Alternative CDF environments for 2(or more)-genomes chips

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Introduction

Let's start by loading the package:

> library(altcdfenvs)

The *Plasmodium / Anopheles* is taken as an example:

> library(plasmodiumanophelescdf)

One will adapt easily the code below for other chips.

How to build a CdfEnvAffy object from the cdfenv package

The first step is to wrap the naked environment in the package plasmodium an object:

> planocdf <- wrapCdfEnvAffy(plasmodiumanophelescdf, 712, 712, "plasmodiumanophelescd

Instance of class CdfEnvAffy:

name : plasmodiumanophelescdf
chip-type: plasmodiumanophelescdf

size : 712 x 712

> print(planocdf)

22769 probe set(s) defined.

The numbers 712 and 712 correspond to the dimension of the array. If you do not know these numbers for your chip, the easiest (for the moment) is to read CEL data in an AffyBatch and call the function print on this object.

How to create a CdfEnvAffy that is a subset of the 2-genomes one

If the identifiers starting with 'Pf' correspond to plasmodium, it is an easy job to find them:

However, this is not that simple: the environment created does not contain all the probe set ids from Plasmodium. Unfortunately, one cannot rely on pattern matching on the probe set id to find all the probe set ids associated with Plasmodium. The list of plasmodium ids included in the package can let us build a Plasmodium-only CdfEnvAffy (contributed by Zhining Wang).

Before we eventually save our environment, we may want to give it an explicit name:

```
> plcdf@envName <- "Plasmodium ids only"
> print(plcdf)
```

Instance of class CdfEnvAffy:
 name : Plasmodium ids only
 chip-type: plasmodiumanophelescdf

size : 712 x 712

5407 probe set(s) defined.

Assign the new Cdf data to an AffyBatch

Handling of AffyCdfEnv directly in within an AffyBatch, or AffyBatch-like, structure is being completed... in the meanwhile, the current mecanism for cdfenvs has to be used. If your CEL files were read into an AffyBatch named abatch.

```
envplcdf <- as(plcdf, "environment")
abatch@cdfName <- "plcdf"</pre>
```

From now on, abatch will only consider Cdf information from plcdf. If you want to save this further use, I would recommend to do:

```
save(abatch, plcdf, envplcdf, file="where/to/save.rda")
```