## WEBVTT

- $1.00:00:00.000 \longrightarrow 00:00:03.000$  (people chattering)
- 2 00:00:10.312 --> 00:00:12.012 <v Man>(indistinct) Biostatistics</v>
- $3\ 00:00:12.930 --> 00:00:15.107$  at the University of Minnesota.
- 4 00:00:15.107 --> 00:00:18.940 And he's currently attending or an associate professor
- 5~00:00:18.940 --> 00:00:23.299 at the University of the Texas Dell Medical School.
- $6~00:00:23.299 \longrightarrow 00:00:27.160~\mathrm{Dr}$ . Hobbes is a library recognized as an expert
- $7~00:00:27.160 \longrightarrow 00:00:31.160$  in clinical oncology and research (indistinct).
- 8 00:00:31.160 --> 00:00:33.743 Among his many accomplishments,
- 9 00:00:35.309 --> 00:00:39.392 in 2017, Dr. Hobbes (indistinct)
- 10 00:00:40.400 --> 00:00:44.039 of The National Cancer Institute, clinical trial
- $11\ 00:00:44.039 \dashrightarrow 00:00:49.039$  (in distinct) a national consensus recommendations
- $12\ 00:00:49.680 \longrightarrow 00:00:50.513$  or (indistinct).
- 13 00:00:54.840 --> 00:00:59.840 In 2019, Dr. Hobbes (indistinct).
- $14\ 00:01:08.424 \longrightarrow 00:01:13.424\ {\rm In}\ 2021,\ {\rm Dr.\ Hobbes}\ ({\rm indistinct})$
- 15 00:01:46.510 --> 00:01:47.650 <-> Brian Hobbes>Thank you, thank you very much,</r>
- $16\ 00:01:47.650 \longrightarrow 00:01:50.893$  and for that long and generous introduction.
- $17~00{:}01{:}53.580 \dashrightarrow 00{:}01{:}55.410$  I'm excited to give this talk today.
- 18 00:01:55.410 --> 00:01:57.740 I wish I could visit in-person.
- $19\ 00:01:57.740 \longrightarrow 00:02:00.260\ I$  was fortunate to have that opportunity a few years ago,
- $20\ 00:02:00.260 \longrightarrow 00:02:02.483$  so thank you for inviting me back.
- $21\ 00:02:04.810 --> 00:02:09.530$  Okay, so I got tired of giving talks
- 22 00:02:09.530 --> 00:02:12.720 that were very technical and very specific
- $23\ 00:02:12.720 \longrightarrow 00:02:14.133$  to a specific problem.
- $24\ 00:02:15.045$  --> 00:02:17.697 Because, if you don't have an understanding of the problem,
- $25\ 00:02:17.697 --> 00:02:19.450$  you don't have an understand that the biomarkers,

- $26\ 00:02:19.450 \longrightarrow 00:02:22.600$  or you don't work in a particular area of methodology,
- $27\ 00:02:22.600 \dashrightarrow 00:02:25.783$  I think it becomes very, you know, what do you wanna say?
- 28 00:02:25.783 --> 00:02:28.640 I think people lose interest pretty quickly.
- $29~00:02:28.640 \longrightarrow 00:02:31.450$  And so I decided to start giving talks
- $30\ 00:02:31.450 \longrightarrow 00:02:34.350$  that had to do overview a subject
- $31\ 00:02:34.350 \longrightarrow 00:02:37.000$  that I think is very relevant in the field right now.
- $32\ 00:02:37.890 \longrightarrow 00:02:42.770$  So recently, I'm an external advisor on a grant
- $33\ 00:02:42.770 --> 00:02:45.690$  that Genentech has got from the FDA
- $34\ 00:02:45.690$  --> 00:02:48.903 for developing clinical trials that use real-world data.
- 35 00:02:49.920 --> 00:02:52.130 I've worked with Flatiron in the last few years,
- $36\ 00:02:52.130 \longrightarrow 00:02:54.380$  as well as the CancerLinQ.
- 37 00:02:54.380 --> 00:02:56.130 And I've been watching this field,
- $38\ 00:02:56.130 --> 00:02:58.070$  sort of the discussions in this field
- $39\ 00:02:58.070 \longrightarrow 00:02:59.250$  about real-world evidence,
- $40\ 00:02:59.250 \longrightarrow 00:03:00.290$  and where does it fit-in?
- 41 00:03:00.290 --> 00:03:03.470 and specifically in the context of cancer drug development.
- $42\ 00:03:03.470 \longrightarrow 00:03:05.600$  So I decided to talk about that today.
- $43\ 00:03:05.600 \longrightarrow 00:03:09.590$  So yeah, I think this is what I'm gonna do.
- $44\ 00:03:09.590 \longrightarrow 00:03:11.330$  So does real-world evidence
- $45\ 00:03:11.330 \longrightarrow 00:03:13.280$  have a role in cancer drug development?
- 46 00:03:14.160 --> 00:03:14.993 So if you'll see,
- $47\ 00{:}03{:}14.993 \dashrightarrow 00{:}03{:}18.010$  there's a question mark at the end of the statement.
- 48 00:03:18.010 --> 00:03:20.450 So I'm gonna talk about this,
- $49~00:03:20.450 \dashrightarrow 00:03:22.660$  and I'm gonna give you the perspective that I have,
- $50\ 00:03:22.660 --> 00:03:25.033$  which comes from a methodologist
- $51\ 00:03:25.033 \longrightarrow 00:03:26.560$  that really wants real-world evidence

- 52 00:03:26.560 --> 00:03:28.470 to have a role in cancer drug development,
- $53\ 00:03:28.470 --> 00:03:31.543$  because, databases are growing.
- $54\ 00:03:32.740 \longrightarrow 00:03:34.370$  Data science becomes more relevant
- $55\ 00:03:34.370 \longrightarrow 00:03:36.800$  if those databases are useful.
- $56\ 00:03:36.800 \longrightarrow 00:03:38.150$  We all want to write algorithms
- 57 00:03:38.150 --> 00:03:41.470 and do, you know, causal inference on database.
- $58\ 00:03:41.470 \longrightarrow 00:03:44.300$  We want to unlock those databases with our intelligence,
- $59\ 00:03:44.300 \longrightarrow 00:03:45.400$  for drug development.
- 60 00:03:45.400 --> 00:03:47.450 Drug development is incredibly expensive.
- $61\ 00:03:48.360 \longrightarrow 00:03:51.610$  Patients need access to the rapies
- $62\ 00:03:51.610 --> 00:03:52.730$  that are gonna save their lives.
- 63~00:03:52.730 --> 00:03:55.680 We have refractory patients enrolling in clinical trials.
- $64~00:03:55.680 \longrightarrow 00:03:58.050$  There's probably not enough clinical trials.
- $65\ 00:03:58.050 \longrightarrow 00:04:00.210$  And we do have advances in biology
- $66\ 00:04:00.210 \dashrightarrow 00:04:05.190$  that have manifest themselves in precision the rapeutics.
- $67\ 00:04:05.190 \longrightarrow 00:04:07.830$  So we want all of this to work together.
- $68\ 00:04:07.830 \longrightarrow 00:04:09.400$  We want this to be true.
- $69\ 00:04:09.400 \longrightarrow 00:04:10.540$  On the other hand,
- $70~00{:}04{:}10.540 \dashrightarrow 00{:}04{:}13.530$  I have designed hundreds of clinical trials
- 71 00:04:13.530 --> 00:04:14.400 and I continue to,
- 72 00:04:14.400 --> 00:04:15.840 most of my collaborations continue
- $73\ 00:04:15.840 \longrightarrow 00:04:17.533$  with MD Anderson in this space.
- $74~00:04:18.580 \dashrightarrow 00:04:22.280$  I've worked with oncologists for over a decade now.
- $75~00:04:22.280 \longrightarrow 00:04:24.840$  I've worked with translational researchers in oncology,
- $76~00:04:24.840 \longrightarrow 00:04:28.840$  and I see the issues that are presented.
- 77 00:04:28.840 --> 00:04:31.400 I mean, maybe I should say the challenges,
- $78\ 00:04:31.400 \longrightarrow 00:04:32.513$  the challenges that we confront

- $79\ 00:04:32.513 \longrightarrow 00:04:34.500$  when we think about this space.
- 80 00:04:34.500 --> 00:04:37.550 So I'm gonna talk about this.
- 81 00:04:37.550 --> 00:04:41.730 And I think if I was the 30-year-old version of myself,
- 82 00:04:41.730 --> 00:04:43.960 Brian Hobbs, the 30-year-old,
- $83~00{:}04{:}43.960 \dashrightarrow 00{:}04{:}46.030$  there would not be a question mark here.
- $84\ 00:04:46.030$  -->  $00:04:48.520\ \mathrm{I}$  would be saying we can use real-world evidence,
- $85\ 00:04:48.520 \longrightarrow 00:04:50.190$  and this is how.
- $86~00{:}04{:}50.190 \dashrightarrow 00{:}04{:}54.280$  But now that I'm 40 years old, there's a question mark.
- 87 00:04:54.280 --> 00:04:56.840 And I think that, you know okay,
- 88 00:04:56.840 --> 00:04:58.650 so when you've seen other talks about this,
- 89 00:04:58.650 --> 00:05:00.870 I don't know if you're experiencing the same thing I have,
- $90\ 00:05:00.870 \dashrightarrow 00:05:05.060$  but, I've seen several talks at seminars, conferences,
- $91\ 00:05:05.060 \longrightarrow 00:05:09.400$  where people are presenting very specific cases.
- 92 00:05:09.400 --> 00:05:12.000 Specific cases where they could use real-world evidence
- 93  $00:05:12.000 \longrightarrow 00:05:12.833$  and it made sense,
- $94\ 00:05:12.833 --> 00:05:16.120$  or it was the only thing that could be done in that context.
- $95\ 00:05:16.120 \longrightarrow 00:05:17.950$  So I've seen a lot of talks like that.
- 96 00:05:17.950 --> 00:05:20.210 I'm gonna take this from the other perspective,
- $97\ 00:05:20.210 --> 00:05:22.410$  I'm gonna talk about what's going on in oncology right now.
- 98 00:05:22.410 --> 00:05:24.090 What are the most important developments happening
- 99  $00:05:24.090 \longrightarrow 00:05:24.923$  in oncology?
- 100 00:05:24.923 --> 00:05:26.660 And then I'm gonna ask the question,
- $101\ 00{:}05{:}26.660 \dashrightarrow 00{:}05{:}30.500$  can we use real-world evidence to help augment

- $102\ 00:05:30.500 \longrightarrow 00:05:33.000$  our trial designs and drug development in general?
- $103\ 00:05:34.600 \longrightarrow 00:05:37.290$  So to begin with what is real-world evidence?
- $104\ 00:05:37.290 \longrightarrow 00:05:40.360$  So there's different definitions of this.
- $105\ 00:05:40.360 \longrightarrow 00:05:43.193$  It tends to be a very broad definition.
- $106\ 00{:}05{:}44.950 \dashrightarrow 00{:}05{:}49.070$  That's, you know, that different people use this.
- $107\ 00:05:49.070 --> 00:05:50.980$  I've taken this diagram from the CancerLinQ,
- $108\ 00{:}05{:}50.980 {\:{\mbox{--}}}{\:{\mbox{--}}} 00{:}05{:}53.580$  which is a nonprofit organization that works
- $109\ 00{:}05{:}53.580 \dashrightarrow 00{:}05{:}57.370$  with the American Society of Clinical Oncology.
- $110\ 00:05:57.370 \longrightarrow 00:06:00.300$  They are massing and organizing a large database,
- 111 00:06:00.300 --> 00:06:02.540 I collaborate with Elizabeth Garrett-Mayer at CancerLinQ
- $112\ 00:06:02.540 \longrightarrow 00:06:05.390$  who's at ASCO, who's great.
- $113\ 00:06:05.390$  --> 00:06:08.350 Who's have a PhD statistician working on this.
- 114 00:06:08.350 --> 00:06:09.946 So this diagram, you know,
- $115\ 00:06:09.946 \longrightarrow 00:06:12.640$  what we often think about as real-world evidence,
- $116\ 00:06:12.640 --> 00:06:15.190$  we think about as the electronic medical health records
- $117\ 00:06:15.190 --> 00:06:19.340$  that are in sort of community hospital systems, right?
- $118\ 00{:}06{:}19.340 \dashrightarrow 00{:}06{:}22.450$  We think about data that's acquired from routine care
- 119 00:06:22.450 --> 00:06:26.860 or from claims that's not on patients
- $120\ 00:06:26.860 \longrightarrow 00:06:28.800$  that are in a clinical study.
- $121\ 00{:}06{:}28.800 \to 00{:}06{:}30.750$  We tend to think about that as real-world evidence.
- $122\ 00:06:30.750 --> 00:06:34.800$  And so CancerLinQ says Real-World Evidence has a capability,
- $123\ 00{:}06{:}34.800 \dashrightarrow 00{:}06{:}37.840$  data tools, processes, organization, underpinning functions

- $124\ 00:06:37.840 \longrightarrow 00:06:39.010$  to drive business intelligence.
- $125~00{:}06{:}39.010 \dashrightarrow 00{:}06{:}41.960$  So that's kind of, you know, very broad.
- $126\ 00:06:41.960 \longrightarrow 00:06:44.850$  They also tell us that there's other things
- $127~00{:}06{:}44.850 \dashrightarrow 00{:}06{:}46.340$  that should count as real-world evidence
- $128\ 00:06:46.340 \longrightarrow 00:06:48.040$  beyond the EMR data.
- 129 00:06:48.040 --> 00:06:49.940 Okay, observational data
- $130\ 00:06:49.940$  --> 00:06:52.300 as well as historical randomized controlled data.
- $131\ 00:06:52.300 \longrightarrow 00:06:53.820$  Okay, that makes sense.
- $132\ 00{:}06{:}53.820 --> 00{:}06{:}57.490$  Pharmacy data, mortality registries, hospital visits,
- 133 00:06:57.490 --> 00:07:02.320 lab values, claim databases, social media,
- $134\ 00:07:02.320 \longrightarrow 00:07:04.620$  they put on this diagram as well.
- 135 00:07:04.620 --> 00:07:07.730 So you know maybe, right?
- $136\ 00:07:07.730 --> 00:07:10.790$  But I think that we're at a place right now
- 137 00:07:10.790 --> 00:07:12.420 where people are excited about using
- 138 00:07:12.420 --> 00:07:14.620 these sources of information in research,
- $139\ 00:07:14.620 --> 00:07:17.060$  but somebody really needs to develop a framework
- $140\ 00:07:17.060 \longrightarrow 00:07:18.030$  for each of these.
- 141 00:07:18.030 --> 00:07:19.730 There's not a single framework that says,
- $142\ 00:07:19.730 \longrightarrow 00:07:22.100$  this is how you use all of these
- $143\ 00{:}07{:}22.100 \dashrightarrow 00{:}07{:}24.150$  in a clinical research program.
- $144\ 00{:}07{:}24.150 \dashrightarrow 00{:}07{:}26.860$  If you're gonna use social media in clinical study
- $145\ 00:07:26.860 --> 00:07:28.420$  for research purposes, you know,
- $146\ 00{:}07{:}28.420 {\:{\mbox{--}}}{>}\ 00{:}07{:}30.100$  there needs to be a framework for how to do it,
- $147\ 00:07:30.100 \longrightarrow 00:07:33.110$  especially in the context of precision oncology.
- 148 00:07:33.110 --> 00:07:33.943 But so we have this,
- $149\ 00:07:33.943 \longrightarrow 00:07:37.552$  and we have groups that are working on these databases,
- $150\ 00:07:37.552 \longrightarrow 00:07:40.350$  they want to make this a realization.

