The TurboBlotter System is a rapid downward blotting device for high-resolution transfer of DNA and RNA. The conventional Northern/Southern transfer stack has been turned upside down in order to take advantage of gravity. No heavy weights are required on the top of the transfer stack, eliminating the messy set-up of standard upward capillary transfers. The TurboBlotter System offers greater speed, target resolution, and convenience versus traditional blotting procedures. Alkaline DNA transfers can be performed in as little as 1 h, while neutral (SSC) transfers of DNA and RNA take only 3 h. TurboBlotter is available in two sizes. The smaller model is 12 × 16 cm, for gels from 7 × 8 cm to 11 × 14 cm. The larger model is 21 × 26 cm, for gels from 12 × 21 cm to 20 × 25 cm. Each size comes in your choice of refill packs. The refill packs are available separately.
Alkaline Transfer of DNA to Nytran™
SuPerCharge Nylon Membranes

Alkaline Transfer Buffers

To make one liter

<table>
<thead>
<tr>
<th>Denaturing Buffer</th>
<th>Transfer Buffer</th>
<th>* Neutralizing Buffer 5x</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 M NaCl, 0.4 M NaOH</td>
<td>3 M NaCl, 8 mM NaOH</td>
<td>1 M phosphate buffer, pH 6.8</td>
</tr>
<tr>
<td>175.5 g NaCl</td>
<td>175.5 g NaCl</td>
<td>79.25 g Na₂HPO₄</td>
</tr>
<tr>
<td>16.0 g NaOH</td>
<td>0.32 g NaOH</td>
<td>60.25 g NaH₂PO₄·H₂O</td>
</tr>
</tbody>
</table>

* This is a 5x concentrate; no pH adjustment is necessary. Make a 1:5 dilution of 5x neutralizing buffer.

To prevent neutralization by CO₂, store the transfer buffer in a tightly closed glass bottle.

Important Note: Alkaline transfers onto Amersham™ Protran™ Supported 0.45 and Amersham Protran Supported 0.2 are not recommended with the TurboBlotter system.

A. Denaturation of DNA gels

1. Incubate the gel in denaturing buffer 2 times, 30 min each. (Shake the dish containing the gel slowly.)
2. Wash the gel in transfer buffer for 15 min.

B. Transfer

1. Soak transfer membrane (Nytran SuPerCharge nylon) in distilled water for 15 min.
2. Place "stack tray" of transfer device on bench, making sure it is level.
3. Place 20 sheets of dry GB003, GB005 or 3MM blotting paper (thick) in stack tray.
4. Place 4 sheets of dry 3MM Chr blotting paper (thin) on top of stack.
5. Place one sheet of prewet 3MM Chr blotting paper in transfer buffer on stack.
6. Place transfer membrane on stack.
7. Cover the membrane with agarose gel; cut the gel to the size of the membrane, making sure there are no air bubbles between the gel and the membrane.

Note: If the gel is smaller than the blotting paper used, cover excess blotting paper with plastic wrap or Parafilm™ to prevent wicking of buffer through the paper rather than the gel.
8. Wet the top surface of the gel with transfer buffer, and place 3 sheets of 3MM Chr blotting paper, presoaked in transfer buffer, on top of the gel.
9. Attach the "buffer tray" of the transfer device to the bottom tray, using the circular alignment buttons to align both trays.
10. Fill the buffer tray with transfer buffer.
   125 ml for 7 × 8 cm to 11 × 14 cm transfers
   200 ml for 12 × 21 cm to 20 × 25 cm transfers
11. Start the transfer by connecting the gel stack with the buffer tray using the precut "buffer wick" (included in each blotter stack), presoaked in transfer buffer. Place the wick across the stack so that the short dimension of the wick completely covers the blotting stack and both ends of the long dimension extend into the buffer tray. Place the "wick cover" on top of the stack to prevent evaporation. Make sure the edges of the wick are immersed in the transfer buffer.
12. Continue the transfer for 1 h. Additional transfer time may be required for gels thicker than 4 mm or larger-sized nucleic acids.

NOTE: Do not place any other weight on top of the "wick cover" during transfer. This is unnecessary and may inhibit transfer by crushing the pore structure of the agarose gel.

C. Neutralization

Following transfer, gently wash the transfer membrane in 1x neutralizing buffer (0.2 M sodium phosphate, pH 6.8) for 5 min.

D. Drying (Immobilization)

1. Place the membrane on a fresh sheet of dry 3MM Chr blotting paper to remove any excess of neutralizing buffer.
2. Bake membrane at 80 °C for 20 min-2 h.

