

# pfasbiomon

August 4, 2023

**Type** Package

**Title** PFAS Biomonitoring and In Vitro PODs to generate Margins of Exposure

**Version** 0.1.0

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**Description** The package takes data from PFAS biomonitoring studies (blood, plasma) and in vitro bioactivity data and calculates margins of exposure (MoEs). This supports the manuscript titled ``A Comparison of In Vitro and In Vivo Points of Departure with Human Biomonitoring Levels for Per- and Polyfluoroalkyl Substances (PFAS)''

**License** GPL-3

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 7.2.3

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ACToR.parse	<i>Parse the ACToR files and fill in the remaining missed data</i>
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**Description**

Parse the ACToR files and fill in the remaining missed data

**Usage**

```
ACToR.parse(dataset = "PFAS_3M")
```

**Arguments**

dir                      The directory where the lists are stored

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ACToR.parse.fillin	<i>Parse the ACToR files</i>
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**Description**

Parse the ACToR files

**Usage**

```
ACToR.parse.fillin(  
  datasets = c("Denmark", "Germany", "HealthCanada", "Japan", "Michigan",  
              "Norway_Nordic", "Ohio", "Sweden")  
)
```

**Arguments**

dir                      The directory where the lists are stored

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dnt.check	<i>check the processing for the DNT PODs</i>
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**Description**

check the processing for the DNT PODs

**Usage**

```
dnt.check(to.file = F)
```

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driver	<i>Run all of the analyzes starting with the hand-edited data file</i>
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**Description**

Run all of the analyzes starting with the hand-edited data file

**Usage**

```
driver()
```

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extract_NHANES	<i>Code to format the NHANES data Note that this will only run under R 4.x</i>
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**Description**

Code to format the NHANES data Note that this will only run under R 4.x

**Usage**

```
extract_NHANES(dir = "../data/", dataset = "PFAS_J")
```

**Arguments**

dir	The directory to look for data
dataset	The name of the NHANES dataset

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pfas.for.mdh	<i>Pull the PFAS data fro MDH</i>
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**Description**

Pull the PFAS data fro MDH

**Usage**

```
pfas.for.mdh(to.file = F)
```

---

```
pfasBiomonitoringMerge
```

*Merge the ACToR data with NHANES and put in a common format*

---

**Description**

Merge the ACToR data with NHANES and put in a common format

**Usage**

```
pfasBiomonitoringMerge()
```

---

```
pfasBiomonitoringOldNewMergeForFiltering
```

*Setup the new filtered input file using the latest assays from Doris*

---

**Description**

Setup the new filtered input file using the latest assays from Doris

**Usage**

```
pfasBiomonitoringOldNewMergeForFiltering()
```

---

```
pfasBiomonitoringOutliers
```

*Find outliers*

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**Description**

Find outliers

**Usage**

```
pfasBiomonitoringOutliers()
```

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pfasBloodLevelxChainLength  
*Plotthe blood levels by chain length*

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### Description

Plotthe blood levels by chain length

### Usage

pfasBloodLevelxChainLength(to.file = F)

### Arguments

to.file	If TRUE, write graphs to a file
data.version	Label of folder where input data sits

---

pfasBloodLevelxChainLengthGG  
*Plotthe blood levels by chain length*

---

### Description

Plotthe blood levels by chain length

### Usage

pfasBloodLevelxChainLengthGG(to.file = F)

### Arguments

to.file	If TRUE, write graphs to a file
data.version	Label of folder where input data sits

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pfasChemicalTable      *Generate a table of the chemicals and the data we have for them*

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### Description

Generate a table of the chemicals and the data we have for them

### Usage

pfasChemicalTable()

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pfasCorrectedMOE	<i>Calculate the TK-corrected MOE values</i>
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**Description**

Calculate the TK-corrected MOE values

**Usage**

```
pfasCorrectedMOE()
```

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pfasCorrectedMOE.noTK	<i>Calculate the TK-corrected MOE values</i>
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**Description**

Calculate the TK-corrected MOE values

**Usage**

```
pfasCorrectedMOE.noTK()
```

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pfasHalfLife	<i>Extract the half-life information from Dawson et al. supplemental file</i>
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**Description**

Extract the half-life information from Dawson et al. supplemental file

**Usage**

```
pfasHalfLife(species = "Human", route = "Oral")
```

**Arguments**

species	The species to use
route	The exposre route

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pfasInVittoVsRat	<i>Compare the in vitro PODs vs the Rat POD concentration</i>
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### Description

Compare the in vitro PODs vs the Rat POD concentration

### Usage

```
pfasInVittoVsRat(to.file = F)
```

### Arguments

to.file	If TRUE, write graphs to a file
data.version	Label of folder where input data sits

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pfasMoeBoxplot	<i>Plot the distribution of PODs</i>
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### Description

Plot the distribution of PODs

### Usage

```
pfasMoeBoxplot(to.file = F)
```

### Arguments

dir	The directory where the lists are stored
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pfasPartitionCoefficients	<i>Get the partition coefficients</i>
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### Description

Get the partition coefficients

### Usage

```
pfasPartitionCoefficients(species = "Human")
```

### Arguments

to.file	If TRUE, write graphs to a file
data.version	Label of folder where input data sits

---

pfasPerChemicalMoeBoxplot  
*Build the raw MoE Boxplots*

---

**Description**

Build the raw MoE Boxplots

**Usage**

```
pfasPerChemicalMoeBoxplot(to.file = F)
```

**Arguments**

to.file            If TRUE, write graphs to a file

---

pfasRawMoeBoxplotsGG    *Build the raw MoE Boxplots*

---

**Description**

Build the raw MoE Boxplots

**Usage**

```
pfasRawMoeBoxplotsGG(to.file = F)
```

**Arguments**

to.file            If TRUE, write graphs to a file

---

printCurrentFunction    *Print the name of the current function*

---

**Description**

Print the name of the current function

**Usage**

```
printCurrentFunction(comment.string = NA)
```

**Arguments**

comment.string    An optional string to be printed



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