

**HAEMONETICS ANTICOAGULANT CITRATE PHOSPHATE DOUBLE DEXTROSE SOLUTION (CP2D)- citric acid monohydrate, trisodium citrate dihydrate, sodium phosphate, monobasic, monohydrate, anhydrous dextrose solution**  
**Haemonetics Corporation**

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**Haemonetics Anticoagulant Citrate Phosphate Double Dextrose Solution (CP2D)**

**Instructions for Use**

For use with Haemonetics Apheresis Devices  
Rx Only

Products: Anticoagulant Citrate Phosphate Double Dextrose (CP2D)  
and Additive Solution Formula 3 (AS-3)

Company:  
Haemonetics Corporation  
400 Wood Road  
Braintree, MA 02184

**I. Indications for Use**

The Haemonetics 250mL Anticoagulant Citrate Phosphate Double Dextrose (CP2D) and 250mL Additive Solution Formula 3 (AS-3) nutrient solution are intended to be used only with automated apheresis devices for collecting human blood and blood components. The anticoagulant solution is metered by the apheresis machine into the collected whole blood. It is not to be infused directly into the donor. After the anticoagulant is used, the bag in which it was contained is discarded. When collecting plasma in the RBCP protocol, the plasma is collected into an empty plasma collection bag. One hundred milliliters (100ml) of AS-3 is transferred into one RBC collection bag when using the RBCP protocol or 2 separate bags when using the 2RBC protocol. AS-3 solution provides nutrients to keep the red blood cells viable for 42 days when refrigerated.

CP2D is also indicated for the collection of FFP and PF24 plasma, collected and stored plasma collected using the 822, 822-2P and 822F-2P disposable sets may be frozen within 8 hours (FFP) or within 24 hours which includes 8 hours room temperature storage and 16 hours refrigeration storage (PF24).

The 300mL AS-3 is used in conjunction with automated red blood cell washing devices. AS-3 serves as the nutrient solution for storage of the red blood cell product after deglycerolization. The red blood cells are washed using the Model 215 System. AS-3 is used for priming the disposable and for the last wash of the red blood cells. The washed cells are then re-suspended in 100mL of the AS-3 before transfer into a product collection bag or storage bag. Additive Solution Formula 3 (AS-3) provides nutrients to keep the washed red blood cells viable for up to 14 days after washing when refrigerated.

Neither the CP2D nor the AS-3 container is used for the storage of blood or blood components.

**Dosage Form**

The container for all CP2D and AS-3 solutions is a flexible polyvinyl chloride (PVC) bag sized for holding the appropriate amount of solution. The bag has a single port that is used for bag filling. The port is sealed after filling with a male luer assembly (CP2D) or female luer assembly (AS-3). Both the male and female luers are gamma irradiated prior to use. The plastic bag is contained in an overwrap which is added prior to sterilization.

The formulations for these products are as follows:

Citrate Phosphate Double Dextrose Solution (CP2D)

Each 100 mL contains:

Citric Acid (Monohydrate), USP 0.327 g  
Sodium Citrate (Dihydrate), USP 2.630 g  
Monobasic Sodium Phosphate (Monohydrate), USP 0.222 g  
Dextrose (Anhydrous), USP 4.640 g

Additive Solution 3 (AS-3)

Each 100mL contains:

Citric Acid (Monohydrate), USP 0.042 g  
Monobasic Sodium Phosphate (Monohydrate), USP 0.276 g  
Sodium Chloride, USP 0.410 g  
Adenine, USP 0.030 g  
Dextrose (Anhydrous), USP 1.000 g  
Sodium Citrate (Dihydrate), USP 0.588 g  
containing 15 mEq of Sodium.

**Dosage and Administration:**

CP2D Anticoagulant Solution and AS-3 Solution may be used with Haemonetics apheresis devices. See the Haemonetics Operation Manual for full operating instructions.

Prior to use of the solutions, check the solutions for leaks by squeezing each of the bags firmly. If leaks are found, discard the solution.

**III. Clinical Data**

Studies of the effectiveness of 250mL CP2D and 250mL AS-3 made by Haemonetics Corporation Union, South Carolina facility included *in vivo* survival data in healthy subjects at one test site and *in vitro* characterization of stored red blood cells at three test sites.

