Beyond Normalization: Incorporating Scale Uncertainty in ALDEx2

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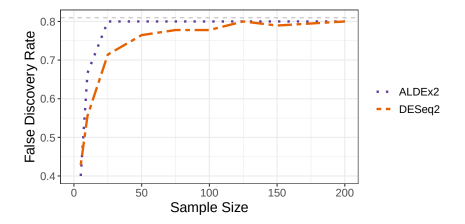
Recap: Sequencing depth can confound conclusions.

Observed data (Y)	Sample 1	Sample 2	Sample 3	
Condition	Pre	Pre	Post	Conclusion
Entity 1	5	10	100	Increase
Entity 2	10	25	3	Decrease
Entity 3	0	1	8	Increase
Entity 4	0	0	19	Increase
Sequencing Depth	15	36	130	

This can mislead analyses.

System data (W)	Sample 1	Sample 2	Sample 3	
Condition	Pre	Pre	Post	Conclusion
Entity 1	227	351	154	Decrease
Entity 2	684	891	3	Decrease
Entity 3	48	32	15	Decrease
Entity 4	43	39	27	Decrease
Scale (W^{\perp})	1,002	1,313	200	

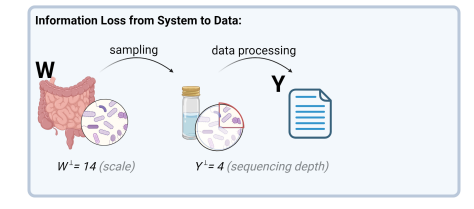
.. and lead to unacknowledged bias.



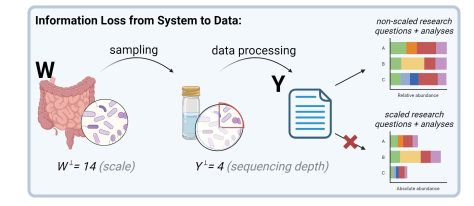
Section 1

Problem Set-Up

Observed Data as a Sample from the System



Observed Data as a Sample from the System



Notation

• Y: a measurement of the underlying system W.

$$\mathbf{W}_{dn} = \underbrace{\mathbf{W}_{dn}^{\parallel}}_{\text{composition}} \times \underbrace{W_{n}^{\perp}}_{\text{scale}}$$

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$$\mathbf{W}_{dn} = \underbrace{\mathbf{W}_{dn}^{\parallel}}_{\mathsf{composition}} \times \underbrace{\mathbf{W}_{n}^{\perp}}_{\mathsf{scale}}$$

- Composition: $\mathbf{W}_{dn}^{\parallel} = \frac{\mathbf{W}_{dn}}{\sum_{d=1}^{D} \mathbf{W}_{dn}}$
- Scale: $W_n^{\perp} = \sum_{d=1}^{D} W_{dn}$

Example: Notation

System data (W^{\parallel})	Sample 1	Sample 2	Sample 3
Condition	Pre	Pre	Post
Entity 1	0.27	0.27	0.77
Entity 2	0.68	0.68	0.02
Entity 3	0.05	0.02	0.08
Entity 4	0.04	0.03	0.13

	Sample 1	Sample 2	Sample 3
Condition	Pre	Pre	Post
Scale (W^{\perp})	1,002	1,313	200

Differential Abundance/Expression Analysis

- **Research Question:** How do entities (e.g., taxa or genes) change between conditions?
- \bullet θ : what we want to estimate.

$$\theta_d = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\,\mathbf{W}_{dn})$$

Step 1: Model Sampling Uncertainty

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(lpha)$

Step 2: Centered Log-Ratio Transformation

$$\log \mathbf{W}_{\cdot n} = \left[\log \mathbf{W}_{1n}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}), ..., \log \mathbf{W}_{Dn}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel})\right]$$

Step 3: Calculate LFCs and Test if Different from Zero.

$$\theta_d = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\,\mathbf{W}_{dn})$$

Implied Assumptions about Scale

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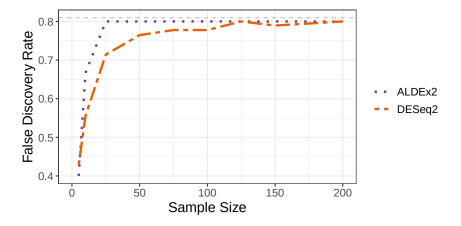
$$\theta_d = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\,\mathbf{W}_{dn})$$

