



Certificate of CE-Registration

Device Directive 98/79/EC, Medical Device Safety Service GmbH (MDSS) agrees to perform all duties and responsibilities as the Authorized Representative for

Instant Technologies, Inc.
883 Norfolk Square
Norfolk, VA. 23502


USA

as stipulated and demanded by the aforementioned Directive. The German Competent Authorities have allocated the In Vitro Diagnostic Medical Devices of the Manufacturer the following registration numbers:

<u>EDMS Code</u>	<u>Class.</u>	<u>EDMS Description</u>	<u>Registration No.</u>
12 70 09 01 00	other	Amphetamines Group - Rapid Test	DE/CA09/0170/IVD/1497
12 09 01 02 00	other	Amphetamine/Methamphetamine Specific(+Ecstasy) - Rapid Test	DE/CA09/0170/IVD/1498
12 70 09 03 00	other	Barbiturates - Rapid Test	DE/CA09/0170/IVD/1499
12 70 09 04 00	other	Benzodiazepines - Rapid Test	DE/CA09/0170/IVD/1500
12 70 09 05 00	other	Cannabinoids - Rapid Test	DE/CA09/0170/IVD/1501
12 70 09 06 00	other	Cocaine + Cocaine Metabolites - Rapid Test	DE/CA09/0170/IVD/1502
12 70 09 07 00	other	Methadone - Rapid Test	DE/CA09/0170/IVD/1503
12 70 09 08 00	other	Opiates - Rapid Test	DE/CA09/0170/IVD/1504
12 70 09 09 00	other	Phencyclidine - Rapid Test	DE/CA09/0170/IVD/1505
12 70 09 10 00	other	Tricyclic Antidepressants – Rapid Test	DE/CA09/0170/IVD/1506
12 70 09 70 00	other	Multiple Drugs of Abuse/Toxicology Rapid Tests	DE/CA09/0170/IVD/1507
12 70 09 90 00	other	Other Drugs of Abuse/Toxicology Rapid Tests	DE/CA09/0170/IVD/1508

The Manufacturer has provided MDSS with the appropriate Declaration(s) of Conformity confirming that the In Vitro Diagnostic Medical Devices fulfil the applicable requirements of Directive 98/79/EC. In compliance with German law, a safety officer has been appointed for Germany.

23 February 2006


 Ludger Möller
 President
 Medical Device Safety Service GmbH