Haemonetics Clinical Evaluation #95012, involved autologous *in vivo* recovery and survival studies, and *in vitro* red blood cell characterization conducted at a single site. RBC results collected with Haemonetics CP2D/AS-3 were compared to crossover controls of manually collected RBCs from the same donors using CP2D/AS-3 made by Medsep Corporation (crossover manual controls). In addition, results were compared to data from other RBC apheresis donors using CP2D/AS-3 solutions manufactured by MedSep (unmatched apheresis controls). Test parameters included hematocrit, hemoglobin, RBC and WBC counts, ATP levels, pH, supernatant potassium, free hemoglobin, supernatant glucose levels, and product weight. The RBC quality of test units after 42-day storage was equivalent or slightly superior to those of the control units. RBCs collected by the two different collection protocols (Red Blood Cells with or without Plasma) showed no significant differences in terms of RBC quality and red blood cells from all groups met FDA and AABB guidelines for 24-hour recovering at 42 days storage.

Study #98001 was performed using 250mL CP2D and 250mL AS-3 to confirm, *in vitro*, the results of Clinical Evaluation #95012. Three test sites each performed three RBCP and three 2RBC apheresis procedures using the MCS+ LN8150 and associated collection sets. In addition, each site collected six manual whole blood units as concurrent (unmatched) controls using Medsep sets and prepared RBCs according to standard procedures.

*In vitro* tests of red blood cell functional and physical integrity were conducted at Days 0 and 42 of storage. All plasma units from the RBCP and manual procedures were evaluated on Day 0 for *in vitro* product characteristics. Group t-tests (two tailed) were performed to examine any statistically significant differences between the properties of apheresis and manual RBCs. Statistically significant differences ( $p < 0.05$ ) were evaluated in terms of clinical significance. Differences in glucose or lactate levels, hematocrit, pH, and supernatant potassium were not considered clinically relevant because these parameters have not been found to correlate significantly with the *in vivo* 24 hour percent recovery.

The results from this study confirmed the observations from Clinical Evaluation #95012. The quality of stored RBC products collected by apheresis is similar to RBC products collected manually. A difference in ATP and hemolysis levels, both after processing and on day 42 of storage, for the apheresis units were not statistically significant when compared to ATP and hemolysis levels from manually collected RBCs and were similar to historical reference values.

AS-3 300ml was used in conjunction with an automated closed system, the Haemonetics Model 215 (now called the ACP) in Clinical Evaluation #97002. Five test sites were used in this study. 100 samples were tested *in vitro* and 30 *in vivo*. Red blood cells derived from CPDA-1 whole blood units were used. AS-1 red blood cells were used as a control. A total of 140 red blood cell units were glycerolized and deglycerolized. The *in vitro* RBC quality and *in vivo* RBC viability data obtained on these units demonstrate that red cell units glycerolized and deglycerolized using the Model 215 System and resuspended in 300ml AS-3 solution manufactured by Haemonetics, Union, South Carolina are processed in a closed system and can be stored for 15 days at 4°C.

Clinical report TR-CLN-100229-A was conducted to support the use of CP2D for the collection of FFP and PF24 {plasma collected and stored Plasma collected using the 822, 822-2P and 822F-2P disposable sets may be frozen within 8 hours (FFP) or within 24 hours which includes 8 hours room temperature storage and 16 hours refrigeration storage (PF24). The following tables provide a summary of the results of this evaluation.

**Note:** Table 1, Table 2, and Table 3 below demonstrate the results of a clinical study conducted for various coagulation factors on plasma that was frozen within 8 hours (FFP) versus plasma frozen within 24 hours, which includes 8 hours room temperature storage and 16 hours refrigeration storage (PF24).

As noted in Table 1 below, Significant differences were observed between PF24 and FFP for assays Prothrombin Time, Partial Thromboplastin Time, FVlla, and FVIII.

Table 2 provides the results for Plasma Sample A, which was an outlier and subsequently excluded from the analysis.

Table 3 provides a summary of how many samples tested in each group differed by >20% when FFP and PF24 were compared.

