



## Create a class Github repository [↗](#)

Version 2

Forked from [Create "abe487" Github repo](#)

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[dx.doi.org/10.17504/protocols.io.wv2fe8e](https://doi.org/10.17504/protocols.io.wv2fe8e)

[Biosys-analytics](#)

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### ABSTRACT

The github repository will serve as a location for you to turn-in your homework, and allow the instructors to push back code modifications and suggestions.

### EXTERNAL LINK

<http://hurwitzlab.org>

### PROTOCOL STATUS

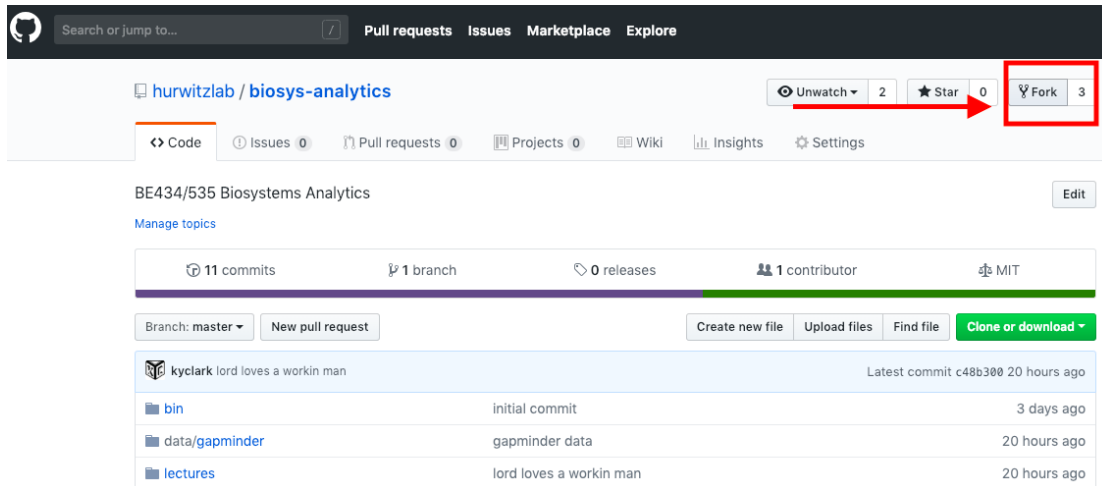
#### **Working**

We use this protocol in our group and it is working

### BEFORE STARTING

Sign up for a free Github account: <http://github.com>

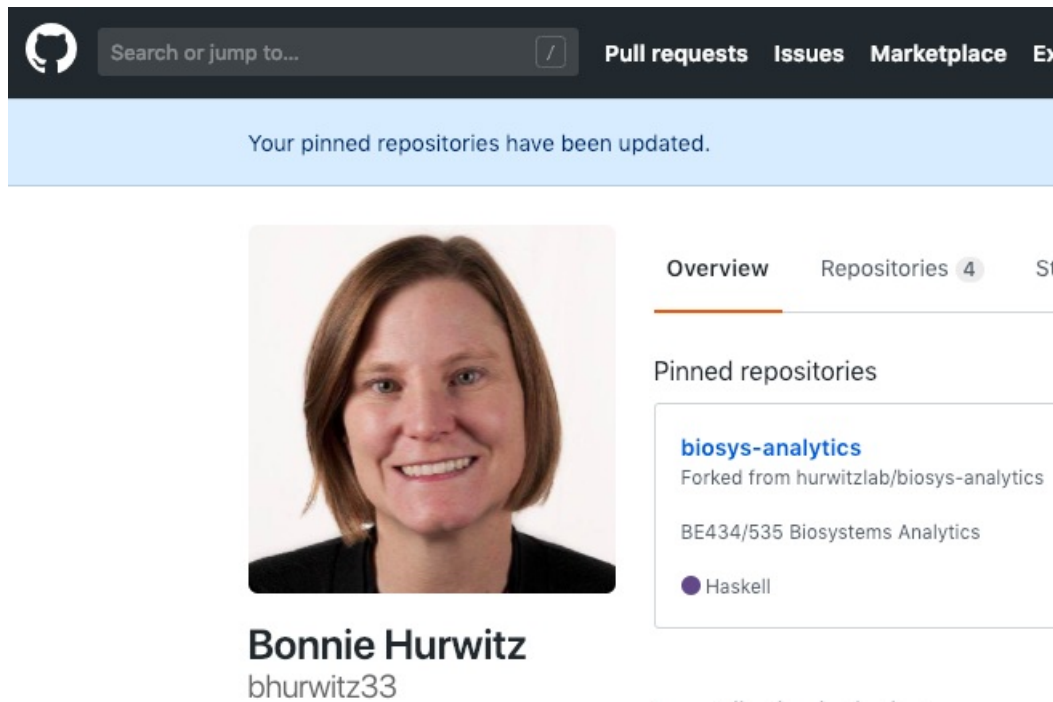
- 1 Login to <http://github.com>
- 2 Go to the class Github repository: <https://github.com/hurwitzlab/biosys-analytics>
- 3 Fork (or "copy") the class repository into your own Github account. This creates a copy of the repository in your own account that you can edit and update. Select your Github account as the place to "fork" the repository.



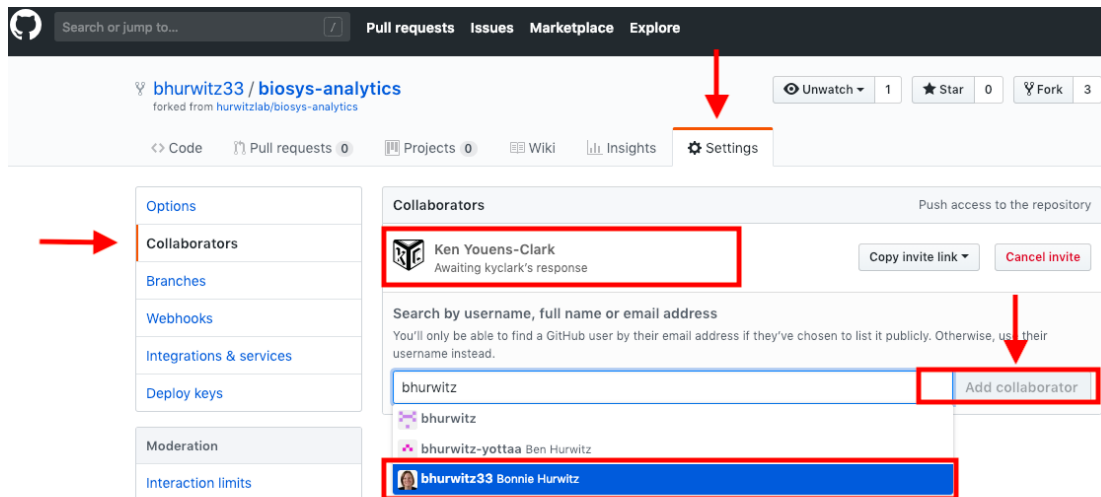
Forking (or copying the biosys-analytics repository). All homework assignments will be in this repository.

- 4 Go to your Github account. You will see a directory called biosys-analytics. This is your copy.

<http://github.com/my-username> . (replace my-username with yours!)



- 5 Click on your "biosys-analytics" repository. Click on the "Settings" tab on the top bar, click on the "Collaborators" tab on the side bar, and search for & add the instructors as collaborators. The instructors Github usernames are: bhurwitz33 and kycklark.



Adding collaborators (the instructors) to YOUR Github repository called biosys-analytics



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