

Rx Only

INTERCEPT® Blood System for Platelets - Package Insert - Large Volume (LV) Processing Set

DESCRIPTION

The INTERCEPT Blood System for Platelets is intended for the ex vivo treatment and storage of platelet components. The system is used to inactivate a broad range of pathogens, including viruses ¹⁻¹⁰, bacteria, and protozoan parasites ¹¹⁻¹⁸, thus reducing the risk of transfusion-transmitted infections. Contaminating donor leukocytes are also inactivated by the INTERCEPT treatment, reducing the risk of transfusion-associated graft versus host disease (TA-GVHD). ¹⁹⁻²³

The INTERCEPT Blood System for platelets is a Class IV medical device consisting of single-use processing sets containing amotosalen solution (INTERCEPT Processing Sets) and an illumination device (INTERCEPT Illuminator INT100, software version 6.1), which provides a controlled dose of ultraviolet A (UVA) light for each treatment. Platelet components treated with the INTERCEPT Blood System are referred to as INTERCEPT Platelets.

The INTERCEPT Blood System for platelets is available in several processing set configurations. This package insert describes information specific for the Large Volume (LV) processing set.

The LV set includes one final platelet storage container and is used for the treatment and storage of single dose platelet concentrates. The Instructions for Use section describes the platelet unit specifications required for treatment using this processing set. The key components of the LV set are as follows:

- Amotosalen container with 17.5 mL of 3 mM amotosalen solution in 0.924% saline
- Illumination container for treatment in INTERCEPT Illuminator
- Compound Adsorption Device (CAD) for removal of residual amotosalen and free photoproducts
- Platelet storage container

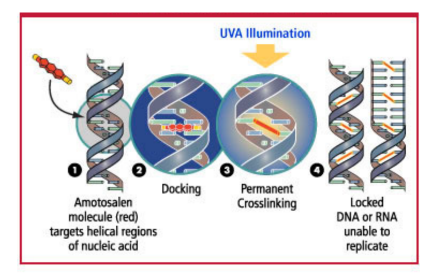
The INTERCEPT Illuminator delivers a controlled amount of UVA light (wavelength 320 to 400 nm) to the illumination container during each treatment cycle. The machine has the capability of illuminating 1 or 2 INTERCEPT processing sets per treatment cycle. Each illumination container rests on a UVA transparent tray that undergoes horizontal agitation during the illumination process. UVA illumination in each chamber is provided by 2 opposing banks of fluorescent lamps mounted above and below the illumination tray, and is monitored by independent photodiode systems which integrate the UVA dose. The INTERCEPT Illuminator delivers a 3 Joules/cm² (J/cm²) UVA treatment within approximately 3 to 4 minutes. Upon completion of the illumination step, the platelets are transferred to the bag containing the CAD and agitated for 6-24 hours, depending on the platelet suspension medium, at $22^{\circ}\text{C} \pm 2^{\circ}\text{C}$. At the completion of the CAD incubation, the platelets are transferred by gravity flow to the storage container and are ready for transfusion.

PRINCIPLE OF OPERATION

The INTERCEPT treatment process utilizes the synthetic psoralen known as amotosalen (also referred to as S-59; structure shown in **Figure 1** below). Psoralens are a class of planar, aromatic molecules that can bind reversibly to nucleic acids by intercalation. Upon illumination with long wavelength UVA light, intercalated psoralens form monoadducts and interstrand crosslinks with RNA and DNA.²⁴ The result is permanent, irreparable adducts in DNA and RNA, which prevent replication of pathogens and leukocytes.²⁴ The intercalation, illumination and crosslinking mechanism of action is shown in **Figure 2**.

Figure 1 Amotosalen: C₁₆H₁₉NO₄

Figure 2 Amotosalen Mechanism of Action



INDICATIONS AND USAGE

The INTERCEPT Blood System for platelets is intended to be used for the ex vivo preparation of pathogen-reduced, whole blood derived buffy coat platelet or apheresis platelet components in order to reduce the risk of transfusion-transmitted infection (TTI).

CONTRAINDICATIONS

- Contraindicated for preparation of platelet components intended for patients with a history of hypersensitivity reaction to amotosalen or other psoralens.
- Contraindicated for preparation of platelet components intended for neonatal patients treated with phototherapy devices that emit a peak energy wavelength less than 425 nm, or have a lower bound of the emission bandwidth <375 nm, due to the potential for erythema resulting from interaction between ultraviolet light and amotosalen.

WARNINGS

Treatment with INTERCEPT Blood System does not replace applicable standards for processing in open and closed systems. If there is a leak in the set during processing, platelet product must be discarded.

Amotosalen in contact with skin may result in photosensitization in the presence of ultraviolet light. If skin exposure occurs, flush exposed skin with water.

The sterile connecting device (SCD) and tube sealer should be used according to manufacturer's instructions for use.

While laboratory studies of amotosalen processing with UVA light have shown a reduction in levels of certain viruses, bacteria, and parasites, there is no pathogen inactivation process that has been shown to eliminate all pathogens.

Neonatal patients who require platelet transfusion during phototherapy for treatment of hyperbilirubinemia should not be treated with phototherapy devices that emit light less than 425 nm or have a lower bound of the emission bandwidth <375 nm, due to the potential for erythema resulting from interaction between ultraviolet light and amotosalen.

Di(2-ethylhexyl) phthalate (DEHP) is known to be released from polyvinyl chloride (PVC) medical devices; increased leaching can occur with extended storage or increased surface area contact. The INTERCEPT processing sets only have tubing components, container ports and an in-line filter that contain PVC; all containers and other parts are PVC-free. During use of this processing set, blood components are in contact with PVC for a brief period of time (approx. <15 minutes). Based on limited surface area contact and minimal contact time, DEHP levels in blood components after use of the processing set are estimated to be well below those resulting from other medical applications containing PVC tubing (e.g., hemodialysis, intravenous fluid administration, extracorporeal membrane oxygenation and cardiopulmonary bypass procedures). The risks associated with DEHP released to the blood components must be weighed against the benefits of therapeutic transfusion and inactivation of harmful viruses, bacteria and other pathogens.

