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Structural Analysis and Visualization of C++ Code Evolution using Syntax Trees

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ABSTRACT

We present a method to detect and visualize evolution patterns in C++ source code. Our method consists of three steps. First, we extract an annotated syntax tree (AST) from each version of a given C++ source code. Next, we hash the extracted syntax nodes based on a metric combining structure and type information, and construct matches (correspondences) between similar-hash subtrees. Our technique detects code fragments which have not changed, or changed little, during the software evolution. By parameterizing the similarity metric, we can flexibly decide what is considered to be identical or not during the software evolution. Finally, we visualize the evolution of the code structure by emphasizing both changing and constant code patterns. We demonstrate our technique on a versioned code base containing a variety of changes ranging from simple to complex.

Categories and Subject Descriptors

I.3.8 [Computer Graphics]: Applications; D.2.8 [Distribution, Maintenance and Enhancement]: Restructuring, reverse engineering, and reengineering

Keywords

code structure evolution, syntax trees, software visualization

1. INTRODUCTION

Software configuration management (SCM) systems, such as CVS, Subversion, ClearCase and Perforce, are an important part of the management of large-scale software development projects. SCM systems support tasks such as checking in and out different file versions in a software project, managing parallel development branches, release scheduling, bug management, and collaborative teamwork. At the lowest level, a SCM system supports these functions by maintaining a history of changes of the software artifacts. Typically, these artifacts are source code (text) or binary files, and the changes are recorded and managed by the SCM system at line or character (byte) level.

In the recent years, several tools and techniques emerged to augment the above basic functionality with more complex features. These tools can be loosely classified into data mining tools, which extract various facts from code repositories, and visualization tools, used to present and navigate both raw data and extracted facts. An overview of such tools is presented in [17]. Such tools support a wide range of tasks, such as identifying the developer network, finding stable software releases, monitoring software quality and debugging activity, and analyzing the software architecture evolution. The primary users of such tools are software architects, who must manage large systems at medium or high abstraction levels, and are not interested in all the minute code details.

In this paper, we focus on a subset of these activities. We are interested to understand how the detailed structure of source code changes during the evolution of a code project consisting of hundreds of source code files, changed by tens of developers over several years. Developers are interested to see which parts of the code changed, and how. At a fine level-of-detail, this can be done by visualizing the evolution of each code line in one, or a few, files of interest across all versions. Tools such as WinDiff and CVSscan support this task, using the diff operator of the underlying SCM system to tell which lines have been changed, added, or deleted [18]. However useful to see small-scale, minute code changes, this approach has several limitations. First, one cannot detect and show more complex code evolution patterns located at coarser levels of details, such as the moving of a whole code block from one function or class to another. Second, the line-based code evolution analysis is sensitive to low-level changes such as reformatting, identifier renaming, expression rewriting, or declaration order change. Such changes are irrelevant to activities such as code refactoring, rearchitecting, and generally when one works with software source code at higher levels.

In this paper, we present a set of techniques to detect and visualize structural code evolution patterns for C and C++ code repositories. We use a C++ parser [13] to extract a full abstract syntax tree (AST) from every version of every file of interest in the repository. Next, we augment an existing technique for detecting structurally similar subtrees in a larger tree to detect structurally similar code fragments in ASTs corresponding to consecutive versions. At the core of our technique, we have a user-specified similarity metric which combines semantic and structural information in order to tell what is constant and what has changed. Finally, we propose a visualization method to present the code structure evolution and emphasize both constant and changing patterns.

This paper is structured as follows. In Section 2, we review related work on detecting similar (evolving) code patterns. Section 3 details our similar code pattern detection method. Section 4 presents our visualization method for code structure evolution. Sec-
tion 5 demonstrates our method on a C++ code base. Section 6 discusses our method. Section 7 concludes the paper and outlines future work directions.

