

Package ‘quantMSImageR’

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Title Processing and Quantification of Targeted Mass Spectrometry Imaging Data

Version 0.99.7

Description Implements tools for processing and quantifying targeted DESI-MRM mass spectrometry imaging (MSI) datasets, extending the Cardinal package. Includes signal-to-noise filtering against background pixels, tissue/background separation, per-feature ion images, quantile heatmaps, quantification against on-slide or on-tissue calibration standards, and batch export of per-feature text images for external viewers.

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URL <https://mjs-708.github.io/quantMSImageR>,
<https://github.com/MJS-708/quantMSImageR>

BugReports <https://github.com/MJS-708/quantMSImageR/issues>

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quantMSImageR-package	<i>quantMSImageR: processing and quantification of targeted mass spectrometry imaging data</i>
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Description

Tools (extending Cardinal) for processing and quantifying targeted DESI-MRM mass spectrometry imaging data: signal-to-noise filtering, tissue/background separation, per-feature ion images, quantile heatmaps and standards-based quantification.

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See Also

Useful links:

- <https://mjs-708.github.io/quantMSImageR>
- <https://github.com/MJS-708/quantMSImageR>
- Report bugs at <https://github.com/MJS-708/quantMSImageR/issues>

alignFeatures	<i>Align two MSI objects to their common features</i>
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Description

Cardinal's cbind (used by [combineMSIs](#)) requires the mz key column to match exactly across objects. When acquisitions come from different instrument methods or ion-library versions, features with the same display name can carry different mz values, causing cbind to fail with a "non-matching key columns" error.

Usage

```
alignFeatures(
  obj1,
  obj2,
  feature_match = c("transition", "name"),
  mz_tolerance = 0.4
)
```

Arguments

obj1	An MSImagingExperiment – a quant_MSImagingExperiment is one. Its feature set and mz values are used as the reference, and its fData() must carry a name column.
obj2	An MSImagingExperiment to align against obj1, likewise carrying fData()\$name.
feature_match	Character. "transition" (default) verifies that same-named features share a precursor and product m/z; "name" matches on the display name alone, without checking what it refers to.
mz_tolerance	Numeric. Half-width in Da within which two precursor or product m/z values are taken to be the same (default 0.4, i.e. nominal mass). MRM selects Q1 and Q3 at unit resolution, so two panels can record the same channel as 308.17 and 308.2; at a tighter tolerance this check would reject them as different transitions. See readMRM .

Details

alignFeatures resolves this by:

1. Subsetting both objects to the *intersection* of feature names.
2. Reordering obj2 to match obj1's feature order.
3. Forcing obj2's mz values to equal obj1's, which permits Cardinal's key check to pass after features have been matched by name. [combineMSIs](#) then restores fData from obj1, so all feature metadata originates from the first object.

Features are paired by display name, and by default that pairing is then *verified* against the transition each name refers to: precursor and product m/z must agree to within mz_tolerance. Two methods can reuse a name while differing in precursor ion, product ion or transition definition, and because this function rewrites the mz key, an unverified pairing would silently merge different measurements into one feature. A disagreement is therefore an error naming the features involved.

feature_match = "name" restores name-only matching for objects that carry no precursor/product metadata. It is not the default: it cannot detect the failure above, and this function is called automatically inside [generateTxtImages](#).

The function is called automatically inside [generateTxtImages](#) before every cross-sample [combineMSIs](#), so mixed-method studies (e.g. 1 Hz + 2 Hz acquisitions) are handled transparently. It is also exported so that it can be used directly in custom scripts or Rmd reports whenever two quant_MSImagingExperiment objects need to be combined.

Value

A named list with two elements:

obj1 Subset of obj1 containing only common features.

obj2 Subset of obj2 reordered and mz-keyed to match obj1, ready for [combineMSIs](#).

See Also

[combineMSIs](#), [generateTxtImages](#)

Other combining acquisitions: [bindPanels\(\)](#), [combineMSIs](#), [MSImagingExperiment-method](#), [stitchAcquisitions\(\)](#)

Examples

```
p1 <- system.file("extdata", "example.raw", "section01.RDS",
                  package = "quantMSImageR")
p2 <- system.file("extdata", "example.raw", "section02.RDS",
                  package = "quantMSImageR")
a1 <- alignFeatures(readRDS(p1), readRDS(p2))
```

```
applySNR,quant_MSImagingExperiment-method
```

Apply SNR mask to intensity values

Description

Sets pixels to NA in `val_slot` wherever the `snr` spectra slot is NA. Typically called after `int2SNR()`: pixels that did not pass the SNR threshold (stored as NA in the `snr` slot) are suppressed in the intensity slot so they are excluded from downstream analysis and visualisation.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
applySNR(MSIobject, val_slot = "intensity", ...)
```

Arguments

<code>MSIobject</code>	A <code>quant_MSImagingExperiment</code> object containing both <code>val_slot</code> and an <code>snr</code> spectra slot (populated by <code>int2SNR()</code>).
<code>val_slot</code>	Character. Name of the intensity slot to mask (default "intensity").
<code>...</code>	Additional arguments (currently unused).

Value

The input object with sub-threshold pixels set to NA in `val_slot`.

Destructive

This modifies `val_slot` in place rather than adding a new layer. Keep a copy of the object if the pre-masking values are needed afterwards.

See Also

[int2SNR\(\)](#)

Other filtering: [back2NA,quant_MSImagingExperiment-method](#), [int2SNR,quant_MSImagingExperiment-method](#), [int2response,quant_MSImagingExperiment-method](#), [zero2NA,quant_MSImagingExperiment-method](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- int2SNR(obj, snr_thresh = 3)
obj <- applySNR(obj, val_slot = "intensity")
```

back2NA,quant_MSImagingExperiment-method

Set background pixel intensities to NA

Description

Replaces the spectral values of background pixels with NA in a chosen spectra slot. Tissue pixels are left unchanged.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
back2NA(
  MSIobject,
  val_slot = "intensity",
  pixel_header = "sample_name",
  background = "background_pixels",
  sample_type = NULL,
  ...
)
```

Arguments

MSIobject	A quant_MSImagingExperiment object whose pData() contains a column identifying pixel type.
val_slot	Character. Name of the spectra slot to modify (default "intensity").
pixel_header	Character. Column of pData() holding the pixel-type labels (default "sample_name", the imaging convention).
background	Character. Value in pixel_header that labels background pixels (default "background_pixels"). The historical "noise_pixels" / "Noise" labels are matched too, so masks made with earlier versions keep working.
sample_type	Deprecated alias for pixel_header, kept for backward compatibility. When supplied it overrides pixel_header.
...	Additional arguments (currently unused).

Value

The input quant_MSImagingExperiment with background pixels set to NA in val_slot. Returns the object unchanged if no background pixels are found.

Destructive

This modifies val_slot in place rather than adding a new layer. Keep a copy of the object if the original values are needed afterwards.

See Also

Other filtering: [applySNR,quant_MSImagingExperiment-method](#), [int2SNR,quant_MSImagingExperiment-method](#), [int2response,quant_MSImagingExperiment-method](#), [zero2NA,quant_MSImagingExperiment-method](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- back2NA(obj)
```

bindPanels	<i>Merge two MSI objects of the same tissue by coordinate-matched rbind</i>
------------	---

Description

Combines two MSImagingExperiment objects acquired on the *same physical sample* into one whose feature set spans both inputs. Pixels are matched by (x, y) coordinates rather than by row order, so the two acquisitions can have different pixel counts (which is the rule, not the exception: different MRM panels or polarities almost always sample at slightly different rates and therefore produce different grids).

Usage

```
bindPanels(
  obj1,
  obj2,
  label = NULL,
  feature_match = c("transition", "name"),
  mz_tolerance = 0.4
)
```

Arguments

obj1, obj2	MSImagingExperiment (or quant_MSImagingExperiment) objects of the same physical tissue. Both must carry x and y in pData(.).
label	Character. Optional. If supplied, written to pData(result)\$run so the merged object combines cleanly downstream.
feature_match	Character. How a feature name appearing in both inputs is treated: "transition" (default) keeps obj1's copy only after confirming both refer to the same precursor and product m/z, and errors otherwise; "name" keeps obj1's copy on the strength of the name alone.
mz_tolerance	Numeric. Half-width in Da within which two precursor or product m/z values count as the same (default 0.4, i.e. nominal mass – MRM selects Q1 and Q3 at unit resolution). See readMRM() .

Details

Pixels present in only one input are dropped – a small percentage of edge coverage in the more densely sampled acquisition typically. Pixels present in both keep real intensities from both inputs, so cross-panel / cross-polarity colocalisation is meaningful at the combined object.

The common use is pairing a positive- and a negative-mode acquisition of the same section, but any two panels of the same physical area work.

Value

A quant_MSImagingExperiment with:

- Features = obj1's features followed by obj2's features (feature names duplicated in both are kept once, with obj1's values).
- Pixels = intersection of (x, y) grids.
- Each pixel carries real intensities from both inputs.
- Every spectra layer both inputs carry – response, snr and calibrated amounts as well as intensity – together with obj1's experiment metadata and calibration/tissue slots.

Processed inputs

Both inputs must carry the same set of spectra layers, since a layer present on only one side cannot be filled in for the other half of the features. Bind panels at the same stage of processing: either both raw, or both after the same steps.

See Also

[combineMSIs\(\)](#), [generateTxtImages\(\)](#)

Other combining acquisitions: [alignFeatures\(\)](#), [combineMSIs, MSImagingExperiment-method](#), [stitchAcquisitions\(\)](#)

Examples

```
# Both inputs must be the SAME physical area measured twice, so the example
# splits one section into two disjoint "panels" rather than using two
# different sections -- binding unrelated sections by coordinate would
# silently pair pixels that are not the same piece of tissue.
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- readRDS(p)
panel_a <- obj[1:4, ]
panel_b <- obj[5:nrow(fData(obj)), ]

merged <- bindPanels(panel_a, panel_b, label = "SampleA_1")
nrow(fData(merged)) # features from both panels
```

buildFeatureMeta

Build per-feature metadata by joining an ion library on m/z

Description

Joins each row of fData(combined) to the ion library on the (precursor_mz, product_mz) pair, within mz_tolerance. Robust to feature renames in the YAML (features.rename) because m/z is stable across name changes.

Usage

```
buildFeatureMeta(
  combined,
  ion_lib_meta,
  mz_tolerance = 0.4,
  ambiguity = c("combine", "error", "warn", "nearest"),
  verbose = TRUE
)
```

Arguments

<code>combined</code>	An MSI object whose <code>fData</code> has columns <code>name</code> , <code>precursor_mz</code> and <code>product_mz</code> . Multi-transition features stored with " "-joined m/z strings are matched on the first token.
<code>ion_lib_meta</code>	A <code>data.frame</code> of ion library metadata (must contain <code>precursor_mz</code> and <code>product_mz</code>). Pass <code>NULL</code> to short-circuit and return <code>NULL</code> .
<code>mz_tolerance</code>	Numeric. Half-width in Da within which a feature's precursor and product – both, not either – are taken to be the library's. Default 0.4, nominal-mass matching, because MRM selects Q1 and Q3 at unit resolution. See readMRM() .
<code>ambiguity</code>	Character. What to do when a feature matches more than one library entry: "combine" (default, name it for all of them joined with " "), "error", "warn" or "nearest". See readMRM() .
<code>verbose</code>	Logical. Emit a <code>message()</code> reporting how many features matched. Default <code>TRUE</code> .

