



September 15, 2023

Zhenjiang Assure Medical Equipment Co., Ltd.  
% Eva Li  
Consultant  
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Room 1401, Dongfang Building, 1500# Century Ave.,  
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Shanghai,  
China

Re: K232199

Trade/Device Name: Bariatric Heavy Duty Wheelchair (YJ-010B 20"DS; YJ-010B 20"DE; YJ-010B 20"DFS; YJ-010B 20"DFE; YJ-010B 20"ADS; YJ-010B 20"ADE; YJ-010B 20"ADFS; YJ-010B 20"ADFE; YJ-010B 22"DS; YJ-010B 22"DE; YJ-010B 22"DFS; YJ-010B 22"DFE; YJ-010B 22"ADS; YJ-010B 22"ADE; YJ-010B 22"ADFS; YJ-010B 22"ADFE; YJ-010B 24"DS; YJ-010B 24"DE; YJ-010B 24"DFS; YJ-010B 24"DFE; YJ-010B 24"ADS; YJ-010B 24"ADE; YJ-010B 22"ADFS; YJ-010B 24"ADFE)

Regulation Number: 21 CFR 890.3850

Regulation Name: Mechanical Wheelchair

Regulatory Class: Class I, reserved

Product Code: IOR

Dated: July 25, 2023

Received: July 25, 2023

Dear Eva Li:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at <https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm> identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of

Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see <https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products>); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to <https://www.fda.gov/medical-devices/medical-device-safety/medical-device-reporting-mdr-how-report-medical-device-problems>.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance>) and CDRH Learn (<https://www.fda.gov/training-and-continuing-education/cdrh-learn>). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (<https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice>) for more information or contact DICE by email ([DICE@fda.hhs.gov](mailto:DICE@fda.hhs.gov)) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

  
**Tushar Bansal -S**

for Heather Dean, PhD  
Assistant Director, Acute Injury Devices Team  
DHT5B: Division of Neuromodulation  
and Physical Medicine Devices  
OHT5: Office of Neurological  
and Physical Medicine Devices  
Office of Product Evaluation and Quality  
Center for Devices and Radiological Health

## Indications for Use

510(k) Number (if known)

K232199

Device Name

Bariatric Heavy Duty Wheelchair (YJ-010B 20''DS; YJ-010B 20''DE; YJ-010B 20''DFS; YJ-010B 20''DFE; YJ-010B 20''ADS; YJ-010B 20''ADE; YJ-010B 20''ADFS; YJ-010B 20''ADFE; YJ-010B 22''DS; YJ-010B 22''DE; YJ-010B 22''DFS; YJ-010B 22''DFE; YJ-010B 22''ADS; YJ-010B 22''ADE; YJ-010B 22''ADFS; YJ-010B 22''ADFE; YJ-010B 24''DS; YJ-010B 24''DE; YJ-0

Indications for Use (Describe)

The YJ-010B Manual Wheelchair is to provide mobility to persons limited to a sitting position.

Type of Use (Select one or both, as applicable)

Prescription Use (Part 21 CFR 801 Subpart D)

Over-The-Counter Use (21 CFR 801 Subpart C)

### CONTINUE ON A SEPARATE PAGE IF NEEDED.

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# 510(k) Summary

K232199

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## 1. Submitter

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Prepared Date: July 18<sup>th</sup>, 2023

## 2. Device

Name of Device: Bariatric Heavy Duty Wheelchair

Common Name: Manual Wheelchair

Model(s):

YJ-010B 20"DS	YJ-010B 20"DE	YJ-010B 20"DFS	YJ-010B 20"DFE	YJ-010B 20"ADS	YJ-010B 20"ADE	YJ-010B 20"ADFS	YJ-010B 20"ADFE
YJ-010B 22"DS	YJ-010B 22"DE	YJ-010B 22"DFS	YJ-010B 22"DFE	YJ-010B 22"ADS	YJ-010B 22"ADE	YJ-010B 22"ADFS	YJ-010B 22"ADFE
YJ-010B 24"DS	YJ-010B 24"DE	YJ-010B 24"DFS	YJ-010B 24"DFE	YJ-010B 24"ADS	YJ-010B 24"ADE	YJ-010B 24"ADFS	YJ-010B 24"ADFE

Regulatory Information

Classification Name: Mechanical Wheelchair

Regulatory Class: I

Product code: IOR

Regulation Number: 890.3850

Review Panel: Physical Medicine

## 3. Predicate device:

K201461

Ningbo Shenyu Medical Equipment Co.,Ltd.

