

AlphaFold 2.3.2

(brief notes about installation)

Python environment

We here assume miniforge is installed in /apl/alphafold/miniforge3 and code of AlphaFold 2.3.2 is in /apl/alphafold/2.3.2.

```
(base) [user@ccfep4 2.3.2]$ conda install -y -c conda-forge openmm=7.5.1 cudatoolkit==11.2.2 cudnn pdbfixer pip python=3.8
(base) [user@ccfep4 2.3.2]$ conda install -y -c bioconda hmmer==3.3.2 hhsuite==3.3.0 kalign2==2.04
(base) [user@ccfep4 2.3.2]$ CONDA_OVERRIDE_CUDA=11.2 conda install jax==0.3.25 jaxlib=0.3.25==*cuda*
(base) [user@ccfep4 2.3.2]$ pip install absl-py==1.0.0 biopython==1.79 chex==0.0.7 dm-haiku==0.0.9 dm-tree==0.1.6 immutabledict==2.0.0 ml-collections==0.1.0 numpy==1.21.6 pandas==1.3.4 scipy==1.7.0 tensorflow-cpu==2.11.0
(base) [user@ccfep4 2.3.2]$ cd /apl/alphafold/miniforge3/lib/python3.8/site-packages/
(base) [user@ccfep4 site-packages]$ patch -p0 < ../../2.3.2/docker/openmm.patch
```

- When we employed CUDA 11.1.1 (according to official one), target version of jax couldn't be installed. CUDA 11.2.2 was tentatively employed.

DB

The latest versions of pdb_mmcif and pdb_seqres were newly installed. For other DBs, those for 2.3.1 installed on Jan 30, 2023 were employed.

AlphaFold

- alphafold/common/stereo_chemical_props.txt was placed as usual.
- patch for alphafold/data/tools/hhblits.py applied (number of threads for hhblits can be modified via env variable)

```
-- hhblits.py.org 2024-02-06 16:27:37.000000000 +0900
+++ hhblits.py 2024-02-07 12:19:54.000000000 +0900
@@ -94,6 +94,14 @@
     self.p = p
     self.z = z

+    n_cpu_env = os.getenv("HHBLITS_NTHREADS")
+    if n_cpu_env:
+        try:
+            n_cpu_env = int(n_cpu_env)
+            self.n_cpu = n_cpu_env
+        except:
+            pass
+
 def query(self, input_fasta_path: str) -> List[Mapping[str, Any]]:
     """Queries the database using HHblits."""
     with utils.tmpdir_manager() as query_tmp_dir:
```

- patch for alphafold/data/tools/jackhmmer.py applied (number of threads for jackhmmer can be modified via env variable)

```
-- jackhmmer.py.org 2024-02-06 16:33:47.000000000 +0900
+++ jackhmmer.py 2024-02-07 12:20:23.000000000 +0900
@@ -87,6 +87,14 @@
     self.get_tblout = get_tblout
     self.streaming_callback = streaming_callback

+    n_cpu_env = os.getenv("JACKHMMER_NTHREADS")
+    if n_cpu_env:
+        try:
+            n_cpu_env = int(n_cpu_env)
+            self.n_cpu = n_cpu_env
+        except:
+            pass
+
 def _query_chunk(self,
```

```
input_fasta_path: str,  
database_path: str,
```

