Introduction
The Agilent 2200 TapeStation system is a complete solution for automated, fast, and reliable electrophoresis. It is made up of three elements: the ScreenTape consumable, the 2200 TapeStation instrument, and the analysis software. The 2200 TapeStation system is straightforward to use; simply place the sample tubes and ScreenTape consumable in the instrument and let it load, separate, image, analyze, and present the results in approximately 1–2 minutes per sample.

The 2200 TapeStation system offers scalable throughput and rapid results, making it an ideal solution for quality control of biological samples in next-generation sequencing (NGS), microarray, and quantitative PCR workflows. During assay preparation, the instructions regarding reagent preparation and instrument maintenance must be strictly followed. Important technical points are described in the TapeStation User Manual and other supporting documentation. This Technical Overview describes techniques for ensuring reliable quantification and sizing results using DNA assays on the 2200 TapeStation system.
**Experimental**

**Materials**

The 2200 TapeStation system (G2964AA/G2965AA), D1000 ScreenTape and Reagents (5067-5582 and 5067-5583), and Genomic DNA ScreenTape and reagents (5067-5365 and 5067-5366), were obtained from Agilent Technologies (Waldborn, Germany). The IKA MS 3 basic vortexer with PCR plate adapter (4674100) was purchased from IKA GmbH & Co. KG (Staufen, Germany www.ika.com).

**Quantification**

Acquiring DNA concentration data is essential for sample QC. To ensure accurate quantification from the 2200 TapeStation system, it is important to note the following:

**Sample mixing**

By taking the D1000 and High Sensitivity D1000 ScreenTape as an example, the protocol states that sample and sample buffer are vortex mixed using the IKA vortexer and adaptor at 2,000 rpm for 1 minute to ensure proper mixing. After vortex mixing, it is recommended that the samples be collected at the bottom of the tube by brief centrifugation. The effects of insufficient mixing were investigated using D1000 ScreenTape. Reagents and results are presented in Figures 1 and 2, as well as Table 1.

It is clearly shown that poor mixing dramatically affects the reported sample concentration, and that following the recommended procedure is the best way to attain accurate quantification results.

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**Figure 1.** A) shows the electropherogram and B) the gel image of the D1000 ScreenTape mixing tests. In both panels the green trace shows recommended protocol of vortex mixing using the IKA vortexer and adaptor at 2,000 rpm for 1 minute followed by brief centrifugation; Blue shows results for pipette mixing only; Red shows the effect of no mixing. Images were taken from the Agilent 2200 TapeStation Analysis Software.

**Figure 2.** Chart of reported concentrations for the D1000 ScreenTape mixing tests. Concentrations are expressed as a percentage of the theoretical for the three mixing methods. As above, the green bar represents the recommended protocol of vortex mixing using the IKA vortexer and adaptor at 2,000 rpm for 1 minute followed by brief centrifugation; Blue - pipette mixing only; Red - no mixing.

<table>
<thead>
<tr>
<th>Measured concentration (ng/µL)</th>
<th>Theoretical concentration (ng/µL)</th>
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<tbody>
<tr>
<td>Vortex mixing then centrifugation</td>
<td>68.45</td>
</tr>
<tr>
<td>Pipette mixing only</td>
<td>21.20</td>
</tr>
<tr>
<td>No mixing</td>
<td>6.54</td>
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</table>

**Table 1.** Agilent D1000 ScreenTape mixing tests. Quantification values obtained from the Agilent 2200 TapeStation Analysis Software and the Agilent D1000 ScreenTape when using the correct protocol (vortex mix followed by brief centrifugation) and two incorrect mixing protocols (pipette mixing only and no mixing).
Peak integration

For D1000 ScreenTape and High Sensitivity D1000 ScreenTape, the concentration values are calculated using the area of the sample peak compared to the known concentration of the top marker. It is, therefore, essential that all peaks are integrated correctly in the 2200 TapeStation Analysis Software. Ensure that both the upper marker and sample peaks are properly integrated by manually adjusting the peak when necessary. Figure 3 shows examples of correct marker peak integration.

Figure 4 demonstrates the effect of peak integration on the quantitative results. Incorrect peak integration can significantly bias the determined DNA sample concentration.