- $151\ 00:07:40.350 \longrightarrow 00:07:42.200$  What are the regulators saying?
- $152\ 00{:}07{:}42.200$  -->  $00{:}07{:}44.410$  Well, so real-world data and real-world evidence
- $153\ 00:07:44.410 --> 00:07:46.820$  really got a boost from the 21st Century Cures Act
- $154\ 00:07:46.820 \longrightarrow 00:07:48.740$  signed into law in 2016.
- $155\ 00:07:48.740 --> 00:07:50.960$  They advocated for the use of real-world evidence
- $156\ 00:07:50.960 \longrightarrow 00:07:53.650$  to support new indications for approved drugs.
- $157\ 00{:}07{:}53.650$  -->  $00{:}07{:}57.457$  Of course, the US Government wants the innovations
- $158\ 00:07:57.457 \longrightarrow 00:08:00.550$  that we have in biology to translate into therapeutics
- $159\ 00:08:00.550 \longrightarrow 00:08:01.383$  for patients.
- 160 00:08:01.383 --> 00:08:03.840 And we have a very, you know,
- $161\ 00:08:03.840 \longrightarrow 00:08:06.350\ I$  think forward looking approach when it comes to that,
- $162\ 00:08:06.350 \longrightarrow 00:08:08.340$  if the drug is relatively safe
- $163\ 00:08:08.340 \longrightarrow 00:08:09.910$  and can demonstrate some efficacy
- 164 00:08:09.910 --> 00:08:12.393 it gets to the market, it gets to patients.
- 165 00:08:13.320 --> 00:08:14.990 So that there's a guidance document
- $166\ 00:08:14.990 \longrightarrow 00:08:16.300$  about the use of real-world evidence
- 167 00:08:16.300 --> 00:08:18.010 to support regulatory decision-making,
- $168\ 00:08:18.010 \longrightarrow 00:08:20.320$  which was initially for devices.
- $169\ 00:08:20.320 \longrightarrow 00:08:22.980$  There's another one for biologics in 2019,
- 170 00:08:22.980 --> 00:08:24.930 there's actually a website you can go to,
- $171\ 00:08:24.930 \longrightarrow 00:08:29.145$  which is the framework they discussed in 2018.
- 172 00:08:29.145 --> 00:08:33.257 If you go to that website, and this was done,
- $173\ 00:08:33.257 \longrightarrow 00:08:36.210$  they have quotes from Scott Gottlieb here.
- 174 00:08:36.210 --> 00:08:38.600 You can see that a little more of a definition,
- $175\ 00{:}08{:}38.600 \dashrightarrow 00{:}08{:}40.930$  real-world data can be used to improve efficiency
- $176\ 00:08:40.930 \longrightarrow 00:08:42.980$  of clinical trials, even if it's not used

- $177\ 00:08:42.980 \longrightarrow 00:08:44.820$  for product effectiveness.
- 178 00:08:44.820 --> 00:08:46.550 So the FDA is still saying,
- $179\ 00{:}08{:}46.550$  -->  $00{:}08{:}49.430$  we don't want to use real-world data as a control arm
- 180 00:08:49.430 --> 00:08:52.630 to replace a randomized control, for example,
- $181~00{:}08{:}52.630 \dashrightarrow 00{:}08{:}55.660$  but we could use it to generate hypothesis, right?
- $182\ 00:08:55.660 \longrightarrow 00:08:57.030$  What is the expected event rate
- 183 00:08:57.030 --> 00:08:59.080 for this population that we're enrolling?
- $184\ 00:09:00.000$  --> 00:09:00.910 How many events do we expected
- 185 00:09:00.910 --> 00:09:03.200 to have in a certain timeframe?
- $186\ 00{:}09{:}03.200 \dashrightarrow 00{:}09{:}06.320$  How likely is it that we can roll that population.
- $187\ 00:09:06.320 \longrightarrow 00:09:09.090$  Trial feasibility and forming prior distributions
- $188\ 00:09:09.090 \longrightarrow 00:09:09.923$  in Bayesian models.
- 189 00:09:09.923 --> 00:09:11.170 So I liked that observation,
- 190 00:09:11.170 --> 00:09:14.610 but, you know, what is our expectation?
- 191 00:09:14.610 --> 00:09:16.490 Maybe we're not starting from nothing.
- 192 00:09:16.490 --> 00:09:18.660 And then prognostic indicators.
- 193 00:09:18.660 --> 00:09:20.420 Are there things we should stratify for
- $194\ 00:09:20.420$  --> 00:09:22.850 or account for an analysis that could be imbalanced,
- $195\ 00:09:22.850 \longrightarrow 00:09:25.410$  especially when we don't randomize?
- $196\ 00:09:25.410 \longrightarrow 00:09:27.310$  So this is what regulators are saying,
- 197 00:09:28.250 --> 00:09:29.640 but they also say the standard
- $198\ 00:09:29.640 --> 00:09:32.300$  for drug approval remains the same.
- $199\ 00:09:32.300 --> 00:09:33.960$  And this is an important statement.
- $200\ 00:09:33.960 \dashrightarrow 00:09:35.510$  The basis of approval remains the same.
- $201\ 00:09:35.510 \longrightarrow 00:09:37.730$  Substantial evidence that the drug will have the effect,
- $202\ 00:09:37.730 \dashrightarrow 00:09:40.700$  and adequate well-controlled clinical investigations.
- $203\ 00:09:40.700 \longrightarrow 00:09:43.970$  So. and I was just at a meeting at UNC

- $204\ 00:09:43.970 \longrightarrow 00:09:47.830$  with Genentech and FDA and people from the EMA,
- $205\ 00:09:47.830 \longrightarrow 00:09:50.533$  and they're standing firm on this.
- 206 00:09:52.000 --> 00:09:53.270 While we're discussing
- $207\ 00:09:53.270$  --> 00:09:55.910 how you could potentially augment a randomized-control
- $208\ 00:09:55.910 \longrightarrow 00:09:57.203$  with real-world controls,
- 209 00:09:58.070 --> 00:10:01.270 there's no sort of interest in replacing
- $210\ 00:10:02.438 \longrightarrow 00:10:04.760$  of randomized-control right now.
- $211\ 00:10:04.760 \longrightarrow 00:10:07.960$  Not unless there's absolutely no ethical way
- $212\ 00:10:07.960 \longrightarrow 00:10:08.960$  you could randomize.
- 213 00:10:10.595 --> 00:10:12.400 So they say that, you know,
- $214\ 00:10:12.400 \longrightarrow 00:10:14.410$  there's more flexibility when the disease is rare,
- $215\ 00:10:14.410 \longrightarrow 00:10:17.163$  and the patient population lacks a suitable control.
- 216 00:10:18.715 --> 00:10:19.947 So what about the CancerLinQ?
- $217\ 00:10:19.947 \longrightarrow 00:10:22.140$  So these slides are a little dated
- $218\ 00:10:22.140 \longrightarrow 00:10:23.810$  as of the last year, March of 2020,
- $219\ 00:10:23.810 \longrightarrow 00:10:25.960$  but they had at that time
- $220\ 00:10:25.960$  --> 00:10:30.050 over two and a half million patients in their database.
- $221\ 00:10:30.050 \longrightarrow 00:10:31.870$  So they have worked on data codes
- 222 00:10:31.870 --> 00:10:36.870 and structuring outcomes, structuring CONMED data.
- 223 00:10:37.010 --> 00:10:38.860 They've done a lot with this database,
- $224\ 00:10:38.860 \longrightarrow 00:10:41.103$  and I used it at Cleveland Clinic.
- $225\ 00:10:42.760 \longrightarrow 00:10:45.463$  So this is growing as a resource.
- 226 00:10:46.430 --> 00:10:50.529 Also what happened is that Flatiron,
- 227 00:10:50.529 --> 00:10:53.660 which has over 2 million active patients in their database.
- 228 00:10:53.660 --> 00:10:56.530 Of course, this is an industry group
- $229\ 00:10:56.530 \longrightarrow 00:10:59.660$  that's partially owned by Roche.

- $230\ 00:10:59.660$  --> 00:11:02.210 They have partnered with Foundation Medicine,
- 231 00:11:02.210 --> 00:11:05.010 and now there's an intersection of Flatiron patients
- $232\ 00:11:05.010 --> 00:11:08.670$  that also have genetic testing from Foundation Medicine.
- $233\ 00{:}11{:}08.670 \dashrightarrow 00{:}11{:}12.390$  And they're calling this the Clinical Genomic Database.
- 234 00:11:12.390 --> 00:11:14.300 And at the time that I took this slide,
- $235\ 00:11:14.300 \longrightarrow 00:11:15.443$  they had over 40,000 patients
- $236\ 00{:}11{:}15.443 \dashrightarrow 00{:}11{:}20.443$  that had the real-world data matched to the molecular data.
- 237 00:11:20.582 --> 00:11:22.370 I think that's very interesting,
- $238\ 00:11:22.370 \longrightarrow 00:11:24.070$  and I think that's very important.
- 239 00:11:25.000 --> 00:11:27.950 One of the main issues with real-world evidence
- $240\ 00:11:27.950 \longrightarrow 00:11:29.610$  in the oncology setting
- $241\ 00{:}11{:}30.450 \dashrightarrow 00{:}11{:}33.680$  is that we don't have a real-world tumor response.
- 242 00:11:33.680 --> 00:11:35.500 So for those of you that work in oncology,
- $243\ 00{:}11{:}35.500 \dashrightarrow 00{:}11{:}39.620$  of course, you know that phase one, phase two trials
- $244\ 00{:}11{:}39.620 \dashrightarrow 00{:}11{:}42.800$  are designed on the basis of endpoints
- $245\ 00:11:42.800 \longrightarrow 00:11:45.223$  that measure reductions in tumor burden.
- $246\ 00:11:46.250 \longrightarrow 00:11:48.610$  So for solid tumors, this is done through scans.
- 247 00:11:48.610 --> 00:11:50.370 So patients are scanned at baseline.
- $248\ 00{:}11{:}50.370 \dashrightarrow 00{:}11{:}52.930$  They're scanned regularly at follow-up intervals
- $249\ 00:11:52.930 \longrightarrow 00:11:56.740$  after every visit or every cycle of therapy.
- 250 00:11:56.740 --> 00:11:59.860 Those scans go for an adjudication process,
- $251\ 00:11:59.860 --> 00:12:01.930$  which is done by more than one person
- $252\ 00{:}12{:}01.930 \dashrightarrow 00{:}12{:}03.780$  where they actually measure how much reduction
- $253\ 00:12:03.780 --> 00:12:06.100$  in tumor burden happens after treatment.

- 254 00:12:06.100 --> 00:12:08.240 And then we look at that longitudinally,
- $255\ 00:12:08.240 \longrightarrow 00:12:10.490$  we take the best reduction
- $256\ 00:12:10.490 \longrightarrow 00:12:12.600$  or the most reduction that we saw,
- $257\ 00:12:12.600 \longrightarrow 00:12:15.610$  we consider did they have distant migration of disease?
- 258 00:12:15.610 --> 00:12:17.770 So for example, if they had a brain tumor,
- $259\ 00:12:17.770 --> 00:12:21.775$  did they also come in with tumors in their liver?
- $260\ 00:12:21.775 \longrightarrow 00:12:25.370$  And then we come up, we have a four point ordinal scale,
- $261\ 00:12:25.370 \longrightarrow 00:12:28.720$  and it tells us whether the patient has a complete response,
- 262 00:12:28.720 --> 00:12:31.120 which means the tumor burden's gone, right?
- $263\ 00:12:31.120 \longrightarrow 00:12:32.110$  The lesions are gone,
- 264 00:12:32.110 --> 00:12:34.520 or the blast counts in their blood are gone,
- $265\ 00:12:34.520 \longrightarrow 00:12:36.290$  if they have leukemia.
- $266\ 00:12:36.290 \longrightarrow 00:12:37.650$  They had a partial response.
- $267\ 00:12:37.650 \longrightarrow 00:12:39.920$  That means there was a reduction in their tumor size,
- $268\ 00:12:39.920 --> 00:12:43.210$  and it was a clinically meaningful reduction.
- 269 00:12:43.210 --> 00:12:44.290 They had stable disease,
- $270\ 00:12:44.290 \longrightarrow 00:12:46.420$  which means that there could have been a reduction,
- 271 00:12:46.420 --> 00:12:48.210 but it wasn't clinically meaningful,
- 272 00:12:48.210 --> 00:12:50.140 and it didn't really increase.
- $273~00{:}12{:}50.140 \dashrightarrow 00{:}12{:}52.920$  And progressive disease, the tumor burden is much higher
- $274\ 00:12:52.920 \longrightarrow 00:12:54.770$  than it was at baseline.
- 275 00:12:54.770 --> 00:12:59.080 So this process is critical for understanding
- 276 00:12:59.080 --> 00:13:01.900 and making decisions in phase two trials,
- $277\ 00:13:01.900 \longrightarrow 00:13:03.427$  as well as now the phase one trials
- $278\ 00:13:03.427 --> 00:13:06.870$  that we have in oncology, which are very large.
- $279\ 00:13:06.870 \longrightarrow 00:13:09.120$  This forms the basis for many go-decisions

- 280 00:13:09.120 --> 00:13:10.700 of whether you continue to develop a drug.
- $281\ 00:13:10.700 \longrightarrow 00:13:14.020$  Did it have a local effect on the tumor burden?
- 282 00:13:14.020 --> 00:13:15.330 It's very expensive to do this.
- $283\ 00:13:15.330 \longrightarrow 00:13:17.680$  It's very difficult to do this.
- $284\ 00:13:17.680 \longrightarrow 00:13:18.630$  So now we have to think
- $285~00:13:18.630 \longrightarrow 00:13:21.913$  about how can we get this information from an EMR?
- $286\ 00:13:22.870 --> 00:13:25.930$  Certainly patients may have scans in an EMR
- $287\ 00:13:25.930 \longrightarrow 00:13:27.510$  that we could use,
- $288\ 00:13:27.510 \longrightarrow 00:13:29.820$  but there's several issues with that.
- $289\ 00:13:29.820 --> 00:13:32.810$  So if we're going to use scans in a database
- 290 00:13:32.810 --> 00:13:35.500 to assess a patient's tumor burden,
- $291\ 00{:}13{:}35.500 \dashrightarrow 00{:}13{:}39.473$  number one, those scans don't go for a central review.
- 292 00:13:40.440 --> 00:13:42.280 The process by which the community
- $293\ 00:13:42.280 \longrightarrow 00:13:45.290$  or the non-trial evaluation of those scans
- $294\ 00:13:45.290 \longrightarrow 00:13:47.843$  is very different than the clinical trial process.
- 295 00:13:48.710 --> 00:13:51.290 They don't really have an ordinal scale
- $296\ 00:13:51.290 \longrightarrow 00:13:52.973$  like this that they use.
- 297 00:13:53.960 --> 00:13:57.520 Certainly, I think you could distinguish progressive disease
- $298\ 00:13:57.520 \longrightarrow 00:13:58.570$  from complete response.
- 299 00:13:58.570 --> 00:13:59.890 I think it'd be very difficult
- $300\ 00{:}13{:}59.890 \dashrightarrow 00{:}14{:}02.870$  to distinguish partial response from stable disease.
- $301~00{:}14{:}02.870 \dashrightarrow 00{:}14{:}05.210$  So we have groups that are saying they can do this, right?
- $302\ 00:14:05.210 \longrightarrow 00:14:07.440$  They're going back to the clinical annotations
- $303\ 00:14:07.440 \longrightarrow 00:14:09.080$  and the writing algorithms that look
- $304\ 00:14:09.080 \longrightarrow 00:14:11.190$  at the clinical annotations that says,
- 305 00:14:11.190 --> 00:14:13.840 well, if the notes say the lesions are all gone,
- 306 00:14:13.840 --> 00:14:16.030 then they had a complete response, right?

- 307 00:14:16.030 --> 00:14:19.980 If there was an increase, overall increase,
- $308\ 00:14:19.980 --> 00:14:22.930$  or there was new lesions, they had progressive disease.
- $309\ 00:14:22.930 \longrightarrow 00:14:24.930$  So if the annotations are good enough,
- 310 00:14:24.930 --> 00:14:27.420 I guess, you could get to progressive disease
- 311 00:14:27.420 --> 00:14:28.773 versus complete response.
- $312\ 00:14:29.870 \longrightarrow 00:14:32.890$  However, there are several issues with this.
- $313\ 00:14:32.890 \longrightarrow 00:14:35.820$  Everything in oncology is based on the line of therapy.
- $314\ 00{:}14{:}35.820 --> 00{:}14{:}39.130$  Patients come in, they get a sequence of treatments.
- $315\ 00:14:39.130 \longrightarrow 00:14:42.850$  Usually, they progress and go to a second line of therapy.
- $316\ 00:14:42.850 \longrightarrow 00:14:43.820$  Or they progress again
- $317\ 00:14:43.820 \longrightarrow 00:14:45.780$  and they go to a third line of therapy.
- $318\ 00:14:45.780 \longrightarrow 00:14:48.700$  The expectations for tumor response as both survival
- 319 00:14:48.700 --> 00:14:51.060 are very different by line of therapy.
- 320 00:14:51.060 --> 00:14:52.290 So if you're gonna go into the EMR,
- $321\ 00:14:52.290 \longrightarrow 00:14:54.550$  you have to now make sure
- $322\ 00{:}14{:}54.550 --> 00{:}14{:}58.180$  that the scans you're getting align with the line of the rapy
- $323\ 00:14:58.180 \longrightarrow 00:15:00.500$  that you're enrolling in your clinical study.
- 324 00:15:00.500 --> 00:15:03.890 So most clinical studies in oncology require
- $325\ 00:15:03.890 \longrightarrow 00:15:05.120$  a specific line of therapy.
- 326 00:15:05.120 --> 00:15:07.070 So first-line therapy means patients
- 327 00:15:07.070 --> 00:15:08.910 that haven't been treated previously.
- 328 00:15:08.910  $\rightarrow$  00:15:10.450 Second-line therapy means patients
- $329\ 00:15:10.450 \longrightarrow 00:15:12.460$  that have progressed on a prior treatment,
- $330\ 00{:}15{:}12.460 \dashrightarrow 00{:}15{:}14.700$  and now they're trying a subsequent treatment.
- $331\ 00:15:14.700 --> 00:15:17.650$  So the expectations are very different for response by that.

