# WHO Prequalification of Diagnostics Programme PUBLIC REPORT

# Product: BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System PQ number: PQDx 0197-045-00

# Abstract

**BD** FACSPresto<sup>™</sup> Near-Patient CD4 Counter, BD FACSPresto<sup>™</sup> Cartridge and BD FACSPresto<sup>™</sup> cartridge kit with product codes 651000, 657681, 655495 and associated product codes, manufactured by Becton, Dickinson and Company, CE-marked regulatory version, was accepted for the WHO list of prequalified in vitro diagnostics and was listed 19 September 2014.

Subsequently, WHO received a change notification related to certain changes in manufacturing process. Documentation related to the changes was reviewed, accepted and this WHO prequalification public report amended on 24 March 2016.

## Intended use:

BD FACSPresto<sup>™</sup> Near-Patient CD4 System consists of BD FACSPresto<sup>™</sup> Counter and BD FACSPresto<sup>™</sup> cartridge which contains dried flourochrome-conjugated antibody reagents. This automated system is intended for in vitro enumeration of CD4 absolute count, CD4 percentage and hemoglobin concentration in human capillary and venous blood specimens. The number of CD4+ T cells in the peripheral blood is currently used to decide when to initiate treatment and monitor response to treatment in HIV infected individuals.

<b>Catalogue Number</b>	Product Description			
651000	BD FACSPresto™ instrument packaging includes:			
	Portable instrument			
	Power supply			
	Adapter Cords			
	Instrument Cover			
	Work station			
	Printer Paper			
	USB Flash Drive			
	<ul> <li>BD FACSPresto<sup>™</sup> Power Supply Adapter</li> </ul>			
	<ul> <li>BD FACSPresto<sup>™</sup> Near-Patient CD4 counter Instruction for</li> </ul>			
	use			
657681	BD FACSPresto™ Cartridge packaging includes			
	<ul> <li>BD FACSPresto<sup>™</sup> Cartridge (100 tests)</li> </ul>			
	<ul> <li>BD<sup>™</sup> Disposable Pipettes (100 pipettes)</li> </ul>			

	• BD FACSPresto <sup>™</sup> Cartridge Instruction for use			
655495	FACSPresto Cartridge Kit packaging includes			
	<ul> <li>BD FACSPresto<sup>™</sup> Cartridge</li> </ul>			
	• BD FACSPresto™ Finger Stick Sample Collection Kit			
	BD Microtainer Contact-Activated Lancet			
	Sterile Alcohol Prep Pads			
	Plastic Adhesive Bandage			
	A sterile Nonwoven Sponge			
658210	<ul> <li>BD FACSPresto<sup>™</sup> Instrument Carrying Case</li> </ul>			
658212	• BD FACSPresto <sup>™</sup> Solar Charge Kit (includes solar panel,			
	solar generator and power supply)			
658885	<ul> <li>BD FACSPresto<sup>™</sup> Solar Generator</li> </ul>			
658860	• BD FACSPresto™ Car Battery Charger Adapter (12V DC			
	power adaptor)			

# Assay principle:

When blood is introduced into the BD FACSPresto<sup>™</sup> Cartridge, the specific antibodies bind to the surface antigens on the T lymphocytes and monocytes during the incubation period. When the stained cartridge is inserted into the counter, the dedicated software identifies and counts the CD4+ T lymphocyte absolute and percentage cells, and calculates the hemoglobin concentration. The BD FACSPresto<sup>™</sup> Cartridge also contains immobilized antibodies as a quality control measure which the instrument uses to ensure that the reagents are present and sufficient blood specimen volume has been added.

The BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System includes:

# Storage:

The BD FACSPresto<sup>™</sup> Cartridge should be stored at 4 °C to 31 °C.

# Shelf-life:

12 months.

# Summary of Prequalification status for BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter with BD FACSPresto<sup>™</sup> Cartridge and BD FACSPresto<sup>™</sup> Cartridge kit

	Initial acceptance		
	Date	Outcome	
PQ amended	23March 2016		
Status on PQ list	18 September 2014	listed	
Dossier assessment	28 August 2014	MR	
Inspection status	13 August 2014	MR	
Laboratory evaluation	25 August 2014	MR	

# MR: Meets Requirements

BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System was accepted for the WHO list of in vitro prequalified diagnostics on the basis of data submitted and publicly available information.

