**ShortBOL tutorial**

**Introduction**

Welcome to the ShortBOL Tutorial. ShortBOL is a scripting language, designed to be easy to use, powerful and extensible. ShortBOL is based around structured text to capture your ideas, and doesn't require any prior coding skills. When these scripts are run, they generate SBOL files which can then be used to derive the DNA sequences for your design from its parts, generate diagrams, and can be loaded into any SBOL-compliant computer-aided genome design tools.

This tutorial will get you up to speed in how to rapidly prototype synthetic biology designs with ShortBOL. It works through several steps to introduce the language, and give you practical experience using it to capture your designs. Our running example is a TetR/LacI toggle switch (see Gardner 2000). By the end of the tutorial, you will be able to represent the toggle switch structure and behaviour in ShortBOL and be able to run this script to generate an SBOL file that can then be used in any SBOL-compliant tooling. We then move on to develop a more complex example using CRISPR as described by Crispr transcriptional repression devices and layered circuits in mammalian cells.

**Downloading and installing ShortBOL**

1. Download or clone the ShortBOL repository:

git clone https://github.com/intbio-ncl/shortbol.git

1. Navigate to your install directory
2. Install dependencies with:  python setup.py install –user
3. Test the installation using the simple example provided

* simple\_example.rdfsh in the /examples folder is a design for a single promoter with its associated sequence
* Compile the simple\_example.rdfsh file with python run.py -s sbolxml examples/simple\_example.rdfsh -o <output-file>
* <output-file>  is the name of the desired SBOL XML-RDF file

**Designing a genetic toggle switch**

1. Adding basic parts

We're going to start by building the ShortBOL for the TetR inverter of the TetR/LacI toggle switch. The TetR inverter couples a tetracycline-repressed promoter with the *lacI* coding sequence, so that in the absence of tetracycline, LacI is produced. We are going to describe the design of the TetR inverter using ShortBOL. Create a text document containing the following:

pTetR\_prom is a Promoter()

lacI\_CDS is a CDS()

These two lines simply declare a promoter called pTetR, and declare a complement determining an open reading frame, or coding sequence (CDS) called lacI. Comments an be added to the script. Any line starting with a pound '#' character is treated as a comment, and ignored. Blank lines can also be added for formatting and are also ignored.

# Declare a promoter named pTetR

pTetR is a Promoter()

lacI\_CDS is a CDS()

2. Adding properties to SBOL components.

So far we have created a promoter and a CDS and named them. In ShortBOL, we call pTetR and lacI\_CDS instances. An instance is any thing that you have named as part of your description of your design. It may be a piece of DNA, or a large biological module, or a reference to a simulation, or perhaps a publication or co-worker. Instance names are case-sensitive, so pTetR and PTetR are different instances, as the case of their leading letter differs. You can choose any name you like for an instance. The name is there to refer to it within your script. However, by choosing meaningful names, you will make the script easier to read and understand.

With ShortBOL we can attach properties and values to instances. ShortBOL uses brackets to make lines ‘properties of' a containing instance. For example, we can add a human-readable description and comments to pTetR like this:

# Declare a promoter named pTetR

pTetR is a Promoter()

(

# give pTetR a description

description = "pTet promoter"

)

lacI\_CDS is a CDS()

Comments *are not* carried through to the final **SBOL** representation, so use them to document your script, for people who will read it in the future (probably you!) and may need some hints. Information needed to understand your design, rather than your script, needs to be added as properties like description, as these *are* available from an **SBOL** design.

You can add any names and values you like to an instance. It is perfectly fine for you to make new ones up as you need them. However, some names mean something special within the SBOL standard. You will frequently use these two SBOL properties for documenting your design:

* **description**: associates a human-readable descriptions with things. This can be an extended block of text, that tells us more about an instance.
* **name**: a human-readable name, possibly including spaces and special characters. For our *pTetR*, a good choice of name would be "pTetR".

**Exercise 1:**

Start with the provided skeleton script and modify it so that lacI has the name "lacI", description "LacI protein coding region. Don't forget your brackets around the property names. The answers can be found at the end of this document.