Figure 3. Correct upper marker integration for the Agilent D1000 ScreenTape and Agilent High Sensitivity D1000 ScreenTape assays.

Figure 4. Example of correct (A) and incorrect (B) sample peak integration, and their effect on reported sample concentration.

NOTE: NGS libraries should be quantified using the region mode of the 2200 TapeStation Analysis software. Please see the following sections for further details.
Use the correct protocol

Each ScreenTape type, for example D1000 ScreenTape and High Sensitivity D1000 ScreenTape, is designed for use with its corresponding reagent kit. It is important that the correct reagents are used with the selected ScreenTape. It is also important that the correct sample preparation protocol is followed exactly, using the correct volumes of sample and sample buffer as seen in the Figure 5. Any variations in the volumes shown can adversely affect the quantification results generated from the system.

In addition, it is important to choose the correct assay based on the concentration of the sample. Using sample concentrations outside the specified quantitative ranges (as detailed in Table 2) will lead to inaccurate quantification.

Use the correct tools for the job

Use pipettes which are calibrated and sufficient for the volume to pipette. A 20-μL pipette will not be as accurate at pipetting 1–2 μL as a 2-μL or 10-μL pipette. Ensuring correct pipetting technique assists in ensuring that the volumes used in each assay are precise, and that the concentrations can be calculated correctly.

Use a vortexer designed for mixing 8-way tube strips, or 96-well plates. TapeStation systems are supplied with an optional IKA MS3 vortexer, which includes a 96-well plate adaptor suitable for both 96-well plates and 8-way strips. This vortexer is recommended for use with the D1000 ScreenTape and High Sensitivity D1000 ScreenTape assays.

It is recommended that TapeStation users obtain the IKA MS3 vortexer for best results. This vortexer can only be obtained directly from IKA (www.ika.com) by quoting the part number 4674100. Agilent Technologies, Inc. will not sell these parts separately. If an IKA MS3 vortexer is not available, ensure thorough mixing by vortex for 10 seconds on maximum speed.

With a 96-well plate, use the 2200 TapeStation foil cover (part number 5067-5154) to prevent the sample from leaving the plate during vortexing.

After vortexing, use an appropriate centrifuge for either 96-well plates, or 8-way strips to ensure that all of the samples are at the bottom of the tube before placing in the 2200 TapeStation.

CAUTION: To avoid damage to the 2200 TapeStation instrument and to ensure correct results, use only the recommended consumables and reagents with the 2200 TapeStation system as listed in the TapeStation User Manual. The use of 96-well plate covers from other suppliers could result in a failure to pierce the film potentially leading to an instrument crash.

<table>
<thead>
<tr>
<th>Quantitative range</th>
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<tbody>
<tr>
<td>Agilent D1000 ScreenTape</td>
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<tr>
<td>0.1 ng/μL–50 ng/μL</td>
</tr>
<tr>
<td>Agilent High Sensitivity D1000 ScreenTape</td>
</tr>
<tr>
<td>10 pg/μL–1,000 pg/μL</td>
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Table 2. Specified quantitative ranges for the Agilent D1000 and Agilent High Sensitivity D1000 assays.
TapeStation analysis in the Agilent SureSelect workflow

The 2200 TapeStation system has been verified for use within the Agilent SureSelect protocol. However, tests have shown that analysis after the purification step, where DNA libraries are purified using AMPure XP beads, can have an effect on quantification. It is postulated that carryover of the beads can cause issues, as the beads would be retained at the top of the separation matrix and contribute to the area of the upper marker (Figure 6). This causes a lower relative reported value for sample concentration. The phenomenon can be identified when there is a visible signal above the upper marker, as seen in Figure 6. A similar profile can also occur when over-amplification of samples causes the signal to run concurrently with the upper marker.

This artifact can be avoided by increasing the time for which the samples are incubated on the magnetic plate to 10 minutes, thereby removing a higher percentage of the beads.

Sizing

Peak maxima versus average molecular weight sizing

The 2200 TapeStation analysis software contains options to display both an Electropherogram view and a Region view. The Region view is designed for analyzing samples that appear as a smear (for example, NGS libraries) and gives slightly different information to that displayed in the electropherogram view (Figure 7). With this in mind, it is important to use the correct function for your samples.

