ICARUS2 Corpus Query Processor Specification

Markus Gärtner

2020

Contents

1	Notations and Definitions				
	1.1	Tree I	$\operatorname{nclusion}$	6	
2	Plain Matching				
3	Sequ	ience M	Iatching	8	
	3.1	Defini	${ m tions} \ldots \ldots \ldots \ldots \ldots \ldots \ldots$. 8	
	3.2	Rules		9	
		3.2.1	Single Node	10	
		3.2.2	Node Repetition	11	
		3.2.3	Existential Negation	11	
		3.2.4	Universal Quantification	11	
		3.2.5	Node Disjunction	11	
		3.2.6	Iterative Scan	. 11	
		3.2.7	Region Interval	12	
		3.2.8	Fixed Spot	12	
	3.3	Algori	thm	13	
4	Tree	Match	ing	14	
Αı	Appendices				

List of Figures

List of Tables

List of Algorithms

3.1	Generic node matching	10
3.2	Single node matching	11
3.3	Iterative scan matching	12
3.4	Fixed spot matching	13
3.5	Adjacent sequence matching	13
3.6	Quantified matching	13

Introduction

The ICARUS2 Corpus Query Processor (ICQP) is a custom evaluation engine for corpus queries that follow the ICARUS2 Query Language (IQL) specification.

Notations and Definitions

```
sequence xxx  \begin{aligned} &\textbf{tree } \textbf{xxx} \\ &\textbf{search node a } search \ node \ n \ \text{is a tuple } (c, \delta_p, \delta_s) \ \text{where } c \ \text{is the local constraint predicate taking} \\ &\textbf{as } \text{argument a target node } n_i \ \text{from a sequence, } \delta_p \ \text{is the minimum distance to the node's} \\ &\text{predecessor and} \end{aligned}   \begin{aligned} &\textbf{tree search node a } tree \ search \ node \ v \ \text{is a tuple } (c, l, ) \end{aligned}   \begin{aligned} &\textbf{sequence } \textbf{xxx} \end{aligned}   \begin{aligned} &\textbf{sequence } \textbf{xxx} \end{aligned}   \begin{aligned} &\textbf{sequence } \textbf{xxx} \end{aligned}   \end{aligned}   \begin{aligned} &\textbf{sequence } \textbf{xxx} \end{aligned}
```

1.1 Tree Inclusion

sequence xxx

Let T be a rooted tree. We say that T is annotated if each node v in T is assigned a set of annotations $a_1(v)...a_n(v)$ where each annotation function a_i provides its own alphabet Σ_i of available annotation values. This definition is similar to the basic notion of a labeled tree where each node is assigned a single label or value from a common alphabet Σ , but to accommodate the nature of multi-layer annotations in linguistic corpora, we extend this definition. T is further ordered if for any node v its children $v_1...v_n$ follow a globally consistent ordering scheme. If not specified otherwise, all trees in this document are rooted and annotated.

A tree P is said to be *included* in T, denoted $P \sqsubseteq T$, if deleting nodes in T can yield P. Deleting a node v in T means replacing v with the sequence of its children. Solving the tree inclusion problem means determining if P actually is included in T and then also returning all (or up to a specified number of) subtrees of T that include P.

Plain Matching

Sequence Matching

Elements participating in sequence matching:

node singular and optionally quantified element

sequence sequence of elements that adhere to given arrangement

grouping sequence of elements that is optionally quantified as a whole

disjunction two or more alternative elements

3.1 Definitions

Let L be a list, then $N_L = |L|$ is its length and $l_i \in L$ denotes the element at position i of the list where $1 \le i \le N_L$. Let T be the list of target elements and S the tree of search nodes. Utility procedures and functions used in the algorithms of this section:

- atom(s) Returns the single wrapped child node for s.
- **child**(s, i) Returns the child node s_i on index i for s.
- eval(s,t) Evaluates the inner constraints of search node s on the target item t. The result is a cache entry with a boolean value indicating whether the evaluation was successful.
- cacheGet(s, t) Retrieve a cached entry for the evaluation of search node s on target element t. If no entry exist, nil is returned.
- cacheSet(s, t, entry) Stores the given entry in the cache for the evaluation of search node s on target element t.
- **finished()** Returns whether or not the state machine has produced a sufficient number of matches for the current target sequence. This function is required for nodes that can produce multiple matches such as *repetition* and *scan*.
- mark(s,t) Stores the fact that s matched t in the preliminary result buffer. Note that each s that models an instance of iql:IqlNode is assigned a stack to manage result candidates, so in the case of quantification that allows repetition multiple hits can be stored.
- minSize(s) Returns the minimum number of elements needed to satisfy a node.