Note: Fixation of DNA requires baking for only 20 min or until the blot has dried completely. As more blots are placed in the oven, more time will be required.
3. As an alternative to drying on Nytran SuPerCharge membrane, the DNA may also be covalently bound to the membrane by cross-linking the molecule to the nylon matrix in the presence of UV light (Church and Gilbert, 1984). Expose the blot to a source of UV light (254 nm) for a total dose of 120 mJ/cm² for a damp membrane.

E. Storage

Store blots desiccated at 4°C for several months.

F. Hybridization

Hybridize and detect using desired method.
Neutral Transfer of DNA or RNA to Amersham Protran Supported 0.45 and Amersham Protran Supported 0.2 or Nytran SuPerCharge Nylon Membranes

**Neutral Transfer Buffers**

To make one liter

<table>
<thead>
<tr>
<th>Denaturing Buffer</th>
<th>20× SSC Transfer Buffer</th>
<th>Neutralizing Buffer</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0.5 M NaOH, 1.5 M NaCl)</td>
<td>3M NaCl, 0.3M Na citrate</td>
<td>1.5M NaCl, 0.5M Tris-HCl pH 7.0</td>
</tr>
<tr>
<td>20 g NaOH</td>
<td>175.5 g NaCl</td>
<td>60.56 g Tris-HCl</td>
</tr>
<tr>
<td>87.66 g NaCl</td>
<td>88.2 g Na3C6H5O7 ·2H2O</td>
<td>87.66 g NaCl</td>
</tr>
<tr>
<td>Bring to 1 L with DI H2O</td>
<td>Bring to 1 L DI H2O</td>
<td>Bring to 1 L DI H2O</td>
</tr>
</tbody>
</table>

**Important Note:** Alkaline transfers onto Amersham Protran Supported 0.45 and Amersham Protran Supported 0.2 are not recommended with the TurboBlotter system.

A. Denaturation of DNA gels

1. Place gel in denaturing buffer for 30 min at room temperature. (Shake the dish containing the gel slowly.)

2. Rinse the gel with distilled water, and transfer to neutralizing buffer. Shake gel slowly for 30 min at room temperature.


   **Note:** If DNA fragments are <500 bp, use 20× SSC transfer buffer. If fragments are >500 bp, 10× SSC transfer buffer is a high-enough ionic strength to retain DNA on nitrocellulose.

B. Treatment of RNA gels

RNA gels that have been run in 2.2 M formaldehyde should be rinsed four times in deionized water and then maintained in low-ionic-strength conditions (DI H2O or low-salt solution) prior to transfer. Alternatively, if the concentration of formaldehyde is 0.41 M (Chomczynski, 1992), no gel washing is required prior to transfer.

C. Transfer

1. Wet and immerse nitrocellulose or nylon membrane (Amersham Protran Supported 0.2 or Nytran SuPerCharge) in distilled water. Then soak membrane in 20× SSC transfer buffer for 5 min.

2. Place “stack tray” of transfer device on bench, making sure it is level.

3. Place 20 sheets of dry GB003 or GB005 blotting paper (thick) in stack tray.

4. Place 4 sheets of dry 3MM Chr blotting paper (thin) on top of stack.

5. Place one sheet of 3MM Chr blotting paper, prewet in transfer buffer on stack.

6. Place transfer membrane on stack.

7. Cover the membrane with agarose gel; cut the gel to the size of the membrane, making sure there are no air bubbles between the gel and the membrane.

   **Note:** If the gel is smaller than the blotting paper used, cover excess blotting paper with plastic wrap or Parafilm to prevent wicking of buffer through the paper rather than the gel.

8. Wet the top surface of the gel with transfer buffer, and place 3 sheets of 3MM Chr blotting paper, presoaked in transfer buffer, on top of the gel.

9. Attach the “buffer tray” of the transfer device to the bottom tray, using the circular alignment buttons to align both trays.

10. Fill the buffer tray with transfer buffer.

11. Start the transfer by connecting the gel stack with the buffer tray, using the precut “buffer wick” (included in each blotter stack), presoaked in transfer buffer. Place the wick across the stack so that the short dimension of the wick completely covers the blotting stack and both ends of the long dimension extend into the buffer tray. Place the “wick cover” on top of the stack to prevent evaporation. Make sure the edges of the wick are immersed in the transfer buffer.

12. Continue the transfer for 3 hr. Additional transfer time may be required for gels thicker than 4 mm or larger-size nucleic acids.

