Fine-Tuning and Retrieval-Augmentation: Teaching LLMs new tricks

Self Learning Tutorial (for AI in Healthcare, Spring 2024) Terence Lim

Website (and 3 Python notebooks):

https://t-lim.github.io

Outline

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Overview

Instruction-tuned LLM's have achieved remarkable natural language processing capabilities with promising applications in healthcare, for example, as AI medical assistants to interact "knowledgeably" or "informatively" with patients or clinical staff, hence allowing scarce, skilled, human labor to be deployed elsewhere.

Open-source LLMs, such as Meta's LLama2 (July 2023) and Google's Gemma (February 2024) can **train and run on consumer hardware**. While these may not be as powerful as the closed-source behemoths such as GPT4, they can be more suitable for use in the healthcare domain due to concerns about data privacy and security, especially if they can be customized for the targeted use-case.

Out-of-the-box, open-source LLMs may not have been instructed for the desired downstream tasks. **Fine-tuning** LLMs can help learn such new tasks. LLMs have also been observed to respond with "hallucinations" if pre-trained with stale or non-relevant data, but fine-tuning with large, new datasets can be computationally expensive: enter **retrieval-augmented generation (RAG)** as a feasible solution.

This tutorial demonstrates how to (1) fine-tune the latest open-source LLMs on mere consumer-grade systems, and (2) implement RAG for medical NLP tasks. All the code and experiments were run locally, on an over-2-year-old personal laptop with a consumer-grade RTX3080 16B GPU.

Open-source LLM's

- Following the impact of the closed-source GPT4 from OpenAI in late 2022, many smaller, opensource, pretrained LLM's were (and are being) released. While less powerful (but improving), these possess advantages that are particularly important for healthcare applications:
 - Transparency and flexibility, with full in-house control over sensitive information hence reducing the risk of data leak or unauthorized access
 - Cost savings because no licensing fees are involved
 - ... Benefit from community contributions and experiments
- One popular open-source LLM with a license that allows commercial use is LlaMa-2, released in July 2023 by Meta AI, with pre-trained and chat-tuned models of 7 to 70 billion parameters.
- In February 2024, Google unveiled two open-source Gemma models designed for smaller tasks such as simple chatbots or summarizations. The 7 billion parameter version is claimed to surpass significantly larger models on key benchmarks and can run on a developer's laptop, while the 2 billion parameter version can run directly on mobile devices.
- Huggingface provides a repository for open-source transformer and large-language models, and standard interfaces for fine-tuning and other usages.
 - ~ Community contributions such as quantization and low-rank adaptation "fit" models in consumer hardware
 - ... The Appendix introduces basic HF functions and parameters for performing fine-tuning and inference

Meta Al's Llama model

Meta Al's Llama-2, released in July 2023, comes in three different sizes (7B, 13B, and 70B parameters) in both chat and base versions.

MODEL SIZE (PARAMETERS)	PRETRAINED	FINE-TUNED FOR CHAT USE CASE
7B	Model architecture:	Data collection for helpfulness and safety:
13B	Pretraining Tokens: 2 Trillion	Supervised fine-tuning: Over 100,000
70B	Context Length: 4096	Human Preferences: Over 1,000,000

Llama-2 is a "gated model", to access on Hugging Face:

- Begin by filling an access form on Meta's website. You should receive an approval email fairly quickly
- Hugging Face's model card shows another license to be accepted: the email used on Meta's access form must be the same as that used in your Hugging Face account.
- To download the models from Hugging Face we must authenticate ourselves. Head to Hugging Face Settings > Access Tokens > New token, then create a new and copy this access token to place into your code later. You can also save the token on your machine using the huggingface-cli login terminal login command.

Google's Gemma model

- Google DeepMind's Gemma models, released in February 2024, come in two sizes: 2B and 7B parameters, with each size offering both a base (pretrained) and an instruction-tuned variant
 - ~ 2B: Trained on 2T tokens. With a memory footprint of approximately 1.5GB, it's well-suited for tasks like text classification and straightforward question answering.
 - 7B: Trained on 6T tokens. Suitable for consumer-grade GPUs and TPUs. 5GB memory requirement enables tackling more intricate tasks such as summarization and code generation.
- Training data primarily consisting of English content sourced from Web Docs, mathematical texts, and code.
- Impressive context length of 8,192 tokens.
- Commercial use allowed

Parameters	2B	7B
d_model	2048	3072
Layers	18	28
Feedforward hidden dims	32768	49152
Num heads	8	16
Num KV heads	1	16
Head size	256	256
Vocab size	256128	256128

Table 1 | Key model parameters.

User:	<start_of_turn>user</start_of_turn>
	<pre>Knock knock.<end_of_turn> <start_of_turn>model</start_of_turn></end_of_turn></pre>
Model:	Who's there? <end_of_turn></end_of_turn>
User:	<start_of_turn>user</start_of_turn>
	Gemma. <end_of_turn></end_of_turn>
	<start_of_turn>model</start_of_turn>
Model:	Gemma who? <end_of_turn></end_of_turn>

Table 4 | Example dialogue with user and model control tokens.

Task 1. Medical NLI

NLI in the medical domain

- Natural language inference (NLI) is the task of determining whether a given hypothesis can be inferred from a given premise. This seemingly simple, but challenging problem, formerly known as recognizing textual entailment (RTE) has long been a popular task among NLP researchers.
- Romanov and Shivade (2018) explored the problem of NLI in the clinical domain, and introduced MedNLI - a new, publicly available, expert annotated dataset.
 - As the source of premise sentences, they used the MIMIC-III v1.3 (Johnson et al., 2016) database. With de-identified records of 38,597 patients, it is the largest repository of publicly available clinical data, and contains 2,078,705 clinical notes written by healthcare professionals in English.

The hypothesis sentences were generated by clinicians. They were asked to write three sentences (hypotheses): 1)

A clearly true statement, 2) A clearly false statement, and 3) A statement that might be true or false. This procedure produces three training pairs of sentences for each initial premise with three different labels: entailment,

contradiction, and neutral.

