Arthrex Angel® Concentrated Platelet Rich Plasma (cPRP) Processing Set

Instructions for Use

DFU-0260-3, rev. 0

SET CONTENTS

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2 Platelet Cuvette
3 Rotating Valve
4 Pump Loop Tubing
5 Threaded Luer Caps (quantity, 3 each)
6 Whole Blood Compartment
7 RBC Compartment
8 PPP Compartment
9 Three-Compartment Reservoir Bag
10 Variable Volume Separation Chamber
11 60 ml Specimen Cups (quantity, 2 each)
12 Syringe Activated PRP Valve Port
13 20 ml Luer Lock Syringe
14 Male-Female Luer Lock (quantity, 7 each)
15 Whole Blood Spike Adapter

Figure 1 - Arthrex Angel® Concentrated Platelet Rich Plasma (cPRP) Processing Set

Variable Volume Separation Chamber: The Arthrex Angel® Concentrated Platelet Rich Plasma (cPRP) Processing Set (Angel® cPRP Processing Set) utilizes a variable volume separation chamber that can process from 40 ml - 180 ml of anticoagulated autologous whole blood or a mixture of blood and bone marrow in a single cycle.

Platelet Cuvette/Valve Assembly: The platelet cuvette/valve assembly contains three major components: (1) the platelet cuvette, (2) the pump loop tubing and (3) the rotating valve.

The top section of the variable volume separation chamber (Figure 1) is the separation chamber plate. The separation chamber plate is used to seat the variable volume separation chamber in the centrifuge.

The platelet cuvette/valve assembly has been designed so that the operator can easily install the platelet cuvette/valve assembly while insuring that the platelet cuvette is properly seated in the platelet sensor and that the rotating valve aligns with the valve assembly driver.
PRP Valve Port: A luer lock syringe is attached to the PRP valve port to collect PRP. At the end of a processing cycle, the PRP valve port can also be used to collect PPP. A Syringe Activated Valve is included as an accessory and can be attached to maintain a closed port when the PRP syringe is removed.

Three-Compartment Reservoir Bag: The three-compartment reservoir bag is used to collect anticoagulated whole blood, the mixture of blood and bone marrow, and the separated blood components. The whole blood compartment is used as a reservoir for collected anticoagulated whole blood/bone marrow from a patient. The clinician may use syringes or whole blood bags to collect anticoagulated whole blood or a mixture of blood and bone marrow from a patient. The RBC Compartment is used to collect the concentrated red cells at the end of the processing cycle. The PPP Compartment is used to collect platelet poor plasma; the PPP is the first blood component collected after separation has been completed.

Syringe activated valves are used to access the PPP, and whole blood compartments of the three-compartment reservoir bag.

Other items in the Angel cPRP Processing Set include:

20 ml Luer Lock Syringe: The 20 ml luer lock syringe is used for the collection of platelet rich plasma. However, the syringe activated PRP valve will accommodate most luer fitting syringes.

60 ml Wrapped Specimen Cups (2 ea.): For use in collecting samples of the separated platelet poor plasma and platelet rich plasma, as needed.

Male/Female Luer Plug: The male/female luer plug can be used during and at the end of the procedure to seal open luer lock connections.

Whole Blood Bag Spike Adapter: The whole blood spike adapter is used to transfer blood from a whole blood bag to the whole blood compartment of the three-compartment reservoir bag.

Labels: Appropriate labels supplied with kit, to label collected whole blood and separated components.

DESCRIPTION
The Angel cPRP Processing Set consists of a pre-connected variable volume separation chamber, a tubing set with a platelet sensor/valve assembly, and a three-compartment reservoir bag for the collection of blood products (whole blood, red blood cells, and platelet poor plasma). The Angel cPRP Processing Set also contains a 20ml luer lock syringe for the collection of platelet rich plasma (PRP), two 60 ml specimen cups for collecting samples of the blood components, as needed, a whole blood bag spike adapter, a male-female luer plug, and labels for collected blood components.

Contents of this set have been sterilized by ethylene oxide gas and have non-pyrogenic fluid pathway.

INDICATIONS FOR USE
The Arthrex Angel® Concentrated Platelet Rich Plasma (cPRP) Set is intended for use with the Angel® Concentrated Platelet Rich Plasma (cPRP) System to separate and collect an autologous plasma fraction rich in platelets and red blood cells from the patient’s whole blood or a small mixture of blood and bone marrow perioperative to a surgical procedure.

