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Specifications

Assay Specifications

Table 1. Assay Specifications

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sizing Range</td>
<td>5 kDa - 80 kDa</td>
</tr>
<tr>
<td>Sizing Resolution&lt;sup&gt;a&lt;/sup&gt;</td>
<td>± 10% 14 - 80 kDa, ± 20% &lt;14 kDa</td>
</tr>
<tr>
<td>Sizing Accuracy</td>
<td>± 20% up to 80 kDa, ± 10% (CAII, BLG)&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Sizing Reproducibility</td>
<td>3% CV (CAII, BLG)</td>
</tr>
<tr>
<td>Linear Concentration Range</td>
<td>30 - 2000 ng/µL (BLG, CAII in PBS)</td>
</tr>
<tr>
<td>Maximum Total Protein</td>
<td>10 mg/mL</td>
</tr>
<tr>
<td>Concentration</td>
<td></td>
</tr>
<tr>
<td>Quantitation Reproducibility</td>
<td>30% CV up to 80 kDa. Above 80 kDa, quantitation is not specified.</td>
</tr>
<tr>
<td>Sample Capacity per Chip</td>
<td>400 samples (four 96-well plates or one 384-well plate)</td>
</tr>
<tr>
<td>Prep</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> Resolution is defined as the height of the valley between two peaks to be no more than 50% of the maximum peak height. Actual separation performance can depend on the sample and application.

<sup>b</sup> CAII = Carbonic Anhydrase, BLG = beta-Lactoglobulin

Sample Conditions

Table 2. Sample Conditions

<table>
<thead>
<tr>
<th>Condition</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buffers, Salts and Additives</td>
<td>Refer to “Buffer, Salt and Additive Compatibility” on page 30 for compatibility with specific buffers, salts and additives. If your conditions are not listed, contact PerkinElmer Technical Support (see page 32) for more information on compatibility.</td>
</tr>
<tr>
<td>Particulates</td>
<td>Sample plates should be spun down prior to analysis. All buffers should be filtered with a 0.22 µm cellulose acetate filter.</td>
</tr>
<tr>
<td>Salt Concentration</td>
<td>Total salt concentration must not exceed 0.5 M</td>
</tr>
</tbody>
</table>
Kit Contents

**Storage:** Prior to use, store chips and reagents refrigerated at 4 °C. After use leave chips at 4 °C for up to 30 days. Store the Low MW Protein Ladder at -20°C.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Vial</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Protein Dye Solution Blue</td>
<td>Blue</td>
<td>1 vial, 0.09 mL</td>
</tr>
<tr>
<td>Sample Buffer</td>
<td>White</td>
<td>2 vials, 1.4 mL each</td>
</tr>
<tr>
<td>Low MW Protein Gel Matrix Red</td>
<td>Red</td>
<td>2 vials, 1.8 mL each</td>
</tr>
<tr>
<td>Low MW Protein Ladder Yellow</td>
<td>Yellow</td>
<td>1 vial, 0.08 mL</td>
</tr>
<tr>
<td>Lower Marker Green</td>
<td>Green</td>
<td>1 vial, 0.5 mL</td>
</tr>
<tr>
<td>Wash Buffer</td>
<td>Purple</td>
<td>4 vials, 1.8 mL each</td>
</tr>
</tbody>
</table>

**Table 4. Consumable Items**

<table>
<thead>
<tr>
<th>Item</th>
<th>Supplier and Catalog Number</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spin Filters</td>
<td>Costar, Cat. # 8160</td>
<td>8</td>
</tr>
<tr>
<td>Centrifuge Tubes</td>
<td>Eppendorf, Cat. # 022363352</td>
<td>5, 2.0 mL</td>
</tr>
<tr>
<td>Ladder Tubes</td>
<td>Genemate, Cat. # C-3258-1</td>
<td>10, 0.2 mL</td>
</tr>
<tr>
<td>Detection Window Cleaning Cloth</td>
<td>VWR, Cat. # 21912-046</td>
<td>1</td>
</tr>
<tr>
<td>Swab</td>
<td>ITW Texwipe®, Cat. # TX758B</td>
<td>3</td>
</tr>
<tr>
<td>Buffer Tubes</td>
<td>E&amp;K Scientific, Cat. # 697075-NC</td>
<td>10, 0.75 mL</td>
</tr>
</tbody>
</table>