#	Premise	Hypothesis	Label
1	ALT , AST , and lactate were elevated as noted above	patient has abnormal lfts	entailment
2	Chest x-ray showed mild congestive heart failure	The patient complains of cough	neutral
3	During hospitalization, patient became progressively more dyspnic requiring BiPAP and then a NRB	The patient is on room air	contradiction
4	She was not able to speak, but appeared to comprehend well	Patient had aphasia	entailment
5	T1DM: x 7yrs, h/o DKA x 6 attributed to poor medication compliance, last A1c [** 3-23 **]: 13.3 % 2	The patient maintains strict glucose control	contradiction
6	Had an ultimately negative esophagogastroduo- denoscopy and colonoscopy	Patient has no pain	neutral
7	Aorta is mildly tortuous and calcified.	the aorta is normal	contradiction

MedNLI dataset

- Clone https://github.com/jgc128/mednli
- Data files are in jsonl format, containing:
- Training, development and test pairs of premises and hypotheses (with gold labels)

Dataset size	
Training pairs	11232
Development pairs	1395
Test pairs	1422
Average sentence length in tokens	
Premise	20.0
Hypothesis	5.8
Maximum sentence length in token	ıs
Premise	202
Hypothesis	20

Get mednli dataset

```
# helper to read jsonl
import json
def read_jsonl(filename, max_samples=None):
    """helper to read jsonl files as pandas dataframe"""
    lines = []
    with open(filename) as f:
        lines = f.read().splitlines()
    line_dicts = [json.loads(line) for line in lines]
    max_samples = max_samples or len(line_dicts)
    return pd.DataFrame(line_dicts).iloc[:max_samples]
```

```
# read in train, dev and test sets
X_train = read_jsonl('mednli/mli_train_v1.jsonl')
y_train = X_train['gold_label']
X_dev = read_jsonl('mednli/mli_dev_v1.jsonl')
y_dev = X_dev['gold_label']
X_test = read_jsonl('mednli/mli_test_v1.jsonl')
y_test = X_test['gold_label']
```

```
# Show premise, hypothesis and label for dev examples
X_dev.iloc[:6][['sentence1', 'sentence2', 'gold_label']]
```

gold_label	sentence2	sentence1	
entailment	Patient has angina	No history of blood clots or DVTs, has never h	0
contradiction	Patient has had multiple PEs	No history of blood clots or DVTs, has never h	1
neutral	Patient has CAD	No history of blood clots or DVTs, has never h	2
entailment	He has been less alert over the past week	Over the past week PTA he has been more somnol	3
contradiction	Over the past week he has been alert and orie	Over the past week PTA he has been more somnol	4
neutral	He is disorientated and complains of weakness	Over the past week PTA he has been more somnol	5

Fine-tune LLMs for Medical NLI

Initialize model and parameters

Load pre-trained model from HF hub by name, or from training "checkpoints" in local folders. Then set TrainingArguments to be used by the Trainer class to configure for fine-tuning downstream tasks (see Appendix for descriptions of more parameters):

- optim The optimizer to use: adamw_hf, adamw_torch, adamw_torch_fused, adamw_apex_fused, adamw_anyprecision or adafactor.
- lr_scheduler_type The scheduler type to use.
- learning_rate The initial learning rate.
- weight_decay The weight decay to apply (if not zero) to all layers except all bias and LayerNorm weights.
- fp16 Whether to use fp16 16-bit (mixed) precision training instead of 32-bit training.
- prediction_loss_only When performing evaluation and generating predictions, only returns the loss.

```
# pre-trained LLM to use
model id = "meta-llama/Llama-2-7b-chat-hf"
model id = "meta-llama/Llama-2-13b-chat-hf"
model id = "google/gemma-2b-it"
model id = "google/gemma-7b-it"
output dir = os.path.join('models', model id)
# select to load model from HF hub, or previously checkpoint-
from checkpoint = model id
#from checkpoint = output dir
#from checkpoint = os.path.join(output dir, 'checkpoint-474')
# whether to train and/or evaluate
do train = True
do eval = True
# define arguments for trainer
training args = TrainingArguments(
    output dir=output dir,
    optim="paged adamw 32bit",
    lr scheduler type="cosine",
    learning rate=2e-4,
    weight decay=0.001,
    bf16=False,
    max grad norm=0.3,
    max steps=-1.
    warmup ratio=0.03,
```

group by length=False,

gradient accumulation steps=8,

fp16=True, # use mixed precision floats

Load and quantize the model

If a model is "too big" to fit in the computer system available, then basic Hugging Face pipelines may be reconfigured to be allow loading, training, or running inference:

Quantization is a technique to represent the weights and activations with lowprecision data types like 8-bit integer (int8) instead of the usual 32-bit floating point (float32). Reducing the computational and memory costs means operations like matrix multiplication can be performed much faster and even allows to run models on embedded devices.

- The bitsandbytes library is a lightweight Python wrapper around CUDA custom functions, in particular 8-bit optimizers, matrix multiplication (LLM.int8()), and 8 & 4-bit quantization functions.
- Create and include a quantization_config in AutoModel when loading the model.

Get quantized pre-trained model

```
# Load model from HF hub or local folder
compute dtype = getattr(torch, "float16")
if do train: # Load and quantize pre-trained model to start fine-tune
    bnb config = BitsAndBytesConfig(
        load in 4bit=True,
        bnb 4bit use double quant=False,
        bnb 4bit quant type="nf4",
        bnb 4bit compute dtype=compute dtype
    model = AutoModelForCausalLM.from pretrained(
        from checkpoint,
        device map=device, # "auto" may be slower because offload to cpu
        quantization config=bnb config
else:
         # Load previously checkpoint-saved model to continue fine-tune or evaluate
    model = AutoPeftModelForCausalLM.from pretrained(
        from checkpoint,
        torch dtype=compute dtype,
        return dict=False,
        low cpu mem usage=True,
        device map=device, # "auto" may be slower because offload to cpu
# Load tokenizer
tokenizer = AutoTokenizer.from pretrained(model id)
# Fix missing pad token if error
if tokenizer.pad token is None:
    tokenizer.add special tokens({'pad token': tokenizer.eos token})
    model.resize token embeddings(len(tokenizer))
model.config.use cache = False
model.config.pretraining tp = 1
```

Train with QLoRA

PEFT (<u>Parameter-Efficient Fine-Tuning</u>) methods enable efficient adaptation of large pretrained models to downstream applications by only fine-tuning a small number of (extra) model parameters instead of all the model's parameters.

- LoRA (<u>Low-Rank Adaptation</u>) works by attaching extra trainable parameters into a model and decomposing a large weight matrix into two smaller, low-rank matrices (called update matrices). These new matrices can be trained to adapt to the new data while keeping the overall number of changes low.
- QLoRA by Tim Dettmers et al (2023) combines quantization of a model to 4-bits and inserting a small set of trainable low-rank adaptation (LoRA) weights to enable efficient training.

