAB_DATE</i>	<i>LAB_VAL</i>	<i>LAB_UNIT</i>
999999	1	2000-01-01	15	U/l
999999	1	2000-01-01	15	U/l

The typical solution to this is to generate an auto-incremented value ? *RECORD\_ID* - for each record in the table:

<i>RECORD_ID</i>	<i>PATIENT</i>	<i>TYPE_ID</i>	<i>LAB_DATE</i>	<i>LAB_VAL</i>	<i>LAB_UNIT</i>
1	999999	1	2000-01-01	15	U/l
2	999999	1	2000-01-01	15	U/l

But it can, and should be dealt with in the design of the study. The presence of duplicate records like in the example shown above might not make any sense; if they did occur it then would be on account of a recording error. If, however, it was important to keep records that show measurements taken the same day but at different times, the format of the *LAB\_DATE* shouldn't be date (YYYY-MM-DD) but instead should be date-time (YYYY-MM-DD hh:mm:ss):

<i>PATIENT</i>	<i>TYPE_ID</i>	<i>LAB_DATE</i>	<i>LAB_VAL</i>	<i>LAB_UNIT</i>
999999	1	2000-01-01 10:00:00	15	U/l
999999	1	2000-01-01 14:00:00	15	U/l

Different and more sophisticated methods to make records unique will be presented later in this document under the definition of the LAB table.

## Considerations for using the format to create a database

### Administrative fields

Sometimes it might be needed to have a fixed value that shows from which visit or merger a value originates, this does not only apply to the *VIS* table but could be applied to all tables. This however does depend on the nature of the database and needs for data management, the field below should be considered an administrative support field for data management.

VISIT
Visit number
Numeric: 0 = Baseline Visit 1 = First follow up visit 2 = Second follow up visit etc.

Often the above field is used for clinical trials databases where there is a need to associate the data directly with a given week's follow-up. Codes could then be the week number e.g. 4, 12, 24 etc or ?1 for screening/randomisation and 0 for baseline visits.

In some cases it might be useful to have a separate field that defines the correct order of the periods. This becomes important where several dates are incomplete (unknown days, unknown months and possibly unknown years). The ordering by date would then not be correct.

One solution to this is use a *PERI\_ID* field that numbers the periods from the 1<sup>st</sup> until N<sup>th</sup> usage:

PERI_ID
Period of usage (1 <sup>st</sup> , 2 <sup>nd</sup> , 3 <sup>rd</sup> etc.)
Numeric

However this is an optional field that for most cohorts may not be needed. It also requires additional maintenance to keep it updated.

For databases that work with double data entry, such as most clinical databases, it becomes necessary to make each data entry unique and backwards traceable. For this to work a field like the above would have to be part of the primary key of each table that requires double data entry.

ENTRY_ID
Number of data entry
Numeric: 1 = first data entry 2 = second data entry 3 = comparison of 1 <sup>st</sup> and 2 <sup>nd</sup> data entry 4 = final approved record including corrections

With respect to performance, it might also be a good design to have 3 copies of each table, one to hold the data while being entered and compared, one for the two data entries to be archived into once a final record has been approved and a table holding the final and approved values. This way it is avoided that queries will have to work on checking for *ENTRY\_ID* = 4 and to select amongst a table holding 3 times the almost same data.

As part of an audit trail in a database a time stamp field could be added for each record to fix the exact time when the record last was inserted or updated. Along with the time stamp name of the user who entered or altered data can be recorded.

T_STAMP	USER_LOG
Date and time of data entry	Username of user that last inserted or updated data
yyyy-mm-dd hh:mm:ss	character

Often it's necessary to keep a log of user action in a separate table. The above suggestion will only be valid for inserts and updates, and only be valid for the most recent action performed.

To record a complete audit trail a logging facility must be implemented. In most database management systems this is done using triggers on the tables. For any insert, update or delete actions performed on the data, the user, time stamp, old value and new value are recorded in the logging table.

The *T\_STAMP* field could also include information about which time zone is relevant for data entry. Depending on database requirements this might in fact be mandatory if the [?FDA's 21 CFR part 11](#) on electronic records and signatures applies.

