

# Package ‘gtexr’

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**Title** Query the GTEx Portal API

**Version** 0.2.0

**Description** A convenient R interface to the Genotype-Tissue Expression (GTEx) Portal API. The GTEx project is a comprehensive public resource for studying tissue-specific gene expression and regulation in human tissues. Through systematic analysis of RNA sequencing data from 54 non-diseased tissue sites across nearly 1000 individuals, GTEx provides crucial insights into the relationship between genetic variation and gene expression. This data is accessible through the GTEx Portal API enabling programmatic access to human gene expression data. For more information on the API, see <<https://gtexportal.org/api/v2/redoc>>.

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**URL** <https://docs.ropensci.org/gtexr/>,  
<https://github.com/ropensci/gtexr>

**BugReports** <https://github.com/ropensci/gtexr/issues>

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**calculate\_expression\_quantitative\_trait\_loci**  
*Calculate Expression Quantitative Trait Loci*

---

## Description

Calculate your own eQTLs

- This service calculates the gene-variant association for any given pair of gene and variant, which may or may not be significant.
- This requires as input a GENCODE ID, GTEx variant ID, and tissue site detail ID.

By default, the calculation is based on the latest GTEx release.

[GTEx Portal API documentation](#).

## Usage

```
calculate_expression_quantitative_trait_loci(  
    tissueSiteDetailId,  
    gencodeId,  
    variantId,  
    datasetId = "gtex_v8",  
    .return_raw = FALSE  
)
```

## Arguments

tissueSiteDetailId  
String. The ID of the tissue of interest. Can be a GTEx specific ID (e.g. "Whole\_Blood"; use [get\\_tissue\\_site\\_detail\(\)](#) to see valid values) or an Ontology ID.

gencodeId  
String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".

<code>variantId</code>	String. A gtex variant ID.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Details

Notes on output:

- Beta and standard error are recorded in columns `nes` and `error` respectively (see [GTEx FAQs](#))
- `variantId` contains (in order) chromosome, position, reference allele, alternative allele and human genome build separated by underscores. The reference and alternative alleles for "chr1\_13550\_G\_A\_b38" for example are "G" and "A" respectively.
- See examples for how to calculate minor and alternative allele frequencies.

Notes on input:

- Argument `variantId` also accepts RSIDs.

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Dynamic Association Endpoints: [calculate\\_ieqtls\(\)](#), [calculate\\_isqtls\(\)](#), [calculate\\_splicing\\_quantitative\\_trait\\_loci\(\)](#)

## Examples

```
## Not run:
# perform request - returns a tibble with a single row
calculate_expression_quantitative_trait_loci(
  tissueSiteDetailId = "Whole_Blood",
  gencodeId = "ENSG00000203782.5",
  variantId = "rs79641866"
)

# unnest list columns with tidyr::unnest()
calculate_expression_quantitative_trait_loci(
  tissueSiteDetailId = "Whole_Blood",
  gencodeId = "ENSG00000203782.5",
  variantId = "rs79641866"
) |>
  tidyr::unnest(c("data", "genotypes"))

# to calculate minor and alternative allele frequencies
calculate_expression_quantitative_trait_loci(
  tissueSiteDetailId = "Liver",
  gencodeId = "ENSG00000237973.1",
  variantId = "rs12119111"
) |>
```

```

dplyr::bind_rows(.id = "rsid") |>
tidyr::separate(
  col = "variantId",
  into = c(
    "chromosome",
    "position",
    "reference_allele",
    "alternative_allele",
    "genome_build"
  ),
  sep = "_"
) |>
# ...then ascertain alternative_allele frequency
dplyr::mutate(
  alt_allele_count = (2 * homoAltCount) + hetCount,
  total_allele_count = 2 * (homoAltCount + hetCount + homoRefCount),
  alternative_allele_frequency = alt_allele_count / total_allele_count
) |>
dplyr::select(
  rsid,
  beta = nes,
  se = error,
  pValue,
  minor_allele_frequency = maf,
  alternative_allele_frequency,
  chromosome:genome_build,
  tissueSiteDetailId
)
## End(Not run)

```

**calculate\_ieqtls***Calculate Ieqtls***Description**

Calculate your own Cell Specific eQTLs.

- This service calculates the gene-variant association for any given pair of gene and variant, which may or may not be significant.
- This requires as input a GENCODE ID, GTEx variant ID, and tissue site detail ID.

By default, the calculation is based on the latest GTEx release.

[GTEx Portal API documentation](#).

**Usage**

```
calculate_ieqtls(
  cellType,
```

```

tissueSiteDetailId,
gencodeId,
variantId,
datasetId = "gtex_v8",
.return_raw = FALSE
)

```

## Arguments

<code>cellType</code>	String. "Adipocytes", "Epithelial_cells", "Hepatocytes", "Keratinocytes", "Myocytes", "Neurons", "Neutrophils".
<code>tissueSiteDetailId</code>	String. The ID of the tissue of interest. Can be a GTEx specific ID (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or an Ontology ID.
<code>gencodeId</code>	String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".
<code>variantId</code>	String. A gtex variant ID.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Dynamic Association Endpoints: [calculate\\_expression\\_quantitative\\_trait\\_loci\(\)](#), [calculate\\_isqtls\(\)](#), [calculate\\_splicing\\_quantitative\\_trait\\_loci\(\)](#)

## Examples

```

## Not run:
# perform request
calculate_ieqtls(
  cellType = "Adipocytes",
  tissueSiteDetailId = "Adipose_Subcutaneous",
  gencodeId = "ENSG00000203782.5",
  variantId = "chr1_1099341_T_C_b38"
)
## End(Not run)

```

---

**calculate\_isqtl**      *Calculate Isqtl*

---

**Description**

Calculate your own Cell Specific sQTLs.

- This service calculates the gene-variant association for any given pair of gene and variant, which may or may not be significant.
- This requires as input a GENCODE ID, GTEx variant ID, and tissue site detail ID.

By default, the calculation is based on the latest GTEx release.

[GTEx Portal API documentation](#).

**Usage**

```
calculate_isqtl(
  cellType,
  tissueSiteDetailId,
  phenotypeId,
  variantId,
  datasetId = "gtex_v8",
  .return_raw = FALSE
)
```

**Arguments**

cellType	String. "Adipocytes", "Epithelial_cells", "Hepatocytes", "Keratinocytes", "Myocytes", "Neurons", "Neutrophils".
tissueSiteDetailId	String. The ID of the tissue of interest. Can be a GTEx specific ID (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or an Ontology ID.
phenotypeId	String. See <a href="#">GTEx portal FAQs</a> for further details.
variantId	String. A gtex variant ID.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Dynamic Association Endpoints: [calculate\\_expression\\_quantitative\\_trait\\_loci\(\)](#), [calculate\\_ieqtls\(\)](#), [calculate\\_splicing\\_quantitative\\_trait\\_loci\(\)](#)

## Examples

```
## Not run:
# perform request
calculate_isqtlis(
  cellType = "Neutrophils",
  tissueSiteDetailId = "Whole_Blood",
  phenotypeId = "chr1:15947:16607:clu_40980:ENSG00000227232.5",
  variantId = "chr1_1099341_T_C_b38"
)
## End(Not run)
```

**calculate\_splicing\_quantitative\_trait\_loci**  
*Calculate Splicing Quantitative Trait Loci*

## Description

GTEX Portal API documentation.

## Usage

```
calculate_splicing_quantitative_trait_loci(
  tissueSiteDetailId,
  phenotypeId,
  variantId,
  datasetId = "gtex_v8",
  .return_raw = FALSE
)
```

## Arguments

<code>tissueSiteDetailId</code>	String. The ID of the tissue of interest. Can be a GTEX specific ID (e.g. "Whole_Blood"; use <code>get_tissue_site_detail()</code> to see valid values) or an Ontology ID.
<code>phenotypeId</code>	String. See <a href="#">GTEX portal FAQs</a> for further details.
<code>variantId</code>	String. A gtex variant ID.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw = TRUE`.

## See Also

Other Dynamic Association Endpoints: [calculate\\_expression\\_quantitative\\_trait\\_loci\(\)](#),  
[calculate\\_ieqtls\(\)](#), [calculate\\_isqtls\(\)](#)

## Examples

```
## Not run:  
# perform request - returns a tibble with a single row  
calculate_splicing_quantitative_trait_loci(  
  tissueSiteDetailId = "Whole_Blood",  
  phenotypeId = "chr1:15947:16607:clu_40980:ENSG00000227232.5",  
  variantId = "chr1_14677_G_A_b38"  
)  
  
## End(Not run)
```

---

download

*Download*

---

## Description

[GTEx Portal API documentation](#)

## Usage

```
download(  
  materialTypes = NULL,  
  tissueSiteDetailIds = NULL,  
  pathCategory = NULL,  
  tissueSampleIds = NULL,  
  sex = NULL,  
  sortBy = "sampleId",  
  sortDirection = "asc",  
  searchTerm = NULL,  
  sampleIds = NULL,  
  subjectIds = NULL,  
  ageBrackets = NULL,  
  hardyScales = NULL,  
  hasExpressionData = NULL,  
  hasGenotype = NULL,  
  .return_raw = FALSE  
)
```

## Arguments

**materialTypes** String, vector. Options: "Cells:Cell Line Viable", "DNA:DNA Genomic", "DNA:DNA Somatic", "RNA:Total RNA", "Tissue:PAXgene Preserved", "Tissue:PAXgene Preserved Paraffin-embedded", "Tissue:Fresh Frozen Tissue".

