Package 'pharmaverseadam'

October 25, 2024

```
Type Package
Title ADaM Test Data for the 'Pharmaverse' Family of Packages
Version 1.1.0
Description A set of Analysis Data Model (ADaM) datasets constructed using the
      Study Data Tabulation Model (SDTM) datasets contained in the 'pharmaversesdtm' package and
      the template scripts from the 'admiral' family of packages. ADaM dataset specifications
      are described in the CDISC ADaM implementation guide, accessible by creating a free ac-
      count on <https://www.cdisc.org/>.
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      https://github.com/pharmaverse/pharmaverseadam/
Encoding UTF-8
Language en-US
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LazyDataCompression bzip2
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Description

Adverse Events Analysis

Usage

adae

Format

A data frame with 105 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

AESEQ Sequence Number

AESPID Sponsor-Defined Identifier

AETERM Reported Term for the Adverse Event

AELLT Lowest Level Term

AELLTCD Lowest Level Term Code

AEDECOD Dictionary-Derived Term

AEPTCD Preferred Term Code

AEHLT High Level Term

AEHLTCD High Level Term Code

AEHLGT High Level Group Term

AEHLGTCD High Level Group Term Code

AEBODSYS Body System or Organ Class

AEBDSYCD Body System or Organ Class Code

AESOC Primary System Organ Class

AESOCCD Primary System Organ Class Code

AESEV Severity/Intensity

AESER Serious Event

AEACN Action Taken with Study Treatment

AEREL Causality

AEOUT Outcome of Adverse Event

AESCAN Involves Cancer

AESCONG Congenital Anomaly or Birth Defect

AESDISAB Persist or Signif Disability/Incapacity

AESDTH Results in Death

AESHOSP Requires or Prolongs Hospitalization

AESLIFE Is Life Threatening

AESOD Occurred with Overdose

AEDTC Date/Time of Collection

AESTDTC Start Date/Time of Adverse Event

AEENDTC End Date/Time of Adverse Event

AESTDY Study Day of Start of Adverse Event

AEENDY Study Day of End of Adverse Event

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

DTHDT Date of Death

EOSDT End of Study Date

ASTDTM Analysis Start Date/Time

ASTDTF Analysis Start Date Imputation Flag

ASTTMF Analysis Start Time Imputation Flag

AENDTM Analysis End Date/Time

AENDTF Analysis End Date Imputation Flag

AENTMF Analysis End Time Imputation Flag

ASTDT Analysis Start Date

AENDT Analysis End Date

ASTDY Analysis Start Relative Day

AENDY Analysis End Relative Day

ADURN Analysis Duration (N)

ADURU Analysis Duration Units

LDOSEDTM End Date/Time of Last Dose

ASEV Analysis Severity/Intensity

AREL Analysis Causality

TRTEMFL Treatment Emergent Analysis Flag

ASEVN Analysis Severity/Intensity (N)

AOCCIFL 1st Max Sev./Int. Occurrence Flag

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_adae.R).

References

None

Examples

data("adae")

adbcva_ophtha

adbcva_ophtha

Description

Best Corrected Visual Acuity Analysis

Usage

adbcva_ophtha

Format

A data frame with 120 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

OESEQ Sequence Number

OECAT Category for Ophthalmic Test or Exam

OESCAT Subcategory for Ophthalmic Test or Exam

OEDTC Date/Time of Collection

VISIT Visit Name

VISITNUM Visit Number

VISITDY Planned Study Day of Visit

OESTRESN Numeric Result/Finding in Standard Units

OESTRESC Character Result/Finding in Std Format

OEORRES Result or Finding in Original Units

OETEST Name of Ophthalmic Test or Exam

OETESTCD Short Name of Ophthalmic Test or Exam

OETSTDTL Ophthalmic Test or Exam Detail

OELAT Laterality

OELOC Location Used for the Measurement

OEDY Study Day of Visit/Collection/Exam

OEMETHOD Method of Test or Examination

OEORRESU Original Units

OESTRESU Standard Units

OESTAT Completion Status

OETPT Planned Time Point Name

OETPTNUM Planned Time Point Number

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

STUDYEYE Study Eye Location

AVAL Analysis Value

AVALU Analysis Value Unit

DTYPE Derivation Type

AFEYE Affected Eye

PARAM Parameter

PARAMCD Parameter Code

AVALC Analysis Value (C)

ADT Analysis Date

ADY Analysis Relative Day

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

BASETYPE Baseline Type

ONTRTFL On Treatment Record Flag

ABLFL Baseline Record Flag

ANL01FL Analysis Flag 01

ANL02FL Analysis Flag 02

WORS01FL Worst Post Baseline Obs

BASE Baseline Value

BASEC Baseline Value (C)

CHG Change from Baseline

PCHG Percent Change from Baseline

ASEQ Analysis Sequence Number

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country/Region

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

CRIT1 Analysis Criterion 1

CRIT1FL Criterion 1 Evaluation Result Flag

CRIT2 Analysis Criterion 2

CRIT2FL Criterion 2 Evaluation Result Flag

CRIT3 Analysis Criterion 3

CRIT3FL Criterion 3 Evaluation Result Flag

CRIT4 Analysis Criterion 4

CRIT4FL Criterion 4 Evaluation Result Flag

CRIT5 Analysis Criterion 5

CRIT5FL Criterion 5 Evaluation Result Flag

CRIT6 Analysis Criterion 6

CRIT6FL Criterion 6 Evaluation Result Flag

CRIT7 Analysis Criterion 7

CRIT7FL Criterion 7 Evaluation Result Flag

CRIT8 Analysis Criterion 8

CRIT8FL Criterion 8 Evaluation Result Flag

AVALCA1N Analysis Value Category 1 (N)

AVALCAT1 Analysis Value Category 1

Source

Generated from admiralophtha package (template ad_adbcva.R).

