

# Words and Echoes: Assessing and Mitigating the Non-Randomness Problem in Word Frequency Distribution Modeling

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## Abstract

Frequency distribution models tuned to words and other linguistic events can predict the number of distinct types and their frequency distribution in samples of arbitrary sizes. We conduct, for the first time, a rigorous evaluation of these models based on cross-validation and separation of training and test data. Our experiments reveal that the prediction accuracy of the models is marred by serious overfitting problems, due to violations of the random sampling assumption in corpus data. We then propose a simple pre-processing method to alleviate such non-randomness problems. Further evaluation confirms the effectiveness of the method, which compares favourably to more complex correction techniques.

## 1 Introduction

Large-Number-of-Rare-Events (LNRE) models (Baayen, 2001) are a class of specialized statistical models that allow us to estimate the characteristics of the distribution of type probabilities in type-rich linguistic populations (such as words) from limited samples (our corpora). They also allow us to extrapolate quantities such as vocabulary size (the number of distinct types) and the number of hapaxes (types occurring just once) beyond a given corpus or make predictions for completely unseen data from the same underlying population.

LNRE models have applications in theoretical linguistics, e.g. for comparing the type richness of morphological or syntactic processes that are attested to

different degrees in the data (Baayen, 1992). Consider for example a very common prefix such as *re-* and a rather rare prefix such as *meta-*. With LNRE models we can answer questions such as: If we could obtain as many tokens of *meta-* as we have of *re-*, would we also see as many distinct types? In other words, is the prefix *meta-* as productive as the prefix *re-*? Practical NLP applications, on the other hand, include estimating how many out-of-vocabulary words we will encounter given a lexicon of a certain size, or making informed guesses about type counts in very large data sets (e.g., *how many typos are there on the Internet?*)

In this paper, after introducing LNRE models (Section 2), we present an evaluation of their performance based on separate training and test data as well as cross-validation (Section 3). As far as we know, this is the first time that such a rigorous evaluation has been conducted. The results show how evaluating on the training set, a common strategy in LNRE research, favours models that overfit the training data and perform poorly on unseen data. They also confirm the observation by Evert and Baroni (2006) that current LNRE models achieve only unsatisfactory prediction accuracy, and this is the issue we turn to in the second part of the paper (Section 4). Having identified the violation of the random sampling assumption by real-world data as one of the main factors affecting the quality of the models, we present a new approach to alleviating non-randomness problems. Further evaluation shows our solution to outperform Baayen's (2001) partition-adjustment method, the former state-of-the-art in non-randomness correction. Section 5 concludes by

pointing out directions for future work.

## 2 LNRE models

Baayen (2001) introduces a family of models for Zipf-like frequency distributions of linguistic populations, referred to as *LNRE models*. Such a linguistic population is formally described by a finite or countably infinite set of types  $\omega_i$  and their occurrence probabilities  $\pi_i$ . Word frequency models are not concerned with the probabilities (i.e., relative frequencies) of specific individual types, but rather the overall distribution of these probabilities.

Numbering the types in order of decreasing probability ( $\pi_1 \geq \pi_2 \geq \pi_3 \geq \dots$ , called a population *Zipf ranking*), we can specify a LNRE model for their distribution as a function that computes  $\pi_i$  from the Zipf rank  $i$  of  $\omega_i$ . For instance, the Zipf-Mandelbrot law<sup>1</sup> is defined by the equation

$$\pi_i = \frac{C}{(i + b)^a} \quad (1)$$

with parameters  $a > 1$  and  $b > 0$ . It is mathematically more convenient to formulate LNRE models in terms of a type density function  $g(\pi)$  on the interval  $\pi \in [0, 1]$ , such that

$$\int_A^B g(\pi) d\pi \quad (2)$$

is the (approximate) number of types  $\omega_i$  with  $A \leq \pi_i \leq B$ . Evert (2004) shows that Zipf-Mandelbrot corresponds to a type density of the form

$$g(\pi) := \begin{cases} C \cdot \pi^{-\alpha-1} & A \leq \pi \leq B \\ 0 & \text{otherwise} \end{cases} \quad (3)$$

with parameters  $0 < \alpha < 1$  and  $0 \leq A < B$ .<sup>2</sup> Models that are formulated in terms of such a type density  $g$  have many direct applications (e.g. using  $g$  as a Bayesian prior), and we refer to them as *proper LNRE models*.

