```
biocompute object example for "antiviral resistance detection[ICD:B17]"
{
    "id": "obj.1001",
    "type": "antiviral resistance detection[ICD:B17]",
    "name": "HCV1a[taxID:31646] ledipasvir[PubChem:67505836] resistance SNP [SO:
           0000694]",
    "version": "1.2",
    "verification": "manual",
    "creation date": "Mon Jun 6 13:25:21 EDT 2016",
    "modification_date": "Thu Jun 9 08:12:14 EDT 2016",
    "publication status": "draft",
    "keywords": [
        "antiviral resistance",
        "SNP"
    ],
    "xref": [
        "DO:1883",
        "ICD:B17",
        "taxID:9606"
    "usability domain": [
        "Identify baseline single nucleotide polymorphisms (SNPs[SO:0000694]),
           insertions[SO:0000667], and deletions[SO:0000045] that correlate with
           reduced ledipasvir[PubChem: 67505836] antiviral drug efficacy in
           Hepatitis C virus subtype 1",
        "Identify treatment emergent amino acid substitutions[SO:0000048] that
           correlate with antiviral drug treatment failure",
        "Determine whether the treatment emergent amino acid substitutions[SO:
           0000048] identified correlate with treatment failure involving other
           drugs against the same virus"
    ],
    "authors": [
        {
                "first name": "Charles",
                "last name": "Darwin",
                "affiliation": "HMS Beagle"
        },
                "first name": "Rosalind",
                "last name": "Franklin",
                "affiliation": "King's College, London"
        }
    "pipeline": "obj.10001",
    "alignment" : {
        "seed": 14,
        "minimum match len": 50,
        "divergence threshold percent": 10
    "variant calling" : {
```

```
"minimum coverage": 20,
       "frequency cutoff": 0.20
       },
   "verification_kit":{
       "reference sequence":[
           {
               "source": "NCBI",
               "db": "genbank",
               "id": ["JQ434001.1"]
           },
               "source": "HIVE",
               "db": "Honeycomb",
               "id": "obj.00123"
           }
       ],
       "sequence_reads": [
           {
               "source": "HIVE",
               "db" : "Honeycomb",
               "id": "obj.00013"
           },
               "source": "HIVE",
               "db": "Honeycomb",
               "id": "obj.10065"
           }
       ],
       "snv profile file": {
           "format": [".csv",".vcf"],
           "filename": "SNPProfile"
       "alignment file": {
           "format": [".sam"],
           "filename": "SNPProfile.sam"
       },
       "aminoacid calls file": {
           "format": [".csv",".vcf"],
           "filename": "AAcalls.csv"
       }
   }
pipeline definition example for "HIVE-Viral Mutation Detection Pipeline"
______
    " id": "obj.10001",
    "_type": "pipeline_definition",
    "name": "HIVE-Viral Mutation Detection Pipeline",
```

}

{

```
"description": "HIVE-Viral Mutation Detection uses two HIVE tools: Hexagon
          for alignment and Heptagon for variant calling. Only the options that
          need to be selected or changed are listed.",
    "workflow": {
       "launcher": "system",
       "script": "obj.123123"
    "tool_used": [
       {
           "name": "HIVE-hexagon",
           "version": "1.3",
           "arguments": [
               {
                   "parameter": "seed",
                   "value": 16
               },
               {
                   "parameter": "divergence threshold percent",
                   "value": 15
               },
               {
                   "parameter": "minimum match len",
                   "value": 100
               }
           1
       },
           "name": "HIVE-heptagon",
           "version": "1.1",
           "arguments": [
               {
                   "parameter": "min coverage",
                   "value": 10
               },
               {
                   "parameter": "freq cutoff",
                   "value": 0.10
               }
           ]
       }
   ]
typedef for "antiviral resistance detection"
______
    " type": "type",
    "name": "antiviral_resistance_detection",
    "description": "This pipeline uses an aligner and then a profiler to detect
          possible mutations involved in drug resistance.",
    " fields": {
```

