WEBVTT

- 1 00:00:00.000 --> 00:00:04.167 (interference drowns out speaker)
- 2 00:00:08.192 --> 00:00:11.279 < v Announcer>Biostatistics, computational.</br/> < v >
- 3 00:00:11.279 --> 00:00:14.670 (interference drowns out speaker)
- $4\ 00:00:14.670 \longrightarrow 00:00:19.486$ So before joining UCLA in 2013.
- $5\ 00:00:19.486 --> 00:00:23.640$ (interference drowns out speaker)
- $6\ 00:00:23.640 \longrightarrow 00:00:24.660$ Production work.
- 7 00:00:24.660 --> 00:00:28.800 (interference drowns out speaker)
- $8~00{:}00{:}28.800$ --> $00{:}00{:}33.242$ On the script is (interference drowns out speaker)
- 9 00:00:33.242 --> 00:00:35.317 include differentiation factors,
- 10 00:00:35.317 --> 00:00:37.316 asymmetric (indistinct) replication,
- 11 00:00:37.316 --> 00:00:39.520 p-value-free false discovery (indistinct),
- $12\ 00:00:39.520 \longrightarrow 00:00:42.540$ and a high dimensional variable selection.
- 13 00:00:42.540 --> 00:00:46.200 And on the bio (indistinct) application side,
- 14 00:00:46.200 --> 00:00:50.415 her research include all single cell (indistinct)
- $15\ 00:00:50.415 --> 00:00:52.031$ for (indistinct) genomics and (indistinct).
- 16 00:00:52.031 --> 00:00:56.198 (interference drowns out speaker)
- 17 00:00:59.610 --> 00:01:01.331 Research published.
- 18 00:01:01.331 --> 00:01:05.498 (interference drowns out speaker)
- 19 00:01:13.971 --> 00:01:14.804 2019
- 20~00:01:16.170 --> 00:01:20.905 She's an MIT Technology Review certified (indistinct)
- $21\ 00:01:20.905 \longrightarrow 00:01:24.679$ in 2020, and she has received from Harvard.
- 22 00:01:24.679 --> 00:01:28.846 (interference drowns out speaker)
- 23 00:01:30.943 --> 00:01:33.360 <
v Jingyi>I couldn't wait for the introduction.
</v>
- 24 00:01:33.360 --> 00:01:36.240 It's my honor here to present my work,
- $25~00:01:36.240 \longrightarrow 00:01:39.810$ and my sabbatical in this fellowship program
- $26\ 00:01:39.810 --> 00:01:41.370$ at Harvard Radcliffe Institute.

- $27\ 00:01:41.370 --> 00:01:46.050$ So it's my pleasure to talk about some of our recent work
- $28\ 00{:}01{:}46.050 \dashrightarrow 00{:}01{:}50.910$ related to how statistic rigor is important in genomics.
- 29 00:01:50.910 --> 00:01:54.390 So I want to say that when I was a student,
- $30\ 00:01:54.390 \longrightarrow 00:01:56.940$ especially I think most of our audience here are students,
- $31\ 00:01:56.940 --> 00:01:59.550$ I want to give you this motivation.
- 32 00:01:59.550 --> 00:02:02.040 When I was a student back in 2007,
- $33\ 00:02:02.040 --> 00:02:05.340$ that was when I just started my PhD
- $34\ 00:02:05.340 \longrightarrow 00:02:08.010$ and I was interested in bioinformatics.
- $35~00{:}02{:}08.010 \dashrightarrow 00{:}02{:}11.280~\mathrm{I}$ had a lot of questions about bioinformatics methods
- $36\ 00:02:11.280 \longrightarrow 00:02:13.740$ after I took statistics classes.
- $37\ 00:02:13.740 \longrightarrow 00:02:16.020$ I think some of the questions I listed here
- 38 00:02:16.020 --> 00:02:18.540 include are P values valid?
- $39\ 00:02:18.540 --> 00:02:20.910$ Because P values are so widely used
- $40\ 00:02:20.910 \longrightarrow 00:02:22.800$ in genomics bioinformatics.
- $41\ 00:02:22.800 \longrightarrow 00:02:24.930$ And also, we have a lot
- $42\ 00:02:24.930 --> 00:02:28.380$ of bio bioinformatics methods developed for data analysis.
- $43~00{:}02{:}28.380 \dashrightarrow 00{:}02{:}31.680$ And I wonder why don't we use classical statistical methods
- 44 00:02:31.680 --> 00:02:32.550 in textbooks?
- 45 00:02:32.550 --> 00:02:33.840 And the third thing is,
- $46~00:02:33.840 \longrightarrow 00:02:38.280$ when we use statistical test to understand the question,
- $47\ 00:02:38.280 \longrightarrow 00:02:39.750$ to answer some pivot question,
- $48\ 00:02:39.750 \longrightarrow 00:02:41.880$ what is the proper null hypothesis?
- $49\ 00:02:41.880 --> 00:02:44.760$ So you will see those questions in the topics
- $50\ 00:02:44.760 --> 00:02:46.710$ I will talk about next.
- 51~00:02:46.710 --> 00:02:51.510 So this talk will focus on the multiple testing problem.

- 52 00:02:51.510 --> 00:02:53.220 See, multiple testing, what it means
- 53 00:02:53.220 --> 00:02:56.730 is that we have multiple hypothesis tests,
- $54\ 00:02:56.730 \longrightarrow 00:03:00.660$ and the criteria we use in this problem are P values,
- $55\ 00:03:00.660 \longrightarrow 00:03:04.170$ which we have one P value per test.
- 56~00:03:04.170 --> 00:03:07.620 So we know that the requirement for a valid P value
- $57\ 00{:}03{:}07.620$ --> $00{:}03{:}11.490$ is that P values should follow the uniform distribution
- $58~00:03:11.490 \dashrightarrow 00:03:14.460$ between zero one under the null hypothesis.
- $59\ 00:03:14.460 --> 00:03:17.640$ Or we may relax this to be super uniform.
- 60 00:03:17.640 --> 00:03:18.900 Just for your information,
- $61\ 00:03:18.900 \longrightarrow 00:03:21.780$ super uniform means that the P values
- $62\ 00:03:21.780 --> 00:03:24.900$ have higher density toward one
- $63\ 00:03:24.900 \longrightarrow 00:03:26.430$ and lower density towards zero.
- 64 00:03:26.430 --> 00:03:29.760 So that's still okay for type one error control,
- $65\ 00:03:29.760 \longrightarrow 00:03:31.290$ even though you may have a larger
- $66\ 00:03:31.290 \longrightarrow 00:03:33.000$ than expected type two error.
- 67 00:03:33.000 --> 00:03:35.010 So given the many, many P values,
- $68~00:03:35.010 \dashrightarrow 00:03:39.900$ we need one criterion to set a cutoff on the P values.
- $69\ 00:03:39.900 \longrightarrow 00:03:41.970$ And the most commonly used criterion
- 70 00:03:41.970 --> 00:03:43.800 for multiple testing correction
- 71 00:03:43.800 --> 00:03:46.950 is called a false discovery rate, short as FPR.
- $72\ 00:03:46.950 \longrightarrow 00:03:51.600$ So the definition here is the expectation of this ratio,
- $73\ 00:03:51.600 \longrightarrow 00:03:55.410$ and this ratio is the number of false discoveries
- $74\ 00:03:55.410 \longrightarrow 00:03:57.270$ over the number of discoveries.
- $75\ 00:03:57.270 --> 00:04:00.090$ So this notation means the maximum
- $76\ 00:04:00.090 \longrightarrow 00:04:02.280$ between the number of discoveries and one.
- $77\ 00:04:02.280 \longrightarrow 00:04:05.640$ In other words, we don't allow the denominator to be zero,
- $78\ 00:04:05.640 \longrightarrow 00:04:07.200$ if we don't make any discovery.

- $79\ 00:04:07.200 --> 00:04:09.750$ So this is to avoid the dividing zero issue.
- $80\ 00{:}04{:}09.750 --> 00{:}04{:}13.650$ And this ratio has a name called false discovery proportion.
- $81\ 00:04:13.650 \longrightarrow 00:04:15.720$ In other words, we can have this proportion
- $82\ 00:04:15.720 \longrightarrow 00:04:18.030$ for one particular data set.
- 83 00:04:18.030 --> 00:04:21.660 However, as you know, we don't observe this ratio
- 84~00:04:21.660 --> 00:04:24.510 because we don't know which discoveries are false.
- $85\ 00{:}04{:}24.510 \dashrightarrow 00{:}04{:}27.780$ So therefore, this ratio is only a hypothetical concept,
- $86\ 00:04:27.780 \longrightarrow 00:04:30.420$ but not really computable.
- $87\ 00:04:30.420 \longrightarrow 00:04:31.920$ And here, the expectation
- $88\ 00:04:31.920 \longrightarrow 00:04:35.100$ is taken over all possible data set
- $89\ 00:04:35.100 \longrightarrow 00:04:38.130$ from the same distribution as our data set.
- 90 00:04:38.130 --> 00:04:40.260 So this is the frequentist concept
- 91 00:04:40.260 --> 00:04:43.980 because we have imaginary potential data sets.
- $92\ 00:04:43.980 --> 00:04:46.950$ So therefore, the phenomena paper
- 93 00:04:46.950 --> 00:04:49.830 by Benjamini and Hochburg gave us a way
- 94 00:04:49.830 --> 00:04:53.280 to control this expectation called FDR
- 95 00:04:53.280 --> 00:04:56.730 under a claimed level, say, 5%,
- $96\ 00:04:56.730 \longrightarrow 00:05:00.600$ even though we couldn't realize this ratio itself.
- $97\ 00:05:00.600 \longrightarrow 00:05:02.400$ But we could control its expectation.
- $98\ 00:05:02.400 \longrightarrow 00:05:04.830$ So that's the magic of statistics.
- 99 $00:05:04.830 \longrightarrow 00:05:07.020$ So Benjamini Hochburg algorithm allows us
- $100\ 00{:}05{:}07.020 \dashrightarrow 00{:}05{:}11.190$ to set a cutoff on the P values to control the FDR.
- 101~00:05:11.190 --> 00:05:14.790 But I want to emphasize that the FDS's only controlled
- 102 00:05:14.790 --> 00:05:17.280 when P values satisfy this assumption,
- $103\ 00:05:17.280 \longrightarrow 00:05:19.320$ otherwise, it may not be.
- $104\ 00{:}05{:}19.320$ --> $00{:}05{:}24.320$ So I want to say three common causes of ill-posed P values,

- $105\ 00:05:24.360 --> 00:05:27.480$ which make P values don't satisfy this assumption
- $106\ 00{:}05{:}27.480 \dashrightarrow 00{:}05{:}30.217$ in genomics, and I'll go through them one by one.
- $107\ 00:05:31.110 \longrightarrow 00:05:34.170$ The first issue is what I call the formulation
- $108\ 00{:}05{:}34.170 \dashrightarrow 00{:}05{:}37.740$ of a two sample test problem as a one sample test.
- $109\ 00:05:37.740 \longrightarrow 00:05:39.060$ What does this mean?
- $110\ 00:05:39.060 --> 00:05:42.090$ So I will use the common genomic analysis
- $111\ 00:05:42.090 --> 00:05:44.670$ of ChIP-seq data as an example.
- 112 00:05:44.670 --> 00:05:45.990 So in ChIP-seq data,
- $113\ 00:05:45.990 \longrightarrow 00:05:50.160$ we want to measure where a protein binds in the genome.
- $114\ 00:05:50.160 --> 00:05:53.520$ So you can consider the X axis as the genome
- $115\ 00:05:53.520 \longrightarrow 00:05:56.790$ and the Y axis as the protein binding intensity
- $116\ 00:05:56.790 \longrightarrow 00:05:58.680$ measured by ChIP-seq.
- 117 00:05:58.680 --> 00:06:02.070 So here, we have experimental sample,
- $118\ 00:06:02.070 --> 00:06:05.550$ the condition of our interest, say, a certain cell line.
- $119\ 00:06:05.550 \longrightarrow 00:06:08.040$ And the background sample is what we know
- 120 00:06:08.040 --> 00:06:09.660 that there's no protein,
- $121\ 00:06:09.660 --> 00:06:11.790$ so there should be no protein binding.
- $122\ 00{:}06{:}11.790 \dashrightarrow 00{:}06{:}15.420$ But we still want to measure the noise from the experiment.
- $123\ 00:06:15.420 \longrightarrow 00:06:17.430$ So we need this contrast.
- $124\ 00:06:17.430 \longrightarrow 00:06:21.900$ And here, we want to say that the region in the red box,
- $125\ 00:06:21.900 \longrightarrow 00:06:25.500$ this interval, we want to call it as a peak,
- $126\ 00{:}06{:}25.500 {\: \hbox{\scriptsize -->}\:} 00{:}06{:}29.550$ if we see the intensity in the experimental sample
- $127\ 00:06:29.550 \longrightarrow 00:06:32.790$ is much larger than the intensity in the background sample.
- $128\ 00{:}06{:}32.790 \dashrightarrow 00{:}06{:}35.940$ So we do the comparison and we want to cut this at a peak.

