Domain-Specific Paraphrase Extraction
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Abstract
The validity of applying paraphrase rules depends on the domain of the text that they are being applied to. We develop a novel method for extracting domain-specific paraphrases. We adapt the bilingual pivoting paraphrase method to bias the training data to be more like our target domain of biology. Our best model results in higher precision while retaining complete recall, giving a 10% relative improvement in AUC.

1 Introduction
Many data-driven paraphrase extraction algorithms have been developed in recent years (Madnani and Dorr, 2010; Androutsopoulos and Malakasiotis, 2010). These algorithms attempt to learn paraphrase rules, where one phrase can be replaced with another phrase which has equivalent meaning in at least some context. Determining whether a paraphrase is appropriate for a specific context is a difficult problem (Bhagat and Hovy, 2013), encompassing issues of syntax (Callison-Burch, 2008), word sense (Apidianaki et al., 2014), and style (Xu et al., 2012; Pavlick and Nenkova, 2015). To date, the question of how domain effects paraphrase has been left unexplored.

Although most paraphrase extraction algorithms attempt to estimate a confidence with which a paraphrase rule might apply, these scores are not differentiated by domain, and instead correspond to the general domain represented by the model’s training data. As illustrated by Table 1, paraphrases that are highly probable in the general domain (e.g. hot = sexy) can be extremely improbable in more specialized domains like biology. Dominant word senses change depending on domain: the verb treat is used in expressions like treat you to dinner in conversational domains versus treat an infection in biology. This domain shift changes the acceptability of its paraphrases.

We address the problem of customizing paraphrase models to specific target domains. We explore the following ideas:

1. We sort sentences in the training corpus based on how well they represent the target domain, and then extract paraphrases from a subsample of the most domain-like data.

2. We improve our domain-specific paraphrases by weighting each training example based on its domain score, instead of treating each example equally.

3. We dramatically improve recall while maintaining precision by combining the subsampled in-domain paraphrase scores with the general-domain paraphrase scores.

2 Background
The paraphrase extraction algorithm that we customize is the bilingual pivoting method (Bannard and Callison-Burch, 2005) that was used to create PPDB, the paraphrase database (Ganitkevitch et al., 2013). To perform the subsampling, we adapt and improve the method that Moore and Lewis (2010) originally developed for domain-specific language models in machine translation.

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2.1 Paraphrase extraction

Paraphrases can be extracted via bilingual pivoting. Intuitively, if two English phrases $e_1$ and $e_2$ translate to the same foreign phrase $f$, we can assume that $e_1$ and $e_2$ have similar meaning, and thus we can "pivot" over $f$ and extract $(e_1, e_2)$ as a paraphrase pair. Since many possible paraphrases are extracted in this way, and since they vary in quality (in PPDB, the verb treat has 1,160 potential paraphrases, including address, handle, deal with, care for, cure him, 'm paying, and 's on the house), it is necessary to assign some measure of confidence to each paraphrase rule. Bannard and Callison-Burch (2005) defined a conditional paraphrase probability $p(e_2|e_1)$ by marginalizing over all shared foreign-language translations $f$:

$$p(e_2|e_1) \approx \sum_f p(e_2|f)p(f|e_1) \tag{1}$$

where $p(e_2|f)$ and $p(f|e_1)$ are translation model probabilities estimated from the bilingual data.

Equation 1 approximates the probability with which $e_1$ can paraphrase as $e_2$, but its estimate inevitably reflects the domain and style of the bilingual training text. If $e_1$ is a polysemous word, the highest probabilities will be assigned to paraphrases of the most frequently occurring sense of $e_1$, and lower probabilities to less frequent senses. This results in inaccurate probability estimates when moving to a domain with different sense distributions compared to the training corpus.

2.2 Sorting by domain specificity

The crux of our method is to train a paraphrase model on data from the same domain as the one in which the paraphrases will be used. In practice, it is unrealistic that we will be able to find bilingual parallel corpora precompiled for each domain of interest. We instead subsample from a large bitext, biasing the sample towards the target domain.

