## WEBVTT

NOTE duration: "00:15:43.5090000"

NOTE language:en-us

NOTE Confidence: 0.8298683

 $00:00:02.950 \longrightarrow 00:00:05.008$  Title of this training is enriching.

NOTE Confidence: 0.8298683

 $00:00:05.010 \longrightarrow 00:00:06.390$  Rare populations of primary

NOTE Confidence: 0.8298683

00:00:06.390 --> 00:00:08.109 human CD 34 positive cells

NOTE Confidence: 0.8298683

 $00:00:08.110 \longrightarrow 00:00:10.462$  by fax and has been prepared by members

NOTE Confidence: 0.8298683

00:00:10.462 --> 00:00:13.391 of the Kraus lab at Yale School of

NOTE Confidence: 0.8298683

00:00:13.391 --> 00:00:15.315 Medicine in partial fulfillment of

NOTE Confidence: 0.8298683

 $00:00:15.315 \longrightarrow 00:00:17.045$  the educational aims outlined in

NOTE Confidence: 0.8298683

00:00:17.045 --> 00:00:19.298 the mission of the Yale Cooperative

NOTE Confidence: 0.8298683

00:00:19.298 --> 00:00:21.468 Center for Excellence in Hematology.

NOTE Confidence: 0.8298683

 $00:00:21.470 \longrightarrow 00:00:24.590$  The purpose of this protocol is to analyze

NOTE Confidence: 0.8298683

 $00:00:24.590 \longrightarrow 00:00:27.440$  an isolate where populations of CD 34,

NOTE Confidence: 0.8298683

00:00:27.440 --> 00:00:29.032 positive matter, poetic stem,

NOTE Confidence: 0.8298683

 $00:00:29.032 \longrightarrow 00:00:31.420$  and progenitor cells from human G,

NOTE Confidence: 0.8298683

00:00:31.420 --> 00:00:33.192 CSF, mobilized peripheral blood,

 $00:00:33.192 \longrightarrow 00:00:35.407$  but maybe more broadly applicable

NOTE Confidence: 0.8298683

 $00{:}00{:}35.407 \dashrightarrow 00{:}00{:}37.967$  to other starting cell populations.

NOTE Confidence: 0.8298683

 $00:00:37.970 \longrightarrow 00:00:40.020$  We will demonstrate the sorting

NOTE Confidence: 0.8298683

00:00:40.020 --> 00:00:41.660 strategy for human megakaryocytic

NOTE Confidence: 0.8298683

 $00:00:41.660 \longrightarrow 00:00:43.597$  over three progenitors in this video

NOTE Confidence: 0.8298683

00:00:43.597 --> 00:00:45.867 as an example of a rare population

NOTE Confidence: 0.8298683

00:00:45.867 --> 00:00:48.105 within the CD 34 positive fraction.

NOTE Confidence: 0.88397837

 $00:00:50.300 \longrightarrow 00:00:52.562$  Human CD 34 positive cells are

NOTE Confidence: 0.88397837

 $00{:}00{:}52.562 \dashrightarrow 00{:}00{:}54.427$ a heterogeneous mixture of stem

NOTE Confidence: 0.88397837

 $00{:}00{:}54.427 \dashrightarrow 00{:}00{:}56.882$  and progenitor cells at various

NOTE Confidence: 0.88397837

 $00{:}00{:}56.882 \dashrightarrow 00{:}00{:}58.394$  stages of differentiation.

NOTE Confidence: 0.88397837

 $00:00:58.400 \longrightarrow 00:01:00.674$  To study a specific population of

NOTE Confidence: 0.88397837

 $00{:}01{:}00.674 \dashrightarrow 00{:}01{:}02.970$  cells like high metabolic stem cells,

NOTE Confidence: 0.88397837

 $00{:}01{:}02.970 \dashrightarrow 00{:}01{:}06.066$  this population must be enriched prior

NOTE Confidence: 0.88397837

 $00:01:06.066 \longrightarrow 00:01:09.578$  to any assets that may take place.

 $00:01:09.580 \longrightarrow 00:01:11.540$  Each progenitor population is

NOTE Confidence: 0.88397837

 $00{:}01{:}11.540 {\:{\mbox{--}}\!>\:} 00{:}01{:}13.990$  defined by a combination of

NOTE Confidence: 0.88397837

 $00:01:13.990 \longrightarrow 00:01:16.599$  surface marker expression profiles.

NOTE Confidence: 0.88397837

 $00:01:16.600 \longrightarrow 00:01:18.692$  Fluorescence activated cell sorting

NOTE Confidence: 0.88397837

 $00{:}01{:}18.692 \dashrightarrow 00{:}01{:}21.830$  or fax uses fluorescent markers to

NOTE Confidence: 0.88397837

 $00:01:21.905 \longrightarrow 00:01:24.995$  detect and separate the cells with

NOTE Confidence: 0.88397837

 $00{:}01{:}24.995 \dashrightarrow 00{:}01{:}27.055$  desired surface marker expression.

NOTE Confidence: 0.88397837

00:01:27.060 --> 00:01:31.386 Building a customized flow cytometry panel.

NOTE Confidence: 0.88397837

00:01:31.390 --> 00:01:34.631 Designing a multi color panel for flow

NOTE Confidence: 0.88397837

 $00:01:34.631 \longrightarrow 00:01:36.980$  cytometry takes advanced preparation.

NOTE Confidence: 0.88397837

 $00:01:36.980 \longrightarrow 00:01:40.697$  Factors that need to be considered include.

NOTE Confidence: 0.88397837

 $00{:}01{:}40.700 \dashrightarrow 00{:}01{:}42.500$  Configuration of the instrument.

NOTE Confidence: 0.88397837

 $00:01:42.500 \longrightarrow 00:01:45.200$  Such as the lasers and filters.

NOTE Confidence: 0.88397837

 $00{:}01{:}45.200 \dashrightarrow 00{:}01{:}49.766$  It is equipped with. The number of

NOTE Confidence: 0.88397837

 $00:01:49.766 \longrightarrow 00:01:53.438$  colors or antibodies to be used.

NOTE Confidence: 0.88397837

 $00:01:53.440 \longrightarrow 00:01:57.070$  And the abundance of the antigens.

