DECLARATION OF CONFORMITY TO COUNCIL DIRECTIVE 93/42/EEC CONCERNING MEDICAL DEVICES

MANUFACTURER: SHANDONG WEIGAO GROUP MEDICAL POLYMER CO., LTD No.18 XINGSHAN ROAD, TORCH HIGH-TECH SCIENCE PARK, 264210 WEIHAI, SHANDONG PROVINCE, PEOPLE' S REPUBLIC OF CHINA MEDICAL DEVICE: STERILE PLASTIC BLOOD BAG FOR SINGLE USE TYPE AND SIZE: S-250ML, S-350ML, S-450ML, D-250ML, D-350ML, D-450ML, T-250ML, T-350ML, T-450ML, Q-250ML, Q-350ML, Q-450ML GMDNS CODE: 10426 BLOOD DONOR SET CLASSIFICATION - ANNEX IX: CLASS IIB, RULE 18 CONFORMITY ASSESSMENT ROUTE: ANNEX II.3 WE, SHANDONG WEIGAO GROUP MEDICAL POLYMER CO., LTD., HEREWITH DECLARE THAT THE STATED MEDICAL DEVICES MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE 93/42/EEC OF 14 JUNE 1993 CONCERNING MEDICAL DEVICES; INCLUDING, AT 21 MARCH 2010, THE AMENDMENTS BY COUNCIL DIRECTIVE 2007/47/EEC. ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURER. THE MANUFACTURER IS EXCLUSIVELY RESPONSIBLE FOR THE DECLARATION OF CONFORMITY. TÜV SÜD PRODUCT SERVICE GMBH NOTIFIED BODY: RIDLERSTR 65, D-80339 MÜNCHEN, GERMANY **CE**₀₁₂₃ IDENTIFICATION NUMBER (EC) CERTIFICATE(S): G1 094273 0003 REV.03 ECREP MEDNET EC-REP GMBH EUROPEAN REPRESENTATIVE: BORKSTRASSE 10, 48163 MUENSTER, GERMANY VALID UNTIL : 2024-05-26 PLACE, DATE OF DECLARATION: WEIHAI, 2021-03-20

SIGNATURE:

NAME: WANG YI POSITION: (RESPONSIBLE SENIOR EXECUTIVE OF MANUFACTURER)