**Table 5. High Resolution LabChip**

<table>
<thead>
<tr>
<th>Item</th>
<th>Catalog Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Resolution Chip for use with GXII</td>
<td></td>
</tr>
<tr>
<td>Touch HT</td>
<td>Cat. # 760524</td>
</tr>
<tr>
<td>High Resolution Chip for use with GXII</td>
<td></td>
</tr>
<tr>
<td>Touch 24</td>
<td>Cat. # CLS138951</td>
</tr>
</tbody>
</table>
Safety Warnings and Precautions

WARNING!

For Research Use Only. Not recommended or intended for diagnosis of disease in humans or animals. Do not use internally or externally in humans or animals.

CAUTION

We recommend that this product and components be handled only by those who have been trained in laboratory techniques and that it is used in accordance with the principles of good laboratory practice. As all chemicals should be considered as potentially hazardous, it is advisable when handling chemical reagents to wear suitable protective clothing, such as laboratory overalls, safety glasses, and gloves. Care should be taken to avoid contact with skin or eyes. In case of contact with skin or eyes, wash immediately with water.

WARNING!

- Dye Solution contains DMSO. Avoid contact with skin and eyes.
- Dye Solution contains SDS. Avoid inhalation and contact with skin and eyes.
- Wash Buffer and Sample Buffer contain LDS. Avoid inhalation and contact with skin and eyes.
- Gel Matrix contains Methyl urea. Avoid contact with skin and eyes.
Preparation Procedures

Additional Items Required

- 0.6 mL centrifuge tubes and/or 96-well plates for denaturing protein samples.
- Means for heating samples to 100°C — 96-well PCR instrument or heating block.

*Note:* Avoid using non-stick lab consumables. They may induce unexpected or erratic assay results caused by surface treatments leaching into dye or gel components.

- 18 megohm, 0.22-µm filtered water (Milli-Q® or equivalent).
- 70% isopropanol solution in DI water.
- Reducing agents: BME (beta-mercaptopethanol), 1M DTT (dithiothreitol) or 100 mM TCEP.

*Note:* Allow the chip and all reagents to equilibrate to room temperature before use (approximately 20 to 30 minutes).

Preparing the Gel-Dye and Gel Destaining Solutions

*Notes:* The Dye solution contains DMSO and **must be thawed completely** before use.

The dye is light sensitive. **Do not expose the Dye solution or Gel-Dye to light for any length of time.** Keep the prepared Gel-Dye solution in the dark.

**Do not exceed 9300 rcf when filtering Gel-Dye solution. Exceeding 9300 rcf will change the properties of the gel.**

1. Vortex the thawed Dye Solution for 20 seconds before use.

2. Transfer 520 µL of Low MW Protein Express Gel Matrix (red cap) using a reverse pipetting technique to the 2.0 mL centrifuge tube provided with the reagent kit. Add 20 µL of Protein Dye Solution (blue cap).

3. Vortex the solution until it is well mixed.

*Note:* Gel matrix is extremely viscous. Make sure the Gel-Dye mixture has an even blue color before transferring to the spin filter. **Insufficient mixing of gel and dye will cause inconsistent assay results.**
4 Transfer the solution into one of the spin filters provided with the reagent kit.

5 Transfer 250 µL of the Low MW Protein Express Gel Matrix (red cap) into a separate spin filter. This will be used as the Destain solution.

6 Centrifuge at 9300 rcf for 5 minutes at RT.

7 Discard filters, label and date the tubes

8 Store in the dark at 4°C. Use within 3 weeks.

9 The volumes of Gel-Dye and Destain solutions prepared are the required amount for one chip prep.

Preparing the Sample Denaturing Solution

1 Pipette 700 µL of Sample Buffer (white cap) into a 2.0 mL centrifuge vial.

2 If samples need to be reduced, add 24.5 µL of BME or 1 M DTT or 3.75 µL of 100 mM TCEP.

3 Vortex for 10 seconds. This volume of sample buffer and reducing agent is sufficient to prepare 96 samples. A smaller volume can be prepared if running fewer than 96 samples.