Details

Matching uses the same rule as [readMRM\(\)](#), so the two cannot disagree about which library entry a feature is. They previously built their m/z keys independently – one rounding to whole numbers, the other not – and a feature matching several library entries silently took the first one's annotation.

Used by `runStudy.R` (via `inst/runStudy.R`) and [runExample\(\)](#) to build a per-feature metadata frame the heatmap report can consume – e.g. for splitting / colour-bar annotation by an ion-library column such as `Met-1`.

Value

A `data.frame` of metadata, one row per feature in `fData(combined)`, with row names set to the feature name and a `name` column appended. Unmatched features become a row of NAs. Returns `NULL` when `ion_lib_meta` is `NULL`.

See Also

[quantileHm\(\)](#), [generateTxtImages\(\)](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
  package = "quantMSImageR")
obj <- readRDS(p)
lib <- read.csv(system.file("extdata", "example_ion_library.csv",
  package = "quantMSImageR"), check.names = FALSE)
fm <- buildFeatureMeta(obj, lib)
```

calibrationInfo-class *Calibration metadata for an imaging experiment*

Description

Holds everything needed to turn a measured response into an estimated amount: the calibration design, the summarised response at each level, the fitted per-analyte models, and their fit quality. It is carried in the calibrationInfo slot of a [quant_MSImagingExperiment](#).

Usage

```
## S4 method for signature 'calibrationInfo'
show(object)
```

Arguments

object A calibrationInfo object.

Details

The slots are filled in workflow order and are empty (zero-row data frames, empty list) until the corresponding step has run:

cal_metadata data.frame. The calibration design supplied by the user: one row per (spot, analyte), with columns identifier, analyte, amount_pg and level. Set by [summariseCalLevels\(\)](#).

cal_response_data data.frame. Summarised response per calibration level, with columns analyte, pg_perpixel, response_perpixel and level. Set by [summariseCalLevels\(\)](#).

cal_list list. One stats::lm per analyte, of response_perpixel ~ pg_perpixel, **named by analyte** so that [int2conc\(\)](#) can match models to features by fData(x)\$name. Set by [createCalCurve\(\)](#).

r2_df data.frame. One row per analyte, with feature and r2; [int2conc\(\)](#) adds an out_of_range column giving the proportion of pixels extrapolated beyond the calibrated range. Set by [createCalCurve\(\)](#).

Invariants checked by the validity method: cal_list must be named if it is non-empty, and every name must appear in r2_df\$feature when r2_df is populated. An unnamed cal_list is accepted only when empty – matching models to features positionally would silently apply one analyte's curve to another.

Use the accessors ([calibrationModels\(\)](#), [calibrationDiagnostics\(\)](#), [calibrationLevels\(\)](#), [calibrationMetadata\(\)](#)) rather than @.

Value

An object of class calibrationInfo.

Slots

cal_metadata data.frame of the calibration design.

cal_list Named list of fitted lm models, one per analyte.

cal_response_data data.frame of summarised response per level.

r2_df data.frame of fit quality per analyte.

See Also

[quant_MSImagingExperiment](#), [createCalCurve\(\)](#), [quantMSImageR-accessors](#)

Other calibration: [createCalCurve](#), [quant_MSImagingExperiment-method](#), [int2conc](#), [quant_MSImagingExperiment-method](#), [plotCalCoverage](#), [quant_MSImagingExperiment-method](#), [summariseCalLevels](#), [quant_MSImagingExperiment-method](#)

Examples

```
# Empty, as attached to a freshly coerced object
ci <- calibrationInfo()
ci

# Populated by the calibration workflow
cal_dir <- system.file("extdata", "cal_example.raw", package = "quantMSImageR")
cal <- as(readRDS(file.path(cal_dir, "cal_MSI.RDS")),
          "quant_MSImagingExperiment")
cal_metadata <- read.csv(file.path(cal_dir, "calibration_metadata.csv"))
cal <- summariseCalLevels(cal, cal_metadata, val_slot = "intensity",
                          cal_label = "Cal", id = "identifier")
cal <- createCalCurve(cal, cal_type = "cal")
calibrationDiagnostics(cal)
```

combineMSIs,MSImagingExperiment-method

Combine MSI experiments across acquisitions

Description

Concatenates two or more acquisitions pixel-wise into a single object, so that a whole study can be filtered, summarised and plotted together. Each input keeps its own run, which is what downstream per-sample summaries group on.

Usage

```
## S4 method for signature 'MSImagingExperiment'
combineMSIs(MSIobject, ...)
```

Arguments

MSIobject	An MSImagingExperiment; the first acquisition, whose fData() defines the shared feature axis.
...	Further MSImagingExperiment objects to combine. All must share the feature axis and the pData() columns of MSIobject – use alignFeatures() first if they do not.

Value

A single quant_MSImagingExperiment holding every input's pixels, with the shared fData() re-stored and one run level per acquisition.

See Also

Other combining acquisitions: [alignFeatures\(\)](#), [bindPanels\(\)](#), [stitchAcquisitions\(\)](#)

Examples

```
p1 <- system.file("extdata", "example.raw", "section01.RDS",
                  package = "quantMSImageR")
p2 <- system.file("extdata", "example.raw", "section02.RDS",
                  package = "quantMSImageR")
combined <- combineMSIs(readRDS(p1), readRDS(p2))
```

contributionHm

Group-mean heatmap with per-sample contribution as opacity

Description

The same heatmap as [quantileHm\(\)](#) – same orientation, same group and feature colour bars, same square cells – with only the cell fill changed. Each cell encodes two things at once:

Usage

```
contributionHm(
  MSIobject,
  quant_val = 0.5,
  scale = c("shared", "feature"),
  heatmap_order = NA,
  heatmap_labs = NA,
  feature_split = NULL,
  feature_split_name = "Pathway",
  group_split_name = "Group",
  cell_size = 8,
  max_aspect = 1.5,
  cell_border = "white",
  fontsize = 8,
  palette = c("heatmap2", "heatmap0"),
  group_palette = "hat",
  feature_palette = "reading",
  alpha_floor = 0.6,
  saturate = 0.04,
  na_col = "grey88"
)
```

Arguments

MSIobject	A quant_MSImagingExperiment object.
quant_val	Numeric in (0, 1). Quantile of pixel intensities summarised per feature per sample (default 0.5, the median).

scale	Character. How the colour scale is set. "shared" (default) puts every feature on ONE scale, capped at a quantile of all the group means, so a strongly separating feature is vivid and a flat one is pale – colours mean the same thing in every column and features can be compared with each other. "feature" normalises each feature by its own extreme, so every column fills the ramp regardless of how far it actually moved. That shows the pattern within a feature and makes comparison BETWEEN features meaningless – a feature varying by a few percent looks exactly like one that doubles. Use "shared" to ask which features changed most, and "feature" to read each feature's own profile. The study report also accepts "both" in <code>parameters\$heatmap_scale</code>, which draws each quantile section twice, once on each scale. That is a report-level setting; this function draws one heatmap and takes only "shared" or "feature".
heatmap_order	Character vector of run names in the desired column order. Must match values in <code>pData(MSIobject)\$run</code> . Defaults to NA (use discovery order).
heatmap_labs	Character vector of display labels, one per entry in <code>heatmap_order</code> . Used to create column-split groups. Defaults to NA (no splitting).
feature_split	Optional character/factor vector of length equal to the number of features, used to split the heatmap columns into labelled groups. Typically derived from an ion library metadata column (e.g. Met-1). Defaults to NULL (no splitting).
feature_split_name	Display name for the feature-group annotation legend (e.g. "Met-1", "Pathway"). Defaults to "Pathway".
group_split_name	Display name for the sample-group annotation legend (e.g. "Treatment", "Group"). Defaults to "Group".
cell_size	Numeric. Side of a heatmap cell, in millimetres (default 8). Giving the body an absolute size is what makes cells square; leaving it to <code>ComplexHeatmap</code> stretches them to fill the device, which produces very oblong cells when there are far more features than samples. Set to NA to restore the fill-the-device behaviour.
max_aspect	Numeric ≥ 1 . Largest cell width-to-height ratio allowed when one dimension has many more entries than the other (default 1.5). Cells stay square until the counts differ by more than four-fold; beyond that the shorter dimension's cells are widened up to this ratio so the plotting area is not reduced to a sliver.
cell_border	Colour for the line drawn around every cell (default "white"), which separates neighbouring cells instead of letting equal colours merge into one block. NA draws no border.
fontsize	Numeric. Point size for the sample and feature labels and the annotation names (default 8). <code>ComplexHeatmap</code> defaults to 12, which crowds the panel once feature names are long enough to need rotating.
palette	Character. Diverging colour ramp for the z-scores: "heatmap2" (default, blue-white-red) or "heatmap0". See quantPalettes() .
group_palette, feature_palette	Character. Qualitative palettes for the sample-group and feature-group colour bars (defaults "hat" and "reading"). Any <code>grDevices::hcl.colors()</code> palette name also works.
alpha_floor	Numeric in $[0, 1)$. Opacity given to a sample that did not contribute (default 0.6). Below roughly this a non-contributing sample starts to look like missing data.

saturate	Numeric in (0, 0.5). Fraction of cells allowed to saturate at the colour limit (default 0.04, i.e. 4%).
na_col	Colour for features that cannot be scored – no variance across samples, or missing values (default "grey88").

Details

Hue the sample's **group** mean for that feature – what the group did.

Opacity that **sample's** contribution to the group mean – who made it do that.

A plain group-mean heatmap cannot show that a group effect rests on one replicate. Here it is visible directly: a uniformly solid block is a real group effect, whereas one opaque tile among faded ones means the mean is carried by a single sample.

Value

A `ComplexHeatmap::Heatmap` (rows = samples, columns = features), carrying the opacity key in `attr(x, "contribution_legend")`. Pass that to `ComplexHeatmap::draw(annotation_legend_list = )` so the opacity channel is labelled; the report does this.

How the two channels are computed

Values are the per-sample quantile of pixel intensities, as in `quantileHm()`, then z-scored **per feature across all samples**, not within group – so a cell reads "high or low for this feature" rather than "abundant or not". Features with zero variance, and missing values, stay NA and are drawn in `na_col`; they are never imputed to the row mean.

The group mean of those z-scores is the hue. The contribution is $z * \text{sign}(\text{group mean})$, so positive means the sample moved with its group and zero or negative means it did not; only the positive part drives opacity, rescaled onto `[alpha_floor, 1]`. `alpha_floor` exists because a nearly transparent tile reads as missing data rather than as weak agreement.

Why there are two colour limits

The hue is a group **mean**, and averaging n replicates shrinks it by roughly $\sqrt{n}$. Reusing the per-sample z limit for the group mean therefore produces a washed-out panel that no amount of alpha tuning fixes. Both limits are computed from the quantity actually drawn, each as the $1 - \text{saturate}$ quantile of its own absolute values, so a similar small fraction of cells saturates on each channel and the bulk of the data gets the full ramp. Expect the two to differ by a factor of two or more. Note this differs from `quantileHm()`, which clips to a fixed `[-1, 1]`.

Both limits above are single numbers covering the whole panel, which is what lets one column be compared with another. `scale = "feature"` replaces them with a limit per feature, so each column is normalised by its own extreme and fills the ramp whatever its actual spread. That is the same reading `quantileHm()` gives, but keeping the group-mean hue and the contribution opacity: use it to look along a single feature, never across features.

Comparing features with each other

One cap or many. That is the whole of it.

`scale = "shared"` computes a single limit for the panel and draws every column on it, so a feature whose group mean reaches the cap saturates and one at a fifth of it stays pale. The pale column is pale because it moved less, and that difference is the information: features can be ranked against each other by how far their groups separate, in units of each feature's own spread.

scale = "feature" gives every column its own limit, its own extreme, so each fills the ramp whatever it actually did. Nothing is pale, so nothing can be ranked. A feature with no group effect at all looks exactly as vivid as the strongest one in the study – its largest group mean might be 0.05, and it is still drawn at full intensity. Use it to read one feature's profile across samples; never to argue that a feature matters.

What is being compared under "shared" is **effect size, not fold change**. The hue is a group separation measured in standard deviations of that feature, so a feature that doubles but varies wildly between replicates can sit below one that moves 20% in every sample. For magnitude, read the fold-change table instead. The two orderings are both correct and answer different questions.

For a balanced two-group design the group-mean z-score is bounded at ± 1 : perfectly separated groups with no within-group scatter put every sample at $z = \pm 1$, so the group means land on the bound, and identical groups put them on 0. A cap of 0.5 therefore means the strongest feature in the panel reaches about half the separation the design can show.

Two limits on the comparison, both worth remembering. The standard deviation is estimated from the samples in hand, so with a handful per group those estimates are noisy and the ranking is coarse – read it as "which features stand out", not as a precise order. And the deviation is pooled across groups, so a large separation inflates its own denominator; the scale compresses at the top and is not linear in effect size there.

See Also

[quantileHm\(\)](#) for the per-sample view of the same matrix.

Other visualisation: [imageR](#), [quant_MSImagingExperiment-method](#), [quantPalettes\(\)](#), [quantileHm\(\)](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
hm <- contributionHm(obj, quant_val = 0.5,
                    heatmap_order = "section01", heatmap_labs = "A")
```

