WEBVTT

 $1~00:00:000.000 \dashrightarrow 00:00:02.613 < v$ Robert>Hi, I'm a Professor McDougal,</v>

 $2\ 00:00:06.358$ --> 00:00:07.990 and Professor Wayne is also in the back.

3 00:00:07.990 --> 00:00:11.060 If you haven't signed in, please make sure that you pass

 $4\ 00:00:11.060 \longrightarrow 00:00:13.310$ this, get a chance to sign the sign in sheet.

 $5\ 00:00:14.590 \dashrightarrow 00:00:19.260$ So today we are very, very privileged to be joined

6 00:00:19.260 --> 00:00:20.810 by Professor Naim Rashid

7 00:00:22.030 --> 00:00:25.360 from the University of North Carolina Chapel Hill,

8 00:00:25.360 --> 00:00:29.890 Professor Rashid got his bachelor's in biology from Duke,

900:00:29.890 --> 00:00:34.890 and his PhD in biostatistics from UNC Chapel Hill.

 $10\ 00:00:34.930$ --> 00:00:39.930 He's the author of 34 publications, and he holds a patent

11 $00:00:39.960 \rightarrow 00:00:44.410$ on methods in composition for prognostic

12 00:00:44.410 --> 00:00:47.313 and/or diagnostic supply chain of pancreatic cancer.

13 00:00:48.170 --> 00:00:50.640 He's currently an associate professor at UNC Chapel Hill's

14 00:00:50.640 --> 00:00:53.710 department of biostatistics, and he's also affiliated

 $15\ 00:00:53.710 \longrightarrow 00:00:56.903$ with their comprehensive cancer center there.

 $16\ 00:00:59.100 \dashrightarrow 00:01:04.100$ With that, Professor Rashid, would you like to take it away?

17 00:01:04.440 --> 00:01:05.920 <v ->Sure.</v>

 $18\ 00:01:05.920 \longrightarrow 00:01:08.470$ It looks like it says host disabled screen sharing.

 $19\ 00:01:10.344 \rightarrow 00:01:12.301$ (chuckling)

20 00:01:12.301 --> 00:01:13.760 <v Robert>All right, give me one second.</v>

21 00:01:13.760 --> 00:01:14.823 Thank you.

22 00:01:16.760 --> 00:01:17.883 I'm trying to do.

 $23\ 00:01:26.781 \longrightarrow 00:01:29.198$ (indistinct)

24 00:01:33.645 --> 00:01:35.901 Okay, you should be, you should be able to come on now.

25 00:01:35.901 --> 00:01:36.984 <v ->All right.</v>

26 00:01:38.584 --> 00:01:39.873 Can you guys see my screen?

27 00:01:43.650 --> 00:01:44.483 All right.

 $28\ 00:01:47.537 \longrightarrow 00:01:48.637$ Can you guys see this?

29 00:01:49.840 --> 00:01:50.913 <v Robert>There we go.</v>

 $30\ 00:01:52.062 \longrightarrow 00:01:52.895$ Perfect. Thank you.

31 00:01:52.895 --> 00:01:53.850 <v ->Okay, great.</v>

32 00:01:53.850 --> 00:01:56.501 So yes, thanks to the department for inviting me to speak

33 00:01:56.501 --> 00:02:00.483 today, and also thanks to Robert and Wayne for organizing.

34~00:02:01.460 --> 00:02:04.420 And today I'll be talking about issues regarding 35~00:02:04.420 --> 00:02:07.500 replicability in terms of clinical prediction models,

 $36\ 00:02:07.500$ --> 00:02:11.830 specifically in the context of genomic prediction models,

 $37\ 00:02:11.830 \longrightarrow 00:02:13.423$ derived from clinical trials.

learning

38 00:02:16.080 --> 00:02:17.870 So as an overview, we'll be talking first a little bit

 $39\ 00:02:17.870 \longrightarrow 00:02:20.670$ about the problems of replicability in general,

40 00:02:20.670 --> 00:02:24.300 in scientific research, and also about specific issues

41 00:02:24.300 --> 00:02:28.040 in genomics itself, and then I'll be moving on to talking

42 00:02:28.040 --> 00:02:31.070 about a method that we've proposed to assist 43 00:02:31.070 --> 00:02:34.380 with issues regarding data integration, and

 $44\ 00:02:34.380$ --> 00:02:37.680 in this environment when you have a heterogeneous data sets.

45 00:02:37.680 --> 00:02:39.860 I'll talk a little bit about a case study

46 $00:02:39.860 \rightarrow 00:02:42.901$ where we apply these practices to subtyping

 $47\ 00:02:42.901 \longrightarrow 00:02:44.670$ pancreatic cancer, touch on some current work

 $48\ 00:02:44.670 \longrightarrow 00:02:46.581$ that we're doing, and then end

 $49\ 00:02:46.581 --> 00:02:47.890$ with some concluding thoughts.

50 00:02:47.890 --> 00:02:49.861 And feel free to interrupt, you know,

 $51\ 00:02:49.861 \longrightarrow 00:02:52.211$ as the talk is long, if you have any questions.

 $52~00{:}02{:}53.540$ --> $00{:}02{:}55.650$ So I'm now an associate professor in the department

 $53\ 00:02:55.650 \longrightarrow 00:02:57.017$ of biostatistics at UNC.

54 00:02:58.160 --> 00:03:00.430 My work generally involves problems

55 00:03:00.430 --> 00:03:04.730 surrounding cancer and genomics, and more recently

 $56\ 00:03:04.730 \longrightarrow 00:03:07.390$ we've been doing work regarding epigenomics.

57 00:03:07.390 --> 00:03:09.370 We just recently published a supply-connected package called

58 00:03:09.370 --> 00:03:13.120 Epigram for a consistence of differential key calling,

 $59\ 00:03:13.120$ --> 00:03:15.480 and we've also done some work in model-based clustering.

6000:03:15.480 $\operatorname{-->}$ 00:03:18.310 We published a package called, FSCSeq,

 $61\ 00:03:18.310 \rightarrow 00:03:21.780$ which helps you derive and discover clusters

62 00:03:21.780 --> 00:03:23.830 from RNA seq data, while also determining

 $63\ 00:03:24.717 \longrightarrow 00:03:25.550$ clusters in specific genes.

64 00:03:25.550 --> 00:03:27.980 And today we'll be talking more about the topic

 $65\ 00:03:27.980 \longrightarrow 00:03:30.340$ of multi-study replicability, which is the topic

 $66\ 00:03:30.340 \longrightarrow 00:03:33.710$ of a paper that we published a year or two ago,

 $67\ 00{:}03{:}33{.}710$ --> $00{:}03{:}36{.}570$ and in our package that we've developed more recently,

 $68\ 00:03:36.570 \longrightarrow 00:03:38.463$ implementing some of the methods.

69 $00{:}03{:}40.090 \dashrightarrow 00{:}03{:}42.660$ So before I get deeper into the talk, one of the things

70 00:03:42.660 \rightarrow 00:03:45.130 I wanted to establish is this definition

 $71\ 00:03:45.130 \longrightarrow 00:03:47.090$ of what we mean by replicability.

 $72\ 00{:}03{:}47.090$ --> $00{:}03{:}49.670$ You might've heard the term reproducibility as well,

73 00:03:49.670 --> 00:03:52.430 and to make the distinction between the two terms,

74 00:03:52.430 $\rightarrow 00:03:54.140$ I'd like to define reproducibility in a way

 $75\ 00:03:54.140 \longrightarrow 00:03:56.910$ that Jeff Leak has defined in the past,

76 $00:03:56.910 \rightarrow 00:03:59.410$ where reproducibility is the ability to take

 $77\ 00{:}03{:}59{.}410$ --> $00{:}04{:}02{.}540$ coding data from a publication, and to rerun the code,

 $78\ 00:04:02.540$ --> 00:04:05.630 and get the same results as the original publication.

 $79\ 00{:}04{:}05{.}630 \dashrightarrow 00{:}04{:}08{.}650$ Where replicability, we're defining as the ability to be run

8000:04:08.650 --> 00:04:10.980 an experiment generating new data, and get results

 $81\ 00:04:10.980 \longrightarrow 00:04:12.780$ that are quote, unquote "consistent"

 $82\ 00:04:14.088 \longrightarrow 00:04:15.560$ with that of the original study.

83 00:04:15.560 --> 00:04:18.720 So in this sort of context, when it comes to replicability,

84 00:04:18.720 --> 00:04:21.890 you might've heard about publications that have come out

 $85\ 00:04:21.890$ --> 00:04:23.773 in the past that talk about how there are issues $86\ 00:04:23.773$ --> 00:04:27.600 regarding replicating the research that's been published

87 00:04:27.600 --> 00:04:29.570 in the scientific literature.

88 00:04:29.570 --> 00:04:32.280 This one paper in PLOS Medicine was published

89 00:04:32.280 --> 00:04:36.150 by, and that is in 2005, and there's been a number

90 00:04:36.150 --> 00:04:37.920 of publications that have come out since,

91 00:04:37.920 --> 00:04:40.880 talking about problems regarding replicability,

 $92\ 00:04:40.880 \longrightarrow 00:04:43.290$ and ways that we could potentially address it.

93 00:04:43.290 --> 00:04:45.820 And the problem has become large enough where it has

94 00:04:45.820 --> 00:04:48.840 its own Wikipedia entry talking about the crisis,

 $95\ 00:04:48.840 \longrightarrow 00:04:51.300$ and has a long list of examples that talks

96 00:04:51.300 --> 00:04:54.170 about issues regarding replicating results

97 00:04:54.170 --> 00:04:55.400 from the scientific studies.

98 00:04:55.400 --> 00:04:57.550 So this is something that has been a known issue

99 00:04:57.550 $\rightarrow 00:05:00.320$ for a while, and these problems also extend

 $100\ 00:05:00.320 \longrightarrow 00:05:03.270$ to situations where you want to, for example,

101 00:05:03.270 --> 00:05:06.300 develop clinical prediction models in genomics.

102 00:05:06.300 --> 00:05:10.280 So to give an example of this, let's say that we wanted to,

103 00:05:10.280 --> 00:05:13.200 in the population of metastatic breast cancer patients,

 $104 \ 00:05:13.200 \longrightarrow 00:05:15.710$ we wanted to develop a model that predicts

105 00:05:15.710 --> 00:05:18.170 some clinical outcome Y, given a set

 $106\ 00:05:18.170 \longrightarrow 00:05:20.530$ of gene expression values X.

107 00:05:20.530 --> 00:05:23.020 And so the purpose of this sort of exercise is

 $108\ 00:05:23.020 \longrightarrow 00:05:26.120$ to hopefully translate this sort of model

109 00:05:26.120 --> 00:05:27.930 that we've developed, and apply it to the clinic,

110 00:05:27.930 --> 00:05:31.030 where we can use it for clinical decision-making.

111 00:05:31.030 --> 00:05:34.653 Now, if we have data from one particular trial 112 00:05:34.653 --> 00:05:36.960 that pertains to this patient population,

 $113\ 00:05:36.960\ -->\ 00:05:39.020$ and the same clinical outcome being measured,

114 $00{:}05{:}39{.}020 \dashrightarrow 00{:}05{:}40{.}640$ in addition to having gene expression data,

 $115\ 00:05:40.640 \longrightarrow 00:05:42.640$ let's say that we derived a model, let's say

116 00:05:42.640 --> 00:05:44.470 that we're modeling some sort of binary outcome,

 $117\ 00:05:44.470 \longrightarrow 00:05:45.800$ let's say tumor response.

 $118\ 00:05:45.800 \longrightarrow 00:05:48.190$ And in this model, we used a cost report,

119 $00:05:48.190 \dashrightarrow 00:05:51.110$ or penalized logistic regression model

120 00:05:51.110 --> 00:05:54.060 that we fit to the data to try and predict the outcome,

121 00:05:54.060 --> 00:05:55.940 given the gene expression values.

 $122\ 00:05:55.940 \longrightarrow 00:05:58.770$ And here we obtained, let's say, 12 genes

123 00:05:58.770 --> 00:06:03.640 after the fitting process, and the internal model 1 UNC

 $124\ 00:06:03.640 \longrightarrow 00:06:05.733$ on the sort of training subjects is 0.9.

125 00:06:06.740 --> 00:06:08.500 But then let's say there's another group at Duke

 $126\ 00:06:08.500 \longrightarrow 00:06:10.870$ that's using data from their clinical trial,

 $127\ 00:06:10.870 \longrightarrow 00:06:13.197$ and they have a larger sample size.

128 00:06:13.197 --> 00:06:15.870 They also found more genes, 65 genes,

129 00:06:15.870 \rightarrow 00:06:18.211 but have a slightly lower training at UNC.

