EC DECLARATION OF CONFORMITY

Manufacturer

Address

Manufacturer: Changsha Renji Medical Equipments Co., Ltd.

Post Add: Building B8, Changsha E Center, No.18 Xiangtai Road.

Liuyang Jingkai District, Changsha City, Hunan

Province, China. 410300

Authorized Representative Name: Lotus NL B.V. Add: Koningin Julianaplein 10, 1e Verd, 2595AA,

European Representative

The Hague, Netherlands.

Product Name: Blood Collection Tube & Vacuum Blood

Product

Collection Tube

Information

No Additive, Clot Activator, Gel & Clot Activator, EDTA+NaF

Lithium Heparin, Sodium Heparin, K2EDTA, K3EDTA,

Na2EDTA, Sodium Citrate, ESR

Specification: 100T/Box, 1200T/Carton

Classification

Others

We, Changsha Renji Medical Equipments Co., Ltd, under our sole

responsibility declare that the above-mentioned products meet

Assessment

Conformity

the provisions of the following EC Council Directives and

Route: Annex III

Standards. All supporting documentations are retained under the

premises of the manufacturer.

In vitro diagnostic medical devices directive:

General **Applicable Directives**

DIRECTIVE 98/79/EC OF THE EUROPEAN PARLLAMENT AND OF THE COUNCIL OF 27 October 1998 on in vitro diagnostic medical

devices.

EN 13612:2002/AC:2002

Standards Applied

EN ISO 14971: 2012

EN ISO 23640:201514 RENJIM

EN ISO 18113-1: 2011

EN ISO 18113-2:2011

EN ISO 15223-1: 2016

EN 13641: 2002

Address: Changsha City, Hunan Province, China.

Date: Nov 8, 2021

Name: Li Renjiang

Position: Managing Direct

Signature: