Conjugate Product Graphs for Globally Optimal 2D-3D Shape Matching
– Appendix –

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A1. Segmentation Pre-Matching

In Fig. A1, we visualise the pre-matching which is used by the approach in [3]. It is obvious that the injection of such information into the objective function makes finding valid solutions substantially easier.

![Figure A1. Visualisation of pre-matched segmentation information on cat from the TOSCA dataset. Different colours encode different segments, while darkest blue encodes the transition between different segments.](image)

For a fair comparison, in addition to our method that does not use this information, we also evaluate "Ours-Seg.", in which we incorporate the above segmentation information as an additional feature descriptor, see [3] for details.

A2. Branch and Bound Algorithm

Algorithm A1 describes our optimisation strategy. We adapt the branch and bound algorithm introduced in [3] to conjugate product graphs and implement runtime improvements by increasing chances of finding tighter upper bounds earlier.

The final goal of the optimisation is to find a cyclic path with minimal cost. However, Dijkstra’s algorithm only finds shortest (but not necessarily cyclic) paths. To that end, we represent the (conjugate) product graph as sequential graph, in which the first and last layers are duplicates, such that a path going from the same vertex in the first and last layer corresponds to a cyclic path.

Thus, the cyclic path with minimal cost can be found by computing the shortest path for every vertex on the first layer to every respective vertex on the last layer, and subsequently choosing among the computed paths the one with minimal cost. In general, this requires to solve a total of \(2|E_X| + |V_X|\) (ordinary) shortest path problems, and is computationally more expensive than the branch-and-bound strategy that we pursue.

The main idea of branch-and-bound is to iteratively subdivide the search space, while tightening upper and lower bounds using the results of previous iterations. In that sense, instead of searching for shortest paths from each vertex on the first layer to each respective vertex on the last layer, we search for the shortest path from a set of vertices \(B \subset V^*\) on the first layer to the respective set of vertices \(B \subset V^*\) on the last layer, see Fig. A2 (left). There is no guarantee that the path \(C = (v_1^1, \ldots, v_{|C|}^1)\) from \(B\) (first layer) to its duplicate \(B\) (last layer) with minimal energy is indeed cyclic, i.e., that the final vertex \(v_{|C|}^1\) in the last layer is indeed the same as the starting vertex \(v_1^1\) in the first layer. If \(C\) is not cyclic, we partition \(B\) into smaller, disjoint subsets \(B_1\) and \(B_2\) (with \(B_1 \cup B_2 = B\) and \(B_1 \cap B_2 = \emptyset\)) until a cyclic path is found (this is the branching strategy, see Fig. A2). The partitioning is done by calculating Voronoi cells around edges \(e_1^N\) and \(e_{|C|}^N\) on 3D shape assuming \(e_1^N\) and \(e_{|C|}^N\) are not identical (where the conjugate product vertex \(v_1^1 = (v_1^1, e_1^N)\) contains edge \(e_1^N\) on 3D shape and conjugate product vertex \(v_{|C|}^1 = (v_{|C|}^1, e_{|C|}^N)\) contains edge \(e_{|C|}^N\) on 3D shape). If \(e_1^N\) and \(e_{|C|}^N\) are identical we partition according to \(B_1 = B \setminus \{v_i^1\}\) and \(B_2 = \{v_i^1\}\).

The path cost \(d_C\) of non-cyclic paths (i.e. \(v_1^1 \neq v_{|C|}^1\)) is a lower bound \(b(\cdot)\) on the path cost of the globally optimal cyclic path. Whenever \(v_1^1\) and \(v_{|C|}^1\) are equal (meaning that \(C\) is a cyclic path), an upper bound \(b_{\text{upper}}\) is found, which might already be the globally optimal path, but can only be identified as such if all other branches do not yield cyclic paths with lower costs. Hence, the algorithm has to explore all other branches, which in the worst case are as many as there are vertices on one layer (i.e. \(2|E_X| + |V_X|\) many).

