In this lecture, we will discuss the question of *graph reachability*: given two vertices $v$ and $w$, does there exist a path from $v$ to $w$?

**Additional Resources**

- Review slides (https://cs.cmu.edu/~15122/handouts/slides/review/24-dfs.pdf)
- Code for this lecture (https://cs.cmu.edu/~15122/handouts/code/24-dfs.tgz)

This maps as follows onto the learning goals for this course:

**Computational Thinking:** We continue learning about graphs, and specifically about paths in a graph. An important question is whether there exists a path between two given nodes. A related problem is to produce this path (if it exists).

**Algorithms and Data Structures:** We explore two classic approaches to answering these questions: depth-first search and breadth-first search. Both rely on the need to remember what we have done already, and to go back and try something else if we get stuck.

**Programming:** We give two implementations of depth-first search, one recursive that uses the call stack of C to remember what we have done, and the other iterative that uses an explicit stack for that purpose. We also see that breadth-first search is the variant of the latter where a queue is used instead of a stack.

As a reminder, we are working with the following minimal graph interface. We will be implementing our search in terms of this interface.
typedef unsigned int vertex;
typedef struct graph_header *graph_t;

graph_t graph_new(unsigned int numvert);
//@
ensures \result != NULL;

void graph_free(graph_t G);
//@
requires G != NULL;

unsigned int graph_size(graph_t G);
//@
requires G != NULL;
//@
requires v < graph_size(G) && w < graph_size(G);

bool graph_hasedge(graph_t G, vertex v, vertex w);
//@
requires G != NULL;
//@
requires v < graph_size(G) && w < graph_size(G);
//@
requires v != w && !graph_hasedge(G, v, w);

typedef struct neighbor_header *neighbors_t;

neighbors_t graph_get_neighbors(graph_t G, vertex v);
//@
requires G != NULL && v < graph_size(G);
//@
ensures \result != NULL;

bool graph_hasmore_neighbors(neighbors_t nbors);
//@
requires nbors != NULL;

vertex graph_next_neighbor(neighbors_t nbors);
//@
requires nbors != NULL;
//@
requires graph_hasmore_neighbors(nbors);

void graph_free_neighbors(neighbors_t nbors);
//@
requires nbors != NULL;
1 Paths in Graphs

A path in a graph is a sequence of vertices where each vertex is connected to the next by an edge. That is, a path is a sequence

$$v_0, v_1, v_2, v_3, \ldots, v_l$$

of some length $l \geq 0$ such that there is an edge from $v_i$ to $v_{i+1}$ in the graph for each $i < l$.

For example, all of the following are paths in the graph above:

$$A - B - E - C - D$$
$$A - B - A$$
$$E - C - D - C - B$$
$$B$$

The last one is a special case: The length of a path is given by the number of edges in it, so a node by itself is a path of length 0 (without following any edges). Paths always have a starting vertex and an ending vertex, which coincide in a path of length 0. We also say that a path connects its endpoints.

The graph reachability problem is to determine if there is a path connecting two given vertices in a graph. If we know the graph is connected, this problem is easy since one can reach any node from any other node. But we might refine our specification to request that the algorithm return not just a boolean value (reachable or not), but an actual path. At that point the problem is somewhat interesting even for connected graphs. In complexity theory it is sometimes said that a path from vertex $v$ to vertex $w$ is a certificate or explicit evidence for the fact that vertex $w$ is reachable from another vertex $v$. It is easy to check whether the certificate is valid, since it is easy to check if each node in the path is connected to the next one by an edge. It is more difficult to produce such a certificate.

For example, the path

$$A - B - E - C - D$$
is a certificate for the fact that vertex $D$ is reachable from vertex $A$ in the above graph. It is easy to check this certificate by following along the path and checking whether the indicated edges are in the graph.

In most of what follows we are not concerned with finding the path, but only with determining whether one exists. It is not difficult to see how to extend the algorithms we discuss to compute the path as well.

## 2 Depth-First Search

The first algorithm we consider for determining if one vertex is reachable from another is called depth-first search.

