

Harmonization Function and Parameter Errors

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Data loading

In this document we harmonize calibration curves between two matrices:

- **SU** (reference matrix)
- **POOL** (target matrix to be scaled and shifted)

We work with two analytes: *anisole* and *heptanone*.

```
## 'data.frame':    28 obs. of  2 variables:  
##   $ concentration: num  10.326 30.326 0.326 20.326 5.326 ...  
##   $ intensity     : num  1331.8 2088.8 44.6 1862 885.7 ...  
  
## 'data.frame':    28 obs. of  2 variables:  
##   $ concentration: num  10 30 0 20 5 10 30 0 20 5 ...  
##   $ intensity     : num  1013 1593.2 21.9 1462.2 642.6 ...
```

Harmonization function

Methodology

The objective is to determine the **scale** and **shift** parameters that optimally transform the SU calibration curve so that it aligns with the POOL measurements.

The procedure is:

1. Fit a **third-degree polynomial model** to the SU calibration data (no intercept).
2. Evaluate this polynomial on a dense concentration grid to obtain a smooth SU curve.
3. For each candidate pair $(\text{scale}, \text{shift})$ in a predefined grid:
 - Transform the SU concentration axis using

$$C \mapsto \frac{C}{\text{scale}} - \text{shift}$$

* Interpolate the transformed SU curve onto the POOL concentration points. * Compute a loss value (with optional variance normalisation). 4. Select the $(\text{scale}, \text{shift})$ pair that **minimises the loss**. 5. Fit a **local quadratic surface (paraboloid)** to the loss landscape around the optimum to estimate parameter uncertainties.

Quadratic model (Paraboloid)

Locally around the optimum $((s_0, t_0))$, the loss function is approximated by a general quadratic form:

$$L(s, t) \approx L_{\min} + a(s - s_0)^2 + b(t - t_0)^2 + c(s - s_0)(t - t_0).$$

Vector form:

$$L(s, t) - L_{\min} \approx \mathbf{x}^T M \mathbf{x}, \quad \mathbf{x} = \begin{pmatrix} s - s_0 \\ t - t_0 \end{pmatrix}.$$

where

$$M = \begin{pmatrix} a & c/2 \\ c/2 & b \end{pmatrix}.$$

This matrix M is **half of the Hessian** of the loss.

Ellipse of constant loss

Parameter uncertainties are obtained from the contour:

$$L(s, t) = 2L_{\min},$$

which leads to the ellipse:

$$\mathbf{x}^T M \mathbf{x} = L_{\min}.$$

Diagonalising $M = V \Lambda V^T$:

- the eigenvalues give the curvature,
- the eigenvectors give the orientation,
- the semi-axes are:

$$\text{semi-axis}_i = \sqrt{\frac{L_{\min}}{\lambda_i}}.$$

Projection onto scale and shift axes

The ellipse describes the joint uncertainty in ((s,t)). However, we often want **individual uncertainties** for each parameter.

Let:

- ($V = [v_1; v_2]$) be the eigenvector matrix,
- (a_1, a_2) be the semi-axes.

Any point on the ellipse can be written as:

$$\mathbf{x} = a_1 v_1 \cos \theta + a_2 v_2 \sin \theta.$$

To obtain the **uncertainty in scale**, we project the ellipse onto the horizontal axis:

$$\Delta s = \max_{\theta} |x_1| = \sqrt{(a_1 v_{1,1})^2 + (a_2 v_{2,1})^2}.$$

Likewise, for **shift**:

$$\Delta t = \sqrt{(a_1 v_{1,2})^2 + (a_2 v_{2,2})^2}.$$

These projected half-widths are reported as

$$\text{scale} \pm \Delta s, \quad \text{shift} \pm \Delta t.$$

Function definition

Principal Function

```
harmonize <- function(df_target, df_sub,
                       scale_range = c(0.2, 2),
                       shift_range = c(0, 5),
                       n_scale = 200,
                       n_shift = 200,
                       n_grid = 200,
                       min_fraction = 0.5,
                       optimize_shift = TRUE,
                       local_scale_win = 0.1,
                       local_shift_win = 1) {

  # 1. Validate & prepare input
  x <- check_and_prepare_input(df_target, df_sub)
  df_target <- x$df_target
  df_sub    <- x$df_sub

  # 2. Fit SU model
  model_su <- fit_su_model(df_sub)

  # 3. Grid SU
```

```

su_grid <- make_su_grid(df_sub, n_grid, model_su)

# 4. Variance normalisation
vn <- compute_variance_normalisation(df_target)
df_target <- vn$df_target

# 5. Grid search
scale_seq <- seq(scale_range[1], scale_range[2], length.out = n_scale)
shift_seq <- if (optimize_shift) seq(shift_range[1], shift_range[2], length.out = n_shift) else 0

loss_grid <- compute_loss_grid(df_target, su_grid, scale_seq, shift_seq, min_fraction)

best <- find_best_parameters(loss_grid)

best_pred <- predict_best_curve(su_grid, df_target, best$best_scale, best$best_shift)

# 6. Local paraboloid with Hessiana
parab <- fit_local_paraboloid(
  loss_grid,
  best_scale = best$best_scale,
  best_shift = best$best_shift,
  best_loss = best$best_loss,
  local_scale_win = local_scale_win,
  local_shift_win = local_shift_win
)

# Output structure
list(
  scale = best$best_scale,
  shift = best$best_shift,
  loss = best$best_loss,

  scale_error = parab$scale_error,
  shift_error = parab$shift_error,
  loss_threshold = parab$loss_threshold,
  paraboloid = parab,

  fitted_points = df_target %>%
    dplyr::mutate(intensity_pred = best_pred),

  su_grid = su_grid,
  su_grid_trans = tibble::tibble(
    concentration = su_grid$concentration / best$best_scale - best$best_shift,
    intensity = su_grid$intensity
  ),
  loss_grid = loss_grid,
  variance_norm = vn$use_var_norm,
  min_fraction = min_fraction,
  optimize_shift = optimize_shift
)
}