2. PREVIOUS WORK

Showing constant and changing patterns in evolving code inherently relates to clone detection and clone tracking methods. Given two files $f_A$ and $f_B$, which can be totally unrelated or two versions of the same file $f$, clone detection methods find a set of code patterns $c_{ij}^A \subseteq f_A$ which closely resemble a set of patterns $c_{ij}^B \subseteq f_B$. Several such methods exist. Baxter et al. extract abstract syntax trees from the code, determine a hash code from the entire tree structure, and compare same-hashcode trees using a bottom-up matching algorithm [6]. Jiang et al. compute fixed-length vector descriptors of syntax tree nodes, recording the number of occurrences of each node type, and hash similar subtrees based on the Euclidean distance between vectors [11]. Koschke et al. use a suffix token tree approach, comparing syntax trees by serializing the tree node types to strings, thereby combining the speed of string approaches with the precision of tree-based approaches [12]. Wåhler et al. use an XML-based syntax trees and database queries to find code clones as frequent item-sets [19]. Ducasse et al. advocate a string-based clone detection, thereby removing the need for heavyweight parsers [9]. Recently, Etoko and Robillard proposed a method to track code clones across several versions of a code base, by reusing the SimScan clone detector atop of a lightweight clone representation combining structural and lexical clone information [8]. Clone detection methods are also implemented in widely-used clone detection software, such as the well-known CCfinder tool.

On a different track, several methods have been proposed to visualize code evolution patterns, at code line level [18, 13], intermediate (group-of-lines) level [1], and file and component level [14, 16].

3. TRACKING SIMILAR CODE

We are interested in detecting code evolution patterns at several levels of detail, ranging from an identifier and expression up to a whole function, class, or namespace. In the following, we consider $N$ versions of a given source code file $f_1, ..., f_N$. We proceed as follows.

For each version $f_i$, we extract its complete annotated syntax tree (AST) $T_i$ using a $\texttt{gcc}$-based C++ parser, following the approach described in [13]. Next, we compare all successive trees $T_i$ with $T_{i+1}$ and identify similar subtrees. For this, we extend the method of Auber et al. to detect structurally similar subtrees [4, 7], by adding semantic (type and code text) information extracted by our parser. This allows us to flexibly specify what we consider ‘similar’ and/or ‘different’ during the code evolution.

Our method consists of two steps. In the first step, we group all extracted AST nodes into equivalence classes, based on structural and semantic information (Sec. 3.1, 3.2). Same-class nodes are likely to be similar, whereas nodes in different classes are not. In the second step, we construct explicit correspondences between similar subtrees rooted in the same class (Sec. 3.4). Finally, we visualize these correspondences (Sec. 4).

3.1 Structural node classification

In the first step, we group AST nodes from version-consecutive syntax trees $T_i$ and $T_{i+1}$ into equivalence classes. Same-class nodes are roots of potentially similar subtrees. In the following, we shall denote the subtree rooted at node $u$ by $R(u)$. We compute equivalence classes using a distance metric $d(u, v)$ which combines structural and semantic information from $R(u)$ and $R(v)$, as follows.

Following [7], we define the structural distance component $d_{str}(u, v)$ as the difference between three topological invariant, graph-theoretic, metrics on $R(u)$ and $R(v)$: The number of direct children, or degree $\delta(u)$, of $u$; the size $\nu(u)$ of $R(u)$; and the Strahler number $\sigma(u)$ of $R(u)$. The first two metrics are well-known. The Strahler number is defined as follows

$$
\sigma(u) = \begin{cases} 
1, & u \text{ is a leaf} \\
\max_{1 \leq k \leq \nu(u) + 1} \sigma(u_i) + k, & u \text{ has } k \text{ children } u_i
\end{cases}
$$

(1)

The Strahler number is used in graph theory to succinctly characterize the topological complexity (e.g. ramification) of a tree or a DAG. For further details, we refer the reader to the literature [15, 3].