- $129\ 00:06:35.940 \longrightarrow 00:06:38.820$ That's the purpose of this analysis.
- 130 00:06:38.820 --> 00:06:41.550 And I wanna say that, in the field,
- $131\ 00:06:41.550 --> 00:06:45.390$ because ChIP-seq has become popular since 2008,
- $132\ 00{:}06{:}45.390 \dashrightarrow 00{:}06{:}49.290$ Macs and Homer are probably the two most popular software
- $133\ 00:06:49.290 \longrightarrow 00:06:50.940$ for cutting peaks.
- $134\ 00{:}06{:}50.940 \dashrightarrow 00{:}06{:}53.850$ Even though they have very complex procedures
- $135\ 00:06:53.850 --> 00:06:56.460$ for processing the sequencing data
- $136\ 00:06:56.460 \longrightarrow 00:06:58.380$ that in a statistical part
- $137\ 00:06:58.380 \longrightarrow 00:07:00.960$ to call a region as a peak or not,
- $138\ 00:07:00.960 \dashrightarrow 00:07:04.140$ I can say, their formulation is as follows.
- $139\ 00:07:04.140 --> 00:07:08.580$ Given a region, we count its number of ChIP-seq reads
- $140\ 00{:}07{:}08.580 \dashrightarrow 00{:}07{:}12.210$ in the background sample and in the experimental sample.
- 141 $00:07:12.210 \longrightarrow 00:07:15.450$ So let's just summarize this intensity as a count,
- $142\ 00:07:15.450 \longrightarrow 00:07:19.140$ a count here, a count here, and both are now negative.
- $143\ 00:07:19.140 --> 00:07:21.270$ So I call the background count as big X,
- $144\ 00:07:21.270 \longrightarrow 00:07:23.580$ experimental count as big Y.
- $145\ 00{:}07{:}23.580 {\:{\mbox{--}}\!>\:} 00{:}07{:}27.120$ And in our data, we have the observations, right?
- $146\ 00:07:27.120 \longrightarrow 00:07:30.180$ We refer to them as small x, small y.
- $147\ 00:07:30.180 --> 00:07:33.330$ Then, the P value in both software
- $148\ 00:07:33.330 \longrightarrow 00:07:36.840$ is essentially this probability, the probability
- $149\ 00:07:36.840 \longrightarrow 00:07:41.840$ that big Y is greater or equal than the observed small y,
- $150\ 00:07:41.910 \longrightarrow 00:07:45.240$ where the big Y follows upon some distribution
- $151\ 00:07:45.240 \longrightarrow 00:07:48.240$ with mean parameter as the small x.
- $152\ 00:07:48.240 --> 00:07:51.090$ Now, when I look at this formula back in 2008,

- $153\ 00:07:51.090 --> 00:07:54.633$ the Macs paper, I wonder whether this is correct.
- 154 00:07:55.620 --> 00:07:57.090 And I don't think so.
- 155 00:07:57.090 --> 00:07:58.950 Because the reason, if you look at it,
- $156\ 00:07:58.950 \longrightarrow 00:08:00.900$ is what is the null hypothesis?
- 157 00:08:00.900 --> 00:08:03.990 The null hypothesis is essentially, okay,
- $158\ 00:08:03.990 --> 00:08:05.700$ let's assume the experimental count
- 159 00:08:05.700 --> 00:08:08.610 is our test statistic, okay?
- $160\ 00:08:08.610 \longrightarrow 00:08:11.310$ We assume it follows a Poisson distribution
- $161\ 00:08:11.310 \longrightarrow 00:08:12.960$ with mean lambda.
- $162\ 00:08:12.960$ --> 00:08:17.960 And here, the null hypothesis is lambda is equal to small x.
- 163 00:08:18.090 --> 00:08:21.150 Alternative is lambda greater than small x.
- $164\ 00:08:21.150 \longrightarrow 00:08:23.160$ So what's the problem with here?
- $165~00{:}08{:}23.160 \dashrightarrow 00{:}08{:}27.240$ Essentially, we are using small x as a fixed parameter
- $166\ 00:08:27.240 \longrightarrow 00:08:29.280$ instead of a random observation.
- $167\ 00:08:29.280 \longrightarrow 00:08:30.270$ So in other words,
- $168\ 00:08:30.270 \longrightarrow 00:08:33.390$ the randomness in the background count is ignored.
- $169\ 00{:}08{:}33.390 --> 00{:}08{:}36.720$ We only consider experimental count as the random variable.
- $170\ 00{:}08{:}36.720 \dashrightarrow 00{:}08{:}39.990$ So in other words, where use the two sample testing problem
- $171\ 00:08:39.990 \longrightarrow 00:08:41.970$ to a one sample testing problem
- $172\ 00:08:41.970 --> 00:08:43.637$ because we only consider the randomness
- $173\ 00:08:43.637 --> 00:08:45.960$ in the experimental sample.
- $174\ 00:08:45.960 --> 00:08:50.040$ But this is not something our textbook teaches us.
- $175\ 00{:}08{:}50.040 \dashrightarrow 00{:}08{:}52.830$ The reason is because if we consider background
- $176\ 00:08:52.830 \longrightarrow 00:08:56.130$ as one condition, experimental has another condition,

 $177\ 00:08:56.130 --> 00:08:59.640$ under each condition, our sample size is only one.

178 00:08:59.640 --> 00:09:02.070 So therefore, the T test will not apply

 $179\ 00:09:02.070 \longrightarrow 00:09:04.950$ because a central limit here clearly doesn't apply.

180 00:09:04.950 --> 00:09:08.733 So how do we calculate P value, any ideas?

181 00:09:09.600 --> 00:09:12.750 I think one possibility that we could still assume

 $182\ 00:09:12.750 \longrightarrow 00:09:15.660$ Poisson distribution for both background X

 $183\ 00:09:15.660 \longrightarrow 00:09:16.830$ and experimental Y.

 $184\ 00:09:16.830 --> 00:09:20.520$ You have two Poisson, under the independence,

 $185\ 00:09:20.520 --> 00:09:22.560$ we can probably derive the distribution

186 00:09:22.560 --> 00:09:25.620 for Y minus X, right, and what's the null distribution.

187 00:09:25.620 --> 00:09:26.760 That's the only way.

188 00:09:26.760 --> 00:09:30.000 But, if you think about it, how can we verify

 $189\ 00{:}09{:}30.000 {\:{\mbox{--}}\!>\:} 00{:}09{:}32.583$ whether the Poisson distribution is reasonable?

190 00:09:32.583 --> 00:09:34.890 You only have one observation from it.

191 00:09:34.890 --> 00:09:37.140 The distribution could be anything, right?

 $192\ 00:09:37.140 --> 00:09:40.950$ So assuming a parametric distribution seems quite,

193 00:09:40.950 --> 00:09:42.300 I will say, aggressive.

 $194~00:09:42.300 \dashrightarrow 00:09:45.090$ So I think P value calculation is challenging here.

195 00:09:45.090 --> 00:09:48.600 And also, I even wonder, in this case,

 $196\ 00:09:48.600 \longrightarrow 00:09:51.000$ for this one versus one comparison,

197 00:09:51.000 --> 00:09:53.100 should we use a P value?

 $198\ 00:09:53.100 \dashrightarrow 00:09:57.060$ Or is this really a testing problem that's feasible?

 $199~00:09:57.060 \dashrightarrow 00:09:59.940$ So I would say, over the years, I gradually realized

200 00:09:59.940 --> 00:10:02.670 that here we looked at many, many regions,

- $201\ 00:10:02.670 \longrightarrow 00:10:04.230$ not just one region.
- $202\ 00{:}10{:}04.230 \dashrightarrow 00{:}10{:}08.160$ So the goal or the criterion that's ultimately used
- 203 00:10:08.160 --> 00:10:09.240 is actually FDR.
- 204 00:10:09.240 --> 00:10:12.180 And in this process,
- 205 00:10:12.180 --> 00:10:15.960 P values are just intermediate for FDR control,
- 206 00:10:15.960 --> 00:10:18.180 instead of our final target.
- 207 00:10:18.180 --> 00:10:21.090 So do we have to stick with P values?
- 208~00:10:21.090 --> 00:10:25.110 This motivated me to write this paper with my students
- $209\ 00{:}10{:}25.110 \dashrightarrow 00{:}10{:}30.110$ to propose a way to achieve p-value-free FDR control
- $210\ 00:10:30.180 \dashrightarrow 00:10:34.230$ by leveraging the theory in Barber and Candes paper,
- 211 00:10:34.230 --> 00:10:35.610 their knockoff paper,
- $212\ 00:10:35.610 --> 00:10:38.580$ so we could actually doing FDR control
- 213 00:10:38.580 --> 00:10:41.190 in this example without using P value.
- 214 00:10:41.190 --> 00:10:43.170 So I will talk about this later in my talk,
- $215\ 00:10:43.170 \longrightarrow 00:10:46.830$ but this is one motivation for the Clipper paper.
- $216\ 00:10:46.830 \longrightarrow 00:10:49.950$ The second issue with P values is that we observe,
- $217\ 00:10:49.950 --> 00:10:51.680$ sometimes, P values are not valid
- $218\ 00:10:51.680$ --> 00:10:56.680 because the parametric model used may not fit the data well.
- 219 00:10:57.000 --> 00:11:00.600 So this is an example for this commonly used
- $220\ 00{:}11{:}00.600 \dashrightarrow 00{:}11{:}04.620$ differential expression analysis on RNA sequencing data.
- 221 00:11:04.620 --> 00:11:06.600 So for this task,
- $222\ 00{:}11{:}06.600 \dashrightarrow 00{:}11{:}09.750$ the two popular softwares are DESeq2 and edgeR.
- $223\ 00:11:09.750 \longrightarrow 00:11:12.420$ So the data usually looks like this.
- $224\ 00:11:12.420 \longrightarrow 00:11:15.390$ So we want to compare two conditions

- $225\ 00{:}11{:}15.390 \dashrightarrow 00{:}11{:}18.780$ and seeing which genes are differentially expressed.
- 226 00:11:18.780 --> 00:11:21.630 So condition one, we have three samples,
- 227 00:11:21.630 --> 00:11:23.400 which we cause to replicate,
- $228\ 00:11:23.400 \longrightarrow 00:11:25.410$ condition two, three replicates.
- 229 00:11:25.410 --> 00:11:29.070 So every row is one replicate,
- 230 00:11:29.070 --> 00:11:31.170 while every column is one gene.
- 231 00:11:31.170 --> 00:11:33.900 So to call a gene as differentially expressed,
- $232\ 00:11:33.900 \longrightarrow 00:11:36.000$ we need to compare its three values
- $233\ 00:11:36.000$ --> 00:11:39.150 from condition one, two, three values from condition two.
- $234\ 00{:}11{:}39.150 \dashrightarrow 00{:}11{:}42.960$ So clearly, we can see the left one may be a D gene,
- 235 00:11:42.960 --> 00:11:45.060 the right one may not be a D gene, right?
- 236 00:11:45.060 --> 00:11:46.170 That's our intuition.
- 237 00:11:46.170 --> 00:11:49.470 And we want to make this more formal
- $238\ 00:11:49.470 \longrightarrow 00:11:51.854$ by doing a statistical test.
- 239 00:11:51.854 --> 00:11:54.660 But in both edgeR and DESeq2,
- $240\ 00:11:54.660 \longrightarrow 00:11:58.590$ you can see that to compensate the small sample size,
- $241\ 00:11:58.590 \longrightarrow 00:12:00.120$ like three versus three,
- $242\ 00:12:00.120 \dots > 00:12:05.120$ they assume a gene follows a negative binomial distribution
- $243\ 00:12:05.460 \longrightarrow 00:12:06.630$ under each condition.
- $244\ 00:12:06.630 \longrightarrow 00:12:09.300$ So essentially, these three values are assumed
- $245\ 00:12:09.300 \longrightarrow 00:12:12.120$ to follow one negative binomial distribution.
- 246 00:12:12.120 --> 00:12:13.380 These three values
- $247\ 00:12:13.380 \longrightarrow 00:12:16.170$ follow another negative binomial distribution.
- 248 00:12:16.170 --> 00:12:17.640 And the null hypothesis
- $249\ 00:12:17.640 \longrightarrow 00:12:20.700$ is the two negative binomial distributions
- $250\ 00:12:20.700 \longrightarrow 00:12:23.550$ have the same mean, that's the problem.
- 251 00:12:23.550 --> 00:12:27.090 Okay, so we actually discovered an issue

- $252\ 00:12:27.090 \longrightarrow 00:12:29.850$ with popular methods from this data set.
- 253~00:12:29.850 --> 00:12:32.306 And thanks to my collaborator Dr. Wei Li
- 254 00:12:32.306 --> 00:12:35.520 who is a computation of biologist at UC Irvine.
- 255 00:12:35.520 --> 00:12:39.480 So actually, from this patient data,
- 256 00:12:39.480 --> 00:12:43.110 we have a much larger sample size, 51 patients
- $257\ 00:12:43.110 \longrightarrow 00:12:46.645$ before the treatment of some immunotherapy medicine,
- 258 00:12:46.645 --> 00:12:49.620 58 patients on treatment.
- $259~00{:}12{:}49.620 \dashrightarrow 00{:}12{:}52.680$ So we want to compare the RNA sequencing data
- $260\ 00:12:52.680 \longrightarrow 00:12:54.930$ of these two groups of patients.
- 261 00:12:54.930 --> 00:12:59.930 So essentially, when we apply DESeq2 or edgeR to this data,
- $262~00{:}13{:}00.840 \dashrightarrow 00{:}13{:}05.253$ the red dots indicate the number of D genes identified.
- $263\ 00:13:06.300 \longrightarrow 00:13:11.300$ To verify whether we can still identify D genes
- 264 00:13:11.640 --> 00:13:12.840 from permuted data,
- $265\ 00:13:12.840 \longrightarrow 00:13:15.150$ because the reason is that we want to see
- $266\ 00:13:15.150 \longrightarrow 00:13:18.780$ whether the permuted data is actually really,
- 267 00:13:18.780 --> 00:13:20.220 because we know the permuted data
- 268 00:13:20.220 --> 00:13:21.840 shouldn't give us any signals.
- $269\ 00:13:21.840 \longrightarrow 00:13:23.820$ If we just disrupt the two groups,
- $270\ 00:13:23.820 \longrightarrow 00:13:25.830$ we shouldn't expect any D genes.
- 271 00:13:25.830 --> 00:13:29.070 But surprisingly, we found that each method
- $272\ 00{:}13{:}29.070 \dashrightarrow 00{:}13{:}33.540$ can identify sometimes even more D genes from permuted data.
- $273\ 00{:}13{:}33.540 \dashrightarrow 00{:}13{:}37.230$ So the bar and the error bars show the distribution
- 274 00:13:37.230 --> 00:13:40.350 of D genes identified from permuted data.
- $275\ 00:13:40.350 \longrightarrow 00:13:43.710$ So this is something quite unexpected.
- 276 00:13:43.710 --> 00:13:46.530 And to look into the reason, our first thought
- 277 00:13:46.530 --> 00:13:49.920 is to check the negative binomial assumption.
- 278 00:13:49.920 --> 00:13:51.780 Because now, under each group,

