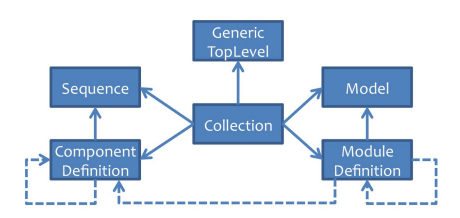
Introduction to the SBOL2 Data Model, the Java Library (libSBOLj), and the Javascript Library (sboljs)

Chris J. Myers

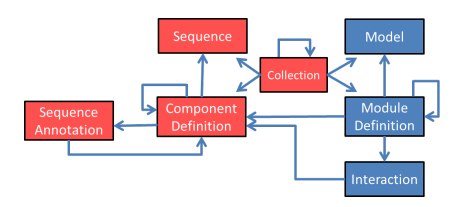
University of Utah

SBOL Workshop August 1, 2018

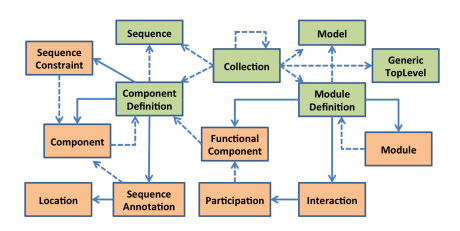
SBOL 2 Data Model



SBOL 2 Data Model



SBOL 2 Data Model



SBOL 2.2 Updates

- Attachment class serves as a general container for data files.
- Implementation class represents a physical instance of a synthetic biological construct.
- Provenance ontology (Prov-O):
 - Activity class something that occurs over a period of time and acts upon or with entities.
 - Agent class something that bears some form of responsibility for an activity taking place.
 - Plan class an entity that represents a set of actions or steps intended by one or more agents to achieve some goals.

SBOL Libraries

- Crucial to the success of a standard is software infrastructure to support developers' integration of the standard within their tools.
- There are several library implementations of the SBOL data structure, which provide an application programmers interface (API) for tool developers to interact with SBOL data objects.
 - libSBOLj native Java library
 - sboljs Javascript library
 - libSBOL C/C++ library
 - pySBOL Python library

SBOL Libraries

- Crucial to the success of a standard is software infrastructure to support developers' integration of the standard within their tools.
- There are several library implementations of the SBOL data structure, which provide an application programmers interface (API) for tool developers to interact with SBOL data objects.
 - libSBOLj native Java library
 - sboljs Javascript library
 - libSBOL C/C++ library
 - pySBOL Python library

libSBOLj - native Java SBOL Library

- Library available from Maven central and Sonatype:
 - Group Id org.sbolstandard
 - Artifact Id libSBOLj
 - Version 2.3.1 (latest release)
 - Version 2.3.2-SNAPSHOT (current version)
- Distribution includes detailed documentation for the class definitions and the methods provided by the API.
- Supports validation/conversion to/from FASTA, GenBank, and SBOL1.
- Includes support to search, fetch, and submit designs to SynBioHub.
- Utilized by an online validator/converter available from the SBOL website, which also provides a webservice to be used by non-java applications.

Example: Converting Cello UCF File to SBOL

- JSON file format including the following sections:
 - Parts basic DNA parts (promoters, rbs, cds, terminators, etc.)
 - Gate parts compositions of DNA parts to form complete functions
 - Gates additional information about these gate parts
 - Other information response functions, gate toxicity, etc.

Cello UCF File Example

```
"collection": "parts",
  "type": "cds",
  "name": "AmtR".
  "dnasequence": "ATGGCAGGCGCAGTTGGTCGTCGCGTCGTAGTGCACCGCGTCGTGCAGGTAAAAATCCGCGTGAAGAAATTC..."
},...
  "collection": "gates".
  "regulator": "AmtR",
  "group name": "AmtR",
  "gate_name": "A1_AmtR",
  "gate type": "NOR",
  "system": "TetR",
  "color_hexcode": "3BA9E0"
  "collection": "gate_parts",
  "gate_name": "A1_AmtR",
  "expression cassettes": [
      "maps to variable": "x",
      "cassette parts": [
        "BydvJ",
        "A1".
        "AmtR".
        "L3S2P55"
  "promoter": "pAmtR"
}, ...
```

Parsing Cello JSON File

```
HashMap<String.JSONObject> partsMap = new HashMap<String.JSONObject>():
HashSet<JSONObject> qate_partsArr = new HashSet<JSONObject>();
HashMap<String, JSONObject> gatesMap = new HashMap<String, JSONObject>();
HashMap<String.JSONObject> responseMap = new HashMap<String.JSONObject>():
JSONParser parser = new JSONParser();
JSONArray a = (JSONArray) parser.parse(new FileReader(pathToUCFFile));
for (Object o : a)
  JSONObject ucf = (JSONObject) o:
  String collection = (String) ucf.get("collection");
  if (collection.equals("parts")) {
    partsMap.put((String)ucf.get("name"),ucf);
  else if (collection.equals("gate parts")) {
    gate partsArr.add(ucf);
  else if (collection.equals("gates"))
    gatesMap.put((String)ucf.get("gate name"),ucf);
  else if (collection.equals("response functions")) {
    responseMap.put((String)ucf.get("gate name"),ucf);
```

SBOL Documents

- libSBOLj organizes all SBOL data within an SBOLDocument.
- Example creating an SBOLDocument:

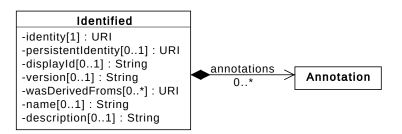
```
String uriPrefix = "http://cellocad.org/";
SBOLDocument document = new SBOLDocument();
document.setDefaultURIPrefix(uriPrefix);
document.setComplete(true);
document.setCreateDefaults(true);
```

- Default URI prefix prefix to use when none provided to create method.
- Complete ensure that all URI references point to valid SBOL objects.
- Create defaults implicitly create ComponentInstances as needed.

Understanding UML Diagrams

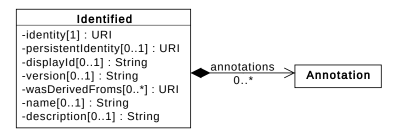
- Classes represented as boxes labeled with their member variables with types and cardinalities.
- Arrows with cardinality indicate associations between classes.
- A hollow diamond at the origin of an arrow represents shared aggregation (i.e., object is referenced and not owned).
- A solid diamond at the origin of an arrow represents composite aggregation (i.e., child object is owned by its parent object).
- Hollow arrows are used to represent inheritance.

Identified (Base Class for All SBOL Objects)



- identity globally unique URI to identify this object (required).
- persistentIdentity identity shared by multiple versions of the same object (optional).
- displayId human-readable id composed of alphanumeric and underscore characters (optional).
- version uses semantic versioning to identify multiple versions of the same object (optional).

Identified (Base Class for All SBOL Objects)



- wasDerivedFroms identities of objects that this is derived from (optional).
- name human-readable String of arbitrary characters (optional).
- description thorough text description of the object (optional).
- annotations additional data about this object (more later).