130 00:06:18.211 --> 00:06:21.910 However, we really need to use external validation

131 00:06:21.910 --> 00:06:25.150 to sort of get an independent assessment of how well

 $132\ 00:06:25.150 \longrightarrow 00:06:27.340$ each one of these alternative models are doing.

133 00:06:27.340 --> 00:06:29.807 So let's say we have data from a similar study from Harvard,

134 00:06:29.807 \rightarrow 00:06:31.740 and we applied both these train models

135 00:06:32.615 --> 00:06:35.260 to the genomic data from that study at Harvard.

136 00:06:35.260 --> 00:06:37.790 We have the outcome information for those patients as well,

137 00:06:37.790 --> 00:06:42.153 so we can calculate how well the model predicts

 $138\ 00:06:42.153 \longrightarrow 00:06:44.487$ on those validation subjects.

 $139\ 00:06:44.487 \longrightarrow 00:06:46.240$ And we find here in this data set,

140 00:06:46.240 --> 00:06:48.740 model 2 seems to be doing better than model 1,

141 00:06:48.740 --> 00:06:50.870 but if you try this again with another data set

142 00:06:50.870 --> 00:06:53.470 from Michigan, you might find that model 1 is doing

143 00:06:53.470 --> 00:06:54.730 better, better than model 2.

144 00:06:54.730 --> 00:06:57.640 So the problem here is where we have researchers

145 00:06:57.640 --> 00:06:58.960 that are pointing fingers at each other,

146 00:06:58.960 --> 00:07:01.470 and it's really hard to know, "Well, who's who's right?"

147 00:07:01.470 --> 00:07:03.580 And why is this even happening in the first place,

148 00:07:03.580 --> 00:07:05.938 in terms of why do we get different genes, numbers of genes,

149 00:07:05.938 --> 00:07:08.797 and each of the models derived from study 1 and study 2?

150 00:07:08.797 --> 00:07:11.770 And why are we seeing very low performance

 $151\ 00:07:11.770 \longrightarrow 00:07:13.620$ in some of these validation datasets?

 $152\ 00:07:15.290 \longrightarrow 00:07:17.330$ So here's an example from 2014,

153 00:07:17.330 --> 00:07:19.600 in the context of ovarian cancer.

154 00:07:19.600 --> 00:07:22.410 The authors basically collected 10 studies,

 $155\ 00:07:22.410 \longrightarrow 00:07:24.063$ all were microarray studies.

 $156\ 00:07:24.920 \longrightarrow 00:07:27.200$ The goal here was to predict overall survival

157 00:07:27.200 --> 00:07:29.550 in this population of ovarian cancer patients,

158 00:07:29.550 $\rightarrow 00:07:31.870$ given gene expression measurements

 $159\ 00:07:31.870 \longrightarrow 00:07:33.800$ from this microarray platform.

 $160\ 00:07:33.800 \longrightarrow 00:07:34.633$ So through a series

161 00:07:34.633 --> 00:07:38.640 of really complicated cross-fertilization approaches,

 $162\ 00:07:38.640 \longrightarrow 00:07:40.430$ the data was normalized, and harmonized

163 00:07:40.430 --> 00:07:43.413 across the studies, using a combination of ComBat

164 00:07:43.413 --> 00:07:45.639 and frozen RNA, and then they took

165 00:07:45.639 --> 00:07:47.640 14 published prediction models in the literature,

 $166\ 00:07:47.640$ --> 00:07:50.970 and they applied each of those models to each $167\ 00:07:50.970$ --> 00:07:53.255 of the subjects from these 10 studies, and they compared

 $168\ 00:07:53.255 \longrightarrow 00:07:57.590$ the model predictions across each subject.

 $169\ 00:07:57.590 \longrightarrow 00:08:00.490$ So each column here in this matrix is a patient,

170 00:08:00.490 --> 00:08:03.060 and each row is a different prediction model,

171 00:08:03.060 \rightarrow 00:08:06.260 and each cell represents the prediction

 $172\ 00:08:06.260 \longrightarrow 00:08:08.090$ from that model on that patient.

 $173\ 00:08:08.090$ --> 00:08:11.700 So an ideal scenario, where we have the models generalizing

174 00:08:11.700 --> 00:08:14.480 and replicating across each of these individuals,

 $175\ 00:08:14.480 \longrightarrow 00:08:15.860$ we would expect to see the column,

 $176\ 00:08:15.860 \longrightarrow 00:08:18.919$ each column here to have the same color value,

 $177\ 00:08:18.919 \longrightarrow 00:08:20.080$ meaning that the predictions are consistent.

178 00:08:20.080 --> 00:08:22.220 But clearly we see here that the predictions are

179 00:08:22.220 --> 00:08:24.310 actually very inconsistent,

 $180\ 00{:}08{:}24.310$ --> $00{:}08{:}26.060$ and very different from each other.

181 00:08:27.230 --> 00:08:28.220 In addition, if you look

 $182\ 00:08:28.220 \longrightarrow 00:08:30.410$ at the individual risk prediction models

 $183\ 00:08:30.410 \longrightarrow 00:08:31.990$ that the authors used, there was also

184 00:08:31.990 --> 00:08:33.770 substantial differences in the genes

 $185\ 00:08:33.770 \longrightarrow 00:08:36.210$ that were selected in each of these models.

186 00:08:36.210 --> 00:08:39.760 So there's a max 2% overlap in terms of common genes

 $187\ 00:08:39.760 \longrightarrow 00:08:41.350$ between each of these approaches.

188 00:08:41.350 --> 00:08:43.150 And one thing to mention here is that each one

 $189\ 00:08:43.150 \longrightarrow 00:08:45.380$ of these risk-prediction models were derived

190 00:08:45.380 --> 00:08:48.270 from separate individual studies.

191 00:08:48.270 --> 00:08:50.631 So the question here is, you know, how exactly,

192
 00:08:50.631 --> 00:08:53.669 if you were a clinician, you're eager to sort of take

 $193\ 00:08:53.669 \longrightarrow 00:08:57.020$ the results that you're seeing here,

 $194\ 00:08:57.020 \longrightarrow 00:08:58.430$ and extend to the clinic,

 $195\ 00:08:58.430 \longrightarrow 00:09:00.860$ which model do you use, which is right?

 $196\ 00:09:00.860 \longrightarrow 00:09:02.610$ Why are you seeing this level of variability?

197 00:09:02.610 --> 00:09:05.840 This is, of course, concerning, if you, if your goal is

198
 $00{:}09{:}05{.}840$ --> $00{:}09{:}08{.}070$ to move things towards the clinic, and this also has

199 $00{:}09{:}08{.}070 \dashrightarrow 00{:}09{:}11.250$ implications in terms of, you know, getting in the way

 $200\ 00:09:11.250 \longrightarrow 00:09:12.980$ of trying to approve the use of some

 $201\ 00:09:12.980 \longrightarrow 00:09:15.453$ of these, and for clinical use.

 $202\ 00:09:17.360 \longrightarrow 00:09:18.950$ So why is this happening?

 $203\;00{:}09{:}18.950 \dashrightarrow 00{:}09{:}21.600$ So there's been a lot of studies have been done

 $204\;00{:}09{:}21.600 \dashrightarrow 00{:}09{:}24.487$ that have tied issues to, obviously, sample size

 $205\ 00:09:24.487 \rightarrow 00:09:27.160$ in the training studies, smaller sample sizes,

 $206\;00{:}09{:}27.160 \dashrightarrow > 00{:}09{:}30.710$ and models trained on them may lead to more unstable models,

 $207\ 00:09:30.710 \longrightarrow 00:09:32.182$ or less accurate models.

 $208\ 00:09:32.182 \longrightarrow 00:09:34.765$ Between different studies, you might have

 $209\ 00:09:34.765 \longrightarrow 00:09:36.080$ different prevalences of the clinical outcome.

210 $00{:}09{:}36.080 \dashrightarrow> 00{:}09{:}38.640$ In some studies, you might have higher levels of response,

211 00:09:38.640 --> 00:09:40.390 and other studies, you might have lower levels of response,

212 00:09:40.390 --> 00:09:42.920 for example, if you have this binary clinical outcome,

 $213\ 00:09:42.920 \longrightarrow 00:09:46.290$ and also there's issues regarding differences

 $214\ 00:09:46.290$ --> 00:09:49.090 in lab conditions, where the genomic data was extracted.

215 00:09:49.090 --> 00:09:51.630 We've seen at Lineberger that, depending on the type

216 00:09:51.630 --> 00:09:54.570 of extraction, RNA extraction kit that you use,

217 00:09:54.570 --> 00:09:57.740 you might see differences in the expression of a gene,

 $218\ 00:09:57.740 \longrightarrow 00:10:00.010$ even from the same original tumor.

219 $00:10:00.010 \dashrightarrow 00:10:01.640$ And also the issue of batch placement,

220 00:10:01.640 --> 00:10:03.730 which has been widely talked about in the literature,

221 00:10:03.730 --> 00:10:06.170 where depending on the day you run the experiment,

 $222\ 00:10:06.170 \longrightarrow 00:10:10.500$ or the technician who's handling the data,

223 00:10:10.500 --> 00:10:12.023 you might see slight differences,

224 00:10:12.023 --> 00:10:14.263 technical differences in expression.

 $225\ 00:10:15.380 \longrightarrow 00:10:16.810$ There's also differences due to protocols.

 $226\ 00:10:16.810 \longrightarrow 00:10:18.460$ Some trials might have different inclusion

227 00:10:18.460 --> 00:10:20.560 and exclusion criteria, so they might be recruiting

 $228\ 00:10:20.560 \longrightarrow 00:10:22.280$ a slightly different patient population,

229 00:10:22.280 --> 00:10:23.640 even though they might be all

230 00:10:23.640 \rightarrow 00:10:25.240 in the context of metastatic breast cancer.

231 00:10:25.240 --> 00:10:29.161 All of these things can help impart heterogeneity

232 00:10:29.161 --> 00:10:33.590 between what the genomic data and the outcome data

233 00:10:33.590 --> 00:10:36.120 across different studies.

234 00:10:36.120 --> 00:10:38.710 In the context of genomic data in particular,

 $235\ 00:10:38.710 \longrightarrow 00:10:41.280$ there's also this aspect of data preprocessing.

236 00:10:41.280 --> 00:10:44.510 For the normalization taking that you use is very important,

 $237\ 00:10:44.510 \longrightarrow 00:10:46.630$ and we'll talk about that in a little bit.

238 00:10:46.630 $\rightarrow 00:10:48.330$ And it's a very critical part when it comes

239 00:10:48.330 --> 00:10:51.680 to training models, and trying to validate your model

240 00:10:51.680 --> 00:10:54.023 on other datasets, and depending on the type 241 00:10:54.023 --> 00:10:57.923 of normalization you use, this could also impact

 $242\ 00:10:57.923 \longrightarrow 00:10:59.623$ how well your model works.

243 00:11:00.480 --> 00:11:03.427 In addition, there's also differences in the potential way

 $244\ 00:11:03.427 \longrightarrow 00:11:04.470$ in which you measure gene expression.

245 00:11:04.470 --> 00:11:07.410 Some trials might use an older technology called microarray.

246 00:11:07.410 --> 00:11:08.940 I know other trials might use something

247 00:11:08.940 --> 00:11:11.490 relatively more recent called RNAC,

248 00:11:11.490 --> 00:11:12.593 or a particular trial might use

249 00:11:12.593 --> 00:11:14.910 a more targeted platform like NanoString.

 $250\ 00{:}11{:}14{.}910$ --> $00{:}11{:}19{.}087$ So the differences in platform also can lead to differences

251 00:11:19.087 --> 00:11:21.470 in your ability to help validate some of these studies.

252 00:11:21.470 --> 00:11:23.870 If you train something in marker rate, it's very difficult

253 00:11:23.870 --> 00:11:26.360 to take that model, and apply it to RNAC,

254 00:11:26.360 --> 00:11:29.900 because the expression values are just are just different.

255 00:11:29.900 --> 00:11:32.450 And so, as I mentioned before, this also impacts

 $256\ 00{:}11{:}32{.}450$ --> $00{:}11{:}37{.}180$ through to normalization on model performance as well.

 $257\ 00:11:37.180 \longrightarrow 00:11:39.660$ So the main thing to remember here is that

258 00:11:39.660 --> 00:11:43.080 the traditional way in which prediction models,

259 00:11:43.080 --> 00:11:46.130 based on genomic data for using the clinical training is

 $260\ 00:11:46.130 \longrightarrow 00:11:49.343$ typically on the results from a single study.

261 00:11:51.760 --> 00:11:53.510 To talk a little bit more about question

262 00:11:53.510 --> 00:11:57.260 of between-study normalization, and the purpose of this is

263 00:11:57.260 --> 00:12:00.360 to put the expression data on basically an even scale,

 $264\ 00:12:00.360 \longrightarrow 00:12:02.330$ which helps facilitate training.

