

---

# Autoprotocol Documentation

*Release 4.0.0*

**Transcriptic**

Aug 02, 2018



---

## Contents

---

|           |  |            |
|-----------|--|------------|
| <b>1</b>  | <b>autoprotocol.protocol module</b>          | <b>1</b>   |
| <b>2</b>  | <b>autoprotocol.instruction module</b>       | <b>61</b>  |
| <b>3</b>  | <b>autoprotocol.container</b>                | <b>75</b>  |
| 3.1       | container.Container . . . . .                | 75         |
| 3.2       | container.Well . . . . .                     | 78         |
| 3.3       | container.WellGroup . . . . .                | 79         |
| <b>4</b>  | <b>autoprotocol.container_type</b>           | <b>83</b>  |
| 4.1       | container_type.ContainerType . . . . .       | 83         |
| 4.2       | Container Types . . . . .                    | 85         |
| <b>5</b>  | <b>autoprotocol.builders</b>                 | <b>107</b> |
| 5.1       | Summary . . . . .                            | 107        |
| <b>6</b>  | <b>autoprotocol.liquid_handle</b>            | <b>121</b> |
| 6.1       | liquid_handle.liquid_handle_method . . . . . | 121        |
| 6.2       | liquid_handle.liquid_class . . . . .         | 124        |
| 6.3       | liquid_handle.transfer . . . . .             | 128        |
| 6.4       | liquid_handle.mix . . . . .                  | 137        |
| 6.5       | liquid_handle.tip_type . . . . .             | 139        |
| <b>7</b>  | <b>autoprotocol support</b>                  | <b>141</b> |
| 7.1       | autoprotocol.unit . . . . .                  | 141        |
| 7.2       | autoprotocol.util . . . . .                  | 142        |
| 7.3       | autoprotocol.harness . . . . .               | 143        |
| <b>8</b>  | <b>Changelog</b>                             | <b>147</b> |
| <b>9</b>  | <b>Credits</b>                               | <b>159</b> |
| <b>10</b> | <b>License</b>                               | <b>161</b> |
| <b>11</b> | <b>Installation</b>                          | <b>163</b> |
| <b>12</b> | <b>Building a Protocol</b>                   | <b>165</b> |

|                            |            |
|----------------------------|------------|
| <b>13 Contributing</b>     | <b>169</b> |
| <b>14 Search the Docs</b>  | <b>171</b> |
| <b>Python Module Index</b> | <b>173</b> |

# CHAPTER 1

## autoprotocol.protocol module

Module containing the main *Protocol* object and associated functions

**copyright** 2018 by The Autoprotocol Development Team, see AUTHORS for more details.

**license** BSD, see LICENSE for more details

**class** autoprotocol.protocol.**Protocol** (*refs=None*, *instructions=None*)

A Protocol is a sequence of instructions to be executed, and a set of containers on which those instructions act.

Initially, a Protocol has an empty sequence of instructions and no referenced containers. To add a reference to a container, use the ref() method, which returns a Container.

```
p = Protocol()
my_plate = p.ref("my_plate", id="ctlxae8jabbe6",
                  cont_type="96-pcr", storage="cold_4")
```

To add instructions to the protocol, use the helper methods in this class

```
p.transfer(source=my_plate.well("A1"),
           dest=my_plate.well("B4"),
           volume="50:microliter")
p.thermocycle(my_plate, groups=[
    { "cycles": 1,
      "steps": [
          { "temperature": "95:celsius",
            "duration": "1:hour"
          }
      ]
  }])
```

Autoprotocol Output:

```
{
  "refs": {
    "my_plate": {
      "id": "ctlxae8jabbe6",
```

(continues on next page)

(continued from previous page)

```

    "store": {
        "where": "cold_4"
    }
},
"instructions": [
{
    "groups": [
    {
        "transfer": [
        {
            "volume": "50.0:microliter",
            "to": "my_plate/15",
            "from": "my_plate/0"
        }
    ]
}
],
"op": "pipette"
},
{
    "volume": "10:microliter",
    "dataref": null,
    "object": "my_plate",
    "groups": [
    {
        "cycles": 1,
        "steps": [
        {
            "duration": "1:hour",
            "temperature": "95:celsius"
        }
    ]
}
],
"op": "thermocycle"
}
]
}
}

```

**absorbance** (*ref*, *wells*, *wavelength*, *dataref*, *flashes=25*, *incubate\_before=None*, *temperature=None*, *settle\_time=None*)

Read the absorbance for the indicated wavelength for the indicated wells. Append an Absorbance instruction to the list of instructions for this Protocol object.

Example Usage:

```

p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-flat",
                     storage="warm_37")

p.absorbance(sample_plate, sample_plate.wells_from(0,12),
             "600:nanometer", "test_reading", flashes=50)

```

Autoprotocol Output:

```

"instructions": [
    {
        "dataref": "test_reading",
        "object": "sample_plate",
        "wells": [
            "A1",
            "A2",
            "A3",
            "A4",
            "A5",
            "A6",
            "A7",
            "A8",
            "A9",
            "A10",
            "A11",
            "A12"
        ],
        "num_flashes": 50,
        "wavelength": "600:nanometer",
        "op": "absorbance"
    }
]

```

## Parameters

- **ref** (*str or Ref*) – Object to execute the absorbance read on
- **wells** (*list(Well) or WellGroup or Well*) – WellGroup of wells to be measured or a list of well references in the form of [“A1”, “B1”, “C5”, …]
- **wavelength** (*str or Unit*) – wavelength of light absorbance to be read for the indicated wells
- **dataref** (*str*) – name of this specific dataset of measured absorbances
- **flashes** (*int, optional*) – number of flashes for the read
- **temperature** (*str or Unit, optional*) – set temperature to heat plate reading chamber
- **settle\_time** (*Unit, optional*) – the time before the start of the measurement, defaults to vendor specifications
- **incubate\_before** (*dict, optional*) – incubation prior to reading if desired

```

{
    "shaking": {
        "amplitude": str or Unit
        "orbital": bool
    },
    "duration": str or Unit
}

```

**Returns** Returns the `autoprotocol.instruction.Absorbance` instruction created from the specified parameters

**Return type** `Absorbance`

**Raises**

- `TypeError` – Invalid input types, e.g. wells given is of type `Well`, `WellGroup` or list of `wells`
- `ValueError` – Wells specified are not from the same container
- `ValueError` – Settle time has to be greater than 0
- `UnitError` – Settle time is not of type `Unit`

### `acoustic_transfer`(*source*, *dest*, *volume*, *one\_source=False*, *droplet\_size='25:nanoliter'*)

Specify source and destination wells for transferring liquid via an acoustic liquid handler. Droplet size is usually device-specific.

Example Usage:

```
p.acoustic_transfer(  
    echo.wells(0, 1).set_volume("12:nanoliter"),  
    plate.wells_from(0, 5), "4:nanoliter", one_source=True)
```

Autoprotocol Output:

```
"instructions": [  
    {  
        "groups": [  
            {  
                "transfer": [  
                    {  
                        "volume": "0.004:microliter",  
                        "to": "plate/0",  
                        "from": "echo_plate/0"  
                    },  
                    {  
                        "volume": "0.004:microliter",  
                        "to": "plate/1",  
                        "from": "echo_plate/0"  
                    },  
                    {  
                        "volume": "0.004:microliter",  
                        "to": "plate/2",  
                        "from": "echo_plate/0"  
                    },  
                    {  
                        "volume": "0.004:microliter",  
                        "to": "plate/3",  
                        "from": "echo_plate/1"  
                    },  
                    {  
                        "volume": "0.004:microliter",  
                        "to": "plate/4",  
                        "from": "echo_plate/1"  
                    }  
                ]  
            }  
        ],  
        "droplet_size": "25:microliter",  
        "op": "acoustic_transfer"  
    }  
]
```

## Parameters

- **source** (`Well` or `WellGroup` or `list(Well)`) – Well or wells to transfer liquid from. If multiple source wells are supplied and one\_source is set to True, liquid will be transferred from each source well specified as long as it contains sufficient volume. Otherwise, the number of source wells specified must match the number of destination wells specified and liquid will be transferred from each source well to its corresponding destination well.
- **dest** (`Well` or `WellGroup` or `list(Well)`) – Well or WellGroup to which to transfer liquid. The number of destination wells must match the number of source wells specified unless one\_source is set to True.
- **volume** (`str` or `Unit` or `list`) – The volume(s) of liquid to be transferred from source wells to destination wells. Volume can be specified as a single string or Unit, or can be given as a list of volumes. The length of a list of volumes must match the number of destination wells given unless the same volume is to be transferred to each destination well.
- **one\_source** (`bool, optional`) – Specify whether liquid is to be transferred to destination wells from a group of wells all containing the same substance.
- **droplet\_size** (`str` or `Unit, optional`) – Volume representing a droplet\_size. The volume of each *transfer* group should be a multiple of this volume.

**Returns** Returns the `autoprotocol.instruction.AcousticTransfer` instruction created from the specified parameters

**Return type** `AcousticTransfer`

## Raises

- `TypeError` – Incorrect input types, e.g. source/dest are not Well or WellGroup or list of Well
- `RuntimeError` – Incorrect length for source and destination
- `RuntimeError` – Transfer volume not being a multiple of droplet size
- `RuntimeError` – Insufficient volume in source wells

**add\_time\_constraint** (`from_dict, to_dict, less_than=None, more_than=None, mirror=False, ideal=None, optimization_cost=None`)

Constraint the time between two instructions

Add time constraints from `from_dict` to `to_dict`. Time constraints guarantee that the time from the `from_dict` to the `to_dict` is less than or greater than some specified duration. Care should be taken when applying time constraints as constraints may make some protocols impossible to schedule or run.

Though autoprotocol orders instructions in a list, instructions do not need to be run in the order they are listed and instead depend on the preceding dependencies. Time constraints should be added with such limitations in mind.

Constraints are directional; use `mirror=True` if the time constraint should be added in both directions. Note that mirroring is only applied to the `less_than` constraint, as the `more_than` constraint implies both a minimum delay between two timing points and also an explicit ordering between the two timing points.

Ideal time constraints are sometimes helpful for ensuring that a certain set of operations happen within some specified time. This can be specified by using the `ideal` parameter. There is an optional `optimization_cost` parameter associated with `ideal` time constraints for specifying the penalization system used for calculating deviations from the `ideal` time. When left unspecified, the `optimization_cost` function defaults to linear. Please refer to the ASC for more details on how this is implemented.

Example Usage:

```
plate_1 = protocol.ref("plate_1", id=None, cont_type="96-flat",
                      discard=True)
plate_2 = protocol.ref("plate_2", id=None, cont_type="96-flat",
                      discard=True)

protocol.cover(plate_1)
time_point_1 = protocol.get_instruction_index()

protocol.cover(plate_2)
time_point_2 = protocol.get_instruction_index()

protocol.add_time_constraint(
    {"mark": plate_1, "state": "start"},
    {"mark": time_point_1, "state": "end"},  
    less_than = "1:minute")
protocol.add_time_constraint(
    {"mark": time_point_2, "state": "start"},  
    {"mark": time_point_1, "state": "start"},  
    less_than = "1:minute", mirror=True)

# Ideal time constraint
protocol.add_time_constraint(
    {"mark": time_point_1, "state": "start"},  
    {"mark": time_point_2, "state": "end"},  
    ideal = "30:second",
    optimization_cost = "squared")
```

Autoprotocol Output:

```
{
  "refs": {
    "plate_1": {
      "new": "96-flat",
      "discard": true
    },
    "plate_2": {
      "new": "96-flat",
      "discard": true
    }
  },
  "time_constraints": [
    {
      "to": {
        "instruction_end": 0
      },
      "less_than": "1.0:minute",
      "from": {
        "ref_start": "plate_1"
      }
    },
    {
      "to": {
        "instruction_start": 0
      },
      "less_than": "1.0:minute",
      "from": {
        "ref_start": "plate_1"
      }
    }
  ]
}
```

(continues on next page)

(continued from previous page)

```

        "instruction_start": 1
    }
},
{
    "to": {
        "instruction_start": 1
    },
    "less_than": "1.0:minute",
    "from": {
        "instruction_start": 0
    }
},
{
    "from": {
        "instruction_start": 0
    },
    "to": {
        "instruction_end": 1
    },
    "ideal": {
        "value": "5:minute",
        "optimization_cost": "squared"
    }
}
]

,
"instructions": [
{
    "lid": "standard",
    "object": "plate_1",
    "op": "cover"
},
{
    "lid": "standard",
    "object": "plate_2",
    "op": "cover"
}
]
}
}

```

## Parameters

- **from\_dict** (*dict*) – Dictionary defining the initial time constraint condition. Composed of keys: “mark” and “state”
  - mark:** int or Container instruction index of container
  - state:** “start” or “end” specifies either the start or end of the “mark” point
- **to\_dict** (*dict*) – Dictionary defining the end time constraint condition. Specified in the same format as from\_dict
- **less\_than** (*str or Unit, optional*) – max time between from\_dict and to\_dict
- **more\_than** (*str or Unit, optional*) – min time between from\_dict and to\_dict
- **mirror** (*bool, optional*) – choice to mirror the from and to positions when time constraints should be added in both directions (only applies to the less\_than constraint)

- **ideal**(*str or Unit, optional*) – ideal time between from\_dict and to\_dict
- **optimization\_cost** (*Enum({ "linear", "squared", "exponential" }), optional*) – cost function used for calculating the penalty for missing the *ideal* timing

#### Raises

- `ValueError` – If an instruction mark is less than 0
- `TypeError` – If mark is not container or integer
- `TypeError` – If state not in ['start', 'end']
- `TypeError` – If any of *ideal*, *more\_than*, *less\_than* is not a Unit of the 'time' dimension
- `KeyError` – If *to\_dict* or *from\_dict* does not contain 'mark'
- `KeyError` – If *to\_dict* or *from\_dict* does not contain 'state'
- `ValueError` – If time is less than '0:second'
- `ValueError` – If *optimization\_cost* is specified but *ideal* is not
- `ValueError` – If *more\_than* is greater than *less\_than*
- `ValueError` – If *ideal* is smaller than *more\_than* or greater than *less\_than*
- `RuntimeError` – If *from\_dict* and *to\_dict* are equal
- `RuntimeError` – If *from\_dict*["marker"] and *to\_dict*["marker"] are equal and *from\_dict*["state"] = "end"

#### `as_dict()`

Return the entire protocol as a dictionary.

Example Usage:

```
from autoprotocol.protocol import Protocol
import json

p = Protocol()
sample_ref_2 = p.ref("sample_plate_2",
                     id="ct1cxaee331kj",
                     cont_type="96-pcr",
                     storage="ambient")
p.seal(sample_ref_2)
p.incubate(sample_ref_2, "warm_37", "20:minute")

print json.dumps(p.as_dict(), indent=2)
```

Autoprotocol Output:

```
{
  "refs": {
    "sample_plate_2": {
      "id": "ct1cxaee331kj",
      "store": {
        "where": "ambient"
      }
    }
  },
  "instructions": [
```

(continues on next page)

(continued from previous page)

```
{
  "object": "sample_plate_2",
  "op": "seal"
},
{
  "duration": "20:minute",
  "where": "warm_37",
  "object": "sample_plate_2",
  "shaking": false,
  "op": "incubate"
}
]
```

**Returns** dict with keys “refs” and “instructions” and optionally “time\_constraints” and “outs”, each of which contain the “refified” contents of their corresponding Protocol attribute.

**Return type** dict

**autopick** (*sources*, *dests*, *min\_abort*=0, *criteria*=None, *dataref*=‘autopick’)

Pick colonies from the agar-containing location(s) specified in *sources* to the location(s) specified in *dests* in highest to lowest rank order until there are no more colonies available. If fewer than *min\_abort* pickable colonies have been identified from the location(s) specified in *sources*, the run will stop and no further instructions will be executed.

Example Usage:

Autoprotocol Output:

#### Parameters

- **sources** (`Well` or `WellGroup` or `list(Well)`) – Reference wells containing agar and colonies to pick
- **dests** (`Well` or `WellGroup` or `list(Well)`) – List of destination(s) for picked colonies
- **criteria** (`dict`) – Dictionary of autopicking criteria.
- **min\_abort** (`int, optional`) – Total number of colonies that must be detected in the aggregate list of *from* wells to avoid aborting the entire run.
- **dataref** (`str`) – Name of dataset to save the picked colonies to

**Returns** Returns the `autoprotocol.instruction.Autopick` instruction created from the specified parameters

**Return type** `Autopick`

#### Raises

- `TypeError` – Invalid input types for sources and dests
- `ValueError` – Source wells are not all from the same container

**batch\_containers** (*containers*, *batch\_in*=True, *batch\_out*=False)

Batch containers such that they all enter or exit together.

Example Usage:

```
plate_1 = protocol.ref("p1", None, "96-pcr", storage="cold_4")
plate_2 = protocol.ref("p2", None, "96-pcr", storage="cold_4")

protocol.batch_containers([plate_1, plate_2])
```

Autoprotocol Output:

```
{
  "refs": {
    "p1": {
      "new": "96-pcr",
      "store": {
        "where": "cold_4"
      }
    },
    "p2": {
      "new": "96-pcr",
      "store": {
        "where": "cold_4"
      }
    }
  },
  "time_constraints": [
    {
      "from": {
        "ref_start": "p1"
      },
      "less_than": "0:second",
      "to": {
        "ref_start": "p2"
      }
    },
    {
      "from": {
        "ref_start": "p1"
      },
      "more_than": "0:second",
      "to": {
        "ref_start": "p2"
      }
    }
  ]
}
```

## Parameters

- **containers** (*list (Container)*) – Containers to batch
- **batch\_in** (*bool, optional*) – Batch the entry of containers, default True
- **batch\_out** (*bool, optional*) – Batch the exit of containers, default False

## Raises

- `TypeError` – If containers is not a list
- `TypeError` – If containers is not a list of Container object

**container\_type** (*shortname*)

Convert a ContainerType shortname into a ContainerType object.

**Parameters** **shortname** (*str*) – String representing one of the ContainerTypes in the \_CONTAINER\_TYPES dictionary.

**Returns** Returns a Container type object corresponding to the shortname passed to the function.

If a ContainerType object is passed, that same ContainerType is returned.

**Return type** *ContainerType*

**Raises** `ValueError` – If an unknown ContainerType shortname is passed as a parameter.

**count\_cells** (*wells*, *volume*, *dataref*, *labels=None*)

Count the number of cells in a sample that are positive/negative for a given set of labels.

Example Usage:

```
p = Protocol()

cell_suspension = p.ref(
    "cells_with_trypan_blue",
    id=None,
    cont_type="micro-1.5",
    discard=True
)
p.count_cells(
    cell_suspension.well(0),
    "10:microliter",
    "my_cell_count",
    ["trypan_blue"]
)
```

Autoprotocol Output:

```
{
  "refs": {
    "cells_with_trypan_blue": {
      "new": "micro-1.5",
      "discard": true
    }
  },
  "instructions": [
    {
      "dataref": "my_cell_count",
      "volume": "10:microliter",
      "wells": [
        "cells_with_trypan_blue/0"
      ],
      "labels": [
        "trypan_blue"
      ],
      "op": "count_cells"
    }
  ]
}
```

## Parameters

- **wells** (`Well` or `list(Well)` or `WellGroup`) – List of wells that will be used for cell counting.
- **volume** (`Unit`) – Volume that should be consumed from each well for the purpose of cell counting.
- **dataref** (`str`) – Name of dataset that will be returned.
- **labels** (`list(string)`, *optional*) – Cells will be scored for presence or absence of each label in this list. If staining is required to visualize these labels, they must be added before execution of this instruction.

**Returns** Returns the `autoprotocol.instruction.CountCells` instruction created from the specified parameters

**Return type** `CountCells`

**Raises** `TypeError` – `wells` specified is not of a valid input type

#### **cover** (`ref, lid=None, retrieve_lid=None`)

Place specified lid type on specified container

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-flat",
                      storage="warm_37")
p.cover(sample_plate, lid="universal")
```

Autoprotocol Output:

```
"instructions": [
    {
        "lid": "universal",
        "object": "sample_plate",
        "op": "cover"
    }
]
```

#### **Parameters**

- **ref** (`Container`) – Container to be covered.
- **lid** (`str, optional`) – Type of lid to cover the container. Must be a valid lid type for the container type.
- **retrieve\_lid** (`bool, optional`) – Flag to retrieve lid from previously stored location (see `uncover`).

**Returns** Returns the `autoprotocol.instruction.Cover` instruction created from the specified parameters

**Return type** `Cover`

**Raises**

- `TypeError` – If `ref` is not of type `Container`.
- `RuntimeError` – If container type does not have `cover` capability.
- `RuntimeError` – If `lid` is not a valid lid type.

- `RuntimeError` – If container is already sealed with a seal.
- `TypeError` – If `retrieve_lid` is not a boolean.

**`dispense`**(*ref*, *reagent*, *columns*, *is\_resource\_id=False*, *step\_size='5:uL'*, *flowrate=None*, *nozzle\_position=None*, *pre\_dispense=None*, *shape=None*, *shake\_after=None*)  
Dispense specified reagent to specified columns.

Example Usage:

```
from autoprotocol.liquid_handle.liquid_handle_builders import *
from autoprotocol.instructions import Dispense
from autoprotocol import Protocol

p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-flat",
                      storage="warm_37")

p.dispense(sample_plate,
            "water",
            Dispense.builders.columns(
                [Dispense.builders.column(0, "10:uL"),
                 Dispense.builders.column(1, "20:uL"),
                 Dispense.builders.column(2, "30:uL"),
                 Dispense.builders.column(3, "40:uL"),
                 Dispense.builders.column(4, "50:uL")]
            ))
        )

p.dispense(
    sample_plate,
    "water",
    Dispense.builders.columns(
        [Dispense.builders.column(0, "10:uL")]
    ),
    Dispense.builders.nozzle_position(
        position_x=Unit("1:mm"),
        position_y=Unit("2:mm"),
        position_z=Unit("20:mm")
    ),
    shape_builder(
        rows=8, columns=1, format="SBS96"
    )
)
```

Autoprotocol Output:

```
"instructions": [
    {
        "reagent": "water",
        "object": "sample_plate",
        "columns": [
            {
                "column": 0,
                "volume": "10:microliter"
            },
            {

```

(continues on next page)

(continued from previous page)

```

        "column": 1,
        "volume": "20:microliter"
    },
    {
        "column": 2,
        "volume": "30:microliter"
    },
    {
        "column": 3,
        "volume": "40:microliter"
    },
    {
        "column": 4,
        "volume": "50:microliter"
    }
],
"op": "dispense"
},
{
    "reagent": "water",
    "object": "sample_plate",
    "columns": [
        {
            "column": 0,
            "volume": "10:microliter"
        }
    ],
    "nozzle_position" : {
        "position_x" : "1:millimeter",
        "position_y" : "2:millimeter",
        "position_z" : "20:millimeter"
    },
    "shape" : {
        "rows" : 8,
        "columns" : 1,
        "format" : "SBS96"
    }
    "op": "dispense"
},
]

```

## Parameters

- **ref** ([Container](#)) – Container for reagent to be dispensed to.
- **reagent** (*str or well*) – Reagent to be dispensed. Use a string to specify the name or resource\_id (see below) of the reagent to be dispensed. Alternatively, use a well to specify that the dispense operation must be executed using a specific aliquot as the dispense source.
- **columns** (*list(dict("column": int, "volume": str/Unit))*) – Columns to be dispensed to, in the form of a list(dict) specifying the column number and the volume to be dispensed to that column. Columns are expressed as integers indexed from 0. [{“column”: <column num>, “volume”: <volume>}, ...]
- **is\_resource\_id** (*bool, optional*) – If true, interprets reagent as a resource ID
- **step\_size** (*str or Unit, optional*) – Specifies that the dispense operation

must be executed using a peristaltic pump with the given step size. Note that the volume dispensed in each column must be an integer multiple of the step\_size. Currently, step\_size must be either 5 uL or 0.5 uL. If set to None, will use vendor specified defaults.

- **flowrate** (str or Unit, optional) – The rate at which liquid is dispensed into the ref in units of volume/time.
- **nozzle\_position** (dict, optional) – A dict represent nozzle offsets from the bottom middle of the plate's wells. see Dispense.builders.nozzle\_position; specified as {"position\_x": Unit, "position\_y": Unit, "position\_z": Unit}.
- **pre\_dispense** (str or Unit, optional) – The volume of reagent to be dispensed per-nozzle into waste immediately prior to dispensing into the ref.
- **shape** (dict, optional) – The shape of the dispensing head to be used for the dispense. See liquid\_handle\_builders.shape\_builder; specified as {"rows": int, "columns": int, "format": str} with format being a valid SBS format.
- **shake\_after** (dict, optional) – Parameters that specify how a plate should be shaken at the very end of the instruction execution. See Dispense.builders.shake\_after.

**Returns** Returns the `autoprotocol.instruction.Dispense` instruction created from the specified parameters

**Return type** `Dispense`

**Raises**

- `TypeError` – Invalid input types, e.g. ref is not of type Container
- `ValueError` – Columns specified is invalid for this container type
- `ValueError` – Invalid step-size given
- `ValueError` – Invalid pre-dispense volume

```
dispense_full_plate(ref, reagent, volume, is_resource_id=False, step_size='5:uL',
                      flowrate=None, nozzle_position=None, pre_dispense=None, shape=None,
                      shake_after=None)
```

Dispense the specified amount of the specified reagent to every well of a container using a reagent dispenser.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-flat",
                     storage="warm_37")

p.dispense_full_plate(sample_plate,
                      "water",
                      "100:microliter")
```

Autoprotocol Output:

```
"instructions": [
  {
    "reagent": "water",
    "object": "sample_plate",
    "columns": [
```

(continues on next page)

(continued from previous page)

```
{  
    "column": 0,  
    "volume": "100:microliter"  
},  
{  
    "column": 1,  
    "volume": "100:microliter"  
},  
{  
    "column": 2,  
    "volume": "100:microliter"  
},  
{  
    "column": 3,  
    "volume": "100:microliter"  
},  
{  
    "column": 4,  
    "volume": "100:microliter"  
},  
{  
    "column": 5,  
    "volume": "100:microliter"  
},  
{  
    "column": 6,  
    "volume": "100:microliter"  
},  
{  
    "column": 7,  
    "volume": "100:microliter"  
},  
{  
    "column": 8,  
    "volume": "100:microliter"  
},  
{  
    "column": 9,  
    "volume": "100:microliter"  
},  
{  
    "column": 10,  
    "volume": "100:microliter"  
},  
{  
    "column": 11,  
    "volume": "100:microliter"  
}  
],  
"op": "dispense"  
}  
]
```

## Parameters

- **ref** ([Container](#)) – Container for reagent to be dispensed to.

- **reagent** (`str or Well`) – Reagent to be dispensed. Use a string to specify the name or resource\_id (see below) of the reagent to be dispensed. Alternatively, use a well to specify that the dispense operation must be executed using a specific aliquot as the dispense source.
- **volume** (`Unit or str`) – Volume of reagent to be dispensed to each well
- **is\_resource\_id** (`bool, optional`) – If true, interprets reagent as a resource ID
- **step\_size** (`str or Unit, optional`) – Specifies that the dispense operation must be executed using a peristaltic pump with the given step size. Note that the volume dispensed in each column must be an integer multiple of the step\_size. Currently, step\_size must be either 5 uL or 0.5 uL. If set to None, will use vendor specified defaults.
- **flowrate** (`str or Unit, optional`) – The rate at which liquid is dispensed into the ref in units of volume/time.
- **nozzle\_position** (`dict, optional`) – A dict represent nozzle offsets from the bottom middle of the plate's wells. see Dispense.builders.nozzle\_position; specified as {"position\_x": Unit, "position\_y": Unit, "position\_z": Unit}.
- **pre\_dispense** (`str or Unit, optional`) – The volume of reagent to be dispensed per-nozzle into waste immediately prior to dispensing into the ref.
- **shape** (`dict, optional`) – The shape of the dispensing head to be used for the dispense. See liquid\_handle\_builders.shape\_builder; specified as {"rows": int, "columns": int, "format": str} with format being a valid SBS format.
- **shake\_after** (`dict, optional`) – Parameters that specify how a plate should be shaken at the very end of the instruction execution. See Dispense.builders.shake\_after.

**Returns** Returns the `autoprotocol.instruction.Dispense` instruction created from the specified parameters

**Return type** `Dispense`

#### `flash_freeze` (`container, duration`)

Flash freeze the contents of the specified container by submerging it in liquid nitrogen for the specified amount of time.

Example Usage:

```
p = Protocol()

sample = p.ref("liquid_sample", None, "micro-1.5", discard=True)
p.flash_freeze(sample, "25:second")
```

Autoprotocol Output:

```
{
  "refs": {
    "liquid_sample": {
      "new": "micro-1.5",
      "discard": true
    }
  },
  "instructions": [
    {
      "duration": "25:second",
      "object": "liquid_sample",
      "type": "flash_freeze"
    }
  ]
}
```

(continues on next page)

(continued from previous page)

```

        "op": "flash_freeze"
    }
}
}
```

## Parameters

- **container** (`Container or str`) – Container to be flash frozen.
- **duration** (`str or Unit`) – Duration to submerge specified container in liquid nitrogen.

**Returns** Returns the `autoprotocol.instruction.FlashFreeze` instruction created from the specified parameters

**Return type** `FlashFreeze`

**flow\_analyze** (`dataref, FSC, SSC, neg_controls, samples, colors=None, pos_controls=None`)

Perform flow cytometry. The instruction will be executed within the voltage range specified for each channel, optimized for the best sample separation/distribution that can be achieved within these limits. The vendor will specify the device that this instruction is executed on and which excitation and emission spectra are available. At least one negative control is required, which will be used to define data acquisition parameters as well as to determine any autofluorescent properties for the sample set. Additional negative positive control samples are optional. Positive control samples will be used to optimize single color signals and, if desired, to minimize bleed into other channels.

For each sample this instruction asks you to specify the *volume* and/or *captured\_events*. Vendors might also require *captured\_events* in case their device does not support volumetric sample intake. If both conditions are supported, the vendor will specify if data will be collected only until the first one is met or until both conditions are fulfilled.

Example Usage:

```

p = Protocol()
dataref = "test_ref"
FSC = {"voltage_range": {"low": "230:volt", "high": "280:volt"},
        "area": True, "height": True, "weight": False}
SSC = {"voltage_range": {"low": "230:volt", "high": "280:volt"},
        "area": True, "height": True, "weight": False}
neg_controls = {"well": "well0", "volume": "100:microliter",
                "captured_events": 5, "channel": "channel0"}
samples = [
    {
        "well": "well0",
        "volume": "100:microliter",
        "captured_events": 9
    }
]

p.flow_analyze(dataref, FSC, SSC, neg_controls,
                samples, colors=None, pos_controls=None)
```

Autoprotocol Output:

```
{
    "channels": {
        "FSC": {
```

(continues on next page)

(continued from previous page)

```

    "voltage_range": {
        "high": "280:volt",
        "low": "230:volt"
    },
    "area": true,
    "height": true,
    "weight": false
},
"SSC": {
    "voltage_range": {
        "high": "280:volt",
        "low": "230:volt"
    },
    "area": true,
    "height": true,
    "weight": false
}
},
"op": "flow_analyze",
"negative_controls": {
    "channel": "channel0",
    "well": "well0",
    "volume": "100:microliter",
    "captured_events": 5
},
"dataref": "test_ref",
"samples": [
    {
        "well": "well0",
        "volume": "100:microliter",
        "captured_events": 9
    }
]
}
}

```

## Parameters

- **dataref** (*str*) – Name of flow analysis dataset generated.
- **FSC** (*dict*) – Dictionary containing FSC channel parameters in the form of:

```
{
    "voltage_range": {
        "low": "230:volt",
        "high": "280:volt"
    },
    "area": true,           //default: true
    "height": true,         //default: true
    "weight": false         //default: false
}
```

- **SSC** (*dict*) – Dictionary of SSC channel parameters in the form of:

```
{
    "voltage_range": {
        "low": <voltage>,
        "high": <voltage>
    }
}
```

(continues on next page)

(continued from previous page)

```
},
  "area": true,           //default: true
  "height": true,         //default: false
  "weight": false         //default: false
}
```

- **neg\_controls** (*list (dict)*) – List of negative control wells in the form of:

```
{
  "well": well,
  "volume": volume,
  "captured_events": integer,    // optional, default infinity
  "channel": [channel_name]
}
```

at least one negative control is required.