Pulmonary events: Acute Respiratory Distress Syndrome (ARDS)

INTERCEPT processed platelets may cause the following adverse reaction: *Acute Respiratory Distress Syndrome (ARDS)*

An increased incidence of ARDS was reported in a randomized trial for recipients of INTERCEPT processed platelets, 5/318 (1.6%), compared to recipients of conventional platelet components (0/327). Monitor patients for signs and symptoms of ARDS. ^{25, 26}

PRECAUTIONS

The INTERCEPT processing sets are intended for single-use. Do not reuse sets or components of sets.

Do not use if: tamper-evident package has been opened; signs of deterioration are visible; fluid path closures are loose or not intact; cannulae are broken; or there is no fluid in the amotosalen solution container.

All the following conditions must be met for pathogen inactivation:

- Platelet collection should be in accordance with blood bank practice.
- Platelets mixed with amotosalen must be exposed to UVA light using the INTERCEPT Illuminator. No
 other source of UVA light may be used. Please refer to the Operator's Manual for the INTERCEPT
 Illuminator.
- Platelets must be incubated with the CAD after illumination and prior to use. Refer to the Instructions for Use section for the appropriate minimum and maximum CAD incubation times.

The INTERCEPT Blood System can accommodate a range of platelet collection and processing technologies. However, regardless of the method of collection or processing, blood establishments need to adjust their practices to conform to defined product guard bands for input volume, treatment window times, and other key processing parameters.

Treat all blood products as though they contain an infectious agent. Follow institutional guidelines regarding the handling of infectious agents. Dispose of all materials used in the procedure as biohazardous waste.

ADVERSE EVENTS

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a product cannot be directly compared to rates in the clinical trials of another product and may not reflect the rates observed in clinical practice.

As anticipated, a large number of adverse events (AEs) were reported in the clinical studies in patients with severe platelet-dependent thrombocytopenia, including patients treated for malignancy and patients undergoing hematopoietic stem cell transplantation (HSCT). Most events were considered to be related to the primary disease treatment and/or the underlying disease and were not reported as related to the use of study platelets.

In the Phase 3/3B randomized controlled blinded trials enrolling patients repeatedly transfused with INTERCEPT Platelets prepared from single donor apheresis or pooled donor buffy coat products and stored for up to 5 days, adverse events were classified as possibly, probably, or related to the exposure to INTERCEPT Platelets. Those related adverse events observed in \geq 5% of patients included transfusion reactions (13%), blood alkaline phosphatase not otherwise specified (NOS) increased (5%), blood urea increased (5%) and petechiae (7%). Adverse reactions (defined as possibly or probably related to study transfusion) that occurred in \geq 5 (1.25%) patients treated with INTERCEPT or conventional (control) platelets in these randomized clinical trials are summarized in **Table 1**.

Table 1 Related Adverse Events Occurring in ≥5 (1.25%) Patients Treated with INTERCEPT or Conventional (Control) Platelets in Randomized Phase 3/3B Clinical Trials

Event	INTERCEPT (n=392)	Control (n=399)	
Any AE	103 (26%)	112 (28%)	
Febrile Neutropenia	5 (1%)	1 (<1%)	
Thrombocytopenia	4 (1%)	7 (2%)	
Transfusion Reaction	51 (13%)	65 (16%)	
Haematemesis	7 (2%)	8 (2%)	
Mouth Haemorrhage	8 (2%)	9 (2%)	
Injection Site Erythema	7 (2%)	8 (2%)	
Oedema Lower Limb	9 (2%)	6 (2%)	
Pyrexia	11 (3%)	8 (2%)	
Rigors	15 (4%)	12 (3%)	
Hyperbilirubinaemia	6 (2%)	8 (2%)	
Hypoproteinaemia	12 (3%)	13 (3%)	
Jaundice NOS	7 (2%)	7 (2%)	
Activated Partial Thromboplastin Time Prolonged	6 (2%)	3 (1%)	
Alanine Aminotransferase Increased	11 (3%)	9 (2%)	
Aspartate Aminotransferase Increased	15 (4%)	12 (3%)	
Bilirubin Conjugated Increased	6 (2%)	5 (1%)	
Blood Alkaline Phosphatase NOS Increased	18 (5%)	13 (3%)	
Blood Bilirubin Increased	5 (1%)	5 (1%)	
Blood Carbon Dioxide Decreased	6 (2%)	5 (1%)	
Blood Chloride Increased	4 (1%)	6 (2%)	
Blood Creatinine Increased	8 (2%)	6 (2%)	

Event	INTERCEPT (n=392)	Control (n=399)	
Blood Lactate Dehydrogenase Increased	6 (2%)	5 (1%)	
Blood Magnesium Decreased	15 (4%)	5 (2%)	
Blood Urea Increased	18 (5%)	11 (3%)	
Faecal Occult Blood Positive	15 (4%)	13 (3%)	
Weight Increased	7 (2%)	7 (2%)	
Hypermagnesaemia	6 (2%)	4 (1%)	
Hyperphosphataemia	12 (3%)	9 (2%)	
Hypocalcaemia	13 (3%)	7 (2%)	
Hypokalaemia	16 (4%)	11 (3%)	
Hyponatraemia	12 (3%)	6 (2%)	
Hypophosphataemia	10 (3%)	9 (2%)	
Headache NOS	1 (<1%)	6 (2%)	
Haematuria	17 (4%)	13 (3%)	
Dyspnoea NOS	6 (2%)	3 (1%)	
Epistaxis	14 (4%)	18 (5%)	
Haemoptysis	7 (2%)	6 (2%)	
Нурохіа	6 (2%)	3 (1%)	
Pleural Effusion	5 (1%)	3 (1%)	
Contusion	16 (4%)	20 (5%)	
Dermatitis NOS	14 (4%)	9 (2%)	
Erythema Not Elsewhere Classified (NEC)	5 (1%)	8 (2%)	
Petechiae	29 (7%)	17 (4%)	
Pruritus NOS	8 (2%)	9 (2%)	
Urticaria NOS	6 (2%)	4 (1%)	
Catheter Site Haemorrhage	7 (2%)	6 (2%)	
Refractoriness To Platelet Transfusion	15 (4%)	6 (2%)	

Hemorrhage: Hemorrhagic AEs occurred in all studies, and generally consisted of mild mucosal-cutaneous bleeding. Clinically severe hemorrhagic AEs were observed at equivalent frequencies following administration of INTERCEPT or conventional platelets across the controlled clinical studies.