CONTRAINDICATIONS
The Angel cPRP Processing Set is not intended to wash blood salvaged from the surgical site, nor is it intended for direct connection to the patient. The risk/benefit ratio of autologous sequestration must be determined on a case by case basis by the qualified medical personnel in charge of the patient’s care; the responsibility for the use of this device belongs solely to the physician in charge.

WARNINGS
1. Only Angel cPRP Processing Sets are approved for patient use with the Angel Concentrated Platelet Rich Plasma (cPRP) System.

2. Do not use the Angel cPRP Processing Set if the sterile packaging barrier has been broken.

3. Carefully examine the Angel cPRP Processing Set for damage, prior to use. Do not use if the Processing Set is damaged.

4. Carefully observe the Angel cPRP Processing Set for leaks during use. Leakage may result in loss of sterility of the device and/or loss of blood product.

5. When collecting and processing autologous blood products or a mixture of blood and bone marrow, it is recommended that the following precautions be followed to insure that the autologous product is not contaminated.
   • Use sterile technique when setting up the Angel cPRP Processing Set
   • Thoroughly clean and disinfect the donation site
   • Use sterile technique whenever handling autologous blood products

6. The whole blood or the mixture of blood and bone marrow must be anticoagulated before it can be processed for separation. Inadequate anticoagulation may result in clotting, interfering with the processing of the blood products.
Blood/ bone marrow containing clots will not pass through the syringe-activated valve located on the Whole Blood compartment of the three-compartment reservoir bag.

7. If centrifugation is discontinued before the completion of a processing cycle, the variable volume separation chamber is pressurized and presents the risk for exposure to blood and blood borne pathogens if the variable volume separation chamber is not properly removed. Please refer to the Angel Concentrated Platelet Rich Plasma (cPRP) System Operator’s Manual for unloading a variable volume separation chamber containing blood.

8. Failure to properly secure the luer lock syringe to the syringe-activated valve on the valve assembly may result in a leakage of fluids.

9. Do not directly connect the patient to the three-compartment reservoir bag. Direct connection to the patient could lead to vascular damage, shock or air embolism.

10. Disposal of Angel cPRP Processing Sets should be in accordance with federal, state, and local regulations. These materials should be considered biohazardous. Universal precautions for blood borne pathogens should be practiced when disposing of these items.

11. Place the Angel Concentrated Platelet Rich Plasma (cPRP) System on a flat, stable surface. Never try to move the Angel Concentrated Platelet Rich Plasma (cPRP) System while the device is in operation.

12. The platelet rich plasma is not intended for transfusion.

**PRECAUTIONS**

1. Carefully read these Instructions for Use before using this product. Refer to the Angel Concentrated Platelet Rich Plasma (cPRP) System Operator’s Manual for complete instructions.

2. Due to the possibility of operator exposure to blood borne pathogens (such as HIV, hepatitis viruses, bacteria, etc.), Universal Precautions for blood borne pathogens should be practiced.

3. When removing the Angel cPRP Processing Set from its packaging, check to ensure that the three (3) Threaded Luer Caps (see Figure 1, item 5) on each Compartment (Whole Blood, RBC and PPP) are securely tightened prior to installation into the Angel System.

4. The Angel cPRP Processing Set is intended for a single use. Do not resterilize any part of this Processing Set.

5. Failure to properly load the Angel cPRP Processing Set per the enclosed instructions may affect the performance of the system.

6. Luer lock syringes should be used with the Angel cPRP Processing Set.

7. This product is intended for use by trained personnel only.

8. Caution: Federal law restricts this device to sale by or on the order of a physician.

9. The safety and effectiveness of this device for in-vivo indications for use has not been established.

**INSTRUCTIONS FOR USE**

**Turning on the Angel cPRP System**

Turn on the Angel Concentrated Platelet Rich Plasma (cPRP) System by pressing the power switch on the back of the machine (see Figure 2). The message “Self test in progress. Please stand by.” will be displayed on the Angel cPRP System’s touch screen display. The machine will then orientate the valve assembly driver to the loading position.

**Initial Setup**

With the Angel cPRP System turned on, perform the following steps:

1. Open the centrifuge lid cover and lift the centrifuge stator arm to lock the centrifuge adapter within the centrifuge well.

2. Remove the Angel cPRP Processing Set from the tray.

3. Lay the Angel cPRP Processing Set on the top of the machine.

4. Check to ensure that the three (3) Threaded Luer Caps (Figure 1, item 5) on each Compartment (Whole Blood, RBC and PPP) are securely tightened prior to proceeding to the next step.
5. Insert the variable volume separation chamber into the centrifuge adapter by aligning the notches in the separation chamber plate down near the location of the position indicator and turn clockwise until the position indicator snaps into place (see Figure 3).