### Further normalisation

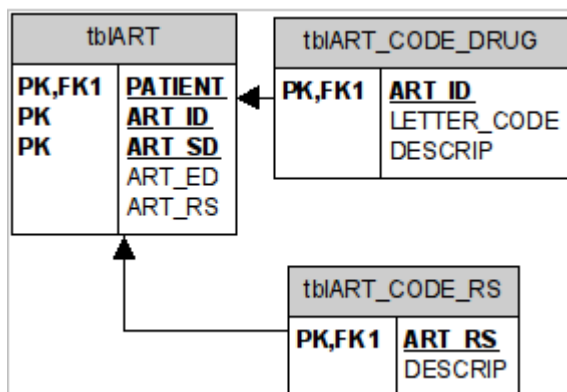
Depending on performance considerations it might be worth looking at how data are queried for data entry and data analysis. A smaller *tbIBAS* table might increase performance: Since processing a smaller table is always faster than processing a larger table, one could put drop-out, death, birthday, date of aids diagnosis, etc. into separate tables and keep the core patient list in a separate master table

But if the database is used e.g. for BMI calculations directly on the running database, performance might be enhanced by keeping the patient list and the height together in the same table so that a query involves 2 tables (*tbIBAS* and *tbIVIS*) rather than perhaps 3 or more.

Another consideration is space. Although it may not be much of an issue, it will be possible to minimise the actual size of the database by putting fields that may be empty for most patients, like death information, into a separate table in a 0-1 to 1 relation to the master table.

### Lookup tables

In a running database the #\_ID fields could be implemented as a foreign key to a linked lookup table containing all possible codes and their corresponding definitions in a text string.



This setup not only enables integrity of the data, but also defines the domain<sup>1</sup> for the #\_ID values and enables data to both become human readable and easily recoded<sup>2</sup>.

An important note on lookup tables is that the performance on a large database can be slowed significantly by using character based keys to link lookup tables with the primary table as it is presented in this document. A work around is to use numeric value for the codes.

<sup>1</sup>: Domain is a term in the definition of the relational database model that defines a set of allowed values for a given set of fields (attributes), the mixing of different domains is not allowed in order to preserve the integrity of a relational and normalised model.

<sup>2</sup>: Easily recoded permanently if the relation is specified as cascade on update or recoded dynamic by selecting a different column from the lookup table when querying the data through SQL

## Performance

As already outlined in the above section, there are also performance issues that may have to be considered.

When using the suggested data types presented in this document for a database implementation, it may be worth looking at the actual data at hand when defining the final data types. The aim of this document is to present a format that will work between cohorts with rather different setups.

If it is at all possible in many cases there may be a large performance gain by using numeric instead of character fields. Character fields have been suggested here for, amongst others, the *PATIENT* field. If the *PATIENT* id is purely numeric it's worth using a numeric data type since it always faster for querying than a character field.

Whenever the field has to be character, make sure that only the needed amount of space is assigned for the field length; there is no need to assign 50 characters of memory if the field in fact only stores a 3-letter code.

## QA checks

This page lists all QA checks currently defined for [HICDEP 1.80](#). The attached QA.csv file also serves as data source for the QA sections in the articles describing the individual tables.

Checks added in so far have been collated from 3 sources:

- CASCADE
- EPPICC FPV/DRV mergers
- PENTA Eurocoord merger

Checks on a table's additional fields have not been included at this stage

Only a few checks for non-HICDEP tables, non-HICDEP fields in HICDEP tables and Study-specific checks have been included at this stage for demo/testing purposes.

## Error Codes

The error codes allow to unambiguously reference a certain problem with a given data set. When merging data from different cohorts, the data manager may report problems for instance as follows:

[ATC006] in tblART/ART\_ID: there is no drug named 'XYZ' in the coding list

**Note:** the use of the error code alone is not sufficient to locate a problem since some of them are fairly generic.

Used like above, error codes can help to clearly communicate issues. This holds especially if the QA requirements are being checked by an automatic system.

## QA checks listed nowhere else

These are QA checks which are either not directly applicable to a specific table or to tables which are not (yet) part of HICDEP.