```

Auxiliar Functions

```
# AUXILIAR FUNCTIONS

# VALIDATE AND PREPARE INPUT
check_and_prepare_input <- function(df_target, df_sub) {

  if (!all(c("concentration", "intensity") %in% colnames(df_target)))
    stop("df_target must contain columns 'concentration' and 'intensity'.")

  if (!all(c("concentration", "intensity") %in% colnames(df_sub)))
    stop("df_sub must contain columns 'concentration' and 'intensity'.")

  df_target <- df_target %>%
    dplyr::mutate(
      concentration = as.numeric(concentration),
      intensity     = as.numeric(intensity)
    )

  df_sub <- df_sub %>%
    dplyr::mutate(
      concentration = as.numeric(concentration),
      intensity     = as.numeric(intensity)
    )

  list(df_target = df_target, df_sub = df_sub)
}

# FIT SU MODEL AND PREDICT

fit_su_model <- function(df_sub) {
  lm(intensity ~ poly(concentration, 3, raw = TRUE) - 1, data = df_sub)
}

predict_su <- function(model_su, C) {
  cf <- coef(model_su)
  cf[1] * C + cf[2] * C^2 + cf[3] * C^3
}

# MAKE SU GRID (sample the polynomial model)
make_su_grid <- function(df_sub, n_grid, model_su) {
  C_grid <- seq(0, max(df_sub$concentration, na.rm = TRUE), length.out = n_grid)
  I_grid <- predict_su(model_su, C_grid)
  tibble::tibble(concentration = C_grid, intensity = I_grid)
}

# CHOOSE IF VARIANCE NORMALIZATION IS APPLIED IN THE LOSS

compute_variance_normalisation <- function(df_target) {

  df_var <- df_target %>%
    dplyr::group_by(concentration) %>%
    dplyr::summarise(
      n           = dplyr::n(),
      n_sq        = n^2,
      n_sq_minus_1 = n_sq - 1,
      mean        = mean(intensity),
      variance   = sum((intensity - mean)^2) / n_sq_minus_1
    )
}
```

```

    var_intensity = if (n() > 1) var(intensity) else NA_real_,
    .groups       = "drop"
  )

use_var_norm <- all(!is.na(df_var$var_intensity)) &&
  all(df_var$var_intensity > 0) &&
  all(df_var$n >= 2)

df_target <- df_target %>%
  dplyr::left_join(df_var, by = "concentration") %>%
  dplyr::mutate(denom = ifelse(use_var_norm, var_intensity, 1))

list(df_target = df_target, use_var_norm = use_var_norm)
}

# LOSS GRID COMPUTATION

compute_loss_grid <- function(df_target, su_grid, scale_seq, shift_seq, min_fraction) {

  C_t <- df_target$concentration
  I_t <- df_target$intensity
  D_t <- df_target$denom
  n_total <- length(C_t)

  C_grid <- su_grid$concentration
  I_grid <- su_grid$intensity

  loss_grid <- expand.grid(scale = scale_seq, shift = shift_seq)
  loss_grid$loss <- NA_real_

  for (i in seq_len(nrow(loss_grid))) {

    sc <- loss_grid$scale[i]
    sh <- loss_grid$shift[i]

    C_trans <- C_grid / sc - sh

    # Strict interpolation (no extrapolation)
    I_pred <- approx(x = C_trans, y = I_grid, xout = C_t, rule = 1)$y

    # Check the fraction of points within range
    valid <- !is.na(I_pred)

    # Not enough coverage + reject this (scale, shift)
    if (sum(valid) < min_fraction * n_total) {
      loss_grid$loss[i] <- Inf
      next
    }

    # Linear extrapolation only above the SU range
    above <- which(is.na(I_pred) & C_t > max(C_trans))
    if (length(above) > 0) {

```

```

x1 <- C_trans[length(C_trans) - 1]
x2 <- C_trans[length(C_trans)]
y1 <- I_grid[length(I_grid) - 1]
y2 <- I_grid[length(I_grid)]
slope <- (y2 - y1) / (x2 - x1)
I_pred[above] <- y2 + slope * (C_t[above] - x2)
}

# Compute loss using all points
res_sq <- (I_t - I_pred)^2
loss_grid$loss[i] <- mean(res_sq / D_t)
}

# Identify non-covered region and report approximate boundary line
bad <- loss_grid[!is.finite(loss_grid$loss), , drop = FALSE]

if (nrow(bad) > 2) {

  fit_line <- lm(shift ~ scale, data = bad)
  b0 <- coef(fit_line)[1]    # intercept
  b1 <- coef(fit_line)[2]    # slope

  # Linear boundary in: a*shift + b*scale >= c
  a <- 1
  b <- -b1
  c <- b0

  cat(
    "Warning: insufficient SU coverage in part of the search grid.\n",
    "Approximate boundary of the uncovered region:\n",
    "  ", a, " * shift + ",
    "  b, " * scale >= ",
    "  c, '\n',
    "Points above this boundary typically fail the minimum coverage criterion.\n",
    "Consider avoiding this region or reducing the 'min_fraction' parameter."
  )
}

loss_grid
}

# OPTIMAL PARAMETERS SELECTION

find_best_parameters <- function(loss_grid) {

  finite <- which(is.finite(loss_grid$loss))

  if (length(finite) == 0) {
    return(list(
      best_scale = NA_real_,
      best_shift = NA_real_,

