# Data Generation for PuMaS

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### 1 Introduction

This introduction to data generation for PuMaS will cover how to specify subjects and populations to use with PuMaS. You will learn how to read data, specify dosage regimens, covariates, observations, and generate populations to use with the software.

### 1.1 Installation

Because PuMaS is still unregistered, you will need to give the Git repository in order to add the package. To do this, use the command <code>ladd https://github.com/UMCTM/PuMaS.jl.</code> Doing it this way, PuMaS and its dependencies will install automatically. If one cannot authenticate for this command (since the repository is currently private!), then first clone the repository how you please, use <code>ladd path/to/PuMaS.jl</code>, then then do <code>lbuild PuMaS.</code> Using the build command will download and install the dependencies.

## 1.2 Subjects

PuMaS unit of observation are subjects.

Each Subject is an individual composed of:

• id: an unique identifier

• cvs: covariates

obs: observations

• evs: events

## 1.3 Population

A Population is a collection of subjects.

### 1.4 NMTRAN data format

The NMTRAN schema uses a single tabular representation to describe a Population

Consider a NMTRAN text file. We can read the data using the CSV package to read it, and the DataFrames package to work with the tabular representation.

Load the packages through using

```
using CSV, DataFrames, PuMaS using DataFrames: head
```

We'll be working with an example NMTRAN data set in PuMaS

```
nmtran_data = example_nmtran_data("data1")
show(nmtran_data)
```

We can read the text file using the CSV and DataFrames packages

```
data = DataFrame(CSV.File(nmtran_data)) # Make a DataFrame from the text file.
head(data) # Show the first six rows
```

	id	time	mdv	evid	$d\mathbf{v}$	amt	sex	wt	$\operatorname{etn}$
1	1	0.0	1	1	0.0	10000.0	1	51.6	1
2	1	0.0	0	0	1.2161	0.0	1	51.6	1
3	1	0.25	0	0	12.565	0.0	1	51.6	1
4	1	0.5	0	0	11.218	0.0	1	51.6	1
5	1	0.75	0	0	17.709	0.0	1	51.6	1
6	1	1.0	0	0	21.925	0.0	1	51.6	1

One can obtain a Population from a NMTRAN table through the process\_nmtran function.

```
using PuMaS
@doc process_nmtran
```

```
process_nmtran(filename, cvs=[], dvs=[:dv])
process nmtran(data, cvs=[], dvs=[:dv])
```

Import NMTRAN-formatted data.

- cvs covariates specified by either names or column numbers
- dvs dependent variables specified by either names or column numbers

Based on the documentation for process\_nmtran it takes a DataFrame, an optional list of covariates, and a list of dependent variables which [:dv] as the default value. In our case, we want to specify sex and wt as covariates.

```
population = process_nmtran(data, [:sex, :wt])
show(population)

Population
   Subjects: 40
   Covariates: sex, wt
   Observables: dv
```

Just like that we have read NMTRAN data and ready it for analysis using PuMaS.

<sup>&</sup>quot;/Users/jbsc/.julia/packages/PuMaS/Zktty/src/../examples/data1.csv"

### 1.5 Interactive

PuMaS offers an alternative to the NMTRAN schema is to generate data

Let us simulate the some values for glucose as out observations

```
using Distributions distribution = Normal(5.5, 1.5) show(distribution)  
Normal{Float64}(\mu=5.5, \sigma=1.5)  
glucose = rand(distribution, 5) # Five draws from the distribution  
obs = DataFrame(time = 0:8:32, # Makes a sequence starting at 0 by 8 until 32 glucose = glucose)  
head(obs)
```

	time	glucose
1	0	6.15155
2	8	5.17305
3	16	4.70792
4	24	7.23497
5	32	4.28062

Next we generate the covariates for our subject

```
cvs = (sex = "male", age = 25)
show(cvs)

(sex = "male", age = 25)
```

Lastly, we can generate a series of dosage events through a DosageRegimen

@doc DosageRegimen

#### DosageRegimen

Lazy representation of a series of Events.

The first argument is an amount. addl specifies additional dosages and ii the interdose interval.

```
evs = DosageRegimen(5, addl = 3, ii = 8u"hr")
head(evs.data)
```

```
time cmt amt evid ii addl rate ss
1 0.0 1 5.0 1 8.0 3 0.0 0
```

This regimen specifies 4 dosage events of five milligrams starting at time = 0 with an interdose interval of 8 hours