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
Appendix	CrossTable	GC001	New patients who were not submitted last time we did this merger	EPPICC	NO
Appendix	CrossTable	GC002	Patients left out who were submitted last time we did this merger	EPPICC	NO
tblAE_NEW	CrossTable	ANC001	tblBAS says AE_FPV=1, yet records in tblAE_NEW whilst on FPV	EPPICC	NO
tblAE_NEW	CrossTable	ANC002	AE_DRUG not reported in tblART or tblMED	EPPICC	NO
tblART	WithinTable	AW003	ART_RS=98 yet ART_OTH is null	EPPICC	NO
tblMED	WithinTable	MW002	MED_ONG=0 and MED_ED null	EPPICC	NO
tblMED	WithinTable	MW003	MED_ONG=1 and MED_ED non-null	EPPICC	NO
Appendix	CrossTable	GC003	Spot-check dates to ensure no Excel date-zero errors	EPPICC	NO
tblAE_NADM	WithinTable	ANDW001	Duplicate records for same EVENT_ID	EPPICC	NO
tblAE_NADM	CrossTable	ANDC001	EVENT_ID doesn't exist in tblAE_NEW	EPPICC	NO
tblAE_NADM	WithinTable	ANDW002	Missing Patient	EPPICC	NO
tblAE_NADM	WithinTable	ANDW003	Missing EVENT_ID	EPPICC	NO
tblAE_NADM	WithinTable	ANDW004	Missing CANTYP_T	EPPICC	NO
tblAE_NADM	WithinTable	ANDW005	Missing PATREP_S	EPPICC	NO
tblAE_NADM	WithinTable	ANDW006	DIAOTH_Y=1 and DIA_S is missing	EPPICC	NO
tblAE_NADM	CrossTable	ANDC002	PATIENT doesn't exist in tblBAS	EPPICC	NO
tblAE_NEW	CrossTable	ANC003	PATIENT has no record in BAS	EPPICC	NO
tblAE_NEW	CrossTable	ANC004	AE happened before that drug was started	EPPICC	NO



tblAE_NEW	WithinTable	ANW001	More than one record for same patient on same date with same AE_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW002	More than one record for same patient with same EVENT_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW003	Missing PATIENT	EPPICC	NO
tblAE_NEW	WithinTable	ANW004	Missing AE_D	EPPICC	NO
tblAE_NEW	WithinTable	ANW005	Missing AE_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW006	Missing EVENT_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW007	Missing AE_RES	EPPICC	NO
tblAE_NEW	WithinTable	ANW008	Missing AE_TEXT	EPPICC	NO
tblAE_NEW	WithinTable	ANW009	Missing AE_DRUG	EPPICC	NO
tblAE_NEW	CrossTable	ANC005	AE_ART=1 but doesn't agree with records in ART table	EPPICC	NO

### All QA checks

A complete list of all QA checks listed in the QA.csv file:

Table	Crosstable	Error Code	Description	Study specific	HICDEP?
AllTables	CrossTable	ATC001	any date in database after DEATH_D in tblLTFU		YES
AllTables	CrossTable	ATC002	any date in database after DROP_D in tblLTFU		YES
AllTables	CrossTable	ATC003	any date in database before BIRTH_D in tblBAS		YES
AllTables	CrossTable	ATC004	any date in database in the future		YES
AllTables	CrossTable	ATC005	patients submitted previously who have been missed out		YES
AllTables	CrossTable	ATC006	Any fields not coded as coding lists on table definition		YES
Appendix	CrossTable	GC001	New patients who were not submitted last time we did this merger	EPPICC	NO
Appendix	CrossTable	GC002	Patients left out who were submitted last time we did this merger	EPPICC	NO
tblAE_NEW	CrossTable	ANC001	tblBAS says AE_FPV=1, yet records in tblAE_NEW whilst on FPV	EPPICC	NO
tblAE_NEW	CrossTable	ANC002	AE_DRUG not reported in tblART or tblMED	EPPICC	NO
tblART	WithinTable	AW001	ART_RS not null, but end date NULL		YES
tblART	WithinTable	AW002	ART_RS null but end-date non NULL		YES
tblART	WithinTable	AW003	ART_RS=98 yet ART_OTH is null	EPPICC	NO
tblART	WithinTable	AW004	Duplicate records for same cohort, patient, art_id and art_sd		YES
tblART	WithinTable	AW005	Dose out of range for those dose units and frequency	PaediatricOnly	YES
tblART	WithinTable	AW006	Missing art_fr	PaediatricOnly	YES
tblART	WithinTable	AW007	Missing art_do	PaediatricOnly	YES
tblART	WithinTable	AW008	Missing patient		YES
tblART	WithinTable	AW009	Missing art_id		YES