Clinical refractoriness to platelet transfusion: Clinical refractoriness to study platelet transfusions, defined as 2 consecutive study platelet transfusions with the 1-hour CCI < 5x10³, was monitored in the prospective phase 3/3B studies. In one study, there was a significantly higher rate of clinical refractoriness to platelet transfusions based on this definition [21.4% (68/318) with INTERCEPT Platelets compared to 7.0% (23/327) with conventional platelets]. Clinical refractoriness was transient and the frequency of persistent refractoriness was not increased with exposure to INTERCEPT Platelets.²⁸ There was no evidence of higher rates of immunologic refractoriness associated with INTERCEPT Platelets, and the higher rates seen in some instances were attributed to lower average platelet doses in INTERCEPT platelet components compared to reference platelet components, with resultant lower post-transfusion count increments. Persistent clinical refractoriness has not been reported in post-marketing studies.³⁰⁻³²

Immunogenicity

2340 post-baseline plasma samples from a total of 412 subjects who received INTERCEPT Platelet transfusions in the prospective clinical studies were tested for the presence of antibodies to amotosalen or amotosalen-associated neoantigens. Four samples from two patients who received INTERCEPT Platelets prepared from buffy coat platelets were initially reactive, but other samples from these subjects were inconsistently reactive above the threshold, thus the reactivity was not considered to represent an immune response to amotosalen or amotosalen-associated neoantigens.³³

NONCLINICAL TOXICOLOGY

Nonclinical studies have been conducted to evaluate the potential toxicity of exposure to amotosalen. There was no indication of a toxicological risk in animal experiments using platelet components treated with the INTERCEPT Blood System. In intravenous 28-day toxicity studies in rats and dogs exposed to amotosalen, the no observed adverse effect levels were at least 75,000- and 25,000-fold, respectively, the clinical exposure from transfusion of a platelet component.³⁴

Amotosalen was rapidly eliminated in mice and rats with an initial plasma $t_{1/2}$ of less than 1 hour. The primary route of excretion of amotosalen and its photoproducts was fecal. There was no evidence of amotosalen accumulation after repeated exposures over periods as long as 13 weeks.

In preliminary single dose studies, transient clinical effects were seen at very high doses of amotosalen (more than 150,000 times in rat and 30,000 times in dog the daily human exposure from a single platelet component) but no target organ toxicity was observed. In 7 day studies (rat and dog) and 14 day studies (rats), there was no evidence of any systemic toxicity following exposure to high doses of amotosalen (more than 75,000 times in rat and 30,000 times in dog the daily human exposure form a single platelet component).³⁴

In fertility studies of embryo-fetal or peri-postnatal development in rats and rabbits as well as in one study with neonatal rats, there was no evidence of reproductive or developmental toxicity of amotosalen.³⁵

There was no *in vivo* evidence of genotoxicity or carcinogenicity of amotosalen, including a study using the validated p53 knock-out mouse model. In the presence of mammalian hepatic metabolic activation, no evidence of *in vitro* mutagenicity of amotosalen was observed.³⁶

CLINICAL STUDIES

The safety and effectiveness of INTERCEPT Platelets were investigated in 11 controlled clinical studies and 2 single arm observational studies ^{27-29, 37-49}. A total of 1267 study subjects received INTERCEPT Platelets in the clinical studies; only the pivotal studies are presented.

INTERCEPT Platelets prepared from apheresis collections stored up to 5 days

The SPRINT Trial was a randomized, controlled, double blind, parallel-group study to compare the therapeutic efficacy and safety of INTERCEPT apheresis platelet components to conventional apheresis platelet components using a non-inferiority design.²⁸ The population selected for the study included patients with severe thrombocytopenia secondary to a spectrum of hematologic and non-hematologic malignancies, as well as other hematologic conditions associated with severe thrombocytopenia, who required platelet transfusion for prevention and/or treatment of active bleeding. The primary endpoint was the proportion of patients with Grade 2 bleeding during a period of platelet transfusion support up to 28 days. A total of 645 patients (318 Test, 327 Reference) were transfused and received a total of 4719 platelet transfusions (2678 Test, 2041 Reference). There were 186/318 (58.5%) patients in the Test group and 188/327 (57.5%) patients in the Reference group with Grade 2 bleeding. The primary efficacy endpoint of this study was met and non-inferiority of INTERCEPT Platelets confirmed using a noninferiority margin of 12.5%. However, differences in the mean number of transfusions, average interval between transfusions, rates of refractoriness, mean days with Grade 2 bleeding and mean count increment and corrected count increment at 1 and 24 hours were all statistically significantly different favoring the control group. The observations of reduced count increments, more PC transfusions, and shorter transfusion intervals in recipients of INTERCEPT PCs may have been partly attributable to an increased frequency of lower platelet doses in INTERCEPT PCs. The targeted platelet transfusion dose was 3.7x10¹¹. Sixty percent (60%) of the subjects in the INTERCEPT processed platelet group received at least one platelet dose of less than 3x10¹¹ compared to 36% in the control group. Additionally, subjects in the INTERCEPT processed platelet group received off-protocol transfusions (32% vs. 15%) more often than the control group because of protocol violations.

There were no differences between treatment groups in the frequencies of Grade 3 and Grade 4 adverse events, serious adverse events, or mortality. Statistically significant differences for increased incidence of four adverse events of Grade 3 and Grade 4 severity were observed among recipients of INTERCEPT processed platelets. These included: pneumonitis not otherwise specified (5 versus 0); ARDS (5 versus 0); hypocalcemia (21 versus 8); and syncope (6 versus 0). A subsequent blinded, retrospective analysis of patients with clinically significant pulmonary adverse events (n = 148) demonstrated no significant increase in incidence of acute lung injury (ALI), including ARDS. However a trend of increased ARDS in the patients receiving INTERCEPT processed platelets remained. The causality of these events remains uncertain and an increased risk of ARDS with the INTERCEPT processed platelets cannot be ruled out.

