Write-Up:

The structures are very different from each other, although the highest 3 scores are very close to each other. This means that sampling has not yet converged. These structures are still quite far from the native structure. The score is still not very low yet, and it is visibility noticeable that the structures are not super similar.

The total energies of the sampled decoys are very far off from the relaxed natives. Although most of the Rosetta score terms are fairly different, the fa_rep, fa_elec weights, omega, and rama_prepro are the most far apart. In regards to the constraints, the atom_pair_constraint's are the most far off from each other.

The energies of the new structures are a little better, but overall, they still look pretty different from the native structures. Even structures with energies in the 100's do not look very similar to the native structures.

For my improvements, I generated protein fragments from the Robetta server (http://robetta.bakerlab.org/fragmentsubmit.jsp) from the fasta sequences. I then made a PyRosetta Fragment mover which, each time the mover is applied, selects a random 3-mer window and inserts the backbone torsion angles from a random matching fragment in the fragment set. Then, I added the constraints and relaxed the pose incrementally as in Part 2. This method results in structures with very low energies. The structures start looking very similar to the native conformations.

Code:

import nglview as nv

```
import pyrosetta
pyrosetta.init('-mute all')

import os

FILES_PATH = "/Users/tgoel/Downloads/Classes/GENOME/GENOME541/hw2/"

def get_pose_from_file(file_name):
    file_n = os.path.join(FILES_PATH, file_name)
    return pyrosetta.pose_from_file(file_n)

def get_pose_from_fasta(file_name):
    file_n = os.path.join(FILES_PATH, file_name)
    with open(file_n) as f:
        sequence = f.read().split("\n")[1]
    return pyrosetta.pose_from_sequence(sequence)
```

```
import numpy as np
def set backbone torsions(pose):
  for i in range(pose.total residue()):
     pose.set phi(i+1, np.random.randint(-180, 180))
     pose.set psi(i+1, np.random.randint(-180, 180))
def set constraints(pose, res1, res2): # set constraints on the target pose between the Cbeta of
residue res1 and residue res2
  id i = pyrosetta.rosetta.core.id.AtomID(pose.residue(res1).atom_index("CB"), res1)
  id j = pyrosetta.rosetta.core.id.AtomID(pose.residue(res2).atom index("CB"), res2)
  ijfunc = pyrosetta.rosetta.core.scoring.constraints.BoundFunc(0.0, 6.0, 1.0, 'cst1');
  cst ij = pyrosetta.rosetta.core.scoring.constraints.AtomPairConstraint(id i, id j, ijfunc)
  pose.add_constraint(cst_ij)
def set constraints with file(pose, file name):
  with open(FILES_PATH + file_name, "r") as f:
     constraints = f.read().strip().split("\n")
  for constraint in constraints:
     res1, res2 = constraint.split()
     set constraints(pose, int(res1), int(res2))
scorefxn =
pyrosetta.rosetta.core.scoring.ScoreFunctionFactory.create_score_function("ref2015_cst.wts")
def relax pose(pose, macrocycles=5):
  fastrelax = pyrosetta.rosetta.protocols.relax.FastRelax(scorefxn, macrocycles)
  fastrelax.apply(pose)
  return scorefxn.score(pose)
def fold protein(protein, show=False):