- $332\ 00:15:17.650 --> 00:15:20.170$  So you would have to know that this is the first,
- 333 00:15:20.170 --> 00:15:22.027 if you're using a first-line therapy study,
- $334\ 00:15:22.027 \longrightarrow 00:15:23.270$  you would have to know
- 335 00:15:23.270 --> 00:15:25.100 that this is this patient's first line of therapy
- $336\ 00:15:25.100 \longrightarrow 00:15:27.070$  and these scans correspond to that.
- $337\ 00:15:27.070 \longrightarrow 00:15:28.540$  Not only that, you'd have to make sure
- $338\ 00{:}15{:}28.540 \dashrightarrow 00{:}15{:}31.400$  the scans reasonably aligned with the time-frame
- $339\ 00:15:31.400 \longrightarrow 00:15:32.410$  by which the clinical trial
- 340 00:15:32.410 --> 00:15:34.513 is actually going to acquire their energy.
- 341 00:15:36.900 --> 00:15:40.062 Beyond that, you'd have to, you know,
- $342\ 00:15:40.062 --> 00:15:43.290$  there are several other issues with that, right?
- $343\ 00:15:43.290 \longrightarrow 00:15:47.210$  Patients may not be scanned in the community setting.
- 344 00:15:47.210 --> 00:15:49.210 And working with oncologists for a long time,
- 345 00:15:49.210 --> 00:15:51.530 I know that there's a certain point
- $346\ 00{:}15{:}51.530 {\:{\mbox{--}}}{>}\ 00{:}15{:}54.063$  where if a patient fails a few lines of the rapy,
- $347\ 00{:}15{:}55{.}000 \dashrightarrow 00{:}15{:}58.230$  they may not wanna risk the patient getting nephrotoxicity
- $348\ 00{:}15{:}58.230 \dashrightarrow 00{:}16{:}00.940$  from the contrast that are used in the scans.
- $349\ 00{:}16{:}00{.}940 \dashrightarrow 00{:}16{:}04{.}440$  So if a patient doesn't have a lot of good treatment options
- 350 00:16:04.440 --> 00:16:06.193 or they're reasonably unhealthy,
- $351\ 00:16:06.193 \longrightarrow 00:16:09.180$  where there's concern about kidney or liver issues,
- $352\ 00:16:09.180 --> 00:16:11.660$  they don't scan the patients in the community.
- $353\ 00:16:11.660 \longrightarrow 00:16:15.900$  So up till now, I think that the consensus has been,
- $354\ 00{:}16{:}15{.}900 \dashrightarrow 00{:}16{:}18{.}560$  there is no real-world tumor response right now.
- 355~00:16:18.560 --> 00:16:19.510 We don't have that.
- $356\ 00:16:19.510 --> 00:16:21.790$  And I think that's difficult because

- $357\ 00{:}16{:}22.910 \dashrightarrow 00{:}16{:}26.130$  we want to use real-world data to sort of augment
- $358\ 00:16:26.130 --> 00:16:27.410$  or supplement the areas
- $359\ 00{:}16{:}27.410 \dashrightarrow 00{:}16{:}29.650$  where we don't have a lot of information, right?
- $360\ 00:16:29.650 \longrightarrow 00:16:32.750$  And that is the early phase studies, right?
- 361 00:16:32.750 --> 00:16:33.830 Once you go to phase three,
- $362\ 00{:}16{:}33.830 \dashrightarrow 00{:}16{:}36.960$  you've kind of established that the drug may be promising
- $363~00:16:36.960 \longrightarrow 00:16:39.240$  and you're gonna run a seven-year trial.
- $364\ 00:16:39.240 \longrightarrow 00:16:40.350$  And over that seven years,
- 365 00:16:40.350 --> 00:16:42.570 you're gonna acquire lots of information,
- $366\ 00:16:42.570 \longrightarrow 00:16:45.030$  and you're gonna follow them for survival.
- $367\ 00:16:45.030 --> 00:16:46.710$  This could, with really the narrative
- 368 00:16:46.710 --> 00:16:47.927 about real-world evidence in oncology,
- $369\ 00:16:47.927 --> 00:16:49.500$  has really been we can supplement
- $370\ 00:16:49.500 \longrightarrow 00:16:51.830$  those early phase decisions.
- 371 00:16:51.830 --> 00:16:52.663 But to do that,
- $372\ 00:16:52.663 --> 00:16:54.710$  we really have to have a real-world tumor response.
- $373\ 00:16:54.710 \longrightarrow 00:16:57.010$  And right now we don't have it.
- 374 00:16:57.010 --> 00:16:59.020 This is a paper from Advanced Therapeutics
- $375\ 00:16:59.020 \longrightarrow 00:17:00.900$  that was published this year.
- 376 00:17:00.900 --> 00:17:03.560 We have the Flatiron group going back
- $377\ 00:17:03.560 \longrightarrow 00:17:05.670$  to the major immunotherapy trials
- $378\ 00:17:05.670 \longrightarrow 00:17:08.160$  that have been implemented in recent years.
- 379 00:17:08.160 --> 00:17:09.760 They're comparing their algorithm
- $380\ 00:17:09.760 \longrightarrow 00:17:11.980$  for real-world response rates
- $381\ 00:17:11.980 \longrightarrow 00:17:16.030$  with the trial confirmed response.
- $382\ 00:17:16.030 \longrightarrow 00:17:16.863$  So they're saying,
- $383\ 00:17:16.863 --> 00:17:19.300$  for each patient, what did we say the response was

- $384\ 00:17:19.300 \longrightarrow 00:17:21.430$  based on our EMR data?
- $385\ 00:17:21.430 \longrightarrow 00:17:23.060$  What did the trial said the response was?
- 386 00:17:23.060 --> 00:17:24.880 And they're looking at sort of coordinates
- $387\ 00:17:24.880 \longrightarrow 00:17:26.480$  between those measures.
- $388\ 00:17:26.480 \longrightarrow 00:17:28.290$  And they're doing this by line of therapy.
- $389\ 00:17:28.290 \longrightarrow 00:17:29.940$  So maybe we'll get there,
- $390\ 00:17:29.940 \longrightarrow 00:17:32.290$  but right now the consensus is we're not there.
- 391 00:17:33.563 --> 00:17:38.210 So we presented this paper at ASCO,
- $392\ 00{:}17{:}38.210 \dashrightarrow 00{:}17{:}41.270$  which is a big cancer meeting in the US last year,
- 393 00:17:41.270 --> 00:17:42.103 talking about,
- $394\ 00:17:42.103 \longrightarrow 00:17:44.210$  can we actually replace randomized controls
- $395\ 00:17:44.210 \longrightarrow 00:17:46.310$  with external real-world controls?
- $396\ 00{:}17{:}46.310 \dashrightarrow 00{:}17{:}50.710$  And we actually built some tools that Genentech is using
- 397 00:17:50.710 --> 00:17:51.820 that actually calculate,
- $398~00:17:51.820 \dashrightarrow 00:17:56.200$  that takes your assumptions about bias, heterogeneity,
- $399\ 00:17:56.200 \longrightarrow 00:17:57.730$  or other things that you might see in a trial
- 400 00:17:57.730 --> 00:18:00.580 and actually tells you how wrong you can go
- 401~00:18:00.580 --> 00:18:04.350 with a go-decision when you use an external control.
- $402\ 00:18:04.350 --> 00:18:06.503$  And of course, I think maybe every body knows this,
- $403\ 00:18:06.503 \longrightarrow 00:18:10.796$  that the reality is that if there's no bias, it's useful.
- $404\ 00{:}18{:}10.796 \dashrightarrow 00{:}18{:}14.010$  If there is bias, things can go really wrong very quickly,
- $405\ 00:18:14.010 \longrightarrow 00:18:15.940$  depending on the direction of the bias.
- $406\ 00:18:15.940 \longrightarrow 00:18:18.220$  And that is really unknown.
- $407\ 00:18:18.220$  --> 00:18:23.220 So we tried to think about this in a very systematic way,
- $408\ 00:18:23.330 \longrightarrow 00:18:25.330$  and I think it's challenging.

- $409\ 00:18:25.330 \longrightarrow 00:18:26.980\ I\ don't\ know\ that\ we\ can\ do\ this.$
- $410\ 00{:}18{:}27.920 \dashrightarrow 00{:}18{:}31.040$  So that leads to, you know, what this discussion was
- $411\ 00{:}18{:}31.040 \dashrightarrow 00{:}18{:}33.950$  at UNC with the FDA, the EMA, and Genentech
- $412\ 00:18:33.950 \longrightarrow 00:18:35.110$  where we're talking
- $413\ 00{:}18{:}35{.}110 \dashrightarrow 00{:}18{:}39{.}130$  about now, can we augment randomized control arms
- 414 00:18:39.130 --> 00:18:42.330 with data from real-world sources?
- $415\ 00:18:42.330 \longrightarrow 00:18:46.283$  So we don't get rid of the randomized control,.
- $416\ 00:18:46.283 \longrightarrow 00:18:47.670$  We keep the randomized control,
- $417\ 00{:}18{:}47.670 \dashrightarrow 00{:}18{:}50.740$  but we supplement it with some external controls.
- $418\ 00:18:50.740 \longrightarrow 00:18:51.597$  How could we do that?
- 419 00:18:51.597 --> 00:18:53.040 And could we even acquire those
- $420\ 00:18:53.040 \longrightarrow 00:18:55.950$  before the trial gets initiated?
- 421 00:18:55.950 --> 00:18:57.940 Of course, it takes a long time for protocols
- $422\ 00:18:57.940 \longrightarrow 00:19:00.140$  to be reviewed and other things to happen.
- 423 00:19:00.140 --> 00:19:03.180 Well, this gets interesting to me
- $424\ 00:19:03.180 \longrightarrow 00:19:07.570$  because while I developed tools to do this a long time ago,
- $425\ 00:19:07.570 \longrightarrow 00:19:10.610$  which I called Multi-source Adaptive Designs.
- 426 00:19:10.610 --> 00:19:14.840 And this was done many years ago
- 427 00:19:14.840 --> 00:19:17.090 before we talked about real-world evidence.
- $428\ 00:19:17.090 \longrightarrow 00:19:20.140$  We were talking about historical controls at that time,
- $429\ 00:19:20.140 \longrightarrow 00:19:21.660$  but of course we can do interesting things
- $430\ 00:19:21.660 \longrightarrow 00:19:22.623$  with modeling here.
- $431\ 00:19:23.640 \longrightarrow 00:19:25.550$  We could take real-world controls,
- $432\ 00:19:25.550 \longrightarrow 00:19:28.860$  we could think about an interim analysis
- 433 00:19:28.860 --> 00:19:29.950 of a randomized trial,
- $434\ 00{:}19{:}29.950 \dashrightarrow 00{:}19{:}33.100$  where we have randomized treated and randomized controls.

- $435\ 00:19:33.100 \longrightarrow 00:19:36.570$  We could do any sort of fancy model that you wanna fit,
- $436\ 00{:}19{:}36.570 \dashrightarrow 00{:}19{:}39.990$  and we could assess how biased are these historical controls
- $437\ 00:19:39.990 --> 00:19:43.020$  or real-world controls in relation to the control data
- $438\ 00:19:43.020$  --> 00:19:45.710 that we're seeing in the actual randomized trial.
- $439\ 00:19:45.710 \longrightarrow 00:19:47.070$  On the basis of this model,
- 440 00:19:47.070 --> 00:19:50.130 we could actually adapt the allocation, right?
- $441\ 00:19:50.130 \longrightarrow 00:19:51.440$  If we don't see a lot of bias,
- $442\ 00:19:51.440 \longrightarrow 00:19:55.370$  so those patients, based on the eligibility of the trial,
- 443 00:19:55.370 --> 00:19:56.710 those patients from the community,
- 444 00:19:56.710 --> 00:19:59.500 they look a lot like the patients in the trial,
- $445\ 00:19:59.500 \dashrightarrow 00:20:01.720$  then you have more information on the control side.
- $446\ 00{:}20{:}01.720 \dashrightarrow 00{:}20{:}04.090$  You need to rebalance the rest of your allocation
- $447\ 00:20:04.090 \longrightarrow 00:20:05.890$  so that you can increase power.
- $448\ 00:20:05.890 \longrightarrow 00:20:07.040$  So this is the only designs
- $449\ 00{:}20{:}07.040 \dashrightarrow 00{:}20{:}09.807$  where you can actually increase statistical power
- $450\ 00:20:09.807 \longrightarrow 00:20:12.010$  with a smaller trial.
- 451 00:20:12.010 --> 00:20:13.100 Because what we're trying to do,
- $452\ 00{:}20{:}13.100 \dashrightarrow 00{:}20{:}15.090$  is we're trying to balance the overall information
- $453\ 00:20:15.090 \longrightarrow 00:20:17.130$  between the treatments, right?
- $454\ 00{:}20{:}17.130 \dashrightarrow 00{:}20{:}19.628$  If you look at the outcome, adaptive randomized studies,
- $455\ 00:20:19.628 \longrightarrow 00:20:21.320$  they required larger trials
- $456\ 00:20:21.320 \longrightarrow 00:20:22.810$  because they're imbalancing.
- 457 00:20:22.810 --> 00:20:24.770 They're imbalancing based on outcomes.
- $458\ 00:20:24.770 \longrightarrow 00:20:27.010$  We're trying to balance based on bias.

- $459\ 00:20:27.010 --> 00:20:28.990$  So we worked out this methodology
- $460\ 00:20:28.990 --> 00:20:31.550$  and you know, ASCO and Flatiron
- $461\ 00:20:31.550 \longrightarrow 00:20:33.083$  are interested in using this.
- $462\ 00:20:34.240 \longrightarrow 00:20:35.510$  We have a paper that describes
- $463\ 00:20:35.510 \longrightarrow 00:20:37.000$  an open-source tool that we have.
- 464 00:20:37.000 --> 00:20:38.700 It's still on MD Anderson's website
- 465~00:20:38.700 --> 00:20:41.763 that I built when I was at MD Anderson with Nan Chen.
- $466\ 00:20:42.660 \longrightarrow 00:20:43.970$  Who is pictured is here.
- $467\ 00:20:43.970 \longrightarrow 00:20:45.503$  So Nan is now at Gilead.
- 468 00:20:46.640 --> 00:20:49.000 But if you interested in this, it's here.
- 469 00:20:49.000 --> 00:20:54.000 So I think based on in oncology setting,
- $470\ 00:20:54.290 \longrightarrow 00:20:56.260$  we need to focus on this area.
- 471 00:20:56.260 --> 00:20:57.930 We need to focus on hybrid controls,
- 472 00:20:57.930 --> 00:21:00.050 not replacing control arms, right?
- $473\ 00:21:00.050 \longrightarrow 00:21:01.680$  At least for most studies.
- $474\ 00{:}21{:}01.680 \dashrightarrow 00{:}21{:}05.700$  Of course, in rare diseases or areas of pediatric cancer,
- 475 00:21:05.700 --> 00:21:08.300 or both, you need to do something else, right?
- 476 00:21:08.300 --> 00:21:09.990 And that's what the FDA is talking about
- 477 00:21:09.990 --> 00:21:11.300 when they talk about flexibility.
- 478 00:21:11.300 --> 00:21:12.133 But I'm talking about
- $479\ 00{:}21{:}12.133 \dashrightarrow 00{:}21{:}16.020$  from kind of the standard drug development program
- 480 00:21:16.020 --> 00:21:17.203 in oncology right now.
- 481 00:21:18.354 --> 00:21:20.840 So I've talked about the issues,
- $482\ 00:21:20.840 \longrightarrow 00:21:23.650$  I've talked about the databases
- $483\ 00:21:23.650 \longrightarrow 00:21:24.750$  and sort of what's going on
- $484\ 00:21:24.750 \longrightarrow 00:21:26.550$  with real-world data in oncology.
- $485\ 00:21:26.550 \longrightarrow 00:21:29.940$  There's another group of sort of players in the space.
- $486\ 00:21:29.940 \longrightarrow 00:21:30.773$  And I would call them

- $487\ 00:21:30.773 --> 00:21:33.233$  kind of the real-world evidence zealots.
- 488 00:21:34.642 --> 00:21:39.642 This guy, Dr. Butte from Stanford has,
- 489 00:21:39.670 --> 00:21:41.620 I think represents one of these people.
- 490 00:21:42.520 --> 00:21:46.540 So he is a strong advocate for using databases
- $491\ 00:21:46.540 \longrightarrow 00:21:49.623$  to replace clinical research.
- 492 00:21:51.910 --> 00:21:54.300 He has at least three TED Talks,
- 493 00:21:54.300 --> 00:21:57.293 and I was going through them yesterday.
- 494 00:21:58.610 --> 00:22:01.220 He has a very strong feeling that we just need
- 495 00:22:01.220 --> 00:22:02.640 to organize these databases,
- $496~00{:}22{:}02.640 \dashrightarrow 00{:}22{:}05.770$  and we can answer any medical or scientific question
- $497\ 00:22:05.770 \longrightarrow 00:22:07.290$  that we want to answer.
- 498 00:22:07.290 --> 00:22:08.461 And in fact, he even says,
- $499\ 00:22:08.461 --> 00:22:12.310$  the problem is there's not enough people asking questions.
- $500\ 00:22:12.310 \longrightarrow 00:22:13.910$  That's the real issue right now.
- 501 00:22:15.240 --> 00:22:17.470 So there's this other group of people
- 502 00:22:17.470 --> 00:22:19.610 that are you know really hyping up
- $503\ 00:22:20.640 \longrightarrow 00:22:22.620$  the fact that it's just a computing problem.
- $504\ 00:22:22.620 \longrightarrow 00:22:23.453$  we have the data,
- 505~00:22:23.453 --> 00:22:26.630 we can use algorithms to answer any question we want.
- 506~00:22:26.630 --> 00:22:31.233 This group of people seems to lack any recognition
- 507 00:22:31.233 --> 00:22:34.840 of the principles of experimental design.
- $508~00{:}22{:}34.840 \dashrightarrow 00{:}22{:}37.840$  They don't seem to acknowledge them anywhere in the process.
- $509~00{:}22{:}39.619 \dashrightarrow 00{:}22{:}43.950$  And Dr. Butte and his TED talks actually says
- $510~00:22:43.950 \dashrightarrow 00:22:46.270$  that we don't need randomized controls after all,
- $511\ 00:22:46.270 \longrightarrow 00:22:49.420$  we just need to build databases.
- $512\ 00:22:49.420 \longrightarrow 00:22:50.489$  So we had these groups,

