The graph2tab library, an introduction

This package is a generic implementation of a method for producing spreadsheets out of pipeline graph. The core of this processing is based on the node splitting approach: all the nodes in the input pipeline are reduced to 'chain nodes', i.e.: nodes that have at most one input and at most one output. This is done by taking those nodes having splittings or pooling, creating copies of them and distributing the excessive inputs or outputs over the copies. See the class ChainsBuilder for details about such procedure.

The package is designed in a way that allows to adapt a particular object model (eg.: ISATAB or MAGETAB) and to reuse classes like TableBuilder to produce spreadsheets describing a particular experimental pipeline (ie.:CSV/TSV files).

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To-do list

In a nutshell, what you've to do to use graph2tab, in most cases is:

- define your node wrappers and the attributes the yield
- define your node factory
- define your table builder
- if needed, define some criteria to sort nodes, e.g.: tells the library that 'Source Name' comes before 'Sample Name'
- invoke your table builder with the right set of nodes coming from your to-be-converted model (such as the sources or the end nodes). Collect the results as a matrix, the SDRF-like table.

Let's see the details below.

Set up

The graph2tab library is available as a Maven Java project. The quickest way to use it is to include it as a dependency in your Maven project:

```
<dependency>
  <groupId>org.isatools</groupId>
  <artifactId>graph2tab</artifactId>
  <version>1.0</version>
</dependency>
```

TODO: will change?

```
<repository>
  <id>ebi-repo</id>
  <name>EBI repo</name>
  <url>http://www.ebi.ac.uk/~maven/m2repo</url>
</repository>
```

You may want to download the sources, especially to look at the examples we describe in this guide. The sources are available via GIT at the address (TODO: will change):

https://github.com/marco-brandizi/ISAvalidator-ISAconverter-BIImanager/tree/master/graph2tab

We have never tried it without using Maven. If that is your case, we suggest that you use the Assembly plug-in from the sources to build a Jar, that can be included in your project:

TODO: this is not ready yet, we need to add the Assembly descriptor for this.

The basics

graph2tab can be used to convert a graph about an experimental pipeline to a tabular view, similar to the SDRF structure in the MAGETAB format, or the Sample/Assay files in the ISATAB format.

In order to do that, you must start from a Java object model that is used to represent the experimental pipeline (e.g.: the BII model, or the model defined in the Limpopo MAGETAB parser). For example, you'll have instances of classes like BioSource or Hybridization and these will have properties like factorValues, characteristics, inputs, outputs.

graph2tab is designed to be generic: apart from few basic elements, the graph/table conversion does not depend neither on the particular graph that is being converted, nor on the particular set of headers that the output consists of. The library ensures this generality by working with a few basic interfaces.

Node is one of the most important of them, it just defines inputs, outputs and attributes. The former point at other instances of the Node interface itself. The attributes are defined in in a tabular-oriented way, that is: every attribute is a pair of strings, the header and the value. These will correspond to headers/values in the spawn table format. The way this is implemented consists of two lists of strings, defined in TabValueGroup, one for the headers and one for the values.

uk/ac/ebi/tablib/export/graph algorithm/Node.java

```
package uk.ac.ebi.tablib.export.graph_algorithm;

public interface Node extends Comparable<Node>
{
    public SortedSet<Node> getInputs ();
    public boolean addInput ( Node input );
    public SortedSet<Node> getOutputs ();
    public SortedSet<Node> getOutputs ();
    public boolean addOutput ( Node output );
    public boolean removeOutput ( Node output );
    public boolean removeOutput ( Node output );
    public List<TabValueGroup> getTabValues ();
    public String getType ();
    public int getOrder ();
    public Node createIsolatedClone ();
}
```

uk/ac/ebi/tablib/export/graph algorithm/TabValueGroup.java

```
package uk.ac.ebi.tablib.export.graph_algorithm;

public interface TabValueGroup
{
    public List<String> getHeaders ();
    public List<String> getValues ();
}
```