• A randomized, double-blind, controlled, parallel-group trial was conducted to evaluate the therapeutic efficacy and safety of apheresis INTERCEPT Platelets prepared with the commercial INTERCEPT system compared to conventional apheresis PCs.²⁹ The study was designed as a difference study to characterize any potential differences between INTERCEPT Platelets and conventional apheresis platelet concentrates. The population selected for the study were patients receiving platelet transfusions for thrombocytopenia due to malignancy or myelosuppressive therapy for malignancy. The primary endpoint was the 1-hour platelet count increment. A total of 43 patients (22 Test, 21 Reference) were enrolled at

3 study centers, and a total of 218 platelet components were transfused (103 Test and 115 Reference). The 1-hour post transfusion platelet count increment was lower, on average by 7.2 x 10¹¹ platelets for INTERCEPT Platelets compared to conventional apheresis platelet concentrates based on a longitudinal regression analysis model.

INTERCEPT Platelets from pooled whole blood derived buffy coat concentrates stored up to 5 days

• The euroSPRITE trial was a randomized, controlled, double blind, parallel-group trial designed to evaluate the therapeutic efficacy and safety of INTERCEPT Platelets in comparison to conventional platelets prepared from whole blood derived buffy coat platelet concentrates.²⁷ The study was designed as a difference study to characterize any potential differences between PCT (Test) and untreated (Reference) buffy coat platelet concentrates. Hematology-oncology patients were enrolled to receive platelet transfusion support up to 56 days. The primary endpoints were the 1-hour platelet count increment. A total of 103 patients were included (52 Test, 51 Reference) and received a total of 676 platelet transfusions (390 for Test group versus 286 for Reference group). The mean per patient 1-hour CI for the Test group (27x10⁹/L) was statistically significantly lower than the mean 1-hour CI for the Reference group (36x10⁹/L; mean difference (95% CI) = 8.34 (1.23 to 15.45); p-value=0.02.

Statistically significant differences in the incidence of the following adverse events were observed: sepsis NOS (Not Otherwise Specified) (6 for Test group versus 0 for Reference group), blood creatinine increase (13 for the Test group versus 5 for Reference group), and phlebitis NOS (0 for Test group versus 5 for Reference group).

Single transfusions of INTERCEPT Platelets stored up to 7-days

This group includes data from two studies using random donor buffy coat or apheresis platelets.

- A randomized, double-blind, single-center, two-treatment, two-period cross-over, non-inferiority pilot study was conducted to compare INTERCEPT Platelets with conventional platelets prepared from pooled random-donor buffy coat platelet concentrates with 7 day storage prior to transfusion. The study was performed in a population of thrombocytopenic patients from a hematology-oncology clinic, and each subject received two transfusions: a single transfusion of INTERCEPT Platelets and a single transfusion of conventional platelets stored for 7 days. The primary endpoint was the 1-hour (10 to 90 min) CCI after 7-day-old platelet transfusion, and the primary efficacy analysis was performed in 20 patients: 9 patients randomized to the Sequence I-R (INTERCEPT platelets followed by Reference platelets) and 11 patients randomized to Sequence R-I. The non-inferiority of INTERCEPT platelets relative to conventional platelets could not be demonstrated for the 1-hour post-transfusion CCI.
- The TESSI study was a randomized, controlled, double-blinded, non-inferiority study designed to compare the safety and efficacy of INTERCEPT Platelets stored for 6-7 days with conventional platelets of a similar age.44 Patients were recruited from the hematology-oncology departments of the participating clinical study centers, and were thrombocytopenic or expected to develop thrombocytopenia requiring platelet transfusion within 30 days of randomization. The primary endpoint was the 1-hour CCI. A total of 211 patients were randomized and received one study platelet transfusion (105 Test, 106 Reference) of platelets stored >5 days (80% of PCs were stored for 7 days). Non-inferiority of 1- hour CCI for INTERCEPT Platelets relative to conventional platelets was not demonstrated based on the full analysis set (FAS) that included all randomized patients who received the study treatment (INTERCEPT or conventional platelets).

Observational Studies with INTERCEPT Platelets

This group includes data from two studies at two sites, one of which used PCs from "double-dose" apheresis collections and one of which prepared PCs from either apheresis or whole blood derived buffy coat platelet concentrates.

INTERCEPT Platelets prepared from apheresis and buffy coat platelets

• This was a single-arm, open label study of 560 INTERCEPT Platelet transfusions in 51 patients with malignant hematology disorders, including HSCT. 45 The study was designed to monitor the safety and efficacy of INTERCEPT Platelets transfused on a routine basis in adult patients. The primary endpoint was the frequency of acute transfusion reactions. Platelet concentrates were prepared from either apheresis collections or whole blood derived buffy coat platelets, treated with INTERCEPT in place of gamma irradiation, and stored up to 5 days prior to transfusion. The frequency of acute transfusion reactions was 1.6% (9/553). This study was small in size and uncontrolled therefore the rate of acute transfusion reactions cannot be compared with that for conventional platelets.

INTERCEPT Platelets prepared from double-dose apheresis platelet collections

• This study was designed as a single-arm, open observational label study of 551 INTERCEPT Platelet transfusions in 46 hematology patients with malignant disorders, including HSCT, to monitor the safety and efficacy of INTERCEPT Platelets transfused on a routine basis in a Swiss Red Cross blood center. The primary endpoint was the frequency of acute transfusion reactions. INTERCEPT Platelets were prepared from apheresis "double unit" collections (5-6 x 10¹¹) without gamma irradiation and stored up to 5 days. The frequency of acute transfusion reactions was 2% (12/551). This study was small in size and uncontrolled therefore the rate of acute transfusion reactions cannot be compared with that for conventional platelets.

Post-Marketing Studies

Safety data were obtained from three hemovigilance (HV) programs in routine use without patient selection: the Cerus hemovigilance program and the regulatory surveillance programs in France and in Switzerland. 30-32, 50-60

The populations monitored in the Cerus hemovigilance studies (CLI-HV 00045, HV1-075-RF-HIV CTR and HV2-489-P-NIV-CTR) included 4,067 patients, where 62 patients were under the age of 1 year and 185 patients were 1-18 years of age. 50% of the patients enrolled in these studies were hematology-oncology patients, of which 12% were HSCT patients. Adverse events within 24 hours and serious adverse events within 7 days of platelet transfusion were reported. The frequencies of adverse events attributed to INTERCEPT processed platelet transfusions were not increased compared to conventional platelet transfusions reported in European regulatory hemovigilance programs.