- $513\ 00:22:50.489 \longrightarrow 00:22:53.220$  so these are the kind of the players pushing this forward.
- 514~00:22:53.220 --> 00:22:54.460 So now I'm gonna transition here.
- 515 00:22:54.460 --> 00:22:56.110 I'm gonna talk about what's going on
- 516 00:22:56.110 --> 00:22:57.453 in precision oncology.
- 517 00:22:58.980 --> 00:23:00.550 Okay, so this is how you learned
- $518\ 00:23:00.550 \longrightarrow 00:23:02.610$  about drug development programs.
- $519\ 00:23:02.610 \longrightarrow 00:23:06.109$  You learned that we chose dose in phase one,
- $520\ 00{:}23{:}06.109 \dashrightarrow 00{:}23{:}11.010$  if the dose was promising and we were able to discover
- $521~00{:}23{:}12.020 \dashrightarrow 00{:}23{:}15.050$  what the MTD was, and we felt like it wasn't toxic
- $522\ 00:23:15.050 \longrightarrow 00:23:15.900$  and we had a good dose,
- $523\ 00:23:15.900 \longrightarrow 00:23:17.640$  we would go to a phase two trial.
- 524 00:23:17.640 --> 00:23:19.850 In oncology, we would look at tumor response.
- 525 00:23:19.850 --> 00:23:21.960 So again, reduction in tumor burden.
- $526\ 00:23:21.960 \longrightarrow 00:23:23.940$  Usually these would be uncontrolled.
- $527\ 00:23:23.940 \longrightarrow 00:23:26.360$  They would be about 50-100 patients.
- $528\ 00:23:26.360 \longrightarrow 00:23:29.660$  If we saw the drug had local activity on tumor burden,
- $529\ 00:23:29.660 \longrightarrow 00:23:31.540$  we would go to a phase three trial.
- $530~00:23:31.540 \longrightarrow 00:23:33.190$  The phase three trial would randomize
- $531\ 00:23:33.190 \longrightarrow 00:23:35.090$  to the existing standard of care.
- $532\ 00:23:35.090 \longrightarrow 00:23:38.073$  And would see if the treatment prolonged survival.
- 533 00:23:38.910 --> 00:23:40.830 This is what you learned about,
- $534\ 00:23:40.830 --> 00:23:44.090$  but oncology has changed very rapidly.
- $535\ 00:23:44.090 --> 00:23:47.000$  Regulatory policy has changed as well.
- $536\ 00{:}23{:}47.000 \dashrightarrow 00{:}23{:}50.770$  So molecular biologists have some victories recently.
- $537\ 00{:}23{:}50.770 --> 00{:}23{:}53.430$  They have really, you know, a lot of the biological models
- $538\ 00:23:53.430 --> 00:23:56.620$  that were discovered a decade ago

- $539\ 00:23:56.620 \longrightarrow 00:23:59.890$  have been translated into the rapeutics.
- $540\ 00:23:59.890 \longrightarrow 00:24:02.570$  So it used to be that we needed one
- $541\ 00:24:02.570 --> 00:24:04.640$  or two well-controlled phase three trials
- $542\ 00:24:04.640 \longrightarrow 00:24:08.150$  before we got regulatory approval.
- $543\ 00:24:08.150 --> 00:24:11.380$  It turns out that cancer biologists
- 544 00:24:11.380 --> 00:24:14.520 have identified very specific cancer subsets
- $545\ 00:24:14.520 --> 00:24:16.893$  based on genetics and based on immunology.
- $546\ 00:24:17.930 \longrightarrow 00:24:21.040$  With those cancer subsets, we have seen very promising,
- $547~00{:}24{:}21.040 \dashrightarrow 00{:}24{:}26.040$  very exciting results in phase two trials without controls.
- $548~00{:}24{:}26.360 \dashrightarrow 00{:}24{:}29.050$  The FDA started to allow conditional approvals
- $549\ 00:24:29.050 \longrightarrow 00:24:30.720$  after phase two on the basis
- $550~00{:}24{:}30.720 \dashrightarrow 00{:}24{:}33.210$  of those biomarker targeted treatments.
- $551\ 00:24:33.210 \longrightarrow 00:24:35.730$  Now we're in kind of stage three here.
- $552~00{:}24{:}35.730 \dashrightarrow 00{:}24{:}38.220$  Now we have the awareness that many of the targets,
- 553 00:24:38.220 --> 00:24:39.390 many of the genetic targets,
- $554\ 00:24:39.390 \longrightarrow 00:24:44.330$  as well as the immune phenotypes that we're interested in,
- $555\ 00:24:44.330 \longrightarrow 00:24:46.710$  they actually exist across several different
- $556\ 00:24:46.710 \longrightarrow 00:24:50.380$  sort of traditionally distinct cancer patients.
- $557~00{:}24{:}50.380 \dashrightarrow 00{:}24{:}54.260$  So patients with pancreatic cancer and lung cancer
- $558~00{:}24{:}54.260 \dashrightarrow 00{:}24{:}56.960$  may be very different from a clinical perspective,
- $559\ 00:24:56.960 --> 00:24:58.710$  but they might share a molecular feature
- $560\ 00:24:58.710 \longrightarrow 00:25:01.380$  that can be targeted by the same drug.
- $561\ 00:25:01.380 \longrightarrow 00:25:02.940$  And we're now in the space
- 562 00:25:02.940 --> 00:25:06.210 of histology-agnostic drug development,
- $563\ 00:25:06.210 --> 00:25:09.048$  where we might be replacing
- $564\ 00:25:09.048 \longrightarrow 00:25:11.220$  traditional classification criteria

- $565\ 00:25:11.220 \longrightarrow 00:25:13.310$  based on molecular features.
- $566\ 00:25:13.310 --> 00:25:17.603$  So we're basically finding new subtypes of cancers as we go.
- 567 00:25:18.820 --> 00:25:20.800 These subtypes are very small,
- $568\ 00:25:20.800 \longrightarrow 00:25:21.940$  and they're becoming smaller
- $569\ 00:25:21.940 --> 00:25:23.943$  as we learn more about cancer biology.
- $570~00:25:24.830 --> 00:25:28.270~\mathrm{But}$  a few of them had had very exceptional results.
- 571 00:25:28.270 --> 00:25:29.650 A few drugs targeting these event,
- $572~00{:}25{:}29.650 \dashrightarrow 00{:}25{:}31.080$  have had very exceptional results,
- 573 00:25:31.080 --> 00:25:32.950 crossing many tumor types.
- $574~00{:}25{:}32.950 \dashrightarrow 00{:}25{:}36.940$  And they have gotten accelerated approval for agnostic drugs
- $575\ 00:25:36.940 --> 00:25:38.330$  and drugs that can be administered
- $576\ 00:25:38.330 \longrightarrow 00:25:40.850$  without regard to the tissue of origin.
- 577 00:25:40.850 --> 00:25:42.653 And this has happened in phase one.
- 578 00:25:43.790 --> 00:25:45.410 So the regulatory landscape has changed,
- 579 00:25:45.410 --> 00:25:50.330 the development landscape has changed.
- $580\ 00:25:50.330 \longrightarrow 00:25:52.840$  So I got to be a part of this review
- 581 00:25:52.840 --> 00:25:55.093 for Nature of Clinical Oncology,
- $582\ 00{:}25{:}57.200 \dashrightarrow 00{:}26{:}00.340$  where we talked about these tissue-agnostic drugs.
- 583 00:26:00.340 --> 00:26:02.480 There's actually four drugs so far
- 584 00:26:02.480 --> 00:26:04.110 that have been approved by the FDA
- $585\ 00:26:04.110 --> 00:26:07.580$  that could be administered based on a marker feature,
- $586\ 00:26:07.580 \longrightarrow 00:26:09.873$  not on the actual cancer tissue.
- $587\ 00:26:11.050 \longrightarrow 00:26:13.883$  So now look at the issues with this.
- 588~00:26:13.883 --> 00:26:17.260 There's four drugs, and there's three different biomarkers
- $589\ 00{:}26{:}17.260 \dashrightarrow 00{:}26{:}19.960$  that have been approved for tissue-agnostic treatment.
- 590 00:26:20.870 --> 00:26:23.190 One of the biomarkers is the NTRK fusion,

- $591\ 00:26:23.190 --> 00:26:25.200$  which we'll talk about little later.
- $592\ 00:26:25.200 \longrightarrow 00:26:27.040$  It's exceedingly rare.
- 593 00:26:27.040 --> 00:26:28.360 You can see that breast cancer,
- $594\ 00:26:28.360 \longrightarrow 00:26:30.602$  we're talking about less than 0.1%
- 595 00:26:30.602 --> 00:26:33.607 of the patients have an NTRK fusion, right?
- $596\ 00:26:33.607 --> 00:26:37.320$  And CRC it's about 1% of patients.
- 597 00:26:37.320 --> 00:26:40.470 There's a few tumors where it's more common,
- 598 00:26:40.470 --> 00:26:42.180 but this becomes very challenging.
- $599\ 00:26:42.180 \longrightarrow 00:26:45.410$  It becomes very challenging to design a study
- 600~00:26:45.410 --> 00:26:48.350 where we can actually study patients with NTRK fusions.
- $601\ 00:26:48.350 --> 00:26:49.890$  And then who are you gonna get in your study?
- $602\ 00:26:49.890 --> 00:26:52.960$  You're going to get a mixture of many different tissues
- 603 00:26:52.960 --> 00:26:54.100 that were traditionally thought
- $604\ 00:26:54.100 \longrightarrow 00:26:55.503$  to be separate cancers.
- $605\ 00:26:56.670 \longrightarrow 00:27:01.670$  So with this transition to tissue-agnostic drug development,
- $606\ 00{:}27{:}03.210 \dashrightarrow 00{:}27{:}05.210$  there's a statistical question that we have to answer,
- $607\ 00:27:05.210 \longrightarrow 00:27:07.331$  and that is who can be averaged?
- $608\ 00{:}27{:}07.331$  -->  $00{:}27{:}12.331$  Which tissue types could be averaged statistically,
- $609\ 00{:}27{:}12.770 {\:{\circ}{\circ}{\circ}}>00{:}27{:}16.510$  when we assess the effectiveness of a biomarker targets
- $610\ 00:27:16.510 \longrightarrow 00:27:18.451$  and a therapeutic?
- $611\ 00:27:18.451 --> 00:27:22.250$  And that's the question of statistical exchangeability.
- 612 00:27:22.250 --> 00:27:24.460 So we have developed patient models
- $613\ 00:27:24.460 \longrightarrow 00:27:28.470$  that actually characterize
- $614~00{:}27{:}28.470 \dashrightarrow 00{:}27{:}32.770$  what subsets of tumors actually respond in a similar way
- $615\ 00:27:32.770 \longrightarrow 00:27:33.950$  to a targeted therapy.

- 616 00:27:33.950 --> 00:27:37.240 And this gives us statistical criteria
- $617\ 00:27:37.240 \longrightarrow 00:27:40.320$  for understanding what is agnostic and what is not.
- 618 00:27:40.320 --> 00:27:42.580 And I got the, you know, this is the first time,
- $619\ 00:27:42.580 \longrightarrow 00:27:44.040\ I$  got to collaborate with Dr. Kane
- 620 00:27:44.040 --> 00:27:46.170 on actually building out tools for this.
- 621 00:27:46.170 --> 00:27:49.920 So I can do the methods, but the tools or something else.
- 622 00:27:49.920 --> 00:27:51.730 So Michael got these incredible tools.
- $623\ 00{:}27{:}51.730 {\:\hbox{--}}{>}\ 00{:}27{:}55.520$  And we have an open source package for fitting these models.
- 624 00:27:55.520 --> 00:27:57.690 Just to give you sort of motivation here.
- $625\ 00:27:57.690 \longrightarrow 00:27:58.990$  This is an actual trial
- $626\ 00{:}27{:}58.990$  -->  $00{:}28{:}03.760$  that was evaluating a drug called Bendroflumeth
- 627~00:28:03.760 --> 00:28:07.670 in BRAF tumors, patients that have BRAF mutations.
- $628\ 00:28:07.670 \longrightarrow 00:28:09.500$  So there is BRAF mutations can occur
- $629\ 00:28:09.500 \longrightarrow 00:28:10.770$  in many different tumors.
- $630\ 00{:}28{:}10.770\ -->\ 00{:}28{:}12.980$  They initially developed this drug in Melanoma,
- 631 00:28:12.980 --> 00:28:15.020 but then they saw BRAF tumors,
- $632~00:28:15.020 \longrightarrow 00:28:17.230~\mathrm{BRAF}$  mutations exist in these other cancers.
- 633 00:28:17.230 --> 00:28:19.810 Histocytosis, thyroid cancer,
- 634 00:28:19.810 --> 00:28:21.940 cholangiocarcinoma, for example.
- $635\ 00:28:21.940 --> 00:28:23.880$  So they ran up, what's known as a Basket Trial,
- $636\ 00:28:23.880 \longrightarrow 00:28:26.220$  where they allowed these different tumor types
- $637\ 00:28:26.220 \longrightarrow 00:28:27.363$  in the same trial.
- 638 00:28:28.386 --> 00:28:31.170 So we show in this nature of these clinical oncology paper,
- $639\ 00:28:31.170 \longrightarrow 00:28:33.530$  how these exchangeability models work.
- $640\ 00{:}28{:}33.530 \dashrightarrow 00{:}28{:}36.110$  We call them multi-source exchangeability models.

- $641\ 00{:}28{:}36.110 --> 00{:}28{:}39.400$  Where we start with an assumption that these tumors
- 642 00:28:39.400 --> 00:28:41.210 are gonna act in the same way, right?
- $643\ 00:28:41.210 \longrightarrow 00:28:42.680$  So the drug target combination
- 644 00:28:42.680 --> 00:28:47.090 is going to be kind of equally efficacious
- $645\ 00:28:47.090 \longrightarrow 00:28:47.960$  among all the tumors.
- $646\ 00:28:47.960 \longrightarrow 00:28:49.590$  So they're exchangeable statistically.
- $647\ 00:28:49.590 \longrightarrow 00:28:51.300$  We can average them.
- 648 00:28:51.300 --> 00:28:54.610 As we start to get data from the trial,
- $649\ 00:28:54.610 --> 00:28:57.670$  we can now start to assess the heterogeneity
- $650\ 00:28:57.670 \longrightarrow 00:28:58.917$  that we see across these tumors.
- 651 00:28:58.917 --> 00:29:00.690 And we can ask the question,
- 652 00:29:00.690 --> 00:29:03.810 is it really agnostic to the tumor type?
- 653 00:29:03.810 --> 00:29:05.220 Now, when it comes to vendor afatinib,
- $654\ 00:29:05.220 \longrightarrow 00:29:08.700$  we had three tumor types that did really well in this trial,
- 655~00:29:08.700 --> 00:29:11.800 histiocytosis, thyroid, and non-small cell lung cancer.
- 656 00:29:11.800 --> 00:29:13.910 Colorectal did not do well.
- 657 00:29:13.910 --> 00:29:15.640 So even though colorectal cancer patients
- 658 00:29:15.640 --> 00:29:17.350 had BRAF mutations,
- $659\ 00:29:17.350 --> 00:29:19.740$  they did not respond to vendor afatinib.
- $660\ 00:29:19.740 \longrightarrow 00:29:22.160$  These tumors did respond.
- $661\ 00:29:22.160 \longrightarrow 00:29:23.770$  We don't know about cholangiocarcinoma.
- $662\ 00:29:23.770 --> 00:29:25.750$  There wasn't enough information in that trial
- $663\ 00:29:25.750 \longrightarrow 00:29:26.583$  to really tell us.
- $664\ 00:29:26.583 \longrightarrow 00:29:28.520$  So they're kind of in the center here.
- 665 00:29:28.520 --> 00:29:31.010 So, you know, this is just to give you a flavor
- 666 00:29:31.010 --> 00:29:33.182 of what's going on in oncology right now,
- 667 00:29:33.182 --> 00:29:35.750 As we start to go towards precision medicine,
- $668~00{:}29{:}35.750 \dashrightarrow 00{:}29{:}38.730$  that means that we have features across traditionally

- $669\ 00:29:38.730 \longrightarrow 00:29:39.790$  very different cancers.
- $670\ 00:29:39.790 \longrightarrow 00:29:40.790$  And we have to understand
- 671 00:29:40.790 --> 00:29:43.570 whether it's actually the feature that's driving,
- $672\ 00:29:43.570 \longrightarrow 00:29:45.582$  what we see in the response.
- $673\ 00:29:45.582 \longrightarrow 00:29:47.930$  Okay, so this is an issue
- 674 00:29:47.930 --> 00:29:52.430 that I don't think is that well understood outside
- $675\ 00:29:52.430 \longrightarrow 00:29:55.200$  of our sort of biostatistical and statistical communities.
- 676 00:29:55.200 --> 00:29:57.260 And that is how, just the extent
- 677 00:29:57.260 --> 00:29:59.518 to which prognostic heterogeneity plays a role
- $678\ 00:29:59.518 --> 00:30:02.730$  in the precision oncology space, or any space
- $679\ 00{:}30{:}02.730 {\:-->\:} 00{:}30{:}06.470$  where you're doing biomarker driven the rapeutics.
- $680\ 00{:}30{:}06.470 \dashrightarrow 00{:}30{:}09.240$  So what I'm showing you here is the cancer immunity cycle
- $681\ 00:30:09.240 \longrightarrow 00:30:11.020$  by Chen and Mellman.
- $682\ 00{:}30{:}11.020 \dashrightarrow 00{:}30{:}13.090$  So this diagram sort of revolutionized
- $683\ 00:30:13.090 --> 00:30:16.025$  how we think about how the immune system identifies
- $684\ 00:30:16.025 \longrightarrow 00:30:19.043$  and counteracts malignant cells.
- $685~00{:}30{:}20.880 \dashrightarrow 00{:}30{:}23.956$  So cancer cells release antigens.
- $686\ 00{:}30{:}23.956 {\:{\mbox{--}}}{>}\ 00{:}30{:}27.883$  They have to be detected by the immune system.
- 687 00:30:29.115 --> 00:30:31.680 If the immune system detects antigens,
- $688~00:30:31.680 \longrightarrow 00:30:33.370$  means your immune system is actually aware
- $689\ 00:30:33.370 \longrightarrow 00:30:35.000$  that you have cancer.
- $690\ 00:30:35.000 --> 00:30:38.550$  They have to produce natural killer cells.
- $691~00{:}30{:}38.550 \dashrightarrow 00{:}30{:}40.620$  So the T cells have to be produced in the lymph nodes.
- $692\ 00:30:40.620 \longrightarrow 00:30:43.250$  They have to infiltrate the tumor.
- $693\ 00:30:43.250 \dashrightarrow 00:30:45.900$  They have to recognize which cells are malignant cells.