A model is prepared for training with LoRA by creating and including a peft_config with the base model in a SFTTrainer (Supervised Finetuning Trainer) class, with parameters:

- r attention dimension (the "rank")
- lora_alpha scaling factor for the weight matrices, a higher alpha assigns more weight to the LoRA activations
- lora_dropout The dropout probability for Lora layers.

```
# Set config for PEFT
peft_config = LoraConfig(
    lora_alpha=16,
    lora_dropout=0.1,
    r=64,
    bias="none",
    task_type="CAUSAL_LM",
)
```

```
# Set config for SFT Trainer
trainer = SFTTrainer(
    model=model,
    train_dataset=train_data,
    eval_dataset=dev_data,
    peft_config=peft_config,
    dataset_text_field="prompt",
    tokenizer=tokenizer,
    args=training_args,
    packing=False,
    max_seq_length=1024,
)

lap: 100%| 11232/11232 [00:00<00:00</pre>
```

```
if do_train:
    for _ in range(10): # reclaim memory b
        with torch.no_grad():
            torch.cuda.empty_cache()
        gc.collect()
        training_stats = trainer.train()
```

lap: 100%

1395/1395 [00:00<00:00,

Generate model predictions

 Create prompt with instruction + context (premise) + query (hypothesis):

```
# format premise and hypothesis as chat prompt
def as test prompt(ex):
    """prompt for response to test example"""
   return f"""
           Use the following context to determine if the factuality of the
           statement enclosed in square brackets at the end is entailment,
            neutral, or contradiction, and return the answer in 1 word as
            "entailment" or "neutral" or "negative":
            {ex['sentence1']}
            [{ex['sentence2']}]
            Answer:
            """.strip()
# format premise and hypothesis with gold label as training example
def as prompt(ex):
   """training example"""
   return f"{as test prompt(ex)} {ex['gold label']}".strip()
```

```
# reformat all examples as prompts
X_train = pd.DataFrame(X_train.apply(as_prompt, axis=1), columns=["prompt"])
train_data = Dataset.from_pandas(X_train)
X_dev = pd.DataFrame(X_dev.apply(as_prompt, axis=1), columns=["prompt"])
dev_data = Dataset.from_pandas(X_dev)
X_test = pd.DataFrame(X_test.apply(as_test_prompt, axis=1), columns=["prompt"])
test_data = Dataset.from_pandas(X_test)
```

Generate response from model:

```
def predict(X test, model, tokenizer):
    """Generate model predictions on test set"""
   v pred = []
    for i in tqdm(range(len(X test))):
        prompt = X test.iloc[i]["prompt"]
        input ids = tokenizer(prompt, return tensors="pt").to("cuda")
        outputs = model.generate(**input ids,
                                 max new tokens=4, #8
                                 do sample=False, # True
                                 temperature=0.00, # 0.01
        result = tokenizer.decode(outputs[0][len(input ids[0]):])
        answer = result.lower() # result.split("=")[-1].lower()
        if "entailment" in answer:
           v pred.append("entailment")
        elif "contradiction" in answer:
            y pred.append("contradiction")
        else:
            y pred.append("neutral")
    return y pred
```

Results before fine-tuning

the factu

Before the pre-trained models were fine-tuned, they failed to respond correctly to the downstream NLI task:

- Given the same prompt instruction, they did not produce an answer as required.
- Overall accuracy was no better than random responses.

```
Accuracy: 0.333
Classification Report:
               precision
                            recall f1-score
                                               support
contradiction
                    0.00
                              0.00
                                        0.00
                                                   474
   entailment
                    0.00
                              0.00
                                        0.00
                                                   474
                    0.33
                              1.00
      neutral
                                        0.50
                                                   474
     accuracy
                                        0.33
                                                  1422
    macro avo
                    0.11
                              0.33
                                        0.17
                                                  1422
 weighted avg
                    0.11
                              0.33
                                        0.17
                                                  1422
Confusion Matrix:
       0 4741
       0 4741
      0 474]]
Prompt: Use the following context to determine if the factuality of the
            statement enclosed in square brackets at the end is entailment,
            neutral, or contradiction, and return the answer in 1 word as
            "entailment" or "neutral" or "negative":
            He could think of what he wanted to say but was having trouble
getting the words out.
            [ The patient is having trouble speaking. ]
            Answer:
Answer:
```

Few-shot prompting before fine-tuning

Again before fine-tuning, few-shot prompting (where in-context learning is enabled by providing demonstrations in the prompt to steer the model) only slightly improved performance.

 With 1-shot (where a total of 3 training examples – one each of "entailment", "neutral" and "contradiction" – were shown) and 3-shot (total of 9 examples) prompting, Llama-13b test set accuracy rose from 0.582 to 0.6. Results were not much better with additional examples or other models.

```
def zero shot(row):
    """Helper to format the zero-shot question part of a prompt"""
    return f"""
Using the premise, answer in one word whether the hypothesis is entailment, neutral, or contradiction.
Premise: {row['sentencel']}
Hypothesis: {row['sentence2']}
Answer:
[/INST]
""".strip()
def few shot helper(row):
    """Helper to format a one-shot example part of a few-shot prompt"""
    return zero shot(row) + f"""
{row['gold label']}</s><s>
[INST]
def few shot(test, train):
    """Helper to format the few-shot examples and question part of a prompt"""
    return "\n".join([few shot helper(train.iloc[row]) for row in range(len(train))]
                     + [zero shot(test)])
```

```
Evaluation of 1-shot prompts for medNLI task:
Accuracy: 0.582
Classification Report:
                             recall f1-score
               precision
                                                support
contradiction
                    0.76
                               0.72
                                         0.74
                                                     474
   entailment
                    0.52
                               0.89
                                         0.66
                                                     474
     neutral
                    0.39
                               0.14
                                         0.20
                                                     474
                                         0.58
                                                    1422
     accuracy
                                         0.53
                                                    1422
    macro avq
                    0.56
                               0.58
weighted avg
                               0.58
                                         0.53
                                                    1422
                    0.56
```

Evaluation of 3-shot prompts for medNLI task: Accuracy: 0.600 Classification Report: recall f1-score precision support contradiction 0.760.80 0.78 474 entailment 0.520.89 0.66 474 neutral 0.450.11 0.18 474 0.60 1422 accuracy 0.58 0.54 1422 macro avo 0.60 weighted avg 0.58 0.60 0.54 1422

Evaluation of fine-tuned models

After fine-tuning on training set examples, the models are evaluated based on accuracy of test set predictions:

- gemma-7b performed best (accuracy = 0.878), slightly ahead of runner-up Llama-2-13b (0.870)
- both pairs of open-source LLMs gemma (2024) and Llama-2 (2023) – outperformed an early encoder-only transformers-model, electra-base-discriminator (2020) – see Appendix for details about the latter.

gemma-7b-it

Accuracy: 0.878				
Classification R	eport: recision	recall	f1-score	support
contradiction	0.92	0.94	0.93	474
entailment	0.86	0.87	0.86	474
neutral	0.85	0.82	0.84	474

gemma-2b-it

Accuracy: 0.81	7			
Classification	Report: precision	recall	f1-score	support
contradiction	0.87	0.88	0.87	474
entailment	0.81	0.78	0.79	474
neutral	0.77	0.80	0.79	474

Llama-2-13b-chat

Accuracy: 0.87	9			
Classification	Report: precision	recall	f1-score	support
contradiction	0.92	0.94	0.93	474
entailment	0.86	0.85	0.85	474
neutral	0.83	0.82	0.83	474