createCalCurve,quant_MSImagingExperiment-method
Fit per-analyte calibration models

Description

Fits a linear response-versus-amount model for each standard, which [int2conc\(\)](#) later inverts to turn a measured response into an estimated amount per pixel.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
createCalCurve(
  MSIobject,
  cal_type,
  level = "level",
  background = "background",
  weighting = c("1/x", "none", "1/x2")
)
```

Arguments

MSIobject	A quant_MSImagingExperiment whose cal_response_data has been populated by summariseCalLevels() .
cal_type	Character, required. "cal" for standards deposited onto the slide, "std_addition" for standard addition on tissue. See Details.
level	Character. Column of cal_response_data holding the level labels (default "level").
background	Character. Value of level marking the background level subtracted when cal_type = "std_addition".
weighting	Character. Regression weights: "1/x" (default), "1/x2" or "none". See the Weighting section.

Details

cal_type has no default and must be chosen explicitly, because the two modes apply materially different processing.

cal_type = "cal" fits the summarised response directly against the amount deposited, appropriate for standards spotted onto the slide beside the tissue (on-slide / external calibration).

cal_type = "std_addition" is classical **standard addition**, for standards deposited onto tissue where endogenous analyte is already present. It:

1. fits an unweighted response $\sim$ deposited amount across all levels;
2. takes the x-intercept of that line – the amount at which predicted response is zero – as an estimate of the endogenous amount already in the tissue (this value is negative when endogenous signal is present);
3. subtracts it from every deposited amount, which shifts the x-axis so it expresses **total** amount present (endogenous + added);
4. drops the rows at the background level and refits on that shifted axis.

Value

The input object with cal_list (one lm per standard, response versus amount in pg per pixel), r2_df (fit R-squared per standard) and the calibration metadata populated.

Weighting

Calibration response in MSI is usually heteroscedastic – absolute scatter grows with amount – so an unweighted fit lets the top standards dominate and the low end, where most tissue pixels sit, is fitted worst. weighting therefore defaults to "1/x"; "1/x2" weights the low end harder still, and "none" is ordinary least squares.

A zero (blank) level has no 1/x weight. It is given the weight of the lowest positive level rather than a weight derived from a substituted tiny amount, which would let a single blank determine the whole regression.

Weighting is an empirical model choice, not a property of the data. Check it against residual structure, back-calculated accuracy at each level and replicate precision rather than against R-squared – see [calibrationDiagnostics\(\)](#) and [plotCalCoverage\(\)](#).

See Also

Other calibration: [calibrationInfo-class](#), [int2conc](#), [quant_MSImagingExperiment-method](#), [plotCalCoverage](#), [quant_MSImagingExperiment-method](#), [summariseCalLevels](#), [quant_MSImagingExperiment-me](#)

Examples

```
cal_dir <- system.file("extdata", "cal_example.raw", package = "quantMSImageR")
cal <- as(readRDS(file.path(cal_dir, "cal_MSI.RDS")),
          "quant_MSImagingExperiment")
cal_metadata <- read.csv(file.path(cal_dir, "calibration_metadata.csv"))
cal <- summariseCalLevels(cal, cal_metadata, val_slot = "intensity",
                        cal_label = "Cal", id = "identifier")
cal <- createCalCurve(cal, cal_type = "cal")
```

```
createMSIDataMatrix,quant_MSImagingExperiment-method
```

Create a feature-by-sample MSI data matrix

Description

Flattens the pixel-level object into tabular form for downstream statistics: one row per pixel, or one row per region of interest when roi_header names a grouping column.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
createMSIDataMatrix(
  MSIobject,
  val_slot = "intensity",
  inputNA = TRUE,
  roi_header = NA
)
```

Arguments

MSIobject	A quant_MSImagingExperiment object.
val_slot	Character. Spectra slot to tabulate (default "intensity").
inputNA	Logical. Convert zeros in the matrix to NA (default TRUE).
roi_header	Character. Column of pData() identifying regions of interest to average over. NA (the default) skips the ROI average and keeps one row per pixel.

Value

The input object with its tissueInfo slot populated: all_pixel_matrix (one row per pixel, one column per feature), roi_average_matrix (one row per ROI, when roi_header is given) and the accompanying sample/ROI metadata.

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- createMSIDataMatrix(obj, val_slot = "intensity", roi_header = NA)
```

generateTxtImages	<i>Load, process and optionally export per-feature text-image matrices</i>
-------------------	--

Description

Reads one or more DESI-MRM acquisitions, applies SNR filtering and tissue/background separation, and writes per-feature text matrices that can be imported by imaging software (e.g. MassLynx QuanOptimise overlay tools). Setting `output_txt = FALSE` runs all processing but skips file writing, which is useful when the caller only needs the processed MSI objects (e.g. for HTML report rendering).

Usage

```
generateTxtImages(
  fns,
  data_path,
  image_dir,
  lib_ion_path,
  snr_thresh = 0,
  thresh = 20,
  perc = 97,
  rot_clockwise = 0,
  average_method = "median",
  output_txt = TRUE,
  exclude = NULL,
  rename = NULL,
  ratios = NULL,
  output_ratios = TRUE,
  snr_overrides = NULL,
  is_name = NULL,
  is_norm_header = "IS_norm",
  is_mode = "line",
  is_window = 15,
  remove_IS = TRUE,
  type_header = "Type"
)
```

Arguments

<code>fns</code>	Character vector of acquisition names (without <code>.raw</code> suffix) for single-polarity studies. For dual-polarity studies pass a list where each element is either a plain string (single acquisition) or a named list with fields <code>pos</code> , <code>neg</code> , <code>label</code> , and optionally <code>prefix_pos</code> / <code>prefix_neg</code> (see bindPanels()). <code>runStudy.R</code> builds this list automatically from the YAML <code>samples</code> section.
<code>data_path</code>	Path to the folder that contains the <code>.raw</code> acquisition directories.
<code>image_dir</code>	Root output directory. One sub-directory per acquisition is created inside it.
<code>lib_ion_path</code>	Full path to the MRM ion-library CSV. Must contain columns: <code>transition_id</code> , <code>precursor_mz</code> , <code>product_mz</code> , <code>collision_eV</code> , <code>cone_V</code> , <code>Polarity</code> , <code>Type</code> .

snr_thresh	Numeric scalar or vector. One or more minimum signal-to-noise thresholds; pixels below each threshold are set to NA. When a vector is supplied, one complete output directory tree is created per threshold value. A value of 0 skips SNR filtering entirely <i>and</i> removes the requirement for a <code>tissue_pixels.csv</code> mask – handy for a smoke-test pass before ROIs are drawn (default 0).
thresh	Numeric. Cold-spot percentile passed to <code>imageR()</code> as threshold (default 20).
perc	Numeric. Hot-spot percentile passed to <code>imageR()</code> as percentile (default 97).
rot_clockwise	Integer (0-3). Number of 90 deg clockwise rotations applied via <code>pracma::rot90()</code> (default 0).
average_method	Character, "mean" or "median". Statistic used to summarise the noise pixel vector in <code>int2SNR()</code> . Applied consistently to every acquisition (default "median").
output_txt	Logical. When FALSE, processing runs but no files are written to disk (default TRUE).
exclude	Character vector of feature names to drop before processing. Names must match <code>fData()\$name</code> exactly (default NULL – keep all).
rename	Named character vector or list mapping old feature names to new display names, e.g. <code>c("old name" = "new name")</code> . Applied to all combined objects after loading; the new names appear in file names and the heatmap (default NULL – no renaming).
ratios	Optional list of metabolite ratio pairs. Each element must be a named list with fields: num Name of the numerator feature (matches <code>fData()\$name</code> after any rename overrides are applied). den Name of the denominator feature. label <i>(optional)</i> Output filename prefix. Defaults to " <code><num>_over_<den></code> ". Ratio images (numerator / denominator, pixel-wise) are written to the same SNR-filtered subdirectories as the individual ion images, immediately after them. Pixels where the denominator is zero or either ion is NA are set to NA (default NULL – no ratios).
output_ratios	Logical. When FALSE, ratio txt files are not written even if ratios pairs are defined. Has no effect when <code>output_txt</code> is FALSE (default TRUE).
snr_overrides	Optional named map of feature name to a feature-specific SNR threshold. Keys may be original ion-library names or post-rename display names. Listed features use their own threshold in every report; all other features use the global <code>snr_thresh</code> . Ignored for any report whose global threshold is 0 (the mask-free smoke-test pass). Default NULL (all features use <code>snr_thresh</code>).
is_name	Character. Value identifying the internal standard in <code>fData()\$feature_type</code> – that is, the ion library's Type column, typically "IS". It is not a transition name. When supplied, each acquisition is normalised with <code>int2response()</code> before SNR filtering, and every downstream step – SNR, background masking, images and ratios – works on the resulting response layer instead of intensity. NULL or "None" (the default) skips normalisation.
is_norm_header	Character. Ion-library column mapping each analyte to the standard that normalises it, needed only when <code>is_name</code> matches more than one feature (default "IS_norm"). See <code>int2response()</code> .
is_mode	Character. Level at which the internal standard is summarised: "line" (default), "sample", "pixel" or "window" (a rolling median over <code>is_window</code> consecutive pixels along the acquisition line). See <code>int2response()</code> .

is_window	Integer. Number of consecutive pixels averaged when is_mode = "window" (default 15).
remove_IS	Logical. Drop the internal-standard feature after normalising (default TRUE).
type_header	Character. Ion-library column holding the feature type, passed to readMRM() and matched by is_name (default "Type").

Value

Invisibly returns a named list:

combined Raw combined object (background pixels retained).

combined_snr SNR-filtered intensity for the **first** threshold in snr_thresh; sub-threshold pixels NA. Provided for backward compatibility.

combined_snr_list Named list of SNR-filtered objects, one per threshold in snr_thresh, named "snr<value>" (e.g. "snr3").

combined_NAbackground Raw intensity with background set to NA.

See Also

[int2SNR\(\)](#), [applySNR\(\)](#), [back2NA\(\)](#), [imageR\(\)](#)

Other workflow: [print.quantValidation\(\)](#), [runExample\(\)](#), [runStudy\(\)](#), [validateConfig\(\)](#)

Examples

