

## AlphaFold 2.2.0 (2022/3/11)

### Webpage (source code)

<https://github.com/deepmind/alphafold>

### Version

2.2.0 (2022/3/11; alphafold params were also downloaded on 2022/3/11. Other databases are the same as [2.1.1](#) case.)

### Installation

- The same procedure as [2.1.1](#) case was employed.
  - AlphaFold params were newly downloaded. (download\_alphafold\_params.sh)

### Update of script

Wrapper script was updated to support new features.

[run\\_alphafold22x\\_rccs.sh](#)

- default AlphaFold code and database setting were updated.
- Options added
  - -g (re-enabled)
    - This is necessary when you want to perform relaxation of the predicted structures using GPU. (You don't need to specify number of GPUs etc.)
  - -R
    - skip running MSA tools and use precomputed one. NOTE: this will not check if sequence/db/conf have changed. (--use\_precomputed\_msas=True of run\_alphafold.py)
  - -s [seeds per model]
    - number of seeds per model for multimer system. (default: 5) (Number of models (usually 5)) \* (number of seeds; this param) predictions will be performed. MSA time will not be affected by this param. (--num\_multimer\_predictions\_per\_model=(num) of run\_alphafold.py)
  - -S
    - skip relaxation of predicted structure (default: do relaxation) (--run\_relax=False of run\_alphafold.py)

### Notes

- Calculation may be very slow if disk load is high...