 $\mathbf{next}(s)$ Returns the next search node in the sequence after s.

size(s) Returns the number of child nodes attached to s.

scopeCreate() Registers and returns a scope marker that can later be used to reset all marked matches in the result buffer, back until this marker.

scopeReset(scope) Removes from the result buffer all matches registered via mark(s,t) that have been added since scope has been created. This is used by nodes that have to explore many alternative or iterative scenarios, such as scan, repetition or branch.

type(s) Returns the type of the given search node, one of the following:¹

single A single atomic node that is existentially quantified.

repetition A quantifier (c_{min}, c_{max}) and an associated atom search node.

negation An embedded node that must **not** match.

universal An embedded node that is expected to match all available target elements. Note that universal quantification is only allowed if the node is the only one in the (resolved) global context.

branch A group of nodes that represent logical alternatives.

scan An embedded node is iteratively checked against all remaining index positions in the target sequence.

region An embedded node and associated index interval $i_{start} = first(s)$ to $i_{end} = last(s)$ of legal positions for matching.

spot An embedded node and a single associated fixed index i = exact(s) for matching.

 $\mathbf{unmark}(s,t)$ Removes the fact that s matched t from the preliminary result buffer.

value(entry) Extracts the actual boolean result value from a cache entry.

3.2 Rules

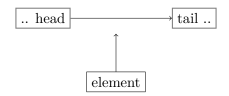
This section describes the recursive rules for constructing the object graph of search nodes from the original elements in the query. All rules assume the existence of already processed nodes that resulted in some *head* part to the left that is connected to a *tail* to the left, as illustrated below:



Initially the *head* node is just a generic entry point and the *tail* simply implements the final acceptance node that ensures that all actual search nodes in the query have been evaluated already.

The elements in the query are effectively processed top-down and left-to-right and for every encountered <IqlElement> instance a new node is inserted between the current *head* and *tail*:

¹Nested nodes can be obtained with atom(s) for single embeddings or child(s,i) for indexed elements where size(s) returns the number of embedded nodes and $1 \le i \le atom(s)$.



This process is recursive and builds the graph of interconnected search nodes used in the algorithms described in the respective sub sections.

Algorithm 3.1 Match a generic node at a specified position. This procedure exists merely for multiplexing to the specialized counterparts depending on the type of s.

```
1: procedure MATCH(s, T, j)
2:
       type_s \leftarrow \text{TYPE}(s)
       if type_s = single then
3:
          return MATCH-NODE(s, T, j)
4:
       else if type_s = repetition then
5:
          return MATCH-REPETITION(s, T, j)
6:
7:
       else if type_s = negation then
          return MATCH-NEGATION(s, T, j)
8:
       else if type_s = universal then
9:
          return MATCH-ALL(s, T, j)
10:
       else if type_s = branch then
11:
12:
          return MATCH-BRANCH(s, T, j)
13:
       else if type_s = scan then
          return MATCH-SCAN(s, T, j)
14:
       else if type_s = region then
15:
          return MATCH-REGION(s, T, j)
16:
17:
       else if type_s = spot then
          return MATCH-SPOT(s, T, j)
18:
       end if
19:
20: end procedure
```

Actual sequence matching is subsequently performed by calling MATCH(root, T, 1) with root being the first node of the state machine to. This function takes care of multiplexing the call to the specialized procedure depending on a node's type (as provided by type(s) in line 2 of Algorithm 3.1).

3.2.1 Single Node

A single instance of iql:IqlNode (IQL, Section 1.5.3) is the most basic constraint fragment in an IQL query. Note that in the presence of quantifiers (IQL, Section 1.14) on a node the evaluation is split into two levels: First the iql:IqlNode content is processed into a search node s of type single and then the quantifiers are handled (which will result in a combination of repetition, branch, negation, universal nodes, depending on the complexity of the quantifiers involved), resulting in a potentially very complex node structure with node s as atom.

Algorithm 3.2 Matching of a single node at a specific position. Local constraints and the tail of s are taken into account. Memoization is employed for evaluation of local constraints to prevent repeatedly executing costly constraint expressions.

```
1: procedure MATCH-NODE(s, T, j)
 2:
        entry \leftarrow \text{CACHEGET}(s, t_i)
        if entry = nil then
 3:
            entry \leftarrow \text{EVAL}(s, t_i)
 4:
            CACHESET(s, t_j, entry)
 5:
        end if
 6:
        matched \leftarrow VALUE(entry)
 7:
        if matched = true then
 8:
            tail \leftarrow next(s)
 9:
            matched \leftarrow MATCH(tail, T, j + 1)
10:
11:
        end if
        if matched = true then
12:
            MARK(s, t_i)
13:
        end if
14:
15:
        return matched
16: end procedure
```

Matching a single node is pretty straightforward, as shown in Algorithm 3.2. The only local evaluation concerns the execution of the internal constrain expression, for which memoization is employed in order to prevent repeatedly executing it for the same target element. Only if the local constraint evaluates to **true** further evaluation is delegated to the tail of the state machine via matching the next(s) node in line 10. A successful match is automatically stored in the associated result buffer. Note that surrounding scan or repetition nodes that can produce multiple matches have the ability to reset the state of this result buffer. So if a match fails, the individual nodes won't have to do any cleanup work.

