Splitting Vertices in 2-Layer Graph Drawings

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Abstract

Bipartite graphs model the relationships between two disjoint sets of entities in several applications and are naturally drawn as 2-layer graph drawings. In such drawings, the two sets of entities (vertices) are placed on two parallel lines (layers), and their relationships (edges) are represented by segments connecting vertices. Methods for constructing 2-layer drawings often try to minimize the number of edge crossings. We use vertex splitting to reduce the number of crossings, by replacing selected vertices on one layer by two (or more) copies and suitably distributing their incident edges among these copies. We study several optimization problems related to vertex splitting, either minimizing the number of crossings or removing all crossings with fewest splits. While we prove that some variants are NP-complete, we obtain polynomial-time algorithms for others. We run our algorithms on a benchmark set of bipartite graphs representing the relationships between human anatomical structures and cell types.

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1 Introduction

Multilayer networks are used in many applications to model complex relationships between different sets of entities in interdependent subsystems [19]. When analyzing and exploring the interaction between two such subsystems S_t and S_b , bipartite or 2-layer networks arise naturally. The nodes of the two subsystem are modeled as a bipartite vertex set $V = V_t \cup V_b$ with $V_t \cap V_b = \emptyset$, where V_t contains the vertices of the first subsystem S_t and V_b those of S_b . The inter-layer connections between S_t and S_b are modeled as an edge set $E \subseteq V_t \times V_b$, forming a bipartite graph $G = (V_t \cup V_b, E)$. Visualizing this bipartite graph G in a clear and understandable way is then a key requirement for designing tools for visual network analysis [21].

A layered graph is a natural representation of the relationships between different genomic components, such as cell types and genes/proteins. Graphs of this type can be obtained using massively parallel sequencing in bulk at the single-cell level [3]. Börner et al. [3] studied different tasks expected to be performed on such graphs (e.g., finding neighbors, finding shared neighbors, etc.).

In a 2-layer graph drawing of a bipartite graph the vertices are drawn as points on two distinct parallel lines ℓ_t and ℓ_b , and edges are drawn as straight-line segments [9]. The vertices in V_t (top vertices) lie on ℓ_t (top layer) and those in V_b (bottom vertices) lie on ℓ_b (bottom layer). In addition to direct applications of 2-layer networks for modeling the relationships between two communities as mentioned above [21], such drawings also occur in tanglegram layouts for comparing phylogenetic trees [23] or as components in layered drawings of directed graphs [24].

Purchase [22] has studied different graph layout properties that have an impact on human understanding. In particular, it is known that edge crossings can significantly impact graph-based tasks, which is why minimizing the number of edge crossings (or removing all edge crossings altogether) is of interest.

The primary optimization goal for 2-layer graph drawings is to find permutations of one or both vertex sets V_t , V_b to minimize the number of edge crossings. While the existence of a crossing-free 2-layer drawing can be tested in linear time [7], the crossing minimization problem is NP-complete even if the permutation of one layer is given [9]. Hence, both fixed-parameter algorithms [17] and approximation algorithms [5] have been published. Further, graph layouts on two layers have also been widely studied in the area of graph drawing beyond planarity [6]. However, from a practical point of view, minimizing the number of crossings in 2-layer drawings may still result in visually complex drawings [15].

Hence, in this paper, as an alternative approach to construct readable 2-layer drawings, we study vertex splitting [8, 10, 16, 18]. The *vertex-split* operation (or *split*, for simplicity) for a vertex v deletes v from G, adds two new copies v_1 and v_2 (in the original vertex subset of G), and distributes the edges originally incident to v among the two new vertices v_1 and v_2 . Placing v_1 and v_2 independently in the 2-layer drawing can in turn reduce the number of crossings.

Vertex splitting has been studied in the context of the *splitting number* of an arbitrary graph G, which is the smallest number of vertex-splits needed to transform G into a planar graph. The splitting number problem is NP-complete, even for cubic graphs [11], but the splitting numbers of complete and complete bipartite graphs are known [12,14]. Vertex splitting has also been studied in the context of *split thickness*, which is the minimum maximum number of splits per vertex to obtain a graph with a certain property, e.g., a planar graph or an interval graph [10].

In this paper we consider reducing or removing edge crossings by vertex splitting. Vertex splitting is a useful technique to visualize complex pathway graphs of biological mechanisms. Nielsen et al. [20] have proposed an approach using Machine Learning to facilitate the visualization of pathway graphs by training a Support Vector Machine with actions taken during manual biocuration. Henry et al. [13] have improved the readability of clustered social networks using vertex splitting. These studies demonstrate the importance of vertex splitting for visualizing complex networks.

We study variations of the algorithmic problem of constructing planar or crossing-minimal 2-layer drawings with vertex splitting. In visualizing graphs defined on anatomical structures and cell types in the human body [1], the two vertex sets of G play different roles and vertex splitting is permitted only on one side of the layout. This motivates our interest in splitting only the bottom vertices. The top vertices may either be specified with a given context-dependent input ordering, e.g., alphabetically, following a hierarchy structure, or sorted according to an important measure, or we may be allowed to arbitrarily permute them to perform fewer vertex splits.

1.1 Contributions

We prove that for a given integer k it is NP-complete to decide whether G admits a planar 2-layer drawing with an arbitrary permutation on the top layer and at most k vertex splits on the bottom layer (see Theorem 3). NP-completeness also holds if at most k vertices can be split, but each an arbitrary number of times (see Theorem 5).

If, however, the vertex order of V_t is given, then we present two linear-time algorithms to compute planar 2-layer drawings, one minimizing the total number of splits (see Theorem 4), and one minimizing the number of split vertices (see Theorem 6). In view of their linear-time complexity, our algorithms may be useful for practical applications; we perform an experimental evaluation of the algorithm for Theorem 4 using real-world data sets stemming from anatomical structures and cell types in the human body [1].

We further study the setting in which the goal is to minimize the number of crossings (but not necessarily remove all of them) using a prescribed total number of splits. For this setting, we prove NP-completeness even if the vertex order of V_t is given (see Theorem 7). On the other hand, we provide an XP-time algorithm parameterized by the number of allowed splits (see Theorem 8), which, in other words, means that the algorithm has a polynomial running time for any fixed number of allowed splits.

2 Preliminaries

We denote the order of the vertices in V_t and V_b in a 2-layer drawing by π_t and π_b , resp. If a vertex u precedes a vertex v, then we denote it by $u \prec v$. Although 2-layer drawings are defined geometrically, their crossings are fully described by π_t and π_b , as in the following folklore lemma.

▶ Lemma 1. Let Γ be a 2-layer drawing of a bipartite graph $G = (V_t \cup V_b, E)$. Let (v_1, u_1) and (v_2, u_2) be two edges of E such that $v_1 \prec v_2$ in π_t . Then, edges (v_1, u_1) and (v_2, u_2) cross each other in Γ if and only if $u_2 \prec u_1$ in π_b .

In the following we formally define the problems we study. For all of them, the input contains a bipartite graph $G = (V_t \cup V_b, E)$ and a split parameter k.

Crossing Removal with k Splits – CRS(k): Decide if there is a planar 2-layer drawing of G after applying at most k vertex-splits to the vertices in V_b .

1:4 Splitting Vertices in 2-Layer Graph Drawings

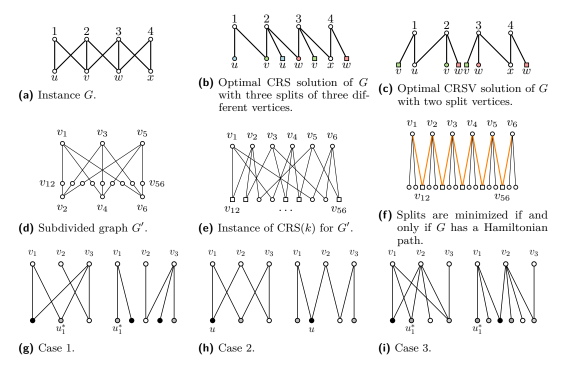


Figure 1 Differences between CRS and CRSV problems (a)–(c); illustrations for the reduction in Theorem 3 (d)–(f); illustrations for the optimization algorithm for CRS(k) with Fixed Order, where vertices in N^+ are colored in shades of gray (g)–(i).