Let’s try to work our way up to this algorithm. Assume we are trying to find a path from $u$ to $w$. We start at $u$. If it is equal to $w$ we are done, because $w$ is reachable by a path of length 0. If not we pick an arbitrary edge leaving $u$ to get us to some node $v$. Now we have “reduced” the original problem to the one of finding a path from $v$ to $w$.

The problem here is of course that we may never arrive at $w$ even if there is a path. For example, say we want to find a path from $A$ to $D$ in our earlier example graph.

We can go $A \rightarrow B \rightarrow E \rightarrow A \rightarrow B \rightarrow E \rightarrow \cdots$ without ever reaching $D$ (or we can go just $A \rightarrow B \rightarrow A \rightarrow B \rightarrow \cdots$), even though there exists a path.

We need to avoid repeating nodes in the path we are exploring. A cycle is a path of length 1 or greater that has the same starting and ending point. So another way to say we need to avoid repeating nodes is to say that we need to avoid cycles in the path. We accomplish this by marking the nodes we have already considered so when we see them again we know not to consider them again.

Let’s go back to the earlier example and play through this idea while trying to find a path from $A$ to $D$. We start by marking $A$ (indicated by hollowing the circle) and go to $B$. We indicate the path we have been following by drawing a double-line along the edges contained in it.
When we are at B we mark B and have three choices for the next step.

1. We could go back to A, but A is already marked and therefore ruled out.

2. We could go to E.

3. We could go to C.

Say we pick E. At this point we have again three choices. We might consider A as a next node on the path, but it is ruled out because A has already been marked. We show this by dashing the edge from A to E to indicate it was considered, but ineligible. The only possibility now is to go to C, because we have been at B as well (we just came from B).

From C we consider the link to D (before considering the link to B) and we arrive at D, declaring success with the path

\[ A - B - E - C - D \]

which, by construction, has no cycles.

There is one more consideration to make, namely what we do when we get stuck. Let’s reconsider the original graph
and the goal to find a path from $E$ to $B$. Let’s say we start $E \rightarrow C$ and then $C \rightarrow D$. At this point, all the vertices we could go to (which is only $C$) have already been marked! So we have to backtrack to the most recent choice point and pursue alternatives. In this case, this could be $C$, where the only remaining alternative would be $B$, completing the path $E \rightarrow C \rightarrow B$. Notice that when backtracking we have to go back to $C$ even though it is already marked.

Depth-first search is characterized not only by the marking, but also that when we get stuck we always return to our most recent choice and follow a different path. When no other alternatives are available, we backtrack further. Let’s consider the following slightly larger graph, where we explore the outgoing edges using the alphabetically last label first. We will trace the search for a path from $A$ to $B$.

We write the current node we are visiting on the left and on the right a stack of nodes we have to return to when we backtrack. For each of these we also remember which choices remain (in parentheses). We annotate marked nodes with an asterisk, which means that we never pick them as the next node to visit.

For example, going from step 4 to step 5 we do not consider $E^*$ but go to $D$ instead. We backtrack when no unmarked neighbors remain for the current node. We are keeping the visited nodes on a stack so we can easily return to the most recent one. The stack elements are separated by | and the lists of neighbors are wrapped in parentheses, e.g., $(B, A^*)$.

<table>
<thead>
<tr>
<th>Step</th>
<th>Current</th>
<th>Stack</th>
<th>Remark</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$A$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>$E$</td>
<td>$A^* (B)$</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>$C$</td>
<td>$E^* (B, A^*)</td>
<td>A^* (B)$</td>
</tr>
<tr>
<td>4</td>
<td>$G$</td>
<td>$C^* (E^*, D)</td>
<td>E^* (B, A^*)</td>
</tr>
<tr>
<td>5</td>
<td>$D$</td>
<td>$C^* ()</td>
<td>E^* (B, A^*)</td>
</tr>
<tr>
<td>6</td>
<td>$F$</td>
<td>$D^* (C^*)</td>
<td>C^* ()</td>
</tr>
<tr>
<td>7</td>
<td>$B$</td>
<td>$E^* (A^*)</td>
<td>A^* (B)</td>
</tr>
</tbody>
</table>
2.1 Recursive Depth-First Search