Preparing the Protein Samples and Ladder

Notes: Samples can be prepared in either a 96-well or 384-well PCR plate or in 0.6 mL microcentrifuge tubes (and subsequently pipetted into a plate). Procedures for both are described here.

The Low MW Protein Ladder should be kept frozen. It is recommended that you aliquot the ladder into 12 µL lots for individual use to avoid freeze-thawing. Store at -20°C.

1 For each sample to be analyzed, pipette 7 µL of denaturing solution into the wells of a microtiter plate or into individual 0.6 mL microcentrifuge tubes.

2 Pipette 2 µL of each protein sample into the wells of the 96-well plate or microcentrifuge tube. When finished, cover the plate with foil to minimize evaporation.
3 Allow the Low MW Protein Express Ladder (yellow cap) to thaw completely followed by gentle vortexing for 10 seconds. Briefly spin the ladder vial. Ensure no precipitate is visible in the solution. If precipitate is present, let the vial sit at room temperature for a little longer then repeat the gentle vortex and spin. Pipette 12 µL of Low MW Protein Express Ladder into a microcentrifuge tube or into the well of a microtiter plate.

**Do not add denaturing solution to the ladder.**

4 Denature samples and ladder at 100°C for 5 minutes. Optimum denaturing conditions may vary by sample type.

5 Tap or spin the sample plate to move the fluid to the bottom of the wells.

6 Spin the Low MW Protein Express Ladder (and sample tubes if used) for 15 seconds using a mini-centrifuge.

7 Add 35 µL of water (Milli-Q® or equivalent) to each sample well or sample tube and mix by pipetting up and down a few times. Avoid creating air bubbles. This step should not be done more than an hour before starting the assay. Vortex the sample tubes (if used) for a few seconds. If using a plate, a pipettor or plate shaker can be used to mix the water with the samples.

8 Add 120 µL of water (Milli-Q® or equivalent) to the Low MW Protein Express Ladder. Vortex the ladder mixture for a few seconds to achieve good mixing.

9 If the samples are prepared in tubes, transfer 44 µL of each sample onto a 96-well plate.

10 Spin the sample plate to eliminate bubbles and move the fluid to the bottom of the wells.

11 Place the sample plate onto the instrument's plate holder.

12 Transfer 120 µL of prepared ladder to the provided 0.2 mL Ladder Tube. Ensure there are no air bubbles in the Ladder Tube.

13 Insert the Ladder Tube into ladder slot on the LabChip GXII Touch instrument.

**Preparing the Buffer Tube**

1 Add 750 µL of Wash Buffer (purple cap) to the 0.75 mL Buffer Tube provided with the reagent kit. Ensure there are no air bubbles in the Buffer Tube.
2 Insert the Buffer Tube into the buffer slot on the LabChip GXII Touch instrument.

**Note:** Replace the Buffer Tube with a freshly prepared tube every 8 hours when the chip and instrument are in use.

![Figure 1. Locations of the Buffer Tube and Ladder Tube in the GXII Touch instrument.](image)

**Preparing the Chip**

**Note:** Use the High Resolution Chip for either Low MW Protein or Glycan Profiling, but not both assays on the same chip.

1 Allow the chip to come to room temperature.

2 Use a pipette tip attached to a vacuum line to thoroughly aspirate all fluid from the chip wells (see Figure 2). For more details on how to set up a vacuum line see page 31.

3 Each active chip well (1, 2, 3, 4, 7, 8, 9 and 10) should be rinsed and completely aspirated twice with water (Milli-Q® or equivalent). Do not allow active wells to remain dry.

4 If any water spilled onto the top or bottom of the chip surfaces during rinsing, aspirate using the vacuum line. DO NOT run the tip over the central region of the detection window. Use the provided Detection Window Cleaning Cloth to clean the chip detection window.
Figure 2. Using a vacuum to aspirate the chip wells is more effective than using a pipette. See page 31 for more details.

5 Using a reverse pipetting technique, add Gel-Dye solution to chip wells 3, 7, 8 and 10 as shown in Figure 3.

Figure 3. Reagent placement.