- **Crossing Removal with k Split Vertices CRSV**(k): Decide if there is a planar 2-layer drawing of G after splitting at most k original vertices of V_b .
- **Crossing Minimization with k Splits** CMS(k, M): Decide if there is a 2-layer drawing of G with at most M crossings after applying at most k vertex-splits to the vertices in V_b , where M is an additional integer specified as part of the input.

Note that in CRSV(k), once we decide to split an original vertex, then we can further split its copies without incurring any additional cost. The example in Figure 1a–1c demonstrates the difference between CRS and CRSV.

For all problems, we refer to the variant where the order π_t of the vertices in V_t is given as part of the input by adding the suffix "with Fixed Order".

The following lemma implies conditions under which a vertex split must occur.

- ▶ **Lemma 2.** Let $G = (V_t \cup V_b, E)$ be a bipartite graph and let $u \in V_b$ be a bottom vertex adjacent to two top vertices $v_1, v_2 \in V_t$, with $v_1 \prec v_2$ in π_t . In any planar 2-layer drawing of G in which u is not split, we have that:
- C.1 A top vertex that appears between v_1 and v_2 in π_t can only be adjacent to u;
- **C.2** In π_b , u is the last neighbor of v_1 and the first neighbor of v_2 .

Proof. If there is a top vertex v' between v_1 and v_2 adjacent to a bottom vertex $u' \neq u$, then (v', u') crosses (v_1, u) or (v_2, u) . If there is a neighbor u'' of v_1 after u in π_b , then the edges (v_1, u'') and (v_2, u) cross. A symmetric argument holds when there is a neighbor of v_2 before u in π_b .

3 Crossing Removal with k Splits

In this section, we prove that the CRS(k) problem is NP-complete in general and linear-time solvable when the order π_t of the top vertices is part of the input.

▶ **Theorem 3.** The CRS(k) problem is NP-complete.

Proof. The problem belongs to NP since, given a set of at most k splits for the vertices in V_b , we can check whether the resulting graph is planar 2-layer [7].

We use a reduction from the Hamiltonian Path problem to show the NP-hardness. Given an instance G = (V, E) of the Hamiltonian Path problem, we denote by G' the bipartite graph obtained by subdividing every edge of G once (Figure 1d). We construct an instance of the CRS(k) problem (Figure 1e) by setting the top vertex set V_t to consist of the original vertices of G, the bottom vertex set V_b to consist of the subdivision vertices of G', and the split parameter to k = |E| - |V| + 1. The reduction can be easily performed in linear time. We prove the equivalence.

Suppose that G has a Hamiltonian path v_1, \ldots, v_n . Set $\pi_t = v_1, \ldots, v_n$, and split all the vertices of V_b , except for the subdivision vertex of the edge (v_i, v_{i+1}) , for each $i = 1, \ldots, n-1$ (Figure 1f). This results in $|V_b| - (n-1)$ splits, which is equal to k, since $|V_b| = |E|$ and n = |V|. We then construct π_b such that, for each $i = 1, \ldots, n-1$, all the neighbors of v_i appear before all the neighbors of v_{i+1} , with their common neighbor being the last neighbor of v_i and the first of v_{i+1} . This guarantees that both conditions of Lemma 2 are satisfied for every vertex of V_b . Together with Lemma 1, this guarantees that the 2-layer drawing is planar.

Suppose now that G' admits a planar 2-layer drawing with at most |E| - |V| + 1 splits. Since $|E| = |V_b|$ and every vertex of V_b has degree exactly 2 (subdivision vertices), there exist at least |V| - 1 vertices in V_b that are not split. Consider any such vertex $u \in V_b$. By C.1 of Lemma 2, the two neighbors of u are consecutive in π_t . Also, these vertices are connected in G by the edge whose subdivision vertex is u. Since this holds for each of the at least |V| - 1 non-split vertices, we have that each of the |V| - 1 distinct pairs of consecutive vertices in V_t (recall that $V_t = V$) is connected by an edge in G. Thus, G has a Hamiltonian path.

Next, we focus on the optimization version of the CRS(k) with the Fixed Order problem. Our recursive algorithm considers a constrained version of the problem, where the first neighbor in π_b of the first vertex in π_t may be prescribed. At the outset of the recursion, there exists no prescribed first neighbor. The algorithm returns the split vertices in V_b and the corresponding order π_b .

In the base case, there is only one top vertex v, i.e., $|V_t| = 1$. Since all vertices in V_b have degree 1, no split takes place. We set π_b to be any order of the vertices in V_b where the first vertex is the prescribed first neighbor of v, if any.

In the recursive case when $|V_t| > 1$, we label the vertices in V_t as $v_1, \ldots, v_{|V_t|}$, according to π_t . If the first neighbor of v_1 is prescribed, we denote it by u_1^* . Also, we denote by N^1 the set of degree-1 neighbors of v_1 , and by N^+ the other neighbors of v_1 . Note that only the vertices in N^+ are candidates to be split for v_1 . In particular, by C.1 of Lemma 2, a vertex in N^+ can avoid being split only if it is also incident to v_2 . Further, since any vertex in N^+ that is not split must be the last neighbor of v_1 in π_b , by C.2 of Lemma 2, at most one of the common neighbors of v_1 and v_2 will not be split. Analogously, if u_1^* is prescribed, then it must be split, unless v_1 has degree 1.

In view of these properties, we distinguish three cases based on the common neighborhood of v_1 and v_2 . In all cases, we will recursively compute a solution for the instance composed

of the graph $G' = (V'_t \cup V'_b, E')$ obtained by removing v_1 and the vertices in N^1 from G, and of the order $\pi'_t = v_2, \ldots, v_{|V_t|}$. We denote by π'_b and s' the computed order and the number of splits for the vertices in V'_b . In the following we specify for each case whether the first neighbor of v_2 in the new instance is prescribed or not, and how to incorporate the neighbors of v_1 into π'_b .

Case 1: v_1 and v_2 have no common neighbor; see Figure 1g. In this case, we do not prescribe the first neighbor of v_2 in the instance composed of G' and π'_t . To compute a solution for the original instance, we split each vertex in N^+ so that one copy becomes incident only to v_1 . We construct π_b by selecting the prescribed vertex u_1^* , if any, followed by the remaining neighbors of v_1 in any order and, finally, by appending π'_b . This results in $s = |N^+| + s'$ splits.

Case 2: v_1 and v_2 have exactly one common neighbor u. If $u = u_1^*$ and v_1 have a degree larger than 1, then u cannot be the last neighbor of v_1 and must be split. Thus, we perform the same procedure as in Case 1. Otherwise, in the instance composed of G' and π'_t , we set u as the prescribed first neighbor of v_2 ; bluesee Figure 1h. To compute a solution for the original instance, we split each vertex in N^+ , except u, so that one copy becomes incident only to v_1 . We construct π_b by selecting the prescribed vertex u_1^* , if any, followed by the remaining neighbors of v_1 different from u in any order and, finally, by appending π'_b . This results in $s = |N^+| -1 + s'$ splits.

Case 3: v_1 and v_2 have more than one common neighbor. If v_1 and v_2 have exactly two common neighbors u, u' and one of them is u_1^* , say $u = u_1^*$, then u cannot be the last neighbor of v_1 , as v_1 has degree larger than 1. Thus, we proceed exactly as in Case 2, using u' as the only common neighbor of v_1 and v_2 .

Otherwise, there are at least two neighbors of v_1 different from u_1^* ; see Figure 1i. We want to choose one of these vertices as the last neighbor of v_1 , so that it is not split. However, the choice is not arbitrary as this may affect the possibility for v_2 to save the split for a neighbor it shares with v_3 . In the instance composed of G' and π'_t , we do not prescribe the first vertex of v_2 . To compute a solution for the original instance, we simply choose as the last neighbor of v_1 any of its common neighbors with v_2 that has not been set as the last neighbor of v_2 in π'_b . Such a vertex, say u, always exists since v_1 and v_2 have at least two common neighbors different from u_1^* , and can be moved to become the first vertex in π'_b . Specifically, we split all the vertices in N^+ , except for u, so that one copy becomes incident only to v_1 . We construct π_b by selecting the prescribed vertex u_1^* , if any, followed by the remaining neighbors of v_1 different from v_1 in any order. We then modify v_2 by moving v_1 to be the first vertex. Note that this operation does not affect planarity, as it only involves reordering the set of consecutive degree-1 vertices incident to v_2 . Finally, we append the modified v_2 . This results in v_1 in v_2 splits.

▶ **Theorem 4.** For a bipartite graph $G = (V_t \cup V_b, E)$ and an order π_t of V_t , the optimization version of CRS(k) with Fixed Order can be solved in O(|E|) time.

Proof. By construction, for each $i=1,\ldots,|V_t|-1$, all neighbors of v_i precede all neighbors of v_{i+1} in π_b . Thus, by Lemma 1, the drawing is planar. The minimality of the number of splits follows from Lemma 2, as discussed before the case distinction. In particular, any minimum-splits solution can be shown to be equivalent to the one produced by our algorithm. The time complexity follows as each vertex only needs to check its neighbors a constant number of times.

We conclude this section by mentioning that the CRS(k) problem had already been considered, under a different terminology, in the context of molecular QCA circuits design [4].

Algorithm 1 CRS(k)-with-Fixed-Order(V_t, V_b, π_t)

```
if there is only one vertex in V_t then

Let N(v) be the neighbor vertices of v
Output an arbitrary order N(v)
else

Let v_1 and v_2 are the first two vertices in V_t
if N(v_1) \cap N(v_2) \neq \emptyset then

Let N^+(v_1) be the set of neighbors of v_1 having degree more than one let \pi'_b be the output of \operatorname{CRS}(k)-with-Fixed-Order(V_t - v_1, V_b, \pi_t - v_1)

Find a common neighbor u of v_1 and v_2 not in \pi'_b
output u, \pi'_b
else

let \pi'_b be the output of \operatorname{CRS}(k)-with-Fixed-Order(V_t - v_1, V_b, \pi_t - v_1)
Output N(v_1), \pi'_b
end if
```

Here, the problem was claimed to be NP-complete, without providing a formal proof. In the same work, when the order π_t of the top vertices is part of the input, an alternative algorithm was proposed based on the construction of an auxiliary graph that has superlinear size. Exploiting linear-time sorting algorithms and observations that allow avoiding explicitly constructing all edges of this graph, the authors were able to obtain a linear-time implementation. We believe that our algorithm of Theorem 4 is simpler and more intuitive, and directly leads to a linear-time implementation.

4 Crossing Removal with k Split Vertices

In this section, we prove that the CRSV(k) problem is NP-complete in general and linear-time solvable when the order π_t of the top vertices is part of the input. Ahmed et al. [2] showed that CRSV(k) is FPT when parameterized by k. To prove the NP-completeness we can use the reduction of Theorem 3. In fact, in the graphs produced by that reduction all vertices in V_b have degree 2. Hence, the number of vertices that are split coincides with the total number of splits.

▶ **Theorem 5.** The CRSV(k) problem is NP-complete.

For the version with Fixed Order, we first use C.1 of Lemma 2 to identify vertices that need to be split at least once, and repeatedly split them until each has degree 1. For a vertex $u \in V_b$, we can decide if it needs to be split by checking whether its neighbors are consecutive in π_t and, if u has degree at least 3, all its neighbors different from the first and last have degree exactly 1.

We first perform all necessary splits. For each $i = 1, ..., |V_t| - 1$, consider the two consecutive top vertices v_i and v_{i+1} . If they have no common neighbor, no split is needed. If they have exactly one common neighbor u, then we set u as the last neighbor of v_i and the first of v_{i+1} , which allows us not to split u, according to C.2. Since u did not participate in any necessary split, if u is also adjacent to other vertices, then all its neighbors have degree 1, except possibly the first and last. Hence, C.2 can be guaranteed for all pairs of consecutive neighbors of u.

Otherwise, v_i and v_{i+1} have at least two common neighbors and thus have degree at least 2. Hence, all common neighbors of v_i and v_{i+1} must be split, except for at most one, namely the one that is set as the last neighbor of v_i and as the first of v_{i+1} . Since all these vertices are incident only to v_i and v_{i+1} , as otherwise they would have been split by C.1, we can arbitrarily choose any of them, without affecting the splits of other vertices.

Algorithm 2 CRSV(k)-with-Fixed-Order(V_t, V_b, π_t)

```
for i = 1, \dots, |V_t| - 1 do
   if v_i and v_{i+1} has no common neighbors then
       No split is needed
   end if
   if v_i and v_{i+1} has exactly one common neighbor then
       Let u be the common neighbor
       if u is also adjacent to other vertices that has degree 1 except the first an last one
then
          No split is needed
       else
          Split u when mandatory
       end if
   else
       Split all common neighbors except at most once
   end if
end for
```

At the end we construct the order π_b so that, for each $i=1,\ldots,|V_t|-1$, all the neighbors of v_i precede all the neighbors of v_{i+1} , and the unique common neighbor of v_i and v_{i+1} , if any, is the last neighbor of v_i and the first of v_{i+1} . By Lemma 1, this guarantees planarity. Identifying and performing all unavoidable splits and computing π_b can be easily done in O(|E|) time. Since we only performed unavoidable splits, as dictated by Lemma 2, we have the following.

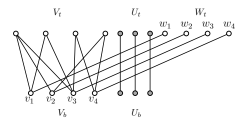
▶ **Theorem 6.** For a bipartite graph $G = (V_t \cup V_b, E)$ and an order π_t of V_t , the optimization version of CRSV(k) with Fixed Order minimizing the number of split vertices can be solved in O(|E|) time.

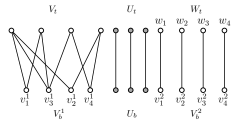
5 Crossing Minimization with k Splits

In this section we consider minimizing crossings (not necessarily removing all), by applying at most k splits. We first prove NP-completeness of the decision problem $\mathrm{CMS}(k,M)$ with Fixed Order and then give a polynomial-time algorithm assuming the integer k is a constant.

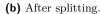
▶ **Theorem 7.** For a bipartite graph $G = (V_t \cup V_b, E)$, an order π_t of V_t , and integers k, M, problem CMS(k, M) with Fixed Order is NP-complete.

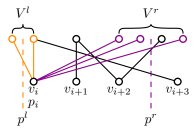
Proof. We reduce from the NP-complete DECISION CROSSING PROBLEM (DCP) [9], where given a bipartite 2-layer graph with one vertex order fixed, the goal is to find an order of the other set such that the number of crossings is at most a given integer M. Given an instance of DCP, i.e., a 2-layer graph $G = (V_t \cup V_b, E)$, with ordering π_t of V_t and integer M, we construct an instance G' of CMS(k, M) where $k = |V_b|$. First let $G' = (V_t' \cup V_b', E')$ be a copy of G. We give an arbitrary ordering π_b to the vertices of V_b' . We then add, respectively,





(a) Instance of CMS(k) constructed from a DCP instance, in light gray the vertices in $U_t \cup U_b$; before splitting.





- (c) Vertex v_i with span 6 and a split in V^l (span 1) and V^r (span 3). The barycenter of V^r is p^r . When moving right from p_i in CR-count we process v_{i+1} (reduces $|V^r|$ crossings) and v_{i+2} (reduces $|V^r|$ but adds 1 crossing), but not v_{i+3} .
- Figure 2 Illustrations for Theorem 7 (a)–(b); illustration for the crossing reduction heuristics (c).

to each vertex set V_t and V_b a set U_t and U_b of M+1 vertices and connect each $u \in U_t$ to exactly one $v \in U_b$, forming a matching of size M+1. We add the vertices of U_t to π_t (resp. U_b to π_b) after all the vertices of V_t (V_b). We lastly add a set W_t of k vertices to V_t' , placed at the end of π_t , such that each $w_i \in W_t$ (i = 1, ..., k) has exactly one neighbor $v_i \in V_b$ and vice versa; see Figure 2a.

