Methods:

Sequencing runs:

ONT 1D R9 runs were carried out in the lab:

Species name	Sample name	Total sequences	Total size (bp)	Longest sequence	Genome size (approx, pg 2C)
Sorbus aria	Database: SCI-	17,928	23,196,528	66,551	0.71
		0160729_Sorbus- or112154.			
Nepenthes	Database:	51,608	68,411,676	95,184	0.28
alata	SCI-FEST-				(N.pervillei)
	A_20160729	_Napenthes-			,
	alata.				
Silene	Database:	22,923	36,943,694	23,031	2.7 (S.latifolia)
uniflora	SCI-FEST-				
	$C_20160802$	$_{\rm Silene}$ 1D $_{\rm sf}$.			
Erycina	Database:	57,699	71,713,785	144,298	1.9
echinata	SCI-FEST-				(E.diaphena)
	$C_20160802$	_Erycina.			
Beta patula	Database:	23,694	40,492,055	117,469	1.25;1.28
	SCI-FEST-				(B.vulgaris/B.maritim
	A_20160802	_beta.			

Science festival sequencing was carried out in the Kew Science Festival marquee (with flowcells later transported back to the lab to continue running in some cases):

	Sci fest	Sci fest		Reads		Time to	First	fasta	<u>.</u>
	dresequenc-	ran-	Sci fest	se-	bp se-	identifica-	se-	file	(11)
Species Name	ing run number	domised letter	guessed ID	quenced at ID	quenced at ID	$ \text{tion} \\ (\text{TTI}) $	quence start	$rac{ ext{write}}{ ext{time}}$	(oldest TTI)
???	Sample_01	?	Nepenthe alata	es1460*	1495967	*	?	?	14:35*
???	Sample_02	?	Arabidop lyrata ssp. petraea**		603677*	*	?	17:26	(next day)
Silene uniflora	$Sample_03$	F	Silene uniflora	1,421	2,725,09	0	?	14:50	15:19
Erycina echinata	Sample_04	Е	Erycina echinata	1,679	2,028,97	5	?	16:56	22:56
Sorbus aria	Sample_05	D	Sorbus aria	210	505,043		?	14:11	14:19
Beta patula	Sample_06	В	Beta patula	3,546	7,226,10	6	?	16:01	16:05

Resampling in silico:

Simulations with 20 replicates were constructed by subsampling (without replacement) from the R4IDs-sequencing and festival-resequenced data sets. For each simulation replicate, the R4ID input and resequencing run were subsampled randomly for 100, 500, 1000, 5000 and 10000 reads, in combination (25 orthogonal combinations per input sample). Each R4IDs dataset was converted to a BLASTN database separately.

Sample ID via BLASTN:

Each simulated sample was matched to every BLASTN database in turn with the following parameters: blastn -outfmt "6 sacc qacc length pident evalue" -evalue 0.01 -num_threads 6 -num_alignments 1 -max_hsps 1 -db \$this_db -query \$this_test_input

Only the top hit for each read is reported in the output file. Output files for each database/sample intensity combination (5 samples * 5 databases * 5 sampling intensities * 5 database intensities; 625 total) were then combined as follows: For each input sample at a given sampling/DB intensity combination, all hits against each of the 5 databases were collected for every read. The following ID statistics were then compiled:

Stat	Definition
TP	Reads only
one-	matching the
way	correct DB
FP one-	Reads only
way	matching an
	incorrect DB
TP	Reads
two-	matching the
way	correct DB,
	and at least
	one other DB,
	but with the
	longest number
	of identities for
	the correct DB
FP two-	Reads
way	matching the
	correct DB,
	and at least
	one other DB,
	but with the
	longest number
	of identities for
	the INcorrect
	DB; or
	matching more
	than one DB
	but NOT the
	correct one at
	all

Stat	Definition
mean	Average of (TP
bias	hit -
	next-best-hit)
	for all reads
	with at least
	two hits, one
	of which is for
	the correct DB
two-	Ratio of TP
way	two-way: total
rate	two-way hits
two-	Ratio of hits
way	with bias $>$
rate	50bp: total
with	two-way hits
cutoff	(e.g., more
	$\mathbf{stringent})$
total	Total number
hits	of reads with
	any hit to any
	DB

Not all replicate / sample combinations generated results owing to random error. In the plots below data were aggregated amongst replicates. 20 replicates were carried out in total.

 $\it Note$: For these the following sample identities were assumed...

