Steps for online MSA generation, afv3

- 1. Make folder for project. For example, make a folder names online_MSA_afv3
 - 1. Mkdir online MSA afv3
- 2. Upload fasta file to "online_MSA_afv3"
 - 1. Vi test models afv3.fasta
- 3. Upload the python scripts (pipeline_CFB) to current folder
 - 1. Scp –r Downloads/ pipeline_colabfold_batch_onlineMSAgen vmischley@login.expanse.sdsc.edu:/expanse/lustre/projects/was136/vmischley/online MSA afv3
- 4. Run the script: run_pipeline_CFB_onlineMSAgen.py. This will split the fasta file into individual fasta files with sequences that will in total take less than 48 hours to complete. It will then create the submit script. Input needed is path to fasta file (use full file path), number of recycles and version of AF
 - 1. Python pipeline_colabfold_batch_onlineMSAgen/run_pipeline_CFB_onlineMSAgen.py /expanse/lustre/projects/was136/vmischley/online MSA afv3/test models afv3.fasta 12 alphafold2 multimer v3
- 5. Change into the model folder where all of the submit scripts are
 - 1. Cd models
- 6. Submit all of the submit scripts
 - 1. Sbatch 1_CFB_test_models_MSA_1_submit.slurm
- 7. Check that all of the models completed within 48 hours by looking at the log.txt files
 - 1. Vi models/test_models_afv3_1_models/log.txt
- 8. Run move_af.py to copy only pdb files and .json files into one folder (optional). If using move_af.py you must run an interactive node, or else you run out of memory. Input is path to model folder. Output is folder named combined_models
 - 1. srun --partition=debug --pty --account=was138 --nodes=1 --ntasks-per-node=4 --mem=32G -t 00:30:00 --wait=0 --export=ALL /bin/bash
 - 2. Python move_af.py /expanse/lustre/projects/was136/vmischley/cluster_MSA_afv2/test_models
- 9. After completion, download folder