**1. What is included with each kit or panel?**

Panel configurations:
- Two 96-well (2 x 96w FAST) plates with optical plate covers
- One 384-well (384w) plate with optical plate covers

Kit configurations:
- Two 96-well (2 x 96w FAST) plates with optical plate covers & TaqMan® Gene Expression Master Mix
- One 384-well (384w) plate with optical plate covers & TaqMan® Gene Expression Master Mix

**Part 1: Sample Generation**

**2. How many cells are required to run one experiment?**

While we recommend that you use about 0.5 million cells per experiment, or the number of cells equivalent to 1 well of a 6-well dish, you can use as few as 100,000 cells when performing RNA purification. If you perform a lysis protocol using Cells-to-CT™ or CellsDirect™ One-Step qPCR kits, as few as 15,000 cells is sufficient. Please note, however, that reducing cell number can compromise the quality of the results.

**3. Can I use cells that are on feeders or in feeder-free conditions?**

Yes. The hPSC Scorecard™ Panel measures the potential for self-renewal and trilineage differentiation of PSCs grown on feeders or in feeder-free conditions.
4. **How do I harvest human PSCs cultured using feeder-dependent culture systems?**

We recommend removing feeders from your culture before proceeding with Scorecard™ analysis. To remove feeders, harvest your PSCs with collagenase, allow the colonies to settle by gravity sedimentation to reduce feeder-carryover, and re-seed cells in feeder-free conditions on Geltrex® matrix-coated dishes and MEF conditioned medium for a 1 to 2 passages. The presence of feeders can contribute to gene expression measured in the assay, thus altering the gene-signature pattern.

5. **Can I use cells on a feeder-free system cultured in novel media?**

The hPSC Scorecard™ Panel measures self-renewal and trilineage differentiation potential of PSCs and is not restricted by culture conditions. You should be aware that using novel media systems may have particular effects on the cells in terms of pluripotency and/or their differentiation. We recommend designing experiments that use novel media by including a control condition that utilizes traditional media or media in which expression patterns have been tested and confirmed. The most recent version of the hPSC Scorecard™ Analysis application offers a differentiation index plot for viewing gene expression in EBs relative to the undifferentiated state.

6. **What is the minimum time required to form EBs and is there a specific protocol?**

You can differentiate cells using any of the established methods. When using suspension embryoid bodies, we recommend that you allow at least 7 days of differentiation prior to analysis.

7. **I am only interested in directed differentiated cells. Do I need to use a particular method?**

You can perform directed differentiation according to your own methods. However, the time point when expression is noticeable will largely depend on the robustness of the methods. We recommend testing a few time points to monitor differentiation with time.

8. **Have you tested cells grown in NIM with Scorecard™ analysis?**

In the data shown for directed differentiation of cells into ectoderm in monolayer culture, the 3-day time point is too early to detect ectoderm expression, but the cells become positive for ectoderm by day 5. H9 cells were differentiated into NSCs using Gibco® PSC Neural Induction Medium (denoted as NIM) and hPSC Scorecard™ analysis was performed at various time points. The control sample is undifferentiated H9 ESC.
9. What is the earliest passage that I can use when I generate iPSCs?

We recommend that you culture iPSC clones to at least passage 8–10 until they are stable and homogeneous, prior to analyzing the hPSC Scorecard™ Panel. Early passage iPSC clones may give low self-renewal (“pluri” in v1.1 of the analysis software) scores or show higher expression of lineage genes. See “Part 6: Examples of results” for more information on how to interpret the hPSC Scorecard™ Analysis Software output.