Given an ordering π_b^* of V_b that results in a drawing of G with at most M crossings, we show that we can solve the $\mathrm{CMS}(k,M)$ instance G'. In G', we split each vertex of V_b to obtain the sets V_b^1 and V_b^2 in which we place exactly one copy of each original vertex. We place V_b^2 after the vertices of U_b in π_b in the same order that the vertices of W_t appear in π_t and draw a single edge between the copies and their neighbor in W_t . We place V_b^1 before the vertices of M_2 in π_b in the same order as in π_b^* . The graph induced by V_b^1 and V_t is the same graph as G, hence it has at most M crossings. Since V_t only has neighbors in V_b and all those neighbors are in V_b^1 , it has no other outgoing edges, similarly, all edges incident to vertices in W_t are assigned to the copies in V_b^2 . The remaining graph is crossing-free as the vertices in U_t and W_t form a crossing-free matching with the vertices in U_b and V_b^2 .

Conversely, let G^* be a 2-layer drawing obtained from G' after k split operations that has at most M crossings. Since each vertex $v \in V_b$ has a neighbor $w \in W_t$, it induces M+1 crossings with edges induced by the vertices in $U_t \cup U_b$. Since the vertices in U_b have a single neighbor, they cannot be split, thus every vertex in V_b is split once, and their neighborhood are partitioned for each copy in the following way: one copy receives the neighbor in W_t and one copy receives the remaining neighbors, which are in V_t (Figure 2b), thus avoiding the at least M+1 crossings induced by $U_t \cup U_b$. Any other split would imply at least M+1 crossings. The graph induced by the copies that receive the neighbors in V_t has at most M crossings, thus, the ordering found for those copies is a solution to the DCP instance G.

Next, we present a simple XP-time algorithm for the crossing minimization version of CMS(,) parameterized by the number k of splits, i.e., the algorithm runs in polynomial time $O(n^{f(k)})$, where n is the input size, k is the parameter, and f is a computable function. Let $G = (V_t \cup V_b, E)$ be a 2-layer graph with vertex orders π_t and π_b and let k be the desired number of splits. Our algorithm executes the following steps. First, it determines a set of splits by choosing k times a vertex from the n vertices in V_b – we enumerate all options. For any vertex $v \in V_b$ split i times in the first step, v is replaced by the set of copies $\{v_1, ..., v_{i+1}\}$. The neighborhood N(v) of a vertex $v \in V_b$ is a subset of V_t ordered by π_t . We partition this ordered neighborhood into i+1 consecutive subsets, i.e., for each subset, all its elements are sequential in N(v) – again, we enumerate all possible partitions. Each set is assigned to be the neighborhood of one of the copies of v. The algorithm then chooses an ordering of all copies of all split vertices and attempts all their possible placements by merging them into the order π_b of the unsplit vertices of V_b . The crossing number of every resulting layout is computed and the graph with minimum crossing number yields the solution to our input. It remains to show that the running time of this algorithm is polynomial for constant k.

▶ Theorem 8. For a 2-layer graph $G = (V_t \cup V_b, E)$ with vertex orders π_t, π_b and a constant $k \in \mathbb{N}$ we can minimize the number of crossings by applying at most k splits in time $O(n^{4k})$.

Proof. Let G^* be a crossing-minimal solution after applying k splits on V_b and let us assume that our algorithm would not find a solution with this number of crossings. As our algorithm considers all possibilities to apply k splits, it also attempts the splits applied in G^* . Similarly, the neighborhood partition of G^* and the copy placement are explicitly considered by the algorithm as it enumerates all possibilities. Hence a solution at least as good as G^* is found, proving correctness.

Let $n_t = |V_t|$ and $n_b = |V_b|$ with $n = n_t + n_b$. The algorithm initially chooses k times from n_b vertices leading to n_b^k possible sets of copies. Since a vertex has degree at most n_t , there are at most n_t^k possible neighborhoods for each copy. Additionally, there are (2k)! orderings of at most 2k copies. Lastly, there are n_b^{2k} possible placement of the 2k ordered copies between the at most n_b unsplit vertices in π_b . This leads to an overall runtime of $O((2k)! \cdot n^{4k}) = O(n^{4k})$ to iterate through all possible solutions and select the one with a minimum number of crossings.

6 Crossing Reduction Heuristics

In this section we present two greedy heuristics to iteratively reduce crossings in a two-layer drawing by selecting and splitting vertices; see Algorithms 3–4 in the supplemental material for the pseudocode. The input to the algorithm is a bipartite graph $G = (V_t \cup V_b, E)$, order π_t of V_t and order π_b of V_b . Here we use the barycenter heuristic for the initial orders, but any initial order computed by a crossing-reduction algorithm can also be used. Additionally, an input parameter k is specified that represents a budget of available split operations. For both heuristics we iteratively perform k splits by selecting the most promising vertex in V_b and a partition of its respective neighbors in V_t into a consecutive set of left neighbors V^l and a consecutive set of right neighbors V^r . After splitting, the original vertex receives the set V^l assigned as neighbors and the copy receives the set V^r . Next we describe how the vertex to be split is selected in each of the heuristics.

Firstly, in the case of the max-span heuristic we select the vertex v with the maximum span, i.e., the maximum distance between its leftmost neighbor and rightmost neighbor in π_t . Then, we process v by iterating in order over its neighbors and assigning them to either

the set of left neighbors V^l or right neighbors V^r depending on the index; see Fig. 2c. In each iteration we compute the sum of the squared span of V^l and V^r . The minimum value indicates the best partition of v's neighbors. The complexity of the heuristic is linear in the number of edges O(|E|).

Secondly, the CR-count heuristic selects promising split vertices by computing crossings that can be reduced and selects the vertex with potentially most reduced crossings. First, we assign each vertex v_i in V_b a position $p_i \in \mathbb{R}$ which is the barycenter of positions in π_t of its neighbors in V_t . Similarly to max-span, we iterate in order over the neighbors of all v_i creating partitions V^l and V^r . In each iteration we compute the barycenter p^l of V^l and p^r of V^r ; see Figure 2c. Next, we start from position p_i and move in ascending order processing all other vertices in V_b until we reach p^r . To process a vertex $v_i \in V_b$ we look at all edges to its neighbors $N(v_i) \subseteq V_t$ and use a case distinction to count how many crossings can be reduced. In the first case a neighbor in V_t is left of the leftmost vertex in V^r . Here, we would reduce $|V^r|$ crossings as the new position of v_r would not cross. In the second case a neighbor is between the leftmost and rightmost vertex in V^r , i.e., we would reduce some crossings but add others. In the third case a neighbor is right of the rightmost vertex in V^r , i.e., no crossings can be reduced. Likewise, we process vertices to the left of v_i up to the barycenter p^l . Finally, after computing all potentially reducible crossings for each vertex and split combination, we select the combination reducing the crossings most, assign V_l to the original vertex and V_r to the new copy. Both vertices are positioned in their respective barycenters p^l and p^r in the order. The complexity of the algorithm is $O(|V_b|^2|V_t|)$.