 $694\ 00:30:45.900 \longrightarrow 00:30:48.350$  and then they have to kill the malignant cells.

 $695\ 00:30:48.350 \longrightarrow 00:30:50.650$  This process is very complicated

 $696\ 00:30:50.650 \longrightarrow 00:30:52.340$  and there are biomarkers

 $697\ 00:30:52.340 \longrightarrow 00:30:56.030$  that can tell us about what's happening with the patient.

698~00:30:56.030 --> 00:30:59.070 What's happening with the patient's innate immune response

 $699\ 00:30:59.070 \longrightarrow 00:31:00.700$  to cancer.

 $700\ 00{:}31{:}00.700 \dashrightarrow 00{:}31{:}04.360$  So the biomarkers that have been most developed recently

701 00:31:04.360 --> 00:31:08.240 are the PD-L1 biomarkers, which is this last step.

702 00:31:08.240 --> 00:31:09.870 So if a patient is expressing

703 00:31:09.870 --> 00:31:12.150 a lot of program death like in one,

 $704\ 00:31:12.150 \longrightarrow 00:31:13.920$  it means that the malignant cells

 $705\ 00:31:13.920 --> 00:31:16.390$  are actually hiding from the T cells.

 $706\ 00{:}31{:}16.390 \dashrightarrow 00{:}31{:}18.750$  So the patients might be producing lymphocytes.

 $707\ 00:31:18.750 \longrightarrow 00:31:22.100$  They might be getting to the tumor, but they can't attach.

 $708\ 00:31:22.100 \longrightarrow 00:31:24.620$  They can't identify which cells are malignant cells,

709 00:31:24.620 --> 00:31:26.490 malignant cells are hiding.

 $710\ 00:31:26.490 --> 00:31:28.140$  So there're very interesting things that happen

 $711\ 00:31:28.140 --> 00:31:30.333$  when you get to a biological perspective.

712 00:31:31.520 --> 00:31:34.490 The immune phenotypes based on these biomarkers.

713 00:31:34.490 --> 00:31:38.320 If we look at T-cell infiltration versus PD-L1 expression.

 $714\ 00:31:38.320 \longrightarrow 00:31:40.280$  Patients that are producing T cells

715 00:31:41.530 --> 00:31:43.720 and that have low PD-L1 expression.

716 00:31:43.720 --> 00:31:45.540 So that means T-cells are being produced,

 $717\ 00:31:45.540 \longrightarrow 00:31:47.250$  they're coming to the tumor

- 718 00:31:47.250 --> 00:31:49.580 and then they're effective when they get to the tumor.
- 719 00:31:49.580 --> 00:31:52.950 These patients have a different immune profile,
- $720\ 00:31:52.950 \longrightarrow 00:31:54.640$  than the opposite case
- 721 00:31:54.640 --> 00:31:57.190 where patients are not producing T cells.
- 722 00:31:57.190 --> 00:31:59.100 So it's like their immune system isn't aware
- $723\ 00:31:59.100 \longrightarrow 00:32:00.260$  that they have cancer.
- 724 00:32:00.260 --> 00:32:02.540 And then even if they did produce T cells,
- $725\ 00:32:02.540$  --> 00:32:05.320 they're not effective once they get to the tumor.
- $726\ 00:32:05.320 \longrightarrow 00:32:09.760$  So there's various things happening in this phase.
- 727 00:32:09.760 --> 00:32:12.450 And so now I go back to Professor Butte
- 728 00:32:13.360 --> 00:32:15.653 and sort of what he's saying,
- $729\ 00:32:16.630 --> 00:32:18.560$  So there's several articles that he's written
- $730\ 00:32:18.560 \longrightarrow 00:32:19.690$  that say things like this,
- $731\ 00:32:19.690 \longrightarrow 00:32:22.810$  precision medicine makes doctors nervous.
- $732\ 00:32:22.810 \longrightarrow 00:32:25.830$  And he says, the reason that makes doctors nervous
- $733\ 00:32:25.830 \longrightarrow 00:32:29.350$  is because they have to admit that what they were doing
- $734\ 00:32:29.350 \longrightarrow 00:32:31.043$  before was not precise.
- $735\ 00:32:31.900 \longrightarrow 00:32:35.245$  So we see these things
- $736\ 00:32:35.245 --> 00:32:37.395$  and we see these kinds of narratives coming
- $737\ 00:32:38.490 \longrightarrow 00:32:40.260$  from the group that's really pushing that
- $738\ 00:32:40.260 \longrightarrow 00:32:42.473$  we just need to analyze these databases.
- 739 00:32:43.860 --> 00:32:46.900 So he's talking about retroactive crowdsourcing, right?
- $740\ 00:32:46.900 \longrightarrow 00:32:49.050$  A high school kid can do it.
- 741 00:32:49.050 --> 00:32:50.270 So if you've listened to his talks,
- $742\ 00{:}32{:}50.270 --> 00{:}32{:}52.320$ he's always saying, a high school kid can do that.
- $743\ 00:32:52.320 \longrightarrow 00:32:53.900$  A high school kid could do this.

 $744\ 00:32:53.900 --> 00:32:57.550$  I think a high school kid could apply a T test to a dataset.

 $745\ 00:32:57.550 \longrightarrow 00:32:59.433$  I don't disagree with that.

746 00:33:00.500 --> 00:33:02.700 But I have a 14 year old at home

 $747\ 00:33:02.700 \longrightarrow 00:33:04.540$  and he has trouble making his bed.

 $748~00{:}33{:}04.540 \dashrightarrow 00{:}33{:}09.510$  So I think that there's a narrative out there

 $749\ 00:33:09.510 \longrightarrow 00:33:13.350$  that doesn't recognize things like this.

750 00:33:13.350 --> 00:33:15.730 So when I was at MD Anderson,

 $751\ 00:33:15.730 \longrightarrow 00:33:17.040$  we spent a lot of time thinking

 $752\ 00:33:17.040 \longrightarrow 00:33:19.040$  about these immune phenotypes.

753 00:33:19.040 --> 00:33:21.490 And I actually developed radiomics models,

 $754\ 00:33:21.490 \longrightarrow 00:33:23.550$  that characterized patterns

 $755\ 00:33:23.550 \longrightarrow 00:33:25.770$  that we saw in images in the tumor

 $756\ 00:33:25.770 \longrightarrow 00:33:28.790$  that reflected these immune phenotypes.

 $757\ 00:33:28.790 \longrightarrow 00:33:30.460$  And the reason we were doing that,

 $758\ 00{:}33{:}30.460 \dashrightarrow 00{:}33{:}33.210$  is because these biomarkers were incredibly unreliable.

759 00:33:34.600 --> 00:33:36.270 What I'm showing you here is a scatter plot,

 $760\ 00:33:36.270 --> 00:33:39.100$  that this came from the Garcia student's lab at MD Anderson,

761 00:33:39.100 --> 00:33:43.330 probably the best immune pathologists

 $762\ 00:33:43.330 \longrightarrow 00:33:45.203$  in the field right now.

763 00:33:46.280  $\rightarrow$  00:33:48.570 These are patients with non-small cell lung cancer.

764 00:33:48.570 --> 00:33:50.540 They all got treated with definitive surgery.

765 00:33:50.540 --> 00:33:52.500 So there was no chemotherapy.

766 00:33:52.500 --> 00:33:55.850 They came in, they could be treated with surgery.

 $767\ 00:33:55.850 \longrightarrow 00:33:58.830$  So we don't have sort of a confounding factor

 $768\ 00:33:58.830 --> 00:34:01.320$  of chemotherapy here with these patients.

769 00:34:01.320 --> 00:34:04.970 We got their tissue microarray staining,

 $770\ 00{:}34{:}04.970 {\:\hbox{--}}{>}\ 00{:}34{:}07.640$  and this was both malignant cells and immune cells,

- 771 00:34:07.640 --> 00:34:10.200 are PD-L1 positivity at biopsy.
- $772\ 00:34:10.200 \longrightarrow 00:34:12.180$  So the patients are coming in, they're getting a biopsy.
- 773  $00:34:12.180 \longrightarrow 00:34:13.820$  The biopsy is taking a needle,
- $774\ 00:34:13.820 \longrightarrow 00:34:16.130$  sticking it in a few different locations.
- $775\ 00:34:16.130 \longrightarrow 00:34:17.780$  We use that tissue and we try to assess
- 776 00:34:17.780 --> 00:34:19.050 how much PD-L1 expression
- 777 00:34:19.050 --> 00:34:21.470 do they have and their lung cancer?
- $778\ 00:34:21.470 \longrightarrow 00:34:23.780$  Then they go in, they had surgery.
- 779 00:34:23.780 --> 00:34:26.760 We took their whole excise tumor.
- $780\ 00:34:26.760 \longrightarrow 00:34:28.880$  And we went back and we did whole section staining,
- $781\ 00:34:28.880 --> 00:34:32.210$  of the excise tumor for PD-L1 expression.
- $782\ 00:34:32.210 \longrightarrow 00:34:33.850$  This is a scatterplot we got.
- $783\ 00:34:33.850 \longrightarrow 00:34:35.550$  So each point is the same patient.
- 784 00:34:36.600 --> 00:34:38.740 So this patient at biopsy,
- $785\ 00:34:38.740 \longrightarrow 00:34:40.090$  just this isn't the worst one,
- 786  $00:34:40.090 \longrightarrow 00:34:43.460$  but this patient at biopsy was over 50%.
- $787\ 00:34:43.460 \longrightarrow 00:34:45.693$  After surgery, they're only at 15%.
- $788\ 00:34:46.970 \longrightarrow 00:34:49.020$  This patient is much worse.
- $789\ 00:34:49.020 \longrightarrow 00:34:51.433$  So what's going on here?
- $790~00:34:52.370 \dashrightarrow 00:34:54.330$  Either the immune system is constantly changing
- 791 00:34:54.330 --> 00:34:56.533 and these biomarkers are not reproducible,
- 792 00:34:56.533 --> 00:34:59.500 in the sense that your state is changing,
- 793 00:34:59.500 --> 00:35:01.610 or when we stick that needle in
- 794 00:35:01.610 --> 00:35:04.920 and we take just a few points,
- $795\ 00:35:04.920 \longrightarrow 00:35:07.720$  we get a very different answer than when we do surgery.
- 796 00:35:07.720 --> 00:35:09.080 Of course, we have to use biopsy
- 797 00:35:09.080 --> 00:35:11.225 if we're gonna make a treatment selection.
- $798\ 00:35:11.225 \longrightarrow 00:35:13.760$  So this is problematic.

 $799\ 00:35:13.760 \longrightarrow 00:35:16.610$  So when I think about, you know, we just need databases,

800 00:35:16.610 --> 00:35:18.120 we don't have to understand the science

 $801\ 00:35:18.120 \longrightarrow 00:35:20.595$  and we can answer all these fundamental questions,

802 00:35:20.595 --> 00:35:22.093 I don't think it's true.

803 00:35:23.810 --> 00:35:25.900 You know, you have,

 $804\ 00:35:25.900 --> 00:35:28.520$  There's issues like this with every biomarker.

 $805\ 00:35:28.520 \longrightarrow 00:35:30.210$  The biomarkers have to be reproducible.

 $806\ 00:35:30.210 \longrightarrow 00:35:32.683$  We have to understand them in a rigorous manner,

807 00:35:33.580 --> 00:35:35.463 if you're going to use scanning data.

808 00:35:37.080 --> 00:35:38.020 So, you know,

 $809\ 00:35:38.020$  --> 00:35:40.190 so we've published this paper in scientific reports.

810 00:35:40.190  $\rightarrow$  00:35:42.150 It has been cited I think almost a hundred times

 $811\ 00:35:42.150 \longrightarrow 00:35:43.260$  in a few years.

812 00:35:43.260 --> 00:35:45.310 Where we actually developed a radiomics model

 $813\ 00:35:45.310 \longrightarrow 00:35:49.010$  for understanding the immune pathology.

 $814\ 00:35:49.010 \longrightarrow 00:35:50.620$  Now, why did we do that?

 $815\ 00:35:50.620 --> 00:35:51.810$  We did that because we didn't think

 $816\ 00:35:51.810 \longrightarrow 00:35:53.760$  these biopsy assessments were reliable.

 $817\ 00:35:54.700 \longrightarrow 00:35:57.230$  So we thought that maybe the scans were more reliable.

 $818\ 00:35:57.230 \longrightarrow 00:35:58.320$  Maybe we could take the scans

 $819~00{:}35{:}58.320 \rightarrow 00{:}36{:}00.550$  and we can understand the patterns in the scans.

820 00:36:00.550 --> 00:36:02.570 And you can see that patients

821 00:36:02.570 --> 00:36:03.980 with different immune phenotypes,

822 00:36:03.980 --> 00:36:06.550 but in terms of T-cell infiltration and PD-L1,

 $823\ 00{:}36{:}06.550 {\:{\mbox{--}}}{>} 00{:}36{:}09.490$  they had very different expectations for survival.

- $824\ 00:36:09.490 \longrightarrow 00:36:11.930$  So this is not a treatment effect.
- $825\ 00:36:11.930 \longrightarrow 00:36:14.800$  This is just simply the impact
- $826\ 00:36:14.800 \longrightarrow 00:36:17.470$  of the fact that the patients have different immune systems.
- 827 00:36:17.470 --> 00:36:22.210 And those immune systems have differential effectiveness
- $828\ 00:36:22.210 \longrightarrow 00:36:24.463$  in fighting the tumor.
- 829 00:36:25.340  $\rightarrow$  00:36:28.440 So patients that have T-cells and low PD-L1 positivity,
- $830\ 00:36:28.440 \longrightarrow 00:36:30.060$  they're doing well.
- 831 00:36:30.060 --> 00:36:31.620 The opposite is true for patients
- 832 00:36:31.620 --> 00:36:33.723 that have high PD-L1 and low T cells.
- 833 00:36:34.840 --> 00:36:37.640 So we developed a radiomics model,
- $834\ 00:36:37.640 \longrightarrow 00:36:41.040$  which take the scans and actually assess these patterns.
- $835\ 00:36:41.040 \longrightarrow 00:36:44.513$  Of course, there's complications with that.
- 836 00:36:45.590 --> 00:36:48.080 If you're to scan any data in oncology,
- $837\ 00:36:48.080 --> 00:36:49.600$  you're probably having contrast.
- $838\ 00:36:49.600 \longrightarrow 00:36:51.330$  You need to understand what the protocol
- $839\ 00:36:51.330 \longrightarrow 00:36:54.050$  for contrast was for that scan.
- $840\ 00:36:54.050 \longrightarrow 00:36:56.610$  Because you need to take the image
- 841 00:36:56.610 --> 00:36:59.020 when the contrast is in the tumor.
- 842 00:36:59.020 --> 00:37:00.710 So of course you can't just go blindly
- $843\ 00:37:00.710 \longrightarrow 00:37:03.121$  and grab a bunch of images from a database.
- 844 00:37:03.121 --> 00:37:05.690 So, I've talked a little bit about
- 845 00:37:05.690 --> 00:37:07.320 what's happening on precision oncology.
- 846 00:37:07.320 --> 00:37:08.780 Where we're developing biomarkers,
- $847\ 00:37:08.780 \longrightarrow 00:37:10.410$  we want to use to guide treatment,
- $848\ 00:37:10.410 \longrightarrow 00:37:12.000$  but it's very complicated.
- 849 00:37:12.000 --> 00:37:13.390 And I don't think doctors are scared
- 850 00:37:13.390 --> 00:37:14.480 because they're not precise,
- $851\ 00:37:14.480 \longrightarrow 00:37:16.183$  they're scared because we need to understand

 $852\ 00:37:16.183 \longrightarrow 00:37:17.440$  that these biomarkers

 $853\ 00:37:17.440$  --> 00:37:20.420 and make sure they're reliable and reproducible.