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>Self-renewal</th>
<th>Ectoderm</th>
<th>Mesoderm</th>
<th>Endoderm</th>
</tr>
</thead>
<tbody>
<tr>
<td>H9-CM-Ctrl</td>
<td>0.61</td>
<td>-0.53</td>
<td>-0.80</td>
<td>-1.57</td>
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<tr>
<td>H9-NIM-Day3</td>
<td>-3.62</td>
<td>-0.11</td>
<td>-1.06</td>
<td>-1.79</td>
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<tr>
<td>H9-NIM-Day5</td>
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<td>1.16</td>
<td>-1.12</td>
<td>-1.99</td>
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<tr>
<td>H9-NIM-Day7</td>
<td>-5.62</td>
<td>0.96</td>
<td>-1.75</td>
<td>-1.73</td>
</tr>
</tbody>
</table>

Table 1. Comparison of hPSC Scorecard™ Panel results at different passage numbers. At passage 5 (P5), the expression profile of cells does not correlate to that of a truly pluripotent cell line. At passage 8 (P8), the expression profile matches the reference standard for the four “good” clones (BS4-iPS C1, BS4-iPS C3, BS4-iPS C7, BS4-iPS C8), but not for the one “poor” clone (BS4-iPS C5). Scores close to zero (0) for “pluri”, endo, meso, and ecto represent qualities of a “good” clone; those shaded green indicate upregulation (>1) and shaded red indicate downregulation (<1).

10. When I use Sendai virus (SEV) to generate iPSCs, can I check for the presence of residual virus?

SEV is included in the panel and can detect the presence of the Sendai virus backbone. Note that this method will not distinguish between the different reprogramming factors but just the presence or absence of the residual virus in the cells. If an unexpected signal is detected, check the amplification curve to determine if it is a false positive. If it’s a false positive, you may ignore the flag.
11. Will the presence of Sendai virus affect my results?

Results indicate that the presence or absence of Sendai virus in established iPSC clones does not have an impact on pluripotency.

12. Does the reprogramming method affect the Scorecard™ results?

Minor differences in gene expression profiles are sometimes observed based on the reprogramming method, but in general the hPSC Scorecard™ results do not change significantly for lines derived using various reprogramming methods. As an example, ESC and iPSC derived using episomal or Sendai-based reprogramming systems were tested before and after seven days of spontaneous differentiation. Cells were grown in KSR-based media on irradiated MEFs prior to removal of FGF for EB formation.

![Graphs showing pass/fail results and scores for each germ layer](image)

Figure 1. The top panel shows the pass/fail results in the main interface of the hPSC Scorecard™ Analysis Software. The graphs on the bottom show the scores for each of the four sub-panels tested. These scores measure how closely the sample aligns to the expression profile of the nine undifferentiated pluripotent stem cell lines in the reference set. In general, scores close to zero indicate similarity to the reference set, and scores greater than about 1 indicate upregulation relative to the reference set and scores less than -1 indicate downregulation relative to the reference set. Notice that after seven days of EB formation, the pluripotency (or self-renewal) markers are significantly downregulated and ecto, meso and endo factors are upregulated. Also note that the scores for each germ layer are independent of the others. For example, the magnitude of the scores for ectoderm cannot be compared to the magnitude of the scores for endoderm. Scores can only be compared among different samples for the same germ layer.
13. Can I analyze somatic non-pluripotent primary cells with the TaqMan® hPSC Scorecard™ Panel?

Somatic non-pluripotent primary cells, such as the parental lines used for iPSC generation, are not pluripotent in nature and their scores will be low. However, the expression of lineage markers will largely rely on homogeneity of the cells. Note that the markers in the panel are designed to evaluate early germ layer specification and not any particular terminally differentiated state.
Part 2: Sample Preparation

14. How long can I store cells harvested in TRIzol® reagent?

You can store homogenized samples at room temperature for a few hours, or at –60 to –70°C for at least one month.

See http://tools.invitrogen.com/content/sfs/manuals/trizol_reagent.pdf for more information.

15. Can I prepare RNA using the RNA kit of my choice?

Yes, but we recommend using Life Technologies products because the hPSC Scorecard™ Panel protocol was developed and optimized using Life Technologies reagents and kits.

16. Can I skip the RNA purification step and use a lysis protocol instead?

Yes, while RNA purification is recommended, it is not strictly required.