References

None

Examples

data("adbcva_ophtha")

10 adce_vaccine

adce_vaccine

adce_vaccine

Description

Clinical Events Analysis for Vaccine

Usage

adce_vaccine

Format

A data frame with 56 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

CESEQ Sequence Number

CELNKID Link ID

CELNKGRP Link Group ID

CETERM Reported Term for the Clinical Event

CEDECOD Dictionary-Derived Term

CELAT Laterality

CELOC Location of Event

CECAT Category for the Clinical Event

CESCAT Subcategory for the Clinical Event

CEPRESP Clinical Event Pre-specified

CEOCCUR Clinical Event Occurrence

CESEV Severity/Intensity

CEREL Causality

CEOUT Outcome of Event

EPOCH Epoch

CEDTC Date/Time of Event Collection

CESTDTC Start Date/Time of Clinical Event

CEENDTC End Date/Time of Clinical Event

CEDUR Duration of Clinical Event

CETPT Planned Time Point Name

CETPTNUM Planned Time Point Number

CETPTREF Time Point Reference

adce_vaccine 11

CERFTDTC Date/Time of Reference Time Point

CEEVINTX Evaluation Interval Text

CESTAT Completion Status

CEREASND Reason Clinical Event Not Collected

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

ASTDT Analysis Start Date

AENDT Analysis End Date

ASTDY Analysis Start Relative Day

AENDY Analysis End Relative Day

APERIOD Period

APERSDT Period Start Date

APEREDT Period End Date

APERSTDY Analysis Sub-period Start Relative Day

AREL Analysis Causality

ASEV Analysis Severity/Intensity

ASEVN Analysis Severity/Intensity (N)

AOCC01FL Event Occurrence Flag

ASEQ Analysis Sequence Number

ADURN Analysis Duration (N)

ADURU Analysis Duration Units

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

AGE Age

AGEU Age Units

SEX Sex

RACE Race

COUNTRY Country

ETHNIC Ethnicity

SITEID Study Site Identifier

SUBJID Subject Identifier for the Study

Source

Generated from admiralvaccine package (template ad_adce.R).

References

None

Examples

data("adce_vaccine")

12 adcm

adcm adcm

Description

Concomitant Medications Analysis

Usage

adcm

Format

A data frame with 95 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

CMSEQ Sequence Number

CMSPID Sponsor-Defined Identifier

CMTRT Reported Name of Drug, Med, or Therapy

CMDECOD Standardized Medication Name

CMINDC Indication

CMCLAS Medication Class

CMDOSE Dose per Administration

CMDOSU Dose Units

CMDOSFRQ Dosing Frequency per Interval

CMROUTE Route of Administration

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

CMDTC Date/Time of Collection

CMSTDTC Start Date/Time of Medication

CMENDTC End Date/Time of Medication

CMSTDY Study Day of Start of Medication

CMENDY Study Day of End of Medication

CMENRTPT undocumented field

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

DTHDT Date of Death

adcm 13

EOSDT End of Study Date

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

ASTDTM Analysis Start Date/Time

ASTDTF Analysis Start Date Imputation Flag

ASTTMF Analysis Start Time Imputation Flag

AENDTM Analysis End Date/Time

AENDTF Analysis End Date Imputation Flag

AENTMF Analysis End Time Imputation Flag

ASTDT Analysis Start Date

AENDT Analysis End Date

ASTDY Analysis Start Relative Day

AENDY Analysis End Relative Day

ADURN Analysis Duration (N)

ADURU Analysis Duration Units

ONTRTFL On Treatment Record Flag

PREFL Pre-treatment Flag

FUPFL Follow-up Flag

ANL01FL Analysis Flag 01

AOCCPFL 1st Occurrence of Preferred Term Flag

APHASE Phase

APHASEN Description of Phase N

TRTP Planned Treatment

TRTA Actual Treatment

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

14 adcm

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Treatment End Datetime Imput Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_adcm.R).

adeg 15

References

None

Examples

data("adcm")

adeg

adeg

Description

Electrocardiogram Tests Analysis

Usage

adeg

Format

A data frame with 108 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

EGSEQ Sequence Number

EGTESTCD ECG Test or Examination Short Name

EGTEST ECG Test or Examination Name

EGORRES Result or Finding in Original Units

EGORRESU Original Units

EGSTRESC Character Result/Finding in Std Format

EGSTRESN Numeric Result/Finding in Standard Units

EGSTRESU Standard Units

EGSTAT Completion Status

EGLOC Lead Location Used for Measurement

EGBLFL Baseline Flag

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

EGDTC Date/Time of ECG

EGDY Study Day of ECG

16 adeg

EGTPT Planned Time Point Name

EGTPTNUM Planned Time Point Number

EGELTM Planned Elapsed Time from Time Point Ref

EGTPTREF Time Point Reference

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

ADTM Analysis Datetime

ATMF Analysis Time Imputation Flag

ADY Analysis Relative Day

PARAMCD Parameter Code

AVAL Analysis Value

AVALC Analysis Value (C)

ADT Analysis Date

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

DTYPE Derivation Type

ONTRTFL On Treatment Record Flag

ANRLO Analysis Normal Range Lower Limit

ANRHI Analysis Normal Range Upper Limit

ANRIND Analysis Reference Range Indicator

BASETYPE Baseline Type

ABLFL Baseline Record Flag

BASE Baseline Value

BASEC Baseline Value (C)

BNRIND Baseline Reference Range Indicator

CHG Change from Baseline

PCHG Percent Change from Baseline

ANL01FL Analysis Flag 01

TRTP Planned Treatment

TRTA Actual Treatment

ASEQ Analysis Sequence Number

AVALCAT1 Analysis Value Category 1

AVALCA1N Analysis Value Category 1 (N)

adeg 17

CHGCAT1 Change from Baseline Category 1

CHGCAT1N Change from Baseline Category 1 (N)

PARAM Parameter

PARAMN Parameter (N)

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_adeg.R).