wrapper script

```
#!/bin/bash  
# Description: AlphaFold non-docker version  
# Author: Sanjay Kumar Srikakulam  
#  
#  
# RCCS notes:  
# This script was customized for RCCS by M. Kamiya (IMS).  
# original: https://github.com/kalininalab/alphafold_non_docker  
  
# This script is for AlphaFold 2.3.2!  
# Former AlphaFold versions may not be compatible with this script!  
  
# RCCS default value  
af2root="/apl/alphafold/2.3.2"  
data_dir="/apl/alphafold/databases/20240206"  
  
max_template_date="2024-02-06"  
benchmark=false  
db_preset="full_dbs"  
model_preset="monomer"  
use_gpu=false  
MYOPTS="" # variable for misc options  
  
usage() {  
    echo ""  
    echo "Usage: $0 <OPTIONS>"  
    echo "Required Parameters:"  
    echo "-o <output_dir> Path to a directory that will store the results."  
    echo "-f <fasta_path> Path to a FASTA file containing one sequence"  
    echo ""  
    echo "Optional Parameters:"  
    echo "-a <alphafolddir> Path to alphafold code"  
    echo "-d <data_dir> Path to directory of supporting data"  
    echo "-t <max_template_date> Maximum template release date to consider (ISO-8601 format - i.e. YYYY-MM-DD). Important if folding historical test sets (default: 2021-11-05)"  
    echo "-Q          show also pTM score etc. (alias of -m monomer_ptm)"  
    echo "-b <benchmark> Run multiple JAX model evaluations to obtain a timing that excludes the compilation time, which should be more indicative of the time required for inferencing many proteins (default: 'False')"  
    echo "-g          Enable NVIDIA runtime to run with GPUs"  
    echo "-a <gpu_devices> Comma separated list of devices to pass to 'CUDA_VISIBLE_DEVICES' (default: "")  
    echo "-r <relax_tgt> Choose relax target from all ('all'), most confidential mode ('best'), or skip relaxation ('none')"  
    echo "-R          Skip running MSA tools and use precomputed one. NOTE: this will not check if sequence/db/conf have changed."  
    echo "-s <seeds per model> Number of seeds per model for multimer system. (Number of models (usually 5)) * (number of seeds; this param predictions will be performed. (default: 5)"  
    echo "-p <db_preset> Choose db preset - no ensembling (full_dbs), reduced version of dbs (reduced_dbs) (default: 'full_dbs')"  
    echo "-m <model_preset> Choose model preset - monomer model (monomer), monomer with extra ensembling (monomer_casp14), monomer model with pTM head (monomer_ptm), or multimer model (multimer) (default: 'monomer')"  
    echo ""  
    exit 1  
}  
  
while getopts ":a:d:o:f:t:a:p:s:m:r:bgQR" i; do  
    case "${i}" in  
        a)  
            echo "INFO: set AF2 root to $OPTARG"  
            af2root=$OPTARG  
            ;;  
        d)  
            echo "INFO: set database root to $OPTARG"  
            data_dir=$OPTARG  
            ;;  
    esac  
done
```

```

o)
    output_dir=$OPTARG
;;
f)
    fasta_path=$OPTARG
;;
t)
    max_template_date=$OPTARG
;;
b)
    benchmark=true
;;
g)
    use_gpu=true
;;
Q)
    echo "INFO: set model_preset=monomer_ptm"
    model_preset="monomer_ptm"
;;
a)
    gpu_devices=$OPTARG
;;
p)
    db_preset=$OPTARG
;;
m)
    model_preset=$OPTARG
;;
r)
    MYOPTS="$MYOPTS --models_to_relax=$OPTARG"
;;
s)
    MYOPTS="$MYOPTS --num_multimer_predictions_per_model=$OPTARG"
;;
R)
    MYOPTS="$MYOPTS --use_precomputed_msas=True"
;;
esac
done

# Parse input and set defaults
if [[ "$data_dir" == "" || "$output_dir" == "" || "$fasta_path" == "" ]]; then
    usage
fi

if [[ "$db_preset" != "full_dbs" && "$db_preset" != "reduced_dbs" ]]; then
    echo "Unknown db_preset! Using default ('full_dbs')"
    db_preset="full_dbs"
fi

if [[ "$model_preset" != "monomer" && "$model_preset" != "monomer_casp14" && "$model_preset" != "monomer_ptm" && "$model_preset" != "multimer" ]]; then
    echo "Unknown model_preset! Using default ('monomer')"
    model_preset="monomer"
fi

alphafold_script="$af2root/run_alphafold.py"
if [ ! -f "$alphafold_script" ]; then
    echo "Alphafold python script $alphafold_script does not exist."
    exit 1
fi

if "$use_gpu"; then
    MYOPTS="$MYOPTS --use_gpu_relax=True"
else
    MYOPTS="$MYOPTS --use_gpu_relax=False"