  display(nv.show file(FILES PATH + protein + ".pdb")) if show else None #
nv.get pose from file(FILES PATH + protein + ".pdb")
  pose = get pose from fasta(FILES PATH + protein + ".fasta")
  set_backbone_torsions(pose)
  display(nv.show rosetta(pose)) if show else None
  set_constraints_with_file(pose, protein + ".contacts")
  score = relax pose(pose)
  display(nv.show_rosetta(pose)) if show else None
```

```
return (pose, score) if not show else score
PROTEIN ID = "6qfi A"
# fold_protein(PROTEIN_ID, show=False)
import time
OUT PATH = "/Users/tgoel/Downloads/"
def fold(prot, fold fn, folds=10, show=False, save=False):
  prot scores = []
  for i in range(folds):
     start time = time.time()
     pose, score = fold_fn(prot)
     prot_scores.append((pose, score))
     end time = time.time()
     print(prot, "fold", i+1, "completed --> score:", f"{score:3f}", "time:",
f"{end time-start time:2f}", "seconds")
  display(nv.show_file(FILES_PATH + prot + ".pdb")) if show else None
  prot scores.sort(key=lambda x: x[1])
  for i, pose_score in enumerate(prot_scores[:3]):
     pose, score = pose score
    if save:
       pose.dump_pdb(OUT_PATH + prot + "_" + str(i) + ".pdb")
    if show:
       display(nv.show_rosetta(pose))
# fold("5gua_A", fold_fn=fold_protein, show=True, save=False)
# fold("5h9h C", fold fn=fold protein, show=True, save=False)
# fold("6ipy A", fold fn=fold protein, show=True, save=False)
# fold("6qfi A", fold fn=fold protein, show=True, save=False)
def relax_native(protein, show=False):
  pose = get pose from file(FILES PATH + protein + ".pdb")
  set constraints with file(pose, protein + ".contacts")
  score = relax_pose(pose)
  print(scorefxn.show(pose))
  display(nv.show_rosetta(pose)) if show else None
```

```
# relax native("5gua A", show=True)
# relax native("5h9h C", show=True)
# relax native("6ipy A", show=True)
# relax native("6qfi A", show=True)
# https://pubmed.ncbi.nlm.nih.gov/11276088
residues data = {
  'G': [(87, 7), (-66, -35), (70, -144), (105, 170), (-171, 177), (-87, 163)],
  'D': [(-140, 165), (-78, 141), (-108, 103), (-97, 5), (-64, -39), (57, 39)],
  'N': [(-140, 165), (-78, 141), (-108, 103), (-97, 5), (-64, -39), (57, 39)],
  'I': [(-132, 153), (-86, 127), (-118, 125), (-91, -9), (-63, -42), (57, 39)],
  'V': [(-132, 153), (-86, 127), (-118, 125), (-91, -9), (-63, -42), (57, 39)],
  'P': [(-64, 145), (-60, -29), (-60, -29), (-77, 161), (-77, 161), (-84, -2)],
  '.': [(-136, 153), (-76, 143), (-112, 119), (-91, -9), (-63, -42), (57, 39)]
}
def set_backbone_torsionsV2(pose):
  angle = np.random.randint(6)
  for i, residue in enumerate(list(pose.residues)):
     amino = residue.name1()
     amino = '.' if amino not in residues data else amino
     phi, psi = residues_data[amino][angle]
     pose.set phi(i+1, phi)
     pose.set_psi(i+1, psi)
def get constraints with file(file name):
  with open(FILES PATH + file name, "r") as f:
     constraints = f.read().strip().split("\n")
     constraints = [x.split() for x in constraints]
     constraints = [(int(x[0]), int(x[1]))] for x in constraints
     distances = [abs(x[0] - x[1]) for x in constraints]
     combined = list(zip(constraints, distances))
     combined.sort(key=lambda x: x[1])
  sorted constraints = [x[0]] for x in combined
  return sorted constraints
```