**Table 1, Summary of FFP and PFZ4 plasma product assays (N=59)**

Assay	Mean (SD)		Median		(Min, Max)		Mean difference (PF24, FFP) (95% confidence interval)
	FFP	PF24	FFP	PF24	FFP	PF24	
Prothrombin Time (sec)	12.0 (0.50)	12.1 (0.59)	11.9	12.1	(11.2, 13.8)	(10.5, 14.0)	0.15 (0.09,0.20)
Activated Partial Thromboplastin Time (sec)	29.4 (3.32)	30.3 (3.34)	28.8	29.8	(23.1, 38.5)	(24.1, 39.5)	0.86 (0.64, 1.08)
Favtor V (%)	120.1 (23.93)	120.0 (23.88)	120.0	120.0	(61.0, 181.0)	(59.0, 182.0)	-0.14 (-2.03, 1.76)
Factor VIIa (mU/mL)*	60.5 (26.95)	45.0 (24.72)	54.0	41.5	(21.0, 131.0)	(12.0, 159.0)	-15 .50 (-19 .23, -11.77)
Factor VIII:C (%)*	94.3 (34.24)	81.7 (27.31)	88.5	82.5	(33.0, 194.0)	(34.0, 152.0)	-12.67 (-15.97, -9.38)
AT-III (%)	97.8 (12.24)	94.7 (15.97)	98.0	95.0	(71.0, 126.0)	(19.0, 134.0)	-3.05 (-6.30, 0.20)
Factor XI (%)	112.2 (17.42)	112.8 (18.46)	113.0	113.0	(81.0,151.0)	(82.0, 155.0)	0.63 (-0.70,1.95)
vWF (R:Co) (% function)	89.5 (35.65)	89.4 (35.58)	85.0	87.0	(35.0, 179.0)	(37.0, 185.0)	-0.15 (-2.21, 1.91)
Protein S (% activity)	74.9 (19.61)	73.4 (20.76)	72.0	71.0	(33.0, 125.0)	(39.0, 126.0)	-1.49 (-3.81, 0.83)
Protein C (% activity)	117.1 (17.59)	117.7 (17.49)	115.0	118.0	(65.0,157.0)	(68.0, 162.0)	0.53 (-0.54,1.59)
Fibrinopeptide F 1.2 (pmol/L)*	136.4 (63.34)	133.1 (55.15)	113.0	114.0	(80.0, 344.0)	(80.0,331.0)	-3.29 (-8.37,1.78)
Thrombln-Anti-thrombin Complex (ng/mL)	2.3 (1.38)	2.1 (0.40)	2.0	2.0	(2.0, 12.4)	(2.0,4.8)	-0.17 (-0 .53, 0.18)

\*Sample size = 58. The outlier pair was excluded from the analysis based on the statistical criteria or laboratory errors.

One sample (Plasma Sample A) showed significantly increased levels for Factor VIIa and F1.2. Another sample (Plasma Sample B) showed significantly decreased level of Factor VIII:C for the FFP sample (Table 2). These data were excluded from Table 1 since the values for VIIa and F1.2 were more than 4 SD from the mean. The factor VIII:C was not more than 4 SD from the mean but was significantly lower in FFP as compared to PF24. While other parameters in this sample did not show significant differences when compared to the control FFP, the FDA could not exclude the possibility that the activation found in the PF24 sample was related to the preparation procedure. The rate of such event occurrence was unknown and, based on review of published values for PF24 products, is most likely very rare. The clinical significance of the elevated markers of clotting activation for transfusion recipients is undetermined.

**Table 2, Three outlier pairs in plasma measurements**

Subject #	Measurement	FFP	PF24	Range	without	the	outliers
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				<b>FFP Min</b>	<b>FFP Max</b>	<b>PF24 Min</b>	<b>PF24 Max</b>
RBCP16	Factor FVIIa (mU/mL)	220	767	21	131	12	159
RBCP16	F1.2 (pmol/L)	702	1940	80	344	80	331
RBCP01	Factor VIII: C(%)	2	63	33	194	34	152

