DreamDrug - A crowdsourced NER dataset for detecting drugs in darknet markets

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Abstract

We present DreamDrug, a crowdsourced dataset for detecting mentions of drugs in noisy user-generated item listings from darknet markets. Our dataset contains nearly 15,000 manually annotated drug entities in over 3,500 item listings scraped from the darknet market platform "DreamMarket" in 2017. We also train and evaluate baseline models for detecting these entities, using contextual language models fine-tuned in a few-shot setting and on the full dataset, and examine the effect of pretraining on in-domain unannotated corpora.

1 Introduction

This paper describes the construction of DreamDrug, a dataset of drug-related product listings from darknet marketplaces with human-annotated drug entities, suitable for training and evaluating named entity recognition (NER) systems. We provide a detailed description of all steps of annotation and dataset construction, and we also optimize and evaluate standard NER architectures on the new corpus. Our experiments include training models in a few-shot setting and using the full dataset, as well as a study of the effect of domain-adaptive pre-training using unannotated datasets of similar topic and genre. All software used in producing DreamDrug and for reproducing our experiments is published under an MIT license along with sample data files¹, the full dataset is available upon request². Our main contributions are the following:

- Specifications of an annotation task for marking mentions of drug entities in darknet market listings
- 2. A detailed description of preprocessing and

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filtering steps used to create the annotation input

- 3. The novel NER corpus DreamDrug, annotated by crowdworkers based on these specifications and reviewed manually by the authors
- 4. Quantitative evaluation of standard supervised learning architectures for NER on the DreamDrug dataset, in a few-shot setting and using the full dataset
- Three unannotated datasets for domainadaptive pretraining and experiments demonstrating their impact on the performance of the NER baselines

The paper is structured as follows. In Section 2 we introduce the task and provide an overview of recent related work on information extraction from noisy user-generated text. Section 3 describes the corpus construction process. Section 4 and Section 5 present our baseline experiments and results, respectively, and Section 6 provides a brief conclusion.

2 Background

Extracting information from text concerning illicit drug consumption or trade is a non-trivial task, since such information is rarely available in a structured form. Patients who misuse their medication are more likely to discuss this on social media than with medical professionals (Bigeard et al., 2018), while drug dealers intentionally misspell names of drugs on social media to prevent automatic detection (Li et al., 2019). Drug-specific named entity recognition (NER) models are most often developed for the medical domain and cannot handle such misspellings, the usage of slang, and other characteristics of noisy user-generated texts such as lack of punctuation and non-standard grammar (Rezaei et al., 2020). Many of these issues are

Inttps://github.com/jbogensperger/
DRUG_CROSSNER

²DreamDrugDataset@gmail.com

peruanse 92 cocain very good snife and for people tghe want cooking out very good smills good frech from the block orginail good good good AAA.

Figure 1: Sample listing from the *DreamMarket* dataset

illustrated by the example listing from a darknet marketplace in Figure 1.

Still, many organizations are dependent on extracting information about drugs or similiar goods from noisy online sources. Public health organizations study drug use patterns in society such as the opioid epidemic in the United States (Ostling et al., 2018), pharmaceutical companies can collect evidence of adverse drug reactions from social media data (Chen et al., 2017), and law enforcement agencies can track illicit trading activities in darknet markets. Each of these use-cases can be supported by the automatic detection of drug entities in noisy user-generated text. To enable the development of such systems we create a dataset for training and evaluation by annotating drug-related item listings of darknet marketplaces. An example of such a listing is shown in Figure 2.

2.1 NER in User-Generated Texts

Much of recent research on named entity recognition in user-generated texts was facilitated by the 2017 W-NUT Shared Task on Novel and Emerging Entity Recognition (Derczynski et al., 2017). The dataset provided by the organizers of the competition was compiled from a variety of social media sources (Twitter, Reddit, Youtube, StackExchange), annotated for 6 categories (person, location, corporation, product, creative work, group) by multiple crowdworkers, and corrected by the authors. The difficulty of the task of detecting novel entities in noisy domains is indicated by the relatively low performance of all competing systems. The top-performing system of Aguilar et al. (2017) achieved an average F-score of 41.86 on the evaluation dataset. Their system combines character- and word-level features with gazetteers for each entity type, and uses them as input to a Bidirectional Long Short-term Memory (BiLSTM) network with a Conditional Random Fields (CRF) classifier. Other top-performing approaches included Jansson and Liu (2017), who use LDA topic modeling for creating word-level features for an LSTM classifier with a CRF output layer, and Lin et al. (2017), who incorporated part-of-speech tags and dependency relations in their word representations and also used

a BiLSTM with CRF output layer. A significant improvement on this benchmark was achieved by Akbik et al. (2019b) using pooled contextualized embeddings, a method to aggregate character-level representations from contextual language models. This method is implemented in the FLAIR framework (Akbik et al., 2019a) and achieves an F-score of 49.59 on the W-NUT 2017 dataset.