Now we can easily write the depth-first search code recursively, letting the call stack keep track of everything we need for backtracking.

```c
bool dfs_helper(graph_t G, bool *mark, vertex start, vertex target) {
    REQUIRES(G != NULL && mark != NULL);
    REQUIRES(start < graph_size(G) && target < graph_size(G));
    REQUIRES(!mark[start]);

    // mark start as seen
    mark[start] = true;

    // there is an edge from start to v and a path from v to target if...
    // target == start, or
    if (target == start) return true;
    // there is an edge from start to v ...
    neighbors_t nbors = graph_get_neighbors(G, start);
    while (graph_hasmore_neighbors(nbors)) {
        vertex v = graph_next_neighbor(nbors); // v is one of start’s neighbors
        // ... and a path from v to target
        if (!mark[v] && dfs_helper(G, mark, v, target)) {
            graph_free_neighbors(nbors);
            return true;
        }
    }
    graph_free_neighbors(nbors);
    return false;
}
```

We shall free the neighbor list before each return, or risk a memory leak.

We’ve named the function dfs_helper because the user of the search should not have to worry about supplying the array of marks. Instead the user calls the function dfs, below, which creates the marks and passes them to the recursive helper function.

```c
bool dfs(graph_t G, vertex start, vertex target) {
    REQUIRES(G != NULL);
    REQUIRES(start < graph_size(G) && target < graph_size(G));
    bool *mark = xmalloc(graph_size(G), sizeof(bool));
    bool connected = dfs_helper(G, mark, start, target);
    free(mark);
    return connected;
}
```
What is the cost of recursive DFS for a graph with \( v \) vertices and \( e \) edges?

The function `dfs` creates an array of marks with all positions initialized to zero (i.e., to `false`) on line ?? This takes \( O(v) \) time. The call to `free` is constant time. Therefore, the asymptotic complexity of `dfs` is equal to the cost of the call to `dfs_helper` on line ??.

The analysis of `dfs_helper` bears similarities to that of `graph_print` in the last chapter, with the novelty that `dfs_helper` is recursive rather than iterative. The call to `graph_get_neighbors` on line ?? has constant cost in the adjacency list representation and costs \( O(v) \) in the adjacency matrix representation. The body of the loop on lines ??–?? runs \( O(e) \) times overall since every edge will be visited exactly twice altogether (once from each direction). In particular, line ?? is executed \( O(e) \) times altogether.

This entails that there will be at most \( 2e \) recursive calls to `dfs_helper`. Furthermore, lines ?? and ??, and the fact that marks are never reset, ensures that this function will be called no more than \( v \) times. Thus, there can be at most \( \min(2e, v) \) recursive calls to `dfs_helper`.

Each call to `graph_free_neighbors` costs \( O(1) \) in the adjacency list representation while they altogether cost \( O(e) \) in the adjacency matrix representation. All other operations in `dfs_helper` have constant cost, and therefore \( O(e) \) overall since they are executed at most \( 2e \) times.

Tallying up all these components, we have an \( O(e) \) worst case complexity for the call to `dfs_helper` on line ?? with an adjacency list representation. This gives `dfs` a worst-case complexity of \( O(v + e) \) using this representation.

The cost of `dfs` is \( O(\min(v^2, ev)) \) with the adjacency matrix representation — the latter is often simplified to \( O(v^2) \) since most graphs encountered in realistic applications have at least \( O(v) \) edges.

### 2.2 Depth-First Search with an explicit stack

When scrutinizing the above example, we notice that the sophisticated data structure of a stack of nodes with their adjacency lists was really quite unnecessary for DFS. The recursive implementation is simple and elegant, but its effect is to make the data management more complex than necessary: all we really need for backtracking is a stack of nodes that have been seen but not yet considered.

This can all be simplified by making the stack explicit. In that case there is a single stack with all the nodes on it that we still need to look at. (In the sample code in Figure ??, we use a stack specialized to hold things of type
vertex just to keep the code simple.)