Compliant Top-Level URIs

$$\langle \mathit{URI}\ \mathit{prefix} \rangle / \langle \mathit{displayId} \rangle / \langle \mathit{version} \rangle$$

- The identity MUST begin with a URI prefix that maps to a domain over which the user has control.
- The persistentIdentity and displayId properties are REQUIRED.
- The persistentIdentity MUST end with a delimiter ('/', '#', or ':') followed by the displayId of the object.
- If an object is not given a version, then its identity and persistentIdentity properties MUST contain the same URI.
- If an object has a version, then its identity property MUST contain a URI of the form \(\langle persistentIdentity \rangle / \langle version \rangle ".

Compliant Child URIs

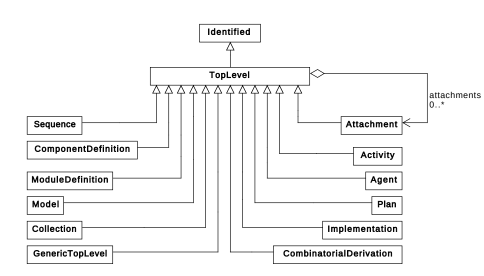
 $\langle parent\ persistent | dentity \rangle / \langle display | d \rangle / \langle parent\ version \rangle$

- The persistentIdentity MUST begin with the persistentIdentity of its parent object and be immediately followed by a delimiter ('/', '#', or ':') and the displayId of the object.
- The version MUST contain the same String as the version property of the object's parent object.

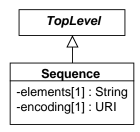
TopLevel Objects

- Each SBOLDocument includes a list of each type of TopLevel object.
- These lists are organized to allow for easy search by their unique reference identifiers (URIs) and validation that they are distinct.
- libSBOLj includes methods for creating, updating, accessing, and removing these data objects, as well as, their child objects.

TopLevel



Sequence¹



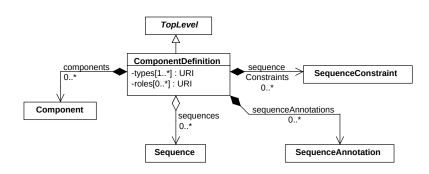
- elements String of characters representing constituents of a biological or chemical molecule.
- encoding URI indicating how elements are to be interpreted.

Encoding	URI	CD Type
IUPAC DNA, RNA	http://www.chem.qmul.ac.uk/iubmb/misc/naseq.html	DNA, RNA
IUPAC Protein	http://www.chem.qmul.ac.uk/iupac/AminoAcid/	Protein
SMILES	http://www.opensmiles.org/opensmiles.html	SmallMolecule

Creating Sequences for Parts

```
for (JSONObject part : partsMap.values()) {
  String name = (String)part.get("name");
  String dnasequence = (String)part.get("dnasequence");
  Sequence sequence = document.createSequence(name + " sequence", version, dnasequence,
    Sequence. IUPAC DNA);
  sequence.setName(name+" sequence");
  sequence.addWasGeneratedBy(activityURI);
  sequence.createAnnotation(new OName(dcTermsNS, "created", "dcTerms"), createdDate);
<sbol:Sequence rdf:about="http://cellocad.org/AmtR sequence/1">
  <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR sequence"/>
  <sbol:displayId> AmtR sequence </sbol:displayId>
  <sbol:version> 1 </sbol:version>
  <dcterms:title> AmtR sequence </dcterms:title>
  <prev:wasGeneratedBy rdf:resource="http://cellocad.org/cello2sbol/1"/>
  <dcterms:created> 2017-08-09T13:54Z </dcterms:created>
  <sbol:elements> ATGGCAGGCGCAGTTGGTCGTCGCGCGTCGTAGTGCACCGCGTCGTGCAGGTAAAAATCCGCGTGAAGAAATTCTGGATGC
  <sbol:encoding rdf:resource="http://www.chem.gmul.ac.uk/iubmb/misc/naseq.html"/>
</sbol:Sequence>
```

ComponentDefinition



ComponentDefinition (Types/Roles)

CD Type	URI for BioPAX Term
DNA	http://www.biopax.org/release/biopax-level3.owl#DnaRegion
RNA	http://www.biopax.org/release/biopax-level3.owl#RnaRegion
Protein	http://www.biopax.org/release/biopax-level3.owl#Protein
Small Molecule	http://www.biopax.org/release/biopax-level3.owl#SmallMolecule
Complex	http://www.biopax.org/release/biopax-level3.owl#Complex

CD Role	URI for SequenceOntology Term	
Promoter	http://identifiers.org/so/SO:0000167	DNA
RBS	http://identifiers.org/so/SO:0000139	DNA
CDS	http://identifiers.org/so/SO:0000316	DNA
Terminator	http://identifiers.org/so/SO:0000141	DNA
Gene	http://identifiers.org/so/SO:0000704	DNA
Operator	http://identifiers.org/so/SO:0000057	DNA
Engineered Gene	http://identifiers.org/so/SO:0000280	DNA
mRNA	http://identifiers.org/so/SO:0000234	RNA

Converting Part Type to SBOL ComponentDefinition Role

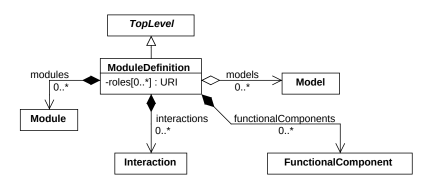
```
public static URI getRole(String type) {
 String so = "http://identifiers.org/so/";
 if (type.equals("ribozyme")) {
    return URI.create(so + "SO:0000374");
  else if (type.equals("scar")) {
    return URI.create(so + "S0:0001953");
  else if (type.equals("cds")) {
    return URI.create(so + "SO:0000316");
  else if (type.equals("promoter")) {
    return URI.create(so + "S0:0000167");
  else if (type.equals("rbs")) {
    return URI.create(so + "SO:0000139");
  else if (type.equals("terminator")) {
    return URI.create(so + "SO:0000141");
  else (
    System.err.println("Part, Type, not, found");
    return null;
```

Converting Parts to SBOL ComponentDefinitions

Converting Parts to SBOL ComponentDefinitions

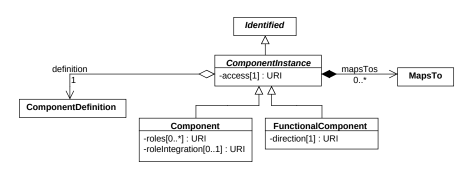
```
<sbol:ComponentDefinition rdf:about="http://cellocad.org/AmtR/1">
 <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR"/>
 <sbol:displayId> AmtR </sbol:displayId>
 <sbol:version> 1 </sbol:version>
 <dcterms:title> AmtR </dcterms:title>
 <dcterms:created> 2017-08-09T13:54Z </dcterms:created>
 <sbol:type rdf:resource="http://www.biopax.org/release/biopax-level3.owl#DnaRegion"/>
 <sbol:role rdf:resource="http://identifiers.org/so/S0:0000316"/>
 <sbol:sequence rdf:resource="http://cellocad.org/AmtR sequence/1"/>
</sbol:ComponentDefinition>
<sbol:ComponentDefinition rdf:about="http://cellocad.org/AmtR protein/1">
 <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR_protein"/>
 <sbol:displayId> AmtR protein </sbol:displayId>
 <shol:version> 1 </shol:version>
 <dcterms:title> AmtR protein </dcterms:title>
 <prev:wasGeneratedBy rdf:resource="http://cellocad.org/cello2sbol/1"/>
 <dcterms:created> 2017-08-09T13:547 </dcterms:created>
 <sbol:type rdf:resource="http://www.biopax.org/release/biopax-level3.owl#Protein"/>
</sbol:ComponentDefinition>
```

ModuleDefinition



Creating ModuleDefinitions for Genetic Production

FunctionalComponent (Child of ModuleDefinition)



Access URI

http://sbols.org/v2#public http://sbols.org/v2#private

Direction URI

http://sbols.org/v2#in http://sbols.org/v2#out http://sbols.org/v2#inout http://sbols.org/v2#none

Description

MAY be referred to by remote **MapsTo** objects.

MUST NOT be referred to by remote **MapsTo** objects.

Description

Indicates that it is an input.

Indicates that it is an output.

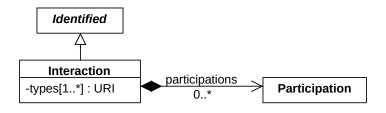
Indicates that it is both an input and output

Indicates that it is neither an input and output.

Creating FunctionalComponents for Genetic Production

```
moduleDefinition.createFunctionalComponent(name, AccessType.PUBLIC,
  componentDefinition.getIdentity(), DirectionType.NONE);
moduleDefinition.createFunctionalComponent(name+" protein", AccessType.PUBLIC,
  proteinComponentDefinition.getIdentity(), DirectionType.NONE);
<sbol:ModuleDefinition rdf:about="http://cellocad.org/AmtR protein production/1">
  <sbol:functionalComponent>
    <sbol:FunctionalComponent rdf:about="http://cellocad.org/AmtR protein production/AmtR/1">
      <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR_protein_production/AmtR"/>
      <sbol:displayId> AmtR </sbol:displayId>
      <sbol:version> 1 </sbol:version>
      <sbol:definition rdf:resource="http://cellocad.org/AmtR/1"/>
      <sbol:access rdf:resource="http://sbols.org/v2#public"/>
      <sbol:direction rdf:resource="http://sbols.org/v2#none"/>
    </sbol:FunctionalComponent>
  </sbol:functionalComponent>
  <sbol:functionalComponent>
    <sbol:FunctionalComponent rdf:about="http://cellocad.org/AmtR_protein_production/AmtR_protein/</pre>
      <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR protein production/AmtR prot</pre>
      <sbol:displayId> AmtR_protein </sbol:displayId>
      <sbol:version> 1 </sbol:version>
      <sbol:definition rdf:resource="http://cellocad.org/AmtR protein/1"/>
      <sbol:access rdf:resource="http://sbols.org/v2#public"/>
      <sbol:direction rdf:resource="http://sbols.org/v2#none"/>
    </sbol:FunctionalComponent>
  </sbol:functionalComponent>
</sbol:ModuleDefinition>
```

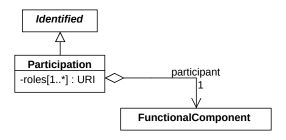
Interaction (Child of ModuleDefinition)



Interaction Type URI for SystemsBiologyOntology Term

Inhibition http://identifiers.org/biomodels.sbo/SBO:0000169
Stimulation http://identifiers.org/biomodels.sbo/SBO:0000170
Biochemical Reaction http://identifiers.org/biomodels.sbo/SBO:0000170
Non-Covalent Binding http://identifiers.org/biomodels.sbo/SBO:0000177
Genetic Production http://identifiers.org/biomodels.sbo/SBO:0000179
Genetic Production http://identifiers.org/biomodels.sbo/SBO:0000188
Control

Participation (Child of Interaction)



Part. Role	URI for SBO Term	Interaction Types
Inhibitor	http://identifiers.org/biomodels.sbo/SBO:0000020	Inhibition
Inhibited	http://identifiers.org/biomodels.sbo/SBO:0000642	Inhibition
Stimulator	http://identifiers.org/biomodels.sbo/SBO:0000459	Stimulation
Stimulated	http://identifiers.org/biomodels.sbo/SBO:0000643	Stimulation
Reactant	http://identifiers.org/biomodels.sbo/SBO:0000010	Non-Covalent Binding, Degradation, Biochemical Reaction
Product	http://identifiers.org/biomodels.sbo/SBO:0000011	Non-Covalent Binding, Genetic Production,
		Biochemical Reaction
Promoter	http://identifiers.org/biomodels.sbo/SBO:0000598	Inhibition, Stimulation, Genetic Production
Modifier	http://identifiers.org/biomodels.sbo/SBO:0000019	Biochemical Reaction, Control

Modified

Template

http://identifiers.org/biomodels.sbo/SBO:0000644

http://identifiers.org/biomodels.sbo/SBO:0000645

Biochemical Reaction, Control

Genetic Production

Creating Interactions for Genetic Production

```
Interaction interaction = moduleDefinition.createInteraction(name+" protein interaction",
  SystemsBiologyOntology.GENETIC PRODUCTION);
interaction.createParticipation(name, name, SystemsBiologyOntology.TEMPLATE);
interaction.createParticipation(name+" protein", name+" protein", SystemsBiologyOntology.PRODUCT);
  <sbol:interaction>
    <sbol:Interaction rdf:about="http://cellocad.org/AmtR protein production/AmtR protein interact</pre>
      <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR_protein_production/AmtR_prot</pre>
      <sbol:displayId> AmtR protein interaction </sbol:displayId>
      <sbol:version> 1 </sbol:version>
      <sbol:type rdf:resource="http://identifiers.org/biomodels.sbo/SB0:0000589"/>
      <sbol:participation>
        <sbol:Participation rdf:about="http://cellocad.org/AmtR protein production/AmtR protein in</pre>
          <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR protein production/AmtR</pre>
          <sbol:displayId> AmtR </sbol:displayId>
          <sbol:version> 1 </sbol:version>
          <sbol:role rdf:resource="http://identifiers.org/biomodels.sbo/SBO:0000645"/>
          <sbol:participant rdf:resource="http://cellocad.org/AmtR protein production/AmtR/1"/>
        </sbol:Participation>
      </sbol:participation>
      <sbol:participation>
        <sbol:Participation rdf:about="http://cellocad.org/AmtR protein production/AmtR protein in</pre>
          <sbol:persistentIdentity rdf:resource="http://cellocad.org/AmtR protein production/AmtR</pre>
          <sbol:displayId> AmtR_protein </sbol:displayId>
          <sbol:version> 1 </sbol:version>
          <sbol:role rdf:resource="http://identifiers.org/biomodels.sbo/SB0:0000011"/>
          <sbol:participant rdf:resource="http://cellocad.org/AmtR protein production/AmtR protein</pre>
        </sbol:Participation>
      </sbol:participation>
    </sbol:Interaction>
  </sbol:interaction>
</sbol:ModuleDefinition>
```