Algorithm 3 max-span heuristic

```
Require: G = (V_t \cup V_b, E), \, \pi_t, \, \pi_b, \, k
Ensure: G', V'_b, \pi'_b
  G' \leftarrow G
  V_b' \leftarrow V_b
  \pi_b' \leftarrow \pi_b
  while k > 0 do
       maxSpan \leftarrow 0
       splitNode \leftarrow None
       for i=1,\cdots,|V_b'| do
           // compute index of leftmost/rightmost neighbor of v_i in \pi_t
           minNeighbor \leftarrow leftmostNeighbor(v_i, N(v_i), \pi_t)
           maxNeighbor \leftarrow rightmostNeighbor(v_i, N(v_i), \pi_t)
           span \leftarrow maxNeighbor - minNeighbor
           if span > maxSpan then
                maxSpan \leftarrow span
                splitNode \leftarrow v_i
           end if
       end for
       if maxSpan == 0 then // no split possible
           return G, V_h', \pi_h'
       else
           N^o \leftarrow orderByIndex(N(splitNode), \pi_t)
           V^l \leftarrow \emptyset
           V^r \leftarrow N^o
           minSpan \leftarrow (computeSpan(V^r, \pi_t))^2
           V^l_s \leftarrow V^l
           V_{s}^{r} \leftarrow V^{r}
           for i = 1, ..., |N^o| do
                V^l \leftarrow V^l \cup \{v_i\}
                V^r \leftarrow V^r \setminus \{v_i\}
                span \leftarrow (computeSpan(V^l, \pi_t))^2 + (computeSpan(V^r, \pi_t))^2
                if span < minSpan then
                    minSpan \leftarrow span
                    V_s^l \leftarrow V^l
                    V_s^r \leftarrow V^r
                end if
           end for
           // update G and \pi_b by removing v_i and adding v_l and v_r with neighbors V_s^l and V_s^r
           //v_l and v_r get the placed in the barycenter of V_s^l and V_s^r in the order \pi_b'
           updateGraphAndOrder(G', V'_b, \pi'_b, V^l_s, V^r_s)
       end if
       k \leftarrow k-1
  end while
  return G, V_b', \pi_b
```

Algorithm 4 CR-count heuristic

```
Require: G = (V_t \cup V_b, E), \pi_t, \pi_b, k
Ensure: G', V_b', \pi_b'
   G' \leftarrow G
  V_b' \leftarrow V_b \\ \pi_b' \leftarrow \pi_b
   while k > 0 do
       V_s^l \leftarrow \emptyset \\ V_s^r \leftarrow \emptyset
        bestSplit \leftarrow 0
        for i = 1, ..., |V_b'| do
            N^o \leftarrow orderByIndex(N(splitNode), \pi_t)
            V^l \leftarrow \emptyset
            V^r \leftarrow N^o
            p_i \leftarrow \text{computeBarycenter}(G, N^o, \pi_t)
            for j = 1, ..., |N^o| do
                 V^l \leftarrow V^l \cup \{v_i\}
                 V^r \leftarrow V^r \setminus \{v_j\}
                 p^l \leftarrow \text{computeBarycenter}(G, V^l, \pi_t)
                 p^r \leftarrow \text{computeBarycenter}(G, V^r, \pi_t)
                 // count avoidable crossings in a range, i.e. between p_i and p^r
                 count^l \leftarrow countCrossingsInRange(G, \pi_t, \pi'_h, p^l, p_i)
                 count^r \leftarrow \text{countCrossingsInRange}(G, \pi_t, \pi_h', p_i, p^r)
                 if bestSplit < count^l + count^r then
                      bestSplit \leftarrow count^l + count^r
                      V_s^l \leftarrow V^l
                      V_c^r \leftarrow V^r
                 end if
            end for
        end for
        if bestSplit == 0 then // no split possible
            return G, V_h', \pi_h'
        else
             // update G and \pi_b by removing v_i and adding v_l and v_r with neighbors V_s^l and V_s^r
            //v_l and v_r get the placed in the barycenter of V_s^l and V_s^r in the order \pi_b'
            updateGraphAndOrder(G', V'_b, \pi'_b, V^l_s, V^r_s)
        end if
        k \leftarrow k-1
   end while
  return G, V_b', \pi_b
```

7 Experimental Results

We have experimentally evaluated four of the five algorithms described earlier with 22 real-world datasets: the exact algorithm of CRS(k) with Fixed Order, the exact algorithm of CRSV(k) with Fixed Order, and two heuristics for crossing reduction. The algorithm behind Thm. 8 is inefficient in practice. We analyze performance w.r.t. the number of crossings in the layouts, number of vertex splits, number of vertices that we split, the maximum number

1:14 Splitting Vertices in 2-Layer Graph Drawings

of splits, and running time.

7.1 Experimental Design

We have mentioned that 2-layer drawings have been applied in visualizing graphs defined on anatomical structures and cell types in the human body [1]. There exists a variety of cell types, genes, and proteins related to different organs of the human body. Hierarchical structures have been used to show the relationship between organs to anatomical structures, anatomical structures to cell types, and cell types to genes/proteins. Cell types and genes/proteins situate on a particular layer, unlike anatomical structures. Hence, we can consider a 2-layer graph G where cell types represent one layer and genes/proteins represent another layer and analyze G before and after splitting. In this section, we consider the real-world 2-layer graphs generated from the dataset of different organs and show the experimental results obtained on those graphs.

7.2 Datasets

We use 22 real-world instances of 2-layer graphs from [1]; see Table 1.

Organ	V	E	Cell types	Genes/proteins	Density	Max degree
Blood	179	461	30	149	0.0289	57
Fallopian Tube	42	32	19	23	0.0371	3
Lung	231	231	69	162	0.008	8
Peripheral Nervous System	3	2	1	2	0.666	2
Thymus	552	658	41	511	0.00432	93
Heart	60	51	15	45	0.028	7
Lymph Nodes	299	491	44	255	0.0110	36
Prostate	43	36	12	31	0.039	3
Ureter	44	53	14	30	0.0560	9
Bone Marrow	343	662	45	298	0.011	25
Kidney	201	237	58	143	0.011	8
Skin	102	90	36	66	0.017	7
Urinary Bladder	46	55	15	31	0.053	9
Brain	381	346	127	254	0.004	5
Large Intestine	124	139	51	73	0.0182	8
Ovary	9	6	3	6	0.166	2
Small Intestine	18	13	5	13	0.084	4
Uterus	61	65	16	45	0.035	9
Eye	145	270	47	98	0.0258	68
Liver	73	57	26	47	0.0216	5
Pancreas	69	100	29	40	0.042	12
Spleen	290	414	65	225	0.009	23

Table 1 Statistics about the organ graphs from the HubMAP dataset [1]. The density of a graph G = (V, E) with $V = V_t \cup V_b$ is defined as 2|E|/(|V|(|V|-1)).

7.3 Interactive Tool Design

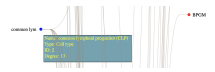
We developed an interactive tool, where the user can upload a dataset and visualize the corresponding 2-layer graph; see Fig. 5. A dataset can be loaded by pasting JSON-formatted text in the input area on the left-hand side of the interface. We use blue and red colors to draw nodes that represent cell types and genes/proteins respectively. There is a legend in the top left corner of the interface to describe the color code. Instead of using top and bottom layers, we use left and right layers, for easier node labeling (i.e., the left and right layers represent V_t and V_b). There are multiple radio buttons to the configuration of the drawing. The user can fix the order of either the blue vertices or the red vertices by selecting one set of radio buttons. The number of vertex splits depends on the initial layout. We consider two types of initial layouts: the vertices in each layer are positioned in alphabetical order, or in barycentric order [24]. The user can select an initial order from the input interface by using another set of radio buttons. There are "Draw" and "Split" buttons in the interface. Once the user selects an order for the left layer and clicks the draw button, then the initial ordered layout will be shown on the right side of the interface. Clicking the split button replaces the initial layout and shows the final layout on the right side of the interface.

The right output interface is interactive; the user can see further details using different interactions. When the graph is large the user can scroll up and down to see different parts of the layout. The user can highlight the adjacent edges by clicking on a particular vertex in case of dense layouts. We keep the label texts less than or equal to ten characters. If a label is longer then we show the first ten characters and truncate the rest. If the user puts the mouse over the label or the corresponding vertex, a pop-up message will show the full label. If the user moves out the mouse, the message will be removed too. Besides showing the full label, we also provide other useful information, e.g., the degree and ID of the vertex; see Figure 3.



(a) The system highlights the adjacent edges when the user clicks on a vertex ("basophil" in this case).

Figure 3 Interacting with the system.



(b) A pop-up message showing the full label, and other related information.