Sample	File	TP label
sample_1	_Volun	nes <u>N.</u> Slata
	FEST-	
	A_sci-	
	fest-	
	$data_2$	0160823_{samp}
	-	retools.fasta
$sample_2$		nes_SCI-
		Arabidopsis
		(omitted $)$
	fest-	
		0160823_samp
	-	retools.fasta
$sample_3$		ne S_&Gf lora
	FEST-	
	A_sci-	
	fest-	04.00000
		0160823_samp
1 4	-	retools.fasta
sample_4		ne £_&©I nata
	FEST-	
	A_sci-	
	fest-	0160000
		0160823_{samp}
	runs.po	retools.fasta

```
Sample
           File
                    TP label
sample 5
             VolumeS.aSGI-
           FEST-
           A sci-
           fest-
           data_20160823_sample_05_basecall_both-
           runs.poretools.fasta
sample\_6
            Volume B. Sattlela
           FEST-
           A sci-
           fest-
           data_20160823_sample_06_basecall_both-
           runs.poretools.fasta
```

Results

```
# set up
#library(ROCR)
library(lattice)
# read in the input for each labelling / subsampling size
\#rep_01 = read.table('~/Documents/all_work/programming/oddjects-sandbox/R4IDs/manuscript-analyses/collegistary)
rep_01 = read.table('~/Documents/all_work/programming/oddjects-sandbox/R4IDs/manuscript-analyses/201908
# cbind them all into a big table, adding the 'training_intensity' and 'query_intensity' cols
# calculate TP rate for a variety of cutoffs
# plot basic
plot(sample_DB_intensity ~ species_DB_intensity, data=rep_01[rep_01$TP_species=='Beta-patula',])
plot(sample_DB_intensity ~ two_way_rate_with_cutoff, data=rep_01[rep_01$TP_species=='Beta-patula',])
# plot heatmap prediction
#rep_01[rep_01$TP_species=='Beta-patula',c(2,4,12)]
levels(rep_01$TP_species)
species = c(
  "Beta-patula",
  "Erycina-echinata",
  "Napenthes-alata",
  "Silene-uniflora",
  "Sorbus-aria")
# bias
levelplot(two_way_rate_with_cutoff ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=rep_01
```

```
levelplot(two_way_rate ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=rep_01[rep_01$TP_s]
levelplot(two_way_rate ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=rep_01, col.region
```

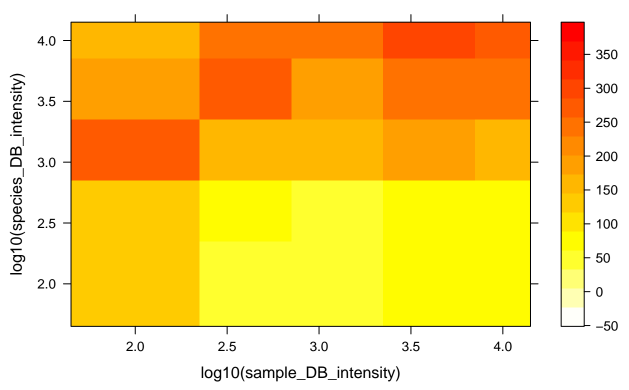
Now plot 2-way rate with cutoff=50:

```
levelplot(two_way_rate_with_cutoff ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=rep_01
```

Now repeat it properly for all reps. Mean bias first:

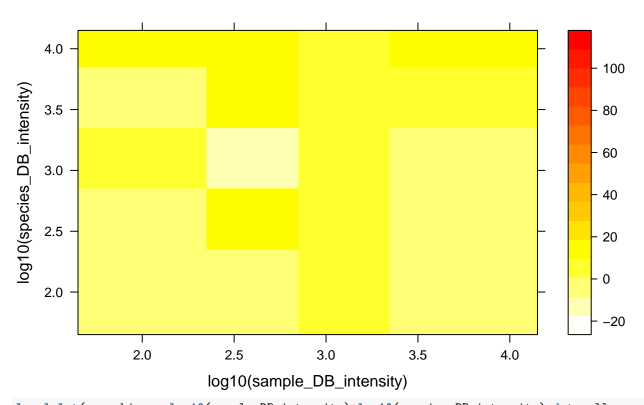
```
# read in the input for each labelling / subsampling size
#all_reps = read.table('~/Documents/all_work/programming/oddjects-sandbox/R4IDs/manuscript-analyses/col
all_reps = read.table('~/Documents/all_work/programming/oddjects-sandbox/R4IDs/manuscript-analyses/2019
# bias
levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps[all_reps$TP_
```

