

Package ‘waydown’

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Type Package

Title Computation of Approximate Potentials for Weakly Non-Gradient Fields

Version 1.1.0

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Description Computation of approximate potentials for both gradient and non gradient fields. It is known from physics that only gradient fields, also known as conservative, have a well defined potential function. Here we present an algorithm, based on the classical Helmholtz decomposition, to obtain an approximate potential function for non gradient fields. More information in Rodríguez-Sánchez (2020) <[doi:10.1371/journal.pcbi.1007788](https://doi.org/10.1371/journal.pcbi.1007788)>.

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Encoding UTF-8

LazyData true

RoxygenNote 7.1.1

VignetteBuilder knitr

Suggests testthat, knitr, rmarkdown, deSolve, dplyr, colorRamps, ggplot2, gridExtra, latticeExtra, bindrcpp

Imports numDeriv, Matrix

NeedsCompilation no

Repository CRAN

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Contents

approxPot1D	2
approxPot2D	3
deltaV	4

Index	6
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`approxPot1D`*Approximate potential in one dimension*

Description

Approximate potential in one dimension

Usage

```
approxPot1D(f, xs, V0 = "auto")
```

Arguments

<code>f</code>	One-dimensional representing the flow (right hand side of differential equation)
<code>xs</code>	Vector of positions to evaluate
<code>V0</code>	(Optional) Value of V at first element of xs. When default, the global minimum is assigned 0

Value

The potential estimated at each point in xs

Author(s)

Pablo Rodríguez-Sánchez (<https://pabrod.github.io>)

References

<https://arxiv.org/abs/1903.05615>

See Also

[approxPot2D](#), [deltaV](#)

Examples

```
# Flow
f = function(x) { sin(x) }

# Sampling points
xs <- seq(0, 2*pi, length.out = 1e3)

# Approximated potential
Vs <- approxPot1D(f, xs)
```

`approxPot2D`*Approximate potential in two dimensions*

Description

Approximate potential in two dimensions

Usage

```
approxPot2D(f, xs, ys, V0 = "auto", mode = "mixed")
```

Arguments

<code>f</code>	Two-dimensional representing the flow (right hand side of differential equation)
<code>xs</code>	Vector xs positions to evaluate
<code>ys</code>	Vector of ys positions to evaluate
<code>V0</code>	(Optional) Value of V at first element of (xs,ys). When default, the global minimum is assigned 0
<code>mode</code>	(Optional) Integration mode. Options are horizontal (default), vertical and mixed

Value

The potential estimated at each point (xs, ys)

Author(s)

Pablo Rodríguez-Sánchez (<https://pabrod.github.io>)

References

<https://arxiv.org/abs/1903.05615>

See Also

[approxPot1D](#), [deltaV](#)

Examples

```
# Flow
f = function(x) {c(-x[1]*(x[1]^2 - 1.1), -x[2]*(x[2]^2 - 1))}

# Sampling points
xs <- seq(-1.5, 1.5, length.out = 10)
ys <- seq(-1.5, 1.5, length.out = 15)

# Approximated potential
Vs <- approxPot2D(f, xs, ys, mode = 'horizontal')
```

deltaV *Approximate potential difference between two points*

Description

Approximate potential difference between two points

Usage

```
deltaV(f, x, x0, normType = "f")
```

Arguments

f	Flow equations (right hand side of differential equation)
x	Position where we want to know the approximate potential
x0	Reference position (center of the Taylor expansion)
normType	(default: 'f') Matrix norm used to compute the error

Value

A list containing the approximate potential difference between x and x0 and the estimated error

Author(s)

Pablo Rodríguez-Sánchez (<https://pabrod.github.io>)

References

<https://arxiv.org/abs/1903.05615>

See Also

[approxPot1D](#), [approxPot2D](#), [norm](#)

Examples

```
# One dimensional flow
f <- function(x) { cos(x) }

# Evaluation points
x0 <- 1
x1 <- 1.02

dV <- deltaV(f, x1, x0)

# Two dimensional flow
f <- function(x) { c(
  -2*x[1]*x[2],
```

```
-x[1]^2 - 1
)}

# Evaluation points
x0 <- matrix(c(1,2), ncol = 1)
x1 <- matrix(c(0.98,2.01), ncol = 1)

dV <- deltaV(f, x1, x0)
```

Index

[approxPot1D](#), [2](#), [3](#), [4](#)

[approxPot2D](#), [2](#), [3](#), [4](#)

[deltaV](#), [2](#), [3](#), [4](#)

[norm](#), [4](#)