Master in Artificial Intelligence

Machine Learning NFRC

Sequence tagging: the B-I-O approach

General

Structure Detailed

Structure Core task

Goals & Deliverables

Advanced Human Language Technologies





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 - Learner
 - Classifier
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Machine Learning NERC

Sequence tagging: the B-I-O approach

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Session 2 - NERC using machine learning

Assignment

Machine Learning NERC

Sequence tagging: the B-I-O approach

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Goals & Deliverables The main program parses all XML files in the folder given as argument and recognizes and classifies drug names, calling a sequence-tagging machine learning algorithm.

```
$ python3 ./ml-NER.py data/Devel/ result.out
$ more result.out
DDI-DrugBank.d278.s0|0-9|Enoxaparin|drug
DDI-DrugBank.d278.s0|93-108|pharmacokinetics|group
DDI-DrugBank.d278.s0|113-124|eptifibatide|drug
DDI-MedLine.d88.s0|15-30|chlordiazepoxide|drug
DDI-MedLine.d88.s0|33-43|amphetamine|drug
DDI-MedLine.d88.s0|49-55|cocaine|drug
DDI-MedLine.d88.s1|82-95|benzodiazepine|drug
...
```

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Sequence tagging: the B-I-O approach

- We want to detect subsequences in a sentence (e.g. drug names).
- To approach this as a ML classification problem, we will classify each token.
- The classes predicted by the classifier must allow the later reconstruction of the target subsequences.
- B-I-O schema: mark each token as Begin of a subsequence,
 Inside a subsequence, or Outside any subsequence.
- If we not only want to recognize the subsequences, but also classify them, we use more informative B-I-O classes:

 Ascorbic acid , aspirin , and the common cold .

 B-drug I-drug 0 B-brand 0 0 0 0 0 0 0
- Different variations of this schernal xist: BIO, BIOS, BIOES (aka BILOU)

Machine Learning NERC

Sequence tagging: the B-I-O approach

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Machine Learning NERC

Sequence tagging: the B-I-O approach

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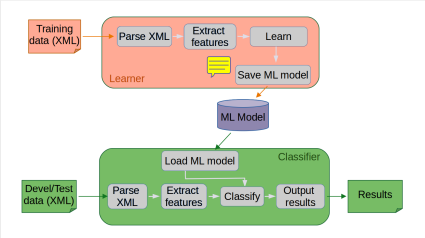
Machine Learning NERC

Sequence tagging: the B-I-O approach

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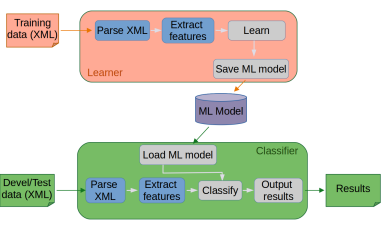
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Sequence tagging: the B-I-O approach

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Extracting features is a costly operation, which we do not want to repeat for every possible experiment or algorithm parametrization.

Machine Learning NERC

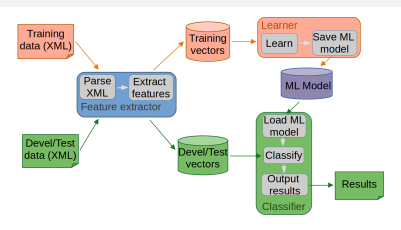
Sequence tagging: the B-I-O approach

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Feature extraction process is performed once, out of learning or predicting processes.

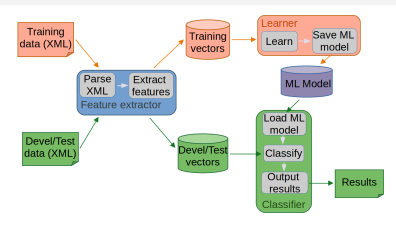
Machine Learning NERC

Sequence tagging: the B-I-O approach

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Feature extraction process is performed once, out of learning or predicting processes.

Thus, we need to write not a single program, but three different components: feature extractor, learner, and classifier.

