We, Sakura Finetek Europe B.V., Flemingweg 10A, 2408 AV, Alphen aan den Rijn, The Netherlands

as Authorized Representative herewith declare that:

Equipment: Tissue Processor

Model name/number: Histo-Tek® VP1™ / 1700

Manufactured by:

Sakura Seiki Co. Ltd., 75-5 Imojiya, Chikuma-shi, Nagano-ken, Japan

in accordance with the following Directives:

applied.
equirements of the
and its amending directives.
e Low Voltage Directive and its
of the Restriction of amending directives.

has been designed and manufactured to the relevant parts of the following standards:

EN ISO13485:2012, EN ISO14971:2012, EN 61010-1:2010, EN 61326-1:2013, EN 50581:2012, EN 61010-2-101:2017 and EN 61326-2-6:2013.

In addition the following internal standard applies:

ISO 9001:2015 Quality Management System requirements.

C.Koeman General Manager

I hereby declare that the equipment named above has been tested and found to comply with the relevant sections of the above referenced specifications. The unit complies with all essential requirements of the Directives.

Signed:

Alphen aan den Rijn, 16 October 2018