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# thinknode™ Examples

These examples provide a starting point for issuing http connections and requests to the dosimetry app on the thinknode™ framework. They are provided as is, and are written in python. Any further dependencies are listed along with the provided scripts.

# **Python**

Please refer to the Python Script Library Readme for dependent python modules and a high level list of what these libraries include.

# **Python: Overview**

The provided python scripts and libraries are meant to be a foundation and starting point for using the astroid apps on the thinknode<sup>m</sup> framework. The provided scripts outline the basics of using ISS to store objects, as well as constructing and making calculation requests to the calculation provider. The below sections detail the basic usage for each script.

**Download:** The python astroid\_script\_library can be downloaded from the .decimal GitHub repository.

# thinknode.cfg

There is a simple configuration file (thinknode.cfg) that is used to store user data for connecting to the astroid app on the thinknode™ framework. This file is required by all scripts in the python astroid\_script\_library to authenticate and use the app. A sample file with no user data is available in the repository and the details of the information to include in the file are provided below.

- basic\_user being a base64 encoded username and password. Refer to the thinknode documentation for more information.
- api url being the connection string to the thinknode™ framework.
- apps
  - app\_name being the current app name (e.g. dosimetry or dicom).
    - app\_version being the current version of the app existing on the thinknode™
      framework being used. If left blank the thinknode\_worker will select the first app's
      version returned by the Realm Versions GET request.
    - branch name not currently implemented
- realm name thinknode realm
- account\_name thinknode account name

### thinknode.cfg

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```
{
    "basic user": "<Base64 encoded thinknode username:password>",
    "api url": "https://<thinknode account>.thinknode.io/api/v1.0",
    "apps":
    {
        "dosimetry":
            "app_version": "1.0.0-beta1",
            "branch name": "master"
        },
        "dicom":
        {
            "app version": "",
            "branch name": "master"
        },
        "rt_types":
            "app_version": "",
            "branch name": "master"
        }
    },
    "realm_name": "<thinknode realm>",
    "account_name": "<thinknode account>"
}
```

# **Python: Immutable Storage**

# **Post Generic ISS Object**

The *post\_iss\_object\_generic.py* is a basic python script that provides an example to post any dosimetry type as an immutable object to the dosimetry app on the thinknode™ framework. This example can be used for any immutable storage post using any datatype by replacing the json iss file. The current example posts an rt\_study DICOM App datatype object that is read in from the study.json data file.

### Dependencies:

- thinknode.cfg
- .decimal Python Libraries
- study.json (or any other prebuilt json file of a dosimetry object as described in the Apps Manifest Guide)

### post\_iss\_object\_generic.py

```
# Copyright (c) 2015 .decimal, Inc. All rights reserved.
```

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```
# Desc: Post an immutable json object to the thinknode framework

from lib import thinknode_worker as thinknode
import requests
import json

iss_dir = "iss_files"
json_iss_file = "study.json"
obj_name = "rt_study"

# Get IAM ids
iam = thinknode.authenticate(thinknode.read_config('thinknode.cfg'))

# App object to post to iss
with open(iss_dir + '/' + json_iss_file) as data_file:
    json_data = json.load(data_file)

# Post immutable object to ISS
res = thinknode.post_immutable_named(iam, "dicom", json_data, obj_name)
```

#### **Returns:**

1. The ID (in json) of the object stored in Immutable Storage.

# **Python: Calculation Request**

# **Generic Calc Request**

The post\_calc\_request\_generic.py is a basic example to post a calculation request to dosimetry. This example can be used for any calculation request using any datatype by replacing the calculation request json file. This request will post a calculation request, check the status using long polling with a specified timeout, and return the calculation result.