- **Electropherogram view** is designed for use with discrete peaks, and the default size reported is that of the highest point of the peak.

- **Region view** calculates data over a whole smear or region, and reports size as that of the center of the regions’ mass. This gives the user an idea of the distribution of sizes within that sample.

![Electropherogram and Region views](image.png)

Figure 6. Enlarged image of the upper marker showing additional signal from AMPure beads.

![Electropherogram and Region views](image.png)

Figure 7. The sizing data obtained in Electropherogram and Region views of the Agilent 2200 TapeStation analysis software.
Identifying the correct markers
It is important to ensure that the correct upper and lower markers are assigned in the 2200 TapeStation analysis software. The markers are used within the software as internal references in order to determine the molecular weight of each sample peak. Incorrect identification can lead to miscalculations in the reported sizing values.

Figure 8 shows an example of incorrect lower marker identification next to the corrected file. In each of the images, lanes 2 and 3 are the same sample at different concentrations. As seen in Figure 8A, incorrect identification of the lower marker has caused misalignment of the sample; therefore all the fragments are reported with the incorrect sizes. Manually assigning the correct lower marker (Figure 8B) provides accurate sizing information.

Flicking the ScreenTape
Due to the nature of the ScreenTape consumables, bubbles can form in the buffer chamber. If bubbles form at the gel/buffer interface, a loss of performance can be observed in that lane. It is important therefore always to ‘flick’ the ScreenTape before placing it into the 2200 TapeStation instrument to move the bubble to the top of the chamber where it will no longer affect sample loading, see Figure 9.

![Figure 8. Screenshots of the Agilent 2200 TapeStation Analysis Software, showing A) incorrect lower marker identification in the middle lane of the gel image and B) corrected lower marker in the middle lane. The lower marker is always highlighted with green.](image)

![Figure 9. Flicking the ScreenTape consumable removes bubbles from the gel interface.](image)
Always use fresh genomic DNA ladder

When using the Genomic DNA ScreenTape Assay, it is important that the Genomic DNA Ladder is freshly prepared for each run. For tube strips or 96-well plate, the ladder must always be the first well selected to run. Failure to do so will affect the sizing results obtained from the assay.

It is also important to note that there is no Genomic DNA software saved ladder in the 2200 TapeStation Analysis Software.

Effect of shaking the genomic DNA ladder vial

The Genomic DNA Ladder must only be vortex mixed as shaking the ladder vial can degrade the top fragment (Figure 11). Therefore, before pipetting the ladder into the TapeStation approved tube strips or 96-well plate, the ladder vial should be handled carefully, then gently vortexed for 5 seconds to maintain accurate performance.

To minimize shaking during transit, the Genomic DNA reagents are shipped frozen, on dry ice. Once received, these should be kept at 2–8 °C in the refrigerator.

Table 3. The effect of room temperature (RT) equilibrated Genomic DNA reagents as well as cold reagents on the sizing accuracy of the Genomic DNA ScreenTape assay.

<table>
<thead>
<tr>
<th>Reagents at 4 °C</th>
<th>Expected</th>
<th>Reagents at RT</th>
</tr>
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<tbody>
<tr>
<td>Accuracy</td>
<td>–</td>
<td>+ 11 %</td>
</tr>
<tr>
<td>MW (bp)</td>
<td>17,000</td>
<td>18,837 bp</td>
</tr>
<tr>
<td></td>
<td></td>
<td>24,369 bp</td>
</tr>
</tbody>
</table>

Figure 10. The effect of bubbles at the gel interface on the DNA separation profile (lanes A2 and B2).

Figure 11. The effect of shaking the Genomic DNA Ladder. A) Genomic DNA Ladder has been vortex mixed for 5 seconds prior to analysis on the Genomic DNA ScreenTape assay. B) the Genomic DNA Ladder vial was shaken by manually inverting the tube 30 times. Degradation of the top fragment (48,500 bp) is clearly shown.
Conclusion

The Agilent 2200 TapeStation system provides scalable throughput and automation, making it an ideal solution for quality control of biological samples in next-generation sequencing (NGS), microarray, and quantitative PCR workflows.

Following the good measurement practices described in this Technical Overview ensures reliable and accurate DNA quantification and sizing.

References