# **Background information**

Becton, Dickinson and Company submitted an application for prequalification of BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System. Based on the established prioritization criteria, BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System was given priority for prequalification.

# Product dossier assessment

Becton, Dickinson and Company submitted a product dossier for BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System as per the Instructions for compilation of a product dossier (PQDx\_018 v1). The information submitted in the product dossier was reviewed by WHO staff and external experts (assessors) appointed by WHO in accordance with the internal report on the screening and assessment of a product dossier (PQDx\_009 v2). Based on the product dossier screening and assessment findings, a recommendation was made to accept the product dossier for BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System for prequalification.

Commitments for prequalification: The current shelf life for the FACSPresto Cartridge is 12 months at 31 °C and is supported by accelerated stability studies. Real-time stability studies are on-going and expected to support the shelf life of 12 months at 31 °C. This will be followed up at the next re-inspection.

## Manufacturing site inspection

A comprehensive inspection was performed at the site of manufacture (2350 Qume Drive, San Jose, 95131 CA, USA) of the BD FACSCount<sup>™</sup> System in March 2011 as per the Information for manufacturers on prequalification inspection procedures for the sites of manufacture of diagnostics (PQDx\_014 v1). The inspection found that the manufacturer had an acceptable quality management system and good manufacturing practices in place that ensured the consistent manufacture of a product of good quality.

A quality management documentation review was performed in August 2014, and it was established that the manufacturer continuously implemented a quality management system in compliance with ISO 13485:2003 and that no significant changes were implemented since the first inspection.

Quality documentation was submitted to support an additional site of manufacture (30 Tuas Ave 2, Singapore, 639461) on 14 September 2015, and accepted on 6 November 2015.

# Laboratory evaluation

The BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System was evaluated in two WHO collaborating laboratories in Antwerp, Belgium and Dar es Salaam, Tanzania between May and July 2014. The evaluation was conducted using the WHO evaluation protocol (PQDx 114) which was also approved by in-country ethical review boards in Belgium and Tanzania. A total of 1630 fresh capillary and venous blood specimens in Tanzania and Belgium were used to study failure rates, reproducibility (intra-assay variation, inter assay variation, inter-instrument variation, instrument precision) and agreement with the FACSCalibur<sup>™</sup> as the reference method. Lastly, ease to use was assessed.

The acceptance criteria are as follows: Specimen failure should be less than 10%. For reproducibility studies, a percentage coefficient of variation (%CV) should be less than 15% for CD4+ T counts of less than or equal to  $200/\mu$ L and %CV should be less than 10% for CD4 counts of more than 200 cells/ $\mu$ L. Compared to the reference method, the bias should be less than 10%.

Specimen failure which was defined as failure of the instrument to provide valid results was between 0-9 % for venous whole blood and between 2.5 and 5.5% for capillary whole blood.

Testing of fresh specimens was conducted to assess the ability of the BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System to provide reproducible results. The results indicated the following: the intra-assay variation on venous blood ranged from 4.7% to 7.0% for CD4 absolute counts and from 4.3% to 8% for CD4 percentages. The inter-instrument variability was below 4.8% for both CD4 absolute counts and CD4 percentages. The inter-assay variation for specimens kept up to 24 hours after collection ranged from 5.2% to 8% for CD4 absolute counts and CD4 percentages. Lastly, BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System had precision of less than 4% for CD4 absolute counts in both venous and capillary whole blood.

Regarding agreement with the reference method and agreement between capillary and venous whole blood specimens, the correlation coefficients were high with minimal bias in both laboratories. The performance of BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System to measure hemoglobin was not assessed in the current evaluation.

Operational characteristics of the BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System were assessed using a structured questionnaire by testing personnel. The BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System was found to be simple to use.

In conclusion, based on the results of evaluations conducted in two laboratories under the instruction of WHO, the performance of the BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter System fulfilled the WHO laboratory performance criteria using both venous and capillary whole blood specimens compared to BD FACSCalibur<sup>™</sup>.

# **Change notification**

In 2015, Becton, Dickinson and Company, submitted a change notification for addition of a new site of assembly for certain components. The change notification was assessed and product was found to meet WHO prequalification requirements.