265 00:12:02.330 --> 00:12:05.510 If there's global shifts, and some of the expression values

 $266\;00{:}12{:}05{.}510$ --> $00{:}12{:}08{.}820$ in one sample versus another, it's very difficult to train

 $267\ 00:12:08.820$ --> 00:12:11.090 an accurate model in that particular scenario.

268 00:12:11.090 --> 00:12:13.213 So normalization helps to align

 $269\ 00:12:13.213 \longrightarrow 00:12:15.600$ the expression you get from different samples,

270 00:12:15.600 --> 00:12:19.020 and hopefully across the between difference as well.

271 00:12:19.020 --> 00:12:23.090 And so the goal here is to eventually predict this outcome

272 00:12:23.090 --> 00:12:25.110 in a new patient, you plug in the genomic data

273 00:12:25.110 --> 00:12:28.190 from a new patient in order to get the predicted outcome

 $274\ 00:12:28.190 \longrightarrow 00:12:30.350$ for that patient based on that training model.

275 00:12:30.350 --> 00:12:33.650 So the, in order to do that, you also have to normalize

276 00:12:33.650 --> 00:12:35.910 the new data to the training data, right?

277 00:12:35.910 --> 00:12:38.151 Because you also want to put the new data on the same scale

 $278\ 00:12:38.151 \longrightarrow 00:12:41.450$ as a training data, and in the ideal scenario,

279 00:12:41.450 --> 00:12:43.610 you would want to make sure that the training samples

 $280\ 00{:}12{:}43.610$ --> $00{:}12{:}47.150$ that you use to train your original model are untouched,

281 00:12:47.150 --> 00:12:49.120 because what some people try to do is they try

 $282\ 00:12:49.120 \longrightarrow 00:12:52.140$ to sort of sidestep this normalization issue,

283 00:12:52.140 --> 00:12:54.644 they would combine the new data with the old training data,

 $284\ 00:12:54.644 \longrightarrow 00:12:57.160$ and renormalize everything at once.

 $285\ 00{:}12{:}57.160 \dashrightarrow 00{:}12{:}58.790$ And the problem with this is that this changes

 $286\ 00:12:58.790 \longrightarrow 00:13:00.727$ your training sample values, and in a sense,

287 00:13:00.727 --> 00:13:03.640 would necessitate the fact that you need to retrain

288 00:13:03.640 --> 00:13:04.473 your old model again.

289 00:13:04.473 --> 00:13:06.950 And this leads to instability, and lack of stability

 $290\ 00:13:06.950 \longrightarrow 00:13:09.333$ over time in terms of the model itself.

291 00:13:10.270 \rightarrow 00:13:12.231 So in the prior example from ovarian cancer,

 $292\ 00:13:12.231 \longrightarrow 00:13:14.950$ this is not as big of an issue, because you have

 $293\ 00:13:14.950 \longrightarrow 00:13:17.590$ all the data you want to work with in hand.

294 00:13:17.590 --> 00:13:19.670 This is a retrospective study, you have 10 data sets,

295 00:13:19.670 --> 00:13:22.450 so you just normalize everything at the same time,

296 00:13:22.450 --> 00:13:23.960 that's in ComBat and frozen RNA.

297 00:13:23.960 --> 00:13:26.950 And so you can split up those studies into separate training

298 00:13:26.950 --> 00:13:30.750 and test studies, and they're all rated on the same scale.

299 00:13:30.750 --> 00:13:34.250 But the problem is that in practice, you're trying to do

300 00:13:34.250 --> 00:13:37.130 a prospective type of analysis, where when you train

301 00:13:37.130 --> 00:13:40.300 your model, you're normalizing all of the available studies

302 00:13:40.300 --> 00:13:43.690 you have, let's say, and then you use that to predict

 $303\ 00{:}13{:}43.690$ --> $00{:}13{:}47.010$ the outcome in a future patient, or a future study.

 $304\;00{:}13{:}47.010 \dashrightarrow 00{:}13{:}51.150$ And so the problem with that is that you have to find

 $305\ 00:13:51.150 -> 00:13:54.610$ a good way to align, as I mentioned before,

 $306~00{:}13{:}54.610 \dashrightarrow 00{:}13{:}56.780$ the data from that future study for your training samples,

 $307\ 00:13:56.780 \longrightarrow 00:14:00.080$ and that may not be an easy task to do,

30800:14:00.080 --> 00:14:02.730 especially for some of the newer platforms like RNAC.

309 00:14:04.160 --> 00:14:06.165 So taking this problem a step further,

310 00:14:06.165 --> 00:14:09.830 what if there's no good cross study normalization approach

 $311\ 00:14:09.830 \longrightarrow 00:14:12.200$ that's available to begin with?

312 00:14:12.200 --> 00:14:15.200 This really is going to make things difficult in terms

313 00:14:15.200 --> 00:14:17.560 of the training in the model in the first place.

314 00:14:17.560 --> 00:14:20.860 Another more complicated problem is that you might have

315 00:14:20.860 --> 00:14:23.770 different types of platforms at that training time.

316 00:14:23.770 --> 00:14:26.040 For example, you might have the only type of data

317 00:14:26.040 --> 00:14:29.160 that's available from one study is NanoString in one case,

318 00:14:29.160 --> 00:14:32.640 and another study it's only RNAC, so what do you do?

319 $00{:}14{:}32.640 \dashrightarrow 00{:}14{:}35.250$ And looking forward, as platforms change,

320 00:14:35.250 --> 00:14:36.382 as technology evolves, you have different ways

321 00:14:36.382 --> 00:14:41.382 of measuring gene expression, for example.

322 00:14:41.950 --> 00:14:44.440 So what do you do with the models that are trained

323 00:14:44.440 --> 00:14:48.060 on old data, because you can't apply them to the new data?

324 00:14:48.060 --> 00:14:49.770 So oftentimes you find this situation

 $325\ 00{:}14{:}49{.}770 \dashrightarrow 00{:}14{:}53{.}470$ where you have to retrain new models on these new platforms,

 $326\ 00:14:53.470 \longrightarrow 00:14:57.000$ and the old models are not able to be applied $327\ 00:14:57.000 \longrightarrow 00:14:58.440$ directly to this new data types.

328 00:14:58.440 --> 00:15:00.690 So that leads to waste here.

329 00:15:00.690 --> 00:15:03.370 So if you take all of these problems together,

 $330\ 00:15:03.370 \longrightarrow 00:15:07.320$ regarding cross-study normalization,

 $331\ 00:15:07.320 \longrightarrow 00:15:09.300$ and changes in platform,

 $332\ 00:15:09.300 \longrightarrow 00:15:11.390$ and a lot of the other issues, you know,

333 00:15:11.390 --> 00:15:13.280 regarding replicability that I mentioned,

 $334\ 00:15:13.280 \longrightarrow 00:15:16.580$ it's no wonder that there's only a small handful

335 00:15:16.580 --> 00:15:21.430 of expression-based clinically applicable assets have been

336 00:15:21.430 --> 00:15:23.777 approved by the FDA, like Oncotype DX, MammaPrint,

337 00:15:23.777 --> 00:15:27.203 and Prosigna, because this is a very, very tough problem.

 $338\;00{:}15{:}29{.}884 {--}{>}\;00{:}15{:}32{.}600$ So I want to move on with that, to an approach 339 $00{:}15{:}32{.}600 {--}{>}\;00{:}15{:}36{.}130$ that we proposed to help tackle this sort of issue

 $340\ 00:15:36.130 \longrightarrow 00:15:39.210$ by using this idea of multi-study learning,

341 00:15:39.210 --> 00:15:43.020 where instead of just using, and deriving, and generating

342 00:15:43.020 --> 00:15:44.810 models from individual studies, we combine data

343 00:15:44.810 --> 00:15:47.790 from multiple studies together, and create a consensus model

344 00:15:47.790 --> 00:15:50.110 that we use for prediction, which will hopefully be

345 00:15:50.110 --> 00:15:54.140 more stable, and more accurate down the road.

 $346\ 00:15:54.140 \longrightarrow 00:15:56.400$ So this approach of combining data is called

347 00:15:56.400 --> 00:15:59.190 horizontal data integration, where we're merging data

348 00:15:59.190 --> 00:16:01.360 from let's say K different studies.

349 00:16:01.360 --> 00:16:04.300 And the pro of this approach is that we get increased power,

 $350\ 00:16:04.300 \longrightarrow 00:16:06.160$ and the ability to reach some sort of consensus

 $351\ 00:16:06.160 \longrightarrow 00:16:08.860$ across these different studies.

 $352\ 00:16:08.860 \longrightarrow 00:16:11.650$ The negative is that the effect of a gene

353 00:16:11.650 --> 00:16:13.710 and its relationship to outcome may actually vary

 $354~00{:}16{:}13.710 \dashrightarrow 00{:}16{:}16.040$ across studies, and also by, you know, depending on,

 $355\;00{:}16{:}16{.}040 \dashrightarrow 00{:}16{:}18{.}940$ and also the way that you normalize the genes may also vary

356 00:16:18.940 --> 00:16:21.178 across studies too if we're using published data

 $357\ 00:16:21.178 \longrightarrow 00:16:23.630$ from some prior publication.

358 00:16:23.630 --> 00:16:25.470 There's also this issue of sample size and balance.

 $359\ 00:16:25.470 \longrightarrow 00:16:27.630$ You might have a study that has 500 subjects,

 $360\ 00:16:27.630 \longrightarrow 00:16:29.860$ and another one that might have 200 subjects.

361 00:16:29.860 --> 00:16:33.820 So there are some methods that were designed to account for

362 00:16:33.820 --> 00:16:36.190 between-study heterogeneity after you do

 $363\ 00:16:36.190 \longrightarrow 00:16:37.830$ horizontal data integration.

364 00:16:37.830 --> 00:16:41.040 One is called the meta-lasso, another is called 365 00:16:41.040 --> 00:16:43.590 the AW statistic, but these two methods don't really have

 $366\ 00:16:43.590 \longrightarrow 00:16:46.370$ any prediction aspect about them.

367 00:16:46.370 --> 00:16:48.496 They're more about feature selection.

368 00:16:48.496 --> 00:16:50.420 Ensembling is one approach that can directly account

 $369\ 00:16:50.420 \longrightarrow 00:16:52.310$ for between-study heterogeneity

 $370\ 00:16:52.310 \longrightarrow 00:16:54.350$ after horizontal data integration, but there's

 $371\ 00:16:54.350 \longrightarrow 00:16:56.870$ no explicit future selection step here.

 $372\ 00:16:56.870 \longrightarrow 00:16:58.800$ But all of these approaches assume

 $373\ 00:16:58.800 \longrightarrow 00:17:01.670$ that the data has been pre-normalized.

 $374\ 00:17:01.670 \longrightarrow 00:17:03.350$ As we talked about before,

375 00:17:03.350 --> 00:17:06.820 for prospective decision-making, based off a train model,

 $376\ 00:17:06.820 \longrightarrow 00:17:10.070$ that might be prohibitive in some cases,

 $377\ 00:17:10.070 \longrightarrow 00:17:13.380$ and we need a strategy also to easily predict $378\ 00:17:13.380 \longrightarrow 00:17:17.153$ and apply these models in new patients.

379 00:17:20.260 --> 00:17:24.080 Okay, so moving on, we're going to talk first 380 00:17:24.080 --> 00:17:26.670 about this issue of how do we integrate data,

381 00:17:26.670 --> 00:17:30.300 and sort of sidestep this normalization problem

382 00:17:30.300 --> 00:17:33.190 at training time, and also at test time where we,

 $383\ 00:17:33.190 \longrightarrow 00:17:35.040$ when we try to predict in new subjects?

384 00:17:35.040 --> 00:17:38.520 So the approach that we put forth is to use

385 00:17:38.520 --> 00:17:40.860 what's called top scoring pairs, which you can think of

386 00:17:40.860 --> 00:17:44.560 as a rank-based transformation of the original set

 $387\ 00:17:44.560 \rightarrow 00:17:47.320$ of gene expression values from a patient.

 $388\ 00:17:47.320 \longrightarrow 00:17:49.510$ So the idea here originally,

389 $00{:}17{:}49{.}510 \dashrightarrow 00{:}17{:}50{.}630$ when top scoring pairs were introduced,

390 00:17:50.630 --> 00:17:53.390 was you're trying to find a pair of genes

391 00:17:53.390 --> 00:17:56.390 where it's such that if the expression of gene A

392 00:17:56.390 --> 00:17:58.908 in the pair is greater than gene B, that would imply

393 00:17:58.908 --> 00:18:02.970 that the, let's say, the subtype for that individual is,

 $394\ 00:18:02.970 \longrightarrow 00:18:05.490$ say, subtype one, and if it's less,

 $395\ 00{:}18{:}05{.}490 \dashrightarrow 00{:}18{:}09{.}080$ then that implies subtype zero with high probability.