 $00:01:57.070 \longrightarrow 00:02:00.514$  We highly recommend utilizing Flora Finder as

NOTE Confidence: 0.88397837

 $00{:}02{:}00.514 \dashrightarrow 00{:}02{:}04.537$  a platform for designing a multi color panel.

NOTE Confidence: 0.88397837

 $00:02:04.540 \longrightarrow 00:02:07.316$  Floor Finder allows the user to enter the

NOTE Confidence: 0.88397837

00:02:07.316 --> 00:02:09.139 exact specifications of their cytometer,

NOTE Confidence: 0.88397837

 $00:02:09.140 \longrightarrow 00:02:11.235$  including which lasers and filter

NOTE Confidence: 0.88397837

 $00:02:11.235 \longrightarrow 00:02:13.980$  sets are installed in the machine.

NOTE Confidence: 0.88397837

00:02:13.980 --> 00:02:14.400 Next,

NOTE Confidence: 0.88397837

 $00:02:14.400 \longrightarrow 00:02:16.920$  users can search and select which

NOTE Confidence: 0.88397837

 $00:02:16.920 \longrightarrow 00:02:19.096$  cell surface antigens and or

NOTE Confidence: 0.88397837

 $00:02:19.096 \longrightarrow 00:02:20.776$  fluorescent proteins they intend

NOTE Confidence: 0.88397837

 $00{:}02{:}20.776 \longrightarrow 00{:}02{:}23.289$  to analyze their cell sample for.

NOTE Confidence: 0.88397837

 $00:02:23.290 \longrightarrow 00:02:25.546$  It is useful to have prior knowledge of

NOTE Confidence: 0.88397837

 $00:02:25.546 \longrightarrow 00:02:27.578$  the antigen abundance on the cells in

NOTE Confidence: 0.88397837

00:02:27.578 --> 00:02:29.710 your sample and cell type of interest,

NOTE Confidence: 0.88397837

 $00:02:29.710 \longrightarrow 00:02:32.470$  but it is not necessary.

 $00:02:32.470 \longrightarrow 00:02:34.455$  Based on the specifications of

NOTE Confidence: 0.88397837

 $00:02:34.455 \longrightarrow 00:02:36.884$  the CYTOMETER and antigens to be

NOTE Confidence: 0.88397837

 $00:02:36.884 \longrightarrow 00:02:38.759$  detected that the user entered,

NOTE Confidence: 0.88397837

00:02:38.760 --> 00:02:41.172 Flora Finder populates a chart of

NOTE Confidence: 0.88397837

 $00:02:41.172 \longrightarrow 00:02:42.378$  commercially available antibodies

NOTE Confidence: 0.88397837

 $00:02:42.378 \longrightarrow 00:02:44.493$  for those antigens categorized by

NOTE Confidence: 0.88397837

00:02:44.493 --> 00:02:46.558 fluorescent channel such that the

NOTE Confidence: 0.88397837

 $00:02:46.558 \longrightarrow 00:02:49.142$  selection of 1 antibody in a particular

NOTE Confidence: 0.88397837

 $00{:}02{:}49.142 \dashrightarrow 00{:}02{:}50.546$  channel automatically blocks the

NOTE Confidence: 0.88397837

 $00:02:50.550 \longrightarrow 00:02:52.122$  user from selecting additional

NOTE Confidence: 0.88397837

 $00:02:52.122 \longrightarrow 00:02:54.087$  antibodies in the same channel,

NOTE Confidence: 0.88397837

 $00{:}02{:}54.090 \dashrightarrow 00{:}02{:}57.122$  thus reducing spectral overlap.

NOTE Confidence: 0.88397837

 $00:02:57.122 \longrightarrow 00:02:59.396$  Titrating staining antibodies.

NOTE Confidence: 0.88397837

 $00{:}02{:}59.400 \dashrightarrow 00{:}03{:}01.435$  To ensure appropriate staining but

NOTE Confidence: 0.88397837

00:03:01.435 --> 00:03:03.063 not compromise antibody specificity,

NOTE Confidence: 0.88397837

 $00:03:03.070 \longrightarrow 00:03:06.136$  it is recommended that you titrate

00:03:06.136 --> 00:03:08.686 your antibodies prior to using

NOTE Confidence: 0.88397837

 $00{:}03{:}08.686 \longrightarrow 00{:}03{:}11.570$  them in a flow analysis or sort.

NOTE Confidence: 0.88397837

00:03:11.570 --> 00:03:13.500 To accomplish this utiliza cell

NOTE Confidence: 0.88397837

 $00:03:13.500 \longrightarrow 00:03:15.842$  sample that contains a mixture of

NOTE Confidence: 0.88397837

 $00:03:15.842 \longrightarrow 00:03:17.900$  cells that are positive for the

NOTE Confidence: 0.88397837

 $00:03:17.900 \longrightarrow 00:03:19.490$  antigen and negative pipette.

NOTE Confidence: 0.88397837

 $00:03:19.490 \longrightarrow 00:03:22.618$  A constant number of cells in a constant

NOTE Confidence: 0.88397837

 $00:03:22.618 \longrightarrow 00:03:25.302$  volume of staining solution into at

NOTE Confidence: 0.88397837

 $00:03:25.302 \longrightarrow 00:03:28.580$  least three but preferably 5 fax chips.

NOTE Confidence: 0.88397837

 $00{:}03{:}28.580 \dashrightarrow 00{:}03{:}31.106$  Keep one tube as the unstained

NOTE Confidence: 0.88397837

 $00{:}03{:}31.106 \dashrightarrow 00{:}03{:}33.280$  control and add increasing volumes

NOTE Confidence: 0.88397837

 $00:03:33.280 \longrightarrow 00:03:36.171$  of the antibody wish to titer to

NOTE Confidence: 0.88397837

00:03:36.171 --> 00:03:37.450 each subsequent tube.

NOTE Confidence: 0.88397837

 $00:03:37.450 \longrightarrow 00:03:39.370$  The range of volumes you choose

NOTE Confidence: 0.88397837

 $00:03:39.370 \longrightarrow 00:03:41.727$  should be based on the recommended

00:03:41.727 --> 00:03:43.735 dilution from the manufacturer.

NOTE Confidence: 0.88397837

 $00:03:43.740 \longrightarrow 00:03:44.764$  For example.