References

None

Examples

data("adeg")

adex adex

Description

Exposure Analysis

Usage

adex

Format

A data frame with 92 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

EXSEQ Sequence Number

EXTRT Name of Treatment

EXDOSE Dose

EXDOSU Dose Units

EXDOSFRM Dose Form

EXDOSFRQ Dosing Frequency per Interval

EXROUTE Route of Administration

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

EXSTDTC Start Date/Time of Treatment

EXENDTC End Date/Time of Treatment

EXSTDY Study Day of Start of Treatment

EXENDY Study Day of End of Treatment

EXADJ Reason for Dose Adjustment

EXPLDOS Planned Dose

TRTSDT Date of First Exposure to Treatment

TRTSDTM Datetime of First Exposure to Treatment

TRTEDTM Datetime of Last Exposure to Treatment

ASTDTM Analysis Start Datetime

ASTDTF Analysis Start Date Imputation Flag

ASTTMF Analysis Start Time Imputation Flag

AENDTM Analysis End Datetime

AENDTF Analysis End Date Imputation Flag

AENTMF Analysis End Time Imputation Flag

ASTDY Analysis Start Relative Day

AENDY Analysis End Relative Day

EXDURD Duration of Treatment (Days)

ASTDT Analysis Start Date

AENDT Analysis End Date

DOSEO Dose O

PDOSEO PDose O

PARAMCD Parameter Code

AVAL Analysis Value

AVALC Analysis Value (C)

PARCAT1 Parameter Category 1

PARAM Parameter

PARAMN Parameter (N)

AVALCAT1 Analysis Value Category 1

ASEQ Analysis Sequence Number

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

TRTSTMF Time of First Exposure Imput. Flag

TRTETMF Time of Last Exposure Imput. Flag

TRTEDT Date of Last Exposure to Treatment

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_adex.R).

References

None

Examples

data("adex")

22 adface_vaccine

adface_vaccine

adface_vaccine

Description

Findings About Clinical Events Analysis

Usage

adface_vaccine

Format

A data frame with 61 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

SUBJID Subject Identifier for the Study

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

SAFFL Safety Population Flag

ARM Description of Planned Arm

ARMCD Planned Arm Code

ACTARM Description of Actual Arm

ACTARMCD Actual Arm Code

TRTSDT Date of First Exposure to Treatment

TRTSDTM Datetime of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRTEDTM Datetime of Last Exposure to Treatment

FATEST Findings About Test Name

FALNKID Link ID

FALNKGRP Link Group ID

FATESTCD Findings About Test Short Name

PARAMCD Parameter Code

PARAM Parameter

PARAMN Parameter (N)

FAOBJ Object of the Observation

adface_vaccine 23

PARCAT1 Parameter Category 1

PARCAT2 Parameter Category 2

AVALC Analysis Value (C)

AVAL Analysis Value

FASTAT Completion Status

FAREASND Reason Not Performed

FAEVAL Evaluator

EPOCH Epoch

ADT Analysis Date

ADTM Analysis Datetime

FAEVINTX Evaluation Interval Text

ADY Analysis Relative Day

ATPT Analysis Timepoint

ATPTN Analysis Timepoint (N)

ATPTREF Analysis Timepoint Reference

EXDOSE Dose

EXTRT Name of Treatment

EXSTDTC Start Date/Time of Treatment

EXENDTC End Date/Time of Treatment

TRTA Actual Treatment

TRTP Planned Treatment

APERIOD Period

APERSDT Period Start Date

APEREDT Period End Date

FAORRES Result or Finding in Original Units

TRT01P Planned Treatment for Period 01

TRT02P Planned Treatment for Period 02

TRT01A Actual Treatment for Period 01

TRT02A Actual Treatment for Period 02

VAX01DT Vaccination Date 01

VAX02DT Vaccination Date 02

EVENTFL Event Value Flag

EVENTDFL Day Event Value Flag

ANL01FL Analysis Flag 01

ANL02FL Analysis Flag 02

ANL03FL undocumented field

24 adis_vaccine

Source

Generated from admiralvaccine package (template ad_adface.R).

References

None

Examples

```
data("adface_vaccine")
```

adis_vaccine

adis_vaccine

Description

Immunogenicity Specimen Assessments

Usage

adis_vaccine

Format

A data frame with 102 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

ISSEQ Sequence Number

ISTESTCD Immunogenicity Test/Exam Short Name

ISTEST Immunogenicity Test or Examination Name

ISCAT Category for Immunogenicity Test

ISORRES Results or Findings in Original Units

ISORRESU Original Units

ISSTRESC Character Result/Finding in Std Format

ISSTRESN Numeric Results/Findings in Std. Units

ISSTRESU Standard Units

ISSTAT Completion Status

ISREASND Reason Not Done

ISNAM Vendor Name

ISSPEC Specimen Type

ISMETHOD Method of Test or Examination

adis_vaccine 25

ISBLFL Baseline Flag

ISLLOQ Lower Limit of Quantitation

VISITNUM Visit Number

EPOCH Epoch

ISDTC Date/Time of Collection

ISDY Study Day of Visit/Collection/Exam

ISULOQ Upper Limit of Quantitation

LOD Limit of Detection

AVISITN Analysis Visit (N)

AVISIT Analysis Visit

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

ATPTREF Analysis Timepoint Reference

ADT Analysis Date

RFSTDTC Subject Reference Start Date/Time

PPROTFL Per-Protocol Population Flag

ADY Analysis Relative Day

PARAMCD Parameter Code

PARAM Parameter

PARAMN Parameter (N)

PARCAT1 Parameter Category 1

CUTOFF02 First Cutoff Value

CUTOFF03 Second Cutoff Value

AVAL Analysis Value

AVALU Analysis Value Unit

SERCAT1 Pre-vaccination seropositivity status

SERCAT1N Pre-vaccination sero status (n)

DTYPE Derivation Type

BASETYPE Baseline Type

BASE Baseline Value

ABLFL Baseline Record Flag

BASECAT1 Baseline Category 1

CHG Change from Baseline

R2BASE Ratio to Baseline

CRIT1FL Criterion 1 Evaluation Result Flag

CRIT1FN Criterion 1 Evaluation Result Flag (N)

CRIT1 Analysis Criterion 1

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APERIOD Period

APERSDT Period Start Date

APEREDT Period End Date

TRTA Actual Treatment

TRTP Planned Treatment

PPSRFL Per-Protocol Record-Level Flag

SUBJID Subject Identifier for the Study

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

INVID Investigator Identifier

INVNAM Investigator Name

BRTHDTC Date/Time of Birth

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

TRT02P Planned Treatment for Period 02

TRT01A Actual Treatment for Period 01

TRT02A Actual Treatment for Period 02

TRTSDTM Datetime of First Exposure to Treatment

TRTEDTM Datetime of Last Exposure to Treatment

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

VAX01DT Vaccination Date 01

VAX02DT Vaccination Date 02

AP01SDT Period 01 Start Date

AP01EDT Period 01 End Date

AP02SDT Period 02 Start Date

AP02EDT Period 02 End Date

Source

Generated from admiralvaccine package (template ad_adis.R).