DEVICE PERFORMANCE

Pathogen Inactivation

The INTERCEPT Blood System has been validated to effectively inactivate a broad spectrum of enveloped and non-enveloped viruses, Gram-positive and Gram-negative bacteria, spirochetes, parasites and leukocytes. Selection of pathogens was intended to provide evidence of the broad capabilities of the inactivation process. Pathogens selected represent those associated with transfusion transmission as well as model viruses of more general applicability. **Table 2** through **Table 5** summarize the pathogen inactivation data.

Table 2 Viral Inactivation (Enveloped) 61,62

	Extent of Inactivation (Log ₁₀ Reduction)				
Pathogen	Platelets in Plasma/Additive Solution	Platelets in 100% Plasma			
HIV-1 IIIB, cell-associated	≥5.4	-			
HIV-1 IIIB, cell-free	≥5.6	≥4.7			
HIV-1 Z84 (clinical isolate)	>3.3	=			
HIV-2 CLB-20 (clinical isolate)	>2.4	-			
DHBV	≥4.8	-			
BVDV	>4.4	>3.3			
HTLV-I	4.7	-			
HTLV-II	5.1	-			
Cytomegalovirus (CMV)	>4.9	-			
PRV (model for CMV)	-	>4.2			
Influenza A virus	≥5.9	- -			

[&]quot;-" indicates not tested

 Table 3
 Viral Inactivation (Non-Enveloped) 61, 62

	Extent of Inactivation (Log ₁₀ Reduction)			
Pathogen	Platelets in Plasma/Additive Solution	Platelets in 100% Plasma		
Calicivirus (model non-enveloped virus)	2.1	-		
Blue Tongue virus (model non-enveloped virus)	4.4	-		

[&]quot;-" indicates not tested

Table 4 Bacterial Inactivation* 61,62

	Extent of Inactivation (Log ₁₀ Reduction)						
Pathogen	Platelets in Plasma/Additive Solution	Platelets in 100% Plasma					
Gı	Gram-Negative Bacteria						
Klebsiella pneumonia	5.8	-					
Escherichia coli	≥6.3	-					
Serratia marcescens	≥6.7	-					
Pseudomonas aeruginosa	≥3.9						
Salmonella choleraesuis	6.2	-					
Enterobacter cloacae	5.5	-					
Yersinia enterocolitica	≥5.9	-					
G	Gram-Positive Bacteria						
Staphylococcus epidermidis	≥6.1	-					
Staphylococcus aureus	≥5.4	-					
Listeria monocytogenes	≥6.3	-					
Corynebacterium minutissimum	≥5.3	-					
Streptococcus pyogenes	≥6.8	-					
Bacillus cereus (vegetative)	≥5.5	-					
Bacillus cereus (spore forming)	≥1.16	-					
Anaerobic Gram-Positive Bacteria and Spirochete Bacteria							
Borrelia burgdorferi	>6.8	-					

[&]quot;-" indicates not tested

Table 5 Protozoa and Leukocyte Inactivation 61,62

	Extent of Inactivation (Log ₁₀ Reduction)				
Pathogen	Platelets in Plasma/Additive Solution	Platelets in 100% Plasma			
Plasmodium falciparum	≥5.6	-			
Human T-cells	>4	-			

[&]quot;-" indicates not tested

^{*}Some of the studies above do not include storage, but refer to immediate inactivation post illumination.

Leukocyte Inactivation

Irradiation of blood components to inactivate leukocytes (T-cells) is frequently performed to prevent TA-GVHD in blood product recipients. The INTERCEPT Blood System has been shown to be effective in inactivating leukocytes, eliminating leukocyte replication and also inhibiting cytokine production. TA-GVHD has not been reported in INTERCEPT Platelet recipients based on approximately 7 years of hemovigilance data, surveying the use of INTERCEPT in over 180,000 platelet units. ⁵⁶⁻⁶⁰

Both INTERCEPT treatment and gamma irradiation inactivate leukocytes. Using a limiting dilution assay (LDA), the INTERCEPT Blood System demonstrates reduction of viable T-cells comparable to a 2,500 cGy treatment of gamma irradiation (5-6 log₁₀ inactivation).^{19, 20}

- INTERCEPT treatment results in approximately one amotosalen adduct per 83 base pairs, a sufficient frequency to ensure inactivation of most genes.²¹
- In an *in vitro* study, both untreated and gamma irradiated platelet samples spiked with high levels of leukocytes had significant levels of interleukin 8 after 5 to 7 days of storage, while cytokine production was substantially inhibited in the sample treated with amotosalen and UVA light.²²
- In an *in vivo* murine transfusion model, mice receiving untreated splenic leukocytes developed TA-GVHD, while mice infused with gamma irradiated (2,500 cGy) or INTERCEPT treated splenic leukocytes remained healthy and did not develop TA-GVHD.²³

In vitro Characterization of Platelet Components Processed with the INTERCEPT Blood System.

In vitro platelet function characteristics were evaluated in paired in vitro studies of INTERCEPT apheresis platelet components and INTERCEPT whole-blood derived buffy coat platelets compared to untreated (Control) platelet components. In vitro platelet function of the platelet components were evaluated on day 5 and day 7 after donation. The physical and metabolic characteristics evaluated for each study are included in **Table 6a**, **Table 6b** and **Table 6c**.

Double-dose Amicus platelets containing 5.4 to 10.6 x10¹¹ platelets in 100% plasma were split evenly to produce an untreated Control and an INTERCEPT treated platelet using the INTERCEPT Blood System for Platelets large volume (LV) platelet processing set and stored for 7 days post-donation.

