The IMS Open Corpus Workbench (CWB)

CQP Query Language Tutorial

— CWB Version 3.4.15 —

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http://cwb.sourceforge.net/

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# CQP Query Language Tutorial

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1 Introduction

1.1 The IMS Open Corpus Workbench (CWB)

History and framework

- Tool development
  - 1993 – 1996: Project on Text Corpora and Exploration Tools (financed by the Land Baden-Württemberg)
  - 1998 – 2004: Continued in-house development (partly financed by various research and industrial projects)
  - Open-source CWB version 3.0 to be released on 1 April 2010 (pre-release versions have been shipping since October 2001)

- Related projects and applications at the IMS
  - 1994 – 1996: DECIDE project (EU programme MLAP-93) (extraction of collocation candidates, macro processor mp)
  - Since 1996: Various commercial and research applications (terminology extraction, dictionary updates)
  - 1999 – 2000: DOT project (Databank Overheidsterminologie)
  - 1999 – 2003: Implementation of YAC chunk parser for German (PhD Kermes)
  - 2001 – 2003: Transferbereich 32 (financed by the DFG)

- Some external applications of the IMS Corpus Workbench (see http://cwb.sourceforge.net/demos.php for a comprehensive list)
  - AC/DC project at the Linguateca centre (SINTEF, Oslo, Norway) (on-line access to a 180 M word corpus of Portuguese newspaper text) http://www.linguateca.pt/ACDC/
  - CorpusEye (user-friendly CQP) in the VISL project (SDU, Denmark) (on-line access to annotated corpora in various languages) http://corp.hum.sdu.dk/
  - SSLMIT Dev Online services (SSLMIT, University of Bologna, Italy) (on-line access to 380 M words of Italian newspaper text and other corpora) http://sslmitdev-online.sslmit.unibo.it/corpora/corpora.php (site no longer online)
  - CucWeb project (UPF, Barcelona, Spain) (Google-style access to 208 million words of text from Catalan Web pages) http://ramsesii.upf.es/cucweb/ (site no longer online)
  - BNCweb (CQP edition) (Web interface to the British National Corpus, ported from SARA to CQP) http://corpora.lancs.ac.uk/BNCweb/

\footnote{Designing and evaluating Extraction Tools for Collocations in Dictionaries and Corpora}
Technical aspects

- CWB uses a bespoke token-based format for corpus storage:
  - binary encoding ⇒ fast access
  - full index ⇒ fast look-up of word forms and annotations
  - specialised data compression algorithms
  - corpus size: up to 500 million words, depending on annotations
  - text data and annotations cannot be modified after encoding
    (but it is possible to add new annotations or overwrite existing ones)
  - assumes Latin-1 encoding, but compatible with other 8-bit ASCII extensions
    (Unicode text in UTF-8 encoding can be processed with some caveats)

- Typical compression ratios for a 100 million word corpus:
  - uncompressed text: ≈ 1 GByte (without index & annotations)
  - uncompressed CWB attributes: ≈ 790 MBytes (ratio: 1.3)
  - word forms & lexical attributes: ≈ 360 MBytes (ratio: 2.8)
  - categorical attributes (e.g. POS tags): ≈ 120 MBytes (ratio: 8.5)
  - binary attributes (yes/no): ≈ 50 MBytes (ratio: 20.5)

- Supported operating systems:
  - GNU Linux 2.6+ (32-bit and 64-bit Intel/AMD processors)
  - Mac OS X 10.4+ (32-bit and 64-bit Intel, 32-bit PPC)
  - SUN Solaris 2.8+ (Sparc processors)
  - experimental support for Microsoft Windows (32-bit)
  - Source code should compile on most recent Unix platforms
  - Corpus data format is platform-independent and compatible with all releases since 2001

Components of the CWB

- tools for encoding, indexing, compression, decoding, and frequency distributions
- global “registry” holds information about corpora (name, attributes, data path)
- corpus query processor (CQP):
  - fast corpus search (regular expression syntax)
  - use in interactive or batch mode
  - results displayed in terminal window
- CWB/Perl interface for post-processing, scripting and web interfaces
- CQPweb: a browser-based web interface to CWB/CQP, with extended analysis tools
1.2 The CWB corpus data model

The following steps illustrate the transformation of textual data with some XML markup into the CWB data format.

1. **Formatted text** (as displayed on-screen or printed)
   
   An easy example. Another *very* easy example. Only the easiest examples!

2. **Text with XML markup** (at the level of texts, words or characters)

   ```
   <text id=42 lang="English"> <s>An easy example.</s><s> Another <i>very</i> easy example.</s> <s><b>O</b>nly the <b>ea</b>siet <b>ex</b>a<b>mpl</b>es!</s> </text>
   ```

3. **Tokenised text** (character-level markup has to be removed)

   ```
   <text id=42 lang="English"> <s> An easy example . </s> <s> Another very easy example . </s> <s> Only the easiest examples ! </s> </text>
   ```

4. **Text with linguistic annotations** (annotations are added at token level)

   ```
   <text id=42 lang="English"> <a> An/DET/a easy/ADJ/easy example/NN/example ./PUN/. </a> <s> Another/DET/another very/ADV/very easy/ADJ/easy example/NN/example ./PUN/. </s> <s> Only/ADV/only the/DET/the easiest/ADJ/easy examples/NN/example !/PUN!/ </s> </text>
   ```

5. **Text encoded as CWB corpus** (tabular format, similar to relational database)

   A schematic representation of the encoded corpus is shown in Figure 1. Each token (together with its annotations) corresponds to a row in the tabular format. The row numbers, starting from 0, uniquely identify each token and are referred to as corpus positions.

   Each (token-level) annotation layer corresponds to a column in the table, called a positional attribute or p-attribute (note that the original word forms are also treated as an attribute with the special name `word`). Annotations are always interpreted as character strings, which are collected in a separate lexicon for each positional attribute. The CWB data format uses lexicon IDs for compact storage and fast access.

   Matching pairs of XML start and end tags are encoded as token regions, identified by the corpus positions of the first token (immediately following the start tag) and the last token (immediately preceding the end tag) of the region. (Note how the corpus position of an XML tag in Figure 1 is identical to that of the following or preceding token, respectively.) Elements of the same name (e.g. `<s>...</s>` or `<text>...</text>`) are collected and referred to as a structural attribute or s-attribute. The corresponding regions must be non-overlapping and non-recursive. Different s-attributes are completely independent in the CWB: a hierarchical nesting of the XML elements is neither required nor can it be guaranteed.

   Key-value pairs in XML start tags can be stored as an annotation of the corresponding s-attribute region. All key-value pairs are treated as a single character string, which has to be “parsed” by a CQP query that needs access to individual values. In the recommended encoding procedure, an additional s-attribute (named `element_key`) is automatically created for each key and is directly annotated with the corresponding value (cf. `<text_id>` and `<text_lang>` in Figure 1).

6. **Recursive XML markup** (can be automatically renamed)

   Since s-attributes are non-recursive, XML markup such as

   ```
   <np>the man <pp>with <np>the telescope</np></pp> </np>
   ```
is not allowed in a CWB corpus (the embedded `<np>` region will automatically be dropped).\footnote{Recall that only the nesting of a `<np>` region within a larger `<np>` region constitutes recursion in the CWB data model. The nesting of `<pp>` within `<np>` (and vice versa) is unproblematic, since these regions are encoded in two independent s-attributes (named `pp` and `np`).}

In the recommended encoding procedure, embedded regions (up to a pre-defined level of embedding) are automatically renamed by adding digits to the element name:

```xml
<np>the man <pp>with <np1>the telescope</np1></pp></np>
```

<table>
<thead>
<tr>
<th>corpus position</th>
<th>word ID</th>
<th>part of ID</th>
<th>lemma ID</th>
<th>form</th>
<th>speech</th>
<th>part of ID</th>
<th>lemma ID</th>
<th>form</th>
<th>speech</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0)</td>
<td><code>&lt;text&gt;</code> value = &quot;id=42 lang=&quot;English&quot;&quot;]&quot;</td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<td>(0)</td>
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<tr>
<td>(0)</td>
<td><code>&lt;text_lang&gt;</code> value = &quot;English&quot;</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>easy</td>
<td>1</td>
<td>ADJ</td>
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<td>easy</td>
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<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>example</td>
<td>2</td>
<td>NN</td>
<td>2</td>
<td>example</td>
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<td>2</td>
<td></td>
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</tr>
<tr>
<td>3</td>
<td>.</td>
<td>3</td>
<td>PUN</td>
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<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(3)</td>
<td><code>&lt;/s&gt;</code></td>
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<tr>
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<td>DET</td>
<td>0</td>
<td>another</td>
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<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>very</td>
<td>5</td>
<td>ADV</td>
<td>4</td>
<td>very</td>
<td>5</td>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>easy</td>
<td>1</td>
<td>ADJ</td>
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<td>easy</td>
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<td>1</td>
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</tr>
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<tr>
<td>8</td>
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<td>9</td>
<td>Only</td>
<td>6</td>
<td>ADV</td>
<td>4</td>
<td>only</td>
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<td>6</td>
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</tr>
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<td>the</td>
<td>7</td>
<td>DET</td>
<td>0</td>
<td>the</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>easiest</td>
<td>8</td>
<td>ADJ</td>
<td>1</td>
<td>easy</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>examples</td>
<td>9</td>
<td>NN</td>
<td>2</td>
<td>example</td>
<td>2</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>!</td>
<td>10</td>
<td>PUN</td>
<td>3</td>
<td>!</td>
<td>8</td>
<td>8</td>
<td></td>
<td></td>
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<tr>
<td>(13)</td>
<td><code>&lt;/s&gt;</code></td>
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<td></td>
</tr>
<tr>
<td>(13)</td>
<td><code>&lt;/text_lang&gt;</code></td>
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<tr>
<td>(13)</td>
<td><code>&lt;/text_id&gt;</code></td>
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<td>(13)</td>
<td><code>&lt;/text&gt;</code></td>
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</tr>
</tbody>
</table>

Figure 1: Sample text encoded as a CWB corpus.
1.3 Corpora used in the tutorial

Pre-encoded versions of these corpora are distributed free of charge together with the IMS Corpus Workbench. Perl scripts for encoding the *British National Corpus* (World Edition) can be provided at request.

**English corpus: DICKENS**

- a collection of novels by Charles Dickens
- ca. 3.4 million tokens
- derived from Etext editions (Project Gutenberg)
- document-structure markup added semi-automatically
- part-of-speech tagging and lemmatisation with TreeTagger
- recursive noun and prepositional phrases from Gramotron parser

**German corpus: GERMAN-LAW**

- a collection of freely available German law texts
- ca. 816,000 tokens
- part-of-speech tagging with TreeTagger
- morphosyntactic information and lemmatisation from IMSLex morphology
- partial syntactic analysis with YAC chunker

See Appendix A.3 for a detailed description of the token-level annotations and structural markup of the tutorial corpora (positional and structural attributes).
2 Basic CQP features

2.1 Getting started

• start CQP by typing
  $ cqp -e
in a shell window (the $ indicates a shell prompt)
• -e flag activates command-line editing features³
• optional -C flag activates colour highlighting (experimental)
• every CQP command must be terminated with a semicolon (;)
• when command-line editing is activated, CQP will automatically add a semicolon at the end of each input line if necessary; explicit ; characters are only necessary to separate multiple commands on a single line in this mode
• change the registry directory (where CQP will look for available corpora)
  > set Registry "/some/path/to/a/directory";
  if the registry is not set, CQP will look in the default location
• list available corpora
  > show corpora;
• get information about corpus (including corpus size in tokens)
  > info DICKENS;
displays information file associated with the corpus, whose contents may vary; ideally, this should give a description of the corpus composition, a summary of the positional and structural annotations, and a brief overview of annotation codes such as the part-of-speech tagset used
• activate corpus for subsequent queries (use TAB key for name completion)
  [no corpus]> DICKENS;
  DICKENS>
in the following examples, the CQP command prompt is indicated by a > character
• list attributes of activated corpus ("context descriptor")
  > show cd;

2.2 Searching for words

• search single word form (single or double quotes are required: ‘...’ or "...")
  > "interesting";
  → shows all occurrences of interesting
• the specified word is interpreted as a regular expression
  > "interest(s|ed|ing)(ly)?";
  → interest, interests, interested, interesting, interestingly, interestingly

³The -e mode is not enabled by default for reasons of backward compatibility. When command-line editing is active, multi-line commands are not allowed, even when the input is read from a pipe.
• see Appendix A.1 for an introduction to the regular expression syntax
• the regular expression “flavour” used by CQP is Perl Compatible Regular Expressions usually known as PCRE; lots of documentation and rexamples can be found on the WWW
• note that special characters have to be “escaped” with backslash (\)

"?" fails; "\?" → ?; "." → . ! ? a b c ...; "\$." → $

“critical” characters are: . ? * + | ( ) [ ] { } ^ $

• CWB 3.0: IATPEX-style escape sequences \", \‘, \` and \“, followed by an appropriate ASCII letter, are used to represent characters with diacritics when they cannot be entered directly

"B\"ar" → Bär;  "d\‘e\ʝ\"a" → déjà

NB: this feature is deprecated; it works only for the Latin-1 encoding and cannot be deactivated

• CWB 3.0: additional special escape sequences:

\"s → β; \,c → c; \,C → C; \¬n → n; \¬N → N;

version 3.0.3 introduces two-digit hex escapes for inserting arbitrary byte values:

\xDF → β in a Latin1-encoded corpus; \xC3\x9F → ÿ in a UTF-8-encoded corpus

• CWB 3.5: full support for PCRE regular expressions, including two- and four-digit hex escapes

• use flags %c and %d to ignore case / diacritics

DICKENS> "interesting" %c;
GERMAN-LAW> "wahrung" %cd;

• if you need to match a word form containing single or double quotes (e.g. 'em or 12"-screen), there are two possibilities:

  − if the string does not contain both single and double quotes, simply pick an appropriate quote character: "'em" vs. "12"-screen"
  − otherwise, double every occurrence of the quote character inside the string; our two examples could also be matched with the queries "'''em' and "12""-screen"

2.3 Display options

• KWIC display (“key word in context”)

15921: ry moment an <interesting> case of spo
17747: appeared to <interest> the Spirit
20189: ge , with an <interest> he had neve
24026: rgetting the <interest> he had in w
35161: require . My <interest> in it , is
35490: require . My <interest> in it was s
35903: ken a lively <interest> in me sever
43031: been deeply <interested> , for I rem