Beta patula



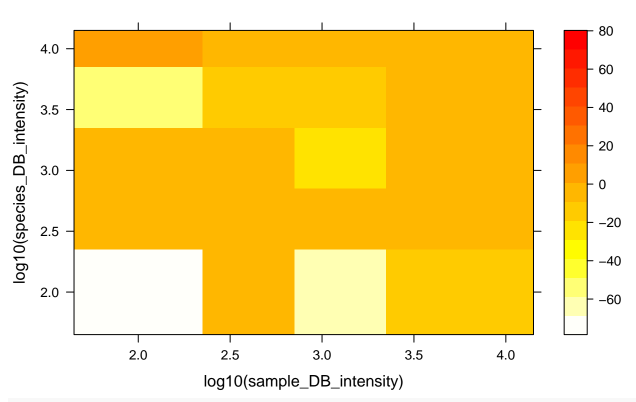
levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps[all_reps\$TP_

Erycina echinata



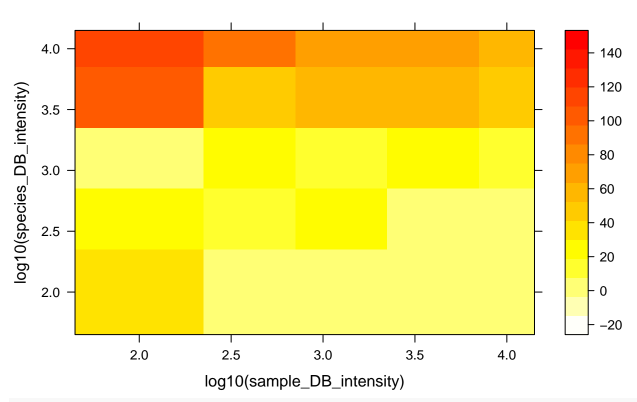
 ${\tt levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity), data=all_reps[all_reps$TP_intensity]}, and the substitution of the substitution of$

Napenthes alata



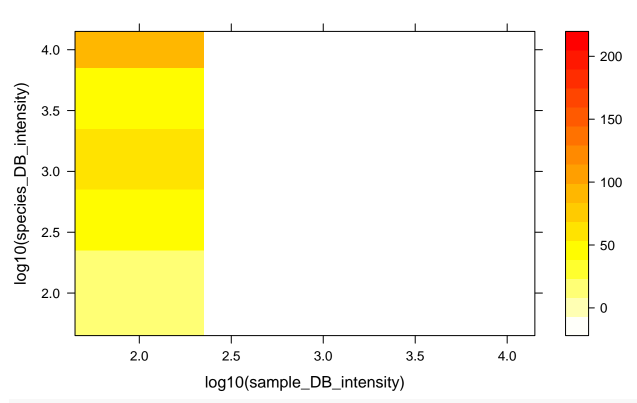
levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps[all_reps\$TP_

Silene uniflora

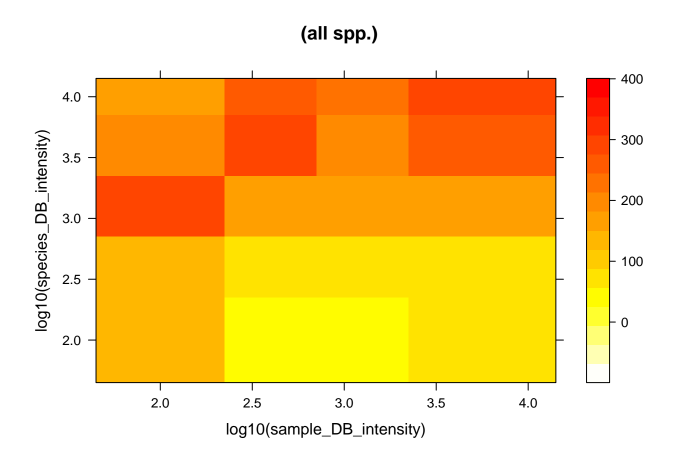


levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps[all_reps\$TP_

Sorbus aria



levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.regions



Now mean bias times total hits, e.g. what we'd actually see in the GUI

```
levelplot(two_way_rate*total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_rep
```

Now plot 2-way rate %:

```
levelplot(two_way_rate ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps[all_reps$
levelplot(two_way_rate ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.regi
```

Now plot 2-way rate with cutoff=50:

```
levelplot(two_way_rate_with_cutoff ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_relevelplot(two_way_rate_with_cutoff ~ log10(sample_DB_intensity)*log10(species_DB_int
```

```
boxplot(mean_bias ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='Sorb
boxplot(two_way_rate ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='S
boxplot(two_way_rate_with_cutoff ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='N
boxplot(two_way_rate ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='N
boxplot(mean_bias ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='N
boxplot(mean_bias ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='Siles
boxplot(mean_bias ~ species_DB_intensity * sample_DB_intensity,data=all_reps[all_reps$TP_species=='Eryc
```

Now try adding 1- and 2-way hits

```
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity)
```

Hits ratios heatmap by species:

```
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(spec
```

So far...