Llama-2-7b-chat

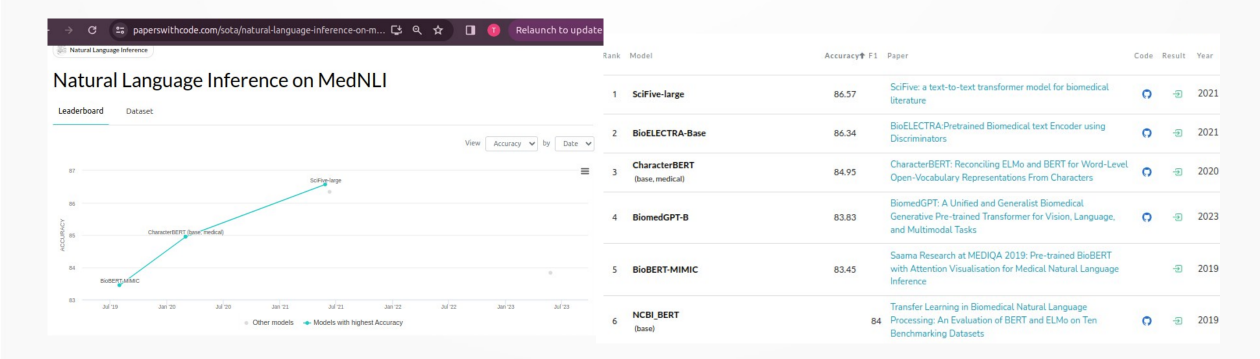
Accuracy: 0.86	1			
Classification	Report: precision	recall	f1-score	support
contradiction	0.91	0.91	0.91	474
entailment	0.83	0.86	0.85	474
neutral	0.84	0.80	0.82	474

electra-base-discriminator

Accuracy: 0.80	9			
Classification	Report: precision	recall	f1-score	support
contradiction	0.87	0.89	0.88	462
entailed	0.78	0.77	0.78	481
neutral	0.77	0.76	0.77	479

Leaderboard

According to the (unofficial) leaderboard in <u>paperswithcode</u>, gemma-7b-it after fine-tuning would have earned the top spot. However, it should be observed from the "Year" column that the webpage (retrieved March 20, 2024) may not (yet) have been updated with most recent research findings.



Task 2: Medical QA

QA in the medical domain

The question answering (QA) NLP task challenges a model to respond to an input question. Closed-domain systems answer the questions from a specific context, while open-domain systems are based on broad unrestricted knowledge. Jin et al (2019) introduced PubMedQA, a biomedical question answering (QA) dataset collected from PubMed abstracts. The task of PubMedQA is to answer research questions with yes/no/maybe using the corresponding abstracts. PubMedQA has 1000 expert-annotated*, 61,200 unlabeled and 211,300 artificially generated QA instances.

* Two annotators (both qualified M.D.candidates) labeled 1000 instances from pre-PQA-U with yes/no/maybe to build PQA-L.

Clone json data files from repo at: https://pubmedqa.github.io

PubMedQA is split into three subsets: labeled, unlabeled and artificially generated. They are denoted as PQA-L(abeled), PQA-U(nlabeled) and PQA-A(rtificial), respectively. We show the architecture of PubMedQA dataset in Fig. 2.

Statistic	PQA-L	PQA-U	PQA-A
Number of QA pairs	1.0k	61.2k	211.3k
Prop. of yes (%)	55.2	-	92.8
Prop. of no (%)	33.8	-	7.2
Prop. of maybe (%)	11.0	-	0.0
Avg. question length	14.4	15.0	16.3
Avg. context length	238.9	237.3	238.0
Avg. long answer length	43.2	45.9	41.0

PubMedQA dataset

• Each example is composed of (1) a question which is either an existing research article title or derived from one, (2) a context which is the corresponding abstract without its conclusion, (3) a long answer which is the conclusion of the abstract, and (4) a yes/no/maybe answer which summarizes the conclusion

Load pubmedga articles and questions

```
pqal = pd.read_json('pubmedqa-master/data/ori_pqal.json', orient='index')
print('CONTEXT:', "\n".join(pqal.iloc[0]['CONTEXTS']))
print('QUESTION:', pqal.iloc[0]['QUESTION'])
print('ANSWER:', pqal.iloc[0]['final_decision'])
pqal['final_decision'].value_counts()
```

CONTEXT: Programmed cell death (PCD) is the regulated death of cells within an organism. The lace plant (Aponogeton madagascariensis) produces perforations in its leaves through PCD. The leaves of the plant consist of a latticework of longitudinal and transverse veins enclosing areoles. PCD occurs in the cells at the center of these areoles and progresses outwards, stopping approximately five cells from the vasculature. The role of mitochondria during PCD has been recognized in animals; however, it has been less studied during PCD in plants.

The following paper elucidates the role of mitochondrial dynamics during developmentally regulated PCD in vivo in A. madagascariensis. A single a reole within a window stage leaf (PCD is occurring) was divided into three areas based on the progression of PCD; cells that will not undergo PCD (NPCD), cells in early stages of PCD (EPCD), and cells in late stages of PCD (LPCD). Window stage leaves were stained with the mitochondrial dynamics were delineated into four categories (M1-M4) based on characteristics including distribution, motility, and membrane potential (ΔΨm). A TUNEL assay showed fragmented nDNA in a gradient over these mitochondrial stages. Chloroplasts and transvacuolar strands were also examined using live cell imaging. The possible importance of mitochondrial permeability transition pore (PTP) formation during PCD was indirectly examined via in vivo cyclosporine A (CsA) treatment. This treatment resulted in lace plant leaves with a sign ificantly lower number of perforations compared to controls, and that displayed mitochondrial dynamics similar to that of non-PCD cells.

QUESTION: Do mitochondria play a role in remodelling lace plant leaves during programmed cell death?

ANSWER: yes

Retrieval-Augmented Generation

RAG

- An LLM may not be aware of specific content, which may be proprietary or recent, absent in its pre-training data.
 - Meta's prompting guide: "Employing RAG is more affordable than fine-tuning, which may be costly and negatively impact the foundational model's capabilities."
- Retrieval Augmented Generation does not require model fine-tuning; it provides an LLM with additional context that is retrieved from relevant data (may be internal or external such as web-scraped) so that it can generate a better-informed response. As illustrated in the diagram:
 - External data is converted into embedding vectors with a separate model and kept in a vector database. Updating the embedding vectors on a regular basis is faster, cheaper, and easier than fine-tuning.
 - Given an unstructured query, retrieve the most relevant (similar) documents (vectors).
 - Pass the original question along with the retrieved context documents to the generation step.
 - With the added context, the same exact LLM model provides a much more relevant and informed answer.