Manual Wheelchair (A011)

## 4. Device description

The YJ-010B series is a mechanical wheelchair which is a manually operated device with wheels that is intended for medical purposes to provide mobility to persons restricted to a

sitting position. It can be folded for transport by bringing the two sides together. The manual wheelchair incorporates a main frame, a seat, two adjustable footplates and four wheels. The larger rear wheels have hand rims of slightly smaller diameter projecting just beyond the tire. These allows the user to maneuver the chair by pushing them on without requiring them to grasp the tires. The manual wheelchairs have brakes that bear on the tires of the rear wheels and two push handles at the upper rear of the frame to allow for manual propulsion by an assistant.

Main Components:

Main frame, back upholstery, seat upholstery, handgrip, armrest, front wheel, rear wheel, hand rim, crossbar, leg rest strap, footplate, brake, leg rest (only device model with "-E").

The device can be operated indoors, or outdoors on dry, level surfaces composed of concrete, blacktop, or asphalt under normal driving conditions.

The specification table is as below:

Model	YJ-010B series				
Overall dimension	-E: 1275mm(L)*1258mm(H)* 20"width 730mm/22"width 780mm/24"width 830mm				
	-S: 1175 mm(L)*1258mm(H) 20"width 730mm/22"width 780mm/ 24"width 830mm				
Folded dimension	Width 310mm				
Seat width	20"	509mm	Backrest width	20"	497mm
	22"	559mm		22"	547mm
	24"	610mm		24"	597mm
Seat depth	485mm		Seat height from floor	547mm	
Backrest angle	10°		Backrest height	420mm	
Footplate length	-E: 245 mm		Footplate-to-seat distance	-E: 448-555mm	
	-S:370 mm			-S:370-520mm	
Footplate clearance	50mm		Footplate-leg-angle	95°	
Front location of armrest structure	-D: 275mm		Front-armrest-to-backrest distance	-D: 275mm	
	-F: 432mm			-F: 432mm	
Front-armrest-to-backrest distance	-D: 275mm -F: 432mm		Armrest length	-D: 260mm -F: 352mm	
Leg-to-seat-surface angle	110°		Armrest angle	25°	
Material	Q235		Armrest-to-seat distance	245mm	
Distance between armrests	20"	510mm	Seat plane angle	3°	
	22"	560mm	Armrest width	55mm	
	24"	610mm			
Hand rim diameter	Φ 570mm		Maximum weight bearing	500LBS	

		capacity of the device	
Horizontal location of axle	65°	Weight of the device(net)	20" 27.3kg 22" 27.5kg 24" 27.7kg
diameter of front wheels	Φ190mm	diameter of rear wheels	Φ613mm
Backpack carrier or permission	N/A	Static stability sideways	11.75°
Static stability uphill	12.8°	Static stability downhill	13.2°
Minimum turning radius		865mm	
<b>Parking brake</b>			
Max slope uphill		7.3°	
Max slope downhill		7.1°	

### 5. Indication for use

The YJ-010B Manual Wheelchair is to provide mobility to persons limited to a sitting position.

### 6. Comparison of technological characteristics with the predicate device

Device	Predicate Device	Proposed Device	Results
510(k) Number	K201461	K232199	---
Manufacturer	Ningbo Shenyu Medical Equipment Co., Ltd.	Zhenjiang Assure Medical Equipment Co., Ltd.	---
Proprietary Name	Manual Wheelchair	Bariatric Heavy Duty Wheelchair	---
Classification	I	I	same
Indications for use	The A011 Manual wheelchair is to provide mobility to persons limited to a sitting position.	The YJ-010B Manual Wheelchair is to provide mobility to persons limited to a sitting position.	same
Design Characteristic	Manual Operation, Four-Wheels, Foldable, Cross-Brace, Pull-to-Lock, Armrest, Backrest, Foot rest	Manual Operation, Four-Wheels, Foldable, Cross-Brace, Pull-to-Lock, Armrest, Backrest, Foot rest, side panel	Similar*1
Brake control	occupant-operated brake only	occupant-operated brake only	same
Operation Environment	For indoor/outdoor use	For indoor/outdoor use	same
Control Mode	Mechanical	Mechanical	same
Size(unfold)	1100 (L) *660 (W) * 910mm (H)	-E: 1275mm(L)*1258mm(H)* 20"width 730mm/22"width 780mm/24"width 830mm -S: 1175 mm(L)*1258mm(H)	Different*2

		20"width 730mm/22"width 780mm/ 24"width 830mm	
Stowage length/width/height	810 (L) X 320 (W) X 930mm (H)	E: 1275mm(L)*1258mm(H)*310mm(W) S: 1175 mm(L)*1258mm(H)*310mm(W)	different*2
Weight (Total)	16kg(35.2lbs)	27.3-27.7Kg	different*2
Weight Capacity	136Kg(300lbs)	500LBS	different*2
Seat Width	480mm	509-610mm	different*2
Seat height	540mm	547mm	different*2
Seat depth	420mm	485mm	different*2
Back type	Fixed	Fixed	Same
Tires	Front: 200mm Rear:610mm	Front: 190mm Rear:613mm	different*2
Armrest	Flip back armrest	Armrest height adjustable (-A)/fixed	Similar*3
Foot rest	Optional/ swing away Optional/ swing away	with elevating leg rest(-E) with swing-away leg rest(-S)	Similar*3
Rear Axle Position	Single	Single	Same
Frame Construction	Foldable frame Push inward from left and right sides to fold	Foldable frame Push inward from left and right sides to fold	Same
Safety Feature	Manual Wheel Lock	Manual Wheel Lock	Same
Performance	Comply with: ISO7176-1 ISO7176-3 ISO7176-5 ISO7176-7 ISO7176-8 ISO7176-11 ISO7176-13 ISO7176-15 ISO7176-16	Comply with: ISO7176-1 ISO7176-3 ISO7176-5 ISO7176-7 ISO7176-8 ISO7176-11 ISO7176-13 ISO7176-15 ISO7176-16	Same
Biocompatibility	Comply with: ISO10993-1 ISO10993-5 ISO10993-10	Comply with: ISO10993-5 ISO10993-10 ISO10993-23	Similar*4

### Discussion:

Similar*1:	Compared to the predicate device, the subject device adds two skirt guards which are installed to the sides of the seat frame under the arm rests to provide a barrier between the occupant and the wheels. This feature can prevent occupant's clothes from getting caught in the wheels. This feature will not raise any new risk of safety or effectiveness.
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Different*2:	Compared to the predicate device, the subject device has different value on the unfold size, stowage size, device weight, capacity, seat width, seat height, seat depth, and tire size. However, the subject has passed the <ISO 7176-7-1998 Part7: Measurement of seating and wheel dimensions > and <ISO 7176-5-2008 Part 5: Determination of dimensions, mass and maneuvering space>, so the above different will not raise any new risk of safety or effectiveness.
Similar*3:	Compared to the predicate device, the subject device model with “-E” adds the elevating leg rest. This component provides convenience for orthopedic surgery patients and the armrest of the subject can’t be flipped. The above differences will not raise any new risk of safety or effectiveness.
Similar*4:	The stimulation test in ISO 10993-23 replaces the stimulation test in ISO 10993-10:2010. It will not raise any new risk of safety or effectiveness.

## 7. Summary of Non-Clinic Performance Testing

### Performance Testing

Non-clinical tests were conducted to verify that the subject device met all design specifications and was Substantially Equivalent (SE) to the predicate device. The test results demonstrated that the proposed device complies with the following standards:

ISO 7176-1:2014, Wheelchairs - Part 1: Determination of static stability

ISO 7176-3:2012 Wheelchairs - Part 3: Determination of effectiveness of brakes

ISO 7176-5: 2008 Wheelchairs - Part 5: Determination of overall dimensions, mass and maneuvering space

ISO 7176-7: 1998 Wheelchairs - Part 7: Measurement of seating and wheel dimensions

ISO 7176-8: 2014 Wheelchairs - Part 8: Requirements and test methods for static, impact and fatigue strengths

ISO 7176-11: 2012 Wheelchairs - Part 11: Test dummies

ISO 7176-13: 1989 Wheelchairs - Part 13: Determination of coefficient of friction of test surfaces

ISO 7176-15: 1996 Wheelchairs - Part 15: Requirements for information disclosure, documentation and labeling

ISO 7176-16:2012 Wheelchairs - Part 16: Resistance to ignition of postural support devices

ISO 7176-22:2014 Wheelchairs — Part 22: Set-up procedures

### Biocompatibility

Biocompatibility testing was conducted in accordance with the 2020 FDA guidance *Use of International Standard ISO 10993-1, “Biological Evaluation of Medical Devices – Part 1: Evaluation and testing within a risk management process.”* Testing included:

- Cytotoxicity (ISO 10993-5:2009)
- Sensitization (ISO 10993-10:2010)
- Irritation (ISO 10993-23:2021)

The testing supports the biocompatibility of the patient-contacting device materials that were shown to be non-cytotoxic, non-irritating, and non-sensitizing.

## 8. Clinical Test Conclusion

No clinical study is included in this submission.



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**9. Conclusion**

The conclusion drawn from the nonclinical tests demonstrates that the subject device in this 510(k) submission is as safe and effective as the legally marketed predicate device cleared under K201461.