6 Using a reverse pipetting technique, add Destain solution to chip wells 2 and 9 as shown in Figure 3.

7 If the chip will be used to analyze multiple 96-well plates or will be in use for up to 8 hours, add 120 µL of Protein Express Lower Marker (green cap) to chip well 4. If the chip will only be used to analyze one 96-well plate or a partial plate and then stored for future use, the marker volume can be reduced to 50 µL. Make sure the marker volume is pipetted accurately. If there is not enough marker in chip well 4, the marker will deplete and will not be added to subsequent samples on-chip. Data collected without marker peaks cannot be analyzed by the software.

8 Make sure the rims of the chip wells are clean and dry.
9 IMPORTANT: Ensure chip well 1 (waste well) is empty before placing the chip into the instrument.

Inserting a Chip into the LabChip GXII Touch Instrument

1 Check that the sample plate, Buffer Tube, and Ladder Tube are placed on the instrument properly.

2 Remove the chip from the chip storage container and inspect the chip window. Clean BOTH sides of the chip window with the PerkinElmer-supplied clean-room cloth dampened with a 70% isopropanol solution in DI water.

3 Touch the Unload Chip button on the Home screen.

4 Insert the chip into the LabChip GXII Touch instrument (Figure 5) and close the chip door securely.
Figure 5. Chip in the LabChip GXII Touch instrument.

5 Touch the Load Plate button on the Home screen (Figure 4) to retract the sample plate and send the sipper to the Buffer Tube.

*Note:* Do not keep the chip door open for any length of time. Dye is sensitive to light and can be photobleached.

Running the Assay

*Note:* Chips can be primed independently from running assays. Select the assay of choice from the insert (see Figure 7). Touch the Prime button on the Home screen. Make sure the Buffer Tube is placed on the instrument.

Figure 6. Chip priming screen.

1 Touch the Run button (see Figure 6).
2 Select the appropriate assay type (see Figure 7), plate name, well pattern, and whether to read wells in columns or rows. Select number of times each well is sampled under Adv. Settings (Figure 8). Touch the green arrow.

![Figure 7. The Assay Choices menu.](image1)

3 In the Setup Run tab, select the operator name, the option to read barcode, the destination of the file, the inclusion of sample names, expected peaks, and excluded peaks and the filename convention. Select Auto Export to export results tables automatically (Figure 9). Touch the green arrow.

![Figure 8. Selecting wells.](image2)
4 Touch **Start** to begin the run.
Storing the Chip

After use, the chip must be cleaned and stored in the chip container.

1 Place the chip into the plastic storage container. The sipper should be submerged in the fluid reservoir.

2 Remove the reagents from each well of the chip using vacuum.

3 Each active well (1, 2, 3, 4, 7, 8, 9 and 10) should be rinsed and aspirated twice with water (Milli-Q® or equivalent).

4 Add 120 µL water (Milli-Q® or equivalent) to the active wells.

5 Cover the wells with Parafilm® to prevent evaporation and store the chip at room temperature until next use. The chip must be used to its lifetime (to the total number of 400 samples) within 30 days of analyzing the first plate of samples.

Chip Cartridge Cleaning

1 Daily

   a Inspect the inside of the chip cartridge and O-rings for debris.

   b Use the provided lint-free swab dampened with water (Milli-Q® or equivalent) to clean the O-rings using a circular motion. If the O-rings stick to the chip or a pressure leak is detected, perform the more extensive monthly cleaning procedure.

2 Monthly

   a To reduce pressure leaks at the chip interface, clean the O-rings frequently. Remove the O-rings from the top plate of the chip interface on the LabChip GXII Touch instrument. Soak O-rings in water (Milli-Q® or equivalent) for a few minutes. Clean the O-ring faces by rubbing between two fingers. Wear gloves.

   b To reduce the occurrence of current leaks, clean the chip interface frequently. Clean the top plate of the chip interface using the provided lint free swab dampened with water (Milli-Q® or equivalent).

   c Allow the O-rings and chip interface to air dry. Reinsert the O-rings into the chip cartridge.
Results

Low MW Protein Express Ladder Result

The electropherogram of a typical Low MW Protein Express ladder is shown in Figure 11. Peaks to the right of the lower marker and system peaks in order of increasing migration time correspond to proteins of increasing size i.e. 6.5 kDa, 15.9 kDa, 20.4 kDa, 28.9 kDa, 48.4 kDa, and 68.4 kDa respectively.

