

Agilent Captiva EMR—Lipid Method Guide

General instructions for 96-well plate and 1 mL cartridge formats:

Agilent Captiva EMR—Lipid cartridges and 96-well plates allow streamlined in-well protein precipitation, filtration, and cleanup for lipid-containing samples. The improved filter design gives easy flow with vacuum or positive pressure, and allows solvent-first protein precipitation without clogging during elution. The novel EMR—Lipid sorbent chemistry provides highly selective and efficient lipid/matrix removal without impacting analyte recovery. Effective matrix removal assures minimal ion suppression or enhancement on target analytes, which significantly improves method reliability and ruggedness.

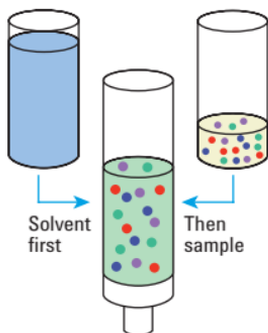
The 96-well plates are ideal for high-throughput workflows, while 1 mL cartridges can accommodate small batch needs.



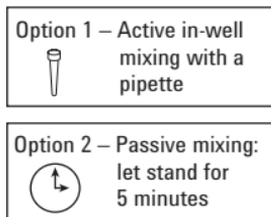
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Operating instructions and tips for Agilent Captiva EMR—Lipid 96-Well Plate and 1 mL Cartridge Products

1. Add crash solvent and sample*



2. Mix to precipitate protein



3. Filter



* Alternatively, protein precipitation (Steps 1 and 2) can be performed off-line (Option 3), at which point the sample can be transferred to step 3.

● Salts ● Proteins ● Lipids ● Analyte



User Tips

Protein precipitation workflow

Product configuration	96-well plate or 1 mL cartridge
Sample size	Between 20–200 μ L
Sample treatment	<p>Crash solvent ratio: between 3:1 and 5:1 ACN + 1 % formic acid to sample. Most commonly 3:1 and 4:1.</p> <p>If total volume is less than 500 μL, add additional 4:1 ACN:H₂O to reach a minimum volume of 500 μL.</p> <p>ACN is preferable to MeOH to maximize protein precipitation and avoid gelation.</p>
Sample addition order	<ol style="list-style-type: none">1) Crash solvent2) Sample
Mixing	<p>Option 1: Active in-well mixing. For in-well protein precipitation, pipette mixing (preferably using wide bore pipette tips) is recommended for 3 to 5 aspiration/dispense cycles.</p> <p>Option 2: Passive mixing. Let stand for 5 minutes to allow for complete protein precipitation to occur.</p> <p>Option 3: Protein precipitation and mixing can be performed in a separate tube, centrifuged, and subsequently transferred to the Captiva EMR—Lipid well/cartridge.</p>
Pass-through filtration and cleanup	<p>Vacuum between 2–5 in Hg initiates flow. Positive pressure (3–4 psi) is also acceptable. For optimal lipid removal, a controlled flow rate of one drop every 3–5 seconds is highly recommended. After elution, apply higher vacuum or positive pressure to ensure maximum sample recovery.</p> <p>Flow rate is dependent on sample type, age, and mixing.</p> <p>An alternative approach to vacuum and positive pressure is centrifugation. For 96-well plates, 500–800 rpm for a minimum of 10 minutes is recommended.</p> <p>Centrifugation speed and time are dependent on the sample volume and matrix.</p>

Agilent Captiva EMR—Lipid ordering information

Part number	Description	Quantity
5190-1000	Agilent Captiva EMR—Lipid 96-well plate*	1 plate
5190-1001	Agilent Captiva EMR—Lipid 96-well plate*	5 plates
5190-1002	Agilent Captiva EMR—Lipid 1 mL cartridge*	100/pk

* Incorporates a non-drip frit to allow in-well protein precipitation

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