We are aware of two annotated NER corpora from the darknet domain. Durrett et al. (2017) created a dataset for identifying products in cybercrime marketplace postings, annotating 1,938 posts across 4 forums. When training NER models on data from the same forum, they achieve F-score performances between 83 and 90, in a cross-forum setting their performance drops by 7-11 points. A more recent darknet dataset is NUTOT (Nabki et al., 2020), containing 851 sentences from darknet sources annotated manually with the categories of the W-NUT challenge. Their experiments involve compensating for the absence of domain-specific gazetteers with Local Distance Neighbor (LDN) features, a method for enriching word representations with nearest neighbors in a word embedding space. Performance of top-performing systems on their dataset is in the range of 40 to 60 across categories (F-scores of entities). The corpus we introduce in this paper is annotated for a single entity type (drug) that is more specific than the categories of W-NUT and NUToT and allows our simple baselines to achieve F-scores above 70 even in a few-shot setting and above 80 when using the full dataset (see Section 5). In addition to enabling higher accuracy, we believe that NER datasets that correspond directly to a narrow information extraction task are better models of real-world NER use-cases than the commonly used datasets labeled for broad categories such as person, location, organization, or even product. While the fine-grained task definiton necessary for consistent annotation of DreamDrug (see Section 3.2) requires considerations specific to a single task, it also provides an example of the narrow and domain-specific entity categories that are typical of real-world NER applications. Besides providing a novel dataset and baseline models, our work also demonstrates that entity recognition tasks for narrow domains and specialized text genres can benefit from even small amounts of annotated data.

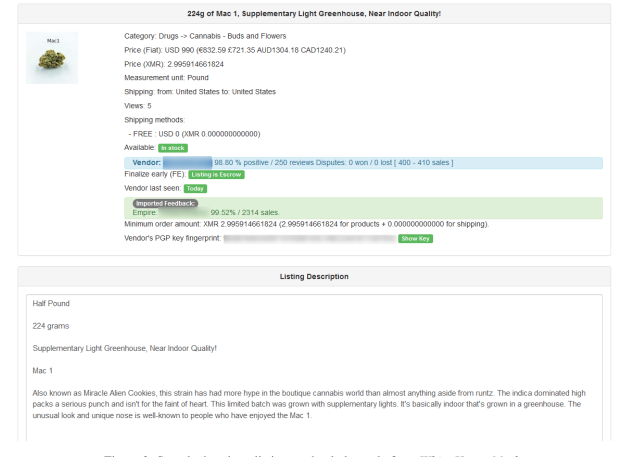


Figure 2: Sample drug item listing on the darknet platform White House Market

2.2 Crowdsourcing for NER

Crowdsourcing, the practice of soliciting nonexpert annotators for assistance in creating datasets on a scale, usually with the help of online platforms, has become a standard practice of researchers aiming to build annotated corpora with limited resources. The most common scheme, where crowd workers are rewarded for each annotation task they complete, is sometimes referred to as mechanized labor (Sabou et al., 2014). In Section 3.3 we shall give a detailed description of the entire process of creating the DreamDrug dataset. For defining the Named Entity Recognition task we consult several recent sets of guidelines for NER tasks (Finin et al., 2010; Nédellec et al., 2006; Benikova et al., 2014). When designing the annotation process we rely on discussions of annotation methodologies in (Sabou et al., 2014) and (Fort et al., 2009). Approaches to measuring inter-annotator agreement are studied in detail by Asheghi et al. (2014), crowdsourcing techniques are discussed by Feyisetan et al. (2018), and we also rely on the work of Gomes et al. (2020) for initial effort estimation, which we see as essential in ensuring the ethical treatment of annotators.

3 Corpus construction

3.1 Data preparation

The starting point for our corpus construction process is a segment of the AZSecure dataset (Du et al., 2018) containing 91,463 darknet market listings scraped from Dream Market, collected between 2013 and 2017. Listing documents typically contain a product name and description along with metadata such as price, quantity, contact information of the vendor, etc. We extracted 45,446 items belonging to categories associated with drugs and performed a series of data cleaning steps. First, deduplication removed all listings containing exactly the same description text as another item listing. The English only step removed all listings with a description text categorized as other than English by the language-detection library (Shuyo, 2010). Next, outlier lengths removed listings with a description text longer than 3000 characters or shorter than 30 characters. While this step also removed some relevant content, it efficiently filtered product listings with long boilerplate sections (e.g. about delivery details),

as well as items that contained no relevant text (e.g. a product description might only state *Free shipping*). Finally, soft deduplication removed listings with a description text whose first 100 characters contained the same non-numeric characters as another listing's description, characteristic of listings of the same product with different quantities. The resulting dataset contains 11,674 items, Table 1 shows the number and percentage of listings removed by each filtering step.

filtering step	# removed	% removed	# kept
deduplication English only outlier lengths soft deduplication	25,012	55.0%	20,434
	2,102	10.3%	18,332
	1,488	8.1%	16,844
	5,170	30.7%	11,674

Table 1: Number and percentage of item listings filtered in each step (see text for description of each step)

For the construction of the DreamDrug dataset we used the product name and product description fields of the 11,674 remaining listings, the final dataset contains 3,507 items randomly selected from this set. Common unique identifiers of vendors in the product description texts were replaced with random values. Telephone numbers were detected using the python library phonenumber³, email addresses and URLs were replaced using regular expressions (see Appendix B) and the faker⁴ package, while vendor names were available as separate fields in the original database. Finally, product description texts were tokenized using Stanza⁵ (Oi et al., 2020), version 1.2.0.

3.2 Task definition

Our definition of which sequences of words constitute drug entities was developed in an initial set of experiments that involved manual annotation of 500 item listings by the authors. Our starting point was the definition used by the Food and Drug Administration (FDA) (U.S. Food and Drug Administration, 2021):

- "articles intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease"
- 2. "articles (other than food) intended to affect the structure or any function of the body of

man or other animals."

The guidelines we developed for consistent annotation clarify that for an entity to be labeled it must uniquely identify a specific article according to the FDA definition. Besides direct mentions of substances (cocaine, THC, etc.), slang terms that clearly identify a drug, such as speed and pep for amphetamine or Mary Jane for cocaine, are also considered drug entities. Phrases that do not uniquely identify a substance, such as Charley's mellow Sleeper Bars or Green Dom Perignon, are not to be annotated. Following the FDA definition we do not annotate substances occurring naturally in the human body such as testosterone or dopamine. Phrases referring to the physical form of a drug only are also not annotated, even if the form is unique to a particular drug, as e.g. blotter is to LSD. The common term mushroom constitutes a corner case, as one may argue that it refers to the form of a drug, but since in this dataset it clearly identifies a particular chemical (psilocybin, the active agent of all types of magic mushrooms), we decided to annotate it as a drug entity. Some phrases describing drugs contain multiple elements that could each be annotated as a separate drug entity based on our definition. Examples include phrases such as Viagra Sildenfanil and Haze weed — we annotate these as two subsequent drug spans. The annotation guidelines provided to crowd workers contain a simplified description of our definition with examples and is presented in Appendix C. All documents annotated by crowd workers were subsequently reviewed and corrected manually by the authors to ensure high quality and consistent treatment of corner cases — details of the annotation and review process are presented in Section 3.3.

3.3 Data annotation

Crowdsourced annotation was performed using the definition of drug entities presented in Section 3.2. The guidelines presented to annotators were revised several times during pilot experiments that involved the implementation of two annotation interface prototypes using the platforms Appen and Amazon Mechanical Turk (AMT), both of which are regularly used for annotating NER datasets (Jalal et al., 2020; Feyisetan et al., 2015; Bontcheva et al., 2017; Feyisetan et al., 2018). Appen was then chosen, primarily because of its built-in capabilities for evaluating annotator performance and for screening annotators using test questions. Due to an unexpected

³https://pypi.org/project/
phonenumbers/

⁴https://github.com/joke2k/faker

⁵https://stanfordnlp.github.io/stanza/

Drug Labelling

In this HIT we want you label all text spans, which clearly point to a specific drug or drug type.

HOW TO DO THIS?

- 1. Read the Context above the text to get an idea of which items could be contained in the text
- Read the provided drug description and label ALL spans which clearly identify a drug
 Are you unsure about what is a drug? We provided a extensive description, just press on "Instructions" and go to "more instructions".

Keep in mind: Each span should Clearly identify a specific drug or drug type!

Practical examples are:

- Clearly represent a specific drug e.g. hash, meth
 Chemical Compounds e.g. "3, 4 methylphendiate"
- Short names e.g. XTC Agents e.g. THC, CBD or Diazepam
- Slang drug names: Coke, Fent
- Drug categories e.g. Indica/hybrid, benzodiazipines, opiates
- Pill imprints e.g. "V-3923"
- Chemical Details which extend the drug e.g. Viagra Sildenafil Citrate
 Legal or partial legal drugs e.g. "Alcohol", "Testosterone Enthalate", "Aspirin" or "Viagra"

DO NOT label tokens like:

- · Description/Adjectives like drug strength, colour, form (powder/pill..) or looks etc.
- Form of Drug: e.g. powder, pill, paste
 Vague references like "this pill", "My Super Chocolate Bars" or "indoor bud"
- Producer of drugs e.g. Pfizer
- · Natural Chemicals in your body e.g. estrogen/dopamin

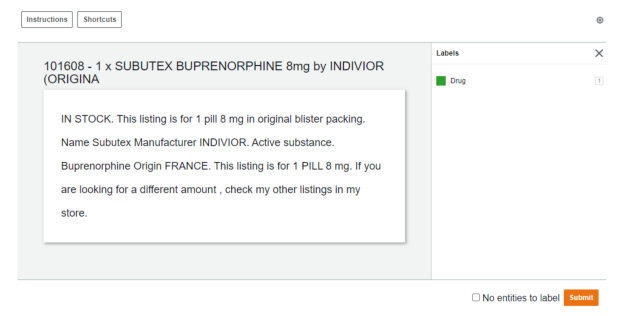


Figure 3: Screenshot of the annotation interface on Amazon Mechanical Turk. The long version of the annotation guidelines (see Appendix C) was accessible via the *Instructions* button.