An example of how these interface can be used is provided in the package uk.ac.ebi.tablib.export.graph_algorithm.simple_biomodel_tests (test/ folder). Here, you've two examples of how a to-be-converted model can be adapted to graph2tab. One is very simple and it's in the package dummy_graphs_tests. Here nodes from the input model are directly implementing out Node interface. While this is so simple, it's not very realistic, because you will typically have your model and you will not want to derive it from classes in the grap2tab package. So, in such cases, what you need is something similar to the example in the package simple_biomodel_tests. Let's describe this in detail. Suppose that the model you've to convert has a node like BioMaterial in this model, which has this shape:

uk/ac/ebi/tablib/export/graph algorithm/simple biomodel tests/model/BioMaterial.java

```
package uk.ac.ebi.tablib.export.graph_algorithm.simple_biomodel_tests.model;

public class BioMaterial extends ExperimentNode
{
    public BioMaterial ( String name )
    {
        super ( name );
    }

    /**
    * Biomaterial characteristics are managed by means of {@link Annotation}.
    */
    public void addCharacteristic ( String type, String value, String termAcc, String termSrc )
    {
        super.addAnnotation ( type, value, termAcc, termSrc );
    }

    /**
    * Biomaterial characteristics are managed by means of {@link Annotation}.
    */
    protected List<Annotation> getCharacteristics () {
        return super.getAnnotations ();
    }
}
```

uk/ac/ebi/tablib/export/graph algorithm/simple biomodel tests/model/Annotation.java

```
public Annotation ( String type, String value, OntoTerm ontoTerm ) {
    ...
}

public Annotation ( String type, String value ) {
    ...
}

public String getType () {
    ...
}

public String getValue () {
    ...
}

public OntoTerm getOntoTerm () {
    ...
}
```

As you can see, Annotation defines the possible attributes for the input model. This is an example quite similar to the kind of attributes the nodes have in MAGE-OM, MAGETAB, ISATAB. In order to be adaptable to concrete specific situations it generic, the graph2tab library sees the model nodes in an abstract generic form, where the node is a list of string pairs, like Source Name = 'source 1', Characteristics[Organism] = 'Mus-musculus', ... Most of the work that you have to do to use the library in your case consists of writing code to convert the attributes from your model, whatever they are, to such string pairs, i.e.: you've to convert them to instances of TabValueGroup. The reason why a Node exposes a list of groups of pairs, instead of a flat list of header/value pairs, is that certain groups of attribute pairs must be kept always together, in order to produce correct results. For example, Characteristics[Organism] = 'Mus-musculus', Term Source REF = 'NCBI-Tax', Term Accession = '123' must be the a successive groups of columns in the resulting table. You may have noted the existence of the methods getType() and getOrder(). This is explained in the section 'Uneven graphs and their layering'.

Model wrapping

The typical approach to implement such an adaptation from your model to the graph2tab interfaces is defining wrappers. Let's go straight to an example about the aforementioned Biomaterial. This derives from another class, written for this particular model, which is the abstract ExpNodeWrapper.

$uk/ac/ebi/tablib/export/graph_algorithm/simple_biomodel_tests/node_wrappers/ExpNodeWrapper.java$

```
package uk.ac.ebi.tablib.export.graph_algorithm.simple_biomodel_tests.node_wrappers;
...
public abstract class ExpNodeWrapper extends DefaultAbstractNode
{
    private ExperimentNode base;

    ExpNodeWrapper ( ExperimentNode base )
    {
        this.base = base;
    }

    /**
    * This is used by the implementation of {@link Node#createIsolatedClone()}. Essentially, copies
    * the wrapped node and makes empty input/output sets.
    **
```

```
*/
protected ExpNodeWrapper ( ExpNodeWrapper original )
       this.base = original.base;
       this.inputs = new TreeSet<Node> ();
       this.outputs = new TreeSet<Node> ();
}
/**
* In this case we're able to write a generic method that is customised by the descendants.
 * The methods shows how to add an ontology term to a free text value. This is done by keeping
 * all into the same {@link TabValueGroup}.
 * @param nameHeader eg, "BioMaterial Name", "Protocol REF"
 * @param annHeaderPrefix eg: "Characteristic", "Parameter Value", here headers will be built
 * with the schema annHeaderPrefix [ type ], eg: Characteristic [ Organism ]
*/
protected List<TabValueGroup> getTabValues ( String nameHeader, String annHeaderPrefix )
       List<TabValueGroup> result = new ArrayList<TabValueGroup> ();
       result.add ( new DefaultTabValueGroup ( nameHeader, base.getName () ) );
       for ( Annotation annotation: base.getAnnotations () )
       {
               DefaultTabValueGroup tbg = new DefaultTabValueGroup (
                 annHeaderPrefix + " [ " + annotation.getType () + " ]", annotation.getValue ()
               OntoTerm ot = annotation.getOntoTerm ();
               if ( ot != null ) {
    tbg.add ( "Term Accession Number", ot.getAcc () );
    tbg.add ( "Term Source REF", ot.getSource () );
               result.add ( tbg );
       }
       return result;
}
* If it's not a clone produced by {@link Node#createIsolatedClone()},
* it uses {@link NodeFactory} to build wrappers
* This is the typical way this method is implemented by.
*/
@Override
public SortedSet<Node> getInputs ()
       if ( inputs != null ) return super.getInputs ();
       inputs = new TreeSet<Node> ();
       NodeFactory nodeFact = NodeFactory.getInstance ();
       for ( ExperimentNode in: base.getInputs () ) inputs.add ( nodeFact.getNode ( in ) );
       return super.getInputs ();
}
@Override
public SortedSet<Node> getOutputs ()
       . . .
}
. . .
```