- 279 00:13:51.780 --> 00:13:54.900 we have 51 and 58 sample sizes,
- $280\ 00:13:54.900 \longrightarrow 00:13:58.680$ so we could check the distribution, and here's what we get.
- $281\ 00{:}13{:}58.680 \dashrightarrow 00{:}14{:}02.753$ You see that for the genes that are frequently identified
- $282\ 00:14:02.753 \dashrightarrow 00:14:06.990$ from permuted data, if we run the goodness-of-fit test,
- 283 00:14:06.990 --> 00:14:09.870 we check the negative binomial distribution,
- 284 00:14:09.870 --> 00:14:12.360 these genes have very small P values,
- $285\ 00:14:12.360 \longrightarrow 00:14:15.090$ indicating that this fit is not good.
- $286\ 00:14:15.090 --> 00:14:16.350$ Well, if you look at the genes
- 287 00:14:16.350 --> 00:14:19.950 that are rarely identified from permuted data,
- $288\ 00:14:19.950 --> 00:14:22.860$ the P values are bigger and the goodness-of-fit is better.
- $289\ 00:14:22.860 \longrightarrow 00:14:25.200$ So we do see this relationship
- $290\ 00{:}14{:}25.200 \dashrightarrow 00{:}14{:}28.740$ between the goodness-of-fit of negative binomial
- $291\ 00:14:28.740 --> 00:14:31.590$ and the frequency that a gene is identified
- $292\ 00:14:31.590 \longrightarrow 00:14:33.240$ from permuted data.
- $293\ 00{:}14{:}33.240 \dashrightarrow 00{:}14{:}36.480$ So negative binomial model seems to not fit well
- $294\ 00:14:36.480 \longrightarrow 00:14:39.030$ on this patient data.
- $295~00{:}14{:}39.030$ --> $00{:}14{:}42.090$ Because here, the 51 patients shouldn't be regarded
- $296\ 00:14:42.090 \longrightarrow 00:14:44.700$ as replicates, they're not experimental replicates,
- $297\ 00:14:44.700 \longrightarrow 00:14:46.110$ they are individuals.
- $298~00{:}14{:}46.110 \dashrightarrow 00{:}14{:}49.590$ So therefore, the theory for deriving negative binomials
- $299~00{:}14{:}49.590 \dashrightarrow 00{:}14{:}52.440$ usually assume as a Gamma-Poisson Mixture model,
- $300~00:14:52.440 \dashrightarrow 00:14:54.180$ Gamma-Poisson Hierarchical model.
- 301 00:14:54.180 --> 00:14:56.580 That one may no longer hold,
- $302\ 00:14:56.580 --> 00:14:59.610$ and that's why we think the parametric model

- 303~00:14:59.610 --> 00:15:03.510 is not applicable to this patient data.
- $304\ 00:15:03.510 \longrightarrow 00:15:05.580$ So what's the consequence, right?
- $305\ 00:15:05.580 \longrightarrow 00:15:07.650$ So we want to convince the scientist
- $306\ 00:15:07.650 \longrightarrow 00:15:10.530$ what's the consequence of doing this analysis
- $307\ 00:15:10.530 \longrightarrow 00:15:12.090$ in this problematic way.
- $308\ 00:15:12.090 \longrightarrow 00:15:14.910$ We show that if we just use the D genes
- $309\ 00:15:14.910 \longrightarrow 00:15:17.070$ found by DESeq2 and edgeR,
- $310\ 00:15:17.070 \longrightarrow 00:15:20.233$ which are the genes corresponding to the red dot,
- 311 00:15:20.233 --> 00:15:23.460 around the so called gene oncology analysis,
- $312\ 00:15:23.460 \longrightarrow 00:15:26.130$ that is to check which functional terms
- $313\ 00:15:26.130 \longrightarrow 00:15:29.370$ are enriched in those two gene sets,
- 31400:15:29.370 --> 00:15:31.350 we can see many functional terms
- $315\ 00:15:31.350 \longrightarrow 00:15:33.510$ are related to immune functions.
- $316\ 00:15:33.510 --> 00:15:35.730$ Which would suggest that if we trust
- $317\ 00{:}15{:}35.730 \dashrightarrow 00{:}15{:}38.820$ these two methods' results, we may conclude that,
- 318 00:15:38.820 --> 00:15:41.400 yes, between the two groups of patients,
- $319\ 00:15:41.400 \longrightarrow 00:15:44.430$ there are differences in immune responses, right?
- $320\ 00{:}15{:}44.430 \dashrightarrow 00{:}15{:}48.030$ That seems to confirm our scientific hypothesis.
- 321 00:15:48.030 --> 00:15:50.610 However, now, we see many of these genes
- 322 00:15:50.610 --> 00:15:53.790 were also identified from permuted data,
- $323\ 00:15:53.790 \longrightarrow 00:15:57.120$ then, that will make the results dubious.
- $324\ 00{:}15{:}57.120 --> 00{:}16{:}01.470$ So what we tried is that, even the sample size is so large,
- 325 00:16:01.470 --> 00:16:03.690 we tried the classical Wilcoxon rank sign test,
- 326 00:16:03.690 --> 00:16:05.240 which everybody learned, right?
- $327\ 00:16:06.119 --> 00:16:08.310$ So non parametric two sample test
- $328\ 00:16:08.310 \longrightarrow 00:16:11.130$ that doesn't assume a parametric distribution.
- 329 00:16:11.130 --> 00:16:13.080 And here, it's self consistent,

- 330 00:16:13.080 --> 00:16:16.590 it doesn't identify D genes from real data,
- 331 00:16:16.590 --> 00:16:20.040 but also, it doesn't identify D genes from permuted data.
- $332\ 00:16:20.040 \longrightarrow 00:16:22.650$ So there's no contradiction here.
- $333\ 00:16:22.650 \longrightarrow 00:16:25.860$ And this result motivated me to ask this question,
- $334\ 00:16:25.860 \longrightarrow 00:16:27.603$ which I had years ago,
- $335\ 00:16:28.590 \longrightarrow 00:16:32.730$ should we always use popular bioinformatics tools?
- 336 00:16:32.730 --> 00:16:35.010 Like, check the citation of these two methods,
- $337\ 00:16:35.010 \longrightarrow 00:16:36.213$ super highly cited.
- 338 00:16:37.080 --> 00:16:39.150 Should I reuse popular method
- $339\ 00{:}16{:}39.150 \dashrightarrow 00{:}16{:}42.990$ or should we consider general statistical methods,
- 340 00:16:42.990 --> 00:16:44.760 like Wilcoxon.
- $341\ 00:16:44.760 --> 00:16:49.650$ So our recommendation is sample size matters, right?
- $342\ 00:16:49.650 \longrightarrow 00:16:52.380$ We may have different methods
- 343 00:16:52.380 --> 00:16:54.660 suitable for different sample sizes,
- $344\ 00:16:54.660 \longrightarrow 00:16:57.510$ and essentially, why statistics has so many methods,
- 345 00:16:57.510 --> 00:16:58.923 paramedic, non parametric,
- $346\ 00:16:59.910$ --> 00:17:02.880 is because we have different scenarios in our data.
- 347 00:17:02.880 --> 00:17:04.740 That's the first thing we should realize.
- 348 00:17:04.740 --> 00:17:07.500 It's not like one method can do all the things.
- $349\ 00:17:07.500 \longrightarrow 00:17:10.140$ And the second thing is sanity check.
- $350\ 00{:}17{:}10.140 \dashrightarrow 00{:}17{:}12.770$ We should always consider doing some sanity check
- $351\ 00:17:12.770 --> 00:17:14.760$ to make sure we trust the results
- $352\ 00:17:14.760 \longrightarrow 00:17:17.460$ instead of just take the results for granted.
- $353\ 00:17:17.460 \longrightarrow 00:17:20.370$ So these things were summarized in our paper
- $354\ 00:17:20.370 \longrightarrow 00:17:22.920$ published earlier this year.

- 355 00:17:22.920 --> 00:17:24.660 And since its publication,
- $356\ 00:17:24.660 \longrightarrow 00:17:27.960$ we have received a lot of discussions on Twitter,
- $357\ 00:17:27.960 \longrightarrow 00:17:29.010$ if you are interested.
- $358\ 00{:}17{:}29.010 \dashrightarrow 00{:}17{:}31.800$ But anyway, so it means that many people are interested
- $359\ 00{:}17{:}31.800 \dashrightarrow 00{:}17{:}35.940$ in this topic, especially many people, users believe
- $360\ 00:17:35.940 \longrightarrow 00:17:39.377$ that popular bioinformatics tools are the state-of-the-art,
- 361 00:17:39.377 --> 00:17:41.985 right, the way, standard methods (indistinct).
- $362\ 00{:}17{:}41.985 --> 00{:}17{:}45.420$ But if you are bio statisticians, you may not like this.
- $363\ 00:17:45.420 \longrightarrow 00:17:47.760$ Because we want to develop new methods.
- 364 00:17:47.760 --> 00:17:49.500 Otherwise, what's our job, right?
- $365\ 00:17:49.500 \longrightarrow 00:17:53.400$ So in this case, we need to really find the loopholes,
- $366\ 00{:}17{:}53.400 {\:\hbox{--}}{>}\ 00{:}17{:}57.090$ or the limitations, or the gap between current approach
- $367\ 00:17:57.090 \longrightarrow 00:17:58.410$ and the data scenarios,
- 368 00:17:58.410 --> 00:18:00.900 and try convinces people that, yes,
- $369~00:18:00.900 \dashrightarrow 00:18:03.570$ we do need careful thoughts when we choose method.
- $370\ 00:18:03.570 \longrightarrow 00:18:06.240$ It's not always one method.
- 371 00:18:06.240 --> 00:18:08.280 And a related question is,
- $372\ 00{:}18{:}08.280 \dashrightarrow 00{:}18{:}12.720$ in Wilcoxon, definitely doesn't have a strong assumption,
- $373\ 00:18:12.720 --> 00:18:15.120$ and (indistinct) have a reasonable power
- $374\ 00:18:15.120 \longrightarrow 00:18:16.920$ when the sample size is large.
- 375 00:18:16.920 --> 00:18:19.770 But what if sample sizes are small, right?
- $376\ 00:18:19.770 \longrightarrow 00:18:21.450$ So when it's small, we know,
- 377 00:18:21.450 --> 00:18:24.750 non parametric tests like Wilcoxon doesn't have power.

 $378\ 00{:}18{:}24.750 \dashrightarrow 00{:}18{:}29.670$ So in this case, we actually proposed Clipper again,

 $379\ 00{:}18{:}29.670 \dashrightarrow 00{:}18{:}34.050$ so it can work as a downstream correction tool

 $380\ 00:18:34.050 \longrightarrow 00:18:36.300$ for DESeq2 and edgeR.

 $381\ 00:18:36.300 --> 00:18:38.700$ Because they are supposed to be quite powerful,

382 00:18:38.700 --> 00:18:41.010 even though they find probably too many.

 $383\ 00:18:41.010 \longrightarrow 00:18:44.190$ So hopefully, we could use that to borrow their power,

 $384\ 00:18:44.190 \longrightarrow 00:18:47.310$ but help them improve the FDR control.

 $385\ 00:18:47.310 \longrightarrow 00:18:50.310$ So I'll show the results later in my talk.

 $386\ 00:18:50.310 \longrightarrow 00:18:51.630$ That's the second cause.

387 00:18:51.630 --> 00:18:53.760 And the third cause for ill-posed P values

 $388\ 00:18:53.760 --> 00:18:55.950$ is a little more complicated.

 $389\ 00:18:55.950 \longrightarrow 00:18:59.670$ And this is the issue commonly observed in single cell data,

 $390\ 00:18:59.670 \longrightarrow 00:19:01.080$ single cell RNA-seq data.

 $391\ 00:19:01.080 \longrightarrow 00:19:02.910$ So I will use this analysis

 $392\ 00:19:02.910$ --> 00:19:07.910 called pseudotime differentially expressed genes as example.

 $393\ 00:19:08.190 \longrightarrow 00:19:09.858$ What is a pseudotime?

394 00:19:09.858 --> 00:19:13.110 Pseudotime means it's not real time, it's pseudo, right?

395 00:19:13.110 --> 00:19:15.720 So it's something we inferred

396 00:19:15.720 --> 00:19:17.670 from single cell RNA-seq data,

 $397\ 00:19:17.670 \longrightarrow 00:19:20.430$ so those cells are measured all at once.

398 00:19:20.430 \rightarrow 00:19:25.430 But we want to infer some time trajectory from the cells.

399 00:19:25.920 --> 00:19:28.830 So I'll just use the screenshot from Slingshot,

 $400\ 00{:}19{:}28.830 \dashrightarrow 00{:}19{:}33.830$ which is a method for inferring pseudotime for explanation.

 $401\ 00:19:34.050 \dashrightarrow 00:19:39.050$ So here, this is a two-dimensional PCA plot of cells,

- $402\ 00:19:39.180 \longrightarrow 00:19:41.280$ and the cells are pre-clustered,
- $403\ 00:19:41.280 \longrightarrow 00:19:44.250$ so each color represents one cluster.
- 404 00:19:44.250 --> 00:19:47.100 So the Slingshot algorithm does the following,
- 405 00:19:47.100 --> 00:19:50.610 first, it takes the cluster means' centers,
- $406\ 00:19:50.610 \longrightarrow 00:19:52.770$ and connect them using the algorithm
- 407 00:19:52.770 --> 00:19:54.450 called minimum spanning tree.
- $408\ 00:19:54.450 \longrightarrow 00:19:55.730$ So if you're not familiar with that,
- 409 00:19:55.730 --> 00:19:59.370 it has an equivalence with hierarchical clustering actually.
- $410\ 00:19:59.370 --> 00:20:02.400$ So with the minimum spanning tree, you get this tree,
- 411 00:20:02.400 --> 00:20:06.840 and then, they smooth out the tree using principle curves.
- $412\ 00:20:06.840 \longrightarrow 00:20:08.130$ So we have two curves,
- $413\ 00:20:08.130 \longrightarrow 00:20:09.810$ and then for every cell,
- $414\ 00:20:09.810$ --> 00:20:13.080 we find the closest curve and project the cell to the curve.
- 415 00:20:13.080 --> 00:20:14.970 So therefore, in each curve,
- $416\ 00:20:14.970 \longrightarrow 00:20:18.090$ the projections are called pseudotime values.
- $417\ 00:20:18.090 \longrightarrow 00:20:21.060$ And usually, it's normalized between zero and one,
- $418\ 00:20:21.060 \longrightarrow 00:20:23.580$ so we need to find the root and call it zero,
- $419\ 00:20:23.580 \longrightarrow 00:20:25.410$ the other end is called one.
- $420\ 00:20:25.410 --> 00:20:28.260$ So this whole process is called pseudotime inference.
- 421 00:20:28.260 --> 00:20:31.680 In other words, after it, we will give every cell
- $422\ 00:20:31.680 \longrightarrow 00:20:35.250$ a pseudotime value in each trajectory.
- $423~00{:}20{:}35.250 \longrightarrow 00{:}20{:}37.650$ Okay, so one thing I want to emphasize
- $424\ 00:20:37.650 \longrightarrow 00:20:40.200$ is that in this pseudotime inference
- $425\ 00:20:40.200 \longrightarrow 00:20:43.470$ we used gene expression values already.
- $426\ 00:20:43.470 --> 00:20:46.860$ So it's not like we observe pseudotime as external variable,
- $427\ 00:20:46.860 \longrightarrow 00:20:48.930$ but it's from the same data.