![Diagram of sample preparation steps](image)

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>Self-renewal</th>
<th>Ectoderm</th>
<th>Mesoderm</th>
<th>Endoderm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hg TRizol</td>
<td>0.25</td>
<td>-0.69</td>
<td>-0.52</td>
<td>-0.86</td>
</tr>
<tr>
<td>Hg CellstoCt</td>
<td>0.04</td>
<td>-0.45</td>
<td>-0.34</td>
<td>-0.62</td>
</tr>
<tr>
<td>Hg CellsDirect</td>
<td>0.52</td>
<td>-0.91</td>
<td>-0.45</td>
<td>-0.94</td>
</tr>
</tbody>
</table>

Figure 2. All samples, regardless of sample size, pass the test for undifferentiated cells. Cells were cultured in KSR-based media on irradiated MEFs in the culture dish formats shown prior to Scorecard™ analysis. Note that negative scores for germ layer markers in undifferentiated samples just indicates lower levels of these factors relative to the undifferentiated reference standard lines, and is a perfectly acceptable result. The difference between scores of the self-renewal (pluripotency) factors in these samples is unlikely to be statistically significant. A minimum of 15,000 undifferentiated cells or 6 embryoid bodies is required to obtain an answer with the hPSC Scorecard™ analysis using the Cells-to-Ct or CellsDirect protocols. When purifying RNA, a minimum of 100,000 cells is recommended.

Visit thermofisher.com/scorecard for more product information, data, protocols, and troubleshooting tips or contact stemcell-help@lifetech.com for support.
17. Do I need to perform DNase treatment?

We recommend that you include DNase treatment of all samples as a good practice, despite the fact that the majority of primers span across an intron and do not amplify (or in some cases, minimally amplify) contaminating genomic DNA.


18. How long can I store RNA after it is prepared?

Properly prepared RNA will last for years when stored properly at –80°C.

19. How much RNA is required for the RT reaction?

We recommend using 1 μg of RNA per sample.
**Part 3: Sample Loading**

20. Why should I set up cDNA synthesis in 8 wells of a 96-well plate or in 8-well PCR strips?

Setting up your cDNA synthesis in 8 wells of a 96-well plate or in 8-well PCR strips facilitates sample loading of the 96-well and 384-well hPSC Scorecard™ Panel plates. Follow the schematic below:

21. How do I load the samples on to my 384-well plate if I don’t have a multichannel pipette?

You can load samples using a single channel pipette, but this method is time-consuming and may increase pipetting error. We strongly recommend that you use an 8- or 16-channel multichannel pipette. You also can dispense samples using automated systems, with the understanding that additional sample will be required to compensate for the dead volume.

22. How do I load the samples onto the 384-well plates if I only have an 8-channel pipette?

The tips of most 8-channel pipettes will align with every other well so you will need to pipette twice to load every well in each column of the plate. The first set of 8 samples can be loaded in rows A, C, E, G, I, K, M, and O. The second set of 8 samples can then be loaded into the alternate rows B, D, F, H, J, L, N, and P. Follow the schematic below:
23. How do I load the samples onto the 384-well plates if I only have a 16-channel pipette?

The tips of most 16-channel pipettes will align with every well in each column of the plate. However, if your cDNA reactions were set up in 8 wells of a 96-well plate or in 8-well PCR strips, additional sample will be required to compensate for the dead volume. When you insert 2 tips of the 16-channel pipette into 1 well, the tips can’t reach the bottom of the well, resulting in a need for additional dead volume.

Visit thermofisher.com/scorecard for more product information, data, protocols, and troubleshooting tips or contact stemcell-help@lifetech.com for support.
Part 4: Gene Expression Assay

24. What qRT-PCR instruments are currently supported?

ViiA™ 7, QuantStudio™ 12K Flex, QuantStudio™ 7 Flex, QuantStudio™ 6 Flex, StepOnePlus®, 7500 Fast, and 7900HT Fast Real-Time PCR systems are compatible with the hPSC Scorecard™ Panel.

25. What if I don’t have access to a compatible qRT-PCR system?

The hPSC Scorecard™ Panel analysis is also available as a service. You can send Life Technologies isolated RNA samples, cDNA samples, or even cells in TRIzol® reagent.

See lifetechnologies.com/cellmodel for more information.

26. Where can I find the run template files for a compatible qRT-PCR system?

Template files are available for download from the product catalog pages and at:

lifetechnologies.com/scorecardhelp.