**Answer 1:**

# Declare a promoter named pTetR

pTetR is a Promoter()

(

# give pTetR a description

description = "pTet promoter"

)

lacI\_CDS is a CDS()

(

# Properties of lacI CDS

name = “lacI”

description = “LacI protein coding region”

)

3. Working with types.

In the previous example, we created instances to represent *pTetR* and *lacI* in the *TetR* inverter device, and gave them names, descriptions and displayIds. When we put it all together, that example looks like this:

# Declare a promoter named pTetR

pTetR is a Promoter()

(

# give pTetR a description

description = "pTetR promoter"

)

lacI\_CDS is a CDS()

(

# Properties of lacI CDS

name = “lacI”

description = “LacI protein coding region”

)

Let's look at this example again. It declares two instances, a Promoter called pTetR and a CDS called lacI. The Promoter and CDS are types. They say what sort of thing pTetR and LacI are. In ShortBOL, whenever you declare an instance, you construct it with a type. The name of the type is linked to the name of the instance with ‘is a’ to denote that the instance is a type of something. A type can be distinguished from an instance since it will have a “()” suffix which indicates the tyoe constructor. A constructor can be used to initialise values of properties in an instance when it is created. More about this later in the section on creating sequences.

SBOL provides a pallet of types that can be used in your designs for all the common types of genetic parts. Here are some of the ones you may use most frequently:

Promoter: A genomic region where transcription is initiated.

CDS: A complement determining sequence; a genomic region that encodes a protein.

Terminator: A genomic region that terminates transcription.

RBS: A ribosome binding region, where the ribosome will bind to a transcript.

Operator: A region where proteins bind to regulate transcription.

You can add any number of these genetic parts to your design. Just give them each a unique name within your script.

**Exercise 2:**

The *TetR* inverter is made of four parts. A promoter, RBS, CDS and terminator. Edit the design above to include additional instances for an RBS instance called lacI\_RBS and a Terminator instance called lac\_term.

**Answer 2:**

# Declare a promoter named pTetR

pTetR is a Promoter()

(

# give pTetR a description

description = "pTet promoter"

)

# Declare a CDS named pTetR

lacI\_CDS is a CDS()

(

# Properties of lacI CDS

name = “lacI”

description = “LacI protein coding region”

)

# Declare a RBS named lacI\_RBS

lacI\_RBS is a RBS()

(

name = “lacI\_RBS”

description = “RBS for the lacI CDS”

)

# Declare a terminator named lacI\_term

lacI\_term is a Terminator()

(

name = “lacI\_term”

description = “Terminator for the lacI CDS”

)

4. Adding sequences

Ultimately, when you build a genetic design, you need the corresponding DNA sequence. Each individual genetic part in your design will have its own sequence, and the sequence of the whole design is composed from these. ShortBOL has a type called **DnaSequence** that lets you specify a DNA sequence, and a property **sequence** that lets you associate this with an instance representing a genetic part. e.g.

lacITSeq is a DnaSequence ("ttcagccaaaaaacttaagaccgccggtct

tgtccactaccttgcagtaatgcggtggacaggatcggcggttttcttttctcttctcaa")

Here we have constructed a DnaSequence named lacITSeq, and rather than setting a property, the DNA sequence string is passed into the DnaSequence constructor. ShortBOL instances are often created by giving the type constructor some values to work with. The constructor will use these to set up properties for you.

Now that we know how to make a sequence, we need to attach it to the corresponding part. This is done in the same way that we set the name, description and displayId for the parts earlier. SBOL defines a property called sequence that links from a genetic part back to the sequence it has. This time, rather than quoting the value, we use the naked value. This tells ShortBOL that we are linking to another instance, rather than capturing some text. Instances are always linked by the name that their ShortBOL instance was declared with, rather than by the value of their name, or any other data property.

lacITSeq is a DnaSequence ("ttcagccaaaaaacttaagaccgccggtct

tgtccactaccttgcagtaatgcggtggacaggatcggcggttttcttttctcttctcaa")

lacIT is a Terminator()

(

sequence = lacITSeq

)

**Exercise 3:** Edit the shortbol above to also include a new promoter pTetR with its own sequence

**Answer 3:**

lacITSeq is a DnaSequence ("ttcagccaaaaaacttaagaccgccggtct

tgtccactaccttgcagtaatgcggtggacaggatcggcggttttcttttctcttctcaa")

pTetRSeq is a DnaSequence (“tccctatcagtgatagagattgacatccctatcagtgatagagatactgagcac”)

