tilibit folding kit basic, type p7249

Store at -20°C

Product No.: K1-1-0

Description:
This kit contains materials for preparing DNA origami self-assembly reactions and a custom gel loading dye for analyzing reaction products using agarose gel electrophoresis. Ultrapure water and staple DNA strands are not included.

Contents:
500 µl 100 nM single-stranded scaffold DNA, type p7249
500 µl tilibit 10x folding buffer XM
500 µl 200 mM MgCl₂ stock solution
1000 µl tilibit 6x gel loading dye

Please refer to individual product data sheets for details.
The gel loading dye does not contain EDTA, SDS, or glycerol and supports straight bands in agarose gel electrophoresis.
Product sheet

Guidelines:
To prepare a ‘standard’ self-assembly reaction with 100 µl volume, mix the following components:

<table>
<thead>
<tr>
<th>vol [µl]</th>
<th>component</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>tilibit 10x folding buffer XM</td>
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</tbody>
</table>

either

| 10       | 200 mM MgCl₂ stock solution (3D DNA origami) |
| or       | 6 200 mM MgCl₂ stock solution (2D DNA origami) |

20 100nM single-stranded scaffold DNA

N staple DNA strand mixture (sold separately)
(for a final staple strand conc. of 200 nM)

either

| 60-N     | ultrapure ddH₂O (3D DNA origami) |
| or       | 64-N ultrapure ddH₂O (2D DNA origami) |

Mix gently, but thoroughly.
Incubate at 65°C for 10 minutes.
Cool from 60°C to 40°C with a rate of 1°C per hour.

The kit is good for 25 reactions.
Gel loading dye: use in 6-fold effective dilution.

Exemplary references for usage:
Rothemund, PWK: “Folding DNA to create nanoscale shapes and patterns” -- Nature. 2006 Mar 16; 440(7082):297-302

Douglas, SM; Dietz, H; Liedl, T; Högberg, B; Graf, F; Shih, WM: “Self-assembly of DNA into nanoscale three-dimensional shapes” -- Nature. 2009 May 21; 459(7245):414-418

Detailed usage recipes: