



CE DECLARATION OF CONFORMITY

Application of Council Directives:
89/336/EEC (EMC), 73/23/EEC (LVD), and 2011/65/EU (RoHS 2)

Manufacturer's Name: Sutter Instrument Company

Manufacturer's Address: One Digital Drive
Novato, CA. 94949 USA
Tel: +1 415 883 0128

Equipment Tested: **XenoWorks Microinjection System**

Model(s): **BRI, BRE110, BRE220**

Conforms to Standards: EMI/EMC:
EN 55011, Class B, CISPR 11, CLASS B,
EN 50082-1:1992, IEC 801-2:1991,
IEC 801-3:1984, IEC 801-4:1991

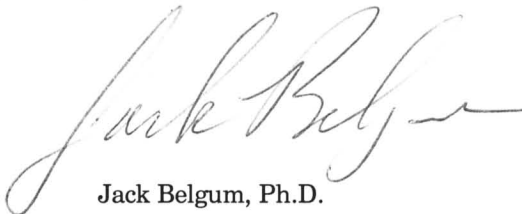
LVD (Safety): EM61010-1:1993

Tested By: TUV Product Service.
10040 Mesa Rim Road
San Diego, CA 92121 USA

Year Tested: 2002, 2015

Sutter Instrument Company hereby declares that the equipment specified above was tested and conforms to the Directives and Standards listed above, and further certifies conformation to the requirements of the European Union's Restriction on Hazardous Substances in Electronic Equipment Directive 2011/65/EU (RoHS 2).

Project Engineer:



Jack Belgum, Ph.D.
Senior Vice President

SUTTER INSTRUMENT

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