EMBRYO CULTURE G-TL™

Day 0



G-TI™



OVOIL™

Prepare micro-droplet culture dishes with 25 μL droplets of G-TL for washing and for culture. Cover with OVOIL and pre-equilibrate at

37°C 6 % CO₂ overnight



 $G\text{-}MOPS^{™} PLUS$

Warm G-MOPS PLUS (for fertilisation assessment) in rinsed tightly capped tubes in a warming incubator without CO₂* at

37°C overnight

* An adequately calibrated warming block can be used for tubes instead of a warming incubator.

Ensure that the denudation and washing procedures are performed at 37°C

Day 1

1. Fertilisation assessment

For inseminated oocytes, transfer the oocytes to a centre well dish with pre-warmed G-MOPS PLUS. If denudation and fertilisation assessment can be performed within 2 minutes, G-IVF PLUS can be used instead of G-MOPS PLUS. Remove cumulus and corona cells from oocytes using a denudation pipette and assess fertilisation at

37°C



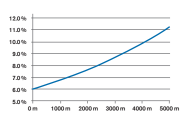
For ICSI oocytes, assess fertilisation in the G-TL micro-droplet culture dish.

2. Culture

Wash the zygotes extensively in the G-TL micro-droplet dishes prepared on Day 0 and transfer the zygotes to 25 µL G-TL culture droplets covered with OVOIL. Culture to the blastocyst stage at

37°C 6 % CO2

If your clinic is located at a higher altitude than sea level, CO_2 percentage should be increased, see graph below.



We recommend G-MOPS PLUS for washing and assessment of embryos before transfer to G-TL or EmbryoGlue®

Day 5

In the morning of day 5

Assess embryo cleavage and move the blastocysts selected for transfer to the equilibrated transfer dish and leave at

37°C 6 % CO₂ until transfer

For blastocyst transfer, see separate Embryo transfer protocol