```

```

        best_loss  = Inf
    ))
}

idx <- finite[which.min(loss_grid$loss[finite])]

list(
  best_scale = loss_grid$scale[idx],
  best_shift = loss_grid$shift[idx],
  best_loss  = loss_grid$loss[idx]
)
}

# OPTIMAL CURVE PREDICTION

predict_best_curve <- function(su_grid, df_target, best_scale, best_shift) {

  C_grid <- su_grid$concentration
  I_grid <- su_grid$intensity
  C_t     <- df_target$concentration

  C_trans <- C_grid / best_scale - best_shift
  approx(x = C_trans, y = I_grid, xout = C_t, rule = 1)$y
}

# PARABOLOID AND HESSIAN MATRIX

fit_local_paraboloid <- function(loss_grid, best_scale, best_shift,
                                   best_loss, local_scale_win, local_shift_win) {

  local <- loss_grid %>%
    dplyr::filter(
      is.finite(loss),
      scale >= best_scale - local_scale_win,
      scale <= best_scale + local_scale_win,
      shift >= best_shift - local_shift_win,
      shift <= best_shift + local_shift_win
    )

  if (nrow(local) < 10) return(NULL)

  x1 <- local$scale - best_scale
  x2 <- local$shift - best_shift
  y   <- local$loss - best_loss

  model <- try(
    lm(y ~ I(x1^2) + I(x2^2) + I(x1*x2) - 1,
       silent = TRUE
    )
  )

  if (inherits(model, "try-error")) return(NULL)
}

```

```

cf <- coef(model)

a <- cf["I(x1^2)"]
b <- cf["I(x2^2)"]
c <- cf["I(x1 * x2)"]

M <- matrix(c(a, c/2, c/2, b), nrow = 2, byrow = TRUE)

eig <- eigen(M)
lambda <- eig$values
V       <- eig$vectors

if (any(lambda <= 0) || best_loss <= 0) {
  return(list(a=a, b=b, c=c, M=M, eigenvalues=lambda,
              eigenvectors=V, semiaxes=NA,
              scale_error=NA, shift_error=NA))
}

semiaxes <- sqrt(best_loss / lambda)

v1 <- V[,1]
v2 <- V[,2]

proj_scale <- sqrt((semiaxes[1] * v1[1])^2 + (semiaxes[2] * v2[1])^2)
proj_shift <- sqrt((semiaxes[1] * v1[2])^2 + (semiaxes[2] * v2[2])^2)

list(
  a=a, b=b, c=c,
  M=M,
  eigenvalues=lambda,
  eigenvectors=V,
  semiaxes=semiaxes,
  scale_error=proj_scale,
  shift_error=proj_shift,
  loss_threshold = 2 * best_loss
)
}

```

Visualization of harmonized curves

We now define a simple plotting function to compare:

- POOL original points
- SU original curve
- SU transformed curve (using the optimized scale and shift)

```

plot_harmonization <- function(h, title = "Calibration Curve Harmonization") {

  # POOL original points
  pool_orig <- h$fitted_points %>%
    dplyr::transmute(
      concentration = concentration,
      intensity     = intensity,
      type          = "POOL original"
}

```

```

)
# SU original curve
su_orig <- h$su_grid %>%
  dplyr::mutate(type = "SU original")

# SU transformed curve
su_trans <- h$su_grid_trans %>%
  dplyr::mutate(type = "SU transformed")

combined_df <- dplyr::bind_rows(
  pool_orig,
  su_orig,
  su_trans
)

ggplot(
  combined_df,
  aes(x = concentration, y = intensity, color = type, linetype = type)
) +
  # POOL points
  geom_point(
    data = dplyr::filter(combined_df, type == "POOL original"),
    size = 2.2,
    alpha = 0.9
  ) +
  # SU curves
  geom_line(
    data = dplyr::filter(combined_df, type != "POOL original"),
    linewidth = 0.8
  ) +
  scale_color_manual(values = c(
    "POOL original" = "darkorange2",
    "SU transformed" = "navy",
    "SU original" = "grey40"
  )) +
  scale_linetype_manual(values = c(
    "POOL original" = "blank",
    "SU transformed" = "solid",
    "SU original" = "dashed"
  )) +
  labs(
    title = title,
    x = "Concentration (ppb)",
    y = "Intensity (a.u.)",
    color = "Series",
    linetype = "Series"
  ) +
  theme_minimal(base_size = 14)
}