NOTE Confidence: 0.88397837

 $00:03:44.764 \longrightarrow 00:03:47.836$  If the recommended volume test for

NOTE Confidence: 0.88397837

 $00:03:47.836 \longrightarrow 00:03:50.299$  the manufacturer is 5 microliters,

NOTE Confidence: 0.88397837

00:03:50.300 --> 00:03:52.990 pipette 5 microliters of antibody

NOTE Confidence: 0.88397837

 $00:03:52.990 \longrightarrow 00:03:54.604$  into one tube.

NOTE Confidence: 0.88397837

 $00:03:54.610 \longrightarrow 00:03:57.050$  Two microliters of antibody into

NOTE Confidence: 0.88397837

 $00:03:57.050 \longrightarrow 00:03:58.514$  the next tube.

NOTE Confidence: 0.88397837

 $00{:}03{:}58.520 {\:{\mbox{--}}\!>}\ 00{:}04{:}00.800$  One microliter of antibody into

NOTE Confidence: 0.88397837

 $00:04:00.800 \longrightarrow 00:04:02.168$  the next tube.

NOTE Confidence: 0.88397837

 $00{:}04{:}02.170 \dashrightarrow 00{:}04{:}05.008$  And half a microliter of antibody

NOTE Confidence: 0.88397837

 $00:04:05.008 \longrightarrow 00:04:06.900$  into the last tube.

NOTE Confidence: 0.88397837

 $00:04:06.900 \longrightarrow 00:04:08.716$  At the Cytometer record,

NOTE Confidence: 0.88397837

 $00:04:08.716 \longrightarrow 00:04:11.930$  at least 10,000 events for each tube.

NOTE Confidence: 0.88397837

 $00:04:11.930 \longrightarrow 00:04:13.935$  Calculate the staining index by

NOTE Confidence: 0.88397837

 $00:04:13.935 \longrightarrow 00:04:16.819$  measuring the MFA of the negative cells

 $00:04:16.819 \longrightarrow 00:04:19.225$  and the standard deviation as well

NOTE Confidence: 0.88397837

 $00:04:19.225 \longrightarrow 00:04:22.039$  as the MFA for the positive cells.

NOTE Confidence: 0.88397837

 $00{:}04{:}22.040 \dashrightarrow 00{:}04{:}23.912$  With these three measurements,

NOTE Confidence: 0.88397837

 $00:04:23.912 \longrightarrow 00:04:25.784$  calculate the staining index

NOTE Confidence: 0.88397837

 $00:04:25.784 \longrightarrow 00:04:28.054$  by subtracting the MFA of the

NOTE Confidence: 0.88397837

 $00:04:28.054 \longrightarrow 00:04:29.634$  negative population from the MFA

NOTE Confidence: 0.88397837

 $00:04:29.634 \longrightarrow 00:04:31.538$  of the positive population.

NOTE Confidence: 0.88397837

 $00:04:31.540 \longrightarrow 00:04:33.634$  And divide this by two times

NOTE Confidence: 0.88397837

 $00:04:33.634 \longrightarrow 00:04:35.030$  the standard deviation of

NOTE Confidence: 0.8527813

 $00{:}04{:}35.103 \dashrightarrow 00{:}04{:}37.365$  the MFA of the negative population.

NOTE Confidence: 0.8527813

 $00:04:37.370 \longrightarrow 00:04:39.476$  Plot the staining index for each

NOTE Confidence: 0.8527813

 $00:04:39.476 \longrightarrow 00:04:42.879$  tube on a graph to identify the best

NOTE Confidence: 0.8527813

 $00:04:42.879 \longrightarrow 00:04:45.279$  dilution for that particular antibody.

NOTE Confidence: 0.8527813

 $00:04:45.280 \longrightarrow 00:04:48.577$  Sawing and staining CD. 34 positive cells.

NOTE Confidence: 0.84104687

 $00:04:50.890 \longrightarrow 00:04:53.767$  For the purposes of this didactic video,

 $00:04:53.770 \longrightarrow 00:04:56.885$  we will demonstrate the staining protocol for

NOTE Confidence: 0.84104687

 $00:04:56.885 \longrightarrow 00:04:58.990$  human megakaryocytic erythroid progenitors.

NOTE Confidence: 0.84104687

 $00:04:58.990 \longrightarrow 00:05:00.490$  You will need to substitute

NOTE Confidence: 0.84104687

 $00:05:00.490 \longrightarrow 00:05:01.690$  your antibodies of interest,

NOTE Confidence: 0.84104687

 $00:05:01.690 \longrightarrow 00:05:04.810$  but the protocol remains the same.

NOTE Confidence: 0.84104687

 $00:05:04.810 \longrightarrow 00:05:06.078$  With stained with CD,

NOTE Confidence: 0.84104687

00:05:06.078 --> 00:05:08.455 34 positive cells with a panel of

NOTE Confidence: 0.84104687

 $00:05:08.455 \longrightarrow 00:05:11.054$  seven antibodies, one of which is

NOTE Confidence: 0.84104687

 $00{:}05{:}11.054 \dashrightarrow 00{:}05{:}13.730$  a cocktail of Lenny edge markers.

NOTE Confidence: 0.84104687

 $00:05:13.730 \longrightarrow 00:05:16.386$  To reduce the number of channels required for

NOTE Confidence: 0.84104687

 $00{:}05{:}16.386 \to 00{:}05{:}19.148$  the sort and thus reduce spectral overlap,

NOTE Confidence: 0.84104687

 $00:05:19.150 \longrightarrow 00:05:21.310$  we use antibodies against the lineages.

NOTE Confidence: 0.84104687

 $00:05:21.310 \longrightarrow 00:05:23.138$  Markers that are directly

NOTE Confidence: 0.84104687

 $00:05:23.138 \longrightarrow 00:05:24.509$  conjugated to biotin.

NOTE Confidence: 0.84104687

 $00:05:24.510 \longrightarrow 00:05:26.405$  So we incorporate a secondary

NOTE Confidence: 0.84104687

 $00:05:26.405 \longrightarrow 00:05:27.921$  stain with streptavidin antibody

 $00:05:27.921 \longrightarrow 00:05:29.539$  directly conjugated to a floor.