- **samples** (*list (dict)*) – List of samples in the form of:

```
{
  "well": well,
  "volume": volume,
  "captured_events": integer    // optional, default infinity
}
```

at least one sample is required

- **colors** (*list (dict)*, *optional*) – Optional list of colors in the form of:

```
[
  {
    "name": "FitC",
    "emission_wavelength": "495:nanometer",
    "excitation_wavelength": "519:nanometer",
    "voltage_range": {
      "low": <voltage>,
      "high": <voltage>
    },
    "area": true,           //default: true
    "height": false,        //default: false
    "weight": false         //default: false
  }, ...
]
```

- **pos\_controls** (*list (dict)*, *optional*) – Optional list of positive control wells in the form of:

```
[
  {
    "well": well,
    "volume": volume,
    "captured_events": integer,    // default: infinity
    "channel": [channel_name],
    "minimize_bleed": [{          // optional
      "from": color,
      "to": [color]
    }, ...
  ]
```

**Returns** Returns the `autoprotocol.instruction.FlowAnalyze` instruction created from the specified parameters

**Return type** `FlowAnalyze`

**Raises**

- `TypeError` – If inputs are not of the correct type.
- `UnitError` – If unit inputs are not properly formatted.
- `AssertionError` – If required parameters are missing.
- `ValueError` – If volumes are not correctly formatted or present.

**fluorescence** (`ref, wells, excitation, emission, dataref, flashes=25, temperature=None, gain=None, incubate_before=None, detection_mode=None, position_z=None, settle_time=None, lag_time=None, integration_time=None`)

Read the fluorescence for the indicated wavelength for the indicated wells. Append a Fluorescence instruction to the list of instructions for this Protocol object.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-flat",
                     storage="warm_37")

p.fluorescence(sample_plate, sample_plate.wells_from(0,12),
               excitation="587:nanometer", emission="610:nanometer",
               dataref="test_reading")
```

Autoprotocol Output:

```
"instructions": [
    {
        "dataref": "test_reading",
        "excitation": "587:nanometer",
        "object": "sample_plate",
        "emission": "610:nanometer",
        "wells": [
            "A1",
            "A2",
            "A3",
            "A4",
            "A5",
            "A6",
            "A7",
            "A8",
            "A9",
            "A10",
            "A11",
            "A12"
        ],
        "num_flashes": 25,
        "op": "fluorescence"
    }
]
```

## Parameters

- **ref** (*str or Container*) – Container to plate read.
- **wells** (*list(Well) or WellGroup or Well*) – WellGroup of wells to be measured or a list of well references in the form of [“A1”, “B1”, “C5”, ...]
- **excitation** (*str or Unit*) – Wavelength of light used to excite the wells indicated
- **emission** (*str or Unit*) – Wavelength of light to be measured for the indicated wells
- **dataref** (*str*) – Name of this specific dataset of measured fluorescence
- **flashes** (*int, optional*) – Number of flashes.
- **temperature** (*str or Unit, optional*) – set temperature to heat plate reading chamber
- **gain** (*float, optional*) – float between 0 and 1, multiplier, gain=0.2 of maximum signal amplification
- **incubate\_before** (*dict, optional*) – incubation prior to reading if desired
- **detection\_mode** (*str, optional*) – set the detection mode of the optics, [“top”, “bottom”], defaults to vendor specified defaults.
- **position\_z** (*dict, optional*) – distance from the optics to the surface of the plate transport, only valid for “top” detection\_mode and vendor capabilities. Specified as either a set distance - “manual”, OR calculated from a WellGroup - “calculated\_from\_wells”. Only one position\_z determination may be specified

```
position_z = {
    "manual": Unit
    - OR -
    "calculated_from_wells": []
}
```

- **settle\_time** (*Unit, optional*) – the time before the start of the measurement, defaults to vendor specifications
- **lag\_time** (*Unit, optional*) – time between flashes and the start of the signal integration, defaults to vendor specifications
- **integration\_time** (*Unit, optional*) – duration of the signal recording, per Well, defaults to vendor specifications

incubate\_before example:

```
{
    "shaking": {
        "amplitude": str or Unit
        "orbital": bool
    },
    "duration": str or Unit
}
```

position\_z examples:

```
position_z = {
    "calculated_from_wells": ["plate/A1", "plate/A2"]
}
```

(continues on next page)

(continued from previous page)

```
-OR-

position_z = {
    "manual": "20:micrometer"
}
```

**Returns** Returns the `autoprotocol.instruction.Fluorescence` instruction created from the specified parameters

**Return type** `Fluorescence`

**Raises**

- `TypeError` – Invalid input types, e.g. wells given is of type Well, WellGroup or list of wells
- `ValueError` – Wells specified are not from the same container
- `ValueError` – Settle time, integration time or lag time has to be greater than 0
- `UnitError` – Settle time, integration time, lag time or position z is not of type Unit
- `ValueError` – Unknown value given for `detection_mode`
- `ValueError` – Position z specified for non-top detection mode
- `KeyError` – For position\_z, only `manual` and `calculated_from_wells` is allowed
- `NotImplementedError` – Specifying `calculated_from_wells` as that has not been implemented yet

### `gel_purify(extracts, volume, matrix, ladder, dataref)`

Separate nucleic acids on an agarose gel and purify according to parameters. If gel extract lanes are not specified, they will be sequentially ordered and purified on as many gels as necessary.

Each element in extracts specifies a source loaded in a single lane of gel with a list of bands that will be purified from that lane. If the same source is to be run on separate lanes, a new dictionary must be added to extracts. It is also possible to add an element to extract with a source but without a list of bands. In that case, the source will be run in a lane without extraction.

Example Usage:

```
p = Protocol()
sample_wells = p.ref("test_plate", None, "96-pcr",
                     discard=True).wells_from(0, 8)
extract_wells = [p.ref("extract_" + str(i.index), None,
                      "micro-1.5", storage="cold_4").well(0)
                 for i in sample_wells]

extracts = [make_gel_extract_params(
            w,
            make_band_param(
                "TE",
                "5:microliter",
                80,
                79,
                extract_wells[i]))
            for i, w in enumerate(sample_wells)]
```

(continues on next page)

(continued from previous page)

```
p.gel_purify(extracts, "10:microliter",
              "size_select(8,0.8%)", "ladder1",
              "gel_purify_example")
```

Autoprotocol Output:

For extracts[0]

```
{
    "band_list": [
        {
            "band_size_range": {
                "max_bp": 80,
                "min_bp": 79
            },
            "destination": Well(Container(extract_0), 0, None),
            "elution_buffer": "TE",
            "elution_volume": "Unit(5.0, 'microliter')"
        }
    ],
    "gel": None,
    "lane": None,
    "source": Well(Container(test_plate), 0, None)
}
```

## Parameters

- **extracts** (*list(dict)*) – Dictionary containing parameters for gel extraction, must be in the form of:

```
[
    {
        "band_list": [
            {
                "band_size_range": {
                    "max_bp": int,
                    "min_bp": int
                },
                "destination": Well,
                "elution_buffer": str,
                "elution_volume": Volume
            }
        ],
        "gel": int or None,
        "lane": int or None,
        "source": Well
    }
]
```

`util.make_gel_extract_params()` and `util.make_band_param()` can be used to create these dictionaries

**band\_list: list(dict)** List of bands to be extracted from the lane

**band\_size\_range: dict** Dictionary for the size range of the band to be extracted

**max\_bp: int** Maximum size for the band

**min\_bp: int** Minimum size for the band  
**destination: Well** Well to place the extracted material  
**elution\_buffer: str** Buffer to use to extract the band, commonly “water”  
**elution\_volume: str or Unit** Volume of elution\_buffer to extract the band into  
**gel: int** Integer identifier for the gel if using multiple gels  
**lane: int** Integer identifier for the lane of a gel to run the source  
**source: Well** Well from which to purify the material

- **volume (str or Unit)** – Volume of liquid to be transferred from each well specified to a lane of the gel.
- **matrix (str)** – Matrix (gel) in which to gel separate samples
- **ladder (str)** – Ladder by which to measure separated fragment size
- **dataref (str)** – Name of this set of gel separation results.

**Returns** Returns the `autoprotocol.instruction.GelPurify` instruction created from the specified parameters

**Return type** `GelPurify`

**Raises**

- `RuntimeError` – If matrix is not properly formatted.
- `AttributeError` – If extract parameters are not a list of dictionaries.
- `KeyError` – If extract parameters do not contain the specified parameter keys.
- `ValueError` – If `min_bp` is greater than `max_bp`.
- `ValueError` – If extract destination is not of type `Well`.
- `ValueError` – If extract elution volume is not of type `Unit`
- `ValueError` – if extract elution volume is not greater than 0.
- `RuntimeError` – If gel extract lanes are set for some but not all extract wells.
- `RuntimeError` – If all samples do not fit on single gel type.
- `TypeError` – If lane designated for gel extracts is not an integer.
- `RuntimeError` – If designated lane index is outside lanes within the gel.
- `RuntimeError` – If lanes not designated and number of extracts not equal to number of samples.

### `gel_separate(wells, volume, matrix, ladder, duration, dataref)`

Separate nucleic acids on an agarose gel.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-flat",
                     storage="warm_37")

p.gel_separate(sample_plate.wells_from(0, 12), "10:microliter",
```

(continues on next page)

(continued from previous page)

```
"agarose(8,0.8%)", "ladder1", "11:minute",
"genotyping_030214")
```

## Autoprotocol Output:

```
"instructions": [
    {
        "dataref": "genotyping_030214",
        "matrix": "agarose(8,0.8%)",
        "volume": "10:microliter",
        "ladder": "ladder1",
        "objects": [
            "sample_plate/0",
            "sample_plate/1",
            "sample_plate/2",
            "sample_plate/3",
            "sample_plate/4",
            "sample_plate/5",
            "sample_plate/6",
            "sample_plate/7",
            "sample_plate/8",
            "sample_plate/9",
            "sample_plate/10",
            "sample_plate/11"
        ],
        "duration": "11:minute",
        "op": "gel_separate"
    }
]
```

## Parameters

- **wells** (*list (Well) or WellGroup or Well*) – List of wells or WellGroup containing wells to be separated on gel.
- **volume** (*str or Unit*) – Volume of liquid to be transferred from each well specified to a lane of the gel.
- **matrix** (*str*) – Matrix (gel) in which to gel separate samples
- **ladder** (*str*) – Ladder by which to measure separated fragment size
- **duration** (*str or Unit*) – Length of time to run current through gel.
- **dataref** (*str*) – Name of this set of gel separation results.

**Returns** Returns the `autoprotocol.instruction.GelSeparate` instruction created from the specified parameters

**Return type** `GelSeparate`

## Raises

- `TypeError` – Invalid input types, e.g. wells given is of type Well, WellGroup or list of wells
- `ValueError` – Specifying more wells than the number of available lanes in the selected matrix

**get\_instruction\_index()**

Get index of the last appended instruction

Example Usage:

```
p = Protocol()
plate_1 = p.ref("plate_1", id=None, cont_type="96-flat",
                 discard=True)

p.cover(plate_1)
time_point_1 = p.get_instruction_index() # time_point_1 = 0
```

**Raises** ValueError – If an instruction index is less than 0

**Returns** Index of the preceding instruction

**Return type** int

**illuminaeq(*flowcell*, *lanes*, *sequencer*, *mode*, *index*, *library\_size*, *dataref*, *cycles*=None)**

Load aliquots into specified lanes for Illumina sequencing. The specified aliquots should already contain the appropriate mix for sequencing and require a library concentration reported in ng/uL.

Example Usage:

```
p = Protocol()
sample_wells = p.ref(
    "test_plate", None, "96-pcr", discard=True).wells_from(0, 8)

p.illuminaeq(
    "PE",
    [
        {"object": sample_wells[0], "library_concentration": 1.0},
        {"object": sample_wells[1], "library_concentration": 5.32},
        {"object": sample_wells[2], "library_concentration": 54},
        {"object": sample_wells[3], "library_concentration": 20},
        {"object": sample_wells[4], "library_concentration": 23},
        {"object": sample_wells[5], "library_concentration": 23},
        {"object": sample_wells[6], "library_concentration": 21},
        {"object": sample_wells[7], "library_concentration": 62}
    ],
    "hiseq", "rapid", 'none', 250, "my_illumina")
```

Autoprotocol Output:

```
"instructions": [
    {
        "dataref": "my_illumina",
        "index": "none",
        "lanes": [
            {
                "object": "test_plate/0",
                "library_concentration": 1
            },
            {
                "object": "test_plate/1",
                "library_concentration": 5.32
            },
            {

```

(continues on next page)

(continued from previous page)

```

        "object": "test_plate/2",
        "library_concentration": 54
    },
    {
        "object": "test_plate/3",
        "library_concentration": 20
    },
    {
        "object": "test_plate/4",
        "library_concentration": 23
    },
    {
        "object": "test_plate/5",
        "library_concentration": 23
    },
    {
        "object": "test_plate/6",
        "library_concentration": 21
    },
    {
        "object": "test_plate/7",
        "library_concentration": 62
    }
],
"flowcell": "PE",
"mode": "mid",
"sequencer": "hiseq",
"library_size": 250,
"op": "illumina_sequence"
}
]

```

## Parameters

- **flowcell** (*str*) – Flowcell designation: “SR” or ” “PE”
- **lanes** (*list (dict)*) –

```

"lanes": [
{
    "object": aliquot, Well,
    "library_concentration": decimal, // ng/uL
},
{...}]

```

- **sequencer** (*str*) – Sequencer designation: “miseq”, “hiseq” or “nextseq”
- **mode** (*str*) – Mode designation: “rapid”, “mid” or “high”
- **index** (*str*) – Index designation: “single”, “dual” or “none”
- **library\_size** (*int*) – Library size expressed as an integer of basepairs
- **dataref** (*str*) – Name of sequencing dataset that will be returned.
- **cycles** (*Enum({ "read\_1", "read\_2", "index\_1", "index\_2" })*) – Parameter specific to Illuminaseq read-length or number of sequenced bases. Refer to the ASC for more details

**Returns** Returns the `autoprotocol.instruction.IlluminaSeq` instruction created from the specified parameters

**Return type** `IlluminaSeq`

**Raises**

- `TypeError` – If index and dataref are not of type str.
- `TypeError` – If library\_concentration is not a number.
- `TypeError` – If library\_size is not an integer.
- `ValueError` – If flowcell, sequencer, mode, index are not of type a valid option.
- `ValueError` – If number of lanes specified is more than the maximum lanes of the specified type of sequencer.
- `KeyError` – Invalid keys specified for cycles parameter

**image\_plate** (*ref*, *mode*, *dataref*)  
Capture an image of the specified container.

Example Usage:

```
p = Protocol()

agar_plate = p.ref("agar_plate", None, "1-flat", discard=True)
bact = p.ref("bacteria", None, "micro-1.5", discard=True)

p.spread(bact.well(0), agar_plate.well(0), "55:microliter")
p.incubate(agar_plate, "warm_37", "18:hour")
p.image_plate(agar_plate, mode="top", dataref="my_plate_image_1")
```

Autoprotocol Output:

```
{
  "refs": {
    "bacteria": {
      "new": "micro-1.5",
      "discard": true
    },
    "agar_plate": {
      "new": "1-flat",
      "discard": true
    }
  },
  "instructions": [
    {
      "volume": "55.0:microliter",
      "to": "agar_plate/0",
      "from": "bacteria/0",
      "op": "spread"
    },
    {
      "where": "warm_37",
      "object": "agar_plate",
      "co2_percent": 0,
      "duration": "18:hour",
      "shaking": false,
      "op": "incubate"
    }
  ]
}
```

(continues on next page)

(continued from previous page)

```

},
{
  "dataref": "my_plate_image_1",
  "object": "agar_plate",
  "mode": "top",
  "op": "image_plate"
}
]
}

```

### Parameters

- **ref** (*str or Container*) – Container to take image of
- **mode** (*str*) – Imaging mode (currently supported: “top”)
- **dataref** (*str*) – Name of data reference of resulting image

**Returns** Returns the `autoprotocol.instruction.ImagePlate` instruction created from the specified parameters

**Return type** `ImagePlate`

**incubate** (*ref, where, duration, shaking=False, co2=0, uncovered=False, target\_temperature=None, shaking\_params=None*)

Move plate to designated thermoisolater or ambient area for incubation for specified duration.

Example Usage:

```

p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-pcr",
                      storage="warm_37")

# a plate must be sealed/covered before it can be incubated
p.seal(sample_plate)
p.incubate(sample_plate, "warm_37", "1:hour", shaking=True)

```

Autoprotocol Output:

```

"instructions": [
  {
    "object": "sample_plate",
    "op": "seal"
  },
  {
    "duration": "1:hour",
    "where": "warm_37",
    "object": "sample_plate",
    "shaking": true,
    "op": "incubate",
    "co2_percent": 0
  }
]

```

### Parameters

- **ref** (`Ref or str`) – The container to be incubated
- **where** (`Enum({ "ambient", "warm_37", "cold_4", "cold_20", "cold_80" })`) – Temperature at which to incubate specified container
- **duration** (`Unit or str`) – Length of time to incubate container
- **shaking** (`bool, optional`) – Specify whether or not to shake container if available at the specified temperature
- **co2** (`int, optional`) – Carbon dioxide percentage
- **uncovered** (`bool, optional`) – Specify whether the container should be uncovered during incubation
- **target\_temperature** (`Unit or str, optional`) – Specify a target temperature for a device (eg. an incubating block) to reach during the specified duration.
- **shaking\_params** (`dict, optional`) – Specify “path” and “frequency” of shaking parameters to be used with compatible devices (eg. thermoshakes)

**Returns** Returns the `autoprotocol.instruction.Incubate` instruction created from the specified parameters

**Return type** `Incubate`

**Raises**

- `TypeError` – Invalid input types given, e.g. ref is not of type Container
- `RuntimeError` – Incubating uncovered in a location which is shaking

**luminescence** (`ref, wells, dataref, incubate_before=None, temperature=None, settle_time=None, integration_time=None`)

Read luminescence of indicated wells.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-flat",
                     storage="warm_37")

p.luminescence(sample_plate, sample_plate.wells_from(0,12),
               "test_reading")
```

Autoprotocol Output:

```
"instructions": [
    {
        "dataref": "test_reading",
        "object": "sample_plate",
        "wells": [
            "A1",
            "A2",
            "A3",
            "A4",
            "A5",
            "A6",
            "A7",
            "A8",
```

(continues on next page)

(continued from previous page)

```

        "A9",
        "A10",
        "A11",
        "A12"
    ],
    "op": "luminescence"
}
]

```

## Parameters

- **ref** (*str or Container*) – Container to plate read.
- **wells** (*list (Well) or WellGroup or Well*) – WellGroup of wells to be measured or a list of well references in the form of ["A1", "B1", "C5", ...]
- **dataref** (*str*) – Name of this dataset of measured luminescence readings.
- **temperature** (*str or Unit, optional*) – set temperature to heat plate reading chamber
- **settle\_time** (*Unit, optional*) – the time before the start of the measurement, defaults to vendor specifications
- **incubate\_before** (*dict, optional*) – incubation prior to reading if desired
- **integration\_time** (*Unit, optional*) – duration of the signal recording, per Well, defaults to vendor specifications

```
{
    "shaking": {
        "amplitude": str or Unit
        "orbital": bool
    },
    "duration": str or Unit
}
```

**Returns** Returns the `autoprotocol.instruction.Luminescence` instruction created from the specified parameters

**Return type** `Luminescence`

## Raises

- `TypeError` – Invalid input types, e.g. wells given is of type Well, WellGroup or list of wells
- `ValueError` – Wells specified are not from the same container
- `ValueError` – Settle time or integration time has to be greater than 0
- `UnitError` – Settle time or integration time is not of type Unit

**mag\_collect** (*head, container, cycles, pause\_duration, bottom\_position=0.0, temperature=None, new\_tip=False, new\_instruction=False*)

Collect beads from a container by cycling magnetized tips in and out of the container with an optional pause at the bottom of the insertion.

Example Usage:

```
p = Protocol()
plate = p.ref("plate_0", None, "96-pcr", storage="cold_20")

p.mag_collect("96-pcr", plate, 5, "30:second", bottom_position=
    0.0, temperature=None, new_tip=False,
    new_instruction=False)
```

Autoprotocol Output:

```
"instructions": [
    {
        "groups": [
            [
                {
                    "collect": {
                        "bottom_position": 0,
                        "object": "plate_0",
                        "temperature": null,
                        "cycles": 5,
                        "pause_duration": "30:second"
                    }
                }
            ]
        ],
        "magnetic_head": "96-pcr",
        "op": "magnetic_transfer"
    }
]
```

## Parameters

- **head** (*str*) – Magnetic head to use for the magnetic bead transfers
- **container** (*Container*) – Container to incubate beads
- **cycles** (*int*) – Number of cycles to raise and lower tips
- **pause\_duration** (*str or Unit*) – Time tips are paused in bottom position each cycle
- **bottom\_position** (*float*) – Position relative to well height that tips are held during pause
- **temperature** (*str or Unit*) – Temperature heat block is set at
- **new\_tip** (*bool*) – Specify whether to use a new tip to complete the step
- **new\_instruction** (*bool*) – Specify whether to create a new `magnetic_transfer` instruction

**Returns** Returns the `autoprotocol.instruction.MagneticTransfer` instruction created from the specified parameters

**Return type** `MagneticTransfer`

**mag\_dry** (*head, container, duration, new\_tip=False, new\_instruction=False*)

Dry beads with magnetized tips above and outside a container for a set time.

Example Usage:

```
p = Protocol()
plate = p.ref("plate_0", None, "96-pcr", storage="cold_20")

p.mag_dry("96-pcr", plate, "30:minute", new_tip=False,
          new_instruction=False)
```

Autoprotocol Output:

```
"instructions": [
    {
        "groups": [
            [
                {
                    "dry": {
                        "duration": "30:minute",
                        "object": "plate_0"
                    }
                }
            ],
            "magnetic_head": "96-pcr",
            "op": "magnetic_transfer"
        }
    ]
]
```

## Parameters

- **head** (*str*) – Magnetic head to use for the magnetic bead transfers
- **container** (*Container*) – Container to dry beads above
- **duration** (*str or Unit*) – Time for drying
- **new\_tip** (*bool*) – Specify whether to use a new tip to complete the step
- **new\_instruction** (*bool*) – Specify whether to create a new `magnetic_transfer` instruction

**Returns** Returns the `autoprotocol.instruction.MagneticTransfer` instruction created from the specified parameters

**Return type** `MagneticTransfer`

**mag\_incubate** (*head, container, duration, magnetize=False, tip\_position=1.5, temperature=None, new\_tip=False, new\_instruction=False*)  
Incubate the container for a set time with tips set at *tip\_position*.

Example Usage:

```
p = Protocol()
plate = p.ref("plate_0", None, "96-pcr", storage="cold_20")

p.mag_incubate("96-pcr", plate, "30:minute", magnetize=False,
               tip_position=1.5, temperature=None, new_tip=False)
```

Autoprotocol Output:

```
"instructions": [
    {
```

(continues on next page)

(continued from previous page)

```

"groups": [
    [
        {
            "incubate": {
                "duration": "30:minute",
                "tip_position": 1.5,
                "object": "plate_0",
                "magnetize": false,
                "temperature": null
            }
        }
    ],
    "magnetic_head": "96-pcr",
    "op": "magnetic_transfer"
}
]

```

## Parameters

- **head** (*str*) – Magnetic head to use for the magnetic bead transfers
- **container** (*Container*) – Container to incubate beads
- **duration** (*str or Unit*) – Time for incubation
- **magnetize** (*bool*) – Specify whether to magnetize the tips
- **tip\_position** (*float*) – Position relative to well height that tips are held
- **temperature** (*str or Unit*) – Temperature heat block is set at
- **new\_tip** (*bool*) – Specify whether to use a new tip to complete the step
- **new\_instruction** (*bool*) – Specify whether to create a new `magnetic_transfer` instruction

**Returns** Returns the `autoprotocol.instruction.MagneticTransfer` instruction created from the specified parameters

**Return type** `MagneticTransfer`

**mag\_mix** (*head, container, duration, frequency, center=0.5, amplitude=0.5, magnetize=False, temperature=None, new\_tip=False, new\_instruction=False*)

Mix beads in a container by cycling tips in and out of the container.

Example Usage:

```

p = Protocol()
plate = p.ref("plate_0", None, "96-pcr", storage="cold_20")

p.mag_mix("96-pcr", plate, "30:second", "60:hertz", center=0.75,
          amplitude=0.25, magnetize=True, temperature=None,
          new_tip=False, new_instruction=False)

```

Autoprotocol Output:

```

"instructions": [
{
    "groups": [

```

(continues on next page)

(continued from previous page)

```
[  
  {  
    "mix": {  
      "center": 0.75,  
      "object": "plate_0",  
      "frequency": "2:hertz",  
      "amplitude": 0.25,  
      "duration": "30:second",  
      "magnetize": true,  
      "temperature": null  
    }  
  }  
]  
,  
"magnetic_head": "96-pcr",  
"op": "magnetic_transfer"  
}  
]
```

## Parameters

- **head** (*str*) – Magnetic head to use for the magnetic bead transfers
- **container** (*Container*) – Container to incubate beads
- **duration** (*str or Unit*) – Total time for this sub-operation
- **frequency** (*str or Unit*) – Cycles per second (hertz) that tips are raised and lowered
- **center** (*float*) – Position relative to well height where oscillation is centered
- **amplitude** (*float*) – Distance relative to well height to oscillate around “center”
- **magnetize** (*bool*) – Specify whether to magnetize the tips
- **temperature** (*str or Unit*) – Temperature heat block is set at
- **new\_tip** (*bool*) – Specify whether to use a new tip to complete the step
- **new\_instruction** (*bool*) – Specify whether to create a new `magnetic_transfer` instruction

**Returns** Returns the `autoprotocol.instruction.MagneticTransfer` instruction created from the specified parameters

**Return type** `MagneticTransfer`

**mag\_release** (*head, container, duration, frequency, center=0.5, amplitude=0.5, temperature=None, new\_tip=False, new\_instruction=False*)

Release beads into a container by cycling tips in and out of the container with tips unmagnetized.

Example Usage:

```
p = Protocol()  
plate = p.ref("plate_0", None, "96-pcr", storage="cold_20")  
  
p.mag_release("96-pcr", plate, "30:second", "60:hertz", center=0.75,  
              amplitude=0.25, temperature=None, new_tip=False,  
              new_instruction=False)
```

## Autoprotocol Output:

```
"instructions": [
    {
        "groups": [
            [
                {
                    "release": {
                        "center": 0.75,
                        "object": "plate_0",
                        "frequency": "2:hertz",
                        "amplitude": 0.25,
                        "duration": "30:second",
                        "temperature": null
                    }
                }
            ],
            "magnetic_head": "96-pcr",
            "op": "magnetic_transfer"
        }
    ]
]
```

**Parameters**

- **head** (*str*) – Magnetic head to use for the magnetic bead transfers
- **container** (*Container*) – Container to incubate beads
- **duration** (*str or Unit*) – Total time for this sub-operation
- **frequency** (*str or Unit*) – Cycles per second (hertz) that tips are raised and lowered
- **center** (*float*) – Position relative to well height where oscillation is centered
- **amplitude** (*float*) – Distance relative to well height to oscillate around “center”
- **temperature** (*str or Unit*) – Temperature heat block is set at
- **new\_tip** (*bool*) – Specify whether to use a new tip to complete the step
- **new\_instruction** (*bool*) – Specify whether to create a new `magnetic_transfer` instruction

**Returns** Returns the `autoprotocol.instruction.MagneticTransfer` instruction created from the specified parameters

**Return type** `MagneticTransfer`

**measure\_concentration** (*wells, dataref, measurement, volume='2:microliter'*)

Measure the concentration of DNA, ssDNA, RNA or protein in the specified volume of the source aliquots.

Example Usage:

```
p = Protocol()

test_plate = p.ref("test_plate", id=None, cont_type="96-flat",
                   storage=None, discard=True)
p.measure_concentration(test_plate.wells_from(0, 3), "mc_test",
                       "DNA")
```

(continues on next page)

(continued from previous page)

```
p.measure_concentration(test_plate.wells_from(3, 3),
    dataref="mc_test2", measurement="protein",
    volume="4:microliter")
```

Autoprotocol Output:

```
{
  "refs": {
    "test_plate": {
      "new": "96-flat",
      "discard": true
    }
  },
  "instructions": [
    {
      "volume": "2.0:microliter",
      "dataref": "mc_test",
      "object": [
        "test_plate/0",
        "test_plate/1",
        "test_plate/2"
      ],
      "op": "measure_concentration",
      "measurement": "DNA"
    }, ...
  ]
}
```

## Parameters

- **wells** (`list(Well)` or `WellGroup` or `Well`) – WellGroup of wells to be measured
- **volume** (`str` or `Unit`) – Volume of sample required for analysis
- **dataref** (`str`) – Name of this specific dataset of measurements
- **measurement** (`str`) – Class of material to be measured. One of [“DNA”, “ssDNA”, “RNA”, “protein”].

**Returns** Returns the `autoprotocol.instruction.MeasureConcentration` instruction created from the specified parameters

**Return type** `MeasureConcentration`

**Raises** `TypeError` – `wells` specified is not of a valid input type

## `measure_mass(container, dataref)`

Measure the mass of a container.

Example Usage:

```
p = Protocol()

test_plate = p.ref("test_plate", id=None, cont_type="96-flat",
    storage=None, discard=True)
p.measure_mass(test_plate, "test_data")
```

Autoprotocol Output:

```
{
    "refs": {
        "test_plate": {
            "new": "96-flat",
            "discard": true
        }
    },
    "instructions": [
        {
            "dataref": "test_data",
            "object": [
                "test_plate"
            ],
            "op": "measure_mass"
        }
    ]
}
```

## Parameters

- **container** (`Container`) – container to be measured
- **dataref** (`str`) – Name of this specific dataset of measurements

**Returns** Returns the `autoprotocol.instruction.MeasureMass` instruction created from the specified parameters

**Return type** `MeasureMass`

**Raises** `TypeError` – Input given is not of type Container

## `measure_volume(wells, dataref)`

Measure the volume of each well in wells.

Example Usage:

```
p = Protocol()

test_plate = p.ref("test_plate", id=None, cont_type="96-flat",
                    storage=None, discard=True)
p.measure_volume(test_plate.from_wells(0,2), "test_data")
```

Autoprotocol Output:

```
{
    "refs": {
        "test_plate": {
            "new": "96-flat",
            "discard": true
        }
    },
    "instructions": [
        {
            "dataref": "test_data",
            "object": [
                "test_plate/0",
                "test_plate/1"
            ],

```

(continues on next page)

(continued from previous page)

```
        "op": "measure_volume"
    }
}
```

### Parameters

- **wells** (`list(Well)` or `WellGroup` or `Well`) – list of wells to be measured
- **dataref** (`str`) – Name of this specific dataset of measurements

**Returns** Returns the `autoprotocol.instruction.MeasureVolume` instruction created from the specified parameters

**Return type** `MeasureVolume`

**Raises** `TypeError` – `wells` specified is not of a valid input type