• if query results do not fit on screen, they will be displayed one page at a time

• press SPC (space bar) to see next page, RET (return) for next line, and q to return to CQP

• some pagers support b or the backspace key to go to the previous page, as well as the use of the cursor keys, PgUp, and PgDn

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• at the command prompt, use cursor keys to edit input (← and →, Del, backspace key) and repeat previous commands (↑ and ↓)

• change context size
  > set Context 20; (20 characters)
  > set Context 5 words; (5 tokens)
  > set Context s; (entire sentence)
  > set Context 3 s; (same, plus 2 sentences each on left and right)

• type “cat;” to redisplay matches

• display current context settings
  > set Context;

• left and right context can be set independently
  > set LeftContext 20;
  > set RightContext s;

• all option names are case-insensitive; most options have abbreviations:
  c for Context, lc for LeftContext, rc for RightContext
  (shown in square brackets when current value is displayed)

• show/hide annotations
  > show +pos +lemma; (show)
  > show -pos -lemma; (hide)

• summary of selected display options (and available attributes):
  > show cd;

• structural attributes are shown as XML tags
  > show +s +np_h;

• hide annotations of XML tags
  > set ShowTagAttributes off;

• hide corpus position
  > show -cpos;

• show annotation of region(s) containing match
  > set PrintStructures "np_h";
  > set PrintStructures "novel_title, chapter_num";
  > set PrintStructures "";

2.4 Useful options

• enter set; to display list of options (abbreviations shown in brackets)

• set <option>; shows current value

• set ProgressBar (on|off);
  to show progress of query execution

• set Timing (on|off);
  to show execution times of queries and some other commands
• set PrintMode (ascii|sgml|html|latex);
to set output format for KWIC display and frequency distributions

• set PrintOptions (hdr|nohdr|num|nonum|...);
to turn various formatting options on (hdr, num, ...) or off (nohdr, nonum, ...)
type set PrintOptions; to display the current option settings
useful options: hdr (display header), num (show line numbers), tbl (format as table in HTML and \LaTeX modes), bdr (table with border lines)

• set (LD|RD) <string>;
change left/right delimiter in KWIC display from the default < and > markers

• set ShowTagAttributes (on|off);
to display key-value pairs in XML start tags (if annotated in the corpus)

• create .cqprc file in your home directory with your favourite settings
(contains arbitrary CQP commands that will be read and executed during startup)

• for a persistent command history, add the lines
set HistoryFile "<home>/.cqphistory";
set WriteHistory yes;
to your .cqprc file (if CQP is run with -e option)
NB: the size of the history file is not limited automatically by CQP

• set AutoShow off;
no automatic KWIC display of query results

• set Optimize on;
enable experimental optimisations (sometimes included in beta versions)

2.5 Accessing token-level annotations

• specify p-attribute/value pairs (brackets are required)
  > [pos = "JJ"];   (find adjectives)
  > [lemma = "go"];  

• "interesting" is an abbreviation for [word = "interesting"]

• the implicit attribute in the abbreviated form can be changed with the DefaultNonbrackAttr
  option; for instance, enter
  > set DefaultNonbrackAttr lemma;
  to search for lemmatised words instead of surface forms

• %c and %d flags can be used with any attribute/value pair
  > [lemma = "pole" %c];

• values are interpreted as regular expressions, which the annotation string must match; add %l
  flag to match literally:
  > [word = "?" %l];

• != operator: annotation must not match regular expression
  [pos != "N.*"] → everything except nouns
matches any token (⇒ matchall pattern)

see Appendix A.2 for a list of useful part-of-speech tags and regular expressions

or find out with the /codist[] macro (more on macros in Sections 6.4 and 6.5):
  > /codist["whose", pos];
  → finds all occurrences of the word whose and computes frequency distribution of the
  part-of-speech tags assigned to it

use a similar macro to find inflected forms of go:
  > /codist[lemma, "go", word];
  → finds all tokens whose lemma attribute has the value go and computes frequency distribution
  of the corresponding word forms

abort query evaluation with Ctrl-C
  (does not always work, press twice to exit CQP immediately)

2.6 Combinations of attribute constraints: Boolean expressions

operators: & (and), | (or), ! (not), -⇒ (implication, cf. Section 4.1)
  > [(lemma="under.+") & (pos="V.*")];
  → verb with prefix under...

attribute/attribute-pairs: compare attributes as strings
  > [(lemma="under.+") & (word!=lemma)];
  → inflected forms of lemmas with prefix under...

complex expressions:
  > [(lemma="go") & !(word="went"%c | word="gone"%c)];

any expression in square brackets [...] describes a single token (⇒ pattern)

2.7 Sequences of words: token-level regular expressions

a sequence of words or patterns matches any corresponding sequence in the corpus
  > "on " and "on|off";
  > "in" "any|every" [pos = "NN"];

modelling of complex word sequences with regular expressions over patterns (i.e. tokens): ev-
  ery [...] expression is treated like a single character (or, more precisely, a character set) in
  conventional regular expressions

token-level regular expressions use a subset of the POSIX syntax

repetition operators:
  ? (0 or 1), * (0 or more), + (1 or more), \{n\} (exactly n), \{n,m\} (n...m)

grouping with parentheses: (...)

disjunction operator: | (separates alternatives)

parentheses delimit scope of disjunction: ( alt₁ | alt₂ | ... )

Figure 2 shows simple queries matching prepositional phrases (PPs) in English and German. The
query strings are spread over multiple lines to improve readability, but each one has to be entered
on a single line in an interactive CQP session.
Figure 2: Simple queries matching PPs in English and German.

2.8 Example: finding “nearby” words

- insert optional matchall patterns between words
  > "right" []? "left";

- repeated matchall for longer distances
  > "no" "sooner" []* "than";

- use the range operator {,} to restrict number of intervening tokens
  > "as" []{1,3} "as";

- avoid crossing sentence boundaries by adding within s to the query
  > "no" "sooner" []* "than" within s;

- order-independent search
  > "left" "to" "right"
    | "right" "to" "left";

2.9 Sorting and counting

- sort matches alphabetically (re-displays query results)
  > [pos = "IN"] "any""every" [pos = "NN"];
  > sort by word;
• add %c and %d flags to ignore case and/or diacritics when sorting
  > sort by word %cd;

• matches can be sorted by any positional attribute; just type
  > sort;
  without an attribute name to restore the natural ordering by corpus position

• query results can also be sorted in random order (to avoid looking only at matches from the first
  part of a corpus when paging through query results):
  > sort randomize;
  more on random sorting and an important application in Section 3.6

• select descending order with desc(ending), or sort matches by suffix with reverse;
  note the ordering when the two options are combined:
  > sort by word descending reverse;

• compute frequency distribution of matching word sequences (or annotations)
  > count by word;
  > count by lemma;

• %c and %d flags normalise case and/or diacritics before counting
  > count by word %cd;

• set frequency threshold with cut option
  > count by lemma cut 10;

• descending option affects ordering of word sequences with the same frequency; use reverse for
  some amusing effects (note that these keywords go before the cut option)

• sort by right or left context (especially useful for keyword searches)
  > "interesting";
  > sort by word %cd on matchend[1] .. matchend[42]; (right context)
  > sort by word %cd on match[-1] .. match[-42]; (left context, by words)
  > sort by word %cd on match[-42] .. match[-1] reverse; (same by characters)

• see Sections 3.2 and 3.3 for an explanation of the syntax used in these examples and more
  information about the sort and count commands
3 Working with query results

3.1 Named query results

- store query result in memory under specified name (should begin with capital letter)
  > Go = [lemma = "go"] "and" [];
  note that query results are not automatically displayed in this case

- list named query results (or NQR for short)
  > show named;

- The fully NQR specifiers shown in the output of this command are qualified with the CWB name of the corpus, e.g. DICKENS:Go for the query above. As a consequence, there can be multiple NQRs with the same name for different corpora. If short query names are used in CQP commands (as in the examples below), they are automatically prefixed with the currently activated corpus.

- result of last query is implicitly named Last; commands such as cat, sort, and count operate on Last by default; note that Last is always temporary and will be overwritten when a new query is executed (or a subset command, cf. Section 3.5)

- display number of results
  > size Go;

- (full or partial) KWIC display
  > cat Go;
  > cat Go 5 9; (6th – 10th match)

- sorting a named query result automatically re-displays the matches
  > sort Go by word %cd;

- the count command also sorts the named query on which it operates:
  > count Go by lemma cut 5;
  implicitly executes the command sort Go by lemma;

- this has the advantage that identical word sequences now appear on adjacent lines in the KWIC display and can easily be printed with a single cat command; the respective line numbers are shown in square brackets at the end of each line in the frequency listing

13 go and see [#128-#140]
10 go and sit [#144-#153]
 9 go and do [#29-#37]
 7 go and fetch [#42-#48]
 7 go and look [#87-#93]
 7 go and play [#107-#113]

to display occurrences of go and see, enter
  > cat Go 128 140;

- If fully qualified NQR are used, a query result can be accessed even if its corpus isn’t currently activated:
  > size DICKENS:Go;
  > count DICKENS:Go by lemma cut 5;
  will work regardless of the current corpus.
• Due to a long-standing bug in CQP, this feature should not be used with cat or any other command that generates KWIC output (such as sort). Doing so will corrupt the context descriptor, which holds information about all available attributes, those selected for printing, and the KWIC context size.

```
> GERMAN-LAW;
> show cd;
> cat DICKENS:Time;
> show cd;
```

• The context descriptor can only be repaired by temporarily activating a different corpus and then re-activating the desired corpus.

```
> DICKENS;
> GERMAN-LAW;
```

3.2 Saving data to disk

• named query results can be stored on disk in the DataDirectory
  
  ```
  > set DataDirectory ".";
  > DICKENS;
  ```
  
  NB: you need to re-activate your working corpus after setting the DataDirectory option

• save named query to disk (in a platform-dependent uncompressed binary format)
  
  ```
  > save Go;
  ```

• md* flags show whether a named query is loaded in memory (m), saved on disk (d), or has been modified from the version saved on disk (*)

  ```
  > show named;
  ```

• discard named query results to free memory
  
  ```
  > discard Go;
  ```

• set DataDirectory to load named queries from disk
  (after discarding, or in a new CQP session)

  ```
  > set DataDirectory ".";
  > show named;
  > cat Go;
  ```
  
  note that the actual data are only read into memory when the query results are accessed

• write KWIC output to text file (use TAB key for filename completion)

  ```
  > cat Go > "go.txt";
  ```

  use set PrintOptions hdr; to add header with information about the corpus and the query (previous CQP versions did this automatically)

• if the filename ends in .gz or .bz2, the file will automatically be compressed (provided that the respective command-line utilities gzip and bzip2 are available)

• append to an existing file with >>; this also works for compressed files

  ```
  > cat Go >> "go.txt";
  ```
you can also write to a pipe (this example saves only matches that occur in questions, i.e. sentences ending in ?)

```bash
> set Context 1 s;
> cat Go | grep '\?$' > go2.txt;
```

- set `PrintMode` and `PrintOptions` for HTML output and other formats (see Section 2.4)
- frequency counts for matches can also be written to a text file
  ```bash
  > count Go by lemma cut 5 > "go.cnt";
  ```
- new in CQP v3.4.14: `cat` can also be used to print an arbitrary string or redirect it to a file; escape sequences `\t` (TAB), `\r` (CR) and `\n` (LF) are interpreted, all other backslashes are passed through verbatim; note that the string is not automatically terminated with a newline
  ```bash
  > cat "Just another\n\tCQP hacker.\n";
  ```
- the new functionality can be combined with output redirection, which is particularly convenient for adding header rows to tabular output files, e.g.
  ```bash
  > cat "f\tmatch [results]\n" > "go.cnt";
  > count Go by lemma cut 5 >> "go.cnt";
  ```

### 3.3 Anchor points

- the result of a (complex) query is a list of token sequences of variable length (⇒ matches)
- each match is represented by two anchor points: `match` (corpus position of first token) and `matchend` (corpus position of last token)
- set additional target anchor with `@` marker in query (prepended to a pattern)
  ```bash
  > "in" @[pos="DT"] [lemma="case"];  
  → shown in bold font in KWIC display
  ```
- only a single token can be marked as target; if multiple `@` markers are used (or if the marker is in the scope of a repetition operator such as `+`), only the rightmost matching token will be marked
  ```bash
  > [pos="DT"] (@[pos="JJ.*"] ","?){2,} [pos="NNS?"];
  ```
- when `targeted` pattern is optional, check how many matches have target anchor set
  ```bash
  > A = [pos="DT"] @[pos="JJ"];  
  > size A;
  > size A target;
  ```
- anchor points allow a flexible specification of sort keys with the general form
  ```bash
  > sort by attribute on start point .. end point ;
  ```
  both `start point` and `end point` are specified as an anchor, plus an optional offset in square brackets; for instance, `match[-1]` refers to the token before the start of the match, `matchend` to the last token of the match, `matchend[1]` to the first token after the match, and `target[-2]` to a position two tokens left from the `target` anchor
  
  NB: the `target` anchor should only be used in the sort key when it is always defined
- example: sort noun phrases by adjectives between determiner and noun
  ```bash
  > [pos="DT"] [pos="JJ"]{2,} [pos="NNS?"];
  > sort by word %cd on match[1] .. matchend[-1];
  ```
• if end point refers to a corpus position before start point, the tokens in the sort keys are compared from right to left; e.g. sort on the left context of the match (by token)
  > sort by word %cd on match[-1] .. match[-42];
  whereas the reverse option sorts on the left context by character
  > sort by word %cd on match[-42] .. match[-1] reverse;

• complex sort operations can sometimes be speeded up by using an external helper program (the standard Unix sort tool)\(^4\)
  > sort by word %cd;
  > set ExternalSort on;
  > sort by word %cd;
  > set ExternalSort off;

• the count command accepts the same specification for the strings to be counted
  > count by lemma on match[1] .. matchend[-1];

• display corpus positions of all anchor points in tabular format
  > A = "behind" @[pos="JJ"]? [pos="NNS?"];
  > dump A;
  > dump A 9 14; (10\(^{\mathrm{th}}\) - 15\(^{\mathrm{th}}\) match)
  the four columns correspond to the match, matchend, target and keyword (see Section 3.7) anchors; a value of -1 means that the anchor has not been set:

  
  | 1019887 | 1019888 | -1 | -1 |
  | 1924977 | 1924979 | 1924978 | -1 |
  | 1986623 | 1986624 | -1 | -1 |
  | 2086708 | 2086710 | 2086709 | -1 |
  | 2087618 | 2087619 | -1 | -1 |
  | 2122565 | 2122566 | -1 | -1 |

  note that a previous sort or count command affects the ordering of the rows (so that the n-th row corresponds to the n-th line in a KWIC display obtained with cat)