<b>tissueSiteDetailIds</b>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
<b>pathCategory</b>	Character vector. Options: "adenoma", "amylacea", "atelectasis", "atherosclerosis", "atherosis", "atrophy", "calcification", "cirrhosis", "clean_specimens", "congestion", "corpora_albicantia", "cyst", "desquamation", "diabetic", "dysplasia", "edema", "emphysema", "esophagitis", "fibrosis", "gastritis", "glomerulosclerosis", "goiter", "gynecomastoid", "hashimoto", "heart_failure_cells", "hemorrhage", "hepatitis", "hyalinization", "hypereosinophilia", "hyperplasia", "hypertrophy", "hypoxic", "infarction", "inflammation", "ischemic_changes", "macrophages", "mastopathy", "metaplasia", "monckeberg", "necrosis", "nephritis", "nephrosclerosis", "no_abnormalities", "nodularity", "pancreatitis", "pigment", "pneumonia", "post_menopausal", "prostatitis", "saponification", "scarring", "sclerotic", "solar_elastosis", "spermatogenesis", "steatosis", "sweat_glands", "tma".
<b>tissueSampleIds</b>	Array of strings. A list of Tissue Sample ID(s).
<b>sex</b>	String. Options: "male", "female".
<b>sortBy</b>	String. Options: "sampleId", "ischemicTime", "aliquotId", "tissueSampleId", "hardyScale", "pathologyNotes", "ageBracket", "tissueSiteDetailId", "sex".
<b>sortDirection</b>	String. Options: "asc", "desc". Default = "asc".
<b>searchTerm</b>	String.
<b>sampleIds</b>	Character vector. GTEx sample ID.
<b>subjectIds</b>	Character vector. GTEx subject ID.
<b>ageBrackets</b>	The age bracket(s) of the donors of interest. Options: "20-29", "30-39", "40-49", "50-59", "60-69", "70-79".
<b>hardyScales</b>	Character vector. A list of Hardy Scale(s) of interest. Options: "Ventilator case", "Fast death - violent", "Fast death - natural causes", "Intermediate death", "Slow death".
<b>hasExpressionData</b>	Logical.
<b>hasGenotype</b>	Logical.
<b>.return_raw</b>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Details

Note: running this request with no filters (i.e. `download()`) raises an error.

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Biobank Data Endpoints: [get\\_sample\\_biobank\\_data\(\)](#)

## Examples

```
## Not run:
download(
  materialTypes = "RNA:Total RNA",
  tissueSiteDetailIds = "Thyroid",
  pathCategory = "clean_specimens",
  sex = "male",
  ageBrackets = "50-59"
)
## End(Not run)
```

**get\_annotation**      *Get Annotation*

## Description

This service returns the list of annotations and allowed values by which a particular dataset can be subsetted. Results may be filtered by dataset.

[GTEx Portal API documentation](#)

## Usage

```
get_annotation(
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Details

Note: the output for this function appears to be incomplete currently.

**Value**

A tibble. Or a list if `.return_raw = TRUE`.

**See Also**

Other Datasets Endpoints: `get_collapsed_gene_model_exon()`, `get_downloads_page_data()`, `get_file_list()`, `get_full_get_collapsed_gene_model_exon()`, `get_functional_annotation()`, `get_linkage_disequilibrium_by_variant_data()`, `get_linkage_disequilibrium_data()`, `get_sample_datasets()`, `get_subject()`, `get_tissue_site_detail()`, `get_variant()`, `get_variant_by_location()`

**Examples**

```
## Not run:
get_annotation()

## End(Not run)
```

## `get_clustered_median_exon_expression`

*Get Clustered Median Exon Expression*

**Description**

Find median transcript expression data along with hierarchical clusters.

- Returns median normalized transcript expression in tissues of all known transcripts of a given gene along with the hierarchical clustering results of tissues and transcripts, based on exon expression, in Newick format.
- The hierarchical clustering is performed by calculating Euclidean distances and using the average linkage method.
- **This endpoint is not paginated.**

By default, this endpoint queries the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
get_clustered_median_exon_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

## Examples

```
## Not run:
get_clustered_median_exon_expression(c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))

# clustering data is stored as an attribute "clusters"
result <- get_clustered_median_exon_expression(c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))
attr(result, "clusters")

# process clustering data with the ape package
# install.packages("ape")
# phylo_tree <- ape::read.tree(text = attr(result, "clusters")$tissue)
# plot(phylo_tree)
# print(phylo_tree)

## End(Not run)
```

---

**get\_clustered\_median\_gene\_expression**  
*Get Clustered Median Gene Expression*

---

## Description

Find median gene expression data along with hierarchical clusters.

- Returns median gene expression in tissues along with The hierarchical clustering results of tissues and genes, based on gene expression, in Newick format.
- Results may be filtered by dataset, gene or tissue, but at least one gene must be provided
- The hierarchical clustering is performed by calculating Euclidean distances and using the average linkage method.
- **This endpoint is not paginated.**

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_clustered_median_gene_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

## Examples

```
## Not run:
get_clustered_median_gene_expression(gencodeIds = c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))

# clustering data is stored as an attribute "clusters"
result <- get_clustered_median_gene_expression(c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))
attr(result, "clusters")

# process clustering data with the ape package
# install.packages("ape")
# phylo_tree <- ape::read.tree(text = attr(result, "clusters")$tissue)
# plot(phylo_tree)
# print(phylo_tree)

## End(Not run)
```

**get\_clustered\_median\_junction\_expression**  
*Get Clustered Median Junction Expression*

## Description

Find median junction expression data along with hierarchical clusters.

- Returns median junction read counts in tissues of a given gene from all known transcripts along with the hierarchical clustering results of tissues and genes, based on junction expression, in Newick format.
- Results may be filtered by dataset, gene or tissue, but at least one gene must be provided.
- The hierarchical clustering is performed by calculating Euclidean distances and using the average linkage method.
- **This endpoint is not paginated.**

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
get_clustered_median_junction_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  .return_raw = FALSE
)
```

**Arguments**

<code>gencodeIds</code>	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

**See Also**

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

**Examples**

```
## Not run:
get_clustered_median_junction_expression(gencodeIds = c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))

# clustering data is stored as an attribute "clusters"
result <- get_clustered_median_junction_expression(c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))
attr(result, "clusters")

# process clustering data with the ape package
# install.packages("ape")
```

```
# phylo_tree <- ape::read.tree(text = attr(result, "clusters")$tissue)
# plot(phylo_tree)
# print(phylo_tree)

## End(Not run)
```

**get\_clustered\_median\_transcript\_expression**  
*Get Clustered Median Transcript Expression*

## Description

Find median transcript expression data of all known transcripts of a gene along with hierarchical clusters.

- Returns median normalized expression in tissues of all known transcripts of a given gene along with the hierarchical clustering results of tissues and genes, based on expression, in Newick format.
- Results may be filtered by dataset, gene or tissue, but at least one gene must be provided.
- The hierarchical clustering is performed by calculating Euclidean distances and using the average linkage method.
- **This endpoint is not paginated.**

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_clustered_median_transcript_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw = TRUE`.

**See Also**

Other Expression Data Endpoints: `getClusteredMedianExonExpression()`, `getClusteredMedianGeneExpression()`, `getClusteredMedianJunctionExpression()`, `getExpressionPCA()`, `getGeneExpression()`, `getMedianExonExpression()`, `getMedianGeneExpression()`, `getMedianJunctionExpression()`, `getMedianTranscriptExpression()`, `getSingleNucleusGex()`, `getSingleNucleusGexSummary()`, `getTopExpressedGenes()`

**Examples**

```
## Not run:
getClusteredMedianTranscriptExpression(gencodeIds = c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))

# clustering data is stored as an attribute "clusters"
result <- getClusteredMedianTranscriptExpression(c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))
attr(result, "clusters")

# process clustering data with the ape package
# install.packages("ape")
# phylo_tree <- ape::read.tree(text = attr(result, "clusters")$tissue)
# plot(phylo_tree)
# print(phylo_tree)

## End(Not run)
```

`getCollapsedGeneModelExon`  
*Get Collapsed Gene Model Exon*

**Description**

This service returns the collapsed exons in the gene model of the given gene. Gene-level and exon-level expression quantification were based on the GENCODE annotation, collapsed to a single transcript model for each gene using an algorithm developed by the GTEx analysis team.

By default, this service queries the models used by the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
getCollapsedGeneModelExon(
  gencodeId,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

gencodeId	String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Datasets Endpoints: [get\\_annotation\(\)](#), [get\\_downloads\\_page\\_data\(\)](#), [get\\_file\\_list\(\)](#), [get\\_full\\_get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_functional\\_annotation\(\)](#), [get\\_linkage\\_disequilibrium\\_by\\_vcf\(\)](#), [get\\_linkage\\_disequilibrium\\_data\(\)](#), [get\\_sample\\_datasets\(\)](#), [get\\_subject\(\)](#), [get\\_tissue\\_site\\_detail\(\)](#), [get\\_variant\(\)](#), [get\\_variant\\_by\\_location\(\)](#)

**Examples**

```
## Not run:
getCollapsedGeneModelExon(gencodeId = "ENSG00000132693.12")

## End(Not run)
```

`get_dataset_info`      *Get Dataset Info*

## Description

[GTEx Portal API documentation](#)

## Usage

```
get_dataset_info(
  datasetId = NULL,
  organizationName = NULL,
  .return_raw = FALSE
)
```

## Arguments

<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>organizationName</code>	String. Options: "GTEx Consortium" "Kid's First".
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## Examples

```
## Not run:
get_dataset_info(datasetId = "gtex_v8", organizationName = "GTEx Consortium")

## End(Not run)
```

`get_downloads_page_data`      *Get Downloads Page Data*

## Description

Retrieves all the files belonging to the given `project_id` for display on the Downloads Page

[GTEx Portal API documentation](#)

## Usage

```
get_downloads_page_data(project_id, .return_raw = FALSE)
```

## Arguments

- project\_id String. Options: "gtex", "adult-gtex", "egtex".  
.return\_raw Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Details

Note: The GTEx Portal API documentation states "GTEx currently has one project available: gtex". However, project\_id values "adult-gtex" and "egtex" both return results, whereas "gtex" does not (see examples).