tblART	WithinTable	AW010	Missing art_sd		YES
tblART	WithinTable	AW011	Overlapping periods of same drug		YES
tblART	WithinTable	AW012	Double reporting - records reported for both combination drugs and their components		YES
tblART	WithinTable	AW013	Periods of overlap of contra-indicated drugs		YES
tblART	WithinTable	AW014	Restart of same drug without a stop		YES
tblART	WithinTable	AW015	ART_SD greater than or equal to ART_ED		YES
tblART	CrossTable	AC001	Patient has no record in table BAS		YES
tblART	CrossTable	AC002	Records exist in tblART yet RECART_Y=0 in tblBAS		YES
tblBAS	WithinTable	BW001	AIDS date < SEROCO_D		YES
tblBAS	WithinTable	BW002	Duplicate patients		YES
tblBAS	WithinTable	BW003	First 3 chars of PATIENT don't form valid cohort code	CascadeOnly	YES
tblBAS	WithinTable	BW004	Missing PATIENT		YES
tblBAS	WithinTable	BW005	Missing CENTER		YES
tblBAS	WithinTable	BW006	Missing BIRTH_D		YES
tblBAS	WithinTable	BW007	Missing FRVIS_D		YES
tblBAS	WithinTable	BW008	Missing ENROL_D		YES
tblBAS	WithinTable	BW009	Missing GENDER		YES
tblBAS	WithinTable	BW010	Missing HEIGH		YES
tblBAS	WithinTable	BW011	Missing MODE		YES
tblBAS	WithinTable	BW012	Missing MODE_OTH if MODE=90		YES
tblBAS	WithinTable	BW013	Missing ORIGIN		YES
tblBAS	WithinTable	BW014	Missing ETHNIC		YES
tblBAS	WithinTable	BW015	Missing SEROCO_D		YES
tblBAS	WithinTable	BW016	Missing RECART_Y		YES
tblBAS	WithinTable	BW017	Missing AIDS_Y		YES
tblBAS	WithinTable	BW018	Missing AIDS_D if AIDS_Y=1		YES
tblBAS	WithinTable	BW019	BIRTH_D out of range (15-85 yrs)	CascadeOnly	YES
tblBAS	WithinTable	BW020	BIRTH_D out of range (<18)	PENTA	YES
tblBAS	CrossTable	BC001	RECART_Y=1 but no records in tblART		YES
tblBAS	CrossTable	BC002	AIDS_Y=0, but AIDS-defining records in tblDIS		YES
tblBAS	CrossTable	BC003	AIDS_Y=1 but no AIDS-defining records in tblDIS table		YES
tblDIS	WithinTable	DW001	Duplicate records for same DIS_ID and same DIS_D		YES
tblDIS	WithinTable	DW002	Miscoded DIS_WD as codes on table definition		YES
tblDIS	WithinTable	DW003	Miscoded DIS_ID - as in code list attached to table definition		YES
tblDIS	WithinTable	DW004	DIS_D missing		YES
tblDIS	WithinTable	DW005	DIS_ID missing		YES