With the above metrics $\delta, \nu$ and $\sigma$, we define the structural distance $d_{str}(u, v)$ as

$$
d(u, v) = \left(\hat{\delta}(u) - \hat{\delta}(v)\right)^2 + \left(\hat{\nu}(u) - \hat{\nu}(v)\right)^2 + \left(\hat{\sigma}(u) - \hat{\sigma}(v)\right)^2
$$

(2)

where $\hat{\delta}$ denotes the normalized version of the metric $\delta$, i.e. $\hat{\delta}(u) = \delta(u) - \delta_{min}$, where $\delta_{min}$ and $\delta_{max}$ are the minimal, respectively maximal values of $\delta$ over all nodes in $T_i$. $T_{i+1}$, and similarly for the metrics $\nu$ and $\sigma$.

3.2 Using type information

The distance $d_{str}(u, v)$ effectively ‘hashes’ structurally similar subtrees [7, 4]. However, when comparing software code, we want to consider the code semantics too. Each AST node $n$ has a type $t(n)$, e.g. declaration, function, assignment, for statement, identifier, and so on. The C++ grammar our parser uses has approximately 150 such types. We use these types to enhance the structural similarity, as follows.

First, it is not meaningful to compare totally unrelated types, e.g. an arithmetic expression $a+b$ with, say, a declaration $\texttt{int } x$, even though such constructs may have structurally similar ASTs. We model this by a second distance $d_{typ}(u, v)$ which tells the difference between two node types $t(u)$ and $t(v)$. The simplest way is to test strict type equality, i.e. use $d_{typ}(u, v) = (t(u) == t(v))$. However, this distance is too strict. Often, one wishes to track ‘fuzzier’ patterns over the code evolution, i.e. patterns which are not exactly the same, but related, syntax types. To do this, we define several classes of type-equivalence $C_i$ for C++ syntax node types, based on what one considers to be a change during evolution. A class $C_i$ contains all types considered similar. Hence, the distance $d_{typ}$ is:

$$
d_{typ}(u, v) = \begin{cases} 
0, & t(u) \text{ and } t(v) \text{ are in the same class } C_i \\
1, & \text{otherwise}
\end{cases}
$$

(3)

Examples of useful type-equivalence classes are: for, while, and do statements (if one is interested to track iterative constructs); and classes, structures, and unions (if one is interested to track aggregate types). Any classes can be defined, depending on what one considers to be the ‘same’ during the code evolution. In the above, a C++ class declaration that changes into a struct, but keeps the same internal structure, is considered to be the same code. We do not detail other classes here, as their exact definition depends on the task at hand, but also on the specific C++ grammar used.

1The ASTs of these constructs are both a tree with two leafs, i.e. the identifiers $a$ and $b$ for the expression $a+b$, and the type $\texttt{int}$ and the declarator $x$ for the declaration $\texttt{int } x$.
The structural distance $d_{str}$ and type distance $d_{typ}$ yield our final distance $d(u, v)$:

$$d(u, v) = [1 + (w(u) + w(v))d_{typ}(u, v)]d_{str}(u, v) \quad (4)$$

In Eqn 4, $w(u)$ is a real-valued weight, or importance, of the type $t(u)$ of a node $u$. If the types of the compared nodes $u$ and $v$ are in the same type-class, then $d(u, v)$ equals the structural distance $d_{str}(u, v)$. If not, i.e. $d_{str}(u, v) = 1$ then $d(u, v)$ is larger if the types $t(u)$ and $t(v)$ are considered to be more important. The importance weight is useful to emphasize certain types of changes during the code evolution. For example, if we are interested in interface evolution, then everything having to do with global declaration node types (e.g. functions, classes, etc) will be important, whereas implementation-related node types (e.g. expressions, iterative statements, assignments, etc) are irrelevant. We can achieve this by setting a high weight $w$ for nodes of the former types and a low weight $w$ for nodes of the latter types. For a neutral standpoint, all weights $w$ for all AST node types are set to 1.