While searching for the optimal path, the algorithm only explores paths with cost \(d_C < b_{\text{upper}}\) and thus performance can be improved if tighter upper bounds \(b_{\text{upper}}\) are found as early as possible. We improve the branch-and-bound algorithm of [3] by computing all paths \(C_{\text{all}}\) of a branch, and then...
Figure A2. Illustration of the branching strategy in Algorithm A1. First, the shortest path from all vertices in $B$ on the first layer to the same vertices on the duplicate first layer (which amounts to the last layer) is computed. The resulting shortest path from $B$ on the first layer to $B$ on the last layer might not start and end on the same vertex (since we are searching for a shortest path from a set of vertices to a set of vertices). Whenever this is the case, $B$ is partitioned into two sets $B_1$ and $B_2$, for which in subsequent iterations shortest paths are computed analogously as for $B$.

search within these for cyclic paths to find lower values of the upper bound $b_{upper}$ earlier. We want to point out that no additional computational effort is required to compute $C_{all}$ using the implementation of [3], since all paths are already available (see Fig. A4 for runtime comparisons).

A3. Number of Conjugate Product Edges

As mentioned in the main paper, the conjugate product graph $P^*$ has 7 times more vertices than $P$ and $c \approx 11$ times more edges. In the following we derive $c$. To this end, we count outgoing edges of each conjugate product vertex (which is sufficient since $P^*$ is cyclic). Further, we assume that on average each vertex $j$ of the 3D shape $\mathcal{N}$ is connected to 6 edges [1]. Thus, each (directed) edge on 3D shape is connected to 5 other directed edges via their shared vertex, see Fig. A3.

In conclusion, each conjugate product vertex is connected to 5 conjugate product vertices on the same layer (reflecting degenerate 2D conjugate product vertices) and

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Input : 2D shape $\mathcal{M} = (\mathcal{V}_M, \mathcal{E}_M)$, 3D shape $\mathcal{N} = (\mathcal{V}_N, \mathcal{E}_N)$

Output: Optimal Path $C_{opt} \subset \mathcal{V}^*$

// First branch is complete first layer
$B_0 \leftarrow \{v^* = (e^\mathcal{M}_i, e^\mathcal{N}_i) \mid i_0 = 0, e^\mathcal{M} = (i_0, i_1)\}$;
// Initialise bounds and branches
$b(B_0) \leftarrow 0$;
$b_{upper} \leftarrow \infty$;
$B_{Branches} \leftarrow B_0$;
// Run until no branches with a gap between lower and upper bound exist
while $\min_{B \in B_{Branches}} b(B) < b_{upper}$ do