```

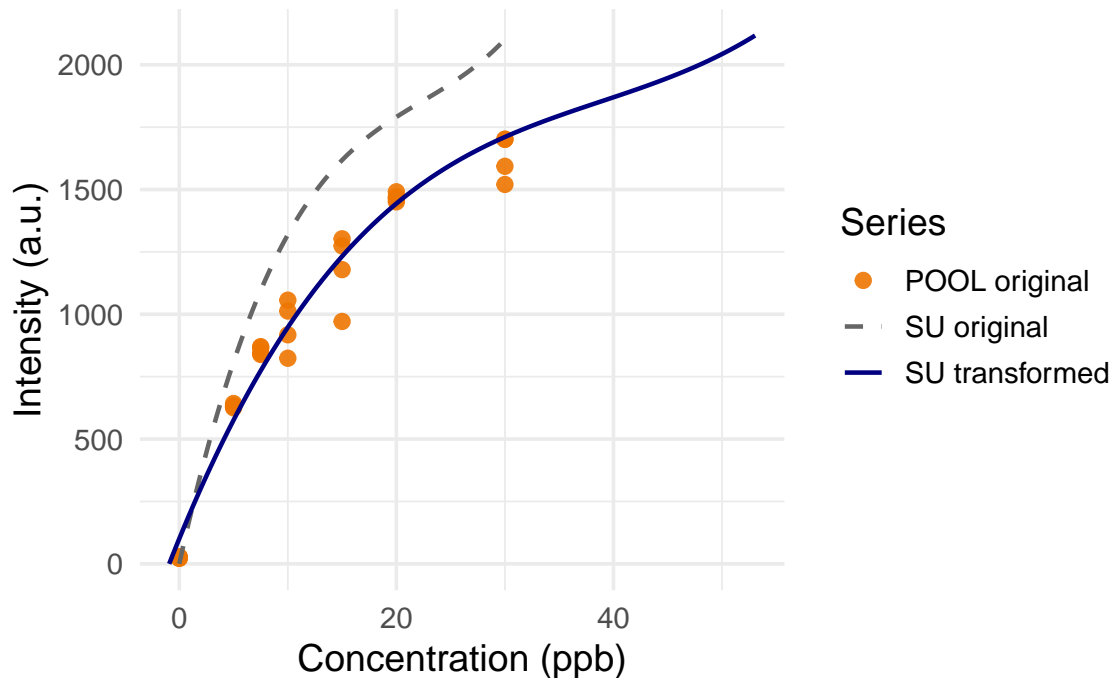
Anisole

```
res_anisole <- harmonize(pool_anisole, su_anisole)
```

Table 1: Harmonization parameters and uncertainty (Anisole)

Parameter	Value	Error	Units
Scale	0.562	± 0.124	–
Shift	0.930	± 2.023	ppb

Calibration Harmonization – Anisole



Heptanone

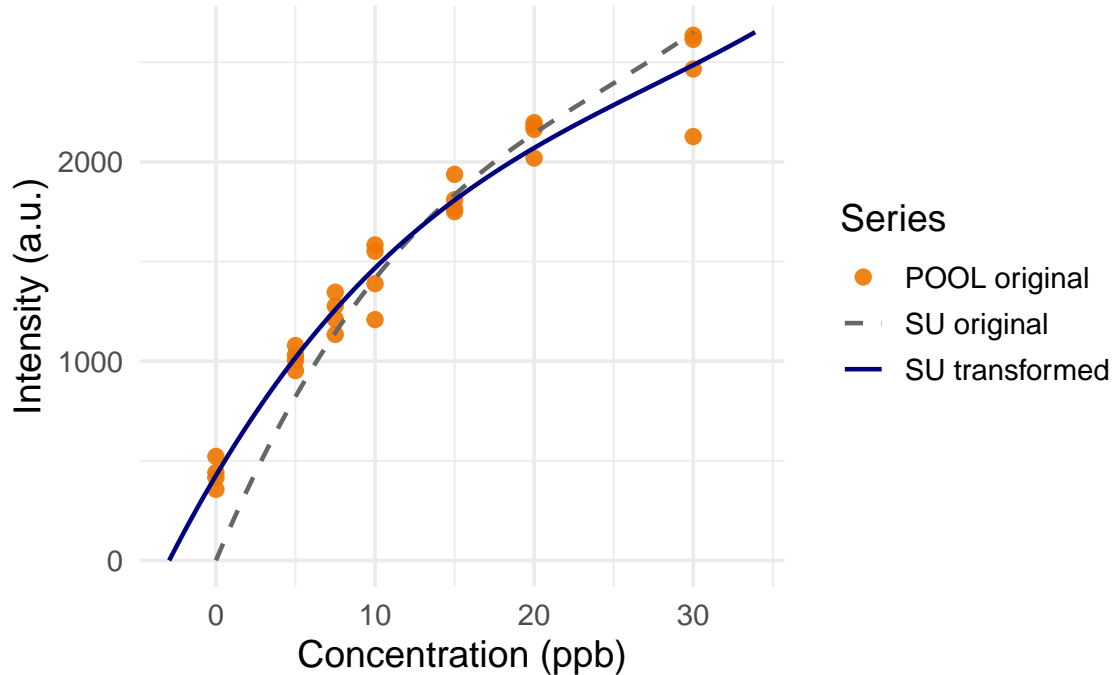
```
res_heptanone <- harmonize(pool_heptanone, su_heptanone)
```

Table 2: Harmonization parameters and uncertainty (Heptanone)

Parameter	Value	Error	Units
Scale	0.815	± 0.156	–
Shift	2.940	± 2.258	ppb

```
plot_harmonization(res_heptanone, title = "Calibration Harmonization - Heptanone")
```

Calibration Harmonization – Heptanone



Loss landscape and paraboloid approximation

In this section we visualize the loss landscape and the paraboloid approximation.

- A 2D heatmap of the loss over (scale, shift).
- A 2D heatmap of the fitted paraboloid on the same grid.
- 1D profiles:
 - Loss vs scale (shift fixed at its optimum).
 - Loss vs shift (scale fixed at its optimum).