Since $\log \mathbf{W}_{dn} = \log \mathbf{W}_{dn}^{\parallel} + \log W_n^{\perp}$, the CLR normalization implies:

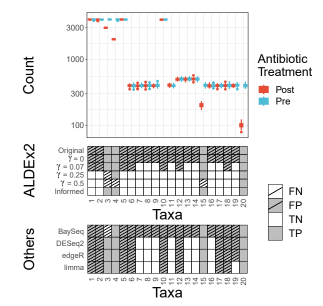
$$\log W_{dn} = \log \mathbf{W}_{dn}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel})$$
$$\log W_{n}^{\perp} = -\operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}).$$

What happens when this is wrong?

Unacknowledged bias!



Adding Uncertainty in Scale can Help.



Section 2

Scale Reliant Inference

Scale Reliant Inference: The Basics

- The CoDA perspective: Research questions that depend on W^{\perp} (scale) cannot be answered rigorously.
- The Normalization perspective: Research questions that depend on W^{\perp} (scale) can be answered after normalization.
- Who is right?

Scale Reliant Inference: The Basics

- The CoDA perspective: Research questions that depend on W^{\perp} (scale) cannot be answered rigorously.
- The Normalization perspective: Research questions that depend on W^{\perp} (scale) can be answered after normalization.
- Who is right?
- The CoDA perspective: Rigourous, but scientifically limiting.
- The Normalization perspective: Practical, but unacknowledged bias.

Scale Reliant Inference: The Basics

• For LFCs, θ depends on W^{\perp} :

$$\begin{split} \theta_d &= \mathsf{mean}_{\mathsf{case}}(\log \mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\log \mathbf{W}_{dn}) \\ &= \dots \\ &= \underbrace{\mathsf{mean}_{\mathsf{case}}(\log \mathbf{W}_{dn}^{\parallel}) - \mathsf{mean}_{\mathsf{control}}(\log \mathbf{W}_{dn}^{\parallel})}_{\theta^{\parallel}} \\ &+ \underbrace{\mathsf{mean}_{\mathsf{case}}(\log W_n^{\perp}) - \mathsf{mean}_{\mathsf{control}}(\log W_n^{\perp})}_{\theta^{\perp}} \end{split}$$

Scale Reliant Inference: Theory Intro

Recall for LFCs:

$$egin{aligned} heta_d &= \mathsf{mean}_\mathsf{case}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_\mathsf{control}(\mathsf{log}\,\mathbf{W}_{dn}) \ &= heta^{\parallel} + heta^{\perp} \end{aligned}$$

• What can we say about θ from θ^{\parallel} alone?

Scale Reliant Inference: Theory Intro

Recall for LFCs:

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- What can we say about θ from θ^{\parallel} alone?
- Statistical perspective: θ is not identifiable without θ^{\perp} .
- Practical issues: unbiased estimators, calibrated confidence sets, and type-I error control **NOT** possible!
- See Nixon et al. (2023) for details.

θ^{\perp} : The Missing Piece

$$\theta^{\perp} = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\ W_n^{\perp}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\ W_n^{\perp})$$

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- The change in scales between conditions matters for estimating LFCs.
- The scale only needs to be known up to a constant (see Nixon et. al (2023)).

$$\theta^{\perp} = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\ W_n^{\perp}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\ W_n^{\perp})$$

- The change in scales between conditions matters for estimating I FCs.
- The scale only needs to be known up to a constant (see Nixon et. al (2023)).
- Each normalization implies a value of θ^{\perp} (e.g., CLR):

$$\theta_{\mathsf{CLR}}^{\perp} = \mathsf{mean}_{\mathsf{case}}(-\log\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel})) - \mathsf{mean}_{\mathsf{control}}(-\log\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel}))$$

Scale Simulation Random Variables

Goal: Estimate $\theta = f(\mathbf{W}^{\parallel}, W^{\perp})$.

- Draw samples of \mathbf{W}^{\parallel} from a measurement model (can depend on Y).
- ② Draw samples of W^{\perp} from a scale model (can depend on W^{\parallel}).
- **3** Estimate samples of $\theta = f(\mathbf{W}^{\parallel}, W^{\perp})$.