# Labelling

1. Labels

2. Instructions for use

# 1. Labels

## A. Cartridge Pouch Label



# B. Cartridge Box Label



C. Cartridge Kit Box Label



D. Instrument Model Plate Label

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Model: 🚭	Model: ⊖ BD FACSPresto™ REF Near-Patient CD4 Counter						
This device complies with Part 15 of the FCC Rules. Operation is subject to the following two conditions: (1) this device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation. This Class A digital apparatus meets all requirements of the Canadian Interference Causing Equipment Regulations. Cet apparell numérique de la classe A respecte toutes les exigences de Réglement sur le matér- ial brouilleur du Canada.							
Mfd:							
SN:							
VDC:	18 V 🚃	3.3 A	Power:	60 W			
MD ( E FC 🗵							
Becton	Becton, Dickinson and Company, BD Biosciences, 2350 Qume Drive, San Jose, CA 95131 USA						
BD, BD Dickins	Logo and all other tr on and Company. © 2	, uun Laoghaire, Co. D ademarks are property 2014 BD	of Becton,	Made in USA 657250 Rev. 02			

	Near-Patie	ent CD4 Counte	r	
This device of conditions: (' any interfere Class A digiti Equipment R	omplies with Part 15 c ) this device may not nce received, includin al apparatus meets all egulations.	of the FCC Rules. Op cause harmful interf g interference that n requirements of the	eration is subject to the erence, and (2) this dev nay cause undesired op Canadian Interference	e following two vice must accept eration. This Causing
Cet appareil ial brouilleur	numérique de la class du Canada.	e A respecte toutes le	es exigences de Réglem	ent sur le matér
Mfd:				
SN:				
VDC:	18 V 🚃	3.3 A	Power:	60 W
		IV		
			EQ Queen Dalue Can lare	CA 95131 USA
Becto	on, Dickinson and Comp	any, BD Biosciences, 23	150 Qume Drive, San Jose	
EC REP Bene	n, Dickinson and Comp x Limited, Pottery Road,	any, BD Biosciences, 23 Dun Laoghaire, Co. Du	ublin, Ireland	
Becto EC REP Bene Asser	n, Dickinson and Comp x Limited, Pottery Road, nbled in Singapore	any, BD Biosciences, 23 Dun Laoghaire, Co. Du	ublin, Ireland	

E. Instrument Shipping box Label

⊖ BD FACSPresto™ Near-Patient CD4 Counter		⊕ BD FACSPresto™ Near-Patient CD4 Counter	technic, Sickhows and Carpany     Bisterioran     239 Quere Onio     Son Area CA 391     Son Area CA 391     Son Area CA 391     Tot 455,554,367     Chold Sept Carbon Wirk-rom
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	Control Open: Proceedings		Examplesis Continues Support Tel 332,4003,9035 Fax > 322,4012,9264 heigh also consideration graduity com because Dicklosses Pay Lid, a Strates and which have
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			kelikissienseurom 80, 80 Loga and 80 FACSPrento are trademarks of Rector, Diskinson
	₿BD		and Companys 0 2013 80 Made in USA 23-1554)-00

⊜BD FACSPresto™ Near-Patient CD4 Counter	11 🗭 🗹 11 🕶 (f 11	⊖ BD FACSPresto™ Near-Patient CD4 Counter	Locine         Discine         of Electronic           10 Electronic         Electronic         Electronic           10 Electronic         Electronic         Electronic           11 Electronic         Electronic         Electronic           11 Electronic         Electronic         Electronic           11 Electronic         Electronic         Electronic           12 Electronic         Electronic         Electronic           12 Electronic         Electronic         Electronic           12 Electronic         Electronic         Electronic           12 Electronic         Electronic         Electronic           13 Electronic         Electronic         Electronic           14 Electronic         Electronic         Electronic
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# **Attachment 5.4**

BD FACSPresto Cartridge

IFU

Appendices are numbered independently.



# BD FACSPresto™ Cartridge

100 Tests-Catalog No. 657681

2/2016

23-12814-01



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EC REP

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#### 1. INTENDED USE

For use only with the BD FACSPresto<sup>™</sup> Near-Patient CD4 Counter.