References

None

Examples

data("adis_vaccine")

adlb

adlb

Description

Laboratory Analysis

Usage

adlb

Format

A data frame with 115 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

LBSEQ Sequence Number

LBTESTCD Lab Test or Examination Short Name

LBTEST Lab Test or Examination Name

LBCAT Category for Lab Test

LBORRES Result or Finding in Original Units

LBORRESU Original Units

LBORNRLO Reference Range Lower Limit in Orig Unit

LBORNRHI Reference Range Upper Limit in Orig Unit

LBSTRESC Character Result/Finding in Std Format

LBSTRESN Numeric Result/Finding in Standard Units

LBSTRESU Standard Units

LBSTNRLO Reference Range Lower Limit-Std Units

LBSTNRHI Reference Range Upper Limit-Std Units

LBNRIND Reference Range Indicator

LBBLFL Baseline Flag

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

LBDTC Date/Time of Specimen Collection

LBDY Study Day of Specimen Collection

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

ADT Analysis Date

ADY Analysis Relative Day

PARAMCD Parameter Code

PARAM Parameter

PARAMN Parameter (N)

PARCAT1 Parameter Category 1

AVAL Analysis Value

AVALC Analysis Value (C)

ANRLO Analysis Normal Range Lower Limit

ANRHI Analysis Normal Range Upper Limit

DTYPE Derivation Type

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

ONTRTFL On Treatment Record Flag

ANRIND Analysis Reference Range Indicator

BASETYPE Baseline Type

ABLFL Baseline Record Flag

BASE Baseline Value

BASEC Baseline Value (C)

BNRIND Baseline Reference Range Indicator

CHG Change from Baseline

PCHG Percent Change from Baseline

ATOXDSCL Analysis Toxicity Description Low

ATOXDSCH Analysis Toxicity Description High

ATOXGRL Analysis Toxicity Grade Low

ATOXGRH Analysis Toxicity Grade High

ATOXGR Analysis Toxicity Grade

BTOXGRL Baseline Toxicity Grade Low

BTOXGRH Baseline Toxicity Grade High

BTOXGR Baseline Toxicity Grade

R2BASE Ratio to Baseline

R2ANRLO Ratio of Analysis Val compared to ANRLO

R2ANRHI Ratio of Analysis Val compared to ANRHI

SHIFT1 Shift from Baseline to Analysis Value

SHIFT2 Shift from Baseline to Overall Grade

ANL01FL Analysis Flag 01

LVOTFL Last Value On Treatment Record Flag

TRTP Planned Treatment

TRTA Actual Treatment

ASEQ Analysis Sequence Number

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

adlbhy 31

Source

Generated from admiral package (template ad_adlb.R).

References

None

Examples

data("adlb")

adlbhy

adlbhy

Description

Analysis of Lab Hy's Law

Usage

adlbhy

Format

A data frame with 14 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

TRT01A Actual Treatment for Period 01

PARAMCD Parameter Code

LBSEQ Sequence Number

ADT Analysis Date

AVISIT Analysis Visit

ADY Analysis Relative Day

AVAL Analysis Value

ANRHI Analysis Normal Range Upper Limit

CRIT1 Analysis Criterion 1

CRIT1FL Criterion 1 Evaluation Result Flag

AVALC Analysis Value (C)

PARAM Parameter

Source

Generated from admiral package (template ad_adlbhy.R).

References

None

Examples

data("adlbhy")

admh

admh

Description

Medical History Analysis

Usage

admh

Format

A data frame with 114 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

MHSEQ Sequence Number

MHSPID Sponsor-Defined Identifier

MHTERM Reported Term for the Medical History

MHLLT Lowest Level Term

MHDECOD Dictionary-Derived Term

MHHLT High Level Term

MHHLGT High Level Group Term

MHCAT Category for Medical History

MHBODSYS Body System or Organ Class

MHSEV Severity/Intensity

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

MHDTC Date/Time of History Collection

MHSTDTC Start Date/Time of Medical History Event

MHDY Study Day of History Collection

MHENDTC End Date/Time of Medical History Event

MHPRESP Medical History Event Pre-Specified

MHOCCUR Medical History Occurrence

MHSTRTPT Start Relative to Reference Time Point

MHENRTPT End Relative to Reference Time Point

MHSTTPT Start Reference Time Point

MHENTPT End Reference Time Point

MHENRF End Relative to Reference Period

MHSTAT Completion Status

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

DTHDT Date of Death

EOSDT End of Study Date

ASTDT Analysis Start Date

AENDT Analysis End Date

ASTDY Analysis Start Relative Day

AENDY Analysis End Relative Day

ADT Analysis Date

ADY Analysis Relative Day

SMQ02NAM SMQ 02 Name

SMQ02CD SMQ 02 Code

SMQ02SC SMQ 02 Scope

SMQ02SCN SMQ 02 Scope (N)

SMQ03NAM SMQ 03 Name

SMQ03CD SMQ 03 Code

SMQ03SC SMQ 03 Scope

SMQ03SCN SMQ 03 Scope (N)

SMQ05NAM SMQ 05 Name

SMQ05CD SMQ 05 Code

SMQ05SC SMQ 05 Scope

SMQ05SCN SMQ 05 Scope (N)

CQ01NAM Customized Query 01 Name

CQ04NAM Customized Query 04 Name

CQ04CD Customized Query 04 Code

AHIST Response of Med Hx (past or current)

AOCCFL 1st Occurrence within Subject Flag

AOCCSFL 1st Occurrence of SOC Flag

AOCCPFL 1st Occurrence of Preferred Term Flag

AOCPFL 1st Occur w/in Trt Prd FL

AOCPSFL 1st Occur of SOC w/in Trt Prd FL

AOCPPFL 1st Occur of PT w/in Trt Prd FL

ANL01FL Analysis Flag 01

TRTP Planned Treatment

TRTA Actual Treatment

APHASE Phase

APHASEN Description of Phase N

MHTERMN Medical History Term (N)

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Treatment End Datetime Imput Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_admh.R).