```
# Section-mode run on the bundled synthetic data (no tissue mask needed at
# snr_thresh = 0), returning the processed objects without writing files.
fns <- list(list(neg = "example", section = "section01", label = "A"))
res <- generateTxtImages(
  fns      = fns,
  data_path = system.file("extdata", package = "quantMSImageR"),
  image_dir = tempfile(),
  lib_ion_path = system.file("extdata", "example_ion_library.csv",
                             package = "quantMSImageR"),
  snr_thresh = 0,
  output_txt = FALSE
)
```

imageR, quant_MSImagingExperiment-method

Draw an ion image for one feature

Description

Renders a single feature's spatial distribution as a ggplot. Because the result is an ordinary ggplot, the layout, theme and scales can be modified with the usual + syntax – for example + `facet_wrap(~ sample, ncol = 3)` to arrange several acquisitions in a grid.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
imageR(
  MSIobject,
  val_slot = "intensity",
  value = "response %",
  scale = "suppress",
  threshold = 1,
  sample_lab = "sample_ID",
  pixels = NA,
  percentile = 99,
  feat_ind = 1,
  perc_scale = FALSE,
  blank_back = TRUE,
  aspect_ratio = 1,
  text_image = FALSE,
  palette = c("heatmap0", "viridis")
)
```

Arguments

MSIobject	A quant_MSImagingExperiment object.
val_slot	Character. Spectra slot to image (default "intensity").
value	Character. Legend label describing what the values represent (e.g. "pg/mm2" for a calibrated layer).
scale	Character. Colour-scale treatment: "suppress" (cap at percentile), "histogram" (contrast-enhance per image) or "sqrt".
threshold	Numeric. Lowest percentile removed from the colour scale, to stop residual background dominating it (default 1).
sample_lab	Character. Column of pData(MSIobject) used to label and facet the images (default "sample_ID").
pixels	Character. Value of pData(MSIobject)\$sample_type selecting which pixels to plot; NA (the default) plots all.
percentile	Numeric. Percentile at which the colour scale is capped when scale = "suppress" (default 99).
feat_ind	Integer. Row index of the feature in fData(MSIobject) to image (default 1).
perc_scale	Logical. Express the colour scale as a percentage of the maximum (TRUE) rather than raw values (FALSE, the default).
blank_back	Logical; when TRUE background/zero pixels are drawn transparent.
aspect_ratio	numeric plot aspect ratio (default 1).
text_image	Logical; when TRUE return the image as a numeric matrix (for text export) rather than a ggplot.
palette	Character. Colour scale for the image: "heatmap0" (the default, matching the YAML colours: ion_image: default) or "viridis" (perceptually uniform and colour-blind safe). See quantPalettes() .

Value

A ggplot object, or a numeric matrix when text_image = TRUE.

See Also

Other visualisation: [contributionHm\(\)](#), [quantPalettes\(\)](#), [quantileHm\(\)](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
gg <- imageR(obj, feat_ind = 1, sample_lab = "run", scale = "suppress")
```

int2conc,quant_MSImagingExperiment-method

Convert response to calibrated amount estimates

Description

Inverts each analyte's calibration model to turn a measured response into an **estimated amount per pixel**. Features without a calibration model are dropped.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
int2conc(
  MSIobject,
  val_slot = "intensity",
  pixel_header = "sample_name",
  pixels = "tissue_pixels",
  max_out_of_range = 0.1
)
```

Arguments

MSIobject	A quant_MSImagingExperiment carrying calibration models fitted by createCalCurve() .
val_slot	Character. Spectra slot holding the measured response (default "intensity").
pixel_header	Character. Column of pData() holding the pixel-type labels (default "sample_name"). Pass pixel_header = "sample_type" for a calibration acquisition.
pixels	Character. Label(s) in pixel_header marking the pixels to quantify (default "tissue_pixels", the imaging convention). Pass pixels = "Tissue" for a calibration acquisition.
max_out_of_range	Numeric in [0, 1]. Largest proportion of a feature's pixels allowed to fall outside the calibrated range, i.e. to be extrapolated rather than interpolated (default 0.1, so at least 90% of pixels must be interpolated between real standards). Exceeding it is an error , naming the features and pointing at plotCalCoverage() : either the standards need to bracket the tissue, or the extrapolation has to be accepted explicitly by raising this value (1 accepts any). Any extrapolation at all is reported by message(); this controls only the threshold at which that becomes fatal.

Details

Mass spectrometry imaging measures analyte response at each pixel rather than analyte amount directly. The values produced here are calibrated *amount* estimates (pg per pixel, and pg per mm-squared where the pixel size is known), not volumetric concentrations. They are conditional on the calibration design, matrix matching, acquisition conditions and the validity of the fitted model – see [plotCalCoverage\(\)](#) for the coverage check that should accompany them.

Value

The object subset to pixels, with two spectra slots added: `pg_pixel` (estimated amount per pixel) and, when `experimentData(MSIObject)$pixelSize` is available, `pg_mm2` (estimated amount per unit area). Features with no calibration model are removed, and `r2_df` gains an `out_of_range` column giving the proportion of each feature's pixels that were extrapolated beyond the calibrated range.

Areal conversion

`pg_pixel` is inverted directly from the calibration model. `pg_mm2` is then derived from the pixel size recorded in the acquisition metadata:

$$pg/mm^2 = (pg/pixel) \times (1000/s)^2$$

where `s` is `experimentData(MSIObject)$pixelSize`, **expected in micrometres**, and the factor 1000 converts micrometres to millimetres. A single scalar is assumed, i.e. square pixels of side `s`; rectangular pixels are not currently supported, and supplying `s` in millimetres would inflate the result by a factor of 10^6 . When `pixelSize` is missing or not a positive number the `pg_mm2` slot is omitted and a warning is issued.

See Also

Other calibration: [calibrationInfo-class](#), [createCalCurve](#), [quant_MSImagingExperiment-method](#), [plotCalCoverage](#), [quant_MSImagingExperiment-method](#), [summariseCalLevels](#), [quant_MSImagingExperiment-me](#)

Examples

```
cal_dir <- system.file("extdata", "cal_example.raw", package = "quantMSImageR")
cal <- as(readRDS(file.path(cal_dir, "cal_MSI.RDS")),
         "quant_MSImagingExperiment")
cal_metadata <- read.csv(file.path(cal_dir, "calibration_metadata.csv"))
cal <- summariseCalLevels(cal, cal_metadata, val_slot = "intensity",
                        cal_label = "Cal", id = "identifier")
cal <- createCalCurve(cal, cal_type = "cal")

# A calibration acquisition labels its pixels in `sample_type`, so the
# imaging defaults are overridden here
cal <- int2conc(cal, pixel_header = "sample_type", pixels = "Tissue")
```

int2response,quant_MSImagingExperiment-method

Normalise pixel intensities to internal-standard response

Description

Divides each feature's intensity by an internal standard measured in the same pixel, line or sample, which suppresses drift and much of the local variation in ionisation efficiency.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
int2response(
  MSIobject,
  val_slot = "intensity",
  normalisation = c("internal_standard", "within_feature"),
  IS_name = NULL,
  is_norm_header = "IS_norm",
  mode = c("line", "sample", "pixel", "window"),
  window = 15,
  remove_IS = TRUE,
  ...
)
```

Arguments

MSIobject	A quant_MSImagingExperiment object.
val_slot	Character. Spectra slot to normalise (default "intensity").
normalisation	Character. "internal_standard" (default) or "within_feature". See Details.
IS_name	Character. Value identifying the internal standard in the feature-type column of fData(MSIobject) – the ion library's Type column, typically "IS". This is a type label, not a transition name: passing a feature name will not match. Required when normalisation = "internal_standard"; a value that matches no feature is an error, since falling back to a different normalisation would silently change the analytical method.
is_norm_header	Character. Feature-metadata column holding the analyte-to-standard mapping (default "IS_norm"). It is consulted only when IS_name matches more than one feature, so "None" (or NULL) is correct for the common case of a single standard shared by every analyte.
mode	Character. Level at which the standard is summarised: "line" Median across a whole acquisition line (the default). A <i>line</i> is one horizontal raster row – constant y, varying x – which is the order DESI acquires in, so it is also the axis along which source drift accumulates. "sample" Median across the whole acquisition. "pixel" The standard in that pixel alone – responsive to local suppression, but carries the standard's own shot noise.

	"window" Rolling median over window consecutive pixels along the same horizontal row, ordered by x. A compromise between "pixel" and "line": it smooths the standard's own shot noise while still tracking drift across the row. The window is truncated at the ends of a row rather than wrapping, so pixels from different rows are never mixed.
window	Integer. Number of consecutive pixels averaged when mode = "window" (default 15). Ignored for the other modes.
remove_IS	Logical. Drop the internal-standard features from the returned object (default TRUE).
...	Additional arguments (currently unused).

Details

normalisation states what is being done, so that no argument value has to be read as an instruction:

"internal_standard" Divide by the standard named in IS_name. This is the quantitative path.

"within_feature" Divide each feature by its own summary at the chosen mode. This removes between-line or between-sample scale differences within a feature, but it is *not* internal-standard normalisation and does not make features comparable to one another.

To leave values untouched, do not call this function. There is deliberately no "none" value here: skipping a step is the caller's decision, not a mode of the step. The YAML workflow exposes normalisation: none for that, and simply does not call this function.

Value

The input object with a response spectra slot holding the normalised values.

Multiple internal standards

When IS_name matches exactly one feature, every analyte is normalised to it. When it matches several – a panel carrying one class-specific standard per lipid class, for instance – an explicit analyte-to-standard mapping is required, because dividing by several standards at once has no defined meaning. Supply it as an ion-library column (named by is_norm_header, default "IS_norm") whose value on each analyte row is the transition name of the standard that normalises it. Any analyte left unmapped is an error rather than a silent fallback.

With a single standard the column is never read, so is_norm_header = "None" is the right setting for a panel where every analyte shares one standard.

See Also

Other filtering: [applySNR, quant_MSImagingExperiment-method](#), [back2NA, quant_MSImagingExperiment-method](#), [int2SNR, quant_MSImagingExperiment-method](#), [zero2NA, quant_MSImagingExperiment-method](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
# No internal standard in the example panel: normalise each feature to
# itself, per acquisition line.
obj <- int2response(obj, val_slot = "intensity",
                   normalisation = "within_feature")
```