```
mix(well,           volume,           rows=1,           columns=1,           liquid=<class
     col.liquid_handle.liquid_class.LiquidClass'>,           method=<class
     col.liquid_handle.mix.Mix'>, one_tip=False)
Generates LiquidHandle instructions within wells
```

Mix liquid in specified wells.

### Parameters

- **well** (`Well` or `WellGroup` or `list(Well)`) – Well(s) to be mixed.
- **volume** (`str` or `Unit` or `list(str)` or `list(Unit)`) – Volume(s) of liquid to be mixed within the specified well(s). The number of volume(s) specified must correspond with the number of well(s).
- **rows** (`int`, `optional`) – Number of rows to be concurrently mixed
- **columns** (`int`, `optional`) – Number of columns to be concurrently mixed
- **liquid** (`LiquidClass` or `list(LiquidClass)`, `optional`) – Type(s) of liquid contained in the Well(s). This affects the aspirate and dispense behavior including the flowrates, liquid level detection thresholds, and physical movements.
- **method** (`Mix` or `list(Mix)`, `optional`) – Method(s) with which Integrates with the specified liquid to define a set of physical movements.
- **one\_tip** (`bool`, `optional`) – If True then a single tip will be used for all operations

**Returns** Returns a list of `autoprotocol.instruction.LiquidHandle` instructions created from the specified parameters

**Return type** `list(LiquidHandle)`

### Raises

- `ValueError` – if the specified parameters can't be interpreted as lists of equal length
- `ValueError` – if `one_tip` is true, but not all mix methods have a `tip_type`
- `ValueError` – if the specified volume is larger than the maximum tip capacity of the available liquid\_handling devices for a given mix

## Examples

Mix within a single well

```
from autoprotocol import Protocol, Unit

p = Protocol()
plate = p.ref("example_plate", cont_type="384-flat", discard=True)

p.mix(plate.well(0), "5:ul")
```

Sequential mixes within multiple wells

```
wells = plate.wells_from(0, 8, columnwise=True)
volumes = [Unit(x, "ul") for x in range(1, 9)]
p.mix(wells, volumes)
```

Concurrent mixes within multiple wells

```
# single-column concurrent mix
p.mix(plate.well(0), "5:ul", rows=8)

# 96-well concurrent mix in the A1 quadrant
p.mix(plate.well(0), "5:ul", rows=8, columns=12)

# 96-well concurrent mix in the A2 quadrant
p.mix(plate.well(1), "5:ul", rows=8, columns=12)

# 384-well concurrent mix
p.mix(plate.well(0), "5:ul", rows=16, columns=24)
```

Mix with extra parameters

```
from autoprotocol.liquid_handle import Mix
from autoprotocol.instruction import LiquidHandle

p.mix(
    plate.well(0), "5:ul", rows=8,
    method=Mix(
        mix_params=LiquidHandle.builders.mix(
            )
    )
)
```

See also:

**Mix()** base LiquidHandleMethod for mix operations

**oligosynthesize**(*oligos*)

Specify a list of oligonucleotides to be synthesized and a destination for each product.

Example Usage:

```
oligo_1 = p.ref("oligo_1", None, "micro-1.5", discard=True)

p.oligosynthesize([{ "sequence": "CATGGTCCCCTGCACAGG",
```

(continues on next page)

(continued from previous page)

```
"destination": oligo_1.well(0),
"scale": "25nm",
"purification": "standard"}])
```

Autoprotocol Output:

```
"instructions": [
{
    "oligos": [
        {
            "destination": "oligo_1/0",
            "sequence": "CATGGTCCCCTGCACAGG",
            "scale": "25nm",
            "purification": "standard"
        }
    ],
    "op": "oligosynthesize"
}
]
```

**Parameters** **oligos** (*list (dict)*) – List of oligonucleotides to synthesize. Each dictionary should contain the oligo's sequence, destination, scale and purification

```
[
{
    "destination": "my_plate/A1",
    "sequence": "GATCRYMKSWHBVDN",
    // - standard IUPAC base codes
    // - IDT also allows rX (RNA), mX (2' O-methyl RNA), and
    //   X*/rX*/mX* (phosphorothioated)
    // - they also allow inline annotations for
    //   modifications,
    //   e.g. "GCGACTC/3Phos/" for a 3' phosphorylation
    //   e.g. "aggg/iAzideN/cgcgc" for an
    //   internal modification
    "scale": "25nm" | "100nm" | "250nm" | "1um",
    "purification": "standard" | "page" | "hplc",
    // default: standard
}, ...
]
```

**Returns** Returns the `autoprotocol.instruction.Oligosynthesize` instruction created from the specified parameters

**Return type** `Oligosynthesize`

**provision** (*resource\_id, dests, volumes*)

Provision a commercial resource from a catalog into the specified destination well(s). A new tip is used for each destination well specified to avoid contamination.

**Parameters**

- **resource\_id** (*str*) – Resource ID from catalog.
- **dests** (`Well` or `WellGroup` or *list (Well)*) – Destination(s) for specified resource.

- **volumes** (*str or Unit or list(str) or list(Unit)*) – Volume(s) to transfer of the resource to each destination well. If one volume of specified, each destination well receive that volume of the resource. If destinations should receive different volumes, each one should be specified explicitly in a list matching the order of the specified destinations.

**Raises**

- **TypeError** – If `resource_id` is not a string.
- **RuntimeError** – If length of the list of volumes specified does not match the number of destination wells specified.
- **TypeError** – If volume is not specified as a string or `Unit` (or a list of either)
- **ValueError** – Volume to provision exceeds max capacity of well

**Returns** Returns the `autoprotocol.instruction.Provision` instruction created from the specified parameters

**Return type** `Provision`

**ref** (*name, id=None, cont\_type=None, storage=None, discard=None, cover=None*)

Add a Ref object to the dictionary of Refs associated with this protocol and return a Container with the id, container type and storage or discard conditions specified.

Example Usage:

```
p = Protocol()

# ref a new container (no id specified)
sample_ref_1 = p.ref("sample_plate_1",
                     cont_type="96-pcr",
                     discard=True)

# ref an existing container with a known id
sample_ref_2 = p.ref("sample_plate_2",
                     id="ct1cxaе33lkj",
                     cont_type="96-pcr",
                     storage="ambient")
```

Autoprotocol Output:

```
{
  "refs": {
    "sample_plate_1": {
      "new": "96-pcr",
      "discard": true
    },
    "sample_plate_2": {
      "id": "ct1cxaе33lkj",
      "store": {
        "where": "ambient"
      }
    },
    "instructions": []
  }
}
```

**Parameters**

- **name** (*str*) – name of the container/ref being created.
- **id** (*str*) – id of the container being created, from your organization’s inventory on <http://secure.transcriptic.com>. Strings representing ids begin with “ct”.
- **cont\_type** (*str or ContainerType*) – container type of the Container object that will be generated.
- **storage** (*Enum({“ambient”, “cold\_20”, “cold\_4”, “warm\_37”}), optional*) – temperature the container being referenced should be stored at after a run is completed. Either a storage condition must be specified or discard must be set to True.
- **discard** (*bool, optional*) – if no storage condition is specified and discard is set to True, the container being referenced will be discarded after a run.
- **cover** (*str, optional*) – name of the cover which will be on the container/ref

### Returns

Container object generated from the id and container type provided.

### Return type *Container*

### Raises

- `RuntimeError` – If a container previously referenced in this protocol (existant in refs section) has the same name as the one specified.
- `RuntimeError` – If no container type is specified.
- `RuntimeError` – If no valid storage or discard condition is specified.

### **sangerseq** (*cont, wells, dataref, type=’standard’, primer=None*)

Send the indicated wells of the container specified for Sanger sequencing. The specified wells should already contain the appropriate mix for sequencing, including primers and DNA according to the instructions provided by the vendor.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-flat",
                     storage="warm_37")

p.sangerseq(sample_plate,
            sample_plate.wells_from(0,5).indices(),
            "seq_data_022415")
```

Autoprotocol Output:

```
"instructions": [
    {
        "dataref": "seq_data_022415",
        "object": "sample_plate",
        "wells": [
            "A1",
            "A2",
            "A3",
            "A4",
            "A5"
        ]
    },
    ...
]
```

(continues on next page)

(continued from previous page)

```

        "op": "sanger_sequence"
    }
]
```

## Parameters

- **cont** ([Container or str](#)) – Container with well(s) that contain material to be sequenced.
- **wells** ([list\(Well\) or WellGroup or Well](#)) – WellGroup of wells to be measured or a list of well references in the form of ["A1", "B1", "C5", ...]
- **dataref** ([str](#)) – Name of sequencing dataset that will be returned.
- **type** ([Enum\({ "standard", "rca" }\)](#)) – Sanger sequencing type
- **primer** ([Container, optional](#)) – Tube containing sufficient primer for all RCA reactions. This field will be ignored if you specify the sequencing type as "standard". Tube containing sufficient primer for all RCA reactions

**Returns** Returns the [autoprotocol.instruction.SangerSeq](#) instruction created from the specified parameters

**Return type** [SangerSeq](#)

## Raises

- [RuntimeError](#) – No primer location specified for rca sequencing type
- [ValueError](#) – Wells belong to more than one container
- [TypeError](#) – Invalid input type for wells

**seal** (*ref, type=None, mode=None, temperature=None, duration=None*)

Seal indicated container using the automated plate sealer.

Example Usage:

```

p = Protocol()
sample_plate = p.ref("sample_plate",
                     None,
                     "96-pcr",
                     storage="warm_37")

p.seal(sample_plate, mode="thermal", temperature="160:celsius")

```

Autoprotocol Output:

```

"instructions": [
    {
        "object": "sample_plate",
        "type": "ultra-clear",
        "mode": "thermal",
        "mode_params": {
            "temperature": "160:celsius"
        }
        "op": "seal"
    }
]

```

## Parameters

- **ref** (`Container`) – Container to be sealed
- **type** (`str, optional`) – Seal type to be used, such as “ultra-clear” or “foil”.
- **mode** (`str, optional`) – Sealing method to be used, such as “thermal” or “adhesive”. Defaults to None, which is interpreted sensibly based on the execution environment.
- **temperature** (`Unit or str, optional`) – Temperature at which to melt the sealing film onto the ref. Only applicable to thermal sealing; not respected if the sealing mode is adhesive. If unspecified, thermal sealing temperature defaults correspond with manufacturer-recommended or internally-optimized values for the target container type. Applies only to thermal sealing.
- **duration** (`Unit or str, optional`) – Duration for which to press the (heated, if thermal) seal down on the ref. Defaults to manufacturer-recommended or internally- optimized seal times for the target container type. Currently applies only to thermal sealing.

**Returns** Returns the `autoprotocol.instruction.Seal` instruction created from the specified parameters

**Return type** `Seal`

## Raises

- `TypeError` – If ref is not of type Container.
- `RuntimeError` – If container type does not have *seal* capability.
- `RuntimeError` – If seal is not a valid seal type.
- `RuntimeError` – If the sealing mode is invalid, or incompatible with the given ref
- `RuntimeError` – If thermal sealing params (temperature and/or duration) are specified alongside an adhesive sealing mode.
- `RuntimeError` – If specified thermal sealing parameters are invalid
- `RuntimeError` – If container is already covered with a lid.

**spectrophotometry** (`dataref, obj, groups, interval=None, num_intervals=None, temperature=None, shake_before=None`)

Generates an instruction with one or more plate reading steps executed on a single plate with the same device. This could be executed once, or at a defined interval, across some total duration.

Example Usage:

```
p = Protocol()
read_plate = p.ref("read plate", cont_type="96-flat", discard=True)

groups = Spectrophotometry.builders.groups(
    [
        Spectrophotometry.builders.group(
            "absorbance",
            Spectrophotometry.builders.absorbance_mode_params(
                wells=read_plate.wells(0, 1),
                wavelength=["100:nanometer", "200:nanometer"],
                num_flashes=15,
                settle_time="1:second"
            )
        ),
        Spectrophotometry.builders.group(
            "reflectance",
            Spectrophotometry.builders.reflectance_mode_params(
                wells=read_plate.wells(2, 3),
                wavelength=[550, 600, 650, 700, 750, 800]
            )
        )
    ]
)
```

(continues on next page)

(continued from previous page)

```

    "fluorescence",
    Spectrophotometry.builders.fluorescence_mode_params(
        wells=read_plate.wells(0, 1),
        excitation=[
            Spectrophotometry.builders.wavelength_selection(
                ideal="650:nanometer"
            )
        ],
        emission=[
            Spectrophotometry.builders.wavelength_selection(
                shortpass="600:nanometer",
                longpass="700:nanometer"
            )
        ],
        num_flashes=15,
        settle_time="1:second",
        lag_time="9:second",
        integration_time="2:second",
        gain=0.3,
        read_position="top"
    )
),
Spectrophotometry.builders.group(
    "luminescence",
    Spectrophotometry.builders.luminescence_mode_params(
        wells=read_plate.wells(0, 1),
        num_flashes=15,
        settle_time="1:second",
        integration_time="2:second",
        gain=0.3
    )
),
Spectrophotometry.builders.group(
    "shake",
    Spectrophotometry.builders.shake_mode_params(
        duration="1:second",
        frequency="9:hertz",
        path="ccw_orbital",
        amplitude="1:mm"
    )
),
]
)

shake_before = Spectrophotometry.builders.shake_before(
    duration="10:minute",
    frequency="5:hertz",
    path="ccw_orbital",
    amplitude="1:mm"
)
)

p.spectrophotometry(
    dataref="test data",
    obj=read_plate,
    groups=groups,
    interval="10:minute",
    num_intervals=2,
)

```

(continues on next page)

(continued from previous page)

```

        temperature="37:celsius",
        shake_before=shake_before
    )

```

Autoprotocol Output:

```
{
  "op": "spectrophotometry",
  "dataref": "test data",
  "object": "read plate",
  "groups": [
    {
      "mode": "absorbance",
      "mode_params": {
        "wells": [
          "read plate/0",
          "read plate/1"
        ],
        "wavelength": [
          "100:nanometer",
          "200:nanometer"
        ],
        "num_flashes": 15,
        "settle_time": "1:second"
      }
    },
    {
      "mode": "fluorescence",
      "mode_params": {
        "wells": [
          "read plate/0",
          "read plate/1"
        ],
        "excitation": [
          {
            "ideal": "650:nanometer"
          }
        ],
        "emission": [
          {
            "shortpass": "600:nanometer",
            "longpass": "700:nanometer"
          }
        ],
        "num_flashes": 15,
        "settle_time": "1:second",
        "lag_time": "9:second",
        "integration_time": "2:second",
        "gain": 0.3,
        "read_position": "top"
      }
    },
    {
      "mode": "luminescence",
      "mode_params": {
        "wells": [
          "read plate/0",

```

(continues on next page)

(continued from previous page)

```

        "read_plate/1"
    ],
    "num_flashes": 15,
    "settle_time": "1:second",
    "integration_time": "2:second",
    "gain": 0.3
}
},
{
    "mode": "shake",
    "mode_params": {
        "duration": "1:second",
        "frequency": "9:hertz",
        "path": "ccw_orbital",
        "amplitude": "1:millimeter"
    }
},
],
"interval": "10:minute",
"num_intervals": 2,
"temperature": "37:celsius",
"shake_before": {
    "duration": "10:minute",
    "frequency": "5:hertz",
    "path": "ccw_orbital",
    "amplitude": "1:millimeter"
}
}
}

```

## Parameters

- **dataref** (*str*) – Name of the resultant dataset to be returned.
- **obj** (*Container or str*) – Container to be read.
- **groups** (*list*) – A list of groups generated by SpectrophotometryBuilders groups builders, any of absorbance\_mode\_params, fluorescence\_mode\_params, luminescence\_mode\_params, or shake\_mode\_params.
- **interval** (*Unit or str, optional*) – The time between each of the read intervals.
- **num\_intervals** (*int, optional*) – The number of times that the groups should be executed.
- **temperature** (*Unit or str, optional*) – The temperature that the entire instruction should be executed at.
- **shake\_before** (*dict, optional*) – A dict of params generated by SpectrophotometryBuilders.shake\_before that dictates how the obj should be incubated with shaking before any of the groups are executed.

**Returns** Returns the `autoprotocol.instruction.Spectrophotometry` instruction created from the specified parameters

**Return type** `Spectrophotometry`

## Raises

- `TypeError` – Invalid num\_intervals specified, must be an int

- `ValueError` – No interval specified but shake groups specified with no duration

**spin**(*ref*, *acceleration*, *duration*, *flow\_direction*=*None*, *spin\_direction*=*None*)

Apply acceleration to a container.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-flat",
                      storage="warm_37")

p.spin(sample_plate, "1000:g", "20:minute", flow_direction="outward")
```

Autoprotocol Output:

```
"instructions": [
    {
        "acceleration": "1000:g",
        "duration": "20:minute",
        "flow_direction": "outward",
        "spin_direction": [
            "cw",
            "ccw"
        ],
        "object": "sample_plate",
        "op": "spin"
    }
]
```

## Parameters

- **ref** (`Container`) – The container to be centrifuged.
- **acceleration** (`str`) – Acceleration to be applied to the plate, in units of *g* or *meter/second*<sup>2</sup>.
- **duration** (`str or Unit`) – Length of time that acceleration should be applied.
- **flow\_direction** (`str`) – Specifies the direction contents will tend toward with respect to the container. Valid directions are “inward” and “outward”, default value is “inward”.
- **spin\_direction** (`list(str)`) – A list of “cw” (clockwise), “cww” (counterclockwise). For each element in the list, the container will be spun in the stated direction for the set “acceleration” and “duration”. Default values are derived from the “flow\_direction” parameter. If “flow\_direction” is “outward”, then “spin\_direction” defaults to [“cw”, “ccw”]. If “flow\_direction” is “inward”, then “spin\_direction” defaults to [“cw”].

**Returns** Returns the `autoprotocol.instruction.Spin` instruction created from the specified parameters

**Return type** `Spin`

## Raises

- `TypeError` – If ref to spin is not of type Container.
- `TypeError` – If spin\_direction or flow\_direction are not properly formatted.

- `ValueError` – If spin\_direction or flow\_direction do not have appropriate values.

**spread**(*source*, *dest*, *volume*=’50:microliter’, *dispense\_speed*=’20:microliter/second’)

Spread the specified volume of the source aliquot across the surface of the agar contained in the object container.

Uses a spiral pattern generated by a set of liquid\_handle instructions.

Example Usage: .. code-block:: python

```
p = Protocol()

agar_plate = p.ref("agar_plate", None, "1-flat", discard=True) bact = p.ref("bacteria", None,
"micro-1.5", discard=True)

p.spread(bact.well(0), agar_plate.well(0), "55:microliter")
```

### Parameters

- **source** (`Well`) – Source of material to spread on agar
- **dest** (`Well`) – Reference to destination location (plate containing agar)
- **volume** (*str or Unit, optional*) – Volume of source material to spread on agar
- **dispense\_speed** (*str or Unit, optional*) – Speed at which to dispense source aliquot across agar surface

**Returns** Returns a `autoprotocol.instruction.LiquidHandle` instruction created from the specified parameters

**Return type** `LiquidHandle`

### Raises

- `TypeError` – If specified source is not of type Well
- `TypeError` – If specified destination is not of type Well

**store**(*container*, *condition*)

Manually adjust the storage destiny for a container used within this protocol.

### Parameters

- **container** (`Container`) – Container used within this protocol
- **condition** (*str*) – New storage destiny for the specified Container

### Raises

- `TypeError` – If container argument is not a Container object
- `RuntimeError` – If the container passed is not already present in self.refs

**thermocycle**(*ref*, *groups*, *volume*=’10:microliter’, *dataref*=None, *dyes*=None, *melting\_start*=None, *melting\_end*=None, *melting\_increment*=None, *melting\_rate*=None, *lid\_temperature*=None)

Append a Thermocycle instruction to the list of instructions, with groups is a list(dict) in the form of:

```
"groups": [ {
    "cycles": integer,
    "steps": [
        {
            "duration": duration,
```

(continues on next page)

(continued from previous page)

```
        "temperature": temperature,
        "read": boolean // optional (default false)
    },
    {
        "duration": duration,
        "gradient": {
            "top": temperature,
            "bottom": temperature
        },
        "read": boolean // optional (default false)
    }
],
}];
```

Thermocycle can also be used for either conventional or row-wise gradient PCR as well as qPCR. Refer to the examples below for details.

## Example Usage:

**To thermocycle a container according to the protocol:**

- **1 cycle:**
    - 95 degrees for 5 minutes
  - **30 cycles:**
    - 95 degrees for 30 seconds
    - 56 degrees for 20 seconds
    - 72 degrees for 30 seconds
  - **1 cycle:**
    - 72 degrees for 10 minutes
  - **1 cycle:**
    - 4 degrees for 30 seconds
  - all cycles: Lid temperature at 97 degrees

```
from instruction import Thermocycle

p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-pcr",
                      storage="warm_37")

# a plate must be sealed before it can be thermocycled
p.seal(sample_plate)

p.thermocycle(
    sample_plate,
    [
        Thermocycle.builders.group(
            steps=[
                Thermocycle.builders.step("95:celsius", "5:minute")
            ]
        )
    ]
)
```

---

(continues on next page)

(continued from previous page)

```

),
Thermocycle.builders.group(
    steps=[
        Thermocycle.builders.step("95:celsius", "30:s"),
        Thermocycle.builders.step("56:celsius", "20:s"),
        Thermocycle.builders.step("72:celsius", "20:s"),
    ],
    cycles=30
),
Thermocycle.builders.group(
    steps=[
        Thermocycle.builders.step("72:celsius", "10:minute")
    ]
),
Thermocycle.builders.group(
    steps=[
        Thermocycle.builders.step("4:celsius", "30:s")
    ]
)
],
lid_temperature="97:celsius"
)

```

## Autoprotocol Output:

```

"instructions": [
{
    "object": "sample_plate",
    "op": "seal"
},
{
    "volume": "10:microliter",
    "dataref": null,
    "object": "sample_plate",
    "groups": [
        {
            "cycles": 1,
            "steps": [
                {
                    "duration": "5:minute",
                    "temperature": "95:celsius"
                }
            ]
        },
        {
            "cycles": 30,
            "steps": [
                {
                    "duration": "30:second",
                    "temperature": "95:celsius"
                },
                {
                    "duration": "20:second",
                    "temperature": "56:celsius"
                },
                {
                    "duration": "20:second",
                    "temperature": "72:celsius"
                }
            ]
        }
    ]
}
]

```

(continues on next page)

(continued from previous page)

```

        "temperature": "72:celsius"
    }
]
},
{
    "cycles": 1,
    "steps": [
        {
            "duration": "10:minute",
            "temperature": "72:celsius"
        }
    ]
},
{
    "cycles": 1,
    "steps": [
        {
            "duration": "30:second",
            "temperature": "4:celsius"
        }
    ]
}
],
"op": "thermocycle"
}
]
```

To gradient thermocycle a container according to the protocol:

- **1 cycle:**

- 95 degrees for 5 minutes

- **30 cycles:**

- 95 degrees for 30 seconds

Top Row: \* 65 degrees for 20 seconds Bottom Row: \* 55 degrees for 20 seconds

- 72 degrees for 30 seconds

- **1 cycle:**

- 72 degrees for 10 minutes

```

p = Protocol()
sample_plate = p.ref("sample_plate",
    None,
    "96-pcr",
    storage="warm_37")

# a plate must be sealed before it can be thermocycled
p.seal(sample_plate)

p.thermocycle(
    sample_plate,
    [
        Thermocycle.builders.group(
            steps=[
```

(continues on next page)

(continued from previous page)

```
        Thermocycle.builders.step("95:celsius", "5:minute")
    ],
),
Thermocycle.builders.group(
    steps=[
        Thermocycle.builders.step("95:celsius", "30:s"),
        Thermocycle.builders.step(
            {"top": "65:celsius", "bottom": "55:celsius"},,
            "20:s"
        ),
        Thermocycle.builders.step("72:celsius", "20:s"),
    ],
    cycles=30
),
Thermocycle.builders.group(
    steps=[
        Thermocycle.builders.step("72:celsius", "10:minute")
    ]
)
)
```

To conduct a qPCR, at least one dye type and the dataref field has to be specified. The example below uses SYBR dye and the following temperature profile:

- **1 cycle:**
    - 95 degrees for 3 minutes
  - **40 cycles:**
    - 95 degrees for 10 seconds
    - 60 degrees for 30 seconds (Read during extension)

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-pcr",
                      storage="warm_37")

# a plate must be sealed before it can be thermocycled
p.seal(sample_plate)

p.thermecycle(
    sample_plate,
    [
        Thermocycle.builders.group(
            steps=[
                Thermocycle.builders.step("95:celsius", "3:minute")
            ]
        ),
        Thermocycle.builders.group(
            steps=[
                Thermocycle.builders.step(
                    "95:celsius",
                    "10:second",
                    read=False
                )
            ]
        )
    ]
)
```

---

(continues on next page)

(continued from previous page)

```
)  
    Thermocycle.builders.step(  
        "95:celsius",  
        "10:second",  
        read=True  
    )  
],  
cycles=40  
)  
]  
dataref = "my_qpcr_data",  
dyes = {"SYBR": sample_plate.all_wells().indices() }  
)
```

## Parameters

- **ref** (`Container`) – Container to be thermocycled.
- **groups** (`list (dict)`) – List of thermocycling instructions formatted as above
- **volume** (`str or Unit, optional`) – Volume contained in wells being thermocycled
- **dataref** (`str, optional`) – Name of dataref representing read data if performing qPCR
- **dyes** (`dict, optional`) – Dictionary mapping dye types to the wells they're used in
- **melting\_start** (`str or Unit, optional`) – Temperature at which to start the melting curve.
- **melting\_end** (`str or Unit, optional`) – Temperature at which to end the melting curve.
- **melting\_increment** (`str or Unit, optional`) – Temperature by which to increment the melting curve. Accepted increment values are between 0.1 and 9.9 degrees celsius.
- **melting\_rate** (`str or Unit, optional`) – Specifies the duration of each temperature step in the melting curve.
- **lid\_temperature** (`str or Unit, optional`) – Specifies the lid temperature throughout the duration of the thermocycling instruction

**Returns** Returns the `autoprotocol.instruction.Thermocycle` instruction created from the specified parameters

**Return type** `Thermocycle`

## Raises

- `AttributeError` – If groups are not properly formatted
- `TypeError` – If ref to thermocycle is not of type Container.
- `ValueError` – Container specified cannot be thermocycled
- `ValueError` – Lid temperature is not within bounds

```
transfer(source, destination, volume, rows=1, columns=1, source_liquid=<class 'autoprotocol.liquid_handle.liquid_class.LiquidClass'>, destination_liquid=<class 'autoprotocol.liquid_handle.liquid_class.LiquidClass'>, method=<class 'autoprotocol.liquid_handle.transfer.Transfer'>, one_tip=False)
```

Generates LiquidHandle instructions between wells

Transfer liquid between specified pairs of source & destination wells.

#### Parameters

- **source** (`Well` or `WellGroup` or `list(Well)`) – Well(s) to transfer liquid from.
- **destination** (`Well` or `WellGroup` or `list(Well)`) – Well(s) to transfer liquid to.
- **volume** (`str` or `Unit` or `list(str)` or `list(Unit)`) – Volume(s) of liquid to be transferred from source wells to destination wells. The number of volumes specified must correspond to the number of destination wells.
- **rows** (`int`, *optional*) – Number of rows to be concurrently transferred
- **columns** (`int`, *optional*) – Number of columns to be concurrently transferred
- **source\_liquid** (`LiquidClass` or `list(LiquidClass)`, *optional*) – Type(s) of liquid contained in the source Well. This affects the aspirate and dispense behavior including the flowrates, liquid level detection thresholds, and physical movements.
- **destination\_liquid** (`LiquidClass` or `list(LiquidClass)`, *optional*) – Type(s) of liquid contained in the destination Well. This affects liquid level detection thresholds.
- **method** (`Transfer` or `list(Transfer)`, *optional*) – Integrates with the specified source\_liquid and destination\_liquid to define a set of physical movements.
- **one\_tip** (`bool`, *optional*) – If True then a single tip will be used for all operations

**Returns** Returns a list of `autoprotocol.instruction.LiquidHandle` instructions created from the specified parameters

**Return type** `list(LiquidHandle)`

#### Raises

- `ValueError` – if the specified parameters can't be interpreted as lists of equal length
- `ValueError` – if one\_tip is true, but not all transfer methods have a tip\_type

## Examples

Transfer between two single wells

```
from autoprotocol import Protocol, Unit

p = Protocol()
source = p.ref("source", cont_type="384-flat", discard=True)
destination = p.ref(
    "destination", cont_type="394-pcr", discard=True
)
p.transfer(source.well(0), destination.well(1), "5:ul")
```

Sequential transfers between two groups of wells

```
sources = source.wells_from(0, 8, columnwise=True)
dests = destination.wells_from(1, 8, columnwise=True)
volumes = [Unit(x, "ul") for x in range(1, 9)]
p.transfer(sources, dests, volumes)
```

Concurrent transfers between two groups of wells

```
# single-column concurrent transfer
p.transfer(
    source.well(0), destination.well(1), "5:ul", rows=8
)

# 96-well concurrent transfer from the A1 to B2 quadrants
p.transfer(
    source.well(0), destination.well(13), "5:ul", rows=8, columns=12
)

# 384-well concurrent transfer
p.transfer(
    source.well(0), destination.well(0), "5:ul", rows=16, columns=24
)
```

Transfer with extra parameters

```
from autoprotocol.liquid_handle import Transfer
from autoprotocol.instruction import LiquidHandle

p.transfer(
    source.well(0), destination.well(0), "5:ul",
    method=Transfer(
        mix_before=True,
        dispense_z=LiquidHandle.builders.position_z(
            reference="well_top"
        )
    )
)
```

Transfer using other built in Transfer methods

```
from autoprotocol.liquid_handle import DryWellTransfer

p.transfer(
    source.well(0), destination.well(1), "5:ul",
    method=DryWellTransfer
)
```

For examples of other more complicated behavior, see the documentation for LiquidHandleMethod.

See also:

**Transfer()** base LiquidHandleMethod for transfer operations

**uncover**(*ref*, *store\_lid=None*)

Remove lid from specified container

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-flat",
                      storage="warm_37")
# a plate must have a cover to be uncovered
p.cover(sample_plate, lid="universal")

p.uncover(sample_plate)
```

Autoprotocol Output:

```
"instructions": [
    {
        "lid": "universal",
        "object": "sample_plate",
        "op": "cover"
    },
    {
        "object": "sample_plate",
        "op": "uncover"
    }
]
```

## Parameters

- **ref** ([Container](#)) – Container to remove lid.
- **store\_lid** (*bool, optional*) – Flag to store the uncovered lid.

**Returns** Returns the [\*autoprotocol.instruction.Uncover\*](#) instruction created from the specified parameters

**Return type** [\*Uncover\*](#)

## Raises

- **TypeError** – If ref is not of type Container.
- **RuntimeError** – If container is sealed with a seal not covered with a lid.
- **TypeError** – If store\_lid is not a boolean.

## **unseal** (*ref*)

Remove seal from indicated container using the automated plate unsealer.

Example Usage:

```
p = Protocol()
sample_plate = p.ref("sample_plate",
                      None,
                      "96-pcr",
                      storage="warm_37")
# a plate must be sealed to be unsealed
p.seal(sample_plate)

p.unseal(sample_plate)
```

Autoprotocol Output:

```
"instructions": [
    {
        "object": "sample_plate",
        "op": "seal",
        "type": "ultra-clear"
    },
    {
        "object": "sample_plate",
        "op": "unseal"
    }
]
```

**Parameters** `ref` (`Container`) – Container to be unsealed.

**Returns** Returns the `autoprotocol.instruction.Unseal` instruction created from the specified parameters

**Return type** `Unseal`

**Raises**

- `TypeError` – If ref is not of type Container.
- `RuntimeError` – If container is covered with a lid not a seal.