We adapt and extend a method developed by Moore and Lewis (2010) (henceforth M-L), which builds a domain-specific sub-corpus from a large, general-domain corpus. The M-L method assigns a score to each sentence in the large corpus based on two language models, one trained on a sample of target domain text and one trained on the general domain. We want to identify sentences which are similar to our target domain and dissimilar from the general domain. M-L captures this notion using the difference in the cross-entropies according to each language model (LM). That is, for a sentence $s_i$, we compute

$$\sigma_i = H_{tgt}(s_i) - H_{gen}(s_i) \tag{2}$$

where $H_{tgt}$ is the cross-entropy under the in-domain language model and $H_{gen}$ is the cross-entropy under the general domain LM. Cross-entropy is monotonically equivalent to LM perplexity, in which lower scores imply a better fit. Lower $\sigma_i$ signifies greater domain-specificity.

3 Domain-Specific Paraphrases

To apply the M-L method to paraphrasing, we need a sample of in-domain monolingual text. This data is not directly used to extract paraphrases, but instead to train an n-gram LM for the target domain. We compute $\sigma_i$ for the English side of every sentence pair in our bilingual data, using the target domain LM and the general domain LM. We sort the entire bilingual training corpus so that the closer a sentence pair is to the top of the list, the more specific it is to our target domain.

We can apply Bannard and Callison-Burch (2005)'s bilingual pivoting paraphrase extraction algorithm to this sorted bitext in several ways:

1. By choosing a threshold value for $\sigma_i$ and discarding all sentence pairs that fall outside of that threshold, we can extract paraphrases from a subsampled bitext that approximates the target domain.

2. Instead of simply extracting from a subsampled corpus (where each training example is equally weighted), we can weight each training example proportional to $\sigma_i$ when computing the paraphrase scores.

3. We can combine multiple paraphrase scores: one derived from the original corpus and one from the subsample. This has the advantage of producing the full set of paraphrases that can be extracted from the entire bitext.

4 Experimental Conditions

Domain data We evaluate our domain-specific paraphrasing model in the target domain of biology. Our monolingual in-domain data is a combination of text from the GENIA database (Kim et al., 2003) and text from an introductory biology textbook. Our bilingual general-domain data is the $10^9$ word parallel corpus (Callison-Burch et al.,
2009), a collection of French-English parallel data covering a mix of genres from legal text (Steinberger et al., 2006) to movie subtitles (Tiedemann, 2012). We use 5-gram language models with Kneser-Ney discounting (Heafield et al., 2013).

**Evaluation** We measure the precision and recall of paraphrase pairs produced by each of our models by collecting human judgments of what paraphrases are acceptable in sentences drawn from the target domain and in sentences drawn from the general domain. We sample 15K sentences from our biology data, and 10K general-domain sentences from Wikipedia. We select a phrase from each sentence, and show the list of candidate paraphrases to 5 human judges. Judges make a binary decision about whether each paraphrase is appropriate given the domain-specific context. We consider a paraphrase rule to be good in the domain if it is judged to be good in at least one context by the majority of judges. See Supplementary Materials for a detailed description of our methodology.

**Baseline** We run normal paraphrase extraction over the entire 10^9 word parallel corpus (which has 828M words on the English side) without any attempt to bias it toward the target domain. We refer this system as General.

**Subsampling** After sorting the 10^9 word parallel corpus by Equation 2, we chose several threshold values for subsampling, keeping only top-ranked \( \tau \) words of the bitext. We train models on for several values of \( \tau \) (1.5M, 7M, 35M, and 166M words). We refer to these model as \( M-L, \tau \).

**M-L, Change Point** We test a model where \( \tau \) is set at the point where \( \sigma_i \) switches from negative to positive. This includes all sentences which look more like the target domain than the general. This threshold is equivalent to sampling 20M words.