int2SNR, quant_MSImagingExperiment-method

Calculate background-referenced signal-to-noise ratios

Description

Expresses each tissue pixel's response as a ratio to the same feature's response in the background pixels of the same acquisition.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
int2SNR(
  MSIobject,
  val_slot = "intensity",
  pixel_header = "sample_name",
  background = "background_pixels",
  tissue = "tissue_pixels",
  snr_thresh = 3,
  average = c("median", "mean"),
  snr_overrides = NULL,
  no_background = c("error", "all_na", "copy"),
  noise = NULL,
  sample_type = NULL,
  ...
)
```

Arguments

MSIobject	A quant_MSImagingExperiment whose pData() carries a background label in the pixel_header column.
val_slot	Character. Spectra slot holding the measured response (default "intensity"; use "response" after int2response()).
pixel_header	Character. Column of pData() holding the pixel-type labels (default "sample_name", the imaging convention).
background	Character. Value in pixel_header marking the background (non-tissue) pixels used to estimate the noise level (default "background_pixels"). The historical "noise_pixels" / "Noise" labels are matched too, so masks made with earlier versions keep working.
tissue	Character. Value in pixel_header marking the tissue pixels SNR is calculated for (default "tissue_pixels").
snr_thresh	Global minimum SNR to accept (below this value SNR = NA). Applied to every feature unless overridden by snr_overrides.
average	Character, "median" (default) or "mean": statistic used to summarise the background-pixel vector per feature.
snr_overrides	Optional named numeric vector mapping feature name (as in fData(MSIobject)\$name) to a feature-specific SNR threshold. A feature listed here uses its own threshold instead of snr_thresh; features not listed fall back to snr_thresh. Default NULL (all features use the global threshold).

no_background	Character. What to do when pixel_header contains no background pixels, so there is nothing to reference the signal against: "error" (default), "all_na" (create the slot, all NA) or "copy" (copy val_slot into the snr slot – note the slot then holds measured response, not a ratio). "error" and "all_na" apply only when snr_thresh > 0; with snr_thresh = 0 no filtering was requested, so the values are copied through regardless.
noise	Deprecated alias for background, kept for backward compatibility. When supplied it overrides background.
sample_type	Deprecated alias for pixel_header, kept for backward compatibility. When supplied it overrides pixel_header.
...	Additional arguments (currently unused).

Details

For feature f and tissue pixel p , with background-pixel set B :

$$SNR_{f,p} = I_{f,p} / \text{summary}_{b \in B}(I_{f,b})$$

where summary is the mean or median selected by average. This is a background-referenced signal ratio, not the classical analytical definition based on the standard deviation of the noise.

median is the default because a background region that clips part of the tissue, or contains a spot of carryover, shifts a mean far more than a median. Zero background values are treated as missing and imputed at one tenth of the smallest non-zero background value for that feature, so a feature whose background is entirely zero or NA is skipped and its snr stays NA.

Run this **after** internal-standard normalisation ([int2response\(\)](#)) when one is used. A threshold of 3 is a detection-level criterion; a stricter threshold is advisable before converting a feature to calibrated amounts.

Value

The input object with an additional snr spectra slot. Values below the selected threshold, and values outside the pixels labelled tissue, are stored as NA in that slot. No existing slot is modified.

See Also

Other filtering: [applySNR, quant_MSImagingExperiment-method](#), [back2NA, quant_MSImagingExperiment-method](#), [int2response, quant_MSImagingExperiment-method](#), [zero2NA, quant_MSImagingExperiment-method](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")

# The defaults match the imaging convention used throughout the package
obj <- int2SNR(obj, snr_thresh = 3)
names(spectraData(obj))
```

plotCalCoverage, quant_MSImagingExperiment-method

Check that measured pixels fall within the calibrated range

Description

Converting a response into a calibrated amount is only trustworthy where the standards actually constrain the fit. A calibration whose levels sit above or below the signal measured in tissue is extrapolation, not quantification. `int2conc()` reports the proportion of extrapolated pixels per feature, but without an explicit coverage check that is still easy to overlook – and the plot shows *where* the mismatch is, not just how much of it there is.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
plotCalCoverage(
  MSIobject,
  features = NULL,
  val_slot = "intensity",
  conc_slot = "pg_pixel",
  log_base = 2,
  bins = 40,
  cal_colour = "red",
  cal_alpha = 0.4,
  cal_linetype = "dotted",
  ...
)
```

Arguments

MSIobject	A quant_MSImagingExperiment that has been through <code>createCalCurve()</code> and <code>int2conc()</code> , so it carries both the fitted models (see <code>calibrationModels()</code>) and a calibrated-amount slot.
features	Features to plot: a character vector of names as in <code>fData(MSIobject)\$name</code> , or numeric row indices. Defaults to every feature that has a calibration model.
val_slot	Character. Spectra slot holding the measured response (default "intensity").
conc_slot	Character. Spectra slot holding the calibrated amount estimate written by <code>int2conc()</code> . Defaults to "pg_pixel"; objects calibrated with earlier versions carry "conc - pg/pixel", which is matched automatically.
log_base	Numeric. Base for both axes (default 2).
bins	Integer. Number of histogram bins for the pixel distribution (default 40).
cal_colour	Colour used for the standards and the level markers (default "red").
cal_alpha	Numeric. Opacity of the calibration-level marker lines (default 0.4), kept faint so they read as a reference grid rather than competing with the data.
cal_linetype	Line type for the calibration-level markers (default "dotted").
...	Additional arguments (currently unused).

Details

This function plots the two together on log axes so the overlap can be judged directly. The lower panel is the calibration itself: the standard spots and the fitted model that `int2conc()` inverts. The upper panel is the distribution of the object's own pixels on the same x-axis, with the calibration levels marked. When the histogram sits inside the span of those marks, every converted pixel is interpolated between real standards.

The standards are read back from each model's own model frame (`stats::model.frame`) rather than from `cal_response_data`. This matters for `cal_type = "std_addition"`, where `createCalCurve()` shifts the amount axis by the estimated endogenous amount before fitting: the stored response data is on the unshifted axis, so plotting it against the fitted line would put the points and the model on different scales. The model frame always holds the points the line was actually fitted to.

The fitted model is linear in amount, so on log axes it is drawn as a curve evaluated across the plotted range rather than as a straight line. Any part of it predicting a non-positive response has no log-scale counterpart and is dropped.

Value

A patchwork object: the pixel histogram stacked above the calibration curve, sharing an x-axis. Printing it draws the figure; it can also be further modified with `ggplot2` layers.

See Also

Other calibration: [calibrationInfo-class](#), [createCalCurve](#), [quant_MSImagingExperiment-method](#), [int2conc](#), [quant_MSImagingExperiment-method](#), [summariseCalLevels](#), [quant_MSImagingExperiment-method](#)

Examples

```
cal_dir <- system.file("extdata", "cal_example.raw", package = "quantMSImageR")
cal <- as(readRDS(file.path(cal_dir, "cal_MSI.RDS")),
         "quant_MSImagingExperiment")
cal_metadata <- read.csv(file.path(cal_dir, "calibration_metadata.csv"))
cal <- summariseCalLevels(cal, cal_metadata, val_slot = "intensity",
                        cal_label = "Cal", id = "identifier")
cal <- createCalCurve(cal, cal_type = "cal")
cal <- int2conc(cal, pixel_header = "sample_type", pixels = "Tissue")

plotCalCoverage(cal, val_slot = "intensity")
```

`print.quantValidation` *Print a validation result*

Description

Print a validation result

Usage

```
## S3 method for class 'quantValidation'
print(x, ...)
```

Arguments

x A quantValidation object.
 ... Ignored.

Value

x, invisibly.

See Also

Other workflow: [generateTxtImages\(\)](#), [runExample\(\)](#), [runStudy\(\)](#), [validateConfig\(\)](#)

quantileHm	<i>Quantile heatmap of MSI features across samples</i>
------------	--

Description

For each feature, computes the nth quantile of pixel intensities within each acquisition run, then z-scores the resulting per-feature profile across samples (clipped to [-1, 1]) and renders a `ComplexHeatmap::Heatmap`.

Usage