Note that the structural distance $d_{str}$ and the type importance $w$ could have been merged in a single function. However, this means defining a $N$ by $N$ matrix of real values, where $N$ is the number of C++ type nodes. For $N = 150$, this is impractical. By decoupling the two concerns as discussed above, we use just two $N$-valued vectors: a set of disjoint type equivalence classes $C$, and a real-valued vector of type weights $w$.

### 3.3 Labeling nodes

We use the distance $d(u, v)$ from Eqn 4 to distribute the nodes in $T_i$ and $T_{i+1}$ in equivalence classes, as stated at the begin of Sec 3.1. Each node $n$ is labeled by its integer class-id $\lambda(n)$, following the algorithm in Fig 1. Essentially, all nodes which are closer than a user threshold $\varepsilon_d$ are put in the same class $\lambda$. The concrete values for $\varepsilon_d$ depend, of course, on the range of the distance metric $d$ (Eqn 4), which further depends on the type weights $w$. If we normalize $d$ between 0 and 1, we found that good values for $\varepsilon_d$ are in the range of 0.005 to 0.01, which put only strongly similar nodes in the same class.

Clearly, the results may depend on the order the nodes are considered in, since not all distance pairs are computed. However, as discussed in [7], the structure-and-type distance function $d$ (Eqn 4) is strongly discriminative, so in practice there are just few cases when many nodes have similar distances to each other. A heuristic that further improves the result is to consider nodes sorted in the decreasing order of their Strahler metric $\sigma$ (Eqn 1) (line 3 in Fig 1). This ensures that larger, more complex and thus more interesting, code fragments are classified first, thereby limiting the impact of the order nodes are treated in to the less interesting code fragments.

```plaintext
Listing 1: Node labeling algorithm

```

### 3.4 Correspondence construction

The first phase of our algorithm discussed so far distributes AST nodes from each two consecutive versions $i$ and $i + 1$ into equivalence classes (Sec. 3.1). The first phase can be seen as a hashing which groups similar subtrees into the same equivalence class. Yet, after the first phase, we do not know for sure which of the subtrees in the same class are truly similar. The second phase performs an in-depth analysis which finds and marks those parts of all subtrees in an equivalence class which are indeed similar.

The correspondence construction works recursively top-down. For two nodes $u$ and $v$ in the same equivalence class, i.e. $\lambda(u) = \lambda(v)$, denote $F(u)$ and $F(v)$ the sets of direct children of $u$ and $v$ in $T_i$ and $T_{i+1}$ respectively. For a given equivalence class $l \in [1, L]$, denote $F_i(u)$ all children of $u$ being in class $l$, i.e. having $\lambda(F_i(u)) = l$, and similarly $F_i(v)$ for $v$’s children. We say that $u$ and $v$ are $D$-similar if

$$D(u, v) = \sum_{l=1}^{L} \text{abs} \left( |F_i(u)| - |F_i(v)| \right) \leq \varepsilon_D \quad (5)$$

Here, $|F|$ denotes the size of set $F$ and ‘abs’ the absolute value. $D(u, v)$ can be seen as a similarity metric based on the direct children of two nodes in the same equivalence class. If we normalize $D$ by the total number of children of a node, good values for $\varepsilon_D$ are around 0.1.

The final step is to extend the similarity metric $D$, defined on the children of $u$ and $v$, to a metric $\Delta$ on the full subtrees $R(u)$ and $R(v)$, as follows. If $u$ and $v$ are not $D$-similar, then we say $R(u)$ and $R(v)$ are also not $\Delta$-similar. If $u$ and $v$ are $D$-similar, then we recursively compare the subtrees rooted by their children within the same equivalence class $F_i(u)$ and $F_i(v)$, for all classes $l \in [1, L]$. Denote the number of such subtrees which are found $\Delta$-similar by $\kappa(u) = \kappa(v)$. We say $u$ and $v$ are $\Delta$-similar if

$$\Delta(u, v) = \frac{\kappa(u) + \kappa(v)}{|R(u)| + |R(v)|} \leq \varepsilon_\Delta \quad (6)$$

i.e. if we found within the trees the $u$ and $v$ a relative fraction of at least $\varepsilon_\Delta$ $\Delta$-similar subtrees. The recursion ends when we compare two leaves. These are always $\Delta$-similar, and always have $\kappa = 1$. In our experiments, we used $\varepsilon_\Delta = 0.8$. Higher values imply a stricter matching, the maximum being 1, when a strict identity is required. Lower values imply that less similar subtrees are considered to match.