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Datasets Endpoints: [get\\_annotation\(\)](#), [get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_file\\_list\(\)](#), [get\\_full\\_get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_functional\\_annotation\(\)](#), [get\\_linkage\\_disequilibrium\\_by\\_location\(\)](#), [get\\_linkage\\_disequilibrium\\_data\(\)](#), [get\\_sample\\_datasets\(\)](#), [get\\_subject\(\)](#), [get\\_tissue\\_site\\_detail\(\)](#), [get\\_variant\(\)](#), [get\\_variant\\_by\\_location\(\)](#)

## Examples

```
## Not run:  
# "adult-gtex" (default `project_id` value) and "egtex" both return results  
get_downloads_page_data("adult-gtex")  
egtex <- get_downloads_page_data("egtex")  
egtex  
  
# ... "gtex" does not  
get_downloads_page_data("gtex")  
  
# get details for whole blood methylation data, including download URL  
purrr::pluck(  
  egtex$children,  
  1,  
  "folders",  
  "Methylation - EPIC Array",  
  "children",  
  "folders",  
  "mQTLs",  
  "children",  
  "files",  
  "WholeBlood.mQTLs.regular.txt.gz"  
)  
  
## End(Not run)
```

---

<code>get_eqtl_genes</code>	<i>Get Eqtl Genes</i>
-----------------------------	-----------------------

---

## Description

Retrieve eGenes (eQTL Genes).

- This service returns eGenes (eQTL Genes) from the specified dataset.
- eGenes are genes that have at least one significant cis-eQTL acting upon them.
- Results may be filtered by tissue. By default, the service queries the latest GTEx release.

For each eGene, the results include the allelic fold change (log2AllelicFoldChange), p-value (pValue), p-value threshold (pValueThreshold), empirical p-value (empiricalPValue), and q-value (qValue).

- The log2AllelicFoldChange is the allelic fold change (in log2 scale) of the most significant eQTL.
- The pValue is the nominal p-value of the most significant eQTL.
- The pValueThreshold is the p-value threshold used to determine whether a cis-eQTL for this gene is significant. For more details see <https://gtexportal.org/home/documentationPage#staticTextAnalysisMethods>.
- The empiricalPValue is the beta distribution-adjusted empirical p-value from FastQTL.
- The qValues were calculated based on the empirical p-values. A false discovery rate (FDR) threshold of <= 0.05 was applied to identify genes with a significant eQTL.

[GTEx Portal API documentation](#).

## Usage

```
get_eqtl_genes(
  tissueSiteDetailIds = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <code>get_tissue_site_detail()</code> to see valid values) or Ontology IDs.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>page</code>	Integer (default = 0).

<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

**See Also**

Other Static Association Endpoints: [get\\_fine\\_mapping\(\)](#), [get\\_independent\\_eqtl\(\)](#), [get\\_multi\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\\_by\\_location\(\)](#), [get\\_significant\\_single\\_tissue\\_ieqlts\(\)](#), [get\\_significant\\_single\\_tissue\\_isqlts\(\)](#), [get\\_significant\\_single\\_sqtl\\_genes\(\)](#)

**Examples**

```
## Not run:
get_eqtl_genes(c("Whole_Blood", "Artery_Aorta"))

## End(Not run)
```

get\_exons

*Get Exons***Description**

This service returns exons from all known transcripts of the given gene.

- A versioned GENCODE ID is required to ensure that all exons are from a single gene.
- A dataset ID or both GENCODE version and genome build must be provided.
- Although annotated exons are not dataset dependent, specifying a dataset here is equivalent to specifying the GENCODE version and genome build used by that dataset.

[GTEx Portal API documentation](#)

**Usage**

```
get_exons(
  gencodeIds,
  gencodeVersion = NULL,
  genomeBuild = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>gencodeIds</code>	A character vector of Versioned GENCODE IDs, e.g. <code>c("ENSG00000132693.12", "ENSG00000203782.5")</code> .
<code>gencodeVersion</code>	String (default = "v26"). GENCODE annotation release. Either "v26" or "v19".
<code>genomeBuild</code>	String. Options: "GRCh38/hg38", "GRCh37/hg19". Default = "GRCh38/hg38".
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Reference Genome Endpoints: [get\\_gene\\_search\(\)](#), [get\\_genes\(\)](#), [get\\_genomic\\_features\(\)](#), [get\\_gwas\\_catalog\\_by\\_location\(\)](#), [get\\_neighbor\\_gene\(\)](#), [get\\_transcripts\(\)](#)

## Examples

```
## Not run:
get_exons(gencodeIds = c("ENSG00000132693.12", "ENSG00000203782.5"))

## End(Not run)
```

`get_expression_pca`      *Get Expression Pca*

## Description

Find gene expression PCA data.

- Returns gene expression PCA (principal component analysis) in tissues.
- Results may be filtered by tissue, sample, or dataset.

By default, the service queries the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
get_expression_pca(
  tissueSiteDetailIds,
  datasetId = "gtex_v8",
  sampleId = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_srnaseq_pilot".
sampleId	String. ^GTEX-[A-Z0-9]{5}-[0-9]{4}-SM-[A-Z0-9]{5}\$
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

**Examples**

```
## Not run:
get_expression_pca(tissueSiteDetailIds = c(
  "Adipose_Subcutaneous",
  "Whole_Blood"
))
get_expression_pca(
```

```

tissueSiteDetailIds = "Adipose_Subcutaneous",
sampleId = "GTEX-1117F-0226-SM-5GZZ7"
)

## End(Not run)

```

**get\_file\_list**      *Get File List*

## Description

Get all the files in GTEx dataset for Download page

[GTEx Portal API documentation](#)

## Usage

```
get_file_list(.return_raw = FALSE)
```

## Arguments

.return\_raw      Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Details

The returned tibble includes a nested list column, "filesets". This details files, sub-categorised by fileset (see examples section).

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Datasets Endpoints: [get\\_annotation\(\)](#), [getCollapsedGeneModelExon\(\)](#), [getDownloadsPageData\(\)](#), [getFullGetCollapsedGeneModelExon\(\)](#), [getFunctionalAnnotation\(\)](#), [getLinkageDisequilibriumByLocation\(\)](#), [getLinkageDisequilibriumData\(\)](#), [getSampleDatasets\(\)](#), [getSubject\(\)](#), [getTissueSiteDetail\(\)](#), [getVariant\(\)](#), [getVariantByLocation\(\)](#)

## Examples

```

## Not run:
# Column "filesets" is a list column
get_file_list()

# Get "GTEx Analysis V9" file list
gtex_v9_files <- get_file_list() |>
  dplyr::filter(name == "GTEx Analysis V9") |>
  dplyr::pull(filesets)

```

```
# "GTEx Analysis V9" filesets
names(gtex_v9_files[[1]])

# "GTEx Analysis V9", "snRNA-Seq Data" fileset files
names(gtex_v9_files[[1]][["snRNA-Seq Data"]])$files

## End(Not run)
```

**get\_fine\_mapping      *Get Fine Mapping***

### Description

Retrieve Fine Mapping Data

- Finds and returns Fine Mapping data for the provided list of genes
- By default, this endpoint fetches data from the latest GTEx version

The retrieved data is split into pages with `items_per_page` entries per page

[GTEx Portal API documentation](#)

### Usage

```
get_fine_mapping(
  gencodeIds,
  datasetId = "gtex_v8",
  variantId = NULL,
  tissueSiteDetailIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

### Arguments

<code>gencodeIds</code>	A character vector of Versioned GENCODE IDs, e.g. <code>c("ENSG00000132693.12", "ENSG00000203782.5")</code> .
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: <code>"gtex_v8"</code> , <code>"gtex_snrnaseq_pilot"</code> .
<code>variantId</code>	String. A gtex variant ID.
<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. <code>"Whole_Blood"</code> ; use <code>get_tissue_site_detail()</code> to see valid values) or Ontology IDs.
<code>page</code>	Integer (default = 0).

<code>.itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

### Value

A tibble. Or a list if `.return_raw` = TRUE.

### See Also

Other Static Association Endpoints: `get_eqtl_genes()`, `get_independent_eqtl()`, `get_multi_tissue_eqtls()`, `get_significant_single_tissue_eqtls()`, `get_significant_single_tissue_eqtls_by_location()`, `get_significant_single_tissue_ieqtls()`, `get_significant_single_tissue_isqlts()`, `get_significant_single_tissue_isqlts_by_location()`, `get_sqtl_genes()`

### Examples

```
## Not run:
# search by gene
get_fine_mapping(gencodeIds = c(
  "ENSG00000132693.12",
  "ENSG00000203782.5"
))

# optionally filter for a single variant and/or one or more tissues
get_fine_mapping(
  gencodeIds = c(
    "ENSG00000132693.12",
    "ENSG00000203782.5"
  ),
  variantId = "chr1_153228363_A_G_b38",
  tissueSiteDetailIds = c(
    "Whole_Blood",
    "Thyroid"
  )
)

## End(Not run)
```

`get_full_get_collapsed_gene_model_exon`  
*Get Full Get Collapsed Gene Model Exon*

### Description

This service allows the user to query the full Collapsed Gene Model Exon of a specific gene by gencode ID

[GTEx Portal API documentation](#)

## Usage

```
get_full_get_collapsed_gene_model_exon(
  gencodeId,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeId	String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Datasets Endpoints: [get\\_annotation\(\)](#), [getCollapsedGeneModelExon\(\)](#), [getDownloadsPageData\(\)](#), [getFileList\(\)](#), [getFunctionalAnnotation\(\)](#), [getLinkageDisequilibriumByVariantData\(\)](#), [getLinkageDisequilibriumData\(\)](#), [getSampleDatasets\(\)](#), [getSubject\(\)](#), [getTissueSiteDetail\(\)](#), [getVariant\(\)](#), [getVariantByLocation\(\)](#)

## Examples

```
## Not run:
get_full_get_collapsed_gene_model_exon(gencodeId = "ENSG00000203782.5")

## End(Not run)
```

---

**get\_functional\_annotation**  
*Get Functional Annotation*

---

**Description**

This endpoint retrieves the functional annotation of a certain chromosome location. Default to most recent dataset release.