In the 1D plots:

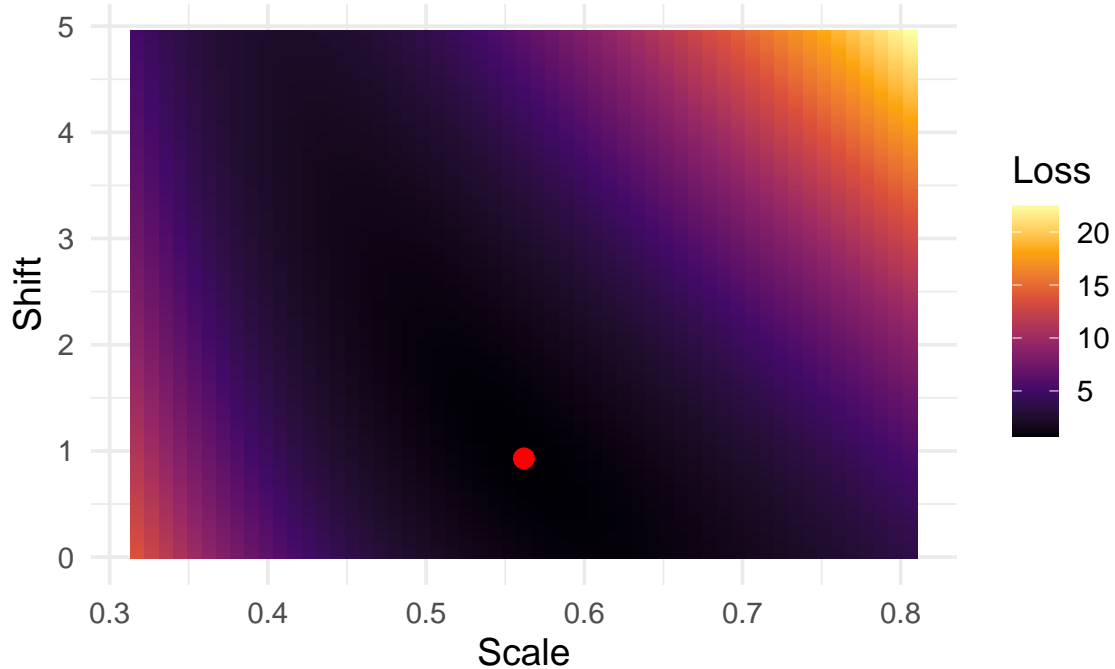
- Real loss: **black**.
- Paraboloid: **navy** (dashed).
- Optimal point: **red** vertical line.

2D heatmaps: real loss vs paraboloid

Anisole

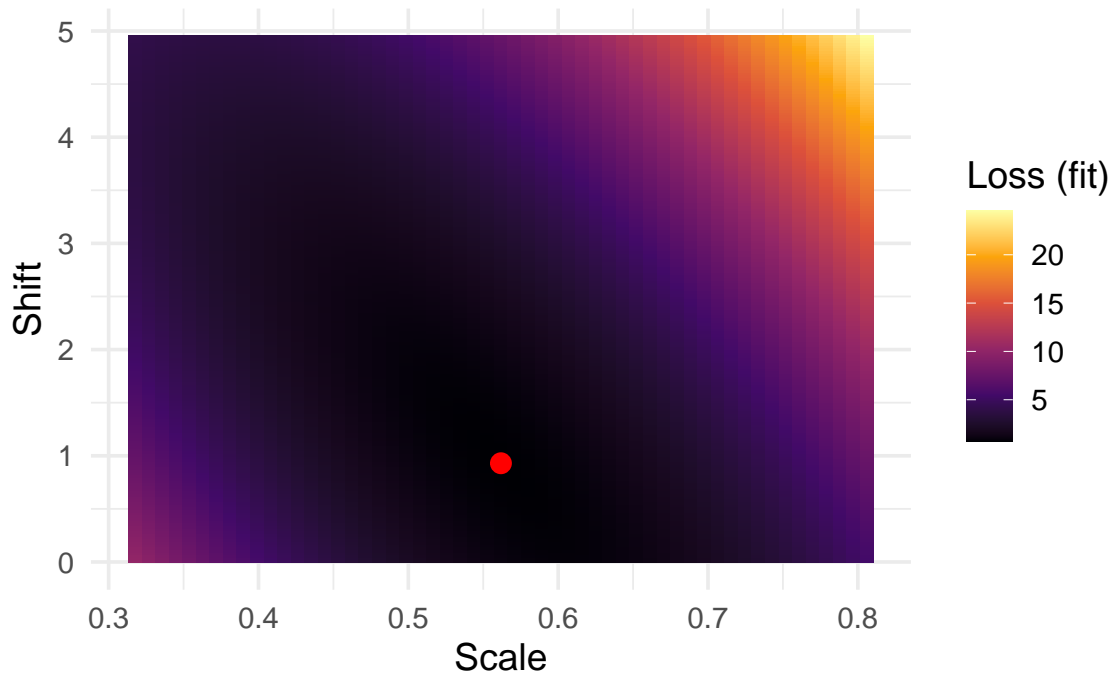
```
hm_anisole <- plot_loss_heatmaps(res_anisole, title_prefix = "Anisole - ", )
hm_anisole$real
```

Anisole – Loss landscape (real)