 $854\ 00:37:20.420 \longrightarrow 00:37:22.203$  And that knowledge is important.

 $855\ 00:37:23.760 \longrightarrow 00:37:26.480$  Not only that, but because of all this complexity,

 $856\ 00:37:26.480$  --> 00:37:29.060 drug development on oncology has changed a lot.

 $857\ 00:37:29.060 \longrightarrow 00:37:33.117$  And we no longer have this, phase one to phase two.

 $858\ 00:37:33.117 --> 00:37:37.677$  This is what early phase drug trials look like now,

 $859\ 00:37:37.677 \longrightarrow 00:37:39.420$  especially for the big companies

 $860\ 00:37:39.420 --> 00:37:41.170$  that have a lot of money to invest.

 $861\ 00:37:42.340 \longrightarrow 00:37:45.370$  They're taking multiple dose levels from dose expansion,

 $862\ 00{:}37{:}45.370 \dashrightarrow 00{:}37{:}48.400$  they're running massive dose expansion cohorts.

 $863\ 00:37:48.400 \longrightarrow 00:37:49.520$  Those dose expansion cohorts,

864 00:37:49.520 --> 00:37:53.780 usually span multiple tumor types.

 $865\ 00:37:53.780 \longrightarrow 00:37:55.770$  And they might randomize across dose level,

 $866\ 00:37:55.770 \longrightarrow 00:37:57.050$  because we don't have

867 00:37:57.050 --> 00:37:59.550 these very clear monotonic relationships

868 00:37:59.550 --> 00:38:01.660 between dose and toxicity anymore.

 $869\ 00:38:01.660 --> 00:38:04.540$  And selecting a dose isn't as simple as it used to be

 $870~00:38:04.540 \longrightarrow 00:38:06.720$  when we did cytotoxic drug development.

 $871\ 00:38:06.720 \longrightarrow 00:38:08.900$  So these non cytotoxic targeted therapies,

 $872\ 00:38:08.900 \longrightarrow 00:38:10.850$  it's hard to select a dose.

 $873\ 00:38:10.850 --> 00:38:13.823$  These dose expansion cohorts can be hundreds of patients.

 $874\ 00:38:14.900 \longrightarrow 00:38:17.260$  They may not even stop for a phase two trial.

875 00:38:17.260 --> 00:38:19.360 They may go straight to phase two

 $876\ 00:38:19.360 \longrightarrow 00:38:21.570$  and expand on the expansion.

- 877 00:38:21.570 --> 00:38:23.430 Or they may skip phase two altogether
- $878\ 00:38:23.430$  --> 00:38:25.380 because they've already acquired so much information
- $879\ 00:38:25.380 \longrightarrow 00:38:27.130$  in their phase one trial.
- 880 00:38:27.130 --> 00:38:28.710 So this is what we see happening now.
- $881\ 00:38:28.710 --> 00:38:32.100$  Of course, the key note trial evaluated in Pembrolizumab
- $882\ 00:38:32.100 --> 00:38:33.680$  had eight expansion cohorts.
- $883\ 00:38:33.680 --> 00:38:35.140$  There was over a thousand patients
- 884 00:38:35.140 --> 00:38:37.860 in this first in human phase one trial.
- $885\ 00:38:37.860 \longrightarrow 00:38:39.240$  This trial is what motivated
- 886 00:38:39.240 --> 00:38:42.380 that NCI Clinical Trial Design Task Force,
- $887\ 00:38:42.380 \longrightarrow 00:38:43.213$  that I got to be a part of,
- 888 00:38:43.213 --> 00:38:45.870 because this was a massive departure
- 889 00:38:45.870 --> 00:38:48.180 from what we saw typically in oncology
- $890~00:38:48.180 \dashrightarrow 00:38:52.190$  and how IRBs would review these studies.
- 891 00:38:52.190 --> 00:38:55.950 More recently, Genentech drug (indistinct)
- $892\ 00:38:55.950 \longrightarrow 00:38:59.160$  had a phase one trial with nine expansion cohort.
- 893 00:38:59.160 --> 00:39:00.630 Looking at the dose, expansions alone,
- 894 00:39:00.630 --> 00:39:03.290 the bladder cancer cohort had 97 patients,
- $895\ 00:39:03.290 \longrightarrow 00:39:05.610$  and they randomized the three dose levels.
- $896\ 00:39:05.610 \longrightarrow 00:39:06.863$  So this is a new world.
- $897\ 00:39:08.510 \longrightarrow 00:39:11.990\ 97$  patients already in their dose expansion.
- $898~00:39:11.990 \longrightarrow 00:39:14.430$  So this is where Master Protocols come in.
- $899\ 00:39:14.430 \longrightarrow 00:39:16.630$  So we have innovations in design
- $900\ 00:39:16.630 \longrightarrow 00:39:18.000$  that are sort of targeting this
- 901 00:39:18.000 --> 00:39:22.787 and there's many, many methodology recommendations.
- 902 00:39:23.730 --> 00:39:24.890 The other thing that's happened in oncology
- $903\ 00:39:24.890 \longrightarrow 00:39:28.060$  is that phase three continues to be poor.
- 904 00:39:28.060 --> 00:39:29.590 So phase three trials continue

- $905\ 00:39:29.590 \longrightarrow 00:39:31.490$  to have a poor track record relative
- $906\ 00:39:31.490 \longrightarrow 00:39:34.040$  to other areas of medicine.
- $907\ 00:39:34.040 \longrightarrow 00:39:36.490$  You can see lots of articles that described this.
- $908\ 00:39:37.860 \longrightarrow 00:39:40.040$  Of course, Gan et al did a review
- 909 00:39:40.040 --> 00:39:43.010 of 235 published randomized controlled trials.
- 910 00:39:43.010 --> 00:39:46.410 Regulatory approval was, you know, less than 38%.
- 911 00:39:46.410 --> 00:39:47.940 And what's happening?
- $912\ 00:39:47.940 --> 00:39:49.730$  While the investigators are not very good
- $913\ 00:39:49.730 \longrightarrow 00:39:52.886$  about making the assumptions for that phase three trial,
- $914\ 00:39:52.886 --> 00:39:55.310$  we see a lot of phase three trials in oncology
- 915 00:39:55.310 --> 00:39:57.463 that have unrealistic expectations.
- 916 00:39:58.990 --> 00:40:02.180 Okay, so now I talked about precision oncology.
- $917\ 00:40:02.180 \longrightarrow 00:40:03.690$  I'm gonna go into some case studies
- 918 00:40:03.690 --> 00:40:05.540 that I think are interesting.
- 919 00:40:05.540 --> 00:40:08.600 And I want you ask the question,
- 920 00:40:08.600 --> 00:40:11.060 how could you have used real-world evidence
- 921 00:40:11.060 --> 00:40:12.563 to change what happens here?
- 922 00:40:13.610  $\rightarrow$  00:40:14.670 So this is coming at it
- 923 00:40:14.670  $\rightarrow$  00:40:15.937 from, these are the high profile trials
- $924\ 00:40:15.937 \longrightarrow 00:40:20.210$  that we have been running in the last few years in oncology.
- 925 00:40:20.210 --> 00:40:21.370 We want to know,
- 926 00:40:21.370 --> 00:40:22.900 how could we have used real-world evidence
- $927\ 00:40:22.900 \longrightarrow 00:40:24.453$  in these settings?
- 928 00:40:25.660 --> 00:40:28.090 So I'm gonna talk about Atezolizumab
- $929\ 00:40:28.090 \longrightarrow 00:40:30.300$  and bladder cancer.
- 930 00:40:30.300 --> 00:40:33.290 So Atezolizumab is another PD-1 inhibitor.
- $931\ 00:40:33.290 --> 00:40:36.283$  So immunotherapy, similar to Pembrolizumab.
- 932 00:40:37.670 --> 00:40:39.820 So it was developed for many different areas.

- 933 00:40:39.820 --> 00:40:41.750 Again, we're talking about tissue-agnostic here.
- 934 00:40:41.750  $\rightarrow$  00:40:45.440 So it's targeting a feature of the immune system,
- $935\ 00:40:45.440 --> 00:40:47.140$  that feature of the immune system can exist
- 936 00:40:47.140 --> 00:40:49.130 across many different tumor types.
- $937\ 00:40:49.130 --> 00:40:51.880$  They evaluated nine in their phase one trial.
- 938 00:40:51.880 --> 00:40:53.177 So after the phase one trial,
- 939 00:40:53.177 --> 00:40:57.030 they ran a bunch of trials and different types of cancers
- 940 00:40:57.030 --> 00:40:58.620 and different lines of therapy.
- 941 00:40:58.620  $\rightarrow$  00:41:01.540 One of them was second-line bladder cancer.
- 942 00:41:01.540 --> 00:41:03.340 So these are patients with bladder cancer
- $943\ 00:41:03.340 \longrightarrow 00:41:06.020$  that have progressed on a prior therapy.
- 944 00:41:06.020 --> 00:41:09.030 So they already progressed on chemotherapy,
- 945 00:41:09.030 --> 00:41:11.130 now they're getting this immunotherapy.
- $946\ 00{:}41{:}11.130 \dashrightarrow 00{:}41{:}15.010$  So they ran this study and the biomarker they're targeting
- $947\ 00:41:15.010 \longrightarrow 00:41:18.080$  is they're calling IC2/3.
- 948 00:41:18.080 --> 00:41:21.130 That is immune cell staining of PD-L1.
- 949 00:41:21.130 --> 00:41:24.864 And those immune cells have 5% or more expression.
- 950 00:41:24.864 --> 00:41:29.864 So 5% of the immune cells that they stained had,
- 951 00:41:29.973 --> 00:41:34.083 at least 5% had Programmed Death Ligand  $^{\rm 1}$
- 952 00:41:34.083 --> 00:41:36.260 That's their target.
- 953 00:41:36.260 --> 00:41:38.160 So, but they enrolled in this phase two trial,
- $954\ 00:41:38.160 \longrightarrow 00:41:39.450$  they enrolled all comers.
- $955\ 00:41:39.450 \longrightarrow 00:41:41.200$  It wasn't restricted to the target.
- $956\ 00:41:42.180 \longrightarrow 00:41:43.070$  They enrolled all comers.
- $957\ 00:41:43.070 \longrightarrow 00:41:45.190$  So the IC2/3 population is their target.

- $958\ 00{:}41{:}45.190 \dashrightarrow 00{:}41{:}47.950$  That's where the mechanism is supposed to work.
- 959 00:41:47.950 --> 00:41:49.420 So among a hundred patients
- $960\ 00:41:49.420 \longrightarrow 00:41:52.930$  with that target they got a 26% response rate.
- 961 00:41:52.930 --> 00:41:54.840 You can see patients that don't have the target,
- $962\ 00:41:54.840 \longrightarrow 00:41:57.020$  there was 11 and there was eight.
- 963 00:41:57.020 --> 00:41:58.790 And if you look back at their paper,
- $964\ 00:41:58.790 \longrightarrow 00:42:01.580$  they told us that they expected 10%.
- 965 00:42:01.580 --> 00:42:04.220 So they said the null hypothesis was 10%
- $966\ 00:42:04.220 \longrightarrow 00:42:05.220$  for this population.
- 967 00:42:05.220  $\rightarrow$  00:42:06.550 We got 26%.
- 968 00:42:06.550 --> 00:42:08.753 This is very exciting, right?
- 969 00:42:09.940 --> 00:42:11.760 This is the survival curves that they present
- 970 00:42:11.760 --> 00:42:13.110 from their phase two trial.
- 971 00:42:13.110 --> 00:42:15.020 Again, this is uncontrolled.
- $972\ 00:42:15.020 \longrightarrow 00:42:17.100$  There's no chemotherapy arm here.
- 973 00:42:17.100 --> 00:42:20.570 This is just the treated arm, Atezolizumub
- $974\ 00:42:20.570 \longrightarrow 00:42:22.690$  by biomarker status.
- 975 00:42:22.690 --> 00:42:23.580 And when you look at this,
- $976\ 00:42:23.580 \longrightarrow 00:42:25.210$  you see this blue Kaplan-Meier curve,
- $977\ 00:42:25.210 \longrightarrow 00:42:26.950$  that's above everybody else.
- 978 00:42:26.950 --> 00:42:29.010 That Kaplan-Meier curve is the target feature.
- 979  $00:42:29.010 \longrightarrow 00:42:31.020$  That's the IC2/3 population.
- 980 00:42:31.020 --> 00:42:32.150 So they're responding,
- 981 00:42:32.150 --> 00:42:34.780 their tumors are shrinking and they're living longer.
- 982 00:42:34.780 --> 00:42:37.600 It looks like this is very promising, right?
- 983 00:42:37.600 --> 00:42:39.890 On the basis of that, they got accelerated approval.
- $984\ 00:42:39.890 \longrightarrow 00:42:42.240$  And that was given in 2016.

985 00:42:42.240 --> 00:42:45.400 And the reason was increased levels of PD-L1 expression

 $986\ 00:42:45.400$  --> 00:42:48.150 on immune cells are associated with increased response.

 $987\ 00:42:49.190 \longrightarrow 00:42:51.370$  Let's go to the phase three trial.

988 00:42:51.370 --> 00:42:53.500 So as a part of the conditional approval

 $989\ 00:42:53.500 \longrightarrow 00:42:54.520$  with accelerated approval,

990 00:42:54.520 --> 00:42:56.287 they have to run a randomized phase three trial

991 00:42:56.287 --> 00:42:58.890 and sort of replicate this result.

992 00:42:58.890 --> 00:43:01.570 So they designed this trial, IMvigor211,

993 00:43:01.570 --> 00:43:04.030 multi-center open-label phase three trial.

994 00:43:04.030 --> 00:43:05.900 They compared to three chemotherapies,

995 00:43:05.900  $\rightarrow$  00:43:07.950 which were standard chemotherapies used at the time.

996  $00:43:07.950 \longrightarrow 00:43:10.550$  So there was a physician's choice.

997 00:43:10.550 --> 00:43:12.230 If the patient was randomized to chemotherapy,

998 00:43:12.230 --> 00:43:13.860 the physician would choose

999 00:43:13.860 --> 00:43:16.300 which among these three chemotherapies.

 $1000\ 00:43:16.300 \longrightarrow 00:43:18.000$  So what happened?

1001 00:43:18.000 --> 00:43:20.860 We had this blockbuster results in phase two,

 $1002\ 00:43:20.860 \longrightarrow 00:43:22.040$  but there was no difference

 $1003\ 00:43:22.040 \longrightarrow 00:43:24.230$  in overall survival in phase three.

 $1004\ 00:43:24.230 \longrightarrow 00:43:26.340$  Not only was there not a difference in overall survival,

 $1005\ 00:43:26.340 \longrightarrow 00:43:28.100$  the objective response rates were similar.

 $1006\ 00:43:28.100 --> 00:43:30.240$  So the tumor responses were similar.

1007 00:43:30.240 --> 00:43:33.280 Moreover they enrolled 931 patients

 $1008\ 00:43:33.280 \longrightarrow 00:43:35.723$  and only 234 actually had the target.

 $1009~00{:}43{:}36.863 --> 00{:}43{:}41.863$  So 24% of the trial was used for the primary analysis.

 $1010\ 00:43:43.110 \longrightarrow 00:43:45.750$  When we look at the data, what happened?