```
class autoprotocol.protocol.Ref(name, opts, container)
Link a ref name (string) to a Container instance.
```

# CHAPTER 2

---

## autoprotocol.instruction module

---

Contains all the Autoprotocol Instruction objects

**copyright** 2018 by The Autoprotocol Development Team, see AUTHORS for more details.

**license** BSD, see LICENSE for more details

```
class autoprotocol.instruction.Absorbance(object, wells, wavelength, dataref, flashes=25,
                                           incubate_before=None, temperature=None, settle_time=None)
```

Read the absorbance for the indicated wavelength for the indicated wells. Append an Absorbance instruction to the list of instructions for this Protocol object.

### Parameters

- **object** (*str or Ref*) – Object to execute the absorbance read on
- **wells** (*list (Well) or WellGroup*) – WellGroup of wells to be measured or a list of well references in the form of [“A1”, “B1”, “C5”, …]
- **wavelength** (*str or Unit*) – wavelength of light absorbance to be read for the indicated wells
- **dataref** (*str*) – name of this specific dataset of measured absorbances
- **flashes** (*int, optional*) – number of flashes for the read
- **incubate\_before** (*dict, optional*) – incubation prior to reading if desired

#### shaking: dict, optional

##### shake parameters if desired

**amplitude:** **str or Unit** amplitude of shaking between 1 and 6:millimeter

**orbital:** **bool** True for orbital and False for linear shaking

**duration:** **str, Unit, optional** time prior to plate reading

- **temperature** (*str or Unit, optional*) – set temperature to heat plate reading chamber

- **settle\_time** (*str or Unit, optional*) – time to pause before each well read

```
class autoprotocol.instruction.AcousticTransfer (groups, droplet_size)
```

Specify source and destination wells for transferring liquid via an acoustic liquid handler. Droplet size is usually device-specific.

#### Parameters

- **groups** (*list (dict)*) – List of *transfer* groups in the form of:

```
{  
    "transfer": [  
        {  
            "to": "foo/A1",  
            "from": "bar/A1",  
            "volume": "1:n1"  
        }  
    ]  
}
```

- **droplet\_size** (*str or Unit*) – Volume representing a droplet\_size. The volume of each transfer should be a multiple of this volume.

```
class autoprotocol.instruction.Autopick (groups, criteria, dataref)
```

Pick colonies from the agar-containing location(s) specified in *sources* to the location(s) specified in *dests* in highest to lowest rank order until there are no more colonies available. If fewer than min\_abort pickable colonies have been identified from the location(s) specified in *sources*, the run will stop and no further instructions will be executed.

#### Parameters

- **groups** (*list (dict)*) – Groups of colonies to pick and where to transport them to
- **criteria** (*dict*) – Dictionary of autopicking criteria.
- **dataref** (*str*) – Name of dataset to save the picked colonies to

```
class autoprotocol.instruction.CountCells (wells, volume, dataref, labels=None)
```

Count the number of cells in a sample that are positive/negative for a given set of labels.

#### Parameters

- **wells** (*WellGroup*) – List of wells that will be used for cell counting.
- **volume** (*Unit*) – Volume that should be consumed from each well for the purpose of cell counting.
- **dataref** (*str*) – Name of dataset that will be returned.
- **labels** (*list (string), optional*) – Cells will be scored for presence or absence of each label in this list. If staining is required to visualize these labels, they must be added before execution of this instruction.

```
class autoprotocol.instruction.Cover (object, lid='standard', retrieve_lid=None)
```

Place specified lid type on specified container

#### Parameters

- **object** (*str*) – Container to be covered
- **lid** (*Enum({'standard', 'universal', 'low\_evaporation'})*, *optional*) – Type of lid to cover container with
- **retrieve\_lid** (*bool*) – Flag to retrieve lid from stored location

---

```
class autoprotocol.instruction.Dispense(object, columns, reagent=None, re-  

source_id=None, reagent_source=None,  

step_size=None, flowrate=None, noz-  

zle_position=None, pre_dispense=None,  

shape=None, shake_after=None)
```

Dispense specified reagent to specified columns. Only one of reagent, resource\_id, and reagent\_source can be specified for a given instruction.

#### Parameters

- **object** (`Container or str`) – Container for reagent to be dispensed to.
- **columns** (`list`) – Columns to be dispensed to, in the form of a list of dicts specifying the column number and the volume to be dispensed to that column. Columns are indexed from 0. [{“column”: <column num>, “volume”: <volume>}, …]
- **reagent** (`str, optional`) – Reagent to be dispensed.
- **resource\_id** (`str, optional`) – Resource to be dispensed.
- **reagent\_source** (`Well, optional`) – Aliquot to be dispensed from.
- **step\_size** (`str or Unit, optional`) – Specifies that the dispense operation must be executed using a pump that has a dispensing resolution of step\_size.
- **flowrate** (`str or Unit, optional`) – The rate at which the peristaltic pump should dispense in Units of flow rate, e.g. microliter/second.
- **nozzle\_position** (`dict, optional`) – A dict represent nozzle offsets from the center of the bottom of the plate’s well. see Dispense.builders.nozzle\_position; specified as {“position\_x”: Unit, “position\_y”: Unit, “position\_z”: Unit}.
- **pre\_dispense** (`str or Unit, optional`) – The volume of reagent to be dispensed per-nozzle into waste immediately prior to dispensing into the ref.
- **shape** (`dict, optional`) – The shape of the dispensing head to be used for the dispense. See liquid\_handle\_builders.shape\_builder; specified as {“rows”: int, “columns”: int, “format”: str} with format being a valid SBS format.
- **shake\_after** (`dict, optional`) – Parameters that specify how a plate should be shaken at the very end of the instruction execution. {“duration”: Unit, “frequency”: Unit, “path”: str, “amplitude”: Unit}

```
class autoprotocol.instruction.FlashFreeze(object, duration)
```

Flash freeze the contents of the specified container by submerging it in liquid nitrogen for the specified amount of time.

#### Parameters

- **object** (`Container or str`) – Container to be flash frozen.
- **duration** (`str or Unit`) – Duration to submerge specified container in liquid nitrogen.

```
class autoprotocol.instruction.FlowAnalyze(dataref, FSC, SSC, negative_controls, samples,  

colors=None, positive_controls=None)
```

Perform flow cytometry. The instruction will be executed within the voltage range specified for each channel, optimized for the best sample separation/distribution that can be achieved within these limits. The vendor will specify the device that this instruction is executed on and which excitation and emission spectra are available. At least one negative control is required, which will be used to define data acquisition parameters as well as to determine any autofluorescent properties for the sample set. Additional negative positive control samples are optional. Positive control samples will be used to optimize single color signals and, if desired, to minimize bleed into other channels.

For each sample this instruction asks you to specify the *volume* and/or *captured\_events*. Vendors might also require *captured\_events* in case their device does not support volumetric sample intake. If both conditions are supported, the vendor will specify if data will be collected only until the first one is met or until both conditions are fulfilled.

Example Usage:

Autoprotocol Output:

### Parameters

- **dataref** (*str*) – Name of flow analysis dataset generated.
- **FSC** (*dict*) – Dictionary containing FSC channel parameters in the form of:

```
{  
    "voltage_range": {  
        "low": "230:volt",  
        "high": "280:volt"  
    },  
    "area": true,           //default: true  
    "height": true,         //default: true  
    "weight": false         //default: false  
}
```

- **SSC** (*dict*) – Dictionary of SSC channel parameters in the form of:

```
{  
    "voltage_range": {  
        "low": <voltage>,  
        "high": <voltage>"  
    },  
    "area": true,           //default: true  
    "height": true,          //default: false  
    "weight": false          //default: false  
}
```

- **negative\_controls** (*list (dict)*) – List of negative control wells in the form of:

```
{  
    "well": well,  
    "volume": volume,  
    "captured_events": integer,      // optional, default infinity  
    "channel": [channel_name]  
}
```

at least one negative control is required.

- **samples** (*list (dict)*) – List of samples in the form of:

```
{  
    "well": well,  
    "volume": volume,  
    "captured_events": integer,      // optional, default infinity  
}
```

at least one sample is required

- **colors** (*list (dict)*, *optional*) – Optional list of colors in the form of:

```
[{
    "name": "FitC",
    "emission_wavelength": "495:nanometer",
    "excitation_wavelength": "519:nanometer",
    "voltage_range": {
        "low": <voltage>,
        "high": <voltage>
    },
    "area": true, //default: true
    "height": false, //default: false
    "weight": false //default: false
}]
```

**positive\_controls** [list(dict), optional] Optional list of positive control wells in the form of:

```
[{
    "well": well,
    "volume": volume,
    "captured_events": integer, // optional, default infinity
    "channel": [channel_name],
    "minimize_bleed": [{ // optional
        "from": color,
        "to": [color]
    }]
}]
```

```
class autoprotocol.instruction.Fluorescence(object, wells, excitation, emission, dataref,  

                                              flashes=25, incubate_before=None,  

                                              temperature=None, gain=None, detection_mode=None, position_z=None,  

                                              settle_time=None, lag_time=None, integration_time=None)
```

Read the fluorescence for the indicated wavelength for the indicated wells. Append a Fluorescence instruction to the list of instructions for this Protocol object.

#### Parameters

- **object** (*str or Container*) – object to execute the fluorescence read on
- **wells** (*list(Well) or WellGroup*) – WellGroup of wells to be measured or a list of well references in the form of [“A1”, “B1”, “C5”, ...]
- **excitation** (*str or Unit*) – wavelength of light used to excite the wells indicated
- **emission** (*str or Unit*) – wavelength of light to be measured for the indicated wells
- **dataref** (*str*) – name of this specific dataset of measured absorbances
- **flashes** (*int, optional*) – number of flashes for this read
- **incubate\_before** (*dict, optional*) – incubation prior to reading if desired

#### shaking: dict, optional

##### shake parameters if desired

**amplitude: str or Unit** amplitude of shaking between 1 and 6:millimeter

**orbital: bool** True for orbital and False for linear shaking

**duration: str, Unit, optional** time prior to plate reading

- **temperature**(*str or Unit, optional*) – set temperature to heat plate reading chamber
- **gain**(*float, optional*) – float between 0 and 1, multiplier of maximum signal amplification
- **detection\_mode**(*str, optional*) – set the detection mode of the optics, [“top”, “bottom”], defaults to vendor specified defaults.
- **position\_z**(*dict, optional*) – distance from the optics to the surface of the plate transport, only valid for “top” detection\_mode and vendor capabilities. Specified as either a set distance - “manual”, OR calculated from a WellGroup - “calculated\_from\_wells”. Only one position\_z determination may be specified

```
position_z = {
    "manual": Unit
    - OR -
    "calculated_from_wells": []
}
```

- **manual**(*str, Unit, optional*) – parameter available within “position\_z” to set the distance from the optics to the plate transport.
- **calculated\_from\_wells**(*list, WellGroup, Well, optional*) – parameter available within “position\_z” to set the distance from the optics to the plate transport. If specified, the average optimal (maximal signal) distance will be chosen from the list of wells and applied to all measurements.
- **settle\_time**(*Unit, optional*) – the time before the start of the measurement, defaults to vendor specifications
- **lag\_time**(*Unit, optional*) – time between flashes and the start of the signal integration, defaults to vendor specifications
- **integration\_time**(*Unit, optional*) – duration of the signal recording, per Well, defaults to vendor specifications

**class** autoprotocol.instruction.**GelPurify**(*objects, volume, matrix, ladder, dataref, extract*)  
Separate nucleic acids on an agarose gel and purify.

#### Parameters

- **objects**(*list or WellGroup*) – WellGroup of wells to be purified
- **volume**(*str or Unit*) – Volume of sample required for analysis
- **dataref**(*str*) – Name of this specific dataset of measurements
- **matrix**(*str*) – Agarose concentration and number of wells on gel used for separation
- **ladder**(*str*) – Size range of ladder to be used to compare band size to
- **dataref** – Name of dataset containing fragment sizes returned
- **extract**(*list(dict)*) –

```
"extract": [
    "elution_volume": volume,
    "elution_buffer": string, "water" | "TE",
    "lane": int,
    "band_size_range": {
```

(continues on next page)

(continued from previous page)

```

        "min_bp": int,
        "max_bp": int,
    },
    "destination": well
},
{...}]

```

**class** autoprotocol.instruction.**GelSeparate**(*objects*, *volume*, *matrix*, *ladder*, *duration*, *dataref*)

Separate nucleic acids on an agarose gel.

#### Parameters

- **objects** (*list or WellGroup or Well*) – List of wells or WellGroup containing wells to be separated on gel.
- **volume** (*str or Unit*) – Volume of liquid to be transferred from each well specified to a lane of the gel.
- **matrix** (*str*) – Matrix (gel) in which to gel separate samples
- **ladder** (*str*) – Ladder by which to measure separated fragment size
- **duration** (*str or Unit*) – Length of time to run current through gel.
- **dataref** (*str*) – Name of this set of gel separation results.

**class** autoprotocol.instruction.**IlluminaSeq**(*flowcell*, *lanes*, *sequencer*, *mode*, *index*, *library\_size*, *dataref*, *cycles*)

Load aliquots into specified lanes for Illumina sequencing. The specified aliquots should already contain the appropriate mix for sequencing and require a library concentration reported in ng/uL.

#### Parameters

- **flowcell** (*str*) – Flowcell designation: “SR” or “PE”
- **lanes** (*list (dict)*) –
 

```

"lanes": [
    {
        "object": aliquot, Well,
        "library_concentration": decimal, // ng/uL
    },
    {...}]
      
```
- **sequencer** (*str*) – Sequencer designation: “miseq”, “hiseq” or “nextseq”
- **mode** (*str*) – Mode designation: “rapid”, “mid” or “high”
- **index** (*str*) – Index designation: “single”, “dual” or “none”
- **library\_size** (*integer*) – Library size expressed as an integer of basepairs
- **dataref** (*str*) – Name of sequencing dataset that will be returned.
- **cycles** (*Enum({ "read\_1", "read\_2", "index\_1", "index\_2"})*) – Parameter specific to Illuminaseq read-length or number of sequenced bases. Refer to the ASC for more details

**class** autoprotocol.instruction.**ImagePlate**(*object*, *mode*, *dataref*)

Capture an image of the specified container.

#### Parameters

- **object** (*str*) – Container to take image of

- **mode** (*str*) – Imaging mode (currently supported: “top”)
- **dataref** (*str*) – Name of data reference of resulting image

```
class autoprotocol.instruction.Incubate(object, where, duration, shaking=False,
                                         co2=0, target_temperature=None, shaking_params=None)
```

Store a sample in a specific environment for a given duration. Once the duration has elapsed, the sample will be returned to the ambient environment until it is next used in an instruction.

#### Parameters

- **object** (*Ref or str*) – The container to be incubated
- **where** (*Enum({“ambient”, “warm\_37”, “cold\_4”, “cold\_20”, “cold\_80”})*) – Temperature at which to incubate specified container
- **duration** (*Unit or str*) – Length of time to incubate container
- **shaking** (*bool, optional*) – Specify whether or not to shake container if available at the specified temperature
- **target\_temperature** (*Unit or str, optional*) – Specify a target temperature for a device (eg. an incubating block) to reach during the specified duration.
- **shaking\_params** (*dict, optional*) – Specify “path” and “frequency” of shaking parameters to be used with compatible devices (eg. thermoshakes)
- **co2** (*int, optional*) – Carbon dioxide percentage

```
class autoprotocol.instruction.Instruction(op, data)
```

Base class for an instruction that is to later be encoded as JSON.

```
json()
```

Return instruction object properly encoded as JSON for Autoprotocol.

**Returns** Instruction object encoded as json string

**Return type** str

```
class autoprotocol.instruction.LiquidHandle(locations, shape=None, mode=None,
                                              mode_params=None)
```

Manipulates liquids within locations

A liquid handle instruction is constructed as a list of locations, where each location consists of the well location and the tip transports carried out within the well.

Each liquid handle instruction corresponds to a single tip or set of tips.

#### Parameters

- **locations** (*list (dict)*) – See Also LiquidHandle.builders.location
- **shape** (*dict, optional*) – See Also LiquidHandle.builders.shape
- **mode** (*str, optional*) – the liquid handling mode
- **mode\_params** (*dict, optional*) – See Also LiquidHandle.builders.instruction\_mode\_params

```
class autoprotocol.instruction.Luminescence(object, wells, dataref, incubate_before=None,
                                              temperature=None, settle_time=None, integration_time=None)
```

Read luminescence of indicated wells

#### Parameters

- **object** (*str or Container*) – object to execute the luminescence read on
- **wells** (*list or WellGroup*) – WellGroup or list of wells to be measured
- **dataref** (*str*) – name which dataset will be saved under
- **incubate\_before** (*dict, optional*) – incubation prior to reading if desired  
shaking: *dict, optional*

#### shake parameters if desired

**amplitude:** *str or Unit* amplitude of shaking between 1 and 6:millimeter

**orbital:** *bool* True for orbital and False for linear shaking

**duration:** *str, Unit, optional* time prior to plate reading

- **temperature** (*str or Unit, optional*) – set temperature to heat plate reading chamber
- **settle\_time** (*str or Unit, optional*) – time to pause before each well read
- **integration\_time** (*Unit, optional*) – duration of the signal recording, per Well, defaults to vendor specifications

**class** autoprotocol.instruction.**MagneticTransfer** (*groups, head\_type*)

A magnetic\_transfer instruction is constructed as a list of lists of groups, executed in order, where each group is a collect, release, dry, incubate, or mix sub-operation. These sub-operations control the behavior of tips which can be magnetized, and a heating platform. Groups in the same list of groups use the same tips.

#### Parameters

- **groups** (*list (dict)*) – dict in the groups should belong to one of the following categories:
  - collect:** Collects beads from the specified “object” by raising and lowering magnetized tips repeatedly with an optional pause at well bottom.
  - release:** Release beads from unmagnetized tips by oscillating the tips vertically into and out of the “object”.
  - dry:** Dry beads on magnetized tips above and outside the “object”.
  - incubate:** Incubate the “object”.
  - mix:** Oscillate the tips into and out of the “object”
- **head\_type** (*str*) – Head-type used for this instruction

**class** autoprotocol.instruction.**MeasureConcentration** (*object, volume, dataref, measurement*)

Measure the concentration of DNA, ssDNA, RNA or Protein in the specified volume of the source aliquots.

#### Parameters

- **object** (*list or WellGroup*) – WellGroup of wells to be measured
- **volume** (*str or Unit*) – Volume of sample required for analysis
- **dataref** (*str*) – Name of this specific dataset of measurements
- **measurement** (*str*) – Class of material to be measured. One of [“DNA”, “ssDNA”, “RNA”, “protein”].

**class** autoprotocol.instruction.**MeasureMass** (*object, dataref*)

Measure the mass of containers

## Parameters

- **object** (`Container`) – Container ref
- **dataref** (`str`) – Name of the data for the measurement

**class** `autoprotocol.instruction.MeasureVolume` (*object*, *dataref*)

Measure the mass of containers

## Parameters

- **object** (`list(Container)`) – list of containers
- **dataref** (`str`) – Name of the data for the measurement

**class** `autoprotocol.instruction.Oligosynthesize` (*oligos*)

**Parameters** **oligos** (`list of dicts`) – List of oligonucleotides to synthesize. Each dictionary should contain the oligo's sequence, destination, scale and purification

```
[  
  {  
    "destination": "my_plate/A1",  
    "sequence": "GATCRYMKSWHBVDN",  
    // - standard IUPAC base codes  
    // - IDT also allows rX (RNA), mX (2' O-methyl RNA), and  
    //   X*/rX*/mX* (phosphorothioated)  
    // - they also allow inline annotations for modifications,  
    //   eg "GCGACTC/3Phos/" for a 3' phosphorylation  
    //   eg "aggg/iAzideN/cgcgc" for an internal modification  
    "scale": "25nm" | "100nm" | "250nm" | "1um",  
    "purification": "standard" | "page" | "hplc",  
    // default: standard  
  },  
  ...  
]
```

**class** `autoprotocol.instruction.Provision` (*resource\_id*, *dests*)

Provision a commercial resource from a catalog into the specified destination well(s). A new tip is used for each destination well specified to avoid contamination.

## Parameters

- **resource\_id** (`str`) – Resource ID from catalog.
- **dests** (`list(dict)`) – Destination(s) for specified resource, together with volume information

## Raises

- `TypeError` – If `resource_id` is not a string.
- `RuntimeError` – If length of the list of volumes specified does not match the number of destination wells specified.
- `TypeError` – If volume is not specified as a string or Unit (or a list of either)

**class** `autoprotocol.instruction.SangerSeq` (*object*, *wells*, *dataref*, *type*, *primer=None*)

Send the indicated wells of the container specified for Sanger sequencing. The specified wells should already contain the appropriate mix for sequencing, including primers and DNA according to the instructions provided by the vendor.

## Parameters

- **object** (`Container or str`) – Container with well(s) that contain material to be sequenced.
- **wells** (`list (str)`) – Well indices of the container that contain appropriate materials to be sent for sequencing.
- **dataref** (`str`) – Name of sequencing dataset that will be returned.
- **type** (`Enum ({ "standard", "rca" })`) – Sanger sequencing type
- **primer** (`Container, optional`) – Tube containing sufficient primer for all RCA reactions. This field will be ignored if you specify the sequencing type as “standard”. Tube containing sufficient primer for all RCA reactions

```
class autoprotocol.instruction.Seal (object, type='ultra-clear', mode=None,  
                                mode_params=None)
```

Seal indicated container using the automated plate sealer.

#### Parameters

- **object** (`Ref or str`) – Container to be sealed
- **type** (`str, optional`) – Seal type to be used (optional)
- **mode** (`str, optional`) – Method used to seal plate (optional). “thermal” or “adhesive”
- **mode\_params** (`dict, optional`) – Thermal sealing parameters  
**temperature** [str, optional] Temperature to seal plate at  
**duration** [str, optional] Duration for which to apply heated sealing plate onto ref

```
class autoprotocol.instruction.Spectrophotometry (dataref, object, groups, interval=None,  
                                         num_intervals=None,  
                                         temperature=None,  
                                         shake_before=None)
```

Execute a Spectrophotometry plate read on the obj.

#### Parameters

- **dataref** (`str`) – Name of the resultant dataset to be returned.
- **object** (`Container or str`) – Container to be read.
- **groups** (`list`) – A list of groups generated by SpectrophotometryBuilders groups builders, any of absorbance\_mode\_params, fluorescence\_mode\_params, luminescence\_mode\_params, or shake\_mode\_params.
- **interval** (`Unit or str, optional`) – The time between each of the read intervals.
- **num\_intervals** (`int, optional`) – The number of times that the groups should be executed.
- **temperature** (`Unit or str, optional`) – The temperature that the entire instruction should be executed at.
- **shake\_before** (`dict, optional`) – A dict of params generated by SpectrophotometryBuilders.shake\_before that dictates how the obj should be incubated with shaking before any of the groups are executed.

```
class autoprotocol.instruction.Spin (object, acceleration, duration, flow_direction=None,  
                                spin_direction=None)
```

Apply the specified amount of acceleration to a plate using a centrifuge.

## Parameters

- **object** (`Ref` or `str`) – Container to be centrifuged.
- **acceleration** (`str`) – Amount of acceleration to be applied to the container, expressed in units of “g” or “meter/second<sup>2</sup>”
- **duration** (`str` or `Unit`) – Amount of time to apply acceleration.
- **flow\_direction** (`str`) – Specifies the direction contents will tend toward with respect to the container. Valid directions are “inward” and “outward”, default value is “inward”.
- **spin\_direction** (`list(str)`) – A list of “cw” (clockwise), “cww” (counterclockwise). For each element in the list, the container will be spun in the stated direction for the set “acceleration” and “duration”. Default values are derived from the “flow\_direction”. If “flow\_direction” is “outward”, then “spin\_direction” defaults to [“cw”, “ccw”]. If “flow\_direction” is “inward”, then “spin\_direction” defaults to [“cw”].

```
class autoprotocol.instruction.Thermocycle(object, groups, volume='25:microliter',
                                             dataref=None, dyes=None, melting_start=None, melting_end=None, melting_increment=None, melting_rate=None,
                                             lid_temperature=None)
```

Append a Thermocycle instruction to the list of instructions, with groups being a list of dicts in the form of:

```
"groups": [
    {
        "cycles": integer,
        "steps": [
            {
                "duration": duration,
                "temperature": temperature,
                "read": boolean // optional (default true)
            },
            {
                "duration": duration,
                "gradient": {
                    "top": temperature,
                    "bottom": temperature
                },
                "read": boolean // optional (default true)
            }
        ]
},
```

To specify a melting curve, all four melting-relevant parameters must have a value.

## Parameters

- **object** (`str` or `Ref`) – Container to be thermocycled
- **groups** (`list(dict)`) – List of thermocycling instructions formatted as above
- **volume** (`str` or `Unit`, `optional`) – Volume contained in wells being thermocycled
- **dataref** (`str, optional`) – Name of dataref representing read data if performing qPCR
- **dyes** (`dict, optional`) – Dictionary mapping dye types to the wells they’re used in
- **melting\_start** (`str or Unit`) – Temperature at which to start the melting curve.
- **melting\_end** (`str or Unit`) – Temperature at which to end the melting curve.

- **melting\_increment** (*str or Unit*) – Temperature by which to increment the melting curve. Accepted increment values are between 0.1 and 9.9 degrees celsius.
- **melting\_rate** (*str or Unit*) – Specifies the duration of each temperature step in the melting curve.
- **lid\_temperature** (*str or Unit*) – Specifies the lid temperature throughout the duration of the instruction

#### Raises

- `ValueError` – If one of dataref and dyes is specified but the other isn't.
- `ValueError` – If all melting curve-related parameters are specified but dyes isn't.
- `ValueError` – If some melting curve-related parameteres are specified but not all of them.
- `ValueError` – If invalid dyes are supplied.

**static convert\_well\_map\_to\_dye\_map (well\_map)**

Take a map of wells to the dyes it contains and returns a map of dyes to the list of wells that contain it.

well\_map - [{well:str}]

**static find\_invalid\_dyes (dyes)**

Take a set or list of dye names and returns the set that are not valid.

dyes - [list or set]

**class autoprotocol.instruction.Uncover (object, store\_lid=None)**

Remove lid from specified container

#### Parameters

- **object** (*str*) – Container to remove lid from
- **store\_lid** (*bool*) – Flag to store the uncovered lid

**class autoprotocol.instruction.Unseal (object)**

Remove seal from indicated container using the automated plate unsealer.

**Parameters object** (*Ref or str*) – Container to be unsealed



# CHAPTER 3

---

## autoprotocol.container

---

### 3.1 container.Container

```
class autoprotocol.container.Container(id, container_type, name=None, storage=None,  
                                      cover=None)
```

A reference to a specific physical container (e.g. a tube or 96-well microplate).

Every Container has an associated ContainerType, which defines the well count and arrangement, amongst other properties.

There are several methods on Container which present a convenient interface for defining subsets of wells on which to operate. These methods return a WellGroup.

Containers are usually declared using the Protocol.ref method.

#### Parameters

- **id** (*str*) – Alphanumeric identifier for a Container.
- **container\_type** (*ContainerType*) – ContainerType associated with a Container.
- **name** (*str, optional*) – name of the container/ref being created.
- **storage** (*str, optional*) – name of the storage condition.
- **cover** (*str, optional*) – name of the cover on the container.

**Raises** `AttributeError` – Invalid cover-type given

#### \_\_repr\_\_()

Return a string representation of a Container using the specified name. (ex. `Container('my_plate')`)

#### **all\_wells** (*columnwise=False*)

Return a WellGroup representing all Wells belonging to this Container.

**Parameters** `columnwise` (*bool, optional*) – returns the WellGroup columnwise instead of rowwise (ordered by well index).

**Returns** WellGroup of all Wells in Container

**Return type** *WellGroup***decompose**(*well\_ref*)

Return a tuple representing the column and row number of the well index given based on the ContainerType of the Container.

Uses the decompose function from the ContainerType class. Refer to *ContainerType.decompose()* for more information.

**discard**()

Set the storage condition of a container to None and container to be discarded if ref in protocol.

**Example**

```
p = Protocol()
container = p.ref("new_container", cont_type="96-pcr",
                  storage="cold_20")
p.incubate(c, "warm_37", "30:minute")
container.discard()

Autoprotocol generated:

.. code-block:: json

    "refs": {
        "new_container": {
            "new": "96-pcr",
            "discard": true
        }
    }
```

**humanize**(*well\_ref*)

Return the human readable representation of the integer well index given based on the ContainerType of the Container.

Uses the humanize function from the ContainerType class. Refer to *ContainerType.humanize()* for more information.

**inner\_wells**(*columnwise=False*)

Return a WellGroup of all wells on a plate excluding wells in the top and bottom rows and in the first and last columns.

**Parameters** *columnwise* (*bool, optional*) – returns the WellGroup columnwise instead of rowwise (ordered by well index).

**Returns** WellGroup of inner wells

**Return type** *WellGroup***is\_covered**()

Check if Container is covered.

**is\_sealed**()

Check if Container is sealed.

**quadrant**(*quad*)

Return a WellGroup of Wells corresponding to the selected quadrant of this Container.

This is only applicable to 384-well plates.

**Parameters** `quad` (*int*) – Specifies the quadrant number of the well (ex. 2)

**Returns** WellGroup of wells for the specified quadrant

**Return type** `WellGroup`

**Raises** `ValueError` – Invalid quadrant specified for this Container type

**robotize** (*well\_ref*)

Return the integer representation of the well index given, based on the ContainerType of the Container.

Uses the robotize function from the ContainerType class. Refer to `ContainerType.robotize()` for more information.

**set\_storage** (*storage*)

Set the storage condition of a container, will overwrite an existing storage condition, will remove discard True.

**Parameters** `storage` (*str*) – Storage condition.

**Returns** Container with modified storage condition

**Return type** `Container`

**Raises** `TypeError` – If storage condition not of type str.

**tube** ()

Checks if container is tube and returns a Well representing the zeroth well.

**Returns** Zeroth well of tube

**Return type** `Well`

**Raises** `AttributeError` – If container is not tube

**well** (*i*)

Return a Well object representing the well at the index specified of this Container.

**Parameters** `i` (*int, str*) – Well reference in the form of an integer (ex: 0) or human-readable string (ex: “A1”).

**Returns** Well for given reference

**Return type** `Well`

**Raises** `TypeError` – index given is not of the right type

**wells** (\**args*)

Return a WellGroup containing references to wells corresponding to the index or indices given.

**Parameters** `args` (*str, int, list*) – Reference or list of references to a well index either as an integer or a string.

**Returns** Wells from specified references

**Return type** `WellGroup`

**Raises** `TypeError` – Well reference is not of a valid input type

**wells\_from** (*start, num, columnwise=False*)

Return a WellGroup of Wells belonging to this Container starting from the index indicated (in integer or string form) and including the number of proceeding wells specified. Wells are counted from the starting well rowwise unless columnwise is True.

**Parameters**

- **start** (`Well` or `int` or `str`) – Starting well specified as a Well object, a human-readable well index or an integer well index.
- **num** (`int`) – Number of wells to include in the Wellgroup.
- **columnwise** (`bool`, *optional*) – Specifies whether the wells included should be counted columnwise instead of the default rowwise.

**Returns** WellGroup of selected wells

**Return type** `WellGroup`

**Raises** `TypeError` – Incorrect input types, e.g. `num` has to be of type `int`

## 3.2 container.Well

**class** `autoprotocol.container.Well(container, index)`

A Well object describes a single location within a container.

Do not construct a Well directly – retrieve it from the related Container object.

### Parameters

- **container** (`Container`) – The Container this well belongs to.
- **index** (`int`) – The index of this well within the container.

**\_\_repr\_\_()**

Return a string representation of a Well.

**add\_properties** (`properties`)

Add properties to the properties attribute of a Well. If any key/value pairs are present in both the old and new dictionaries, they will be overwritten by the pairs in the new dictionary.

**Parameters** `properties` (`dict`) – Dictionary of properties to add to a Well.

**Returns** Well with modified properties

**Return type** `Well`

**available\_volume** ()

Returns the available volume of a Well. This is calculated as nominal volume - container\_type dead volume

**Returns** Volume in well

**Return type** `Unit(volume)`

**Raises** `RuntimeError` – Well has no volume

**humanize** ()

Return the human readable representation of the integer well index given based on the ContainerType of the Well.

Uses the humanize function from the ContainerType class. Refer to `ContainerType.humanize()` for more information.

**Returns** Index of well in Container (in human readable form)

**Return type** `str`

**set\_name** (`name`)

Set a name for this well for it to be included in a protocol's "outs" section

**Parameters** `name` (`str`) – Well name.

**Returns** Well with modified name

**Return type** `Well`

**set\_properties** (`properties`)

Set properties for a Well. Existing property dictionary will be completely overwritten with the new dictionary.

**Parameters** `properties` (`dict`) – Custom properties for a Well in dictionary form.

**Returns** Well with modified properties

**Return type** `Well`

**set\_volume** (`vol`)

Set the theoretical volume of liquid in a Well.

**Parameters** `vol` (`str, Unit`) – Theoretical volume to indicate for a Well.

**Returns** Well with modified volume

**Return type** `Well`

**Raises**

- `TypeError` – Incorrect input-type given
- `ValueError` – Volume set exceeds maximum well volume

### 3.3 container.WellGroup

**class** `autoprotocol.container.WellGroup` (`wells`)

A logical grouping of Wells.

Wells in a WellGroup do not necessarily need to be in the same container.

**Parameters** `wells` (`list`) – List of Well objects contained in this WellGroup.

**Raises** `TypeError` – Wells is not of the right input type

**\_\_add\_\_** (`other`)

Append a Well or Wells from another WellGroup to this WellGroup.

**Parameters** `other` (`Well, WellGroup`) –

**Returns** WellGroup with appended wells

**Return type** `WellGroup`

**Raises** `TypeError` – Input given is not of type Well or WellGroup

**\_\_getitem\_\_** (`key`)

Return a specific Well from a WellGroup.

**Parameters** `key` (`int`) – Position in a WellGroup in robotized form.

**Returns** Specified well from given key

**Return type** `Well`

**\_\_len\_\_** ()

Return the number of Wells in a WellGroup.

**\_\_repr\_\_()**

Return a string representation of a WellGroup.

**\_\_setitem\_\_(key, item)**

Set a specific Well in a WellGroup.

**Parameters**

- **key** (*int*) – Position in a WellGroup in robotized form.
- **item** (*Well*) – Well or WellGroup to be added

**Raises** `TypeError` – Item specified is not of type *Well*

**add\_properties(properties)**

Add the same properties for each Well in a WellGroup.

**Parameters** **properties** (*dict*) – Dictionary of properties to set on Well(s).

**Returns** WellGroup with modified properties

**Return type** *WellGroup*

**append(other)**

Append another well to this WellGroup.

**Parameters** **other** (*Well*) – Well to append to this WellGroup.

**Returns** WellGroup with appended well

**Return type** *WellGroup*

**Raises** `TypeError` – other is not of type *Well*

**extend(other)**

Extend this WellGroup with another WellGroup.

**Parameters** **other** (*WellGroup or list of Wells*) – WellGroup to extend this WellGroup.

**Returns** WellGroup extended with specified WellGroup

**Return type** *WellGroup*

**Raises** `TypeError` – Input WellGroup is not of the right type

**indices()**

Return the indices of the wells in the group in human-readable form, given that all of the wells belong to the same container.

**Returns** List of humanized indices from this WellGroup

**Return type** `list(str)`

**insert(i, well)**

Insert a well at a given position.

**Parameters**

- **i** (*int*) – index to insert the well at
- **well** (*Well*) – insert this well at the index

**Returns** WellGroup with inserted wells

**Return type** *WellGroup*

**Raises** `TypeError` – index or well defined does not have right input type

**pop**(*index=-1*)

Removes and returns the last well in the wellgroup, unless an index is specified. If index is specified, the well at that index is removed from the wellgroup and returned.

**Parameters** **index** (*int, optional*) – the index of the well you want to remove and return

**Returns** Well with selected index from WellGroup

**Return type** *Well*

**set\_group\_name**(*name*)

Assigns a name to a WellGroup.

**Parameters** **name** (*str*) – WellGroup name

**Returns** Name of wellgroup

**Return type** str

**set\_properties**(*properties*)

Set the same properties for each Well in a WellGroup.

**Parameters** **properties** (*dict*) – Dictionary of properties to set on Well(s).

**Returns** WellGroup with modified properties

**Return type** *WellGroup*

**set\_volume**(*vol*)

Set the volume of every well in the group to vol.

**Parameters** **vol** (*Unit, str*) – Theoretical volume of each well in the WellGroup.

**Returns** WellGroup with modified volume

**Return type** *WellGroup*

**wells\_with**(*prop, val=None*)

Returns a wellgroup of wells with the specified property and value

**Parameters**

- **prop** (*str*) – the property you are searching for
- **val** (*str, optional*) – the value assigned to the property

**Returns** WellGroup with modified properties

**Return type** *WellGroup*

**Raises** `TypeError` – property or value defined does not have right input type



# CHAPTER 4

---

## autoprotocol.container\_type

---

### 4.1 container\_type.ContainerType

**class** autoprotocol.container\_type.ContainerType

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
  - **shortname** (*str*) – Short name used to refer to a ContainerType.
  - **col\_count** (*int*) – Number of columns a ContainerType contains.
  - **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.

- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

**decompose** (*idx*)

Return the (col, row) corresponding to the given well index.

**Parameters** **idx** (*str or int*) – Well index in either human-readable or integer form.

**Returns** tuple containing the column number and row number of the given well\_ref.

**Return type** tuple

**Raises** `TypeError` – Index given is not of the right parameter type

**humanize** (*well\_ref*)

Return the human readable form of a well index based on the well format of this ContainerType.

Example Usage:

```
>>> p = Protocol()
>>> my_plate = p.ref("my_plate", cont_type="6-flat", discard=True)
>>> my_plate.humanize(0)
'A1'
>>> my_plate.humanize(5)
'B3'
>>> my_plate.humanize('0')
'A1'
```

**Parameters** **well\_ref** (*int, str, list[int or str]*) – Well reference to be humanized in integer or string form. If string is provided, it has to be parseable into an int. Also accepts lists of int or str

**Returns** **well\_ref** – Well index passed as human-readable form.

**Return type** str

**Raises**

- `TypeError` – If well reference given is not an accepted type.
- `ValueError` – If well reference given exceeds container dimensions.

**robotize** (*well\_ref*)

Return a robot-friendly well reference from a number of well reference formats.

Example Usage:

```
>>> p = Protocol()
>>> my_plate = p.ref("my_plate", cont_type="6-flat", discard=True)
>>> my_plate.robotize("A1")
```

(continues on next page)

(continued from previous page)

```

0
>>> my_plate.robotize("5")
5
>>> my_plate.robotize(my_plate.well(3))
3
>>> my_plate.robotize(["A1", "A2"])
[0, 1]

```

**Parameters** `well_ref` (`str, int, Well, list[str or int or Well]`) – Well reference to be robotized in string, integer or Well object form. Also accepts lists of str, int or Well.

**Returns** `well_ref` – Single or list of Well references passed as rowwise integer (left-to-right, top-to-bottom, starting at 0 = A1).

**Return type** int, list

**Raises**

- `TypeError` – If well reference given is not an accepted type.
- `ValueError` – If well reference given exceeds container dimensions.

#### `row_count()`

Return the number of rows of this ContainerType.

## 4.2 Container Types

`autoprotocol.container_type.FLAT384 = ContainerType(name='384-well UV flat-bottom plate', ...)`  
The ContainerType class holds the capabilities and properties of a particular container type.

**Parameters**

- `name` (`str`) – Full name describing a ContainerType.
- `is_tube` (`bool`) – Indicates whether a ContainerType is a tube (container with one well).
- `well_count` (`int`) – Number of wells a ContainerType contains.
- `well_depth_mm` (`float`) – Depth of well(s) contained in a ContainerType in millimeters.
- `well_volume_ul` (`Unit`) – Maximum volume of well(s) contained in a ContainerType in microliters.
- `well_coating` (`str`) – Coating of well(s) in container (ex. collagen).
- `sterile` (`bool`) – Indicates whether a ContainerType is sterile.
- `cover_types` (`list`) – List of valid covers associated with a ContainerType.
- `seal_types` (`list`) – List of valid seals associated with a ContainerType.
- `capabilities` (`list`) –  
**List of capabilities associated with a ContainerType (ex. ["spin", "incubate"]).**
- `shortname` (`str`) – Short name used to refer to a ContainerType.
- `col_count` (`int`) – Number of columns a ContainerType contains.

- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.PCR384 = ContainerType(name='384-well PCR plate', is_tube=False)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** ([Unit](#)) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –

#### List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).

- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.

- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.ECHO384 = ContainerType(name='384-well Echo plate', is_tube=False)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.FLAT384WHITELV = ContainerType(name='384-well flat-bottom low ...')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.

- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.FLAT384WHITETC = ContainerType(name='384-well flat-bottom low ...')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –

**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**

- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.FLAT384CLEAR = ContainerType(name='384-well fully clear high b...')

The ContainerType class holds the capabilities and properties of a particular container type.
```

**Parameters**

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –

**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**

- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some

ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)

- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.FLAT96 = ContainerType(name='96-well flat-bottom plate', is_tube=False)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
  - **shortname** (*str*) – Short name used to refer to a ContainerType.
  - **col\_count** (*int*) – Number of columns a ContainerType contains.
  - **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
  - **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
  - **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
  - **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
  - **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
  - **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.FLAT96UV = ContainerType(name='96-well flat-bottom UV transparent plate', is_tube=False)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.PCR96 = ContainerType(name='96-well PCR plate', is_tube=False,
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).

- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.DEEP96 = ContainerType(name='96-well extended capacity plate',  
The ContainerType class holds the capabilities and properties of a particular container type.
```

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.

- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.V96KF = ContainerType(name='96-well v-bottom King Fisher plate
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** ([Unit](#)) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
  - **shortname** (*str*) – Short name used to refer to a ContainerType.
  - **col\_count** (*int*) – Number of columns a ContainerType contains.
  - **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
  - **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
  - **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
  - **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
  - **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
  - **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.DEEP96KF = ContainerType(name='96-well extended capacity King F
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
  - **shortname** (*str*) – Short name used to refer to a ContainerType.
  - **col\_count** (*int*) – Number of columns a ContainerType contains.
  - **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
  - **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
  - **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
  - **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
  - **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
  - **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.DEEP24 = ContainerType(name='24-well extended capacity plate',
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.

- **well\_volume\_ul** ([Unit](#)) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.MICRO2 = ContainerType(name='2mL Microcentrifuge tube', is_tube=True)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** ([Unit](#)) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.

- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.MICRO15 = ContainerType(name='1.5mL Microcentrifuge tube', is_tube=True)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –

List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).

- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.

- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

`autoprotocol.container_type.FLAT6 = ContainerType(name='6-well cell culture plate', is_tube=True)`  
The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, ”incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

`autoprotocol.container_type.FLAT1 = ContainerType(name='1-well flat-bottom plate', is_tube=True)`  
The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).

- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.FLAT6TC = ContainerType(name='6-well TC treated plate', is_tube=True)
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.

- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.RESSW96HP = ContainerType(name='96-well singlewell highprofile')
The ContainerType class holds the capabilities and properties of a particular container type.
```

### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.

- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.RESMW8HP = ContainerType(name='8-row multiwell highprofile rese...
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** ([Unit](#)) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
  
List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.RESMW12HP = ContainerType(name='12-column multiwell highprofile...
```

The ContainerType class holds the capabilities and properties of a particular container type.

## Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

`autoprotocol.container_type.FLAT96CLEARTC = ContainerType(name='96-well flat-bottom TC treat')`  
The ContainerType class holds the capabilities and properties of a particular container type.

## Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.

- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.V384CLEAR = ContainerType(name='384-well v-bottom polypropylene')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.

- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str*, *optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.ROUND384CLEAR = ContainerType(name='384-well round-bottom plate')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** ([Unit](#)) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –

#### List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).

- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str*, *optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str*, *optional*) – ContainerType vendor catalog number, if available.

- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.RESSW384LP = ContainerType(name='384-well singlewell lowprofile')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
  - **shortname** (*str*) – Short name used to refer to a ContainerType.
  - **col\_count** (*int*) – Number of columns a ContainerType contains.
  - **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
  - **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
  - **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
  - **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
  - **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
  - **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.ECHO384LDV = ContainerType(name='384-well Echo low dead volume')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.

- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –  
**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**
- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** (*Unit*) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** (*Unit*) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** (*Unit, optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

```
autoprotocol.container_type.ECHO384LDVPLUS = ContainerType(name='384-well Echo low dead vol')
```

The ContainerType class holds the capabilities and properties of a particular container type.

#### Parameters

- **name** (*str*) – Full name describing a ContainerType.
- **is\_tube** (*bool*) – Indicates whether a ContainerType is a tube (container with one well).
- **well\_count** (*int*) – Number of wells a ContainerType contains.
- **well\_depth\_mm** (*float*) – Depth of well(s) contained in a ContainerType in millimeters.
- **well\_volume\_ul** (*Unit*) – Maximum volume of well(s) contained in a ContainerType in microliters.
- **well\_coating** (*str*) – Coating of well(s) in container (ex. collagen).
- **sterile** (*bool*) – Indicates whether a ContainerType is sterile.
- **cover\_types** (*list*) – List of valid covers associated with a ContainerType.
- **seal\_types** (*list*) – List of valid seals associated with a ContainerType.
- **capabilities** (*list*) –

**List of capabilities associated with a ContainerType (ex. [“spin”, “incubate”]).**

- **shortname** (*str*) – Short name used to refer to a ContainerType.
- **col\_count** (*int*) – Number of columns a ContainerType contains.
- **dead\_volume\_ul** ([Unit](#)) – Volume of liquid that cannot be aspirated from any given well of a ContainerType via liquid-handling.
- **safe\_min\_volume\_ul** ([Unit](#)) – Minimum volume of liquid to ensure adequate volume for liquid-handling aspiration from any given well of a ContainerType.
- **true\_max\_vol\_ul** ([Unit](#), *optional*) – Maximum volume of well(s) in microliters, often same value as well\_volume\_ul (maximum working volume), however, some ContainerType(s) can have a different value corresponding to a true maximum volume of a well (ex. echo compatible containers)
- **vendor** (*str, optional*) – ContainerType commercial vendor, if available.
- **cat\_no** (*str, optional*) – ContainerType vendor catalog number, if available.
- **prioritize\_seal\_or\_cover** (*str, optional*) – “seal” or “cover”, determines whether to prioritize sealing or covering defaults to “seal”

# CHAPTER 5

---

## autoprotocol.builders

---

Builders Module containing builders, which help build inputs for Instruction parameters

**copyright** 2018 by The Autoprotocol Development Team, see AUTHORS for more details.

**license** BSD, see LICENSE for more details

### 5.1 Summary

These builder methods are used to generate and validate complex data structures used in Autoprotocol specification. Each of them is capable of using their own output as input. Therefore these builders are also used as inline checks in Protocol methods.

---

#### Notes

Generally these builders should not be called from this file directly. They're more easily accessible by referencing a specific Instruction's builders attribute (e.g. *Spectrophotometry.Builders.mode\_params*).

---

#### See also:

**Instruction** Instructions corresponding to each of the builders

**class** autoprotocol.builders.**DispenseBuilders**

These builders are meant for helping to construct arguments in the *Protocol.dispense* method.

**\_\_delattr\_\_**  
x.\_\_delattr\_\_(‘name’) <==> del x.name

**\_\_format\_\_()**  
default object formatter

**\_\_getattribute\_\_**  
x.\_\_getattribute\_\_(‘name’) <==> x.name

**\_\_hash\_\_**

**\_\_reduce\_\_** ()  
helper for pickle

**\_\_reduce\_ex\_\_** ()  
helper for pickle

**\_\_repr\_\_**

**\_\_setattr\_\_**  
x.\_\_setattr\_\_('name', value) <==> x.name = value

**\_\_sizeof\_\_** () → int  
size of object in memory, in bytes

**\_\_str\_\_**

**static column** (*column*, *volume*)  
Generates a validated column parameter.

**Parameters**

- **column** (*int*) –
- **volume** (*str*, [Unit](#)) –

**Returns** Column parameter of type {“column”: int, “volume”: Unit}

**Return type** dict

**columns** (*columns*)  
Generates a validated columns parameter.

**Parameters** **columns** (*list* ({“column”: *int*, “volume”: *str*, [Unit](#)}))

**Returns** List of columns of type ({“column”: int, “volume”: str, Unit})

**Return type** list

**Raises**

- **ValueError** – No *column* specified for columns
- **ValueError** – Non-unique column indices

**static nozzle\_position** (*position\_x=None*, *position\_y=None*, *position\_z=None*)  
Generates a validated nozzle\_position parameter.

**Parameters**

- **position\_x** ([Unit](#), *optional*) –
- **position\_y** ([Unit](#), *optional*) –
- **position\_z** ([Unit](#), *optional*) –

**Returns** Dictionary of nozzle position parameters

**Return type** dict

**shake\_after** (*duration*, *frequency=None*, *path=None*, *amplitude=None*)  
Generates a validated shake\_after parameter.

**Parameters**

- **duration** ([Unit](#), *str*) –
- **frequency** ([Unit](#), *str*, *optional*) –

- **path** (*str, optional*) –
- **amplitude** (*Unit, str, optional*) –

**Returns** Shake after dictionary of type {“duration”: Unit, “frequency”: Unit, “path”: str, “amplitude”: Unit}

**Return type** dict

**Raises** ValueError – Invalid shake path specified

### **shape** (*rows=1, columns=1, format=None*)

Helper function for building a shape dictionary

**Parameters**

- **rows** (*int, optional*) – Number of rows to be concurrently transferred
- **columns** (*int, optional*) – Number of columns to be concurrently transferred
- **format** (*str, optional*) – Plate format in String form. e.g. “SBS96” or “SBS384”

**Returns** shape parameters

**Return type** dict

**Raises**

- TypeError – If rows/columns aren’t ints
- ValueError – If an invalid row/column count is given
- ValueError – If an invalid shape is given
- ValueError – If rows/columns are greater than what is allowed for the format

## **class autoprotocol.builders.InstructionBuilders**

General builders that apply to multiple instructions

- \_\_delattr\_\_**  
x.\_\_delattr\_\_('name') <==> del x.name
- \_\_format\_\_()**  
default object formatter
- \_\_getattribute\_\_**  
x.\_\_getattribute\_\_('name') <==> x.name
- \_\_hash\_\_**
- \_\_reduce\_\_()**  
helper for pickle
- \_\_reduce\_ex\_\_()**  
helper for pickle
- \_\_repr\_\_**
- \_\_setattr\_\_**  
x.\_\_setattr\_\_('name', value) <==> x.name = value
- \_\_sizeof\_\_()** → int  
size of object in memory, in bytes
- \_\_str\_\_**

**shape** (*rows=1, columns=1, format=None*)  
Helper function for building a shape dictionary

#### Parameters

- **rows** (*int, optional*) – Number of rows to be concurrently transferred
- **columns** (*int, optional*) – Number of columns to be concurrently transferred
- **format** (*str, optional*) – Plate format in String form. e.g. “SBS96” or “SBS384”

**Returns** shape parameters

**Return type** dict

#### Raises

- `TypeError` – If rows/columns aren’t ints
- `ValueError` – If an invalid row/column count is given
- `ValueError` – If an invalid shape is given
- `ValueError` – If rows/columns are greater than what is allowed for the format

**class** autoprotocol.builders.**LiquidHandleBuilders**

Builders for LiquidHandle Instructions

#### \_\_delattr\_\_

x.\_\_delattr\_\_(‘name’) <==> del x.name

#### \_\_format\_\_()

default object formatter

#### \_\_getattribute\_\_

x.\_\_getattribute\_\_(‘name’) <==> x.name

#### \_\_hash\_\_

#### \_\_reduce\_\_()

helper for pickle

#### \_\_reduce\_ex\_\_()

helper for pickle

#### \_\_repr\_\_

#### \_\_setattr\_\_

x.\_\_setattr\_\_(‘name’, value) <==> x.name = value

#### \_\_sizeof\_\_() → int

size of object in memory, in bytes

#### \_\_str\_\_

**blowout** (*volume, initial\_z, flowrate=None*)

Helper for building blowout params for LiquidHandleMethods

#### Parameters

- **volume** ([Unit](#) or str) – the volume of the blowout step
- **initial\_z** (*dict*) – the position that the tip should move to prior to blowing out  
See Also LiquidHandle.builders.position\_z

- **flowrate** (*dict, optional*) – the flowrate of the blowout See Also LiquidHandle.builders.flowrate

**Returns** blowout params for a LiquidHandleMethod

**Return type** dict

**static flowrate** (*target, initial=None, cutoff=None, acceleration=None, deceleration=None*)

Helper for building flowrates

**Parameters**

- **target** (*Unit or str*) – Target flowrate
- **initial** (*Unit or str, optional*) – Initial flowrate
- **cutoff** (*Unit or str, optional*) – Cutoff flowrate
- **acceleration** (*Unit or str, optional*) – Volumetric acceleration for initial to target (in ul/s<sup>2</sup>)
- **deceleration** (*Unit or str, optional*) – Volumetric deceleration for target to cutoff (in ul/s<sup>2</sup>)

**Returns** flowrate parameters for a LiquidHandle instruction

**Return type** dict

**static instruction\_mode\_params** (*tip\_type=None*)

Helper for building instruction mode\_params

**Parameters** **tip\_type** (*str, optional*) – the string representation of a tip\_type See Also tip\_type.py

**Returns** mode\_params for a LiquidHandle instruction

**Return type** dict

**location** (*location=None, transports=None*)

Helper for building locations

**Parameters**

- **location** (*Well or str, optional*) – Location refers to the well location where the transports will be carried out
- **transports** (*list(dict), optional*) – Transports refer to the list of transports that will be carried out in the specified location See Also LiquidHandle.builders.transport

**Returns** location parameters for a LiquidHandle instruction

**Return type** dict

**Raises**

- **TypeError** – If locations aren't str/well
- **ValueError** – If transports are specified, but empty

**mix** (*volume, repetitions, initial\_z, asp\_flowrate=None, dsp\_flowrate=None*)

Helper for building mix params for Transfer LiquidHandleMethods

**Parameters**

- **volume** (*Unit or str*) – the volume of the mix step
- **repetitions** (*int*) – the number of times that the mix should be repeated

- **initial\_z** (*dict*) – the position that the tip should move to prior to mixing See Also LiquidHandle.builders.position\_z
- **asp\_flowrate** (*dict, optional*) – the flowrate of the aspiration portions of the mix See Also LiquidHandle.builders.flowrate
- **dsp\_flowrate** (*dict, optional*) – the flowrate of the dispense portions of the mix See Also LiquidHandle.builders.flowrate

**Returns** mix parameters for a LiquidHandleMethod

**Return type** dict

**Raises** TypeError – If repetitions is not an int

**mode\_params** (*liquid\_class=None, position\_x=None, position\_y=None, position\_z=None, tip\_position=None*)

Helper for building transport mode\_params

Mode params contain information about tip positioning and the liquid being manipulated

**Parameters**

- **liquid\_class** (*Enum({ "default", "air" })*, *optional*) – The name of the liquid class to be handled. This affects how vendors handle populating liquid handling defaults.
- **position\_x** (*dict, optional*) – Target relative x-position of tip in well. See Also LiquidHandle.builders.position\_xy
- **position\_y** (*dict, optional*) – Target relative y-position of tip in well. See Also LiquidHandle.builders.position\_xy
- **position\_z** (*dict, optional*) – Target relative z-position of tip in well. See Also LiquidHandle.builders.position\_z
- **tip\_position** (*dict, optional*) – A dict of positions x, y, and z. Should only be specified if none of the other tip position parameters have been specified.

**Returns** mode\_params for a LiquidHandle instruction

**Return type** dict

**Raises**

- ValueError – If liquid\_class is not in the allowed list
- ValueError – If both position\_xlylz and tip\_position are specified

**static move\_rate** (*target=None, acceleration=None*)

Helper for building move\_rates

**Parameters**

- **target** (*Unit or str, optional*) – Target velocity. Must be in units of
- **acceleration** (*Unit or str, optional*) – Acceleration. Must be in units of

**Returns** move\_rate parameters for a LiquidHandle instruction

**Return type** dict

**position\_xy** (*position=None, move\_rate=None*)

Helper for building position\_x and position\_y parameters

**Parameters**

- **position** (*Numeric, optional*) – Target relative x/y-position of tip in well in unit square coordinates.
- **move\_rate** (*dict, optional*) – The rate at which the tip moves in the well See Also LiquidHandle.builders.move\_rate

**Returns** position\_xy parameters for a LiquidHandle instruction

**Return type** dict

**Raises**

- TypeError – If position is non-numeric
- ValueError – If position is not in range

**position\_z** (*reference=None, offset=None, move\_rate=None, detection\_method=None, detection\_threshold=None, detection\_duration=None, detection\_fallback=None, detection=None*)

Helper for building position\_z parameters

**Parameters**

- **reference** (*str, optional*) – Must be one of “well\_top”, “well\_bottom”, “liquid\_surface”, “preceding\_position”
- **offset** (*Unit or str, optional*) – Offset from reference position
- **move\_rate** (*dict, optional*) – Controls the rate at which the tip moves in the well See Also LiquidHandle.builders.move\_rate
- **detection\_method** (*str, optional*) – Must be one of “tracked”, “pressure”, “capacitance”
- **detection\_threshold** (*Unit or str, optional*) – The threshold which must be crossed before a positive reading is registered. This is applicable for capacitance and pressure detection methods
- **detection\_duration** (*Unit or str, optional*) – The contiguous duration where the threshold must be crossed before a positive reading is registered. This is applicable for pressure detection methods
- **detection\_fallback** (*dict, optional*) – Fallback option which will be used if sensing fails See Also LiquidHandle.builders.position\_z
- **detection** (*dict, optional*) – A dict of detection parameters. Should only be specified if none of the other detection parameters have been specified.

**Returns** position\_z parameters for a LiquidHandle instruction

**Return type** dict

**Raises**

- ValueError – If reference was not in the allowed list
- ValueError – If both detection\_method|duration|threshold|fallback and detection are specified
- ValueError – If detection\_method is not in the allowed list
- ValueError – If detection parameters were specified, but the reference position doesn’t support detection

**shape** (*rows=1, columns=1, format=None*)  
Helper function for building a shape dictionary

**Parameters**

- **rows** (*int, optional*) – Number of rows to be concurrently transferred
- **columns** (*int, optional*) – Number of columns to be concurrently transferred
- **format** (*str, optional*) – Plate format in String form. e.g. “SBS96” or “SBS384”

**Returns** shape parameters

**Return type** dict

**Raises**

- `TypeError` – If rows/columns aren’t ints
- `ValueError` – If an invalid row/column count is given
- `ValueError` – If an invalid shape is given
- `ValueError` – If rows/columns are greater than what is allowed for the format

**transport** (*volume=None, pump\_override\_volume=None, flowrate=None, delay\_time=None, mode\_params=None*)  
Helper for building transports

**Parameters**

- **volume** (*Unit or str, optional*) – Volume to be aspirated/dispensed. Positive volume -> Dispense. Negative -> Aspirate
- **pump\_override\_volume** (*Unit or str, optional*) – Calibrated volume, volume which the pump will move
- **flowrate** (*dict, optional*) – Flowrate parameters See Also `LiquidHandle.builders.flowrate`
- **delay\_time** (*Unit or str, optional*) – Time spent waiting after executing mandrel and pump movement
- **mode\_params** (*dict, optional*) – Mode parameters See Also `LiquidHandle.builders.mode_params`

**Returns** transport parameters for a `LiquidHandle` instruction

**Return type** dict

**class** `autoprotocol.builders.SpectrophotometryBuilders`

These builders are meant for helping to construct arguments for the `Spectrophotometry` instruction.

**\_\_delattr\_\_**  
x.**\_\_delattr\_\_**(‘name’) <==> del x.name

**\_\_format\_\_**()  
default object formatter

**\_\_getattribute\_\_**  
x.**\_\_getattribute\_\_**(‘name’) <==> x.name

**\_\_hash\_\_**

**\_\_reduce\_\_**()  
helper for pickle

---

```

__reduce_ex__()           helper for pickle
__repr__
__setattr__               x.__setattr__('name', value) <==> x.name = value
__sizeof__()              → int
                         size of object in memory, in bytes
__str__

static absorbance_mode_params (wells, wavelength, num_flashes=None, settle_time=None)

```

**Parameters**

- **wells** (`iterable(Well)` or `WellGroup`) – Wells to be read.
- **wavelength** (`Unit` or `str`) – The wavelengths at which to make absorbance measurements.
- **num\_flashes** (`int`, *optional*) – The number of discrete reads to be taken and then averaged.
- **settle\_time** (`Unit` or `str`, *optional*) – The time to wait between moving to a well and reading it.

**Returns** Formatted mode\_params for an absorbance mode.**Return type** dict**Raises**

- `TypeError` – Invalid type specified for input parameters, e.g. `num_flashes` not of type `int`
- `ValueError` – Invalid wells specified

```
fluorescence_mode_params (wells, excitation, emission, num_flashes=None, settle_time=None,
                           lag_time=None,      integration_time=None,      gain=None,
                           read_position=None)
```

**Parameters**

- **wells** (`iterable(Well)` or `WellGroup`) – Wells to be read.
- **excitation** (`list(dict)`) – A list of SpectrophotometryBuilders.wavelength\_selection to determine the wavelength(s) of excitation light used.
- **emission** (`list(dict)`) – A list of SpectrophotometryBuilders.wavelength\_selection to determine the wavelength(s) of emission light used.
- **num\_flashes** (`int`, *optional*) – The number of discrete reads to be taken and then combined.
- **settle\_time** (`Unit` or `str`, *optional*) – The time to wait between moving to a well and reading it.
- **lag\_time** (`Unit` or `str`, *optional*) – The time to wait between excitation and reading.
- **integration\_time** (`Unit` or `str`, *optional*) – Time over which the data should be collected and integrated.

- **gain** (*int, optional*) – The amount of gain to be applied to the readings.
- **read\_position** (*str, optional*) – The position from which the wells should be read.

**Returns** Formatted mode\_params for a fluorescence mode.

**Return type** dict

**Raises**

- `TypeError` – Invalid input types, e.g. `settle_time` is not of type `Unit(second)`
- `ValueError` – Invalid wells specified
- `ValueError` – Gain is not between 0 and 1

**group** (*mode, mode\_params*)

**Parameters**

- **mode** (*str*) – A string representation of a valid spectrophotometry mode.
- **mode\_params** (*dict*) – A dict of mode\_params corresponding to the mode.

**Returns** A spectrophotometry group.

**Return type** dict

**Raises** `ValueError` – Invalid mode specified

**groups** (*groups*)

**Parameters** **groups** (*list(dict)*) – A list of spectrophotometry groups.

**Returns** A list of spectrophotometry groups.

**Return type** list(dict)

**static luminescence\_mode\_params** (*wells, num\_flashes=None, settle\_time=None, integration\_time=None, gain=None*)

**Parameters**

- **wells** (*iterable(Well) or WellGroup*) – Wells to be read.
- **num\_flashes** (*int, optional*) – The number of discrete reads to be taken and then combined.
- **settle\_time** (*Unit or str, optional*) – The time to wait between moving to a well and reading it.
- **integration\_time** (*Unit or str, optional*) – Time over which the data should be collected and integrated.
- **gain** (*int, optional*) – The amount of gain to be applied to the readings.

**Returns** Formatted mode\_params for a luminescence mode.

**Return type** dict

**Raises**

- `TypeError` – Invalid input types, e.g. `settle_time` is not of type `Unit(second)`
- `ValueError` – Gain is not between 0 and 1

**shake\_before** (*duration, frequency=None, path=None, amplitude=None*)

**Parameters**

- **duration** (`Unit or str`) – The duration of the shaking incubation.
- **frequency** (`Unit or str, optional`) – The frequency of the shaking motion.
- **path** (`str, optional`) – The name of a shake path. See the spectrophotometry ASC for diagrams of different shake paths.
- **amplitude** (`Unit or str, optional`) – The amplitude of the shaking motion.

**Returns** Formatted mode\_params for a shake mode.

**Return type** dict

**shake\_mode\_params** (`duration=None, frequency=None, path=None, amplitude=None`)

**Parameters**

- **duration** (`Unit or str, optional`) – The duration of the shaking incubation, if not specified then the incubate will last until the end of read interval.
- **frequency** (`Unit or str, optional`) – The frequency of the shaking motion.
- **path** (`str, optional`) – The name of a shake path. See the spectrophotometry ASC for diagrams of different shake paths.
- **amplitude** (`Unit or str, optional`) – The amplitude of the shaking motion.

**Returns** Formatted mode\_params for a shake mode.

**Return type** dict

**shape** (`rows=1, columns=1, format=None`)

Helper function for building a shape dictionary

**Parameters**

- **rows** (`int, optional`) – Number of rows to be concurrently transferred
- **columns** (`int, optional`) – Number of columns to be concurrently transferred
- **format** (`str, optional`) – Plate format in String form. e.g. “SBS96” or “SBS384”

**Returns** shape parameters

**Return type** dict

**Raises**

- `TypeError` – If rows/columns aren’t ints
- `ValueError` – If an invalid row/column count is given
- `ValueError` – If an invalid shape is given
- `ValueError` – If rows/columns are greater than what is allowed for the format

**static wavelength\_selection** (`shortpass=None, longpass=None, ideal=None`)

Generates a representation of a wavelength selection by either filters (using shortpass/longpass) or monochromators (using ideal)

**Parameters**

- **shortpass** (`Unit, str, optional`) –

- **longpass** (`Unit`, `str`, `optional`) –
- **ideal** (`Unit`, `str`, `optional`) –

**Returns** Wavelength selection parameters.

**Return type** dict

```
class autoprotocol.builders.ThermocycleBuilders
```

These builders are meant for helping to construct the `groups` argument in the `Protocol.thermocycle` method

```
__delattr__
    x.__delattr__('name') <==> del x.name

__format__()
    default object formatter

__getattribute__
    x.__getattribute__('name') <==> x.name

__hash__

__reduce__()
    helper for pickle

__reduce_ex__()
    helper for pickle

__repr__

__setattr__
    x.__setattr__('name', value) <==> x.name = value

__sizeof__() → int
    size of object in memory, in bytes

__str__
```

**group** (`steps`, `cycles=1`)

Helper function for creating a thermocycle group, which is a series of steps repeated for the number of cycles

#### Parameters

- **steps** (`list (ThermocycleBuilders.step)`) – Steps to be carried out. At least one step has to be specified. See `ThermocycleBuilders.step` for more information
- **cycles** (`int`, `optional`) – Number of cycles to repeat the specified steps. Defaults to 1

**Returns** A thermocycling group

**Return type** dict

#### Raises

- `TypeError` – Invalid input types, i.e. `cycles` is not of type `int` and `steps` is not of type `list`
- `ValueError` – `cycles` is not positive
- `ValueError` – `steps` does not contain any elements

**shape** (`rows=1`, `columns=1`, `format=None`)

Helper function for building a shape dictionary

#### Parameters

- **rows** (*int, optional*) – Number of rows to be concurrently transferred
- **columns** (*int, optional*) – Number of columns to be concurrently transferred
- **format** (*str, optional*) – Plate format in String form. e.g. “SBS96” or “SBS384”

**Returns** shape parameters

**Return type** dict

**Raises**

- `TypeError` – If rows/columns aren’t ints
- `ValueError` – If an invalid row/column count is given
- `ValueError` – If an invalid shape is given
- `ValueError` – If rows/columns are greater than what is allowed for the format

### **static step** (*temperature, duration, read=None*)

Helper function for creating a thermocycle step.

**Parameters**

- **temperature** (*Unit or dict(str, Unit)*) – Block temperature which the contents should be thermocycled at.

If a gradient thermocycle is desired, specifying a dict with “top” and “bottom” keys will control the desired temperature at the top and bottom rows of the block, creating a gradient along the column.

..code-block:: python

```
temperature = {"top": "50:celsius", "bottom": "45:celsius"}
```

- **duration** (*str or Unit*) – Duration where the specified temperature parameters will be applied
- **read** (*Boolean, optional*) – Determines if a read at wavelengths specified by the dyes in the parent *thermocycle* instruction will be enabled for this particular step. Useful for qPCR applications.

**Returns** A thermocycling step

**Return type** dict

**Raises**

- `TypeError` – Invalid input types, e.g. `read` is not of type bool
- `ValueError` – Invalid format specified for `temperature` dict
- `ValueError` – Duration is not greater than 0 second



# CHAPTER 6

---

## autoprotocol.liquid\_handle

---

### 6.1 liquid\_handle.liquid\_handle\_method

LiquidHandleMethod

Base class for generating complex liquid handling behavior.

#### 6.1.1 Summary

LiquidHandleMethods are passed as arguments to Protocol methods along with LiquidClasses to specify complex series of liquid handling behaviors.

---

#### Notes

Methods in this file should not be used directly, but are intended to be extended by other methods depending on desired behavior.

When creating a vendor-specific library it's likely desirable to monkey patch *LiquidHandleMethod.\_get\_tip\_types* to reference TipTypes that the vendor supports.

---

**class** autoprotocol.liquid\_handle.liquid\_handle\_method.**LiquidHandleMethod**(tip\_type=None, blowout=True)

Base LiquidHandleMethod

General framework for liquid handling abstractions and helpers for building a series of liquid\_handle transports.

---

#### \_shape

*dict* – the SBS shape and number of rows and columns of the liquid\_handle

#### \_transports

*list* – tracks transports to be added to the LiquidHandle instruction

---

## Notes

There is a hierarchy of logic to all LiquidHandleMethods that abstracts a complex set of liquid handling behavior into smaller, discrete steps.

**For step  $x$  (aspirate, dispense, mix) and parameter  $y$  (e.g. blowout):**

- **Protocol method:**
    - calls LiquidHandleMethod.\_‘x‘\_transports
  - **LiquidHandleMethod.\_‘x‘\_transports method:**
    - clears the \_transports list
    - walks through all \_transport methods including \_transport\_‘y‘
    - returns the \_transports lists
  - **LiquidHandleMethod.\_transport\_‘y‘ method:**
    - checks parameter  $y$  in addition to the default\_‘y‘ method
    - possibly generates a series of transports based on the two values
    - calls lower level helper methods
  - **LiquidHandleMethod lower level helper methods:**
    - generate transports and append them to \_transports
- 

## Examples

For specifying a single, global liquid handling behavior across all volumes the easiest way is to specify parameters when instantiating a LiquidHandleMethod.

```
from autoprotocol import Unit
from autoprotocol.instruction import LiquidHandle
from autoprotocol.liquid_handle import LiquidHandleMethod

lhm = LiquidHandleMethod(
    blowout=LiquidHandle.builders.blowout(volume=Unit(10, "uL"))
)
```

For behavior that relies on more liquid handling parameters or even defines new behavior you can define your own LiquidHandleMethod.

```
from autoprotocol import Unit
from autoprotocol.instruction import LiquidHandle
from autoprotocol.liquid_handle import LiquidHandleMethod

class NewLHM(LiquidHandleMethod):
    def default_blowout(self, volume):
        if volume < Unit(10, "uL"):
            blowout_volume = Unit(1, "uL")
        else:
            blowout_volume = Unit(10, "uL")
        return LiquidHandle.builders.blowout(
            volume=blowout_volume
        )
```

**See also:**

**Transfer** method for handling liquid between two locations  
**Mix** method for handling liquid within locations  
**LiquidClass** contain properties that are intrinsic to specific liquids  
**Protocol** contains methods that accept LiquidHandleMethods as arguments

**\_\_delattr\_\_**  
x.\_\_delattr\_\_('name') <==> del x.name

**\_\_format\_\_()**  
default object formatter

**\_\_getattribute\_\_**  
x.\_\_getattribute\_\_('name') <==> x.name

**\_\_hash\_\_**

**\_\_reduce\_\_()**  
helper for pickle

**\_\_reduce\_ex\_\_()**  
helper for pickle

**\_\_repr\_\_**

**\_\_setattr\_\_**  
x.\_\_setattr\_\_('name', value) <==> x.name = value

**\_\_sizeof\_\_()** → int  
size of object in memory, in bytes

**\_\_str\_\_**

**default\_blowout(volume)**  
Default blowout behavior

**Parameters** **volume** ([Unit](#)) –

**Returns** blowout\_params

**Return type** dict

**See also:**

**blowout()** holds any user specified blowout parameters

**\_transport\_blowout()** generates the actual blowout transports

**static default\_lld\_position\_z(liquid)**  
Default lld position\_z

**Returns** position\_z for sensing the liquid surface

**Return type** dict

**static default\_tracked\_position\_z()**  
Default tracked position\_z

**Returns** position\_z for tracking the liquid surface

**Return type** dict

```
static default_well_bottom_position_z()
Default well bottom position_z
```

**Returns** position\_z for the well bottom

**Return type** dict

```
static default_well_top_position_z()
```

Default well top position\_z

**Returns** position\_z for the well top

**Return type** dict

## 6.2 liquid\_handle.liquid\_class

### LiquidClass

Base class for defining the portions of liquid handling behavior that are intrinsic to specific types of liquids.

```
class autoprotocol.liquid_handle.liquid_class.LiquidClass(calibrated_volume=None,
                                                               aspi-
                                                               rate_flowrate=None,
                                                               dis-
                                                               pense_flowrate=None,
                                                               delay_time=None,
                                                               clld_threshold=None,
                                                               plld_threshold=None)
```

Contains properties intrinsic to individual LiquidClasses

#### **name**

*str* – the name of the liquid\_class may be used by vendors to generate more sensible defaults for unspecified behavior

#### **volume\_calibration\_curve**

*dict(str, VolumeCalibration)* – a calibration curve describing the relationship between tip\_type, volume bins, and volume calibration parameters See Also VolumeCalibration

#### **aspirate\_flowrate\_calibration\_curve**

*dict(str, VolumeCalibration)* – a calibration curve describing the relationship between tip\_type, volume bins, and aspirate flowrate calibration parameters See Also VolumeCalibration

#### **dispense\_flowrate\_calibration\_curve**

*dict(str, VolumeCalibration)* – a calibration curve describing the relationship between tip\_type, volume bins, and dispense flowrate calibration parameters See Also VolumeCalibration

#### **\_safe\_volume\_multiplier**

*Numeric* – a multiplier used by LiquidHandleMethods to estimate safe pump buffers for volume calibration without any prior knowledge about tip\_type See Also LiquidHandleMethod.\_estimate\_calibrated\_volume

### Examples

For specifying a single, global liquid handling behavior across all volumes the easiest way is to specify parameters when instantiating a LiquidClass. If the following LiquidClass is specified then the pump\_override\_volume will always be set to 10: $\mu$ L and the flowrate for all aspirate steps will have a target of 10: $\mu$ L/s, regardless of the stated volume to be transferred.

```

from autoprotocol import Unit
from autoprotocol.instruction import LiquidHandle
from autoprotocol.liquid_handle import LiquidClass

lc = LiquidClass(
    aspirate_flowrate=LiquidHandle.builders.flowrate(
        target=Unit(10, "uL/s")
    ),
    calibrated_volume=Unit(10, "uL")
)

```

For behavior that differs between volumes you can define your own LiquidClass.

```

from autoprotocol import Unit
from autoprotocol.instruction import LiquidHandle
from autoprotocol.liquid_handle.liquid_class import (
    LiquidClass, VolumeCalibration, VolumeCalibrationBin
)

vol_curve = {
    "generic_1_50": VolumeCalibration(
        (Unit(5, "uL"), VolumeCalibrationBin(
            slope=1.1, intercept=Unit(0.1, "uL")
        )),
        (Unit(10, "uL"), VolumeCalibrationBin(
            slope=0.9, intercept=Unit(0.2, "uL")
        ))
    )
}
asp_flow_curve = {
    "generic_1_50": VolumeCalibration(
        (Unit(5, "uL"), LiquidHandle.builders.flowrate(
            target=Unit(50, "uL/s")
        )),
        (Unit(15, "uL"), LiquidHandle.builders.flowrate(
            target=Unit(200, "uL/s")
        ))
    )
}

class NewLC(LiquidClass):
    def __init__(self, *args, **kwargs):
        super(NewLC, self).__init__(*args, **kwargs)
        self.volume_calibration_curve = vol_curve
        self.aspirate_flowrate_calibration_curve = asp_flow_curve

```

See also:

[VolumeCalibration](#) used to specify calibration\_curves

[LiquidHandleMethod](#) used to specify liquid handling movement behavior

[Protocol.transfer](#) accepts LiquidClass arguments to determine behavior

[Protocol.mix](#) accepts a LiquidClass argument to determine behavior

```

__delattr__
x.__delattr__('name') <==> del x.name

```

```
__format__()
    default object formatter

__getattribute__
    x.__getattribute__('name') <==> x.name

__hash__

__reduce__()
    helper for pickle

__reduce_ex__()
    helper for pickle

__repr__

__setattr__
    x.__setattr__('name', value) <==> x.name = value

__sizeof__() → int
    size of object in memory, in bytes

__str__

class autoprotocol.liquid_handle.liquid_class.VolumeCalibration(*args)
Wrapper for a volume-binned calibration curve A data structure that represents a calibration curve for either volumes or flowrates that are binned by upper bounded volume ranges.

__delattr__
    x.__delattr__('name') <==> del x.name

__format__()
    default object formatter

__getattribute__
    x.__getattribute__('name') <==> x.name

__hash__

__reduce__()
    helper for pickle

__reduce_ex__()
    helper for pickle

__repr__

__setattr__
    x.__setattr__('name', value) <==> x.name = value

__sizeof__() → int
    size of object in memory, in bytes

__str__

binned_calibration_for_volume(volume)
Gets the smallest suitable bin in the calibration curve Finds the smallest point on the calibration curve that has a bin that's greater than or equal to the size of the specified value.

Parameters volume (Unit or int or float) – the value to be binned
Returns target_bin
Return type dict
Raises RuntimeError – No suitably large calibration bin
```

**class** autoprotocol.liquid\_handle.liquid\_class.**VolumeCalibrationBin**

Wrapper for slope and intercept parameters for linear fitting Holds information required to calibrate a volume for liquid handle step assuming a linear relationship between volume and calibrated volume.

**\_\_add\_\_**

x.\_\_add\_\_(y) <==> x+y

**\_\_contains\_\_**

x.\_\_contains\_\_(y) <==> y in x

**\_\_delattr\_\_**

x.\_\_delattr\_\_('name') <==> del x.name

**\_\_eq\_\_**

x.\_\_eq\_\_(y) <==> x==y

**\_\_format\_\_()**

default object formatter

**\_\_ge\_\_**

x.\_\_ge\_\_(y) <==> x>=y

**\_\_getattribute\_\_**

x.\_\_getattribute\_\_('name') <==> x.name

**\_\_getitem\_\_**

x.\_\_getitem\_\_(y) <==> x[y]

**\_\_getslice\_\_**

x.\_\_getslice\_\_(i, j) <==> x[i:j]

Use of negative indices is not supported.

**\_\_gt\_\_**

x.\_\_gt\_\_(y) <==> x>y

**\_\_hash\_\_****\_\_iter\_\_****\_\_le\_\_**

x.\_\_le\_\_(y) <==> x<=y

**\_\_len\_\_****\_\_lt\_\_**

x.\_\_lt\_\_(y) <==> x<y

**\_\_mul\_\_**

x.\_\_mul\_\_(n) <==> x\*n

**\_\_ne\_\_**

x.\_\_ne\_\_(y) <==> x!=y

**\_\_reduce\_\_()**

helper for pickle

**\_\_reduce\_ex\_\_()**

helper for pickle

**\_\_rmul\_\_**

x.\_\_rmul\_\_(n) <==> n\*x

**\_\_setattr\_\_**

x.\_\_setattr\_\_('name', value) <==> x.name = value

**\_\_sizeof\_\_()** → int  
size of object in memory, in bytes

**\_\_str\_\_**

**calibrate\_volume**(*volume*)  
Calibrates the volume using slope and intercept

**Parameters** **volume** ([Unit](#)) – the volume to be calibrated

**Returns** calibrated volume

**Return type** [Unit](#)

**count**(*value*) → integer – return number of occurrences of value

**index**(*value*[, *start*[, *stop*]]) → integer – return first index of value.  
Raises ValueError if the value is not present.

**intercept**

Alias for field number 1

**slope**

Alias for field number 0

## 6.3 liquid\_handle.transfer

Transfer LiquidHandleMethod

Base LiquidHandleMethod used by Protocol.transfer to generate a series of movements between pairs of wells.

```
class autoprotocol.liquid_handle.transfer.DryWellTransfer(tip_type=None,
                                                       blowout=True,
                                                       prime=True,
                                                       transit=True,
                                                       mix_before=False,
                                                       mix_after=False,
                                                       aspirate_z=None,
                                                       dispense_z=None)
```

Dispenses while tracking liquid without mix\_after

**\_\_delattr\_\_**

x.\_\_delattr\_\_('name') <==> del x.name

**\_\_format\_\_()**

default object formatter

**\_\_getattribute\_\_**

x.\_\_getattribute\_\_('name') <==> x.name

**\_\_hash\_\_**

**\_\_reduce\_\_()**

helper for pickle

**\_\_reduce\_ex\_\_()**

helper for pickle

**\_\_repr\_\_**

**\_\_setattr\_\_**

x.\_\_setattr\_\_('name', value) <==> x.name = value

**\_\_sizeof\_\_()** → int  
size of object in memory, in bytes

**\_\_str\_\_**

**default\_aspirate\_z(volume)**  
Default aspirate\_z parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** aspirate position\_z

**Return type** dict

See also:

**aspirate\_z()** holds any user defined aspirate\_z parameters

**\_transport\_aspirate\_target\_volume()** generates actual aspirate transports

**default\_blowout(volume)**

Default blowout behavior

**Parameters** **volume** ([Unit](#)) –

**Returns** blowout\_params

**Return type** dict

See also:

**blowout()** holds any user specified blowout parameters

**\_transport\_blowout()** generates the actual blowout transports

**default\_dispense\_z(volume)**

Default aspirate\_z parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** dispense position\_z

**Return type** dict

See also:

**dispense\_z()** holds any user defined dispense\_z parameters

**\_transport\_dispense\_target\_volume()** generates actual dispense transports

**static default\_lld\_position\_z(liquid)**

Default lld position\_z

**Returns** position\_z for sensing the liquid surface

**Return type** dict

**default\_mix\_after(volume)**

Default mix\_after parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** mix\_after params

**Return type** dict

See also:

`mix_after()` holds any user defined mix\_after parameters  
`_transport_mix()` generates the actual mix\_after transports

`default_mix_before(volume)`  
Default mix\_before parameters

**Parameters** `volume (Unit)` –  
**Returns** mix\_before params  
**Return type** dict

See also:

`mix_before()` holds any user defined mix\_before parameters  
`_transport_mix()` generates the actual mix\_before transports

`default_prime(volume)`  
Default prime volume

**Parameters** `volume (Unit)` –  
**Returns** priming volume  
**Return type** Unit

See also:

`prime()` holds any user defined prime volume  
`_transport_aspirate_target_volume()` generates actual aspirate transports

`static default_tracked_position_z()`  
Default tracked position\_z  
**Returns** position\_z for tracking the liquid surface  
**Return type** dict

`default_transit(volume)`  
Default transit volume  
**Parameters** `volume (Unit)` –  
**Returns** transit volume  
**Return type** Unit

See also:

`transit()` holds any user defined transit volume  
`_transport_aspirate_transit()` generates the actual transit transports  
`_transport_dispense_transit()` generates the actual transit transports

`static default_well_bottom_position_z()`  
Default well bottom position\_z  
**Returns** position\_z for the well bottom

**Return type** dict

```
static default_well_top_position_z()
    Default well top position_z
```

**Returns** position\_z for the well top

**Return type** dict

```
class autoprotocol.liquid_handle.transfer.PreMixBlowoutTransfer(tip_type=None,
    blowout=True,
    prime=True,
    transit=True,
    mix_before=False,
    mix_after=True,
    aspirate_z=None,
    dispense_z=None,
    pre_mix_blowout=True)
```

Adds an additional blowout before the mix\_after step

- \_\_delattr\_\_**  
x.\_\_delattr\_\_('name') <==> del x.name
- \_\_format\_\_()**  
default object formatter
- \_\_getattribute\_\_**  
x.\_\_getattribute\_\_('name') <==> x.name
- \_\_hash\_\_**
- \_\_reduce\_\_()**  
helper for pickle
- \_\_reduce\_ex\_\_()**  
helper for pickle
- \_\_repr\_\_**
- \_\_setattr\_\_**  
x.\_\_setattr\_\_('name', value) <==> x.name = value
- \_\_sizeof\_\_()** → int  
size of object in memory, in bytes
- \_\_str\_\_**

**default\_aspirate\_z(volume)**  
Default aspirate\_z parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** aspirate position\_z

**Return type** dict

**See also:**

- aspirate\_z()** holds any user defined aspirate\_z parameters
- \_transport\_aspirate\_target\_volume()** generates actual aspirate transports

**default\_blowout** (*volume*)

Default blowout behavior

**Parameters** **volume** ([Unit](#)) –

**Returns** blowout\_params

**Return type** dict

See also:

**blowout** () holds any user specified blowout parameters

**\_transport\_blowout** () generates the actual blowout transports

**default\_dispense\_z** (*volume*)

Default aspirate\_z parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** dispense position\_z

**Return type** dict

See also:

**dispense\_z** () holds any user defined dispense\_z parameters

**\_transport\_dispense\_target\_volume** () generates actual dispense transports

**static default\_lld\_position\_z** (*liquid*)

Default lld position\_z

**Returns** position\_z for sensing the liquid surface

**Return type** dict

**default\_mix\_after** (*volume*)

Default mix\_after parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** mix\_after params

**Return type** dict

See also:

**mix\_after** () holds any user defined mix\_after parameters

**\_transport\_mix** () generates the actual mix\_after transports

**default\_mix\_before** (*volume*)

Default mix\_before parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** mix\_before params

**Return type** dict

See also:

**mix\_before** () holds any user defined mix\_before parameters

`_transport_mix()` generates the actual mix\_before transports

**default\_pre\_mix\_blowout** (*volume*)

Default pre\_mix\_blowout parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** pre\_mix\_blowout params

**Return type** dict

See also:

`pre_mix_blowout()` holds any user defined pre\_mix\_blowout parameters

`_transport_pre_mix_blowout()` generates the actual blowout transports

**default\_prime** (*volume*)

Default prime volume

**Parameters** **volume** ([Unit](#)) –

**Returns** priming volume

**Return type** [Unit](#)

See also:

`prime()` holds any user defined prime volume

`_transport_aspirate_target_volume()` generates actual aspirate transports

**static default\_tracked\_position\_z** ()

Default tracked position\_z

**Returns** position\_z for tracking the liquid surface

**Return type** dict

**default\_transit** (*volume*)

Default transit volume

**Parameters** **volume** ([Unit](#)) –

**Returns** transit volume

**Return type** [Unit](#)

See also:

`transit()` holds any user defined transit volume

`_transport_aspirate_transit()` generates the actual transit transports

`_transport_dispense_transit()` generates the actual transit transports

**static default\_well\_bottom\_position\_z** ()

Default well bottom position\_z

**Returns** position\_z for the well bottom

**Return type** dict

**static default\_well\_top\_position\_z** ()

Default well top position\_z

**Returns** position\_z for the well top

**Return type** dict

```
class autoprotocol.liquid_handle.transfer.Transfer(tip_type=None,    blowout=True,
                                                    prime=True,           trans-
                                                    sit=True,            mix_before=False,
                                                    mix_after=True,       aspir-
                                                    rate_z=None,         dispense_z=None)
```

LiquidHandleMethod for generating transfers between pairs of wells

LiquidHandleMethod for transferring volume from one well to another.

#### \_source\_liquid

*LiquidClass* – used to determine calibration, flowrates, and sensing thresholds

#### \_destination\_liquid

*LiquidClass* – used to determine calibration, flowrates, and sensing thresholds

---

## Notes

The primary entry points that for this class are:

- `_aspirate_transports` : generates transports for a source location
- `_dispense_transports` : generates transports for a destination location

---

## See also:

`LiquidHandleMethod` base LiquidHandleMethod with reused functionality

`Protocol.transfer` the standard interface for interacting with Transfer

#### `__delattr__`

`x.__delattr__('name')` <==> `del x.name`

#### `__format__()`

default object formatter

#### `__getattribute__`

`x.__getattribute__('name')` <==> `x.name`

#### `__hash__`

#### `__reduce__()`

helper for pickle

#### `__reduce_ex__()`

helper for pickle

#### `__repr__`

#### `__setattr__`

`x.__setattr__('name', value)` <==> `x.name = value`

#### `__sizeof__()` → int

size of object in memory, in bytes

#### `__str__`

#### `default_aspirate_z(volume)`

Default aspirate\_z parameters

**Parameters** `volume` ([Unit](#)) –

**Returns** aspirate position\_z

**Return type** dict

See also:

`aspirate_z()` holds any user defined aspirate\_z parameters

`_transport_aspirate_target_volume()` generates actual aspirate transports

`default_blowout(volume)`

Default blowout behavior

**Parameters** `volume` ([Unit](#)) –

**Returns** blowout\_params

**Return type** dict

See also:

`blowout()` holds any user specified blowout parameters

`_transport_blowout()` generates the actual blowout transports

`default_dispense_z(volume)`

Default dispense\_z parameters

**Parameters** `volume` ([Unit](#)) –

**Returns** dispense position\_z

**Return type** dict

See also:

`dispense_z()` holds any user defined dispense\_z parameters

`_transport_dispense_target_volume()` generates actual dispense transports

`static default_lld_position_z(liquid)`

Default lld position\_z

**Returns** position\_z for sensing the liquid surface

**Return type** dict

`default_mix_after(volume)`

Default mix\_after parameters

**Parameters** `volume` ([Unit](#)) –

**Returns** mix\_after params

**Return type** dict

See also:

`mix_after()` holds any user defined mix\_after parameters

`_transport_mix()` generates the actual mix\_after transports

**default\_mix\_before**(*volume*)

Default mix\_before parameters

**Parameters** **volume** ([Unit](#)) –

**Returns** mix\_before params

**Return type** dict

See also:

**mix\_before()** holds any user defined mix\_before parameters

**\_transport\_mix()** generates the actual mix\_before transports

**default\_prime**(*volume*)

Default prime volume

**Parameters** **volume** ([Unit](#)) –

**Returns** priming volume

**Return type** [Unit](#)

See also:

**prime()** holds any user defined prime volume

**\_transport\_aspirate\_target\_volume()** generates actual aspirate transports

**static default\_tracked\_position\_z()**

Default tracked position\_z

**Returns** position\_z for tracking the liquid surface

**Return type** dict

**default\_transit**(*volume*)

Default transit volume

**Parameters** **volume** ([Unit](#)) –

**Returns** transit volume

**Return type** [Unit](#)

See also:

**transit()** holds any user defined transit volume

**\_transport\_aspirate\_transit()** generates the actual transit transports

**\_transport\_dispense\_transit()** generates the actual transit transports

**static default\_well\_bottom\_position\_z()**

Default well bottom position\_z

**Returns** position\_z for the well bottom

**Return type** dict

**static default\_well\_top\_position\_z()**

Default well top position\_z

**Returns** position\_z for the well top

**Return type** dict

## 6.4 liquid\_handle.mix

Mix LiquidHandleMethod

Base LiquidHandleMethod used by Protocol.mix to generate a series of movements within individual wells.

```
class autoprotocol.liquid_handle.mix.Mix(tip_type=None, blowout=True, repetitions=None, position_z=None)
```

LiquidHandleMethod for generating transfers within wells

LiquidHandleMethod for moving volume within a single well.

liquid

*LiquidClass* – used to determine calibration, flowrates, and sensing thresholds

### Notes

The primary entry points that for this class are:

- `_mix_transports` : generates transports within a single location

### See also:

`LiquidHandleMethod` base LiquidHandleMethod with reused functionality

`Protocol.mix` the standard interface for interacting with Mix

\_\_delattr\_\_

`x.__delattr__('name')` <==> `del x.name`

\_\_format\_\_()

default object formatter

\_\_getattribute\_\_

`x.__getattribute__('name')` <==> `x.name`

\_\_hash\_\_

\_\_reduce\_\_()

helper for pickle

\_\_reduce\_ex\_\_()

helper for pickle

\_\_repr\_\_

\_\_setattr\_\_

`x.__setattr__('name', value)` <==> `x.name = value`

\_\_sizeof\_\_() → int

size of object in memory, in bytes

\_\_str\_\_

`default_blowout(volume)`

Default blowout behavior

Parameters `volume` ([Unit](#)) –

**Returns** blowout\_params

**Return type** dict

See also:

`blowout()` holds any user specified blowout parameters

`_transport_blowout()` generates the actual blowout transports

**static default\_lld\_position\_z(liquid)**

Default lld position\_z

**Returns** position\_z for sensing the liquid surface

**Return type** dict

**default\_position\_z(volume)**

Default position\_z

**Parameters** volume ([Unit](#)) –

**Returns** mix position\_z

**Return type** dict

See also:

`position_z()` holds any user defined position\_z parameters

`_transport_mix()` generates the actual mix transports

**default\_repetitions(volume)**

Default mix repetitions

**Parameters** volume ([Unit](#)) –

**Returns** number of mix repetitions

**Return type** int

See also:

`repetitions()` holds any user defined repetition parameters

`_transport_mix()` generates the actual mix transports

**static default\_tracked\_position\_z()**

Default tracked position\_z

**Returns** position\_z for tracking the liquid surface

**Return type** dict

**static default\_well\_bottom\_position\_z()**

Default well bottom position\_z

**Returns** position\_z for the well bottom

**Return type** dict

**static default\_well\_top\_position\_z()**

Default well top position\_z

**Returns** position\_z for the well top

**Return type** dict

## 6.5 liquid\_handle.tip\_type

Generic tip type and device class mappings for LiquidHandleMethods



---

## autoprotocol support

---

### 7.1 autoprotocol.unit

#### 7.1.1 unit.Unit

`class autoprotocol.unit.Unit(value, units=None)`

A representation of a measure of physical quantities such as length, mass, time and volume. Uses Pint's Quantity as a base class for implementing units and inherits functionalities such as conversions and proper unit arithmetic. Note that the magnitude is stored as a double-precision float, so there are inherent issues when dealing with extremely large/small numbers as well as numerical rounding for non-base 2 numbers.

#### Example

```
vol_1 = Unit(10, 'microliter')
vol_2 = Unit(10, 'liter')
print(vol_1 + vol_2)

time_1 = Unit(1, 'second')
speed_1 = vol_1/time_1
print (speed_1)
print (speed_1.to('liter/hour'))
```

#### Returns

unit object

```
10000010.0:microliter
10.0:microliter / second
0.036:liter / hour
```

**Return type** *Unit*

`__str__(ndigits=12)`

**Parameters** `ndigits` (`int`, *optional*) – Number of decimal places to round to, useful for numerical precision reasons

**Returns** This rounds the string presentation to 12 decimal places by default to account for the majority of numerical precision issues

**Return type** `str`

`ceil()`

Equivalent of `math.ceil(Unit)` for python 2 compatibility

**Returns** `ceil` of `Unit`

**Return type** `Unit`

`floor()`

Equivalent of `math.floor(Unit)` for python 2 compatibility

**Returns** `floor` of `Unit`

**Return type** `Unit`

`round(ndigits)`

Equivalent of `round(Unit)` for python 2 compatibility

**Parameters** `ndigits` (`int`) – number of decimal places to be rounded to

**Returns** rounded unit

**Return type** `Unit`

## 7.2 autoprotocol.util

### 7.2.1 util.incubate\_params()

`autoprotocol.util.incubate_params(duration, shake_amplitude=None, shake_orbital=None)`

Create a dictionary with incubation parameters which can be used as input for instructions. Currently supports plate reader instructions and could be extended for use with other instructions.

**Parameters**

- `shake_amplitude` (`str` or `Unit`) – amplitude of shaking between 1 and 6:milimeter
- `shake_orbital` (`bool`) – True for orbital and False for linear shaking
- `duration` (`str` or `Unit`) – time for shaking

**Returns** Dictionary of incubate parameters

**Return type** `dict`

**Raises** `RuntimeError` – Specifying only shake amplitude or shake orbital

## 7.2.2 util.make\_band\_param()

```
autoprotocol.util.make_band_param(elution_buffer, elution_volume, max_bp, min_bp, destination)
```

Support function to generate gel extraction parameters The *Protocol.gel\_purify()* instruction requires band parameters for extraction, which this function will generate.

### Parameters

- **elution\_buffer** (*str*) – Elution buffer to use to retrieve band
- **elution\_volume** (*str or Unit*) – Volume to elute band into
- **max\_bp** (*int*) – Max basepairs of band
- **min\_bp** (*int*) – Min basepairs of band
- **destination** (*Well*) – Well to place extracted band into

**Returns** Dictionary of band parameters

**Return type** dict

## 7.2.3 util.make\_gel\_extract\_params()

```
autoprotocol.util.make_gel_extract_params(source, band_list, lane=None, gel=None)
```

Support function to generate gel extraction parameters The *Protocol.gel\_purify()* instruction requires a list of extraction parameters, which this function helps to generate.

### Parameters

- **source** (*well*) – Source well for the extraction
- **band\_list** (*list or dict*) – List of bands to collect from the source (use `make_band_param` to make a band dictionary)
- **lane** (*int, optional*) – Lane to load and collect the source. If not set, lane will be auto-generated
- **gel** (*int, optional*) – Gel to load and collect the source. If not set, gel will be auto-generated

**Returns** Dictionary of gel extract parameters

**Return type** dict

## 7.3 autoprotocol.harness

### 7.3.1 harness.run()

```
autoprotocol.harness.run(fn, protocol_name=None, seal_after_run=True)
```

Run the protocol specified by the function.

If `protocol_name` is passed, use preview parameters from the protocol with the matching “name” value in the `manifest.json` file to run the given function. Otherwise, take configuration JSON file from the command line and run the given function.

### Parameters

- **fn** (*function*) – Function that generates Autoprotocol

- **protocol\_name** (*str, optional*) – str matching the “name” value in the manifest.json file
- **seal\_after\_run** (*bool, optional*) – Implicitly add a seal/cover to all stored refs within the protocol using seal\_on\_store()

### 7.3.2 harness.seal\_on\_store()

autoprotocol.harness.**seal\_on\_store**(*protocol*)

Implicitly adds seal/cover instructions to the end of a run for containers that do not have a cover. Cover type applied defaults first to “seal” if its within the capabilities of the container type, otherwise to “cover”.

Example Usage:

```
def example_method(protocol, params):  
    cont = params['container']  
    p.transfer(cont.well("A1"), cont.well("A2"), "10:microliter")  
    p.seal(cont)  
    p.unseal(cont)  
    p.cover(cont)  
    p.uncover(cont)
```

Autoprotocol Output:

```
{  
    "refs": {  
        "plate": {  
            "new": "96-pcr",  
            "store": {  
                "where": "ambient"  
            }  
        }  
    },  
    "instructions": [  
        {  
            "groups": [  
                {  
                    "transfer": [  
                        {  
                            "volume": "10.0:microliter",  
                            "to": "plate/1",  
                            "from": "plate/0"  
                        }  
                    ]  
                }  
            ],  
            "op": "pipette"  
        },  
        {  
            "object": "plate",  
            "type": "ultra-clear",  
            "op": "seal"  
        },  
        {  
            "object": "plate",  
            "op": "unseal"  
        },  
    ]  
}
```

(continues on next page)

(continued from previous page)

```
{
    "lid": "universal",
    "object": "plate",
    "op": "cover"
},
{
    "object": "plate",
    "op": "uncover"
},
{
    "type": "ultra-clear",
    "object": "plate",
    "op": "seal"
}
]
```

### 7.3.3 harness.Manifest

**class** autoprotocol.harness.Manifest(*json\_dict*)

Object representation of a manifest.json file

**Parameters** **object** (*JSON object*) – A manifest.json file with the following format:

```
{
    "format": "python",
    "license": "MIT",
    "description": "This is a protocol.",
    "protocols": [
        {
            "name": "SampleProtocol",
            "version": 1.0.0,
            "command_string": "python sample_protocol.py",
            "preview": {
                "refs": {},
                "parameters": {},
                "inputs": {},
                "dependencies": []
            }
        }
    ]
}
```



# CHAPTER 8

---

## Changelog

---

- #161: shift *op* as an official attribute of Instruction
- #161: add ideal time constraints which can be specified by *add\_time\_constraint* (ASC-037)
- #161: add *batch\_containers*, for controlling containers entering/exiting together
- #161: modify *acoustic\_transfer* to no longer proactively group consecutive instructions. Please use *WellGroup* explicitly instead
- #161: add *util.parse\_unit*, a helper for parsing and checking an unit input
- #161: add *util.check\_unit*, a helper for checking the units in bounds
- #161: all protocol methods now return the Instruction
- #161: add parameters to *p.dispense*, including *flowrate*, *nozzle\_position*, *step\_size*, *reagent\_source*, *dispense\_speed*, *pre\_dispense*, *shape*, *shake\_after* options (ASC-027, ASC-029, ASC-036, ASC-039)
- #161: add *lid\_temperature* to *p.thermocycle* (ASC-035)
- #161: add *settle\_time* to *p.absorbance* (ASC-026)
- #161: add parameters to *p.fluorescence*, including *detection\_mode*, *position\_z*, *settle\_time*, *lag\_time*, *integration\_time* (ASC-026)
- #161: add parameters to *p.luminescence*, including *settle\_time*, *integration\_time* (ASC-026)
- #161: add parameters to *p.seal*, including *mode*, *temperature*, *duration* (ASC-034)
- #161: add *retrieve\_lid* to *p.cover* (ASC-040)
- #161: add *store\_lid* to *p.uncover* (ASC-040)
- #161: change *measure\_mass* instruction to take in a single container instead (ASC-030)
- #161: add *count\_cells* instruction (ASC-033)
- #161: add *spectrophotometry* instruction (ASC-038)
- #161: added builtin support for *ceil* and *floor* and changed py2 compatibility *Unit.floor* and *Unit.ceil* methods to use them

- #161: converted all Unit internals to use Decimals in place of other Numbers
- #161: deprecate `util.make_dottable_dict` and `util.deep_merge_params`
- #161: deprecate `newpick` in `p.autopick`
- #161: deprecate support for generating multiple GelSeparate instructions using `p.gel_separate`
- #161: deprecate support for `p.append` in favor of `p._append_and_return`
- #163: deprecated Pipette, Stamp, Consolidate, Distribute, and Spread instructions
- #163: deprecated the `p.consolidate` and `p.distribute` protocol methods
- #163: replaced the internals of `p.spread` with a new implementation that generates a liquid\_handle instruction
- #163: replaced `p.stamp` & `p.transfer` with a new implementation of `p.transfer` that generates a liquid\_handle instruction
- #163: add LiquidHandleMethods and corresponding protocol methods to represent generic liquid handling abstractions
- #163: add liquid\_handle instruction (ASC-032)
- #168: improved pruning of empty data structures from ‘Instruction.data’ field
- #160: change default linter to pylint and update tox
- #161: cleaned up references of `Unit.fromstring` and `Unit._magnitude`
- #162: fix and update docstrings so that sphinx can be executed with no warnings
- #164: update `docs/requirements.txt` for rtd to build properly
- : fix documentation typos
- : add new containers, `true_max_vol_ul` in `_CONTAINER_TYPES`
- : add prioritize\_seal\_or\_cover allow priority selection
- : allow breathable seals on 96-deep and 24-deep
- : add PerkinElmer 384-well optiplate to container\_type (cat# 6007299), `container-type-384-flat-white-white-optiplate`
- : add support for `more_than` in `add_time_constraint`
- : add ability to specify `x_cassette` for dispense and dispense\_full\_plate methods
- : add ability to specify a well as reagent source for dispense and dispense\_full\_plate methods
- : add `step_size` to dispense and dispense\_full\_plate methods
- : add shaking capabilities to `protocol.incubate()`
- : add `ceil` and `floor` methods to `Unit`
- : remove cover prior to mag steps where applicable
- : convert test suite to py.test
- : docstring cleanup, linting
- : update default lid types for `container-type-384-echo`, `container-type-96-flat`, `container-type-96-flat-uv`, and `container-type-96-flat-clear-clear-tc`
- : update pint requirements, update error handling on UnitError
- : new plate types `container-type-384-v-clear-clear`, `container-type-384-round-clear-clear`, ‘384-flat-white-white-nbs’

- : allow incubation of containers at ambient without covers
- : prevent invalid incubate parameters in *protocol-absorbance*
- : respect incubate conditions where uncovered=True
- : fix Well.set\_properties() so that it completely overwrites the existing properties dict
- : fix name of *container-type-384-round-clear-clear*
- : more descriptive error message in ref protocol
- : add functions and tests to enable use of *-dye\_test* flag
- : Container method: *container-tube*
- : new plate type *container-type-96-flat-clear-clear-tc*
- : Unit validations from str in *protocol-flow-analyze* instruction
- : update documentation for *harness-seal-on-store*
- : cover\_types and seal\_types to \_CONTAINER\_TYPES
- : validations of input types before cover check
- : new plate types *container-type-384-flat-clear-clear*, *container-type-384-flat-white-white-lv*, *container-type-384-flat-white-white-tc*
- : validations before implicit cover or seal
- : WellGroup.extend(wells) can now take in a list of wells
- : WellGroup methods: *wellgroup-group-name*, *wellgroup-pop*, *wellgroup-insert*, *wellgroup-wells-with*
- : assertions and tests for *protocol-flow-analyze*
- : autocover before *protocol-incubate*
- : *is\_resource\_id* added to *protocol-dispense* and *protocol-dispense-full-plate* instructions
- : plate type *container-type-6-flat-tc* to ContainerType
- : unit conversion to microliters in *protocol-dispense* instruction
- : documentation
- : *protocol-dispense* instruction tests
- : using release for changelog and integration into readthedocs documentation
- : convert pipette operations to microliters
- : #128: cover\_types on *container-type-96-deep-kf* and *container-type-96-deep*
- : #127: convert pipette operations to microliters
- : dispense\_speed and distribute\_target in *protocol-distribute* instruction
- : plate type *container-type-6-flat-tc* to ContainerType
- : auto-uncover before *protocol-provision* instructions
- : WellGroup.extend(wells) can now take in a list of wells
- : WellGroup methods: *wellgroup-group-name*, *wellgroup-pop*, *wellgroup-insert*, *wellgroup-wells-with*
- : assertions and tests for *protocol-flow-analyze*
- : autocover before *protocol-incubate*

- : *is\_resource\_id* added to *protocol-dispense* and *protocol-dispense-full-plate* instructions
- : compatibility with py3 in *protocol-flow-analyze*
- : *protocol-spin* auto-cover
- : removed capability ‘cover’ from *container-type-96-pcr* and *container-type-384-pcr* plates
- : *protocol-dispense* instruction json outputs
- : documentation
- : new plate types *container-type-384-flat-clear-clear*, *container-type-384-flat-white-white-lv*, *container-type-384-flat-white-white-tc*
- : validations before implicit cover or seal
- : cover\_types and seal\_types to \_CONTAINER\_TYPES
- : validations of input types before cover check
- : string input types for source, destination wells for Instructions *protocol-consolidate*, *protocol-autopick*, *protocol-mix*
- : track plate cover status - Container objects now have a *cover* attribute, implicit plate unsealing or uncovering prior to steps that require the plate to be uncovered.
- : *protocol-stamp* separates row stamps with more than 2 containers
- : *protocol-mix* allows one\_tip=True
- : *unit-unit* specific error handling
- : *protocol-illuminaseq* allows cycle specification
- : *protocol-add-time-constraint* added
- : harness.py returns proper boolean for thermocycle types
- : *unit-unit* specific error handling
- : thermocycle gradient steps in harness.py
- : *protocol-mix* allows one\_tip=True
- : *protocol-acoustic-transfer* handling of droplet size
- : *protocol-spin* instruction takes directional parameters
- : *protocol-gel-purify* parameters improved
- : *protocol-illuminaseq* instruction
- : *protocol-measure-volume* instruction
- : *protocol-measure-mass* instruction
- : Compatibility of Unit for acceleration
- : fix harness to be python3 compatible
- : Concatenation of Well to WellGroup no longer returns None
- : WellGroup checks that all elements are wells
- : gel string in documentation
- : support for list input type for humanize and robotize (container and container\_type)
- : *protocol-gel-purify* instruction to instruction.py and protocol.py

- :`:ref:container-discard`` and `and container-set-storage` methods for containers
- : csv-table input type to harness.py
- : magnetic transfer instructions to now pass relevant inputs through units
- : Unit support for *molar*
- : disclaimer to README.md on unit support
- : `Unit(Unit(...))` now returns a Unit
- : checking for valid plate read incubate parameters
- : additional parameter, *gain*, to *protocol-fluorescence*
- : documentation for magnetic transfer instructions correctly uses hertz
- : adding magnetic transfer functions to documentation
- : support for a new instruction for *protocol-measure-concentration*
- : helper function in util.py to create incubation dictionaries
- : additional parameters to spectrophotometry instructions (*protocol-absorbance*, *protocol-luminescence*, *protocol-fluorescence*) to instruction.py and protocol.py
- : Updated Unit package to default to *Autoprotocol* format representation for temperature and speed units
- : Updated maximum tip capacity for a transfer operation to 900uL instead of 750uL
- : Updated handling of multiplication and division of Units of the same dimension to automatically resolve when possible
- : *unit-unit* now uses Pint's Quantity as a base class
- : update *container-type-6-flat* well volumes
- : release versioning has been removed in favor of protocol versioning in harness.py
- : kf container types *container-type-96-v-kf* and *container-type-96-deep-kf* in container\_type.py
- : *magnetic\_transfer* instruction to instruction.py and protocol.py
- : *container+* input type to harness.py
- : Update container\_test.py and container\_type\_test.py to include safe\_min\_volume\_ul
- : default versioning in manifest\_test.json
- : updated dead\_volume\_ul values in \_CONTAINER\_TYPES
- : safe\_min\_volume\_ul in \_CONTAINER\_TYPES
- : *protocol-stamp* smartly calculates max\_tip\_volume using residual volumes
- : *protocol-autopick* now conforms to updated ASC (**not backwards compatible**)
- : Allow single Well reading for Absorbance, Fluorescence and Luminescence
- : *wellgroup-extend* method to WellGroup
- : Include well properties in outs
- : *protocol-transfer* respects when *mix\_after* or *mix\_before* is explicitly False
- : Protocol.stamp() allows one\_tip=True when steps use a *mix\_vol* greater than “31:microliter” even if transferred volumes are not all greater than “31:microliter”
- : Protocol.plate\_to\_magblock() and Protocol.plate\_from\_magblock()

- : *protocol-stamp* has been reformatted to take groups of transfers. This allows for `one_tip=True`, `one_source=True`, and `WellGroup` source and destinations
- : `one_tip = True` transfers > 750:microliter are transferred with single tip
- : volume tracking for *protocol-stamp* ing to/from 384-well plates
- : functionality to harness.py for naming aliquots
- : `UserError` exception class for returning custom errors from within protocol scripts
- : more recursion in `make_dottable_dict`, a completely unnecessary function you shouldn't use
- : Better handling of default `append=true` behavior for *protocol-stamp*
- : Small bug for transfer with `one_source=true` fixed
- : Transfers with `one_source true` does not keep track of the value of volume less than  $10^{-12}$
- : *protocol-stamp* transfers are not combinable if they use different tip volume types
- : unit conversion from milliliters or nanoliters to microliters in `Well.set_volume()`, `protocol-provision`, `protocol-transfer`, and `protocol-distribute`
- : “outs” section of protocol. Use `well-set-name` to name an aliquot
- : name property on Well
- : volume tracking to *protocol-stamp* and associated helper functions in `autoprotocol.util`
- : Arguments to *protocol-transfer* for `mix_before` and `mix_after` are now part of **mix\_kwargs** to allow for specifying separate parameters for `mix_before` and `mix_after`
- : Better error handling in harness.py and accompanying tests
- : Test for more complicated *transfer'ing with 'one\_source=True*
- : manually change storage condition destiny of a Container
- : `Protocol.store()`
- : Storage attribute on Container
- : *protocol-stamp* now support selective (row-wise and column-wise) stamping (see docstring for details)
- : semantic versioning fail
- : Arguments to *protocol-transfer* for `mix_before` and `mix_after` are now part of **mix\_kwargs** to allow for specifying separate parameters for `mix_before` and `mix_after`
- : Better error handling in harness.py and accompanying tests
- : Test for more complicated *transfer'ing with 'one\_source=True*
- : manually change storage condition destiny of a Container
- : `Protocol.store()`
- : Storage attribute on Container
- : Error with *transfer'ing with 'one\_source=True*
- : unit conversion from milliliters or nanoliters to microliters in `Well.set_volume()`, `protocol-provision`, `protocol-transfer`, and `protocol-distribute`
- : “outs” section of protocol. Use `well-set-name` to name an aliquot
- : name property on Well
- : volume tracking to *protocol-stamp* and associated helper functions in `autoprotocol.util`

- : Unit scalar multiplication
- : Error when *protocol-transfer* ing over 750 $\mu$ L
- : Error with *protocol-provision* ing to multiple wells of the same container
- : semantic versioning fail
- : *protocol-stamp* now utilizes the new Autoprotocol *stamp* instruction instead of *protocol-transfer*
- : `__repr__` override for Unit class
- : volume tracking to destination wells when using `Protocol.dispense()`
- : *Stamp* class in `autoprotocol.instruction`
- : better error handling for *protocol-transfer* and *protocol-distribute*
- : refactored `Protocol` methods: *protocol-ref*, *protocol-consolidate*, *protocol-transfer*, *protocol-distribute*
- : fixed indentation
- : warnings for `_mul_` and `_div_` scalar Unit operations
- : *protocol-dispense-full-plate*
- : *protocol-dispense* Instruction and accompanying `Protocol` method for using a reagent dispenser
- : aliquot++, integer, boolean input types to `harness.py`
- : properties attribute to `Well`, along with *well-set-properties* method
- : melting keyword variables and changes to conditionals in `Thermocycle`
- : default input value and group and group+ input types in `harness.py`
- : `Protocol.pipette()` is now a private method `_pipette()`
- : *protocol-sangerseq* Instruction and method
- : seal takes a “type” parameter that defaults to ultra-clear
- : more tests
- : *protocol-stamp* Protocol method for using the 96-channel liquid handler
- : Added `pipette_tools` module containing helper methods for the extra pipetting parameters
- : Additional keyword arguments for *protocol-transfer* and *protocol-distribute* to customize pipetting
- : *protocol-oligosynthesize* Instruction
- : *protocol-autopick* Instruction
- : *protocol-spread* Instruction
- : *protocol-flow-analyze* Instruction
- : co2 parameter in *protocol-incubate*
- : 6-flat container type in `_CONTAINER_TYPES`
- : 96-flat-uv container type in `_CONTAINER_TYPES`
- : *container-quadrant* returns a WellGroup of the 96 wells representing the quadrant passed
- : *well-add-properties*
- : `Well.properties` is an empty hash by default
- : At least some Python3 compatibility

- : *protocol-stamp* ing to or from multiple containers now requires that the source or dest variable be passed as a list of [{“container”: <container>, “quadrant”: <quadrant>}, … ]
- : *protocol-gel-separate* generates instructions taking wells and matrix type passed
- : tox for testing with multiple versions of python
- : *new\_group* keyword parameter on *protocol-transfer* and *protocol-distribute* to manually break up *Pipette()* Instructions
- : Thermocycle input type in *harness.py*
- : specify *Wells* on a container using *container.wells(1,2,3)*‘or ‘*container.wells([1,2,3])*
- : More Python3 Compatibility
- : More Python3 Compatibility
- : Additional type-checks in various functions
- : *protocol-provision* Protocol method
- : support for *choice* input type in *harness.py*
- : allow transfer from multiple sources to one destination
- : brought back recursively transferring volumes over 900 microliters
- : *container-type-1-flat* plate type to *\_CONTAINER\_TYPES*
- : support for container names with slashes in them in *harness.py*
- : add *protocol-consolidate* Protocol method and accompanying tests
- : *ImagePlate()* class and *protocol-image-plate* Protocol method for taking images of containers
- : volume adjustment when *protocol-spread* ing
- : collapse *protocol-provision* instructions if they’re acting on the same container
- : *protocol-sangerseq* now accepts a sequencing type of “rca” or “standard” (defaults to “standard”)
- : *criteria* and *dataref* fields to *protocol-autopick*
- : *protocol-flash-freeze* Protocol method and Instruction
- : *protocol-ref* behavior when specifying the *id* of an existing container
- : type check in Container.wells
- : README.rst
- : a wild test appeared!
- : link to library documentation at [readthedocs.org](http://readthedocs.org) to README
- : autoprotocol and JSON output examples for almost everything in docs
- : improved documentation tree
- : documentation for *plate\_to\_mag\_adapter* and *plate\_from\_mag\_adapter* **subject to change in near future**
- : documentation punctuation and grammar
- : check that a well already exists in a WellGroup
- : Added folder for sublime text snippets
- : Protocol.serial\_dilute\_rowwise()
- : Protocol.thermocycle\_ramp()

- : More Python3 Compatibility
- : Additional type-checks in various functions
- : *protocol-provision* Protocol method
- : support for *choice* input type in *harness.py*
- : allow transfer from multiple sources to one destination
- : brought back recursively transferring volumes over 900 microliters
- : *container-type-1-flat* plate type to *\_CONTAINER\_TYPES*
- : support for container names with slashes in them in *harness.py*
- : add *protocol-consolidate* Protocol method and accompanying tests
- : *ImagePlate()* class and *protocol-image-plate* Protocol method for taking images of containers
- : volume adjustment when *protocol-spread* ing
- : typo in *protocol-sangerseq* instruction
- : documentation punctuation and grammar
- : check that a well already exists in a WellGroup
- : Added folder for sublime text snippets
- : *protocol-stamp* ing to or from multiple containers now requires that the source or dest variable be passed as a list of [{“container”: <container>, “quadrant”: <quadrant>}, … ]
- : *protocol-gel-separate* generates instructions taking wells and matrix type passed
- : tox for testing with multiple versions of python
- : *new\_group* keyword parameter on *protocol-transfer* and *protocol-distribute* to manually break up *Pipette()* Instructions
- : Thermocycle input type in *harness.py*
- : specify *Wells* on a container using *container.wells(1,2,3)*‘or ‘*container.wells([1,2,3])*
- : More Python3 Compatibility
- : Transferring liquid from *one\_source* actually works now
- : references to specific reagents for *protocol-dispense*
- : documentation for *plate\_to\_mag\_adapter* and *plate\_from\_mag\_adapter* **subject to change in near future**
- : *Protocol.pipette()* is now a private method *\_pipette()*
- : *protocol-sangerseq* Instruction and method
- : seal takes a “type” parameter that defaults to ultra-clear
- : more tests
- : *protocol-stamp* Protocol method for using the 96-channel liquid handler
- : Added *pipette\_tools* module containing helper methods for the extra pipetting parameters
- : Additional keyword arguments for *protocol-transfer* and *protocol-distribute* to customize pipetting
- : *protocol-oligosynthesize* Instruction
- : *protocol-autopick* Instruction
- : *protocol-spread* Instruction

- : *protocol-flow-analyze* Instruction
- : co2 parameter in *protocol-incubate*
- : 6-flat container type in *\_CONTAINER\_TYPES*
- : 96-flat-uv container type in *\_CONTAINER\_TYPES*
- : *container-quadrant* returns a WellGroup of the 96 wells representing the quadrant passed
- : *well-add-properties*
- : Well.properties is an empty hash by default
- : At least some Python3 compatibility
- : *protocol-gel-separate* generates number of instructions needed for number of wells passed
- : recursion to deal with transferring over 900uL of liquid
- : references to specific matrices and ladders in *protocol-gel-separate*
- : refactoring of type checks in *unit-unit*
- : improved documentation tree
- : melting keyword variables and changes to conditionals in Thermocycle
- : default input value and group and group+ input types in *harness.py*
- : a wild test appeared!
- : link to library documentation at [readthedocs.org](http://readthedocs.org) to README
- : autoprotocol and JSON output examples for almost everything in docs
- : warnings for *\_mul\_* and *\_div\_* scalar Unit operations
- : *protocol-dispense-full-plate*
- : *protocol-dispense* Instruction and accompanying Protocol method for using a reagent dispenser
- : aliquot++, integer, boolean input types to *harness.py*
- : properties attribute to Well, along with *well-set-properties* method
- : spelling of luminescence :(
- : *well\_type* from *\_CONTAINER\_TYPES*
- : “speed” parameter in *protocol-spin* to “acceleration”
- : README.rst
- : “one\_tip” option on *protocol-transfer*
- : volume tracking upon *protocol-transfer* and *protocol-distribute*
- : dead\_volume\_ul in *\_CONTAINER\_TYPES*
- : *wellGroup-indices* returns a list of string well indices
- : 3-clause BSD license, contributor info
- : *container-inner-wells* method to exclude edges
- : *harness.py* for parameter conversion
- : static methods *Pipette.transfers()* and *Pipette.\_transferGroup()*
- : NumPy style docstrings for most methods

- : initializing ap-py



# CHAPTER 9

---

## Credits

---

Autoprotocol-Python is currently maintained by:

- Vanessa Biggers - [polarpine](#) - [vanessa@transcriptic.com](mailto:vanessa@transcriptic.com)
- Yang Choo - [yangchoo](#) - [yang@transcriptic.com](mailto:yang@transcriptic.com)
- Jim Culver - [drjimmypants](#) - [jim@transcriptic.com](mailto:jim@transcriptic.com)
- Donald Dalton - [dbdalton2000](#) - [donald@transcriptic.com](mailto:donald@transcriptic.com)
- Varun Kanwar - [VarunKanwar](#) - [varun@transcriptic.com](mailto:varun@transcriptic.com)
- Peter Lee - [pleaderlee](#) - [peter@transcriptic.com](mailto:peter@transcriptic.com)
- Rhys Ormond - [rhysormond](#) - [rhys@transcriptic.com](mailto:rhys@transcriptic.com)

[See all Github contributors](#)

For more information about Autoprotocol and its specification, visit [autoprotocol.org](#)



# CHAPTER 10

---

## License

---

Copyright (c) 2018, Transcryptic Inc All rights reserved. Redistribution and use in source and binary forms, with or without modification, are permitted provided that the following conditions are met:

- Redistributions of source code must retain the above copyright notice, this list of conditions and the following disclaimer.
- Redistributions in binary form must reproduce the above copyright notice, this list of conditions and the following disclaimer in the documentation and/or other materials provided with the distribution.
- Neither the name of autoprotocol-python nor the names of its contributors may be used to endorse or promote products derived from this software without specific prior written permission.

THIS SOFTWARE IS PROVIDED BY THE COPYRIGHT HOLDERS AND CONTRIBUTORS “AS IS” AND ANY EXPRESS OR IMPLIED WARRANTIES, INCLUDING, BUT NOT LIMITED TO, THE IMPLIED WARRANTIES OF MERCHANTABILITY AND FITNESS FOR A PARTICULAR PURPOSE ARE DISCLAIMED. IN NO EVENT SHALL TRANSCRIPTIC BE LIABLE FOR ANY DIRECT, INDIRECT, INCIDENTAL, SPECIAL, EXEMPLARY, OR CONSEQUENTIAL DAMAGES (INCLUDING, BUT NOT LIMITED TO, PROCUREMENT OF SUBSTITUTE GOODS OR SERVICES; LOSS OF USE, DATA, OR PROFITS; OR BUSINESS INTERRUPTION) HOWEVER CAUSED AND ON ANY THEORY OF LIABILITY, WHETHER IN CONTRACT, STRICT LIABILITY, OR TORT (INCLUDING NEGLIGENCE OR OTHERWISE) ARISING IN ANY WAY OUT OF THE USE OF THIS SOFTWARE, EVEN IF ADVISED OF THE POSSIBILITY OF SUCH DAMAGE.

Use the sidebar to navigate specific module documentation.

Autoprotocol is a standard way to express experiments in life science. The [autoprotocol-python](#) repository contains a python library for generating Autoprotocol.



# CHAPTER 11

---

## Installation

---

```
$ git clone https://github.com/autoproto/autoproto-python  
$ cd autoproto-python  
$ python setup.py install
```

or, alternatively:

```
$ pip install autoproto
```



# CHAPTER 12

## Building a Protocol

A basic protocol object has empty “refs” and “instructions” stanzas. Various helper methods in the Protocol class are then used to append instructions and refs to the object such as in the simple protocol below:

```
import json
from autoprotocol.protocol import Protocol

#instantiate new Protocol object
p = Protocol()

#append refs (containers) to Protocol object
bacteria = p.ref("bacteria", cont_type="96-pcr", storage="cold_4")
medium = p.ref("medium", cont_type="micro-1.5", storage="cold_4")
reaction_plate = p.ref("reaction_plate", cont_type="96-flat", storage="warm_37")

#distribute medium from 1.5mL tube to reaction wells
p.distribute(medium.well(0).set_volume("1000:microliter"), reaction_plate.wells_
    ↪from(0,4), "190:microliter")
#transfer bacteria from source wells to reaction wells
p.transfer(bacteria.wells_from(0,4), reaction_plate.wells_from(0,4),
    ["10:microliter", "20:microliter", "30:microliter", "40:microliter"])
#incubate bacteria at 37 degrees for 5 hours
p.incubate(reaction_plate, "warm_37", "5:hour")
#read absorbance of the first four wells on the reaction plate at 600 nanometers
p.absorbance(reaction_plate, reaction_plate.wells_from(0,4).indices(), "600:nanometer
    ↪",
    "OD600_reading_01092014")

print json.dumps(p.as_dict(), indent=2)
```

The script above produces the following autoprotocol:

```
{
  "refs": {
    "medium": {
```

(continues on next page)

(continued from previous page)

```

    "new": "micro-1.5",
    "store": {
        "where": "cold_4"
    }
},
"bacteria": {
    "new": "96-pcr",
    "store": {
        "where": "cold_4"
    }
},
"reaction_plate": {
    "new": "96-flat",
    "store": {
        "where": "warm_37"
    }
}
},
"instructions": [
{
    "groups": [
    {
        "distribute": {
            "to": [
            {
                "volume": "190.0:microliter",
                "well": "reaction_plate/0"
            },
            {
                "volume": "190.0:microliter",
                "well": "reaction_plate/1"
            },
            {
                "volume": "190.0:microliter",
                "well": "reaction_plate/2"
            },
            {
                "volume": "190.0:microliter",
                "well": "reaction_plate/3"
            }
        ],
        "from": "medium/0"
    }
},
{
        "transfer": [
        {
            "volume": "10.0:microliter",
            "to": "reaction_plate/0",
            "from": "bacteria/0"
        }
    ]
},
{
        "transfer": [
        {
            "volume": "20.0:microliter",
            "to": "reaction_plate/0"
        }
    ]
}
]
}

```

(continues on next page)

(continued from previous page)