Cello UCF File Example

```
"collection": "parts",
  "type": "cds",
  "name": "AmtR".
  "dnasequence": "ATGGCAGGCGCAGTTGGTCGTCGCGTCGTAGTGCACCGCGTCGTGCAGGTAAAAATCCGCGTGAAGAAATTC..."
},...
  "collection": "gates".
  "regulator": "AmtR",
  "group name": "AmtR",
  "gate_name": "A1_AmtR",
  "gate type": "NOR",
  "system": "TetR",
  "color_hexcode": "3BA9E0"
  "collection": "gate_parts",
  "gate_name": "A1_AmtR",
  "expression cassettes": [
      "maps to variable": "x",
      "cassette parts": [
        "BydvJ",
        "A1".
        "AmtR".
        "L3S2P55"
  "promoter": "pAmtR"
}, ...
```

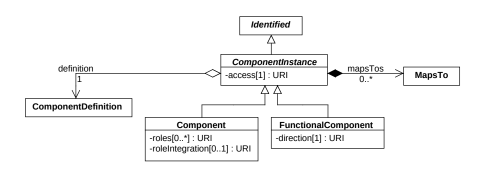
Converting Gate Parts to SBOL ComponentDefinitions

```
for (JSONObject gate : gate partsArr) {
 String gate_name = (String)gate.get("gate_name");
 ComponentDefinition componentDefinition =
   document.createComponentDefinition(gate name, version, ComponentDefinition.DNA);
 componentDefinition.setName(gate name);
 componentDefinition.addRole(SequenceOntology.ENGINEERED_REGION);
 componentDefinition.addWasGeneratedBy(activityURI);
 componentDefinition.createAnnotation(new QName(dcTermsNS, "created", "dcTerms"), createdDate);
 componentDefinition.createAnnotation(new OName(celloNS, "family", "cello"),
    (String) gatesMap.get(gate_name).get("system"));
 componentDefinition.createAnnotation(new QName(celloNS, "gate_type", "cello"),
    (String) gatesMap.get(gate_name).get("gate_type"));
 componentDefinition.createAnnotation(new QName(celloNS, "group name", "cello"),
    (String) gatesMap.get(gate_name).get("group_name"));
 componentDefinition.createAnnotation(new QName(celloNS, "color hexcode", "cello"),
    (String) gatesMap.get(gate name).get("color hexcode"));
```

Adding Response Functions to Gate Parts

```
componentDefinition.createAnnotation(new QName(celloNS, "response_function", "cello"),
   (String) responseMap.get(gate_name).get("equation"));
JSONArray parameters = (JSONArray) responseMap.get(gate_name).get("parameters");
for (Object obj : parameters) {
   String name = (String)((JSONObject)obj).get("name");
   componentDefinition.createAnnotation(new QName(celloNS, name, "cello"),
   (Double)((JSONObject)obj).get("value"));
}
JSONArray variables = (JSONArray) responseMap.get(gate_name).get("variables");
for (Object obj : variables) {
   String name = (String)((JSONObject)obj).get("name");
   componentDefinition.createAnnotation(new QName(celloNS, name+"_off_threshold", "cello"),
   (Double)((JSONObject)obj).get("off_threshold"));
   componentDefinition.createAnnotation(new QName(celloNS, name+"_on_threshold", "cello"),
   (Double)((JSONObject)obj).get("on_threshold"));
```

Component (Child of ComponentDefinition)



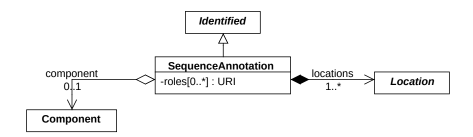
Access URI

http://sbols.org/v2#public http://sbols.org/v2#private

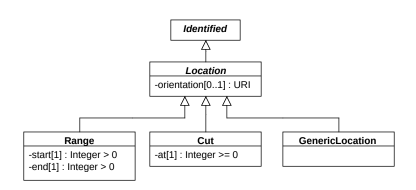
Description

MAY be referred to by remote **MapsTo** objects. MUST NOT be referred to by remote **MapsTo** objects.

SequenceAnnotation (Child of ComponentDefinition)



Location (Child of SequenceAnnotation)



Orientation URI

http://sbols.org/v2#inline http://sbols.org/v2#reverseComplement

Description

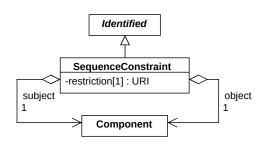
The region is inline with the sequence.

The region is on the reverse-complement translation.

Creating Composite Gates Using SequenceAnnotations

```
JSONArray expression_cassettes = (JSONArray) gate.get("expression_cassettes");
String seg = "";
for (Object obj : expression_cassettes) {
 int annotationCount = 0;
 int start = 1:
 JSONObject expression cassette = (JSONObject) obj;
 JSONArray cassette parts = (JSONArray)expression cassette.get("cassette parts");
 for (Object obj2 : cassette parts) {
    String partId = (String)obj2;
    ComponentDefinition partComponentDefinition =
     document.getComponentDefinition(partId, version);
    String cass_seg = document.getSequence(partId+"_sequence",version).getElements();
    seq += cass seq;
    componentDefinition.createComponent(partId, AccessType.PUBLIC, partId, version);
    SequenceAnnotation sa =
     componentDefinition.createSequenceAnnotation("annotation"+annotationCount,
     "range", start, start + cass_seq.length() - 1, OrientationType.INLINE);
    sa.setComponent(partId);
    start += cass_seq.length();
    annotationCount ++:
```

SequenceConstraint (Child of ComponentDefinition)



Restriction URI

http://sbols.org/v2#precedes http://sbols.org/v2#sameOrientationAs http://sbols.org/v2#oppositeOrientationAs

Description

subject MUST precede object Component. subject & object MUST have same orientation. subject & object MUST have opposite orientations.

