README.md 1/22/2022

# Advanced Web Development Midterm

## How to run the application?

Python verson 3 is required to run this project. I tested it using Python version 3.9.6 so in case of any issue, please ensure Python version is up to date. I also used Python vitual environments to isolate the packages that the application relied on.

In order to run the application, please perform the following actions and commands in your OS terminal in the given order:

- Highly recommended but not required: Create a Python virtual environment by running command python -m venv <my\_env> and then activate the virtual environment by running my\_env/Scripts/activate.
- 2. Install the required packages by running pip install -r /path/to/requirements.txt
- 3. Navigate to directory /midterm and then run python manage.py runserver

The app should now be running at default host of localhost with IP address of 127.0.0.1 and Port 8000. To app may now be accessed using a browser or a REST client at the following address: http://127.0.0.1:8000.

In order to run the app at a different address and port, please run the following command:

```
python manage.py runserver localhost:8080
```

Or to only change the port:

```
python manage.py runserver 8080
```

The links in this guide assume port 8000 at localhost has been used.

#### Admin

To login to Django admin, please go to http://127.0.0.1:8000/admin and login using username: admin and password: admin.

# How to import data?

All the migrations have been run and seed data preloaded in the included db.sqlite3 file. However, if anything goes wrong, the data maybe reimported by taking the following steps:

- 1. Delete the file db.sqlite3.
- 2. Run python manage.py makemigrations proteins
- 3. Run python manage.py migrate
- 4. Run cd scripts
- 5. Run python populate-proteins-db.py

#### Models

I imported the provided data into the following models/tables:

README.md 1/22/2022

- 1. Organism
- 2. Pfam
- 3. Domain
- 4. Protein
- 5. ProteinDomainMapping (exists solely to map domains to proteins)

In order to ensure the data is normalized and miniminal redundant data is present in the tables, I separated out the organisms data into their own model: Organism. Since each Protein belongs to an organism, it uses a foreign key taxonomy to reference Organsim model. I noted that length property of Protein is computable doesn't need to be saved in the database table.

I also separated out the Domain and ProteinDomainMapping tables because this way, the mapping table didn't have extra columns which would be tricky to include with the relations. Keeping the mapping table separate from the Domain data made the process really simple with Django rest framework doing the include resolution for us.

I prefered to explicitly declare the primary keys for most of my tables because the IDs in Protein, Organism and Pfam table are already unique.

I felt that domain description should belong to Pfam model and not Domain because it changes with Pfam ID and doesn't change with each Domain instance. However, I kept it with Domain just to conform with the coursework requirements. The data in Domain table is not therefore normalized.

## **REST** endpoints

I implemented the following REST endpoints

```
POST http://127.0.0.1:8000/api/protein/
GET http://127.0.0.1:8000/api/protein/{protein_id}
GET http://127.0.0.1:8000/api/pfam/{pfam_id}
GET http://127.0.0.1:8000/api/proteins/{taxa_id}
GET http://127.0.0.1:8000/api/pfams/{taxa_id}
GET http://127.0.0.1:8000/api/coverage/{protein_id}
```

I used Django Rest Framework to create the above endpoints. Since the endpoints above are simple I didn't want to reinvent the wheel and I was able to achieve the required functionality using the generic API views that Django Rest Framework provides. In the following lines I briefly describe any notable features of my endpoints implementations:

In POST /api/protein/, I create new proteins and accept basic properties such as protein\_id, sequence and taxonomy. I don't create nested objects because I wanted to keep the API RESTful.

In GET /api/protien/{protein\_id}, I use generic RetrieveAPIView and in ProteinRetreiveSerializer, I compute the protein length using len() which I have overriden in Protien model and it simply calculates the number of characters in the protein sequence property. Also I used DomainSerializer to inline domains into protein and also use depth = 1 in Meta which inlines taxonomy.

Again, as I stated above, protein.domains[].description field is not normalized but only because I wanted to follow the coursework requirements. Ideally, this description will be better placed in Pfam table,

README.md 1/22/2022

even though we will end up with two description fields in Pfam table.

### **Tests**

Tests are located in /midterm/proteins/model\_factories.py and /midterm/proteins/tests.py.

To run these tests, navigated to /midterm in the OS terminal, and then run the following command:

python manage.py test