• the output of a dump command can be written (>) or appended (>>) to a file, if the first character of the filename is |, the output is sent to the pipe consisting of the following command(s); use the following trick to display the distribution of match lengths in the query result A:
  > A = [pos="DT"] [pos="JJ.*"]* [pos="NNS?"];
  > dump A > " | gawk '{print $2 - $1 + 1}' | sort -nr | uniq -c | less";

• see Section 7.2 for an opposite to the dump command, which may be useful for certain tasks such as locating a specific corpus position

3.4 Frequency distributions

• frequency distribution of tokens (or their annotations) at anchor points
  > group Go matchend pos;
set cutoff threshold with cut option to reduce size of frequency table

> NP = [pos="DT"]@[pos="JJ"]? [pos="NNS?"];
> group NP target lemma cut 50;

- add optional offset to anchor point, e.g. distribution of words preceding matches
  > group NP match[-1] lemma cut 100;

- frequencies of token/annotation pairs (using different attributes or anchor points)
  > group NP matchend word by target lemma;
  > group Go matchend lemma by matchend pos;

NB: despite what the command syntax and output format suggest, results are sorted by pair frequencies (not grouped by the second item); also note that the order of the two items in the output is opposite to the order in the group command

- you can write the output of the group command to a text file (or pipe)
  > group NP target lemma cut 10 > "adjectives.go";
  (in CQP v.3.4.11 and newer, the file is automatically compressed if it ends in .gz or .bz2)

- new in CQP v3.4.9: use group by instead of by for nested frequency counts
  > group Go matchend lemma group by matchend pos;

where an optional cut clause applies to the individual pairs

3.5 Set operations with named query results

- named queries can be copied, especially before destructive modification (see below)
  > B = A;
  > C = Last;

- compute subset of named query result by constraint on one of the anchor points
  > PP = [pos="IN"] [pos="JJ"]+ [pos="NNS?"];
  > group PP matchend lemma by match word;
  > PP1 = subset PP where match: "in";
  > PP2 = subset PP1 where matchend: [lemma = "time"];
  → PP2 contains instances of in ... time(s)

- set operations on named query results
  > A = intersection B C;  \( A = B \cap C \)
  > A = union B C;  \( A = B \cup C \)
  > A = difference B C;  \( A = B \setminus C \)

intersection (or inter) yields matches common to B and C; union (or join) matches from either B or C; difference (or diff) matches from B that are not in C

- cut query result to first n matches
  > cut A 50; (first 50 matches)

or select a range of matches (as with the restricted cat command)

> cut A 50 99; (51st - 100th match)

NB: cut A 50; is exactly the same as cut A 0 49;
• The modifier `cut n` can also be appended to a query:
  ```
  > "time" cut 50;
  ```
  The main purpose of this usage is to reduce memory consumption and processing time in Web interfaces and similar applications by stopping query execution early if a sufficient number of matches has been found. For internal reasons, this optimization cannot be applied to queries with alignment constraints (see Sec. 5.2); but the `cut` modifier still guarantees that only the first `n` matches will be returned.

### 3.6 Random subsets

• when there are a lot of matches, e.g.
  ```
  > A = "time";
  > size A;
  ```
  it is often desirable to look at a random selection to get a quick overview (rather than just seeing matches from the first part of the corpus); one possibility is to do a `sort randomize` and then go through the first few pages of random matches:
  ```
  > sort A randomize;
  ```
  however, this cannot be combined with other sort options such as alphabetical sorting on match or left/right context; it also doesn’t speed up frequency lists, `set target` and other post-processing operations

• as an alternative to randomized ordering, the `reduce` command randomly selects a given number or proportion of matches, deleting all other matches from the named query; since this operation is destructive, it may be necessary to make a copy of the original query results first (see above)
  ```
  > reduce A to 10%;
  > size A;
  > sort A by word %cd on match .. matchend[42];
  > reduce A to 100;
  > size A;
  > sort A by word %cd on match .. matchend[42];
  ```
  this allows arbitrary further operations to be carried out on a representative sample rather than the full query result

• set random number generator seed before `reduce` for reproducible selection
  ```
  > randomize 42; (use any positive integer as seed)
  ```

• a second method for obtaining a random subset of a named query result is to sort the matches in random order and then take the first `n` matches from the sorted query; the example below has the same effect as `reduce A to 100`; (though it will not select exactly the same matches)
  ```
  > sort A randomize;
  > cut A 100; (NB: this restores corpus order, as with the reduce command)
  ```
  reproducible subsets can be obtained with a suitable `randomize` command before the `sort`; the main difference from the `reduce` command is that `cut` cannot be used to select a percentage of matches (i.e., you have to determine the number of matches in the desired subset yourself)

• the most important advantage of the second method is that it can produce *stable* and *incremental* random samples
for a stable random ordering, specify a positive seed value directly in the sort command:

```cqp
> sort A randomize 42;
```

different seeds give different, reproducible orderings; if you randomize a subset of A with the same seed value, the matches will appear exactly in the same order as in the randomized version of A:

```cqp
> A = "interesting" cut 20; (just for illustration)
> B = A;
> reduce B to 10; (an arbitrary subset of A)
> sort A randomize 42;
> sort B randomize 42;
```

in order to build incremental random samples from a query result, sort it randomly (but with seed value to ensure reproducibility) and then take the first \( n \) matches as sample \#1, the next \( n \) matches as sample \#2, etc.; unlike two subsets generated with `reduce`, the first two samples are disjoint and together form a random sample of size \( 2n \):

```cqp
> A = "time";
> sort A randomize 7;
> Sample1 = A;
> cut Sample1 0 99; (random sample of 100 matches)
> Sample2 = A;
> cut Sample2 100 199; (random sample of 100 matches)
```

Note that the `cut` removes the randomized ordering; you can reapply the stable randomization to achieve full correspondence to the randomized query result A:

```cqp
> sort Sample2 randomize 7;
> cat Sample2;
> cat A 100 199;
```

stability of the randomization ensures that random samples are reproducible even after the initial query has been refined or spurious matches have been deleted manually.

### 3.7 The set target command

- additional `keyword anchor` can be set after query execution by searching for a token that matches a given `search pattern` (see Figure 3)

```
set <named query>
  (keyword | target) (anchor to set)
  (leftmost | rightmost | nearest | farthest) (search strategy)
  [<pattern>] (search pattern)
  within
  (left | right)? (search direction)
  <n> (words | \( s \) | ...) (window)
  from (match | matchend | keyword | target)
  (inclusive)? ; (include start token in search)
```

Figure 3: The set target command.
example: find noun near adjective *modern*

```
A = [(pos="JJ") & (lemma="modern")];
set A keyword nearest [pos="NNS?"] within right 5 words from match;
```

- keyword should be **underlined** in KWIC display (may not work on some terminals)
- search starts from the given anchor point (excluding the anchored token itself), or from the left and right boundaries of the match if `match` is specified
- with **inclusive**, search includes the anchored token, or the entire match, respectively
- `from match` is the default and can be omitted
- the `match` and `matchend` anchors can also be set, modifying the actual matches\(^5\)
- anchors can be copied:
  ```
  set A target match;
  set A matchend keyword;
  ```

\(^5\)The `keyword` and `target` anchors are set to undefined (-1) when no match is found for the search pattern, while the `match` and `matchend` anchors retain their previous values. In this way, a `set match` or `set matchend` command may only modify some of the matches in a named query result.
4 Labels and structural attributes

4.1 Using labels

- patterns can be labelled (similar to the target marker @)
  \>
  > adj:[pos = "JJ.*"] ... ;
  the label adj then refers to the corresponding token (i.e. its corpus position)

- label references are usually evaluated within the global constraint introduced by ::
  > adj:[pos = "ADJ."] :: adj < 500;
  \> adjectives among the first 500 tokens

- annotations of the referenced token can be accessed as adj.word, adj.lemma, etc.

- labels are not part of the query result and must be used within the query expression (otherwise, CQP will abort with an error message)

- labels set to optional patterns may be undefined
  > [pos="DT"] a:[pos="JJ"]? [pos="NNS?"] :: a;
  \> global constraint a is true iff match contains an adjective

- to avoid error messages, test whether label is defined before accessing attributes
  > [pos="DT"] a:[]? [pos="NNS?"] :: a -> a.pos="JJ";
  \> is the logical implication operator →, cf. Section 2.6

- labels are used to specify additional constraints that are beyond the scope of ordinary regular expressions
  > a:[] "and" b:[] :: a.word = b.word;

- labels allow modelling of long-distance dependencies
  > a:[pos="PP"] []{0,5} b:[pos = "VB.*"]
     :: b.pos = "VBZ" -> a.lemma = "he|she|it";
  \> this query ensures that the pronoun preceding a 3rd-person singular verb form is he, she or it; an additional constraint could exclude these pronouns for other verb forms

- labels can be used within patterns as well
  > a:[] [pos = a.pos]{3};
  \> sequences of four identical part-of-speech tags

- however, a label cannot be used within the pattern it refers to; use the special this label represented by a single underscore (_) instead to refer to the current corpus position
  \>
  > [...pos = "NPS"] \iff [pos = "NPS"]

- the built-in functions distance() and distabs() compute the (absolute) distance between 2 tokens (referenced by labels)
  > a:[pos="DT"] [pos="JJ"]* b:[pos="NNS?"] :: distabs(a,b) >= 5;
  \> simple NPs containing 6 or more tokens

- the standard anchor points (match, matchend, and target) are also available as labels (with the same names)
  > [pos="DT"] [pos="JJ"]* [pos="NNS?"] :: distabs(match, matchend) >= 5;
4.2 Structural attributes

- XML tags match start/end of s-attribute region (shown as XML tags in Figure 1)
  
  > <s> [pos = "VBG"];
  > [pos = "VBG"] [pos = "SENT"]? </s>;
  → present participle at start or end of sentence

- pairs of start/end tags enclose single region (if StrictRegions option is enabled)
  
  > <np> []* ([pos="JJ.*"] [^*]{3,} </np>;
  → NP containing at least 3 adverbs

  (when StrictRegions are switched off, XML tags match any region boundaries and may skip intervening boundaries as well as material outside the corresponding regions)

- /region[] macro matches entire region
  
  /region[np]; ⇐⇒ <np> []* </np>;

- different tags can be mixed
  
  > <s><np> []* </np> []* <np> []* </np></s>;
  → sentence that starts and ends with a noun phrase (NP)

- the name of a structural attribute (e.g. np) used within a pattern evaluates to true iff the corresponding token is contained in a region of this attribute (here, a <np> region)
  
  > [(pos = "NNS?") & !np];
  → noun that is not contained in a noun phrase (NP)

- built-in functions lbound() and rbound() test for start/end of a region
  
  > [(pos = "VBG") & lbound(s)];
  → present participle at start of sentence

- new in CQP v3.4.13: Built-in functions lbound_of() and rbound_of() return the corpus positions of the start/end of a region. Because of technical limitations, the anchor position has to be specified explicitly as a second argument, which will often be the this label:
  
  > [(word = "\d+") & (lbound_of(s, _) = lbound_of(chapter, _))];
  → a number in the first sentence of a chapter

  The same query could also be written with an explicit label or anchor reference in a global constraint (which is perhaps easier to read):
  
  > "\d+" :: lbound_of(s, match) = lbound_of(chapter, match);

  If the referenced position is not contained in a suitable s-attribute region, the functions return an undefined value, which evaluates to false in most contexts (in particular, all comparisons with this value will be false).

- The lbound_of() and rbound_of() functions are mainly used in connection with distance() or distabs(). For example, to find occurrences of the word end within the first 40 tokens of a chapter:
  
  > [word = "end"%c & distabs(_, lbound_of(chapter, _)) < 40];

- use within to restrict matches of a query to a single region
  
  > [pos="NN"] []* [pos="NN"] within np;
  → sequence of two singular nouns within the same NP
• most linguistic queries should include the restriction within s to avoid crossing sentence boundaries; note that only a single within clause may be specified

• query matches can be expanded to containing regions of s-attributes
  > A = [pos="JJ.*"] (\[* [pos="JJ.*"]\){2} within np;  
  > B = A expand to np;
  one-sided expansion is selected with the optional left or right keyword
  > C = B expand left to s;

• the expansion can be combined with a query, following all other modifiers  
  > [pos="JJ.*"] ([\]* [pos="JJ.*"]\){2} within np cut 20 expand to np;

4.3 Structural attributes and XML

• XML markup of NPs and PPs in the DICKENS corpus (cf. Appendix A.3)

```xml
<s len=9>
  <np h="it" len=1> It </np>
  is
  <np h="story" len=6> the story
  <pp h="of" len=4> of
    <np h="man" len=3> an old man </np>
  </pp>
</np>
</s>
```

• key-value pairs within XML start tags are accessible in CQP as additional s-attributes with annotated values (marked [A] in the show cd; listing): s_len, np_h, np_len, pp_h, pp_len (cf. Section 1.2)

• s-attribute values can be accessed through label references
  > <np> a:\[] \[* </np> :: a.np_h = "bank";
  → NPs with head lemma bank
  an equivalent, but shorter version:
  > /region[np,a] :: a.np_h="bank";
  or use the match anchor label automatically set to the first token of the match
  > <np> \[* </np> :: match.np_h="bank";

• constraints on key-value pairs can also directly be tested in start tags, using the appropriate auto-generated s-attribute (make sure to use a matching end tag)
  > <np_h = "bank"> \[* </np_h>;
  comparison operators = and != are supported, together with the %c and %d flags;
  = is the default and may be omitted

• constraints on multiple key-value pairs require multiple start tags
  > <np_h="bank">\[<np_len="[1-6]"> \[* </np_len></np_h>;
  (or access the value of np_len through a label reference)
• `<np>` and `<pp>` tags are usually shown without XML attribute values; they can be displayed explicitly as `<np_h>`, `<np_len>`, ... tags:

  > show +np +np_h +np_len;
  > cat;

  (other corpora may show XML attributes in start tags)

• use this label for direct access to s-attribute values within pattern

  > [(pos="NNS?") & (lemma = _.np_h)];

  (recall that np_h would merely return an integer value indicating whether the current token is contained in a `<np>` region, not the desired annotation string)

• typecast numbers to `int()` for numerical comparison

  > /region[np,a] :: int(a.np_len) > 30;

• NB: s-attribute annotations can only be accessed with label references:

  > [np_h="bank"];  does not work!

