

# Package: metabolomicsWorkbenchR (via r-universe)

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**Type** Package

**Title** Metabolomics Workbench in R

**Version** 1.22.0

**Description** This package provides functions for interfacing with the Metabolomics Workbench RESTful API. Study, compound, protein and gene information can be searched for using the API. Methods to obtain study data in common Bioconductor formats such as SummarizedExperiment and MultiAssayExperiment are also included.

**License** GPL-3

**Encoding** UTF-8

**RoxygenNote** 7.3.1

**Collate** 'parse\_fcns.R' 'generics.R' 'class\_def.R' 'constants.R'  
'helper\_fcns.R' 'metabolomicsWorkbenchR.R'

**Imports** data.table, httr, jsonlite, methods, MultiAssayExperiment, struct, SummarizedExperiment, utils

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**VignetteBuilder** knitr

**biocViews** Software, Metabolomics

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\$,mw_base-method	<i>Get slot value from mw_base objects</i>
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## Description

Gets the value of a slot from mw\_base objects, provided they are not listed as 'private'.

## Usage

```
## S4 method for signature 'mw_base'
x$name
```

## Arguments

x	An object derived from mw_base.
name	The name of the slot to access.

## Value

The assigned to the slot.

## Examples

```
# an object derived from mw_base object
C = context$study
# access the name slot
C$name
```

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check_pattern	<i>Check input against acceptable input pattern</i>
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**Description**

Checks an input\_value against a regex pattern to determine if the input\_value is valid. This method is used internally and not for intended for general use.

**Usage**

```
check_pattern(I, input_value, match)

## S4 method for signature 'mw_input_item'
check_pattern(I, input_value, match)
```

**Arguments**

I	An mw_input_item.
input_value	The value for the input item (character).
match	The type of match. One of "exact" or "partial".

**Value**

TRUE if input matches the pattern, or throws an error.

**Examples**

```
check_pattern(input_item$study_id, 'ST000001', 'exact')
```

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check_puts	<i>Check inputs/outputs match</i>
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**Description**

Checks that the provided inputs and output are compatible. This method is used internally and not for intended for general use.

**Usage**

```
check_puts(input_item, output_item)

## S4 method for signature 'mw_input_item'
check_puts(input_item, output_item)
```

**Arguments**

input\_item      An mw\_input\_item.  
output\_item     An mw\_output\_item.

**Value**

TRUE if the items are compatible or throws an error if not.

**Examples**

```
check_puts(input_item$study_id,output_item$summary)
```

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context	<i>Contexts</i>
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**Description**

A predefined list of mw\_context objects. The context have been created to mirror the metabolomics workbench API documentation contexts as closely as possible.

**Usage**

```
context
```

**Format**

An object of class list of length 7.

**Examples**

```
# list available contexts  
names(context)  
  
# get the context 'study'  
context$study
```

---

context_inputs	<i>Valid inputs</i>
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**Description**

Get a list of valid input\_items for a context.

**Usage**

```
context_inputs(context)
```

**Arguments**

context	The name of a valid context (character)
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**Value**

A list of input item names for a context

**Examples**

```
# list of input items for the "study" context
context_inputs("study")
```

---

context_outputs	<i>Valid outputs</i>
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**Description**

Get a list of valid output\_items for a context.

**Usage**

```
context_outputs(context)
```

**Arguments**

context	The name of a valid context (character)
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**Value**

A list of output item names for a context

**Examples**

```
# list of output items for the "study" context
context_outputs("study")
```

---

`do_query`*Query the Metabolomics Workbench database*

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**Description**

Sends a query to the metabolomics database and returns the result. Note that while objects derived from `mw_base` can be used the recommended approach is to use character inputs.

**Usage**

```
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'character,character,character,character'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_moverz_context,list,character,mw_output_item'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_moverz_context,list,character,character'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_moverz_context,list,character,missing'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_moverz_context,character,character,missing'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'character,character,character,missing'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_exactmass_context,list,character,mw_output_item'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_exactmass_context,list,character,character'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_exactmass_context,list,character,missing'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_exactmass_context,character,character,missing'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'character,character,character,missing'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_context,mw_input_item,character,mw_output_item'
do_query(context, input_item, input_value, output_item)
```

```

## S4 method for signature 'mw_context,mw_input_item,character,mw_SE_item'
do_query(context, input_item, input_value, output_item)

## S4 method for signature
## 'mw_context,mw_input_item,character,mw_untarg_SE_item'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_context,mw_input_item,character,mw_DE_item'
do_query(context, input_item, input_value, output_item)

## S4 method for signature
## 'mw_context,mw_input_item,character,mw_untarg_DE_item'
do_query(context, input_item, input_value, output_item)

## S4 method for signature 'mw_context,mw_input_item,character,mw_MAE_item'
do_query(context, input_item, input_value, output_item)

```

### Arguments

context	A valid context name (character)
input_item	A valid input_item name (character)
input_value	The value for the input item (character).
output_item	A valid output_item (character).

### Value

A data.frame, or other output appropriate to the output\_item.

### Examples

```

# Get a summary of all studies with "diabetes" in the title

df = do_query(
  context = 'study',
  input_item = 'study_title',
  input_value = 'diabetes',
  output_item = 'summary'
)

```

---

input_example	<i>Valid input_value for input_item</i>
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### Description

Displays a valid input\_value for an input\_item and returns an example that matches the required input pattern.

**Usage**

```
input_example(input_item)
```

**Arguments**

input\_item      The name of a valid input\_item (character)

**Value**

An example input value matching the pattern required for the chosen input item.

**Examples**

```
# example input_value for input item "study_id"
input_example('study_id')
```

---

input_item	<i>Input items</i>
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**Description**

A predefined list of mw\_input\_item objects. The items have been created to mirror the Metabolomics Workbench API documentation input items as closely as possible.

**Usage**

```
input_item
```

**Format**

An object of class list of length 41.

**Examples**

```
# list available input_items
names(input_item)

# get the input item 'study_id'
input_item$study_id
```

---

`is_valid`*Check validity of context and inputs/outputs*

---

### Description

Compares the input and output items to the expected values for a context and returns TRUE if the inputs/puts are valid for the provided context or an error if not. This method is used internally and not for intended for general use.

### Usage

```
is_valid(context, input_item, input_value, output_item)

## S4 method for signature 'mw_context,character,character,character'
is_valid(context, input_item, input_value, output_item)

## S4 method for signature 'mw_moverz_context,character,character,missing'
is_valid(context, input_item, input_value)

## S4 method for signature 'mw_exactmass_context,character,character,missing'
is_valid(context, input_item, input_value)
```

### Arguments

<code>context</code>	An <code>mw_context</code> object.
<code>input_item</code>	An <code>mw_input_item</code> object, or the name of one.
<code>input_value</code>	The value for the input item (character).
<code>output_item</code>	An <code>mw_output_item</code> , or the name of one.

### Value

TRUE or an error.

### Examples

```
is_valid(context$study,
  input_item$study_id$name,
  'ST000001',
  output_item$summary$name
)
```

---

output_item	<i>Output items</i>
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**Description**

A predefined list of mw\_output\_item objects. The items have been created to mirror the Metabolomics Workbench API documentation output items as closely as possible.

**Usage**

```
output_item
```

**Format**

An object of class list of length 31.

**Examples**

```
# list available output_items
names(output_item)

# get the output item 'summary'
output_item$summary
```

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