Figure 11. Low MW Protein Express ladder electropherogram.
Troubleshooting

Note: Some of the data examples shown in this section were generated with assays other than the assay described in this user guide.

Symptom: No ladder or sample peaks but marker peaks detected.

Note: The lower marker peak height will most likely be greater than normal height.

Possible causes:

1 Air bubble in sipper introduced during chip priming.

What to do:

1 Reprime the chip. See “LabChip Kit Essential Practices” on page 24 for instructions on how to reprime the chip.

Symptom: Missing sample, ladder and marker peaks.

Possible causes:

1 Clog in sipper or marker channel of chip.

What to do:

1 Reprime the chip. See “LabChip Kit Essential Practices” on page 24 for instructions on how to reprime the chip.

Symptom: Ladder detected but no sample peaks.

Possible causes:

1 The sipper is not reaching the sample due to low sample volume in the well of the plate.

2 If the missing sample peaks occurred only in a few wells of the plate, check those wells for air bubbles.

3 The sipper is not reaching the sample due to an incorrect capillary height setting or incorrect plate definition.

4 If the plate has been uncovered for some time, sample evaporation might have occurred.

5 Debris from the sample or sample prep is clogging the sipper.

What to do:

1 Add more sample to the well.
2 Manually insert a larger volume pipette tip (~100 µL) into the sample well and dislodge the bubble. Rerun these sample wells.

3 Check the plate definitions.

4 Check the sample wells, especially around the edge of the plate where evaporation is fastest, and make a fresh plate if volumes are low.

5 If you suspect there may be debris in your samples, spin the sample plate down in a centrifuge (e.g., 3000 rcf for 5 minutes). Unclog the sipper by repriming the chip. See “LabChip Kit Essential Practices” on page 24 for instructions on how to reprime the chip.

Symptom: No ladder peaks but sample peaks and marker peaks are present.

Possible causes:

1 Low or no ladder volume in the Ladder Tube.

What to do:

1 Add more ladder to the Ladder Tube and restart the run. Recommended standard ladder volume is 120 µL (minimum volume is 100 µL).

Symptom: No marker peaks but sample peaks are present.

Possible causes:

1 No marker added to chip well 4.

2 If there is marker solution in chip well 4, the problem may be due to a marker channel clog.

What to do:

1 This may be due to not filling marker well or chip remaining idle on instrument for extended period of time. Add or replenish the marker solution in the chip using the following procedure:
   • Touch the Unload Chip button on the Home screen to open the chip door.
   • Return the chip to the chip container ensuring the sipper is immersed in fluid.
   • Thoroughly aspirate all fluid from chip well 4 using a vacuum line.
   • Ensure that chip well 4 is rinsed and completely aspirated twice with water (Milli-Q® or equivalent).
   • Add Marker Solution (green cap) to chip well 4.
Troubleshooting

- Reinsert the chip back into the instrument.
- Restart the run.

2 Perform a marker channel unclogging procedure by repriming the chip. See “LabChip Kit Essential Practices” on page 24 for instructions on how to reprim the chip.

Symptom: Ladder traces show up in the lanes following the ladders (delayed sip).

Possible causes:
1 Separation channel overloaded with sample.
2 Partial clog in the separation channel.

What to do:
1 Lower the starting sample concentration.
2 Reprime the chip. See “LabChip Kit Essential Practices” on page 24 for instructions on how to reprim the chip.
Symptom: Unexpected sharp peaks.

![Figure 13. Unexpected sharp peak.](image)

Possible causes:
- Dust or other particulates introduced through sample or reagents.

What to do:
1. Do one or all of the following:
   - Replace the 18 megohm, 0.22-µm filtered water (Milli-Q® or equivalent) water used for chip preparation.
   - Replace the buffer used for sample and reagent preparation.
   - Use a 0.22-micron filter for all water and buffers used for chip, sample, and reagent preparation.
   - Spin down sample plate to pellet any particulates.

Symptom: Humps in several electropherograms which do not correspond to sample data.

![Figure 14. Humps in several electropherograms.](image)
Possible causes:

1. Electrode 7 is dirty and has contaminated the Gel-Dye mixture in well 7.

2. High concentrations of detergent in the sample buffer can sometimes cause humps in the electropherogram.

What to do:

1. Before restarting the run, clean electrode 7. Remove the chip and follow the electrode cleaning procedure. We recommend using the provided swab and isopropanol to manually clean electrode 7.

