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=====
biocompute object example for "Huntington's_disease[DO:12858]_detection"
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{
  "_id": "obj.3001",
  "_type": "Huntingtons_disease[DO:12858]_detection",
  "name": "Human[tax:9606] CAG repeats[SO:0000726] detection for Huntington's
    disease[DOID:12858].",
  "version": "1.0",
  "verification": "manual",
  "creation_date": "Thu Aug 4 10:16:32 EDT 2016",
  "modification_date": "Thu Aug 4 10:16:32 EDT 2016",
  "publication_status": "draft",
  "keywords": [
    "disease",
    "human disease"
  ],
  "xref": [
    "DOID:12858",
    "GOID:GO272344",
    "ICD:G10",
    "SO:0000726",
    "tax:9606"
  ],
  "usability_domain": [
    "Determine CAG repeats[SO:0000726] in a sequence for the diagnosis of
      Huntington's disease[ICD:G10] for htt gene[GOID:GO272344]: 10 - 35 CAG
      trinucleotide repeats: No Risk, 36 - 40 CAG trinucleotide repeats:
      Moderate Risk, 40+ CAG trinucleotide repeats: High Risk"
  ],
  "authors": [
    {
      "first_name": "George",
      "last_name": "Huntington",
      "affiliation": "Columbia University"
    }
  ],
  "pipeline": "obj.10003",
  "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",

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        "id": [
            "NM_0002111.8",
            "SRR2028290"
        ]
    },
    ],
    "sequence_reads": [
        {
            "source": "HIVE",
            "db": "Honeycomb",
            "id": ["obj.00001"]
        }
    ],
    "snp_profile_file": {
        "format": [".csv"],
        "filename": "SNPProfile.csv"
    }
}
}

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pipeline definition example for "HIVE Huntington's Disease Detection Pipeline"

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{
    "_id": "obj.20003",
    "_type": "pipeline_definition",
    "name": "HIVE Huntington's Disease Detection Pipeline",
    "description": ["This pipeline uses an aligner and then a profiler to detect
        possible mutations indicating risk of huntington's disease in a gene
        sequence. "],
    "workflow": {
        "launcher": "system",
        "script": "013324"
    },
    "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
                }
            ]
        }
    ]
}

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    ]
  },
  {
    "name": "HIVE-heptagon",
    "version": "1.1",
    "arguments": [
      {
        "parameter": "min_coverage",
        "value": 10
      },
      {
        "parameter": "freq_cutoff",
        "value": 0.10
      }
    ]
  }
]
}

=====
typedef for "HIVE_huntingtons_disease_detection"
=====
{
  "_type": "type",
  "name": "HIVE_huntingtons_disease_detection",
  "description": "Pipeline to determine the risk of huntington's disease in a
    gene sequence",
  "_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": "Non-default arguments for the alignment, HIVE
        Hexagon",
      "title": "HIVE Hexagon Alignment Engine",

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    "seed": {
        "_type": "integer",
        "_default": 11,
        "_limit": {
            "eval-js": "$seed>8 && $seed<12"
        }
    },
    "minimum_match_len": {
        "_type": "integer",
        "_default": 75
    },
    "divergence_threshold_percent": {
        "_type": "real",
        "_default": 15,
        "_limit": {
            "eval-js": ["$divergence_threshold_percent>10 &&
                        $divergence_threshold_percent<40"]
        }
    },
    "variant_calling": {
        "description": "arguments for the engine to perform coverage
                        analysis, base-calling, SNP calling",
        "title": "Sequence Profiling Engine",
        "minimum_coverage": {
            "_type": "integer",
            "_default": 10,
            "_limit": {
                "eval-js": ["$minimum_coverage>=0"]
            }
        },
        "frequency_cutoff": {
            "_type": "real",
            "_default": 0.10,
            "_limit": {
                "eval-js": ["$frequency_cutoff>=0 && $frequency_cutoff<=1.0"]
            }
        }
    },
    "snv_profile": {
        "description": "The output file containing the SNP Profile",
        "title": "SNPProfile Output File",
        "_role": "output",
        "_type": "outfile"
    },
    "_inherits": ["base_biocompute_type"]
}

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=====
typedef for "pipeline_definition_type"
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{
  "_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
        }
      ]
    }
  ]
}

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=====
typedef for "base_biocompute_type"
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{
  "_type": "type",

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"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"
            ]
        }
    },
    "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"]
}

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=====
typedef for "base_metadata_type"
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{
    "_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"
        }
    }
}

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},
"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 insure the confidentiality 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",

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    "_plural": "true"  
  }  
}
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