}

$uk/ac/ebi/tablib/export/graph_algorithm/simple_biomodel_tests/node_wrappers/BioMaterialWrapper.java$

```
package uk.ac.ebi.tablib.export.graph_algorithm.simple_biomodel_tests.node_wrappers;
...
public class BioMaterialWrapper extends ExpNodeWrapper
{
    BioMaterialWrapper ( BioMaterial base ) {
        super ( base );
    }

    private BioMaterialWrapper ( ExpNodeWrapper original ) {
        super ( original );
    }

    public List<TabValueGroup> getTabValues () {
        return getTabValues ( "Biomaterial Name", "Characteristic" );
    }

    public Node createIsolatedClone () {
        return new BioMaterialWrapper ( this );
    }
}
```

Details

You're free to directly implement the Node interface for defining your wrappers. However, for common cases, extending DefaultAbstractNode is probably quite easier. The latter defines common functionality, such as the collection class members for maintaining input/output nodes.

Differently than DefaultAbstractNode, which is a general class and part of the main package, ExpNodeWrapper makes sense in this example only (it's in the test/ folder too) and define a few common elements that apply to the simple model used for this particular case.

Every node type in your original model has a corresponding wrapper and the two are linked by the base object received by the constructor.

getInputs(), getOutputs(). Your custom implementation will typically behave like (for getInput()): if the inputs collection != null returns it, else initialise it by getting wrappers for each node that is input of the base node for the current wrapper. Wrappers are usually created via a custom wrapper factory (more later on that). In short, what we're doing is creating a graph of wrappers that is isomorphic to the graph of nodes from your to-be converted model.

getTabValues(). In this particular model (yours may be more complicated), there is only one type of attribute that can be attached to nodes: the Annotation class. So the getTabValues() method converts the attribute into the required list of headers/values. In this foo example, each header is typed, ie.: it is like 'Characteristics[Organism]', every attribute has a string value and can have an optional ontology term, which, in turn is made of an accession and a source. So, the wrapper defines the node type and name as first TableValueGroup (eg.: 'BioMaterial'/'source1'), followed by one TableGroup per Annotation, which contains the annotation type, its value and, optionally, additional pairs like 'Term Accession Number'/'123', 'Term Source REF'/'MO'.

createIsolatedClone() and ExpNodeWrapper(ExpNodeWrapper original). The conversion algorithm needs to duplicate nodes. That is, given a node, it needs to create another wrapper node that has the same base (points to the same node in your original model), but different edges (ie: different inputs/outputs), taken from the original node. For instance, a 2-input/2-output node is split into two nodes, the second node will get one input and one output from the original, which will be left with the other two edges. The algorithm decides which edges to remove/add, so the duplicated node is initially isolated (no inputs/outputs). Now, createIsolatedClone() provides such a duplicate of the current node (ie: of this).

The typical implementation of such method is something like return new ExpNodeWrapper(this). So the constructor of this type (the ones that receive an instance of the same type the class is), as you can see, actually creates the isolated duplicate. Indeed, it's quite simple: the new wrapper gets the base from the original one and initialises inputs/outputs with empty sets (see ExpNodeWrapper above).