Rotate the centrifuge to a position so that the interlock mechanism shown in Figure 3 does not interfere with the stator arm. If the interlock mechanism interferes with the stator arm, the separation chamber plate will not load properly.

Note: Loading the variable volume separate chamber should always be the first step in the setup process. Loading the variable volume separation chamber and pressing down on the separation chamber plate will remove the excess air volume from the chamber. If the excess air is not removed, the separation chamber plate will not load properly.

6. Place the tube leading from the variable volume separation chamber through the slot on the rim of the centrifuge well.

7. Lower the centrifuge stator arm and align it with the raised tab on the top of the rotating seal of the variable volume separation chamber (see Figure 4).
8. Close the centrifuge lid. After closing the centrifuge lid, make sure that the tubing remains in the slot on the rim of the centrifuge and is not occluded / pinched by the centrifuge lid.

9. Place the pump loop tubing over the pump rotor. The pump loop will automatically load when the processing cycle is initiated. Seat the platelet cuvette/valve assembly by aligning the platelet cuvette and the valve assembly with the platelet sensor body and the valve assembly driver. Press down firmly on the back side of the platelet cuvette/valve assembly, at position A near the pump loop, until the assembly is snapped in place (see label A, Figure 5).

Note: It is essential that the platelet cuvette/valve assembly seats fully on the machine, to obtain proper sensing of the blood components.

10. Hang the three-compartment reservoir bag on the two support pins located on the side of the Angel Concentrated Platelet Rich Plasma (cPRP) System.
11. Remove the breather cap from the PRP valve port located on the valve assembly. If desired, attach the Syringe Activated Valve to the PRP valve port. Attach the 20 ml luer lock syringe (or alternate syringe, if desired), to the PRP valve port.

Note: The luer on the PRP valve port will accommodate most luer lock syringes.

12. After set-up, inspect the tube set to make sure there are no kinks or occlusions.

Blood Collection

The *Angel Concentrated Platelet Rich Plasma (cPRP)* System utilizes a variable volume separation chamber that is capable of processing between 40 ml and 180 ml of anticoagulated whole blood or a mixture of blood and bone marrow (bone marrow aspirate [BMA]) in a single cycle.

The *Angel Concentrated Platelet Rich Plasma (cPRP)* System can accommodate anticoagulated whole blood or anticoagulated bone marrow aspirate that has been collected in syringes or blood collection bags. In either situation, the whole blood or BMA should be collected with citrate anticoagulant (ACD-A) in a 7:1 ratio (seven parts whole blood to one part citrate anticoagulant). The following table defines the appropriate mixture of whole blood or BMA and citrate anticoagulant (ACD-A):

<table>
<thead>
<tr>
<th>Total Volume of Anticoagulated Whole Blood/ Bone Marrow Aspirate (mL)</th>
<th>Volume of ACD-A (mL)</th>
<th>Total Volume of Whole Blood/ Bone Marrow Aspirate Drawn (mL)</th>
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40 ml anticoagulated whole blood or bone marrow aspirate volumes require a patient hematocrit of 30% or greater. The recommended minimum patient hematocror for anticoagulated whole blood or bone marrow aspirate volumes of 50 ml or greater is 28%.

During and after collection, gently mix the whole blood or bone marrow aspirate with the citrate anticoagulant for a thorough distribution of the anticoagulant. Failure to properly mix the collected blood or bone marrow aspirate with anticoagulant may cause blood clot formation. Blood clot formation may interfere with the loading of blood/ bone marrow aspirate into the Whole Blood compartment of the three-compartment reservoir bag and/or may interfere with the processing of the blood or bone marrow aspirate.

If a syringe is used to collect blood or bone marrow aspirate, attach the syringe to the syringe-activated valve located on the Whole Blood compartment of the three-compartment reservoir bag and inject the blood or bone marrow aspirate.

If using a whole blood collection bag to collect blood or bone marrow aspirate, ensure that the blood-citrate/ BMA-citrate ratio is correct by weighing the bag as the blood/ bone marrow aspirate is collected according to AABB standard methods. Place the citrated (ACD-A) blood bank bag on a standard metric scale and zero it prior to beginning to withdraw the blood or bone marrow aspirate. Refer to the instructions for the specific bag that you are using and allow the blood/ bone marrow aspirate to gravity drain into the bag until its weight equals the volume of the bag (1 ml of blood weighs approximately 1.053 grams).