Assuming that a corpus of  $N$  tokens is a random sample from such a population, we can make predictions about lexical statistics such as the number

<sup>1</sup>The Zipf-Mandelbrot law is an extension of Zipf's law (which has  $a = 1$  and  $b = 0$ ). While the latter originally refers to type frequencies in a given sample, the Zipf-Mandelbrot law is formulated for type probabilities in a population.

<sup>2</sup>In this equation,  $C$  is a normalizing constant required in order to ensure  $\int_0^1 \pi g(\pi) d\pi = 1$ , the equivalent of  $\sum_i \pi_i = 1$ .

$V(N)$  of different types in the corpus (the *vocabulary size*), the number  $V_1(N)$  of *hapax legomena* (types occurring just once), as well as the further distribution of type frequencies  $V_m(N)$ . Since the precise values would be different from sample to sample, the model predictions are given by expectations  $E[V(N)]$  and  $E[V_m(N)]$ , which can be computed with relative ease from the type density function  $g$ .

By comparing expected and observed values of  $V$  and  $V_m$  (for the lowest frequency ranks, usually up to  $m = 15$ ), the parameters of a LNRE model can be estimated (we refer to this as *training* the model), allowing inferences about the population (such as the total number of types in the population) as well as further applications of the estimated type density (e.g. for Good-Turing smoothing). Since we can calculate expected values for samples of arbitrary size  $N$ , we can use the trained model to predict how many new types would be seen in a larger corpus, how many hapaxes there would be, etc. This kind of vocabulary growth *extrapolation* has become one of the most important applications of LNRE models in linguistics and NLP.

A detailed account of the mathematics of LNRE models can be found in Baayen (2001, Ch. 2). Baayen describes two LNRE models, *lognormal* and *GIGP*, as well as several other approaches (including a version of *Zipf's law* and the *Yule-Simon* model) that are not based on a type density and hence do not qualify as proper LNRE models. Two LNRE models based on Zipf's law, *ZM* and *fZM*, are introduced by Evert (2004).

In the following, we will only consider proper LNRE models because of their considerably greater utility, and because their performance in extrapolation tasks appears to be better than, or at least comparable to, the other models (Evert and Baroni, 2006). In addition, we exclude the lognormal model because of its computational complexity and numerical instability.<sup>3</sup> In initial evaluation experiments, the performance of lognormal was also inferior to the remaining three models (ZM, fZM and GIGP). Note that ZM is the most simplistic model, with only 2 parameters and assuming an infinite population vocabulary, while fZM and GIGP have 3 parameters

<sup>3</sup>There are no closed form equations for the expectations of the lognormal model, which have to be calculated by numerical integration.

and can model populations of different sizes.

### 3 Evaluation of LNRE models

LNRE models are traditionally evaluated by looking at how well expected values generated by them fit empirical counts extracted from the same dataset used for parameter estimation, often by visual inspection of differences between observed and predicted data in plots. More rigorously, Baayen (2001) and Evert (2004) compare the frequency distribution observed in the training set to the one predicted by the model with a multivariate chi-squared test. As we will show below, evaluating standard LNRE models on the same data that were used to estimate their parameters favours overfitting, which results in poor performance on unseen data.

Evert and Baroni (2006) attempt, for the first time, to evaluate LNRE models on unseen data. However, rather than splitting the data into separate training and test sets, they evaluate the models in an extrapolation setting, where the parameters of the model are estimated on a *subset* of the data used for testing. Evert and Baroni do not attempt to cross-validate the results, and they do not provide a quantitative evaluation, relying instead on visual inspection of empirical and observed vocabulary growth curves.

#### 3.1 Data and procedure

We ran our experiments with three corpora in different languages and representing different textual typologies: the British National Corpus (BNC), a “balanced” corpus of British English of about 100 million tokens illustrating different communicative settings, genres and topics; the deWaC corpus, a Web-crawled corpus of about 1.5 billion German words; and the la Repubblica corpus, an Italian newspaper corpus of about 380 million words.<sup>4</sup>

From each corpus, we extracted 20 non-overlapping samples of randomly selected documents, amounting to a total of 4 million tokens each (punctuation marks and entirely non-alphabetical tokens were removed before sampling, and all words were converted to lowercase). Each of these samples was then split into a training set of 1 million tokens (the *training size*  $N_0$ ) and a test set of 3 million

tokens. The documents in the la Repubblica samples were ordered chronologically before splitting, to simulate a typical scenario arising when working with newspaper data, where the data available for training precede, chronologically, the data one wants to generalize to.