### Dependencies:

- thinknode.cfg
- .decimal Python Libraries
- compute\_aperture.json (or any other prebuilt json file of a dosimetry object as described in the Dosimetry Manifest Guide)

### post calc request generic.py

```
# Copyright (c) 2015 .decimal, Inc. All rights reserved.
# Desc: Post a json calculation request to the thinknode framework
```

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```
request_dir = "request_files"
json_calc_file = "compute_aperture.json"

# Get IAM ids
iam = thinknode.authenticate(thinknode.read_config('thinknode.cfg'))

# App calculation request
with open(request_dir + '/' + json_calc_file) as data_file:
    json_data = json.load(data_file)

# Send calc request and wait for answer
res = thinknode.do_calculation(iam, json_data)
dl.data("Calculation Result: ", str(res))
```

#### **Returns:**

1. The calculation result (in json) of the API function called.

### **SOBP Dose Calculation**

The  $post\_calc\_request\_sobp\_dose.py$  and  $post\_calc\_request\_sobp\_dose\_with\_shifter.py$  are more complete examples that create input data and perform an sobp dose calculation function request to the dosimetry app on the thinknode<sup>TM</sup> framework.

The post\_calc\_request\_sobp\_dose.py example creates the entire calculation request inline using thinknode structure, array, and function requests. The entire dose calculation request is performed using one thinknode calculation provider call. While this structure of a request is a little more complicated to setup and perform, it removes the need to post to ISS or perform ancillary calculations separately.

The post\_calc\_request\_sobp\_dose\_with\_shifter.py adds in the complication of adding a degrader to the sobp calculation. This example performs three separate calculation requests. The first two requests are used to construct the proton degrader\_geometry and the third performs the actual dose calculation request using the previously constructed degrader. The entire example could be condensed into a single more complicated thinknode calculation structure, eliminating the need to perform the separate requests, but in some instances it can be more straight-forward to perform some of the calculations separately as shown. As seen in the example, the first two calculation results for the proton degrader are what is placed into the sobp calculation request, instead of the actual function calls as was done in the case of the aperture in the previous example.

### Dependencies:

- thinknode.cfg
- .decimal Python Libraries

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### **Example**

Below is an abbreviated version of the *post\_calc\_request\_sobp\_dose\_with\_shifter.py* file. The abbreviated sections are denoted as "...". In the below sample, the *dose\_calc* variable is a thinknode function request that is made of individually constructed arguments. Notice how some of the elements, like degrader, can be built upon seperate calculation requests.

- Modules used and explanation:
  - The thinknode\_worker (thinknode) module is a library that provides worker functions for performing and building the authentication, iss, and calculation requests to the thinknode framework.
  - The *dosimetry\_worker* (dosimetry) module is a library that provides simplified common dosimetry tasks.
  - The decimal\_logger (dl) module is a library that provides nicely formatted log output. This
    includes optional file logging, timestamps, and message coloring (when run through
    command windows).

Refer to the .decimal Libraries section for more information on the provided decimal libraries.

```
import json
from lib import thinknode worker as thinknode
from lib import dosimetry worker as dosimetry
from lib import decimal logging as dl
# Get IAM ids
iam = thinknode.authenticate(thinknode.read config('thinknode.cfg'))
def make dose points(pointCount):
. . .
def make layers(sad, range, mod):
    return \
        thinknode.function(iam["account_name"], "dosimetry",
"compute double scattering layers",
                thinknode.reference("55f70f5000c0a247563a909b6087ada0"), #
SOBP Machine from ISS
                thinknode.value(sad),
                thinknode.value(range),
                thinknode.value(mod)
            1)
def make_target():
    return \
        thinknode.function("dosimetry", "make_cube",
                thinknode.value([-32, -20, -30]),
```

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```
thinknode.value([16, -10, 30])
            1)
def compute aperture():
    return dosimetry.compute aperture(iam, make_target(), beam_geometry, 20.0,
0.0, 250.5)
beam geometry = \
. . .
# Get degrader geometry as calculation result
degrade geom = \
    thinknode.function(iam["account name"], "dosimetry", "make shifter",
            thinknode.value(18), # thickness
            thinknode.value("mm"), # units
            thinknode.value(200) # downstream edge
res geom = thinknode.do calculation(iam, degrade geom, True)
degrader = \
    thinknode.function(iam["account name"], "dosimetry", "make degrader",
            thinknode.value(res_geom),
            thinknode.reference("56030a9500c036a0c6393f984b25e303") # Material
spec from ISS
proton degr = thinknode.do calculation(iam, degrader)
# Call compute sobp pb dose2
dose calc = \
    thinknode.function("dosimetry", "compute sobp pb dose2",
            dosimetry.make_image_3d(iam, [-100, -100, -100], [200, 200, 200],
[2, 2, 2], 1), #stopping power image
            thinknode.value(make dose points(181)), # dose points
            beam geometry, #beam geometry
            dosimetry.make grid(iam, [-75, -75], [150, 150], [2, 2]), #
bixel grid
            make layers (2270.0, 152.0, 38.0),
            compute aperture(), # aperture based on targets
            thinknode.value([proton degr]) # degraders
        1)
# Perform calculation
res = thinknode.do calculation(iam, dose calc)
dl.data("Calculation Result: ", res)
```