<table>
<thead>
<tr>
<th>Step</th>
<th>Current</th>
<th>Neighbors</th>
<th>New stack</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td></td>
<td></td>
<td>(A*)</td>
</tr>
<tr>
<td>1</td>
<td>A*</td>
<td>(E, B)</td>
<td>(E*, B*)</td>
</tr>
<tr>
<td>2</td>
<td>E*</td>
<td>(C, B*, A*)</td>
<td>(C*, B*)</td>
</tr>
<tr>
<td>3</td>
<td>C*</td>
<td>(G, E*, D)</td>
<td>(G*, D*, B*)</td>
</tr>
<tr>
<td>4</td>
<td>G*</td>
<td>(C*)</td>
<td>(D*, B*)</td>
</tr>
<tr>
<td>5</td>
<td>D*</td>
<td>(F, C*)</td>
<td>(F*, B*)</td>
</tr>
<tr>
<td>6</td>
<td>F*</td>
<td>(D*)</td>
<td>(B*)</td>
</tr>
<tr>
<td>7</td>
<td>B*</td>
<td>(E*, A*)</td>
<td>()</td>
</tr>
</tbody>
</table>

```c
bool dfs_explicit_stack(graph_t G, vertex start, vertex target) {
    REQUIRES(G != NULL);
    REQUIRES(start < graph_size(G) && target < graph_size(G));
    if (start == target) return true;

    // mark is an array containing only the start
    bool *mark = xcalloc(graph_size(G), sizeof(bool));
    mark[start] = true;

    // Work list initially containing only start
    stack_t S = stack_new();
    push(S, start);

    while (!stack_empty(S)) {
        // Loop invariants to prove correctness go here
        vertex v = pop(S);  // v is the current node
        if (v == target) {
            // if v is the target return true
            stack_free(S);
            free(mark);
            return true;  // Success!
        }

        neighbors_t nbors = graph_get_neighbors(G, v);
        while (graph_hasmore_neighbors(nbors)) {
            vertex w = graph_next_neighbor(nbors);  // w is one of v's neighbors
            if (!mark[w]) {  // if w was not seen before
                mark[w] = true;  // Mark it as known
                push(S, w);  // Add to work list
            }
        }
    }

    free(mark);
    stack_free(S);
    free(S);
    return false;
}
```
In the code in Figure ??, we mark the starting node and push it on the stack. Then we iteratively pop the stack, check if it is our target, and otherwise examine each neighbor of the node we popped. If the neighbor is not already marked, we push it on the stack to make sure we look at it eventually. If the stack is empty then we’ve explored all possibilities without finding the target, so we return false.

While convincing, this explanation comes short of a proof that our implementation is correct, i.e., that it returns true when there is a path between start and target and returns false otherwise. We will now develop a more solid argument, although we will stop short of a formal proof. The function dfs_explict_stack returns in exactly three places. The first is on line ??, when start is equal to target. By definition, there is a degenerate path between these two nodes in this case.

The other two places where the function returns, lines ?? and ??, have us go through loops. To reason about loops, we need to develop loop invariants that we will squeeze in the placeholder on line ??: we know that w (which is equal to target by line ??) is a neighbor of vertex v, but how do we know that there is a path from start to v? Two invariants will prove helpful here:

1. Every marked vertex (i.e., a vertex u such that mark[u] == true) is connected to start.

2. Every vertex in the stack is marked.

These two invariants hold initially since start is the only marked vertex and the only item in the stack before the loop is run the first time. They are preserved by an arbitrary iteration of the loop since only neighbors of v (which is assumed to be reachable from start) are marked on line ?? and they are immediately pushed on the stack on line ???. Therefore, if the loop exits at line ??, we know that there is a path from start to target.

These two invariants are not sufficient to prove that there is no path from start to target if the function returns false on line ???. For this, we need a new concept and a new invariant involving it. The new concept is that of frontier of the search. The frontier is a set of vertices that we know are connected to start but that we have not explored yet. At any point in
the loop on lines ??–??, the frontier is the contents of the stack. The new invariant is the following:

3 Every path from start to target passes through a vertex in the frontier.

It is clearly true initially when the stack (the frontier) only contains start. It is preserved by the loop because intuitively a frontier element is replaced by all of its neighbors that have not been explored already. More interesting is why this invariant allows us to prove that there is no path to target if the function returns on line ??: for this to happen, we must have exited the loop in lines ??–??, which entails that the negation of its loop guard is true: the stack (our frontier) is empty. By our third invariant (which still holds at this point), every path from start to target must go through a vertex in the frontier. But the frontier is empty, so it contains no vertex through which such a pass can go: thus, there cannot be any path from start to target. Only in this way can the invariant be true, if the frontier is empty: if all of zero paths from start to target pass through one of the (zero) vertices in the empty frontier.