- $1011\ 00:43:45.750 \longrightarrow 00:43:49.180\ 23\%$  of the IC2/3 population responded.
- $1012\ 00:43:49.180 \longrightarrow 00:43:50.740$  So that's close to 26%.
- $1013\ 00:43:50.740 \longrightarrow 00:43:52.740$  It looks like that was replicated.
- $1014\ 00:43:52.740 \longrightarrow 00:43:54.800$  When you look at the intention to treat populations,
- $1015\ 00{:}43{:}54.800$  -->  $00{:}43{:}57.370$  that's everybody here, regardless of biomarker,
- $1016\ 00:43:57.370 \longrightarrow 00:43:59.820$  it's 13 and 13.
- $1017~00{:}43{:}59.820 \dashrightarrow 00{:}44{:}02.180$  So it was also lower without the target.
- $1018~00{:}44{:}02.180 \dashrightarrow 00{:}44{:}05.160$  But what's happening with chemotherapy with the target?
- 1019 00:44:05.160 --> 00:44:07.420 It's 22\%, right?
- $1020\ 00{:}44{:}07.420$  -->  $00{:}44{:}11.160$  So chemotherapy is doing great with this biomarker.
- $1021\ 00:44:11.160 \longrightarrow 00:44:13.450$  So this biomarker profile
- $1022\ 00:44:13.450 \longrightarrow 00:44:17.180$  is doing just as well as the targeted therapy,
- $1023\ 00:44:17.180 \longrightarrow 00:44:19.330$  when the patients get the standard of care.
- $1024\ 00:44:20.660 \longrightarrow 00:44:23.240$  Here's the survival curve.
- $1025\ 00{:}44{:}23.240 \dashrightarrow 00{:}44{:}26.020$  Okay, proportional hazards is probably violated.
- 1026 00:44:26.020 --> 00:44:28.210 There is a heavy tail here for the Atezo group.
- $1027\ 00:44:28.210 \longrightarrow 00:44:29.450$  Maybe there's, it looks like
- 1028 00:44:29.450 --> 00:44:31.610 there's some long-term stable disease,
- 1029 00:44:31.610 --> 00:44:33.531 people that are benefiting.
- 1030 00:44:33.531 --> 00:44:36.190 But overall, this is not significant.
- 1031 00:44:36.190 --> 00:44:39.220 And on the basis of this, actually this year,
- $1032\ 00{:}44{:}39.220 \dashrightarrow 00{:}44{:}43.810$  this drug was with drawn from accelerated approval.
- $1033\ 00:44:43.810 \longrightarrow 00:44:45.930$  So it got the accelerated approval,
- $1034\ 00:44:45.930 --> 00:44:47.740$  which was for very exciting drugs
- $1035\ 00{:}44{:}47.740 \dashrightarrow 00{:}44{:}52.150$  that need an accelerated pathway for regulatory.
- 1036 00:44:52.150 --> 00:44:53.670 And then this phase three,

- $1037\ 00:44:53.670 \longrightarrow 00:44:55.020$  on the basis of this phase three trial,
- $1038\ 00:44:55.020 \longrightarrow 00:44:56.523$  they had to withdraw from that.
- $1039\ 00:44:57.380 \longrightarrow 00:44:58.803$  So the question is,
- $1040\ 00:44:59.740 \longrightarrow 00:45:02.890$  how do we use real-world evidence to change this?
- $1041\ 00:45:02.890 \longrightarrow 00:45:05.423$  At the end, there were flaws in this design.
- $1042~00{:}45{:}07.940 \dashrightarrow 00{:}45{:}11.103$  They didn't understand the biomarker.
- 1043 00:45:12.550 --> 00:45:14.280 They didn't understand the biomarker profile
- $1044\ 00:45:14.280 \longrightarrow 00:45:16.650$  on the basis of the standard of care.
- $1045\ 00:45:16.650 --> 00:45:19.880$  So when I first got involved in sort of,
- $1046~00{:}45{:}19.880 \dashrightarrow 00{:}45{:}22.610$  well, over this past year, I've been thinking about
- 1047 00:45:22.610 --> 00:45:23.887 how could we have used real-world evidence?
- 1048 00:45:23.887 --> 00:45:26.720 Here's the case where, you know, there's,
- 1049 00:45:26.720 --> 00:45:29.830 it's kind of a failure of the system here
- $1050\ 00{:}45{:}29.830 --> 00{:}45{:}32.320$  that we had this drug with drawn from accelerated approval.
- 1051 00:45:32.320 --> 00:45:34.550 And it's not the only one, by the way.
- $1052\ 00:45:34.550 \longrightarrow 00:45:38.210$  Is there something in the historical data
- $1053\ 00{:}45{:}38.210 \dashrightarrow 00{:}45{:}40.700$  or the real-world data that we could have used
- $1054\ 00:45:40.700 \longrightarrow 00:45:44.910$  that could have informed us to design a better trial,
- 1055 00:45:44.910 --> 00:45:46.700 or could have told us something
- $1056\ 00:45:46.700 \longrightarrow 00:45:50.033$  about the fact that this biomarker may be prognostic?
- $1057\ 00:45:51.290 --> 00:45:52.960$  Now it gets complicated
- $1058\ 00{:}45{:}52.960 {\: -->\:} 00{:}45{:}56.040$  because actually it's not prognostic for surgery.
- $1059~00{:}45{:}56.040 \dashrightarrow 00{:}45{:}59.340$  Patients that have surgery that have IC2/3 status,
- $1060\ 00:45:59.340 \longrightarrow 00:46:01.730$  they're going to die sooner
- 1061 00:46:01.730 --> 00:46:05.030 than patients that have IC1, IC0,

 $1062\ 00{:}46{:}05.030 {\: -->\:} 00{:}46{:}07.840$  So this marker seems to be a predictive marker

 $1063\ 00:46:07.840 \longrightarrow 00:46:10.930$  for both chemotherapy and for Atezo.

 $1064\ 00:46:10.930 \longrightarrow 00:46:12.330$  So, but we didn't know.

 $1065\ 00:46:12.330 \longrightarrow 00:46:13.550\ I\ didn't\ know\ if\ that\ was\ true.$ 

 $1066~00{:}46{:}13.550 \dashrightarrow 00{:}46{:}16.850$  So my postdoc and I went back and we did a meta-analysis.

1067 00:46:16.850 --> 00:46:20.090 We went and we extracted all of the trials

 $1068\ 00{:}46{:}20.090 \dashrightarrow 00{:}46{:}23.480$  that had enrolled second-line bladder cancer patients

 $1069\ 00:46:23.480 \longrightarrow 00:46:25.150$  in a prospective study

1070 00:46:25.150 --> 00:46:26.880 that evaluated the three chemotherapies

 $1071\ 00:46:26.880 \longrightarrow 00:46:29.160$  that were used in the control arm.

 $1072\ 00:46:29.160 \longrightarrow 00:46:30.980$  So those are given here.

 $1073\ 00:46:30.980 \longrightarrow 00:46:34.590$  So I think back to Dr. Butte saying, you know,

 $1074\ 00:46:34.590 \longrightarrow 00:46:35.810$  the real problem in research

 $1075~00{:}46{:}35.810 \dashrightarrow 00{:}46{:}38.685$  is you don't have enough people asking questions.

 $1076\ 00:46:38.685 \longrightarrow 00:46:42.894$  When we did this literature search,

 $1077\ 00:46:42.894 \longrightarrow 00:46:46.623$  there were like 200 papers on second-line bladder cancer,

 $1078\ 00:46:48.020 --> 00:46:50.770$  Most of them were retrospective reviews and case studies.

 $1079\ 00:46:50.770 \longrightarrow 00:46:52.430$  the overwhelming majority.

 $1080\ 00{:}46{:}52.430 {\: --> \:} 00{:}46{:}57.400$  There were only 11 that were actual perspective studies

 $1081\ 00:46:57.400 \longrightarrow 00:46:59.060$  that we could use in this population.

1082 00:46:59.060 --> 00:47:01.080 So there's a lot of people writing papers

1083 00:47:01.080 --> 00:47:03.610 on retrospective databases, there's lots,

 $1084\ 00{:}47{:}03.610 \dashrightarrow 00{:}47{:}06.916$  but what we actually need are prospective studies.

 $1085\ 00:47:06.916 \longrightarrow 00:47:10.000$  So we see here, we have these 11 trials.

 $1086\ 00:47:10.000 \longrightarrow 00:47:11.730$  We're looking at the overall response rate

- $1087\ 00:47:11.730 \longrightarrow 00:47:12.680$  from these 11 trials,
- 1088 00:47:12.680 --> 00:47:14.813 we're doing a standard meta-analysis.
- 1089~00:47:15.690 --> 00:47:20.150 You can see that Genentech said 10% was their null, right?
- $1090\ 00:47:20.150 \longrightarrow 00:47:22.050$  And really the case for real-world evidence
- $1091\ 00{:}47{:}22.050 \dashrightarrow 00{:}47{:}25.050$  is you can do a better job specifying your null hypothesis.
- 1092 00:47:25.050 --> 00:47:26.770 Your null hypothesis can be specified better
- $1093\ 00:47:26.770 --> 00:47:29.800$  because you know what to expect for control.
- $1094\ 00{:}47{:}29.800 \dashrightarrow 00{:}47{:}33.340$  So based on our meta-analysis of the objective response,
- $1095\ 00:47:33.340 \longrightarrow 00:47:35.390\ 10\%$  is really good estimate.
- $1096~00{:}47{:}35.390 \dashrightarrow 00{:}47{:}38.950$  And 10% is like the hierarchal mean of this meta-analysis.
- $1097\ 00:47:38.950 \longrightarrow 00:47:40.710$  So now we go to the chemotherapy arms
- $1098\ 00:47:40.710 \longrightarrow 00:47:43.167$  that we saw in the Atezo trial.
- $1099\ 00:47:43.167 \longrightarrow 00:47:46.490$  We see the ICO/1 population is right at 10%.
- 1100 00:47:46.490 --> 00:47:47.910 But look at this,
- $1101\ 00:47:47.910 \longrightarrow 00:47:49.660$  this IC2/3 population.
- $1102\ 00:47:49.660 \longrightarrow 00:47:52.090$  Again, this is with chemotherapy.
- $1103\ 00:47:52.090 \longrightarrow 00:47:54.770$  They're statistically significantly better
- $1104\ 00:47:54.770 \longrightarrow 00:47:58.160$  than the hierarchal mean that we estimated
- $1105\ 00:47:58.160 \longrightarrow 00:48:00.180$  from meta-analysis.
- $1106\ 00:48:00.180 \longrightarrow 00:48:01.330$  So what does this mean?
- $1107\ 00:48:02.460 --> 00:48:05.453$  This means that this profile has not been studied before.
- $1108\ 00{:}48{:}06.700 \dashrightarrow 00{:}48{:}11.300$  These trials are mixtures of different immune phenotypes.
- $1109\ 00{:}48{:}11.300 --> 00{:}48{:}14.412$  So we don't know which mean phenotype they're studying.
- $1110\ 00:48:14.412 \longrightarrow 00:48:16.820$  They have a different distribution.
- 1111 00:48:16.820 --> 00:48:19.500 Maybe this one has more IC2/3 population
- $1112\ 00:48:19.500 \longrightarrow 00:48:20.750$  because it's pulled over.

- $1113\ 00:48:22.260 \longrightarrow 00:48:24.640$  But the reality is the information
- $1114\ 00:48:24.640 \longrightarrow 00:48:26.510$  in these historical studies
- 1115 00:48:26.510 --> 00:48:28.843 doesn't tell us about immune staining.
- $1116\ 00:48:29.980 \longrightarrow 00:48:33.083$  So this is a biomarker that wasn't studied before.
- 1117 00:48:34.060 --> 00:48:35.540 And certainly that's goNNA be the case
- $1118\ 00:48:35.540 \longrightarrow 00:48:37.450$  in the community databases.
- 1119 00:48:37.450 --> 00:48:39.230 Because there's only a few institutions
- 1120 00:48:39.230 --> 00:48:41.590 that really can have the infrastructure
- 1121 00:48:41.590 --> 00:48:45.420 to quickly stain these patients
- $1122\ 00:48:45.420 \longrightarrow 00:48:47.363$  as these biomarkers are developing.
- 1123 00:48:48.410 --> 00:48:51.720 So we extracted the Kaplan-Meier curves
- $1124\ 00:48:51.720 \longrightarrow 00:48:53.220$  from these historical studies.
- $1125\ 00{:}48{:}54.210 \dashrightarrow 00{:}48{:}57.370$  And we did a meta-analysis of these Kaplan-Meier curves.
- 1126 00:48:57.370 --> 00:48:58.340 And that's given here,
- $1127\ 00:48:58.340 \longrightarrow 00:49:01.980$  we have a piece-wise exponential model and a Weibull model.
- $1128\ 00{:}49{:}01.980 \dashrightarrow 00{:}49{:}04.010$  When we put the Kaplan-Meier curves together
- $1129\ 00:49:04.010 \longrightarrow 00:49:05.053$  with the survival.
- $1130\ 00:49:05.890 \longrightarrow 00:49:07.420$  Oh, sorry, when you put the overall response
- $1131\ 00:49:07.420 \longrightarrow 00:49:08.290$  with the survival data,
- $1132\ 00{:}49{:}08.290 \dashrightarrow 00{:}49{:}11.970$  we see this purple line is the chemotherapy arm
- $1133\ 00:49:11.970 --> 00:49:16.630$  with this targeted biomarker from the phase three study,
- $1134\ 00:49:16.630 \longrightarrow 00:49:18.400$  that was implemented by Genentech.
- $1135\ 00:49:18.400 --> 00:49:22.730$  So responses is better, survival is significantly better
- $1136\ 00:49:22.730 \longrightarrow 00:49:26.317$  than what our expectation was based on historical evidence.
- $1137\ 00{:}49{:}26.317 \dashrightarrow 00{:}49{:}29.270$  And we actually went back and did simulation studies

- $1138\ 00:49:29.270$  --> 00:49:32.260 where we fit piece-wise exponential and Weibull curves,
- 1139 00:49:32.260 --> 00:49:33.463 till all of these Kaplan-Meier curves
- $1140\ 00:49:33.463 \longrightarrow 00:49:36.568$  that we extracted from the web digitize the tool.
- $1141\ 00:49:36.568 \longrightarrow 00:49:37.920$  When we actually simulated,
- $1142\ 00:49:37.920 \longrightarrow 00:49:39.850$  was it the probability of success
- $1143\ 00:49:39.850 \longrightarrow 00:49:41.343$  for the design implements?
- $1144\ 00:49:42.287 --> 00:49:44.680$  And we looked at that for the PDL-1 population,
- $1145\ 00:49:44.680 \longrightarrow 00:49:46.060$  as well as the ITT population.
- 1146 00:49:46.060 --> 00:49:48.960 We only give this trial 20% chance of success
- $1147\ 00:49:48.960 \longrightarrow 00:49:52.260$  based on the extent to which chemo is interacting
- 1148 00:49:52.260 --> 00:49:53.630 with PD-L1.
- $1149\ 00{:}49{:}53.630 {\: \mbox{--}}{\:>} 00{:}49{:}55.900$  If you like, if you wanna account for the heavy tail
- $1150\ 00:49:55.900 \longrightarrow 00:49:58.410$  that we see in the model, and do piece of exponential,
- $1151\ 00:49:58.410 \longrightarrow 00:50:00.230$  it goes up to 24%.
- $1152\ 00:50:00.230 \longrightarrow 00:50:02.280$  So another case of a phase three trial
- $1153\ 00{:}50{:}02.280$  -->  $00{:}50{:}05.253$  there was, had unrealistic expectations, right?
- $1154\ 00:50:06.140 \longrightarrow 00:50:07.730$  And it's a case
- $1155\ 00:50:07.730 \longrightarrow 00:50:11.090$  where we didn't understand the biomarker profile.
- $1156\ 00:50:11.090 --> 00:50:14.010$  That biomarker profile had not been characterized
- $1157\ 00:50:14.010 \longrightarrow 00:50:16.220$  in the historical evidence.
- $1158\ 00:50:16.220$  --> 00:50:18.640 It's not only Atezolizumab, this happened to Durvalumab.
- $1159\ 00:50:19.584$  --> 00:50:22.340 It happened in bladder cancer for Durvalumab again as well.
- $1160\ 00:50:22.340 --> 00:50:25.393$  Also a PD-1 inhibitor from AstraZeneca.

- 1161 00:50:30.664 --> 00:50:33.747 So Precision Oncology is hard, right?
- $1162\ 00:50:34.610 \longrightarrow 00:50:35.443$  It's hard.
- 1163 00:50:35.443 --> 00:50:39.210 It's not, what I presented here,
- $1164\ 00:50:39.210 \longrightarrow 00:50:43.750$  was not really about the lack of having information,
- $1165\ 00{:}50{:}43.750 --> 00{:}50{:}47.040$  it was a lack of having the biomarker target characterized
- $1166\ 00:50:47.040 \longrightarrow 00:50:48.683$  in prior research studies.
- $1167\ 00:50:49.710 --> 00:50:50.950$  And without the understanding
- $1168\ 00:50:50.950 \longrightarrow 00:50:55.950$  that profile could be predictive for the standard of care,
- $1169\ 00:50:56.340 --> 00:51:00.030$  we have these drugs with drawing from accelerated approval.
- $1170\ 00:51:00.030 \longrightarrow 00:51:01.310$  There's other issues when we look
- 1171 00:51:01.310 --> 00:51:03.210 at tissue-agnostic development.
- $1172~00{:}51{:}03.210 \dashrightarrow 00{:}51{:}06.140$  So I worked with Bayer and MD Anderson last year
- 1173 00:51:06.140 --> 00:51:09.360 to investigate and NTRK fusions.
- 1174 00:51:09.360 --> 00:51:12.260 This is that rare biomarker profile
- $1175\ 00:51:12.260 \longrightarrow 00:51:16.810$  that has led to two drugs getting tissue-agnostic approval,
- $1176\ 00:51:16.810 \longrightarrow 00:51:18.850$  larotrectinib and entrectnib
- $1177\ 00:51:20.410 \longrightarrow 00:51:24.330$  So Bayer bought larotrectinib from LAKSO
- 1178 00:51:25.520 --> 00:51:28.173 and Roche bought entrectnib from Igniter.
- 1179 00:51:29.120 --> 00:51:31.330 So they wanted to understand,
- $1180\ 00{:}51{:}31.330 \dashrightarrow 00{:}51{:}35.690$  and this was used actually in the Canadian approval process.
- $1181\ 00:51:35.690 \longrightarrow 00:51:37.540$  The Canadian approval process is different.
- 1182 00:51:37.540 --> 00:51:39.160 You have a higher level of threshold
- 1183 00:51:39.160 --> 00:51:41.660 that you have to characterize
- $1184\ 00:51:41.660 \longrightarrow 00:51:44.750$  for biomarker targeted therapies.
- 1185 00:51:44.750 --> 00:51:46.600 And they wanted to know specifically,

 $1186\ 00:51:47.570 --> 00:51:50.565$  what is the evidence that NTRK is a prognostic marker?

1187 00:51:50.565 --> 00:51:52.168 How do we know the drugs working

 $1188\ 00:51:52.168 --> 00:51:55.283$  and when it may just be the profile is favorable?

 $1189~00{:}51{:}56.151 \dashrightarrow 00{:}51{:}59.303$  And that's kind of exactly what happened with Ateza.