References

None

Examples

data("admh")

36 adoe_ophtha

adoe_ophtha

adoe_ophtha

Description

Exam Analysis for Ophthalmology

Usage

adoe_ophtha

Format

A data frame with 102 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

OESEQ Sequence Number

OECAT Category for Ophthalmic Test or Exam

OESCAT Subcategory for Ophthalmic Test or Exam

OEDTC Date/Time of Collection

VISIT Visit Name

VISITNUM Visit Number

VISITDY Planned Study Day of Visit

OESTRESN Numeric Result/Finding in Standard Units

OESTRESC Character Result/Finding in Std Format

OEORRES Result or Finding in Original Units

OETEST Name of Ophthalmic Test or Exam

OETESTCD Short Name of Ophthalmic Test or Exam

OETSTDTL Ophthalmic Test or Exam Detail

OELAT Laterality

OELOC Location Used for the Measurement

OEDY Study Day of Visit/Collection/Exam

OEMETHOD Method of Test or Examination

OEORRESU Original Units

OESTRESU Standard Units

OESTAT Completion Status

OETPT Planned Time Point Name

OETPTNUM Planned Time Point Number

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TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

STUDYEYE Study Eye Location

AVAL Analysis Value

AVALC Analysis Value (C)

AVALU Analysis Value Unit

DTYPE Derivation Type

AFEYE Affected Eye

PARAM Parameter

PARAMCD Parameter Code

ADT Analysis Date

ADY Analysis Relative Day

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

BASETYPE Baseline Type

ONTRTFL On Treatment Record Flag

ABLFL Baseline Record Flag

ANL01FL Analysis Flag 01

ANL02FL Analysis Flag 02

WORS01FL Worst Post Baseline Obs

BASE Baseline Value

BASEC Baseline Value (C)

CHG Change from Baseline

PCHG Percent Change from Baseline

ASEQ Analysis Sequence Number

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

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DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiralophtha package (template ad_adoe.R).

References

None

Examples

```
data("adoe_ophtha")
```

adpc

adpc

Description

Pharmacokinetic Concentrations

Usage

adpc

Format

A data frame with 127 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

NFRLT Nom. Rel. Time from Analyte First Dose

PCTESTCD Pharmacokinetic Test Short Name

PCTEST Pharmacokinetic Test Name

PCORRES Result or Finding in Original Units

PCORRESU Original Units

PCSTRESC Character Result/Finding in Std Format

PCSTRESN Numeric Result/Finding in Standard Units

PCSTRESU Standard Units

PCNAM Vendor Name

PCSPEC Specimen Material Type

PCLLOQ Lower Limit of Quantitation

VISIT Visit Name

VISITNUM Visit Number

PCDTC Date/Time of Specimen Collection

PCDY Actual Study Day of Specimen Collection

PCTPT Planned Time Point Name

PCTPTNUM Planned Time Point Number

TRTSDT Date of First Exposure to Treatment

TRTSDTM Datetime of First Exposure to Treatment

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

ADTM Analysis Datetime

ATMF Analysis Time Imputation Flag

ADT Analysis Date

ATM Analysis Time

ADY Analysis Relative Day

FANLDTM First Datetime of Dose for Analyte

AVISITN Analysis Visit (N)

AVISIT Analysis Visit

ASTDT Analysis Start Date

ASTDTM Analysis Start Datetime

AENDT Analysis End Date

AENDTM Analysis End Datetime

ASTTM Analysis Start Time

AENTM Analysis End Time

AFRLT Act. Rel. Time from Analyte First Dose

ARRLT Actual Rel. Time from Ref. Dose

PCRFTDTM Reference Datetime of Dose for Analyte

FANLDT First Date of Dose for Analyte

FANLTM First Time of Dose for Analyte

PCRFTDT Reference Date of Dose for Analyte

PCRFTTM Reference Time of Dose for Analyte

NRRLT Nominal Rel. Time from Ref. Dose

PARCAT1 Parameter Category 1

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

ATPTREF Analysis Timepoint Reference

ABLFL Baseline Record Flag

BASETYPE Baseline Type

DOSEA Actual Treatment Dose

DOSEP Planned Treatment Dose

DOSEU Treatment Dose Units

FRLTU Rel. Time from First Dose Unit

RRLTU Rel. Time from Ref. Dose Unit

PARAMCD Parameter Code

ALLOQ Analysis Lower Limit of Quantitation

AVAL Analysis Value

AVALU Analysis Value Unit

AVALCAT1 Analysis Value Category 1

SRCDOM Source Data

SRCVAR Source Variable

SRCSEQ Source Sequence Number

DTYPE Derivation Type

MRRLT Modified Rel. Time from Ref. Dose

ANL01FL Analysis Flag 01

ANL02FL Analysis Flag 02

BASE Baseline Value

CHG Change from Baseline

ASEQ Analysis Sequence Number

PARAM Parameter

PARAMN Parameter (N)

HTBL Numeric Result/Finding in Standard Units

HTBLU Standard Units

WTBL Numeric Result/Finding in Standard Units

WTBLU Standard Units

BMIBL Baseline Body Mass Index (kg/m2)

BMIBLU BMI at Baseline (Unit)

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTEDT Date of Last Exposure to Treatment

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Under 30 Group

DTHA30FL Over 30 Group

DTHB30FL Over 30 plus 30 days Group

adpp 43

Source

Generated from admiral package (template ad_adpc.R).

