

When your kit arrives

The Acrosolv is delivered as a dry powder. When your kit arrives, add DNA-free water as follows.

Kit size (Rxn)	Code	Volume of water to add
100	XSC0100	1.1 ml
500	XSC0500	5.5 ml
1000	XSC1000	11.0 ml

Technical tips for sample management

- The device must be kept clean and DNA-free. The UV cycle will assist in removing DNA but the wells and the top and base of the heating block should be cleaned regularly using dilute HClO bleach on a swab. Wipe residual bleach off the surfaces using a second swab soaked with DNA-free water.
- The device creates DNA that can be used for SNiPs, STRs, quantitative, multiplex and routine PCR applications.
- OD₂₆₀ methods for yield estimation are unsuitable for the DNA produced by the PDQeX.
- For accurate yield assessment, a qPCR is recommended. The DNA produced by the PDQeX is approximately 90% single-stranded. If standard fluorescent chelating dyes are to be used for quantification, then this factor should be taken into consideration.
- For best results prepare and manage samples at 4°C, or on ice before and after extraction.
- For long term storage of the extracted DNA, add TE buffer to 1x and store at -20°C.

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Automated DNA Extraction



PDQeX forensicGEM SexCrime
(for use with PDQeX XSC- kits)

... simple

Spermatozoa Lysis Protocol

Sample Types:

- Sperm or samples suspected of containing sperm
- Touch samples
- Challenging buccal swabs or applications where a high DNA concentration (>10 ng/μL) is required from a buccal swab.

Stains and Swabs:

The processing of the sample will vary dependent on sample type. For liquid samples, try to keep the volume of the liquid below 10 μL. With cotton swabs, add 1/4 of the swab directly to the extraction cocktail. Stained fabric can be swabbed or small portions added directly.

Before you start:

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Reagent Storage:

- Acrosolv -20°C (Limit freeze/thaw cycles)
- forensicGEM -20°C
- ORANGE Plus Buffer 4°C

Method:

1. Completely thaw *forensicGEM* and Acrosolv reagent and mix by gently inverting the tubes repeatedly. Remove ORANGE Plus buffer from refrigerator and mix.
2. Prepare a master mix for sperm lysis. Per 100 μL extraction:

10 μL	Acrosolv
10 μL	ORANGE Plus Buffer
2 μL	<i>forensicGEM</i>
DNA-free water to 100 μL	
3. Dispense 100 μL of master mix into each PDQeX cartridge. Flick the liquid to the bottom of the tube ensuring there are no bubbles.
4. Add the sample, making sure that it is completely submerged in the reagents.

5. Put the cap on the PDQeX cartridge by inserting the tapered peg into the tube.
6. Load a 24-well plate or 8-strip cartridges in the collection drawer and put the drawer in place.
7. Insert the PDQeX cartridges into the heating block.

MAKE SURE THE PDQeX CARTRIDGES CORRESPOND WITH A COLLECTION TUBE OR WELL
8. Cover the tubes with the hinged flap and close the sliding door.
9. Select the "Spermatozoa" program.

Default spermatozoa program:

52°C	5 mins.
75° C	3 mins.
95° C	3 mins.
110° C	2 mins.

- Times may be adjusted by internal laboratory validations.
- Changes to the default temperatures are not recommended.

PRECAUTIONS

1. Do not load the machine if the control screen indicates a temperature above 50°C.
2. Open the sliding door, and ensure the collection drawer and heating block are clean and DNA-free.
3. Ensure the collection drawer is inserted as far as possible, and that it is straight.
4. If less than 24 reactions are planned, ensure that the PCR tubes are placed in the holes in the drawer corresponding to the channels to be used in the heating block.

For more information, visit: www.zygem.com