The problem

Evaluating Software Clustering What is a good software decomposition?

- How do we know that a particular decomposition of a software system is good?
- What does "good" mean anyway?
- · We can compare against a
 - Mental model
 - Benchmark standard
- · Can be done either manually or automatically

Manual evaluation

- Have experts evaluate automatic decompositions
- Very time-consuming, impractical
- Also quite subjective
- Need an automatic, objective way of doing it

Automatic evaluation

- Usually measures the similarity of an automatic decomposition A to an authoritative decomposition B (prepared manually)
- Major drawback: Assumes there exists one "correct" decomposition
- Other evaluation approaches are possible, such as measuring the stability or the extremity distribution of a clustering algorithm

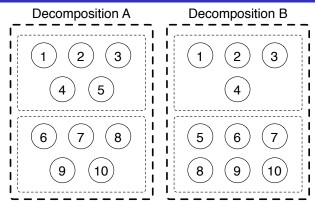
Precision/Recall

- Standard Information Retrieval measures
- Were applied in a software clustering context by Anquetil
- Definitions:
 - Intra pair: A pair of software entities in the same cluster
 - Inter pair: A pair of entities in different clusters

Precision/Recall

- Precision: Percentage of intra pairs in A which are also intra in B
- Recall: Percentage of intra pairs in B also found in A
- A good algorithm should exhibit high values in both measures

Precision / Recall example



Pair 1-5: Intra pair in A but not in B Precision: 16/20 = 80% Recall: 16/21 = 76.2%

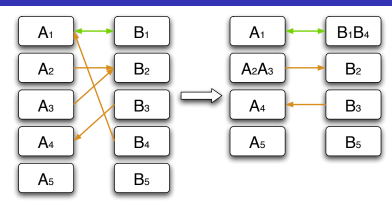
Problems with Precision / Recall

- Sensitive to the size and number of clusters
 - Differences are exaggerated if you have small/many clusters
- Two values makes comparison harder
- The two values are interchangeable if there is no authoritative decomposition

Koschke - Eisenbarth measure (KE)

- · Loosely based on Precision/Recall
- Attempts to be less strict
- Definitions:
 - GOOD match: Two clusters (one in A, one in B) with both precision and recall larger than a threshold p (typical value 70%)
 - OK match: Two clusters with only one of the measures larger than p

Koschke - Eisenbarth metric



$$\frac{\sum_{(a,b)\in\mathsf{GOOD}}\mathsf{overlap}(a,b) + \sum_{(a,b)\in\mathsf{OK}}\mathsf{overlap}(a,b)}{|\mathsf{GOOD}| + |\mathsf{OK}| + |\mathsf{true}| \mathsf{negatives}|}$$

Koschke - Eisenbarth metric

- Does not take edges into account
- In extreme situations (each cluster contains only one element) may provide strange results (similarity of 100%)
- No penalty for joining clusters

Assignment tool: ke

- Takes two .kos files containing different decompositions of the same set of entities
- Example:

ke -cand dec1.kos -ref dec2.kos

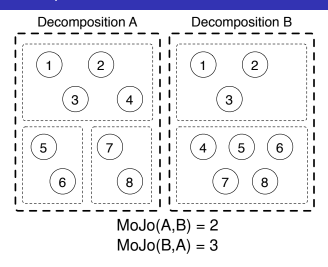
- Produces lots of output. We're interested in the recall rate
- Transform an RSF file with contain facts to the KE format with

unitrans input.rsf output.kos

MoJo distance

- The distance between two different partitions of the same software system is defined as the minimum number of Move and Join operations to transform one to the other
 - Move: Remove an object from a cluster and put it in a different cluster
 - Join: Merge two clusters into one
 - Split: Has to be simulated by Move operations

MoJo example



Why only Move and Join?

- Two clusters can be joined in only one way. One cluster can be split in two in an exponential number of ways
- Joining two clusters only means that we performed more detailed clustering than required
- Splitting is effectively assigned a weight equal to the cardinality of the smaller of the two resulting clusters

Computing MoJo distance

- MoJo distance can be computed in polynomial time
- Worst case complexity is $O(n^3)$ but with real data it is no worse than $O(n\log n)$

Assignment tool: mojo

- Takes two .rsf files containing different decompositions of the same set of entities
- Example: mojo dec1.rsf dec2.rsf
- Output: 383
- If the two decompositions do not refer to the same set of entities, only the intersection of the two sets is considered.

MoJoFM Effectiveness Metric

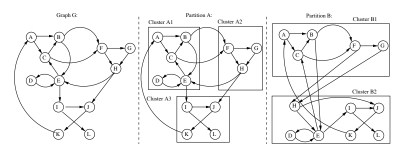
$$\textit{MoJoFM}(\textit{A}) = (1 - \frac{\textit{MoJo}(\textit{A}, \textit{B})}{\textit{max}_{\forall \textit{C}}(\textit{MoJo}(\textit{C}, \textit{B}))}) \times 100\%$$

- The denominator is the maximum possible MoJo distance to decomposition B
 - Can be computed by construction

EdgeSim

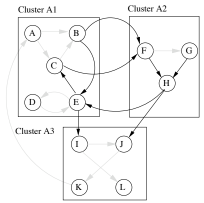
- Other measures do not consider edges
- · Edges might convey important information
- EdgeSim penalizes clustering algorithms for changing the edge types

EdgeSim



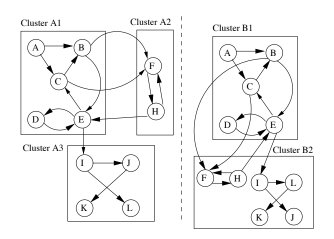
Inter-edge: Edge between clusters
Intra-edge: Edge within a cluster
Y: set of edges that are of the same type in both A
and B

EdgeSim example



EdgeSim(\overline{A} ,B) = $\frac{weight(Y)}{weight(E)}$ x 100% In this example, EdgeSim(\overline{A} ,B) = 52.6%

EdgeSim counterexample



EdgeMoJo philosophy

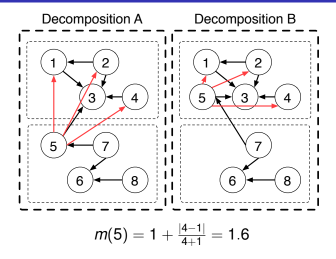
- A similarity measure cannot make assumptions as to what cluster a particular object should belong to
- Dissimilarity between decompositions should increase if the misplacement of an object results in the misplacement of a large number of edges

EdgeMoJo calculation

- Apply MoJo and obtain a series of Move and Join operations
- Perform all Join operations
- The cost of each Move operations increases from 1 to

$$m(o) = 1 + \frac{|(W(o, A_{new}) - W(o, A_{old})|}{W(o, A_{new}) + W(o, A_{old})}$$

EdgeMoJo example



EdgeMoJo in practice

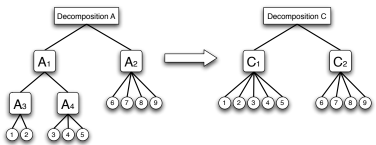
- Experiments with real and synthetic data indicated that EdgeMoJo distance is usually MoJo distance multiplied by a constant factor
- The usefulness of edges in measuring similarity between partitions is still an open question

What about nested decompositions?

- All measures we discussed so far assume a flat decomposition
 - No nested clusters
- Clustering algorithms typically create nested decompositions
- One needs to flatten decompositions to use these measures

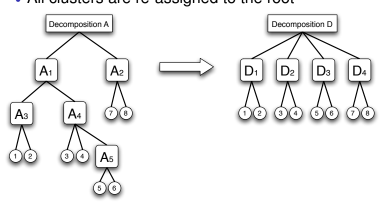
Creating compact flat decompositions

 All objects are re-assigned to the first level cluster they are transitively contained in



Creating detailed flat decompositions

· All clusters are re-assigned to the root



The END framework

- Each decomposition is transformed into a sequence of decompositions
- · One for each level in the containment tree
- Compute the value of a flat evaluation measure M at each level
- Obtain a vector of values S_{Mi}

The END framework

A different solution: UpMoJo

• Compute the combined similarity/distance S as

$$\sqrt{\sum(\textit{w}_{\textit{i}}{\mathcal{S}_{\textit{M}_{\textit{i}}}}^2)}$$
 where $\sum \textit{w}_{\textit{i}}=1$

• Weights need to be assigned to each level

- An extension of MoJo distance that includes an Up operation as well
 - Moves an object one level higher in the containment tree
- A series of Up operations ensures that the top level of the containment tree contains the same set of objects
- MoJo rearranges the top level
- The process repeats for each subtree of the top level