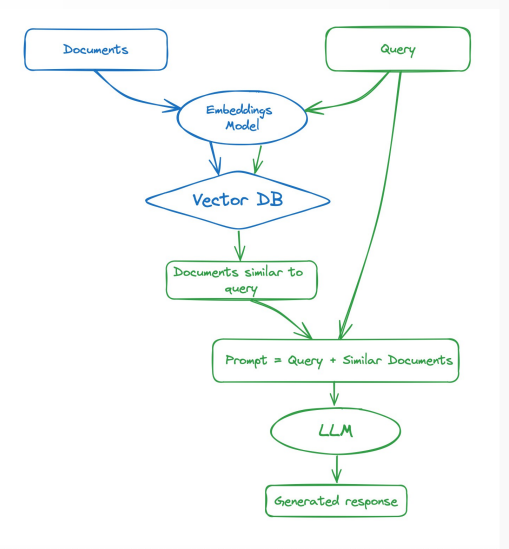


Image: https://huggingface.co/learn/cookbook/rag_zephyr_langchain

Chroma vector database

A Chroma database comprises one or more collections, each of which consists of many individual text documents stored as embedding vectors.

Load chroma client and its default embedding model

```
# Initialize ChromaDB client
db_path = "./database"  # local path to save database
chroma_client = chromadb.PersistentClient(path=db_path)

cprefix = 'pubmedqa_'
for col in chroma_client.list_collections():  # delete old collections
    if col.name.startswith(cprefix):
        chroma_client.delete_collection(col.name)
collections = dict()  # dict for the new collections to create
```

Create database collection with default embedding model

```
# embedding function that embeds any input text
# metadata states how this database should compute similarities (l2 is default)
# default embedding model is all-MiniLM-L6-v2
default_name = "all-MiniLM-L6-v2"
collections[cprefix + default_name] = chroma_client.get_or_create_collection(
    name=f"{cprefix}{default_name}",
    metadata={"hnsw:space": "cosine"}, # l2 is the default
```

Load documents and query

Use chromadb module and its Collection class methods for:

- converting text documents as well as queries to word embedding vectors
- fast retrieval of the top documents that are most similar to the query

See Chroma API reference at https://docs.trychroma.com/api-reference

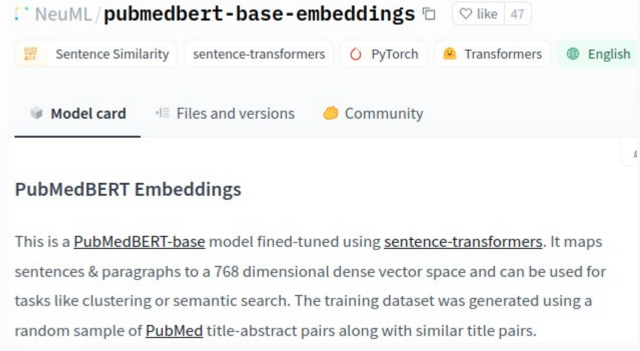
Load data into database

```
: # Each document is a context text string
 documents = pgal['CONTEXTS'].apply(lambda x: "\n".join(x)).to list()
  np.max([len(doc) for doc in documents])
: 2725
                                                                     Query and retrieve an example
: # Each id is the row number
 ids = [str(i) for i in range(len(pgal))]
                                                                     # Each query is a question text string
  #metadatas = [{'row': i} for i in ids]
                                                                     queries = pqal['QUESTION'].to list()
                                                                     gold = 13
: # Load all documents
                                                                     result = collections[cprefix + default name].query(query texts=queries[gold],
 collections[cprefix + default name].add(
                                                                                                                        n results=1)
      documents=documents,
                                                                     print('Gold:', gold, ' Retrieved:', result['ids'])
     ids=ids.
                                                                    Gold: 13
                                                                                Retrieved: [['13']]
```

Word embeddings

- Chroma uses as its default embeddings model: "all-MiniLM-L6-v2"
- Can be any SentenceTransformers model, such as one that is domain-specific:

Can be any sentence-transformer model see https://www.sbert.net/docs/pretrained_models.html # Select other sentence-transformer model by name other_name = 'NeuML/pubmedbert-base-embeddings' other_id = other_name.split('/')[-1] other_model = embedding_functions.SentenceTransformerEmbeddingFunction(model_name=other_name) # Create collection with custom embedding function collections[cprefix + other_id] = chroma_client.get_or_create_collection(name=f"{cprefix}{other_id}", metadata={"hnsw:space": "cosine"}, # l2 is the default embedding_function=other_model,)



Or implement custom embeddings (inherit from EmbeddingFunction class)

Retrieval accuracy

For each question, compare whether the embeddings vector for a query sentence retrieves, as the closest, its true paired PubMed document from the vector database. The results:

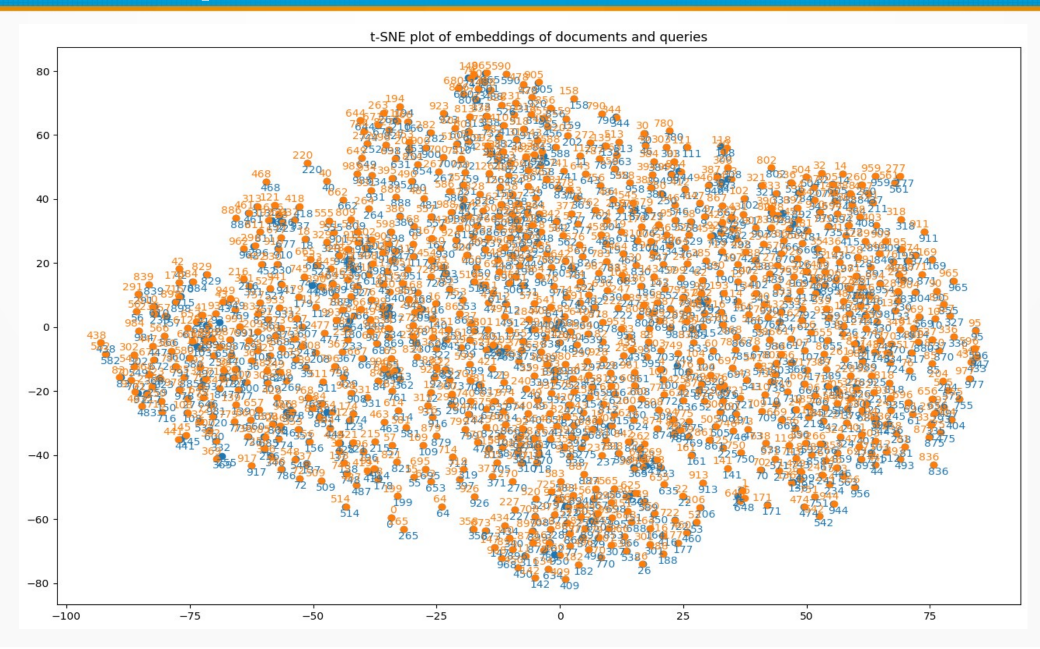
- "pubmedbert-base-embeddings" was more accurate than the default model "all-MiniLM-L6-v2"
- "gte-base" embeddings model was most accurate