)

lacIT is a Terminator()

(

sequence = lacITSeq

)

pTetR is a promoter()

(

sequence = pTetRSeq

)

**Designing a CRISPR logic gate**

1. Composition

A core principle of synthetic biology design is that larger designs are built up from smaller, well-validated components. This paradigm is exemplified by [BioBricks](http://biobricks.org/" \t "_blank" \o "The BioBricks Foundation), an assembly standard and parts registry of genomic parts. The SBOL data standard provides a lot of tooling for describing how a design is composed.

In this tutorial, we are going to look at several strategies for using ShortBOL to compose a larger design from smaller ones, by building up the *TetR* inverter from its component parts using the approach specified in the SBOL specification document.

In the tutorial exercise 2 above, we made instances for the four parts of the *TetR* inverter device. However, we stopped short of assembling them into a composite device. The SBOL type for a composite DNA device is a type of Component  called a DnaComponent. Components are used to compose objects into a structural hierarchy of a DnaComponent

To place the genetic parts we've made within a larger DnaComponent, we create a Component from each DnaComponent to be composed and then send each of these SubComponent the SubComponent property of DnaComponent as shown below in example 1:

**Example 1**

# The genetic parts of the TetR inverter

pTetR is a Promoter()

lacI\_RBS is a RBS()

lacI\_CDS is a CDS()

lacI\_term is a Terminator()

# The composite device for the TetR inverter

tetRInverter is a DnaComponent()

(

# include the child components

hasSubComponent(pTetR\_c)

hasSubComponent(lacI\_RBS\_c)

hasSubComponent(lacI\_CDS\_c)

hasSubComponent(lacI\_term\_c)

)

Because we are adding four sub-components, we set the component property four times. When you assign to a property multiple times, you add new values rather than over-writing previous ones.

We have built a *pTetR* inverter device that contains its four genetic parts as sub-components. However, we haven't specified anything about how these parts are to be assembled. There are two complementary ways to specify this. Firstly, we can attach constraints on their relative positions. Secondly, we can say exactly where the sub-components are located within the composite component.

1. Composition using constraints and locations

Constraints

In this section we are going to explore constraints. Sequence constraints are declared using the sequenceConstraint property. The values of this property are sequenceConstraint instances. In this version of ShortBOL (v1.0) we are true to the SBOL data model and so there is a bit of setting up to do.

SBOL currently defines three types of constraints. These are precedes, sameOrientationAs and differentOrientationAs. These last two tell you if the two components share the same orientation or have different orientations, but not what the orientation of either component is.

The constraint we need in this design is precedes. This says that one component comes before the other in the design. In this way, we can place the genetic parts, left-to-right. In order to do this we need to create a precedes relationship for pairs of Component sand then include them in a Component to form the correct ordering as shown below:

**Example 2**

# Example2

# The genetic parts of the TetR inverter

pTetR is a Promoter()

lacI\_RBS is a RBS()

lacI\_CDS is a CDS()

lacI\_term is a Terminator()

# The composite device for the TetR inverter

tetRInverter is a FunctionalEntity()

(

# relative positions of child components

precedes(pTetR, lacI\_RBS)

precedes(lacI\_RBS, lacI\_CDS)

precedes(lacI\_CDS, lacI\_term)

)

Locations and ranges

In the previous section we saw how ShortBOL can describe the relative positions of children within a parent design. Here we will see how it can give them exact positions. Below we state that each subcomponents location is the length of the full sequence.