```

```

fi

if [[ "$gpu_devices" ]]; then
  export CUDA_VISIBLE_DEVICES=$gpu_devices
fi

export TF_FORCE_UNIFIED_MEMORY='1'
export XLA_PYTHON_CLIENT_MEM_FRACTION='4.0'

# Binary path (change me if required)
hhblits_binary_path=$(which hhblits)
hhsearch_binary_path=$(which hhsearch)
jackhmmer_binary_path=$(which jackhmmer)
kalign_binary_path=$(which kalign)

MYOPTS="$MYOPTS --hhblits_binary_path=$hhblits_binary_path"
MYOPTS="$MYOPTS --hhsearch_binary_path=$hhsearch_binary_path"
MYOPTS="$MYOPTS --jackhmmer_binary_path=$jackhmmer_binary_path"
MYOPTS="$MYOPTS --kalign_binary_path=$kalign_binary_path"

# uniref30 path
uniref_new=$(find $data_dir -maxdepth 1 -name 'UniRef*')
if [ ! -z "$uniref_new" ]; then
  uniref_name=$(basename $uniref_new)
  uniref30_database_path="$data_dir/$uniref_name/$uniref_name"
  elif [ -d "$data_dir/uniref30" ]; then
    uniref30_database_path="$data_dir/uniref30/UniRef30_2021_03"
fi

# bfd path
if [[ "$db_preset" == "reduced_dbs" ]]; then
  small_bfd_database_path="$data_dir/small_bfd/bfd-first_non_consensus_sequences.fasta"
  MYOPTS="$MYOPTS --small_bfd_database_path=$small_bfd_database_path"
  # uniref30 not necessary
else
  bfd_database_path="$data_dir/bfd/bfd_metaclust_clu_complete_id30_c90_final_seq.sorted_opt"
  MYOPTS="$MYOPTS --bfd_database_path=$bfd_database_path"
  # uniref30 required
  MYOPTS="$MYOPTS --uniref30_database_path=$uniref30_database_path"
fi

# Path and user config (change me if required)
if [ -f $data_dir/mgnify/mgy_clusters_2022_05.fa ]; then
  mgnify_database_path="$data_dir/mgnify/mgy_clusters_2022_05.fa"
else
  mgnify_database_path="$data_dir/mgnify/mgy_clusters.fa"
fi
template_mmcif_dir="$data_dir/pdb_mmcif/mmcif_files"
obsolete_pdbs_path="$data_dir/pdb_mmcif/obsolete.dat"
uniref90_database_path="$data_dir/uniref90/uniref90.fasta"

MYOPTS="$MYOPTS --mgnify_database_path=$mgnify_database_path"
MYOPTS="$MYOPTS --template_mmcif_dir=$template_mmcif_dir"
MYOPTS="$MYOPTS --obsolete_pdbs_path=$obsolete_pdbs_path"
MYOPTS="$MYOPTS --uniref90_database_path=$uniref90_database_path"

# for multimer (pdb70 must not be specified this case)
if [[ "$model_preset" == "multimer" ]]; then
  echo "INFO: appending database paths for multimer model..."
  uniprot_database_path="$data_dir/uniprot/uniprot.fasta"
  MYOPTS="$MYOPTS --uniprot_database_path=$uniprot_database_path"
  pdb_seqres_database_path="$data_dir/pdb_seqres/pdb_seqres.txt"
  MYOPTS="$MYOPTS --pdb_seqres_database_path=$pdb_seqres_database_path"
else
  pdb70_database_path="$data_dir/pdb70/pdb70"
  MYOPTS="$MYOPTS --pdb70_database_path=$pdb70_database_path"

```

```
fi

#echo $MYOPTS

# Run AlphaFold with required parameters
$(python $alphafold_script --data_dir=$data_dir --output_dir=$output_dir --fasta_paths=$fasta_path --max_template_date=$max_template_date --
db_preset=$db_preset --model_preset=$model_preset --benchmark=$benchmark --logtostderr $MYOPTS)
```

Notes

- The CUDA version is different from official one due to the dependency problem.
- Number of threads for hhblits and jackhmmer can be modified by HHBLITS_NTHREADS and JACKHMMER_NTHREADS environment variables, respectively.
 - In some cases, better performance can be achieved by using slightly higher number of threads.
 - (This performance improvement might rely on the performance on lustre filesystem.)
 - Decreasing number of threads is not recommended. Using too many threads is not a wise idea.