In this paper, we investigate the problem of building a metaDictionary which reflects variance in a language in space and time. This is done by exploiting the annotation of electronic dictionaries; for obtaining a fine grain annotation, we use declarative parsing techniques.

We are developing a graphical tool for decomposing and annotating the entries/morphemes of an electronic dictionary. The goals of the segmentation are to obtain a list of basic morphemes, to construct a network showing which morphemes can be combined to obtain complex morphemes, and to analyze variance in space and time. Based on the metaDictionary and the annotated morpheme decomposition, a network analysis of morpheme decompositions becomes possible, which can take variance and grammatical properties into account.

We can manage and analyze grids of dictionaries and huge text corpora: dictionaries are especially interesting for our analyses, since they contain settled morpheme cores, and the text corpora document which morphemes can be combined lexicographically. Our dictionary–based morphological analysis has to be tested in the context of huge network corpus data best accessible via grid structures.
1. Introduction

Information and knowledge can be encoded by combining elementary basic components according to special rules. From this point of view, genomes and language code may have structural properties in common which contribute substantially to evolution on the one hand and to the change of language on the other hand. Research in this field with a focus on interdependencies between science and the humanities seems promising. In bioinformatics, research data on genomes have already been generated and are publicly available. On a much smaller scale, we try to build up a comparable research environment in the humanities by collecting empirical data in the field of language change in German within the last 500 years and beyond, based on dictionary information and to be used for the research into mutual relations between data structures and data properties in genomes and language. Our team project combines bioinformatics, informatics, philology and corpus linguistics and has been funded by the German Federal Ministry of Education and Research since 2008 [11].
The starting point of the project is a collection of digitized dictionaries made publicly available by our project partners at the university of Trier (Trierer Wörterbuchnetz [10]): synchronic dictionaries like the Middle High German Dictionaries (Benecke/Müller/Zarncke and Lexer), early High German Dictionaries around 1800 (Adelung and Campe), a selection of dictionaries on regional dialects (Rheinisches, Pfälzisches and Luxemburger Wörterbuch, Wörterbuch der elsässischen bzw. der deutsch–lothringischen Mundarten) and supporting materials of the diachronic Deutsches Wörterbuch by Jacob and Wilhelm Grimm.

For modern German, we can use Klappenbach/Steinitz, Wörterbuch der deutschen Gegenwartssprache (WDG), digitized by the Academy of Sciences of Berlin and Brandenburg [4]. Based on the broad and representative data base, the goal is to develop and test methods and algorithms for detecting and understanding variance. Ideally, biological processes could then be modelled using concepts detected by analyzing language structures, and, vice versa, variance in language might be described using models developed for bioinformatics.

The rest of this paper is organized as follows: In Section 2, we outline some project objectives investigating variance in language and genome. Section 3 explains a prerequisite of the project: how digitized print dictionaries can be annotated in Tei using a declarative grammar formalism. Section 4 presents a tool for analyzing the morphological structure of dictionary entries and annotating their parts. Some conclusions are given in Section 5.

2. Variance in Language and Genome

Our project goals are the compilation of a metaDictionary – based on a fine grain annotation of dictionaries – and the computation of network structures and properties for the comparison with genomes – based on morphological analyses of dictionary entries into basic lexical units (base morphemes). As early as 1934, Karl Bühler discussed the importance of these units for the understanding of variety in language in his language theory and gave an estimate on the dimensions for German – more than 2000 units –, setting out with a morpheme count of 30 pages of Goethe’s novel Wahlverwandtschaften [1]. In this section, we will briefly sketch the project goals and the techniques from computer science that we are using; more details about the techniques will be given in the subsequent sections.

2.1 The metaDictionary

The usage of words in languages varies in space and time, cf. Figure 1. A metaDictionary summarizes different representations of the same lexical units, taken from the dictionaries spread in space and time, in lists of metaLemmas, cf. Figure 2. The metaLemma uses present–day standard German as keyword, with special provision made for units that are no longer used today.

