

Declaration of Conformity

Manufacturer

Vitrolife Sweden AB Gustaf Werners gata 2 SE-421 32 Västra Frölunda Sweden

Product

G-TL

Intended Use

Medium for culture of embryos from fertilization to the blastocyst stage.

Product category

Medical Device in Risk Class III

Attestation

We hereby declare that the product G-TL fulfills the Essential Requirements as stated in Annex I to the Council Medical Devices Directive 93/42/EEC as amended by 2007/47/EC.

Place and Date

Göteborg on 26 March 2015

Signature

On behalf of Vitrolife Sweden AB,

Hans Lehmann

Director QA/QC

some other