The BD FACSPresto Near-Patient CD4 Counter is an automated system for in vitro diagnostic use in performing the direct enumeration of CD4 absolute count, CD4 percentage of lymphocytes, and hemoglobin concentration in human whole blood.

#### **Clinical Applications**

CD4 counts and CD4 percentages (%CD4) have been used to evaluate the immune status of patients diagnosed with, or suspected of developing, immune deficiencies such as acquired immune deficiency syndrome (AIDS).<sup>1,2</sup>

The CD4 antigen is the receptor for the human immunodeficiency virus (HIV).<sup>3</sup> The absolute number and percentage of CD4 T lymphocytes are the cellular parameters most closely associated with HIV disease progression and patient prognosis.<sup>4</sup> The number of CD4 T lymphocytes declines in HIV infection.<sup>5-7</sup>

Hemoglobin is a protein in red blood cells that carries oxygen from the lungs to the body. Low or declining hemoglobin concentration is an indicator of anemia, a hematological abnormality frequently associated with HIV.<sup>8-10</sup>

#### 2. PRINCIPLES OF THE PROCEDURE

The BD FACSPresto<sup>™</sup> cartridge<sup>\*</sup>, the CD4/%CD4/Hb cartridge, contains dried fluorochrome-conjugated antibody reagents. When blood reacts with the reagents, the antibodies in the reagents bind to the surface antigens on the

<sup>\*</sup> BD FACSPresto Cartridge: US Patent 8,248,597

lymphocytes and monocytes. After the incubation period, the cells are analyzed on the BD FACSPresto Near-Patient CD4 Counter (the instrument). The software identifies the cell populations of interest and calculates CD4 absolute counts, CD4 percentages of lymphocytes, and hemoglobin concentration. The system measures total hemoglobin by a spectrophotometric method, using absorbance at an isobestic point for oxyhemoglobin and deoxy-hemoglobin, with correction for scatter.

#### 3. REAGENT COMPOSITION

The cartridge contains CD4 PE -Cy<sup>™</sup>5<sup>†</sup>/ CD3 APC/CD45RA APC/CD14 PE dried antibody reagents. The dried antibody reagents include inert ingredients such as buffer, bovine serum albumin (BSA), and ProClin<sup>®‡</sup> as a preservative. CD3 APC and CD45RA APC enumerate total lymphocytes. Lymphocytes labeled with CD4 PE-Cy5 are designated CD4+ lymphocytes. CD14 PE identifies monocytes, which are excluded from the analysis.

The CD4 antigen<sup>11,12</sup> (55 kDa)<sup>13</sup> is present on the helper/inducer Tlymphocyte subset<sup>14,15</sup> CD3+CD4+, which consists of 28% to 58%<sup>16</sup> of lymphocytes in normal peripheral

† Cy™ is a trademark of GE Healthcare. This product is subject to proprietary rights of GE Healthcare and Carnegie Mellon University, and is made and sold under license from GE Healthcare. This product is licensed for sale only for in vitro diagnostics. It is not licensed for any other use. If you require any additional license to use this product and do not have one, return this material, unopened, to BD Biosciences, 2350 Qume Drive, San Jose, CA 95131, and any money paid for the material will be refunded.

 ProClin is a registered trademark of Rohm and Haas Company. blood,<sup>12</sup> and is present in low density on the surface and in the cytoplasm of monocytes. The CD4 antibody, clone SK3,<sup>11</sup> is derived from the hybridization of mouse myeloma cells with spleen cells from BALB/c mice immunized with human peripheral blood T lymphocytes. The CD4 antibody is composed of mouse IgG<sub>1</sub> heavy chains and kappa light chains.

The CD3 antibody, clone SK7, is derived from the hybridization of NS-1 mouse myeloma cells with spleen cells from BALB/c mice immunized with human thymocytes. CD3 is composed of mouse  $IgG_1$  heavy chains and kappa light chains.

The CD45RA antigen is present on approximately 50% of CD4+ T lymphocytes, approximately 75% of CD8+ T lymphocytes, and on essentially all B lymphocytes and natural killer (NK) lymphocytes.<sup>17</sup> The helper/inducer Tlymphocyte subset expresses the phenotype CD4+CD45RA+.<sup>17</sup> The CD45RA antigen is expressed on naive T lymphocytes. Antigen density decreases upon in vitro activation.<sup>18</sup> A selective loss of the CD4+CD45RA+ subset during active multiple sclerosis has been demonstrated.<sup>17,19</sup>

The CD45RA antibody, clone HI100,<sup>20</sup> is derived from the hybridization of mouse myeloma cells with spleen cells isolated from mice immunized with human whole blood cells (WBCs).