If bone marrow aspirate is to be processed using the *Angel Concentrated Platelet Rich Plasma (cPRP)* System, samples of a patient's bone marrow aspirate should be obtained using the techniques and procedures as practiced at each individual hospital or health-care setting. Other than the capacity of the Whole Blood Compartment, there are no volume limitations of anti-coagulated whole blood and anticoagulated bone marrow aspirate that can be transferred into the whole blood compartment of the three-compartment reservoir bag using the syringe activated valve or Whole Blood Bag Spike Adapter.

Use the Whole Blood Bag Spike Adapter to transfer the blood or the bone marrow aspirate to the whole blood compartment of the three-compartment reservoir bag. Connect the Whole Blood Bag Spike Adapter to the syringe activated valve located on the Whole Blood compartment of the three-compartment reservoir bag and spike the blood collection bag to drain the blood or the bone marrow aspirate into the whole blood compartment. After the blood or bone marrow aspirate has drained into the Whole Blood compartment of the three-compartment reservoir bag, remove the Whole Blood Bag Spike Adapter and recap the syringe-activated valve.
Refer to the Angel Concentrated Platelet Rich Plasma (cPRP) System Operator’s Manual for processing instructions.


RETURN OF USED PRODUCT

For Customers Within the United States

If for any reason this product must be returned to the manufacturer or to Arthrex, Inc., a returned goods authorization (RGA) number is required prior to shipping.

If the product has been in contact with blood or body fluids, it must be thoroughly cleaned and disinfected before packing. It should be shipped in either the original carton, or an equivalent carton, to prevent damage during shipment; and it should be properly labeled with an RGA number and an indication of the biohazardous nature of the contents of the shipment.

Instructions for cleaning and materials, including appropriate shipping containers, proper labeling, and an RGA number may be obtained from Arthrex Customer Service:

Phone: 800-934-4404
Email: cs@arthrex.com

⚠️ It is the responsibility of the health care institution to adequately prepare and identify the product for return shipment. Do not return products that have been exposed to blood borne infectious diseases.

For Customers Outside the United States

If for any reason this product must be returned, please contact your sales representative for specific instructions.

If the product has been in contact with blood or body fluids, it must be thoroughly cleaned and disinfected before packing. It should be shipped in either the original carton, or an equivalent carton, to prevent damage during shipment.

⚠️ It is the responsibility of the health care institution to adequately prepare and identify the product for return shipment. Do not return products that have been exposed to blood borne infectious diseases.

TECHNICAL SUPPORT

Call the Arthrex Technical Support Hotline at 1-(888) 420-9393, Monday through Friday from 9:00 AM to 5:00 PM EST; at +49 89 909005 8800 or techsupport@arthrex.de from 8:00 AM to 5:00 PM CET.

This is not a warranty document. For all warranty information, including disclaimers, exclusions, terms, conditions and related provisions, refer to the “Arthrex U.S. Product Warranty” section of the Arthrex, Inc. website, found at www.arthrex.com whose provisions are incorporated herein by reference.
DEFINITION OF SYMBOLS (as used in product labeling)

FOR SINGLE USE ONLY (DO NOT REUSE)

LOT
BATCH CODE (NUMBER)
(REFERENCE FOR PRODUCT TRACEABILITY)

USE BY - YEAR/MONTH/DAY

DATE OF MANUFACTURE

STERILE EO
STERILE - ETHYLENE OXIDE

NON PYROGENIC

CONTAINS PHTHALATE

DOES NOT CONTAIN NATURAL RUBBER LATEX

WARNING: DO NOT RESTERILIZE.

CONTENTS STERILE ONLY IF PACKAGE IS NOT OPENED, DAMAGED OR BROKEN

REF
CATALOG NUMBER

QTY
QUANTITY

CONSULT INSTRUCTIONS FOR USE

THIS WAY UP

CAUTION

FRAGILE; HANDLE WITH CARE

KEEP AWAY FROM HEAT

KEEP DRY

MANUFACTURER

R2 ONLY
FEDERAL LAW (USA) RESTRICTS THIS DEVICE TO SALE BY, OR ON THE ORDER OF A PHYSICIAN

THIS PRODUCT MEETS THE ESSENTIAL REQUIREMENTS OF MEDICAL DEVICE DIRECTIVE 93/42/EEC

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