Table 6a In Vitro Platelet Function Characteristics of INTERCEPT and Control Apheresis Platelets in 100% plasma After 7 Days of Storage (Mean +/- SD, [range])

Parameter	Day 5 mean ±SD and [range]		Day 7 mean ±SD and [range]			
(Units)	INTERCEPT Treated Test PC	Untreated Control PC	N	INTERCEPT Treated Test PC	Untreated Control PC	N
	Apheresis Plat	elet Components Su	spended in	100% Plasma (SUD 0	0666)	
Component Volume (mL)	294 ± 6 [284-300]	297 ± 3 [294-303]	7	283 ± 4 [275-287]	289 ± 4 [285-297]	6
Platelet Count (x10 ⁶ /mL)	1108 ±215 [790-1373]	1255 ±246 [906-1611]	7	1286 ±266 [895-1647]	1153 ± 167 [912 - 1353]	6
Platelet Dose (x10 ¹¹ /unit)	3.3 ±0.6 [2.3-3.9]	3.7 ±0.7 [2.7-4.8]	7	3.6 ± 0.7 [2.6-4.5]	3.3 ±0.4 [2.7-3.9]	6
pH (22°C)	7.30 ± 0.15 [7.13-7.54]	7.38 ± 0.19 [7.03-7.65]	7	7.13 ± 0.31 [6.59-7.49]	7.06 ± 0.23 [6.86-7.35]	6
pO ₂ (mmHg)	160 ±5 [154-168]	141 ±13 [117-161]	7	159 ±12 [139-175]	164 ±3 [159-169]	6
pCO ₂ (mm Hg)	14 ±1 [12-15]	17 ±2 [15-20]	7	16 ±3 [11-19]	13 ±1 [12-15]	6
HCO ₃ (mM)	6.4 ±3.7 [2.3-12.2]	9.2 ±5.1 [0-16.3]	7	4.5 ±4.5 [0-12]	2.5 ±3.8 [0-7.6]	6
Glucose (mg/dL)	199 ±16 [179-225]	222 ±32 [165-264]	7	173 ±45 [102-231]	159 ±27 [130-191]	6
Lactate (mM)	13.0 ±2.2 [10.4-15.7]	12.2 ±3.8 [8.2-19.4]	7	16.6 ±4.9 [10.8-25.1]	16.9 ±3.0 [13-20.4]	6
LDH (U/L)	420 ±69 [316-535]	373 ±56 [285-441]	7	373 ±54 [304-426]	430 ±81 [305-545]	6
Swirling	+	+	7	+	+	6
MPV (μm³)	7.3 ±0.6 [6.8-8.6]	7.4 ±0.7 [6.8-8.7]	7	7.3 ±0.4 [6.7-7.6]	7.3 ±0.4 [6.7-7.8]	6
p-selectin (ng/mL)	210 ±18 [184-235]	225 ±19 [195-248]	7	261 ±22 [224-276]	270 ±39 [217-313]	6

Apheresis-Derived Platelets in Additive Solution

Single-dose and double-dose Amicus platelets containing 2.9 to 8.0 x10¹¹ platelets in 35% plasma/65% PAS-3 were either stored untreated (Control) or treated with the INTERCEPT Blood System for Platelets dual-storage (DS) platelet processing set and stored for 7 days post-donation.

Table 6b In Vitro Platelet Function Characteristics of INTERCEPT and Control Apheresis Platelets in PAS-3 After 7 Days of Storage (Mean +/- SD, [range])

Donomoton (Unite)	Day 5 mean ±SD and [range] INTERCEPT Treated Test PC Untreated Control PC N		Day 7 mean ±SD and [range]			
Parameter (Units)			INTERCEPT Treated Test PC	Untreated Control PC	N	
Double and Sin	gle Dose Apheresis	Platelet Componen	ts Suspen	ded in Platelet Additi	ve Solution (CLI 00	094)
Platelet Component	Characteristics					
Component volume (mL)	279±74 [162-382]	288±74 [152-400]	63	262±65 [147-360]	272±66 [150-382]	63
Platelet Count (x10 ⁶ /mL)	1383 ±341 [717-2002]	1482 ±375 [730-2250]	63	1350 ±310 [720-1945]	1445 ±373 [678-2323]	63
Platelet Dose (x10 ¹¹ /unit)	3.7 ± 0.8 [2.5-6.1]	4.0 ±0.6 [2.3-5.5]	63	3.4 ± 0.7 [2.4-5.7]	3.7 ± 0.5 [2.0-4.7]	63
MPV (fL)	8.1±0.8 [6.7-10.6]	8.1±0.9 [6.7-10.4]	58	8.3±1.0 [6.5-11.3]	8.2±1.0 [6.6-10.4]	55
Indices of Platelet M	etabolism					
pH (22 °C)	7.02 ± 0.11 [6.78-7.20]	7.03 ±0.12 [6.73-7.25]	63	7.06 ±0.09 [6.86-7.24]	7.10 ± 0.10 [6.83-7.35]	63
pO ₂ (mm Hg)	130±2 [66-168]	124±20 [70-154]	63	133±17 [90-174]	123±21 [70-160]	63
pCO ₂ (mm Hg)	20±6 [10-32]	24±6 [13-35]	63	18±5 [10-26]	21±5 [10-30]	63
HCO ₃ ⁻ (mmol/L)	3.2±1.1 [1.0-5.3]	3.8±1.3 [2.0-7.0]	60	3.1±0.9 [2.0-5.0]	3.9±1.2 [1.0-6.0]	56
Supernatant glucose (mg/dL) ³	21.5±24.0 [0.0-127.8]	15.5±23.3 [0.0-120.6]	61	6.6±15.3 [0.0-102.6]	6.1±15.1 [0.0-99.0]	61
Supernatant lactate (mmol/L)	11±2 [7-15]	12±3 [7-19]	63	12±2 [8-15]	13±2 [9-18]	63
Total ATP (nmol/10 ⁸ Plts)	4.70±1.95 [1.07-9.39]	4.47±1.72 [1.20-10.25]	63	4.34±1.76 [0.12-8.45]	4.22±1.51 [1.21-10.08]	63
Indices Correlating v	vith In Vivo Recove	ry, Survival and Fu	nction			
Morphology	257±47 [176-360]	251±45 [186-360]	63	235±44 [158-335]	233±44 [124-340]	63
Extent of Shape Change (ESC; %)	12.2±8.1 [1.0-49.0]	11.9±9.1 [0.0-32.6]	63	10.8±6.2 [0.0-20.7]	11.7±7.2 [0.0-27.0]	63
Hypotonic Shock Response (HSR; %)	28±18 [0-56]	30±19 [0-62]	63	25±14 [0-51]	28±18 [0-55]	62
Indices of Granular	and Cytoplasmic Co	ntent Retention				
LDH activity (IU/L)	149±92 [10-581]	142±83 [60-460]	63	198±104 [74-586]	177±83 [74-549]	63
LDH (IU/10 ¹² Plts)	110±54 [8-329]	97±49 [35-270]	63	155±85 [39-382]	127±60 [40-322]	63
P-selectin (%)	36.98±16.22 [7.93-83.31]	33.70±13.96 [9.28-74.47]	60	40.96±13.39 [9.47-78.83]	35.40±13.56 [11.06-77.28]	60