Overall, retrieval accuracy scores are high (>96%), and consistent with the t-SNE plot (next slide) where the embedding vectors of true pairs of documents and queries appear clustered

close together.

```
# Check accuracy of id of retrieved document given query
 accuracy = {}
 for name, collection in collections.items():
     y pred = []
     for row, query text in tqdm(enumerate(queries)):
         result = collection.query(query texts=query text, n results=1)
         #print(row, result['ids'][0])
         y pred.append(int(result['ids'][0][0]))
     accuracy[name] = accuracy score(y true=list(range(len(queries))), y pred=y pred)
 pd.DataFrame.from dict(accuracy, orient='index', columns=['Accuracy'])
1000it [00:24, 40.94it/s]
1000it [00:27, 36.22it/s]
1000it [00:27, 36.71it/s]
                                     Accuracy
            pubmedga_all-MiniLM-L6-v2
                                        0.966
 pubmedga_pubmedbert-base-embeddings
                                        0.977
                   pubmedga_gte-base
                                        0.988
```

t-SNE plot



Prompt = Query + Similar Documents

Prompts

Construct prompt with instruction + retrieved document + question (Llama-2 and Gemma worked better with slightly differently worded instructions).

instructions), and ask for answer.

```
def prepare prompt llama(question, context=''):
    """Create prompt for llama model"""
    if context: # with context
        return f"""
Use the following text to return a concise answer to the question at the end
Text:
{context}
Ouestion:
{question}
Answer:
        """.strip()
                  # without context
        return f"""
Ouestion:
{question}
Answer:
        """.strip()
```

```
def prepare prompt gemma(question, context):
    """Create prompt for gemma model"""
    if context: # with context
        return f"""
Use the following text to return an answer to the question.
Return a concise answer to the question in 1 word.
Text:
{context}
Ouestion:
{question}
Answer:
        """.strip()
    else:
                  # without context
        return f"""
Return a concise answer to the question in 1 word.
Ouestion:
{question}
Answer:
        """.strip()
```

Generate model response

To generate response

```
DEFAULT ROLE = "You are a helpful AI assistant."
DEFAULT ROLE = "You are a helpful medical knowledge assistant."
MAX NEW TOKENS = 8
TEMPERATURE = None
def chat(prompt, role, model, tokenizer, max new tokens=MAX NEW TOKENS,
         verbose=False, skip special tokens=True):
   """Prompt the model and return response"""
   # always prepend with a system prompt
   input ids = tokenizer(f"{role}\n{prompt}\n".strip(),
                         return tensors="pt").to("cuda")
   input len = len(input ids[0])
   with torch.no grad():
        outputs = model.generate(**input ids,
                                 max new tokens=max new tokens,
                                 top p=None,
                                 do sample=bool(TEMPERATURE),
                                 temperature=TEMPERATURE,
   if verbose:
        print('Input tokens:', input len,
             ' Output tokens:', len(outputs[0]))
    answer = tokenizer.decode(outputs[0][input len:],
                              skip special tokens=skip special tokens)
   del [input ids, outputs] # reclaim memory
    empty cache()
    return answer.lower()
```

Example of a prompt, and model response

```
# Example of a prompt and response
row = 13
question, context, gold = queries[row], documents[row], y true[row]
print(f"Model={model name}. Example #{row} with gold:", {gold})
role = ""
# without context
prompt = prepare prompt(question, context="", model name=model name)
print('NO CONTEXT:')
print(chat(prompt, role=role, model=model, tokenizer=tokenizer, verbose=True))
print('-----')
# with context
print('WITH CONTEXT:')
prompt = prepare prompt(question, context=context, model name=model name)
print(chat(prompt, role=role, model=model, tokenizer=tokenizer, verbose=True))
print('----')
       Model=google/gemma-7b-it. Example #13 with gold: {'no'}
       NO CONTEXT:
       Input tokens: 40 Output tokens: 43
       yes
       WITH CONTEXT:
       Input tokens: 367 Output tokens: 375
       no
       the text does not describe
```

Evaluate accuracy

Finally, evaluate the accuracy scores of all the models' responses:

- Without any context paragraph, performance was poor Gemma-7b-it (0.551), Llama-7b-chat (0.316). This experiment corresponds to an "open-domain QA" task.
- Given the true context paragraph, Gemma-7b-it (0.679) was slightly more accurate than Llama-7b-chat (0.661). This experiment corresponds to a "closed-domain QA" task.
- When context documents that are most similar to the query string are retrieved with RAG from a vector database: the "gte-base" embeddings model supported the best accuracy (almost equal to with-gold context), followed closely by "pubmedbert-base-embeddings".

	google/gemma-7b-it	meta-llama/Llama-2-7b-chat-hf
pubmedqa_all-MiniLM-L6-v2	0.672	0.652
pubmedqa_pubmedbert-base-embeddings	0.676	0.658
pubmedqa_gte-base	0.678	0.661
with-gold	0.679	0.661
no-context	0.551	0.316

Leaderboard

According to the (unofficial) leaderboard in <u>paperswithcode</u>, Gemma-7b-it with "gte-base" embeddings would earn a Top-20 spot, alongside much larger models.

nk	Model	Accuracy 1	Paper	Code	Result	Yea
1	Meditron-70B (CoT + SC)	81.6	MEDITRON-70B: Scaling Medical Pretraining for Large Language Models	0	Ð	20
2	BioGPT-Large (1.5B)	81.0	BioGPT: Generative Pre-trained Transformer for Biomedical Text Generation and Mining	O	Ð	20
3	Med-PaLM 2 (5-shot)	79.2	Towards Expert-Level Medical Question Answering with Large Language Models		Ð	20
4	Flan-PaLM (540B, Few-shot)	79	Large Language Models Encode Clinical Knowledge		→	20
5	BioGPT (345M)	78.2	BioGPT: Generative Pre-trained Transformer for Biomedical Text Generation and Mining	O	Ð	20
6	Codex 5-shot CoT	78.2	Can large language models reason about medical questions?	0	Ð	20
7	Human Performance (single annotator)	78.0	PubMedQA: A Dataset for Biomedical Research Question Answering	0	Ð	20
8	GAL 120B (zero-shot)	77.6	Galactica: A Large Language Model for Science	0	Ð	20
9	Flan-PaLM (62B, Few-shot)	77.2	Large Language Models Encode Clinical Knowledge		Ð	20
10	MediSwift-XL	76.8	MediSwift: Efficient Sparse Pre-trained Biomedical Language Models		Ð	20
11	BioMedGPT-10B	76.1	BioMedGPT: Open Multimodal Generative Pre-trained Transformer for BioMedicine	0	Ð	20
12	Flan-PaLM (540B.SC)	75.2	Large Language Models Encode Clinical Knowledge		Ð	20