tblDIS	WithinTable	DW006	Same event recorded twice - 2 records, same DIS_ID, DIS_D within 6 months		YES
tblDIS	WithinTable	DW007	DIS_ED present but before DIS_D		YES
tblDIS	CrossTable	DC001	Patient has no record in BAS		YES
tblDIS	CrossTable	DC002	AIDS-defining records, yet AIDS=0 in tblBAS		YES
tblDIS	CrossTable	DC003	First AIDS-defining DIS_D not equal to AIDS_D in tblBAS		YES
tblLAB	WithinTable	LW001	Duplicate records		YES
tblLAB	WithinTable	LW002	Missing LAB_D		YES
tblLAB	WithinTable	LW003	Missing LAB_V		YES
tblLAB	WithinTable	LW004	Missing LAB_U		YES
tblLAB	WithinTable	LW005	Missing LAB_ID		YES
tblLAB	WithinTable	LW008	Missing LAB_FA		YES
tblLAB	WithinTable	LW009	Missing LAB_ST		YES
tblLAB	WithinTable	LW010	LAB_V out of range for unit LAB_U		YES
tblLAB	CrossTable	LC001	Patient has no record in BAS table		YES
tblLAB_CD4	WithinTable	CW001	CD4 value out of range		YES
tblLAB_CD4	WithinTable	CW002	Duplicate records for same date		YES
tblLAB_CD4	WithinTable	CW003	Missing CD4_D		YES
tblLAB_CD4	WithinTable	CW004	Missing CD4_V		YES
tblLAB_CD4	WithinTable	CW005	Missing CD4_U		YES
tblLAB_CD4	WithinTable	CW006	Miscoded CD4_U as coding list on table definition		YES
tblLAB_CD4	WithinTable	CW007	CD4_U=2 (percentage) and CD4_V>100		YES
tblLAB_CD4	WithinTable	CW008	CD4_U=(1 or 3) and CD4_V>3000		YES
tblLAB_CD4	WithinTable	CW009	CD4 counts spike up or down suddenly (large change in less than a year)		YES
tblLAB_CD4	CrossTable	CC001	Patient has no record in BAS		YES
tblLAB_RES	WithinTable	LRW001	Duplicate records for same patient on same date		YES
tblLAB_RES	WithinTable	LRW002	Missing PATIENT		YES
tblLAB_RES	WithinTable	LRW003	Missing TEST_ID		YES
tblLAB_RES	WithinTable	LRW004	Missing SAMPLE_D		YES
tblLAB_RES	WithinTable	LRW005	Missing SEQ_DT		YES
tblLAB_RES	WithinTable	LRW006	Missing LAB		YES
tblLAB_RES	WithinTable	LRW007	Missing LIBRARY		YES
tblLAB_RES	WithinTable	LRW008	Missing REFSEQ		YES
tblLAB_RES	WithinTable	LRW009	Missing KIT		YES
tblLAB_RES	WithinTable	LRW010	Missing SOFTWARE		YES
tblLAB_RES	WithinTable	LRW011	Missing TESTTYPE		YES
tblLAB_RES	WithinTable	LRW012	Missing SUBTYPE		YES
tblLAB_RES	WithinTable	LRW013	SEQ_DT has no time part		YES
tblLAB_RES	WithinTable	LRW014	Missing VIRUSTYPE		YES
tblLAB_RES	CrossTable	LRC001	PATIENT has no record in tblBAS		YES

tblLAB_RES	CrossTable	LRC002	This TEST_ID has both LVL_1 and LVL_2 records		YES
tblLAB_RES	CrossTable	LRC003	TESTTYPE in (1,9) (i.e. not phenotype) yet no records in LVL_1 or LVL_2		YES
tblLAB_RES_LVL_1	WithinTable	L1W001	Duplicate records per TEST_ID and SEQTYPE		YES
tblLAB_RES_LVL_1	WithinTable	L1W002	SEQ_START > SEQ_STOP		YES
tblLAB_RES_LVL_1	WithinTable	L1W003	SEQ_NUC contains invalid IUPAC character		YES
tblLAB_RES_LVL_1	WithinTable	L1W004	Missing TEST_ID		YES
tblLAB_RES_LVL_1	WithinTable	L1W005	Missing SEQ_START		YES
tblLAB_RES_LVL_1	WithinTable	L1W006	Missing SEQ_STOP		YES
tblLAB_RES_LVL_1	WithinTable	L1W007	Missing SEQ_NUC		YES
tblLAB_RES_LVL_1	CrossTable	L1C001	TEST_ID not in tblLAB_RES.TEST_ID		YES
tblLAB_RES_LVL_2	WithinTable	L2W001	AA_FOUND_x but nothing in AA_FOUND(x-1)		YES
tblLAB_RES_LVL_2	WithinTable	L2W002	Duplicate records per TEST_ID, GENE, AA_POS and AA_POS_SUB		YES
tblLAB_RES_LVL_2	WithinTable	L2W003	Missing TEST_ID		YES
tblLAB_RES_LVL_2	WithinTable	L2W004	Missing GENE		YES
tblLAB_RES_LVL_2	WithinTable	L2W005	Missing AA_POS		YES
tblLAB_RES_LVL_2	WithinTable	L2W006	Missing AA_POS_SUB		YES
tblLAB_RES_LVL_2	WithinTable	L2W007	Missing AA_FOUND_1		YES
tblLAB_RES_LVL_2	CrossTable	L2C001	TEST_ID not in tblLAB_RES.TEST_ID		YES
tblLAB_RES_LVL_3	WithinTable	L3W001	Duplicate records for same TEST_ID and ATC_CODE		YES
tblLAB_RES_LVL_3	WithinTable	L3W002	Missing TEST_ID		YES
tblLAB_RES_LVL_3	WithinTable	L3W003	Missing ATC_CODE		YES
tblLAB_RES_LVL_3	WithinTable	L3W004	Missing RES_SCOR		YES
tblLAB_RES_LVL_3	CrossTable	L3C001	TEST_ID not in tblLAB_RES.TEST_ID		YES
tblLAB_RNA	WithinTable	RW001	Duplicate records for same RNA_D		YES
tblLAB_RNA	WithinTable	RW002	RNA_V < 0 and RNA_L missing		YES
tblLAB_RNA	WithinTable	RW003	RNA_V > 10 000 and (RNA_V modulo 1000) = 1 and RNA_UL missing		YES
tblLAB_RNA	WithinTable	RW004	Missing RNA_D		YES
tblLAB_RNA	WithinTable	RW005	Missing RNA_V		YES
tblLAB_RNA	WithinTable	RW006	RNA_V > 10 million		YES
tblLAB_RNA	WithinTable	RW007	RNA_V < RNA_L and RNA_V >= 0		YES
tblLAB_RNA	WithinTable	RW008	RNA_V > RNA_UL		YES
tblLAB_RNA	CrossTable	RC001	Patient has no record in BAS		YES
tblLAB_VIRO	CrossTable	LVC001	Patient doesn't have a record in BAS		YES
tblLAB_VIRO	WithinTable	LVW002	Missing patient		YES
tblLAB_VIRO	WithinTable	LVW003	Missing VS_ID		YES
tblLAB_VIRO	WithinTable	LVW004	Missing VS_D		YES