**Example 3.**

# Example 3

# The genetic parts of the TetR inverter

pTetR is a Promoter()

(

hasDNASequence(pTetR\_seq)

)

lacI\_RBS is a RBS()

(

hasDNASequence(lacI\_RBS\_seq)

)

lacI\_CDS is a CDS()

(

hasDNASequence(lacI\_CDS\_seq)

)

lacI\_term is a Terminator()

(

hasDNASequence(lacI\_term\_seq)

)

pTetR\_c is a SubComponent(pTetR)

(

hasEntireSequence(pTetR\_seq)

)

lacI\_RBS\_c is a SubComponent(lacI\_RBS)

(

hasEntireSequence(lacI\_RBS\_seq)

)

lacI\_CDS\_c is a SubComponent(lacI\_CDS)

(

hasEntireSequence(lacI\_CDS\_seq)

)

lacI\_term\_c is a SubComponent(lacI\_term)

(

hasEntireSequence(lacI\_term\_seq)

)

tetRInverter is a DNAComponent()

(

# include the child components

hasFeature = pTetR\_c

hasFeature = lacI\_RBS\_c

hasFeature = lacI\_CDS\_c

hasFeature = lacI\_term\_c

)

**Exercise 4:** In the previous *pTetR* inverter positions example, we specified the positions of the four parts as the full sequence. However, instead we could specify only sub-parts of the sequence by a specific index (Start and End position).

**Answer 4:**

# Answer 4

# The genetic parts of the TetR inverter

pTetR is a Promoter()

(

hasDNASequence("tccctatcagtggtgatagagatactgagcac")

)

lacI\_RBS is a RBS()

(

hasDNASequence("aaggaggtg")

)

lacI\_CDS is a CDS()

(

hasDNASequence("gtgaaaccagtaacgttatacga")

)

lacI\_term is a Terminator()

(

hasDNASequence("ttcagccaaaaaacctcaa")

)

#Build components for each of the DNAComponents

pTetR\_c is a SubComponent(pTetR)

(

hasInlineRange(pTetR\_seq,1,55)

)

lacI\_RBS\_c is a SubComponent(lacI\_RBS)

(

hasInlineRange(lacI\_RBS\_seq,56,68)

)

lacI\_CDS\_c is a SubComponent(lacI\_CDS)

(

hasInlineRange(lacI\_CDS\_seq,169,1197)

)

lacI\_term\_c is a SubComponent(lacI\_term)

(

hasInlineRange(lacI\_term\_seq,1197,1240)

)

tetRInverter is a DNAComponent()

(

# include the child components

hasFeature = pTetR\_c

hasFeature = lacI\_RBS\_c

hasFeature = lacI\_CDS\_c

hasFeature = lacI\_term\_c

)

**Interactions**

In the functional design of the TetR inverter, the TetR protein represses the expression of the LacI protein. We add an interaction to say that TetR represses LacI. Here we are true to the SBOL data model and so we need to create SubComponents which in turn become Participants in an Interaction as described below in Example 4.

**Example 4**

# Answer 5

# The TetR and LacI proteins

TetR is a Protein()

LacI is a Protein()

# The LacI inverter module

LacI\_inverter is a FunctionalEntity()

(

description = "LacI inverter"

inhibition(TetR,LacI)

)

**Exercise 5:** The LacI inverter is very similar, but in this module LacI represses TetR. Write a script to include this interaction.

**Answer 5:**

# Answer 5

# The TetR and LacI proteins

TetR is a Protein()

LacI is a Protein()

# The LacI inverter module

LacI\_inverter is a FunctionalEntity()

(

description = "LacI inverter"

inhibition(TetR, LacI)

inhibition(LacI, TetR)

)

**Composing Modules**

In the previous section, we have built two modules, one for TetR inverter and one for the LacI inverter. The next step is to combine these into a toggle-switch module.

This composite module contains all of the behaviour of both the TetR and LacI inverter modules. However, at the moment both of the inverters are 'black box', with completely independent behaviour. What we want to do is glue them together, so that they are using the same pool of TetR and LacI molecules. This will cause them to repress one-another, flip-flopping between repressing TetR levels and LacI levels.

To achieve this, we need to wire components in the sub-modules. This is done using the references property.

We are essentially wiring together the FunctionalEntities by referencing together TetR and LacI.

# Example 4

# The TetR and LacI proteins

TetR is a ProteinComponent()

LacI is a ProteinComponent()

# The TetR inverter module

TetR\_inverter is a FunctionalEntity()

(

description = "TetR inverter"

inhibition(TetR,LacI)

)

# The LacI inverter module

LacI\_inverter is a FunctionalEntity()

(

description = "LacI inverter"

inhibition(TetR,LacI)

references(LacI)

references(TetR)

)