

Diffusion-Map-recap

immediate

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A short version to achieve the above is by using the preprocessed version of the dataset provided with this package. `data(guo)` is already preprocessed (using the method first mentioned), has its threshold set to a constant 15 and is ready to use. Since the platform's maximum amplification cycles are 40, that number can be used as upper border of the uncertainty range.

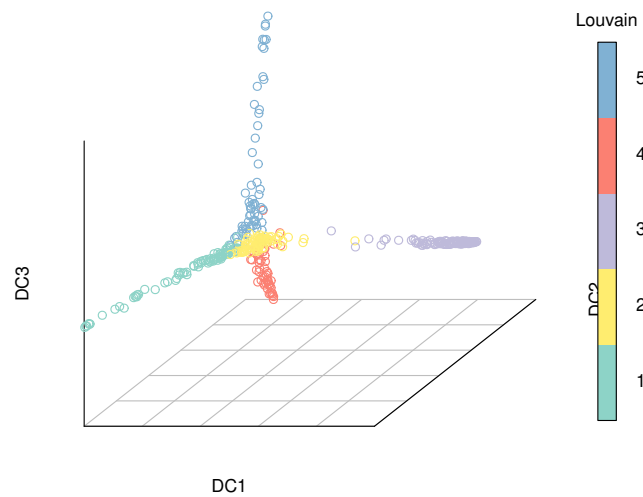
```
In [2]: library(destiny)
        data(guo)
```

It can be used directly for diffusion map creation:

```
In [3]: dm_guo <- DiffusionMap(guo, verbose = FALSE,
                               censor_val = 15, censor_range = c(15, 40))
        dm_guo
```

```
DiffusionMap (20 Diffusion components and 428 observations)
eigenvalues:   num [1:20] 0.954 0.846 0.798 0.776 0.703 ...
eigenvectors:  num [1:428, 1:20] -0.0576 -0.0574 -0.0495 -0.0495 -0.0511 ...
  ..colnames:  chr [1:20] "DC1" "DC2" "DC3" "DC4" ...
optimal_sigma: num [1:428] 7.69 7.63 9.19 6.76 6.35 ...
distance:      chr "euclidean"
```

```
In [4]: plot(dm_guo)
```



using the annotation shows that the approximation worked

```
In [5]: palette(cube_helix(6))
        plot(dm_guo, col_by = 'num_cells',
              legend_main = 'Cell stage')
```