We estimate parameters of the ZM, fZM and GIGP models on each training set, using the zipfR toolkit.<sup>5</sup> The models are then used to predict the expected number of distinct types, i.e., vocabulary size  $V$ , at sample sizes of 1, 2 and 3 million tokens, equivalent to 1, 2 and 3 times the size of the training set (we refer to these as the *prediction sizes*  $N_0$ ,  $2N_0$  and  $3N_0$ , respectively). Finally, the expected vocabulary size  $E[V(N)]$  is compared to the observed value  $V(N)$  in the test set for  $N = N_0$ ,  $N = 2N_0$  and  $N = 3N_0$ . We also look at  $V_1(N)$ , the number of hapax legomena, in the same way.

Our main focus is  $V$  prediction, since this is by far the most useful measure in practical applications, where we are typically interested in knowing how many types (or how many types belonging to a certain category) we will see as our sample size increases (How many typos are there on the Web? How many types with prefix *meta-* would we see if we had as many types of *meta-* as we have of *re-*?) Hapax legomena counts, on the other hand, play a central role in quantifying morphological productivity (Baayen, 1992) and they give us a first insight into how good the models are at predicting frequency distributions, besides vocabulary size (as we will see, a model’s success in predicting  $V$  does not necessarily imply that the model is also capturing the right frequency distribution).

For all models, corpora and prediction sizes, goodness-of-fit of the model on the training set is measured with a multivariate chi-squared test (Baayen, 2001, 118-122). Performance of the models in prediction of  $V$  is assessed via *relative error*, computed for each of the 20 samples from a corpus and the 3 prediction sizes as follows:

$$e = \frac{E[V(N)] - V(N)}{V(N)}$$

where  $N = k \cdot N_0$  is the prediction size (for  $k = 1, 2, 3$ ),  $V(N)$  is the observed  $V$  in the relevant test

<sup>4</sup>See [www.natcorp.ox.ac.uk](http://www.natcorp.ox.ac.uk), <http://wacky.sslmit.unibo.it> and [http://sslmit.unibo.it/la\\_repubblica](http://sslmit.unibo.it/la_repubblica)

<sup>5</sup><http://purl.org/stefan.evert/zipfR>

set at size  $N$ , and  $E[V(N)]$  is the corresponding expected  $V$  predicted by a model.<sup>6</sup>

For each corpus and prediction size we obtain 20 values  $e_i$  (viz.,  $e_1, \dots, e_{20}$ ). As a summary measure, we report the square root of the mean square relative error (rMSE) calculated according to

$$\sqrt{\text{rMSE}} = \sqrt{\frac{1}{20} \cdot \sum_{i=1}^{20} (e_i)^2}$$

This gives us an overall assessment of prediction accuracy (we take the square root to obtain values on the same scale as relative errors, and thus easier to interpret). We complement rMSEs with reports on the average relative error (indicating whether there is a systematic under- or overestimation bias) and its asymptotic 95% confidence intervals, based on the empirical standard deviation of the  $e_i$  across the 20 trials (the confidence intervals are usually somewhat larger than the actual range of values found in the experiments, so they should be seen as “pessimistic estimates” of the actual variance).

### 3.2 Results

The panels of Figure 1 report rMSE values for the 3 corpora and for each prediction size. For now, we focus on the first 3 histograms of each panel, that present rMSEs for the 3 LNRE models introduced above: ZM, fZM and GIGP (the remaining histograms will be discussed later).<sup>7</sup>

For all corpora and all extrapolation sizes beyond  $N_0$ , the simple ZM model outperforms the more sophisticated fZM and GIGP models (which seem to be very similar to each other). Even at the largest prediction size of  $3N_0$ , ZM’s rMSE is well below 10%, whereas the other models have, in the worst case (BNC  $3N_0$ ), a rMSE above 15%. Figure 2 presents plots of average relative error and its empirical confidence intervals (again, focus for now on the ZM, fZM and GIGP results; the rest of the figure is discussed later). We see that the poor performance

<sup>6</sup>We normalize by  $V(N)$  rather than (a function of)  $E[V(N)]$  because in the latter case we would favour models that overestimate  $V$ , compared to ones that are equally “close” to the correct value but underestimate  $V$ .

