Galaxy servers for command-line users

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We have launchable GVL images with

- Galaxy (and its workflow engine)
- SGE (with a scalable CloudMan cluster)
- CloudBioLinux bioinformatics tools
- Toolshed-installed bioinformatics tools
- Galaxy-installed reference genomes
- ...and indices of reference genomes for all sorts of tools
- NGINX and PostgreSQL

These are useful from outside Galaxy too!
Utilities and setup scripts

https://github.com/claresloggettgvl_commandline_utilities

These utilities will configure the current image; some of what they do will be built into future images.
Features for command-line

What is available?

- non-sudo account(s) for ordinary use
- SGE job submission
- environment modules for toolshed tools
- symlinks to genomes
- public_html
- ipython notebook (password-protected, encrypted)

Also...

- BioBlend! (use on instance, or your local machine)