NOTE Confidence: 0.84104687

 $00{:}05{:}29.540 \dashrightarrow 00{:}05{:}31.580$  For that has limited spectral overlap

NOTE Confidence: 0.84104687

 $00:05:31.580 \longrightarrow 00:05:34.198$  with the rest of our staining panel.

NOTE Confidence: 0.63781935

00:05:37.090 --> 00:05:41.030 Required reagents include FBZ.

NOTE Confidence: 0.63781935

 $00:05:41.030 \longrightarrow 00:05:44.578$  Buffer one. Fax buffer.

NOTE Confidence: 0.63781935

 $00:05:44.578 \longrightarrow 00:05:46.750$  And titrated antibodies.

NOTE Confidence: 0.813827

 $00:05:50.840 \longrightarrow 00:05:52.920$  Prepare labeled facts tubes

NOTE Confidence: 0.813827

00:05:52.920 --> 00:05:55.000 for each compensation control.

NOTE Confidence: 0.813827

00:05:55.000 --> 00:05:56.880 Fluorescence minus one control.

NOTE Confidence: 0.813827

 $00:05:56.880 \longrightarrow 00:05:59.700$  As well as the sample tube.

NOTE Confidence: 0.8699426

 $00{:}06{:}03.120 \dashrightarrow 00{:}06{:}05.288$  Compensation controls are cells

NOTE Confidence: 0.8699426

 $00:06:05.288 \longrightarrow 00:06:07.998$  stained with a single antibody.

NOTE Confidence: 0.8699426

 $00:06:08.000 \longrightarrow 00:06:10.185$  Compensation controls are required for

NOTE Confidence: 0.8699426

 $00:06:10.185 \longrightarrow 00:06:13.413$  every floor floor and allow the cytometry

NOTE Confidence: 0.8699426

 $00:06:13.413 \longrightarrow 00:06:15.833$  software to calculate and subtract

 $00:06:15.833 \longrightarrow 00:06:18.150$  spectral overlap between flora force.

NOTE Confidence: 0.8699426

 $00{:}06{:}18.150 \dashrightarrow 00{:}06{:}19.690$  Fluorescence minus one controls

NOTE Confidence: 0.8699426

 $00:06:19.690 \longrightarrow 00:06:22.375$  are cells stained with all but one

NOTE Confidence: 0.8699426

 $00:06:22.375 \longrightarrow 00:06:24.235$  of the antibodies in the panel.

NOTE Confidence: 0.8699426

 $00:06:24.240 \longrightarrow 00:06:27.536$  It identifies the true negative of a floor

NOTE Confidence: 0.8699426

 $00:06:27.536 \longrightarrow 00:06:29.786$  for considering all the interference

NOTE Confidence: 0.8699426

 $00:06:29.786 \longrightarrow 00:06:32.486$  from the other staining floor force.

NOTE Confidence: 0.8699426

 $00:06:32.490 \longrightarrow 00:06:34.650$  These controls help determine the

NOTE Confidence: 0.8699426

 $00{:}06{:}34.650 {\:{\circ}{\circ}{\circ}}>00{:}06{:}37.656$  location of gates that separates cells not

NOTE Confidence: 0.8699426

 $00:06:37.656 \longrightarrow 00:06:40.358$  expressing the antigen from cells that are.

NOTE Confidence: 0.8699426

 $00{:}06{:}40.360 \dashrightarrow 00{:}06{:}42.880$  They are required if the expression

NOTE Confidence: 0.8699426

 $00:06:42.880 \longrightarrow 00:06:45.467$  pattern of the antigen in your

NOTE Confidence: 0.8699426

 $00:06:45.467 \longrightarrow 00:06:47.532$  starting population is a continuum

NOTE Confidence: 0.8699426

 $00:06:47.532 \longrightarrow 00:06:50.009$  of negative to high expressing.

NOTE Confidence: 0.8699426

 $00:06:50.010 \longrightarrow 00:06:52.827$  Or the expression of the antigen is very low,

NOTE Confidence: 0.8699426

 $00:06:52.830 \longrightarrow 00:06:54.372$  so the positive signal is just

 $00:06:54.372 \longrightarrow 00:06:55.850$  slightly higher than the background

NOTE Confidence: 0.8699426

 $00{:}06{:}55.850 \dashrightarrow 00{:}06{:}57.840$  fluorescence of the negative population.

NOTE Confidence: 0.8290179

 $00:06:59.710 \longrightarrow 00:07:01.760$  We quickly thaw cryopreserved CD

NOTE Confidence: 0.8290179

 $00:07:01.760 \longrightarrow 00:07:04.629$  34 positive cells in a 37 degree

NOTE Confidence: 0.8290179

 $00:07:04.629 \longrightarrow 00:07:07.233$  water bath until just a few ice

NOTE Confidence: 0.8290179

 $00:07:07.233 \longrightarrow 00:07:09.546$  crystals are left in the cryotube.

NOTE Confidence: 0.8290179

 $00:07:09.550 \longrightarrow 00:07:11.566$  Take care not to submerge the

NOTE Confidence: 0.8290179

 $00{:}07{:}11.566 \dashrightarrow 00{:}07{:}14.214$  O-ring and cap of the cryo vial

NOTE Confidence: 0.8290179

 $00:07:14.214 \longrightarrow 00:07:16.269$  to reduce risk of contamination.

NOTE Confidence: 0.8290179

 $00:07:16.270 \longrightarrow 00:07:18.058$  When the cells are nearly thought,

NOTE Confidence: 0.8290179

 $00:07:18.060 \longrightarrow 00:07:20.160$  spray the cry of I'll with 70%

NOTE Confidence: 0.8290179

00:07:20.160 --> 00:07:22.434 ethanol and wipe it dry before

NOTE Confidence: 0.8290179

 $00{:}07{:}22.434 \dashrightarrow 00{:}07{:}24.998$  bringing it into the hood to open.

NOTE Confidence: 0.8290179

 $00{:}07{:}25.000 \dashrightarrow 00{:}07{:}26.810$  Be sure to practice strict

NOTE Confidence: 0.8290179

00:07:26.810 --> 00:07:29.036 as eptic technique in a bio safety

 $00:07:29.036 \longrightarrow 00:07:31.430$  cabinet rated for BSL 2 for the

NOTE Confidence: 0.8290179

 $00:07:31.430 \longrightarrow 00:07:33.140$  duration of this protocol.