The complexity considerations we developed for the recursive version of DFS apply here as well — possibly more explicitly. The above code has cost $O(v + e)$ with an adjacency list representation: initializing the array of marks has cost $O(v)$, and the body of the inner loop will run $O(e)$ times, twice for each edge. The cost is $O(\min(v^2, ev)) = O(v^2)$ in most situations — with an adjacency matrix representation as we have to account for the $O(v)$ cost of graph_get_neighbors.

3 Breadth-First Search

The iterative DFS algorithm managed its agenda, i.e., the list of nodes it still had to look at, using a stack. But there’s no reason to insist on a stack for that purpose. What happens if we replace the stack by a queue? All of a sudden, we will no longer explore the most recently found neighbor first as in depth-first search, but, instead, we will look at the oldest neighbor first. This corresponds to a breadth-first search (BFS) where you explore the graph layer by layer. So BFS completes a layer of the graph before proceeding to the next layer. The code for that and many other interesting variations of graph search can be found on the course web page.

Here’s an illustration using our running example of search for a path from A to B in the graph
We find the path much faster this way. But this is just one example. Try to think of another search in the same graph that would cause breadth-first search to examine more nodes than depth-first search would.

The code looks the same as our iterative depth-first search, except for the use of a queue instead of a stack. Therefore we do not include it here. You could write it yourself, and if you have difficulty, you can find it in the code folder that goes with this lecture. Note that our correctness and complexity analysis for DFS never relied on using a stack. Thus, it remains sound once we swap the stack for a queue. Correctness also hold for any other implementation of work list, but complexity may need to be revisited if these implementation cannot provide constant-time insertion and retrieval operations.

4 Conclusion

Breadth-first and depth-first search are the basis for many interesting algorithms as well as search techniques for artificial intelligence.

One potentially important observation about breadth-first versus depth-first search concerns search when the graph remains implicit, for instance in game search. In this case there might be infinite paths in the graph. Once embarked on such a path depth-first search will never backtrack, but will pursue the path endlessly. Breadth-first search, on the other hand, since it searches layer by layer, is not subject to this weakness (every node in a graph is limited to a finite number of neighbors). In order to get some benefits of both techniques, a technique called iterative deepening is sometimes used.
5 Exercises

Exercise 1 (sample solution on page ??). In the following graph, we are searching for vertex G starting from vertex A. Assuming neighbors are visited in alphabetical order, in what order will BFS examine the vertices of this graph in order to find G? What about DFS?

Exercise 2 (sample solution on page ??). Using BFS as inspiration, write a function that takes in a vertex v, and returns an array such that \( arr[u] \) is the shortest distance from v to u, or -1 if they are not connected. Your function should have the following type

\[
\text{int \ *} \text{shortest_distances}(\text{graph_t G, vertex v})
\]

Exercise 3 (sample solution on page ??). Come up with a graph, start, and target for which BFS explores more vertices than DFS, assuming that we visit neighbors in alphabetical order. Then, do the same where DFS explores more vertices than BFS.