   **Note:** Do not place any other weight on top of the “wick cover” during transfer. This is unnecessary and may inhibit transfer by crushing the pore structure of the agarose gel.

D. Neutralization

Following transfer, gently wash the transfer membrane in 2× SSC for 5 min.

E. Drying (Immobilization)

1. Place the membrane on a fresh sheet of dry 3MM Chr blotting paper to remove any excess of 2× SSC buffer.

2. Bake membrane at 80 °C for 20 min-2 h.

   **Note:** A vacuum oven is recommended for baking of nitrocellulose membranes. Fixation of DNA or RNA requires baking for only 20 min or until the blot has dried completely. As more blots are placed in the oven, more time will be required.

3. As an alternative to drying on Nytran SuPerCharge nylon or Amersham Protran supported nitrocellulose membranes, the DNA or RNA may also be covalently bound to the membrane by cross-linking the molecule to the matrix in the presence of UV light (Church and Gilbert, 1984). Expose the blot to a source of UV light (254 nm) for a total dose of 120 mJ/cm² for a damp membrane.

F. Storage

Store blots desiccated at 4 °C for several months.
G. Hybridization
Hybridize and detect, using desired method.

To make TurboBlotter RNase-free

1. Wash the TurboBlotter device in detergent.
2. Rinse the device completely with DI H2O.
3. Dry with ethanol (EtOH).
4. Soak TurboBlotter device in 3% hydrogen peroxide (H2O2) for 10 min at room temperature.
5. Rinse completely with DI H2O that has been treated with 0.1% DEPC.

Note: Do not treat the TurboBlotter device directly with DEPC or autoclave. We recommend purchasing two separate devices for doing Northern and Southern transfers.

Stacking TurboBlotter trays for multiple transfers
For convenience and to save space, the large TurboBlotter systems can be stacked crisscross for multiple transfers.
We recommend that the stacked trays remain stationary on the bench and not be transported from lab to lab while assembled.

Notes and references
1. It has been demonstrate that proteins can be transferred from agarose gels using the TurboBlotter system. B. Nagy et al. 1995.
2. If target DNA fragment is >15kb, then acid depurination may improve transfer. This step precedes the denaturation step. Soak the gel in 0.25M HCl for 30 min.
3. If DNA fragments are <500bp, use 20x SSC transfer buffer. If fragments are >500bp, 10xSSC transfer buffer is a high enough ionic strength to retain DNA on nitrocellulose.

Ordering details
Each transfer uses one sheet of Nytran SPC, one wick of 3MM Chr, and a blotter stack composed of eight sheets of 3MM Chr and twenty sheets of filter paper. Each TurboBlotter Kit or TurboBlotter refill contains five sets of these materials, enough for five transfers.

<table>
<thead>
<tr>
<th>Description</th>
<th>Product No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>TurboBlotter Kit, large, Nytran SPC 0.45 µm, 15 × 20 cm</td>
<td>10416316</td>
</tr>
<tr>
<td>TurboBlotter Kit, small, Nytran SPC 0.45 µm, 10 × 15 cm</td>
<td>10416300</td>
</tr>
<tr>
<td>TurboBlotter Refill, small, Nytran SPC 0.45 µm, 10 × 15 cm</td>
<td>10416302</td>
</tr>
<tr>
<td>TurboBlotter Kit, small, Nytran SPC 0.45 µm, 11 × 14 cm</td>
<td>10416304</td>
</tr>
<tr>
<td>TurboBlotter Refill, small, Nytran SPC 0.45 µm, 11 × 14 cm</td>
<td>10416306</td>
</tr>
<tr>
<td>TurboBlotter Refill, large, Nytran SPC 0.45 µm, 12 × 21 cm</td>
<td>10416310</td>
</tr>
<tr>
<td>TurboBlotter Refill, large, Nytran SPC 0.45 µm, 15 × 15 cm</td>
<td>10416314</td>
</tr>
<tr>
<td>TurboBlotter Refill, large, Nytran SPC 0.45 µm, 15 × 20 cm</td>
<td>10416318</td>
</tr>
<tr>
<td>TurboBlotter Kit, large, Nytran SPC 0.45 µm, 20 × 25 cm</td>
<td>10416324</td>
</tr>
<tr>
<td>TurboBlotter Refill, large, Nytran SPC 0.45 µm, 20 × 25 cm</td>
<td>10416326</td>
</tr>
<tr>
<td>TurboBlotter Kit, small, Nytran SPC 0.45 µm, 7 × 10 cm</td>
<td>10416328</td>
</tr>
</tbody>
</table>

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