[GTEx Portal API documentation](#)

**Usage**

```
get_functional_annotation(
  datasetId = "gtex_v8",
  chromosome,
  start,
  end,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
chromosome	String. One of "chr1", "chr2", "chr3", "chr4", "chr5", "chr6", "chr7", "chr8", "chr9", "chr10", "chr11", "chr12", "chr13", "chr14", "chr15", "chr16", "chr17", "chr18", "chr19", "chr20", "chr21", "chr22", "chrM", "chrX", "chrY".
start	Integer.
end	Integer.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Datasets Endpoints: [get\\_annotation\(\)](#), [getCollapsedGeneModelExon\(\)](#), [getDownloadsPageData\(\)](#), [getFileList\(\)](#), [getFullGetCollapsedGeneModelExon\(\)](#), [getLinkageDisequilibriumByVariantData\(\)](#), [getLinkageDisequilibriumData\(\)](#), [getSampleDatasets\(\)](#), [getSubject\(\)](#), [getTissueSiteDetail\(\)](#), [getVariant\(\)](#), [getVariantByLocation\(\)](#)

## Examples

```
## Not run:  
getFunctionalAnnotation(chromosome = "chr1", start = 192168000, end = 192169000)  
  
## End(Not run)
```

---

get\_genes

*Get Genes*

---

## Description

This service returns information about reference genes. A genome build and GENCODE version must be provided.

- Genes are searchable by gene symbol, GENCODE ID and versioned GENCODE ID.
- Versioned GENCODE ID is recommended to ensure unique ID matching.
- By default, this service queries the genome build and GENCODE version used by the latest GTEx release.

[GTEx API Portal documentation](#)

## Usage

```
get_genes(  
  geneIds,  
  gencodeVersion = "v26",  
  genomeBuild = "GRCh38/hg38",  
  page = 0,  
  itemsPerPage = getOption("gtexr.itemsPerPage"),  
  .verbose = getOption("gtexr.verbose"),  
  .return_raw = FALSE  
)
```

## Arguments

geneIds	A character vector of gene symbols, versioned gencodeIds, or unversioned gencodeIds.
gencodeVersion	String (default = "v26"). GENCODE annotation release. Either "v26" or "v19".
genomeBuild	String. Options: "GRCh38/hg38", "GRCh37/hg19". Default = "GRCh38/hg38".

<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

**See Also**

Other Reference Genome Endpoints: [get\\_exons\(\)](#), [get\\_gene\\_search\(\)](#), [get\\_genomic\\_features\(\)](#), [get\\_gwas\\_catalog\\_by\\_location\(\)](#), [get\\_neighbor\\_gene\(\)](#), [get\\_transcripts\(\)](#)

**Examples**

```
## Not run:
get_genes(c("CRP", "IL6R"))

## End(Not run)
```

**get\_gene\_expression     Get Gene Expression****Description**

Find normalized gene expression data.

- Returns normalized gene expression in tissues at the sample level.
- Results may be filtered by dataset, gene or tissue, but at least one gene must be provided.

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
get_gene_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  attributeSubset = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
attributeSubset	String. Examples include but are not limited to "sex", "ageBracket"
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with <a href="#">options(list(gtexpr.itemsPerPage = 100000))</a> .
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with <a href="#">options(list(gtexpr.verbose = FALSE))</a> .
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

## Examples

```
## Not run:
# multiple genes, selected tissues
get_gene_expression(
  gencodeIds = c(
    "ENSG00000132693.12",
    "ENSG00000203782.5"
  ),
  tissueSiteDetailIds = c("Thyroid", "Whole_Blood")
)

# single gene, selected (single) tissue
get_gene_expression(
  gencodeIds = "ENSG00000132693.12",
  tissueSiteDetailIds = "Whole_Blood"
)
```

```

# subset by sex
get_gene_expression(
  gencodeIds = "ENSG00000132693.12",
  tissueSiteDetailIds = "Whole_Blood",
  attributeSubset = "sex"
)

# subset by age bracket
get_gene_expression(
  gencodeIds = "ENSG00000132693.12",
  tissueSiteDetailIds = "Whole_Blood",
  attributeSubset = "ageBracket"
)

## End(Not run)

```

**get\_gene\_search**      *Get Gene Search*

### Description

Find genes that are partial or complete match of a gene\_id

- gene\_id could be a gene symbol, a gencode ID, or an Ensemble ID
- Gencode Version and Genome Build must be specified

[GTEx Portal API documentation](#)

### Usage

```

get_gene_search(
  geneId,
  gencodeVersion = "v26",
  genomeBuild = "GRCh38/hg38",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)

```

### Arguments

geneId	String. A gene symbol, a gencode ID, or an Ensemble ID.
gencodeVersion	String (default = "v26"). GENCODE annotation release. Either "v26" or "v19".
genomeBuild	String. Options: "GRCh38/hg38", "GRCh37/hg19". Default = "GRCh38/hg38".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).

.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

### Value

A tibble. Or a list if .return\_raw = TRUE.

### See Also

Other Reference Genome Endpoints: [get\\_exons\(\)](#), [get\\_genes\(\)](#), [get\\_genomic\\_features\(\)](#), [get\\_gwas\\_catalog\\_by\\_location\(\)](#), [get\\_neighbor\\_gene\(\)](#), [get\\_transcripts\(\)](#)

### Examples

```
## Not run:  
get_gene_search("CRP")  
  
## End(Not run)
```

---

## get\_genomic\_features *Get Genomic Features*

---

### Description

[GTEx API Portal documentation](#)

### Usage

```
get_genomic_features(.featureId, datasetId = "gtex_v8", .return_raw = FALSE)
```

### Arguments

.featureId	String. A genomic feature e.g. GENCODE ID, RSID or GTEx Variant ID.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

### Details

This endpoint takes a path parameter "featureId".

### Value

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Reference Genome Endpoints: `get_exons()`, `get_gene_search()`, `get_genes()`, `get_gwas_catalog_by_location()`, `get_neighbor_gene()`, `get_transcripts()`

**Examples**

```
## Not run:
# gene symbol
get_genomic_features("brca1")

# GENCODE ID
get_genomic_features("ENSG00000132693.12")

# RSID
get_genomic_features("rs1815739")

# GTEx variant ID
get_genomic_features("chr11_66561023_G_GTTA_b38")

## End(Not run)
```

### `get_gwas_catalog_by_location`

*Get Gwas Catalog By Location*

**Description**

Find the GWAS Catalog on a certain chromosome between start and end locations.

[GTEx API Portal documentation](#)

**Usage**

```
get_gwas_catalog_by_location(
  start,
  end,
  chromosome,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

<code>start</code>	Integer.
<code>end</code>	Integer.

chromosome	String. One of "chr1", "chr2", "chr3", "chr4", "chr5", "chr6", "chr7", "chr8", "chr9", "chr10", "chr11", "chr12", "chr13", "chr14", "chr15", "chr16", "chr17", "chr18", "chr19", "chr20", "chr21", "chr22", "chrM", "chrX", "chrY".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Reference Genome Endpoints: [get\\_exons\(\)](#), [get\\_gene\\_search\(\)](#), [get\\_genes\(\)](#), [get\\_genomic\\_features\(\)](#), [get\\_neighbor\\_gene\(\)](#), [get\\_transcripts\(\)](#)

**Examples**

```
## Not run:
get_gwas_catalog_by_location(start = 1, end = 10000000, chromosome = "chr1")

## End(Not run)
```

get\_image

Get Image

**Description**

GTEEx Portal API documentation

**Usage**

```
get_image(
  tissueSampleIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>tissueSampleIds</code>	Array of strings. A list of Tissue Sample ID(s).
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## Examples

```
## Not run:
get_image()

# filter by `tissueSampleId`
result <- get_image(tissueSampleIds = "GTEX-1117F-0526")
print(result)

# note that `pathologyNotesCategories` (if present) is a list column
print(result$pathologyNotesCategories)

## End(Not run)
```

`get_independent_eqtl`   *Get Independent Eqtl*

## Description

Retrieve Independent eQTL Data

- Finds and returns Independent eQTL Data data for the provided list of genes
- By default, this endpoint fetches data from the latest GTEx version

The retrieved data is split into pages with `items_per_page` entries per page

[GTEx portal API documentation](#)

## Usage

```
get_independent_eqtl(
  gencodeIds,
  tissueSiteDetailIds = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Static Association Endpoints: [get\\_eqtl\\_genes\(\)](#), [get\\_fine\\_mapping\(\)](#), [get\\_multi\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\\_by\\_location\(\)](#), [get\\_significant\\_single\\_tissue\\_ieqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_isqlts\(\)](#), [get\\_significant\\_single\\_tissue\\_isqlts\\_by\\_location\(\)](#), [get\\_sqtl\\_genes\(\)](#)

## Examples

```
## Not run:
# search by gene
get_independent_eqtl(gencodeIds = c(
  "ENSG00000132693.12",
  "ENSG00000203782.5"
))
```

```
# optionally filter for a single variant and/or one or more tissues
get_independent_eqtl(
  gencodeIds = c(
    "ENSG00000132693.12",
    "ENSG00000203782.5"
  ),
  tissueSiteDetailIds = c(
    "Whole_Blood",
    "Thyroid"
  )
)

## End(Not run)
```

## get\_linkage\_disequilibrium\_by\_variant\_data

*Get Linkage Disequilibrium By Variant Data*

### Description

Find linkage disequilibrium (LD) data for a given variant

[GTEx Portal API documentation](#)

### Usage

```
get_linkage_disequilibrium_by_variant_data(
  variantId,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

### Arguments

variantId	String. A gtex variant ID.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

### Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Datasets Endpoints: [get\\_annotation\(\)](#), [getCollapsedGeneModelExon\(\)](#), [getDownloadsPageData\(\)](#), [getFileList\(\)](#), [getFullGetCollapsedGeneModelExon\(\)](#), [getFunctionalAnnotation\(\)](#), [getLinkageDisequilibriumData\(\)](#), [getSampleDatasets\(\)](#), [getSubject\(\)](#), [getTissueSiteDetail\(\)](#), [getVariant\(\)](#), [getVariantByLocation\(\)](#)

## Examples

```
get_linkage_disequilibrium_by_variant_data("chr1_159245536_C_T_b38")
```

**get\_linkage\_disequilibrium\_data**  
*Get Linkage Disequilibrium Data*

## Description

Find linkage disequilibrium (LD) data for a given gene.

This endpoint returns linkage disequilibrium data for the cis-eQTLs found associated with the provided gene in a specified dataset. Results are queried by gencode ID. By default, the service queries the latest GTEx release. Specify a dataset ID to fetch results from a different dataset.

[GTEx Portal API documentation](#)

## Usage

```
get_linkage_disequilibrium_data(
  gencodeId,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeId	String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw = TRUE`.