tblLAB_VIRO	WithinTable	LVW005	Missing VS_R		YES
tblLAB_VIRO	WithinTable	LVW006	Missing VS_V		YES
tblLAB_VIRO	WithinTable	LVW007	Missing VS_U		YES
tblLAB_VIRO	WithinTable	LVW008	More or less than exactly 1 positive HIV test per patient		YES
tblLAB_VIRO	WithinTable	LVW009	More than 1 negative HIV test for a patient		YES
tblLAB_VIRO	WithinTable	LVW010	Date of negative test after date of positive test		YES
tblLAB_VIRO	WithinTable	LVW011	Duplicate records		YES
tblLTFU	WithinTable	LFW001	DROP_Y and DEATH_Y both non-null		YES
tblLTFU	WithinTable	LFW002	DEATH_Y and DROP_RS both non-null		YES
tblLTFU	WithinTable	LFW003	Any of DEATH_Rx or DEATH_RCx non-null but DEATH_Y=0		YES
tblLTFU	WithinTable	LFW004	R2/RC2 non-null but R1/RC1 null		YES
tblLTFU	WithinTable	LFW005	R3/RC3 non-null but R2/RC2 null		YES
tblLTFU	WithinTable	LFW006	Duplicate patients		YES
tblLTFU	WithinTable	LFW007	Missing DROP_Y		YES
tblLTFU	WithinTable	LFW008	Missing DROP_D if DROP_Y=1		YES
tblLTFU	WithinTable	LFW009	Missing DROP_RS if DROP_Y=1		YES
tblLTFU	WithinTable	LFW010	Missing DEATH_Y		YES
tblLTFU	WithinTable	LFW011	Missing DEATH_D if DEATH_Y=1		YES
tblLTFU	WithinTable	LFW012	DEATH_D non null but DEATH_Y=0		YES
tblLTFU	CrossTable	LFC001	Patient not found in tblBAS		YES
tblLTFU	CrossTable	LFC002	patient in tblBAS hasn't got a record in tblLTFU		YES
tblLTFU	CrossTable	LFC003	tblBAS has AIDS=0 but DEATHRx =8.1		YES
tblMED	WithinTable	MW001	Duplicate records		YES
tblMED	WithinTable	MW002	MED_ONG=0 and MED_ED null	EPPICC	NO
tblMED	WithinTable	MW003	MED_ONG=1 and MED_ED non-null	EPPICC	NO
tblMED	WithinTable	MW004	Missing MED_ID		YES
tblMED	WithinTable	MW005	Missing MED_SD		YES
tblMED	WithinTable	MW006	Mising MED_ED		YES
tblMED	WithinTable	MW007	Overlapping periods of same drug		YES
tblMED	WithinTable	MW008	MED_ED < MED_SD		YES
tblMED	WithinTable	MW009	MED_RS not null, but end date NULL		YES
tblMED	CrossTable	MC001	Patient has no records in tblBAS		YES
tblOVERLAP	WithinTable	OW001	Invalid other cohort		YES
tblOVERLAP	CrossTable	OC001	PATIENT not found in tblBAS for that cohort		YES
tblOVERLAP	CrossTable	OC002	PAT_OTH not found in tblBAS for that overlapping cohort		YES