<sup>7</sup>A table with the full numerical results is available upon request; we find, however, that graphical summaries such as those presented in this paper make the results easier to interpret.

of fZM and GIGP is due to their tendency to underestimate the true vocabulary size  $V$ , while variance is comparable across models.

The rMSEs of  $V_1$  prediction are reported in Figure 3.  $V_1$  prediction performance is poorer across the board, and ZM is no longer outperforming the other models. For space reasons, we do not present relative error and variance plots for  $V_1$ , but the general trends are the same observed for  $V$ , except that the bias of ZM towards  $V_1$  overestimation is much clearer than for  $V$ .

Interestingly, goodness-of-fit on the training data is not a good predictor of  $V$  and  $V_1$  prediction performance on unseen data. This is shown in Figure 4, which plots rMSE for prediction of  $V$  against goodness-of-fit (quantified by multivariate  $X^2$  on the training set, as discussed above) for all corpora and LNRE models at the  $3N_0$  prediction size (but the same patterns emerge at other prediction sizes and with  $V_1$ ). The larger  $X^2$ , the poorer the training set fit; the larger rMSE, the worse the prediction. Thus, ideally, we should see a positive correlation between  $X^2$  and rMSE. Focusing for now on the circles (pinpointing the ZM, fZM and GIGP models), we see that there is instead a *negative* correlation between goodness of fit on the training set and quality of prediction on unseen data.<sup>8</sup>

First, these results indicate that, if we take goodness of fit on the training set as a criterion for choosing the best model (as done by Baayen and Evert), we end up selecting the *worst* model for actual prediction tasks. This is, we believe, a very strong case for applying the split train-test cross-validation method used in other areas of statistical NLP to frequency distribution modeling. Second, the data suggest that the more sophisticated models are *overfitting* the training set, leading to poorer performance than the simpler ZM on unseen data. We turn now to what we think is the main cause for this overfitting.

## 4 Non-randomness and echoes

The results in the previous section indicate that the  $V$ s predicted by LNRE models are at best “ballpark estimates” (and  $V_1$  predictions, with a relative error that is often above 20%, do not even qualify as plau-

<sup>8</sup>With correlation coefficients of  $r < -.8$ , significant at the 0.01 level despite the small sample size.

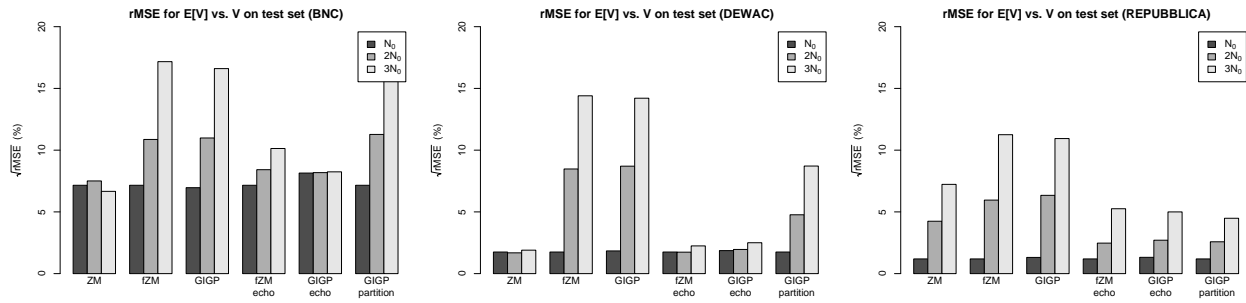


Figure 1: rMSEs of predicted  $V$  on the BNC, deWAC and la Repubblica data-sets

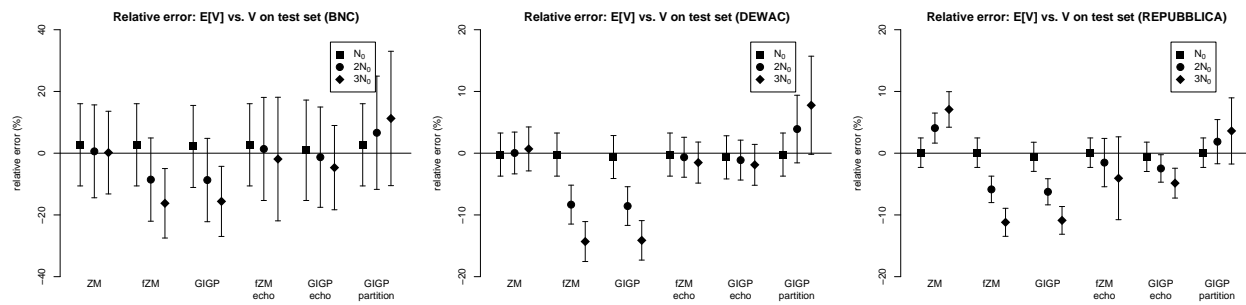


Figure 2: Average relative errors and asymptotic 95% confidence intervals of  $V$  prediction on BNC, deWAC and la Repubblica data-sets