```
quantileHm(
  MSIobject,
  quant_val,
  heatmap_order = NA,
  heatmap_labs = NA,
  feature_split = NULL,
  feature_split_name = "Pathway",
  group_split_name = "Group",
  cell_size = 8,
  max_aspect = 1.5,
  cell_border = "white",
  fontsize = 8,
  palette = c("heatmap2", "heatmap0"),
  group_palette = "hat",
  feature_palette = "reading",
  row_split = NULL,
  row_split_name = NULL,
  column_split_name = NULL
)
```

Arguments

MSIobject A quant_MSIImagingExperiment object.
 quant_val Numeric in (0, 1). Quantile to summarise per feature per sample (e.g. 0.95 for the 95th percentile).
 heatmap_order Character vector of run names in the desired column order. Must match values in `pData(MSIobject)$run`. Defaults to NA (use discovery order).

heatmap_labs	Character vector of display labels, one per entry in heatmap_order. Used to create column-split groups. Defaults to NA (no splitting).
feature_split	Optional character/factor vector of length equal to the number of features, used to split the heatmap columns into labelled groups. Typically derived from an ion library metadata column (e.g. Met-1). Defaults to NULL (no splitting).
feature_split_name	Display name for the feature-group annotation legend (e.g. "Met-1", "Pathway"). Defaults to "Pathway".
group_split_name	Display name for the sample-group annotation legend (e.g. "Treatment", "Group"). Defaults to "Group".
cell_size	Numeric. Side of a heatmap cell, in millimetres (default 8). Giving the body an absolute size is what makes cells square; leaving it to ComplexHeatmap stretches them to fill the device, which produces very oblong cells when there are far more features than samples. Set to NA to restore the fill-the-device behaviour.
max_aspect	Numeric ≥ 1 . Largest cell width-to-height ratio allowed when one dimension has many more entries than the other (default 1.5). Cells stay square until the counts differ by more than four-fold; beyond that the shorter dimension's cells are widened up to this ratio so the plotting area is not reduced to a sliver.
cell_border	Colour for the line drawn around every cell (default "white"), which separates neighbouring cells instead of letting equal colours merge into one block. NA draws no border.
fontsize	Numeric. Point size for the sample and feature labels and the annotation names (default 8). ComplexHeatmap defaults to 12, which crowds the panel once feature names are long enough to need rotating.
palette	Character. Diverging colour ramp for the z-scores: "heatmap2" (default, blue-white-red) or "heatmap0". See quantPalettes() .
group_palette, feature_palette	Character. Qualitative palettes for the sample-group and feature-group colour bars (defaults "hat" and "reading"). Any grDevices::hcl.colors() palette name also works.
row_split, row_split_name, column_split_name	Deprecated aliases for feature_split, feature_split_name and group_split_name. They were named for the axis each grouping landed on before samples and features swapped axes; supplying them still works.

Details

Rows are samples and columns are features, in the order given by heatmap_order. Studies normally have more samples than features, so this puts the long dimension vertically where there is room for it and keeps sample names horizontally readable.

Value

A `ComplexHeatmap::Heatmap` object (rows = samples, columns = features), with an absolutely-sized body unless `cell_size` is NA.

What the colour means

Each feature is z-scored **on its own**, across samples: the column is centred on that feature's mean and divided by its own standard deviation. The colour therefore says where a sample sits *within that feature's* range, and the ramp is anchored at 0 so white is the feature's mean.

Colours are not comparable between features. Because every column is scaled by its own spread, a feature whose samples differ by a few percent fills the same blue-to-red range as one that doubles. Reading across a row to compare two features is the one thing this heatmap will not support. Use it to ask "which samples are high for this feature", not "which feature changed most" – for the latter, see the fold-change table, or `contributionHm()`, which puts every feature on one shared scale.

The z-scores are clipped to $[-1, 1]$ and the ramp spans exactly that, so any sample more than one standard deviation from its feature's mean is drawn at full intensity. With a handful of samples per group a clear separation saturates readily, which is deliberate – the panel is meant to show the pattern, not the magnitude – but it does mean two saturated cells can sit at very different z-scores.

A feature with no variance across samples has an undefined z-score. It is currently drawn at the bottom of the ramp rather than as missing, so a perfectly flat feature reads as uniformly low; treat an entirely single-coloured column with suspicion and check the underlying values.

Why there is no scale argument here

`contributionHm()` takes `scale = "shared"` or `"feature"`, choosing whether features can be compared with each other. This function has no such option, and the reason is structural rather than an omission.

Comparability needs the coloured quantity to still carry cross-feature information when the colour scale is applied. Here it does not. Every column is z-scored to mean 0 and standard deviation 1 *before* anything is drawn, so a feature whose samples differ by two percent and one that doubles arrive at the ramp with the same spread. The fixed $[-1, 1]$ limits are shared by every column, but sharing limits cannot restore what standardising already removed – the flattening happens first.

`contributionHm()` escapes this because it colours the group **mean** of those z-scores, which is not renormalised per feature: it sits near 0 when the groups do not separate and approaches 1 when they separate cleanly, and that range survives to the colour scale. A shared cap on it therefore means something.

So: this heatmap for one feature's profile across samples, and `contributionHm(scale = "shared")` when the question is which features separate the groups. Adding `scale` here would offer a comparison the arithmetic cannot support.

See Also

Other visualisation: `contributionHm()`, `imageR`, `quant_MSImagingExperiment-method`, `quantPalettes()`

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
  package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
hm <- quantileHm(obj, quant_val = 0.5,
  heatmap_order = "section01", heatmap_labs = "A")
```

Description

Extract and replace the contents of a `quant_MSImagingExperiment`'s `calibrationInfo` and `tissueInfo` slots. Prefer these to reaching into the slots with `@`: the internal representation may change, but these functions will not.

Usage

```
calibrationData(x)

## S4 method for signature 'quant_MSImagingExperiment'
calibrationData(x)

calibrationData(x) <- value

## S4 replacement method for signature 'quant_MSImagingExperiment'
calibrationData(x) <- value

calibrationModels(x)

## S4 method for signature 'quant_MSImagingExperiment'
calibrationModels(x)

calibrationModels(x) <- value

## S4 replacement method for signature 'quant_MSImagingExperiment'
calibrationModels(x) <- value

calibrationDiagnostics(x)

## S4 method for signature 'quant_MSImagingExperiment'
calibrationDiagnostics(x)

calibrationR2(x)

## S4 method for signature 'quant_MSImagingExperiment'
calibrationR2(x)

calibrationLevels(x)

## S4 method for signature 'quant_MSImagingExperiment'
calibrationLevels(x)

calibrationMetadata(x)

## S4 method for signature 'quant_MSImagingExperiment'
calibrationMetadata(x)

tissueData(x)

## S4 method for signature 'quant_MSImagingExperiment'
tissueData(x)
```

```
tissueMatrix(x, which = c("roi", "pixel"))

## S4 method for signature 'quant_MSImagingExperiment'
tissueMatrix(x, which = c("roi", "pixel"))
```

Arguments

<code>x</code>	A <code>quant_MSImagingExperiment</code> object.
<code>value</code>	Replacement value.
<code>which</code>	For <code>tissueMatrix()</code> , whether to return the per-ROI average matrix ("roi", the default) or the per-pixel matrix ("pixel").

Details

Note the naming: `calibrationInfo()` and `tissueInfo()` are the *class constructors*, so the slot accessors are `calibrationData()` and `tissueData()`.

The calibration accessors follow the workflow. `summariseCalLevels()` populates `calibrationLevels()`, `createCalCurve()` fills `calibrationModels()` and `calibrationR2()`, and `int2conc()` reads the models back out. `calibrationData()` moves the whole block at once, which is what carrying fitted models from a standards acquisition to a study object requires.

Value

`calibrationModels()` a named list of `lm` objects, one per calibrated analyte. `calibrationDiagnostics()` a data frame of fit quality, one row per analyte: feature, r2, and an `out_of_range` column once `int2conc()` has run. `calibrationR2()` the feature/r2 columns of that table only. `calibrationLevels()` the summarised response per calibration level. `calibrationMetadata()` the calibration design supplied by the user. `calibrationData()` the whole `calibrationInfo` object. `tissueMatrix()` a data frame of features by pixels or by ROI. The replacement forms return the modified object.

Examples

```
cal_dir <- system.file("extdata", "cal_example.raw", package = "quantMSImageR")
cal <- as(readRDS(file.path(cal_dir, "cal_MSI.RDS")),
         "quant_MSImagingExperiment")
cal_metadata <- read.csv(file.path(cal_dir, "calibration_metadata.csv"))

cal <- summariseCalLevels(cal, cal_metadata, val_slot = "intensity",
                        cal_label = "Cal", id = "identifier")
cal <- createCalCurve(cal, cal_type = "cal")

names(calibrationModels(cal))
calibrationDiagnostics(cal)
head(calibrationLevels(cal))

# Carry the fitted models onto a study acquisition
p <- system.file("extdata", "example.raw", "section01.RDS",
                package = "quantMSImageR")
study <- as(readRDS(p), "quant_MSImagingExperiment")
calibrationData(study) <- calibrationData(cal)
```

quantPalettes

*Colour palettes used by quantMSImageR***Description**

The palettes available to `imageR()`, `quantileHm()` and the metadata colour bars, returned as named character vectors of hex colours.

Usage

```
quantPalettes(name = NULL, n = NULL)
```

Arguments

name	Optional palette name. When NULL (default) the whole list is returned.
n	Optional number of colours. Qualitative palettes are truncated, continuous ones are interpolated. NULL (default) returns the palette at its native length.

Details

Four of these – heatmap0, heatmap2, hat and reading – are taken from the `ltc` package by Loukas Theodosiou (<https://github.com/loukesio/ltc-color-palettes>), which is MIT licensed. heatmap2 and hat are re-ordered relative to `ltc` – see below – and hat substitutes a grey for `ltc`'s black; the remaining colours are unchanged. They are reproduced here rather than depended on: `ltc` brings in 34 recursive dependencies, which is a large amount of installation surface for four colour vectors in a package that is otherwise light.

Which palette suits what:

`viridis` Sequential, perceptually uniform and colour-blind safe. The default for ion images, where the quantity is one-directional.

`heatmap0` Nine-colour sequential ramp running dark blue to teal to sand to red. An alternative for ion images.

`heatmap2` Five-colour diverging ramp, blue to white to red. Suits z-scores, where the midpoint is meaningful.

`hat`, `reading` Qualitative, for categorical metadata such as study group or pathway class. `hat` carries 10 well-separated hues, re-sequenced from `ltc`'s ordering so that the first few are maximally distinct, with `ltc`'s black replaced by a neutral grey placed last; `reading` is a softer 8-colour set.

Value

A named list of character vectors, or a single character vector of hex colours when name is given.

See Also

Other visualisation: `contributionHm()`, `imageR`, `quant_MSImagingExperiment-method`, `quantileHm()`

Examples

```
names(quantPalettes())
quantPalettes("heatmap2")
quantPalettes("hat", n = 3)
```

quant_MSImagingExperiment

Quantifiable MS imaging experiment

Description

Extends `Cardinal::MSImagingExperiment` with the metadata `quantMSImageR` needs to filter against background pixels and to convert a measured response into an estimated amount. Nearly every function in the package takes one of these, so a `Cardinal` object is normally coerced on the way in.

Details

The object is **features x pixels**: one row per MRM transition, one column per pixel. Derived layers are added as additional spectra slots rather than overwriting `intensity`, so the raw values survive the whole workflow:

`intensity` Raw measured response, as read from the acquisition.

`response` Internal-standard-normalised, added by [int2response\(\)](#).

`snr` Background-referenced signal-to-noise, added by [int2SNR\(\)](#).

`pg_pixel`, `pg_mm2` Calibrated amount estimates, added by [int2conc\(\)](#).

Inherited structure comes from `Cardinal`: `fData()` for feature metadata (m/z, name, transition m/z, ion-library columns), `pData()` for pixel metadata (x, y, run, and the tissue mask in `sample_name`) and `experimentData()` for acquisition metadata – `pixelSize` in particular, without which [int2conc\(\)](#) cannot produce `pg_mm2`.

Several acquisitions are held in a single object, distinguished by the `run` column of `pData()`; this is what [combineMSIs\(\)](#) produces and what the per-sample summaries group on.