• regions of structural attributes are non-recursive

  ⇒ embedded XML regions are renamed to `<np1>`, `<np2>`, ... `<pp1>`, `<pp2>`, ... 

• embedding level must be explicitly specified in the query:

  > [pos="CC"] <np1> []* </np1>;

  will only find NPs contained in exactly one larger NP

  (use show +np +np1 +np2; to experiment)

• regions representing the attributes in XML start tags are renamed as well:

  ⇒ `<np_h1>`, `<np_h2>`, ..., `<pp_len1>`, `<pp_len2>`, ...

  > /region[np1, a] :: a.np_h1 = a.np_h within np;

• CQP queries typically use maximal NP and PP regions (e.g. to model clauses)

• find any NP (regardless of embedding level):

  > (<np>|<np1>|<np2>) []* (<np2>|</np1>|</np>);

  CQP ensures that a matching pair of start and end tag is picked from the alternatives

• observe how results depend on matching strategy (see Section 6.1 for details)

  > set MatchingStrategy shortest;
  > set MatchingStrategy longest;
  > set MatchingStrategy standard;

  (re-run the previous query after each `set` and watch out for "duplicate" matches)

• when the query expression shown above is embedded in a longer query, the matching strategy usually has no influence

• annotations of a region at an arbitrary embedding level can only be accessed through constraints on key-value pairs in the start tags:

  > (<np_h "bank">|<np_h1 "bank">|<np_h2 "bank">) []* 
  (</np_h2>|</np_h1>|</np_h>);
4.4 XML document structure

- XML document structure of DICKENS:

```xml
<novel title="A Tale of Two Cities">
  <titlepage> ... </titlepage>
  <book num=1>
    <chapter num=1 title="The Period">
      ...
    </chapter>
    ...
  </book>
  ...
</novel>
```

- Use `set PrintStructures` command to display novel, chapter, ... for each match

  ```
  > set PrintStructures "novel_title, chapter_num";
  > A = [lemma = "ghost"];
  > cat A;
  ```

- Find matches in a particular novel

  ```
  > B = [pos = "NP"] [pos = "NP"] :
     match.novel_title = "David Copperfield";
  > group B matchend lemma by match lemma;
  ```

  (Note that `<novel_title = "...">` cannot be used in this case because the XML start tag of the respective `<novel>` region will usually be far away from the match)

- Frequency distributions can also be computed for `s-attribute` values

  ```
  > group A match novel_title;
  ```
5 Working with aligned corpora

All examples in this section are based on EuroParl v3, a parallel corpus of debates of the European Parliament that can be downloaded in pre-indexed form from the CWB Web site.

5.1 Displaying aligned sentences

- CWB can encode information about sentence-level alignment between parallel corpora in its index. For each pair of source and target corpus, only a single alignment may be defined; the name of the corresponding alignment attribute (\texttt{a-attribute}) is a lowercase version of the CWB name of the target corpus.

- For example, the English component of the EuroParl corpus
  \begin{verbatim}
  > EUROPARL-EN;
  \end{verbatim}
is aligned to the French (\texttt{EUROPARL-FR}) and German (\texttt{EUROPARL-DE}) components \textit{inter alia}.

- The available alignment attributes are listed as “Aligned Corpora:” in the output of \texttt{show cd};.

- One or more alignments can be displayed in the KWIC output produced by \texttt{cat}. For example, in order to find out how the idiom \textit{take the biscuit} can be expressed in French and German, we activate the corresponding a-attributes:
  \begin{verbatim}
  > show +europarl-fr +europarl-de;
  > [lemma="take"] "the" "biscuit";
  \end{verbatim}
  Note that the target languages are always printed in the same order as in the \texttt{show cd}; output. In this example, the German translation will be shown first, followed by the French translation.

- It is recommended to set the KWIC context for the source language to a full sentence
  \begin{verbatim}
  > set Context 1 s;
  \end{verbatim}
  However, translations are always displayed as complete \textit{alignment beads}, which can be confusing if multiple sentences in the source language are translated into a single target sentence (a 2:1 bead) or divided in a different way in the target language (a 2:2 bead). For example, the single hit of the query
  \begin{verbatim}
  > "price-tag";
  \end{verbatim}
is translated 1:1 into French, but combined with the previous sentence in the German translation (resulting in a 2:1 bead).

- An unofficial feature allows setting the KWIC context to an a-attribute, ensuring that a complete alignment bead (for the selected target corpus) is displayed.
  \begin{verbatim}
  > set Context europarl-de;
  > cat;
  \end{verbatim}
  Keep in mind that this has been implemented as a special case: a-attributes cannot be used as context specifiers elsewhere (e.g. in a \texttt{within} clause).

- You will find that some sentences have no translation into the target language, e.g.
  \begin{verbatim}
  > "cats" cut 6;
  \end{verbatim}
  Notice that the alignment KWIC context shows only the matching string without surrounding words in this case.
In order to exclude matches outside alignment beads (i.e. without a translation), you can add a trivial alignment constraint to the query (cf. Sec. 5.2). The example below shows that out of 49 occurrences of *cats*, only 46 have a translation into French:

```cqp
> AllCats = "cats";
> size AllCats;
> GoodCats = "cats" :EUROPARL-FR [];
> size GoodCats;
```

### 5.2 Querying aligned corpora

This section explains how alignment information can be used as a filter in CQP queries. As a first example, let us consider the word *nuclear power*, which can be translated into German as Kernkraft, Kernenergie, Atomkraft or Atomenergie.

```cqp
> Nuke = "nuclear"%c "power"%c;
```

Instead of manually perusing the translations of all 1417 hits, we can directly search for hits that contain one of the relevant words in the German translation.

```cqp
> "nuclear"%c "power"%c :EUROPARL-DE [lemma = "Kernkraft"];
> "nuclear"%c "power"%c :EUROPARL-DE [lemma = "Kernenergie "];
etc.
```

An alignment constraint consists of the marker :TARGET-CORPUS followed by an arbitrary query expression. CQP will scan the region aligned to each match of the main query and keep only those for which a match of the alignment constraint is found.

- The alignment constraint is always specified after the main query (including the within clause), but before a cut statement (which applies to the filtered query results). Multiple alignment constraints can be chained and must all be satisfied.
- Alignment constraints can be negated by placing ! immediately after the marker. In this case, only those matches are kept for which the alignment constraint is not satisfied.

```cqp
> Other = "nuclear"%c "power"%c :EUROPARL-DE ! "(Kern|Atom).*";
```

Can you figure out how *nuclear power* is translated in these examples?

- By chaining negated constraints, we can identify cases where the French translation is also different from the expected nucléaire:
  ```cqp
  > "nuclear"%c "power"%c :EUROPARL-DE! "(Kern|Atom).*" :EUROPARL-FR! "nucléaire.*" %cd;
  ```
- An alignment constraint can never be satisfied for a match that has no translation into the target corpus, regardless of whether it is negated or not. Therefore,
  ```cqp
  > "cats" :EUROPARL-FR ! [];
  ```

returns no results at all. If you need to find unaligned instances of *cats*, you can only do so in a two-step process. Using the NQR *AllCats* and *GoodCats* (with translation) from above:

```cqp
> BadCats = diff AllCats GoodCats;
> size BadCats;
```

- Note that alignment constraints can only be added to regular CQP queries, not to MU queries (which are an undocumented feature anyway).
5.3 “Translating” query results

- A named query result can be “translated” to an aligned corpus, which allows more flexible display of the aligned regions, access to metadata, etc. (new in CQP v3.4.9).

- Consider the following example:
  > EUROPARL-DE;
  > set Context 1 s;
  > Zeit = [lemma = "Zeit"];

- The NQR Zeit now contains all occurrences of the German word for time in the German part of EuroParl. The following command “translates” the NQR to the English part of EuroParl, i.e. it replaces each match by the complete aligned region in the target corpus (as would be displayed with `show +europarl-en;`):
  > Time = from Zeit to EUROPARL-EN;

- This creates a new NQR EUROPARL-EN:Time containing the aligned regions. You can now e.g. tabulate or count metadata:
  > tabulate EUROPARL-EN:Time match text_date;
  > group EUROPARL-EN:Time match text_date;

- The somewhat arcane syntax of the command avoids introduction of a new reserved keyword
  - while it looks similar to a corpus query or set operation, the assignment to a new NQR is mandatory (otherwise the parser won't accept the syntax)
  - note that the new NQR must be specified as a short name; the name of the target corpus is implied and added automatically with the assignment

- Some important details:
  - matching ranges that are not aligned to the target corpus are silently discarded; you cannot expect the new NQR to contain the same number of hits as the original NQR
  - if there are multiple matches in the same alignment head, they will not be collapsed in the target corpus; i.e. the new NQR will contain several identical ranges
  - in order to collate source matches with the aligned regions, make sure to discard unaligned hits from the original NQR first:
    > Zeit = [lemma = "Zeit"] :EUROPARL-EN [];
    or post-hoc as a subquery filter
    > Zeit;
    > ZeitAligned = <match> [] :EUROPARL-EN [] !;

- Do not `cat` the translated query directly (`cat EUROPARL-EN:Time;`) without first activating the target corpus, as this would corrupt the context descriptor (cf. Sec. 3.1). The correct procedure is
  > EUROPARL-EN;
  > cat Time;

  You can now customize the KWIC display as desired.

- It is safe to apply `dump`, `tabulate`, `group`, `count` and similar operations. Only commands that auto-print the NQR (including a bare `sort` or a set operation) will trigger the bug.

- The problem is mentioned in this section because users are most likely to be tempted to do this when working with a set of aligned corpora.
• As a second example, we will return to German translations of nuclear power.

  > EUROPARL-DE;
  > Other = from EUROPARL-EN:Other to EUROPARL-DE;

• We can now run a subquery on the aligned regions in the German part of EuroParl in order search for possible translations other than Kern- and Atom-. One possibility is that nuclear power plant has been translated into the acronym AKW (for Atomkraftwerk).

  > Other;
  > [lemma = "AKW"];

• Further translation candidates can be found by computing a frequency breakdown of all nouns in the aligned sentences:

  > N = [pos = "N.*"];
  > group N match word;

• We could have applied the same strategy to the NQR Nuke in order to determine the frequencies of different translation equivalents:

  > Nuke = from EUROPARL-EN:Nuke to EUROPARL-DE;
  > Nuke;
  > TEs = "(Atom|Kern|AKW).*";
  > group TEs match lemma;
6 Advanced CQP features

6.1 The matching strategy

- set MatchingStrategy (shortest | standard | longest);

- in shortest mode, ?, *, and + operators match smallest number of tokens possible (refers to regular expressions at token level)
  ⇒ finds shortest sequence matching query,
  ⇒ optional elements at the start or end of the query will never be included

- in longest mode, ?, *, and + operators match as many tokens as possible

- in the default standard mode, CQP uses an “early match” strategy: optional elements at the start of the query are included, while those at the end are not

- the somewhat inconsistent matching strategy of earlier CQP versions is currently still available in the traditional mode, and can sometimes be useful (e.g. to extract cooccurrences between multiple adjectives in a noun phrase and the head noun)

  > [pos="JJ"]+[pos="NNS?"];
  > group Last matchend lemma by match lemma;
  only gives the intended frequency counts in traditional mode

- Figure 4 shows examples for all four matching strategies

- new in CQP v3.4.12: The matching strategy can be set temporarily with an embedded modifier at the start of a CQP query, e.g.

  > (?longest) [pos = "NP.*"]+;

  Currently, only these four modifiers are supported: (?shortest), (?standard), (?longest) and (?traditional). Embedded modifiers are particularly useful for Web interfaces that do not give users direct control over the matching strategy. Since they are part of the CQP query syntax, no modifications to existing Web interfaces are required.

6.2 Word lists

- word lists can be stored in variables

  > define $week = "Monday Tuesday Wednesday Thursday Friday";
  and used instead of regular expressions in the attribute/value pairs

  > [lemma = $week];

  (word lists are not allowed in XML start tags, though)

- add/delete words with += and -=

  > define $week += "Saturday Sunday";

- show list of words stored in variable

  > show $week;
  use show var; to see all variables
search pattern:

```
DET? ADJ* NN (PREP DET? ADJ* NN)*
```

input:

```
the old book on the table in the room
```

shortest match strategy: (3 matches)

```
⊿ book
⊿ table
⊿ room
```

longest match strategy: (1 match)

```
⊿ the old book on the table in the room
```

standard matching strategy: (3 matches)

```
⊿ the old book
⊿ the table
⊿ the room
```

traditional matching strategy: (7 overlapping matches)

```
⊿ the old book
⊿ old book
⊿ book
⊿ the table
⊿ table
⊿ the room
⊿ room
```

Figure 4: CQP matching strategies.
• read word list from file (one-word-per-line format)
  > define $week < "/home/weekdays.txt";

new in CQP v3.4.11: files ending in .gz or .bz2 are automatically decompressed, and word lists
   can be read from a shell pipe indicated by a | character at the start of the filename; for example,
   to read a file with whitespace-delimited words (and multiple entries per line):
  > define $week < "| perl -pe 's/\s+\n/g' words.txt";

• use TAB key to complete word list names (e.g. type "show $we" + TAB)

• word lists can be used to simulate type hierarchies, e.g. for part-of-speech tags
  > define $common_noun = "NN NNS";
  > define $proper_noun = "NP NPS";
  > define $noun = $common_noun;
  > define $noun += $proper_noun;

• %c and %d flags can not be used with word lists

• use lists of regular expressions with RE() operator (compile regex)
  > define $pref="under.+ over.+";
  > [(lemma=RE($pref)) & (pos="VBG")];

• flags can be appended to RE() operator
  > [word = RE($pref) %cd];

6.3 Subqueries

• queries can be limited to the matching regions of a previous query (⇒ subqueries)

• activate named query instead of system corpus (here: sentences containing interest)
  DICKENS> First = [lemma = "interest"] expand to s;
  DICKENS> First;
  DICKENS:First[624]>

NB: matches of the activated query must be non-overlapping.

• the matches of the named query First now define a virtual structural attribute on the corpus
  DICKENS with the special name match

• all following queries are evaluated with an implicit within match clause
  (an additional explicit within clause may be specified as well)

• re-activate system corpus to exit subquery mode
  DICKENS:First[624] > DICKENS;
  DICKENS>

• XML tag notation can also be used for the temporary match regions
  > <match> [pos = "W.*"];

6 Overlapping matches may result from the traditional matching strategy, set operations, or modification of
   the matching word sequences with expand, set match, or set matchend. When a named query with overlapping matches
   is activated, a warning message is issued and some of the matches will be automatically deleted.
• if target/keyword anchors are set in the activated query result, the corresponding XML tags
  (\textlt{<target>}, \textlt{<keyword>}, ...) can be used, too

\begin{verbatim}
> </target> [ ]* </match>
\end{verbatim}

\begin{itemize}
\item range from target anchor to end of match, but excluding target
\item \textlt{<target>} and \textlt{<keyword>} regions always have length 1!
\end{itemize}

• a subquery that \textit{starts} with an anchor tag is evaluated very efficiently

• appending the \textit{keep} operator \textit{!} to a subquery returns full matches from the activated query result
  (equivalent to an implicit \textit{expand to match})

6.4 The CQP macro language

• complex queries (or parts of queries) can be stored as macros and re-used

• define macros in text file (e.g. \textit{macros.txt}):

\begin{verbatim}
# this is a comment and will be ignored
MACRO np(0)
  \[pos = "DT"]  # another comment
  ([pos = "RB.*"]? [pos = "JJ.*"])*
  [pos = "NNS?"]
;
\end{verbatim}

(defines macro “np” with no arguments)

• load macro definitions from file

\begin{verbatim}
> define macro < "macros.txt";
\end{verbatim}

• macro invocation as part of a CQP command (use \texttt{TAB} key for macro name completion)

\begin{verbatim}
> <s> /np[] @[pos="VB.*"] /np[];
\end{verbatim}

• list all defined macros or those with given prefix

\begin{verbatim}
> show macro;
> show macro region;
\end{verbatim}

• show macro definition
  (you must specify the number of arguments)

\begin{verbatim}
> show macro np(0);
\end{verbatim}

• re-define macro interactively (must be written as a single line)

\begin{verbatim}
> define macro np(0) '[pos="DT"] [pos="JJ.*"]+ [pos="NNS?"]';
\end{verbatim}

or re-load macro definition file

\begin{verbatim}
> define macro < "macros.txt";
\end{verbatim}

• macros are interpolated as plain strings (not as elements of a query expression) and may have to
  be enclosed in parentheses for proper scoping

\begin{verbatim}
> <s> (/np[])+ [pos="VB.*"];
\end{verbatim}

• it is safest to put parentheses around macro definitions:

\begin{verbatim}
> <s> (/np[])* [pos="VB.*"];
\end{verbatim}
MACRO np(0)
(
    [pos = "DT"]
    ([pos = "RB.*"]? [pos = "JJ.*"])*
    [pos = "NNS?"]
)
;

NB: The start (MACRO ...) and end (;) markers must be on separate lines in a macro definition file.

- macros accept up to 10 arguments; in the macro definition, the number of arguments must be specified in parentheses after the macro name
- in the macro body, each occurrence of $0$, $1$, ... is replaced by the corresponding argument value (escapes such as \$1 will not be recognised)
- e.g. a simple PP macro with 2 arguments: the initial preposition and the number of adjectives in the embedded noun phrase
  
  MACRO pp(2)
  
  ([pos = "IN"] & (word="$0")]
  [pos = "DT"]
  [pos = "JJ.*"]{$1}
  [pos = "NNS?"]
  ;

- invoking macros with arguments
  > /pp["under", 2];
  > /pp["in", 3];

- macro arguments are character strings and must be enclosed in (single or double) quotes; they may be omitted around numbers and simple identifiers
- the quotes are not part of the argument value and hence will not be interpolated into the macro body; nested macro invocations will have to specify additional quotes
- define macro with prototype ⇒ named arguments
  MACRO pp ($0=Prep $1=N_Adj)
  ...;

- argument names serve as reminders; they are used by the show command and the macro name completion function (TAB key)
- argument names are not used during macro definition and evaluation
- in interactive definitions, prototypes must be quoted
  > define macro pp(’$0=Prep $1=N_Adj’) ...;

- CQP macros can be overloaded by the number of arguments (i.e. there can be several macros with the same name, but with different numbers of arguments)
• this feature is often used for unspecified or “default” values, e.g.
  MACRO pp($0=Prep, $1=N\_Adj)
  ...
  MACRO pp($0=Prep) (any number of adjectives)
  ...
  MACRO pp() (any preposition, any number of adjs)
  ...

• macro calls can be nested (non-recursive) ⇒ macro file defines a context-free grammar (CFG)
  without recursion (see Figure 5)

```
MACRO adjp()
  [pos = "RB.*"]?
  [pos = "JJ.*"]
;

MACRO np($0=N\_Adj)
  [pos = "DT"]
  ( /adjp[] ){$0}
  [pos = "NNS?"]
;

MACRO np($0=Noun $1=N\_Adj)
  [pos = "DT"]
  ( /adjp[] ){$1}
  [(pos = "NN") & (lemma = "$0")]
;

MACRO pp($0=Prep $1=N\_Adj)
  [(word = "$0") & (pos = "IN|TO")]
  /np[$1]
;
```

Figure 5: A sample macro definition file.

• Macro definition files can import other macro definition files using statements of the form
  IMPORT other_macros.txt

  Each import statement must be written on a separate line. It is recommended (but not required)
  to collect all IMPORTs at the top of the file. Note that files are searched relative to the CQP
  working directory, not the location of the current macro file.

• note that string arguments need to be quoted when they are passed to nested macros (since
  quotes from the original invocation are stripped before interpolating an argument)

• single or double quote characters in macro arguments should be avoided whenever possible; while
  the string ‘s can be enclosed in double quotes (“’s”) in the macro invocation, the macro body
  may interpolate the value between single quotes, leading to a parse error

• in macro definitions, use double quotes which are less likely to occur in argument values
6.5 CQP macro examples

- use macros for easier access to embedded noun phrases (NP)
- write and load the macro definition file shown in Figure 6

```cqp
MACRO np_start()
    (<np>|<np1>|<np2>)
;
MACRO np_end()
    (</np2>|</np1>|</np>)
;
MACRO np()
    ( /np_start[] []* /np_end[] )
;
```

Figure 6: Macro definition file for accessing embedded noun phrases.

- then use /np_start[] and /np_end[] instead of <np> and </np> tags in CQP queries, as well as /np[] instead of /region[np]
  > /np_start[] /np[] "and" /np[] /np_end[];
- CQP ensures that the “generalised” start and end tags nest properly
  (if the StrictRegions option is enabled, cf. Sections 4.2 and 4.3)
- extending built-in macros: view definitions
  > show macro region(1);
  > show macro codist(3);
- extend /region[] macro to embedded regions:

```cqp
MACRO anyregion($0=Tag)
    (<$0>|<$01>|<$02>)
    []*
    (</$02>|</$01>|</$0>)
;
```

- extend /codist[] macro to two constraints:

```cqp
MACRO codist($0=Att1 $1=V1 $2=Att2 $3=V2 $4=Att3)
    _Results = [($0 = "$1") & ($2 = "$3")];
group _Results match $4;
discard _Results;
;
```

- usage examples:
  > "man" /anyregion[pp];
  > /codist[lemma, "go", pos, "V.*", word];

- the simple string interpolation of macros allows some neat tricks,
  e.g. a region macro with constraints on key-value pairs in the start tag
6.6 Feature set attributes (GERMAN-LAW)

- feature set attributes use special notation, separating set members by | characters
- e.g. for the alemma (ambiguous lemma) attribute
  | Zeug | Zeuge | Zeugen | (three elements)
  | Baum | (unique lemma)
  |        (not in lexicon)
- ambiguity() function yields number of elements in set (its cardinality)
  > [ambiguity(alemma) > 3];
- use contains operator to test for membership
  > [alemma contains "Zeuge"];
  → words which can be lemmatised as Zeuge
- test non-membership with not contains
  (alemma not contains "Zeuge")
  ⇔ !(alemma contains "Zeuge")
- also used to annotate phrases with sets of properties
  > /region[np, a] :: a.np_f contains "quot";
- see Appendix A.3 for lists of properties annotated in the GERMAN-LAW corpus
- define macro for easy experimentation with property features
  > define macro find('"$0=Tag $1=Property"')
    <$0_f contains "$1"> []* </$0_f> ;
  > /find[np, brac];
  > /find[advp, temp];
  etc.
- nominal agreement features of determiners, adjectives and nouns are stored in the agr attribute, using the pattern shown in Figure 7 (see Figure 8 for an example)

  "case : gender : number : determination"

  case         Nom, Gen, Dat, Akk
  gender       M, F, N
  number       Sg, Pl
  determination Def, Ind, Nil

Figure 7: Annotation of noun agreement features in the GERMAN-LAW corpus.
• match set members against regular expression
  > [ (pos = "NN") & (agr matches ".*:Pl:*") ];
  → nouns which are uniquely identified as plurals

• both contains and matches use regular expressions and accept the %c and %d flags

• unification of agreement features ⇐⇒ intersection of feature sets

• use built-in /unify[] macro:
  /unify[agr, <label1>, <label2>, ...]

• undefined labels will automatically be ignored
  > a:[pos="ART"] b:[pos="ADJA"]? c:[pos="NN"]
     :: /unify[agr, a,b,c] matches "Gen:*";
  → (simple) NPs uniquely identified as genitive

• use ambiguity() function to find number of possible analyses
  > ... :: ambiguity(/unify[agr, a,b,c]) >= 1;
  → to check agreement within NP

• in the GERMAN-LAW corpus, NPs and other phrases are annotated with partially disambiguated agreement information; these features sets can also be tested with the contains and matches operators, either indirectly through label references or directly in XML start tags
  > /region[np, a] :: a.np_agr matches "Dat:.Pl:*";
  > <np_agr matches "Dat:.Pl:*"> []* </np_agr>;

• in order to improve computational efficiency, /unify[] expects features sets in canonical format, with members sorted according to CWB’s internal sort order; this is usually ensured with the -m option to cwb-s-encode

• even if an attribute hasn’t explicitly been defined as a feature set (and converted to canonical format), ambiguity(), contains and matches are guaranteed to work as long as the 1-separated set notation is used correctly and consistently

• however, the /unify[] macro cannot be used unless the features within each set are in canonical sorted order. The members of a set are sorted at indexing-time only when a feature set is explicitly declared.

• feature sets should not be used to encode ordered lists of values; if you need to distinguish between a first, second, ... alternative, you might add this information explicitly as a feature component, e.g.
7 Interfacing CQP with other software

7.1 Running CQP as a backend

- CQP is a useful tool for interactive work, but many tasks become tedious when they have to be carried out by hand; macros can be used as templates, providing some relief; however, full scripting is still desirable (and in some cases essential)

- similarly, the output of CQP requires post-processing at times: better formatting of KWIC lines (especially for HTML output), different sort options for frequency tables, frequency counts on normalised word forms (or other transformations of the values)

- for both purposes, an external scripting tool or programming language is required, which has to interact dynamically with CQP (which acts as a query engine)

- CQP provides some support for such interfaces: when invoked with the -c flag, it switches to child mode (which could also be called “slave” mode):
  - the init file ~/.cqprc is not automatically read at startup
  - CQP prints its version number after initialisation
  - all interactive features are deactivated (paged display and highlighting)
  - query results are not automatically displayed (set AutoShow off;)
  - after the execution of a command, CQP flushes output buffers (so that the interface will not hang waiting for output from the command)
  - in case of a syntax error, the string PARSE ERROR is printed on stderr
  - the special command .EOL.; inserts the line
    -::-EOL-::- as a marker into CQP’s output
  - when the ProgressBar option is activated, progress messages are not echoed in a single screen line (using carriage returns) on stderr, but rather printed in separate lines on stdout; these lines have the standardized format
    -::-PROGRESS-::- TAB pass TAB no. of passes TAB progress message

- the CWB/Perl interface makes use of all these features to provide an efficient and robust interface between a Perl script and the CQP backend

- the output of many CQP commands is neatly formatted for human readers; this pretty printing feature can be switched off with the command
  > set PrettyPrint off;
the output of the show, group and count commands now has a simple and standardized format that can more easily be parsed by the invoking program; output formats for the different uses of the show command are documented below; see Section 7.3 for the output formats of group and count

  - show corpora; prints the names of all available corpora on separate lines, in alphabetical order
  - show named; lists all named query results on separate lines in the format

  flags TAB query name TAB no. of matches
flags is a three-character code representing the flags m = stored in memory, d = saved to disk, * = modified since last saved; flags that are not set are shown as - characters; query name is the long name of the query result, i.e. it has the form corpus:name; when a query result has not yet been loaded from disk, the no. of matches cannot be determined and is reported as 0

- show; concatenates the output of show corpora; and show named; without any separator; it is recommended to invoke the two commands separately when using CQP as a backend

- show active; prints the name of the currently active corpus on a line on its own (this is in fact available when using CQP interactively, albeit useless because the active corpus is displayed in the CQP command prompt!)

- show cd; lists all attributes that are defined for the currently active corpus; each attribute is printed on a separate line with the format

  attribute type TAB attribute name [ TAB [ -V ] ]

  attribute type is one of the strings p-Att (positional attribute), s-Att (structural attribute)
  or a-Att (alignment attribute), so the attribute type can easily be recognized from the first
  character of the output line; the third column is only printed for s-attributes and is either
  an empty string (no annotations) or -V (regions have annotated values)

- the CWB/Perl interfaces automatically deactivates pretty printing

- running CQP as a backend can be a security risk, e.g. when queries submitted to a Web server
  are passed through to the CQP process unaltered; this may allow malicious users to execute
  arbitrary shell commands on the Web server; as a safeguard against such attacks, CQP provides
  a query lock mode, which allows only queries to be executed, while all other commands (including
  cat, sort, group, etc.) are blocked

- the query lock mode is activated with the command

  > set QueryLock n;
  
  where n is a randomly chosen integer number; it can only be de-activated with
  > unlock n;
  
  using the same number n

7.2 Exchanging corpus positions with external programs

- An important aspect of interfacing CQP with other software is to exchange the corpus positions
  of query matches (as well as target and keyword anchors). This is a prerequisite for the extraction
  of further information about the matches by direct corpus access, and it is the most efficient
  way of relating query matches to external data structures (e.g. in a SQL database or spreadsheet
  application).