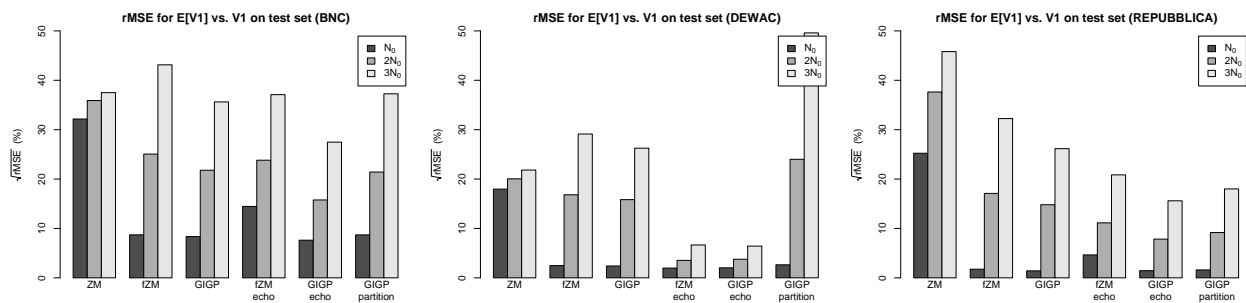


Figure 3: rMSEs of predicted  $V_1$  on the BNC, deWAC and la Repubblica data-sets

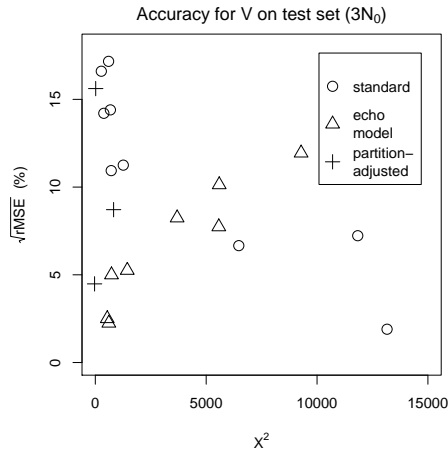


Figure 4: Correlation between  $X^2$  and  $V$  prediction rMSE across corpora and models

sible ballpark estimates). Although such rough estimates might be more than adequate for many practical applications, is it possible to further improve the quality of LNRE predictions?

A major factor hampering prediction quality is that real texts massively violate the *randomness assumption* made in LNRE modeling: words, rather obviously, are not picked at random on the basis of their population probability (Evert and Baroni, 2006; Baayen, 2001). The topic-driven “clumpiness” of low frequency content words reduces the number of hapax legomena and other rare events used to estimate the parameters of LNRE models, leading the models to underestimate the type richness of the population. Interestingly (but unsurprisingly), ZM with its assumption of an infinite population, is less prone to this effect, and thus it has a better prediction performance than the more sophisticated fZM and GIGP models, despite its poor goodness-of-fit.

The effect of non-randomness is illustrated very clearly for the BNC (but the same could be shown for the other corpora) by Figure 5, a comparison of rMSE for prediction of  $V$  from our experiments above to results obtained on versions of the BNC samples with words scrambled in random order, thus forcibly removing non-randomness effects. We see from this figure that the performance of both fZM and GIGP improves dramatically when they are trained and tested on randomized sequences of

words. Interestingly, randomization has instead a *negative* effect on ZM performance.