27. How do I set up my experimental run?

The template files contain all of the necessary experimental details for you to run your instrument. There is no need to modify any of the experimental details if you are using the template file (.edt). Step-by-step guides on how to run the qPCR instruments are available at:

lifetechnologies.com/scorecardinstrument.

28. Do I need to use the template files?

We highly recommend that you use the template file because it contains all of the necessary experimental details for your qPCR run. If the template file is not working for you, please send an email to our stem cell technical support specialists at stemcell-help@lifetech.com.

29. How do I access the hPSC Scorecard™ Analysis Software?

Please visit lifetechnologies.com/scorecarddata for access to the analysis software.

30. I want to know the list of genes before I run the samples? Where I can find it?

To see the gene names, you’ll need to create a demo project.

a. Log in to the hPSC Scorecard™ Analysis Software at lifetechnologies.com/scorecarddata with your Life Technologies username and password.
b. Click on the encircled link to create your demo project.

![Image of software interface](image)

c. Click the Demo Project name in the Project list.

![Project list image](image)

d. Click on the checkbox next to the name of the project above the results summary panel.

![Results Summary image](image)

e. Click the dark gray “Reports” Button at the upper right of the software.

f. Click the “Excel (XLS)” tab, select the results to include (“Scorecard results”, “Scorecard values”, “CT values”, “Fold change values”, and the Excel file type as either “XLS” or “XLSX”) and then click “Download Data.”

![Excel download image](image)

g. Open the spreadsheet file, and click on the “CT Values” worksheet tab to see the list of gene targets.
31. How many genes are tested per sample?

93. There are 9 self-renewal genes, 74 lineage specific genes, and 10 housekeeping and control genes.

32. I accidentally ran a comparative Ct experiment type instead of a standard curve experiment type. Can I still use my data with the hPSC Scorecard™ Analysis Software?

The hPSC Scorecard™ Analysis Software currently only accepts data collected with the “Standard Curve” experiment type. To change your experiment type, open your data in the instrument software. In the “Set-up” menu, change your experiment type to “Standard Curve.” You may receive a warning message, but that is okay. After you change the experiment type, return to the Analysis menu and reanalyze the data by pressing the green “Analyze” button. After you save your data, you can import the file into the hPSC Scorecard™ Analysis Software.
Part 5: Data Analysis

33. What web browsers are compatible with the analysis software?

The hPSC Scorecard™ Analysis Software is compatible with the following web browsers:
- Internet Explorer® 9
- Safari®
- Google® Chrome™

34. How do I access the hPSC Scorecard™ Analysis Software?

Please visit lifetechnologies.com/scorecarddata for access to the analysis software.

35. I can’t sign-in to my Life Technologies account, what should I do?

Retrieve your username or password using the appropriate links on the sign-in page. If you don’t have a username and password, click “Register Now” in the New Customer section of the log-in page.

36. Where do I find more information about how to use the software?

The user manual is available from the help menu within the software, and at lifetechnologies.com/scorecardsoftware.

37. How do I create a project and upload my data?

Click “Create a Project” to identify its Name, Instrument Type, Block, and Description, click “OK”, and then click “Upload Data” to start using the analysis software.
38. My data isn’t being uploaded, what should I do?

The analysis software validates the data file upon import. If your file is not imported (or you receive an error), click the pencil icon 📏 to edit the project, confirm that you have the correct Instrument Type and Block designated in the project window, and verify that you have used one of the following TaqMan® hPSC Scorecard catalog products: A15870, A15876, A15872, or A15871. The analysis software is compatible only with data obtained with those specific panels.

If you see an error message that identifies that the experiment was not analyzed by the Ct method, follow the steps outlined below.

a. Open the .eds file in your data collection software.
b. Click to open Analysis Settings dialog (from the menu, go to Analyze -> Analysis Settings).
c. Check to see if Ct algorithm is selected.
   i. If Ct is selected, close Analysis Settings dialog.
   ii. If Crt is selected, change it to Ct, and then save and close the Analysis Settings dialog.
d. Click the Analyze button.
e. Save it.
f. Click “Upload Data” to import the new file.