References

None

Examples

data("adpc")

adpp

adpp

Description

Pharmacokinetic Parameters

Usage

adpp

Format

A data frame with 79 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

PPTESTCD Parameter Short Name

PPTEST Parameter Name

PPCAT Parameter Category

PPORRES Result or Finding in Original Units

PPORRESU Original Units

PPSTRESU Standard Units

PPSPEC Specimen Material Type

PPRFDTC Date/Time of Reference Point

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

DTHDT Date of Death

EOSDT End of Study Date

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

ADT Analysis Date

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ADY Analysis Relative Day

PARAMCD Parameter Code

PARCAT1 Parameter Category

AVAL Numeric Result/Finding in Standard Units

AVALU Standard Units

SRCDOM Domain Abbreviation

SRCVAR Source Variable

SRCSEQ Sequence Number

AVISITN Analysis Visit (N)

AVISIT Analysis Visit

VISITNUM Visit Number

VISIT Visit Name

TRTP Planned Treatment

TRTA Actual Treatment

AVALCAT1 Analysis Value Category 1

AVALCA1N Analysis Value Category 1 (N)

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

adpp 45

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_adpp.R).

References

None

Examples

data("adpp")

46 adppk

adppk

adppk

Description

Population Pharmacokinetic

Usage

adppk

Format

A data frame with 61 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

EVID Event ID

NFRLT Nom. Rel. Time from Analyte First Dose

AFRLT Act. Rel. Time from Analyte First Dose

APRLT Actual Rel Time from Previous Dose

NPRLT Nominal Rel Time from Previous Dose

DOSEA Actual Treatment Dose

DOSEP Planned Treatment Dose

PARAMCD Parameter Code

ALLOQ Analysis Lower Limit of Quantitation

CMT Compartment

BLQFL Below Lower Limit of Quant Flag

BLQFN Below Lower Limit of Quant Flag (N)

AMT Actual Amount of Dose Received (unit)

DV Dependent Variable Result

AVAL Analysis Value

DVL Log DV

MDV Missing Dependent Variable Result

AVALU Analysis Value Unit

UDTC Date/Time

II Dosing Interval (unit)

SS Steady State

ASEQ Analysis Sequence Number

PARAM Parameter

adppk 47

PARAMN Parameter (N)

PROJID Project Identifier

PROJIDN Project Identifier (N)

STUDYIDN Study Identifier (N)

SITEID Study Site Identifier

SITEIDN Study Site Identifier (N)

USUBJIDN Unique Subject Identifier (N)

SUBJID Subject Identifier for the Study

SUBJIDN Subject Identifier for the Study (N)

AGE Age

SEX Sex

SEXN Sex (N)

COHORT Cohort Subject Enrolled Into

COHORTC Description of Planned Arm

ROUTE Route of Administration

ROUTEN Route of Administration (N)

RACE Race

RACEN Race (N)

ETHNIC Ethnicity

ETHNICN Ethnicity (N)

FORM Drug Formulation

FORMN Drug Formulation (N)

COUNTRY Country

COUNTRYN Country (N)

COUNTRYL Country Name

HTBL Numeric Result/Finding in Standard Units

WTBL Numeric Result/Finding in Standard Units

ALTBL Numeric Result/Finding in Standard Units

ASTBL Numeric Result/Finding in Standard Units

TBILBL Numeric Result/Finding in Standard Units

CREATBL Numeric Result/Finding in Standard Units

BMIBL Baseline Body Mass Index (kg/m2)

BSABL Numeric Result/Finding in Standard Units

CRCLBL Baseline Creatinine Clearance

EGFRBL Age

RECSEQ Record Sequence

48 adrs_onco

Source

Generated from admiral package (template ad_adppk.R).

References

None

Examples

data("adppk")

adrs_onco

adrs_onco

Description

Tumor Response Analysis

Usage

adrs_onco

Format

A data frame with 79 columns:

DOMAIN Domain Abbreviation

STUDYID Study Identifier

USUBJID Unique Subject Identifier

VISITNUM Visit Number

VISIT Visit Name

RSTESTCD Assessment Short Name

RSTEST Assessment Name

RSORRES Result or Finding in Original Units

RSSTRESC Character Result/Finding in Std Format

RSEVAL Evaluator

RSEVALID Evaluator Identifier

RSACPTFL Accepted Record Flag

RSDTC Date/Time of Assessment

RSSEQ Sequence Number

RANDDT Date of Randomization

PARAMCD Parameter Code

PARAM Parameter

adrs_onco 49

PARCAT1 Parameter Category 1

PARCAT2 Parameter Category 2

PARCAT3 Parameter Category 3

ADT Analysis Date

ADTF Analysis Date Imputation Flag

AVISIT Analysis Visit

AVALC Analysis Value (C)

AVAL Analysis Value

ANL01FL Analysis Flag 01

ANL02FL Analysis Flag 02

ASEQ Analysis Sequence Number

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

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TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

DTHDT Date of Death

DTHDTF Date of Death Imputation Flag

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS Cause of Death

DTHDOM Domain for Date of Death Collection

DTHCGR1 Cause of Death Reason 1

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiralonco package (template ad_adrs.R).

References

None

Examples

data("adrs_onco")

adsl 51

adsl adsl

Description

Subject Level Analysis

Usage

adsl

Format

A data frame with 54 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

52 adsl

TRT01A Actual Treatment for Period 01

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF Date of Death Imputation Flag

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiral package (template ad_adsl.R).