2.2 Network Analysis of Morpheme Decompositions

The project partner from bioinformatics is interested in comparing the combinability of (basic) morphemes in words, cf. Figure 3 with the combinability of amino sequences
in genomes. Preliminary tests have shown that the corresponding networks have similar properties. This could be due to the fact that the generative processes behind the evolution of language and genome might be the comparable.

2.3 Techniques from Computer Science

We want to develop a generic e–Infrastructure for analyzing the inhomogenous dictionary entries of 18th, 19th and 20th century lexicographers and for transforming the information to a kind of baseline encoding keeping as much of the valuable dictionary information as possible, e.g., part of speech, gender, and inflectional detail, encoded in lots
of different patterns. The annotated dictionary data are stored in an XML data format conformant with the TEI P5 Guidelines for Electronic Text Encoding and Interchange [8] for researching into variance comparable to genome structures, and they will be publicly available for others researchers.

We use declarative programming in PROLOG for parsing, querying, and transforming the linguistic data [3, 6, 7]. There exists quite a variety of well-known declarative languages for different areas: SQL for relational databases, XQuery and XSLT for XML processing, PROLOG for declarative programming, and rules for decision support systems and grammars. Their advantages are, that they are compact, rapidly programmable, clear, less error-prone, and flexibly extensible. We have tested declarative knowledge extraction by parsing dictionary entries with definite clause grammars (DCG) on the Adelung Dictionary and – within the framework of the TextGrid–Project [9] – on the Campe Dictionary [2] for a fine grain lexicographic analysis. For compiling the metaDictionary, we have split the lexemes from the electronic dictionaries into morpheme terms. We have built a PROLOG–based tool for a declarative, rule–based control and annotation of morpheme segmentations, which we test on WDG data at present. We can extend our approach by alignment methods, which can yield a more precise alignment for the metaDictionary.

3. Annotation of Digitized Print Dictionaries in TEI

Researching morpheme structures requires a fine grain analysis and detailed accounting of the basic linguistic units of the German vocabulary. Using dictionary entries as the basis, helps to minimize influences of inflectional specialties and the variance of morphological combination patterns, irrespective of the permanent change of language in history. The period we can cover with our dictionary approach dates from Middle High German (mass of the texts extant from that period date from the 13th and 14th century) up to present German. And we can apply the results of our dictionary analyses in a second step to text
corpora of middle High German texts and early new High German texts – starting with Luther and the mass of German literary texts – available soon in the TextGrid digital repository [9].

For automatically extracting the keyword (lexeme) and some meta-data, such as word class and inflexion, we use a compact grammar formalism, which we have developed using the declarative programming language PROLOG. Parsing of dictionary entries is a complex task, since there is a lot of structural variance; if the desired data could not be extracted, then we can flexibly and quickly adapt the grammar rules without breaking other cases. The parsed dictionary data are stored in a TEI data format for researching into variance comparable to genome structures.

3.1 Encoding Dictionaries in TEI

With our parsing approach, we can obtain a fine grain annotation of electronic dictionaries in a valid TEI format. E.g., the dictionary entry for Aal from Figure 4 is structured as shown in Figure 5:

Figure 4: Entry of an Electronic Dictionary.

Figure 5: Fine Grain Structuring.
XML is a common data format for modelling, managing, and exchanging semi-structured data. There exist powerful query, transformation and update languages for XML. We can also gain performance compared to relational databases.

### 3.2 Grammar–Based Parsing

Parsing with grammars yields a higher precision compared to regular expressions and statistical parsers. We use a DCG extension, called Extended Definite Clause Grammar (EDCG) rules, which is even more compact and directly, generically generates XML [6]. E.g., for generating the entry elements, we can use the following EDCG rules:

```
entry ===> 
  form:[type:lemma], ..., sense.
form:[type:lemma] ===> 
  sequence(*, form:[type:determiner]),
  form:[type:headword].

sense ===> ...
```

The attribute `xml:id` of the `entry` element, which depends on the position of the entry in the dictionary (and on the headword), is determined in a further processing step. EDCGs offer meta-predicates such as sequence: the call `sequence(*, form:[type:determiner])` generates a sequence of zero or more `form` elements.