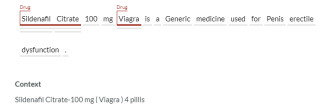


Figure 4: Screenshot of the annotation interface on Appen.

licensing issue we were forced to switch platforms during the annotation process, and the final dataset was constructed using annotations from both Appen and AMT, agreement figures and dataset statistics in this section are provided for both platforms. Crowdworkers were presented with the text of one listing at a time and asked to mark the beginning and end of sequences of words that identify a drug. The title of the listing was provided as context. Screenshots of the annotation interfaces on Amazon Mechanical Turk and Appen are shown in Figures 3 and 4, respectively. On Appen, instructions were provided in a separate window and contained the full annotation guidelines shown in Appendix C. These longer guidelines were available to AMT workers when they clicked the *Instructions* button.

Annotation quality was monitored by measuring Inter-Annotator Agreement (IAA) and by conducting manual reviews of annotation samples from each worker. Parameters of the annotation process such as reward per task, maximum time per assignment, and number of annotations solicited for each listing, were updated during the annotation process based on our experience — a detailed description of this process is available in the Appendix of (Bogensperger, 2021). We obtained over 4,000 valid annotations from Appen and over 7,400 from AMT. Appen annotators received a fixed payment of either \$0.06 or \$0.08 per annotated document (the exact rate changing across batches), on AMT rates also varied across annotators and payments per row ranged between \$0.09 and \$0.30, while hourly rates varied between \$12 and \$18. Our total incurred costs were \$548 on Appen and \$1505 on AMT, in the former case this included payment for 995 additional annotations that were invalidated based on annotators' performance on the test questions. Table 2 shows inter-annotator agreement (IAA) values for annotations obtained via each of the two platforms. Following Viera and Garrett (2005) and Hripcsak and Rothschild (2005) we used F1-Agreement as the primary IAA measure,

IAA	AMT	Appen
Cohen's Kappa	0.76	0.43
Micro F1	0.79	0.55
Macro F1	0.78	0.60

Table 2: Inter-Annotator Agreement measures for the overall annotations by crowd workers of Amazon MTurk and Appen. Macro F1 is the average of F1 scores across pairs of annotators.

	MTurk	Appen
% of characters added	8.56	25.81
% of characters deleted	3.51	0.89
% of spans added	13.19	27.49
% of spans deleted/altered	8.65	10.45

Table 3: Average effort of the review process.

Cohen's Kappa is also calculated for comparison. Spans annotated by at least two annotators were reviewed and corrected manually by the authors, then all annotated documents were reviewed once again by the authors to find missing annotations. The review process followed the principles described in Section 3.2. Table 3 shows the amount of editing performed for each portion of the annotations. Both of these figures and the agreement scores show a considerable difference between the quality of annotations from the two platforms, and without further study we can only speculate that this may be attributed to the continuous improvement of the annotator instructions based on our experiences, the different presentations of these instructions (MTurk annotators had access to a short and long version), or to the different incentive systems of the two platforms (AMT workers were required to hold a "master" qualification on the platform, which they could lose when providing low-quality annotations, while Appen workers' performance was only controlled on the test questions).

The final dataset contains 3,507 item listings with 364,003 tokens, 14,934 annotated spans and 3,048 unique drug entities. The dataset was split into training, validation, and test portions of 2244, 561, and 702 listings, respectively. Table 4 provides basic descriptive statistics about the DreamDrug dataset, further statistics as well as a list of the most frequent spans are presented in Appendix A. Manual evaluation of a sample of rare spans reveals that while some of them refer to

	Min	Max	Mean	Median
words / doc	3	534	103.8	61
chars / doc	25	2,930	572.6	334
words / entity	1	20	1.9	2
chars / entity	1	80	12.1	10
entity / doc	0	101	4.26	2

Table 4: Descriptive Statistics of dataset

rare entities, such as names of cannabis strains (e.g. *alien technology*) or specific chemical descriptors (e.g. *n-acetyl-p-aminophenol*), most of them are spelling variations of common drugs (e.g. "Oxycodone hydrochloride" instead of ... "Oxycodone HCL") and misspellings ("Oxymorophone" instead of "Oxymorphone").

4 Experiments

We train a set of baseline transformer-based models, examine the effects of domain-adaptive pretraining using multiple datasets, and also evaluate top-performing models in a few-shot training scenario.