}

{

```
"reference sequence": {
    "description": "reference sequences can be either FASTA from known
        sources or identifiers from HIVE database. These must be saved as
        a 'genome' type in HIVE to use in this field",
    "title": "Reference Sequences",
    " type": "xref",
    " role": "input",
    __vital": "true",
    "plural": "true"
},
"sequence reads": {
    "description": "sequences can be either FASTA from experimental
        sources or identifiers from HIVE database. These must be saved as
        a 'nuc read' type in HIVE to use in this field",
    "title": "Nucleotide Read",
    "_type": "xref",
    " role": "input",
    " vital": "true",
    " plural": "true"
},
"alignment": {
    "description": "Arguments for the alignment, HIVE Hexagon",
    "title": "HIVE Hexagon Alignment Engine",
    "seed": {
        "_type": "integer",
        " default": 11,
        " limit": {
            "eval-js" : {"$seed>8 && $seed<12"}</pre>
        }
    },
    "minimum match len,": {
        " type": "integer",
        " default": 75
    "divergence threshold percent": {
        " type": "real",
        default: 15
        __limit:" {
            "eval-js" : {"$divergence threshold percent>10 &&
                   $divergence threshold percent<40"}</pre>
        }
    }
},
"variant calling": {
    "description": "arguments for the engine to perform coverage
        analysis, base-calling, SNP calling",
    "title": "Sequence Profiling Engine",
    "minimum coverage": {
        " type": "integer",
        " default": 10,
        " limit": {
            "eval-js" : {"$minimum coverage>=0"}
```

```
}
           },
            "frequency cutoff": {
               " type": "real",
               " default": 0.10,
               " limit": {
                   "eval-js" : {"$frequency cutoff>=0 &&
                          $frequency cutoff<=1.0"}</pre>
               }
           }
       },
        "snv profile": {
           "description": "The output file containing the SNP Profile",
           "title": "SNPProfile Output File",
           " role": "output",
           " type": "outfile"
        "alignment file": {
           "description": "The resulting alignment file from the Hexagon
                computation",
           "title": "Alignment File",
           " type": "outfile"
       },
        "aminoacid calls": {
           "description": "the amino acid calls from Heptagon",
           "title": "Amino Acid Calls",
           " role": "output",
           " type": "outfile"
       }
    " inherits": [
        "base biocompute type"
    ]
}
typedef for "pipeline definition type"
_____
{
    " id": "obj.10001",
    " type": "pipeline definition",
   "name": "HIVE-Viral Mutation Detection Pipeline",
    "description": "HIVE-Viral Mutation Detection uses two HIVE tools: Hexagon
          for alignment and Heptagon for variant calling. Only the options that
          need to be selected or changed are listed.",
    "workflow": {
        "launcher": "system",
       "script": "obj.123123"
    "tool used": [
           "name": "HIVE-hexagon",
```

```
"version": "1.3",
            "arguments": [
                {
                    "parameter": "seed",
                    "value": 16
                },
                {
                    "parameter": "divergence_threshold percent",
                    "value":
                              15
                },
                {
                    "parameter": "minimum match len",
                    "value": 100
                }
            ]
        },
            "name": "HIVE-heptagon",
            "version": "1.1",
            "arguments": [
                {
                    "parameter": "min coverage",
                    "value": 10
                },
                {
                    "parameter": "freq cutoff",
                    "value": 0.10
                }
            ]
       }
    ]
}
typedef for "base biocompute type"
{
    " type": "type",
    "name": "base_biocompute_type",
    "description": "Provides the base fields to populate in a biocompute object",
    " fields": {
        "validation": {
            "description": "Describes the position of an object in the validation
                process. The unreviewed flag indicates that the object has been
                submitted, but no further curation has occurred. The in progress
                flag indicates that validation is underway. manual indicates that
                the object has been manually validated. The suspended flag
                indicates an object that was once valid is no longer considered
                valid. The error flag indicates that an error was detected with
                the biocompute object.",
            "_type": "string",
            " limit": {
```

```
"choice": [
                   "unreviewed",
                   "in progress",
                   "manual",
                   "suspended",
                   "error"
               1
           }
       },
       "usability domain": {
           "description": "Provides a space for the author to define the
               usability
               domane of an object.",
           "_type": "string",
"_vital": "true",
           " plural": "true"
       "pipeline" : {
           "description": "defines the fields that have to be populated for a
               tool description in the pipeline of a biocompute object",
           " type": "pipeline definition"
       }
    " inherits": ["base metadata type"]
}
                          ______
typedef for "base metadata type"
______
   " type": "datatype",
   "name": "base metadata type",
   "description": "Definition of the base metadata type providing fields for
       the inclusion of outside links, submissions, and information pertinent
       to the experiment but not critical for the computation.",
   " fields": {
       "name": {
           " type": "string"
       },
       "creation date": {
           "description": "The time and date that the object is
               created/computation was run",
           " type": " datetime"
       },
       "modification date": {
           "description": "This field will list all the dates and time an
               object ismodified.",
           " type": " datetime",
           " plural": "true"
       },
       "publication status": {
           "description": "This is a choice field with four options. The draft
```

```
flag indicates that an object is in draft form and is still
        being added to or changed. The open access flag indicates that
        an object has been published and is freely available to anyone.
        The objects with the private flag have restrictions on who can
        view and access them. This is a way for researchers using
        restricted data or metadata to ensure the confidentially is
        maintained.",
    "_type": "string",
    " limit": {
        "choice": ["draft", "open access", "private", "recalled"]
},
"authors": {
    "description": "This is the list of authors submitting this object.
        First and last name are required, middle name and affiliation are
        not. The affiliation field is for listing the academic,
        governmental, or private entity that supported the project
        responsible for generating the biocompute object.",
    " plural": "true",
    "first name": {
        __type":"string",
        " vital": "true"
    },
    "middle name": {
        " type": "string"
    },
    "last name": {
        " type": "string",
        "vital": "true"
    "affiliation": {
        " type": "string"
    }
},
"keywords": {
    "description": "Keywords to aid in search-ability and description of
        the object. These are a controlled vocabulary.",
    "_type": "string",
    " plural": "true"
}
```

}

}