NOTE Confidence: 0.7869187

 $00:07:40.600 \longrightarrow 00:07:42.530$  Add one milliliter of 100%

NOTE Confidence: 0.7869187

 $00:07:42.530 \longrightarrow 00:07:45.218$  FBS to a 50 mil conical tube.

NOTE Confidence: 0.7208567

 $00:07:50.700 \longrightarrow 00:07:53.557$  Add one milliliter of 100% FBS to the

NOTE Confidence: 0.7208567

 $00{:}07{:}53.557 \dashrightarrow 00{:}07{:}55.711$  cryotube and gently transfer the full

NOTE Confidence: 0.7208567

00:07:55.711 --> 00:07:57.877 contents of the cryotube into the

NOTE Confidence: 0.7208567

00:07:57.877 --> 00:07:59.930 50 mil conical tube containing FBS.

NOTE Confidence: 0.7640226

 $00:08:13.040 \longrightarrow 00:08:15.525$  Add up to 50 milliliters of buffer.

NOTE Confidence: 0.7640226

00:08:15.530 --> 00:08:16.954 One dropwise cell suspension

NOTE Confidence: 0.7640226

 $00:08:16.954 \longrightarrow 00:08:19.090$  in the 50 mil conical tube.

NOTE Confidence: 0.7640226

 $00:08:19.090 \longrightarrow 00:08:22.294$  To delete out the DMSO in the freezing media.

NOTE Confidence: 0.8578139

00:08:25.410 --> 00:08:28.170 Spin at 1200 RPM 4 degrees

NOTE Confidence: 0.8578139

 $00:08:28.170 \longrightarrow 00:08:30.010$  Celsius for 10 minutes.

NOTE Confidence: 0.8364337

 $00:08:32.860 \longrightarrow 00:08:34.540$  When the spin is done,

NOTE Confidence: 0.8364337

 $00{:}08{:}34.540 \dashrightarrow 00{:}08{:}35.880$  carefully discard the supernatant.

 $00:08:40.300 \longrightarrow 00:08:42.610$  Resuspend the pellet with 20

NOTE Confidence: 0.7796592

 $00:08:42.610 \longrightarrow 00:08:44.458$  milliliters of fax buffer.

NOTE Confidence: 0.7796592

00:08:44.460 --> 00:08:46.315 Take a representative 10 microliter

NOTE Confidence: 0.7796592

 $00:08:46.315 \longrightarrow 00:08:48.783$  aliquot for cell counting and repeat the

NOTE Confidence: 0.7796592

 $00:08:48.783 \longrightarrow 00:08:50.975$  spin of the conical tube at 1200 RPM,

NOTE Confidence: 0.7796592

 $00:08:50.980 \longrightarrow 00:08:52.936$  4 degrees Celsius for 10 minutes.

NOTE Confidence: 0.78864664

00:08:54.970 --> 00:08:57.930 Add 10 microliters of two extra pen blue

NOTE Confidence: 0.78864664

 $00:08:57.930 \dashrightarrow 00:09:01.588$  to the aliquot of cells and pipette to mix.

NOTE Confidence: 0.78864664

 $00:09:01.590 \longrightarrow 00:09:03.360$  Calculate the viable cell number

NOTE Confidence: 0.78864664

 $00:09:03.360 \longrightarrow 00:09:05.130$  in the sample control tubes.

NOTE Confidence: 0.78864664

00:09:05.130 --> 00:09:08.883 Each require 5 \* 10 to the four cells.

NOTE Confidence: 0.78864664

 $00:09:08.890 \longrightarrow 00:09:10.732$  FMO tubes require as few as

NOTE Confidence: 0.78864664

00:09:10.732 --> 00:09:14.880 2 \* 10 to the 4th cells.

NOTE Confidence: 0.78864664

 $00{:}09{:}14.880 \dashrightarrow 00{:}09{:}16.956$  When the second spin is complete,

NOTE Confidence: 0.78864664

 $00:09:16.960 \longrightarrow 00:09:18.565$  resuspend the pellet in 500

00:09:18.565 --> 00:09:20.170 microliters of Facs buffer an

NOTE Confidence: 0.78864664

 $00:09:20.233 \longrightarrow 00:09:22.063$  aliquot the necessary volume of

NOTE Confidence: 0.78864664

 $00:09:22.063 \longrightarrow 00:09:23.893$  cells required for control tubes.

NOTE Confidence: 0.78864664

 $00:09:23.900 \longrightarrow 00:09:25.665$  Add additional facts buffer to

NOTE Confidence: 0.78864664

00:09:25.665 --> 00:09:27.816 bring the sample tube volume back

NOTE Confidence: 0.78864664

 $00:09:27.816 \longrightarrow 00:09:29.108$  up to 500 microliters.

NOTE Confidence: 0.78864664

 $00:09:29.110 \longrightarrow 00:09:31.672$  Take care not to exceed a cell

NOTE Confidence: 0.78864664

 $00:09:31.672 \longrightarrow 00:09:34.276$  concentration of 20 \* 10 to the six

NOTE Confidence: 0.78864664

 $00:09:34.276 \dashrightarrow 00:09:36.740$  cells per milliliter in the sample tube.

NOTE Confidence: 0.78377306

 $00{:}09{:}39.390 \dashrightarrow 00{:}09{:}41.058$  Aliquot the titrated amount

NOTE Confidence: 0.78377306

 $00{:}09{:}41.058 \mathrel{--}{>} 00{:}09{:}42.726$  of primary staining antibodies

NOTE Confidence: 0.78377306

 $00:09:42.726 \longrightarrow 00:09:44.869$  into the corresponding tubes.

NOTE Confidence: 0.78377306

 $00:09:44.870 \longrightarrow 00:09:46.920$  Be especially careful when allocating

NOTE Confidence: 0.78377306

 $00:09:46.920 \longrightarrow 00:09:49.405$  antibodies in the FMO tube so

NOTE Confidence: 0.78377306

00:09:49.405 --> 00:09:51.230 you don't accidentally add the

NOTE Confidence: 0.78377306

 $00:09:51.230 \longrightarrow 00:09:53.210$  antibody that should be absent.