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Machine Learning NERC

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Feature Extractor

The feature extractor:

- Independent program, separated from learner and classifier
- Receives as argument the directory with the XML files to encode.
- Prints the feature vectors to stdout



```
$ python3 ./feature-extractor.py data/devel > devel.feat
$ more devel.feat
DDI-DrugBank.d658.s0 When 0 3 0 form=When formlower=when suf3=hen
suf4=When isTitle BoS formNext=administered
formlowerNext=administered suf3Next=red suf4Next=ered
DDI-DrugBank d658 s0 administered 5 16 0 form=administered
```

formlowerNext=administered suf3Next=red suf4Next=ered
DDI-DrugBank.d658.s0 administered 5 16 0 form=administered
formlower=administered suf3=red suf4=ered formPrev=When
formlowerPrev=when suf3Prev=hen suf4Prev=When isTitlePrev
formNext=concurrently formlowerNext=concurrently suf3Next=tly
suf4Next=ntly

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Sequence tagging: the B-I-O approach

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Feature Extractor

```
# process each file in directory
for f in listdir(datadir) :
    # parse XML file, obtaining a DOM tree
    tree = parse(datadir + "/" + f)
    # process each sentence in the file
    sentences = tree.getElementsBvTagName("sentence")
    for s in sentences :
        sid = s.attributes["id"].value # get sentence id
        stext = s.attributes["text"].value # get sentence text
        # load ground truth entities.
        gold=[]
        entities = s.getElementsByTagName("entity")
        for e in entities :
            # for discontinuous entities, we only get the first span
            offset = e.attributes["charOffset"].value
            (start,end) = offset.split(";")[0].split("-")
            gold.append((int(start), int(end), e.attributes["type"].value))
        # tokenize text
        tokens = tokenize(stext)
        # extract features for each word in the sentence
        features = extract features(tokens)
        # print features in format suitable for the learner/classifier
        for i in range (0, len(tokens)):
            # see if the token is part of an entity, and which part (B/I)
            tag = get_tag(tokens[i], gold)
            print (sid, tokens[i][0], tokens[i][1], tokens[i][2],
                   tag, "\t".join(features[i]), sep='\t')
        # blank line to separate sentences
        print()
```

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Feature Extractor

 $Core\ task$

Feature Extractor Functions - Tokenize text

```
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NERC
```

Sequence tagging: the B-I-O approach

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Feature extraction for NLP

- In most ML applications, the feature space is finite and known (e.g. credit scoring, medical diagnose prediction, churn prevention, fraud detection, etc).
- Also, most of the used features are numerical or categorial (income, age, sex, colestherol level, number of receipts returned, etc.)
- Thus, in these ML applications, feature vectors are usually exhaustive lists of pairs feature-value.

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Sequence tagging: the B-I-O approach

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Feature extraction for NLP

- In most ML applications, the feature space is finite and known (e.g. credit scoring, medical diagnose prediction, churn prevention, fraud detection, etc).
- Also, most of the used features are numerical or categorial (income, age, sex, colestherol level, number of receipts returned, etc.)
- Thus, in these ML applications, feature vectors are usually *exhaustive* lists of pairs feature-value.

BUT...

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Sequence tagging: the B-I-O approach

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Feature extraction for NLP

- In most ML applications, the feature space is finite and known (e.g. credit scoring, medical diagnose prediction, churn prevention, fraud detection, etc).
- Also, most of the used features are numerical or categorial (income, age, sex, colestherol level, number of receipts returned, etc.)
- Thus, in these ML applications, feature vectors are usually exhaustive lists of pairs feature-value.

BUT...

- In most NLP applications, features are related to appearing words, suffixes, prefixes, lemmas, etc. Thus, the feature space is huge.
- Moreover, features are usually binary-valued (a word appears or not, a suffix appears or not, etc).
- Thus, in NLP applications, feature vectors are usually intensive lists of strings (i.e. listing the names for features with value true, and ommiting all the rest), and are stored as sparse vectors.

Machine Learning NERC

Sequence tagging: the B-I-O approach

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Core task

Feature Extractor Functions - Extract features

Machine

Learning

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approach

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Deliverables

Feature Extractor

tagging: the

NFRC

B-I-O

```
def extract features(s) :
 Task:
   Given a tokenized sentence, return a feature vector for each token
 Input:
   s: A tokenized sentence (list of triples (word, offsetFrom, offsetTo) )
 Output:
   A list of feature vectors, one per token.
   Features are binary and vectors are in sparse representation (i.e. only
     active features are listed)
 Example:
   >>> extract features([("Ascorbic".0.7), ("acid".9.12), (".".13.13),
          ("aspirin",15,21), (",",22,22), ("and",24,26), ("the",28,30),
          ("common", 32, 37), ("cold", 39, 42), (".", 43, 43)])
   [ "form=Ascorbic", "suf4=rbic", "next=acid", "prev= BoS ", "
    capitalized" 1.
   "form=acid", "suf4=acid", "next=,", "prev=Ascorbic"],
    "form=.". "suf4=.". "next=aspirin". "prev=acid". "punct" ].
      [ "form=aspirin", "suf4=irin", "next=,", "prev=,"],
      . . .
```