Pools of 10 to 12 whole-blood derived buffy coat platelets in 36 to 42% plasma/58-64% PAS-3 were split evenly to produce an untreated Control and an INTERCEPT treated platelet using the INTERCEPT Blood System for Platelets large volume (LV) platelet processing set and stored for 7 days post-donation.

Table 6c In Vitro Platelet Function Characteristics of INTERCEPT and Control Whole Blood Derived Buffy Coat Platelets in PAS-3 After 7 Days of Storage (Mean +/- SD)

Donometon (Unite)	Parameter (Units) INTERCEPT Untreated Control PC N		Day 7 mean ±SD			
rarameter (Umis)			N	INTERCEPT Treated Test PC	Untreated Control PC	N
Whole-Blood Derive	ed, Single Dose Platel	et Components Susp	ended in I	Platelet Additive Solu	tion (148-REP-NIV	-FE-505)
Parameter (Units)	INTERCEPT Treated Test PC	Untreated Control PC	N	INTERCEPT Treated Test PC	Untreated Control PC	N
Component Volume (mL)	331 ± 23	346 ± 20	6	324 ± 24	338 ± 23	6
Platelet Count (x10 ⁶ /mL)	870 ±68	928 ±77	6	852 ±90	922 ±72	6
Platelet Dose (x10 ¹¹ /unit)	2.9 ±0.3	3.2 ±0.3	6	2.8 ±0.4	3.1 ±0.3	6
pH (22°C)	7.25 ± 0.02	7.32 ± 0.03	6	7.20 ± 0.03	7.28 ± 0.08	6
pO ₂ (mmHg)	18.8 ±2.1	19.7 ± 1.8	6	18.5 ±2.0	16.1 ±3.6	6
pCO ₂ (mm Hg)	3.0 ±0.2	3.3 ±0.2	6	2.9 ±0.3	3.4 ±0.2	6
HCO ₃ (mmol/L)	5.7 ±0.6	6.9 ±0.6	6	4.6 ±0.8	6.0 ±0.7	6
Glucose (mmol/L)	4.8 ±0.5	5.2 ±0.6	6	3.3 ±0.5	3.6 ± 0.5	6
Lactate (mmol/L)	12.3 ±0.9	12.6 ±0.9	6	15.4 ±1.1	15.7 ±1.0	6
ATP	4.9 ±0.3	5.3 ±0.5	6	4.7 ±0.3	5.2 ±0.4	6

Together these studies support the comparability of INTERCEPT-treated apheresis platelets, whether in platelet additive solution or in 100% plasma, with untreated platelets through 5 days of storage. In addition, these studies support the comparability of INTERCEPT-treated whole blood-derived buffy coat platelets in platelet additive solution with untreated platelets through 5 days of storage. All units met CSA/CAN-Z902-10 for pH (22° C) of 6.4-7.8 and local, or study design, requirements for platelet dose.

INSTRUCTIONS FOR USE

- Do not use if: tamper-evident package has been opened; signs of deterioration or damage are visible; fluid path closures are loose or not intact; cannulae are broken; there is no fluid in amotosalen solution container; CAD is bent or broken; base labels are not legible, accurate, or are missing.
- Do not freeze. Do not store above 25°C. Do not vent. Protect the pack and tubing from sharp objects.
- Sets removed from the aluminum foil must be used within **8 hours**.
- Record the Date Opened on the foil pouch label in space provided.
- Unused sets may be kept **20 days** at room temperature in open aluminum foil by folding and securing open end of aluminum foil. Record the Use By date on the foil pouch label in space provided.
- Keep set in light-protective package until time of use. Protect from direct sunlight and strong UVA light source.
- Set is single use only.

This process is designed to be a closed system. Treatment with INTERCEPT Blood System does not replace applicable standards for processing in open and closed systems. If there is a leak in the set during processing, platelet product must be discarded.

MATERIALS AND EQUIPMENT

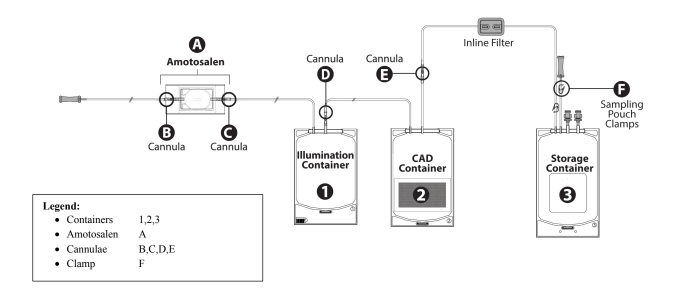
Materials Provided: One (1) INTERCEPT Large Volume Processing Set

Equipment Provided Separately: INTERCEPT Illuminator, Platelet Additive Solution (PAS 3, InterSol (InterSol Solution Platelet Additive Solution 3, manufactured by Fenwal, Inc.) is the only platelet additive solution currently approved in Canada for use with the INTERCEPT Blood System, and is supplied by Cerus upon request)

Equipment Required but Not Provided: Sterile Connecting Device (SCD), Tube Sealer, Flatbed Agitator, Manual Tube Clamp (e.g., Hemostat)

Temperatures should be controlled to meet applicable regulations for platelet processing. Used and unused INTERCEPT sets should be discarded like any used blood containers, as biohazardous waste.

Figure 3 INTERCEPT Large Volume Processing Set



Performing the INTERCEPT Treatment Process

Instructions refer to Figure 3 for labeling and identification of set components.