- $428\ 00{:}20{:}48.930 \dashrightarrow 00{:}20{:}53.130$ So I want to show what we could do after the pseudotime.
- 429 00:20:53.130 --> 00:20:55.560 So a typical analysis is to identify
- 430 00:20:55.560 --> 00:20:57.870 which genes are differentially expressed
- $431\ 00:20:57.870 \longrightarrow 00:20:59.250$ along the pseudotime.
- $432\ 00:20:59.250 \longrightarrow 00:21:03.360$ Like the left one, we see, it has this upward trajectory,
- 433 00:21:03.360 --> 00:21:05.910 so we may call it differentially expressed.
- $434\ 00:21:05.910 \longrightarrow 00:21:08.970$ And here, we want to say the pseudotime
- $435\ 00:21:08.970 --> 00:21:11.910$ represent some cell immune response,
- 436 00:21:11.910 --> 00:21:13.560 and this is an immuno-related gene,
- $437\ 00:21:13.560 \longrightarrow 00:21:16.740$ so we expect to see the upward trajectory.
- $438\ 00:21:16.740 \longrightarrow 00:21:20.340$ For the right gene, we expect to see something constant,
- 439 00:21:20.340 --> 00:21:23.340 so we don't want to come right (indistinct) a D gene,
- $440\ 00:21:23.340 \longrightarrow 00:21:25.350$ that's the intuition.
- 441 00:21:25.350 --> 00:21:28.320 And I want to say that we must realize,
- 442 00:21:28.320 --> 00:21:31.050 pseudotime values are random
- $443\ 00{:}21{:}31.050 \dashrightarrow 00{:}21{:}34.770$ simply because the cells is a random sample, right?
- $444\ 00:21:34.770 \longrightarrow 00:21:36.600$ We need to consider randomness,
- $445\ 00{:}21{:}36.600 \dashrightarrow 00{:}21{:}40.770$ and we want to show this to people by doing subsampling.
- 446 00:21:40.770 --> 00:21:43.290 So you can see that sampling variation
- 447 00:21:43.290 --> 00:21:45.810 would get into pseudotime values.
- $448\ 00:21:45.810 \longrightarrow 00:21:47.880$ Here, every row is a cell.
- 449 00:21:47.880 --> 00:21:49.680 If I randomly subsample,
- $450\ 00:21:49.680 \longrightarrow 00:21:53.310$ say, 80% of cells from the left cells
- 451 00:21:53.310 --> 00:21:56.760 and redo the pseudotime trajectory inference,
- $452\ 00:21:56.760 \longrightarrow 00:22:00.600$ we can see that for the cells in the subsamples
- $453\ 00{:}22{:}00.600 \dashrightarrow 00{:}22{:}04.380$ that include it, its values will vary to some degree.

- $454\ 00:22:04.380 \longrightarrow 00:22:06.630$ So it's not a constant.
- $455\ 00:22:06.630 --> 00:22:09.690$ Okay, so realizing this, we should consider
- $456\ 00:22:09.690 \longrightarrow 00:22:12.810$ the randomness of pseudotime from the data.
- $457\ 00{:}22{:}12.810 \dashrightarrow 00{:}22{:}15.930$ However, existing methods all treat pseudotime
- $458\ 00:22:15.930 \longrightarrow 00:22:17.790$ as an observed covariate.
- 459 00:22:17.790 --> 00:22:21.690 So our goal here is to fix this,
- $460\ 00:22:21.690 \longrightarrow 00:22:24.870$ and we proposed this method called PseudotimeDE,
- 461 00:22:24.870 --> 00:22:27.240 which actually does the inference,
- $462\ 00:22:27.240 \longrightarrow 00:22:29.460$ which infers whether one gene
- $463\ 00:22:29.460 \longrightarrow 00:22:32.310$ is differentially expressed along pseudotime,
- $464\ 00:22:32.310 \longrightarrow 00:22:36.450$ and by considering pseudotime inference uncertainty.
- 465 00:22:36.450 --> 00:22:40.620 So what we did exactly is that, here,
- $466\ 00:22:40.620 \longrightarrow 00:22:43.830$ to see whether a gene changes with pseudotime,
- $467\ 00:22:43.830 \longrightarrow 00:22:45.480$ what's the intuition?
- $468\ 00:22:45.480 --> 00:22:48.270$ We should do regression, right, do a regression analysis
- $469\ 00:22:48.270 --> 00:22:52.530$ by treating a gene's expression value as Y,
- 470~00:22:52.530 --> 00:22:54.660 pseudotime as X, and regular regression.
- $471\ 00:22:54.660 --> 00:22:57.540$ Yeah, this is exactly what existing methods did.
- $472\ 00:22:57.540 \longrightarrow 00:22:59.430$ And to make sure the regression
- $473\ 00:22:59.430 \longrightarrow 00:23:01.740$ is not restricted to be linear,
- $474\ 00{:}23{:}01.740 \dashrightarrow 00{:}23{:}04.920$ and also account for that the gene expression values
- $475\ 00:23:04.920 \longrightarrow 00:23:06.570$ are non negative counts.
- $476\ 00:23:06.570 \longrightarrow 00:23:11.570$ So actually, we choose the generalized additive model,
- 477 00:23:11.610 --> 00:23:14.100 which is also used in an existing method,
- 478 00:23:14.100 --> 00:23:15.750 which I will show very soon.

- $479\ 00{:}23{:}15.750 --> 00{:}23{:}20.400$ So this is a very flexible and interpretable model.
- $480~00:23:20.400 \longrightarrow 00:23:24.060$ So generalized means Y can be non Gaussian
- $481\ 00:23:24.060 \longrightarrow 00:23:25.230$ and the other distribution,
- 482 00:23:25.230 --> 00:23:27.510 just like generalized linear model.
- $483\ 00:23:27.510 \longrightarrow 00:23:28.740$ But additive means
- $484\ 00:23:28.740 \longrightarrow 00:23:32.340$ that we make the linear model more general,
- $485\ 00:23:32.340 \longrightarrow 00:23:36.553$ so every feature can be non linearly transformed,
- $486\ 00{:}23{:}37.860 \dashrightarrow 00{:}23{:}41.550$ but the features after transformations are still added.
- $487\ 00:23:41.550 \longrightarrow 00:23:44.340$ So that's additive, short as GAM.
- 488 00:23:44.340 --> 00:23:47.310 So essentially, once we have a set of cells,
- $489\ 00:23:47.310 \longrightarrow 00:23:49.350$ we first infer the pseudotime,
- 490 00:23:49.350 --> 00:23:52.170 so we order the cells along the pseudotime,
- $491\ 00:23:52.170 \longrightarrow 00:23:53.430$ and for gene J,
- $492\ 00{:}23{:}53.430 \dashrightarrow 00{:}23{:}56.550$ we check how the gene changes with pseudotime,
- $493\ 00:23:56.550 --> 00:23:59.760$ so we run the generalized additive model
- $494\ 00:23:59.760 \longrightarrow 00:24:01.500$ to obtain a test statistic.
- $495~00{:}24{:}01.500 \dashrightarrow 00{:}24{:}05.190$ Please know that generalized additive model has its theory,
- $496\ 00:24:05.190 \longrightarrow 00:24:08.250$ so we could use the theory to calculate
- $497\ 00:24:08.250 --> 00:24:11.817$ to use the null distribution and calculate P value.
- $498\ 00:24:11.817 \longrightarrow 00:24:14.670$ And that was done in an existing method.
- $499\ 00:24:14.670 --> 00:24:17.520$ We want say that this may be problematic
- $500\ 00:24:17.520 --> 00:24:19.784$ because this whole null distribution
- 501 00:24:19.784 --> 00:24:22.440 considers pseudotime to be fixed.
- 502 00:24:22.440 --> 00:24:23.820 So to address this,
- $503\ 00:24:23.820 --> 00:24:26.370$ we need to consider pseudotime inference
- $504\ 00:24:26.370 --> 00:24:30.060$ as part of our test statistic calculation.
- $505\ 00:24:30.060 \longrightarrow 00:24:33.090$ So to do this, we use the top part.

- $506\ 00:24:33.090 --> 00:24:35.023$ We actually do subsapling of the cells.
- $507\ 00:24:36.270 --> 00:24:38.430$ The reason we didn't do bootstrap
- $508~00{:}24{:}38.430 \dashrightarrow 00{:}24{:}41.280$ is simply because we want the method to be flexible
- $509\ 00:24:41.280 \longrightarrow 00:24:43.290$ for pseudotime inference method.
- $510\ 00:24:43.290 --> 00:24:47.070$ Like I show here, there are Slingshot, Monocle3,
- $511\ 00:24:47.070 \longrightarrow 00:24:48.300$ and a few others.
- 512 00:24:48.300 --> 00:24:49.800 We want it to be flexible,
- $513~00{:}24{:}49.800 \dashrightarrow 00{:}24{:}53.520$ and some methods don't allow cells to be repetitive,
- $514\ 00:24:53.520 \longrightarrow 00:24:55.710$ so bootstrap doesn't apply here.
- $515\ 00:24:55.710 \longrightarrow 00:24:59.370$ And we use subsampling with percentage pretty high,
- $516\ 00:24:59.370 --> 00:25:03.330$ like 80%, 90%, and we did a robustness analysis.
- $517\ 00:25:03.330 \longrightarrow 00:25:08.010$ And then, on each subsample, we do pseudotime inference.
- $518\ 00:25:08.010 --> 00:25:11.130$ With this, how do we get a null distribution
- $519\ 00:25:11.130 \longrightarrow 00:25:12.330$ of the test statistic?
- 520 00:25:12.330 --> 00:25:14.700 What we did is to permute the cells,
- 521 00:25:14.700 --> 00:25:17.730 so any relationship between the gene J
- $522\ 00:25:17.730 --> 00:25:19.830$ and the pseudotime is disrupted.
- 523 00:25:19.830 --> 00:25:21.563 So this can be considered from the null,
- 524 00:25:21.563 --> 00:25:25.410 and then, we did the same GAM model,
- $525\ 00{:}25{:}25.410 \dashrightarrow 00{:}25{:}29.520$ and then, we calculate the values of the test statistic
- $526\ 00:25:29.520 \longrightarrow 00:25:31.500$ on these permuted subsamples,
- $527\ 00:25:31.500 \longrightarrow 00:25:33.210$ that gave us a null distribution.
- 528~00:25:33.210 --> 00:25:36.540 So together, we can get a P value, this is what we did.
- 529 00:25:36.540 --> 00:25:38.490 And we can show that this approach
- 530 00:25:38.490 --> 00:25:41.370 indeed can control the P values,

- $531~00{:}25{:}41.370 \dashrightarrow 00{:}25{:}44.820$ make the P values uniformly distributed on the null,
- 532 00:25:44.820 --> 00:25:47.190 while the existing method that uses GAM,
- $533\ 00:25:47.190 \longrightarrow 00:25:50.130$ but only the theoretical distribution called tradeSeq,
- $534~00:25:50.130 \longrightarrow 00:25:53.160$ they have some distortion for P values.
- $535\ 00:25:53.160 --> 00:25:56.126$ And then, you may wonder, what's the consequence?
- 536 00:25:56.126 --> 00:25:58.161 We can show that, oh, and I should say,
- $537~00{:}25{:}58.161 \dashrightarrow 00{:}26{:}03.161$ Monocle3 uses generalized linear model and not uncertainty.
- $538\ 00:26:03.450 \longrightarrow 00:26:06.683$ So you can see that even though it's not as bad as tradeSeq,
- $539\ 00:26:06.683 \longrightarrow 00:26:08.670$ still, some distortion.
- $540\ 00:26:08.670 \longrightarrow 00:26:09.540$ So we wanna show
- 541 00:26:09.540 --> 00:26:13.170 that by calibrating the P value using our way
- $542\ 00:26:13.170 \longrightarrow 00:26:16.740$ we can actually discover more functional terms
- 543 00:26:16.740 --> 00:26:18.510 in our differentially expressed genes.
- $544\ 00:26:18.510 --> 00:26:21.780$ It means that we can find some new biological functions
- $545\ 00:26:21.780 \longrightarrow 00:26:23.730$ that were missed by this new method.
- $546~00{:}26{:}23.730 \dashrightarrow 00{:}26{:}28.080$ Which shows that FDR control not just help with FDR control
- 547 00:26:28.080 --> 00:26:29.160 of P value calibration,
- 548 00:26:29.160 --> 00:26:31.110 not just help with FDR control,
- $549\ 00:26:31.110 \longrightarrow 00:26:33.033$ but may also boost some power.
- $550~00{:}26{:}34.230$ --> $00{:}26{:}37.290~\mathrm{So}$ I just quickly talk about this Pseudotime DE,
- $551\ 00:26:37.290 --> 00:26:40.200$ but I want to say that its computational time
- $552\ 00:26:40.200 \longrightarrow 00:26:42.150$ is the biggest limitation.
- $553~00{:}26{:}42.150 \dashrightarrow 00{:}26{:}46.485$ Because here, our P value calculation requires many rounds
- $554~00{:}26{:}46.485 \dashrightarrow 00{:}26{:}50.430$ of subsampling, pseudotime inference, and permutation.

- $555\ 00:26:50.430 \longrightarrow 00:26:54.630$ So let's say we want the P value with resolution 0.001,
- $556\ 00:26:54.630 --> 00:26:58.230$ we need at least 1000 rounds of such things, right?
- $557\ 00:26:58.230 \longrightarrow 00:26:59.580$ That will take time.
- $558\ 00:26:59.580 \longrightarrow 00:27:00.900$ So the natural question
- 559 00:27:00.900 --> 00:27:04.410 is can we reduce the number of rounds, right,
- 560 00:27:04.410 --> 00:27:06.330 and still achieve FDR control?
- $561\ 00:27:06.330 \longrightarrow 00:27:08.127$ That becomes similar to my first goal.
- $562\ 00{:}27{:}08.127 \dashrightarrow 00{:}27{:}10.920$ Can we get rid of the higher resolution P values,
- $563\ 00:27:10.920 --> 00:27:14.460$ control the FDR, and then, we will use Clipper again.
- 564 00:27:14.460 --> 00:27:15.360 So you can see,
- $565\ 00:27:15.360 \longrightarrow 00:27:18.120$ Clipper is used throughout all the motivations,
- 566 00:27:18.120 --> 00:27:19.740 that's why we proposed it,
- $567\ 00:27:19.740 \longrightarrow 00:27:22.140$ and I'll talk about it in the next minute.
- $568\ 00:27:22.140 --> 00:27:24.330$ And the second question we didn't address
- $569\ 00:27:24.330 \longrightarrow 00:27:28.740$ is that what if the cells don't follow a trajectory at all?
- 570 00:27:28.740 --> 00:27:31.590 So clearly in our null hypothesis,
- $571\ 00:27:31.590 --> 00:27:34.050$ we are assuming there is a trajectory,
- $572~00{:}27{:}34.050 \dashrightarrow 00{:}27{:}38.100$ it's just that gene J doesn't change with the trajectory.
- 573 00:27:38.100 --> 00:27:40.320 But what if the trajectory doesn't exist?
- $574~00{:}27{:}40.320 \dashrightarrow 00{:}27{:}44.580$ So this whole idea of this trajectory pseudotime inference
- $575\ 00:27:44.580 \longrightarrow 00:27:45.810$ doesn't make sense, right?
- $576\ 00:27:45.810 \longrightarrow 00:27:47.190$ We need to consider that.
- $577\ 00:27:47.190 --> 00:27:50.460$ But I don't think we have a good way to do it,
- $578\ 00:27:50.460 \longrightarrow 00:27:53.640$ unless we can change the cells to have a null
- 579 00:27:53.640 --> 00:27:56.490 where the cells don't follow a trajectory.