2. Lower the detergent concentration in the sample (see “Buffer, Salt and Additive Compatibility” on page 30).

Symptom: Peaks migrating much faster or slower than expected.

Note: Some migration time variance between chips or within a plate is considered normal chip performance. All chips are QC tested at PerkinElmer prior to shipment.

Normal migration time windows for the markers are:

- **Low MW Protein Express Assay Lower Marker:** 15 - 17.5 seconds
- **Upper Ladder Protein on the first plate:** 31.5 - 34 seconds
- **Upper Ladder Protein on the third plate:** 29.5 - 32 seconds

Possible causes:

1. Incorrect Gel-Dye ratio. Migration time is sensitive to dye concentration and peaks will migrate too fast or too slow if the dye concentration in the gel is too low or too high, respectively.

   Note: Excess dye within the separation channel will slow down migration, and less dye in the separation channel will make peaks migrate faster.

2. Particulates from the samples may be clogging the separation channel (this will slow down migration).

3. Gel-Dye was not primed properly into the chip.
What to do:

1 Prepare a fresh Gel-Dye solution. Wash and reprime the chip with the new Gel-Dye mixture. See “LabChip Kit Essential Practices” on page 24 for instructions on how to wash and reprime the chip.

2 If fast or slow migration is observed repeatedly on a new chip, contact technical support to arrange return of the chip to PerkinElmer. Please send a data file showing the failure along with the return request.

3 Minimize the loading of particulates in the sample by performing a centrifuge spin of the sample plate (e.g. 3000 rcf for 5 minutes) and/or ensuring the Sip 4 mm plate type is selected in the Select Wells screen before starting a new run. The debris may be flushed out of the chip by washing and re-priming the chip. See “LabChip Kit Essential Practices” on page 24 for instructions on how to wash and reprime the chip.

4 Check the O-rings on the top surface of the chip interface and clean if necessary.

Symptom: High baseline fluorescence (e.g., greater than 1000 counts).

Possible causes:

1 The destain wells (2 and 9) do not contain Destain solution (gel matrix with no dye).

2 The destain wells (2 and 9) may have been contaminated with dye either because the well was improperly flushed after priming or because dye was mistakenly pipetted into the well.

What to do:

1 Prepare a fresh Destain solution. Wash and reprime the chip with the new Destain solution. See “LabChip Kit Essential Practices” on page 24 for instructions on how to wash and reprime the chip.

Symptom: Lower than expected signal for ladders and samples.

Possible causes:

1 Improper SDS concentration in Gel-Dye matrix.
What to do:

1. Ensure that Dye Concentrate is completely thawed and mixed. Prepare a fresh Gel-Dye solution. Wash and reprime the chip with the new Gel-Dye mixture. See “LabChip Kit Essential Practices” on page 24 for instructions on how to wash and reprime the chip.
LabChip Kit Essential Practices

To ensure proper assay performance, please follow the important handling practices described below. Failure to observe these guidelines may void the LabChip Kit product warranty.¹

Note: It is important to keep particulates out of the chip wells, channels and capillary. Many of the following guidelines are designed to keep the chips particulate-free.

For assay and instrument troubleshooting, refer to the LabChip GX Touch Software Help file or call PerkinElmer Technical Support at 1-800-762-4000.

General

- Allow the chip, sample plate and all reagents to equilibrate to room temperature before use (approximately 20 to 30 minutes).
- Clean the O-rings in the chip interface weekly and the electrodes daily. Refer to the Instrument Users Guide Maintenance and Service section for procedures.
- Avoid use of powdered gloves. Use only non-powdered gloves when handling chips, reagents, sample plates, and when cleaning the instrument electrodes and electrode block.
- Calibrate laboratory pipettes regularly to ensure proper reagent dispensing.
- Only the PerkinElmer-supplied clean room cloth can be used on the chip to clean the detection window.
- Water used for chip preparation procedures must be 18 megohm, 0.22-µm filtered water (Milli-Q® or equivalent).
- Using the “Reverse Pipetting Technique” (described next) will help avoid introducing bubbles into the chip when pipetting the gel.