References

None

Examples

data("adsl")

adsl_vaccine 53

adsl_vaccine

adsl_vaccine

Description

Subject Level Analysis for Vaccine

Usage

adsl_vaccine

Format

A data frame with 46 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

INVID Investigator Identifier

INVNAM Investigator Name

BRTHDTC Date/Time of Birth

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country/Region

54 adsl_vaccine

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

TRT02P Planned Treatment for Period 02

TRT01A Actual Treatment for Period 01

TRT02A Actual Treatment for Period 02

TRTSDTM Datetime of First Exposure to Treatment

TRTEDTM Datetime of Last Exposure to Treatment

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

SAFFL Safety Population Flag

PPROTFL Per-Protocol Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

VAX01DT Vaccination Date 01

VAX02DT Vaccination Date 02

AP01SDT Period 01 Start Date

AP01EDT Period 01 End Date

AP02SDT Period 02 Start Date

AP02EDT Period 02 End Date

Source

Generated from admiralvaccine package (template ad_adsl.R).

References

None

Examples

```
data("adsl_vaccine")
```

adtr_onco 55

adtr_onco

adtr_onco

Description

Tumor Results Analysis for Oncology

Usage

adtr_onco

Format

A data frame with 99 columns:

DOMAIN Domain Abbreviation

STUDYID Study Identifier

USUBJID Unique Subject Identifier

TRGRPID Group ID

TRLNKID Link ID

TRTESTCD Tumor/Lesion Assessment Short Name

TRTEST Tumor/Lesion Assessment Test Name

TRORRES Result or Finding in Original Units

TRORRESU Original Units

TRSTRESC Character Result/Finding in Std Format

TRSTRESN Numeric Result/Finding in Standard Units

TRSTRESU Standard Units

VISITNUM Visit Number

VISIT Visit Name

TREVAL Evaluator

TREVALID Evaluator Identifier

TRACPTFL Accepted Record Flag

TRDTC Date/Time of Tumor/Lesion Measurement

TRSEQ Sequence Number

RANDDT Date of Randomization

TULOC Location of the Tumor/Lesion

TULOCGR1 Tumor Site Group 1

LSEXP Lesion IDs Expected

LSASS Lesion IDs Assessed

ADT Analysis Date

36 adtr_onco

ADTF Analysis Date Imputation Flag

ADY Analysis Relative Day

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

PARAMCD Parameter Code

PARAM Parameter

PARCAT1 Parameter Category 1

PARCAT2 Parameter Category 2

PARCAT3 Parameter Category 3

AVAL Analysis Value

ANL01FL Analysis Flag 01

ABLFL Baseline Record Flag

BASE Baseline Value

NADIR NADIR

CHG Change from Baseline

PCHG Percent Change from Baseline

CHGNAD Change from NADIR

PCHGNAD Percent Change from NADIR

PDFL Pharmacodynamic Analysis Set Flag

ANL02FL Analysis Flag 02

ANL03FL Analysis Flag 03

ANL04FL Analysis Flag 04

ASEQ Analysis Sequence Number

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

adtr_onco 57

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRT01P Planned Treatment for Period 01

TRT01A Actual Treatment for Period 01

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

DTHDT Date of Death

DTHDTF Date of Death Imputation Flag

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS Cause of Death

DTHDOM Domain for Date of Death Collection

DTHCGR1 Cause of Death Reason 1

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

58 adtte_onco

Source

Generated from admiralonco package (template ad_adtr.R).

References

None

Examples

```
data("adtr_onco")
```

adtte_onco

 $adtte_onco$

Description

Time to Event Analysis for Oncology

Usage

adtte_onco

Format

A data frame with 20 columns:

STUDYID Study Identifier

USUBJID Unique Subject Identifier

ADT Analysis Date

EVNTDESC Event or Censoring Description

SRCDOM Source Data

SRCVAR Source Variable

SRCSEQ Source Sequence Number

CNSR Censor

CNSDTDSC Censor Date Description

STARTDT Time-to-Event Origin Date for Subject

PARAMCD Parameter Code

PARAM Parameter

AVAL Analysis Value

ASEQ Analysis Sequence Number

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

AGE Age

SEX Sex

advfq_ophtha 59

Source

Generated from admiralonco package (template ad_adtte.R).

References

None

Examples

data("adtte_onco")

advfq_ophtha

 $advfq_ophtha$

Description

Visual Function Questionnaire Analysis

Usage

advfq_ophtha

Format

A data frame with 89 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

QSSEQ Sequence Number

QSTESTCD Question Short Name

QSTEST Question Name

QSCAT Category of Question

QSSCAT Subcategory for Question

QSORRES Finding in Original Units

QSORRESU Original Units

QSSTRESC Character Result/Finding in Std Format

QSSTRESN Numeric Finding in Standard Units

QSSTRESU Standard Units

QSBLFL Baseline Flag

QSDRVFL Derived Flag

VISITNUM Visit Number

VISIT Visit Name

60 advfq_ophtha

VISITDY Planned Study Day of Visit

QSDTC Date/Time of Finding

QSDY Study Day of Finding

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

ADT Analysis Date

ADY Analysis Relative Day

PARAMCD Parameter Code

AVAL Analysis Value

AVALC Analysis Value (C)

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

ONTRTFL On Treatment Record Flag

ABLFL Baseline Record Flag

BASE Baseline Value

CHG Change from Baseline

PCHG Percent Change from Baseline

ANL01FL Analysis Flag 01

ASEQ Analysis Sequence Number

PARAM Parameter

PARCAT1 Parameter Category 1

PARCAT2 Parameter Category 2

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

advfq_ophtha 61

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

DTHB30FL Death Within 30 Days of First Trt Flag

Source

Generated from admiralophtha package (template ad_advfq.R).