Med-PaLM 2 (ER)	75.0	Towards Expert-Level Medical Question Answering with Large Language Models		Ð	2023
Med-PaLM 2 (CoT+SC)	74.0	Towards Expert-Level Medical Question Answering with Large Language Models		Ð	2023
BLOOM (zero-shot)	73.6	Galactica: A Large Language Model for Science	O	Ð	2022
CoT-T5-11B (1024 Shot)	73.42	The CoT Collection: Improving Zero-shot and Few- shot Learning of Language Models via Chain-of- Thought Fine-Tuning	0	Ð	2023
BioLinkBERT (large)	72.2	LinkBERT: Pretraining Language Models with Document Links	0	Ð	2022
BioLinkBERT (base)	70.2	LinkBERT: Pretraining Language Models with Document Links	C	Ð	2022
OPT (zero-shot)	70.2	Galactica: A Large Language Model for Science	C	Ð	2022
Flan-PaLM (8B, Few-shot)	67.6	Large Language Models Encode Clinical Knowledge		Ð	2022
BioELECTRA uncased	64.2	BioELECTRA:Pretrained Biomedical text Encoder using Discriminators	C	Ð	2021
PaLM (62B, Few-shot)	57.8	Large Language Models Encode Clinical Knowledge		Ð	2022
PubMedBERT uncased	55.84	Domain-Specific Language Model Pretraining for Biomedical Natural Language Processing		Ð	2020
PaLM (540B, Few-shot)	55	Large Language Models Encode Clinical Knowledge		Ð	2022
PaLM (8B. Few-shot)	34	Large Language Models Encode Clinical Knowledge		Ð	2022
	Med-PaLM 2 (CoT+SC) BLOOM (zero-shot) CoT-T5-11B (1024 Shot) BioLinkBERT (large) BioLinkBERT (base) OPT (zero-shot) Flan-PaLM (8B, Few-shot) BioELECTRA uncased PaLM (62B, Few-shot) PubMedBERT uncased	(ER) 75.0 Med-PaLM 2 (CoT + SC) 74.0 BLOOM (zero-shot) 73.6 CoT-T5-11B (1024 Shot) 73.42 BioLinkBERT (large) 72.2 BioLinkBERT (base) 70.2 OPT (zero-shot) 70.2 Flan-PaLM (8B, Few-shot) 67.6 BioELECTRA uncased 64.2 PaLM (62B, Few-shot) 57.8 PubMedBERT uncased 55.84 PaLM (540B, Few-shot) 55 PaLM (540B, Few-shot) 34	Med-PaLM 2	Med-PaLM 2	(ER) 75.0 with Large Language Models Med-PaLM 2 (coT+SC) 74.0 Towards Expert-Level Medical Question Answering with Large Language Models (coT+SC) 74.0 With Large Language Models BLOOM (zero-shot) 73.6 Galactica: A Large Language Model for Science CoT-T5-11B (1024 Shot) 73.42 The CoT Collection: Improving Zero-shot and Few-shot Learning of Language Models via Chain-of-Thought Fine-Tuning BioLinkBERT (large) 72.2 LinkBERT: Pretraining Language Models with Document Links •• BioLinkBERT (base) 70.2 LinkBERT: Pretraining Language Models with Document Links •• OPT (zero-shot) 70.2 Galactica: A Large Language Model for Science •• Flan-PaLM (8B, Few-shot) 67.6 Large Language Models Encode Clinical Knowledge •• BioELECTRA uncased 64.2 BioELECTRA Pretrained Biomedical text Encoder using Discriminators •• Biomedical Natural Language Model Pretraining for Biomedical Natural Language Processing •• PubMedBERT uncased 55 Large Language Models Encode Clinical Knowledge •• PalM (5408, Few-shot) 55 Large Language Models Encode Clinical Knowledge ••

Conclusion

Conclusion

This tutorial explored the use of open-source LLMs for healthcare NLP tasks, specifically for PubMed question answering and Medical natural language inference, and demonstrated how to overcome the following challenges:

- 1) If the model has not been instructed for the required downstream task, then fine-tuning can enhance its performance.
- 2) When the model was pre-trained with stale or non-relevant data for the task, then retrieval-augmented generation (RAG) can be an efficient solution.
- 3)On consumer devices like laptops, techniques such as quantization and low-rank adaption can be used to load, train and run the models.

We showed how to apply these methods using the popular Llama-2 (Meta AI, 2023) as well as the recently-released Gemma (Google, February 2024) models – the 2B version of the latter has a footprint small enough to potentially fit in mobile devices. Such approaches towards customizing open-source LLM's are particularly important for conversational and textual applications in the healthcare domain, where privacy, security, and mobility are critical user concerns.

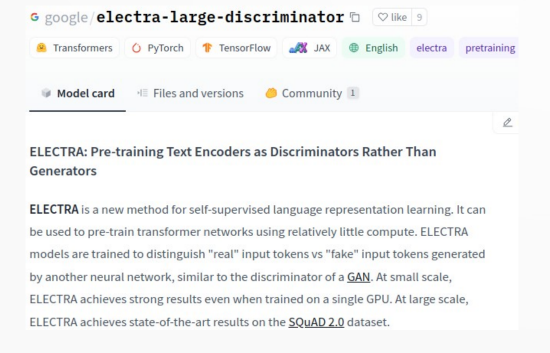
Appendix

Introduction to Hugging Face Transformers

Hugging Face is an online platform with over 350k models and 75k datasets, all open source and publicly available:

- Download pre-trained models from its Model Hub
- Import its Transformers library for training or inference

Example: ELECTRA (2020) is an early encoder-only transformer model, suitable for simple tasks that require understanding input such as sentence classification.