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# **Python: decimal Libraries**

## rt\_types

The *rt\_types* module is a reconstruction of all astroid types in python class format. This includes interdependencies between types (e.g. the class "polyset" requires the class "polygon2").

Each data type detailed in the astroid Manifest Guide has a corresponding class in this python module.

Below you will see a snippet from the rt\_types module that shows the class for the *polyset* rt\_type along with its default initialization, *expand\_data* and *from\_json* functions.

```
class polygon2(object):
    #Initialize
    def init (self):
        blob = blob type()
        self.vertices = blob.toStr()
    def expand data(self):
        data = \{\}
        data['vertices'] =
parse bytes 2d(base64.b64decode(self.vertices['blob']))
        return data
    def from_json(self, jdict):
        for k, v in jdict.items():
            if hasattr(self,k):
                setattr(self, k, v)
class polyset(object):
    #Initialize
    def init (self):
        self.polygons = []
        self.holes = []
    def expand data(self):
        data = \{\}
        polygon = []
        for x in self.polygons:
            s = polygon2()
            s.from json(x)
            polygon.append(s.expand data())
        data['polygons'] = polygon
        hole = []
```

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```
for x in self.holes:
    s = polygon2()
    s.from_json(x)
    hole.append(s.expand_data())
    data['holes'] = hole
    return data

def from_json(self, jdict):
    for k, v in jdict.items():
        if hasattr(self, k):
            setattr(self, k, v)
```

- **Interdependence:** When rt\_types are constructed of other or multiple named types, they will be constructed as such in each class as seen in the *polygons* parameter of the *polyset* in the above example.
- **expand\_data function:** Each class's **expand\_data** function returns a python dictionary containing each of the values in the class, with all data values expanded out to remove compression or other encodings (i.e. providing results in a format more useful for send to other applications or for human-readability).
- **from json function:** Each class's *from json* function provides a method to turn a raw json string (e.g. a result from a thinknode calculation or ISS object) into an rt\_type data type. Proper use is to first construct an empty class instance, then to call the *from json* method on that instance, passing in the desired json data string.

Below is an example usage of getting a thinknode dose image (image\_3d data type in the astroid manifest) and turning it into a rt\_types image\_3d data type, so that it can be expanded and then used to output the image into a VTK graphics file:

```
def dose_to_vtk(dose_id):
    img_data = json.loads(thinknode.get_immutable(iam, 'dicom', dose_id))

img = rt_types.image_3d()
    img.from_json(img_data)
    img2 = img.expand_data()

vtk.write_vtk_image3('E:/dicom/dose.vtk', img2)
```

# thinknode worker

The *thinknode\_worker* module is the main work horse for communication with the astroid app and thinknode. The module will handle authentication, posting objects to ISS, creating most of the common calculation request structures, and posting the calculation request.

Refer to the .decimal GitHub repository for the complete module. Below are a few of the more common thinknode\_worker functions and their intended usages:

```
# Authenticate with thinknode and store necessary ids.
```

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```
# Gets the context id for each app detailed in the thinknode config
# Gets the app version (if non defined) for each app in the realm
   param config: connection settings (url and unique basic user
authentication)
def authenticate(config):
# Send calculation request to thinknode and wait for the calculation to
perform. Caches locally calculation results so if the same
# calculation is performed again, the calculation
# does not have to be repeatedly pulled from thinknode. Saves one calculation
time and bandwidth.
   note: see post_calculation if you just want the calculation ID and don't
need to wait for the calculation to finish or get results
   param config: connection settings (url, user token, and ids for context
and realm)
   param json_data: calculation request in json format
   param return data: When True the data object will be returned, when false
the thinknode id for the object will be returned
   param return error: When False the script will exit when error is found,
when True the sciprt will return the error
def do_calculation(config, json_data, return_data=True, return_error=False):
# Post immutable named type object to ISS
   param config: connection settings (url, user token, and ids for context
and realm)
   param app name: name of the app to use to get the context id from the iam
   param ison data: immutable object in ison format
   param obj name: object name of app to post to
def post_immutable_named(config, app_name, json_data, obj_name):
    scope = '/iss/named/' + config["account name"] + '/rt types' + '/' +
obj_name
    return post immutable(config, app name, json data, scope)
# Post immutable object to ISS
   param config: connection settings (url, user token, and ids for context
and realm)
   param app_name: name of the app to use to get the context id from the iam
confia
    param obj id: thinknode iss reference id for object to get
def get immutable(config, app name, obj id):
```

# dosimetry worker

The dosimetry\_worker module provides high-level functions for building data types and calculation requests for common dosimetry tasks. This library is constantly growing as more routine tasks are

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programmed in python.

Refer to the .decimal GitHub repository for the complete module. Some basic examples of provided functionality are:

- 1. Aperture creation (using structures/beams or basic geometric)
- 2. Dose comparison
- 3. Grid creation
- 4. Image creation
- 5. PBS Spot functions

## vtk\_worker

The VTK worker provides a means to write out common rt\_types to a vtk file format (The Visualization TooKit) that can be visualized in Paraview. It's most useful for displaying and post-processing image, mesh, and other primitive object data types.

Below is an example of turning a dose image 3d into a vtk file for visualization in Paraview:

```
def dose_to_vtk(dose_id):
    img_data = json.loads(thinknode.get_immutable(iam, 'dicom', dose_id))

img = rt_types.image_3d()
    img.from_json(img_data)
    img2 = img.expand_data()

vtk.write_vtk_image3('E:/dicom/dose.vtk', img2)
```

# decimal\_logging

The decimal logging module provides formatted and detailed output window messages and file logging.

The following settings are available in the decimal\_logging.py file: **display\_timestamps:** display timestamps in the output window/logfile **display\_types:** display message types (e.g. debug, data, alert) in the output window/logfile **log\_file:** sets the logfile name and location

### **Debugging**

When debugging, use the dl.debug() function and set the *isDebug* flag in the decimal\_logging library to True. This toggles on the output for each of the dl.debug calls. By default we keep debugging off, but it can be turned on as needed.

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### Other Flags

The following image shows the logging settings for each message type as:

```
    Timestamps = True; Types = True
    Timestamps = False; Types = True
    Timestamps = False; Types = False
```

```
2015-09-28 11:43:14 -- MESSAGE: decimal message
2015-09-28 11:43:14 -- DEBUG: >>> decimal debug <<<
2015-09-28 11:43:14 --
2015-09-28 11:43:14 !! WARNING: decimal warning
2015-09-28 11:43:14
2015-09-28 11:43:14 -- EVENT: decimal event
2015-09-28 11:43:14 -- DATA: decimal debug_data
data
-- MESSAGE: decimal message
-- DEBUG: >>> decimal debug <<<
-- ALERT: decimal alert !! WARNING: decimal warning
  EVENT: decimal event
-- DATA: decimal debug_data
data
decimal message
decimal debug <<<
decimal warning
decimal event
decimal debug_data
data
```

### **File Logging**

The decimal\_logging library also provides simple file logging. The *log\_file* variable at the top of the library sets the log file. By using any of the following functions, you can easily log data to the specified file:

- log(message)
- log\_debug\_data(message,data)
- log\_data(data)

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