The complete correspondence computation algorithm for two trees $T_i$ and $T_{i+1}$ is shown in listings 2 and 3. For all equivalence classes $l$, we check if the node pairs $(s, s')$ in the class $l$ are $\Delta$-similar (Fig 2). We only check trees rooted in the same class, since only these have a fair chance to be structurally similar (Sec. 3.1). The function $\text{checkDelta}$ checks the $\Delta$-similarity. If two $\Delta$-similar subtrees are found, then all their nodes are marked as being in the same pattern. We store the values of $\kappa(u)$, which store for each node $u$ the number of $\Delta$-similar subtrees rooted at $u$’s descendants, in an auxiliary field. $\kappa(u)$ is set to zero before each tree-pair comparison (line 10, listing 2) and updated in bottom-up order during the correspondence construction, following Eqn 6 (line 27, listing 3).

A simple heuristic that improves the chance of finding good correspondences is as follows. When computing the $\Delta$-similarity, we first sort the children $F_i$ and $F'_i$ of the current nodes $u$ and $u'$ under scrutiny before matching their subtrees (lines 13-14, listing 3). We sort nodes in order of their C++ node types, and in order of their code text for identical types. This ensures that nodes as similar as possible are matched first.
4. VISUALIZATION

After computing the correspondences between code fragments, represented as AST sub-trees, of a number of versions, we visualize these. The goal is to visually emphasize important evolution events, such as code refactoring, code drift, or interface and implementation changes.

4.1 Basic design

Our visualization uses the design sketched in Fig. 1. Each syntax tree $T_i$ for each version $i$ in the evolution of a considered code file is drawn using a space-filling technique similar with so-called icicle plot [5]. A tree appears as a vertical strip, with the root to the left and the leaves to the right. Figure 2 shows the principle for a syntax tree for a small function. The syntax tree is shown using a classical tree layout (a) and using our space-filling visualization (b). ASTs of typical C++ source files of thousands of lines are very broad (tens of thousands of nodes) but not too deep (10..40 levels). Hence, trees appear as thin, tall vertical strips. The varying syntax tree depth (not to be mismatched for the code indentation depth) is shown by the strip’s width (see Figs. 3 and 4 for actual snapshots). Wider strip portions indicate deeper, thus typically more complex, syntactic constructs, while thin strip portions show constructs which are shallow nested, i.e. close to the C++ global file scope. Additionally, we use shaded cushions as an effective means of emphasizing the structure, similar to other software visualization applications [18, 10]. The cushions, implemented as one-dimensional textures storing a parabolic luminance profile, are blended atop of the rectangular nodes, an operation which is very efficiently done in OpenGL. The entire visualization is implemented using the Tulip graph visualization system which offers sophisticated mechanisms for customized layout, rendering, interactive navigation, and level-

Listing 2: Correspondence construction algorithm

The result of the entire algorithm is a labeling $\alpha$ of all nodes in the input trees $T_i$ and $T_{i+1}$. A value $\alpha(n) = 0$ means that node $n$ is not in a pattern which has a correspondence in the other tree. A value $\alpha(n) > 0$ means that node $n$ is in the pattern with id $\alpha(n)$. Finally, by applying the algorithm on all tree pairs $(T_i, T_{i+1})$, we obtain the evolution patterns of all code fragments on the entire version set considered.