Value

An object of class `quant_MSImagingExperiment`, extending `Cardinal::MSImagingExperiment` with `calibrationInfo` and `tissueInfo` slots.

Slots

`calibrationInfo` A [calibrationInfo](#) object.

`tissueInfo` A [tissueInfo](#) object.

See Also

[calibrationInfo](#), [tissueInfo](#), [quantMSImageR-accessors](#), [combineMSIs\(\)](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
  package = "quantMSImageR")

# Cardinal object in, quant_MSImagingExperiment out
obj <- as(readRDS(p), "quant_MSImagingExperiment")

dim(obj)           # features x pixels
names(spectraData(obj)) # available layers
head(as.data.frame(pData(obj)), 3)

# Derived layers accumulate alongside intensity
obj <- int2SNR(obj, snr_thresh = 3)
names(spectraData(obj))
```

readMRM

Read a Waters DESI-MRM acquisition into an MSImagingExperiment

Description

Reads one or more analyte text files from `<.raw>/imaging/`, matches transitions to the ion library on rounded (precursor, product) m/z , and constructs a [Cardinal::MSImagingExperiment](#). Handles both the single-file case (one combined Analyte 1.txt, the modern QuanOptimise default) and the multi-file case (older exports that split transitions across several Analyte N.txt files) transparently.

Usage

```
readMRM(
  name,
  folder,
  lib_ion_path,
  overwrite = TRUE,
  type_header = "Type",
  is_norm_header = "IS_norm",
  mz_tolerance = 0.4,
  ambiguity = c("combine", "error", "warn", "nearest")
)
```

Arguments

name	Acquisition name (without the .raw suffix).
folder	Path to the directory that contains the <name>.raw/ folder.
lib_ion_path	Full path to the ion-library CSV. Must contain columns transition_id, precursor_mz, product_mz, collision_eV, cone_V, Polarity, Type ("Analyte" for in-tissue analytes, "IS" for internal standards). Additional metadata columns are preserved by downstream code via buildFeatureMeta() . Measured transitions are matched to it on precursor and product m/z within <code>mz_tolerance</code> ; a transition the library does not describe is kept, named by its instrument index and typed "Unknown".

overwrite	Logical. When TRUE (default) the raw text files are re-parsed; when FALSE and a cached MSImagingExperiment.rds exists inside the .raw folder, it is returned instead.
type_header	Character. Name of the ion-library column holding the feature type – the values that mark internal standards versus analytes (default "Type"). Its contents become fData()\$feature_type, which is what <code>int2response()</code> matches IS_name against.
is_norm_header	Character. Name of the optional ion-library column that maps each analyte to the internal standard normalising it (default "IS_norm"). Carried through to fData() when present, so that <code>int2response()</code> can use it when a panel has more than one standard. "None" (or NULL), or simply omitting the column, is correct for a single-standard panel.
mz_tolerance	Numeric. Half-width, in Da, within which a measured precursor and product are taken to be the library's. Both must agree; a row matching on precursor alone is not a match. The default 0.4 is nominal-mass matching, because that is what the instrument does: a triple quadrupole running MRM selects Q1 and Q3 at unit resolution, so a product ion written as 308.1 and one written as 308.3 are the same measurement and no acquisition can separate them. Matching more tightly than the instrument resolves invents a distinction that is not in the data, and makes annotation depend on how many decimal places somebody typed into the library. The consequence is that genuinely isobaric compounds now collide – which is correct, and is what ambiguity = "combine" is for. Tighten it only for a high-resolution method where Q3 really is selective.
ambiguity	Character. What to do when a measured transition matches more than one library entry within mz_tolerance, which m/z alone cannot resolve. Isomers sharing a nominal precursor and product are the usual cause, and at unit resolution they are indistinguishable by definition. "combine" (default) names the feature for every entry it matched, joined with " " – "LTC4 14_15-LTC4". The joined name says the measurement is one of these, which is what the data supports; naming it for one of them asserts more than was measured. Annotation columns come from the closest entry. "error" refuses, which is right when a collision means the library is wrong rather than the chemistry ambiguous; "warn" takes the closest and says so; "nearest" takes the closest quietly. Two library entries for the <i>same</i> compound are redundancy, not ambiguity. "combine" reports those in its message so they can be removed from the library rather than carried as a joined name forever.

Details

Results are cached to <.raw>/MSImagingExperiment.rds; pass overwrite = TRUE (the default) to re-parse from the raw text files.

Value

An MSImagingExperiment with feature metadata joined to the ion library, ready to pass into `generateTxtImages()` or `selectTissuePixels()`.

See Also

Other acquisition: [removeBlankMzs,quant_MSImagingExperiment-method](#), [selectTissuePixels\(\)](#), [trimMSI\(\)](#)

Examples

```
# Return a previously parsed acquisition from its cached .rds
folder <- system.file("extdata", package = "quantMSImageR")
lib <- system.file("extdata", "ion_library_pos04.csv",
                  package = "quantMSImageR")
obj <- readMRM("pos04_test", folder = folder, lib_ion_path = lib,
              overwrite = FALSE)
```

removeBlankMzs,quant_MSImagingExperiment-method

Remove features without observed signal

Description

Drops any feature whose intensities are entirely zero or NA – typically a transition present in the acquisition method but never detected, or a panel row padded in when acquisitions were aligned. Reporting these alongside real measurements is how empty transitions reach a figure.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
removeBlankMzs(MSIobject)
```

Arguments

MSIobject A quant_MSImagingExperiment object.

Value

The input object with features carrying no data removed.

See Also

Other acquisition: [readMRM\(\)](#), [selectTissuePixels\(\)](#), [trimMSI\(\)](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- removeBlankMzs(obj)
```

runExample

Run the quantMSImageR example study

Description

Executes the full DESI-MRM workflow on a small synthetic dataset bundled with the package. The dataset contains two samples – SampleA (circular tissue, sections 01-03) and SampleB (square tissue, sections 04-06) – acquired on a panel of eight oxylipin features (seven analytes + one internal standard, all negative-ion mode).

Usage

```
runExample(
  render_report = TRUE,
  output_txt = FALSE,
  snr_thresh = 3,
  shapes = NULL,
  calibrate = TRUE
)
```

Arguments

render_report	Logical. Render and open the HTML report in the RStudio viewer or default browser? Requires pandoc. Default TRUE.
output_txt	Logical. Write per-feature .txt image matrices to a temporary directory? Default FALSE.
snr_thresh	Numeric. SNR threshold passed to generateTxtImages() . Default 3.
shapes	Character vector of tissue shapes to include – any of "circle", "square". NULL (the default) uses both, i.e. the full two-group example (group A circle + group B square).
calibrate	Logical. Also run the quantification demo on the bundled synthetic calibration standards (summariseCalLevels() -> createCalCurve() -> int2conc())? In a real study this step is driven by the YAML calibration: block. Default TRUE.

Details

Distinct tissue shapes per sample make the rendered ion images visually easy to distinguish, and the bundled ion library carries a Met-1 column (COX, LOX, IS) that drives the row-split colour bar in the report's main heatmap and the diagonal colour blocks in the correlation triangles.

All file paths are resolved automatically via [system.file\(\)](#), so the example works on any machine without editing a YAML.

Value

Invisibly returns the list produced by [generateTxtImages\(\)](#), with an added calibrated element when `calibrate = TRUE`: the study sections with their tissue pixels converted to `pg_pixel` and `pg_mm2` layers, restricted to the analytes that have a standard. When `render_report = TRUE` a report element gives the path to the rendered HTML.

See Also

Other workflow: `generateTxtImages()`, `print.quantValidation()`, `runStudy()`, `validateConfig()`

Examples

```
# The workflow itself, without rendering or opening the HTML report

res <- runExample(render_report = FALSE, shapes = "circle")
names(res)

# The full demo: renders the report and opens it in the viewer/browser
if (interactive()) {
  res <- runExample()
  res$report      # path to the rendered HTML
}
```

runStudy

Run a full DESI-MRM study from a YAML configuration

Description

Executes the whole workflow for a study described by a single YAML file: loads every acquisition, applies the tissue masks, runs SNR filtering at each requested threshold, optionally builds calibration models and converts tissue pixels to estimated amounts, writes per-feature .txt images, and renders one HTML report per SNR threshold.

Usage

```
runStudy(config_file)
```

Arguments

`config_file` Path to the study YAML configuration.

Details

The configuration format is documented by the template shipped with the package:

```
file.show(system.file("config_template.yaml", package = "quantMSImageR"))
```

A study can also be run from the command line, which uses this function internally:

```
Rscript -e 'quantMSImageR::runStudy("config.yaml")'
```

Value

Invisibly, the list produced by `generateTxtImages()` for the study, with a calibrated element added when the configuration's `calibration:` block is enabled. Called for its side effects: .txt images, the calibrated .RDS and the HTML reports written under the configured output paths.

See Also

Other workflow: [generateTxtImages\(\)](#), [print.quantValidation\(\)](#), [runExample\(\)](#), [validateConfig\(\)](#)

Examples

```
# A minimal study driven from a config, using the bundled synthetic data.
# snr_thresh = 0 skips SNR filtering and the tissue-mask requirement, and
# both outputs are switched off, so this runs without writing a report.

out <- file.path(tempdir(), "qmsi_study")
dir.create(out, showWarnings = FALSE)

cfg <- list(
  study = "example_study",
  paths = list(
    data_path = system.file("extdata", package = "quantMSImageR"),
    out_path = out,
    image_dir = file.path(out, "images"),
    lib_ion_path = system.file("extdata", "example_ion_library.csv",
                              package = "quantMSImageR")
  ),
  samples = list(
    list(neg = "example", sections = list("section01"),
        run_id = "S1", label = "A")
  ),
  parameters = list(snr_thresh = 0),
  output = list(render_report = FALSE, output_txt = FALSE)
)

cfg_file <- file.path(out, "config.yaml")
yaml::write_yaml(cfg, cfg_file)

res <- runStudy(cfg_file)
names(res)
```

selectTissuePixels	<i>Interactively select tissue pixels for an acquisition</i>
--------------------	--

Description

Loads a DESI-MRM acquisition, displays ion images of all features so the user can identify which best discriminates tissue from background, then opens an interactive ROI-selection window on a chosen feature. The resulting logical mask is saved as `tissue_pixels.csv` inside the acquisition's `.raw` folder, where [generateTxtImages\(\)](#) will find it automatically.

Usage