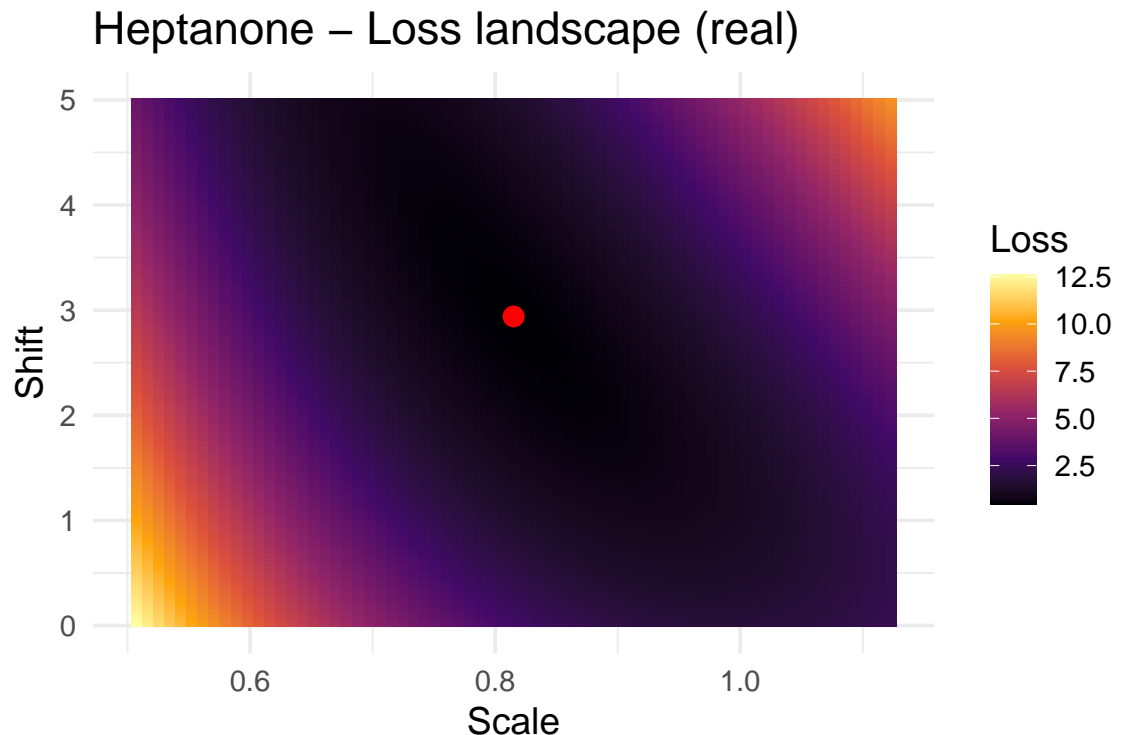
```
hm_anisole$paraboloid
```

Anisole – Loss landscape (paraboloid fit)



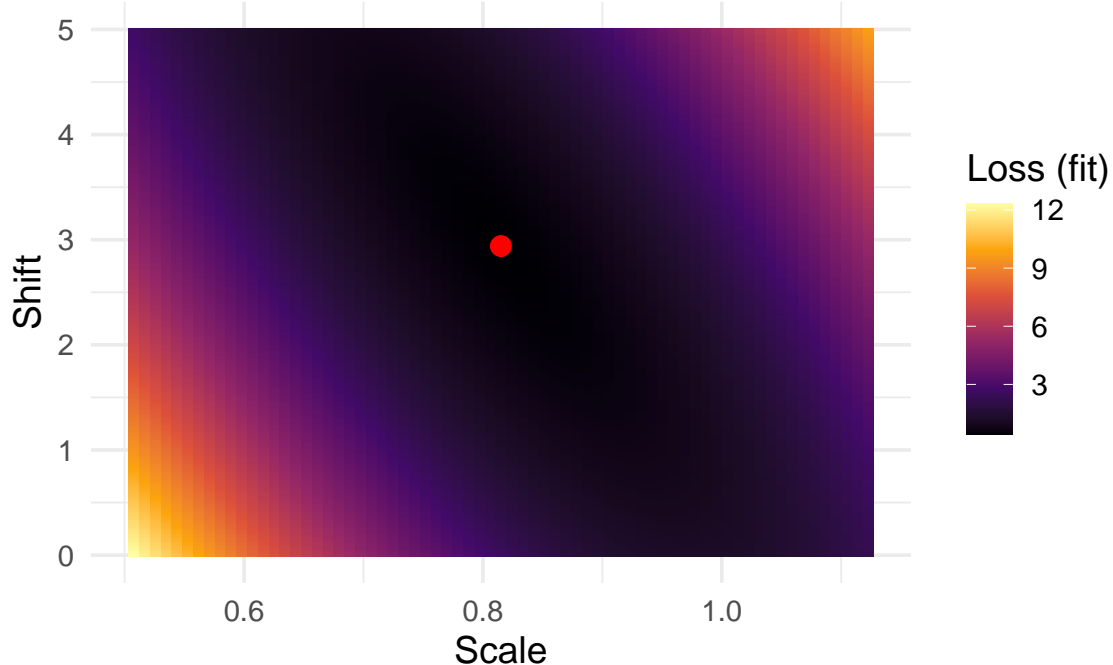
Heptanone

```
hm_heptanone <- plot_loss_heatmaps(res_heptanone, title_prefix = "Heptanone - ")  
hm_heptanone$real
```



```
hm_heptanone$paraboloid
```

Heptanone – Loss landscape (paraboloid fit)



1D profiles: loss vs scale and shift

We now plot the loss as a function of a single parameter:

- **Loss vs scale**, with shift fixed at its optimal value.
- **Loss vs shift**, with scale fixed at its optimal value.