Creating Composite Gates Using SequenceConstraints

```
JSONArray expression_cassettes = (JSONArray) gate.get("expression_cassettes");
for (Object obj : expression_cassettes) {
 int constraintCount = 0;
 Component previousComponent = null:
 Component currentComponent = null:
 JSONObject expression cassette = (JSONObject) obj;
 JSONArray cassette_parts = (JSONArray)expression_cassette.get("cassette_parts");
 for (Object obj2 : cassette parts) {
    String partId = (String)obj2;
    ComponentDefinition partComponentDefinition =
     document.getComponentDefinition(partId, version);
    currentComponent = componentDefinition.createComponent(partId, AccessType.PUBLIC,
     partId, version);
    if (previousComponent != null) {
     componentDefinition.createSequenceConstraint("constraint"+constraintCount,
        RestrictionType.PRECEDES,
        previousComponent.getIdentity(), currentComponent.getIdentity());
     constraintCount ++:
    previousComponent = currentComponent;
```

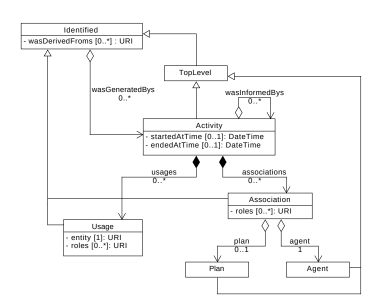
Creating ModuleDefinitions for Inhibition

```
if (partComponentDefinition.getRoles().contains(SequenceOntology.CDS)) {
 String promoter = (String)gate.get("promoter");
 if (document.getModuleDefinition(partId+"_"+promoter+"_repression", version) == null) {
   ModuleDefinition moduleDefinition =
     document.createModuleDefinition(partId+"_"+promoter+"_repression", version);
    moduleDefinition.addWasGeneratedBy(activityURI);
    moduleDefinition.createAnnotation(new QName(dcTermsNS, "created", "dcTerms"), createdDate);
    Interaction interaction =
     moduleDefinition.createInteraction(partId+" "+promoter+" repression",
     SystemsBiologyOntology.INHIBITION);
    interaction.createParticipation(partId+"_protein_participation", partId+"_protein",
     SystemsBiologyOntology.INHIBITOR);
    interaction.createParticipation(promoter+"_promoter_participation", promoter,
     SystemsBiologyOntology.INHIBITED):
```

Creating the Composite Sequence

```
Sequence sequence = document.createSequence(gate_name+"_sequence", version, seq,
    Sequence.IUPAC_DNA);
sequence.setName(gate_name+"_sequence");
sequence.addWasDerivedFrom(derivedFrom);
sequence.addWasGeneratedBy(activityURI);
sequence.createAnnotation(new QName(dcTermsNS,"created","dcTerms"), createdDate);
componentDefinition.addSequence(sequence);
```

Provenance Ontology Classes



Creating the CelloUCF2SBOL Agent

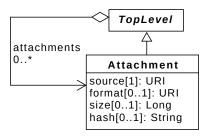
agent.setDescription("A_script_to_convert_Cello_UCF_parts_and_metadata_to_SBOL_2_documents.");
agent.createAnnotation(new OName(dcNS, "source", "dc"), URI.create("https://github.com/MyersResearch

agent.createAnnotation(new QName(dcNS,"creator","dc"), "Prashant_Vaidyanathan");
agent.createAnnotation(new OName(dcNS,"creator","dc"), "Chris J. Myers");

Agent agent = document.createAgent("CelloUCF2SBOL", version);

agent.setName("Cello_UCF_to_SBOL");

Attachment



Creating the UCF File Attachment

<sbol:source rdf:resource="https://github.com/MyersResearchGroup/UCF2SBOL/blob/master/UCF2SBOL/s</pre>

Attachment attachment = document.createAttachment("Eco1C1G1T1_UCF", version, URI.create("https://github.com/MyersResearchGroup/UCF2SBOL/blob/master/" +

<sbol:persistentIdentity rdf:resource="http://cellocad.org/EcolC1G1T1_UCF"/>

"UCF2SB0L/src/main/resources/Eco1C1G1T1.UCF.json"));

<sbol:Attachment rdf:about="http://cellocad.org/Eco1C1G1T1 UCF/1">

<sbol:displayId>Eco1C1G1T1 UCF</sbol:displayId>

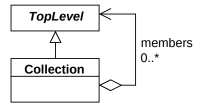
<sbol:version>1</sbol:version>

</sbol:Attachment>

Creating the CelloUCF2SBOL Actvity

```
Activity activity = document.createActivity("CelloUCF2sbol Activity", version);
activity.setName("Cello_UCF.to_SBOL_conversion");
activity.setDescription("Conversion.of.the.Cello.UCF.parts.and.metadata.to.SBOL.2.documents.");
activity.setEndedAtTime(DateTime.now());
activityURI = activity.getIdentity();
activity.createAssociation("association", agent.getIdentity());
activity.createUsage("UCF file", attachment.getIdentity());
<sbol:persistentIdentity rdf:resource="http://cellocad.org/CelloUCF2sbol Activity"/>
   <sbol:displayId>CelloUCF2sbol_Activity</sbol:displayId>
   <sbol:version>1</sbol:version>
   <dcterms:title> Cello UCF to SBOL conversion/dcterms:title>
   <dcterms:description> Conversion of the Cello UCF parts and metadata to SBOL 2 documents.
   <sbol:persistentIdentity rdf:resource="http://cellocad.org/CelloUCF2sbol_Activity/associatio">
           <sbol:displayId>association</sbol:displayId>
           <sbol:version>1</sbol:version>
           </prov:qualifiedAssociation>
       corov:gualifiedUsage>
           < "delay to the content of the conte
               <sbol:persistentIdentity rdf:resource="http://cellocad.org/CelloUCF2sbol Activity/UCF File</pre>
               <sbol:displayId>UCF File</sbol:displayId>
               <sbol:version>1</sbol:version>
```

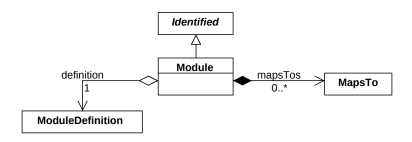
Collection



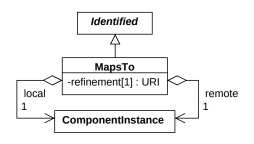
Creating CDS Part Collection

```
Collection cdsCollection = document.createCollection("cdsCollection", version);
for (ComponentDefinition cd : document.getComponentDefinitions()) {
  if (cd.containsRole(SequenceOntology.CDS)) {
    cdsCollection.addMember(cd.getIdentity());
<sbol:Collection rdf:about="http://cellocad.org/cdsCollection/1">
  <sbol:persistentIdentity rdf:resource="http://cellocad.org/cdsCollection"/>
  <sbol:displayId> cdsCollection </sbol:displayId>
  <sbol:version> 1 </sbol:version>
  <sbol:member rdf:resource="http://cellocad.org/SrpR/1"/>
  <sbol:member rdf:resource="http://cellocad.org/AmeR/1"/>
  <sbol:member rdf:resource="http://cellocad.org/HlvIIR/1"/>
  <sbol:member rdf:resource="http://cellocad.org/AmtR/1"/>
  <sbol:member rdf:resource="http://cellocad.org/PhlF/1"/>
  <sbol:member rdf:resource="http://cellocad.org/BM3R1/1"/>
  <sbol:member rdf:resource="http://cellocad.org/BetI/1"/>
</shol:Collection>
```

Module (Child of ModuleDefinition)



MapsTo (Child of ComponentInstance)



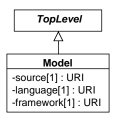
Refinement URI

http://sbols.org/v2#useRemote http://sbols.org/v2#useLocal http://sbols.org/v2#verifyIdentical http://sbols.org/v2#merge

Description

All references MUST dereference to the *remote* CI.
All references MUST dereference to the *local* CI.
The *definition* of the *local* and *remote* MUST be same CD.
All references MUST dereference to both objects.