 $1190\ 00:51:59.303 --> 00:52:04.303$  So we thought that maybe we could interrogate this

1191 00:52:04.840 --> 00:52:06.650 by matching.

1192 00:52:06.650 --> 00:52:08.870 So we had 77 patients from MD Anderson

1193 00:52:08.870 --> 00:52:10.812 that had NTRK fusions.

1194 00:52:10.812 --> 00:52:12.750 Where MD Anderson did the staining,

 $1195\ 00:52:12.750 --> 00:52:15.350$  we knew they had NTRK fusions and we followed them.

 $1196\ 00:52:15.350 \longrightarrow 00:52:17.900$  And some of these patients were on clinical trials.

 $1197\ 00{:}52{:}18.800$  -->  $00{:}52{:}21.420$  So we thought, okay, real-world evidence, right?

1198 00:52:21.420 --> 00:52:24.233 We could match these patients to TCGA data.

 $1199\ 00{:}52{:}25.260 {\:{\mbox{--}}}\!> 00{:}52{:}28.580$  And we could use TCGA data kind of as a control.

 $1200\ 00:52:28.580 \longrightarrow 00:52:29.670$  And we could compare them.

 $1201\ 00:52:29.670 \longrightarrow 00:52:32.440$  We can kind of get a sense of what the expectation was

 $1202\ 00:52:32.440 \longrightarrow 00:52:34.160$  based on TCGA data.

 $1203\ 00:52:34.160 --> 00:52:38.200$  TCGA doesn't have NTRK fusion as one of the mutations.

 $1204\ 00:52:38.200$  --> 00:52:41.340 But they have these indications that were enrolled.

1205 00:52:41.340 --> 00:52:43.100 So among these 77 patients,

1206 00:52:43.100 --> 00:52:46.190 we have like 14 different tumor types.

1207 00:52:46.190 --> 00:52:47.820 So we did this study,

 $1208\ 00:52:47.820 \longrightarrow 00:52:50.070$  we went to TCGA,

- $1209\ 00:52:50.070 \longrightarrow 00:52:51.279$  here are the different tumor types
- $1210\ 00:52:51.279 \longrightarrow 00:52:53.033$  that we had in this trial.
- 1211 00:52:54.630 --> 00:52:56.560 So we're talking about breast cancer
- 1212 00:52:56.560 --> 00:53:01.560 adenocarcinoma, cholangiocarcinoma, GBM.
- 1213 00:53:01.770 --> 00:53:06.400 What I'm showing you here is we extracted the TCGA data
- $1214\ 00:53:06.400 \longrightarrow 00:53:08.680$  from these different tumor types.
- $1215\ 00:53:08.680 \longrightarrow 00:53:10.290$  We matched on stage.
- $1216\ 00:53:10.290 \longrightarrow 00:53:12.180$  We matched on sort of performance status.
- 1217 00:53:12.180 --> 00:53:15.330 We matched on gender or sex, I should say,
- $1218\ 00{:}53{:}15.330 \to 00{:}53{:}18.570$  we matched on all these factors that are relevant
- $1219\ 00{:}53{:}18.570 \dashrightarrow 00{:}53{:}20.510$  for understanding whether patient's expectation
- $1220\ 00:53:20.510 \longrightarrow 00:53:22.230$  is for survival.
- 1221 00:53:22.230 --> 00:53:27.110 And look at the tumor driven heterogeneity.
- 1222 00:53:27.110 --> 00:53:28.700 Thyroid cancers is way up here.
- $1223\ 00:53:28.700 \dashrightarrow 00:53:30.530$  The patients with thyroid cancer that are matched
- 1224 00:53:30.530 --> 00:53:32.130 to these patients at MD Anderson,
- $1225\ 00:53:32.130 \longrightarrow 00:53:33.830$  they're living a really long time.
- $1226\ 00{:}53{:}34.870 \dashrightarrow 00{:}53{:}39.240$  Down here we have patients with GBM, Glioblastoma.
- $1227\ 00:53:39.240 \longrightarrow 00:53:41.050$  This is pancreas.
- $1228\ 00:53:41.050 \longrightarrow 00:53:43.430$  So even though these patients share
- 1229 00:53:43.430 --> 00:53:45.720 a common biomarker profile,
- 1230 00:53:45.720 --> 00:53:47.730 they have tissue types that are very different,
- $1231\ 00:53:47.730 \dashrightarrow 00:53:50.750$  and have very different expectations for survival.
- $1232\ 00:53:50.750 --> 00:53:54.330$  Putting this all together to try to understand
- 1233 00:53:54.330 --> 00:53:56.900 whether NTRK was prognostic or not,
- $1234\ 00:53:56.900 \longrightarrow 00:53:58.533$  was almost impossible to do.
- 1235 00:54:01.060 --> 00:54:04.720 So, you know, conceptually, we have the idea,

- 1236 00:54:04.720 --> 00:54:06.140 we have The Cancer Genome Atlas,
- $1237\ 00:54:06.140 --> 00:54:06.973$  we should be using it.
- $1238\ 00:54:06.973 \longrightarrow 00:54:08.530$  We can use it to do these things.
- $1239\ 00:54:08.530 \longrightarrow 00:54:10.920$  But when it comes down to actually doing it,
- 1240 00:54:10.920 --> 00:54:12.240 it's a real challenge,
- $1241\ 00:54:12.240 \longrightarrow 00:54:14.843$  and it may not provide the information that we need.
- $1242\ 00{:}54{:}15.870 --> 00{:}54{:}19.640$  Okay, so I have two conclusions, very simple ones.
- $1243\ 00:54:19.640 \longrightarrow 00:54:23.500$  Okay, so real-world evidence in precision oncology,
- $1244\ 00:54:23.500 \longrightarrow 00:54:24.740$  how do we use it?
- $1245\ 00:54:24.740 \longrightarrow 00:54:25.860$  Can we use it?
- $1246\ 00:54:25.860 \longrightarrow 00:54:27.350$  The reason we wanna use it,
- $1247\ 00:54:27.350 \longrightarrow 00:54:30.253$  again is because it's very expensive to do,
- $1248\ 00:54:31.510 \longrightarrow 00:54:33.850$  to run trials in oncology.
- $1249\ 00:54:33.850 \longrightarrow 00:54:35.490$  We have these biomarker profiles.
- $1250\ 00:54:35.490 \longrightarrow 00:54:37.360$  Patients have to be stained repeatedly.
- $1251\ 00:54:37.360 \longrightarrow 00:54:39.350$  They have to get imaging,
- $1252\ 00:54:39.350 --> 00:54:42.520$  it's it's burdensome for the patient, and it's expensive.
- $1253\ 00{:}54{:}42.520 \dashrightarrow 00{:}54{:}46.620$  So we wanna make better decisions in early phase
- $1254\ 00{:}54{:}46.620 {\: --> \:} 00{:}54{:}48.980$  because we have all these failures in phase three.
- $1255\ 00:54:48.980 \longrightarrow 00:54:50.020$  We want to do a better job
- $1256\ 00:54:50.020 --> 00:54:52.490$  of designing our phase three trials as well.
- $1257\ 00:54:52.490 \longrightarrow 00:54:53.660$  So what we really wanna know
- $1258~00{:}54{:}53.660 \dashrightarrow 00{:}54{:}56.030$  is can we use real-world evidence to do a better job
- $1259\ 00:54:56.030 \longrightarrow 00:54:57.850$  of setting our null.
- $1260\ 00:54:57.850 \longrightarrow 00:54:59.203$  Where our expectation is.

- $1261\ 00:55:00.038 \longrightarrow 00:55:01.980$  In that case, in that way we can run
- 1262 00:55:01.980 --> 00:55:04.320 these uncontrolled trials in early phase
- 1263 00:55:04.320 --> 00:55:06.680 and, you know, save all the patients
- $1264\ 00:55:06.680 \longrightarrow 00:55:10.300$  to be treated on the potentially promising therapies.
- $1265\ 00{:}55{:}10.300 \dashrightarrow 00{:}55{:}12.770$  Because we'll have a better idea of what our expectation is
- 1266 00:55:12.770 --> 00:55:14.880 and whether this is really promising or not.
- $1267\ 00:55:14.880 \longrightarrow 00:55:17.010$  That is the promise that you hear
- 1268 00:55:17.010 --> 00:55:19.580 about real-word evidence in this setting.
- $1269\ 00:55:19.580 \longrightarrow 00:55:21.893$  So it's really about, can we define the null?
- 1270 00:55:23.270 --> 00:55:25.470 So I think I showed you two examples here
- $1271\ 00:55:25.470 \longrightarrow 00:55:27.860$  where we really couldn't.
- $1272\ 00:55:27.860 \longrightarrow 00:55:29.473$  We tried to.
- 1273 00:55:29.473 --> 00:55:31.890 Like the case is second-line bladder cancer,
- $1274\ 00:55:31.890 \dashrightarrow 00:55:35.060$  we went back to the randomized control trial evidence
- $1275\ 00{:}55{:}35.060 {\: -->\:} 00{:}55{:}38.070$  and the null hypothesis was exactly null hypothesis
- $1276\ 00:55:38.070 \longrightarrow 00:55:39.710$  that Genentech used.
- $1277\ 00:55:39.710$  --> 00:55:42.333 It's just that, that profile was not steady before.
- $1278\ 00:55:43.260 \longrightarrow 00:55:44.800$  So we couldn't do it there.
- 1279 00:55:44.800 --> 00:55:46.620 When we went to the NTRK studies,
- 1280 00:55:46.620 --> 00:55:50.370 the TCGA data didn't characterize NTRK,
- $1281\ 00:55:50.370 --> 00:55:53.100$  but NTRK is so rare that didn't bother us.
- $1282\ 00:55:53.100 --> 00:55:54.410$  So those patients are a mixture
- $1283\ 00:55:54.410 \longrightarrow 00:55:57.020$  of different other mutations.
- $1284\ 00:55:57.020 \longrightarrow 00:55:58.080$  We matched them based
- 1285 00:55:58.080 --> 00:56:00.270 on the clinical prognostic characteristics,
- $1286\ 00:56:00.270 \longrightarrow 00:56:02.670$  but the tumors are so different.
- $1287\ 00:56:02.670$  --> 00:56:05.790 The expectations are so different across the tumors.

- $1288\ 00:56:05.790$  --> 00:56:08.723 It's really hard to understand it from the TCGA data.
- 1289 00:56:10.180 --> 00:56:13.950 In fact, this draws in the question,
- $1290\ 00:56:13.950 \longrightarrow 00:56:17.470$  can you really say, if a patient has GBM
- 1291 00:56:17.470 --> 00:56:19.200 and they also have thyroid cancer,
- $1292\ 00:56:19.200 --> 00:56:21.070$  but they share a mutation,
- $1293\ 00:56:21.070 --> 00:56:24.463$  can we really say something that mutation is the target?
- $1294\ 00{:}56{:}26.180 --> 00{:}56{:}29.730$  Can you really treat those cancers as one cancer type?
- $1295\ 00:56:29.730 \longrightarrow 00:56:33.190$  Which is what the tissue-agnostic model says you can.
- $1296\ 00:56:33.190 \longrightarrow 00:56:35.740$  Right, so the biology is that important.
- $1297\ 00:56:35.740 \longrightarrow 00:56:37.287$  In some cases it has been,
- $1298\ 00:56:37.287 \longrightarrow 00:56:39.210$  in the Pembrolizumab it is,
- $1299\ 00:56:39.210$  --> 00:56:42.270 like immunotherapy, the immune phenotypes really seem
- 1300 00:56:42.270 --> 00:56:45.540 to transcend these cancer tissues,
- $1301\ 00:56:45.540 \longrightarrow 00:56:47.330$  but for other genetic markers,
- $1302\ 00:56:47.330 \longrightarrow 00:56:48.920$  it doesn't seem to be the case.
- 1303 00:56:48.920 --> 00:56:51.920 So I guess my conclusion is retrospectively,
- $1304\ 00:56:51.920 \longrightarrow 00:56:53.370$  we really can't.
- $1305\ 00:56:53.370 \longrightarrow 00:56:55.093$  We can't use it right now.
- $1306\ 00:56:56.060 \longrightarrow 00:56:57.780$  It doesn't seem like we can
- $1307\ 00:56:57.780 \longrightarrow 00:57:01.080$  because we are now in the precision oncology setting.
- 1308 00:57:01.080 --> 00:57:02.300 And yes, of course,
- $1309\ 00:57:02.300 \longrightarrow 00:57:03.930$  if you're in rare disease setting
- $1310\ 00:57:03.930 \longrightarrow 00:57:07.830$  or you're in a non something else, that's unique.
- $1311\ 00:57:07.830 \longrightarrow 00:57:09.880$  You may have to, and do the best you can.
- 1312 00:57:10.880 --> 00:57:12.590 But for the trials I showed you here,
- $1313\ 00:57:12.590 \longrightarrow 00:57:14.050\ I\ don't\ see\ a\ solution\ here$

- $1314\ 00:57:14.050 \longrightarrow 00:57:16.570$  based on retrospective real-world evidence.
- 1315 00:57:16.570 --> 00:57:18.470 I think you could do it prospectively.
- $1316\ 00:57:19.900 \longrightarrow 00:57:22.770$  But I think if you're gonna do it prospectively,
- $1317\ 00:57:22.770 \longrightarrow 00:57:25.380$  there has to be a commitment
- $1318\ 00:57:25.380 \longrightarrow 00:57:27.950$  that right when you start the phase one trial,
- $1319\ 00:57:27.950 \longrightarrow 00:57:29.520$  you need to start staining patients
- $1320\ 00:57:29.520 \longrightarrow 00:57:31.660$  and following them for survival.
- $1321\ 00:57:31.660 \longrightarrow 00:57:33.390$  We don't have a real-world tumor response right now,
- $1322\ 00:57:33.390 \longrightarrow 00:57:35.320$  we can't use that.
- $1323\ 00{:}57{:}35.320 \dashrightarrow 00{:}57{:}40.320$  But you need to have a sort of prospective cohort study
- $1324\ 00:57:40.520 \longrightarrow 00:57:42.400$  that enrolls patients from the community.
- $1325\ 00:57:42.400 \longrightarrow 00:57:45.460$  You need to pay for them to get their assays.
- $1326\ 00{:}57{:}45.460 {\: --> \:} 00{:}57{:}49.030$  You need to understand that the assays may change
- $1327\ 00:57:49.030 \longrightarrow 00:57:51.490$  or develop but to store some information.
- $1328\ 00:57:51.490 \longrightarrow 00:57:53.000$  And then I think later on
- $1329\ 00:57:53.000 \longrightarrow 00:57:54.000$  when you're coming to a decision
- $1330\ 00:57:54.000 \longrightarrow 00:57:55.390$  about phase three or phase two,
- $1331\ 00:57:55.390 \longrightarrow 00:57:57.760$  you go back to that prospective cohort.
- $1332\ 00{:}57{:}57.760 {\longrightarrow} 00{:}58{:}01.260$  And you look for patterns based on the relationships
- $1333\ 00:58:01.260 \dashrightarrow 00:58:04.060$  between the biomarker and the standard of care.
- $1334\ 00:58:04.060 \longrightarrow 00:58:05.380$  So I think you can do it prospectively.
- $1335\ 00{:}58{:}05.380 {\:\dashrightarrow\:} 00{:}58{:}06.800$  That's not what people wanna do though,
- $1336\ 00:58:06.800\ -->\ 00:58:08.987$  they want to use these retrospective databases.
- 1337 00:58:08.987 --> 00:58:13.987 So yeah, I guess that's the end of my talk.
- 1338 00:58:14.590 --> 00:58:16.790 I didn't leave very much time for questions,
- $1339\ 00:58:17.740 \longrightarrow 00:58:19.890$  but I'm happy to take a few if there's any.

- 1340 00:58:24.079 --> 00:58:26.496 (indistinct)
- 1341 00:59:26.340 --> 00:59:28.760 So it's somebody asking a question?
- $1342\ 00:59:28.760 \longrightarrow 00:59:30.760\ I\ can't\ really\ hear.$
- 1343 00:59:30.760 --> 00:59:33.423 I'm sorry, I can't hear it at all, actually.
- 1344 00:59:35.900 --> 00:59:38.900 (people chattering)
- 1345 00:59:45.810 --> 00:59:48.140 <<br/>v Man>Professor Hobbes, can you hear us?</br/>/v>
- 1346 00:59:48.140 --> 00:59:49.780 <v Brian Hobbes>Yeah, I can kind of hear you,</v>
- $1347\ 00:59:49.780 \longrightarrow 00:59:51.543$  but there's a lot of noise.
- 1348 00:59:52.520 --> 00:59:54.826 <v Man>We also have people trying to get in the room</v>
- $1349\ 00:59:54.826 \longrightarrow 00:59:56.646$  so (indistinct)
- 1350 00:59:56.646 --> 00:59:59.813 (students chattering)
- $1351\ 01:00:07.500 --> 01:00:09.502$  Thank you so much, Professor Hobbes.
- 1352 01:00:09.502 --> 01:00:10.827 (students clapping)
- 1353 01:00:10.827 --> 01:00:12.533 <v ->All right, thank you very much.</v>
- $1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Have a great time, thank you.} < / v > 1354\ 01:00:19.030 \longrightarrow 01:00:20.780 < v \text{ Man} > \text{Man} > \text{Man}$
- 1355 01:00:20.780 --> 01:00:23.780 (people chattering)