- $580\ 00:27:56.490 --> 00:27:59.070$ So this motivated us to generate cells
- $581~00{:}27{:}59.070 \dashrightarrow 00{:}28{:}02.250$ that don't follow a trajectory, and we used a simulator.
- $582\ 00:28:02.250 \longrightarrow 00:28:05.730$ So which it will be the last part I will talk about today.
- 583 00:28:05.730 --> 00:28:08.970 Okay, PseudotimeDE is one such a problem
- $584~00{:}28{:}08.970 \dashrightarrow 00{:}28{:}11.820$ where pseudotime is inferred from the same data.
- $585~00:28:11.820 \dashrightarrow 00:28:16.820$ Another common problem is to do clustering on single cells
- 586 00:28:17.370 --> 00:28:19.110 to identify cell clusters,
- 587 00:28:19.110 --> 00:28:21.060 and between cell clusters,
- $588\ 00:28:21.060 \longrightarrow 00:28:23.400$ we identify differentially expressed genes.
- $589\ 00:28:23.400 \longrightarrow 00:28:25.710$ We call this problem ClusterDE.
- 590 00:28:25.710 --> 00:28:29.430 But this is also using the data twice, right?
- $591~00{:}28{:}29.430 \dashrightarrow 00{:}28{:}32.400$ So people have called this term double dipping,
- 592 00:28:32.400 --> 00:28:36.330 meaning that the same data used for twice.
- 593 00:28:36.330 --> 00:28:37.830 To tackle this problem,
- $594\ 00:28:37.830 \longrightarrow 00:28:41.301$ we need to consider the uncertainty in cell clustering,
- $595\ 00:28:41.301 --> 00:28:43.590$ and there are three existing papers
- $596\ 00:28:43.590 --> 00:28:45.690$ that try to address this problem
- $597\ 00:28:45.690 --> 00:28:48.480$ that they either need to assume a distribution,
- 59800:28:48.480 --> 00:28:52.680 like genes follow Gaussian distribution in every cluster
- $599~00{:}28{:}52.680 \dashrightarrow 00{:}28{:}56.550$ or every gene follows a Poisson distribution here
- $600\ 00:28:56.550 \longrightarrow 00:28:58.530$ and they need to do count splitting.
- 601 00:28:58.530 --> 00:29:01.530 So I won't talk into the couple of details here,
- $602\ 00:29:01.530 \longrightarrow 00:29:02.430$ but I just want to say
- $603\ 00{:}29{:}02.430 \dashrightarrow 00{:}29{:}05.340$ that the count splitting approach in my opinion
- $604\ 00:29:05.340 \longrightarrow 00:29:07.200$ tackles a different problem.

- 605 00:29:07.200 --> 00:29:10.710 It is conditional on the observed data matrix,
- $606\ 00:29:10.710 \longrightarrow 00:29:12.660$ rather than considered to be random.
- 607 00:29:12.660 --> 00:29:14.580 But I will not talk about the detail here.
- $608\ 00:29:14.580 \longrightarrow 00:29:16.600$ So motivated by the challenge in this problem,
- $609\ 00:29:16.600$ --> 00:29:21.600 and we want to propose something not distribution-specific.
- $610\ 00:29:22.200$ --> 00:29:27.200 We want to use our simulator to generate the null data
- $611\ 00:29:27.960 \longrightarrow 00:29:31.470$ and then use Clipper to achieve the FDR control.
- 612 00:29:31.470 --> 00:29:34.050 So we want to do this non parametrically.
- $613\ 00:29:34.050 \longrightarrow 00:29:36.480$ So I think the idea was motivated
- $614~00{:}29{:}36.480 \dashrightarrow 00{:}29{:}39.390$ by two phenomenal statistical papers.
- $615\ 00:29:39.390 \longrightarrow 00:29:41.520$ One is the gap statistic paper,
- $616\ 00{:}29{:}41.520 {\:{\mbox{--}}\!>\:} 00{:}29{:}45.260$ which was proposed to find the number of clusters
- $617\ 00:29:45.260 \longrightarrow 00:29:46.800$ in the clustering problem.
- $618~00{:}29{:}46.800 --> 00{:}29{:}49.440$ And if you read a paper, I think the smart idea there
- $619\ 00{:}29{:}49.440 \dashrightarrow 00{:}29{:}53.940$ is they try to generate data points without clusters
- $620\ 00:29:53.940 \longrightarrow 00:29:55.590$ as the negative control.
- 621 00:29:55.590 --> 00:29:59.160 Then, you can control your number of clusters
- 622 00:29:59.160 --> 00:30:00.870 with some statistic,
- 623 00:30:00.870 --> 00:30:03.027 versus what if there's no clusters, right,
- $624\ 00:30:03.027 \longrightarrow 00:30:04.920$ and do the comparison and find the gap.
- $625\ 00:30:04.920 \longrightarrow 00:30:06.180$ That's the gap statistic.
- $626\ 00:30:06.180 \longrightarrow 00:30:08.760$ And knockoffs gave the theoretical foundation
- 627 00:30:08.760 --> 00:30:12.393 for FDR control without using high resolution P values.
- $628\ 00:30:13.230 \longrightarrow 00:30:15.600$ Okay, so the halftime summary
- $629~00:30:15.600 \longrightarrow 00:30:17.970$ is that I talked about three common causes
- 630 00:30:17.970 --> 00:30:19.470 of ill-posed P values.

- 631 00:30:19.470 --> 00:30:20.970 Hopefully, I have convinced you
- $632\ 00:30:20.970 \longrightarrow 00:30:24.600$ that we need something to avoid this problem.
- 633 00:30:24.600 --> 00:30:26.220 So I talked about Clipper,
- $634\ 00{:}30{:}26.220 \dashrightarrow 00{:}30{:}29.730$ the p-value-free FDR control for genomic feature screening.
- $635\ 00:30:29.730 \longrightarrow 00:30:33.030$ And as I said, it was motivated and enabled
- $636\ 00:30:33.030 \longrightarrow 00:30:36.240$ by the FDR control procedure from this paper.
- $637\ 00:30:36.240 \longrightarrow 00:30:39.120$ But the difference here is that we focus
- $638\ 00:30:39.120 \longrightarrow 00:30:42.030$ on marginal screening of interesting features.
- $639\ 00:30:42.030$ --> 00:30:45.450 So in other words, we look at one feature at a time.
- 640 00:30:45.450 --> 00:30:47.190 In my previous examples,
- $641\ 00:30:47.190 \longrightarrow 00:30:50.760$ a feature could be a region or a gene.
- 642 00:30:50.760 --> 00:30:53.220 So in the original knockoff paper,
- $643\ 00:30:53.220 --> 00:30:57.240$ their goal is to generate knockoff data
- $644\ 00:30:57.240 \longrightarrow 00:31:01.170$ just like fake data for multiple features jointly.
- 645 00:31:01.170 --> 00:31:02.940 And that's the very challenging part.
- $646\ 00:31:02.940 \longrightarrow 00:31:05.190$ But in our case, we don't need that
- $647\ 00:31:05.190 --> 00:31:07.440$ because we are looking at one feature at a time,
- 648 00:31:07.440 --> 00:31:09.600 so it's not a multi-varied problem,
- $649\ 00:31:09.600 \longrightarrow 00:31:11.610$ but it's a marginal screening problem.
- $650\ 00:31:11.610 --> 00:31:15.420$ So our goal is to get rid of high resolution P values.
- $651\ 00:31:15.420 \longrightarrow 00:31:16.910$ So the advantage of this
- $652\ 00{:}31{:}16.910 \dashrightarrow 00{:}31{:}20.340$ is we don't need parametric distribution assumptions,
- $653\ 00:31:20.340 \longrightarrow 00:31:22.410$ or we don't need large sample sizes
- $654\ 00{:}31{:}22.410 \dashrightarrow 00{:}31{:}25.890$ to enable non parametric tests, these are not needed.
- $655\ 00:31:25.890 --> 00:31:29.070$ We just need to summarize every feature
- 656 00:31:29.070 --> 00:31:31.470 into a contrast score,
- $657\ 00:31:31.470 --> 00:31:34.500$ and then, set a cutoff on the contrast scores.

- $658\ 00:31:34.500 \longrightarrow 00:31:36.900$ So what do I mean by contrast score?
- 659 00:31:36.900 --> 00:31:39.630 So every feature, say, I have total d features,
- 660 00:31:39.630 --> 00:31:43.380 they have C, D, sorry, d contrast scores
- $661\ 00:31:43.380 \longrightarrow 00:31:45.240$ shown as C1 to Cd,
- $662\ 00:31:45.240 \longrightarrow 00:31:47.479$ so I'm calling the histogram
- $663\ 00:31:47.479 \longrightarrow 00:31:49.650$ of the distribution of contrast scores.
- 664 00:31:49.650 --> 00:31:53.790 So if the theoretical assumption is satisfied,
- $665\ 00:31:53.790 \longrightarrow 00:31:56.550$ then the features that are null features
- $666~00{:}31{:}56.550 \dashrightarrow 00{:}32{:}00.210$ should follow a symmetrical distribution
- $667\ 00:32:00.210 \longrightarrow 00:32:01.950$ around the zero, okay?
- $668\ 00:32:01.950 \longrightarrow 00:32:04.200$ And for the features that are interesting
- $669\ 00:32:04.200 \longrightarrow 00:32:05.610$ and should be discovered,
- $670\ 00:32:05.610 \longrightarrow 00:32:08.610$ should be large and positive on the right tail.
- 671 00:32:08.610 --> 00:32:12.090 So the theory of the FDR control just says,
- $672\ 00:32:12.090 \longrightarrow 00:32:15.930$ we can find the contrast score cutoff as t,
- 673 00:32:15.930 --> 00:32:20.070 such that this ratio is controlled under q.
- $674~00:32:20.070 \longrightarrow 00:32:22.620$ We ought to find the minimum t for this.
- 675~00:32:22.620 --> 00:32:25.902 What this means is can you can consider this ratio
- $676\ 00:32:25.902 \longrightarrow 00:32:29.017$ as a rough estimator of FDR.
- $677\ 00:32:29.999 \longrightarrow 00:32:33.177$ So the denominator is just the left tail,
- 678 00:32:33.177 --> 00:32:35.163 the red part plus one,
- $679\ 00:32:36.060 --> 00:32:38.910$ sorry, the numerator is the right tail plus one,
- $680\ 00{:}32{:}38.910 \dashrightarrow 00{:}32{:}43.166$ the denominator is the, sorry, the left tail is, sorry,
- 681 00:32:43.166 --> 00:32:45.420 the numerator is the left tail plus one,
- $682\ 00{:}32{:}45.420 \dashrightarrow 00{:}32{:}49.290$ the denominator is the right tail with maximum with one.
- $683\ 00:32:49.290 --> 00:32:52.470$ So in other words, still trying to avoid dividing zero.
- $684~00{:}32{:}52.470 \dashrightarrow 00{:}32{:}56.130$ And the idea is that we want to find a threshold t.

- $685\ 00:32:56.130 \longrightarrow 00:32:59.190$ so that the right tail will be called discoveries
- $686\ 00{:}32{:}59.190 \dashrightarrow 00{:}33{:}03.330$ and the left tail represent false discoveries.
- $687\ 00:33:03.330 \longrightarrow 00:33:04.680$ That's the intuition.
- 688 00:33:04.680 --> 00:33:07.770 Because we know, if the feature's null,
- $689\ 00:33:07.770 \longrightarrow 00:33:11.340$ then it will be randomly positive or negative.
- $690\ 00{:}33{:}11.340 \dashrightarrow 00{:}33{:}14.700$ And the sign is independent of the absolute value.
- $691\ 00:33:14.700 \longrightarrow 00:33:18.330$ So that just replaces
- $692\ 00:33:18.330 --> 00:33:21.840$ the uniform distribution requirement for P values,
- $693\ 00:33:21.840 \longrightarrow 00:33:23.580$ we change that to symmetry.
- $694\ 00:33:23.580 \longrightarrow 00:33:26.100$ And another thing is that the feature,
- $695~00{:}33{:}26.100 \dashrightarrow 00{:}33{:}29.490$ if it's large positive, we want to discover it, right?
- $696\ 00:33:29.490 \longrightarrow 00:33:31.440$ So this will be the discovery set
- $697~00{:}33{:}31.440 \dashrightarrow 00{:}33{:}36.060$ and this represents the negative, false discovery set.
- $698~00{:}33{:}36.060 \longrightarrow 00{:}33{:}40.620$ So that's the idea intuition behind this approach.
- $699\ 00:33:40.620 \longrightarrow 00:33:42.480$ But the theory to really prove it,
- $700\ 00:33:42.480 --> 00:33:45.630$ we need to use Martingale in probability to prove it.
- $701\ 00:33:45.630 \longrightarrow 00:33:46.980$ And some of the technique was used
- 702 00:33:46.980 --> 00:33:48.990 for the Benjamini Hochburg procedure
- 703 00:33:48.990 --> 00:33:50.460 still based on Martingale.
- $704~00{:}33{:}50.460 \dashrightarrow 00{:}33{:}54.360$ So anyway, this allows us to really control the FDR
- 705 00:33:54.360 --> 00:33:55.860 just using contrast scores.
- 706 00:33:55.860 --> 00:33:58.110 And another thing I found as appealing
- $707\ 00:33:58.110 \longrightarrow 00:34:01.020$ is that if you visually inspect the contract scores,
- 708~00:34:01.020 --> 90:34:05.100 you can see whether the assumption seems to be reasonable

709 00:34:05.100 --> 00:34:07.680 because you expect to see something symmetrical

 $710\ 00:34:07.680 \longrightarrow 00:34:09.810$ plus a heavy right tail.

 $711\ 00:34:09.810 \longrightarrow 00:34:13.800$ Okay, so we are currently writing to make this more formal,

 $712\ 00:34:13.800 \longrightarrow 00:34:15.000$ so we could actually check

 $713\ 00:34:15.000 \longrightarrow 00:34:18.150$ whether the assumption is reasonably holding.

 $714\ 00:34:18.150 \longrightarrow 00:34:19.470$ So with this approach,

 $715\ 00:34:19.470 \longrightarrow 00:34:24.470$ we can make a lot of the comparison analysis easier

 $716\ 00:34:25.560 --> 00:34:29.550$ because the key is to find a reasonable contrast score

 $717\ 00:34:29.550 \longrightarrow 00:34:31.830$ that satisfies this assumption.