396 00:18:09.080 --> 00:18:11.760 Now, in this case, this sort of approach was developed

 $397\ 00:18:11.760 \longrightarrow 00:18:14.100$ with when one has a binary outcome variable $398\ 00:18:14.100 \longrightarrow 00:18:15.070$ that you care about.

399 00:18:15.070 --> 00:18:17.430 In this case, we're talking about subtype,

400 00:18:17.430 --> 00:18:20.040 but it could also be tumor response or something else.

401 00:18:20.040 --> 00:18:22.070 So essentially what you're doing is that you're taking

 $402\ 00{:}18{:}22.070$ --> $00{:}18{:}25.270$ these continuous measurements in terms of gene expression,

 $403\ 00{:}18{:}25{.}270 \dashrightarrow 00{:}18{:}30{.}270$ or integer, and you are converting that, transforming

 $404\ 00:18:30.600 \longrightarrow 00:18:32.230$ that into basically a binary predictor,

 $405\ 00:18:32.230 \longrightarrow 00:18:34.457$ which takes on the value of the zero or one.

406 00:18:34.457 --> 00:18:38.210 And the hope is that that particular transformed value is

407 00:18:38.210 --> 00:18:41.300 going to be associated with this binary outcome.

408 00:18:41.300 --> 00:18:43.760 So the simple assumption in this scenario is

409 00:18:43.760 --> 00:18:46.100 that the relative rank of these genes

 $410\;00{:}18{:}46{.}100 \dashrightarrow 00{:}18{:}50{.}810$ in a given sample is predictive of subtype, and that's it.

411 00:18:50.810 --> 00:18:54.490 And so the example here I have on the right is an example

 $412\ 00:18:54.490 \longrightarrow 00:18:57.790$ of two genes, GSTP1 and ESR1.

 $413\ 00:18:57.790 \longrightarrow 00:18:59.928$ And so you can see here that if you're

414 00:18:59.928 --> 00:19:02.300 in the upper left quadrant, this is where this gene is

 $415\ 00:19:02.300 \longrightarrow 00:19:04.860$ greater than this gene expression, it's implying

416 $00:19:04.860 \rightarrow 00:19:07.648$ the triangle subtype with high probability,

417 $00:19:07.648 \rightarrow 00:19:10.900$ and otherwise it implies the circle subtype.

418 00:19:10.900 --> 00:19:14.350 So that's the general idea of what we're going for here.

419 $00{:}19{:}14.350 \dashrightarrow 00{:}19{:}16.480$ It's a sort of a rank-based transformation

 $420\ 00:19:16.480 \longrightarrow 00:19:19.643$ of the original continuous predictor space.

 $421\ 00:19:20.750 \longrightarrow 00:19:22.100$ So the nice thing about this approach,

 $422\ 00{:}19{:}22{.}100$ --> $00{:}19{:}24{.}643$ because we're only based on the simple assumption, right?

 $423\ 00:19:24.643 \longrightarrow 00:19:26.710$ That we're only caring about the relative rank

 $424\ 00:19:26.710 \longrightarrow 00:19:28.880$ within a subject, this makes

 $425\ 00:19:28.880 \longrightarrow 00:19:32.450$ this particular new transformed predictor

 $426\ 00{:}19{:}32{.}450$ --> $00{:}19{:}35{.}710$ relatively invariant to batch effects, prenormalization,

427 00:19:35.710 --> 00:19:39.410 and it also most importantly, simplifies merging data

 $428\ 00:19:39.410 \longrightarrow 00:19:40.580$ from different studies.

429 00:19:40.580 --> 00:19:43.090 Everything is now on the same scale, zero to one,

 $430\ 00:19:43.090 \longrightarrow 00:19:44.987$ so it's very easy to paste together the data

431 00:19:44.987 --> 00:19:49.910 from different studies, and we can sidestep this problem

432 00:19:49.910 --> 00:19:52.870 of trying to pick a cross-normalization approach,

433 00:19:52.870 --> 00:19:55.803 and then work in this sort of transformed space.

434 00:19:56.840 --> 00:19:59.130 The other nice thing is that this is easily computable

 $435\ 00:19:59.130 \longrightarrow 00:20:00.690$ for new patients as well.

436 00:20:00.690 --> 00:20:02.670 If you have a new patient that comes into clinic,

437 00:20:02.670 --> 00:20:04.220 you just check to see whether the gene A is

438 00:20:04.220 $-\!>$ 00:20:06.290 greater than gene B in terms of expression,

439 00:20:06.290 --> 00:20:11.290 and then you have your value for this top scoring pair,

440 00:20:11.350 --> 00:20:14.430 and we don't have to worry as much about normalizing

441 00:20:14.430 --> 00:20:17.740 this patient's raw gene spectrum data

442 00:20:17.740 $\rightarrow 00:20:21.470$ to the training sample expression values.

443 00:20:21.470 --> 00:20:23.360 So essentially what we're doing here is that we're,

444 00:20:23.360 --> 00:20:25.700 let's enumerate all possible gene pairs for us, 445 00:20:25.700 --> 00:20:28.200 instead of a candidate genes, and each column here

446 00:20:28.200 -> 00:20:30.530 in this matrix shown on the right pertains

447 00:20:30.530 --> 00:20:33.867 to the zero one values for a particular gene pair J.

448 00:20:33.867 --> 00:20:37.960 And so this value takes the value of one, it is greater

449 00:20:37.960 --> 00:20:41.200 than B, in sample I, in pair j, and zero otherwise.

450 00:20:41.200 --> 00:20:44.603 And then we merge over the common top scoring pairs.

 $451\ 00{:}20{:}46.070$ --> $00{:}20{:}49.050$ So in this example have data from four different studies,

452 00:20:49.050 --> 00:20:50.420 each indicator by a different color here

453 00:20:50.420 --> 00:20:53.750 in the first track, and this data pertains to data

454 00:20:53.750 --> 00:20:54.900 from two different platforms,

 $455\ 00:20:54.900 \longrightarrow 00:20:56.437$ and three different cancer types.

456 00:20:56.437 --> 00:20:59.220 And so the clinical outcome here is binary subtype,

457 00:20:59.220 --> 00:21:02.220 which is given by the orange and the blue color here.

458 00:21:02.220 --> 00:21:05.350 So you can see here that we enumerated the TSPs,

459 00:21:05.350 --> 00:21:07.190 we merged the data together, and now we have

 $460\ 00:21:07.190 \longrightarrow 00:21:09.340$ this transformed predictor agents.

 $461\ 00:21:09.340 \longrightarrow 00:21:10.430$ And the interesting thing is

462 00:21:10.430 --> 00:21:12.620 that you can definitely see some patterning here.

463 00:21:12.620 --> 00:21:15.290 With any study where you have a particular set of TSPs

464 00:21:15.290 --> 00:21:18.950 that had taken a value of one, when the sub-type is blue,

 $465\ 00:21:18.950 \longrightarrow 00:21:20.850$ and it flips when it's orange.

466 00:21:20.850 --> 00:21:24.230 And we see the same general pattern seem to replicate

467 00:21:24.230 --> 00:21:25.380 across different studies,

468 00:21:25.380 --> 00:21:29.168 but not every top scoring pair changes the same way

 $469\ 00:21:29.168 \longrightarrow 00:21:31.700$ across different studies.

 $470\ 00:21:31.700 \longrightarrow 00:21:34.970$ So if we cluster the rows here, we can also see

471 00:21:34.970 --> 00:21:38.120 some patterns sort of persist where we see

472 00:21:38.120 --> 00:21:39.770 some clustering by subtype,

 $473\ 00:21:39.770 \longrightarrow 00:21:41.830$ but also some clustering by study as well.

474 00:21:41.830 --> 00:21:44.620 And so what this implies is that there's a relationship

475 00:21:44.620 --> 00:21:47.108 between TSPs and subtypes, and that can vary across studies,

476 00:21:47.108 --> 00:21:50.107 which is not too different from what we've talked

 $477\ 00:21:50.107 - 00:21:51.380$ about regarding the issues we've seen

 $478\ 00:21:51.380 \longrightarrow 00:21:53.339$ in replicability in the past.

479 00:21:53.339 --> 00:21:57.460 So ideally we would like to see a particular gene pair,

 $480\ 00:21:57.460 \longrightarrow 00:22:00.810$ or TSP vector here take on a value of one,

 $481\ 00:22:00.810 \longrightarrow 00:22:02.500$ only when there's the orange subtype,

 $482\ 00:22:02.500 \longrightarrow 00:22:04.940$ and zero in the blue subtype, or vice versa.

 $483\ 00:22:04.940 \longrightarrow 00:22:06.670$ And we wanted to see this pattern replicated

484 00:22:06.670 --> 00:22:09.680 across patients in studies, but we see obviously

 $485\ 00:22:09.680 \longrightarrow 00:22:11.840$ that that's not the case.

486 00:22:11.840 --> 00:22:14.650 So the question now that we've sort of introduced,

 $487\ 00:22:14.650 \longrightarrow 00:22:16.530$ or proposed is this sort of approach to simplify $488\ 00:22:16.530 \longrightarrow 00:22:18.520$ data merging in normalization.

 $489\ 00:22:18.520 \longrightarrow 00:22:20.020$ The question now that we're sort of dealing

 $490\ 00:22:20.020 -> 00:22:22.066$ with is well, how do we actually now find

491 00:22:22.066 --> 00:22:25.830 features that are consistent across different studies

492 00:22:25.830 --> 00:22:28.560 in their relationship with outcome, and also estimate

 $493\ 00:22:28.560 \dashrightarrow 00:22:31.793$ their study-level effect, and then use them for prediction?

494 00:22:32.860 --> 00:22:35.408 So that leads us to the second part of our paper,

49500:22:35.408 --> 00:22:39.227 where we developed a model to help select

 $496\ 00:22:39.227 \longrightarrow 00:22:42.027$ these particular study-consistent features

497 00:22:42.027 --> 00:22:47.027 while accounting for study-level heterogeneity.

498 $00{:}22{:}47.100 \dashrightarrow 00{:}22{:}49.410$ So to sort of illustrate the idea behind this,

499 00:22:49.410 \rightarrow 00:22:51.700 let's just start with a simple simulation

500 00:22:51.700 --> 00:22:54.130 where we're not doing any normalization,

501 00:22:54.130 --> 00:22:56.310 we're not worrying about resuming, everything's fine

 $502\ 00:22:56.310 \longrightarrow 00:22:58.730$ in terms of the expression values,

 $503\ 00:22:58.730 \longrightarrow 00:23:00.170$ and we're not doing any selection,

 $504\ 00:23:00.170 \longrightarrow 00:23:02.900$ no TSP transmission either.

 $505\ 00:23:02.900 \longrightarrow 00:23:04.760$ So we're going to assimilate data pertaining

 $506\ 00:23:04.760 \longrightarrow 00:23:06.380$ to two, let's say, known biomarkers

507 00:23:06.380 $\rightarrow 00:23:08.550$ that are associated with binary subtype.

508 00:23:08.550 --> 00:23:10.607 We're going to generate K datasets,

 $509\ 00:23:10.607 \longrightarrow 00:23:12.200$ and we're going to try three different strategies $510\ 00:23:12.200 \longrightarrow 00:23:14.690$ for learning a prediction model two to these data sets.

511 00:23:14.690 --> 00:23:18.070 And at the end, we're going to validate each of those models

512 00:23:18.070 --> 00:23:18.903 on an externally-generated data set

513 00:23:18.903 $\rightarrow 00:23:21.610$ to compare their prediction performance.

514 00:23:21.610 --> 00:23:25.390 So to do this, we're going to fit and assume for each study

515 00:23:25.390 --> 00:23:27.790 that we can fit it with a logistic regression model

516 00:23:27.790 --> 00:23:30.640 to model by our outcome with these two predictors,

517 00:23:30.640 --> 00:23:32.410 and in generating these K data sets,

 $518\ 00{:}23{:}32{.}410$ --> $00{:}23{:}34{.}940$ we're going to vary the number of with respect to K.

519 00:23:34.940 --> 00:23:37.690 So we might generate two trained data sets five or 10,

 $520\ 00{:}23{:}37.690$ --> $00{:}23{:}39.770$ and also change the total sample size of each one,

521 00:23:39.770 --> 00:23:41.830 and make sure that the sample sizes are in balanced

 $522\ 00:23:41.830 \longrightarrow 00:23:44.790$ across the different studies, and then assume

523 00:23:44.790 --> 00:23:49.510 values for the coefficients for each of these predictors

524 00:23:49.510 --> 00:23:52.750 to be these values here, and lastly, to induce some sort

 $525\ 00{:}23{:}52{.}750$ --> $00{:}23{:}55{.}787$ of heterogeneity across the different training datasets,

526 00:23:55.787 --> 00:23:59.410 we're gonna add in sort of like a random value drop

527 00:23:59.410 --> 00:24:01.910 from the normal distribution, where we're assuming

528 00:24:02.786 --> 00:24:04.610 this level of variance for this value.