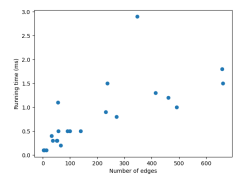
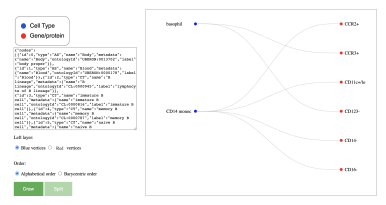
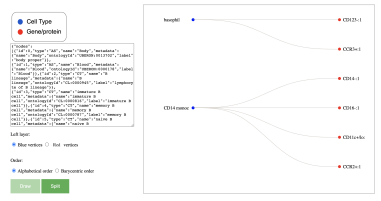


Figure 4 Showing the running time w.r.t. the number of edges. The vertices V_t are the cell types and the vertices V_b are the genes/proteins. The vertices are alphabetically ordered. This is corresponding to the exact algorithm of CRS(k) with Fixed Order. We can see as the number of edges increase the running time increases as expected.



(a) The input layout on the right side appears after inserting the dataset into the text area and clicking the draw button.



(b) The output layout appears on the right side after clicking the split

Figure 5 The user interface. Datasets of the human body [1] can be uploaded, processed with the presented algorithms, and visualized.

7.4 Evaluation Results

We first evaluate the exact algorithms for CRS(k) and CRSV(k) with Fixed Order. We have run experiments on 22 organ graphs on four settings: (1) the blue vertices (cell type) are fixed and the initial layout is generated using alphabetical order, (2) the red vertices (gene/protein) are fixed and the initial layout is generated using alphabetical order, (3) the blue vertices are fixed and initial layout is generated using barycentric heuristic, and (4) the red vertices are fixed and initial layout is generated using the barycentric heuristic. For each setting, we provide the initial number of crossings, number of vertices in the top (or left) layers that have fixed order, the number of bottom (or right) vertices, the number of splits, the number of split vertices, and the maximum number of splits; see Tables 2-5 in the supplemental material. The number of crossings in the initial layouts generated from alphabetical order is 2.7 times larger in total than in layouts generated by the barycentric heuristic. The number of splits is 1.58 times larger in total when we fix the gene/protein vertices (Tables 3 and 5). Note that for all organ graphs, the number of gene/protein vertices is relatively larger compared to the cell type vertices. When the cell-type vertices are fixed, there is more flexibility for splitting. Hence the numbers of splits in Tables 2 and 4 are smaller than in Tables 3 and 5. Similarly, the maximum number of splits is 2.5 times larger in total when the gene/protein vertices are fixed.

Organ	Crossings	117	$ V_b $	CR	S(k) with	Fixed O	rder	$\mathbf{CRSV}(k)$ with Fixed Order				
Organ	Crossings	Vt	V	Splits	Split ver- tices	Max splits	Time (ms)	Splits	Split ver- tices	Max splits	Time (ms)	
Bone Mar- row	111440	45	298	343	136	16	1.5	343	134	17	1.5	
Brain	28345	127	254	78	63	4	2.9	78	63	4	2.9	
Heart	504	15	45	6	6	1	0.3	6	6	1	0.4	
Kidney	13347	58	143	85	52	4	1.5	85	52	4	1.5	
Large intest- ine	4778	51	73	58	25	6	0.5	59	24	6	0.6	
Lung	11654	69	162	63	39	6	0.9	63	39	6	0.9	
Lymph nodes	59709	44	255	213	100	10	1.0	213	100	10	1.0	
Skin	2066	36	66	19	11	3	0.5	19	11	3	0.4	
Spleen	40565	65	225	165	75	10	1.3	166	75	11	1.4	
Thymus	102067	41	511	135	100	5	1.8	135	100	5	1.8	
Eye	17046	47	98	164	78	5	0.8	166	78	5	0.9	
Fallopian Tube	153	19	23	6	5	2	0.4	6	5	2	0.4	
Liver	625	26	47	9	8	2	0.5	9	8	2	0.5	
Pancreas	2510	29	40	56	32	6	0.5	56	32	6	0.6	
Peripheral Nervous System	0	1	2	0	0	0	0.1	0	0	0	0.1	
Prostate	405	12	31	3	3	1	0.3	3	3	1	0.3	
Ovary	8	3	6	0	0	0	0.1	0	0	0	0.1	
Small Intest- ine	28	5	13	0	0	0	0.1	0	0	0	0.1	
Ureter	512	14	30	21	19	2	0.3	21	19	2	0.4	
Urinary Bladder	628	15	31	21	21	1	1.1	21	21	1	1.1	
Uterus	1147	16	45	20	12	4	0.2	20	12	4	0.2	
Blood	49071	30	149	288	131	11	1.2	290	131	14	1.3	

Table 2 The vertices V_t are the cell types and the vertices V_b are the genes/proteins. The vertices are alphabetically ordered.

1:18 Splitting Vertices in 2-Layer Graph Drawings

Organ	Crossings	$ V_t $	1771	CR	S(k) with	Fixed O	rder	CRS	V(k) with) with Fixed O			
Organi	Crossings	$ V_t $	$ V_b $	Splits	Split ver- tices	Max splits	Time (ms)	Splits	Split ver- tices	Max splits	Time (ms)		
Bone Mar- row	111440	298	45	569	45	22	1.7	569	45	22	1.6		
Brain	28345	254	127	214	124	4	3.1	214	124	4	3.1		
Heart	504	45	15	30	14	4	0.3	30	14	4	0.3		
Kidney	13347	143	58	164	56	7	1.4	164	56	7	1.3		
Large in- testine	4778	73	51	76	32	6	0.5	76	32	6	0.4		
Lung	11654	162	69	151	64	4	1.1	151	64	4	1.1		
Lymph nodes	59709	255	44	378	42	26	1.0	379	42	27	0.9		
Skin	2066	66	36	48	32	2	0.6	48	32	2	0.6		
Spleen	40565	225	65	311	58	20	1.5	312	58	19	1.5		
Thymus	102067	511	41	514	36	69	2.1	514	36	69	2.1		
Eye	17046	98	47	172	29	40	0.9	179	29	37	0.9		
Fallopian Tube	153	23	19	12	8	2	0.5	12	8	2	0.5		
Liver	625	47	26	27	12	4	0.7	27	12	4	0.6		
Pancreas	2510	40	29	53	22	7	0.6	53	22	7	0.6		
Peripheral Nervous System	0	2	1	0	0	0	0.1	0	0	0	0.1		
Prostate	405	31	12	20	12	2	0.3	20	12	2	0.3		
Ovary	8	6	3	3	3	1	0.1	3	3	1	0.1		
Small Intestine	28	13	5	7	3	3	0.1	7	3	3	0.1		
Ureter	512	30	14	28	13	6	0.5	28	13	6	0.5		
Urinary Bladder	628	31	15	30	14	5	1.1	30	14	5	1.2		
Uterus	1147	45	16	44	16	6	0.2	44	16	6	0.2		
Blood	49071	149	30	379	30	40	1.4	379	30	40	1.4		

Table 3 The vertices V_t are the genes/proteins and the vertices V_b are the cell types. The vertices are alphabetically ordered.