More notes

- The order the attributes are defined with in the wrappers is reflected in the final output table (this also depends on how nodes of the same types are merged, see below).
- As explained above, those header/value pairs that go together have to be put in the same TableGroup object, so that they are never split apart during the conversion. For instance, if you have a factor value with an ontology term, you have to put the 'Factor Value', 'Term Accession', 'Term Source' headers into the same TableGroup (and add the corresponding values of course, which may be "", as long as getValues().size() is always = getHeaders().size()).
- Two nodes of the same type (eg: two sample nodes) may have different lists of attributes, eg: one has a 'Characteristics[cell line]' attribute and the other doesn't, you just don't define any cell line or value for the second one. The converter takes care automatically of 'merging' the two (or more) samples, creating the right set of columns and populating the lines with proper values or empty cells. **However**, do not play with attribute groups of the same type (having the same initial header), for example, if you define 'Factor Value [X]' + ontology accession/source headers once, just redefine the same three headers every time you need to export the factor value of type X. Fill missing accessions or sources with empty strings if needed (should the factor value be blank, then accession and source will be empty too => you don't need this TabValueGroup at all in such a case). This is necessary, because it's hard to manage table groups with the same initial header but different lists of following headers. You may have unpredictable results if you don't keep them uniform for the same node types.
- getInputs() and getOutputs() return SortedSet(s). This has a few implications. First, it's a bonus that allows you to establish some order on the output lines. Typically, nodes are compared on the basis of attributes like sample name or hybridization name. Second, you need to implement proper versions of compareTo(), equals() and hashCode(). DefaultAbstractNode has sensible default implementations for these methods that take the value of the first element in the first TableGroup returned by getTabValues(), i.e., exactly things like 'Source Name' or 'Sample Name'. This is usually what you're happy with. The implementation in DefaultAbstractNode also ensures a fundamental property: that two physically different node wrappers are considered different (e.g., by collection-related methods), even when they are duplicated and point at the same base (the same node from your to-be-converted model, which also means the same source or alike). This is really important, because unwanted behaviours would be observed otherwise. For example, without such an implementation, duplicated nodes would be considered the same when you attempted to store them in the same set structure (so one of them would be discarded). So, the message is: extend DefaultAbstractNode or, should you have particular needs why this implementation doesn't suit you well, at least give a look to the source of this class.

Node Factory

If you adapt mage2tab to your model with the wrappers approach, you typically will use an extension of AbstractNodeFactory either. This just keeps a map of base nodes (nodes from your model) => wrappers. It uses it in the method NodeWrapper getNode(Base node), which creates a new wrapper and links it to the the given base, but only if this is not created yet and already inside the node/wrapper map. Obviously, it stores the new wrapper in the map, to return it in subsequent calls that have the same base as parameter (instead of creating a new wrapper). This ensures that you create a 1-1 graph of wrappers from the original graph is based on your to-be-converted model. Indeed, you don't need to care about these details, you only need to implement createNewNode(), which will be like this:

uk/ac/ebi/tablib/export/graph algorithm/simple biomodel tests/node wrappers/NodeFactory.java

This method is what the getNode() above will call, when it decides it needs a new wrapper.

Table Builder

The TableBuilder class coordinates the conversion job. We suggest you extend it and define a method that receives nodes from your input graph, as it is done in the constructor SimpleModelTableBuilder(
Set<ExperimentNode> nodes):

$uk/ac/ebi/tablib/export/graph_algorithm/simple_biomodel_tests/node_wrappers/SimpleModelTableBuilder.java$

```
public class SimpleModelTableBuilder extends TableBuilder
{
    public SimpleModelTableBuilder ( Set<ExperimentNode> nodes )
    {
        this.nodes = new HashSet<Node> ();
        NodeFactory nodeFact = NodeFactory.getInstance ();
        for ( ExperimentNode node: nodes ) this.nodes.add ( nodeFact.getNode ( node ) );
    }
}
```

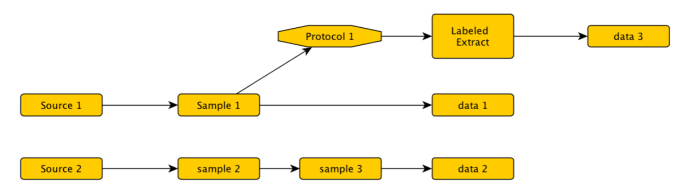
As you can see, this method just populates the nodes class member, containing node wrappers and does it by using your node factory. As for the nodes that should be passed to the table builder, any set of nodes that allow one to reach all the graph by walking from such nodes upstream (following inputs up to the sources) and downstream (following outputs up to the sinks/end nodes) is fine. For a discussion about optimal choices, see the next section about the layering algorithm.