39. What sample type should I select?

The software allows you to easily evaluate pluripotency and/or inductive differentiation one germ layer at a time. To make it a little easier, a set of common sample types have been already built into the software to allow you to get to your results faster:

- **All Pluripotency**: Quickly confirm the self-renewal capacity of undifferentiated samples.
- **All Ectoderm / All Endoderm / All Mesoderm**: Evaluate the inductive differentiation specified to one or more germ layers.
- **All Trilineage**: Evaluate all three germ layers simultaneously (e.g. of EB samples).
- **Multiple Sample Types**: Allows you the flexibility to mix and match samples types for analysis.

40. Do I need to run a new sample if I want to look at a different sample type?

No. The software analyzes and retains all of the qPCR data from the plate upon import. If you want to look at a different sample type, simply select the desired sample type from the drop down list, and the analysis software will automatically update the results.
41. My sample has a red flag on it, or is not showing any results. What does that mean?

The hPSC Scorecard™ Analysis Software performs a set of quality controls checks upon data import to ensure the quality of the results. For more information about “hPSC Scorecard assay results” and how to “Interpret results”, refer to the hPSC Scorecard™ Analysis Software Help system.

You may see one of the following red flag warnings:

- **Internal positive control**: The expression levels of the housekeeping genes were low, signifying that there was an issue with the qRT-PCR. The sample is excluded from analysis and must be repeated.
- **Bad Rox**: The ROX™ dye is used as a passive reference during the qPCR. If the ROX™ dye is not detected, it is suggestive that some of the assay wells may not have received the proper amount of Master Mix. This may explain why a particular assay is not showing the anticipated gene expression level.
- **Sendai Virus Detected (SEV)**: The hPSC Scorecard™ Panel includes a control to look for Sendai virus expression. A flag will appear if a Ct value of greater than 30 is observed for this control. On some rare occasions, the SEV oligonucleotides will cross react with other genes resulting in a false SEV flag in differentiated cells. If you see this flag, and did not anticipate Sendai virus detection, you may want to repeat the sample or look at the amplification plot (In the 96 well plate: well B9; in the 384 well plate: wells D3, D9, D15, and D21) to see if the automated Ct value call is real, or just an error.
- **Insufficient Data**: The analysis software requires a minimum number of genes to be amplified in each the four categories (i.e. self-renewal (“pluri”), ectoderm, mesoderm, and endoderm) to accurately analyze your data. If the minimum number of genes in each category does not amplify, or if data has been omitted by the user, the Insufficient Data flag will appear and the sample is excluded from analysis. The sample should be repeated.

42. What lines were used to generate the reference standard?

Thirteen pluripotent stem cell lines were used to generate the reference standard including:

- H9 ESC P28
- iPS BS3C P35
- iPS18C P29
- HUES9 P28
- HUES13 P56
- HUES28 P28
- HUES44 P25
- HUES48 P21
- HUES49 P21
- HUES53 P24
- HUES63 P46
- HUES64 P30
- HUES65 P29

43. How are the scores calculated?

A proprietary algorithm compares the Ct values for each marker set to the values in the reference database and calculates the score based on how well the expression correlates. In general, scores close to 0 indicate comparable expression to that of the reference standard using undifferentiated cells. Scores higher than 1 indicates up regulation relative to undifferentiated cells and less than -1 indicate down regulation relative to undifferentiated cells.
44. **How do I export Ct values in the software?**

   The only way to view the Ct values in the current version is through the expression plot. We expect the software’s Excel export in include the gene names and Ct values in a future version of the software.

45. **My results don’t look right, how do I get help?**

   Please send an email to our stem cell technical support specialists at stemcell-help@lifetech.com.

46. **Will I need to update my software?**

   No software update is ever required. Because the hPSC Scorecard™ Analysis Software is cloud based, you will always have access to the most up-to-date version of the software every time you log in.
Part 6: Examples of Results

The hPSC Scorecard™ Analysis Software algorithm compares the data in the user’s file to the data in the reference cloud and assigns a score based on how closely the data align with the reference set of nine undifferentiated pluripotent stem cell lines. The plus or minus sign in the user interface indicates whether the sample’s score met the criteria for either self-renewal (pluri) or trilineage differentiation (ecto, endo, meso).