3.2.2 Node Repetition

3.2.3 Existential Negation

3.2.4 Universal Quantification

This special node can only occur as either the root node or as direct element within a disjunction (Section 3.2.5) that in turn is the root. It effectively makes the embedded atom the only node that is allowed to match and it is required to all targets in the current unit-of-interest (UoI). The implementation is very similar to that of scanning (Section 3.2.6) in that it moves the atom through the target sequence and aborting as soon as it fails to match a single element.

3.2.5 Node Disjunction

3.2.6 Iterative Scan

Normally matching a node is done on a single specific position in the target sequence. This covers situations such as node groups with the ADJACENT arrangement. But groups that

are merely $\mathsf{ORDERED}$ (also the default for sequences in IQL when no explicit arrangement is defined) need some sort of scanning mechanism, as individual nodes can be matched at any index position after their predecessor (if one exists). Therefore the scan node takes the index argument j of the matcher function as entry point for an iterative search forward that incrementally tries positions until running out of search space or if the result buffer is filled.

Algorithm 3.3 Match an embedded search node at any of the remaining spots in the target sequence, beginning at j.

```
1: procedure MATCH-SCAN(s, T, j)
        atom \leftarrow ATOM(s)
 2:
 3:
        fence \leftarrow N_T - \text{MINSIZE}(s) + 1
 4:
        result \leftarrow false
 5:
        while i < fence \land \neg FINISHED() do
                                                                     > result limit can end the loop early
 6:
            matched \leftarrow MATCH(atom, T, i)
 7:
 8:
            if matched then
                result \leftarrow true
 9:
            end if
10:
            i \leftarrow i + 1
11:
        end while
12:
13:
        return result
14: end procedure
```

Algorithm 3.3 displays the simple algorithm for iterative scanning. Note that this version is not using caching to speed up exploration in the nested *atom* and also only implements the default left-to-right traversal direction. A specialized alternative exists that shifts *atom* through the search space right-to-left and another one that uses caching to skip known deadends when matching the nested *atom* (line 7). It is not possible to skip known positive subresults, as we must still allow nested nodes to register with the result buffer for any such "new" sub-result in the context of a new host node s. When traversing backwards, the algorithm uses the same boundaries, but simply reverses the direction of the loop variable.

3.2.7 Region Interval

All position markers (IQL, ??) that effectively declare intervals are translated into region nodes. Each region node manages one or more index intervals of legal values. Those intervals are refreshed for every UoI at most once. When matching, membership of the current index j to at least one interval is checked first. If this check fails, the entire match call is aborted before the next node is even considered.

3.2.8 Fixed Spot

Node implementation of the markers IsFirst, IsLast and IsAt that all define a fixed index for the target, only influenced by the size of the UoI during matching.

Algorithm 3.4 Match search node only at a fixed location in the target sequence

```
1: procedure MATCH-SPOT(s,T,j)

2: spot \leftarrow \text{EXACT}(s)

3: if j \neq spot then

4: return false

5: end if

6: node \leftarrow \text{ATOM}(s)

7: return MATCH(node,T,j)

8: end procedure
```

3.3 Algorithm

Algorithm 3.5 Adjacent sequence matching

```
1: procedure MATCH-ADJACENT(C, T, j_{start}, j_{max})
                                                                                        \triangleright j_{start} \leq j_{max}, C \subseteq S
 2:
        j \leftarrow j_{start}
        while j <= j_{max} do
3:
            if LOCAL_MATCH(s, t_i) then
 4:
 5:
                 return j
            end if
 6:
            j \leftarrow j + 1
 7:
        end while
 8:
        return -1
10: end procedure
```

Algorithm 3.6 Quantified matching

```
1: procedure MATCH-QUANTIFIED(s, T, j_{start}, j_{max})
                                                                                               \triangleright j_{start} \leq j_{max}
 2:
        j \leftarrow j_{start}
        while j <= j_{max} do
3:
            if LOCAL_MATCH(s, t_j) then
 4:
                return j
 5:
            end if
 6:
            j \leftarrow j + 1
 7:
        end while
 8:
        return -1
9:
10: end procedure
```

Tree Matching

Appendices