4.1 Baseline models

We train the standard BERT-based architecture for Named Entity Recognition (Devlin et al., 2018), which consists of a transformer layer and a linear layer with dropout, using the PyTorch-based implementations from Huggingface (Wolf et al., 2020), similar to the system in (Liu et al., 2021). We train systems based on the bert-base-cased and roberta-base models with default parameters, using the Adam optimizer, a learning rate of $5 \cdot 10^{-5}$, and a batch size of 4. For the few-shot learning setup we used only the first 100 documents from the training set. In both setups we trained our models for 10 epochs. We also train the FLAIR⁶ system (Akbik et al., 2019a) using default hyperparameters and training for 50 epochs. We use BERT (bert-base-cased) as the input representation of the FLAIR system because it yields better results than FLAIR's own contextualized embedding (Akbik et al., 2019b). For evaluation we use standard NER metrics as defined by the CoNLL 2003 shared task (Tjong Kim Sang and De Meulder, 2003), computing precision, recall, and F-score, where a predicted named entity is considered correct only if it is an exact match of a

gold standard entity. While Named Entity Recogntion is generally implemented on the sentence level, the relatively short average length of listings (see Table 4) allows us to train the baselines on full documents, limited only by the maximum input length of BERT, 512 (subword) tokens. While the average number of words per document in our dataset is 103.79, the average number of subword tokens output by BERT's tokenizer is 152.18, and 4.1% of documents are longer than the maximum input length of 512, these were clipped at 512 tokens for training and evaluation of each of our baselines. A final preprocessing step involved removing all punctuation except commas from item descriptions to reduce noise, the regular expressions we used are presented in Appendix B.

4.2 Domain adaptation

We improve the performance of our baselines by pretraining on unannotated text representing the domain and/or genre of the DreamDrug dataset. We compile three datasets that can be used separately or in combination. Dreammarket contains Dreammarket product listings of drugs that were not used in creating DreamDrug. The size of this dataset is slightly above 800 000 words. The Grams dataset is constructed from the Darknet Market Archives (Branwen et al., 2015), using the subsets Abraxas, Agora, Alpha, ME, and Oxygen, and contains listings from a variety of domains. This dataset contains nearly 1.7 million words. The third dataset used for domain adaptation, Wikipedia, was constructed using 3,210 drug-related articles downloaded from Wikipedia⁷ using its Python API⁸. Sections of Wikipedia articles that do not typically contain running text (References, See also, and External Links) were removed, and the remaining text was searched for possible mentions of drug entities by annotating each sentence for mentions of entities in the knowledge graph DBPedia (Auer et al., 2007) using the annotate function of the Spotlight API⁹, which performs fuzzy matching. Drug entities were detected based on the presence of the DBPedia attributes dbp:legalUs or dbp:legalUn, sentences containing entities with either of these attributes were added to the dataset used for domain adaptation. The size of the resulting dataset is

⁶https://github.com/flairNLP/flair

⁷https://en.wikipedia.org/

⁸https://pypi.org/project/wikipedia/

⁹https://pypi.org/project/
spacy-dbpedia-spotlight/

Base	DAPT	D	F1	Prec.	Rec.
FLAIR	None	NA	59.72	60.29	59.16
BERT	None	0	63.80	67.84	60.21
BERT	All	0	69.86	72.91	67.06
BERT	None	0.5	62.93	70.59	56.77
BERT	DM	0.5	69.43	74.20	65.23
BERT	Grams	0.5	68.49	71.87	65.41
BERT	Wiki	0.5	64.32	68.76	60.42
BERT	All	0.5	70.98	76.73	66.04
RoBERTa	All	0.35	69.11	73.54	65.18

Table 5: Performance of top baseline configurations in the few-shot scenario. The column DAPT indicates the dataset(s) used for domain-adaptive pretraining, D indicates dropout rate.

more than 880 000 words, the three datasets for domain adaptation together contain nearly 3.4 million words.

5 Results

Tables 5 and 6 present the perfomance of top baseline configurations in the few-shot setting and when using the full training set. The BERT-based systems achieved higher scores in the few-shot setup, while RoBERTa-based setups performed better when trained on the full dataset. For both setups we include the performance of the top system with and without dropout as well as with domain-adaptive pretraining (DAPT) on each of the additional datasets. For both the few-shot setup and the full training set we also include the best-performing configuration of the other language model (RoBERTa for the few-shot setting and BERT for the full dataset) and the performance of the best FLAIR model. The effect of domain adaptation using in-domain unannotated text is especially pronounced in the few-shot setting, where it achieves an 8 point increase in F-score, 6.5 of which can be achieved using only the DM dataset, which is the one most similar to the listings in DreamDrug. We believe this shows the strong potential of pretraining on unannotated text that is characteristic of both the domain and the genre of a novel task, especially in settings where the availability of human-annotated data is limited. Finally, Table 7 contains the results of our top models on the test set.