We overlay:

- Real loss (black).
- Paraboloid approximation (navy, dashed).
- Optimal point (red vertical line).

```
plot_loss_1d <- function(h,
                           param = c("scale", "shift"),
                           range_limit = NULL,
                           loss_limits = NULL,
                           tol = 1e-6,
                           title = NULL) {

  param <- match.arg(param)

  if (is.null(h$paraboloid))
    stop("No paraboloid fit available.")

  # Extract optimum
  if (param == "scale") {
    a <- h$paraboloid$a
    if (is.na(a)) stop("Paraboloid coefficient 'a' is NA.")

    x0 <- h$scale
  }
}
```

```

err <- h$scale_error

lg <- h$loss_grid %>%
  dplyr::filter(
    is.finite(loss),
    abs(.data$shift - h$shift) < tol
  ) %>%
  dplyr::arrange(.data$scale) %>%
  dplyr::mutate(
    x = .data$scale,
    loss_parab = h$loss +
      a * (x - h$scale)^2
  )

} else { # param == shift

  b <- h$paraboloid$b
  if (is.na(b)) stop("Paraboloid coefficient 'b' is NA.")

  x0 <- h$shift
  err <- h$shift_error

  lg <- h$loss_grid %>%
    dplyr::filter(
      is.finite(loss),
      abs(.data$scale - h$scale) < tol
    ) %>%
    dplyr::arrange(.data$shift) %>%
    dplyr::mutate(
      x = .data$shift,
      loss_parab = h$loss +
        b * (x - h$shift)^2
    )
}

# Default X-limits = ±3×error
if (is.null(range_limit)) {
  range_limit <- c(x0 - 2*err, x0 + 2*err)
}

# Subset
lg_local <- lg %>% dplyr::filter(x >= range_limit[1], x <= range_limit[2])

# Default Y-limits
if (is.null(loss_limits)) {
  ymin <- min(lg_local$loss, lg_local$loss_parab, na.rm = TRUE)
  ymax <- max(lg_local$loss, lg_local$loss_parab, na.rm = TRUE)

  pad <- 0.10 * (ymax - ymin)
  loss_limits <- c(ymin - pad, ymax + pad)
}

if (is.null(title)) {

```

```

    title <- paste0("Loss vs ", param)
}

# Error bounds
x_lower <- x0 - err
x_upper <- x0 + err

# Plot
p <- ggplot(lg_local, aes(x = x)) +
  geom_line(aes(y = loss, linetype = "Real loss"),
            color = "black", linewidth = 1) +
  geom_line(aes(y = loss_parab, linetype = "Paraboloid"),
            color = "navy", linewidth = 0.9) +
  geom_vline(aes(xintercept = x0, linetype = "Optimum"),
             color = "red", linewidth = 0.9) +
  geom_point(aes(x = x_lower, y = 2 * h$loss),
             color = "red", shape = 4, size = 2, stroke = 1.2) +
  geom_point(aes(x = x_upper, y = 2 * h$loss),
             color = "red", shape = 4, size = 2, stroke = 1.2) +
  scale_linetype_manual(values = c(
    "Real loss" = "solid",
    "Paraboloid" = "dashed",
    "Optimum" = "dotted"
  )) +
  labs(
    title = title,
    x = param,
    y = "Loss",
    linetype = "Legend"
  ) +
  theme_minimal(base_size = 14) +
  theme(legend.position = "right") +
  xlim(range_limit) +
  ylim(loss_limits)

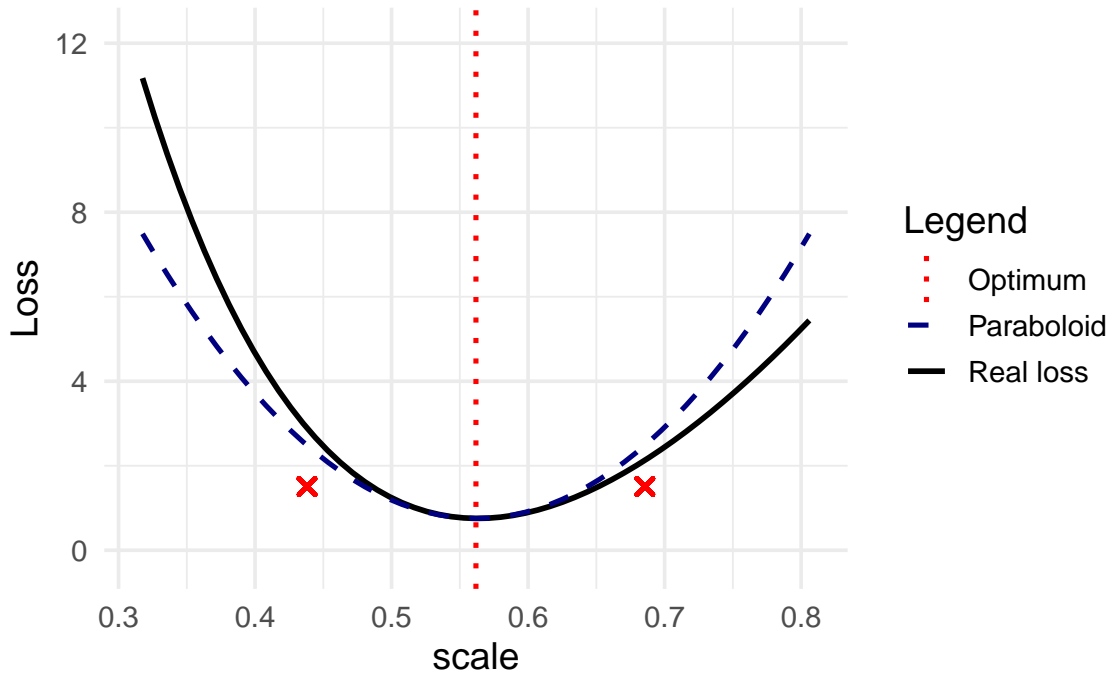
p
}