For more information about “hPSC Scorecard assay results” and how to “Interpret results”, refer to the hPSC Scorecard™ Analysis Software Help system.

EB time course: H9 ESCs randomly differentiated using EB formation show the appearance of markers that are representative of the 3 germ layers as early as Day 4 of differentiation.

H9 ESCs were culture in KSR-based media on irradiated MEFs prior to removal of FGF. H9 EBs were cultured in suspension over the course of 14 days prior to scorecard analysis. The summary shows that the cells are positive for pluripotency. EB formation at day 2 is positive for ectoderm but negative for meso and endoderm. As the EB formation progresses from day 4 to day 14, the scores for all three germ layers continue to increase, whereas the scores for the self-renewal factors decrease. The table with the scores also reflect the same pattern. You may notice a tiny red flag (💃) in one of boxes, which appears because the software includes quality control checks and lets you know that feedback and additional information is waiting for you.

Random differentiation via EB formation versus monolayer: H9 ESCs differentiated for four days either using EB formation or monolayer differentiation in EB media show appearance of markers for the 3 germ layers by Day 4.
When a culture is allowed to differentiated by removing FGF or not changing media, the expression of SSEA4 drops and the scorecard fails pluripotency but shows trilineage differentiation.

**Early versus late passage iPSC:** BS3C iPSC, an established iPSC clone, is scored as pluripotent and the parental fibroblast displays mesoderm positive expression. The 5 clones derived from BJ fibroblasts using CytoTune reprogramming at passage 5 and 8 are marked pluri negative at early passage but marked pluri positive by P8. This may be due to heterogeneity of the cultures at early stages and therefore iPSC at closer to passage 10 is recommended for the hPSC Scorecard™ Panel analysis. Red flag indicates the presence of SEV that is detected at P5 in some clones but is diluted and not detected at P8. 4 of the 5 clones pass the self-renewal test ("pluri") at P8 while 1 clone fails at P8 and hence is not ideal to be chosen for further culturing.

RNA was isolated from early and late passage iPS clones that were cultured in KSR-based media on irradiated MEFs, then analyzed using the hPSC Scorecard. CytoTune v1 is detected at passage 5 in these clones. They do not clear the Sendai virus ([the tiny red flags ()]) indicate SeV+] and pass pluripotency until passage 8.

Visit [thermofisher.com/scorecard](https://thermofisher.com/scorecard) for more product information, data, protocols, and troubleshooting tips or contact [stemcell-help@lifetech.com](mailto:stemcell-help@lifetech.com) for support.
**Impact of SEV presence on pluripotency:** Clones derived using CytoTune® reprogramming kits that have residual presence of SEV do not affect pluripotency or trilineage differentiation.

Examples of results obtained primary cells can be highly variable since the panel was not designed for use with terminally differentiated cells. In addition, donor-to-donor variation is to be expected.

**Skewed lineage and non-pluripotent somatic cells:** NSCs derived from H9 ESCs show expression of ecto without endo or meso genes. The efficiency and specificity of the differentiation method largely dictates the expression pattern. Non-pluripotent stem cells, such as BJ fibroblast-derived human foreskin cells, show expression in ecto. However, prolonger culture of these cells can create a drift and BJs with ecto and meso positive pattern have also been observed.

**Non human cells and genomic DNA:** Because the primers are human specific and most assays are not amplified, murine embryonic feeders show up as insufficient data (left most data). However, cells harvested along with MEFs may result in an incorrect self-renewal read out since the 10% genes that are amplified can impact the results. It is therefore recommended to harvest ESCs with minimal MEF contamination. Genomic DNA harvested from cells also are called out as insufficient data because most
primers span an intron to prevent amplification of contaminating gDNA. However, as best practice, DNase treatment is recommended.