$B \leftarrow \arg\min_{B \in B_{Branches}} b(B)$;
$B_{Branches} \leftarrow B_{Branches} \setminus B$;
Compute all paths $C_{all} = \{C_1, C_2, \ldots\}$ with path cost $d_C < b_{upper}$ starting and ending in $B$;
if $C_{all} = \emptyset$ then
  // No path which meets $d_C < b_{upper}$
  continue;
$C \leftarrow \arg\min_{C \in C_{all}} d_C$;
// Check if current path is cyclic
if $v^*_1 = v^*_|C|$ then
  if $d_C < b_{upper}$ then
    $b_{upper} \leftarrow d_C$;
    $C_{opt} \leftarrow C$;
  else
    // Cut current branch into two parts
    if $e^\mathcal{N}_1 = e^\mathcal{N}_{|C|}$ then
      $B_1 \leftarrow B \setminus \{v^*_|C|\}$;
      $B_2 \leftarrow \{v^*_|C|\}$;
    else
      Compute $B_1, B_2$ as Voronoi cells around $e^\mathcal{N}_1$ and $e^\mathcal{N}_{|C|}$ respectively;
      // Add new branches
      $B_{Branches} \leftarrow B_{Branches} \cup \{B_1, B_2\}$;
      // Update lower bounds
      $b(B_1) = b(B_2) = d_C$;
      // Try to tighten upper bound
      for $C \leftarrow C_{all}$ do
        if $v^*_1 = v^*_|C|$ then
          if $d_C < b_{upper}$ then
            $b_{upper} \leftarrow d_C$;
            $C_{opt} \leftarrow C$;
```

Algorithm A1: Branch and Bound for Cyclic Shortest Path on (Conjugate) Product Graph

6 conjugate product vertices on the next layer (reflecting 5 non-degenerate conjugate product vertices and 1 degenerate 3D conjugate product vertex). In total, each conjugate
product vertex is connected to $c \approx 11$ other conjugate product vertices. In combination with the number of vertices of the conjugate product graph $|V^*| = |V_M| \cdot (2|E_X| + |V_X|)$ we obtain the number of edges of $P^*$.

**A4. Runtime**

**A4.1. Runtime Analysis**

In the following we estimate runtime complexity of our branch-and-bound algorithm for conjugate product graphs. To this end, we use $|E_X| \approx 3|V_X|$ [1] to obtain $|V^*| \approx 7 \cdot |V_M||V_X|$ and $|E^*| \approx c \cdot 7 \cdot |V_M||V_X|$.

The runtime of Dijkstra on an arbitrary graph $G = (V_G, E_G)$ is $O((|E_G| + |V_G|) \cdot \log(|V_G|))$ where $(|E_G| + |V_G|)$ indicates the number of update steps to be made, and $\log(|V_G|)$ indicates the complexity to access the priority heap that is used to keep track of the next nodes to be updated.

In our case, the number of update steps is $(|E^*| + |V^*|) \approx c \cdot 14 \cdot |V_M||V_X|$ (with $c \approx 11$). We make use of the strictly directed order of the $|V_M|$ layers of $P^*$, which allows to use a heap that scales with the number of vertices of one layer $O(|V_X|)$ (also see [3]). In summary, the runtime complexity of a single Dijkstra run in our conjugate product graph $P^*$ is $O(|V_M||V_X| \cdot \log(|V_X|))$.

To find the optimal cyclic path among all possible cyclic paths, we run Dijkstra not just once but at most $O(|V_X|)$ times (without any branch-and-bound optimisation), which leads to a final runtime complexity of $O(|V_M||V_X|^2 \cdot \log(|V_X|))$.

**A4.2. Runtime Comparison**

In Fig. A4, we show the median runtime for the approach by Lähner et al. [3] and our approach. The plot shows that both approaches have the same asymptotic behaviour. Due to the use of the larger conjugate product graph $P^*$ in comparison to product graph $P$ (see also A3), our approach takes by a constant factor more time to compute results. For a fair comparison with equal graph sizes, we additionally include computation times of our approach on the product graph $P$ which shows the improved performance when using Algorithm A1. Nevertheless, we emphasise that our approach (on $P^*$) still requires polynomial time while being the only one that is able to compute 2D-3D matchings without the need for pre-matching.

**A5. 2D to 3D Deformation Transfer**

We compute 2D to 3D deformation transfer by applying the following steps:

**2D-3D Matching** We find a matching between 2D and 3D shape using our approach.

**2D Deformation** We deform the 2D shape by using a skeleton which allows for different articulation of arms, legs and head. In combination with biharmonic weights [2, 7], we obtain a smooth deformation of the 2D shape (we tessellate the interior of the contour for biharmonic weight computation [4]).

**2D-3D Alignment** We find the optimal alignment $T^{3D}_{2D}$ of 2D shape and matched vertices on 3D shape by introducing a third, constant coordinate for 2D vertices and solving the (orthogonal) Procrustes problem [6].

**3D Deformation** We apply the deformation to the 3D shape by transforming the deformation on the 2D shape using $T^{3D}_{2D}$, applying the transformed deformation to a small subset of 3D vertices (chosen by furthest distance) and using their new positions as a constraint when deforming all other vertices of the 3D shape with the as-rigid-as-possible method of [5].

**A6. Ablation Studies**

**A6.1. Cost Function**

We evaluate the performance of different parts of our cost function in Tab. A1 as well as the performance of local rigidity when using multidimensional spectral features.