¹ PerkinElmer warrants that the LabChip Kit meets specification at the time of shipment, and is free from defects in material and workmanship. LabChip Kits are warranted for 60 days from the date of shipment. All claims under this warranty must be made within thirty days of the discovery of the defect.
Reverse Pipetting Technique

Figure 15. Reverse pipetting.

1. Depress the pipette plunger to the second stop.
2. Aspirate the selected volume plus an excess amount from the tube.
3. Dispense the selected volume into the corner of the well by depressing plunger to the first stop.
4. Withdraw the pipette from the well.

Reagents

- Store reagents at 4°C when not in use.
- Protect the dye, Gel-Dye mixture and marker from light. Store in dark and at 4°C when not in use.
- The Gel-Dye mixture expires 3 weeks after preparation.
- For optimal performance, use one reagent kit per chip. The Low MW Protein Express Reagent Kit contains the reagents to run four 96-well plates or four chip preparations, whichever comes first.

Chips

Repriming Chips

- Touch the Unload Chip button on the Home screen to open the instrument door. Place the chip into the instrument.
- Close the chip door securely and choose the corresponding assay.
- Touch the Prime button on the Home screen to reprime the chip.
Washing Chips

**Important Note:** Wash chips only with water (Milli-Q® or equivalent). Use of any other reagents (including Wash Buffer) is likely to cause even more artifacts in subsequent data.

**Notes:** Some protein samples may have components which produce data with extra peaks, spikes or other artifacts. When these artifacts are present, washing chips on the LabChip GXII Touch immediately before the next use can often restore data quality.

Chips should only be washed on the LabChip GXII Touch immediately before they are prepared with fresh reagents and primed on the instrument. Chips should not be washed and left with water in the chip channels for any extended period of time.

For most protein samples, the only chip cleaning protocol that is required is to rinse and aspirate the active wells twice with water (Milli-Q® or equivalent), and store the chip with 120 µL of water in each active well.

- Thoroughly aspirate all fluid from the chip wells using a vacuum line.
- Each active well (1, 2, 3, 4, 7, 8, 9 and 10) should be rinsed and completely aspirated twice with water (Milli-Q® or equivalent). Do not allow active wells to remain dry.
- Add 120 µL of water (Milli-Q® or equivalent) to each active well (1, 2, 3, 4, 7, 8, 9 and 10).
- Touch the *Unload Chip* button on the *Home* screen and place the chip into the instrument.
- Close the chip door securely.
- Transfer 750 µL of water (Milli-Q® or equivalent) into the Buffer Tube. Install into the instrument.
- Touch the *Wash* button (*Figure 16*).
Figure 16. Wash screen.

- After completion of the wash cycle, open the chip cartridge and return the chip to the chip container ensuring the sipper is immersed in fluid.
- Thoroughly aspirate all fluid from the chip wells using a vacuum line.
- Replace fluid in the wells with freshly made reagents as described in “Preparing the Chip” on page 9. Do not let wells remain dry.
- Transfer 750 µL of Wash Buffer (purple cap) into a clean Buffer Tube. Install into the instrument.
- Install the Ladder Tube, sample plate and chip into the instrument and run the assay.
- If air bubbles are not dislodged after a reprime, apply a vacuum to the sipper. Perform this by filling all active wells with 100 µL water (Milli-Q® or equivalent). Then suction the sipper with a vacuum line, as shown in Figure 17, until droplets of fluid flow out from the sipper. When suctioning the sipper, be careful not to bend or break the sipper. To facilitate this, cut the end of the pipette tip attached to the vacuum line to widen the mouth.
Figure 17. Removing an air bubble or clog by suctioning the sipper with a vacuum line.

Other Considerations:

- New chips should be stored refrigerated prior to first use.
- After running the first set of samples, chips must be stored at room temperature and used within 30 days.
- Do not allow the liquid in the chip container to freeze, as this may lead to poor chip performance. Do not submerge the chip in any solution.
- The entire chip surface must be thoroughly dry before use.
- The sipper must be kept immersed in fluid at all times and should not be exposed to an open environment for long periods of time.
- Use care in chip handling to prevent sipper damage. Damage to the sipper can result in inconsistent sampling.
- Avoid exposing the chips to dust by keeping them in a closed environment such as in the chip container or in the instrument before and after chip preparation.
- Chips can be prepared and left in the instrument for extended periods of time so that samples can be run as needed throughout the day. PerkinElmer recommends the chip be re-prepared after it has been idle for 8 hours, but the chip can be used continually over an 8-hour work day as long as the maximum recommended idle time of 8 hours and total chip lifetime of 400 samples are not exceeded.
Samples