The CD14 antigen is present on the majority of normal peripheral blood monocytes.<sup>21</sup> The CD14 antibody recognizes a human monocyte/ macrophage antigen of 55 kDa.<sup>22</sup> The CD14 antibody, clone M\perpensition for the hybridization of Sp2/0 mouse myeloma cells with spleen cells from BALB/c mice immunized with peripheral

blood monocytes from a patient with rheumatoid arthritis. The CD14 antibody is composed of mouse  $IgG_{2b}$  heavy chains and kappa light chains.

#### 4. PACKAGE CONTENTS

Each box contains 100 cartridges and 100 pipets.

## 5. STORAGE AND HANDLING

Store the cartridge:

- In its original foil pouch. Do not use the cartridge if the pouch has been opened for more than 30 minutes.
- At 4°C–31°C (39°F–88°F).
- In 10%–95% non-condensing humidity.
- Until the expiration date. Do not use the cartridge after the expiration date on the package.

Incubate the cartridge at 10°C–40°C (50°F–104°F).

#### 6. MATERIALS REQUIRED BUT NOT PROVIDED

- BD FACSPresto<sup>TM</sup> instrument
- BD FACSPresto<sup>TM</sup> work station
- BD FACSPresto<sup>™</sup> finger stick sample collection kit
- BD Vacutainer<sup>®</sup> EDTA blood collection tubes

# 7. CARTRIDGE QC

Cartridge Quality Control (QC) uses immobilized antibodies. The instrument verifies that the reagent is present and that there is sufficient sample in the cartridge. Cartridge QC runs automatically at every cartridge run.

## 8. PATIENT SPECIMENS

The assay is designed to be used only with peripheral whole blood collected by venipuncture into EDTA tubes or by finger stick.

Capillary<sup>23</sup> or venous blood samples are transferred directly into the cartridge and incubated. Samples are run on the instrument after incubation.

Follow these guidelines for handling your samples:

- Do not dilute whole blood before adding it to the cartridge.
- Do not refrigerate the whole blood specimen before sample preparation.
- Store whole blood collected in EDTA tubes at 20°C–25°C (68°F–77°F) up to 24 hours before applying the blood to the cartridge.
- Minimize exposure of the cartridges to light.
- Do not remove the channel protector on the cartridge until just before you insert the cartridge into the instrument.

**WARNING** Do not use previously fixed and stored samples. Whole blood specimens refrigerated before staining can give incorrect results. Specimens from patients taking immunosuppressive drugs can yield poor resolution.<sup>24</sup> Do not test hemolyzed specimens.

**WARNING** The reagents contain antibodies of mouse and rat origin.

**WARNING** All biological specimens and materials coming in contact with them are considered biohazards. Handle as if capable of transmitting infection<sup>25,26</sup> and dispose of with proper precautions in accordance with federal, state, and local regulations. Never pipette by mouth.

Wear suitable protective clothing, eyewear, and gloves.

**WARNING** Each cartridge is for single use only. Use one cartridge per specimen.

#### 9. PROCESS CONTROLS

For information about process controls or external quality assessment products, contact your local BD Biosciences office or representative.

#### **10. PROCEDURE**

See the following for instructions on preparing and running samples.

- BD FACSPresto Near-Patient CD4 Counter Instructions for Use (IFU)
- BD FACSPresto Near-Patient CD4 Counter Quick Reference Guide

#### **11. REFERENCE INTERVALS**

The reference intervals for the BD FACSPresto cartridge shown in Table 1 are representative for hematologically normal adults.

Subset	Gender	Ν	Mean	Reference interval
CD427	Male	77	811	462–1,306 cells/µL
	Female	83	866	440–1,602 cells/µL
%CD4 <sup>27</sup>	Male	77	41	29%-54%
	Female	83	44	32%-55%
Hb <sup>28</sup>	Male	NA	NA	13.5–18.0 g/dL
	Female	NA	NA	12.0–16.0 g/dL

Table 1 Representative reference intervals

We recommend that laboratories and other users establish their own reference intervals for their patient populations using the BD FACSPresto system to reflect potential sources of variability, such as patient gender, race, age, and preparation techniques.

#### **12. EXPECTED RESULTS**

Performance of the BD FACSPresto cartridge (the cartridge) was established by the testing at the BD Biosciences laboratories in San Jose, CA, USA and at one clinical laboratory in Kisumu, Kenya, Africa.

#### **Method Comparison**

Absolute counts of CD4-positive cells, the percentage of CD4 positive cells in the lymphocyte population, and total hemoglobin concentration in whole blood from HIV-infected patients were determined using the BD FACSPresto system. Results were compared with results from the BD Tritest<sup>™</sup> CD3 FITC/ CD4 PE/CD45 PerCP reagent, in BD Trucount<sup>™</sup> tubes, on the BD FACSCalibur<sup>™</sup> flow cytometer, with the BD FACS™ Loader, using BD Multiset<sup>TM</sup> software and the Sysmex<sup>®§</sup> KX-21 hematology analyzer. Whole blood samples were collected at the clinical laboratories. Regression statistics reported in Table 2 and Table 3 indicate the results are substantially equivalent. Details are shown in Figure 1 through Figure 5.

Table 2 Method comparison for venous blood

Parameter	Ν	R <sup>2</sup>	Slope	Intercept	Range
CD4	189	0.98	0.97	7.37	55– 2,478 cells/μL
%CD4	188	0.96	1.03	0.13	5.06%- 53.77%
Hb	190	0.96	0.94	0.18	3–18.9 g/dL

§ Sysmex is a registered trademark of Sysmex America Inc.

R<sup>2</sup> Parameter Ν Slope Intercept Range 69-CD4 162 0.97 1.03 13.47 2,474 cells/µL 5.9%-56.2% %CD4 161 0.96 1.02 -0.26

Table 3 Method comparison for capillary blood

Regression analysis is not applicable for hemoglobin capillary samples. A total of 163 samples were collected with a range of 4.7–17.1 g/dL. The average bias around the medical decision level (9.5 g/ dL–11.5 g/dL) for capillary hemoglobin specimens was calculated against Sysmex as 2.02%.

**Figure 1** Scatter plot with weighted Deming fit (y = 0.97x + 7.37) for venous blood (CD4)



**Figure 2** Scatter plot with Deming fit (y = 1.03x + 0.13) for venous blood (%CD4)



**Figure 3** Scatter plot with weighted Deming fit (y = 1.03x + 13.47) for capillary blood (CD4)



**Figure 4** Scatter plot with Deming fit (y = 1.02x - 0.26) for capillary blood (%CD4)



**Figure 5** Scatter plot with Deming fit (y = 0.94x + 0.18) for venous blood (Hb)



#### Within-Specimen Reproducibility and Precision

Twenty replicates were created by one operator in one day, using two concentration levels of process controls, one lot of cartridges, and one instrument. Results are shown in Table 4. Estimates of precision were determined at one site, BD Biosciences, using CD4 cellular and hemoglobin process controls. Two replicates of each CD4 control (normal and low) and two replicates of each of the 3 levels of hemoglobin controls were analyzed in each run, and two runs were performed per day for a total of 21 days. Three different instruments and three cartridge lots with three different operators were used, each for seven of the 21 days. Coefficients of variation (CVs) and standard deviations (SDs) are provided for CD4 absolute counts and CD4 percentages for withinrun precision and total precision in Table 5 through Table 7, respectively.

Table 4 Repeatability study

	Level	Mean	с۷	SDa	N
CD4	Low	155.10	5.78		20
	Normal	926.60	2.59		20
%CD4	Low	12.77		0.73	20
	Normal	43.99	1.53		20
Hb	Low	6.95	2.26		20
	Normal	12.99	1.09		20

 Standard deviations (SDs) are reported instead of coefficients of variation (CVs) for %CD4 low level.