 $529\ 00:24:04.610 \longrightarrow 00:24:06.660$ So basically we're just injecting heterogeneity $530\ 00:24:06.660 \longrightarrow 00:24:08.403$ into this data generation process.

531 00:24:09.310 \rightarrow 00:24:10.880 So after we generate the training studies,

 $532\ 00:24:10.880 \longrightarrow 00:24:12.940$ then we're going to apply three different ways

 $533\ 00:24:12.940 \longrightarrow 00:24:15.370$ or strategies to the training data.

534 00:24:15.370 --> 00:24:17.330 The first is the individual study approach,

535 00:24:17.330 --> 00:24:19.730 which we've talked about before, where you train

536 00:24:19.730 --> 00:24:22.390 a generalized model separately for each study. 537 00:24:22.390 --> 00:24:24.600 The second approach is where you merge the data.

538 00:24:24.600 --> 00:24:26.430 Again, we're ignoring the normalization problem here

539 00:24:26.430 --> 00:24:29.770 in simulation, obviously, and then train a single GLMM

 $540\ 00:24:29.770 \longrightarrow 00:24:31.870$ for the combined data, and then lastly,

 $541\ 00:24:31.870 \longrightarrow 00:24:33.660$ we're going to merge the data, and train

542 00:24:33.660 --> 00:24:35.120 a generalized linear mixed model,

543 00:24:35.120 --> 00:24:38.047 where we explicitly account for a random intercept,

 $544\ 00:24:38.047 \longrightarrow 00:24:40.610$ and a random slope for each predictor,

545 00:24:40.610 --> 00:24:44.500 assuming, you know, a study-level random effect.

546 00:24:44.500 --> 00:24:48.490 So after we do that, we'll generate a validation dataset

547 00:24:48.490 --> 00:24:52.224 from the same approach above, and then predict outcome

548 00:24:52.224 $\rightarrow 00:24:54.500$ in this validation dataset with respect

549 00:24:54.500 --> 00:24:57.400 to the models derived from each of these three strategies.

 $550\ 00{:}24{:}59{.}180$ --> $00{:}25{:}01{.}460$ So if we look at the individual strategy performance,

 $551\ 00{:}25{:}01{.}460 {\mbox{--}} > 00{:}25{:}03{.}820$ where we fit a GLM logistical regression model

 $552\ 00:25:03.820 \longrightarrow 00:25:06.010$ separately for each study, and then apply it

 $553\ 00:25:06.010 \longrightarrow 00:25:07.710$ to this validation data set, we can check

 $554\ 00:25:07.710 \longrightarrow 00:25:10.580$ the prediction accuracy, we can find that,

 $555\ 00:25:10.580 \longrightarrow 00:25:13.860$ due to the induced level of heterogeneity

 $556\ 00:25:13.860 \longrightarrow 00:25:15.800$ between studies in predictor effects,

 $557\ 00:25:15.800 \longrightarrow 00:25:18.060$ in one study, we do really poorly,

 $558\ 00:25:18.060 \longrightarrow 00:25:20.070$ and another study we do really well,

 $559\ 00:25:20.070 \longrightarrow 00:25:24.060$ and this variation is entirely due to variations $560\ 00:25:24.060 \longrightarrow 00:25:26.580$ in the gene subtype relationship.

561 00:25:26.580 --> 00:25:28.830 And these predictions obviously vary as a result

 $562\ 00:25:28.830 \longrightarrow 00:25:30.080$ across the different studies.

563 00:25:30.080 --> 00:25:32.440 And this will reflect a little bit of what we see 564 00:25:32.440 --> 00:25:35.030 in some of the examples that we showed earlier,

565 00:25:35.030 --> 00:25:38.003 studies that were trained on different data sets.

566 00:25:40.410 --> 00:25:42.550 And then the second approach is where we combine

567 00:25:42.550 --> 00:25:45.560 the data sets, and train a single logistical question model

 $568\ 00:25:45.560 \longrightarrow 00:25:46.430$ to predict outcome.

569 00:25:46.430 --> 00:25:48.530 And so we see what the median prediction error is better

570 00:25:48.530 --> 00:25:51.630 than most of the models here, but if we fit the GLMM,

571 00:25:51.630 --> 00:25:53.640 the median prediction (indistinct) gets better

572 00:25:53.640 $\rightarrow 00:25:55.800$ than some of the other approaches here.

 $573\ 00:25:55.800 \longrightarrow 00:25:57.890$ So this is basically just one example.

574 00:25:57.890 --> 00:26:00.120 So we did this over and over a hundred times 575 00:26:00.120 --> 00:26:02.640 for every single possible simulation condition, 576 00:26:02.640 --> 00:26:07.130 varying K, and the heterogeneity across different studies.

577 00:26:07.130 --> 00:26:09.560 And some of the things that we found was that

578 00:26:09.560 --> 00:26:12.110 the individual study approach had, as you can see,

579 00:26:12.110 --> 00:26:14.460 the worst prediction error overall,

 $580\ 00:26:14.460 \rightarrow 00:26:16.610$ combining the data improved this a little bit,

 $581\ 00:26:16.610 \longrightarrow 00:26:20.720$ but the estimates for the coefficients

 $582\ 00:26:20.720 \longrightarrow 00:26:23.210$ from the combined GLMM were still biased.

583 00:26:23.210 --> 00:26:26.720 There's supposed to be two in this extreme scenario.

584 00:26:26.720 --> 00:26:30.660 And a kind of heterogeneity with the GLMM mixed model had

 $585\ 00:26:30.660 \longrightarrow 00:26:32.460$ the best performance out of the rest,

 $586\ 00:26:32.460 \longrightarrow 00:26:35.004$ and also had the lowest bias in terms

 $587\ 00:26:35.004 \rightarrow 00:26:38.630$ of the regression coefficients as well.

 $588\ 00:26:38.630 \longrightarrow 00:26:42.150$ So this is great, but we also have a lot

 $589\ 00:26:42.150 \longrightarrow 00:26:43.888$ of potential types of pairs.

 $590\ 00:26:43.888 \longrightarrow 00:26:46.700$ We can't really estimate them all

591 00:26:46.700 --> 00:26:49.800 with a GLMM mixed model, so we need to find a way

592 00:26:49.800 --> 00:26:52.030 where we can, at least in reasonable dimension,

593 00:26:52.030 --> 00:26:54.610 figure out a way which fixed effects are non-zero,

594 00:26:54.610 --> 00:26:56.100 while accounting for, you know,

595 00:26:56.100 --> 00:26:58.850 this sort of study-level heterogeneity for each effect.

596 $00{:}27{:}00{.}460 \dashrightarrow 00{:}27{:}05{.}126$ So this led us to develop a pGLMM, which is basically

597 00:27:05.126 --> 00:27:08.310 a high-dimensional generalized intermixed model,

598 00:27:08.310 --> 00:27:10.770 where we are able to select fixed and random effects

599 00:27:10.770 --> 00:27:13.420 simultaneously using a penalization framework.

 $600\ 00{:}27{:}13.420$ --> $00{:}27{:}16.740$ So essentially here, we're assuming that all the predictors

 $601\ 00:27:16.740 \longrightarrow 00:27:18.740$ in the model, we assume a random effect,

 $602\ 00{:}27{:}19.606$ --> $00{:}27{:}23.046$ a random slope for each one, and so we were aiming to select

 $603\ 00:27:23.046 \longrightarrow 00:27:27.750$ the features that have non-zero fixed effects

60400:27:27.750 --> 00:27:29.540 in this particular approach, and indeed we're assuming

 $605\ 00{:}27{:}29{.}540$ --> $00{:}27{:}31{.}550$ these are going to be study-consistent.

606 00:27:31.550 --> 00:27:34.820 And to do this, we're going to reorganize

 $607\ 00:27:34.820 \longrightarrow 00:27:38.040$ the linear predictor from the standard GLMM,

60800:27:38.040 --> 00:27:41.110 so basically we're starting with the same general likelihood

609 00:27:41.110 --> 00:27:44.220 for, you know, the generalized mixed model.

610 00:27:44.220 --> 00:27:49.024 Here, Y is our outcome, X is our predictor,

 $611\ 00:27:49.024 \longrightarrow 00:27:53.040$ alpha is the, alpha K is the random effect

612 00:27:53.040 --> 00:27:58.040 for the case study, fi here is typically assumed to be

61300:27:58.150 --> 00:28:02.130 multi, very normal, means zero, and a covariant

61400:28:02.130 --> 00:28:05.140 on some sort of unstructured covariance matrix typically.

 $615\ 00:28:05.140 \rightarrow 00:28:08.930$ And so to sort of simplify this, we factor out

 $616\ 00:28:08.930 \longrightarrow 00:28:10.390$ the random effects covariance matrix,

 $617\ 00:28:10.390 \longrightarrow 00:28:12.110$ and incorporate into the linear predictor.

618 00:28:12.110 --> 00:28:15.950 And with some more reorganizing, now we're able to select

619 00:28:15.950 --> 00:28:20.950 the fixed effects and determine which random effects have

 $620\ 00:28:21.420 \longrightarrow 00:28:23.600$ true non-covariance, using this sort

621 00:28:23.600 --> 00:28:25.580 of joint penalization framework.

62200:28:25.580 --> 00:28:27.540 If you want more detail, you can check out the publication

62300:28:27.540 --> 00:28:31.340 that I linked above, and I also forgot to send out

 $624\ 00:28:31.340 \longrightarrow 00:28:33.010$ the link to this talk here.

625 00:28:33.010 --> 00:28:35.470 I'll do that right now, in case you want to check out

62600:28:35.470 --> 00:28:38.283 some of the publications that I'm linking in this talk.

627 00:28:40.660 --> 00:28:42.330 Okay, so how do we do this estimation?

628 00:28:42.330 --> 00:28:44.270 And we use that penalized NCM algorithm,

 $629~00{:}28{:}44{.}270$ --> $00{:}28{:}46{.}510$ where in each step we're drawing from the posterior

 $630\ 00:28:46.510 \longrightarrow 00:28:47.990$ with respect to the random effects, given

 $631\ 00:28:47.990 \longrightarrow 00:28:50.070$ the current aspects of the parameters,

 $632\ 00{:}28{:}50.070$ --> $00{:}28{:}55.070$ and the observed data, using Metropolis point of Gibbs.

633 00:28:55.180 --> 00:28:58.262 In the R packets, I'm going to talk about in a little bit,

634 00:28:58.262 --> 00:29:03.000 we update this to using a Hamiltonian Monte Carlo,

 $635\ 00:29:03.000 \longrightarrow 00:29:03.980$ but in the original version,

636 00:29:03.980 --> 00:29:06.270 we use Metropolis point of Gibbs, where we skipped

637 00:29:07.120 --> 00:29:09.360 components that had zero variance from the M-STEP.

638 00:29:09.360 --> 00:29:11.938 And then we use, in the M-step,

 $639\ 00:29:11.938 \longrightarrow 00:29:13.940$ two conditional maximization steps

 $640\ 00:29:13.940 \longrightarrow 00:29:17.110$ where we first update data, given the draws

641 00:29:17.110 --> 00:29:20.200 from the E-step, and the prior estimates for gamma here,

642 00:29:20.200 --> 00:29:23.740 and then up to gamma using a group penalty.

 $643\ 00:29:23.740 \longrightarrow 00:29:25.400$ So we use a couple of other tricks

 $644\ 00:29:25.400 \longrightarrow 00:29:27.060$ to speed up performance here.

645 00:29:27.060 --> 00:29:28.530 I won't go too much into the details there,

646 00:29:28.530 --> 00:29:31.713 but you can check out the paper for more detail on that.

 $647\ 00{:}29{:}33{.}330 \dashrightarrow 00{:}29{:}34{.}570$ But with this approach, one of the things

648 00:29:34.570 --> 00:29:36.579 that we were able to show was that we have

649 00:29:36.579 --> 00:29:39.290 similar conclusions regarding bias and prediction error,

 $650\ 00:29:39.290 \longrightarrow 00:29:41.420$ as in the simple setup we had before,

651 00:29:41.420 --> 00:29:43.390 where in this particular situation, we're simulating

652 00:29:43.390 --> 00:29:46.920 a bunch of predictors that do not have any association

 $653\ 00:29:46.920 \longrightarrow 00:29:50.760$ with outcome, either 10 to 50 extra predictors,

65400:29:50.760 --> 00:29:53.410 or there's only two that are actually truly relevant.