- The dump command (Section 3.3) prints the required information in a tabular ASCII format that
  can easily be parsed by other tools or read into a SQL database. Each row of the resulting table
  corresponds to one match of the query, and the four columns give the corpus positions of the
  match, matchend, target and keyword anchors, respectively. The example below is reproduced
  from Section 3.3

  Since this command dumps the matches of a named query in their current sort order, the natural order should first be restored by calling sort without a by clause. One exception is a CGI interface that uses the dumped corpus positions for a KWIC display of the query results in their sorted order.
Undefined target anchors are represented by -1 in the third column. Even though no keywords were set for the query, the fourth column is included in the dump table, but all values are set to -1.

- The table created by the dump command is printed on stdout by default, where it can be captured by a program running CQP as a backend (e.g. the CWB/Perl interface, cf. Sec. 7.1). The dump table can also be redirected to a file:
  > dump A > "dump.tbl";
  which is automatically compressed if the filename ends in .gz or .bz2 (new in CQP v3.4.11).

- Alternatively, the output can also be redirected to a pipe, e.g. to create a dump file without the superfluous keyword column
  > dump A > "| cut -f 1-3 > dump.tbl";
  In earlier versions of CQP, such pipes were also needed to compress the dump file on the fly
  > dump A > "| gzip > dump.tbl.gz";

- Sometimes it is desirable to reload a dump file into CQP after it has been modified by an external program (e.g. a database may have filtered the matches against a metadata table). The undump command creates a new named query result (B in the example below) for the currently activated corpus from a dump file (which may be a compressed file in CQP v3.4.11 and newer):
  > undump B < "mydump.tbl";
  Note that B is silently overwritten if it already exits.

- The format of the file mydump.tbl is almost identical to the output of dump, but it contains only two columns for the match and matchend positions (in the default setting). The example below shows a valid dump file for the DICKENS corpus, which can be read with undump to create a query result containing 5 matches:

```
20681 20687
379735 379741
1915978 1915983
2591586 2591591
2591593 2591598
```

Save these lines to a text file named dickens.tbl, then enter the following commands:

  > DICKENS;
  > undump Twas < "dickens.tbl";
  > cat Twas;

- Further columns for the target and keyword anchors (in this order) can optionally be added. In this case, you must append the modifier with target or with target keyword to the undump command:
  > undump B with target keyword < "mydump.tbl";
• Dump files can also be read from a pipe or from standard input. In the latter case the table of corpus positions has to be preceded by a header line that specifies the total number of matches:

```
5
20681 20687
379735 379741
1915978 1915983
2591586 2591591
2591593 2591598
```

CQP uses this information to pre-allocate internal storage for the query result, as well as to validate the file format. This format can also be used as a more efficient alternative if the dump is read from a regular file. CQP automatically detects which of the two formats is used.

• Pipes can e.g. be used to read a dump table generated by another program. They are indicated by a pipe symbol (|) at the start of the filename (new in CQP v3.4.11) or at the end of the filename (earlier versions). Before CQP v3.4.11, pipes were also needed to read a dump table from a compressed file:

```
> undump B < "gzip -cd mydump.tbl.gz |";
```

• In an interactive CQP session, the input file can be omitted and the undump table can then be entered directly on the command line. This feature works only if command-line editing support is enabled with the -e switch. Since the dump table is read from standard input here, only the second format is allowed, i.e. you have to enter the total number of matches first. Try entering the example table above after typing

```
> undump B;
```

• If the rows of the undump table are not sorted in their natural order (i.e. by corpus position), they have to be re-ordered internally so that CQP can work with them. However, the original sort order is recorded automatically and will be used by the cat and dump commands (until it is reset by a new sort command). If you sort a query result A, save it with dump to a text file, and then read this file back in as named query B, then A and B will be sorted in exactly the same order.

• In many cases, overlapping or unsorted matches are not intentional but rather errors in an automatically generated dump table. In order to catch such errors, the additional keyword ascending (or asc) can be specified before the < character:

```
> undump B with target ascending < "mydump.tbl";
```

This command will abort with an error message (indicating the row number where the error occurred) unless the corpus matches in mydump.tbl are non-overlapping and sorted in corpus order.

• A typical use case for dump and undump is to link CQP queries to corpus metadata stored in an external SQL database. Assume that a corpus consists of a large collection of transcribed dialogues, which are marked as <dialogue> regions. A rich amount of metadata (about the speakers, setting, topic, etc.) is available in a SQL database. The database entries can be linked directly to the <dialogue> regions by recording their start and end corpus positions in the database. The following commands generate a dump table with the required information, which can easily be loaded into the database (ignoring the third and fourth columns of the table):
Corpus queries will often be restricted to a subcorpus by specifying constraints on the metadata. Having resolved the metadata constraints in the SQL database, they can be translated to the corresponding regions in the corpus (again represented by start and end corpus position). The positions are then sorted in ascending order and saved to a TAB-delimited text file. Now they can be loaded into CQP with the `undump` command, and the resulting query result can be activated as a subcorpus for following queries. It is recommended to specify the `ascending` option in order to ensure that the loaded query result forms a valid subcorpus:

```bash
> undump SubCorpus ascending < "subcorpus.tbl";
> SubCorpus;
Subcorpus[..]> A = ... ;
```

### 7.3 Generating frequency tables

- For many applications it is important to compute frequency tables for the matching strings, tokens in the immediate context, attribute values at different anchor points, different attributes for the same anchor, or various combinations thereof.

- Frequency tables for the matching strings, optionally normalised to lowercase and extended or reduced by an offset, can easily be computed with the `count` command (cf. Sections 2.9 and 3.3); when pretty-printing is deactivated (cf. Section 7.1), its output has the form

  \[
  \text{frequency TAB first line TAB string (type)}
  \]

- Advantages of the `count` command:
  - Strings of arbitrary length can be counted
  - Frequency counts can be based on normalised strings (%cd flags)
  - The instances (tokens) for a given string type can easily be identified, since the underlying query result is automatically sorted by the `count` command, so that these instances appear as a block starting at match number `first line`

- An alternative solution is the `group` command (cf. Section 3.4), which computes frequency distributions over single tokens (i.e. attribute values at a given anchor position) or pairs of tokens (recall the counter-intuitive command syntax for this case); when pretty-printing is deactivated, its output has the form

  \[
  [ \text{attribute value TAB } ] \text{attribute value TAB frequency}
  \]

- Advantages of the `group` command:
  - Can compute joint frequencies for non-adjacent tokens
  - Faster when there are relatively few different types to be counted
  - Supports frequency distributions for the values of s-attributes

- The advantages of these two commands are for the most part complementary (e.g., it is not possible to normalise the values of s-attributes, or to compute joint frequencies of two non-adjacent multi-token strings); in addition, they have some common weaknesses, such as relatively slow execution, no options for filtering and pooling data, and limitations on the types of frequency distributions that can be computed (only simple joint frequencies, no nested groupings)
• new in CQP v3.4.9: The group command has been re-implemented with a hash-based algorithm. It is very fast now, even for large frequency tables. The other limitations still apply, though.

• therefore, it is often necessary (and usually more efficient) to generate frequency tables with external programs such as dedicated software for statistical computing or a relational database; these tools need a data table as input, which lists the relevant feature values (at specified anchor positions) and/or multi-token strings for each match in the query result; such tables can often be created from the output of cat (using suitable PrintOptions, Context and show settings)

• this procedure involves a considerable amount of re-formatting (e.g. with Unix command-line tools or Perl scripts) and can easily break when there are unusual attribute values in the data; both cat output and the re-formatting operations are expensive, making this solution inefficient when there is a large number of matches

• in most situations, the tabulate command provides a more convenient, more robust and faster solution; the general form is

  > tabulate A column spec, column spec, ...;

  this will print a TAB-separated table where each row corresponds to one match of the query result A and the columns are described by one or more column spec(ification)s

• just as with dump and cat, the table can be restricted to a contiguous range of matches, and the output can be redirected to a file or pipe

  > tabulate A 100 119 column spec, column spec, ...;
  > tabulate A column spec, column spec, ... > "data.tbl";

• each column specification consists of a single anchor (with optional offset) or a range between two anchors, using the same syntax as the sort and count commands; without an attribute name, this will print the corpus positions for the selected anchor:

  > tabulate A match, matchend, target, keyword;

  produces exactly the same output as dump A; when target and keyword anchors are defined for the query result A; otherwise, it will print an error message (and you need to leave out the column specs target and/or keyword)

• when an attribute name is given after the anchor, the values of this attribute for the selected anchor point will be printed; both positional and structural attributes with annotated values can be used; the following example prints a table of novel title, book number and chapter title for a query result from the DICKENS corpus

  > tabulate A match novel_title, match book_num, match chapter_title;

  note that undefined values (for the book_num and chapter_title attributes) are represented by the empty string

• if an anchor point is undefined or falls outside the corpus (because of an offset), tabulate prints an empty string or the corpus position -1 (correct behaviour implemented in v3.4.10)

• a range between to anchor points prints the values of the selected attribute for all tokens in the specified range; usually, this only makes sense for positional attributes; the following example prints the lemma values of 5 tokens to the left and right of each match, which can be used to identify collocates of the matching string(s)


  note that the attribute values for tokens within each range are separated by blanks rather than TABs, in order to avoid ambiguities in the resulting data table
any items in the range that fall outside the bounds of the corpus are printed as empty strings or corpus positions -1; if either the start or end of the range is an undefined anchor, a single empty string or cpos -1 is printed for the entire range (correct behaviour implemented in v3.4.10)

the end position of a range must not be smaller than its start position, so take care to order items properly and specify sensible offsets; in particular, a range specification such as match .. target must not be used if the target anchor might be to the left of the match; the behaviour of CQP in such cases is unspecified

attribute values can be normalised with the flags %c (to lowercase) and %d (remove diacritics); the command below uses Unix shell commands to compute the same frequency distribution as count A by word %c; in a much more efficient manner

> tabulate A match .. matchend word %c > "| sort | uniq -c | sort -nr";

note that in contrast to sort and count, a range is considered empty when the end point lies before the start point and will always be printed as an empty string
8 Undocumented CQP

8.1 Zero-width assertions

- Constraints involving labels have to be tested either in the global constraint or in one of the token patterns; this means that macros cannot easily specify constraints on the labels they define: such a macro would have to be interpolated in two separate places (in the sequence of token patterns as well as in the global constraint).

- Zero-width assertions allow constraints to be tested during query evaluation, i.e. at a specific point in the sequence of token patterns; an assertion uses the same Boolean expression syntax as a pattern, but is delimited by [: ... :] rather than simple square brackets ([ ... ]); unlike an ordinary pattern, an assertion does not “consume” a token when it is matched; it can be thought of as a part of the global constraint that is tested in between two tokens.

- With the help of assertions, NPs with agreement checks can be encapsulated in a macro

```
DEFINE MACRO np_agr(0)
a: [pos="ART"]
b: [pos="ADJA"]*
c: [pos="NN"]
[: ambiguity(/unify[agr, a, b, c]) >= 1 :]
;
```

(this in simple case, the constraint could also have been added to the last pattern)

- When the this label (_) is used in an assertion, it refers to the corpus position of the following token; the same holds for direct references to attributes.

- In this way, assertions can be used as look-ahead constraints, e.g. to match maximal sequences of tokens without activating longest match strategy:

```
> [pos = "NNS?"]{2,} [:pos != "NNS?":];
```

- Assertions also allow the independent combination of multiple constraints that are applied to a single token; for instance, the region(5) macro from Section 6.5 could also have been defined as

```
MACRO region($0=Att $1=Key1 $2=Val1 $3=Key2 $4=Val2)
  <$0> [: _.$0_$1="$2" :] [: _.$0_$3="$4" :] []* </$0>
;
```

- Like the matchall pattern [], the matchall assertion [::] is always satisfied; since it does not “consume” a token either, it is a no-op that can freely be inserted at any point in a query expression; in this way, a label or target marker can be added to positions which are otherwise not accessible, e.g. an XML tag or the start/end position of a disjunction:

```
> ... @[::] /region[np] ... ;
> ... a:[:::] ( ... | ... | ... ) b:[:::] ...;
```

Starting a query with a matchall assertion is extremely inefficient: use the match anchor or the implicit match label instead.

8.2 Labels and scope

- Returning to the np_agr macro from Section 8.1, we note a problem with this query:

```
> A = /np_agr [] [pos = "VVFIN"] /np_agr[];
```
when the second NP does not contain any adjectives but the first does, the b label will still point to an adjective in the first NP; consequently, the agreement check may fail even if both NPs are really valid

- in order to solve this problem, the two NPs should use different labels; for his purpose, every macro has an implicit $$ argument, which is set to a unique value for each interpolation of the macro; in this way, we can construct unique labels for each NP:

  ```
  DEFINE MACRO np_agr(0)
  $$._a:[pos="ART"]
  $$._b:[pos="ADJA"]*
  $$._c:[pos="NN"]
  [: ambiguity(/unify[agr, $$._a,$$._b,$$._c]) >= 1 :]
  ;
  ```

  a comparison with the previous results shows that this version of the /np_agr[] macro finds additional matches that were incorrectly rejected by the first implementation

  ```
  > B = /np_agr[] [pos = "VVFIN"] /np_agr[];
  > diff B A;
  ```

  - however, the problem still persists in queries where the macro is interpolated only once, but may be matched multiple times

  ```
  > A = (/np_agr[]) (3);
  ```

  here, a solution is only possible when the scope of labels can be limited to the body of the macro in which they are defined; i.e., the labels must be reset to undefined values at the end of the macro block; this can be achieved with the built-in /undef[] macro, which resets the labels passed as arguments and returns a true value

  ```
  DEFINE MACRO np_agr(0)
  a:[pos="ART"]
  b:[pos="ADJA"]*
  c:[pos="NN"]
  [: ambiguity(/unify[agr, a,b,c]) >= 1 :]
  [: /undef[a,b,c] :]
  ;
  > B = (/np_agr[]) (3);
  > diff B A;
  ```

  - note that it may still be wise to construct unique label names (either in the form np_agr_a etc., or with the implicit $$ argument) in order to avoid conflicts with labels defined in other macros or in the top-level query