Model



- source URI reference to the source file for the model.
- language URI that species language in which the model is implemented.
 Model Language URI for EDAM Term

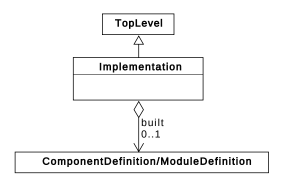
SBML http://identifiers.org/edam/format_2585 CellML http://identifiers.org/edam/format_3240 BioPAX http://identifiers.org/edam/format_3156

• framework - URI that species modeling framework used.

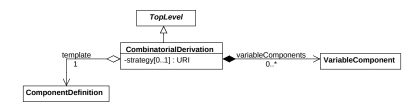
Framework URI for SBO Term

Continuous http://identifiers.org/biomodels.sbo/SBO:0000062
Discrete http://identifiers.org/biomodels.sbo/SBO:0000063

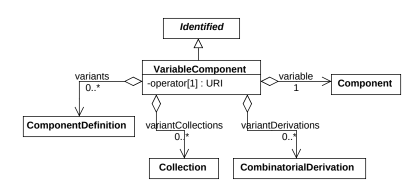
Implementation



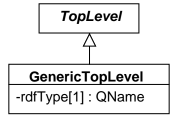
CombinatorialDerivation



VariableComponent



Generic TopLevel



GenericTopLevel Objects and Annotations

- Software tools that need to store data that is not encoded within SBOL can do so using GenericTopLevel objects and custom Annotations.
- When the library reader encounters a tag for a TopLevel object that it does not recognize, this data is stored within a GenericTopLevel object.
- Within TopLevel objects, when a tag is not recognized the data is stored within a custom Annotation object.
- Tools using our library that do not recognize this data will round-trip it unmodified when writing an SBOL file.
- Tools that would like to make use of this data can interpret and manipulate the raw data, which is stored in a tree-like data structure.

Serialization Methods

SBOLWriter class:

```
SBOLWriter.setKeepGoing(true);
SBOLWriter.write(document, String OR File OR OutputStream);
SBOLWriter.write(document, String OR File OR OutputStream, fileType);
fileType = SBOLDocument.RDF, RDFV1, GEMBANK, FASTAformat
document.write(String OR File OR OutputStream, fileType);
SBOLWriter.clearErrors();
SBOLWriter.getNumErrors();
SBOLWriter.getErrors();
```

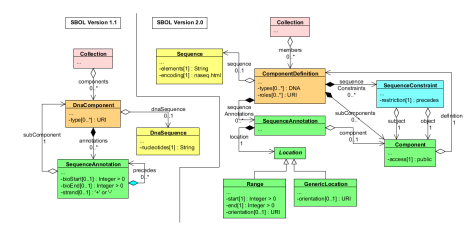
SBOLReader class:

```
SBOLReader.setURIPrefix("http://cellocad.org/");
SBOLReader.setVersion("!");
SBOLReader.setKeepGoing(true);

SBOLReader.getSBOLVersion(String OR File OR InputStream);
SBOLDocument document = SBOLReader.read(String OR File OR InputStream);
document.read(String OR File OR InputStream);
SBOLReader.clearErrors();
SBOLReader.getErrors();
SBOLReader.getErrors();
```

Conversion

- Supports conversion to/from GenBank and FASTA formats.
- Reads/Writes SBOL 1.1 data files.



Validation

Many validation rules checked when documents are read.

```
SBOLReader.setKeepGoing(true);
SBOLDocument document = SBOLReader.read(file);
if (SBOLReader.getNumErrors() > 0) {
    for(String error : SBOLReader.getErrors()) {
        System.out.println(error);
    }
}
```

• Remaining validation rules checked by validateSBOL method.

Method to compare to SBOLDocuments.

```
SBOLValidate.compareDocuments (file, document, file2, document2);
```

SynBioHub Repository Support

Uploading SBOL to SynBioHub:

Other SynBioHub functions:

```
public String getAttachment(URI attachmentURI, String path)
public String getAttachment(URI attachmentURI, OutputStream outputStream)
public int getCount(String objectType)
public ArrayList
JidentifiedMetadata> getRootCollectionMetadata()
public ArrayList
JidentifiedMetadata> getSubCollectionMetadata(URI parentCollectionUri)
public ArrayList
JidentifiedMetadata> getMatchingComponentDefinitionMetadata(String name,
Set<URI> roles, Set<URI> types, Set<URI> collections, Integer offset, Integer limit)
public ArrayList
JidentifiedMetadata> search(SearchQuery query)
public String sparqlQuery(String query)
```

More Information

- libSBOLj is open source under the Apache 2.0 License.
- More information: http://sbolstandard.org/libsbol/.
 - Current snapshot (2.3.2-SNAPSHOT) on GitHub and Sonatype.
 - Latest release (Version 2.3.1) on GitHub and Maven.
 - Issue tracker for reporting bugs and feature requests.
 - JavaDocs for all public methods.
 - A brief getting started tutorial.
 - A detailed code example and sample project for a CRISPR circuit.
 - Several example code files.

sboljs - Javascript SBOL library

- Developed by James McLaughlin at Newcastle University and myself.
- Prerequisites: node.js and npm.
- Install: npm install sboljs

Creating an SBOL Document

```
var SBOLDocument = require('sboljs');
...
var sbol = new SBOLDocument();
```

Creating a GenericTopLevel

```
var actVersion = 1;
var actDisplayId = 'cello2sbol';
var actPersistantIdentity = urlprefix + actDisplayId;
var activityURI = actPersistantIdentity + '/' + actVersion;

const activity = sbol.genericTopLevel(activityURI, provNS + 'Activity');
activity.version = actVersion;
activity displayId = actDisplayId;
activity.name = 'Cello UCF to SBOL conversion';
activity.description = 'Conversion of the Cello UCF parts and metadata to SBOL2';
activity.addStringAnnotation(dcNS + 'creator', 'Prashant Vaidyanathan');
activity.addStringAnnotation(dcNS + 'creator', 'Chris J. Myers');
// TODO: Need to change to PROV endedAtTime
activity.addStringAnnotation('http://purl.org/dc/terms/created', today.toISOString() + '');
activity.persistentIdentity = actPersistantIdentity;
activity.uri = activityURI;
```

Parsing Cello UCF JSON File

```
var ucf = JSON.parse(fs.readFileSync(ucfFilepath) + '');
var gate partsArr = [];
var partsMap = {};
var partsSBOL = {}
var gatesMap = {};
//Go through the UCF file to store collections in local data structures (arrays and maps)
ucf.forEach(function (collection) {
    switch (collection.collection) {
        case 'parts':
            //partsArr.push(collection);
            partsMap[collection.name] = collection;
            break:
        case 'gate parts':
            gate_partsArr.push(collection);
            break:
       case 'gates':
            gatesMap[collection.gate name] = collection;
            break:
}, this);
```