65500:29:54.480 --> 00:29:55.920 And so the prediction error in this model

 $656\ 00:29:55.920 \longrightarrow 00:29:58.650$ after this penalized selection process is

 $657\ 00:29:58.650 \longrightarrow 00:30:01.320$ generally the same, if not a little bit worse.

 $658\ 00:30:01.320 \longrightarrow 00:30:03.440$ And one thing that we find here is that

 $659\ 00:30:03.440 \longrightarrow 00:30:04.940$ the parameters are selected

66000:30:05.782 --> 00:30:07.570 by the individual study approach we're applying now

 $661\ 00{:}30{:}07{.}570 \dashrightarrow 00{:}30{:}09{.}960$ at penalized distribution regression model has

 $662\ 00:30:09.960 \rightarrow 00:30:12.859$ a low sensitivity to detect the true predictors,

663 00:30:12.859 --> 00:30:15.542 and a higher false positive rate in terms of selecting

 $664\ 00:30:15.542 \longrightarrow 00:30:17.210$ predictors that aren't associated

 $665\ 00:30:17.210 \longrightarrow 00:30:18.880$ with outcome and simulation.

666 00:30:18.880 --> 00:30:22.660 And what we find here also is that the approach

667 00:30:22.660 --> 00:30:26.050 that we developed had a much better sensitivity

 $668\ 00:30:26.050 \longrightarrow 00:30:27.800$ compared to other approaches for selecting

 $669\ 00:30:27.800 \longrightarrow 00:30:29.850$ the true predictors when accounting

670 00:30:29.850 --> 00:30:31.723 for study-level homogeneity,

 $671\ 00:30:31.723 \longrightarrow 00:30:33.183$ and the lower false positive rate as well.

672 00:30:36.060 --> 00:30:39.080 The example data sets that I talked about before,

673 00:30:39.080 $-\!>$ 00:30:43.160 the four ones that I showed a figure up earlier,

674 00:30:43.160 --> 00:30:45.030 we did a whole data study analysis where we trained

67500:30:45.030 --> 00:30:48.110 on three studies and held out one of the studies.

676 00:30:48.110 --> 00:30:50.970 We found that, you know, the approach that we put forward

677 00:30:50.970 --> 00:30:53.730 that put combining the data using our TSP approach,

678 00:30:53.730 --> 00:30:58.060 and then training a model using the pGLM had

 $679\ 00:30:58.060 \longrightarrow 00:31:00.100$ the lowest overall holdout study error

680 00:31:00.100 --> 00:31:02.420 compared to the approach using just

681 00:31:02.420 --> 00:31:05.800 a regular generalized linear model,

 $682\ 00{:}31{:}05{.}800$ --> $00{:}31{:}08{.}400$ and then also the individual study approach as well.

683 00:31:09.320 --> 00:31:11.739 And we also compared it to another post called

68400:31:11.739 --> 00:31:14.179 the Meta-Lasso, which we were able to adapt 68500:31:14.179 --> 00:31:15.760 to do prediction, and we didn't see that much improvement

 $686\ 00:31:15.760 \longrightarrow 00:31:17.000$ of performance as well.

687 00:31:17.000 --> 00:31:20.640 But in general, the result that we saw here was

 $688\ 00:31:20.640 \longrightarrow 00:31:23.259$ that the individual study approach had

689 00:31:23.259 --> 00:31:26.570 bad prediction error also across the different studies.

690 00:31:26.570 --> 00:31:29.060 So again, this sort of takes what we've already seen

 $691\ 00:31:29.060 \longrightarrow 00:31:31.190$ in the literature in terms of inconsistency,

 $692\ 00{:}31{:}31{.}190 \dashrightarrow 00{:}31{:}33{.}330$ in terms of the number of genes that are being selected

 $693\ 00:31:33.330 \dashrightarrow 00:31:35.140$ in each of these models, and also the variations

 $694\ 00:31:35.140 \longrightarrow 00:31:38.450$ in the prediction accuracy, this sort of reflects $695\ 00:31:38.450 \longrightarrow 00:31:41.523$ what we've been seeing in some of this prior work.

696 00:31:43.730 --> 00:31:45.663 So in order to you implement this approach 697 00:31:45.663 --> 00:31:49.070 in a more systematic way, my student and I, 698 00:31:49.070 --> 00:31:51.427 Hillary worked, put together an R package called

699 00:31:51.427 --> 00:31:53.880 The GLMMPen R Package.

700 $00:31:53.880 \rightarrow 00:31:56.050$ So this was just recently submitted

701 00:31:56.050 --> 00:31:58.960 to Journal of Statistical Software, but if you want to track

702 00:31:58.960 --> 00:32:01.610 the code, it's available on Github right here,

70300:32:01.610 --> 00:32:05.170 and we're in the process of submitting this to CRAN as well.

704 00:32:05.170 --> 00:32:07.880 This was sort of like a nice starter project that I gave

70500:32:07.880 --> 00:32:12.030 to Hillary to, you know, get her feet wet with coding,

706 00:32:12.030 --> 00:32:14.523 and she's done a really great job, you know,

 $707\ 00:32:14.523 \longrightarrow 00:32:16.280$ in terms of putting this together.

708 00:32:16.280 --> 00:32:19.163 And some of the distinct differences between this

709 $00{:}32{:}19.163 \dashrightarrow 00{:}32{:}21.360$ and what we put forth in the paper is the use

710 00:32:21.360 --> 00:32:23.994 of Hamiltonian Monte Carlo and the east app,

 $711\ 00:32:23.994 \longrightarrow 00:32:25.842$ instead of the Metropolis Gibbs.

712 00:32:25.842 --> 00:32:26.980 It's much faster, much more efficient.

 $713\ 00:32:26.980 \longrightarrow 00:32:28.674$ We also have added helper functions

714 00:32:28.674 --> 00:32:32.978 for the (indistinct) tuning parameters, and also making

715 00:32:32.978 --> 00:32:35.773 some diagnostic plots as well, after convergence.

716 00:32:36.640 --> 00:32:38.670 And we've also implemented some speed

717 00:32:38.670 --> 00:32:41.470 and memory improvements as well, to help with usability.

 $718\ 00:32:44.170 \longrightarrow 00:32:47.060$ Okay, so we talked about some issues

719 $00:32:47.060 \rightarrow 00:32:49.850$ regarding data integration, and then issues

 $720\ 00{:}32{:}49.850 \dashrightarrow 00{:}32{:}52.490$ with normalization, how that impedes, or can impede

 $721\ 00{:}32{:}52{.}490$ --> $00{:}32{:}55{.}730$ validation in future patients, and then we introduced

 $722\ 00:32:55.730 \longrightarrow 00:32:58.680$ a way to sidestep the normalization problem,

723 $00:32:58.680 \rightarrow 00:33:00.890$ using this sort of rank-based transformation,

 $724\ 00:33:00.890 \rightarrow 00:33:03.394$ and an approach to select consistent predictors

725 00:33:03.394 --> 00:33:06.970 in the presence of between-study heterogeneity.

726 00:33:06.970 --> 00:33:09.250 So next, I'm going to talk about a case study 727 00:33:09.250 --> 00:33:12.820 in pancreatic cancer, where we took a lot of these tools,

 $728\ 00{:}33{:}12.820$ --> $00{:}33{:}16.450$ and applied them to a problem that some collaboratives

729 00:33:16.450 --> 00:33:20.150 of mine were having, you know, at the cancer center at UNC.

730 00:33:20.150 --> 00:33:23.370 And to give a brief overview of pancreatic cancer,

 $731\ 00:33:23.370 \longrightarrow 00:33:25.850$ it has a really poor prognosis.

732 00:33:25.850 --> 00:33:29.870 Five-year survival is very low, you know, typically 5%.

733 00:33:29.870 --> 00:33:32.480 The median survival tends to be less than 11 months,

734 00:33:32.480 --> 00:33:35.260 and the main reason why this is the case is that

735 00:33:35.260 --> 00:33:37.280 early detection is very difficult,

 $736\ 00:33:37.280 -> 00:33:39.890$ and so when patients show up to the clinic,

737 00:33:39.890 --> 00:33:43.850 they're often
times in later stages, or gone metastatic.

738 00:33:43.850 --> 00:33:48.030 So for those reasons, it's really important to place

739 00:33:48.030 --> 00:33:51.040 patients on optimal the
rapies upfront, and choosing

740 00:33:51.040 --> 00:33:53.980 the best the
rapies, specifically for a patient, you know,

741 $00:33:53.980 \rightarrow 00:33:55.920$ when after they're diagnosed.

742 00:33:55.920 --> 00:33:58.850 So breast and colorectal cancers have

743 00:33:58.850 --> 00:34:02.350 long-established subtyping systems that are oftentimes used.

744 $00{:}34{:}02{.}350 \dashrightarrow 00{:}34{:}04{.}130$ Again, an example of a few of them in breast

745 $00:34:04.130 \rightarrow 00:34:05.770$ that have actually been approved by the FDA

746 00:34:05.770 --> 00:34:09.190 for clinical use, but there's nothing available for,

747 00:34:09.190 --> 00:34:11.480 in terms of precision medicine for pancreatic cancer,

748 00:34:11.480 $\rightarrow 00:34:14.260$ except for a couple of targeted therapies

749 00:34:14.260 --> 00:34:15.543 for specific mutations.

750 00:34:17.430 --> 00:34:19.870 So in 2015, the Yeh Lab at UNC,

751 00:34:19.870 --> 00:34:23.890 using a combination of non-negative matrix factorization

752 00:34:23.890 --> 00:34:27.480 and consensus clustering, where it was able to discover

 $753\ 00:34:27.480 \longrightarrow 00:34:29.996$ two potentially clinically applicable subtypes

754 00:34:29.996 --> 00:34:33.070 in pancreatic cancer, which they call basallike,

755 00:34:33.070 --> 00:34:37.036 the orange line here, which has a much worse survival

 $756\ 00:34:37.036 \longrightarrow 00:34:40.890$ compared to this classical subtype in blue,

757 00:34:40.890 --> 00:34:43.677 where patients seem to do a little bit better.

 $758\ 00:34:43.677 \longrightarrow 00:34:44.940$ And so with this approach, they used

759 00:34:44.940 --> 00:34:48.140 this unsupervised learning, set of learning techniques

 $760\ 00:34:48.140 \longrightarrow 00:34:51.010$ to derive these novel subtypes.

761 00:34:51.010 --> 00:34:54.010 And so when they took these subtypes and overlaid them

 $762\ 00:34:54.010 \longrightarrow 00:34:55.640$ from data from a clinical trial where they had

763 00:34:55.640 --> 00:34:57.540 treatment response information, they found that

764 00:34:57.540 --> 00:35:02.280 largely patients who with basal-like subtype tended to have

 $765\ 00:35:02.280 \longrightarrow 00:35:03.650$ tumors that did not respond

766 $00:35:03.650 \rightarrow 00:35:06.317$ to common first-line therapy, Folfirinox.

767 $00:35:06.317 \rightarrow 00:35:08.260$ Their tumors tended to grow from baseline.

768 00:35:08.260 --> 00:35:11.920 Whereas patients that were the classical subtype tended

769 00:35:11.920 --> 00:35:15.640 to respond better on average compared to the basal samples.

 $770\ 00:35:15.640$ --> 00:35:19.580 So the implications here are that if you are,

771 $00:35:19.580 \rightarrow 00:35:22.680$ subtype is basal, you should avoid Folfirinox

772 00:35:22.680 --> 00:35:25.020 at baseline entry with an alternative type drug,

773 00:35:25.020 --> 00:35:27.387 typically Gemcitabine and nab-paclitaxel Abraxane.

774 00:35:27.387 --> 00:35:28.740 And then for classical patients,

 $775\ 00:35:28.740 \longrightarrow 00:35:30.290$ they should receive Folfirinox.

776 00:35:32.114 --> 00:35:34.130 But the problem here is that subtyping clearly is

777 00:35:34.130 $\rightarrow 00:35:35.540$ an unsupervised learning approach, right?

 $778\ 00:35:35.540 \longrightarrow 00:35:36.750$ It's not a prediction tool.

779 $00:35:36.750 \dashrightarrow 00:35:41.750$ So it's, this approach is quite limited if it,

 $780\ 00:35:42.240 \longrightarrow 00:35:44.970$ when you have to do, assign a subtype

781 00:35:44.970 --> 00:35:47.710 in a small number of patients, it just doesn't work.

782 $00:35:47.710 \rightarrow 00:35:49.610$ So what some people have done in the past,

 $783\ 00{:}35{:}49.610$ --> $00{:}35{:}52.220$ so they simply take new patients, and recluster them

784 00:35:52.220 --> 00:35:54.570 with existing, their existing training samples. 785 00:35:54.570 --> 00:35:58.140 The problem with that is that the subtype assignments

786 00:35:58.140 --> 00:36:00.100 for those original training samples might change

 $787\ 00:36:00.100 \longrightarrow 00:36:01.110$ when they recluster it.