Organ	Crossings	$ V_t $	1771	CR	S(k) with	Fixed O	rder	$\mathbf{CRSV}(k)$ with Fixed Order				
Organ	Crossings		$ V_b $	Splits	Split ver- tices	Max splits	Time (ms)	Splits	Split ver- tices	Max splits	Time (ms)	
Bone Marrow	32599	45	298	323	130	17	1.5	323	124	17	1.6	
Brain	4773	127	254	71	60	3	3.2	72	59	3	3.1	
Heart	211	15	45	6	6	1	0.4	6	6	1	0.3	
Kidney	1207	58	143	69	46	4	1.4	70	45	4	1.5	
Large in- testine	1878	51	73	49	25	6	0.7	50	24	7	0.6	
Lung	1970	69	162	64	37	7	1.3	64	37	7	1.2	
Lymph nodes	33315	44	255	214	104	10	1.3	216	104	10	1.2	
Skin	339	36	66	14	9	3	0.8	14	9	3	0.7	
Spleen	24833	65	225	171	76	9	1.7	172	75	9	1.8	
Thymus	34863	41	511	136	102	5	2.3	137	101	5	2.4	
Eye	8576	47	98	154	73	5	1.1	155	72	5	1.2	
Fallopian Tube	47	19	23	7	7	1	0.7	7	7	1	0.6	
Liver	84	26	47	8	7	2	0.9	8	7	2	0.9	
Pancreas	925	29	40	46	29	5	0.8	46	29	5	0.7	
Peripheral Nervous System	0	1	2	0	0	0	0.1	0	0	0	0.1	
Prostate	64	12	31	3	3	1	0.4	3	3	1	0.4	
Ovary	0	3	6	0	0	0	0.1	0	0	0	0.1	
Small Intestine	0	5	13	0	0	0	0.1	0	0	0	0.1	
Ureter	126	14	30	18	18	1	0.6	18	18	1	0.7	
Urinary Bladder	151	15	31	18	18	1	1.3	18	18	1	1.3	
Uterus	135	16	45	13	10	3	0.3	14	10	3	0.3	
Blood	19308	30	149	284	130	13	1.7	284	130	13	1.6	

Table 4 The vertices V_t are the cell types and the vertices V_b are the genes/proteins. The vertices are ordered by repeatedly applying the barycentric heuristic on both sides.

1:20 Splitting Vertices in 2-Layer Graph Drawings

Organ	Crossings	$ V_t $	$ V_b $	CR	S(k) with	Fixed O	rder	CRS	$\overline{\mathbf{V}(k)}$ with	Fixed C	rder
organi	Crossings	$ V_t $		Splits	Split ver- tices	Max splits	Time (ms)	Splits	Split ver- tices	Max splits	Time (ms)
Bone Marrow	32599	298	45	412	45	15	1.8	412	45	14	1.7
Brain	4773	254	127	86	78	3	3.2	86	70	3	3.2
Heart	211	45	15	16	11	3	0.4	16	11	3	0.4
Kidney	1207	143	58	94	49	5	1.5	94	47	4	1.5
Large in- testine	1878	73	51	47	24	5	0.8	50	24	5	0.7
Lung	1970	162	69	74	49	3	1.3	74	47	3	1.3
Lymph nodes	33315	255	44	298	41	24	1.4	299	41	23	1.3
Skin	339	66	36	22	19	2	0.8	22	19	2	0.8
Spleen	24833	225	65	223	59	14	1.8	227	59	15	1.8
Thymus	34863	511	41	320	35	46	2.4	320	34	48	2.5
Eye	8576	98	47	146	25	34	1.1	149	25	35	1.1
Fallopian Tube	47	23	19	5	5	1	0.6	5	5	1	0.6
Liver	84	47	26	3	7	2	1.1	3	7	2	1.0
Pancreas	925	40	29	42	21	6	0.8	42	20	6	0.8
Peripheral Nervous System	0	2	1	0	0	0	0.1	0	0	0	0.1
Prostate	64	31	12	13	12	2	0.5	13	12	2	0.4
Ovary	0	6	3	0	0	0	0.1	0	0	0	0.1
Small Intestine	0	13	5	3	3	1	0.1	3	3	1	0.1
Ureter	126	30	14	18	13	4	0.7	18	9	4	0.7
Urinary Bladder	151	31	15	19	14	4	1.4	19	14	4	1.3
Uterus	135	45	16	23	13	4	0.3	25	13	4	0.4
Blood	19308	149	30	300	30	31	1.8	300	30	31	1.7

Table 5 The vertices V_t are the genes/proteins and the vertices V_b are the cell types. The vertices are ordered by repeatedly applying the barycentric heuristic on both sides.

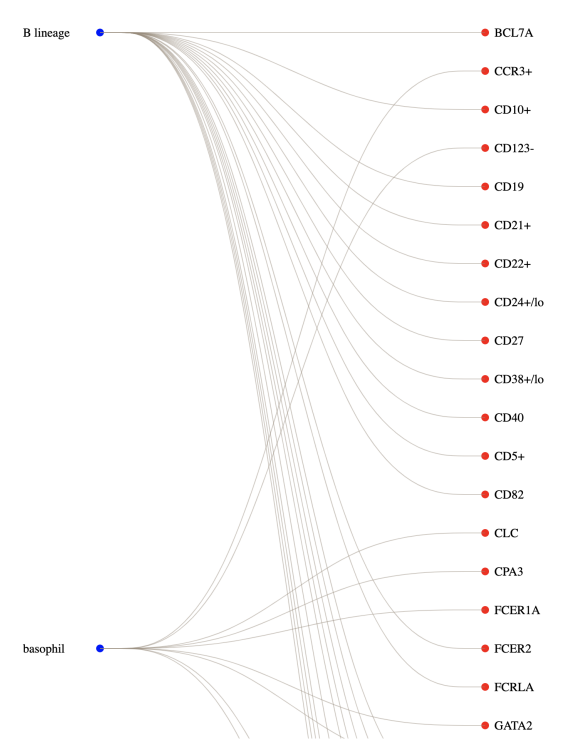


Figure 6 Part of the input layout of the 2-layer graph corresponding to the anatomical structure of blood computed by the exact algorithm for CRS(k) with Fixed Order. The graph is large, hence we show a part of the layout.

1:22 Splitting Vertices in 2-Layer Graph Drawings

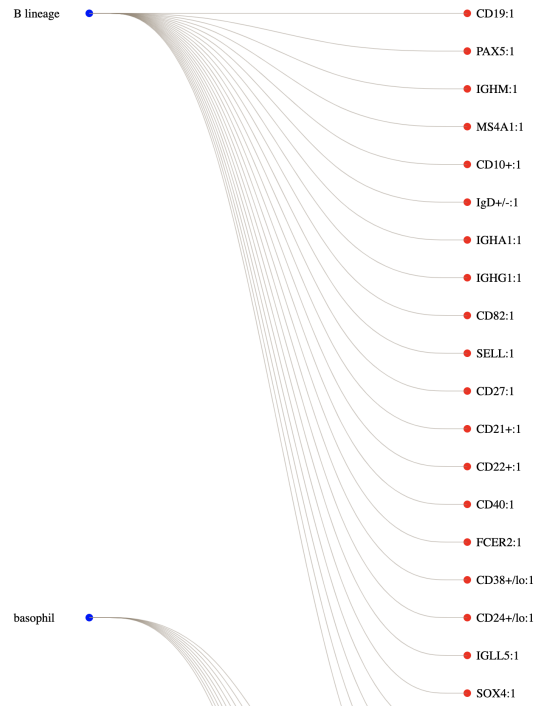


Figure 7 Part of the output layout of the 2-layer graph corresponding to the anatomical structure of blood computed by the exact algorithm for CRS(k) with Fixed Order.

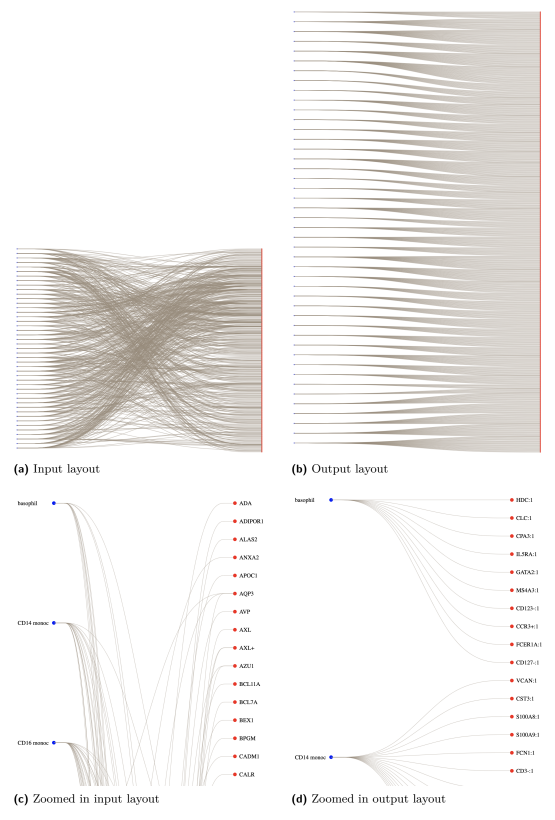


Figure 8 Input and output layouts of the exact algorithm of CRS(k) with Fixed Order for Bone Marrow.