<table>
<thead>
<tr>
<th></th>
<th>MEFs</th>
<th>H9 ESC + MEFs</th>
<th>H9 ESC</th>
<th>H9 ESC Genomic DNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Insufficient Data</td>
<td><img src="image1" alt="Insufficient Data" /></td>
<td><img src="image2" alt="Insufficient Data" /></td>
<td><img src="image3" alt="Insufficient Data" /></td>
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<tr>
<td></td>
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<td><img src="image6" alt="Insufficient Data" /></td>
<td><img src="image7" alt="Insufficient Data" /></td>
<td><img src="image8" alt="Insufficient Data" /></td>
</tr>
</tbody>
</table>

**Cells in feeder-free media system:** H9 ESC and CytoTune® kit-derived iPSC cultured in feeder-free media system such as StemPro® SFM on Geltrex or Essential 8™ Medium on Vitronectin are scored positive for self-renewal (“pluri”).

Visit thermofisher.com/scorecard for more product information, data, protocols, and troubleshooting tips or contact stemcell-help@lifetech.com for support.
## Part 7: Troubleshooting Information

<table>
<thead>
<tr>
<th>Troubleshooting Guide</th>
<th>Probable Cause</th>
<th>Remedy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pluripotent Failure: My cells are pluripotent but are called out Pluri Fail</strong></td>
<td>Pluripotent lines may have marginal differentiation.</td>
<td>Ensure that the cells are high quality with little to no differentiation at the time of harvest for gene expression analysis.</td>
</tr>
<tr>
<td>Pluri scores are high; Lineage scores are high</td>
<td>Cells may be contaminated with MEF feeders. While most primers are specific for human cells, a few lineage primers cross react with mouse cells and may skew results, especially for the ectoderm layer.</td>
<td>We recommend that you eliminate contamination from the cells cultured under feeder-dependent conditions by performing at least one feeder-free passage.</td>
</tr>
<tr>
<td>Pluri scores are low; Lineage scores are high</td>
<td>There may have been extensive differentiation during culture or clones may have reprogrammed partially or incompletely.</td>
<td>We recommend using stable iPSC clones that show minimal spontaneous differentiation (&lt;P8). If you use lower passage clones, ensure the homogeneity of the culture based on morphology and growth characteristics.</td>
</tr>
<tr>
<td>All values are negative; my cells are negative for pluri and lineage</td>
<td>Cells may be contaminated with genomic DNA (gDNA). Although most primers span across introns, some of the control genes cross react with gDNA. This cross-reactivity is factored in during normalization, resulting in scores that shift toward the negative.</td>
<td>We strongly recommend using DNase treatment. If you use a method of RNA isolation or sample preparation that is not recommended by the provided protocol, it is best to confirm that the sample resulting from the method is of a sufficiently high quality.</td>
</tr>
<tr>
<td>A260/280 and A260/230 values are within range, indicating high RNA quality, and DNase treatment was not performed</td>
<td>Because DNA and RNA each have absorbance at A260, the A260/280 ratio is not a good indicator of the presence of contaminating genomic DNA. The ratio instead can be used to identify the absence of contaminants, such as protein or organic solvents.</td>
<td>We recommend that you design your experiment with a “no RT control” using a recommended test primer (ACTB Hs01060665_g1) to test for gDNA contamination. In the absence of gDNA contamination, the resulting CT value would be close to 40. If gDNA contamination is present, you will observe lower CT values.</td>
</tr>
</tbody>
</table>
**Value Range Definitions**

<table>
<thead>
<tr>
<th>High</th>
<th>Zero</th>
<th>Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>(positive Scorecard value)</td>
<td>(Scorecard value close to 0)</td>
<td>(negative Scorecard value)</td>
</tr>
</tbody>
</table>

**Pluripotent**

When compared to the Reference standard, pluripotent markers are:

- Upregulated
- Comparable
- Downregulated

Depending on germ layer markers, sample may be pluripotent. Sample is not pluripotent.

**Germ Layers: Ectoderm, Mesoderm, Endoderm**

When compared to the Reference standard, germ layer markers are:

- Upregulated
- Comparable
- Downregulated

Sample is differentiated, not pluripotent. Depending on pluripotent markers, sample may be pluripotent.

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