### 8.3 CQP built-in functions

The CQP query language offers a number of built-in functions that can be applied to attribute values within query constraints (but not anywhere else, e.g. in group or tabulate commands). While some of these functions have been available for a long time and are documented in this tutorial, others have been added more recently and may be unsupported or experimental. The list below shows all built-in functions that are currently available. Those marked as experimental are not guaranteed to function correctly and may be changed or disabled in future releases of CQP.
f(att): frequency of the current value of p-attribute att (cannot be used with s-attributes or literal values); e.g. \[\text{word} = \".*able\" \& f(\text{word}) < 10\]

dist(a, b), distance(a, b): signed distance between two tokens referenced by labels a and b; explicit numeric corpus positions may be specified instead of labels; computes the difference \(b - a\); e.g. \[... :: \text{dist(\text{matchend}, \text{match})} \geq 10;\]

distabs(a, b): unsigned distance between two tokens; e.g. \[\text{dist(_, 1000)} \leq 10\]
as an inefficient way to match 10 tokens to the left and right of corpus position 1000

int(str): cast str to a signed integer number so numeric comparisons can be made; raises an error if \(str\) is not a number string; e.g. \[... :: \text{int(\text{match.text_year})} \leq 1900;\]

lbound(att), rbound(att): evaluates to true if current corpus position is the first or last token in a region of s-attribute att, respectively

lbound_of(att, a), rbound_of(att, a): returns the corpus position of the start or end of the region of s-attribute att containing the token referenced by label a, suitable for use with dist(); if a is not within a region of att, an undefined value is returned, which evaluates to false in most contexts [new in v3.4.13; experimental]

unify(fs1, fs2): compute the intersection of two sorted feature sets specified as strings fs1 and fs2, corresponding to a unification of feature bundles; if the first argument is an undefined value, fs2 is returned; see Sec. 6.6 for details

ambiguity(fs): compute the size of a feature set specified as string fs, i.e. the number of elements; if fs is an undefined value, a size of 0 is returned (same as for |); see Sec. 6.6 for details

add(x, y), sub(x, y), mul(x, y): simple arithmetic on integer values x and y, which can also be corpus positions specified as labels; in order to make computations on corpus annotations, they have to be typecast with int() first [experimental]

prefix(str1, str2): returns longest common prefix of strings str1 and str2; warning: this function operates on bytes and may return an incomplete UTF-8 character [experimental]

is_prefix(str1, str2): returns true if string str1 is a prefix of str2; e.g. \[\text{is_prefix(\text{lemma}, \text{word})}\]

minus(str1, str2): removes the longest common prefix of str1 and str2 from the string str1 and returns the remaining suffix; warning: this function operates on bytes and may return an incomplete UTF-8 character [experimental]

ignore(a): ignore the label a and always return true; for internal use by the /undef[] macro, see Sec. 8.2 for details

normalize(str, flags): apply case-folding and/or diacritic folding to the string str and return the normalized value; flags must be a literal string "c", "d" or "cd" (with an optional %, e.g. "%cd")
; e.g. \[\text{normalize(\text{word}, \"cd\")} \neq \text{normalize(\text{lemma}, \"cd\")}\] to find non-trivial differences between word form and lemma; [new in v3.4.11; experimental]

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8.4 MU queries

- CQP offers search-engine like “Boolean” queries in a special meet-union (MU) notation. This feature goes back to the original developer of CWB and is not supported officially. In particular, there is no precise specification of the semantics of MU queries and the original implementation does not produce consistent results.

- New in v3.4.12: Recently, MU queries have found more widespread use as proximity queries in the CEQL simple query syntax of BNCweb and CQPweb, giving them a semi-official status. For this reason, the implementation was modified to ensure a consistent and well-defined behaviour, although it may not always correspond to what is desired intuitively. The new MU implementation is documented here.

- Warning: both the syntax and the semantics of MU queries are subject to fundamental revisions in the next major release of CWB (version 4.0).

- A meet-union query consists of nested meet and union operations forming a binary-branching tree that is written in LISP-like prefix notation. MU queries always start with the keyword MU and are completely separate from standard CQP syntax.

- The simplest form of a MU query specifies a single token pattern, which may also be given in shorthand notation for the default p-attribute. These queries are fully equivalent to the corresponding standard queries.

\[ \text{> MU} \ [\text{lemma} \ = \ "light" \ \& \ \text{pos} \ = \ "V.*"] ; \]
\[ \text{> MU} \ "lights" \ %c ; \]

- A meet clause matches two token patterns within a specified distance of each other. More precisely, instances of the first pattern are filtered, keeping only those where the second pattern occurs within the specified window. For example, the following query finds nouns that co-occur with the adjective lovely:

\[ \text{> MU} (\text{meet} \ [\text{pos} \ = \ "NN.*"] \ [\text{lemma} \ = \ "lovely"] \ -2 \ 2) ; \]
This query returns all nouns for which lovely occurs within two tokens to the left (window starting at offset -2) or right (window ending at offset +2)). The adjective lovely is not included in the match, nor marked in any other way.

- In order to match only prenominal adjectives, we change the window to three tokens preceding the noun (i.e. offsets -3 ... -1):

\[ \text{> MU} (\text{meet} \ [\text{pos} \ = \ "NN.*"] \ [\text{lemma} \ = \ "lovely"] \ -3 \ -1) ; \]

- Since a meet clause returns only occurrences of the first token pattern, we need to change the ordering in order to focus on the adjective rather than the nouns. Don’t forget to adjust the window offsets accordingly!

\[ \text{> MU} (\text{meet} \ [\text{lemma} \ = \ "lovely"] \ [\text{pos} \ = \ "NN.*"] \ 1 \ 3) ; \]
Note that meet operations are not symmetric: this query returns fewer matches than the previous one (viz. those cases where multiple nouns occur near the same instance of lovely).

- Alternatively, we can search for co-occurrence within sentences or other s-attribute regions. Again, the ordering of the token constrains determines whether we focus on tea or cakes.

\[ \text{> MU} \ "tea" \%c \ "cakes" \%c \ s ; \]

- A union clause simply combines the matches of two token patterns into a set union, corresponding to a disjunction (logical or) of the constraints. The following three queries are fully equivalent:
MU queries are relatively powerful because the two elements of a meet or union clause can themselves be complex clauses. For example, the trigram in due course can be found by nesting two meet conditions:

> MU(meet (meet "in" "due" 1 1) "course" 2 2);

The inner clause returns all instances of in that are immediately followed by due; the outer clause requires that the following token (i.e. at an offset of +2 from in) must be course. We can obtain exactly the same result with this query:

> MU(meet "in" (meet "due" "course" 1 1) 1 1);

Now the inner clause determines all occurrences of the bigram due course, but returns only the corpus positions of due, which must appear immediately after in. Can you find two other MU formulations that produce exactly the same results?

> MU(meet (meet "course" "due" -1 -1) "in" -2 -2);

MU queries are less flexible than standard CQP queries, but can be much more efficient for determining co-occurrences at relatively large distances and for combinations of one or more very frequent elements followed by a rare item. For example,

> MU(meet (meet [pos="NN.*"] "virtue" 2 2) "of" 1 1);

is considerably faster than

> [pos = "NN.*"] "of" "virtue";

This query finds sentences that contain both one hand and other hand. The MU query returns only the position of one, which is then expanded to the complete sentence:

> MU(meet "one" "hand" 1 1) (meet "other" "hand" 1 1) s) expand to s;

Combinations of meet and union clauses offer additional flexibility. The following query finds nouns occurring close to a superlative adjective, which can either be synthetic (strangest) or analytic (most extravagant).

> MU(meet [pos="NNS?"] (union [pos="JJS"] (meet [pos="JJ"] "most" -1 -1)) -2 4);

Like standard queries, MU queries can be used as subquery filters (followed by !) or combined with a cut and/or expand clause. However, other elements of standard queries are not supported: labels, target markers (@), zero-width assertions (obviously), global constraints (after ::), alignment constraints and within clauses.

8.5 TAB queries

- new in v3.4.12: tabular (TAB) queries are an obscure undocumented feature of CQP. They were dysfunctional for a long time, but have now been resurrected. The implementation is still considered experimental and might be changed or retired without notice.

- A tab query starts with the keyword TAB and matches a sequence of one or more token patterns with optional flexible gaps. In its simplest form, it corresponds to a standard query matching a fixed sequence of tokens, but is often executed faster. Compare
"in" "due" "course";

This query is both simpler and faster than the MU version shown in Sec. 8.4.

- The most substantial performance gains are achieved for sequences that start with very frequent items and end in a selective token pattern, e.g.

  > TAB [pos = "DT"] [pos = "JJ.*"] "tea";

  TAB queries cannot be used as a general optimization for standard queries, though, because the individual elements cannot be made optional or repeated with a quantifier. It is also not possible to specify alternatives (|) within a TAB query (but you can run multiple queries and take the union of the results).

- The main purpose of tabular queries is to match sequences with flexible gaps. The following two-word TAB query finds cats followed by dogs with a gap of up to two intervening tokens:

  > TAB "cats" {0,2} "dogs";

  It is equivalent to the standard query

  > "cats" []{0,2} "dogs";

  but keep in mind that TAB "cats" []{0,2} "dogs"; would mean something entirely different\(^{11}\)

- Gaps can be specified using any of the repetition operators familiar from standard queries

  \[
  \begin{array}{ll}
    \text{op.} & \text{gap size} \\
    ? & 0 \text{ or 1 token} \\
    * & 0 \text{ or more tokens} \\
    + & 1 \text{ or more tokens} \\
    \{n\} & \text{exactly } n \text{ tokens} \\
    \{n,k\} & \text{between } k \text{ and } n \text{ tokens} \\
    \{n,\} & \text{at least } n \text{ tokens} \\
    \{,k\} & \text{up to } k \text{ tokens (same as } \{0,k\}\} \\
  \end{array}
  \]

  All gap specifications behave as if the repetition operator had been applied to a matchall (\[\]) in a standard query.

- TAB queries can additionally be restricted by a \textit{within} clause. For example, the query

  > TAB "girl" {2} "girl";

  finds a repetition of the noun girl after exactly two intervening tokens, but many of the matches cross a sentence boundary. In order to discard these matches, change the query to

  > TAB "girl" {2} "girl" within s;

- Note that TAB queries do not support different matching strategies, but always use an early match principle similar to the default setting of standard CQP queries (regardless of the value of the \texttt{MatchingStrategy} option). For example, the query

  > TAB [pos = "JJ"] {,5} [pos = "NN"];

  will only match the underlined part of the phrase \textit{a small and very old train station}. It cannot be configured to return the shortest (\textit{old train}) or longest (\textit{small and very old train station}) match.

\(^{11}\)Namely, \textit{cats} followed by an arbitrary token, followed by a gap of up to two tokens, followed by \textit{dogs}. Entering this command will print an error message because matchall patterns are not allowed in TAB queries.
TAB queries always return the full range of tokens containing the specified items. Individual items cannot be marked in any way (i.e. neither as target pattern nor with labels), due to limitations of the current CQP implementation.

For more complex TAB queries, it is important to understand how the greedy matching algorithm works, since its results may be different from the corresponding standard CQP queries:

- for every possible start position, i.e. each match of the first token pattern
- scan for a match of the second token pattern within the specified range
- greedily fix the first such match that is encountered
- starting from this position, scan for a match of the third token pattern
- greedily fix the first such match that is encountered
- etc.

If a complete match is found, CQP continues with the next possible start position, so there can be at most one match for each start position. In addition, nested matches are discarded as in standard CQP queries (hence old train above is actually matched by the algorithm, but then discarded as a nested match).

Always keep in mind that CQP does not perform an expensive combinatorial search to consider other matches that might also fall within the specified ranges!

As a concrete example, consider the sentence

\begin{quote}
Fortunately, we had time\textsubscript{A} for\textsubscript{B1} delicious pastries and for\textsubscript{B2} coffee\textsubscript{C}.
\end{quote}

The TAB query
\begin{verbatim}
> TAB "time" * "for" ? "coffee" within s;
\end{verbatim}

would not match this sentence because "for" is greedily fixed to the first available token B1, for which C is not in range. The corresponding standard query
\begin{verbatim}
> "time" [] * @"for" [] ? "coffee" within s;
\end{verbatim}

considers both options, on the other hand, and matches the range A...C, with the target anchor (@) set to B2.

There are two special cases in which TAB queries are guaranteed to find every early match that satisfies the gap specifications:

1. All gaps have a fixed size (\{n\}), which can be different for each gap. This includes in particular the case where token patterns are directly adjacent.
   \begin{verbatim}
   > TAB "Mr" {1} "Mrs" [pos = "N.*"]; > TAB "in" "due" "course";
   \end{verbatim}

2. All gaps are specified as * and the search range is only restricted by a within clause. Note that * and fixed-size gaps (even direct adjacency) must not be mixed in this case.
   \begin{verbatim}
   > TAB "one" * "two" * "three" within s;
   \end{verbatim}

8.6 Easter eggs

- starting with version 3.0 of the Corpus Workbench, CQP comes with a built-in regular expression optimiser; this optimiser detects simple regular expressions commonly used for prefix, suffix or infix searches such as
> "under.+";
> ".+ment";
> ".+time.+";

and replaces the normal regexp evaluation with a highly efficient Boyer-Moore search algorithm

• the optimiser will also recognise some slightly more complex regular expressions; if you want to test whether a given expression can be optimised or not, switch on debugging output with
  > set CLDebug on;

• some beta releases of CQP may contain hidden optimisations and/or other features that are disabled by default because they have not been tested thoroughly; such hidden features will usually be documented in the release notes and can be activated with the option
  > set Optimize on;

• in particular, beta versions leading up to releases v3.0 and v3.5 disable the regexp optimiser by default until it has been tested thoroughly

• the official LTS releases v3.0 and v3.5 of CQP have no hidden features
A Appendix

A.1 Summary of regular expression syntax

At the character level, CQP supports regular expressions using one of two regex libraries:

**CWB 3.0**: Uses POSIX 1003.2 regular expressions (as provided by the system libraries). A full description of the regular expression syntax can be found on the `regex(7)` manpage.

**CWB 3.5**: Uses PCRE (Perl Compatible Regular Expressions). A full description of the regular expression syntax can be found on the `pcrepattern(3)` manpage; see also [http://www.pcre.org/](http://www.pcre.org/).

Various books such as *Mastering Regular Expressions* give a gentle introduction to writing regular expressions and provide a lot of additional information. There are also many tutorials to be found online using Your Favourite Web Search Engine™.

- A regular expression is a concise description of a set of character strings (which are called *words* in formal language theory). Note that only certain sets of words with a relatively simple structure can be represented in such a way. Regular expressions are said to *match* the words they describe. The following examples use the notation:

  \[
  \text{\texttt{<reg.exp.}> \rightarrow word_1, word_2, \ldots}
  \]

  In many programming languages, it is customary to enclose regular expressions in slashes (/). CQP uses a different syntax where regular expressions are written as (single- or double-quoted) strings. The examples below omit any delimiters.