Feature Extractor Functions - Ground truth tag

```
def get tag(token, gold) :
                 Task:
                    Given a token and a list of ground truth entites in a sentence, decide
                     which is the B-I-O tag for the token
Machine
Learning
                 Input:
NFRC
                    token: A token, i.e. one triple (word, offsetFrom, offsetTo)
                    gold: A list of ground truth entities, i.e. a list of triples (
Sequence
                     offsetFrom, offsetTo, type)
tagging: the
B-I-O
                 Output:
approach
                   The B-I-O ground truth tag for the given token ("B-drug", "I-drug", "B-
                     group", "I-group", "0", ...)
General
Structure
                 Example:
                    >>> get_tag(("Ascorbic",0,7), [(0, 12, "drug"), (15, 21, "brand")])
Detailed
                   B-drug
Structure
                   >>> get_tag(("acid",9,12), [(0, 12, "drug"), (15, 21, "brand")])
Feature Extractor
                   I-drug
                   >>> get_tag(("common",32,37), [(0, 12, "drug"), (15, 21, "brand")])
Core task
                    n
Goals &
                    >>> get_tag(("aspirin",15,21), [(0, 12, "drug"), (15, 21, "brand")])
Deliverables
                   B-brand
```

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Machine Learning NERC

Sequence tagging: the B-I-O approach

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Learner

General

Core task

Learner - Option 1: CRF

Machine Learning NERC

Sequence tagging: the B-I-O approach

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Core task

- Install and import pycrfsuite \$ pip install python-crfsuite
- Use provided train-crf.py to learn a model.
 \$ python3 train-crf.py model.crf < train.feat
 You may modify learner parameters (loss function, thresholds, learning rates, etc).

Learner - Option 2: Maximum Entropy

Use megam to train a ME model:

- megam does not expect the extra information in the features file,
 so you need to remove the first 4 fields (sent_id, token,
 span_start, span_end) and the blank lines between sentences:
 \$ python3 extract-features.py data/train > train.feat
 \$ cat train.feat | cut -f5- | grep -v ^\$ > train.mem.feat
- Then you can use the fixed feature-encoded dataset to train a MEM model:
 - \$./megam-64.opt -quiet -nc -nobias multiclass train.mem.feat >
 model.mem

```
Detailed
Structure
Learner
```

Machine

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Sequence

B-I-O approach

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Structure

tagging: the

Core task

Learner - Option 3: Your choice

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Sequence tagging: the B-I-O approach

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Core task

- Select a ML algorithm of your choice (DT, SVM, RF, ...) and a python library implementing it.
- Adapt the feature file format to the needs of the selected algorithm.
- Train a classification model for the task of predicting B-I-O tags for each token.
- Create a module XXX.py with a constructor and a predict method, following the structure of CRF.py and MEM.py.
 Add your new classifier to the constructor in ML_model.py
- DO NOT use neural network approaches, we'll do that later in the course.

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Sequence tagging: the B-I-O approach

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Classifier

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Core task

Classifier - All options

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Detailed Structure Classifier

Core task

Goals & Deliverables

You can apply the learned models to new data:

- \$ python3 extract-features.py data/devel > devel.feat
- \$ python3 predict.py model.mem <devel.feat >devel.out
- \$ python3 util/evaluator.py data/devel devel.out
- \$ python3 extract-features.py data/test > test.feat
- \$ python3 predict.py model.mem <test.feat >test.out
- \$ python3 util/evaluator.py data/test test.out

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Machine Learning NERC

Sequence tagging: the B-I-O approach

General Structure Detailed Structure

Core task

Strategy to follow:

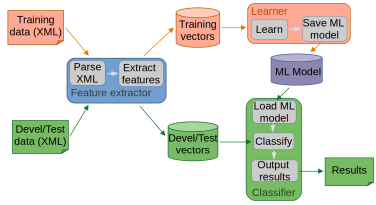
Machine Learning NERC

Sequence tagging: the B-I-O approach

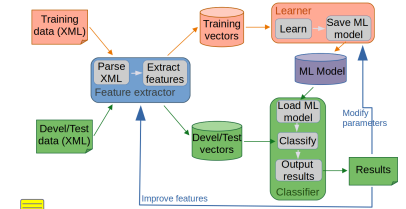
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Strategy to follow:



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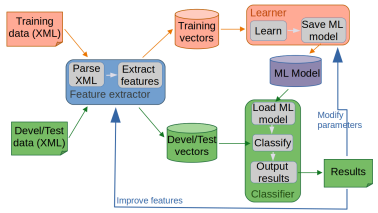
Sequence tagging: the B-I-O approach

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Core task

Strategy to follow:



 Repeat training – evaluation cycle on devel dataset to find out which is the best parameterization for the used algorithm.