6 Conclusion

We have constructed the novel dataset DreamDrug for the task of recognizing drug entities in darknet market listings, used it

Base	DAPT	D	F1	Prec.	Rec.
FLAIR	None	N/A	81.42	80.83	82.01
RoBERTa	None	0	80.84	81.13	80.56
RoBERTa	Wiki	0	80.83	83.65	78.20
RoBERTa	All	0	81.21	86.63	76.43
RoBERTa	None	0.1	80.55	81.43	79.69
RoBERTa	DM	0.1	80.41	83.19	77.80
RoBERTa	Grams	0.1	80.52	84.99	76.49
RoBERTa	Wiki	0.1	82.16	84.86	79.62
RoBERTa	All	0.1	81.51	85.37	77.98
BERT	All	0.45	81.64	85.71	77.94

Table 6: Performance of top baseline configurations using the full training set. The column DAPT indicates the dataset(s) used for domain-adaptive pretraining, D indicates dropout rate.

Model	F1	Prec.	Rec.		
few-shot training					
FLAIR	62.13	65.18	59.35		
Our best model	73.55	78.38	69.28		
full training set					
FLAIR	81.82	82.75	80.92		
Our best model	83.86	84.83	82.92		

Table 7: Performance of top baseline configurations on the test set

to train and evaluate baseline NER models, and demonstrated the effectiveness of domain-adaptive pretraining on large unannotated datasets. As one of the first entity-annotated corpora of darknet text, and the first task-specific one, <code>DreamDrug</code> will facilitate the study of named entity recognition and related information extraction tasks in noisy user-generated texts. Additionally, it can also be extended with several types of annotation, including entities from multiple classes, relations such as "drug-quantity" or "drug-strength", or links from entity mentions to a knowledge base.

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Ethical considerations

An analysis of potential project internal and external impacts of the research activity was conducted as part of the COPKIT project. The relevance of data protection principles, such as lawfulness, fairness, and transparency, purpose limitations, data minimization, accuracy, storage limitation, integrity and confidentiality, and accountability has been analyzed for this dataset publication. The activity was led by the project internal ethical, legal, and privacy team. Regarding concrete measures, the dataset was pseudonymized to remove any personal data, and the base dataset for groundtruth creation was approved for use by the ethical, legal, and privacy team. It was decided to provide access to the data only on request to prevent misuse of the dataset for purposes other than research.

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A Dataset statistics

Figure 6 shows a histogram of the number of tokens per document, Figure 7 and Table 8 present the most frequent drug entities. Figure 5 contains the entity frequency histogram.

Entity	Frequency
cocaine	476
thc	435
mdma	428
cannabis	365
indica	350
sativa	336
hash	239
weed	215
lsd	214
hybrid	175
xanax	154
cbd	151
marijuana	150
og kush	120
xtc	116
sour diesel	116
alprazolam	104
heroin	97
oxycodone	96
speed	93
adderall	93

Table 8: Most frequent drug entities in the DreamDrug dataset (lowercased)

B Custom preprocessing

Listing 1: Preprocessing function for the removal of links and special characters from text

```
def remove_unwanted_elements(text):
    ft = re.sub(
        r'https?:\/\\S*[\r\n]*', '', text)
    ft = re.sub(
        r'\S*.onion\S*[\r\n]*', '', ft)
    ft = re.sub(
        r'[\+!~@#$%^&*()={}\[\]:;<.>?\`"]',
        '', ft)

    ft = re.sub(',', '&#44', ft)

    ft = re.sub(r'[-]+', '-', ft)
    ft = re.sub(r'[_]+', '_', ft)
    return ft
```

Listing 2: Preprocessing functions for replacing email addresses

```
from faker import Faker
# create random name
def get_fake_name():
    # can create fake names
    fake = Faker()
    fake_name = fake.name().replace(
    if random.randint (1, 10) \% 2 == 0:
         fake_name = fake_name.lower()
    if random.randint (1, 10) \% 3 == 0:
         fake_name = "%s%d" % (
             fake_name.lower(),
             random.randint(10, 99))
    return fake_name
def replace_mails(text):
    domain = [
         '.com ; , '.de ', '.ru ', '.org ']
    vendor = [
         'gmail', 'air', 'wing', 'microsoft', 'hotmail',
         'outlook']
    mail = get_fake_name() + '@' +
         vendor [random.randint(
             0, len(vendor)-1)] +
         domain [random.randint(
             0, len(domain)-1)
    return re.sub(
         r'\S+@\S+\scalebox{$\cdot$} s?', mail, text)
```

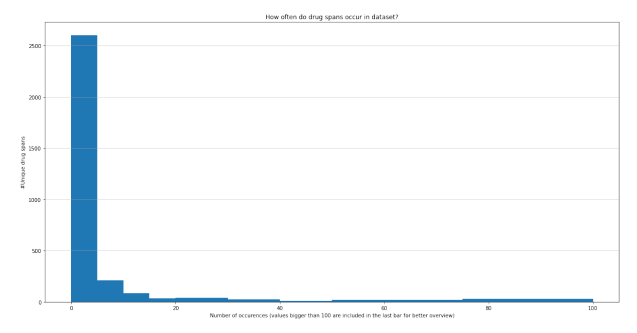


Figure 5: Frequency distribution of drug entities

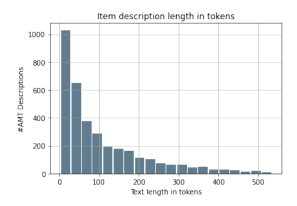


Figure 6: Distribution of document length (number of tokens) in the ${\tt DreamDrug}$ dataset