def fold proteinV2(protein, show=False):

```
display(nv.show file(FILES PATH + protein + ".pdb")) if show else None #
nv.get_pose_from_file(FILES_PATH + protein + ".pdb")
  pose = get_pose_from_fasta(FILES_PATH + protein + ".fasta")
  set backbone torsionsV2(pose)
  display(nv.show rosetta(pose)) if show else None
  constraints = get constraints with file(protein + ".contacts")
  threshold = 5
  for constraint in constraints:
    if abs(constraint[1] - constraint[0]) > threshold:
       relax pose(pose, macrocycles=1)
       threshold += 5
    set_constraints(pose, constraint[0], constraint[1])
  score = relax pose(pose, macrocycles=1)
  display(nv.show_rosetta(pose)) if show else None
  return (pose, score) if not show else score
# fold proteinV2(PROTEIN ID, show=False)
# fold("5gua A", fold fn=fold proteinV2, show=True, save=False)
# fold("5h9h C", fold fn=fold proteinV2, show=True, save=False)
# fold("6ipy_A", fold_fn=fold_proteinV2, show=True, save=False)
# fold("6qfj A", fold fn=fold proteinV2, show=True, save=False)
with open(FILES PATH + PROTEIN ID + ".fasta", "r") as f:
  print(f.read().strip().split("\n")[1])
print(get pose from fasta(FILES PATH + PROTEIN ID + ".fasta").sequence())
print("".join([i.name1() for i in get pose from fasta(FILES PATH + PROTEIN ID +
".fasta").residues]))
# http://bioinf.cs.ucl.ac.uk/psipred/&psipred_uuid=57f6d54a-d58b-11ed-b4fb-00163e100d53
coil_helix_strand = {1: (-140, 130), # helix
            2: (-60, -50)} # strand
# http://bioinf.cs.ucl.ac.uk/psipred/&uuid=3be4ad06-d58f-11ed-b4fb-00163e100d53
2, 2, 2, 2, 2, 2, 2, 2, 2, 2, 0, 0, 2, 2, 2, 2, 2, 2, 2,
      2, 2, 2, 0, 0, 0, 0, 0, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,
      1, 1, 1, 1, 1, 1, 0, 0, 0, 0, 0, 2, 2, 2, 2, 2, 2, 2, 2, 0, 0]
```

```
def set_backbone_torsionsV3(pose, psipreds):
  for i, chs in enumerate(psipreds):
     phi, psi = coil helix strand[chs] if chs != 0 else (np.random.randint(-180, 180),
np.random.randint(-180, 180))
    pose.set phi(i+1, phi)
    pose.set psi(i+1, psi)
def fold proteinV3(protein, show=False):
  display(nv.show_file(FILES_PATH + protein + ".pdb")) if show else None #
nv.get_pose_from_file(FILES_PATH + protein + ".pdb")
  pose = get_pose_from_fasta(FILES_PATH + protein + ".fasta")
  set backbone torsionsV3(pose, psipred)
  display(nv.show_rosetta(pose)) if show else None
  constraints = get constraints with file(protein + ".contacts")
  threshold = 5
  for constraint in constraints:
     if abs(constraint[1] - constraint[0]) > threshold:
       relax_pose(pose, macrocycles=1)
       threshold += 5
     set constraints(pose, constraint[0], constraint[1])
  score = relax_pose(pose, macrocycles=1)
  display(nv.show rosetta(pose)) if show else None
  return (pose, score) if not show else score
# fold_proteinV3(PROTEIN_ID, show=False)
# http://robetta.bakerlab.org/fragmentsubmit.jsp
# http://old.robetta.org/downloads/fragments/79542/
# save http://old.robetta.org/downloads/fragments/79542/aat000 09 05.200 v1 3 to
fragments.txt
fragset = pyrosetta.rosetta.core.fragment.ConstantLengthFragSet(3)
fragset.read_fragment_file(OUT_PATH + "fragments.txt")
def fold_proteinV4(protein, show=False):
  display(nv.show file(FILES PATH + protein + ".pdb")) if show else None #
nv.get_pose_from_file(FILES_PATH + protein + ".pdb")
```

```
pose = get_pose_from_fasta(FILES_PATH + protein + ".fasta")
  movemap = pyrosetta.MoveMap()
  movemap.set bb(True)
  mover_3mer = pyrosetta.rosetta.protocols.simple_moves.ClassicFragmentMover(fragset,
movemap)
  pmm = pyrosetta.PyMOLMover()
  pmm.send_movemap(pose, movemap)
  for i in range(500):
    mover 3mer.apply(pose)
     pmm.apply(pose)
  display(nv.show_rosetta(pose)) if show else None
  constraints = get_constraints_with_file(protein + ".contacts")
  threshold = 5
  for constraint in constraints:
    if abs(constraint[1] - constraint[0]) > threshold:
       relax pose(pose, macrocycles=1)
       threshold += 5
    set constraints(pose, constraint[0], constraint[1])
  score = relax pose(pose, macrocycles=1)
  display(nv.show_rosetta(pose)) if show else None
  return (pose, score) if not show else score
# fold proteinV4(PROTEIN ID, show=True)
```