tblVIS	WithinTable	VW001	Duplicate records, same VIS_D		YES
tblVIS	WithinTable	VW002	Height decreasing over time		YES
tblVIS	WithinTable	VW003	Height out of acceptable range		YES
tblVIS	WithinTable	VW004	Weight out of acceptable range		YES
tblVIS	CrossTable	VC001	patient has no record in BAS table		YES
tblVIS	CrossTable	VC002	No weights within 3 mths of starting FPV/DRV	EPPICC	YES
Appendix	CrossTable	GC003	Spot-check dates to ensure no Excel date-zero errors	EPPICC	NO
tblAE_NADM	WithinTable	ANDW001	Duplicate records for same EVENT_ID	EPPICC	NO
tblAE_NADM	CrossTable	ANDC001	EVENT_ID doesn't exist in tblAE_NEW	EPPICC	NO
tblAE_NADM	WithinTable	ANDW002	Missing Patient	EPPICC	NO
tblAE_NADM	WithinTable	ANDW003	Missing EVENT_ID	EPPICC	NO
tblAE_NADM	WithinTable	ANDW004	Missing CANTYP_T	EPPICC	NO
tblAE_NADM	WithinTable	ANDW005	Missing PATREP_S	EPPICC	NO
tblAE_NADM	WithinTable	ANDW006	DIAOTH_Y=1 and DIA_S is missing	EPPICC	NO
tblAE_NADM	CrossTable	ANDC002	PATIENT doesn't exist in tblBAS	EPPICC	NO
tblAE_NEW	CrossTable	ANC003	PATIENT has no record in BAS	EPPICC	NO
tblAE_NEW	CrossTable	ANC004	AE happened before that drug was started	EPPICC	NO
tblAE_NEW	WithinTable	ANW001	More than one record for same patient on same date with same AE_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW002	More than one record for same patient with same EVENT_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW003	Missing PATIENT	EPPICC	NO
tblAE_NEW	WithinTable	ANW004	Missing AE_D	EPPICC	NO
tblAE_NEW	WithinTable	ANW005	Missing AE_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW006	Missing EVENT_ID	EPPICC	NO
tblAE_NEW	WithinTable	ANW007	Missing AE_RES	EPPICC	NO
tblAE_NEW	WithinTable	ANW008	Missing AE_TEXT	EPPICC	NO
tblAE_NEW	WithinTable	ANW009	Missing AE_DRUG	EPPICC	NO
tblAE_NEW	CrossTable	ANC005	AE_ART=1 but doesn't agree with records in ART table	EPPICC	NO
tblDELIVERY_CHILD	CrossTable	DCC001	MOTHER_ID doesn't exist in tblPREG		YES
tblDELIVERY_CHILD	CrossTable	DCC002	CHILD_ID doesn't exist in tblPREG_OUT		YES
tblDELIVERY_CHILD	WithinTable	DCW001	Duplicate records for same Mother_ID, CHILD_ID		YES
tblDELIVERY_CHILD	WithinTable	DCW002	B_SEQ >4		YES
tblDELIVERY_CHILD	WithinTable	DCW003	B_SEQ>1, but no record exists for same Mother, Child with B_SEQ=1		YES
tblDELIVERY_CHILD	WithinTable	DCW005	DELIV_D < MEMRUP_D		YES
tblDELIVERY_CHILD	WithinTable	DCW004	LABOUR_P >72		YES