```
        "to": "reaction_plate/1",
        "from": "bacteria/0"
    }
]
},
{
    "transfer": [
        {
            "volume": "30.0:microliter",
            "to": "reaction_plate/2",
            "from": "bacteria/0"
        }
    ]
},
{
    "transfer": [
        {
            "volume": "40.0:microliter",
            "to": "reaction_plate/3",
            "from": "bacteria/0"
        }
    ]
}
],
"op": "pipette"
},
{
    "duration": "5:hour",
    "where": "warm_37",
    "object": "reaction_plate",
    "shaking": false,
    "op": "incubate"
},
{
    "dataref": "OD600_reading_01092014",
    "object": "reaction_plate",
    "wells": [
        "A1",
        "A2",
        "A3",
        "A4"
    ],
    "num_flashes": 25,
    "wavelength": "600:nanometer",
    "op": "absorbance"
}
]
```



# CHAPTER 13

---

## Contributing

---

The easiest way to contribute is to fork this repository and submit a pull request. You can also write an email to us if you want to discuss ideas or bugs.

- Autoprotocol Curators: [autoprotocol-curators@transcriptic.com](mailto:autoprotocol-curators@transcriptic.com)

autoprotocol-python is BSD licensed (see LICENSE). Before we can accept your pull request, we require that you sign a CLA (Contributor License Agreement) allowing us to distribute your work under the BSD license. Email one of the authors listed above for more details.



# CHAPTER 14

---

Search the Docs

---

- genindex
- search

**copyright** 2018 by The Autoprotocol Development Team, see AUTHORS for more details.

**license** BSD, see LICENSE for more details



---

## Python Module Index

---

### a

autoprotocol.builders, 107  
autoprotocol.instruction, 61  
autoprotocol.liquid\_handle.liquid\_class,  
    124  
autoprotocol.liquid\_handle.liquid\_handle\_method,  
    121  
autoprotocol.liquid\_handle.mix, 137  
autoprotocol.liquid\_handle.tip\_type, 139  
autoprotocol.liquid\_handle.transfer, 128  
autoprotocol.protocol, 1



### Symbols

\_\_add\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127  
\_\_add\_\_() (autoprotocol.container.WellGroup method), 79  
\_\_contains\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127  
\_\_delattr\_\_ (autoprotocol.builders.DispenseBuilders attribute), 107  
\_\_delattr\_\_ (autoprotocol.builders.InstructionBuilders attribute), 109  
\_\_delattr\_\_ (autoprotocol.builders.LiquidHandleBuilders attribute), 110  
\_\_delattr\_\_ (autoprotocol.col.builders.SpectrophotometryBuilders attribute), 114  
\_\_delattr\_\_ (autoprotocol.builders.ThermocycleBuilders attribute), 118  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.liquid\_class.LiquidClass attribute), 125  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 126  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod attribute), 123  
\_\_delattr\_\_ (autoprotocol.liquid\_handle.mix.Mix attribute), 137  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.transfer.DryWellTransfer attribute), 128  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.transfer.PreMixBlowoutTransfer attribute), 131  
\_\_delattr\_\_ (autoprotocol.col.liquid\_handle.transfer.Transfer attribute), 134  
\_\_eq\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127  
\_\_format\_\_() (autoprotocol.builders.DispenseBuilders method), 107  
\_\_format\_\_() (autoprotocol.builders.InstructionBuilders method), 109  
\_\_format\_\_() (autoprotocol.col.builders.LiquidHandleBuilders method), 110  
\_\_format\_\_() (autoprotocol.col.builders.SpectrophotometryBuilders method), 114  
\_\_format\_\_() (autoprotocol.col.builders.ThermocycleBuilders method), 118  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.liquid\_class.LiquidClass method), 125  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.liquid\_class.VolumeCalibrationBin method), 126  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.liquid\_class.VolumeCalibrationBin method), 127  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod method), 123  
\_\_format\_\_() (autoprotocol.liquid\_handle.mix.Mix method), 137  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.transfer.DryWellTransfer method), 128  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.transfer.PreMixBlowoutTransfer method), 131  
\_\_format\_\_() (autoprotocol.col.liquid\_handle.transfer.Transfer method), 134

—ge\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 110  
    attribute), 127

—getattribute\_\_ (autoprotocol.builders.DispenseBuilders attribute), 107

—getattribute\_\_ (autoprotocol.builders.InstructionBuilders attribute), 109

—getattribute\_\_ (autoprotocol.builders.LiquidHandleBuilders attribute), 110

—getattribute\_\_ (autoprotocol.builders.SpectrophotometryBuilders attribute), 114

—getattribute\_\_ (autoprotocol.builders.ThermocycleBuilders attribute), 118

—getattribute\_\_ (autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 126

—getattribute\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibration attribute), 126

—getattribute\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

—getattribute\_\_ (autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandle attribute), 123

—getattribute\_\_ (autoprotocol.liquid\_handle.mix.Mix attribute), 137

—getattribute\_\_ (autoprotocol.liquid\_handle.transfer.DryWellTransfer attribute), 128

—getattribute\_\_ (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer attribute), 131

—getattribute\_\_ (autoprotocol.liquid\_handle.transfer.Transfer attribute), 134

—getitem\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

—getitem\_\_( autoprotocol.container.WellGroup method), 79

—getslice\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

—gt\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

—hash\_\_ (autoprotocol.builders.DispenseBuilders attribute), 107

—hash\_\_ (autoprotocol.builders.InstructionBuilders attribute), 109

—hash\_\_ (autoprotocol.builders.LiquidHandleBuilders

    \_\_hash\_\_ (autoprotocol.builders.SpectrophotometryBuilders attribute), 114

    \_\_hash\_\_ (autoprotocol.builders.ThermocycleBuilders attribute), 118

    \_\_hash\_\_ (autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 126

    \_\_hash\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibration attribute), 126

    \_\_hash\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

    \_\_hash\_\_ (autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandle attribute), 123

    \_\_hash\_\_ (autoprotocol.liquid\_handle.mix.Mix attribute), 137

    \_\_hash\_\_ (autoprotocol.liquid\_handle.transfer.DryWellTransfer attribute), 128

    \_\_hash\_\_ (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer attribute), 131

    \_\_hash\_\_ (autoprotocol.liquid\_handle.transfer.Transfer attribute), 134

    \_\_len\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

    \_\_len\_\_( autoprotocol.container.WellGroup method), 79

    \_\_lt\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

    \_\_mul\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

    \_\_ne\_\_ (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 127

    \_\_reduce\_\_( autoprotocol.builders.DispenseBuilders method), 107

    \_\_reduce\_\_( autoprotocol.builders.InstructionBuilders method), 109

    \_\_reduce\_\_( autoprotocol.builders.LiquidHandleBuilders method), 110

    \_\_reduce\_\_( autoprotocol.builders.SpectrophotometryBuilders method), 114

    \_\_reduce\_\_( autoprotocol.builders.ThermocycleBuilders method), 118

    \_\_reduce\_\_( autoprotocol.liquid\_handle.liquid\_class.LiquidClass method), 126

    \_\_reduce\_\_( autoprotocol.liquid\_handle.liquid\_class.VolumeCalibration method), 126

```

__reduce__() (autoproto- __repr__ (autoprotocol.builders.DispenseBuilders at-
  col.liquid_handle.liquid_class.VolumeCalibrationBin tribute), 108
  method), 127 __repr__ (autoprotocol.builders.InstructionBuilders at-
  __reduce__() (autoproto- tribute), 109
  col.liquid_handle.liquid_handle_method.LiquidHandleMethod(autoprotocol.builders.LiquidHandleBuilders
  method), 123 attribute), 110
  __reduce__() (autoprotocol.liquid_handle.mix.Mix __repr__ (autoprotocol.builders.SpectrophotometryBuilders
  method), 137 attribute), 115
  __reduce__() (autoprotocol.liquid_handle.transfer.DryWellTransfer __repr__ (autoprotocol.builders.ThermocycleBuilders at-
  method), 128 tribute), 118
  __reduce__() (autoprotocol.liquid_handle.transfer.PreMixBlowoutTransfer __repr__ (autoprotocol.liquid_handle.liquid_class.LiquidClass
  method), 131 attribute), 126
  __reduce__() (autoprotocol.liquid_handle.transfer.Transfer __repr__ (autoprotocol.liquid_handle.liquid_handle_method.LiquidHandle
  method), 134 attribute), 123
  __reduce_ex__() (autoprotocol.builders.DispenseBuilders __repr__ (autoprotocol.liquid_handle.mix.Mix attribute),
  method), 108 137
  __reduce_ex__() (autoprotocol.builders.InstructionBuilders __repr__ (autoprotocol.liquid_handle.transfer.DryWellTransfer
  method), 109 attribute), 128
  __reduce_ex__() (autoprotocol.builders.LiquidHandleBuilders __repr__ (autoprotocol.liquid_handle.transfer.PreMixBlowoutTransfer
  method), 110 attribute), 131
  __reduce_ex__() (autoprotocol.builders.SpectrophotometryBuilders __repr__ (autoprotocol.liquid_handle.transfer.Transfer
  method), 114 attribute), 134
  __reduce_ex__() (autoprotocol.builders.ThermocycleBuilders __repr__ () (autoprotocol.container.Container method), 75
  method), 118 __repr__ () (autoprotocol.container.Well method), 78
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_class.LiquidClass __repr__ () (autoprotocol.container.WellGroup method),
  method), 126 79
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_class.VolumeCalibration __rmul__ (autoprotocol.liquid_handle.liquid_class.VolumeCalibrationBin
  method), 126 attribute), 127
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_class.VolumeCalibrationBin __setattr__ (autoprotocol.builders.DispenseBuilders at-
  method), 127 __attribute), 108
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_handle_method.LiquidHandleMethod __setattr__ (autoprotocol.builders.InstructionBuilders at-
  method), 123 __attribute), 109
  __reduce_ex__() (autoprotocol.liquid_handle.mix.Mix __setattr__ (autoprotocol.builders.LiquidHandleBuilders
  method), 137 attribute), 110
  __reduce_ex__() (autoprotocol.liquid_handle.transfer.DryWellTransfer __setattr__ (autoprotocol.builders.SpectrophotometryBuilders
  method), 128 attribute), 115
  __reduce_ex__() (autoprotocol.liquid_handle.transfer.PreMixBlowoutTransfer __setattr__ (autoprotocol.builders.ThermocycleBuilders
  method), 131 __attribute), 118
  __reduce_ex__() (autoprotocol.liquid_handle.transfer.Transfer __setattr__ (autoprotocol.liquid_handle.liquid_class.LiquidClass
  method), 134 __attribute), 126
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_handle_method.LiquidHandle __setattr__ (autoprotocol.liquid_handle.liquid_class.VolumeCalibration
  method), 123 __attribute), 127
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_class.VolumeCalibration __setattr__ (autoprotocol.liquid_handle.liquid_class.VolumeCalibrationBin
  attribute), 126 __attribute), 127
  __reduce_ex__() (autoprotocol.liquid_handle.liquid_class.VolumeCalibrationBin __setattr__ (autoprotocol.liquid_handle.liquid_handle_method.LiquidHandle
  attribute), 127 __attribute), 123
  __reduce_ex__() (autoprotocol.liquid_handle.mix.Mix __setattr__ (autoprotocol.liquid_handle.mix.Mix attribute),
  method), 137 137
  __reduce_ex__() (autoprotocol.liquid_handle.transfer.DryWellTransfer __setattr__ (autoprotocol.liquid_handle.transfer.DryWellTransfer
  method), 128 attribute), 128
  __reduce_ex__() (autoprotocol.liquid_handle.transfer.Transfer __setattr__ (autoprotocol.liquid_handle.transfer.PreMixBlowoutTransfer
  method), 134 attribute), 131

```

\_\_setattr\_\_(autoprotocol.liquid\_handle.transfer.Transfer attribute), 134  
\_\_setitem\_\_(autoprotocol.container.WellGroup method), 80  
\_\_sizeof\_\_(autoprotocol.builders.DispenseBuilders method), 108  
\_\_sizeof\_\_(autoprotocol.builders.InstructionBuilders method), 109  
\_\_sizeof\_\_(autoprotocol.builders.LiquidHandleBuilders method), 110  
\_\_sizeof\_\_(autoprotocol.builders.SpectrophotometryBuilders method), 115  
\_\_sizeof\_\_(autoprotocol.builders.ThermocycleBuilders method), 118  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.liquid\_class.LiquidClass method), 126  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.liquid\_class.VolumeCalibration\_shape method), 126  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin method), 127  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod), 123  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.mix.Mix method), 137  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.transfer.DryWellTransfer method), 128  
\_\_sizeof\_\_(autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer method), 131  
\_\_str\_\_(autoprotocol.builders.DispenseBuilders attribute), 108  
\_\_str\_\_(autoprotocol.builders.InstructionBuilders attribute), 109  
\_\_str\_\_(autoprotocol.builders.LiquidHandleBuilders attribute), 110  
\_\_str\_\_(autoprotocol.builders.SpectrophotometryBuilders attribute), 115  
\_\_str\_\_(autoprotocol.builders.ThermocycleBuilders attribute), 118  
\_\_str\_\_(autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 126  
\_\_str\_\_(autoprotocol.liquid\_handle.liquid\_class.VolumeCalibration attribute), 126  
\_\_str\_\_(autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 126  
attribute), 128  
\_\_str\_\_(autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod attribute), 123  
\_\_str\_\_(autoprotocol.liquid\_handle.mix.Mix attribute), 137  
\_\_str\_\_(autoprotocol.liquid\_handle.transfer.DryWellTransfer attribute), 129  
\_\_str\_\_(autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer attribute), 131  
\_\_str\_\_(autoprotocol.liquid\_handle.transfer.Transfer attribute), 134  
\_\_str\_\_(autoprotocol.unit.Unit method), 141  
\_destination\_liquid (autoprotocol.liquid\_handle.transfer.Transfer attribute), 134  
\_liquid (autoprotocol.liquid\_handle.mix.Mix attribute), 137  
\_safe\_volume\_multiplier (autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 124  
\_source\_liquid (autoprotocol.liquid\_handle.transfer.Transfer attribute), 134  
\_transports (autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod attribute), 121

## A

Absorbance (class in autoprotocol.instruction), 61  
absorbance() (autoprotocol.protocol.Protocol method), 2  
absorbance\_mode\_params() (autoprotocol.builders.SpectrophotometryBuilders static method), 115  
acoustic\_transfer() (autoprotocol.protocol.Protocol method), 4  
AcousticTransfer (class in autoprotocol.instruction), 62  
add\_properties() (autoprotocol.container.Well method), 78  
add\_properties() (autoprotocol.container.WellGroup method), 80  
add\_time\_constraint() (autoprotocol.protocol.Protocol method), 5  
all\_wells() (autoprotocol.container.Container method), 75  
append() (autoprotocol.container.WellGroup method), 80  
as\_dict() (autoprotocol.protocol.Protocol method), 8  
aspire\_flowrate\_calibration\_curve (autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 124  
Autopick (class in autoprotocol.instruction), 62  
autopick() (autoprotocol.protocol.Protocol method), 9  
autoprotocol.builders (module), 107  
autoprotocol.instruction (module), 61  
autoprotocol.liquid\_handle.liquid\_class (module), 124

|  |   |
|--|---|
| autoprotocol.liquid_handle.liquid_handle_method (module), 121  | default_aspirate_z() (autoproto-col.liquid_handle.transfer.PreMixBlowoutTransfer method), 131                     |
| autoprotocol.liquid_handle.mix (module), 137   | default_aspirate_z() (autoproto-col.liquid_handle.transfer.Transfer method), 134                                  |
| autoprotocol.liquid_handle.tip_type (module), 139  | default_blowout() (autoproto-col.liquid_handle.liquid_handle_method.LiquidHandleMethod method), 123               |
| autoprotocol.liquid_handle.transfer (module), 128  | default_blowout() (autoproto-col.liquid_handle.transfer.DryWellTransfer method), 129                              |
| autoprotocol.protocol (module), 1  | default_blowout() (autoproto-col.liquid_handle.transfer.PreMixBlowoutTransfer method), 131                        |
| available_volume() (autoproto-col.container.Well method), 78   | default_blowout() (autoproto-col.liquid_handle.transfer.Transfer method), 135                                     |
| <b>B</b>   |   |
| batch_containers() (autoproto-col.protocol.Protocol method), 9   | default_dispense_z() (autoproto-col.liquid_handle.transfer.DryWellTransfer method), 129                           |
| binned_calibration_for_volume() (autoproto-col.liquid_handle.liquid_class.VolumeCalibration method), 126 | default_dispense_z() (autoproto-col.liquid_handle.transfer.PreMixBlowoutTransfer method), 132                     |
| blowout() (autoproto-builders.LiquidHandleBuilders method), 110  | default_dispense_z() (autoproto-col.liquid_handle.transfer.Transfer method), 135                                  |
| <b>C</b>   |   |
| calibrate_volume() (autoproto-col.liquid_handle.liquid_class.VolumeCalibrationBinh method), 128          | default_lld_position_z() (autoproto-col.liquid_handle.liquid_handle_method.LiquidHandleMethod static method), 123 |
| ceil() (autoproto-unit.Unit method), 142   | default_lld_position_z() (autoproto-col.liquid_handle.transfer.PreMixBlowoutTransfer static method), 132          |
| column() (autoproto-builders.DispenseBuilders static method), 108  | default_lld_position_z() (autoproto-col.liquid_handle.transfer.Transfer method), 135                              |
| columns() (autoproto-builders.DispenseBuilders method), 108  | default_lld_position_z() (autoproto-col.liquid_handle.liquid_handle_method.LiquidHandleMethod static method), 123 |
| Container (class in autoprotocol.container), 75  | default_lld_position_z() (autoproto-col.liquid_handle.transfer.Transfer static method), 138                       |
| container_type() (autoproto.protocol.Protocol method), 10  | default_lld_position_z() (autoproto-col.liquid_handle.transfer.DryWellTransfer static method), 129                |
| ContainerType (class in autoprotocol.container_type), 83   | default_lld_position_z() (autoproto-col.liquid_handle.transfer.PreMixBlowoutTransfer static method), 132          |
| convert_well_map_to_dye_map() (autoproto-col.instruction.Thermocycle static method), 73                  | default_lld_position_z() (autoproto-col.liquid_handle.transfer.Transfer static method), 135                       |
| count() (autoproto.liquid_handle.liquid_class.VolumeCalibrationBinh method), 128                         | default_mix_after() (autoproto-col.liquid_handle.transfer.DryWellTransfer method), 129                            |
| count_cells() (autoproto.protocol.Protocol method), 11   | default_mix_after() (autoproto-col.liquid_handle.transfer.PreMixBlowoutTransfer method), 132                      |
| CountCells (class in autoprotocol.instruction), 62   | default_mix_after() (autoproto-col.liquid_handle.transfer.Transfer method), 135                                   |
| Cover (class in autoprotocol.instruction), 62  | default_mix_before() (autoproto-col.liquid_handle.transfer.Transfer method), 135                                  |
| cover() (autoproto.protocol.Protocol method), 12   |   |
| <b>D</b>   |   |
| decompose() (autoproto-col.container.Container method), 76   |   |
| decompose() (autoproto-col.container_type.ContainerType method), 84                                      |   |
| DEEP24 (in module autoprotocol.container_type), 94   |   |
| DEEP96 (in module autoprotocol.container_type), 92   |   |
| DEEP96KF (in module autoprotocol.container_type), 93   |   |
| default_aspirate_z() (autoproto-col.liquid_handle.transfer.DryWellTransfer method), 129                  |   |

col.liquid\_handle.transfer.DryWellTransfer  
method), 130

default\_mix\_before() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
method), 132

default\_mix\_before() (autoprotocol.liquid\_handle.transfer.Transfer  
method), 135

default\_position\_z() (autoprotocol.liquid\_handle.mix.Mix method), 138

default\_pre\_mix\_blowout() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
method), 133

default\_prime() (autoprotocol.liquid\_handle.transfer.DryWellTransfer  
method), 130

default\_prime() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
method), 133

default\_prime() (autoprotocol.liquid\_handle.transfer.Transfer  
method), 136

default\_repetitions() (autoprotocol.liquid\_handle.mix.Mix method), 138

default\_tracked\_position\_z() (autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod  
static method), 123

default\_tracked\_position\_z() (autoprotocol.liquid\_handle.mix.Mix static method), 138

default\_tracked\_position\_z() (autoprotocol.liquid\_handle.transfer.DryWellTransfer  
static method), 130

default\_tracked\_position\_z() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
static method), 133

default\_tracked\_position\_z() (autoprotocol.liquid\_handle.transfer.Transfer  
static method), 136

default\_transit() (autoprotocol.liquid\_handle.transfer.DryWellTransfer  
method), 130

default\_transit() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
method), 133

default\_transit() (autoprotocol.liquid\_handle.transfer.Transfer  
method), 136

default\_well\_bottom\_position\_z() (autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod  
static method), 123

default\_well\_bottom\_position\_z() (autoprotocol.liquid\_handle.mix.Mix static method), 138

default\_well\_bottom\_position\_z() (autoprotocol.liquid\_handle.transfer.DryWellTransfer  
static method), 130

default\_well\_bottom\_position\_z() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
static method), 133

default\_well\_bottom\_position\_z() (autoprotocol.liquid\_handle.transfer.Transfer  
static method), 136

default\_well\_top\_position\_z() (autoprotocol.liquid\_handle.liquid\_handle\_method.LiquidHandleMethod  
static method), 124

default\_well\_top\_position\_z() (autoprotocol.liquid\_handle.mix.Mix static method), 138

default\_well\_top\_position\_z() (autoprotocol.liquid\_handle.transfer.DryWellTransfer  
static method), 131

default\_well\_top\_position\_z() (autoprotocol.liquid\_handle.transfer.PreMixBlowoutTransfer  
static method), 133

default\_well\_top\_position\_z() (autoprotocol.liquid\_handle.transfer.Transfer  
static method), 136

discard() (autoprotocol.container.Container method), 76

DispenseMethod (class in autoprotocol.instruction), 62

dispense() (autoprotocol.protocol.Protocol method), 13

dispense\_flowrate\_calibration\_curve (autoprotocol.liquid\_handle.liquid\_class.LiquidClass  
attribute), 124

dispense\_full\_plate() (autoprotocol.protocol.Protocol  
method), 15

DispenseBuilders (class in autoprotocol.builders), 107

DryWellTransfer (class in autoprotocol.liquid\_handle.transfer), 128

**E**

ECHO384 (in module autoprotocol.container\_type), 87

ECHO384LDV (in module autoprotocol.container\_type), 104

ECHO384LDVPLUS (in module autoprotocol.col.container\_type), 105

extend() (autoprotocol.container.WellGroup method), 80

**F**

find\_invalid\_dyes() (autoprotocol.col.instruction.Thermocycle static method), 73

flash\_freeze() (autoprotocol.protocol.Protocol method), 17

FlashFreeze (class in autoprotocol.instruction), 63

FLAT1 (in module autoprotocol.container\_type), 97

FLAT384 (in module autoprotocol.container\_type), 85

FLAT384CLEAR (in module col.container\_type), 89  
 FLAT384WHITELV (in module col.container\_type), 87  
 FLAT384WHITETC (in module col.container\_type), 88  
 FLAT6 (in module autoprotocol.container\_type), 97  
 FLAT6TC (in module autoprotocol.container\_type), 98  
 FLAT96 (in module autoprotocol.container\_type), 90  
 FLAT96CLEARTC (in module autoprotocol.container\_type), 101  
 FLAT96UV (in module autoprotocol.container\_type), 90  
 floor() (autoprotocol.unit.Unit method), 142  
 flow\_analyze() (autoprotocol.protocol.Protocol method), 18  
 FlowAnalyze (class in autoprotocol.instruction), 63  
 flowrate() (autoprotocol.builders.LiquidHandleBuilders static method), 111  
 Fluorescence (class in autoprotocol.instruction), 65  
 fluorescence() (autoprotocol.protocol.Protocol method), 21  
 fluorescence\_mode\_params() (autoprotocol.builders.SpectrophotometryBuilders method), 115

**G**

gel\_purify() (autoprotocol.protocol.Protocol method), 23  
 gel\_separate() (autoprotocol.protocol.Protocol method), 25  
 GelPurify (class in autoprotocol.instruction), 66  
 GelSeparate (class in autoprotocol.instruction), 67  
 get\_instruction\_index() (autoprotocol.protocol.Protocol method), 26  
 group() (autoprotocol.builders.SpectrophotometryBuilders method), 116  
 group() (autoprotocol.builders.ThermocycleBuilders method), 118  
 groups() (autoprotocol.builders.SpectrophotometryBuilders method), 116

**H**

humanize() (autoprotocol.container.Container method), 76  
 humanize() (autoprotocol.container.Well method), 78  
 humanize() (autoprotocol.container\_type.ContainerType method), 84

**I**

IlluminaSeq (class in autoprotocol.instruction), 67  
 illuminaseq() (autoprotocol.protocol.Protocol method), 27  
 image\_plate() (autoprotocol.protocol.Protocol method), 29  
 ImagePlate (class in autoprotocol.instruction), 67

autoproto-  
 incubate (class in autoprotocol.instruction), 68  
 incubate() (autoprotocol.protocol.Protocol method), 30  
 incubate\_params() (in module autoprotocol.util), 142  
 index() (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin method), 128  
 indices() (autoprotocol.container.WellGroup method), 80  
 inner\_wells() (autoprotocol.container.Container method), 76  
 insert() (autoprotocol.container.WellGroup method), 80  
 Instruction (class in autoprotocol.instruction), 68  
 instruction\_mode\_params() (autoprotocol.builders.LiquidHandleBuilders static method), 111  
 InstructionBuilders (class in autoprotocol.builders), 109  
 intercept (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 128  
 is\_covered() (autoprotocol.container.Container method), 76  
 is\_sealed() (autoprotocol.container.Container method), 76

**J**

json() (autoprotocol.instruction.Instruction method), 68

**L**

LiquidClass (class in autoprotocol.liquid\_handle.liquid\_class), 124  
 LiquidHandle (class in autoprotocol.instruction), 68  
 LiquidHandleBuilders (class in autoprotocol.builders), 110  
 LiquidHandleMethod (class in autoprotocol.liquid\_handle.liquid\_handle\_method), 121  
 location() (autoprotocol.builders.LiquidHandleBuilders method), 111  
 Luminescence (class in autoprotocol.instruction), 68  
 luminescence() (autoprotocol.protocol.Protocol method), 31  
 luminescence\_mode\_params() (autoprotocol.builders.SpectrophotometryBuilders static method), 116

**M**

mag\_collect() (autoprotocol.protocol.Protocol method), 32  
 mag\_dry() (autoprotocol.protocol.Protocol method), 33  
 mag\_incubate() (autoprotocol.protocol.Protocol method), 34  
 mag\_mix() (autoprotocol.protocol.Protocol method), 35  
 mag\_release() (autoprotocol.protocol.Protocol method), 36  
 MagneticTransfer (class in autoprotocol.instruction), 69  
 make\_band\_param() (in module autoprotocol.util), 143

make\_gel\_extract\_params() (in module autoprotocol.util), 143  
Manifest (class in autoprotocol.harness), 145  
measure\_concentration() (autoprotocol.protocol.Protocol method), 37  
measure\_mass() (autoprotocol.protocol.Protocol method), 38  
measure\_volume() (autoprotocol.protocol.Protocol method), 39  
MeasureConcentration (class in autoprotocol.instruction), 69  
MeasureMass (class in autoprotocol.instruction), 69  
MeasureVolume (class in autoprotocol.instruction), 70  
MICRO15 (in module autoprotocol.container\_type), 96  
MICRO2 (in module autoprotocol.container\_type), 95  
Mix (class in autoprotocol.liquid\_handle.mix), 137  
mix() (autoprotocol.builders.LiquidHandleBuilders method), 111  
mix() (autoprotocol.protocol.Protocol method), 40  
mode\_params() (autoprotocol.builders.LiquidHandleBuilders method), 112  
move\_rate() (autoprotocol.builders.LiquidHandleBuilders static method), 112

## N

name (autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 124  
nozzle\_position() (autoprotocol.builders.DispenseBuilders static method), 108

## O

Oligosynthesize (class in autoprotocol.instruction), 70  
oligosynthesize() (autoprotocol.protocol.Protocol method), 41

## P

PCR384 (in module autoprotocol.container\_type), 86  
PCR96 (in module autoprotocol.container\_type), 91  
pop() (autoprotocol.container.WellGroup method), 80  
position\_xy() (autoprotocol.builders.LiquidHandleBuilders method), 112  
position\_z() (autoprotocol.builders.LiquidHandleBuilders method), 113  
PreMixBlowoutTransfer (class in autoprotocol.liquid\_handle.transfer), 131  
Protocol (class in autoprotocol.protocol), 1  
Provision (class in autoprotocol.instruction), 70  
provision() (autoprotocol.protocol.Protocol method), 42

## Q

quadrant() (autoprotocol.container.Container method), 76

## R

Ref (class in autoprotocol.protocol), 60  
ref() (autoprotocol.protocol.Protocol method), 43  
RESMW12HP (in module autoprotocol.container\_type), 100  
RESMW8HP (in module autoprotocol.container\_type), 100  
RESSW384LP (in module autoprotocol.container\_type), 104  
RESSW96HP (in module autoprotocol.container\_type), 99  
robotize() (autoprotocol.container.Container method), 77  
robotize() (autoprotocol.container\_type.ContainerType method), 84  
round() (autoprotocol.unit.Unit method), 142  
ROUND384CLEAR (in module autoprotocol.container\_type), 103  
row\_count() (autoprotocol.container\_type.ContainerType method), 85  
run() (in module autoprotocol.harness), 143

## S

SangerSeq (class in autoprotocol.instruction), 70  
sangerseq() (autoprotocol.protocol.Protocol method), 44  
Seal (class in autoprotocol.instruction), 71  
seal() (autoprotocol.protocol.Protocol method), 45  
seal\_on\_store() (in module autoprotocol.harness), 144  
set\_group\_name() (autoprotocol.container.WellGroup method), 81  
set\_name() (autoprotocol.container.Well method), 78  
set\_properties() (autoprotocol.container.Well method), 79  
set\_properties() (autoprotocol.container.WellGroup method), 81  
set\_storage() (autoprotocol.container.Container method), 77  
set\_volume() (autoprotocol.container.Well method), 79  
set\_volume() (autoprotocol.container.WellGroup method), 81  
shake\_after() (autoprotocol.builders.DispenseBuilders method), 108  
shake\_before() (autoprotocol.builders.SpectrophotometryBuilders method), 116  
shake\_mode\_params() (autoprotocol.builders.SpectrophotometryBuilders method), 117  
shape() (autoprotocol.builders.DispenseBuilders method), 109  
shape() (autoprotocol.builders.InstructionBuilders method), 109

shape() (autoprotocol.builders.LiquidHandleBuilders method), 113  
 shape() (autoprotocol.builders.SpectrophotometryBuilders method), 117  
 shape() (autoprotocol.builders.ThermocycleBuilders method), 118  
 slope (autoprotocol.liquid\_handle.liquid\_class.VolumeCalibrationBin attribute), 128  
 Spectrophotometry (class in autoprotocol.instruction), 71  
 spectrophotometry() (autoprotocol.protocol.Protocol method), 46  
 SpectrophotometryBuilders (class in autoprotocol.builders), 114  
 Spin (class in autoprotocol.instruction), 71  
 spin() (autoprotocol.protocol.Protocol method), 50  
 spread() (autoprotocol.protocol.Protocol method), 51  
 step() (autoprotocol.builders.ThermocycleBuilders static method), 119  
 store() (autoprotocol.protocol.Protocol method), 51

## T

Thermocycle (class in autoprotocol.instruction), 72  
 thermocycle() (autoprotocol.protocol.Protocol method), 51  
 ThermocycleBuilders (class in autoprotocol.builders), 118  
 Transfer (class in autoprotocol.liquid\_handle.transfer), 134  
 transfer() (autoprotocol.protocol.Protocol method), 56  
 transport() (autoprotocol.builders.LiquidHandleBuilders method), 114  
 tube() (autoprotocol.container.Container method), 77

## U

Uncover (class in autoprotocol.instruction), 73  
 uncover() (autoprotocol.protocol.Protocol method), 58  
 Unit (class in autoprotocol.unit), 141  
 Unseal (class in autoprotocol.instruction), 73  
 unseal() (autoprotocol.protocol.Protocol method), 59

## V

V384CLEAR (in module autoprotocol.container\_type), 102  
 V96KF (in module autoprotocol.container\_type), 93  
 volume\_calibration\_curve (autoprotocol.liquid\_handle.liquid\_class.LiquidClass attribute), 124  
 VolumeCalibration (class in autoprotocol.liquid\_handle.liquid\_class), 126  
 VolumeCalibrationBin (class in autoprotocol.liquid\_handle.liquid\_class), 126

## W

wavelength\_selection() (autoproto-

col.builders.SpectrophotometryBuilders static method), 117  
 Well (class in autoprotocol.container), 78  
 well() (autoprotocol.container.Container method), 77  
 WellGroup (class in autoprotocol.container), 79  
 wells() (autoprotocol.container.Container method), 77  
 wells\_from() (autoprotocol.container.Container method), 77  
 wells\_with() (autoprotocol.container.WellGroup method), 81