**Table 3, Proportion of Paired Sample:PF24 is greater than or less than FFP by more than 20 %**

<b>Assay</b>	<b>PFZ4&gt;FFP by more than 20 %</b>	<b>PF24&lt;FFP by more than 20 %</b>
Prothrombine Time (sec)	0% (0/59)	0% (0/59)
Partial Thromboplastin Time (sed	0% (0/59)	0% (0/59)
Factor V (%)	1.7% (1/59)	0% (0/59)
Factor Vila (mU/mL)	3.4% (2/58)	89.7% (52/58)
AT-III (%)	1.7% (1/59)	8.5% (5/59)
Factor VIII :C (%)	1.7% (1/58)	22.4% (13/58)
Factor XI (%)	0% (0/59)	0% (0/59)
Fibrinopeptide 1.2 (pmol/L)	0%(0/58)	1.7% (1/58)
Protein C (% activity)	0% (0/59)	0% (0/59)
Protein S (% activity)	5.1% (3/59)	1.7% (1/59)
Thrombin-Antithrombin Complex (ng/mL)	1.7% (1/59)	3.4% (2/59)
vWF (% activity)	3.4% (2/59)	1.7% (1/59)

**Note:** Plasma collected may be labeled as FFP or Plasma Frozen within 24 Hours (PF24).

**Note:** For FFP, the plasma should be placed in a freezer within 8 hours from the completion of collection. The plasma should be frozen at -18°C or colder.

For product intended to be labeled as PF24, the plasma should be refrigerated within 8 hours of collection at 1 – 6°C and frozen within 24 hours of collection.

FFP and PF24 should be frozen at -18°C or colder.

#### **IV. Warnings and Cautions**

##### **Contraindications:**

None known

##### **Precautions for use:**

- Solutions are NOT INTENDED FOR DIRECT INFUSION.
- Do not use if particulate matter is present or if the solution is cloudy  
*Note:* Some opacity of the plastic container due to moisture absorbance by the PVC during the sterilization process may be observed. This is normal and does not affect the quality or the safety of the product. This white, hazy appearance will diminish gradually over time. This advice only applies to the plastic container. Do not use if the solution itself is cloudy or contains particulate matter.
- Do not use if the container is damaged, leaking or if there is any visible sign of deterioration.

*Note:* The over wrap serves as a moisture barrier. Upon removing the solution bag, small droplets of condensation may be present in the

packaging. This is considered normal. After removing the over wrap gently squeeze the inner bag which protects the sterility of the solution. Discard if any leaks are observed or if an excessive amount of solution is noted within the over wrap.

- Do not reuse solution. Discard any unused or partially used solutions.
- Protect from sharp objects.
- Verify that solutions have been appropriately connected to avoid leaks.
- The set should be loaded and primed for use within 8 hours of the start of the collection.
- Carry out the apheresis procedure in accordance with the detailed instructions of the manufacturer of the apheresis device.

### **Adverse Reactions:**

Donors being reinfused with citrated blood or blood components may experience side effects due to the presence of citrate. Patients being transfused with the blood components could also experience a reaction to the citrated blood components. The major symptom experienced by donors is paraesthesia. Should this occur the reinfusion should be stopped or the reinfusion rate decreased. In the event of inadvertent bolus of the product administer calcium gluconate.

### **Drug Abuse / Dependence:**

Both solutions are intended to be used only with automated apheresis devices for collecting human blood and blood components. CP2D is used as an anticoagulant solution and AS-3 is intended for use a nutrient solution. Neither solution is intended for direct infusion; neither solution produces a pharmacological effect.

### **Overdosage:**

Both solutions are intended to be used only with automated apheresis devices for collecting human blood and blood components. CP2D is used as an anticoagulant solution and AS-3 is intended for use a nutrient solution. Neither solution is intended for direct infusion.

### **How Supplied:**

250mL CP2D Anticoagulant Solution is supplied in flexible PVC containers with a sterile, non-pyrogenic fluid path.

250mL and 300mL AS-3 Nutrient Solution is supplied in flexible PVC containers with a sterile, non-pyrogenic fluid path.

### **Storage and Handling:**

Room temperature (25°C/77°F). Avoid excessive heat. Protect from freezing.

113692-00(AB)

### **Package Label**

HAEMONETICS®  
ANTICOAGULANT CITRATE PHOSPHATE  
DOUBLE DEXTROSE SOLUTION

(CP2D)

Intended for use only with automated red cell apheresis devices to collect red blood cells and plasma (FFP and PF24). See the appropriate Operator's manual for additional information and complete usage instructions.

Each 100 mL contains:

Citric Acid (Monohydrate), USP	0.327 g
Sodium Citrate (Dihydrate), USP	2.630 g
Monobasic Sodium Phosphate (Monohydrate), USP	0.222 g
Dextrose (Anhydrous), USP	4.640 g

CAUTION: Not for direct intravenous infusion.

The pouch is a moisture barrier.

Do not use unless solution is clear and no leaks detected.

Single use container. Discard unused portion.

Sterile, nonpyrogenic fluid path.

Rx Only

RECOMENDED STORAGE: Room Temperature (25°C / 77°F)

Avoid excessive heat. Protect from freezing.

L065 DA

Haemonetics Corporation

Braintree, MA 02184 USA

Product code 455A

Lot No.

Exp Date

**Product Label - Package**

HAEMONETICS®

**ANTICOAGULANT CITRATE PHOSPHATE  
DOUBLE DEXTROSE SOLUTION  
(CP2D)**

250 mL

Intended for use only with automated red cell apheresis devices. See Operator's Manual for additional information and complete usage instructions.

Each 100 mL contains:

Citric Acid (Monohydrate), USP	0.327 g
Sodium Citrate (Dihydrate), USP	2.630 g
Monobasic Sodium Phosphate (Monohydrate), USP	0.222 g
Dextrose (Anhydrous), USP	4.640 g

**CAUTION:** Not for direct intravenous infusion.  
The pouch is a moisture barrier.  
Do not use unless solution is clear and no leaks detected.  
Single use container. Discard unused portion.  
Sterile, nonpyrogenic fluid path.  
Rx Only.

**RECOMMENDED STORAGE:** Room temperature (25 °C/77 °F).  
Avoid excessive heat. Protect from freezing.

L065, Rev. D-10	Product code 455A
Haemonetics Corporation	Lot No.
Braintree, MA 02184 USA	Exp. Date

**HAEMONETICS ANTICOAGULANT CITRATE PHOSPHATE DOUBLE  
DEXTROSE SOLUTION (CP2D)**

citric acid monohydrate, trisodium citrate dihydrate, sodium phosphate, monobasic, monohydrate, anhydrous dextrose solution

**Product Information**

<b>Product Type</b>	HUMAN PRESCRIPTION DRUG	<b>Item Code (Source)</b>	NDC:57826-455
<b>Route of Administration</b>	EXTRACORPOREAL		

**Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
<b>CITRIC ACID MONOHYDRATE</b> (UNII: 2968PHW8QP) (ANHYDROUS CITRIC ACID - UNII:XF417D3PSL)	ANHYDROUS CITRIC ACID	32.7 mg in 1 mL
<b>TRISODIUM CITRATE DIHYDRATE</b> (UNII: B22547B95K) (ANHYDROUS CITRIC ACID - UNII:XF417D3PSL)	ANHYDROUS CITRIC ACID	263 mg in 1 mL
<b>SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE</b> (UNII: 593YOG76RN) (PHOSPHATE ION - UNII:NK08V8K8HR)	SODIUM PHOSPHATE, MONOBASIC, MONOHYDRATE	22.2 mg in 1 mL
<b>ANHYDROUS DEXTROSE</b> (UNII: 5SL0G7R0OK) (ANHYDROUS DEXTROSE - UNII:5SL0G7R0OK)	ANHYDROUS DEXTROSE	464 mg in 1 mL

**Inactive Ingredients**

Ingredient Name	Strength
<b>WATER</b> (UNII: 059QF0K00R)	

**Packaging**

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:57826-455-02	250 mL in 1 BAG; Type 0: Not a Combination Product		

**Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	BN000127	01/10/2013	

**Labeler** - Haemonetics Corporation (057827420)**Establishment**

Name	Address	ID/FEI	Business Operations
Haemonetics Manufacturing Inc.		078598396	manufacture(57826-455)

**Establishment**

Name	Address	ID/FEI	Business Operations
Haemonetics Corporation		942344649	manufacture(57826-455)

Revised: 9/2020

Haemonetics Corporation