1:24 Splitting Vertices in 2-Layer Graph Drawings

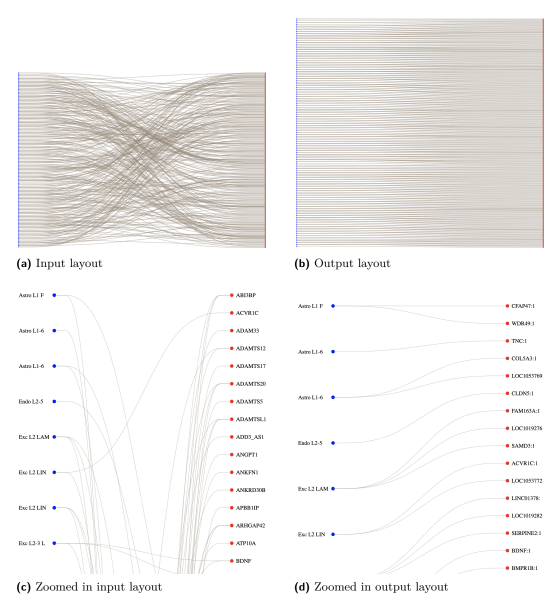


Figure 9 Input and output layouts of the exact algorithm of CRS(k) with Fixed Order for Brain.

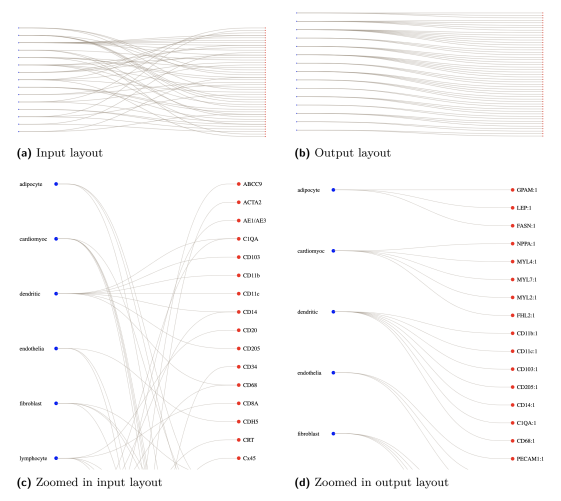
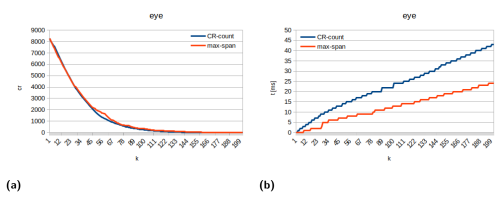


Figure 10 Input and output layouts of the exact algorithm of CRS(k) with Fixed Order for Heart.

A second set of experiments was conducted on the same 22 organ graphs to evaluate the crossing minimization heuristics. We set the maximum budget k of splits to 200 and computed the number of remaining crossings after each split. Additionally, we measured wall clock time after each iteration. Figure 11 shows the number of crossings in regards to k for one example graphs. Examples of other graphs, as well as runtime plots can be found in the supplemental material; see Fig. 6–11. For both algorithms we observed a similar performance regarding crossing reduction. In some cases one algorithm slightly outperformed the other, but no clear trend is visible in the data. Intuitively, it seems that in the case of max-span the length of edges correlates with the number of crossings. Furthermore, the number of crossings declines steeply at the beginning and for some graphs nearly 30% of crossings are removed by the first 10 splits.

The runtime experiments confirmed the asymptotic runtime analysis. max-span outperforms CR-count on every datasets in regards to total runtime and scalability.



■ **Figure 11** max-span and CR-count heuristic applied to the eye dataset. Both algorithms have nearly identical performance regarding crossing reduction as seen in (a). However, max-span outperforms CR-count regarding runtime (b).

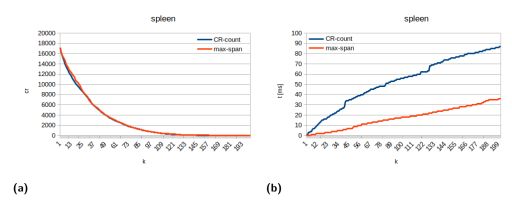


Figure 12 max-span and CR-count heuristic applied to the spleen dataset. Both algorithms have nearly identical performance regarding crossing reduction as seen in (a). However, max-span outperforms CR-count regarding runtime (b).

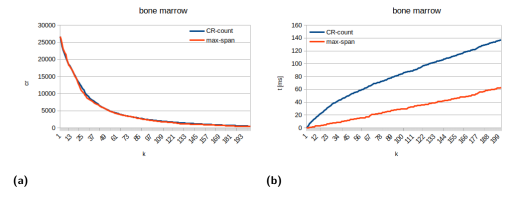


Figure 13 max-span and CR-count heuristic applied to the bone marrow dataset. Both algorithms have nearly identical performance regarding crossing reduction as seen in (a). However, max-span outperforms CR-count regarding runtime (b).

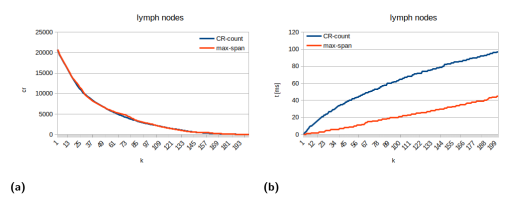


Figure 14 max-span and CR-count heuristic applied to the lymph nodes dataset. Both algorithms have nearly identical performance regarding crossing reduction as seen in (a). However, max-span outperforms CR-count regarding runtime (b).

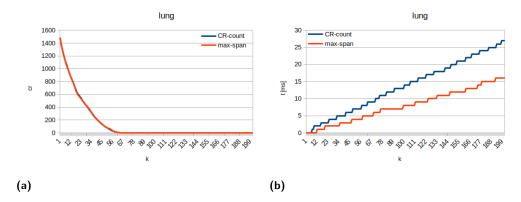
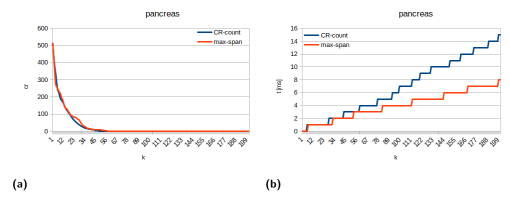


Figure 15 max-span and CR-count heuristic applied to the lung nodes dataset. Both algorithms have nearly identical performance regarding crossing reduction as seen in (a). However, max-span outperforms CR-count regarding runtime (b).



■ Figure 16 max-span and CR-count heuristic applied to the pancrease nodes dataset. Both algorithms have nearly identical performance regarding crossing reduction as seen in (a). However, max-span outperforms CR-count regarding runtime (b).

All codes for user interface, algorithms, experimental data, and analysis are available on

Github at https://github.com/abureyanahmed/split_graphs.

8 Open Problems

Minimizing the total number of splits, or the number of split vertices are natural problems. Other variants include minimizing the maximum number of splits per vertex and considering the case where splits are allowed in both layers. Vertex splits can also be used to improve other quality measures of a 2-layer layout (besides crossings). When visualizing large bipartite graphs, a natural goal is to arrange the vertices so that a small window can capture all the neighbors of a given node, i.e., minimize the maximum distance between the first and last neighbors of a top vertex in the order of the bottom vertices.

Since a great deal of vertex splitting can dramatically change the structure of the graph, it is desirable to have a bound on the number of splits. Hence, in our problems, we take a parameter k that limits the number of splits. One can also consider the problem of finding the minimum number of splits to generate a planar graph. This is known as the Planar Split Thickness of Graphs [10]. It remains an interesting future research direction to find the trade-off between the number of splits and the number of crossings as the graph size changes.

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