# Two Graph Vertex Partitioning Algorithms for Part Consolidation in Axiomatic Design

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## 1 Graph Theory

This section presents the concepts, definitions, and theorems from the field of graph theory that are needed in the development of the two algorithms. This material is primarily taken from the textbooks used [1] and class notes compiled by the author during his undergraduate and graduate graph theory classes at SJSU.

## 1.1 Simple Graphs

The problem of part consolidation is best served by a class of graphs called *simple graphs*:

#### **Definition: Simple Graph**

A simple graph is a mathematical object represented by a tuple  $G=(V,E,\ldots)$  consisting of a non-empty and finite set of vertices (also called nodes) V(G), a finite and possibly empty set of edges E(G), and zero of more relations. Each edge is represented by a two-element subset of V(G) called the *endpoints* of the edge:

$$E(G) \subseteq \mathcal{P}_2(V(G))$$

Each relation has V(G) or E(G) as its domain and is used to associated vertices or edges with problem-specific attributes.

For the remainder of this work, the use of the term "graph" implies a "simple graph."

The choice of two-element subsets of V(G) for the edges has certain ramifications that are indeed characteristics that differentiate a simple graph from other classes of graphs:

- 1. Every two vertices of a graph are the endpoints of at most one edge; there are no so-called *multiple* edges between two vertices.
- 2. The two endpoint vertices of an edge are always distinct; there are no so-called *loop* edges on a single vertex.
- 3. The two endpoint vertices are unordered, suggesting that an edge provides a bidirectional connection between its endpoint vertices.

Thus, a part consolidation problem can be represented by a graph whose vertices are the FRs and whose edges discourage combining their endpoint FRs into a single part: in the case of the first algorithm, each edge is given a numerical score (weight) indicating the magnitude of the desire to not combine the endpoint FRs into a single part, and in the case of the second algorithm, each edge indicates that the endpoint FRs should never be combined into a single part.

Graphs are often portrayed visually using labeled or filled circles for the vertices and lines for the edges such that each edge line is drawn between its two endpoint vertices. An example graph is shown in Figure 1.1.

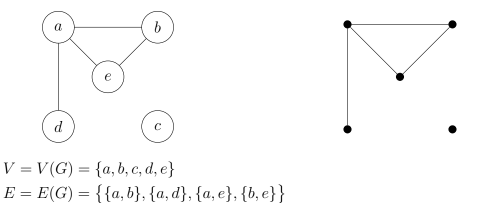


Figure 1: An Example Graph (labeled and unlabeled)

When referring to the edges in a graph, the following common notation will be used:

## **Notation: Edge**

The edge  $\{u, v\}$  is represented by the simple juxtaposition uv or vu.

Note that there is no requirement that every vertex in a graph be an endpoint to some edge:

#### **Definition: Isolated Vertex**

Let G be a graph and let  $u \in V(G)$ . To say that u is an *isolated* vertex means that it is not an endpoint for any edge in E(G):

$$\forall vw \in E(G), u \neq v \text{ and } u \neq w$$

In the example graph of Figure 1.1, notice that vertex c is an isolated vertex.

When two vertices are the endpoints of the same edge the vertices are said to be *adjacent*:

#### **Definition: Adjacent Vertices**

Let G be a graph and let  $u, v \in V(G)$ . To say that u and v are *adjacent* vertices, also called *neighbors*, means that they are the endpoints of some edge  $e \in E(G)$ :

$$\exists e \in E(G), e = uv$$

The edge e is said to *join* its two endpoint vertices u and v. Furthermore, the edge e is said to be *incident* to its endpoint vertices u and v.

In the example graph of Figure 1.1, notice that vertex a is adjacent to vertices b, e, and d; and vertex b is adjacent to vertex e.

We can also speak of adjacent edges, which are edges that share an endpoint:

## **Definition: Adjacent Edges**

Let G be a graph and left  $e, f \in E(G)$ . To say that e and f are adjacent edges means that they share some endpoint  $v \in E(G)$ :

$$\exists v \in V(G), e \cap f = \{v\}$$

or similarly:

$$|e \cap f| = 1$$

Note that two edges in a simple graph can only share one endpoint; otherwise, the two edges would be multiple edges, which are not allowed in simple graphs.

In the example graph of Figure 1.1, notice that ab is adjacent to ad, ae, and be; and ae is adjacent to be.

#### 1.2 Order and Size

Two of the most important characteristics of a graph are the number of vertices in the graph, called the *order* of the graph, and the number of edges in the graph, called the *size* of the graph:

#### **Definition: Order**

Let G be a graph. The *order* of G, denoted by n(G), is the number of vertices in G:

$$n = n(G) = |V(G)|$$

#### **Definition: Size**

Let G be a graph. The *size* of G, denoted by m(G), is the number of edges in G:

$$m = m(G) = |E(G)|$$

In the example graph of Figure 1.1, notice that n=5 and m=4.

Since every two vertices can have at most one edge between them, the number of edges has an upper bound:

#### **Theorem**

Let G be a graph of order n and size m:

$$m \le \frac{n(n-1)}{2}$$

*Proof.* Since each pair of distinct vertices in V(G) can have zero or one edges joining them, the maximum number of possible edges is  $\binom{n}{2}$ , and so:

$$m \le \binom{n}{2} = \frac{n!}{2!(n-2)!} = \frac{n(n-1)}{2}$$

Some choices of graph order and size lead to certain degenerate cases that serve as important termination cases for the two algorithms:

#### **Definition: Degenerate Cases**

- The *null* graph is the non-graph with no vertices (n = m = 0).
- The *trivial* graph is the graph with exactly one vertex and no edges (n=1,m=0). Otherwise, the graph is *non-trivial*.
- An *empty* graph is a graph with possibly some isolated vertices but with no edges (m=0).

Hence, both the null and trivial graphs are empty.

## 1.3 Labeled Graphs

One of the possible relations in a graph tuple is a function that assigns each vertex an identifying label:

#### **Definition: Labeled Graph**

To say that a graph G is *labeled* means that its vertices are considered to be distinct and are assigned identifying names (labels) by adding a bijective labeling function to the graph tuple:

$$\ell:V(G)\to L$$

where L is a set of labels (names). Otherwise, the vertices are considered to be identical (only the structure of the graph matters) and the graph is *unlabeled*.

The vertices in a labeled graph are typically draw as open circles containing the corresponding labels, whereas the vertices in an unlabeled graph are typically drawn as filled circles. This is demonstrated in the example graph of Figure 1.1: the graph on the left is labeled and the graph on the right is unlabeled.

Since the labeling function  $\ell$  is bijective, a vertex  $v \in V(G)$  with label "a" can be identified by v or  $\ell^{-1}(a)$ . In practice, the presence of a labeling function is assumed for a labeled graph and so a vertex is freely identified by its label. This is important to note when a proof includes a phrase such as, "let  $v \in V(G)$  . . ." since v may be a reference to any vertex in V(G) or may call out a specific vertex by its label; the intention is usually clear from the context.

Thus, the part consolidation graphs that act as the inputs to the two algorithms are labeled graphs, where the labels represent the various functional requirements:

$$FR_1, FR_2, FR_3, \ldots, FR_n$$

## References

[1] G. Chartrand and P. Zhang. *A First Course in Graph Theory*. Dover Publications, Mineola, New York, 2012.