718 00:34:31.830 --> 00:34:35.070 And I can say that there may be multiple contrast scores

 $719\ 00:34:35.070 \longrightarrow 00:34:37.410$ that satisfy, not just the unique one.

720 00:34:37.410 --> 00:34:39.550 Then the difference is power, right?

 $721\ 00:34:39.550 \longrightarrow 00:34:41.160$ So we may have a better power

722 00:34:41.160 --> 00:34:44.250 if you have a heavier right tail.

723 00:34:44.250 --> 00:34:47.040 Okay, so for a ChIP-seq peak calling analysis,

 $724\ 00:34:47.040 \longrightarrow 00:34:49.230$ we can say that the contrast score

 $725\ 00:34:49.230 \longrightarrow 00:34:51.870$ will be comparing the target data

 $726\ 00:34:51.870 \longrightarrow 00:34:54.630$ from experimental condition to the null data,

 $727\ 00:34:54.630 \longrightarrow 00:34:56.370$ which is the background condition.

728 00:34:56.370 --> 00:34:59.280 They serve a natural pair of contrast,

 $729\ 00:34:59.280 --> 00:35:03.390$ and we could apply any pipeline to each data,

730 00:35:03.390 --> 00:35:05.633 the same pipeline and then do the contrast, right?

731 00:35:05.633 --> 00:35:08.130 You can imagine, if there's no peak,

 $732\ 00:35:08.130 \longrightarrow 00:35:09.690$ then these two values will be,

 $733\ 00:35:09.690 \longrightarrow 00:35:12.690$ which one is bigger is equally likely.

734 00:35:12.690 --> 00:35:15.720 And for the RNA-seq analysis,

 $735\ 00{:}35{:}15.720 \dashrightarrow 00{:}35{:}20.280$ here, I showed we could use permuted data as the null data

 $736\ 00:35:20.280 \longrightarrow 00:35:21.840$ actual data as a target data.

 $737\ 00:35:21.840 \longrightarrow 00:35:25.020$ So if we run some test on actual data

738 00:35:25.020 --> 00:35:26.550 to get a test statistic,

 $739\ 00:35:26.550 \longrightarrow 00:35:28.560$ we use the same test on permuted data

 $740\ 00:35:28.560$ --> 00:35:32.190 to get a test statistic, and they serve as a contrast.

741 00:35:32.190 --> 00:35:34.890 And finally, for the PseudotimeDE and ClusterDE,

 $742\ 00:35:34.890 \longrightarrow 00:35:36.660$ the single cell problem,

743 00:35:36.660 --> 00:35:40.050 actual data will give us some comparison,

744 00:35:40.050 --> 00:35:41.670 either PseudotimeDE

745 00:35:41.670 --> 00:35:45.750 or the between ClusterDE test statistic.

 $746\ 00:35:45.750 \longrightarrow 00:35:48.150$ And if we have some similar data

 $747\ 00:35:48.150 \longrightarrow 00:35:49.440$ that represents the null,

748 00:35:49.440 --> 00:35:51.900 like null trajectory, null cluster,

749 00:35:51.900 --> 00:35:55.290 we could run the same pipeline and then do the contrast.

 $750\ 00:35:55.290 \longrightarrow 00:35:57.180$ So you see, this actually free us

 $751\ 00:35:57.180 --> 00:36:00.237$ from saying we need to derive P values

 $752\ 00:36:00.237 --> 00:36:02.160$ and we need to know the distribution

753 00:36:02.160 --> 00:36:05.340 by either theory or by numerical simulation, right?

 $754\ 00:36:05.340 \longrightarrow 00:36:06.450$ These are all relieved

 $755\ 00:36:06.450 --> 00:36:08.250$ because we just need to do a contrast.

 $756\ 00:36:08.250 --> 00:36:11.670$ And the power is gained from the many, many tests.

 $757\ 00:36:11.670 \longrightarrow 00:36:12.930$ we look at them together.

758 00:36:12.930 --> 00:36:13.763 So that's why

759 00:36:13.763 --> 00:36:15.137 this idea (background noise drowns out speaker).

- 760 00:36:16.080 --> 00:36:19.950 Okay, so as I said, we tried to implement Clipper
- 761 00:36:19.950 --> 00:36:22.380 as a way to improve FDR control,
- $762\ 00:36:22.380 \longrightarrow 00:36:23.880$ and we did achieve this
- $763\ 00:36:23.880 --> 00:36:27.240$ for the popular software Macs and Homer
- 764 00:36:27.240 --> 00:36:29.070 for ChIP-seq peak calling
- $765~00{:}36{:}29.070$ --> $00{:}36{:}32.820$ and DESeq2 to edgeR for RNA-seq DEG identification.
- 766 00:36:32.820 --> 00:36:36.660 So you see that they did have inflated FDR,
- $767\ 00:36:36.660 \longrightarrow 00:36:39.300$ so the Y axis is the actual FDR,
- 768 00:36:39.300 --> 00:36:41.940 X axis is the target FDR threshold.
- $769\ 00:36:41.940 \longrightarrow 00:36:43.733$ There are inflations,
- $770\ 00:36:43.733 \longrightarrow 00:36:46.410$ but with our Clipper as an add-on
- 771 00:36:46.410 --> 00:36:48.750 to be used downstream of what they output
- $772\ 00:36:48.750 \longrightarrow 00:36:50.430$ and do the contrast,
- $773\ 00:36:50.430 \longrightarrow 00:36:53.610$ we can largely reduce the FDR to the target
- 774 00:36:53.610 --> 00:36:56.340 and still maintain quite good power.
- $775\ 00:36:56.340 \longrightarrow 00:36:59.073$ So that's the usage of Clipper as and add-on.
- 776 00:36:59.910 --> 00:37:01.890 And for the single cell part,
- $777\ 00:37:01.890 --> 00:37:04.710\ I\ didn't\ finish\ about\ the\ null\ data\ generation.$
- 778 00:37:04.710 --> 00:37:06.000 How do we do it?
- 779 00:37:06.000 --> 00:37:09.510 Our simulator was proposed partly for this reason,
- $780\ 00:37:09.510 \longrightarrow 00:37:11.520$ but it has more uses.
- $781\ 00:37:11.520 --> 00:37:14.730\ So\ I$ just want to say that it's called scDesign3
- $782\ 00:37:14.730 --> 00:37:18.570$ because we have scDesign and scDesign2 as two previous work.
- $783\ 00:37:18.570 \longrightarrow 00:37:19.830\ Now, focus on scDesign2$
- $784\ 00:37:19.830 \longrightarrow 00:37:23.580$ because it is the direct predecessor of scDesign3.
- 785 00:37:23.580 --> 00:37:25.650 So what scDesign2 two does
- $786\ 00{:}37{:}25.650 {\: -->\:} 00{:}37{:}30.480$ is it tries to fit a multi-gene probabilistic model

 $787\ 00:37:30.480 \longrightarrow 00:37:32.370$ for each cell type,

 $788\ 00:37:32.370 --> 00:37:35.430$ and then, every gene assumes to follow

 $789\ 00:37:35.430 \longrightarrow 00:37:39.000$ a parametric distribution within the cell type.

 $790\ 00:37:39.000 \longrightarrow 00:37:40.950$ And the major contribution

 $791\ 00:37:40.950 --> 00:37:43.650$ is that we capture gene-gene correlations

792 00:37:43.650 --> 00:37:45.150 using Gaussian copula.

 $793\ 00:37:45.150 --> 00:37:47.430$ That will make the data more realistic.

 $794\ 00:37:47.430 \longrightarrow 00:37:48.990$ Here is the comparison.

 $795\ 00:37:48.990 \longrightarrow 00:37:51.780$ This is the real data used for fitting the model.

 $796\ 00:37:51.780 \longrightarrow 00:37:54.960$ This is the lab (indistinct) test data used for validation.

797 00:37:54.960 --> 00:37:58.860 and this is the synthetic cells using copula.

798 00:37:58.860 \rightarrow 00:38:01.710 If we remove the copula, the cells will look like this.

 $799\ 00:38:01.710 \longrightarrow 00:38:03.600$ So not realistic at all.

 $800\ 00{:}38{:}03.600 \longrightarrow 00{:}38{:}07.770$ And our data is more realistic than other simulators

 $801\ 00:38:07.770 --> 00:38:12.030$ that did not explicitly capture gene-gene correlation.

802 00:38:12.030 --> 00:38:14.340 Although, they have some implicit mechanism,

 $803\ 00:38:14.340 \longrightarrow 00:38:16.710$ but the model is different.

 $804~00{:}38{:}16.710 \dashrightarrow 00{:}38{:}21.300$ Okay, so we realize that scDesign2 is doing a good job

 $805\ 00:38:21.300 \longrightarrow 00:38:22.830$ for displaying cell types,

806 00:38:22.830 --> 00:38:24.750 but it cannot generate data like this

807 00:38:24.750 --> 00:38:26.940 from a continuous trajectory.

 $808~00:38:26.940 \dashrightarrow 00:38:30.960$ What we could do is to force the cells to be divided

 $809\ 00:38:30.960 \longrightarrow 00:38:32.400$ and then use scDesign2.

 $810\ 00:38:32.400 \longrightarrow 00:38:35.490$ But then, you can see the cells are kind of in clusters,

811 $00:38:35.490 \longrightarrow 00:38:36.930$ right, not in real data.

 $812\ 00:38:36.930 \longrightarrow 00:38:40.620$ But with our generalization to the version three,

 $813\ 00:38:40.620 \longrightarrow 00:38:45.090$ we now can generate cells from a continuous trajectory.

 $814\ 00:38:45.090 \longrightarrow 00:38:48.480$ And I can quickly say that we basically generalize this,

815 00:38:48.480 --> 00:38:51.180 this count distribution per cell type

 $816~00{:}38{:}51.180 \dashrightarrow 00{:}38{:}54.600$ to a generalized additive model, which I already said.

 $817\ 00:38:54.600 --> 00:38:57.270$ So we could make it more flexible in general,

 $818~00{:}38{:}57.270 \dashrightarrow 00{:}39{:}01.530$ and scDesign2 becomes a special case of scDesign3.

 $819\ 00:39:01.530 \longrightarrow 00:39:03.020$ And one more thing we could do

820 00:39:03.020 --> 00:39:06.270 is we actually use the technique vine copula,

 $821\ 00{:}39{:}06.270 \dashrightarrow 00{:}39{:}11.070$ so we could get the likelihood of how the model fits

 $822\ 00:39:11.070 \longrightarrow 00:39:15.180$ to the real data, so we can get the likelihood of the model,

823 00:39:15.180 \rightarrow 00:39:18.060 which can also give us more information.

824 00:39:18.060 --> 00:39:21.420 So besides the single cell trajectory data,

825 00:39:21.420 --> 00:39:24.720 we can also use the idea to generate spatial data.

 $826\ 00{:}39{:}24.720 \dashrightarrow 00{:}39{:}27.660$ So here, the modification is that for every gene

 $827\ 00:39:27.660 --> 00:39:31.680$ we assume a Gaussian process in the 2D space,

 $828\ 00:39:31.680 \longrightarrow 00:39:33.630$ so it can have a smooth function

 $829\ 00:39:33.630 \longrightarrow 00:39:35.580$ for (indistinct) expression (indistinct).

 $830\ 00{:}39{:}35.580 \dashrightarrow 00{:}39{:}40.020$ And also, my other student help with making the simulator

 $831\ 00:39:40.020$ --> 00:39:44.220 to generate reads, sequencing reads, not just counts.

 $832\ 00:39:44.220 \longrightarrow 00:39:46.080$ So we can go from counts to reads,

 $833\ 00:39:46.080 --> 00:39:48.450$ and this will give us more functionality

 $834\ 00:39:48.450 \longrightarrow 00:39:51.240$ to benchmark some low level tools.

 $835\ 00:39:51.240 \longrightarrow 00:39:52.380$ So in short,

 $836\ 00:39:52.380 --> 00:39:55.920$ the scDesign3 simulator has two functionalities.

 $837\ 00:39:55.920 \longrightarrow 00:39:58.590$ One is to do, of course, simulation.

838 00:39:58.590 --> 00:40:02.070 We can generate single cell data from cell types,

839 00:40:02.070 --> 00:40:04.740 discrete, continuous trajectories,

 $840\ 00:40:04.740 \longrightarrow 00:40:06.990$ or even in the spatial domain.

 $841\ 00:40:06.990 --> 00:40:09.172$ We could generate feature modalities

 $842\ 00:40:09.172 --> 00:40:11.617$ we call multi-omics, including RNA-seq,

843 00:40:11.617 --> 00:40:13.920 ATAC-seq, which is a technology

844 00:40:13.920 --> 00:40:16.020 for open chromatin measurement,

845~00:40:16.020 --> 00:40:19.350 CITE-seq, which includes both protein and RNA,

 $846\ 00:40:19.350 \longrightarrow 00:40:21.030$ and also DNA methylation.

 $847\ 00:40:21.030 \longrightarrow 00:40:24.120$ These are the examples we tried, but we could do even more.

 $848\ 00{:}40{:}24.120 \dashrightarrow 00{:}40{:}27.960$ We could allow it to generate data with experimental designs

 $849\ 00:40:27.960 --> 00:40:32.960$ including sample covariate, conditions, or even batches.

 $850\ 00:40:33.120 \longrightarrow 00:40:36.150$ So these can make us generate cases

851 00:40:36.150 --> 00:40:38.760 for more types of benchmarking.

852 00:40:38.760 --> 00:40:41.160 And for interpreting real data,

 $853\ 00:40:41.160 --> 00:40:44.730\ scDesign3\ can give us model parameters,$

 $854\ 00{:}40{:}44.730 \dashrightarrow 00{:}40{:}47.400$ so we can know whether a gene has different means

 $855\ 00:40:47.400 \longrightarrow 00:40:48.990$ in two cell types,

 $856~00:40:48.990 \longrightarrow 00:40:51.900$ whether a gene has a certain change on a pseudotime,

 $857\ 00:40:51.900 \longrightarrow 00:40:54.930$ or a gene has a certain change in two dimensional space.