```

Anisole

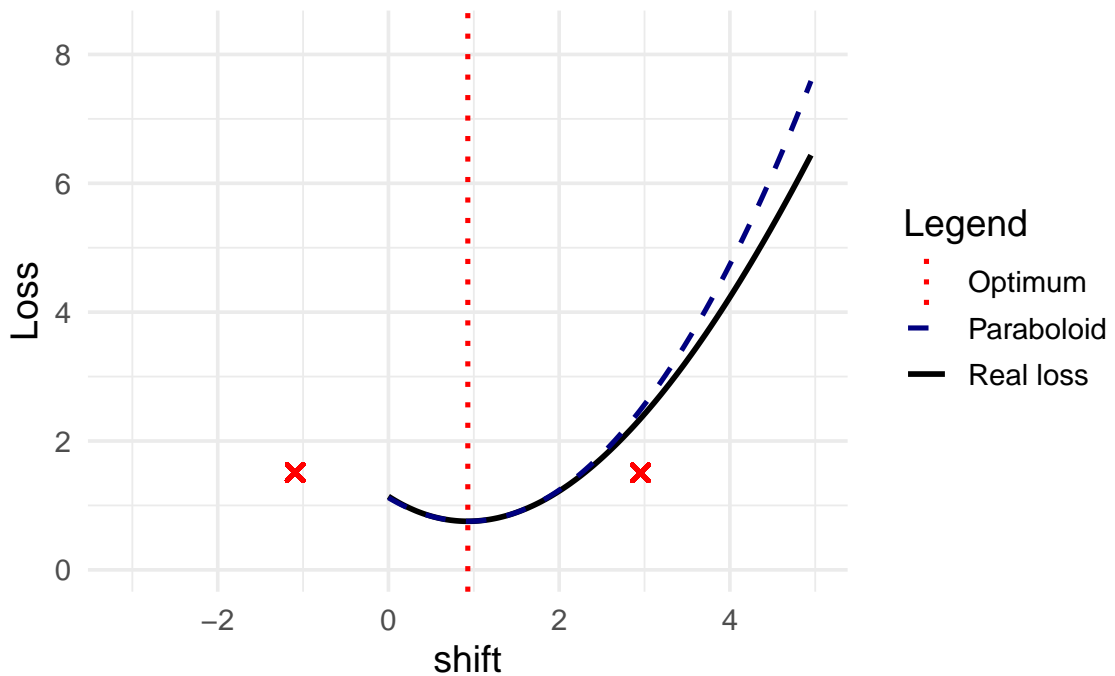
```
plot_loss_1d(res_anisole, param = "scale")
```

Loss vs scale



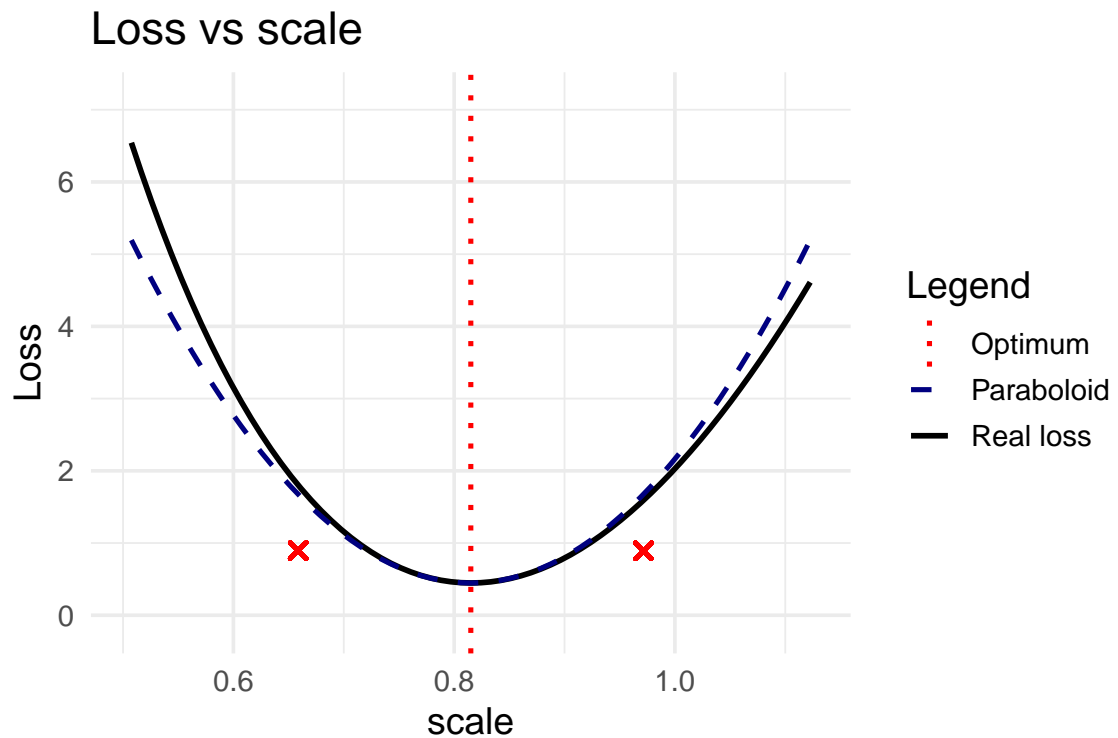
```
plot_loss_1d(res_anisole, param = "shift")
```

Loss vs shift



Heptanone

```
plot_loss_1d(res_heptanone, param = "scale")
```



```
plot_loss_1d(res_heptanone, param = "shift")
```

Loss vs shift