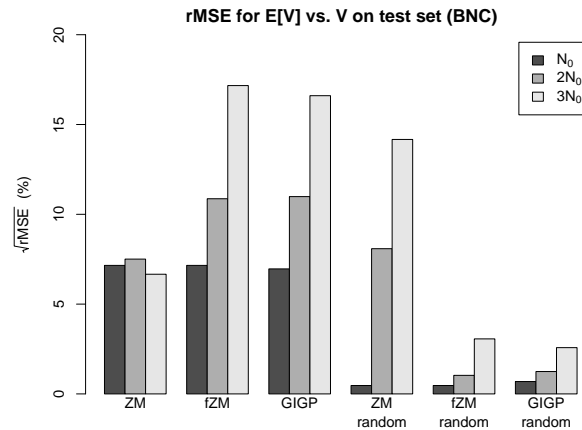


Figure 5: rMSEs of predicted  $V$  on unmodified vs. randomized versions of the BNC sets

#### 4.1 Previous approaches to non-randomness

While non-randomness is widely acknowledged as a serious problem for the statistical analysis of corpus data, very few authors have suggested correction strategies. The key problem of non-random data seems to be that the occurrence frequencies of a type in different documents do not follow the binomial distribution assumed by random sampling models. One approach is therefore to model this distribution explicitly, replacing the binomial with its single parameter  $\pi$  by a more complex distribution that has additional parameters (Church and Gale, 1995; Katz, 1996). However, these distributions are currently not applicable to LNRE modeling, which is based on the overall frequencies of types in a corpus rather than their frequencies in individual documents. The overall frequencies can only be calculated by summation over all documents in the corpus, resulting in a mathematically and numerically intractable model. In addition, the type density  $g(\pi)$  would have to be extended to a multi-dimensional function, requiring a large number of parameters to be estimated from the data.

Baayen (2001) suggests a different approach, which partitions the population into “normal” types that satisfy the random sampling assumption, and “totally underdispersed” types, which are assumed to concentrate all occurrences in the corpus into a

single “burst”. Using a standard LNRE model for the normal part of the population and a simple linear growth model for the underdispersed part, adjusted values for  $E[V]$  and  $E[V_m]$  can easily be calculated. These so-called *partition-adjusted* models (which introduce one additional parameter) are thus the only viable models for non-randomness correction in LNRE modeling and have to be considered the state of the art.

## 4.2 Echo adjustment

Rather than making more complex assumptions about the population distribution or the sampling model, we propose that non-randomness should be tackled as a *pre-processing* problem. The issue, we argue, is really with the way we count occurrences of types. The fact that a rare topic-specific word occurs, say, four times in a single document does not make it any less a hapax legomenon for our purposes than if the word occurred once (this is the case, for example, of the word *chondritic* in the BNC, which occurs 4 times, all in the same scientific document).

We operationalize our intuition by proposing that, for our purposes, each content word (at least each rare, topic-specific content word) occurs maximally once in a document, and all other instances of that word in the document are really instances of a special “anaphoric” type, whose function is that of “echoing” the content words in the document. Thus, in the BNC document mentioned above, the word *chondritic* is counted only once, whereas the other three occurrences are considered as tokens of the *echo* type. Thus, we are counting what in the information retrieval literature is known as *document frequencies*. Intuitively, these are less susceptible to topical clumpiness effects than plain token frequencies. However, by replacing repeated words with echo tokens, we can stick to a sampling model based on random word token sampling (rather than document sampling), so that the LNRE models can be applied “as is” to echo-adjusted corpora.

Echo-adjustment does not affect the sample size  $N$  nor the vocabulary size  $V$ , making the interpretation of results obtained with echo-adjusted models entirely straightforward.  $N$  does not change because repeated types are replaced with echo tokens, not deleted.  $V$  does not change because only repeated types are replaced. Thus, no type present in

the original corpus disappears (more precisely,  $V$  increases by 1 because of the addition of the echo type, but given the large size of  $V$  this can be ignored for all practical purposes). Thus, the expected  $V$  computed for a specified sample size  $N$  with a model trained on an echo-adjusted corpus can be directly compared to observed values at  $N$ , and to predictions made for the same  $N$  by models trained on an unprocessed corpus. The same is not true for the prediction of the frequency distribution, where, for the same  $N$ , echo-based models predict the distribution of *document* frequencies.

We are proposing echoes as a model for the usage of (rare) content words. It would be difficult to decide where the boundary is between topical words that are inserted once in a discourse and then anaphorically modulated and “general-purpose” words that constitute the frame of the discourse and can occur multiple times. Luckily, we do not have to make this decision when estimating a LNRE model, since model fitting is based on the distribution of the lowest frequencies. For example, with the default zipfR model fitting setting, only the lowest 15 spectrum elements are used to fit the models. For any reasonably sized corpus, it is unlikely that function words and common content words will occur in less than 16 documents, and thus their distribution will be irrelevant for model fitting. Thus, we can ignore the issue of what is the boundary between topical words to be echo-adjusted and general words, as long as we can be confident that the set of lowest frequency words used for model fitting belong to the topical set.<sup>9</sup> This makes practical echo-adjustment extremely simple, since all we have to do is to replace all repetitions of a word in the same document with echo tokens, and estimate the parameters of a plain LNRE model with the resulting version of the training corpus.