Exercise 4 (sample solution on page ??). What is the complexity of the call \( \text{ tester}(G) \) for a graph G with \( v \) vertices and \( e \) edges. Give the complexity for both the adjacency list and the adjacency matrix representations of graphs.

```c
int mystery(graph_t G, bool* mark, vertex start, vertex target) {
    REQUIRES(G != NULL && mark != NULL);
    REQUIRES(start < graph_size(G));
    REQUIRES (target >= graph_size(G));
    mark[start] = true;
    queue_t Q = queue_new();
    enq(Q, start);
    // More code here...
}
```
int acc = 0;
while (!queue_empty(Q)) {
    vertex v = deq(Q);
    if (v == target) {
        queue_free(Q);
        free(mark);
        return -1;
    }
    neighbors_t nbors = graph_get_neighbors(G, v);
    while (graph_hasmore_neighbors(nbors)) {
        vertex w = graph_next_neighbor(nbors);
        if (!mark[w]) {
            mark[w] = true;
            enq(Q, w);
        }
    }
    graph_free_neighbors(nbors);
    for (unsigned int i = 0; i < graph_size(G); i++) {
        if (!mark[i])
            acc++;
    }
    queue_free(Q);
    return acc;
}

int tester(graph_t G) {
    bool *mark = xmalloc(sizeof(bool), graph_size(G));
    int x = mystery(G, mark, 0, graph_size(G)+1);
    free(mark);
    return x;
}
Sample Solutions

**Solution of exercise ??** For BFS, the visit order is:

\[A, B, C, D, E, F, G\]

For DFS, it is:

\[A, B, D, E, G\]

**Solution of exercise ??** We just need modify BFS to keep track of the distance from the starting vertex as we traverse the graph. Since BFS already explores paths from shortest to longest, we know that when we visit a vertex we have found the shortest possible path to get to it. Therefore, we can simply replace the `mark` array (an array of booleans) with an array of integers (`dist`) where `dist[v]` is either the distance of `v` from `start` or `-1` to indicate that `v` has not been reached yet.
```c
int *shortest_distances(graph_t G, vertex start) {
    REQUIRES(G != NULL);
    REQUIRES(start < graph_size(G));

    int *dist = xmalloc(sizeof(int) * graph_size(G));
    for (unsigned int i = 0; i < graph_size(G); i++) {
        dist[i] = -1; // -1 means unvisited
    }
    dist[start] = 0; // start is 0 edges away from start

    queue_t Q = queue_new();
    enq(Q, start);

    while (!queue_empty(Q)) {
        vertex v = deq(Q); // v is the current node
        ASSERT(dist[v] != -1); // v has distance dv from start
        neighbors_t nbors = graph_get_neighbors(G, v);
        while (graph_hasmore_neighbors(nbors)) {
            vertex w = graph_next_neighbor(nbors); // w is one of v’s neighbors
            if (dist[w] == -1) { // if w has not been reached yet
                dist[w] = dist[v] + 1; // its distance from start is dv+1
                enq(Q, w); // Enqueue w onto the queue
            }
        }
        graph_free_neighbors(nbors);
    }
    queue_free(Q);
    return dist;
}
```

**Solution of exercise**

There are many possible solutions to this task. The general idea is that on graphs with many long paths, DFS can get lucky and find the target at the end of one of them quickly. On the other hand, it can waste time exploring long paths when the target is a shallow node that is examined later.

Consider the following graph:
Let’s use $A$ as the start vertex and pick $F$ as the target. Under our assumptions DFS will find $F$ after examining exactly three vertices: $A$, $B$ and $F$. By contrast, BFS will have to visit all six vertices: $A$, $B$, $C$, $D$, $E$ and finally $F$.

Using the same graph, let’s set $C$ as our target. In this case, DFS will find $C$ after backtracking from $D$, i.e., after visiting the four vertices $A$, $B$, $F$ and $C$. Instead, BFS will only need to examine three vertices: $A$, $B$ and $C$.

**Solution of exercise** ?? This code differs from BFS by the presence of the loop on lines ??–??. This snippet of code has cost $O(v)$ at each iteration of the outer loop, or $O(v^2)$ altogether since at most $v$ vertices can be pushed on the queue.

Recall that, with the adjacency list representation, the cost of BFS was $O(v + e)$, with the $v$ part coming from the number of deq’s (and from initializing the mark array) and the $e$ part from the processing of the neighbors over all iterations. We need to add $v^2$ to this. The complexity is therefore $O(v + e + v^2)$ which simplifies to $O(v^2)$ since $e \in O(v^2)$.

With the adjacency matrix representation, the cost of BFS was $O(v^2)$. Adding $v^2$ as a new term does not change this complexity.