tblDELIVERY_CHILD	WithinTable	DCW006	2 children, same mother, same MEMRUP_D, DELIV_D > DELIV_D, yet B_SEQ < B_SEQ		YES
tblDELIVERY_MUM	CrossTable	DMC001	MOTHER_ID doesn't exist in tblPREG		YES
tblDELIVERY_MUM	CrossTable	DMC002	MOTHER_ID doesn't exist in tblDELIVERY_CHILD with same MEMRUP_D		YES
tblDELIVERY_MUM	WithinTable	DMW001	PREG_SEQ invalid (PREG_SEQ > 1 and PREG_SEQ = x, but PREG_SEQ = x-1 doesn't exist)		YES
tblDELIVERY_MUM	WithinTable	DMW002	DISCHA_D < MEMRUP_D		YES
tblDELIVERY_MUM	WithinTable	DMW004	INTERV=0-3 but INTERV_O non null		YES
tblDELIVERY_MUM	WithinTable	DMW003	INTERV=90,91, 92 or 93 but INTERV_O null		YES
tblNEWBORN	CrossTable	NC001	CHILD_ID doesn't exist in tblDELIVERY_CHILD		YES
tblNEWBORN	CrossTable	NC002	ABNORM_Y = 1, yet no records in tblNEWBORN_ABNORM		YES
tblNEWBORN	CrossTable	NC003	ABNORM_Y = 0 or 9, yet records in tblNEWBORN_ABNORM		YES
tblNEWBORN	WithinTable	NW001	BRFEED_SD > BRFEED_ED		YES
tblNEWBORN	WithinTable	NW002	APGARM_x's out of order (e.g APGARM_3 < APGARM_2)		YES
tblNEWBORN	WithinTable	NW003	ICU_Y=1, but ICU_S or ICS_D null		YES
tblNEWBORN	WithinTable	NW004	ICU_Y=0 or 9, but ICU_S or ICU_D non null		YES
tblNEWBORN_ABNORM	CrossTable	NAC001	CHILD_ID doesn't exist in NEWBORN		YES
tblNEWBORN_ABNORM	WithinTable	NAW001	ABNORM_T=90 but ABNORM_S null		YES
tblPREG_OBS	CrossTable	POC001	MOTHER_ID+PREG_SEQ doesn't exist in tblPREG		YES
tblPREG_OBS	WithinTable	POW001	PROB_T=99 but PROB_S null		YES
tblPREG_OBS	WithinTable	POW002	PROB_T <> 99 but PROB_S non null		YES
tblPREG_OBS	WithinTable	POW003	PROB_T=2 but CERVIX_S null		YES
tblPREG_OBS	WithinTable	POW003	PROB_T <> 2 but CERVIX_S non null		YES
tblPREG_OUT	CrossTable	PTC001	MOTHER_ID+PREG_SEQ doesn't exist in tblPREG		YES
tblPREG_OUT	CrossTable	PTC002	OUTCOM=1,2 or 3 and CHILD_ID doesn't exist in tblNEWBORN or tblDELIVERYCHILD		YES
tblPREG_OUT	CrossTable	PTC003	OUTCOM=10,11,20 or 21 and CHILD_ID exists in tblNEWBORN or tblDELIVERYCHILD		YES
tblPREG_OUT	WithinTable	PTW001	OUTCOME=20 or 21 and OUTCOM_R null		YES
tblPREG_OUT	WithinTable	PTW002	OUTCOME not 20 or 21 and OUTCOM_R non null		YES
tblPREG	WithinTable	PW001	ANC_D		YES

tblPREG	WithinTable	PW002	INPREG_Y=2,3,4 but INHIST_Y null		YES
tblPREG	WithinTable	PW003	INPREG_Y=1 but INHIST_Y non null		YES
tblPREG	WithinTable	PW004	INHIST_Y=1 but INHIST_S null		YES
tblPREG	WithinTable	PW004	INHIST_Y=0 or 9 but INHIST_S non null		YES
tblPREG	WithinTable	PW005	KARYO_T=2 but KARYO_A null		YES
tblPREG	WithinTable	PW006	KARYO_T<>2 but KARYO_A non null		YES
tblPREG	WithinTable	PW007	ULTRA_x =2, but ULTR_A_x null		YES
tblPREG	WithinTable	PW008	ULTRA_x <>2, but ULTR_A_x non null		YES
tblPREG	CrossTable	PC001	PROB_Y = 1 but no records in tblPREG_OBS		YES
tblPREG	CrossTable	PC002	PROB_Y = 0 or 9 but records in tblPREG_OBS		YES
tblREFILL	CrossTable	PRC001	Patient doesn't have a record in BAS		YES
tblREFILL	WithinTable	PRW002	Missing PATIENT		YES
tblREFILL	WithinTable	PRW003	Missing REFILL_D		YES
tblREFILL	WithinTable	PRW004	Missing DRUG_ID		YES
tblREFILL	WithinTable	PRW005	Missing SUPPLY		YES
tblREFILL	WithinTable	PRW006	SUPPLY < 1		YES