## 4.3 Experiments with echo adjustment

Using the same training and test sets as in Section 3.1, we train the partition-adjusted GIGP model

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<sup>9</sup>The issue becomes more delicate if we want to predict the frequency spectrum rather than  $V$ , since a model trained on echo-adjusted data will predict echo-adjusted frequencies across the board. However, in many theoretical and practical settings only the lowest frequency spectrum elements are of interest, where, again, it is safe to assume that words are highly topic-dependent, and echo-adjustment is appropriate.

implemented in the LEXSTATS toolkit (Baayen, 2001). We estimate the parameters of echo-adjusted ZM, fZM and GIGP models on versions of the training corpora that have been pre-processed as described above. The performance of the models is evaluated with the same measures as in Section 3.1 (for prediction of  $V_1$ , echo-adjusted versions of the test data are used).

Figure 1 reports the performance of the echo-adjusted fZM and GIGP models and of partition-adjusted GIGP (echo-adjusted ZM performed systematically much worse than the other echo-adjusted models and typically worse than uncorrected ZM, and it is not reported in the figure). Both correction methods lead to a dramatic improvement, bringing the prediction performance of fZM and GIGP to levels comparable to ZM (with the latter outperforming the corrected models on the BNC, but being outperformed on la Repubblica). Moreover, echo-adjusted GIGP is as good as partitioned GIGP on la Repubblica, and better on both BNC and deWaC, suggesting that the much simpler echo-adjustment method is at least as good and probably better than Baayen's partitioning. The mean error and confidence interval plots in Figure 2 show that the echo-adjusted models have a much weaker underestimation bias than the corresponding unadjusted models, and are comparable to, if not better than, ZM (although they might have a tendency to display more variance, as clearly illustrated by the performance of echo-adjusted fZM on la Repubblica at  $3N_0$  prediction size). Finally, the echo-adjusted models clearly stand out with respect to ZM when it comes to  $V_1$  prediction (Figure 3), indicating that echo-adjusted versions of the more sophisticated fZM and GIGP models should be the focus of future work on improving prediction of the full frequency distribution, rather than plain ZM. Moreover, echo-adjusted GIGP is outperforming partitioned GIGP, and emerging as the best model overall.<sup>10</sup> Reassuringly, for the echoed models there is a very strong *positive* correlation between goodness-of-fit on the training set and quality of prediction, as illustrated for  $V$  prediction at  $3N_0$  by the triangles in Figure 4 (again, the patterns in this

<sup>10</sup>In looking at the  $V_1$  data, it must be kept in mind, however, that  $V_1$  has a different interpretation when predicted by echo-adjusted models, i.e., it is the number of *document-based* hapaxes, the number of types that occur in one document only.

figure represent the general trend for echo-adjusted models found in all settings).<sup>11</sup> This indicates that the over-fitting problem has been resolved, and for echo-adjusted models goodness-of-fit on the training set is a reliable indicator of prediction accuracy.

## 5 Conclusion

Despite the encouraging results we reported, much work, of course, remains to be done. Even with the echo-adjusted models, prediction of  $V_1$  suffers from large errors and prediction of  $V$  quickly deteriorates with increasing prediction size  $N$ . If the models' estimates for 3 times the size of the training set have acceptable errors of around 5%, for many applications we might want to extrapolate to  $100N_0$  or more (recall the example of estimating type counts for the entire Web). Moreover, echo-adjusted models make predictions pertaining to the distribution of document frequencies, rather than plain token frequencies. The full implications of this remain to be investigated. Finally, future work should systematically explore to what extent different textual typologies are affected by the non-randomness problem (notice, e.g., that non-randomness seems to be a greater problem for the BNC than for the more uniform la Repubblica corpus).

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<sup>11</sup>With significant correlation coefficients of  $r = .76$  for  $2N_0$  ( $p < 0.05$ ) and  $r = .94$  for  $3N_0$  ( $p \ll 0.01$ ).