**See Also**

Other Datasets Endpoints: [get\\_annotation\(\)](#), [getCollapsedGeneModelExon\(\)](#), [getDownloadsPageData\(\)](#), [getFileList\(\)](#), [getFullGetCollapsedGeneModelExon\(\)](#), [getFunctionalAnnotation\(\)](#), [getLinkageDisequilibriumByVariantData\(\)](#), [getSampleDatasets\(\)](#), [getSubject\(\)](#), [getTissueSiteDetail\(\)](#), [getVariant\(\)](#), [getVariantByLocation\(\)](#)

**Examples**

```
get_linkage_disequilibrium_data(gencodeId = "ENSG00000132693.12")
```

---

`get_maintenance_message`

*Get Maintenance Message*

---

**Description**

Getting all the maintenance messages from the database that are enabled.

[GTEx Portal API documentation](#).

**Usage**

```
get_maintenance_message(  
  page = 0,  
  itemsPerPage = getOption("gtexr.itemsPerPage"),  
  .verbose = getOption("gtexr.verbose"),  
  .return_raw = FALSE  
)
```

**Arguments**

<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Details**

Note this typically returns an empty tibble.

## Value

A tibble. Or a list if `.return_raw = TRUE`.

## See Also

Other Admin Endpoints: [get\\_news\\_item\(\)](#)

## Examples

```
## Not run:  
get_maintenance_message()  
  
## End(Not run)
```

---

get\_median\_exon\_expression  
*Get Median Exon Expression*

---

## Description

Find median exon expression data.

- Returns median exon read counts, in tissues, of a collapsed gene model.
- Results may be filtered by dataset, gene or tissue, but at least one gene must be provided

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_median_exon_expression(  
  gencodeIds,  
  datasetId = "gtex_v8",  
  tissueSiteDetailIds = NULL,  
  page = 0,  
  itemsPerPage = getOption("gtexr.itemsPerPage"),  
  .verbose = getOption("gtexr.verbose"),  
  .return_raw = FALSE  
)
```

## Arguments

<code>gencodeIds</code>	A character vector of Versioned GENCODE IDs, e.g. <code>c("ENSG00000132693.12", "ENSG00000203782.5")</code> .
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: <code>"gtex_v8"</code> , <code>"gtex_snrnaseq_pilot"</code> .

tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <code>get_tissue_site_detail()</code> to see valid values) or Ontology IDs.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Expression Data Endpoints: `get_clustered_median_exon_expression()`, `get_clustered_median_gene_expression()`, `get_clustered_median_junction_expression()`, `get_clustered_median_transcript_expression()`, `get_expression_pca()`, `get_gene_expression()`, `get_median_gene_expression()`, `get_median_junction_expression()`, `get_median_transcript_expression()`, `get_single_nucleus_gex()`, `get_single_nucleus_gex_summary()`, `get_top_expressed_genes()`

## Examples

```
## Not run:
# median exon expression values for CRP, filtered for whole blood
get_median_exon_expression(
  gencodeIds = "ENSG00000132693.12",
  tissueSiteDetailIds = "Whole_Blood"
)
## End(Not run)
```

## get\_median\_gene\_expression

*Get Median Gene Expression*

## Description

Find median gene expression data along with hierarchical clusters.

- Returns median gene expression in tissues.
- By default, this endpoint queries the latest GTEx release.

## Usage

```
get_median_gene_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

## Examples

```
## Not run:
get_median_gene_expression(gencodeIds = "ENSG00000132693.12")

## End(Not run)
```

`get_median_junction_expression`  
*Get Median Junction Expression*

## Description

Find junction gene expression data.

- Returns median junction read counts in tissues of a given gene from all known transcripts.
- Results may be filtered by dataset or tissue.

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_median_junction_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>gencodeIds</code>	A character vector of Versioned GENCODE IDs, e.g. <code>c("ENSG00000132693.12", "ENSG00000203782.5")</code> .
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: <code>"gtex_v8"</code> , <code>"gtex_snrnaseq_pilot"</code> .
<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. <code>"Whole_Blood"</code> ; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw = TRUE`.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

## Examples

```
## Not run:
get_median_junction_expression(gencodeIds = "ENSG00000132693.12")

## End(Not run)
```

**get\_median\_transcript\_expression**  
*Get Median Transcript Expression*

## Description

Find median transcript expression data of all known transcripts of a gene.

- Returns median normalized expression in tissues of all known transcripts of a given gene.
- Results may be filtered by dataset or tissue.

By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_median_transcript_expression(
  gencodeIds,
  datasetId = "gtex_v8",
  tissueSiteDetailIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snRNAseq_pilot".

<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <code>get_tissue_site_detail()</code> to see valid values) or Ontology IDs.
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

### Value

A tibble. Or a list if `.return_raw` = TRUE.

### See Also

Other Expression Data Endpoints: `get_clustered_median_exon_expression()`, `get_clustered_median_gene_expression()`, `get_clustered_median_junction_expression()`, `get_clustered_median_transcript_expression()`, `get_expression_pca()`, `get_gene_expression()`, `get_median_exon_expression()`, `get_median_gene_expression()`, `get_median_junction_expression()`, `get_single_nucleus_gex()`, `get_single_nucleus_gex_summary()`, `get_top_expressed_genes()`

### Examples

```
## Not run:
get_median_transcript_expression(gencodeIds = "ENSG00000132693.12")

## End(Not run)
```

## `get_multi_tissue_eqtls` *Get Multi Tissue Eqtls*

### Description

Find multi-tissue eQTL Metasoft results.

- This service returns multi-tissue eQTL Metasoft results for a given gene and variant in a specified dataset.
- A Versioned GENCODE ID must be provided.
- For each tissue, the results include: m-value (mValue), normalized effect size (nes), p-value (pValue), and standard error (se).
- The m-value is the posterior probability that an eQTL effect exists in each tissue tested in the cross-tissue meta-analysis (Han and Eskin, PLoS Genetics 8(3): e1002555, 2012).

- The normalized effect size is the slope of the linear regression of normalized expression data versus the three genotype categories using single-tissue eQTL analysis, representing eQTL effect size.
- The p-value is from a t-test that compares observed NES from single-tissue eQTL analysis to a null NES of 0.

By default, the service queries the latest GTEx release. The retrieved data is split into pages with items\_per\_page entries per page

[GTEx Portal API documentation](#)

## Usage

```
get_multi_tissue_eqtls(
  gencodeId,
  variantId = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeId	String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".
variantId	String. A gtex variant ID.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Static Association Endpoints: [get\\_eqtl\\_genes\(\)](#), [get\\_fine\\_mapping\(\)](#), [get\\_independent\\_eqtl\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\\_by\\_location\(\)](#), [get\\_significant\\_single\\_tissue\\_ieqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_isqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_sqtl\(\)](#), [get\\_sqtl\\_genes\(\)](#)

## Examples

```
## Not run:
# search by gene
get_multi_tissue_eqtls(gencodeId = "ENSG00000132693.12")

# note that 'tissues' is a list-column
x <- get_multi_tissue_eqtls(gencodeId = "ENSG00000132693.12",
                           variantId = "chr1_159476920_T_C_b38")

x$tissues[[1]]

## End(Not run)
```

**get\_neighbor\_gene**      *Get Neighbor Gene*

## Description

Find all neighboring genes on a certain chromosome around a position with a certain window size.

[GTEx API Portal documentation](#)

## Usage

```
get_neighbor_gene(
  pos,
  chromosome,
  bp_window,
  page = 0,
  gencodeVersion = "v26",
  genomeBuild = "GRCh38/hg38",
  itemsPerPage =getOption("gtexr.itemsPerPage"),
  .verbose =getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>pos</code>	Integer.
<code>chromosome</code>	String. One of "chr1", "chr2", "chr3", "chr4", "chr5", "chr6", "chr7", "chr8", "chr9", "chr10", "chr11", "chr12", "chr13", "chr14", "chr15", "chr16", "chr17", "chr18", "chr19", "chr20", "chr21", "chr22", "chrM", "chrX", "chrY".
<code>bp_window</code>	Integer.
<code>page</code>	Integer (default = 0).
<code>gencodeVersion</code>	String (default = "v26"). GENCODE annotation release. Either "v26" or "v19".
<code>genomeBuild</code>	String. Options: "GRCh38/hg38", "GRCh37/hg19". Default = "GRCh38/hg38".

<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

**See Also**

Other Reference Genome Endpoints: `get_exons()`, `get_gene_search()`, `get_genes()`, `get_genomic_features()`, `get_gwas_catalog_by_location()`, `get_transcripts()`

**Examples**

```
## Not run:
get_neighborgene(pos = 1000000, chromosome = "chr1", bp_window = 10000)

## End(Not run)
```

`get_news_item`      *Get News Item*

**Description**

Getting all the news items from the database that are current.

[GTEx Portal API documentation.](#)

**Usage**

```
get_news_item(
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw = TRUE`.

**See Also**

Other Admin Endpoints: [get\\_maintenance\\_message\(\)](#)

**Examples**

```
## Not run:  
get_news_item()  
  
## End(Not run)
```

---

`get_sample_biobank_data`  
*Get Sample (Biobank Data)*

---

**Description**

[GTEx Portal API documentation](#)

**Usage**

```
get_sample_biobank_data(  
  draw = NULL,  
  materialTypes = NULL,  
  tissueSiteDetailIds = NULL,  
  pathCategory = NULL,  
  tissueSampleIds = NULL,  
  sex = NULL,  
  sortBy = "sampleId",  
  sortDirection = "asc",  
  searchTerm = NULL,  
  sampleIds = NULL,  
  subjectIds = NULL,  
  ageBrackets = NULL,  
  hardyScales = NULL,  
  hasExpressionData = NULL,  
  hasGenotype = NULL,  
  page = 0,  
  itemsPerPage = getOption("gtexr.itemsPerPage"),  
  .verbose = getOption("gtexr.verbose"),  
  .return_raw = FALSE  
)
```

## Arguments

draw	Integer.
materialTypes	String, vector. Options: "Cells:Cell Line Viable", "DNA:DNA Genomic", "DNA:DNA Somatic", "RNA:Total RNA", "Tissue:PAXgene Preserved", "Tissue:PAXgene Preserved Paraffin-embedded", "Tissue:Fresh Frozen Tissue".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
pathCategory	Character vector. Options: "adenoma", "amylacea", "atelectasis", "atherosclerosis", "atherosis", "atrophy", "calcification", "cirrhosis", "clean_specimens", "congestion", "corpora_albicania", "cyst", "desquamation", "diabetic", "dysplasia", "edema", "emphysema", "esophagitis", "fibrosis", "gastritis", "glomerulosclerosis", "goiter", "gynecomastoid", "hashimoto", "heart_failure_cells", "hemorrhage", "hepatitis", "hyalinization", "hypereosinophilia", "hyperplasia", "hypertrophy", "hypoxic", "infarction", "inflammation", "ischemic_changes", "macrophages", "mastopathy", "metaplasia", "monckeberg", "necrosis", "nephritis", "nephrosclerosis", "no_abnormalities", "nodularity", "pancreatitis", "pigment", "pneumonia", "post_menopausal", "prostatitis", "saponification", "scarring", "sclerotic", "solar_elastosis", "spermatogenesis", "steatosis", "sweat_glands", "tma".
tissueSampleIds	Array of strings. A list of Tissue Sample ID(s).
sex	String. Options: "male", "female".
sortBy	String. Options: "sampleId", "ischemicTime", "aliquotId", "tissueSampleId", "hardyScale", "pathologyNotes", "ageBracket", "tissueSiteDetailId", "sex".
sortDirection	String. Options: "asc", "desc". Default = "asc".
searchTerm	String.
sampleIds	Character vector. GTEx sample ID.
subjectIds	Character vector. GTEx subject ID.
ageBrackets	The age bracket(s) of the donors of interest. Options: "20-29", "30-39", "40-49", "50-59", "60-69", "70-79".
hardyScales	Character vector. A list of Hardy Scale(s) of interest. Options: "Ventilator case", "Fast death - violent", "Fast death - natural causes", "Intermediate death", "Slow death".
hasExpressionData	Logical.
hasGenotype	Logical.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw = TRUE`.

**See Also**

Other Biobank Data Endpoints: [download\(\)](#)

**Examples**

```
## Not run:
get_sample_biobank_data(tissueSiteDetailIds = "Whole_Blood")

## End(Not run)
```

`get_sample_datasets`    *Get Sample (Datasets)*

**Description**

This service returns information of samples used in analyses from all datasets. Results may be filtered by dataset ID, sample ID, subject ID, sample metadata, or other provided parameters. By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
get_sample_datasets(
  datasetId = "gtex_v8",
  sampleIds = NULL,
  tissueSampleIds = NULL,
  subjectIds = NULL,
  ageBrackets = NULL,
  sex = NULL,
  pathCategory = NULL,
  tissueSiteDetailIds = NULL,
  aliquotIds = NULL,
  autolysisScores = NULL,
  hardyScales = NULL,
  ischemicTimes = NULL,
  ischemicTimeGroups = NULL,
  rins = NULL,
  uberonIds = NULL,
  dataTypes = NULL,
  sortBy = "sampleId",
  sortDirection = "asc",
  page = 0,
  itemsPerPage =getOption("gtexr.itemsPerPage"),
```

```

  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)

```

## Arguments

datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
sampleIds	Character vector. GTEx sample ID.
tissueSampleIds	Array of strings. A list of Tissue Sample ID(s).
subjectIds	Character vector. GTEx subject ID.
ageBrackets	The age bracket(s) of the donors of interest. Options: "20-29", "30-39", "40-49", "50-59", "60-69", "70-79".
sex	String. Options: "male", "female".
pathCategory	Character vector. Options: "adenoma", "amylacea", "atelectasis", "atherosclerosis", "atherosis", "atrophy", "calcification", "cirrhosis", "clean_specimens", "congestion", "corpora_albicania", "cyst", "desquamation", "diabetic", "dysplasia", "edema", "emphysema", "esophagitis", "fibrosis", "gastritis", "glomerulosclerosis", "goiter", "gynecomastoid", "hashimoto", "heart_failure_cells", "hemorrhage", "hepatitis", "hyalinization", "hypereosinophilia", "hyperplasia", "hypertrophy", "hypoxic", "infarction", "inflammation", "ischemic_changes", "macrophages", "mastopathy", "metaplasia", "monckeberg", "necrosis", "nephritis", "nephrosclerosis", "no_abnormalities", "nodularity", "pancreatitis", "pigment", "pneumonia", "post_menopausal", "prostatitis", "saponification", "scarring", "sclerotic", "solar_elastosis", "spermatogenesis", "steatosis", "sweat_glands", "tma".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
aliquotIds	Character vector.
autolysisScores	Character vector. Options: "None", "Mild", "Moderate", "Severe".
hardyScales	Character vector. A list of Hardy Scale(s) of interest. Options: "Ventilator case", "Fast death - violent", "Fast death - natural causes", "Intermediate death", "Slow death".
ischemicTimes	Integer.
ischemicTimeGroups	Character vector. Options: "<= 0", "1 - 300", "301 - 600", "601 - 900", "901 - 1200", "1201 - 1500", "> 1500".
rins	Integer, vector.
uberonIds	Character vector of Uberon IDs (e.g. "UBERON:EFO_0000572"; use <a href="#">get_tissue_site_detail()</a> to see valid values).
dataTypes	Character vector. Options: "RNASEQ", "WGS", "WES", "OMNI", "EXCLUDE".

<code>sortBy</code>	String. Options: "sampleId", "ischemicTime", "aliquotId", "tissueSampleId", "hardyScale", "pathologyNotes", "ageBracket", "tissueSiteDetailId", "sex".
<code>sortDirection</code>	String. Options: "asc", "desc". Default = "asc".
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

**See Also**

Other Datasets Endpoints: [get\\_annotation\(\)](#), [getCollapsedGeneModelExon\(\)](#), [getDownloadsPageData\(\)](#), [getFileList\(\)](#), [getFullGetCollapsedGeneModelExon\(\)](#), [getFunctionalAnnotation\(\)](#), [getLinkageDisequilibriumByVariantData\(\)](#), [getLinkageDisequilibriumData\(\)](#), [getSubject\(\)](#), [getTissueSiteDetail\(\)](#), [getVariant\(\)](#), [getVariantByLocation\(\)](#)

**Examples**

```
## Not run:
get_sample_datasets()

## End(Not run)
```

**get\_service\_info      *Get Service Info*****Description**

General information about the GTEx service.

[GTEx Portal API documentation](#).

**Usage**

```
get_service_info(.return_raw = FALSE)
```

**Arguments**

<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE
--------------------------	---

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

## Examples

```
## Not run:
get_service_info()

## End(Not run)
```

**get\_significant\_single\_tissue\_eqtls**  
*Get Significant Single Tissue Eqtls*

## Description

Find significant single tissue eQTLs.

- This service returns precomputed significant single tissue eQTLs.
- Results may be filtered by tissue, gene, variant or dataset.
- To search by gene, use the versioned GENCODE ID.
- To search by variant, use the dbSNP rs ID (snpId).

By default, the service queries the latest GTEx release and the retrieved data is split into pages with items\_per\_page entries per page

[GTEx Portal API documentation](#).

## Usage

```
get_significant_single_tissue_eqtls(
  gencodeIds = NULL,
  variantIds = NULL,
  tissueSiteDetailIds = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
variantIds	Character vector. Gtex variant IDs.
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.

<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Details

**Note:** although the GTEx Portal API documentation says to use the dbSNP rsID when searching by variant, this returns no results. Instead use gtex variant IDs e.g. use "chr1\_153209640\_C\_A\_b38" instead of "rs1410858".

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Static Association Endpoints: [get\\_eqtl\\_genes\(\)](#), [get\\_fine\\_mapping\(\)](#), [get\\_independent\\_eqtl\(\)](#), [get\\_multi\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\\_by\\_location\(\)](#), [get\\_significant\\_single\\_tissue\\_isqlts\(\)](#), [get\\_significant\\_single\\_tissue\\_sqtls\(\)](#), [get\\_sqtl\\_genes\(\)](#)

## Examples

```
## Not run:
# search by gene
get_significant_single_tissue_eqtls(gencodeIds = c(
  "ENSG00000132693.12",
  "ENSG00000203782.5"
))

# search by variant - must be variantId (not rsid)
get_significant_single_tissue_eqtls(variantIds = "chr1_153209640_C_A_b38")

# filter by gene/variant and tissue site - either `gencodeIds` or `variantIds`
# should be supplied as a minimum
get_significant_single_tissue_eqtls(
  gencodeIds = c(
    "ENSG00000132693.12",
    "ENSG00000203782.5"
  ),
  variantIds = "chr1_153209640_C_A_b38",
  tissueSiteDetailIds = "Whole_Blood"
)

## End(Not run)
```

---

`get_significant_single_tissue_eqtls_by_location`  
*Get Significant Single Tissue eQTLs By Location*

---

## Description

Find significant single tissue eQTLs using Chromosomal Locations.

- This service returns precomputed significant single tissue eQTLs.
- Results may be filtered by tissue, and/or dataset.

By default, the service queries the latest GTEx release. Since this endpoint is used to support a third party program on the portal, the return structure is different from other endpoints and is not paginated.

[GTEx Portal API documentation](#)

## Usage

```
get_significant_single_tissue_eqtls_by_location(  
  tissueSiteDetailId,  
  start,  
  end,  
  chromosome,  
  datasetId = "gtex_v8",  
  .return_raw = FALSE  
)
```

## Arguments

<code>tissueSiteDetailId</code>	String. The ID of the tissue of interest. Can be a GTEx specific ID (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or an Ontology ID.
<code>start</code>	Integer.
<code>end</code>	Integer.
<code>chromosome</code>	String. One of "chr1", "chr2", "chr3", "chr4", "chr5", "chr6", "chr7", "chr8", "chr9", "chr10", "chr11", "chr12", "chr13", "chr14", "chr15", "chr16", "chr17", "chr18", "chr19", "chr20", "chr21", "chr22", "chrM", "chrX", "chrY".
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw = TRUE`.

## See Also

Other Static Association Endpoints: `get_eqtl_genes()`, `get_fine_mapping()`, `get_independent_eqtl()`, `get_multi_tissue_eqtls()`, `get_significant_single_tissue_eqtls()`, `get_significant_single_tissue_ieqtls()`, `get_significant_single_tissue_isqtls()`, `get_significant_single_tissue_sqtls()`, `get_sqtl_genes()`

## Examples

