From Boston to San Francisco: A Survey of Shortest Paths Algorithms in Planar Graphs

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Abstract

- 1 Introduction
- 2 Background
- 3 Single source shortest paths

We call an edge uv relaxed if $d(v) \leq d(u) + c(u, v)$. We call the assignment

$$d(v) \leftarrow \min\{d(v), d(u) + c(u, v)\}$$

the relaxation of vertex v. We know that the labels give a correct shortest path distances if the shortest-path conditions are satisfied:

- d(s) = 0,
- ullet every label d(v) is an upper bound on the s-v distance,
- $\bullet\,$ every edge is relaxed.

3.1 Nonnegative edge weights

3.1.1 Simple algorithm

For a planar graph with nonnegative edge weights, Dijkstra's algorithm runs in $O(n \log n)$ as $m \leq 3n - 6$. It is possible to improve this to O(n). To get there, recall that and r-division of a planar graph is a partition of the graph into $\Theta(n/r)$ regions of size O(r) with boundary size $O(\sqrt{r})$. An r-division of a planar graph can be computed in linear time.

A simple $O(n\sqrt{\log n \log \log n})$ emerges quite beautifully just from the r-division if we set $r = \frac{\log n}{\log \log n}$. The algorithm follows a divide-and-conquer approach in which each region is processed first, followed by a clean-up phase where the results are merged.

Algorithm 1 Shortest paths in each region R

```
for all Regions R do

for all Boundary nodes v \in R do

Compute SSSP from v in R

Store (u, v) distances for any two boundary nodes u, v

end for

end for
```

The first step is to compute the single-source shortest paths for each boundary node in each region R (algorithm 1). We can now replace each region R by a complete graph on R's boundary nodes with shortest paths distances between any two nodes. Call this auxiliary graph G'. The second phase of the algorithm is to compute the SSSP from s in G'. This gives the true shortest paths from s to all the boundary nodes. Finally, we must tidy up by finding the distances from s to the nodes inside each region (algorithm 2).

Algorithm 2 Clean up: shortest paths from s to inside of each region R

```
for all Regions R do

for all Boundary nodes v \in R do

Set d(v) = d_{G'}(s, v)

Compute SSSP from v in R

end for

end for
```

To analyze the algorithm, we will need a few pieces of information:

- Total number of boundary nodes is $O(\sqrt{r})O(n/r) = O(n/\sqrt{r})$.
- Number of nodes in G' is $O(n/r)O(\sqrt{r}) = O(n/\sqrt{r})$.
- Number of edges in G' is O(n/r)O(r) = O(n).

Using $r = \frac{\log n}{\log \log n}$, the first phase is bounded above by

$$O\left(n\frac{\sqrt{\log\log n}}{\sqrt{\log n}}\log n\right) = O(n\sqrt{\log n\log\log n}),$$

the second phase is an SSSP in a size $O(n\frac{\sqrt{\log\log n}}{\sqrt{\log n}})$ graph, and thus also $O(n\sqrt{\log n\log\log n})$, while the tidying up is a series of SSSPs in each of the regions, and has the same bound as the first phase— $O(n\sqrt{\log n\log\log n})$. The total time bound ends up $O(n\sqrt{\log n\log\log n})$.

3.1.2 Recursion

The key idea that can improve the running time to linear is to go deeper inside the recursive world. Instead of just using an r-division of the graph, we recursively subdivide each region until we reach edges which are atomic regions.

The simple algorithm shown above is an improvement over Dijkstra's algorithm, but one can do better. In fact, we can achieve linear time by recursively subdividing the graph. Without loss of generality, assume the graph is directed, and that each node has at most two incoming and two outgoing edges. We call a region atomic if it contains only one edge uv. A nonatomic region will have as children subregions that are contained within it.

For each region R, we maintain a priority queue Q(R) that stores the subregions of R if R is nonatomic, or the single arc uv is R is atomic. The algorithm ensures that for every region R, the minimum element of Q(R) is the minimum label d(v) over all edges vw in R that remain to be processed.

Algorithm 3 Process region

```
procedure PROCESS

if R contains only uv then

if d(v) > d(u) + c(u, v) then

d(v) \leftarrow d(u) + c(u, v)
for each outgoing edge vw of v, call \text{UPDATE}(R(vw), vw, d(v))
end if
Q(R).updateKey(uv, \infty)
else
repeat
R' \leftarrow Q(R).getMin()
PROCESS(()R')
Q(R).updateKey(R', Q(R').getMinKey())
until Q(R).getMinKey() is infinity or if repeated \alpha_{h(R)} times end if
end procedure
```

Algorithm 4 Update region

```
procedure UPDATE(R,x,k)
Q(R).updateKey(x,k)
if updateKey reduced the value of Q(R).getMinKey() then
UPDATE(()parent(R),R,k)
end if
end procedure
```

Algorithm 5 Faster SSSP

```
Find recursive subdivision R(G), R(P_i), \dots, R(uv)

Allocate queue Q for each region d(v) \leftarrow \infty, \forall v

d(s) \leftarrow 0

for all sv \in E(G) do

UPDATE(R(sv), sv, 0)

end for

while Q(R(G)).getMinKey() < \infty do

PROCESS(R(G))

end while
```

The procedures are used in algorithm 3.1.2.

By playing around with the number of levels, number of nodes per region per level, and the α_i , we can achieve linear time. To give an intuition, we present here a $O(n \log \log n)$ algorithm and briefly comment on how to change it to get rid of the $\log \log n$ factor.

3.2 Arbitrary edge weights

4 Multiple source shortest paths

5 Extensions to higher genus