Listing 3: $\Delta$-similarity computation algorithm

```plaintext
int alpha := 0;
for (n in T_i U T_{i+1}) alpha(n) := 0;
for (l := 1; l < L; l++)
{
    S_l := \{ n in T_i | \lambda(n) = l \}
    S'_l := \{ n in T_{i+1} | \lambda(n) = l \}
    for ((s, s') in S_l x S'_l)
    {
        for (n in R(s) U R(s')) alpha(n) := 0;
        if (deltaSim(s, s', \epsilon_D, \epsilon_\Delta))
        {
            alpha := alpha + 1;
            for (n in R(s)) alpha(n) := alpha
            for (n in R(s')) alpha(n) := alpha
            S_l := S_l \ { n in S_l | alpha(n) = alpha }
            S'_l := S'_l \ { n in S'_l | alpha(n) = alpha }
        }
    }
}
```
of-detail techniques for large trees and graphs up to hundreds of thousands of nodes [2].

After drawing all trees for all considered file versions, we draw the correspondences. As explained in Sec. 3, given $N$ trees $T_1, ..., T_N$ for the file versions $f_1, ..., f_N$, we compute only correspondences between subtrees in consecutive versions $T_k$ and $T_{k+1}$. For each correspondence, we connect the rectangles of the corresponding (matched) nodes from the neighboring tree strips using tube-like shapes, as shown in Fig. 1. The tubes are easy to interpret. Straight, horizontal tubes show code regions which stay constant during the evolution. Inclined tubes indicate code fragments which change places in the syntax tree. Crossing tubes indicate code block swapping. Since the top-to-bottom order of nodes in the tree view matches the order of code fragments in a program listing, long tubes indicate fragments which moved considerably during the evolution. Inserted and deleted fragments appear as regions in the tree which are not connected with tubes. Finally, the tubes’ thickness indicate the amount of code in the corresponding pattern.

Tubes can be drawn in two different ways. In the first way, shown for nodes $F, ..., I$ in Fig. 1, tubes are drawn from the right edge of the node in version $i$ to the left edge of the parent of its corresponding node in version $i+1$. In the second way, shown for node $E$, the tube is drawn to cover the two matched nodes and all their children ($K, L$), using a constant transparency value. We found the second method better as it clearly indicates node grouping in patterns. The second way for drawing the tubes has a more subtle advantage. If we look at the two nodes marked $H$ in Fig. 1, the tube connecting them seems to end too low at its right end. Actually, this is a rather surprising optical illusion due to the shaded cushions. The second way for drawing the tubes, using superimposed cushions, corrects this problem.

Figure 3 shows the visualization design on two versions of a file of approximately 850 C++ lines. In the left image, no cushions are used. The nodes are colored with hues that indicate their C++ construct types, using different user-selected hues for classes, functions, methods, expressions, declarations, iterative statements, and conditional statements. The tubes are colored using a slightly darker hue of the correspondent nodes they connect. The right image shows the same dataset. This time, both nodes and tubes are colored using a different hue for each identified pattern (Sec. 3.4). Tubes are drawn using the transparency pattern explained earlier in this section. Deleted or inserted, thus unpaired, code fragments are colored in gray.

Figure 3 b clearly shows several evolution events: Fragment $C$ is deleted, except a tiny code piece indicated by the red filament in the middle. Fragments $A$ and $B$ are swapped in an unchanged state. Fragments $D$ and $E$ stays constant.

We see here a limitation of our method. Although fragments $D$ and $E$ stay constant, they are identified (and drawn) as a multitude of blocks instead of a single block, as in the case of $A$ or $B$. The reason is that $A$ and $B$ are single subtrees in the syntax tree, whereas $D$ and $E$ consist of sets of subtrees which do not share a constant parent. Actually, $A$ and $B$ are two global C++ scope methods, while $D$ and $E$ are sets of sibling nodes representing several same-scope declarations. Since our method only matches single subtree roots, some amount of fragmentation occurs. Note that, if desired, this can be easily removed by applying a postprocessing pass that groups sets of contiguous matched sibling nodes into the same pattern. We preferred to show the individual node correspondences, as these represent actual syntactic code fragments.