Comparison to ALDEx2

The ALDEx2 Model

Step 1: Model Sampling Uncertainty

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

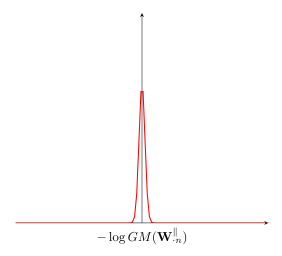
$$\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(\alpha)$$

Step 2: Centered Log-Ratio Transformation

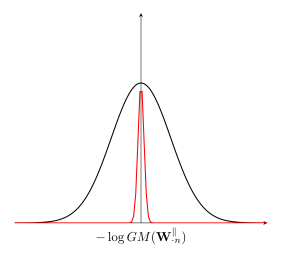
$$\log \mathbf{W}_{\cdot n} = \left[\log \mathbf{W}_{1n}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}), ..., \log \mathbf{W}_{Dn}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel})\right]$$

Step 3: Calculate LFCs and Test if Different from Zero.

The Original Scale Model



Extending the Original Scale Model



ALDEx2 as an SSRV

Step 1: Model Sampling Uncertainty

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(\alpha)$

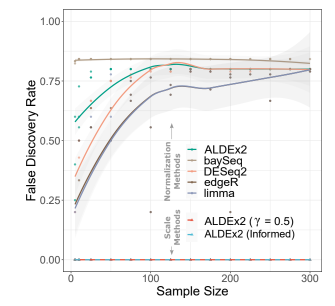
Step 2: Draw Samples from a Scale Model

$$\begin{split} \log W_n^{\perp} &= -\mathsf{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}) + \epsilon, \ \epsilon \sim \mathit{N}(0, \gamma^2) \\ \log \mathbf{W}_{\cdot n} &= \log \mathbf{W}_{\cdot n}^{\parallel} + \log W_n^{\perp} \end{split}$$

Step 3: Calculate LFCs and Test if Different from Zero.

$$\theta_d = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\,\mathbf{W}_{dn})$$

Benefits of Moving Past Normalizations to Scale



Section 3

Updated ALDEx2 Model

ALDEx2 as an SSRV

Step 1: Model Sampling Uncertainty

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(\alpha)$

Step 2: Draw Samples from a Scale Model

$$\log W_n^{\perp} \sim Q$$
 $\log \mathbf{W}_{\cdot n} = \log \mathbf{W}_{\cdot n}^{\parallel} + \log W_n^{\perp}$

Step 3: Calculate LFCs and Test if Different from Zero.

$$\theta_d = \text{mean}_{\text{case}}(\log \mathbf{W}_{dn}) - \text{mean}_{\text{control}}(\log \mathbf{W}_{dn})$$

Intro to Scale Models

There are no restrictions on what scale models can be, although there are some helpful options:

- Based on normalizations. (Stochastic normalizations)
- Based on biological knowledge.
- Based on outside measurements.

Scale Models based on Biological Knowledge

What do past studies or biological mechanisms tell about the scale of the system?

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 A past study showed that a certain disease (e.g., Crohn's disease) leads to lower microbial load in the gut. What do past studies or biological mechanisms tell about the scale of the system?

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$$\log W_{\rm Healthy}^{\perp} \sim N(1, \gamma^2) \\ \log W_{\rm Crohn's}^{\perp} \sim N(0.7, \gamma^2)$$

Scale Models based on Outside Measurements

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- These measurements can be used if they relate to your scale of interest.
- Examples include flow cytometry, qPCR, etc.
- Scale models can incorporate measurement uncertainty.

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$$\log W_n^{\perp} \sim N(\log \mu_{FC,n}, \sigma_{FC,n}^2)$$

Section 4

Changes to the ALDEx2 Interface

Including scale

The new ALDEx2 model removes normalizations in lieu of scale models.

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Major updates:

- A new argument gamma which makes it easy to incorporate scale uncertainty (aldex and aldex.clr functions).
 - gamma can either be a single numeric or a matrix.
 - Single numeric: controls the noise on the default scale model.
 - 2 Matrix: A $N \times S$ matrix of samples of W^{\perp} .
- A new function aldex.senAnalysis to see how analysis results change as a function of scale uncertainty.