 Table 5 Within-run and within-device precision

 CD4 absolute counts

	Low control (CV)	Normal control (CV)
Within-run	6.79	2.18
Within-device	6.79	3.30

# Table 6 Within-run and within-device precision CD4 percentages

	Low control (SD <sup>a</sup> )	Normal control (CV)
Within-run	0.75	1.58
Within-device	0.75	1.74

 Standard deviations (SDs) are reported instead of coefficients of variation (CVs) for %CD4 low level.

# Table 7 Within-run and within-device precision hemoglobin

	Low control (CV)	Medium control (CV)	High control (CV)
Within-run	2.42	1.46	1.07
Within-device	2.42	1.52	1.14

#### Stability

A stability study was conducted at the clinical laboratory in Kisumu to assess the following:

- Changes associated with the storage of whole blood before addition into the cartridge
- Changes as a result of time between addition of blood into the cartridge and data acquisition
- The combined effect of both

Whole blood samples were tested up to 24 hours post draw, and samples were tested up to 2 hours post addition of blood into the cartridge. All samples were maintained at room temperature (20°C–25°C, 68°F–77°F) before addition of blood into the cartridge or acquisition. Based on the results of this study, cartridges should be prepared with whole blood samples within 24 hours of draw, and then run within 2 hours of adding blood into the cartridge.

Linearity was assessed in triplicate measurements of multiple concentrations of CD4<sup>+</sup> cells, lymphocyte cells, and hemoglobin across the reportable range of the assay for CD4 absolute counts, total lymphocytes, and hemoglobin on the instrument. Results are linear in the CD4 range (50–4,000 cells/µL), absolute lymphocyte range (200–10,000 cells/µL), and hemoglobin range (2–20 g/dL).

## **13. CARTRIDGE SPECIFICATIONS**

Item	Description
Blood stability	Up to 24 hours after draw if stored in an EDTA tube at 20°C–25°C (68°F– 77°F)
Sample stability	Up to 2 hours after addition of sample to cartridge
Sample throughput	More than 10 patient results per hour when run in batch mode
Validated range	CD4 count: 50–4,000 cells/µL %CD4: 5%–60% Hb concentration: 2.0–20 g/dL

## **14. LIMITATIONS**

- Use the cartridge only with the BD FACSPresto instrument.
- The cartridge has no user-serviceable parts.
- Performance characteristics outside the validated range have not been established.
- Interfering substances in the sample may result in an inaccurate result.
- Follow the instructions in the IFU on preparing capillary and venous blood samples to ensure accurate results.

## Linearity

#### **15. INTERFERING CONDITIONS**

Table 8 lists the substances that were tested for interference with the reagents in the cartridge. Testing for interference was performed in accordance with EP7<sup>29</sup>. There was no detectable interference at the following concentrations.

Table 8	Interfering	substances
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Analyte	Max concentration
Acetaminophen	20 mg/dL
Albumin	6 g/dL
Amodiaquine	60 ng/mL
Artesunate	600 ng/mL
Ascorbic acid	6 mg/dL
Conjugated bilirubin	5 mg/dL
Creatinine	5 mg/dL
Efavirenz	16 μg/mL
Ethambutol	12 μg/mL
Gamma Globulin	40 mg/mL
Glucose	120 mg/dL
Hemolysis	20%
Ibuprofen	500 μg/mL
Iron	150 μg/dL
Isoniazid	40 µg/mL
Lipemia (intralipid)	2,400 mg/dL
Magnesium	6.3 µg/dL
Methemoglobin	14%
Nevirapine	7 μg/mL
Quinine	48 μg/mL
Rifampicin	64 μg/mL
Salicylic Acid	200 µg/mL
Tenofovir	1,000 ng/mL
Tetracycline	150 μg/mL
Thrombocytes (Platelets)	1.541 x 106 cells/µL

#### Table 8 Interfering substances

Analyte	Max concentration	
Urea	40 mg/dL	
Uric acid	9 mg/dL	
White Blood Cells	25 x 10 <sup>3</sup> cells/µL	
Zidovudine	1,000 ng/mL	
Disease Condition		
Rouleaux Formation		
Cold Agglutinin		

#### WARRANTY

Unless otherwise indicated in any applicable BD general conditions of sale for non-US customers, the following warranty applies to the purchase of these products.

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#### TROUBLESHOOTING

See the troubleshooting section in the *BD FACSPresto Near-Patient CD4 Counter Instructions For Use.* 

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