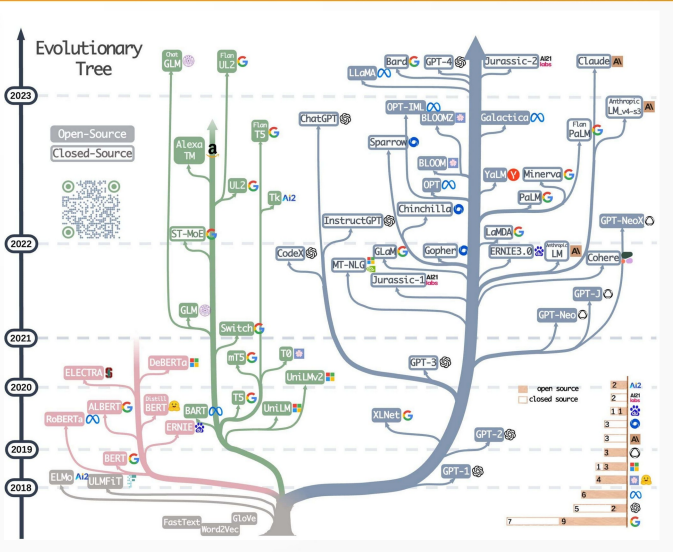


Image: https://www.baeldung.com/cs/bert-vs-gpt-3-architecture

Datasets and AutoTokenizer

- Datasets is a Hugging Face module for accessing and sharing datasets, and for data preprocessing (e.g. tokenizing, resampling and transformations).
- The AutoTokenizer.from_pretrained() class method will be instantiated with one of the tokenizer classes for preparing inputs for a model. The library contains tokenizers for all the models.

```
nli labels = {'entailment': 0, 'neutral': 1, 'contradiction': 2}
if args.dataset == 'snli':
    dataset = datasets.load dataset('snli')
else:
    dataset = datasets.DatasetDict()
    def prepare label(ex):
       """To pre-process mednli dataset examples"""
        lab = ex['label']
       ex['label'] = nli labels[lab] if lab in nli labels else -1
        return ex
    for split in args.dataset.keys():
       print('Loading', split, ':', args.dataset[split], '...')
        # By default, "json" loader places all examples in the "train" split
        dataset[split] = datasets\
            .load dataset('json', data files=args.dataset[split])['train']\
            .rename columns({'gold label': 'label',
                             'sentencel': 'premise',
                             'sentence2': 'hypothesis'})\
            .select columns(['label', 'premise', 'hypothesis'])\
            .map(prepare label)
```

```
# Prepare train and dev set splits
if training args.do train:
    train dataset = dataset['train']
    if args.max train samples > 0:
        train dataset = train dataset.select(range(args.max train samples))
    train dataset tokenized = \
        train dataset.map(prepare dataset,
                          batched=True.
                          remove columns=train dataset.column names)
    dev dataset = dataset['dev']
    if args.max dev samples > 0:
        dev dataset = dev dataset.select(range(args.max dev samples))
    dev dataset tokenized = \
        dev dataset.map(prepare dataset,
                        batched=True,
                        remove columns=dev dataset.column names)
```

<u>https://huggingface.co/docs/datasets/en/index</u>
<u>https://huggingface.co/docs/transformers/main_classes/tokenizer</u>

Auto Classes and TrainingArguments

Set TrainingArguments to configure the Trainer class:

- num_train_epochs Total number of training epochs to perform
- per_device_train_batch try to make this as large as you can without getting CUDA out-of-memory errors.
- eval_steps Interval between two evaluations.
 Should be an integer as number of update steps, or a float in range [0,1) as ratio of total training steps.
- save_steps Number of update steps before two checkpoint saves.
- logging_steps Number of steps between two logs
- report_to Platform to report the results and logs to, such as "azure_ml", "clearml", "codecarbon", "comet_ml", "dagshub", "dvclive", "flyte", "mlflow", "neptune", "tensorboard", and "wandb".

AutoModel subclasses automatically retrieve the relevant architecture with the right model fine-tuning head.

```
# Parameters for the Trainer
training_args = TrainingArguments(
    output_dir=args.output_dir,
    do_train=True,
    do_eval=True,
    num_train_epochs=8.0,
    per_device_train_batch_size=32,
    evaluation_strategy='steps',
    save_steps= 0.2,  # checkpoint interval
    logging_steps = 0.1,  # logging interval
    eval_steps = 0.1,  # evaluation interval
    report_to="tensorboard",
)
```

Load the model and tokenizer

Select to load model from HF hub, or previously checkpoint-saved folder This should either be a HuggingFace model ID (see https://huggingface.co/models) or a path to a saved model checkpoint (a folder containing config.json and model.save_tensors)

```
# Select the model architecture
model_class = AutoModelForSequenceClassification

# Where to load model from
from_checkpoint = args.model_id  # load pre-trained from HF hub
#from_checkpoint = args.output_dir  # load from local folder
#from_checkpoint = os.path.join(args.output_dir, "checkpoint-6740")
```

```
In [ ]:
model = model_class.from_pretrained(from_checkpoint, num_labels=3)
```

Trainer class

Trainer is a complete training and evaluation loop implemented in the Transformers library. You only need to pass it the necessary pieces for training: model, tokenizer, dataset, evaluation function, and hyperparameters

If you want to use custom evaluation metrics, provide your own compute_metrics() function

```
# If you want to use custom metrics, define your own "compute metrics" function.
def compute metrics(eval prediction: EvalPrediction):
    """computes sentence-classification accuracy"""
    return {'accuracy': (np.argmax(eval prediction.predictions, axis=1) ==
                         eval prediction.label ids).astype(np.float32).mean().item()}
# If you want to change how predictions are computed, you should
# subclass Trainer and override the "prediction step" method
# (see https://huggingface.co/transformers/ modules/transformers/trainer.html#Trainer.prediction step).
# If you do this your custom prediction step should probably start by
# calling super().prediction step and modifying the values that it returns
trainer = Trainer(model=model,
                  args=training args,
                  train dataset=train dataset tokenized,
                  eval dataset=dev dataset tokenized,
                  tokenizer=tokenizer,
                  compute metrics=compute metrics)
if training args.do train:
    print('Training the model...')
    trainer.train()
    trainer.save model(args.output dir)
```

Evaluation with scikit-learn

To evaluate classification tasks with scikit-learn:

accuracy score()

- Accuracy: fraction of correct predictions
- Precision: number of true positives divided by the number of true positives plus the number of false positives
- Recall: number of true positives divided by the number of true positives plus the number of false negatives
- F1-score: harmonic mean of precision and recall

classification report()

- Macro average: averages the unweighted mean per label
- Weighted average: averages the support-weighted mean per label

confusion matrix()

• C_{ij} is equal to the number of observations known to be in group i and predicted to be in group j

The electra-base-discriminator model, after fine-tuning with training examples, achieved MedNLI test set accuracy of 0.809

```
# Calculate accuracy
accuracy = accuracy_score(y_true=y_true, y_pred=y_pred)
print(f'Accuracy: {accuracy:.3f}')

class_report = classification_report(y_true=y_true, y_pred=y_pred)
print('\nClassification Report:')
print(class_report)

# Generate confusion matrix
conf_matrix = confusion_matrix(y_true=y_true, y_pred=y_pred)
print('\nConfusion Matrix:')
print(conf_matrix)
```

```
Accuracy: 0.809
Classification Report:
                             recall f1-score
               precision
                                                support
contradiction
                    0.87
                              0.89
                                         0.88
                                                    462
     entailed
                    0.78
                              0.77
                                         0.78
                                                    481
      neutral
                    0.77
                              0.76
                                         0.77
                                                    479
                                         0.81
                                                   1422
     accuracy
    macro avq
                    0.81
                              0.81
                                         0.81
                                                   1422
 weighted avg
                    0.81
                              0.81
                                         0.81
                                                   1422
Confusion Matrix:
[[412 23 27]
 [ 28 372 81]
 [ 34 79 366]]
```

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