Option 1: Default Scale Model

The default scale model is based on errors in the CLR normalization.

$$\log \hat{W}_n^{\perp(s)} = -\text{mean}\left(\log \hat{W}_n^{\parallel(s)}\right) + \Lambda^{\perp} x_n$$
$$\Lambda^{\perp} \sim N(0, \gamma^2).$$

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- When $\gamma = 0$, behavior matches the original ALDEx2 model.
- 2 For any value of $\gamma > 0$, it models potential error in the CLR assumption (false positives will decrease compared to the CLR normalization.)
- 1 It has a concrete interpretation to contextualize scale assumptions.

$$\begin{split} \theta_{\mathsf{Default Scale}}^{\perp} &= \mathsf{mean}_{\mathsf{case}}(-\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel})) - \mathsf{mean}_{\mathsf{control}}(-\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel})) + \epsilon \\ &= \theta_{\mathsf{CLR}}^{\perp} + \epsilon \\ &\epsilon \sim \mathit{N}(0, \gamma^2) \end{split}$$

Interpreting the Default Scale Model

$$\begin{split} \theta_{\mathsf{Default Scale}}^{\perp} &= \mathsf{mean}_{\mathsf{case}}(-\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel})) - \mathsf{mean}_{\mathsf{control}}(-\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel})) + \epsilon \\ &= \theta_{\mathsf{CLR}}^{\perp} + \epsilon \\ &\epsilon \sim \mathit{N}(0, \gamma^2) \end{split}$$

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The default scale model implies that:

• With 95% certainty, the value of θ^{\perp} is within $\pm 2\gamma$ of the value of θ_{CLR}^{\perp} .

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The default scale model implies that:

- With 95% certainty, the value of θ^{\perp} is within $\pm 2\gamma$ of the value of θ_{CLR}^{\perp} .
- With 95% certainty, the true difference in scales falls within the the range $2^{\theta_{CLR}^{\perp} \pm 2\gamma}$.

Example: Interpreting the Default Scale Model

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• Suppose that we are performing differential abundance in a case/control study where $\theta_{\rm CLR}=0.04$.

Example: Interpreting the Default Scale Model

With 95% certainty, the true difference in scales falls within the the range $2^{\theta_{\text{CLR}}^{\pm} \pm 2\gamma}$.

- Suppose that we are performing differential abundance in a case/control study where $\theta_{\rm CLR}=0.04$.
- Suppose we set $\gamma = 0.5$.

With 95% certainty, the true difference in scales falls within the the range $2^{\theta_{CLR}^{\perp}\pm 2\gamma}$.

- Suppose that we are performing differential abundance in a case/control study where $\theta_{CLR} = 0.04$.
- Suppose we set $\gamma = 0.5$.
- Then, this implies that, with 95% certainty, we believe that the scale of the case condition is within a factor of $[2^{0.04-2\times0.5}, 2^{0.04+2\times0.5}] = [0.51, 2.05]$ of the control condition.

Using the Default Scale Model

```
## Adding noise via the default scale model
mod.defaultScale <- aldex(Y, conds, gamma = 0.5)</pre>
```

Option 2: More Complex Scale Models

Alternatively, can pass a matrix of scale samples to gamma so long as:

- **1** The dimension is $N \times S$.
- ② They are samples of W^{\perp} not $\log W^{\perp}$.

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Reasons to do this:

- Biological beliefs: Scale is guided by the biological system or the researcher's prior beliefs.
- Outside Measurements: These can be used in building a scale model if they are informative on the scale of interest (e.g., qPCR, flow cytometry).

Sensitivity Analyses

• Instead of picking γ , why not test over a range instead?