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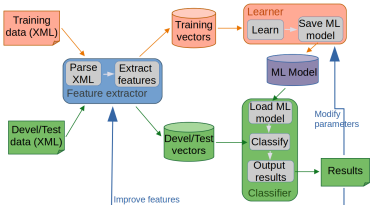
Sequence tagging: the B-I-O approach

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Core task

Strategy to follow:



- Repeat training evaluation cycle on devel dataset to find out which is the best parameterization for the used algorithm.
- Repeat feature extraction training evaluation cycle on devel dataset to find out which features are useful.

Machine Learning NFRC

Sequence tagging: the B-I-O approach

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Core task

Choosing useful features

- Used models are token classifiers, so there is a feature vector per token.
- Features about a token should allow its classification, so they should encode information about *both* the token itself and its context (i.e. nearby words).
- Feature names must be *unambiguous*. E.g., a feature named sufx=azole may not be enough if one wants to encode also context word suffixes. In that case, different feature names are needed (e.g.: sufx=azole for the focus word, plus e.g. sufx-2=azole, sufx-2=azole, sufx+1=azole, sufx+2=azole for nearby words).
- As in the rule-based approach, including features encoding information from external dictionaries will largely improve results.

Machine Learning NERC

Sequence tagging: the B-I-O approach

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Choosing useful features

Machine Learning NFRC

Sequence tagging: the B-I-O approach

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Goals & Deliverables ■ REMEMBER: Feature names such as sufx=azole are not key-value pairs (i.e. not a sufx feature with value azole), but just a string naming a binary (true/false) feature. The feature name could be any (sufxisazole, wordendsinazole, ...) as long as it is active (i.e. present in the sparse vector) only when that property holds.

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Exercise Goals

What you should do:

- Work on your feature extractor. It is the component of the process where you have most control.
- Experiment with different parameterizations of the chosen learner. You may try different learning algorithms if you feel up to. Note that the same feature vectors can be fed to different learners (maybe with some format adaptation).
- Keep track of tried features and parameter combinations, and results produced by each.

What you should **NOT** do:

- Use neural network learners. We'll do that later on the course.
- Alter the provided code structure.

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Sequence tagging: the B-I-O approach

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Exercise Goals

Orientative results:

- Provided initial version achieves 55%(CRF)/47%(MEM) macro average F1 on devel with 2 simple feature templates.
- A set of 10 feature templates is enough to get a macroaverage F1 about 74%(CRF)/66%(MEM). Used information includes (for current, previous, and next tokens)
 - word forms, original and lowercase
 - suffixes (of different lenghts)
 - capitalization pattern (all upper, title, camelcase,...)
 - presence of numbers, dashes, etc
 - existence of the token in external lists
 - **..**

Machine Learning NFRC

Sequence tagging: the B-I-O approach

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Deliverables

Write a report describing the work carried out on NERC tasks. The report must be a **single self-contained PDF document**, under 10 pages, containing:

- *Introduction:* What is this report about. What is the goal of the presented work.
- Rule-based baseline
 - Ruleset construction: What did you observe in the data exploration. Which rules did you wrote according to those observations.
 - Code: Include your extract_entities function (and any other function it may call), properly formatted and commented. Do not include any other code.
 - Experiments and results: Results obtained on the devel and test datasets, for different rule combinations you deem relevant.

Machine Learning NERC

Sequence tagging: the B-I-O approach

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Deliverables (continued)

Machine learning NERC

- Selected algorithm: Which classifier/s did you select or try.
 Reasons of the choice. Comparison if you tried more than one.
- Feature extraction: Tried/discarded/used features. Impact of different feature combinations.
- Code: Include your extract_features function (and any other function it may call), properly formatted and commented. Do not include any other code.
- Experiments and results: Results obtained on the devel and test datasets, for different algorithms, feature combinations, parameterizations you deem relevant.
- Conclusions: Final remarks and insights gained in this task.

Keep result tables in your report in the format produced by the evaluator module. Do not reorganize/summarize/reformat the tables or their content.

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