OK: something a bit odd is going on here, but basically it looks like a) it's all working OK more or less b) species DB intensity seems to make a bigger difference than sample effort/intensity.

To show this, collect all the aggregates in one:

```
par(mfrow=c(2,3))
levelplot(two_way_rate ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.regi
levelplot(two_way_rate_with_cutoff ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.regions
levelplot(mean_bias ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.regions
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP) ~ log10(sample_DB_intensity)*log10(species_DB_intensity)
levelplot((one_way_TP + two_way_TP) / total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity)
levelplot(total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.region
```

checking headline stuff; how does blast DB intensity affect outputs?:

```
par(mfrow=c(2,3))
boxplot((one_way_TP + two_way_TP) /(one_way_FP + two_way_FP) ~ log10(species_DB_intensity),data=all_rep
boxplot((one_way_TP + two_way_TP) /(total_hits) ~ log10(species_DB_intensity),data=all_reps, col.region
boxplot(mean_bias ~ log10(species_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(he
boxplot(two_way_rate_with_cutoff ~ log10(species_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length
boxplot(total_hits ~ log10(species_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(heat.colors)]
```

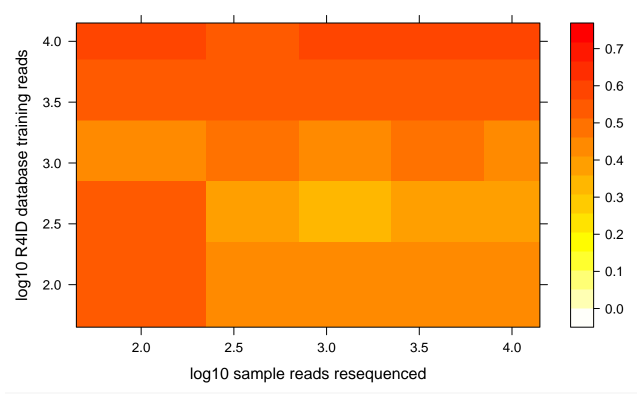
how does resequencing (sampling) intensity affect the same?:

```
par(mfrow=c(2,3))
boxplot((one_way_TP + two_way_TP) /(one_way_FP + two_way_FP) ~ log10(sample_DB_intensity),data=all_reps
boxplot((one_way_TP + two_way_TP) /(total_hits) ~ log10(sample_DB_intensity),data=all_reps, col.regions
boxplot(mean_bias ~ log10(sample_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(heaboxplot(two_way_rate ~ log10(sample_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(boxplot(two_way_rate_with_cutoff ~ log10(sample_DB_intensity),data=all_reps, col.regions = heat.colors(boxplot(total_hits ~ log10(sample_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(heaboxplot(total_hits ~ log10(sample_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(heaboxplot(total_hits ~ log10(sample_DB_intensity),data=all_reps, col.regions = heat.colors(100)[length(heaboxplot(total_hits) ~ log10(sample_DB_i
```

Final summary; ploting expected positive reads vs total sampling intensity:

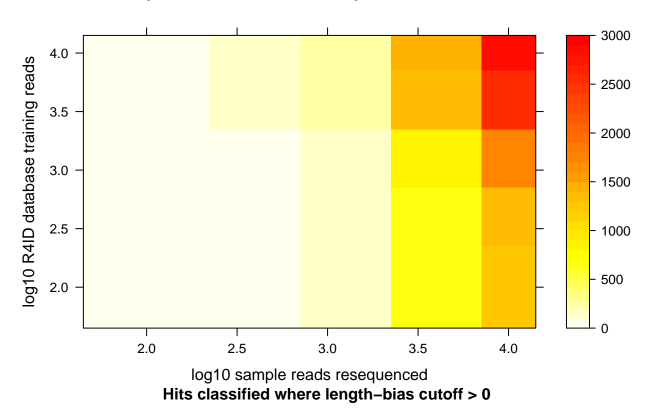
```
# Supplemenary Figure 1
levelplot((one_way_TP + two_way_TP) / (one_way_FP + two_way_FP+one_way_TP + two_way_TP) ~ log10(sample_
```