Sensitivity Analyses

Step 1: Model Sampling Uncertainty

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(\alpha)$

Step 2: Draw Samples from a Scale Model For a given γ :

$$\log W_n^{\perp,\gamma} = -\text{mean}\left(\log \hat{W}_n^{\parallel(s)}\right) + \Lambda^{\perp} x_n$$

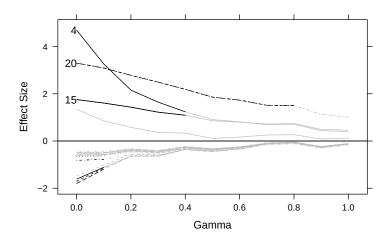
$$\Lambda^{\perp} \sim N(0, \gamma^2)$$

$$\log \mathbf{W}_{\cdot n}^{\gamma} = \log \mathbf{W}_{\cdot n}^{\parallel} + \log W_n^{\perp,\gamma}$$

Step 3: Calculate LFCs and Test if Different from Zero.

Step 4: Repeat for all desired values of γ .

Example: Sensitivity Analyses



Section 5

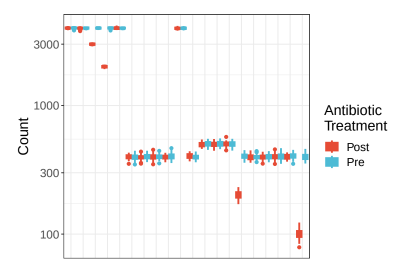
Data Examples

Simulation Study

Consider a simple study of the microbiome pre/post antibiotic administration.

- Research question: Which taxa change in absolute abundance after taking an antibiotic?
- 100 study participants, 50 in each condition (pre/post antibiotics).
- 20 taxa total with 4 taxa truly changing (decreasing)

Data



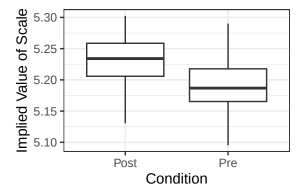
Adding Scale is Easy

```
## Adding noise via the default scale model
mod.ss.high <- aldex(Y, conds, gamma = 0.5)</pre>
```

```
## Looking at the implied scale
clr <- aldex.clr(Y, conds, gamma = 1e-3)</pre>
clr@scaleSamps[1:6, 1:4]
```

```
[,1] [,2] [,3] [,4]
##
## [1,] 5.174279 5.124890 5.199780 5.175163
   [2,] 5.175705 5.144470 5.184953 5.167715
   [3,] 5.178751 5.171188 5.130795 5.100749
   [4,] 5.158594 5.195139 5.164371 5.145696
   [5,] 5.120674 5.175533 5.189581 5.171154
## [6,] 5.208741 5.273464 5.207085 5.162631
```

Investigating Assumptions about Scale, cont.



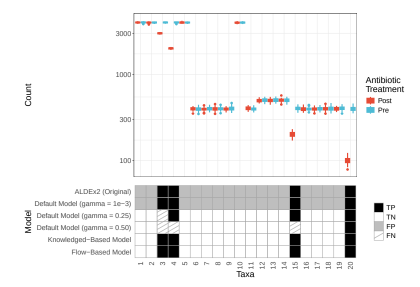
Scale Model based on Biology

```
## Creating an informed model using biological
reasoning
scales \leftarrow c(rep(1, 50), rep(0.9, 50))
scale samps <- aldex.makeScaleMatrix(</pre>
  gamma = .15,
  mu = scales,
  conditions = conds,
  log = FALSE
mod.know <- aldex(Y, conds, gamma = scale_samps)</pre>
```

Scale Model based on Outside Measurements

```
scale_samps <- matrix(NA,</pre>
  nrow = nrow(flow_data_collapse),
  ncol = 128
for (i in 1:nrow(scale_samps)) {
  scale_samps[i, ] <- rnorm(</pre>
    n = 128,
    mean = flow data collapse$mean[i],
    sd = flow data collapse$stdev[i]
mod.flow <- aldex(Y, conds, gamma = scale samps)</pre>
```

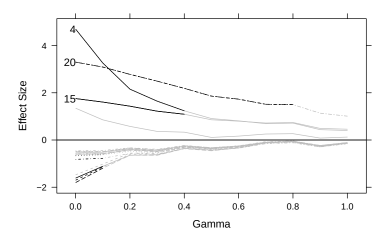
Plotting Results



Sensitivity Analyses

```
## First, specifying different values for the noise
in the scale
gamma to test <-c(1e-3, seq(0.1, 1, by = .1))
## Run the CLR function
clr <- aldex.clr(Y. conds)</pre>
## Run sensitivity analysis function
sen res <- aldex.senAnalysis(clr,
  gamma = gamma_to_test
plotGamma(sen_res,
 thresh = .1.
  blackWhite = TRUE, taxa to label = 3
```

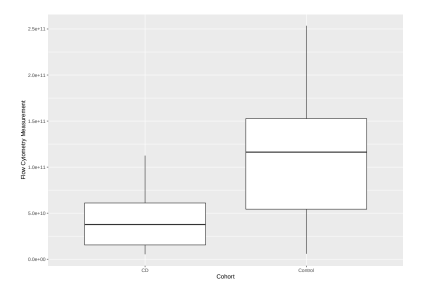
Sensitivity Analyses, cont.