788 00:36:01.110 -> 00:36:02.660 So there's not a stable, it's not really

 $789\ 00:36:02.660 \longrightarrow 00:36:04.930$ a stable approach to really do this.

790 00:36:04.930 --> 00:36:07.938 So the goal here was to leverage the existing training data

791 00:36:07.938 --> 00:36:11.517 that's available to the lab, which come

792 00:36:11.517 --> 00:36:14.855 from different platforms to come up with an approach,

793 00:36:14.855 --> 00:36:17.677 a classifier to predict subtype, given

794 00:36:17.677 --> 00:36:19.930 new subtypes information, genomic,

795 00:36:19.930 --> 00:36:23.394 a new patient's genomic data, to get subtype,

796 $00:36:23.394 \rightarrow 00:36:24.890$ a predicted subtype for that individual.

797 00:36:24.890 --> 00:36:28.410 So of course, in that scenario, we also want to make sure

798 00:36:28.410 --> 00:36:30.670 that that process is simplified, and that we make

799 00:36:30.670 \rightarrow 00:36:32.760 this prediction process as easy as possible,

 $800\ 00{:}36{:}32.760$ --> $00{:}36{:}36.157$ in the face of all these issues we talked about regarding

801 00:36:36.157 --> 00:36:39.780 normalization and the training data to each other,

80200:36:39.780 --> 00:36:42.440 and also normalization of the new patient data

 $803\ 00:36:42.440 \longrightarrow 00:36:43.940$ to the existing training data.

80400:36:45.260 --> 00:36:48.820 So using some of the techniques that we just talked about,

 $805\ 00{:}36{:}48.820$ --> $00{:}36{:}50.760$ we came up with a classifier that we call PurIST,

 $806\ 00:36:50.760 --> 00:36:53.430$ which was published in the CCR last year,

 $807\ 00:36:53.430 \longrightarrow 00:36:56.270$ where essentially we were able to do that.

80800:36:56.270 --> 00:36:59.170 We take in the genomic data for a previous patient,

 $809\ 00:36:59.170$ -> 00:37:04.170 and able to predict subtype based off of that,

 $810\ 00:37:04.180 \longrightarrow 00:37:05.800$ the train model that we developed.

811 00:37:05.800 --> 00:37:08.754 And in this particular paper, we had nine data sets

812 00:37:08.754 --> 00:37:10.750 that we curated from the literature, three of which

 $813\ 00:37:10.750 \longrightarrow 00:37:12.578$ that we used for training,

 $814\ 00:37:12.578 \longrightarrow 00:37:13.540$ the rest we used for validation.

815 00:37:13.540 --> 00:37:16.400 And we did consensus clustering on all of them,

 $816\ 00:37:16.400 \longrightarrow 00:37:18.110$ using the gene list that was derived

 $817\ 00:37:18.110 \longrightarrow 00:37:19.623$ from the original publication,

818 00:37:20.978 --> 00:37:22.800 where the subtypes were discovered to get labels,

 $819\ 00:37:22.800 \longrightarrow 00:37:25.180$ subject labels for each one of the subjects

 $820\ 00:37:25.180 \longrightarrow 00:37:26.820$ in each one of these studies.

821 00:37:26.820 --> 00:37:30.370 So once we had those labels from consensus clustering,

 $822\ 00:37:30.370$ --> 00:37:33.170 we then merged the data from our three largest studies,

 $823\ 00:37:33.170 \longrightarrow 00:37:34.970$ which are our training studies.

824 00:37:34.970 --> 00:37:37.340 We did some sample for filtering based on quality,

 $825\ 00{:}37{:}37{.}340$ --> $00{:}37{:}40{.}070$ and we filtered some genes based off of, you know,

 $826\ 00:37:40.070$ --> 00:37:42.440 expression levels and things like that.

827 00:37:42.440 --> 00:37:45.010 And then we applied our previous training approach

 $828\ 00{:}37{:}45.010$ --> $00{:}37{:}49.917$ to get a small subset of top scoring pairs from the data.

 $829\ 00{:}37{:}49{.}917$ --> $00{:}37{:}51{.}230$ And in this case, we have eight that we selected,

 $830\ 00:37:51.230 \longrightarrow 00:37:55.430$ each with their own study-level coefficient.

831 00:37:55.430 --> 00:37:57.580 And then for prediction, the process is very simple,

832 00:37:57.580 --> 00:38:00.300 we just check in that patient, whether gene A is greater

833 00:38:00.300 --> 00:38:02.130 than gene D for each of these pairs,

83400:38:02.130 --> 00:38:05.240 and that gives us their binary vector of ones and zeros.

 $835\ 00{:}38{:}05{.}240$ --> 00:38:08.630 We multiply that by the coefficients from the train model.

836 00:38:08.630 --> 00:38:11.460 This is basically just calculating a linear predictor

 $837\ 00:38:11.460 \longrightarrow 00:38:13.750$ from this logistic regression model.

838 00:38:13.750 --> 00:38:14.850 And then we can convert that

 $839\ 00:38:14.850 \longrightarrow 00:38:18.130$ to a predicted probability of being basal.

 $840\ 00:38:18.130 \longrightarrow 00:38:23.130$ So using this approach, we were able to select

 $841\ 00:38:23.130 \longrightarrow 00:38:25.170\ 16$ genes pertaining to eight subtypes,

 $842\ 00:38:25.170 \longrightarrow 00:38:27.210$ but we can find here that the predictions

 $843\ 00:38:27.210 \longrightarrow 00:38:30.760$ from this model tends to coincide very strongly

 $844\ 00:38:30.760 \longrightarrow 00:38:32.930$ with the labels that were collected

 $845\ 00:38:32.930 \longrightarrow 00:38:33.980$ using consensus clusters.

846 $00{:}38{:}33{.}980 \dashrightarrow 00{:}38{:}36{.}498$ So that gives us some confidence that reproducing

847 00:38:36.498 --> 00:38:41.070 in some way, you know, this, the result that we got

 $848\ 00:38:41.070 \longrightarrow 00:38:43.100$ using this clustering approach.

849 $00{:}38{:}43.100$ --> $00{:}38{:}46.100$ You can also clearly see here that as the subtype changes,

 $850\ 00{:}38{:}46{.}100$ --> $00{:}38{:}48{.}620$ that you see flips in the expression in each one

 $851\ 00:38:48.620 \longrightarrow 00:38:51.760$ of the pairs of genes that we collected

 $852\ 00:38:51.760 \longrightarrow 00:38:53.680$ in this particular study.

 $853\ 00:38:53.680 \longrightarrow 00:38:55.010$ And then when we applied this model

854 00:38:55.010 --> 00:38:58.740 to six external validation dataset, we found that it had

 $855\ 00{:}38{:}58{.}740$ --> 00:39:01.330 a very good performance in terms of recapitulating subtype,

 $856\ 00:39:01.330 \longrightarrow 00:39:03.660$ where we had a relatively good sensitivity

 $857\ 00{:}39{:}03.660$ --> $00{:}39{:}07.090$ and specificity in each case, which we owe part

858 00:39:07.090 --> 00:39:08.185 to the fact that we don't have to worry as much

85900:39:08.185 --> 00:39:13.185 about this sort of cross-study normalization training time

 $860\ 00{:}39{:}13.218$ --> $00{:}39{:}16.570$ or test time, and also the fact that we lever-aged

 $861\ 00:39:17.407 \longrightarrow 00:39:18.620$ multiple data sets when selecting

862 00:39:20.570 --> 00:39:21.690 the predictors for this model.

863 00:39:21.690 --> 00:39:23.870 And so when we looked at the predictive values

86400:39:23.870 --> 00:39:26.510 in these holdout studies, the predictive subtypes,

865 00:39:26.510 --> 00:39:29.660 we recapitulated the differences in survival

 $866\ 00:39:29.660 \longrightarrow 00:39:31.850$ that we observed in other studies as well,

 $867\ 00:39:31.850 \longrightarrow 00:39:34.354$ where basal-like patients do a lot worse

 $868\ 00:39:34.354 \longrightarrow 00:39:36.700$ compared to classical patients.

869 00:39:36.700 --> 00:39:38.690 If you want to look a little bit more at the details

 $870\ 00:39:38.690$ --> 00:39:41.100 in this paper, you can check out this link here, $871\ 00:39:41.100$ --> 00:39:43.720 and if you want to access the code that we used

 $872\ 00:39:43.720 \longrightarrow 00:39:45.460$ to make these predictions, that's available

 $873\ 00:39:45.460 \longrightarrow 00:39:48.453$ on this Github page at this link right here.

874 00:39:50.380 --> 00:39:53.310 Another thing that we were able to show is that for patients

875 00:39:53.310 --> 00:39:56.450 that had samples that are collected through different modes

 $876\ 00{:}39{:}56{.}450$ --> 00:40:00.070 of collection, whether it was bulk, FNA, FFPE,

877 00:40:00.070 --> 00:40:03.020 we found that the predictions in these patients tend to be

 $878\ 00:40:03.020 \longrightarrow 00:40:06.430$ highly consistent, and this is basically deriving

87900:40:06.430 --> 00:40:08.820 itself, again, from the simple assumption behind TSPs,

 $880\ 00{:}40{:}08.820$ --> 00:40:13.060 where the relative rank within the subject of the expression

881 00:40:13.060 --> 00:40:14.990 of these genes is predicted.

 $882\ 00:40:14.990 \longrightarrow 00:40:17.310$ So as long as that is being preserved,

883 00:40:17.310 --> 00:40:21.440 then you should be able to have the model predict well

884 00:40:21.440 --> 00:40:23.289 in different scenarios.

 $885\ 00{:}40{:}23.289 \dashrightarrow 00{:}40{:}27.630$ So when we also went through CLIA validation for this tool,

886 00:40:27.630 --> 00:40:31.154 we also confirmed 95% agreement between replicated runs

 $887\ 00{:}40{:}31.154$ --> $00{:}40{:}36.154$ in other platforms, and we also confirmed concordance

88800:40:37.950 --> 00:40:42.770 between NanoString and RNAC, also through different modes

 $889\ 00:40:42.770 \longrightarrow 00:40:43.603$ of sample collection.

890 00:40:43.603 --> 00:40:46.690 So right now this is the first clinically applicable test

89100:40:46.690 --> 00:40:50.610 for a prospect of first line treatment selection in PDAC.

892 00:40:50.610 --> 00:40:54.250 And right now we do have a study that just recently opened

893 00:40:54.250 --> 00:40:56.390 at the Medical College of Wisconsin that's using PurIST

 $894\ 00:40:56.390 \longrightarrow 00:40:58.390$ for prospect of treatment selection,

 $895\ 00{:}40{:}58.390$ --> 00:41:01.970 and we have another one opening at University of Rochester,

 $896\ 00:41:01.970 \longrightarrow 00:41:06.320$ and also at UNC soon as well.

897 00:41:06.320 --> 00:41:09.510 So this is just an example about how you can take

89800:41:09.510 --> 00:41:14.040 a problem, you know, in, from the literature, 89900:41:14.040 --> 00:41:17.570 from your collaborators, come up with a method,

900 00:41:17.570 --> 00:41:22.150 and some theory behind it, and really be able to come up

 $901\ 00:41:22.150 \longrightarrow 00:41:24.310$ with a good solution that is robust,

 $902\ 00{:}41{:}24{.}310$ --> $00{:}41{:}27{.}440$ and that can really help your collaborative

 $903\ 00:41:27.440 \longrightarrow 00:41:29.763$ at your institution and elsewhere.

 $904\ 00:41:31.850 \longrightarrow 00:41:33.510$ Okay, so that was the case study.

905 00:41:33.510 --> 00:41:34.560 To talk about some current work

 $906\ 00:41:34.560 \longrightarrow 00:41:36.150$ that we're doing just briefly.

907 00:41:36.150 --> 00:41:39.350 So we wanted to think about how we can also scale up the,

90800:41:39.350 --> 00:41:42.200 this particular framework that we developed for the pGLMM,

909 00:41:42.200 --> 00:41:44.190 and one idea that we're pursuing right now

 $910\ 00:41:44.190 \longrightarrow 00:41:46.400$ with my student Hillary, is that we're thinking

911 00:41:47.773 --> 00:41:49.751 about using, borrowing ideas from factor analysis

912 00:41:49.751 --> 00:41:52.570 to decompose, do a deep, deterministic decomposition

913 00:41:52.570 --> 00:41:56.370 of the random effects to a lower dimensional space,

 $914\ 00:41:56.370 \longrightarrow 00:41:59.690$ where essentially, we can essentially map

915 00:41:59.690 --> 00:42:02.780 between the lower dimensional space (indistinct) factors,

916 00:42:02.780 --> 00:42:05.220 which is r-dimensional, to this higher dimensional space,

917 00:42:05.220 --> 00:42:10.220 using some by matrix B, which is q by r,

918 00:42:11.920 --> 00:42:16.050 and essentially in doing so, this reduces the dimension

919 00:42:16.050 --> 00:42:19.243 of the integral in the Monte Carlo EM algorithm.