BLAST hit accuracy (TP:FP+TP)



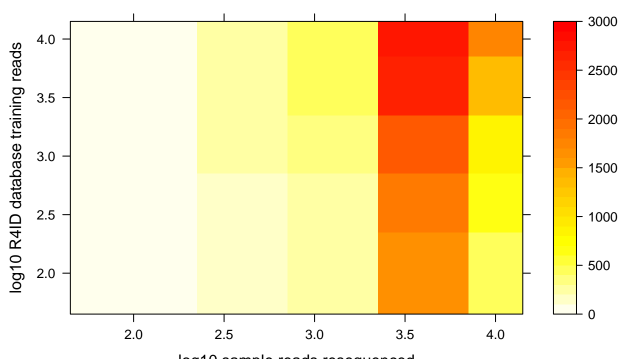
#levelplot(total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_reps, col.regio
Supplementary figure S2a
levelplot(two_way_rate*total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity),data=all_rep

Expected number of true-positive BLAST hits



Supplementary figure S2b
levelplot(two_way_rate_with_cutoff*total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensity),

Expected number of true-positive BLAST hits



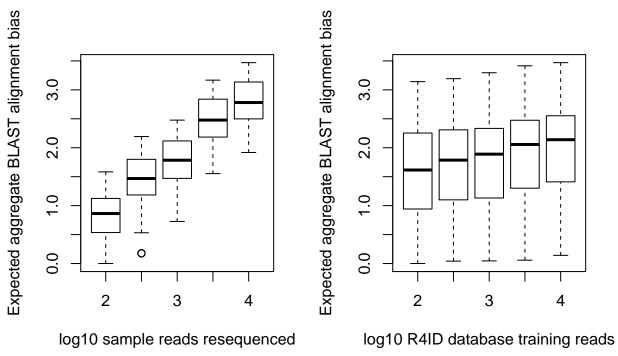
log10 sample reads resequenced

Hits classified where length-bias cutoff > 50

```
levelplot((one_way_FP + two_way_FP) / (one_way_FP + two_way_FP+one_way_TP + two_way_TP) ~ log10(sample_
levelplot(((one way TP + two way TP) / (one way FP + two way FP+one way TP + two way TP)*total hits) ~
levelplot(((one_way_FP + two_way_FP) / (one_way_FP + two_way_FP+one_way_TP + two_way_TP)*total_hits) ~
#levelplot((one_way_TP + two_way_TP) / total_hits ~ log10(sample_DB_intensity)*log10(species_DB_intensi
#xlab='log10 sample reads resequenced'
#ylab='log10 R4ID database training reads'
# boxplots by intensity to see confidence intervals
par(mfrow=c(1,2),oma=c(0,0,2,0))
boxplot(log10(two_way_rate*total_hits) ~ log10(sample_DB_intensity),data=all_reps, xlab='log10 sample r
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 1 is not drawn
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 2 is not drawn
boxplot(log10(two_way_rate*total_hits) ~ log10(species_DB_intensity),data=all_reps, xlab='log10 R4ID da
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 1 is not drawn
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 2 is not drawn
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 3 is not drawn
```

```
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 4 is not drawn
## Warning in bplt(at[i], wid = width[i], stats = z$stats[, i], out =
## z$out[z$group == : Outlier (-Inf) in boxplot 5 is not drawn
title("Expected aggregate BLAST alignment bias",outer=T)
```

Expected aggregate BLAST alignment bias



Conclusion

Overall it seems to be working fine, e.g. we get hits and length differences that tend to correct at low intensities (10E4 is a bloody small amount of sequencing..!)

The biggest determinant of overall success seems to be R4IDs database sampling effort, not resequencing effort. This is interesting.

The length bias cutoff (hits' length difference greater than 50 in favour of TP database for a positive hit) seems to greatly improve accuracy at the expense of total number of hits. An optimal tradeoff will depend on the expected taxonomic distance between likely ID candidates.

Unsure what is driving the between-species differences, but the quality of the runs in terms of total yield and also q-scores is probably to blame, as well as genome size (since this affects coverage). Our target genomes here range from ~ 0.28 to 3.0Gbp, and our maximum R4IDs sequencing yields are in the 20Mbp (0.02Gbp) to 70Mbp zone; so we are sequencing much less than a 1x-coverage genome. Insofar as this works at all it is probably partly (largely?) due to bias; in the **A. thaliana** data we observed that plastid depths were ~ 350 x even though total genome depth $\sim 2x$ (max).

Note will need to know genome sizes more accurately if at all possible - only have C-values.