### 4.2 Visual enhancements

Several enhancements can be added atop of the above basic visualization. First, the user can choose the level of detail at which the syntax trees are shown. Given a level of detail, only syntax nodes up to that level, and associated correspondences, are drawn. This lets one simplify the visualization by removing fine code details which appear as those vertical strips at the right of the version visualizations (e.g. nodes $F, G, ..., L$ in version 2 in Fig. 1). A similar effect can be achieved by adapting the structural distance function $d_{str}$ (Sec. 3.1). The difference is that changing the distance function potentially determines finding different evolution patterns, whereas culling finer detail from the visualization simply shows less correspondences from an already computed set.

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5. EXAMPLE APPLICATION

We applied our structural code evolution visualization on several versioned C++ code bases. Due to space limitations, we present here just one example. The considered versioned code base is a complete C++ image processing program of approximately 5000 lines of code. From discussions with the developers, we know the versions correspond to successive small-scale system refactorings. After parsing each version, we keep in its AST all nodes originating from application code sources and headers, but eliminate nodes coming from system headers, as we are not interested to see changes at that level (if any). Figure 4 a shows a visualization of six versions of one file containing about 1500 lines from our C++ code base. Node colors indicate syntax construct types. Edges have the colors of the matched nodes they connect.

We quickly recognize a few zones in the code where important changes occurred (marked b-f in Fig. 4). For the rest of the code, the skewed parallel tubes running left-to-right in the overview image (Fig. 4 a) indicate constant or near-constant code regions. Over-
The closest method to ours seems to be that of Baxter et al., who also extract and hash syntax nodes prior to tree comparison in their clone detection [6]. While they use only the node types in their hash function, we use a combination of structural parameters (node degree, tree size, and Strahler number) and node types (Sec. 3.1, 3.2). Our hash function seems to be weaker than the one in [6]. We actually cannot tell this for sure as [6] does not include full implementation details. This allows our method match subtrees more freely, a step which Baxter et al. solve by adding a separate "clone sequence" processing phase. Jiang et al. compute one fixed-length vector of type occurrences per subtree and use locally sensitive hashing (LSH) to group similar vectors, which is conceptually similar but differently implemented from our structure+type hashing. Their Euclidean distance used to compare vectors is, again, conceptually similar to our distance function $d_{st}$ and importance weights (Sec. 3.2), but has a quite different implementation.

Let us stress again that our aim is to identify small up to medium-scale evolution patterns in consecutive file versions, which is a very specific case of finding similar code fragments in files of moderate size (up to thousands of lines/file) exhibiting a moderate amount of change. While our method could be in principle used to detect and visualize code clones, this was not our purpose. Code cloning detectors stress scalability, performance, detection of certain pattern types, and do not typically focus on visualizing the detected correspondences. Whether our method can be further optimized and/or parameterized in the context of clone detection is a subject of future research.

Considering performance: The matching of six versions of a code base of approximately 5000 code lines per version took about 12 minutes on a 2 GHz PC running Linux. We are aware that our current implementation could be further optimized to yield the higher speed needed to analyze large code bases. However, since our target scenario is to interactively visualize small-scale, minute code modifications, typically at the level of a single file across several versions, we are less constrained by high performance requirements as e.g. code clone detection methods that run in automatic mode on full industry-size repositories.

7. CONCLUSIONS

We have presented a method to compute and visualize evolution patterns in C++ source code. At the center of our method is a structure-and-type matching technique running on the abstract syntax trees of consecutive versions. The matched subtrees are visualized using dense pixel layout and rendering methods. Our method is useful to detect and browse small to medium-scale changes during source code evolution, such as function and class-level refactorings. To visualize small-scale, minute code modifications, typically at the level of a single file across several versions, we are less constrained by high performance requirements as e.g. code clone detection methods that run in automatic mode on full industry-size repositories.

8. REFERENCES

Figure 4: Visualization of six versions of a C++ software system. a) overview; b-f) details. See the individual captions and text for more details.