920 00:42:20.253 --> 00:42:21.730 So rather than having to do approximate integral

921 00:42:21.730 \rightarrow 00:42:23.560 and q dimensions, which can be difficult,

922 00:42:23.560 --> 00:42:26.870 you can work in a much lower space in terms of integral,

923 00:42:26.870 --> 00:42:28.710 and then have this additional problem

924 00:42:28.710 --> 00:42:30.590 of trying to estimate this matrix,

 $925\ 00:42:30.590 \longrightarrow 00:42:33.170$ and not back to the original dimension cube.

926 00:42:33.170 --> 00:42:34.840 So that's something that we're just starting to work on

927 00:42:34.840 --> 00:42:38.550 right now, and another thing that we're starting to work on

928 00:42:38.550 --> 00:42:41.229 is the idea of trying to extend some of the work

929 00:42:41.229 --> 00:42:42.860 in variational autoencoders

 $930\ 00:42:42.860 \longrightarrow 00:42:45.200$ that my student David is working on now.

 $932\ 00{:}42{:}48.253 \dashrightarrow 00{:}42{:}51.350$ when trying to train these sort of deep learning models,

933 00:42:51.350 --> 00:42:55.170 the VAEs unsupervised learning model's oftentimes used

 $934\ 00:42:55.170 \longrightarrow 00:42:56.010$ for dimensional reduction.

935 00:42:56.010 --> 00:42:57.020 You might've heard of it

 $936\ 00:42:57.020 \longrightarrow 00:43:01.330$ in single cells sequencing applications.

937 00:43:01.330 --> 00:43:02.850 But the question that we wanted to address is, well,

938 00:43:02.850 --> 00:43:04.990 what if you have missing data, you know,

939 00:43:04.990 --> 00:43:08.197 in your input features X, which might be (indistinct)?

940 $00:43:09.529 \rightarrow 00:43:14.260$ So essentially we were able to develop input.

941 00:43:14.260 --> 00:43:17.280 So we have a pre-print up right now, it's the code,

942 00:43:17.280 --> 00:43:20.240 and we're looking to extend this, where essentially,

943 00:43:20.240 --> 00:43:22.680 rather than worrying about this latent space Z,

944 00:43:22.680 $\rightarrow 00:43:24.640$ which we're assuming that that encodes a lot 945 00:43:24.640 $\rightarrow 00:43:26.910$ of the information in the original data,

946 00:43:26.910 \rightarrow 00:43:28.910 we replaced that with learning the posterior

947 00:43:28.910 \rightarrow 00:43:31.550 of the random effect, given the observed data.

948 00:43:31.550 --> 00:43:34.260 And then in the second portion here, we replaced

949 00:43:34.260 --> 00:43:38.820 this generative model with the general model of y given X

 $950\ 00:43:38.820 \longrightarrow 00:43:40.680$ in the random effects.

951 00:43:40.680 \rightarrow 00:43:42.880 So that's another avenue that can allow us

 $952\ 00:43:42.880 \longrightarrow 00:43:44.650$ to hopefully account for non-linearity,

 $953\ 00:43:44.650 \longrightarrow 00:43:47.100$ and arbitrator action between features as well.

954 00:43:47.100 --> 00:43:49.179 And also it might be an easier way to scale up

 $955\ 00:43:49.179 \longrightarrow 00:43:52.570$ some of the analysis we've done too,

 $956\ 00:43:52.570 \longrightarrow 00:43:55.330$ which I've already mentioned.

957 00:43:55.330 --> 00:43:58.361 Okay, so in terms of some concluding thoughts,

958 00:43:58.361 --> 00:44:02.762 I talked a lot about how the original subtypes were derived

959 00:44:02.762 --> 00:44:05.930 for this pancreatic cancer case study using NMF

 $960\ 00:44:05.930 \longrightarrow 00:44:09.310$ and consensus clustering to get two subtypes.

961 00:44:09.310 --> 00:44:12.310 But there were also other groups that are published,

962 00:44:12.310 --> 00:44:15.540 subtyping systems, that in one, they found

 $963\ 00{:}44{:}15{.}540$ --> $00{:}44{:}19{.}150$ three subtypes, and in another one they found four subtypes.

964 00:44:19.150 --> 00:44:22.042 So the question is, well, you know, well,

 $965\ 00:44:22.042 \longrightarrow 00:44:23.270$ which one do we use?

966 00:44:23.270 --> 00:44:26.130 Again, this is also confusing for practitioners 967 00:44:26.130 --> 00:44:28.950 about which approach might be more meaningful

 $968\ 00:44:28.950 \longrightarrow 00:44:30.110$ in the clinical setting.

969 00:44:30.110 --> 00:44:31.840 And each of these approaches were also derived 970 00:44:31.840 --> 00:44:35.480 using NMF and consensus clustering, and they were done

971 00:44:35.480 --> 00:44:37.540 separately on different patient cohorts

972 00:44:37.540 --> 00:44:39.140 at different institutions.

973 00:44:39.140 --> 00:44:41.460 So you can see that this is another reflection

974 00:44:41.460 --> 00:44:44.930 of heterogeneity in single-study learning,

975 00:44:44.930 --> 00:44:48.680 and how we can get these different or discrepant results

97600:44:48.680 --> 00:44:52.170 from applying the same technique to 200 genus datasets

977 00:44:52.170 \rightarrow 00:44:54.400 that were generated at different places.

978 00:44:54.400 --> 00:44:57.000 So of course this creates another problem, you know,

 $979\ 00:44:57.000 \longrightarrow 00:44:59.730$ who's right, which approach do we use?

980 00:44:59.730 --> 00:45:03.350 And it's kind of like a circular argument here.

981 00:45:03.350 --> 00:45:06.870 So in the paper that I mentioned before with PurIST,

 $982\ 00:45:06.870 \longrightarrow 00:45:09.260$ another thing that we did is we overlaid

 $983\ 00:45:09.260 \longrightarrow 00:45:11.839$ the others subtype system calls

984 00:45:11.839 --> 00:45:14.790 with the observed clinical outcomes

985 00:45:14.790 --> 00:45:16.650 for the studies that we collected.

 $986\ 00:45:16.650 \longrightarrow 00:45:19.120$ And one of the things that we found was that,

987 00:45:19.120 $\operatorname{-->}$ 00:45:21.920 and these other subtyping systems,

988 00:45:21.920 --> 00:45:23.840 each of them also had something,

989 00:45:23.840 --> 00:45:26.990 something that was very similar to the basal-like subtype,

990 00:45:26.990 --> 00:45:29.860 and for the remaining subtypes, they had survival

991 00:45:29.860 $\rightarrow 00:45:32.650$ that was similar to the classical subtype.

992 00:45:32.650 --> 00:45:35.210 So one of the arguments that we made was that,

 $993\ 00:45:35.210 \longrightarrow 00:45:36.813$ well, if the clinical outcomes are the same

 $994\ 00:45:36.813 \longrightarrow 00:45:39.570$ for the other subtypes, you know,

 $995\ 00:45:39.570 \longrightarrow 00:45:41.500$ are they exactly right necessary

 $996\ 00:45:41.500 \longrightarrow 00:45:43.250$ for clinical decision-making?

997 00:45:43.250 $\rightarrow 00:45:45.540$ That was one argument that we put forth.

998 00:45:45.540 --> 00:45:48.420 And when we looked at the response data, again,

999 00:45:48.420 --> 00:45:51.410 we saw that one of the subtypes in the other approaches

1000 00:45:51.410 --> 00:45:56.020 also overlapped the basal-like subtype in terms of response.

 $1001\ 00:45:56.020 \longrightarrow 00:45:57.430$ And then for the remaining subtypes,

 $1002\ 00{:}45{:}57{.}430$ --> $00{:}46{:}00{.}900$ they were just kind of randomly dispersed at the other end,

 $1003\ 00{:}46{:}00{.}900$ --> $00{:}46{:}05{.}280$ you know, of the spectrum here in terms of tumor present,

 $1004\ 00:46:05.280 \longrightarrow 00:46:06.730$ tumor change after treatment.

 $1005 \ 00:46:06.730 \longrightarrow 00:46:09.310$ So the takeaway here is that heterogeneity

 $1006\ 00{:}46{:}09{.}310$ --> $00{:}46{:}13{.}660$ between studies also impacts tasks in unsupervised learning,

 $1007\ 00:46:13.660 \longrightarrow 00:46:16.330$ like the NMF+ consensus clustering approach

 $1008 \ 00:46:16.330 \longrightarrow 00:46:18.000$ to discover subtypes.

1009 00:46:18.000 --> 00:46:20.770 And what this also does is, as you can imagine,

1010 00:46:20.770 --> 00:46:23.690 this injects a lot of confusion into the literature,

1011 00:46:23.690 --> 00:46:27.119 and can also slow down the process of translating

 $1012 \ 00:46:27.119 \longrightarrow 00:46:29.980$ some of these approaches to the clinic.

 $1013 \ 00:46:29.980 \longrightarrow 00:46:31.960$ So this also underlies the need

1014 00:46:31.960 --> 00:46:35.280 for replicable cross-study sub discovery approaches,

1015 00:46:35.280 --> 00:46:40.280 for replicable approaches for unsupervised learning.

101600:46:40.580 --> 00:46:42.980 That's something that, you know, something that we might,

 $1017 \ 00:46:42.980 \longrightarrow 00:46:45.630$ we hope to be working on in the future,

 $1018 \ 00:46:45.630 \longrightarrow 00:46:47.623$ and we hope to see more work on as well.

 $1019\ 00{:}46{:}48.660 \dashrightarrow 00{:}46{:}52.640$ So to summarize the, one of the major points

1020 00:46:52.640 --> 00:46:55.470 of this talk was to introduce and discuss, you know,

1021 00:46:55.470 --> 00:46:58.100 replicability issues in genomic prediction models,

1022 00:46:58.100 --> 00:47:01.080 supervised learning, that stems from technical,

 $1023 \ 00:47:01.080 \longrightarrow 00:47:03.420$ and also non-technical sources.

1024 00:47:03.420 --> 00:47:06.770 We also introduced a new approach to facilitate

1025 00:47:06.770 --> 00:47:08.840 data integration and multistory learning

1026 00:47:08.840 --> 00:47:12.426 in a way that captures between-study heterogeneity,

1027 00:47:12.426 --> 00:47:15.400 and showed how this can be used for the prediction

1028 00:47:15.400 --> 00:47:20.360 of subtype for pancreatic cancer, and also introduced

1029 00:47:20.360 --> 00:47:22.522 some scalable methods and future direction

1030 00:47:22.522 --> 00:47:24.933 in replicable subtype discovery.

1031 00:47:26.350 --> 00:47:28.180 So that's it for me.

1032 00:47:28.180 --> 00:47:30.140 I just want to thank some of my faculty crowd,

1033 00:47:30.140 --> 00:47:33.050 collaboratives, Quefeng Li, Junier Oliva

1034 00:47:33.050 --> 00:47:36.750 from UNC computer science, Jen Jen Yeah

1035 00:47:36.750 --> 00:47:40.010 from surgical oncology at Lineberger,

1036 00:47:40.010 --> 00:47:42.550 Joe Ibrahim as well, UNC biostatistics,

1037 00:47:42.550 --> 00:47:45.100 and also my students, Hilary, who's done a lot of work

1038 00:47:45.100 --> 00:47:47.821 in this area, and also David Lim, who's doing

1039 00:47:47.821 --> 00:47:49.840 some of the deep learning work in our group.

 $1040\ 00:47:49.840 \longrightarrow 00:47:51.283$ And that's it, thank you.

1041 00:47:57.800 --> 00:47:59.290 <v Robert>So does any
body here have</v>

 $1042 \ 00:47:59.290 \longrightarrow 00:48:01.830$ any questions for the professor?

 $1043\ 00{:}48{:}09.063$ --> $00{:}48{:}14.063$ Or any body on the, on Zoom, any questions you want to ask?

1044 00:48:25.900 --> 00:48:27.383 <v ->It looks like I'm off the hook.</v>

1045 00:48:28.750 --> 00:48:30.240 <v Robert>All right, well, thank you so much.</v>

1046 00:48:30.240 --> 00:48:31.813 Really appreciated your talk.

 $1047 \ 00:48:33.390 \longrightarrow 00:48:34.490$ Have a good afternoon.

1048 00:48:36.030 --> 00:48:37.880 <v ->All right, thank you for having me.</v>