62 advs

References

None

Examples

data("advfq_ophtha")

advs

advs

Description

Vital Signs Analysis

Usage

advs

Format

A data frame with 105 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

VSSEQ Sequence Number

VSTESTCD Vital Signs Test Short Name

VSTEST Vital Signs Test Name

VSPOS Vital Signs Position of Subject

VSORRES Result or Finding in Original Units

VSORRESU Original Units

VSSTRESC Character Result/Finding in Std Format

VSSTRESN Numeric Result/Finding in Standard Units

VSSTRESU Standard Units

VSSTAT Completion Status

VSLOC Location of Vital Signs Measurement

VSBLFL Baseline Flag

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

VSDTC Date/Time of Measurements

VSDY Study Day of Vital Signs

advs 63

VSTPT Planned Time Point Name

VSTPTNUM Planned Time Point Number

VSELTM Planned Elapsed Time from Time Point Ref

VSTPTREF Time Point Reference

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

ADT Analysis Date

ADY Analysis Relative Day

PARAMCD Parameter Code

AVAL Analysis Value

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

DTYPE Derivation Type

ONTRTFL On Treatment Record Flag

ANRLO Analysis Normal Range Lower Limit

ANRHI Analysis Normal Range Upper Limit

A1LO Analysis Range 1 Lower Limit

A1HI Analysis Range 1 Upper Limit

ANRIND Analysis Reference Range Indicator

BASETYPE Baseline Type

ABLFL Baseline Record Flag

BASE Baseline Value

BNRIND Baseline Reference Range Indicator

CHG Change from Baseline

PCHG Percent Change from Baseline

ANL01FL Analysis Flag 01

TRTP Planned Treatment

TRTA Actual Treatment

ASEQ Analysis Sequence Number

AVALCAT1 Analysis Value Category 1

AVALCA1N Analysis Value Category 1 (N)

PARAM Parameter

PARAMN Parameter (N)

64 advs

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

SEX Sex

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

SCRFDT Screen Failure Date

EOSDT End of Study Date

EOSSTT End of Study Status

FRVDT Final Retrievel Visit Date

RANDDT Date of Randomization

DTHDT Date of Death

DTHDTF undocumented field

DTHADY Relative Day of Death

LDDTHELD Elapsed Days from Last Dose to Death

DTHCAUS undocumented field

DTHDOM undocumented field

DTHCGR1 undocumented field

LSTALVDT Date Last Known Alive

SAFFL Safety Population Flag

RACEGR1 Pooled Race Group 1

AGEGR1 Pooled Age Group 1

REGION1 Geographic Region 1

LDDTHGR1 Last Dose to Death - Days Elapsed Grp 1

DTH30FL Death Within 30 Days of Last Trt Flag

DTHA30FL Death After 30 Days from Last Trt Flag

Source

Generated from admiral package (template ad_advs.R).

DTHB30FL Death Within 30 Days of First Trt Flag

References

None

Examples

data("advs")

advs_peds

advs_peds

Description

Vital Signs Analysis for Pediatrics

Usage

advs_peds

Format

A data frame with 80 columns:

STUDYID Study Identifier

DOMAIN Domain Abbreviation

USUBJID Unique Subject Identifier

VSSEQ Sequence Number

VSTESTCD Vital Signs Test Short Name

VSTEST Vital Signs Test Name

VSPOS Vital Signs Position of Subject

VSORRES Result or Finding in Original Units

VSORRESU Original Units

VSSTRESC Character Result/Finding in Std Format

VSSTRESN Numeric Result/Finding in Standard Units

VSSTRESU Standard Units

VSSTAT Completion Status

VSLOC Location of Vital Signs Measurement

VSBLFL Baseline Flag

VISITNUM Visit Number

VISIT Visit Name

VISITDY Planned Study Day of Visit

VSDTC Date/Time of Measurements

VSDY Study Day of Vital Signs

VSTPT Planned Time Point Name

VSTPTNUM Planned Time Point Number

VSELTM Planned Elapsed Time from Time Point Ref

VSTPTREF Time Point Reference

VSEVAL Evaluator

EPOCH Epoch

SEX Sex

BRTHDTC Date/Time of Birth (Character)

TRTSDT Date of First Exposure to Treatment

TRTEDT Date of Last Exposure to Treatment

TRT01A Actual Treatment for Period 01

TRT01P Planned Treatment for Period 01

BRTHDT Date/Time of Birth

ADT Analysis Date

ADY Analysis Relative Day

AAGECUR Current Analysis Age (Days)

AAGECURU Current Analysis Age Units

PARAMCD Parameter Code

AVAL Analysis Value

ATPTN Analysis Timepoint (N)

ATPT Analysis Timepoint

AVISIT Analysis Visit

AVISITN Analysis Visit (N)

HGTTMP Temporary Height at Timepoint

HGTTMPU Temporary Height at Timepoint Units

PARAM Parameter

PARAMN Parameter (N)

ABLFL Baseline Record Flag

BASE Baseline Value

CHG Change from Baseline

PCHG Percent Change from Baseline

ONTRTFL On Treatment Record Flag

ANL01FL Analysis Flag 01

SUBJID Subject Identifier for the Study

RFSTDTC Subject Reference Start Date/Time

RFENDTC Subject Reference End Date/Time

RFXSTDTC Date/Time of First Study Treatment

RFXENDTC Date/Time of Last Study Treatment

RFICDTC Date/Time of Informed Consent

RFPENDTC Date/Time of End of Participation

DTHDTC Date/Time of Death

DTHFL Subject Death Flag

SITEID Study Site Identifier

AGE Age

AGEU Age Units

RACE Race

ETHNIC Ethnicity

ARMCD Planned Arm Code

ARM Description of Planned Arm

ACTARMCD Actual Arm Code

ACTARM Description of Actual Arm

COUNTRY Country

DMDTC Date/Time of Collection

DMDY Study Day of Collection

TRTSDTM Datetime of First Exposure to Treatment

TRTSTMF Time of First Exposure Imput. Flag

TRTEDTM Datetime of Last Exposure to Treatment

TRTETMF Time of Last Exposure Imput. Flag

TRTDURD Total Treatment Duration (Days)

ASEQ Analysis Sequence Number

Source

Generated from admiralpeds package (template $ad_advs.R$).

References

None

Examples

```
data("advs_peds")
```

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