# Package 'MuPETFlow'

January 7, 2025

Title Multiple Ploidy Estimation Tool for all Species Compatible with Flow Cytometry

Version 0.1.0

**Description** A graphical user interface tool to estimate ploidy from DNA cells stained with fluorescent dyes and analyzed by flow cytometry, following the methodology of Gómez-Muñoz and Fischer (2024) <doi:10.1101/2024.01.24.577056>. Features include multiple file uploading and configuration, peak fluorescence intensity detection, histogram visualizations, peak error curation, ploidy and genome size calculations, and easy results export.

**License** GPL ( $\geq 3$ )

Encoding UTF-8

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RoxygenNote 7.3.2

**Imports** BiocManager, dplyr, DT, flowCore, ggplot2, ggrepel, gridExtra, markdown, shiny, shinythemes, tidyr, zoo

Suggests knitr, rmarkdown

VignetteBuilder knitr

NeedsCompilation no

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**Repository** CRAN

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runMuPETFlow

#### Description

This function launches the Shiny app included in MuPETFlow. Once the application is launched, you can either:

- 1. Load your experimental data.
- 2. Run an in-app example by clicking the 'Example' button.

#### Usage

runMuPETFlow()

#### Details

After launching the app, you can follow the app flow, which is divided into three tabs: **Peaks**, **Regression** and **Summary**. Below is a general description of the options available in each tab:

#### Peaks:

- Select a sample (optional): Allows visual exploration of individual samples if desired.
- Adjust smoothing (optional): Adjusts the histogram curve for noisy samples.
- Adjust window width (optional): Defines the interval where the app will look for peaks.
- Select minimum cell count to call a peak (optional): Useful for samples with a low number of events.
- Select maximum number of peaks to plot (optional): Useful for samples with heterogeneous populations where more peaks are present.

#### **Regression:**

- Select type of analysis: Choose between "Ploidy" or "Genome size" analysis.
- Select number of standards: A minimum of two different standards is required, but more are recommended.
- Select standard samples and values: This is the ploidy or genome size of your standards.

#### **Summary:**

- Results preview: Creates a compiled figure with histograms for all samples.
- **Save plot:** Saves the histograms in either PNG or TIFF format with customizable size and quality. Optionally, you can control the grid layout.
- Save table: Exports the parameters used and the estimated ploidy or genome size as a CSV file.

#### Value

No return value, called for side effects.

### runMuPETFlow

## Examples

```
if (interactive()) {
    # Example: Check that the function exists and runs
    runMuPETFlow()
} else {
    message("This is a Shiny app wrapper. Run interactively to use.")
}
```

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