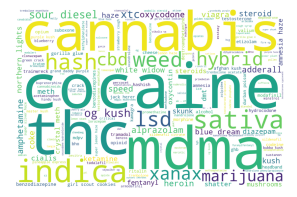


Figure 7: Word-cloud of the most frequent entities in the ${\tt DreamDrug}$ dataset

C Annotation guidelines

The following pages show the long version of the annotation guidelines as they were presented to the annotators of Appen and to AMT workers clicking on the *Instructions* button on the interface shown in Figure 3 (Section 3.3).

Annotator Guidelines - Appen

Instructions -

Overview

In this task you will use the text annotation tool to highlight drugs in texts from the Darknet. This highlighting means you label pieces of text (called "token") in product descriptions of Darknet Markets. The items have a name and a description. We want you to label all DRUGS you can find in the description.

HOW TO DO THIS?

1. Read the Context below the text to get an idea of which items could be contained in the text



- 2. Read the provided drug description and label ALL drugs --> Not just the one in the context.
- 3. Are you unsure about what is a drug? No problem Click on the info of the "Drug" Class or re-open this text.



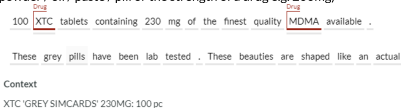
Annotation Guidelines

Label every unique drug name you can find! Typical Drug Examples are:

- Marijuana Strains or types s e.g. "Gorilla Glue", "Indica / Sativa / Hybrid", "Super silver Haze" or "Girl Scout Cookies"
- Types of Drugs or Agents e.g. "Cocaine", "Crystal Meth", "MDMA" or "THC"
- Legal or partial legal drugs e.g. "Alcohol", "Testosterone Enthalate", "Aspirin" or "Viagra"
- Slang names e.g. Coke, Meth, mushrooms, hash, weed

Label ONLY the drug - Descriptions or Adjectives are not part of the drug

(like "Colour, Form or If its powder / oil / paste / pill or the strength of a drug e.g. 200mg)



We want you to label ALL drugs in the text - Pay attention to label drugs with multiple words as one drug. (Strawberry is not drug without amnesia)

If there is a "or '-' contained in the middle of a name label it as well. "Strawberry . Amnesia" or "Strawberry - Amnesia" would also be a drug.



Chemical Details are part of the drug.

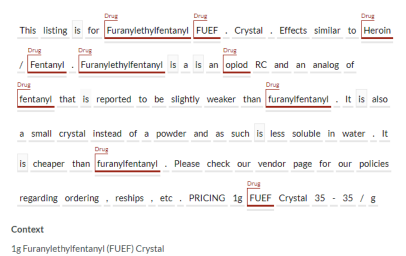
• Sildenfanil Citrate is a single drug and need to be labelled together (not separate)



• Drug Chemicals can also be drugs on their own. - AND Include "-" or "." in the labelling



Abbreviations / Short names of drugs shall be labelled. e.g. XTC, BTH or



Active Agents (The ingredient which makes you high) shall be labelled as drugs. e.g. CBD, THC, Diazepam or DMT

```
100 x. 05g cartridges filled with the highest quality distillate oil , 16 each - ORGANIC - Solvent FREE .

These are the most potent cartridges . Testing at approximately 90 THC , these carts deliver the best vaping experience on the market 500 THC in 1 cartridge . Available flavors . - Strawberry Cheesecake - Guava
```

Official Marketing names ARE drugs

This includes all kinds of marketing names in different countries or commercial marijuana strain names



Drug Classes are Drugs

• Bezondiazipines, Opiates, Steroids, Indica / Sativa / Hybrid - ARE Drugs



Slang names ARE drugs e.g. ganja or dope



Legal Drugs ARE Drugs

• In this example its alcohol, but it could also be Caffeine, Aspirin or tabacco.

```
LIQUID GHB SYRUP THIS GHB IS READY FOR USE - NO PREPPING REQUIRED

STRONG PURE . 99 ,95 PURE BASE GHB . ULTIMATE STEALTH AAA .

DELIVERY GUARANTEED . NEVER COMBINE WITH ALCOHOL CHECK FAQ .

BEFORE USE .

Context

GHB LIQUID SYRUP 250ml GHB (BASF)
```

If a drug is mentioned but doesn't clearly indicate which drug it is, do NOT label it. (e.g. Orange Red bull Pills)

"Fishscale Powder", "These Mitsubishi Pills" or "This strain" will refer to a drug but its not clear, which drug it is exactly. In this case only XTC IS a drug and pills is NOT a drug.