```
## Not run:
get_significant_single_tissue_eqtls_by_location(
  tissueSiteDetailId = "Artery_Aorta",
  start = 10000,
  end = 250000,
  chromosome = "chr11"
)
## End(Not run)
```

`get_significant_single_tissue_ieqtls`  
*Get Significant Single Tissue Ieqtls*

## Description

Retrieve Interaction eQTL Data.

- This service returns cell type interaction eQTLs (ieQTLs), from a specified dataset.
- Results may be filtered by tissue
- By default, the service queries the latest GTEx release.

The retrieved data is split into pages with `items_per_page` entries per page

[GTEx Portal API documentation](#)

## Usage

```
get_significant_single_tissue_ieqtls(
  gencodeIds,
  variantIds = NULL,
  tissueSiteDetailIds = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
variantIds	Character vector. Gtex variant IDs.
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Static Association Endpoints: [get\\_eqtl\\_genes\(\)](#), [get\\_fine\\_mapping\(\)](#), [get\\_independent\\_eqtl\(\)](#), [get\\_multi\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\\_l\(\)](#), [get\\_significant\\_single\\_tissue\\_isqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_sqtls\(\)](#), [get\\_sqtl\\_genes\(\)](#)

## Examples

```
## Not run:
get_significant_single_tissue_ieqtls(c(
  "ENSG00000132693.12",
  "ENSG00000203782.5"
))
## End(Not run)
```

**get\_significant\_single\_tissue\_isqtls**  
*Get Significant Single Tissue Isqtls*

## Description

Retrieve Interaction sQTL Data.

- This service retrieves cell type interaction sQTLs (isQTLs), from a specified dataset.
- Results may be filtered by tissue
- By default, the service queries the latest GTEx release.

The retrieved data is split into pages with items\_per\_page entries per page

[GTEx Portal API documentation](#).

## Usage

```
get_significant_single_tissue_isqtl(
  gencodeIds,
  variantIds = NULL,
  tissueSiteDetailIds = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
variantIds	Character vector. Gtex variant IDs.
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Static Association Endpoints: [get\\_eqtl\\_genes\(\)](#), [get\\_fine\\_mapping\(\)](#), [get\\_independent\\_eqtl\(\)](#), [get\\_multi\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_ieqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_sqtls\(\)](#), [get\\_sqtl\\_genes\(\)](#)

## Examples

```
## Not run:  
get_significant_single_tissue_isqtlis(gencodeIds = c(  
  "ENSG00000065613.9",  
  "ENSG00000203782.5"  
)  
  
## End(Not run)
```

---

**get\_significant\_single\_tissue\_sqtls**  
*Get Significant Single Tissue Sqtls*

---

## Description

Retrieve Single Tissue sQTL Data.

- This service returns single tissue sQTL data for the given genes, from a specified dataset.
- Results may be filtered by tissue
- By default, the service queries the latest GTEx release.

The retrieved data is split into pages with `items_per_page` entries per page

[GTEx Portal API documentation](#).

## Usage

```
get_significant_single_tissue_sqtls(  
  gencodeIds = NULL,  
  variantIds = NULL,  
  tissueSiteDetailIds = NULL,  
  datasetId = "gtex_v8",  
  page = 0,  
  itemsPerPage = getOption("gtexr.itemsPerPage"),  
  .verbose = getOption("gtexr.verbose"),  
  .return_raw = FALSE  
)
```

## Arguments

<code>gencodeIds</code>	A character vector of Versioned GENCODE IDs, e.g. <code>c("ENSG00000132693.12", "ENSG00000203782.5")</code> .
<code>variantIds</code>	Character vector. Gtex variant IDs.
<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <code>get_tissue_site_detail()</code> to see valid values) or Ontology IDs.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_srnaseq_pilot".
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gttxr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gttxr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Static Association Endpoints: `get_eqtl_genes()`, `get_fine_mapping()`, `get_independent_eqtl()`, `get_multi_tissue_eqtls()`, `get_significant_single_tissue_eqtls()`, `get_significant_single_tissue_eqtls_k()`, `get_significant_single_tissue_ieqtls()`, `get_significant_single_tissue_isqls()`, `get_sqtl_genes()`

## Examples

```
## Not run:
# search by gene
get_significant_single_tissue_sqtls(gencodeIds = c(
  "ENSG00000065613.9",
  "ENSG00000203782.5"
))

## End(Not run)
```

## get\_single\_nucleus\_gex

*Get Single Nucleus Gex*

## Description

Retrieve Single Nucleus Gene Expression Data for a given Gene.

[GTEx Portal API documentation](#)

**Usage**

```
get_single_nucleus_gex(
  gencodeIds,
  datasetId = "gtex_snrnaseq_pilot",
  tissueSiteDetailIds = NULL,
  excludedataArray = TRUE,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

gencodeIds	A character vector of Versioned GENCODE IDs, e.g. c("ENSG00000132693.12", "ENSG00000203782.5").
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
excludedataArray	String. Options are TRUE or FALSE
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

**Examples**

```
## Not run:
# Search for one or more genes - returns a tibble with one row per tissue.
```

```

# Column "cellTypes" now contains a tibble of expression summary data, with
# one row for each cell type
get_single_nucleus_gex(gencodeIds = c(
  "ENSG00000203782.5",
  "ENSG00000132693.12"
))

# `excludedataArray = FALSE` - expression values are stored under "celltypes"
# in an additional column called "data"
response <- get_single_nucleus_gex(
  gencodeIds = "ENSG00000132693.12",
  excludedataArray = FALSE,
  itemsPerPage = 2
)

response

# "cellTypes" contains a tibble of data with one row for each
# cell type e.g. for Breast_Mammary_Tissue
response$cellTypes[[2]]

# when `excludedataArray = FALSE`, expression values are stored in "data"
# e.g. for Breast_Mammary_Tissue, Epithelial cell (luminal):
response$cellTypes[[2]]$data[[1]]

## End(Not run)

```

**get\_single\_nucleus\_gex\_summary**  
*Get Single Nucleus Gex Summary*

### Description

Retrieve Summarized Single Nucleus Gene Expression Data.

[GTEx Portal API documentation](#)

### Usage

```

get_single_nucleus_gex_summary(
  datasetId = "gtex_snrnaseq_pilot",
  tissueSiteDetailIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)

```

## Arguments

datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
tissueSiteDetailIds	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexpr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexpr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_top\\_expressed\\_genes\(\)](#)

## Examples

```
## Not run:
# all tissues
get_single_nucleus_gex_summary()

# filter for specific tissue
get_single_nucleus_gex_summary(tissueSiteDetailIds = c(
  "Breast_Mammary_Tissue",
  "Skin_Sun_Exposed_Lower_leg"
))
## End(Not run)
```

## Description

Retrieve sGenes (sQTL Genes).

- This service returns sGenes (sQTL Genes) from the specified dataset.
- Results may be filtered by tissue.
- By default, the service queries the latest GTEx release.

The retrieved data is split into pages with `items_per_page` entries per page

[GTEx Portal API documentation](#).

## Usage

```
get_sqtl_genes(
  tissueSiteDetailIds = NULL,
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

<code>tissueSiteDetailIds</code>	Character vector of IDs for tissues of interest. Can be GTEx specific IDs (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or Ontology IDs.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw = TRUE`.

## See Also

Other Static Association Endpoints: [get\\_eqtl\\_genes\(\)](#), [get\\_fine\\_mapping\(\)](#), [get\\_independent\\_eqtl\(\)](#), [get\\_multi\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\(\)](#), [get\\_significant\\_single\\_tissue\\_eqtls\\_l\(\)](#), [get\\_significant\\_single\\_tissue\\_ieqlts\(\)](#), [get\\_significant\\_single\\_tissue\\_isqlts\(\)](#), [get\\_significant\\_single\\_tissue\\_isqlts\\_l\(\)](#)

## Examples

```
## Not run:
get_sqtl_genes("Whole_Blood")

## End(Not run)
```

get\_subject

*Get Subject*

## Description

This service returns information of subjects used in analyses from all datasets. Results may be filtered by dataset ID, subject ID, sex, age bracket or Hardy Scale. By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_subject(
  datasetId = "gtex_v8",
  sex = NULL,
  ageBrackets = NULL,
  hardyScale = NULL,
  subjectIds = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
sex	String. Options: "male", "female".
ageBrackets	The age bracket(s) of the donors of interest. Options: "20-29", "30-39", "40-49", "50-59", "60-69", "70-79".
hardyScale	String A Hardy Scale of interest. Options: "Ventilator case", "Fast death - violent", "Fast death - natural causes", "Intermediate death", "Slow death".
subjectIds	Character vector. GTEx subject ID.
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Datasets Endpoints: [get\\_annotation\(\)](#), [get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_downloads\\_page\\_data\(\)](#), [get\\_file\\_list\(\)](#), [get\\_full\\_get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_functional\\_annotation\(\)](#), [get\\_linkage\\_disequilibrium\\_by\\_variant\\_data\(\)](#), [get\\_linkage\\_disequilibrium\\_data\(\)](#), [get\\_sample\\_datasets\(\)](#), [get\\_tissue\\_site\\_detail\(\)](#), [get\\_variant\(\)](#), [get\\_variant\\_by\\_location\(\)](#)

**Examples**

```
## Not run:
get_subject()

## End(Not run)
```

**get\_tissue\_site\_detail**

*Get Tissue Site Detail*

**Description**

Retrieve all tissue site detail information in the database

[GTEx Portal API documentation](#)

**Usage**

```
get_tissue_site_detail(
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Datasets Endpoints: [get\\_annotation\(\)](#), [get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_downloads\\_page\\_data\(\)](#), [get\\_file\\_list\(\)](#), [get\\_full\\_get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_functional\\_annotation\(\)](#), [get\\_linkage\\_disequilibrium\\_by\\_variant\\_data\(\)](#), [get\\_linkage\\_disequilibrium\\_data\(\)](#), [get\\_sample\\_datasets\(\)](#), [get\\_subject\(\)](#), [get\\_variant\(\)](#), [get\\_variant\\_by\\_location\(\)](#)

**Examples**

```
## Not run:  
# returns a tibble with one row per tissue  
get_tissue_site_detail()  
  
# `eqtlSampleSummary` and `rnaSeqSampleSummary` are list columns  
bladder_site_details <- get_tissue_site_detail() |>  
  dplyr::filter(tissueSiteDetailId == "Bladder")  
  
purrr::pluck(bladder_site_details, "eqtlSampleSummary", 1)  
  
purrr::pluck(bladder_site_details, "rnaSeqSampleSummary", 1)  
  
## End(Not run)
```

---

**get\_top\_expressed\_genes**

*Get Top Expressed Genes*

---

**Description**

Find top expressed genes for a specified tissue.