```
selectTissuePixels(
  name,
  data_path,
  lib_ion_path,
```

```

    feature = NULL,
    enhance = "histogram",
    overwrite = FALSE,
    preview = TRUE
  )

```

Arguments

name	Character. Acquisition name, without the .raw suffix (e.g. "14Dec_AntibodyStudy_1").
data_path	Character. Path to the folder that contains the .raw acquisition directory.
lib_ion_path	Character. Full path to the MRM ion-library CSV.
feature	Integer index or character name of the feature to use for tissue selection. When NULL (default), all features are displayed first and the user is prompted to choose one.
enhance	Character. Contrast enhancement passed to <code>Cardinal::image()</code> . Default "histogram".
overwrite	Logical. If TRUE, an existing <code>tissue_pixels.csv</code> will be overwritten. Default FALSE.
preview	Logical. Draw the saved mask (tissue versus background over the pixel coordinates) once selection finishes, so it can be checked before the workflow uses it. Default TRUE.

Details

This function must be run **before** the YAML workflow for each acquisition that does not yet have a `tissue_pixels.csv`. It requires an interactive R session with a graphics device (i.e. not inside `Rscript --no-save`).

Value

Invisibly returns a data frame with columns `x`, `y`, `tissue_pixels` and `background_pixels` (logical), one row per pixel, in acquisition order.

See Also

Other acquisition: [readMRM\(\)](#), [removeBlankMzs](#), [quant_MSImagingExperiment-method](#), [trimMSI\(\)](#)

Examples

```

# Requires a graphics device and user input, so it only runs interactively.
if (interactive()) {
  # Step 1 (once per acquisition, before running the study):
  selectTissuePixels(
    name      = "my_acquisition",
    data_path = "path/to/raw",
    lib_ion_path = "path/to/ion_library.csv"
  )

  # Step 2: run the full workflow from the study config
  runStudy("path/to/study.yaml")
}

```

stitchAcquisitions	<i>Stitch acquisitions that are pieces of one tissue into a single sample</i>
--------------------	---

Description

A tissue larger than the stage range is sometimes acquired in two passes – top then bottom – producing two `.raw` folders that are *one* sample. This concatenates their pixels into a single run.

Usage

```
stitchAcquisitions(..., label = NULL)
```

Arguments

<code>...</code>	Two or more <code>MSImagingExperiment</code> objects, or a single list of them. All must carry the same features and the same spectra layers.
<code>label</code>	Character. Run identifier for the stitched sample. Defaults to the first object's run.

Details

It is not the same operation as either `neighbour`, and picking the wrong one is quiet rather than loud:

`bindPanels()` two MRM *panels* over one tissue: different transitions, shared pixel grid. It **intersects** pixels on (x, y), so given two spatial halves it keeps only coordinates present in both – nothing, if the halves do not overlap.

`combineMSIs()` several *samples* into one object, each keeping its own run identifier. It refuses duplicate run identifiers, precisely to stop two sections silently becoming one sample.

`stitchAcquisitions()` several *pieces of one sample* into one run. The union of pixels, one run identifier.

Why this matters beyond the picture: two halves of one tissue are not two replicates. Left as separate samples they count twice in every group summary – the group mean, the box/violin plots, any downstream model – and because the halves agree with each other they look like a reproducible effect rather than one measurement.

Value

One object holding the union of the input pixels, with run set to `label` throughout.

Coordinates

No offset is applied. Waters records absolute stage coordinates, so the pieces already sit in a common frame and shifting them would move the tissue apart. Pixels appearing at the same (x, y) in more than one piece are therefore an error, not something to silently average: it means the acquisitions genuinely overlap, or that they are not pieces of one tissue.

See Also

`bindPanels()`, `combineMSIs()`

Other combining acquisitions: `alignFeatures()`, `bindPanels()`, `combineMSIs`, `MSImagingExperiment-method`

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- readRDS(p)
# Split by pixel, then stitch back: one sample, all pixels, one run.
half1 <- obj[, 1:50]
half2 <- obj[, 51:ncol(obj)]
whole <- stitchAcquisitions(half1, half2, label = "section01")
ncol(whole) == ncol(obj)
```

summariseCalLevels,quant_MSImagingExperiment-method

Summarise the response at each calibration level

Description

Averages the signal across the pixels of each calibration spot, giving one response value per standard per level. This is the input [createCalCurve\(\)](#) fits its models to.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
summariseCalLevels(
  MSIobject,
  cal_metadata,
  val_slot = "intensity",
  cal_header = "sample_type",
  cal_label = "Cal",
  id = "identifier"
)
```

Arguments

MSIobject	A quant_MSImagingExperiment whose pData() carries the calibration labels and spot identifiers.
cal_metadata	Data frame of calibration metadata, with columns analyte (feature name as in fData()\$name), identifier (maps to pData()), amount_pg (amount of standard deposited at each spot) and level (dilution level). A legacy lipid column is read as analyte.
val_slot	Character. Spectra slot to summarise (default "intensity").
cal_header	Character. Column of pData() used to select calibration pixels (default "sample_type").
cal_label	Character. Value of cal_header marking calibration pixels (default "Cal").
id	Character. Column present in both cal_metadata and pData() that identifies a unique calibration spot (default "identifier").

Details

Every spot is also summarised for dispersion, because a mean and a pixel count cannot show whether a calibration level was measured reproducibly. The returned table carries `sd_response` and `cv_response` alongside the mean, so an imprecise level is visible rather than averaged away.

Problems in the calibration design are errors rather than quiet NAs: an analyte that is not in the object, an identifier that matches no pixel, a spot with no usable response, and duplicated identifiers that disagree about amount or level all stop the function. Each of these would otherwise produce a curve that looks fitted but is not anchored to the data it claims.

Value

The input object with `cal_response_data` populated: one row per (standard, calibration spot) pair holding `response_perpixel` (the mean), `median_response`, `sd_response`, `cv_response`, `pixels`, `n_nonmissing` and `pg_perpixel`. Amounts are expressed in **pg per pixel**.

See Also

Other calibration: [calibrationInfo-class](#), [createCalCurve](#), [quant_MSImagingExperiment-method](#), [int2conc](#), [quant_MSImagingExperiment-method](#), [plotCalCoverage](#), [quant_MSImagingExperiment-method](#)

Examples

```
cal_dir <- system.file("extdata", "cal_example.raw", package = "quantMSImageR")
cal <- as(readRDS(file.path(cal_dir, "cal_MSI.RDS")),
         "quant_MSImagingExperiment")
cal_metadata <- read.csv(file.path(cal_dir, "calibration_metadata.csv"))
cal <- summariseCalLevels(cal, cal_metadata, val_slot = "intensity",
                        cal_label = "Cal", id = "identifier")
```

tissueInfo-class

Tissue-level summaries for an imaging experiment

Description

Holds the tabular views of an imaging experiment produced by [createMSIDataMatrix\(\)](#): one row per pixel, and optionally one row per region of interest, together with the accompanying metadata. It is carried in the `tissueInfo` slot of a [quant_MSImagingExperiment](#).

Usage

```
## S4 method for signature 'tissueInfo'
show(object)
```

Arguments

`object` A `tissueInfo` object.

Details

`all_pixel_matrix` `data.frame`, one row per pixel and one column per feature.

`roi_average_matrix` `data.frame`, one row per region of interest, populated only when `createMSIDataMatrix(roi_h)` is given.

`sample_metadata` `data.frame` describing the rows of the matrices above (sample, run, ROI identifier).

`feature_metadata` `data.frame` describing their columns.

All four are zero-row `data.frames` until `createMSIDataMatrix()` has run. Access them with `tissueMatrix()` and `tissueData()` rather than `@`.

Value

An object of class `tissueInfo`.

Slots

`roi_average_matrix` `data.frame` of per-ROI averages.

`all_pixel_matrix` `data.frame` of per-pixel values.

`sample_metadata` `data.frame` describing the matrix rows.

`feature_metadata` `data.frame` describing the matrix columns.

See Also

[quant_MSImagingExperiment](#), [createMSIDataMatrix\(\)](#), [quantMSImageR-accessors](#)

Examples

```
ti <- tissueInfo()

p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- createMSIDataMatrix(obj, val_slot = "intensity", roi_header = NA)
dim(tissueMatrix(obj, which = "pixel"))
```

trimMSI

Remove pure-background border rows and columns from an MSI object

Description

Drops any x-column or y-row in which **every** pixel is labelled as "background_pixels" (or the historical "noise_pixels") in `pData(MSI_data)$sample_name`. This trims empty scan borders that arise from the Waters acquisition geometry without affecting tissue or mixed-content rows/columns.

Usage

```
trimMSI(MSI_data)
```

Arguments

MSI_data A quant_MSImagingExperiment object whose pData() contains columns x, y, and sample_name (populated by makeFactor()).

Value

A subset quant_MSImagingExperiment with pure-background border pixels removed.

See Also

Other acquisition: [readMRM\(\)](#), [removeBlankMzs](#), [quant_MSImagingExperiment-method](#), [selectTissuePixels\(\)](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImager")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- trimMSI(obj)
```

validateConfig

Validate a study YAML configuration

Description

Structural and cross-file checks on a study config before anything is processed: required blocks present, referenced files and ion-library columns actually exist, enumerated values spelled correctly, and the calibration and internal-standard settings coherent with each other.

Usage

```
validateConfig(config, check_paths = TRUE)
```

Arguments

config A config list, or a path to a study YAML file.

check_paths Logical. Also check that the referenced data directory, ion library and calibration files exist on disk (default TRUE). Set FALSE to validate a config's structure away from its data.

Details

Every check runs, so one call reports every problem rather than stopping at the first. [runStudy\(\)](#) calls this first and refuses to start if there are errors – these are mistakes that would otherwise surface as an obscure failure halfway through a long run, or worse, as a plausible-looking result computed from the wrong column.

Value

A quantValidation object with errors, warnings and a summary data frame of every check.

See Also

Other workflow: [generateTxtImages\(\)](#), [print.quantValidation\(\)](#), [runExample\(\)](#), [runStudy\(\)](#)

Examples

```
# The shipped template is structurally valid; its placeholder paths are not
# meant to exist, so the path checks are skipped here.
cfg <- system.file("config_template.yaml", package = "quantMSImageR")
validateConfig(cfg, check_paths = FALSE)
```

```
zero2NA,quant_MSImagingExperiment-method
```

Replace zero intensities with NA

Description

Zeros in an MRM acquisition mean "nothing recorded" rather than "measured as zero", and left in place they bias means and compress colour scales. This converts them to NA so they are excluded from summaries.

Usage

```
## S4 method for signature 'quant_MSImagingExperiment'
zero2NA(MSIobject, val_slot = "intensity")
```

Arguments

MSIobject	A quant_MSImagingExperiment object.
val_slot	Character. Spectra slot to modify (default "intensity").

Value

The input object with zero values in val_slot replaced by NA.

Destructive

This modifies val_slot in place rather than adding a new layer. Keep a copy of the object if the original values are needed afterwards.

See Also

Other filtering: [applySNR,quant_MSImagingExperiment-method](#), [back2NA,quant_MSImagingExperiment-method](#), [int2SNR,quant_MSImagingExperiment-method](#), [int2response,quant_MSImagingExperiment-method](#)

Examples

```
p <- system.file("extdata", "example.raw", "section01.RDS",
                 package = "quantMSImageR")
obj <- as(readRDS(p), "quant_MSImagingExperiment")
obj <- zero2NA(obj, val_slot = "intensity")
```

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