## **November 8 - November 22 Summary**

I did two things in these two weeks:

- I was able to create a pipeline in Python. This pipeline performs all the (modified) steps in the final report document shared last week, with two caveats:
  - It is currently a (very) stupid pipeline, and will form pairs out of pretty much every noun it finds.
  - It currently assumes we're using the FLAIR model (which needs to be trained), rather than BioBERT. This assumption was made for two reasons:
    - At the time of programming the pipeline, I still hadn't gotten a reply from the authors of BioBERT on the issue, and it was getting too late.
    - FLAIR offers a Python API along with their command-line program, which easily fits into a pipeline (BioBERT does not do this).

The pipeline has been pushed to the GitHub repository, shared with Saad.

- Another person commented on the issue, and I now understand the format of the test.tsv file. Each line should contain a word, followed by a tab, followed by the class label. I found this odd, but on asking, the owner closed the issue as out of scope of that particular issue. I updated the generate-tsv.py file to conform to this standard. For class labels, it simply sets every word to the B class (beginning of a term). The idea is that at this stage we merely want the predictions, and we're less interested in the performance itself.
  - I am currently running this model on the assigned VM. If that does not work, this repository seems to have code for BioFLAIR. We would simply need to replace the test.txt file they use, discarding the middle (part of speech) column, and modify their fine-tuning code to only work with two columns (this is configured in line 4).