858 00:40:54.930 --> 00:40:56.100 And also, as I said,

- 859~00:40:56.100 --> 00:40:58.980 we can output a likelihood that can give us a way
- $860\ 00:40:58.980 \longrightarrow 00:41:02.580$ to calculate the basic information criterion BIC,
- $861\ 00:41:02.580 \longrightarrow 00:41:03.960$ so we could evaluate
- $862\ 00:41:03.960 \longrightarrow 00:41:07.230$ whether some pseudotime describes data well,
- $863\ 00{:}41{:}07.230 \dashrightarrow 00{:}41{:}09.050$ whether the algorithm for pseudotime inference
- 864 00:41:09.050 --> 00:41:10.800 does a good job,
- $865\ 00:41:10.800 \longrightarrow 00:41:13.260$ or whether the clusters explain data well.
- $866\ 00:41:13.260 \longrightarrow 00:41:14.850$ So these are the things we could do.
- 867 00:41:14.850 --> 00:41:17.910 And finally, to generate the null data
- 868 00:41:17.910 --> 00:41:19.770 for the Clipper (indistinct),
- $869\ 00:41:19.770 \longrightarrow 00:41:22.200$ we can alter the model parameters.
- $870\ 00:41:22.200 \longrightarrow 00:41:25.080$ Like this is what we fit from real data,
- $871\ 00:41:25.080 \longrightarrow 00:41:27.300$ we could change the model parameters
- $872\ 00{:}41{:}27.300 \dashrightarrow 00{:}41{:}30.360$ to make the gene no longer differentially expressed,
- $873\ 00:41:30.360 \longrightarrow 00:41:32.580$ have the same mean in two subtypes.
- $874\ 00:41:32.580 \longrightarrow 00:41:34.650$ Or, after we fit a real data
- 875 00:41:34.650 --> 00:41:36.900 with two cell types or two clusters,
- $876~00{:}41{:}36.900 \dashrightarrow 00{:}41{:}39.870$ we could change the cluster parameter
- $877\ 00:41:39.870 \longrightarrow 00:41:42.450$ to make sure the cells come from one cluster
- $878\ 00:41:42.450 \longrightarrow 00:41:43.920$ instead of two clusters.
- $879\ 00{:}41{:}43.920 \longrightarrow 00{:}41{:}46.680$ So these are the things we could do with the model.
- 880 00:41:46.680 --> 00:41:49.140 And so this is how our paper,
- $881\ 00:41:49.140 --> 00:41:52.230$ but more details are in our paper, which has been posted,
- $882\ 00:41:52.230 \longrightarrow 00:41:53.910$ if you are interested.
- 883 00:41:53.910 --> 00:41:55.710 And I want to just quickly show
- $884\ 00:41:55.710 --> 00:41:58.830$ how the ClusterDE analysis could be done.
- $885\ 00:41:58.830 \longrightarrow 00:42:01.350$ This is the real data with two clusters.

- 886 00:42:01.350 --> 00:42:03.030 I want to say that this is the case
- 887 00:42:03.030 --> 00:42:04.830 where permutation wouldn't work.
- 888 00:42:04.830 --> 00:42:07.080 If you just permute the cluster labels,
- 889 00:42:07.080 --> 00:42:09.750 the cells will look like the same cells, right?
- $890\ 00:42:09.750 \longrightarrow 00:42:11.340$ They're still two clusters.
- 891 00:42:11.340 --> 00:42:12.690 But if you use our simulator,
- $892\ 00:42:12.690 --> 00:42:15.000$ we could generate cells from one cluster
- $893\ 00{:}42{:}15.000$ --> $00{:}42{:}18.720$ that reflects the complete null, there's no cluster.
- $894\ 00:42:18.720 --> 00:42:22.590$ And the use of this can be shown in this example.
- 895 00:42:22.590 --> 00:42:24.450 There's only one cluster,
- 896 00:42:24.450 --> 00:42:27.270 but if we use clustering algorithms,
- 897 00:42:27.270 --> 00:42:30.810 like these two choices, Seurat is a popular pipeline,
- 898 00:42:30.810 --> 00:42:33.990 Kmeans is the standard classical algorithm,
- 899 00:42:33.990 --> 00:42:38.100 using either to force the cells into two clusters,
- $900\ 00:42:38.100 \longrightarrow 00:42:39.990$ we are using gene expression data.
- 901 00:42:39.990 --> 00:42:43.560 So no wonder that if you look at a gene's expression
- 902 00:42:43.560 --> 00:42:46.470 between the two clusters, you may call it DE,
- $903\ 00{:}42{:}46.470 \dashrightarrow 00{:}42{:}50.010$ but that's not interesting, since there's no clusters.
- 904 00:42:50.010 --> 00:42:52.460 So if we use our scDesign3 to generate null data,
- $905\ 00{:}42{:}54.390 \dashrightarrow 00{:}42{:}58.313$ in this case, null data should be very similar to real data.
- 906 00:42:58.313 --> 00:43:00.300 It still has only one cluster.
- 907 00:43:00.300 --> 00:43:03.960 Then, if we run Seurat or Kmeans,
- 908 00:43:03.960 --> 00:43:05.970 similarly, on null data,
- $909\ 00:43:05.970 \longrightarrow 00:43:08.880$ we would divide the cell in a similar way,
- 910 00:43:08.880 --> 00:43:12.150 and then, if you do a contrast of the two sets of results.

- 911 00:43:12.150 --> 00:43:13.800 you should see no big difference.
- 912 00:43:13.800 --> 00:43:16.200 That's the idea for controlling FDR.
- 913 00:43:16.200 --> 00:43:20.730 So indeed, in that example, if we're just naively wrong,
- $914\ 00:43:20.730 \dashrightarrow 00:43:25.110$ the Seurat pipeline clustering followed by some tests
- 915 00:43:25.110 --> 00:43:27.750 like t, Wilcoxon, bimodal,
- $916\ 00:43:27.750 \longrightarrow 00:43:30.480$ yeah, you will see FDR is one.
- 917 00:43:30.480 --> 00:43:32.730 The reason is you keep finding D genes,
- $918\ 00:43:32.730 \longrightarrow 00:43:34.200$ even though there's no cluster.
- 919 00:43:34.200 --> 00:43:35.430 But using our approach,
- $920\ 00:43:35.430 \longrightarrow 00:43:38.280$ we could control the FDR reasonably well.
- 921 00:43:38.280 --> 00:43:41.520 So that's the predominant results for this purpose
- 922 00:43:41.520 --> 00:43:45.870 for this task, so that summarizes my talk today.
- 923 00:43:45.870 --> 00:43:48.150 And finally, I just want to make a few notes
- $924\ 00:43:48.150 \longrightarrow 00:43:50.370$ to give some messages.
- 925 00:43:50.370 --> 00:43:52.350 I talk about multiple testing,
- 926 00:43:52.350 --> 00:43:53.910 but in many scientific problems,
- 927 00:43:53.910 --> 00:43:57.240 I think the key is whether it should be formulated
- 928 00:43:57.240 --> 00:43:58.860 as a multiple testing problem.
- 929 00:43:58.860 --> 00:44:00.930 So actually, to address this question,
- 930 00:44:00.930 --> 00:44:02.910 I wrote a prospective article
- 931 00:44:02.910 --> 00:44:06.330 with my collaborator Xin Tong at USC.
- $932\ 00:44:06.330 \longrightarrow 00:44:10.470$ We try to clarify statistical hypothesis testing
- 933 $00:44:10.470 \longrightarrow 00:44:12.810$ from machine learning binary classification.
- $934\ 00:44:12.810 \longrightarrow 00:44:13.950$ They seem similar
- $935\ 00{:}44{:}13.950 \dashrightarrow 00{:}44{:}17.010$ because both would give you a binary decision, right?
- 936 00:44:17.010 \rightarrow 00:44:20.490 But I can say that testing is an inference problem,

- $937\ 00:44:20.490 --> 00:44:22.830$ classification is a prediction problem.
- 938 00:44:22.830 --> 00:44:24.690 So if you really think about it,
- 939 $00:44:24.690 \longrightarrow 00:44:27.000$ their fundamental concepts are different.
- $940\ 00:44:27.000 \longrightarrow 00:44:30.900$ So that's why we wrote this to really talk with biologists,
- 941 $00:44:30.900 \longrightarrow 00:44:34.530$ for computational people who use this simultaneously.
- 942 00:44:34.530 --> 00:44:37.230 So if you're interested, you can check it out.
- 943 00:44:37.230 --> 00:44:39.780 And finally, I wanna say that,
- $944\ 00:44:39.780 \longrightarrow 00:44:42.691$ so if it's a multiple testing problem,
- 945 00:44:42.691 --> 00:44:47.580 I talked about three common causes of illposed P values,
- 946 00:44:47.580 --> 00:44:50.340 and I propose a solution, Clipper,
- $947\ 00:44:50.340 \longrightarrow 00:44:54.630$ for simplifying this problem by just using contrast scores,
- $948\ 00:44:54.630 \longrightarrow 00:44:56.160$ and then, set a cutoff.
- 949 00:44:56.160 --> 00:44:58.680 And the simulator, which we hope to be useful
- $950~00{:}44{:}58.680 \dashrightarrow 00{:}45{:}01.080$ for the single cell and spatial omics field
- 951 00:45:01.080 --> 00:45:03.030 because this field is so popular,
- $952\ 00:45:03.030 \longrightarrow 00:45:04.890$ we have more than 1000 methods already.
- $953\ 00{:}45{:}04.890 \dashrightarrow 00{:}45{:}08.250$ So benchmarking seems to be something quite necessary.
- 954 00:45:08.250 --> 00:45:10.650 Because if there's no benchmarking,
- $955\ 00:45:10.650 --> 00:45:13.710$ then maybe new methods wouldn't have much of a chance
- $956\ 00:45:13.710 --> 00:45:16.170$ because people may still use the older method
- 957 00:45:16.170 --> 00:45:18.030 that are better cited.
- 958 00:45:18.030 \rightarrow 00:45:22.950 Okay, so these are the papers related to my talk today.
- 959 00:45:22.950 --> 00:45:25.920 And so, finally, I want to say that,
- $960\ 00:45:25.920 \longrightarrow 00:45:28.950$ so if you're interested, you want to check them out,
- 961 00:45:28.950 --> 00:45:30.990 and let me know if you have any questions.

- 962 00:45:30.990 --> 00:45:32.550 So finally, I'll just say this,
- $963\ 00:45:32.550 \longrightarrow 00:45:34.050$ this is something quite interesting.
- 964 00:45:34.050 --> 00:45:36.930 It's another paper we just recently wrote,
- 965 00:45:36.930 --> 00:45:37.763 and I can say,
- 966 00:45:37.763 --> 00:45:40.290 you should be online in genome biology very soon.
- $967\ 00:45:40.290 \longrightarrow 00:45:43.110$ So we actually did this benchmark
- 968 00:45:43.110 --> 00:45:47.220 for the so called QTL analysis in genetics, right?
- 969 00:45:47.220 --> 00:45:49.770 Quantitative Trait Locus mapping.
- 970 00:45:49.770 --> 00:45:51.330 So in this analysis,
- 971 00:45:51.330 --> 00:45:55.320 a common procedure is to infer hidden variables
- 972 00:45:55.320 --> 00:45:57.360 from the data, like genes expression matrix,
- 973 00:45:57.360 --> 00:46:00.060 want to do hidden variable improvements.
- 974 00:46:00.060 \rightarrow 00:46:03.390 Besides the most part, (indistinct) has the classical PCA,
- $975\ 00:46:03.390 \longrightarrow 00:46:06.690$ several methods propose specific (indistinct).
- $976~00:46:06.690 \longrightarrow 00:46:09.990$ And my student Heather, actually gave her the full credit,
- 977 00:46:09.990 --> 00:46:12.930 she was so careful and she really wanted to understand
- 978 00:46:12.930 --> 00:46:14.400 the method before using it,
- 979 00:46:14.400 --> 00:46:16.560 then that lead to this project.
- 980 00:46:16.560 --> 00:46:19.350 She wants to see, huh, do I really see advantages
- 981 00:46:19.350 --> 00:46:22.290 of this new method even compared to PCA?
- 982 00:46:22.290 --> 00:46:23.880 But that's what she found, right?
- 983 00:46:23.880 --> 00:46:26.370 PCA still seems to be the most stable,
- 984 00:46:26.370 --> 00:46:29.610 robust, and also faster algorithm,
- $985\ 00:46:29.610 \longrightarrow 00:46:32.400$ but this is one of the reviewer's comments
- 986 00:46:32.400 --> 00:46:34.050 I wanna share with you.
- 987 00:46:34.050 --> 00:46:36.520 These results may come as a surprise to some,

988 00:46:36.520 --> 00:46:39.450 given the nearly un-contestable status

989 00:46:39.450 --> 00:46:42.060 that method A has achieved within the community.

990 00:46:42.060 --> 00:46:43.800 But sadly, they reflect the fact

991 00:46:43.800 --> 00:46:46.740 that computational biology methods can rise to fame

 $992\ 00:46:46.740 \longrightarrow 00:46:50.280$ almost by accident rather than by sound statistic arguments.

993 00:46:50.280 --> 00:46:51.570 So if you're interest,

994 00:46:51.570 --> 00:46:53.910 you can check out this paper, it's on bio archive.

995 00:46:53.910 --> 00:46:56.580 But anyway, I think it says how important it is

996 $00:46:56.580 \longrightarrow 00:46:59.880$ for statisticians to convey our message, right?

997 00:46:59.880 --> 00:47:02.793 Why do we need statistical rigor, why does it matter?

998 00:47:03.720 --> 00:47:05.310 So for our students,

999 00:47:05.310 --> 00:47:07.776 if you want to know more about GAM and copulas,

 $1000\ 00:47:07.776 \longrightarrow 00:47:09.630$ there are two books I want to recommend.

 $1001\ 00:47:09.630 \longrightarrow 00:47:12.270$ So they're very good introductory textbooks,

 $1002\ 00:47:12.270 \longrightarrow 00:47:14.490$ so you can know the (indistinct).

 $1003\ 00{:}47{:}14.490 \dashrightarrow 00{:}47{:}18.900$ Finally, I want to thank my collaborator at UC Irvine,

 $1004~00{:}47{:}18.900 \dashrightarrow 00{:}47{:}22.740$ my students for all their tremendous work I talk about today

 $1005\ 00{:}47{:}22.740 \dashrightarrow 00{:}47{:}25.710$ and also the funding agencies for giving us the support.

 $1006\ 00:47:25.710 \longrightarrow 00:47:26.970$ So thank you very much.

1007 00:47:36.226 --> 00:47:38.247 <v Attendee>A question?</v>

 $1008\ 00:47:38.247 \longrightarrow 00:47:39.080 < v \ Jingyi>Yes.</v>$

 $1009\ 00:47:39.080 --> 00:47:40.110 < v \ Attendee>So I was really curious </v>$

 $1010\ 00{:}47{:}40.110 \dashrightarrow 00{:}47{:}44.850$ about the analysis of like the large patient sample.