00:09:53.210 --> 00:09:54.690 Additionally, it is important to

NOTE Confidence: 0.78377306

 $00:09:54.690 \longrightarrow 00:09:56.617$  note that the volume of antibodies

NOTE Confidence: 0.78377306

00:09:56.617 --> 00:09:58.567 added cannot exceed the maximum

NOTE Confidence: 0.78377306

 $00:09:58.567 \longrightarrow 00:10:00.326$  standing volume, which is.

NOTE Confidence: 0.78377306

 $00:10:00.326 \longrightarrow 00:10:02.278$  100 microliters for controls

NOTE Confidence: 0.78377306

 $00:10:02.278 \longrightarrow 00:10:04.880$  and one milliliter for sample.

NOTE Confidence: 0.78377306

 $00:10:04.880 \longrightarrow 00:10:06.008$  In other words,

NOTE Confidence: 0.78377306

00:10:06.008 --> 00:10:08.264 if you allocated 50 microliters of

NOTE Confidence: 0.78377306

 $00:10:08.264 \longrightarrow 00:10:10.458$  control cells to the control tubes,

NOTE Confidence: 0.78377306

00:10:10.460 --> 00:10:12.445 the volume of staining antibody

NOTE Confidence: 0.78377306

 $00:10:12.445 \longrightarrow 00:10:14.430$  cannot exceed 50 microliters so

NOTE Confidence: 0.78377306

00:10:14.494 --> 00:10:16.659 that the total staining volume

NOTE Confidence: 0.78377306

 $00{:}10{:}16.659 \dashrightarrow 00{:}10{:}17.958$  remains 100 microliters.

NOTE Confidence: 0.78377306

 $00:10:17.960 \longrightarrow 00:10:19.808$  If the volume of the staining

NOTE Confidence: 0.78377306

 $00:10:19.808 \longrightarrow 00:10:21.970$  antibodies is less than 50 microliters,

00:10:21.970 --> 00:10:23.968 then add additional facts buffer to

NOTE Confidence: 0.78377306

 $00{:}10{:}23.968 \dashrightarrow 00{:}10{:}26.051$  bring the final staining volume to

NOTE Confidence: 0.78377306

 $00:10:26.051 \longrightarrow 00:10:28.115$  100 microliters for control tubes and

NOTE Confidence: 0.78377306

 $00:10:28.115 \longrightarrow 00:10:29.979$  one milliliter for the sample tube.

NOTE Confidence: 0.80317324

 $00:10:33.140 \longrightarrow 00:10:35.798$  Once all cells Facs buffer an,

NOTE Confidence: 0.80317324

 $00:10:35.800 \longrightarrow 00:10:38.894$  antibodies are added to the staining tubes,

NOTE Confidence: 0.80317324

 $00:10:38.900 \longrightarrow 00:10:41.224$  incubate according to the

NOTE Confidence: 0.80317324

00:10:41.224 --> 00:10:42.386 manufacturer's recommendations.

NOTE Confidence: 0.80317324

 $00{:}10{:}42.390 \dashrightarrow 00{:}10{:}43.842$  Most antibodies require incubation

NOTE Confidence: 0.80317324

 $00:10:43.842 \longrightarrow 00:10:46.640$  for 30 minutes on ice in the dark.

NOTE Confidence: 0.83693075

 $00{:}10{:}49.390 \dashrightarrow 00{:}10{:}51.406$  After the primary stain is complete,

NOTE Confidence: 0.83693075

 $00:10:51.410 \longrightarrow 00:10:53.492$  add one milliliter of fax buffer

NOTE Confidence: 0.83693075

 $00:10:53.492 \longrightarrow 00:10:55.636$  to the control tubes and two

NOTE Confidence: 0.83693075

 $00{:}10{:}55.636 \dashrightarrow 00{:}10{:}57.682$  milliliters of fax buffer to the

NOTE Confidence: 0.83693075

 $00:10:57.682 \longrightarrow 00:10:59.807$  sample tube and spin at 1200 RPM,

NOTE Confidence: 0.83693075

 $00{:}10{:}59.810 \dashrightarrow 00{:}11{:}01.820$ 4 degrees Celsius for 10 minutes.

 $00:11:06.140 \longrightarrow 00:11:07.610$  When the spin is complete,

NOTE Confidence: 0.8434153

 $00{:}11{:}07.610 \longrightarrow 00{:}11{:}08.870$  discard the supernatant, taking

NOTE Confidence: 0.8434153

 $00:11:08.870 \longrightarrow 00:11:11.071$  great care not to discard the cell

NOTE Confidence: 0.8434153

00:11:11.071 --> 00:11:12.877 pellets at the bottom of the tubes.

NOTE Confidence: 0.7804666

 $00:11:21.790 \longrightarrow 00:11:24.086$  If any of the antibodies used in the

NOTE Confidence: 0.7804666

 $00:11:24.086 \longrightarrow 00:11:26.279$  first stain were conjugated to biotin,

NOTE Confidence: 0.7804666

 $00:11:26.280 \longrightarrow 00:11:27.828$  a secondary stain with

NOTE Confidence: 0.7804666

 $00{:}11{:}27.828 \to 00{:}11{:}29.376$  streptavidin conjugated to a

NOTE Confidence: 0.7804666

 $00:11:29.376 \longrightarrow 00:11:31.367$  unique floor 4 will be required.

NOTE Confidence: 0.7804666

00:11:31.370 --> 00:11:33.260 Similarly to the primary stain,

NOTE Confidence: 0.7804666

 $00:11:33.260 \longrightarrow 00:11:35.435$  add the appropriate amount of

NOTE Confidence: 0.7804666

 $00:11:35.435 \longrightarrow 00:11:37.175$  streptavidin according to the

NOTE Confidence: 0.7804666

 $00{:}11{:}37.175 \dashrightarrow 00{:}11{:}39.250$  titration to the appropriate tubes.

NOTE Confidence: 0.7804666

 $00{:}11{:}39.250 \dashrightarrow 00{:}11{:}41.714$  You don't have to add this to single

NOTE Confidence: 0.7804666

 $00:11:41.714 \longrightarrow 00:11:43.900$  color control tubes that were not

00:11:43.900 --> 00:11:45.780 stained with a biotin conjugated

NOTE Confidence: 0.7804666

 $00{:}11{:}45.780 \dashrightarrow 00{:}11{:}48.588$  antibody or to an FMO tube in which the

NOTE Confidence: 0.7804666

 $00:11:48.588 \longrightarrow 00:11:51.358$  biotin conjugated antibody was omitted.

