



## Long term storage of your parallel CE system

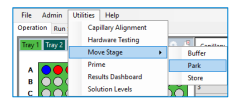
## Preparing the system for prolonged downtime or laboratory closure

For prolonged storage of the system while out of the laboratory the capillary array should be left installed in the system.

- 1. Ensure gel is in the instrument reservoir;
  - a. 5200/5300/5400 Fragment Analyzer, Femto Pulse, ZAG DNA Analyzer: Perform a Full Conditioning.
  - b. Oligo Pro II: Run a Step 3 for 15 minutes with gel only.
- 2. Select **Park** to place the storage plate in the drawer and move the stage to the bottom of the instrument;
  - a. 5200/5300/5400 Fragment Analyzer, Femto Pulse, ZAG DNA Analyzer



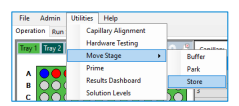
b. Oligo Pro II



- 3. Replace storage solution plate with a new plate
  - a. 5200 Fragment Analyzer, Femto Pulse: Place 1.0mL/well of storage solution into Row H of buffer plate.
  - b. 5300/5400 Fragment Analyzer, ZAG DNA Analyzer, Oligo Pro II systems: Place 200 μL/well of storage solution in fresh sample plate in drawer 3.
    - i) 48 capillary unit rows A-D only
    - ii) 96 capillary unit rows A-H
- 4. Select **Store** to move storage plate under the capillaries.
  - a. 5200/5300/5400 Fragment Analyzer, Femto Pulse, ZAG DNA Analyzer



b. Oligo Pro II



5. Close software, turn off PC and instrument.





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## Powering on system after prolonged downtime or laboratory closure

After prolonged storage of the system it is recommended to perform a Method B - Hot Water Soak of the capillaries followed by a Method C - 0.5N NaOH flush

- 1. Power on the PC, instrument and software.
- 2. Replace the storage solution as described in steps 2-4 above.
- 3. Perform a Method B Hot Water Soak as outlined in the system guide appendix.\*
- 4. Perform a Method C 0.5N NaOH flush as outlined in the system guide appendix.\*
- 5. Prepare fresh conditioning solution and separation gel according to the Kit Guide or Quick Guide, update solution levels in the software and prime lines.

<sup>\*</sup>The Oligo Pro II system does not have appendices for Method B or Method C. Place a buffer tray filled with 1.0 mL/well of hot water (150 °F to 200 °F) in the buffer position and send to this location for 30 minutes.