- Returns top expressed genes for a specified tissue in a dataset, sorted by median expression.
- When the optional parameter filterMtGene is set to true, mitochondrial genes will be excluded from the results. By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

**Usage**

```
get_top_expressed_genes(  
  tissueSiteDetailId,  
  datasetId = "gtex_v8",  
  filterMtGene = TRUE,  
  page = 0,  
  itemsPerPage =getOption("gtexr.itemsPerPage"),
```

```

  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)

```

## Arguments

<code>tissueSiteDetailId</code>	String. The ID of the tissue of interest. Can be a GTEx specific ID (e.g. "Whole_Blood"; use <a href="#">get_tissue_site_detail()</a> to see valid values) or an Ontology ID.
<code>datasetId</code>	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_srnaseq_pilot".
<code>filterMtGene</code>	Logical. Exclude mitochondrial genes.
<code>page</code>	Integer (default = 0).
<code>itemsPerPage</code>	Integer (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	Logical. If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if `.return_raw` = TRUE.

## See Also

Other Expression Data Endpoints: [get\\_clustered\\_median\\_exon\\_expression\(\)](#), [get\\_clustered\\_median\\_gene\\_expression\(\)](#), [get\\_clustered\\_median\\_junction\\_expression\(\)](#), [get\\_clustered\\_median\\_transcript\\_expression\(\)](#), [get\\_expression\\_pca\(\)](#), [get\\_gene\\_expression\(\)](#), [get\\_median\\_exon\\_expression\(\)](#), [get\\_median\\_gene\\_expression\(\)](#), [get\\_median\\_junction\\_expression\(\)](#), [get\\_median\\_transcript\\_expression\(\)](#), [get\\_single\\_nucleus\\_gex\(\)](#), [get\\_single\\_nucleus\\_gex\\_summary\(\)](#)

## Examples

```

## Not run:
get_top_expressed_genes(tissueSiteDetailId = "Artery_Aorta")

## End(Not run)

```

---

**get\_transcripts**      *Get Transcripts*

---

**Description**

Find all transcripts of a reference gene.

- This service returns information about transcripts of the given versioned GENCODE ID.
- A genome build and GENCODE version must be provided.
- By default, this service queries the genome build and GENCODE version used by the latest GTEx release.

[GTEx API Portal documentation](#)

**Usage**

```
get_transcripts(  
  gencodeId,  
  gencodeVersion = "v26",  
  genomeBuild = "GRCh38/hg38",  
  page = 0,  
  itemsPerPage = getOption("gtexr.itemsPerPage"),  
  .verbose = getOption("gtexr.verbose"),  
  .return_raw = FALSE  
)
```

**Arguments**

gencodeId	String. A Versioned GENCODE ID of a gene, e.g. "ENSG00000065613.9".
gencodeVersion	String (default = "v26"). GENCODE annotation release. Either "v26" or "v19".
genomeBuild	String. Options: "GRCh38/hg38", "GRCh37/hg19". Default = "GRCh38/hg38".
page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if .return\_raw = TRUE.

**See Also**

Other Reference Genome Endpoints: [get\\_exons\(\)](#), [get\\_gene\\_search\(\)](#), [get\\_genes\(\)](#), [get\\_genomic\\_features\(\)](#), [get\\_gwas\\_catalog\\_by\\_location\(\)](#), [get\\_neighbor\\_gene\(\)](#)

## Examples

```
## Not run:
get_transcripts(gencodeId = "ENSG00000203782.5")

## End(Not run)
```

**get\_variant**

*Get Variant*

## Description

This service returns information about a variant, including position, dbSNP RS ID, the reference allele, the alternative allele, and whether the minor allele frequency is  $\geq 1\%$ . For GTEx v6p, there is also information about whether the whole exome sequence and chip sequencing data are available. Results may be queried by GTEx variant ID (variantId), dbSNP RS ID (snpId) or genomic location (chromosome and pos). Variants are identified based on the genotype data of each dataset cohort, namely, are dataset-dependent. Each variant is assigned a unique GTEx variant ID (i.e. the primary key). Not all variants have a mappable dbSNP RS ID. By default, this service queries the latest GTEx release.

[GTEx Portal API documentation](#)

## Usage

```
get_variant(
  snpId = NULL,
  variantId = NULL,
  datasetId = "gtex_v8",
  chromosome = NULL,
  poss = NULL,
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

## Arguments

snpId	String
variantId	String. A gtex variant ID.
datasetId	String. Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
chromosome	String. One of "chr1", "chr2", "chr3", "chr4", "chr5", "chr6", "chr7", "chr8", "chr9", "chr10", "chr11", "chr12", "chr13", "chr14", "chr15", "chr16", "chr17", "chr18", "chr19", "chr20", "chr21", "chr22", "chrM", "chrX", "chrY".
poss	Integer, vector.

page	Integer (default = 0).
itemsPerPage	Integer (default = 250). Set globally to maximum value 100000 with options(list(gtexr.itemsPerPage = 100000)).
.verbose	Logical. If TRUE (default), print paging information. Set to FALSE globally with options(list(gtexr.verbose = FALSE)).
.return_raw	Logical. If TRUE, return the raw API JSON response. Default = FALSE

## Value

A tibble. Or a list if .return\_raw = TRUE.

## See Also

Other Datasets Endpoints: [get\\_annotation\(\)](#), [get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_downloads\\_page\\_data\(\)](#), [get\\_file\\_list\(\)](#), [get\\_full\\_get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_functional\\_annotation\(\)](#), [get\\_linkage\\_disequilibrium\\_by\\_variant\\_data\(\)](#), [get\\_linkage\\_disequilibrium\\_data\(\)](#), [get\\_sample\\_datasets\(\)](#), [get\\_subject\(\)](#), [get\\_tissue\\_site\\_detail\(\)](#), [get\\_variant\\_by\\_location\(\)](#)

## Examples

```
# search by rsid
get_variant(snpId = "rs1410858")

# search by variantId
get_variant(variantId = "chr1_153209640_C_A_b38")

# search by chromosome and position
get_variant(
  chromosome = "chr1",
  pos = 153209600:153209700
)
```

---

get\_variant\_by\_location  
Get Variant By Location

---

## Description

This service allows the user to query information about variants on a certain chromosome at a certain location.

[GTEx Portal API documentation](#)

**Usage**

```
get_variant_by_location(
  start,
  end,
  chromosome,
  sortBy = "pos",
  sortDirection = "asc",
  datasetId = "gtex_v8",
  page = 0,
  itemsPerPage = getOption("gtexr.itemsPerPage"),
  .verbose = getOption("gtexr.verbose"),
  .return_raw = FALSE
)
```

**Arguments**

<code>start</code>	<code>Integer.</code>
<code>end</code>	<code>Integer.</code>
<code>chromosome</code>	<code>String.</code> One of "chr1", "chr2", "chr3", "chr4", "chr5", "chr6", "chr7", "chr8", "chr9", "chr10", "chr11", "chr12", "chr13", "chr14", "chr15", "chr16", "chr17", "chr18", "chr19", "chr20", "chr21", "chr22", "chrM", "chrX", "chrY".
<code>sortBy</code>	<code>String.</code> Options: "sampleId", "ischemicTime", "aliquotId", "tissueSampleId", "hardyScale", "pathologyNotes", "ageBracket", "tissueSiteDetailId", "sex".
<code>sortDirection</code>	<code>String.</code> Options: "asc", "desc". Default = "asc".
<code>datasetId</code>	<code>String.</code> Unique identifier of a dataset. Usually includes a data source and data release. Options: "gtex_v8", "gtex_snrnaseq_pilot".
<code>page</code>	<code>Integer</code> (default = 0).
<code>itemsPerPage</code>	<code>Integer</code> (default = 250). Set globally to maximum value 100000 with <code>options(list(gtexr.itemsPerPage = 100000))</code> .
<code>.verbose</code>	<code>Logical.</code> If TRUE (default), print paging information. Set to FALSE globally with <code>options(list(gtexr.verbose = FALSE))</code> .
<code>.return_raw</code>	<code>Logical.</code> If TRUE, return the raw API JSON response. Default = FALSE

**Value**

A tibble. Or a list if `.return_raw` = TRUE.

**See Also**

Other Datasets Endpoints: [get\\_annotation\(\)](#), [get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_downloads\\_page\\_data\(\)](#), [get\\_file\\_list\(\)](#), [get\\_full\\_get\\_collapsed\\_gene\\_model\\_exon\(\)](#), [get\\_functional\\_annotation\(\)](#), [get\\_linkage\\_disequilibrium\\_by\\_variant\\_data\(\)](#), [get\\_linkage\\_disequilibrium\\_data\(\)](#), [get\\_sample\\_datasets\(\)](#), [get\\_subject\(\)](#), [get\\_tissue\\_site\\_detail\(\)](#), [get\\_variant\(\)](#)

**Examples**

```
get_variant_by_location(  
    start = 153209600,  
    end = 153209700,  
    chromosome = "chr1"  
)
```

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