- Prepared sample plates should be free of gas bubbles and particulate debris, both of which may inhibit sipper flow.
- Sample plates containing gas bubbles and/or particulate debris should be spun down at 3000 rpm (1250 rcf) prior to analysis.
- Up to four 96-well plates (400 samples) can be run with a single chip preparation when running the GXII Touch HT instrument. Up to 48 samples can be run with a single chip preparation when running the GXII Touch 24 instrument.
## Buffer, Salt and Additive Compatibility

### Table 6. Compatible Buffers, Salts and Additives

<table>
<thead>
<tr>
<th>Buffer &amp; Salts</th>
<th>Concentration Limit</th>
<th>Additives</th>
<th>Concentration Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris Chloride</td>
<td>250 mM</td>
<td>Octyl Glucoside</td>
<td>2.5%</td>
</tr>
<tr>
<td>Tris Glycine</td>
<td>250 mM</td>
<td>Pluronic F68</td>
<td>0.1%</td>
</tr>
<tr>
<td>HEPES</td>
<td>500 mM</td>
<td>Sarcosyl</td>
<td>1.25%</td>
</tr>
<tr>
<td>PBS</td>
<td>2 X</td>
<td>CHAPS</td>
<td>0.25%</td>
</tr>
<tr>
<td>Sodium Citrate</td>
<td>150 mM</td>
<td>Tween 20</td>
<td>0.4%</td>
</tr>
<tr>
<td>Sodium Phosphate</td>
<td>250 mM</td>
<td>Triton X-100</td>
<td>0.6%</td>
</tr>
<tr>
<td>Sodium Acetate</td>
<td>600 mM</td>
<td>SDS</td>
<td>2%</td>
</tr>
<tr>
<td>Sodium Chloride</td>
<td>500 mM</td>
<td>Zwittergent 3-14</td>
<td>0.4%</td>
</tr>
<tr>
<td>Sodium Azide</td>
<td>1.5%</td>
<td>PEG 3350</td>
<td>1%</td>
</tr>
<tr>
<td>Sodium Hydroxide</td>
<td>125 mM</td>
<td>Glycerol</td>
<td>30%</td>
</tr>
<tr>
<td>Potassium Chloride</td>
<td>225 mM</td>
<td>Urea</td>
<td>8 M</td>
</tr>
<tr>
<td>Ammonium Bicarbonate</td>
<td>1000 mM</td>
<td>Sucrose</td>
<td>1 M</td>
</tr>
<tr>
<td>Magnesium Chloride</td>
<td>37.5 mM</td>
<td>DMSO</td>
<td>25%</td>
</tr>
<tr>
<td>Imidazole</td>
<td>900 mM</td>
<td>EDTA</td>
<td>50 mM</td>
</tr>
<tr>
<td>PhosphoSafe</td>
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<td>Ethanol</td>
<td>50%</td>
</tr>
<tr>
<td>BugBuster</td>
<td>1 X</td>
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</tr>
<tr>
<td>BPER</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>POP Culture</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insect POP Culture</td>
<td>0.25 X</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Table 7. Incompatible Buffers, Salts and Additives

<table>
<thead>
<tr>
<th>Buffer &amp; Salts</th>
<th>Concentration Limit</th>
<th>Additives</th>
<th>Concentration Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>RIPA</td>
<td>All</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Chip Well Aspiration Using a Vacuum

Aspirating with a pipette can leave used reagents in the chip wells. For this reason, PerkinElmer recommends vacuuming the wells instead. This can be accomplished by attaching a permanent pipette tip to a house vacuum line with trap (Figure 18). To avoid contamination, use a new disposable pipette tip over the permanent tip for each chip aspirated (Figure 19).

Figure 18. A: Permanent pipette tip attached to a house vacuum line; B: vacuum line with trap.

Figure 19. Replacing the disposable pipette tip.
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For additional assay and instrument troubleshooting, refer to the LabChip GX Touch Software Help file.
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