- Basic syntax of regular expressions

  - letters and digits are matched literally (including all non-ASCII characters)
    \[
    \text{\texttt{word} \rightarrow \texttt{word}; C3PO \rightarrow C3PO; déjà \rightarrow déjà}
    \]
  - . matches any single character ("matchall")
    \[
    \text{\texttt{rg} \rightarrow \texttt{ring, rung, ranga, r3ng, r3ng, \ldots}}
    \]
  - character set: \[[\ldots]\] matches any of the characters listed
    \[
    \text{\texttt{moderni[sz]e} \rightarrow \texttt{modernise, modernize}}
    \]
  - repetition of the preceding element (character or group):
    \[
    \text{\texttt{?} (0 or 1), \texttt{*} (0 or more), \texttt{+} (1 or more), \texttt{n} \texttt{\{n\}} (exactly n), \texttt{n,m} \texttt{\{n..m\}}}
    \]
    \[
    \text{\texttt{color} \rightarrow \texttt{color, colour}; \texttt{go\{2,4\}d} \rightarrow \texttt{good, goed, goood}}
    \]
  - grouping with parentheses: \((\ldots)\)
    \[
    \text{\texttt{(bla)}+ \rightarrow \texttt{bla, blabla, blablabla, \ldots}}
    \]
    \[
    \text{(school){bus(es)}? \rightarrow \texttt{bus, bases, schoolbus, schoolbuses}}
    \]
  - | separates alternatives (use parentheses to limit scope)
    \[
    \text{\texttt{mouse|mice} \rightarrow \texttt{mouse, mice}; \texttt{corp(us|ora)} \rightarrow \texttt{corpus, corpora}}
    \]

- Complex regular expressions can be used to model (regular) inflection:

  - \text{\texttt{ask(s|ed|ing)?} \rightarrow \texttt{ask, asks, asked, asking}}
    (equivalent to the less compact expression \texttt{ask|asks|asked|asking})
  - \text{\texttt{sa(y|ing)?|id}} \rightarrow \texttt{say, says, saying, said}
  - \text{\texttt{[a-z]+i[sz]e[sd]?iing}} \rightarrow \texttt{any form of a verb with -ise or -ize suffix}
• Backslash (\) “escapes” special characters, i.e. forces them to match literally
  - \? → ?; \(\) → (); .{3} → ...; \$\→ $.
  - ^ and \$ must be escaped although ^ and $ anchors are not useful in CQP
A.2 Part-of-speech tags and useful regular expressions

The English PENN tagset (DICKENS)

- **NN**: Common noun, singular or mass noun
- **NNS**: Common noun, plural
- **NP, NPS**: Proper noun, singular/plural
- **N.***: Matches any common or proper noun
- **PP.***: Matches any pronoun (personal or possessive)
- **JJ**: Adjective
- **JJR, JJS**: Adjective, comparative/superlative
- **VB.***: Matches any verbal form
- **VBG, VGN**: Present/past participle
- **RB**: Adverb
- **RBR, RBS**: Adverb, comparative/superlative
- **MD**: Modal
- **DT**: Determiner
- **PDT**: Predeterminer
- **IN**: Preposition, subordinating conjunction
- **CC**: Coordinating conjunction
- **TO**: Any use of “to”
- **RP**: Particle
- **WP**: Wh-pronoun
- **WDT**: Wh-determiner
- **SENT**: Sentence-final punctuation

The German STTS tagset (GERMAN-LAW)

- **NN**: Common noun (singular or plural)
- **NE**: Proper noun (singular or plural)
- **N.**: Matches any nominal form
- **PP.***: Matches any pronoun (personal or possessive)
- **ADJA**: Attributive adjective
- **ADJD**: Predicative adjective (also when used adverbially)
- **ADJ.**: Matches any adjectival form
- **VV.***: Matches any full verb
- **VA.***: Matches any auxiliary verb
- **VM.**: Matches any modal verb
- **V.***: Matches any verbal form
- **ADV**: Adverb
- **ART**: Determiner
- **APPR**: Preposition
- **APPRART**: Fused preposition and determiner
- **KO.**: Matches any conjunction
- **TRUNC**: Truncated word (e.g. “unter-“)
- **\$\$.**: Sentence-final punctuation
- **\$,**: Sentence-internal punctuation
A.3 Annotations of the tutorial corpora

**English corpus: DICKENS**

- Positional attributes (token annotations)
  - `word` word forms ("plain text")
  - `pos` part-of-speech tags (Penn Treebank tagset)
  - `lemma` base forms (lemmata)

- Structural attributes (XML tags)
  - `novel` individual novels
  - `novel_title` title of the novel
  - `book` when text is subdivided into books
  - `book_num` number of the book
  - `chapter` chapters
  - `chapter_num` number of the chapter
  - `chapter_title` optional title of the chapter
  - `title` encloses title strings of novels, books, and chapters
  - `p` paragraphs
  - `p_len` length of the paragraph (in words)
  - `s` sentences
  - `s_len` length of the sentence (in words)
  - `np` noun phrases
  - `np_h` head lemma of the noun phrase
  - `np_len` length of the noun phrase (in words)
  - `pp` prepositional phrases
  - `pp_h` functional head of the PP (preposition)
  - `pp_len` length of the PP (in words)

**German corpus: GERMAN-LAW**

- Positional attributes (token annotations)
  - `word` word forms ("plain text")
  - `pos` part-of-speech tag (STTS tagset)
  - `lemma` base forms (lemmatised forms)
  - `alemma` ambiguous lemmatisation (feature set, see examples in Section 6.6)
  - `agr` noun agreement features (feature set, see examples in Section 6.6)

  Each agreement feature has the form `ccc: g:nn:ddd` with

  - `ccc` = case (Nom, Gen, Dat, Akk)
  - `g` = gender (M, F, N)
  - `nn` = number (Sg, Pl)
  - `ddd` = determination (Def, Ind, Nil)
XML elements representing syntactic structure

• XML elements representing syntactic structure
  
  `<s>` sentences
  `<pp>` prepositional phrases
  `<np>` noun phrases
  `<ap>` adjectival phrases
  `<advp>` adverbial phrases
  `<vc>` verbal complexes
  `<cl>` subclauses

• Key-value pairs in XML start tags
  
  `<s len=".."` >
  `<pp f=".." h=".." agr=".." len=".."` >
  `<np f=".." h=".." agr=".." len=".."` >
  `<ap f=".." h=".." agr=".." len=".."` >
  `<advp f=".." len=".."` >
  `<vc f=".." len=".."` >
  `<cl f=".." h=".." vlem=".." len=".."` >

  `len` = length of region (in tokens)
  `f` = properties (feature set, see next page)
  `h` = lexical head of phrase (`<pp_h>`: “prep: noun”)
  `agr` = nominal agreement features (feature set, partially disambiguated)
  `vlem` = lemma of main verb

• Properties of syntactic structures (f key in start tags)
  
  `<np_f>` norm (“normal” NP), ne (named entity),
  rel (relative pronoun), wh (wh-pronoun), pron (pronoun),
  refl (reflexive pronoun), es (es), sich (sich),
  nodet (no determiner), quot (in quotes), brac (in parentheses),
  numb (list item), trunc (contains truncated nouns),
  card (cardinal number), date (date string), year (specifies year),
  temp (temporal), meas (measure noun),
  street (address), tel (telephone number), news (news agency)

  `<pp_f>` same as `<np_f>` (features are projected from NP)
  + nogen (no genitive modifier)

  `<ap_f>` norm (“normal” AP), pred (predicative AP),
  invar (invariant adjective), vder (deverbal adjective),
  quot (in quotes), pp (contains PP complement),
  hypo (uncertain, AP was conjectured by chunker)

  `<advp_f>` norm, temp (temporal adverbial), loc (locative adverbial),
  dirfrom (directional source), dirto (directional path)

  `<vc_f>` norm, inf (infinitive), zu (zu-infinitive)

  `<cl_f>` rel (relative clause), subord (subordinate clause),
  fin (finite), inf (infinitive), comp (comparative clause)
A.4 Reserved words in the CQP language

Reserved words cannot be used as identifiers (i.e. corpus handles, attribute names, query names or labels) in CQP queries and interactive commands.

- new in CQP v3.4.13: Reserved words can now be quoted between backticks.
  
  ```
  > show +`no`;
  > `MU` = [lemma = "meeting|union"];
  > group `MU` match lemma;
  > ['size' = "$d\{5,\}$);
  ```

- Quotes are neither required nor allowed in label definitions and qualified label references.
  
  ```
  > left: [pos = "NN"] "after" right: [pos = "NN"] :: left.lemma = right.lemma;
  ```

- The usual rules for identifiers still apply, so e.g. `size `007`; will not be accepted.

a: asc ascending
b: by
c: cat cd collocate contains cut
d: def define delete desc descending diff difference discard dump
e: exclusive exit expand
f: farthest foreach
g: group
h: host
i: inclusive info inter intersect intersection
j: join
k: keyword
l: left leftmost
m: macro maximal match matchend matches meet MU
n: nearest no not NULL
o: off on
r: randomize reduce RE reverse right rightmost
s: save set show size sleep sort source subset
t: TAB tabulate target target[0-9] to
u: undump union unlock user
w: where with within without
y: yes
A.5 Full list of CQP options

This appendix lists all the CQP options that can be changed using the `set` command during a CQP session. There are many more configurable settings, but they cannot be set by the user during a session. Instead, they must be set when CQP is invoked (see `cqp` -h for more).

A.5.1 Boolean options

Boolean (true/false) options are set as `on` or `off`. Their present value is displayed as `yes` or `no`.

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>Option</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>as</td>
<td>AutoSave</td>
<td>Automatically save subcorpora/query results to disk</td>
</tr>
<tr>
<td>sub</td>
<td>AutoSubquery</td>
<td>Automatically enter subquery mode by activating new subcorpus/query result on creation</td>
</tr>
<tr>
<td>col</td>
<td>Colour</td>
<td>Enable colour highlighting</td>
</tr>
<tr>
<td>es</td>
<td>ExternalSort</td>
<td>Use external helper program to sort queries</td>
</tr>
<tr>
<td>h</td>
<td>Highlighting</td>
<td>Highlight hits (and target/keyword anchors) within KWIC output</td>
</tr>
<tr>
<td>o</td>
<td>Optimize</td>
<td>Enable experimental optimisations</td>
</tr>
<tr>
<td>p</td>
<td>Paging</td>
<td>Use external pager program to display KWIC</td>
</tr>
<tr>
<td>pp</td>
<td>PrettyPrint</td>
<td>Format output neatly for human readers</td>
</tr>
<tr>
<td>pb</td>
<td>ProgressBar</td>
<td>Show the progress of query execution</td>
</tr>
<tr>
<td>sa</td>
<td>SaveOnExit</td>
<td>Save all unsaved subcorpora/query results when CQP exits</td>
</tr>
<tr>
<td>sta</td>
<td>ShowTagAttributes</td>
<td>Display key-value pairs in XML tags (in KWIC)</td>
</tr>
<tr>
<td>st</td>
<td>ShowTargets</td>
<td>Print identifier numbers for <code>target(0)</code> and <code>keyword(1)</code> in KWIC; same effect as <code>show +targets</code></td>
</tr>
<tr>
<td>sr</td>
<td>StrictRegions</td>
<td>Make XML start/end tags within query match a single region</td>
</tr>
<tr>
<td>wh</td>
<td>WriteHistory</td>
<td>Write all commands entered to a history file</td>
</tr>
</tbody>
</table>

A.5.2 String options

String options contain a line of data that has some effect on or role in CQP’s operation. When setting a string option, it must be enclosed in quote marks - which will not form part of the actual option value.

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>Option</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>dd</td>
<td>DataDirectory</td>
<td>Directory to be used for saving/loading query results</td>
</tr>
<tr>
<td>col</td>
<td>DefaultNonbrackAttr</td>
<td>P-attribute used to match regular expressions which appear alone outside [...]</td>
</tr>
<tr>
<td>esc</td>
<td>ExternalSortCommand</td>
<td>Shell command to invoke external sort program</td>
</tr>
<tr>
<td>hf</td>
<td>HistoryFile</td>
<td>Location where command history will be saved</td>
</tr>
<tr>
<td>ld</td>
<td>LeftKWICDelim</td>
<td>Delimiter before the “hit” on the KWIC line</td>
</tr>
<tr>
<td>pg</td>
<td>Pager</td>
<td>Shell command to invoke external pager program</td>
</tr>
<tr>
<td>ps</td>
<td>PrintStructures</td>
<td>List of s-attributes to print within KWIC (delimited by space, comma, or full stop)</td>
</tr>
<tr>
<td>r</td>
<td>Registry</td>
<td>Delimiter before the “hit” on the KWIC line</td>
</tr>
<tr>
<td>rl</td>
<td>RightKWICDelim</td>
<td>Delimiter before the “hit” on the KWIC line</td>
</tr>
</tbody>
</table>
A.5.3 Enumerated options

Enumerated options - a sub-type of string - can only be set to one of a fixed list of values. In the case of PrintOptions, each item on the list sets one of a set of Boolean output formatting options either on or off, and some items are synonyms.

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>Option</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>ms</td>
<td>MatchingStrategy</td>
<td>Match strategy used for token-level regular expressions. One of: traditional, shortest, standard, longest.</td>
</tr>
<tr>
<td>pm</td>
<td>PrintMode</td>
<td>Print format used to display KWIC output. One of: ascii, sgml, html, latex.</td>
</tr>
<tr>
<td>po</td>
<td>PrintOptions</td>
<td>Operation to perform on the print options setup. One of: wrap, nowrap, table.tbl, notable/notbl, header/hdr, noheader/nohdr, border/bdr, noborder/nobdr, number/num, nonumber/nonum.</td>
</tr>
</tbody>
</table>

A.5.4 Context options

Context options - a sub-type of string - set the width of the left or right context in the KWIC display (or both) in units of characters, words, or s-attribute regions.

<table>
<thead>
<tr>
<th>Abbr.</th>
<th>Option</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>c</td>
<td>Context</td>
<td>Pseudo-option to set LeftContext and RightContext to the same value</td>
</tr>
<tr>
<td>lc</td>
<td>LeftContext</td>
<td>Set width of left context</td>
</tr>
<tr>
<td>rc</td>
<td>RightContext</td>
<td>Set width of right context</td>
</tr>
</tbody>
</table>