The *-notion is well-known from extended Backus–Naur form (EBNF) of context free grammars. Compared to DCGs and EBNF, an important extension of EDCGs is, that they generically generate XML. When working with DCGs in PROLOG or EBNF in other tools, this generation has to be coded explicitly and thus blows up the code drastically.

### 4. Annotating Morpheme Decompositions

For annotating the large numbers of dictionary entries (which can exceed 100,000 units), one needs linguistic knowledge and suitable tools from computer science: a reliable morphological analyzer, a suitable, compact knowledge representation, inference methods for automatic annotation rules, and a graphical user interface. The architecture of our annotation tool, which we have built using PROLOG and relational databases, is shown in Figure 6.
4.1 Annotated Morpheme Terms

We use a compact knowledge representation of the annotated morpheme decompositions as suitable term structures, which can be managed and extended elegantly in PROLOG.

Using unique abbreviations, such as \texttt{bm} for basic morphem and \texttt{ge} for gap element, the morpheme term \((\text{craft} + \text{s}) + \text{man} + \text{ship}\) of the English word \texttt{craftsmanship}, which is visualized in Figure 7, can be annotated as follows:

\[
((\text{craft*bm} + \text{s*ge}) + \text{man})*\text{noun} + \text{ship}
\]
For persistently storing the morpheme terms and for efficiently accessing the morpheme terms based on their text form, a relational database is used. For every morpheme term, it contains further information, such as the text form, the identifier from the WDG, and the name of the user who annotated the term as well as the timestamp of the annotation.

### 4.2 Annotation Rules

The processing of the morpheme terms is completely done on the Prolog side. In the background, a set of Prolog rules is managed, which can automatically infer further annotations from the annotations and decompositions given by the user. With the following logical annotation rule, the term \((\text{craft s} + \text{man})\text{noun} + \text{ship}\) is recognized as a noun and can be further annotated to \((((\text{craft s} + \text{man})\text{noun} + \text{ship})\text{noun}\):

\[
\text{has_word_class}(X, \text{noun}) : - \\
\text{mc}(X, A, B), \text{has_word_class}(A, \text{noun}), \text{text_form}(B, [\text{ship, ...}]).
\]

### 4.3 The Morpheme Annotation Tool

The morpheme annotation tool supports the user in checking all pre-annotated morpheme decompositions. The user can annotate parts of a morpheme decomposition. Based on a set of annotation rules – which can be extended dynamically – the morpheme decomposition is further annotated as detailed as possible. The graphical user interface, which is also implemented in Prolog, consists of the morpheme editor, a visualization of the morpheme terms as tree structures, and a database browser, cf. Figure 8.

![Figure 8: The Graphical User Interface of the Morpheme Annotation Tool.](image)
5. Conclusions

The metaDictionary of the German language, based on the analyses of a network of dictionaries, forms the core part of the generic e–Infrastructure designed to identify the basic morphemes used in the German language within the last 500 years. The next step will be to test the data using text corpora: first to check for basic morphemes not covered by dictionaries, and second to find out all combinations of basic morphemes used in texts. Here we expect new insights into the combinability of basic units, quantitatively as well as qualitatively, because dictionaries of the German language normally do not register complex morpheme structures representing semantically regular patterns (e.g., German Haus–dach); they only list entries with complex structures showing specific additional semantic features (e.g., German Haus–tür). We will compare our results with the observations in Culturomics, that 52% of the English lexicon – the majority of the words used in English books – consists of lexical dark matter undocumented in standard references [5]. We expect to contribute with our e–Infrastructure in a Grid environment to the development of test methods and algorithms for detecting and understanding variance in language that ideally are applicable to genome structures as well.

References