NOTE Confidence: 0.7804666

 $00:11:51.360 \longrightarrow 00:11:53.405$  Bring the total staining volume

NOTE Confidence: 0.7804666

00:11:53.405 --> 00:11:55.937 up to 100 microliters for control

NOTE Confidence: 0.7804666

00:11:55.937 --> 00:11:58.391 teams or one milliliter for sample

NOTE Confidence: 0.7804666

 $00:11:58.391 \longrightarrow 00:12:00.200$  tubes with fax buffer.

NOTE Confidence: 0.7804666

 $00:12:00.200 \longrightarrow 00:12:02.452$  Incubate according to the

NOTE Confidence: 0.7804666

 $00:12:02.452 \longrightarrow 00:12:03.578$  manufacturer's recommendation.

NOTE Confidence: 0.7804666

 $00:12:03.580 \longrightarrow 00:12:04.892$  Most antibodies require incubation

NOTE Confidence: 0.7804666

 $00{:}12{:}04.892 \dashrightarrow 00{:}12{:}07.419$  for 30 minutes on ice in the dark.

NOTE Confidence: 0.8323322

00:12:11.860 --> 00:12:13.540 Once secondary staining is complete,

NOTE Confidence: 0.8323322

 $00:12:13.540 \longrightarrow 00:12:16.074$  repeat the wash and spin steps as

NOTE Confidence: 0.8323322

 $00{:}12{:}16.074 \dashrightarrow 00{:}12{:}18.600$  was done after the primary stain.

NOTE Confidence: 0.8323322

00:12:18.600 --> 00:12:20.570 Resuspend the control pellets in

NOTE Confidence: 0.8323322

 $00:12:20.570 \longrightarrow 00:12:22.540$  200 microliters of Facs buffer

 $00:12:22.608 \longrightarrow 00:12:24.423$  and 300 microliters of Facs

NOTE Confidence: 0.8323322

 $00:12:24.423 \longrightarrow 00:12:26.238$  buffer for the sample palette.

NOTE Confidence: 0.8323322

 $00:12:26.240 \longrightarrow 00:12:27.860$  Filter these cell suspensions through

NOTE Confidence: 0.8323322

 $00:12:27.860 \longrightarrow 00:12:30.274$  a 40 or 100 Micron filter depending

NOTE Confidence: 0.8323322

 $00{:}12{:}30.274 \dashrightarrow 00{:}12{:}32.464$  on the expected size of yourselves

NOTE Confidence: 0.8323322

 $00:12:32.464 \longrightarrow 00:12:34.238$  and proceed to the cytometer.

NOTE Confidence: 0.78835535

 $00:12:40.540 \longrightarrow 00:12:42.400$  Prime just sorting the cells.

NOTE Confidence: 0.78835535

 $00{:}12{:}42.400 \dashrightarrow 00{:}12{:}44.255$  Prepare collection tubes or plates

NOTE Confidence: 0.78835535

 $00:12:44.255 \longrightarrow 00:12:46.110$  to receive the sorted cells.

NOTE Confidence: 0.78835535

 $00:12:46.110 \longrightarrow 00:12:48.138$  Confirmed that Uber plate you wish

NOTE Confidence: 0.78835535

 $00{:}12{:}48.138 \dashrightarrow 00{:}12{:}49.971$  to use fits the specification

NOTE Confidence: 0.78835535

 $00:12:49.971 \longrightarrow 00:12:52.407$  of the cytometer to be used.

NOTE Confidence: 0.78835535

 $00{:}12{:}52.410 \dashrightarrow 00{:}12{:}54.254$  Typically, growth media supplemented

NOTE Confidence: 0.78835535

 $00:12:54.254 \longrightarrow 00:12:57.020$  with survival cytokines can be used

NOTE Confidence: 0.78835535

00:12:57.087 --> 00:12:59.235 to collect certain cells into and

 $00:12:59.235 \longrightarrow 00:13:01.699$  results in better post sort viability.

NOTE Confidence: 0.78835535

00:13:01.700 --> 00:13:02.600 Of special note,

NOTE Confidence: 0.78835535

 $00:13:02.600 \longrightarrow 00:13:04.700$  be sure to follow all safety policies

NOTE Confidence: 0.78835535

 $00:13:04.763 \longrightarrow 00:13:06.459$  and procedures when transporting

NOTE Confidence: 0.78835535

 $00:13:06.459 \longrightarrow 00:13:08.155$  yourselves to the cytometer.

NOTE Confidence: 0.78835535

00:13:08.160 --> 00:13:08.912 For example,

NOTE Confidence: 0.78835535

 $00{:}13{:}08.912 \dashrightarrow 00{:}13{:}11.168$ unfixed human CD 34 positive cells

NOTE Confidence: 0.78835535

00:13:11.168 --> 00:13:13.083 are required to be transported

NOTE Confidence: 0.78835535

 $00{:}13{:}13.083 \dashrightarrow 00{:}13{:}15.267$  in a secondary container that is

NOTE Confidence: 0.78835535

00:13:15.267 --> 00:13:17.136 labeled with a BSL two sticker.

NOTE Confidence: 0.7975483

00:13:19.290 --> 00:13:20.834 Once at the cytometer,

NOTE Confidence: 0.7975483

 $00:13:20.834 \longrightarrow 00:13:22.764$  set the color parameters that

NOTE Confidence: 0.7975483

 $00:13:22.764 \longrightarrow 00:13:24.990$  will be used to sort the cells.

NOTE Confidence: 0.7975483

 $00:13:24.990 \longrightarrow 00:13:26.830$  Create compensation tubes for each

NOTE Confidence: 0.7975483

00:13:26.830 --> 00:13:29.064 single color control tube you have

NOTE Confidence: 0.7975483

 $00{:}13{:}29.064 \dashrightarrow 00{:}13{:}31.038$  and create a new specimen entitled,

 $00:13:31.040 \longrightarrow 00:13:35.144$  FMO and generate tubes for each FML you have.

NOTE Confidence: 0.7975483

 $00{:}13{:}35.150 \dashrightarrow 00{:}13{:}38.776$  Also create a tube for your sample.

NOTE Confidence: 0.7975483

00:13:38.780 --> 00:13:40.610 Using the unstained control cells,

NOTE Confidence: 0.7975483

00:13:40.610 --> 00:13:42.806 adjust the voltages for each channel,

NOTE Confidence: 0.7975483

00:13:42.810 --> 00:13:44.254 including FC and SC,

NOTE Confidence: 0.7975483

 $00:13:44.254 \longrightarrow 00:13:46.819$  so that your cells are mostly clustered

NOTE Confidence: 0.7975483

 $00:13:46.819 \longrightarrow 00:13:49.744$  in the center of the FC SC dot plot

NOTE Confidence: 0.7975483

 $00{:}13{:}49.817 \dashrightarrow 00{:}13{:}52.199$  and the background signal in each

NOTE Confidence: 0.7975483

 $00:13:52.199 \longrightarrow 00:13:55.770$  color channel is well below 2:50.

NOTE Confidence: 0.7975483

00:13:55.770 --> 00:13:56.382 Before recording,

NOTE Confidence: 0.7975483

 $00:13:56.382 \longrightarrow 00:13:58.218$  check those voltages on each of

NOTE Confidence: 0.7975483

 $00{:}13{:}58.218 \dashrightarrow 00{:}13{:}59.680$  the single color control tubes

NOTE Confidence: 0.7975483

 $00:13:59.680 \longrightarrow 00:14:01.276$  as well as the sample tube.

NOTE Confidence: 0.7975483

 $00:14:01.280 \longrightarrow 00:14:02.960$  Adjust voltages for each channel

NOTE Confidence: 0.7975483

 $00:14:02.960 \longrightarrow 00:14:05.234$  so that the negative peak is below

 $00:14:05.234 \longrightarrow 00:14:07.096$  10 to the third and the positive

NOTE Confidence: 0.7975483

 $00:14:07.096 \longrightarrow 00:14:08.820$  peak is below 10 to the five.

NOTE Confidence: 0.879604

 $00:14:10.870 \longrightarrow 00:14:13.397$  Once you are satisfied with the voltages,

NOTE Confidence: 0.879604

 $00:14:13.400 \longrightarrow 00:14:15.210$  begin by recording single color

NOTE Confidence: 0.879604

 $00:14:15.210 \longrightarrow 00:14:17.020$  controls and calculate the compensation.

NOTE Confidence: 0.879604

 $00:14:17.020 \longrightarrow 00:14:20.200$  Once all have been recorded.

NOTE Confidence: 0.879604

 $00:14:20.200 \longrightarrow 00:14:21.532$  Then record the FML's.

NOTE Confidence: 0.879604

 $00:14:21.532 \longrightarrow 00:14:23.530$  Use the recorded data from the

NOTE Confidence: 0.879604

 $00{:}14{:}23.594 \dashrightarrow 00{:}14{:}25.514$  Fmo's to help draw and position

NOTE Confidence: 0.879604

 $00:14:25.514 \longrightarrow 00:14:27.490$  gates based on true negatives.

NOTE Confidence: 0.8112558

 $00{:}14{:}29.960 \dashrightarrow 00{:}14{:}31.927$  Once all of the gates of the

NOTE Confidence: 0.8112558

00:14:31.927 --> 00:14:33.449 sorting strategy have been drawn,

NOTE Confidence: 0.8112558

 $00:14:33.450 \longrightarrow 00:14:35.935$  load the sample onto the cytometer and

NOTE Confidence: 0.8112558

 $00{:}14{:}35.935 \dashrightarrow 00{:}14{:}38.129$  set the population you wish to sort.

NOTE Confidence: 0.8112558

 $00:14:38.130 \longrightarrow 00:14:40.105$  Also specify what kind of

NOTE Confidence: 0.8112558

00:14:40.105 --> 00:14:41.290 collection container you're

 $00:14:41.290 \longrightarrow 00:14:43.587$  using and how many cells to sort.

NOTE Confidence: 0.8112558

 $00{:}14{:}43.590 \dashrightarrow 00{:}14{:}45.155$  Then begin sorting your sample

NOTE Confidence: 0.8112558

00:14:45.155 --> 00:14:46.720 tube and collecting your enriched

NOTE Confidence: 0.8112558

 $00:14:46.777 \longrightarrow 00:14:48.689$  population for downstream applications.

NOTE Confidence: 0.8187878

 $00:15:00.820 \dashrightarrow 00:15:03.262$  After the service complete, use the

NOTE Confidence: 0.8187878

 $00{:}15{:}03.262 \to 00{:}15{:}05.760$  cells for your downstream application.

NOTE Confidence: 0.8187878

00:15:05.760 --> 00:15:08.748 If sorted cells were collected in a fax tube,

NOTE Confidence: 0.8187878

 $00{:}15{:}08.750 \dashrightarrow 00{:}15{:}10.969$  add 2 milliliters of I MDM media

NOTE Confidence: 0.8187878

 $00:15:10.969 \longrightarrow 00:15:13.354$  to wash the cells in the sorted

NOTE Confidence: 0.8187878

 $00:15:13.354 \longrightarrow 00:15:15.388$  tube and spent at 1200 RPM,

NOTE Confidence: 0.8187878

 $00:15:15.390 \longrightarrow 00:15:17.376$  4 degrees Celsius for 10 minutes.

NOTE Confidence: 0.77992374

 $00{:}15{:}20.680 \dashrightarrow 00{:}15{:}22.750$  Resuspend the cell pellet in the

NOTE Confidence: 0.77992374

 $00{:}15{:}22.750 \dashrightarrow 00{:}15{:}25.050$  appropriate growth media or freezing media.

NOTE Confidence: 0.75087214

 $00:15:28.480 \longrightarrow 00:15:30.905$  This concludes the protocol for

NOTE Confidence: 0.75087214

 $00{:}15{:}30.905 \dashrightarrow 00{:}15{:}33.650$  Multicolor Facs sorting of CD 34.

 $00{:}15{:}33.650 \dashrightarrow 00{:}15{:}36.470$  Positive amount Aquatic Center gender cells.