Converting Parts to SBOL (1)

```
Object.keys(partsMap).forEach(function (partkey) {
    var part = partsMap[partkey];
    var partName = part.name;
    if (!(partName in partsSBOL)) {
        const componentDefinition = sbol.componentDefinition()
        componentDefinition.version = version;
        componentDefinition.displayId = part.name;
        componentDefinition.name = part.name;
        componentDefinition.persistentIdentity = urlprefix + componentDefinition.displayId;
        componentDefinition.uri = componentDefinition.persistentIdentity + '/' +
          componentDefinition.version;
        componentDefinition.wasDerivedFrom = derivedFrom;
        componentDefinition.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
        componentDefinition.addRole(getPartType(part.type));
        componentDefinition.addType(SBOLDocument.terms.dnaRegion);
        const sequence = sbol.sequence()
        sequence.displayId = part.name + ' sequence';
        sequence.name = part.name + '_sequence';
        sequence.version = version;
        sequence.elements = part.dnasequence;
        sequence.persistentIdentity = urlprefix + sequence.displayId;
        sequence.uri = sequence.persistentIdentity + '/' + sequence.version;
        sequence.wasDerivedFrom = derivedFrom;
        sequence.encoding = SBOLDocument.terms.dnaSequence;
        sequence.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
        componentDefinition.addSequence(sequence)
```

Converting Parts to SBOL (2)

```
// Create a Protein
if (part.type === 'cds') {
  const proteinComponentDefinition = sbol.componentDefinition();
  proteinComponentDefinition.version = version;
  proteinComponentDefinition.displayId = componentDefinition.displayId + ' protein';
  proteinComponentDefinition.name = componentDefinition.displayId + ' protein':
  proteinComponentDefinition.persistentIdentity = urlprefix +
    proteinComponentDefinition.displayId;
  proteinComponentDefinition.uri = proteinComponentDefinition.persistentIdentity + '/' +
    proteinComponentDefinition.version;
  proteinComponentDefinition.addType(SBOLDocument.terms.protein);
  proteinComponentDefinition.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
  partsSBOL[partName + '_protein'] = proteinComponentDefinition.uri;
  const moduleDefinition = shol.moduleDefinition():
  moduleDefinition.name = proteinComponentDefinition.displayId + ' production';
  moduleDefinition.version = version;
  moduleDefinition.displayId = proteinComponentDefinition.displayId + ' production';
  moduleDefinition.persistentIdentity = urlprefix + moduleDefinition.displayId;
  moduleDefinition.uri = moduleDefinition.persistentIdentity + '/' + moduleDefinition.version;
  moduleDefinition.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
```

Converting Parts to SBOL (4)

```
//Functional Component for Coding sequence
const functionalComponentCDS = sbol.functionalComponent();
functionalComponentCDS.version = version;
functionalComponentCDS.displayId = componentDefinition.displayId + '_functionalComponent';
functionalComponentCDS.name = componentDefinition.name + ' functionalComponent';
functionalComponentCDS.persistentIdentity = moduleDefinition.persistentIdentity + '/' +
  functionalComponentCDS.displayId;
functionalComponentCDS.uri = functionalComponentCDS.persistentIdentity + '/' +
  functionalComponentCDS.version;
functionalComponentCDS.definition = componentDefinition;
//Functional Component for Protein
const functionalComponentProt = sbol.functionalComponent();
functionalComponentProt.version = version;
functionalComponentProt.displayId = proteinComponentDefinition.displayId + ' functionalComponent';
functionalComponentProt.name = proteinComponentDefinition.name + ' functionalComponent';
functionalComponentProt.persistentIdentity = moduleDefinition.persistentIdentity + '/' +
  functionalComponentProt.displayId;
functionalComponentProt.uri = functionalComponentProt.persistentIdentity + '/' +
  functionalComponentProt.version;
functionalComponentProt.definition = proteinComponentDefinition;
```

Converting Parts to SBOL (5)

```
//CDS to Protein interaction
const interaction = sbol.interaction();
interaction.displayId = proteinComponentDefinition.displayId + ' interaction';
interaction.name = interaction.displayId;
interaction.version = version;
interaction.persistentIdentity = moduleDefinition.persistentIdentity + '/' +
 interaction.displayId;
interaction.uri = interaction.persistentIdentity + '/' + interaction.version;
interaction.addType(productionSBO);
const participationCDS = sbol.participation();
participationCDS.version = version;
participationCDS.name = componentDefinition.displayId + ' participation';
participationCDS.displayId = componentDefinition.displayId + ' participation';
participationCDS.persistentIdentity = interaction.persistentIdentity + '/' +
  participationCDS.displayId;
participationCDS.uri = participationCDS.persistentIdentity + '/' + participationCDS.version;
participationCDS.addRole(templateSBO);
participationCDS.participant = functionalComponentCDS;
const participationProt = sbol.participation();
participationProt.version = version;
participationProt.name = proteinComponentDefinition.displayId + ' participation';
participationProt.displayId = proteinComponentDefinition.displayId + '_participation';
participationProt.persistentIdentity = interaction.persistentIdentity + '/' +
  participationProt.displayId;
participationProt.uri = participationProt.persistentIdentity + '/' + participationProt.version;
participationProt.addRole(productSBO);
participationProt.participant = functionalComponentProt;
interaction.addParticipation(participationCDS);
interaction.addParticipation(participationProt);
```

Converting Parts to SBOL (6)

```
moduleDefinition.addFunctionalComponent(functionalComponentCDS);
    moduleDefinition.addFunctionalComponent(functionalComponentProt);
    moduleDefinition.addInteraction(interaction);
}
componentDefinition.addStringAnnotation('http://purl.org/dc/terms/created',
    datecreated.toISOString() + '');
    var uriVal = componentDefinition.uri;
    partsSBOL[partName] = uriVal;
}
}, this);
```

Converting Gate Parts to SBOL (1)

```
gate_partsArr.forEach(function (gpart) {
    var gpartName = gpart.gate_name;
    const componentDefinition = sbol.componentDefinition();
    componentDefinition.version = version;
    componentDefinition.displayId = gpartName;
    componentDefinition.name = gpartName;
    componentDefinition.persistentIdentity = urlprefix + componentDefinition.displayId;
    componentDefinition.uri = componentDefinition.persistentIdentity + '/' +
        componentDefinition.version;
    componentDefinition.wasDerivedFrom = derivedFrom;
    componentDefinition.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
    componentDefinition.addUriAnnotation(tetR_family, gatesMap[gpartName].system);
    componentDefinition.addUriAnnotation(group_name, gatesMap[gpartName].group_name);
    componentDefinition.addUriAnnotation(gate_type, gatesMap[gpartName].gate_type);
    componentDefinition.addUriAnnotation(color_hexcode, gatesMap[gpartName].color_hexcode);
```