Go!

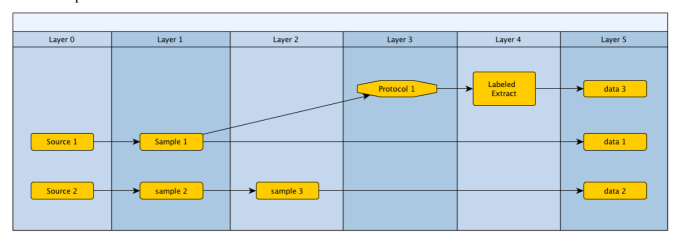
Once all is in place, you can go with the conversion, basically invoking <code>getTable()</code> from your table builder. An invocation example can be found in <code>SimpleBioModelTest</code>.

Uneven graphs and their layering

What we have explained so far works correctly only upon the condition that the graph to be converted is 'even', which means that all the paths from the sources (e.g., bio-sources) to the end nodes (e.g., data) have the same lengths. graph2tab supports 'uneven' graphs as well, e.g., a graph where you may have paths of different lengths, with some experimental workflow stages omitted. An example is this:



This kinds of graphs are converted into tables by means of the same approach described above, which transforms the initial graph into a set of chains. However, in order to cope with such uneven paths, the input graph is pre-processed by a 'layering' component (if you are curious, this is implemented by the LayersBuilder class, but you don't need to know anything about it in order to use the library to convert your experimental workflows). This component distributes the graph nodes into a set of layers, where every layer reflect walks from sources to end nodes and the layering algorithm arranges the nodes into the layers in a way such that all the nodes in a given layer have the same type. In the example above, this would produce:



Once the graph is re-arranged this way (i.e.: every node gets a layer index), it is easy to apply the chaining procedure above and to generate a table that has gaps:

Source Name	Sample Name	Sample Name	Protocol REF	Labeled	Data File
				Extract Name	

Source 1	Sample 1		Protocol 1	Labeled Extract	Data 3
Source 1	Sample 1				Data 1
Source 2	Sample 2	Sample 3			Data 2

The layering algorithm implemented in the library is essentially based on the method briefly described in the MAGE-TAB specification¹, with several necessary details (not explicitly considered in that description) added by us. Once a graph is adapted to the Node interface, there isn't much else that you have to care about in order to make graph2tab to compute the node layers. However, because of the generic nature of the library, there is some information that you are required to provide with. This is done via the Node.getOrder() method. It works like this: layer indexes are initially assigned to the graph based on the highest distance from their sources (e.g.: 'Protocol 1' and 'Sample 3' in the example above would get 2). After that, for each layer, all the nodes in that layer are compared each other. If two nodes in the same layer are of a different type, getOrder() is invoked to decide which one is supposed to shift on the right (the one having a bigger or equal order). If you expect to work with uneven graphs, you do have to provide an implementation of getOrder() in your custom implementation. Two typical ways to do this are:

- An array of header types is defined (of strings), such as 'Source Name', 'Sample Name', 'Extract Name' ... and the order of a node is computed by considering its first header (actually Node.getType(), see below) and taking the index of this header in such array. This is equivalent to providing a list of types which of order is known in advance because of the experimental stage they represent
- An alternative is possible for the case the graph to be exported was initially generated from an existing tabular format (e.g.: loaded from a MAGE-TAB submission) and stored to a database. In such a case it is possible to track and store the column index the initial header of a node comes from and use such index as return value for getOrder(). This kind of solution is interesting for those who want to be able to re-export originally submitted data and reflect the initial submission as much as possible. For instance, in the example above the column for 'Protocol REF' may have been determined according to the first criterion, assuming the input for the (labelling) protocol is omitted and the output is specified (as it is more often the case). The opposite may have been true and you could track this by using this second criterion for setting the order.

The use of the order approach is very flexible and can be used for many different custom cases. For example, you may decide to combine the two criteria above, using a predefined array for all the fields but the protocol references, for which you may want to apply the column position. Another more advanced example consists of considering the type of protocol a 'Protocol REF' node refers to and assigning the order on the basis of this. Then additional fields would be configured in the list of possible fields, e.g.: 'Sampling Protocol', 'Extraction Protocol' etc. This is the reason why the Node interface has the method <code>getType()</code>. Normally the type is the same as the first header (and this is the default implementation in <code>DefaultAbstractNode</code>). However, in a case like the one at issue, it may be that the first header is 'Protocol REF', while the node type is something like 'Extraction Protocol', achieved from the referred protocol. <code>getType()</code> could be useful when you have entities reported with the same header, but of slightly different subtype, e.g., you want to separate "treatment protocol" and "extraction protocol".

There will be certain fields that will have an undefined order (which corresponds to -1 in the graph2tab library), for instance 'Protocol REF'. Where one of such fields have to be shifted, the layering procedure considers the order of the nodes on its left and on its right and the other node being compared, if this has

¹ Section 3.3.2 in http://annotare.googlecode.com/files/MAGE-TABv1.1.pdf

an order different than -1. For instance, in the example above, the protocol 1 is shifted on the right, since sample 3, the node that initially shares the layer with such protocol reference, has an order that is closer to the one of sample 1 and sample 2 than the lowest order found on its right, which is the one of Labelled Extract (order(sample 3) – order(sample 1/2) is 0, being the nodes of the same type). The rationale for this is to prefer a right shift of nodes like protocol references in such situations, because in real biological cases it is more frequent to report a protocol output together with a named protocol and omit the input, rather than the opposite. See the source of LayersBuilder for details about the implemented shifting criteria.

Layering example in the source code

The graph2tab package contains a 'layered implementation' in the package layering_tests. Please see the Junit tests in the class LayeringTests. The class ExpNodeWrapper, which was reported above in reality contains an implementation of the getOrder() method that uses the first approach described before (the array of possible headers). The package layering_tests.model extends simple_biomodel_tests.model with a few nodes that makes it possible to build examples of uneven graphs.

This is a fragment from ExpNodeWrapper, showing how getOrder() is implemented there:

```
public abstract class ExpNodeWrapper extends DefaultAbstractNode
{
    ...
    public static final Map<String, Integer> TYPE_ORDER = new HashMap<String, Integer> ()
    {{
        put ( "Source Name", 0 );
        put ( "Sample Name", 1 );
        put ( "Extract Name", 2 );
        put ( "Labeled Extract Name", 3 );
        put ( "Assay Name", 4 );
        put ( "Data File Name", 5 );
    }};
    ...
    public int getOrder ()
    {
        String header = getType();
        Integer order = TYPE_ORDER.get ( header );
        return order == null ? -1 : order;
    }
    ...
}
```

Disabling the layering stage for even graphs

By default the TableBuilder class creates and instance of the LayersBuider class and invokes the layering procedure, before doing the the rest of the conversion process. If you are absolutely sure that your custom implementation will deal with even graphs only, this layering step, while not affecting the results in your case, is not necessary. Therefore we have provided an option to disable it, so that you can save some processing time (and a little of RAM too) if you don't need this step. To disable the layering stage, see the TableBuilder constructor that accepts the parameter isLayeringRequired. This parameter is true by default.

Choosing the initial node set to be passed to the table builder

We have already shown that TableBuilder needs a certain subset of nodes of your input graph, in order to reach all the nodes and work out the conversion to a table. A simple working choice for this consists of all the nodes the graph is composed of. However, you can speed things up a little by using these criteria:

- If the layering stage is enabled, as it is by default (see above), the best choice is to pass all the

graph end-nodes, i.e.: all the nodes that have no output. This is because the layering stage needs to start working from the rightmost side of the graph and to compute the initial values for the layer indexes recursing from such nodes back to the sources. Hence, the LayersBuilder class will first of all walk the initial nodes you pass to the table builder toward right, seeking for such sink nodes. If you pass this type of nodes directly, this initial search will produce a minimal overhead.

- If the layering stage is disabled, the table builder is in a symmetric situation: the initial nodes are passed directly to ChainsBuilder, the class that transforms the graph into chains. Because in this case the graph has to be scanned in left-to-right direction, the chain builder will initially seek all the source nodes, doing a walk from initial nodes to the leftmost nodes that can be reached from them (i.e., all the nodes without any input). Consequently, in this case the fastest option is to pass your graph sources to the table builder.
- Note that when the layering is active and you pass sinks, as suggested above, LayersBuilder will compute the graph sources as part of its job and these, not the initial node set, will be those that will be passed to ChainsBuilder in this case. This confirms that the sinks is the best choice in such a case.