We can now fully specify a subject bringing the covariates, observations, and regimen

```
subject = Subject(cvs = cvs, obs = obs, evs = evs)
show(subject)
```

```
Subject
  Events: 4
  Observations: 5
  Covariates:
    sex: male
    age: 25
  Observables: glucose
We can inspect the observations through
for obs in subject.observations
    println(obs)
end
Observation
  time of measurement = 0.0
  compartment = 1
  measurements
    glucose = 6.151553163484906
Observation
  time of measurement = 8.0
  compartment = 1
  measurements
    glucose = 5.1730487195611286
Observation
  time of measurement = 16.0
  compartment = 1
  measurements
    glucose = 4.707920798752679
Observation
  time of measurement = 24.0
  compartment = 1
  measurements
    glucose = 7.2349733134410945
Observation
  time of measurement = 32.0
  compartment = 1
  measurements
    glucose = 4.280616885016562
Likewise for the dosage events
for evs in subject.events
    println(evs)
Dose event
  dose amount = 5.0
  dose time = 0.0
  compartment = 1
  {\tt instantaneous}
  interdose interval = 8.0
  infusion start time = 0.0
Dose event
  dose amount = 5.0
```

```
dose time = 8.0
  compartment = 1
  instantaneous
  interdose interval = 8.0
  infusion start time = 8.0
Dose event
  dose amount = 5.0
  dose time = 16.0
  compartment = 1
  instantaneous
  interdose interval = 8.0
  infusion start time = 16.0
Dose event
 dose amount = 5.0
  dose time = 24.0
 compartment = 1
  instantaneous
  interdose interval = 8.0
  infusion start time = 24.0
```

We can build more complex regimens by passing a collection of arguments

```
regimens = DosageRegimen([0.005u"g", 0.0025u"g"], time = [0, 90u"minute"])
head(regimens.data)
```

	time	$\operatorname{cmt}$	amt	evid	ii	addl	rate	ss
	0.0							
2	1.5	1	2.5	1	0.0	0	0.0	0

Given that we have specified the amount in grams, these are converted to milligrams Likewise, for time which was specified in minutes, these are converted to hours.

This Subject does not have any observations or covariates.

The dosage regimen can be inspected through

```
for evs in subject2.events
    println(evs)
end

Dose event
    dose amount = 5.0
    dose time = 0.0
    compartment = 1
    instantaneous
    interdose interval = 0.0
    infusion start time = 0.0
```

Dose event

```
dose amount = 2.5
dose time = 1.5
compartment = 1
instantaneous
interdose interval = 0.0
infusion start time = 1.5
```

One may construct a Population by passing a collection of subjects

```
@doc Population
```

```
Population(::AbstractVector{<:Subject})</pre>
```

A Population is a set of Subjects. It can be instanced passing a collection or Subject.

There will be six subjects

```
ids = 1:6 # There will be 6 subject 1, 2, 3, 4, 5, 6
```

We will simulate the covariates for these six subjects

choose\_sex\_age will randomly choose a sex and an age between 21-25 years

We can make a list with covariates for six subjects through a list comprehension

```
cvs = [ choose_sex_age() for i in 1:6 ]
show(cvs)

NamedTuple{(:sex, :age),Tuple{String,Int64}}[(sex = "male", age = 21), (sex = "female", age = 21), (sex = "female", age = 22), (sex = "female", age = 21)
```

We can generate a table with the time and observations for the dependent variable for each of the six subjects

```
obs = [ DataFrame(time = 0:4:8, dv = rand(3)) for id in 1:6 ]
head.(obs[1:2])
2-element Array{DataFrame,1}:
3{\times}2 DataFrame
Row time dv
      Int64 Float64
 1
      0
             0.158492
             0.545382
     8
            0.866296
3 \times 2 DataFrame
Row time dv
      Int64 Float64
     Ω
            0.691227
1
            0.360029
2
     4
```

0.914054

21), (sex = "female", age = 24), (sex = "male", age = 25)]

We will assign one regimen to the first three and another to the last three

```
regimen1 = DosageRegimen(15, addl = 3, ii = 4)
regimen2 = DosageRegimen(10, addl = 4, ii = 3)
regimens = vcat(repeat([regimen1], 3),
                repeat([regimen2], 3))
We can now generate the six subjects using a
subjects = Subject[]
for (id, cvs, evs) in zip(ids, cvs, regimens)
    global subjects
    push!(subjects, Subject(id = id, cvs = cvs, evs = evs))
end
show(subjects)
Subject [Subject
 Events: 4
  Observations: 0
  Covariates:
    sex: male
    age: 21
, Subject
  Events: 4
  Observations: 0
  Covariates:
    sex: female
    age: 21
, Subject
 Events: 4
  Observations: 0
  Covariates:
   sex: female
    age: 22
, Subject
 Events: 5
 Observations: 0
  Covariates:
    sex: female
    age: 21
, Subject
 Events: 5
  Observations: 0
  Covariates:
    sex: female
    age: 24
, Subject
 Events: 5
 Observations: 0
  Covariates:
    sex: male
    age: 25
]
Lastly, we can make a population with comprised of these subjects
population = Population(subjects)
show(population)
Population
  Subjects: 6
  Covariates: sex, age
```

```
for subject in population
   println(subject)
end
Subject
  Events: 4
  Observations: 0
  Covariates:
    sex: male
    age: 21
Subject
  Events: 4
  Observations: 0
  Covariates:
    sex: female
   age: 21
Subject
 Events: 4
  Observations: 0
  Covariates:
    sex: female
   age: 22
Subject
  Events: 5
  Observations: 0
  Covariates:
    sex: female
    age: 21
Subject
  Events: 5
  Observations: 0
  Covariates:
    sex: female
    age: 24
Subject
  Events: 5
  Observations: 0
  Covariates:
   sex: male
    age: 25
```