**A6.2. Discretisation**

In Tab. A2 we evaluate the robustness of our method w.r.t. to different discretisations. For all our experiments in the main paper we reduce influence of discretisation by decimating 3D shapes to half of their original resolution, which results in more uniform edge lengths [2]. Additionally, we re-sample 2D shapes with edge lengths according to the mean edge length of the decimated 3D shape.

In Fig. A5, we compare the performance of our method w.r.t. to different discretised 3D shapes. We observe that our method is relatively robust to different discretisation.
Table A1. Various cost functions on FAUST. The score is the area under the curve (AUC) of the cumulative segmentation errors. All introduced components increase performance. Our one-dimensional local thickness outperforms the multi-dimensional spectral features due to different intrinsic properties of 2D and 3D shapes.

<table>
<thead>
<tr>
<th>Method</th>
<th>AUC ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local Rigidity &amp; Spectral</td>
<td>0.95</td>
</tr>
<tr>
<td>Local Rigidity</td>
<td>0.76</td>
</tr>
<tr>
<td>Local Thickness</td>
<td>0.92</td>
</tr>
<tr>
<td>Local Rigid. &amp; Local Thick., ((\psi_1(x) = \psi_2(x) =</td>
<td>x</td>
</tr>
<tr>
<td>Ours</td>
<td><strong>0.98</strong></td>
</tr>
</tbody>
</table>

Table A2. Ablation study on the sensitivity of our approach to different discretisations. The score is the area under the curve (AUC) of the cumulative segmentation errors. We fix the discretisation of 3D shape and vary edge lengths of 2D shape. \(\bar{e}\) depicts the mean edge length of 3D shape.

<table>
<thead>
<tr>
<th>Mean Edge Length 2D Shape</th>
<th>AUC ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5-(e)</td>
<td>0.96</td>
</tr>
<tr>
<td>0.75-(e)</td>
<td>0.97</td>
</tr>
<tr>
<td>1-(e)</td>
<td><strong>0.98</strong></td>
</tr>
<tr>
<td>1.25-(e)</td>
<td>0.97</td>
</tr>
<tr>
<td>1.5-(e)</td>
<td>0.95</td>
</tr>
</tbody>
</table>

In Fig. A5 we show results for matchings across different resolution of 2D and 3D shapes. The experiment confirms that our method is robust in reasonable settings. In addition, we can see that matchings on the coarsest and on the finest level result in similar correspondences (cf. shape visualisations on the right of Fig. A6).

In Fig. A6 we show results for matchings across different resolution of 2D and 3D shapes. The experiment confirms that our method is robust in reasonable settings. In addition, we can see that matchings on the coarsest and on the finest level result in similar correspondences (cf. shape visualisations on the right of Fig. A6).

A6.3. Shape Discrepancies

In Fig. A7, we show results of cross-category matchings.

In Fig. A8, we evaluate our method’s robustness to noise when the 3D shape is disturbed by Gaussian noise. We plot the AUC (↑) of % of points in correct segment for each noise level.

| \(|V_M|\) | 39 | 79 | 159 | 318 |
|---|---|---|---|---|
| \(|V_N|\) | 432 | 863 | 1724 | 3446 |
| \(\bar{e}\) | 87.1 | 118.5 | 124.8 | |
| \(\bar{e}\) | 87.1 | 118.5 | 124.8 | |
| \(\bar{e}\) | 87.1 | 118.5 | 124.8 | |

Figure A6. Mesh resolution. We downsample various 2D and 3D shapes (FAUST). The values with green and yellow background are AUC (↑) of % of points in correct segment. The values with grey background are the number of vertices of respective 2D and 3D shapes. On the right we show that matchings from coarsest (\(|V_N| = 432, |V_M| = 39\)) and finest resolution (\(|V_N| = 3446, |V_M| = 318\)) are consistent.

Figure A7. Cross-category matching. We match the 2D shape of woman to the 3D shape of a woman and man, as well as to the 3D shape of dog. We can see that matching of woman to man results in a plausible matching, while matching of woman to dog does not yield meaningful results (as expected). The values are the resulting path-costs for each pair.

A6.4. Noise

In Fig. A8, we evaluate our method’s robustness to noise when the 3D shape is disturbed by Gaussian noise. We plot the AUC (↑) of % of points in correct segment for each noise level.

Figure A8. Robustness w.r.t. Noise. We apply Gaussian noise with \(\sigma = \{0.001, ..., 0.008\}\) to 3D shapes and subsequently match them to respective 2D shapes. Even under severe noise (see qualitative example on the left) our method is able to compute meaningful results which is confirmed by the plot on the right which shows AUC (↑) of % of points in correct segment (vertical axis) for increasing standard deviations (horizontal axis).
A7. Qualitative Results on FAUST

In Fig. A9 we show additional qualitative results.

Figure A9. Qualitative results on instances of FAUST dataset. We can see that left-right-flips occur (indicated with ↖) which nevertheless are plausible matchings.

References