The form / strength / adjectives of the drug shall NOT be labelled.

• At Ecstasy(XTC) it doesn't matter if they are Green Mitsubishi with 200mg

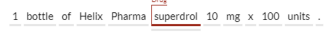


• Or that they have the imprint "OT-20" or "V-3605" on them. Those are NOT drugs.

The Producer or Manufacturer of the drug, is NOT a Drug

This also excludes creations from Drugdealers like Drcokes megasniff or MegaBelt Isolator Vapes. None of those are drugs.

In the example below Helix Pharma is only the producer and superdrol is the drug.



Context

HELIX PHARMA SUPERDROL 10MG X 100 UNITS

Natural chemicals which are part of your body are NOT a drug.

• Dopamin or estrogen do something to your body, but are NOT produced as drug and therefore we DON'T label them.

Processed Items like edibles are not always a drug.

Only the words which clearly refer to a drug are drugs. Brownies, Vapes or Candies are by itself not a drug, only the form!

NO DRUG in the TEXT?? - IMPORTANT

- If you cannot find any drug in the text, just annotate the first word with the tag "NONE" and continue.



HOW TO LABEL? - IMPORTANT

We want to label tokens which belong together as ONE single span:

Two tokens which are part of the same drug e.g. the "Master Yoda" Cannabis strain shall be labelled as single span. (Same goes for OG Kush or Master Kush etc.)



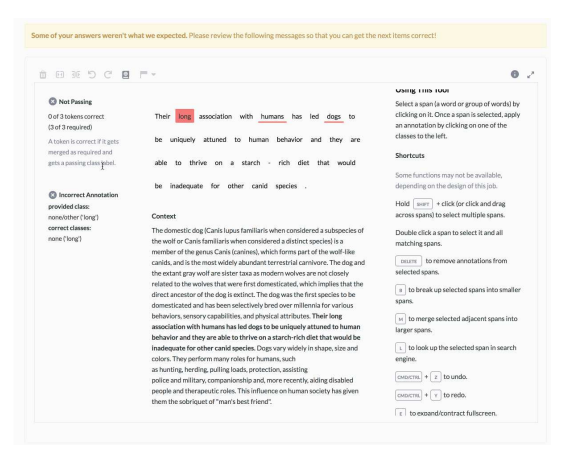
If there is a character (e.g. ". or '-') inside of drug name -- include it in the labelling. The labels are not allowed to be separated!

Are you unsure about what is a drug? No problem - RE-OPEN this wholesome description for details or check the info of the "Drug" Class.

Drug

Test Question Review

When you get a test question wrong, you'll be able to review your mistakes in a special version of the tool. The errors you made will be underlined in red.



Annotator Guidelines Amazon Mechanical Turk

Drug Labelling

In this HIT we want you label all text spans, which clearly point to a specific drug or drug type.

HOW TO DO THIS?

- 1. Read the Context above the text to get an idea of which items could be contained in the text
- 2. Read the provided drug description and label ALL spans which clearly identify a drug
- 3. Are you unsure about what is a drug? We provided a extensive description, just press on "Instructions" and go to "more instructions".

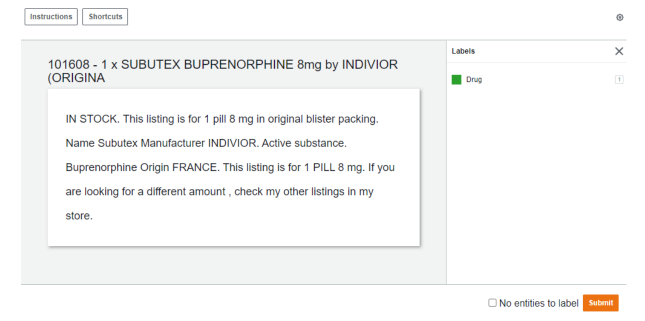
Keep in mind: Each span should Clearly identify a specific drug or drug type!

Practical examples are:

- · Clearly represent a specific drug e.g. hash, meth
- · Chemical Compounds e.g. "3, 4 methylphendiate"
- Short names e.g. XTC
 Agents e.g. THC, CBD or Diazepam
- · Slang drug names: Coke, Fent
- · Drug categories e.g. Indica/hybrid, benzodiazipines, opiates
- Pill imprints e.g. "V-3923"
- Chemical Details which extend the drug e.g. Viagra Sildenafil Citrate
 Legal or partial legal drugs e.g. "Alcohol", "Testosterone Enthalate", "Aspirin" or "Viagra"

DO NOT label tokens like:

- Description/Adjectives like drug strength, colour, form (powder/pill..) or looks etc.
- Form of Drug: e.g. powder, pill, paste Vague references like "this pill", "My Super Chocolate Bars" or "indoor bud"
- Producer of drugs e.g. Pfizer
- · Natural Chemicals in your body e.g. estrogen/dopamin



Note: The long version of the annoation guidelines was accessible over the "Instructions" button. The long version was equal to the long Annotation guidelines from Appen.