- $1011\ 00:47:44.850 \longrightarrow 00:47:46.410\ I$ know that there has in fact
- 1012 00:47:46.410 --> 00:47:47.980 been extensive discussion on it.
- $1013\ 00:47:47.980 --> 00:47:49.870 < v -> Yeah, yeah. < / v >< v -> Which is < / v >$
- $1014\ 00:47:52.080 \longrightarrow 00:47:54.690$ interesting, to say the least, how it's gone down.
- 1015 00:47:54.690 --> 00:47:56.220 But I was kinda curious,
- $1016\ 00:47:56.220 \longrightarrow 00:48:00.570$ the way that it was presented here made me think about like,
- $1017\ 00{:}48{:}00.570 \dashrightarrow 00{:}48{:}04.803$ apologies, if this is like a path that's already been tread,
- $1018\ 00:48:06.810 \longrightarrow 00:48:10.890$ so, yeah, the bar graph.
- $1019\ 00:48:10.890 \longrightarrow 00:48:12.270 < V \ Jingyi> Yeah. < /v>$
- $1020\ 00:48:12.270 \longrightarrow 00:48:15.750 < v \text{ Attendee} > \text{Yeah, so it sort of}, </v>$
- $1021\ 00:48:15.750 --> 00:48:18.870$ it makes me wonder about the application
- $1022\ 00{:}48{:}18.870 \dashrightarrow 00{:}48{:}22.530$ of the term false discovery in different contexts.
- 1023 00:48:22.530 --> 00:48:25.980 And taking patients, you can imagine,
- 102400:48:25.980 --> 00:48:29.490 there can be like unintended structure
- $1025\ 00:48:29.490 \longrightarrow 00:48:32.370$ within those populations.
- $1026\ 00:48:32.370 --> 00:48:34.230$ And by (interference drowns out speaker) chance,
- $1027\ 00:48:34.230 \longrightarrow 00:48:38.321$ if there is 30,000 potential transcripts
- $1028\ 00:48:38.321$ --> 00:48:40.710 that you're looking at, there might actually be,
- 1029 00:48:40.710 --> 00:48:43.890 between individuals who are not isogenic,
- $1030\ 00:48:43.890 --> 00:48:46.920$ truly differentially expressed genes
- $1031\ 00:48:46.920 --> 00:48:50.220$ between even permuted groups.
- $1032\ 00{:}48{:}50{.}220$ --> $00{:}48{:}52{.}950$ And so I'm wondering if there's a useful distinction
- 1033 00:48:52.950 --> 00:48:56.280 between a false discovery and a true,
- $1034\ 00:48:56.280 \longrightarrow 00:48:58.173$ but uninteresting discovery.
- 1035 00:49:00.240 --> 00:49:03.630 <
v Jingyi>I think it depends on how you define truth.
</v>

- $1036\ 00:49:03.630 \longrightarrow 00:49:04.680$ I think that's the key.
- $1037\ 00:49:04.680 \longrightarrow 00:49:07.860$ But what is the definition of D genes?
- $1038\ 00:49:07.860 \longrightarrow 00:49:09.870\ I$ wanna say, to be exact,
- 1039 00:49:09.870 --> 00:49:13.997 the definition of D genes in DESeq2,
- $1040\ 00:49:13.997 \longrightarrow 00:49:17.550\ edgeR$, and that of Wilcoxon is different.
- 1041 00:49:17.550 --> 00:49:21.330 Because in Wilcoxon, the D gene is defined,
- 1042 00:49:21.330 --> 00:49:24.270 okay, if a gene, it has two distributions,
- $1043\ 00:49:24.270 \longrightarrow 00:49:26.520$ one under each condition,
- $1044\ 00:49:26.520 \longrightarrow 00:49:29.280$ and if I randomly take one observation
- $1045\ 00:49:29.280 \longrightarrow 00:49:31.890$ from each distribution from each condition,
- $1046\ 00{:}49{:}31.890 {\:{\mbox{--}}}{\:{\mbox{-}}} 00{:}49{:}34.650$ is the chance that one is bigger than the other
- 1047 00:49:34.650 --> 00:49:35.850 equal to 0.5?
- $1048\ 00:49:35.850 \longrightarrow 00:49:38.070$ That's the Wilcoxon question.
- $1049~00{:}49{:}38.070 --> 00{:}49{:}41.520$ While DESeq2 and edgeR, their D gene definition
- $1050\ 00:49:41.520 --> 00:49:45.090$ is the negative binomial means are different.
- 1051 00:49:45.090 --> 00:49:48.060 But clearly, you can see, it only depends
- $1052\ 00:49:48.060 \longrightarrow 00:49:51.360$ on that negative binomial is a reasonable distribution,
- $1053\ 00:49:51.360 \longrightarrow 00:49:52.193$ that's the key.
- 1054 00:49:52.193 --> 00:49:53.026 So that's why in theory,
- $1055\ 00:49:53.026$ --> 00:49:57.510 if negative binomial is no longer valid or reasonable,
- $1056\ 00:49:57.510 \longrightarrow 00:50:00.090$ then why should we define a D gene
- $1057\ 00{:}50{:}00.090$ --> $00{:}50{:}02.550$ based on negative binomial mean in difference?
- $1058~00{:}50{:}02.550$ --> $00{:}50{:}05.550$ I think that's kind of my answer to your question.
- $1059\ 00{:}50{:}05.550 \dashrightarrow 00{:}50{:}09.150$ But the tricky thing about statistical inference
- 1060 00:50:09.150 --> 00:50:10.770 compared to supervised learning

- $1061\ 00:50:10.770 \longrightarrow 00:50:14.370$ is that we don't observe the truth, that's always the case.
- 1062 00:50:14.370 --> 00:50:16.110 So we're making a guess.
- 1063 00:50:16.110 --> 00:50:19.650 Frequentist people have one way to guess,
- $1064\ 00:50:19.650 --> 00:50:21.390$ Poisson people have another way of guess.
- $1065\ 00{:}50{:}21.390 {\: -->\:} 00{:}50{:}23.700$ And so one issue I've seen in the Twitter discussion
- $1066\ 00:50:23.700 \longrightarrow 00:50:26.490$ is that several people try to,
- 1067 00:50:26.490 --> 00:50:28.200 maybe not intentionally,
- $1068\ 00:50:28.200 \dashrightarrow 00:50:31.500$ confuse frequent ist concept with Poisson concept,
- 1069 00:50:31.500 --> 00:50:33.660 but they're not really comparable, right?
- $1070\ 00{:}50{:}33.660 {\:{\mbox{--}}}{>} 00{:}50{:}35.880$ You cannot talk about them in the same ground.
- 1071 00:50:35.880 --> 00:50:39.270 That's a problem, and here, our criterion,
- 1072 00:50:39.270 --> 00:50:42.360 false discovery rate is a frequentist criteria,
- 1073 00:50:42.360 --> 00:50:43.590 it relies on P values, right?
- $1074\ 00:50:43.590 \dashrightarrow 00:50:46.410$ So therefore, you cannot use Poisson arguments
- 1075 00:50:46.410 --> 00:50:49.350 to argue against such frequentist way.
- 1076 00:50:49.350 --> 00:50:51.630 Because you are doing frequentist, right?
- $1077\ 00:50:51.630 \longrightarrow 00:50:53.700$ But whether frequentist makes sense or not,
- $1078\ 00:50:53.700 \longrightarrow 00:50:55.290$ that's a different topic.
- $1079\ 00:50:55.290 --> 00:50:56.730$ Hopefully, that answers your question.
- 1080 00:50:56.730 --> 00:50:58.520 <v ->Yeah, thank you.</v> <v ->Thank you.</v>
- 1081 00:51:00.870 --> 00:51:01.703 Yes. <v ->Hello,</v>
- 1082 00:51:01.703 --> 00:51:03.443 thank you much for your talk,
- 1083 00:51:03.443 --> 00:51:05.220 and I think that is very interesting.
- $1084\ 00:51:05.220 --> 00:51:09.410$ However, I have a question on slide 26 actually..
- 1085 00:51:12.507 --> 00:51:15.007 It's about what you said that,
- $1086\ 00:51:17.180 \longrightarrow 00:51:18.013$ maybe 26.

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1087\ 00:51:19.707 --> 00:51:22.695 < v \ Jingyi>26, okay, yeah. </v>
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 $1088\ 00:51:22.695 --> 00:51:23.837 < v \ Attendee> Yeah, you said < /v>$

 $1089\ 00:51:23.837 \longrightarrow 00:51:28.170$ that like it is a multi-gene probabilistic model

 $1090\ 00:51:28.170 \longrightarrow 00:51:29.640$ for cell type.

 $1091\ 00:51:29.640 \longrightarrow 00:51:33.300$ However, I'm a little bit confused

 $1092\ 00:51:33.300 \longrightarrow 00:51:35.593$ about how you define the cell type.

1093 00:51:36.840 --> 00:51:40.410 But basically, from my own understandings,

 $1094\ 00:51:40.410 \longrightarrow 00:51:44.103$ that after you get, for example, the single cell rise data,

 $1095\ 00:51:45.210$ --> 00:51:47.670 for example, you will use the route to get the cluster.

1096 00:51:47.670 --> 00:51:48.503 <v Jingyi>Yeah.</v>

1097 00:51:48.503 --> 00:51:51.996 <
v Attendee>And you will annotate this cluster</br/>/v>

 $1098\ 00:51:51.996 \longrightarrow 00:51:53.079$ based on the-

1099 00:51:54.276 --> 00:51:56.305 <v ->Knowledge, yeah.</v> <v ->Gene.</v>

 $1100\ 00:51:56.305 \longrightarrow 00:52:00.697$ And then, if this model based on your

1101 00:52:04.090 --> 00:52:05.757 annotation of, okay.

1102 00:52:08.190 --> 00:52:09.840 <v Jingyi>Yeah, I see you point.</v>

 $1103\ 00:52:09.840 --> 00:52:12.660$ Essentially, yeah, we need cell cluster to be pre-defined.

1104 00:52:12.660 --> 00:52:15.200 So if it's not reasonable, then, yes,

 $1105\ 00:52:15.200 \longrightarrow 00:52:17.250$ it will affect the results for sure.

 $1106\ 00{:}52{:}17.250 \dashrightarrow 00{:}52{:}19.710$ Because the key is that you need to make sure

 $1107\ 00:52:19.710 \longrightarrow 00:52:22.590$ it is reasonable to assume a gene follows

 $1108\ 00:52:22.590 \dashrightarrow 00:52:26.070$ one of the four distribution within a cluster, right?

 $1109\ 00:52:26.070 \longrightarrow 00:52:27.900$ So that's why there are methods

1110 00:52:27.900 --> 00:52:30.240 that try to refine clustering

 $1111\ 00:52:30.240 \dashrightarrow 00:52:33.660$ by checking the negative binomial distribution.

- $1112\ 00:52:33.660 \longrightarrow 00:52:35.970$ So there are several research on that,
- $1113\ 00:52:35.970 \longrightarrow 00:52:37.197$ and they're trying to refine that.
- $1114\ 00:52:37.197 --> 00:52:40.050$ But basically, we are sitting on those methods
- $1115\ 00:52:40.050 \longrightarrow 00:52:42.600$ to do the simulation, that's what we do.
- $1116\ 00:52:42.600 \longrightarrow 00:52:46.287$ But again, so that's why this is the problem with scDesign2,
- $1117\ 00:52:46.287 \dashrightarrow 00:52:49.950$ but scDesign3 sort of tries to address this problem
- $1118\ 00:52:49.950 \longrightarrow 00:52:51.960$ by providing the BIC.
- 1119 00:52:51.960 --> 00:52:54.570 So if the input clusters are bad,
- $1120\ 00:52:54.570 \longrightarrow 00:52:56.700$ then you can see that in the BIC.
- $1121\ 00:52:56.700 --> 00:52:59.220$ Because the likelihood will not be there, yeah.
- $1122\ 00:52:59.220 --> 00:53:00.623 < v \ Attendee > A \ similar \ question. < /v >$
- $1123\ 00:53:02.130 \longrightarrow 00:53:03.789$ I have another question
- $1124\ 00:53:03.789 \longrightarrow 00:53:07.547$ is that basically I assumed (indistinct) about
- 1125 00:53:08.632 --> 00:53:12.900 the experiments have duplicates,
- 1126 00:53:12.900 --> 00:53:16.300 however, in some situations,
- $1127\ 00:53:16.300 \longrightarrow 00:53:20.130$ maybe we do not have the replication.
- $1128\ 00:53:20.130 \longrightarrow 00:53:24.103$ But in this situation, how could we control the FDR,
- $1129\ 00:53:25.735 \longrightarrow 00:53:27.724$ if we do not have replicates,
- $1130\ 00:53:27.724 \longrightarrow 00:53:29.970$ then we cannot get the P value.
- 1131 00:53:29.970 --> 00:53:31.497 <
v Jingyi>That's exactly the point of this talk.
</v>
- $1132\ 00:53:31.497 --> 00:53:34.770$ The only part that has replicates is the RNA-seq part.
- $1133\ 00:53:34.770 --> 00:53:37.200$ The second part, that's the only part we have replicates.
- 1134 00:53:37.200 --> 00:53:39.037 In the first part, when we do the ChIP-seq,
- 1135 00:53:39.037 --> 00:53:41.880 it's just one replicate per condition, right?
- 1136 00:53:41.880 --> 00:53:44.700 That's why I said P value calculation would be helpful.

- $1137\ 00{:}53{:}44.700 \dashrightarrow 00{:}53{:}47.490$ Right, so the reason we could control the FDR
- $1138\ 00:53:47.490 --> 00:53:49.350$ without using P values
- $1139\ 00:53:49.350 \longrightarrow 00:53:51.810$ is just because we have many, many tests.
- $1140\ 00{:}53{:}51.810 --> 00{:}53{:}56.370$ So that's why we're doing this large scale testing.
- 1141 00:53:56.370 --> 00:53:58.816 I think the idea, if you check it out,
- $1142\ 00:53:58.816 \longrightarrow 00:54:03.780$ Bran Efron has talked about it extensively in his book,
- $1143\ 00:54:03.780 \longrightarrow 00:54:06.600$ it's called, so his idea of Empirical Bayes
- $1144\ 00:54:06.600 \longrightarrow 00:54:07.980$ is very similar to this.
- $1145\ 00:54:07.980 --> 00:54:10.710$ We try to borrow information across tests
- $1146\ 00:54:10.710 \longrightarrow 00:54:12.505$ to set a threshold.
- $1147\ 00:54:12.505 \longrightarrow 00:54:14.758$ Yeah, hopefully that answers your question.
- 1148 00:54:14.758 --> 00:54:15.591 Yeah?
- 1149 00:54:15.591 --> 00:54:19.758 (interference drowns out speaker)
- $1150\ 00:54:20.821 --> 00:54:22.657$ Yeah, sounds good, thank you.
- 1151 00:54:22.657 --> 00:54:25.649 (interference drowns out speaker)
- $1152\ 00:54:25.649 \longrightarrow 00:54:26.482$ Thank you.