Converting Gate Parts to SBOL (2)

```
gpart.expression_cassettes.forEach(function (expression_cassettesArr) {
  var seq = "";
  var annotationCount = 0;
  var start = 1;

  expression_cassettesArr.cassette_parts.forEach(function (cassette) {
    const component = sbol.component();
    component.version = version;
    component.displayId = cassette;
    component.name = cassette;
    component.persistentIdentity = componentDefinition.persistentIdentity + '/' +
    component.uri = component.persistentIdentity + '/' + component.version;
    component.definition = sbol.lookupURI(partsSBOL[cassette]);
    componentDefinition.addComponent(component);
```

Converting Gate Parts to SBOL (3)

```
var cass seg = partsMap[cassette].dnasequence;
  seq += cass seq;
  const sa = sbol.sequenceAnnotation();
  sa.displayId = 'annotation' + annotationCount;
  annotationCount++:
  sa.name = cassette;
  sa.version = version;
  sa.persistentIdentity = componentDefinition.persistentIdentity + '/' + sa.displayId;
  sa.uri = sa.persistentIdentity + '/' + sa.version;
  sa.component = component;
  sa.description = partsMap[cassette].type;
  const range = sbol.range();
  range.displayId = 'range';
  range.persistentIdentity = sa.persistentIdentity + '/' + range.displayId;
  range.version = version;
  range.uri = range.persistentIdentity + '/' + range.version;
  range.start = start;
  var end = start + cass seg.length - 1;
  range.end = end;
  range.orientation = 'http://sbols.org/v2#inline';
  sa.addLocation(range);
  componentDefinition.addSequenceAnnotation(sa);
```

Converting Gate Parts to SBOL (4)

```
//Create Protein - Promoter repression Module definition
if (partsMap[cassette].type === 'cds') {
  if (!(cassette in moduleDefnMap)) {
     const moduleDefinition = sbol.moduleDefinition();
      moduleDefinition.version = version;
      moduleDefinition.name = cassette + ' ' + gpart.promoter + ' repression';
      moduleDefinition.displayId = cassette + '_' + gpart.promoter + '_repression';
      moduleDefinition.persistentIdentity = urlprefix + moduleDefinition.displayId;
      moduleDefinition.uri = moduleDefinition.persistentIdentity + '/' +
        moduleDefinition.version:
      moduleDefinition.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
      const functionalComponentCDS = sbol.functionalComponent();
      functionalComponentCDS.version = version;
      functionalComponentCDS.name = cassette + '_protein_functionalComponent';
      functionalComponentCDS.displayId = cassette + '_protein_functionalComponent';
      functionalComponentCDS.persistentIdentity = moduleDefinition.persistentIdentity + '/' +
        functionalComponentCDS.displayId;
      functionalComponentCDS.uri = functionalComponentCDS.persistentIdentity + '/' +
        functionalComponentCDS.version;
      functionalComponentCDS.definition = sbol.lookupURI(partsSBOL[cassette + ' protein'l);
      const functionalComponentProm = sbol.functionalComponent();
      functionalComponentProm.version = version;
      functionalComponentProm.name = gpart.promoter + '_functionalComponent';
      functionalComponentProm.displayId = gpart.promoter + ' functionalComponent';
      functionalComponentProm.persistentIdentity = moduleDefinition.persistentIdentity + '/' +
        functionalComponentProm.displayId;
      functionalComponentProm.uri = functionalComponentProm.persistentIdentity + '/' +
        functionalComponentProm.version;
      functionalComponentProm.definition = sbol.lookupURI(partsSBOL[gpart.promoter]);
```

Converting Gate Parts to SBOL (5)

```
//Protein - Promoter Interaction
const interaction = sbol.interaction();
interaction.version = version;
interaction.name = cassette + '_' + qpart.promoter + '_interaction';
interaction.displayId = cassette + '_' + qpart.promoter + '_interaction';
interaction.persistentIdentity = moduleDefinition.persistentIdentity + '/' +
 interaction.displayId:
interaction.uri = interaction.persistentIdentity + '/' + interaction.version;
interaction.addType(inhibitionSO);
//Protein participation
const participantProt = sbol.participation();
participantProt.version = version;
participantProt.name = cassette + '_protein_participation';
participantProt.displayId = cassette + '_protein_participation';
participantProt.persistentIdentity = interaction.persistentIdentity + '/' +
  participantProt.displayId;
participantProt.uri = participantProt.persistentIdentity + '/' + participantProt.version;
participantProt.addRole(inhibitorSO);
participantProt.participant = functionalComponentCDS;
//Promoter participation
const participantProm = sbol.participation();
participantProm.version = version;
participantProm.name = gpart.promoter + '_participation';
participantProm.displayId = gpart.promoter + ' participation';
participantProm.persistentIdentity = interaction.persistentIdentity + '/' +
  participantProm.displayId;
participantProm.uri = participantProm.persistentIdentity + '/' + participantProm.version;
participantProm.addRole(inhibitedSO);
participantProm.participant = functionalComponentProm;
```

Converting Gate Parts to SBOL (6)

```
interaction.addParticipation(participantProt);
interaction.addParticipation(participantProm);

moduleDefinition.addFunctionalComponent(functionalComponentCDS);
moduleDefinition.addTunctionalComponent(functionalComponentProm);
moduleDefinition.addInteraction(interaction);

moduleDefnMap[cassette] = gpart.promoter;
}
start += cass_seq.length;
});
```

Converting Gate Parts to SBOL (7)

```
const sequence = sbol.sequence()
    sequence.displayId = gpartName + ' sequence';
    sequence.name = gpartName + '_sequence';
    sequence.version = version;
    sequence.elements = seq;
    sequence.persistentIdentity = urlprefix + sequence.displayId;
    sequence.uri = sequence.persistentIdentity + '/' + sequence.version;
    sequence.wasDerivedFrom = derivedFrom;
    sequence, encoding = SBOLDocument, terms, dnaSequence;
    sequence.addUriAnnotation(provNS + 'wasGeneratedBy', activityURI);
    componentDefinition.addSequence(sequence);
 }, this);
 componentDefinition.addType(SBOLDocument.terms.dnaRegion);
 componentDefinition.addRole(gate_parts_so);
 componentDefinition.addStringAnnotation('http://purl.org/dc/terms/created', datecreated.toISOStr
}, this);
```

Writing the SBOL File

```
var fs = require('fs');
fs.writeFile(resultSBOL, sbol.serializeXML(), function (err) {
    if (err) {
        return console.log(err);
    }
    console.log("Cello_SBOL_file_created!");
});
```

Conclusion (SBOL Compliant Software)

- Can either support all classes or only a subset.
- Can support import of SBOL, export of SBOL, or both (lossy/lossless).
- SBOL Test Suite is currently under development here: https://github.com/SynBioDex/SBOLTestSuite
- Validate SBOL files generated using the SBOL Validator found here: http://www.async.ece.utah.edu/sbol-validator/
- Report SBOL-compliant software by filling out the survey found here: http://sbolstandard.org/applications/