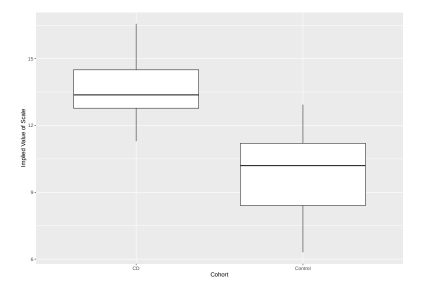
Real Example: Vandputte

- Comparison study of 29 Crohn's disease patients and 66 healthy controls.
- Por each patient, they sequenced the fecal sample and obtained flow cytometry measurements.
- Proposed an approach that supplemented sequence count data with flow cytometry measurements.

Difference in Scale Implied by Flow Cytometry



Difference in Scale Implied by CLR



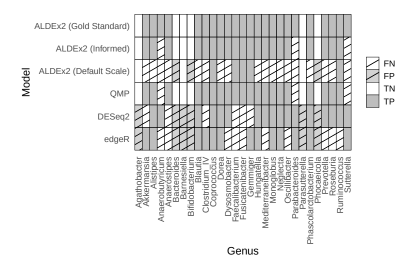
Creating a Gold Standard Model

```
scale_mean <- log2(sample_data(phylo)$CellCount)</pre>
scale var \leftarrow rep(0.7, 95)
scale samples <- matrix(NA, nrow = 95, ncol = 1000)
for (i in 1:95) {
  scale samples[i, ] <- 2^rnorm(</pre>
    1000,
    scale mean[i],
    scale_var[i]
```

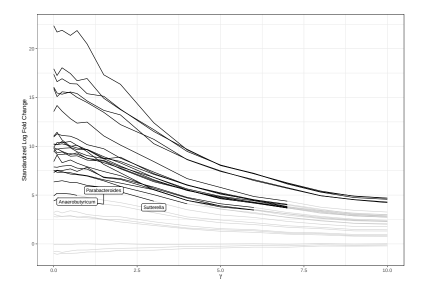
```
Creating an Informed Model
```

```
scale.cd <- 2^matrix(rnorm(1000 * 29,</pre>
  mean = log2(.7), sd = .125
), nrow = 29)
scale.control <- 2^matrix(rnorm(1000 * 66,
  mean = log2(1), sd = .125
), nrow = 66)
scale.informed <- rbind(scale.cd, scale.control)</pre>
aldex informed <- aldex(Y, X,
  mc.samples = 1000,
  gamma = scale.informed
```

Comparing to Other Methods



Sensitivity Analyses



References

Scale Reliant Inference/Updates to ALDEx2:

- Nixon, et. al. (2023) "Scale Reliant Inference." ArXiv Preprint 2201.03616.
- Gloor, Nixon, and Silverman. (2023) "Scale is Not What You Think; Explicit Scale Simulation in ALDEx2." BioRXiv Preprint 2023.10.21.563431.
- Nixon, Gloor, and Silverman. (2024) "Beyond Normalizations: Incorporating Scale Uncertainty in ALDEx2." BioRXiv Preprint 2024.04.01.587602.
- Fernandes et. al. (2014). "Unifying the analysis of high-throughput sequencing datasets: characterizing RNA-seq, 16S rRNA gene sequencing and selective growth experiments by compositional data analysis." *Microbiome*.

References

Data Sources:

- McMurrough et. al. (2014)."Control of catalytic efficiency by a co-evolving network of catalytic and non-catalytic residues." PNAS.
- Vandputte et. al. (2017). "Quantitative microbiome profiling links gut community variation to microbial load." *Nature*.