**Weighted Counts** Instead of weighting each subsampled sentence equally, we test a novel extension of M-L in which we weight each sentence proportional to \( \sigma_i \) when computing the paraphrase scores. Specifically, we weight the counting during the bilingual pivoting so that

the target domain to set the regression weights. This tuning set is disjoint from the test set.

**5 Experimental Results**

What is the effect of subsampling? Figure 1 compares the precision and recall of the different subsampling methods against the baseline of training on everything, when they are evaluated on manually labeled test paraphrases from the biology domain. All of subsampled models have a higher precision than the baseline General model, except for the largest of the subsampled models (which was trained on sentence pairs with 166M words - many of which are more like the general domain than the biology domain).

The subsampled models have reduced recall since many of the paraphrases that occur in the full 10^9 word bilingual training corpus do not occur in the subsamples. As we increase \( \tau \) we improve recall at the expense of precision, since we are including training data that is less and less like our target domain. The highest precision model based on the vanilla M-L method is M-L Change Point, which sets the subsample size to include exactly those sentence pairs that look more like the target domain than the general domain.

Our novel extension of the M-L model (M-L Weighted) provides further improvements. Here, we weight each sentence pair in the bilingual training corpus proportional to \( \sigma_i \) when computing the paraphrase scores. Specifically, we weight the counting during the bilingual pivoting so that

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\(^1\)The candidates paraphrases constitute the full set of paraphrases that can be extracted from our training corpus.
rather than each occurrence counting as 1, each occurrence counts as the ratio of the sentence’s cross-entropies: \( \frac{H_{\text{gen}}}{H_{\text{tgt}}} \). The top-ranked sentence pairs receive an exaggerated count of 52, while the bottom ones receive a tiny factional count of 0.0068. Thus, paraphrases extracted from sentence pairs that are unlike the biology domain receive very low scores. This allows us to achieve higher recall by incorporating more training data, while also improving the precision.

**What is the benefit of combining models?** We have demonstrated that extracting paraphrases from subsampled data results in higher precision domain-specific paraphrases. But these models extract only a fraction of the paraphrases that are extracted by a general model trained on the full bitext, resulting in a lower recall.

We dramatically improve the recall of our domain-specific models by combining the small subsampled models with the large general-domain model. We use binary logistic regression to combine the \( p(e_2 | e_1) \) estimate of the general model with that of each domain-specific model. Figure 2(a) shows that we are able to extend the recall of our domain-specific models to match the recall of the full general-domain model. The precision scores remain higher for the domain-specific models. Our novel M-L Weighted model performs the best. Table 3 gives the area under the curve (AUC). The best combination improves AUC by more than 4 points absolute (>10 points relative) in the biology domain. Table 2 provides examples of paraphrases extracted using our domain-specific model for biology versus the baseline model.

**6 Related Work**

Domain-specific paraphrasing has not received previous attention, but there is relevant prior work on domain-specific machine translation (MT). We build on the Moore-Lewis method, which has been used for language models (Moore and Lewis, 2010) and translation models (Axelrod et al., 2011). Similar methods use LM perplexity to rank sentences (Gao et al., 2002; Yasuda et al., 2008), rather than the difference in cross-entropy. Within MT, Foster and Kuhn (2007) used log-linear weightings of translation probabilities to combine models trained in different domains, as we do here. Relevant to our proposed method of
fractional counting, (Madnani et al., 2007) used introduced a count-centric approach to paraphrase probability estimation. Matsoukas et al. (2009) and Foster et al. (2010) explored weighted training sentences for MT, but set weights discriminatively based on sentence-level features.

7 Conclusion

We have discussed the new problem of extracting domain-specific paraphrases. We adapt a method from machine translation to the task of learning domain-biased paraphrases from bilingual corpora. We introduce two novel extensions to this method. Our best domain-specific model dramatically improves paraphrase quality for the target domain.

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