All the following conditions must be met for pathogen inactivation:

- Platelet source, content, volume, and red blood cell (RBC) count must be within the ranges specified in **Table 7**.
- Follow apheresis device instructions for platelet handling and resting procedures prior to INTERCEPT treatment.
- Platelets should meet institutional specifications for aggregates prior to INTERCEPT treatment.
- Platelets mixed with amotosalen must be exposed to UVA light dose from INTERCEPT Illuminator. No other source of UVA light may be used.
- Platelets collected on Day 0 must be exposed to UVA light by end of Day 1.
- After illumination, platelets must be agitated in CAD container in accordance with the duration specified in **Table 8.**

I. Preparation of Platelets

Platelet products within the following ranges have been shown to be acceptable for use with this processing set (**Table** 7).

Table 7 Platelet Collection Specifications for INTERCEPT
Large Volume Processing Set

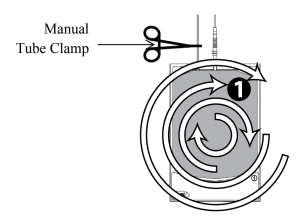
	Suspension Medium			
	PAS	100% Plasma		
Platelet Source	Apheresis, Whole Blood-Derived Buffy Coat Platelets	Apheresis		
PAS	InterSol® PAS 3*	Not Applicable		
Platelet Input Volume	300 – 420 mL	255 – 420 mL		
Platelet Dose	$2.5 - 7.0 \times 10^{11}$	$2.5 - 7.0 \times 10^{11}$		
PAS Content	53 - 68%	0%		
Plasma Content	32 - 47%	100%		
RBC Content	< 4x10 ⁶ RBC/mL	< 4x10 ⁶ RBC/mL		

^{*} InterSol is trademark of Fenwal, Inc.

II. Amotosalen Addition to Platelets

- 1. Remove set from package. Unwrap only illumination container (**1**) from organizer.
- 2. Weld tubing from platelet collection container to amotosalen container (**②**) tubing using SCD.
- 3. Label storage container (3) using appropriate donation identification.
- 4. Hang platelet collection container and break cannula (**G**) below amotosalen container (**A**) to let amotosalen flow into illumination container (**O**); visually verify amotosalen is present.
- 5. Break cannula (**B**) above amotosalen container (**A**) to let platelets flow through amotosalen container (**A**) into illumination container (**D**).
- 6. Ensure the platelets drain completely from initial platelet collection container into illumination container (①) by expressing air from the illumination container (①) into amotosalen container (②).
- 7. When air is removed and platelets have fully drained back into illumination container (**1**), manually clamp tubing above illumination container. Mix illumination container thoroughly by gentle agitation to ensure complete mixing of amotosalen and platelets (**Figure 4**).

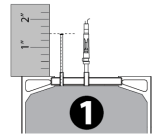
Figure 4 Gently Agitate the Filled Illumination Container



- 8. Open manual tube clamp and expel remainder of air in illumination container (**1**) into amotosalen container (**1**).
- 9. Express a small amount of platelet and amotosalen mixture into tubing, filling about 1.5 inches of tubing. Close manual clamp.
- 10. Seal tubing between illumination container (**1**) and amotosalen container (**2**) so that tubing is no longer than 1.5 inches from the illumination container (**1**) (See Figure 5 below; also refer to INTERCEPT Illuminator Operator's Manual for further details on loading the processing set into the Illuminator).

Warning: During illumination, tubing must be held within large compartment of illumination tray.

Figure 5 Heat Seal Filled Tubing Within 1.5 Inches



11. Remove and discard empty platelet collection container, amotosalen container (**②**) and excess tubing.

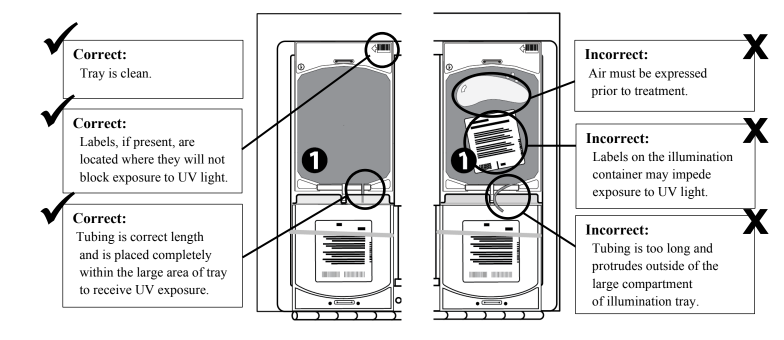
III. Illumination

Illuminate platelets.

Refer to INTERCEPT Illuminator Operator's Manual for complete illumination instructions for use.

<u>Warning</u>: All platelets, both in illumination container and tubing, must be within large compartment of illuminator tray in order for inactivation to occur. The process requires unimpeded light transmission through tray and illumination container with platelets. **No labels or other material should be on this area.** Tray must be clean. Labels should be placed on illumination container flap only. Illumination container should lay flat in order to ensure complete illumination.

Figure 6 The Correct and Incorrect Way to Load a Processing Set Into the Illuminator Tray



IV. Agitation with Compound Adsorption Device (CAD)

Warning: Do not fold or bend CAD.

- 1. Unwrap set from organizer.
- 2. Hang platelets in illumination container (**1**), break cannula (**1**) and allow platelets to flow into CAD container (**2**).
- 3. Express air from CAD container (**2**) into illumination container (**1**). Do not fold or bend CAD.
- 4. Seal tubing close to inlet port of CAD container (**2**).
- 5. Remove and discard empty illumination container (**1**) and excess tubing.
- 6. Place CAD container (2) flat on flatbed agitator with attached storage containers for duration specified in Table 8. Agitation rate must be at least 60 RPM.
- 7. Do not fold or place storage container (3), in line filter, or any tubing under CAD container (2) during agitation.

Suspension Medium CAD Agitation Duration
PAS 6 to 16 hours

16 to 24 hours

Table 8 CAD Agitation Duration

V. Transfer to Storage Container

- 1. Remove platelets from agitator and hang platelets in CAD container (**2**).
- 2. Expel air from platelet sampling pouch and close clamp (**9**).

100% Plasma

- 3. Break cannula (3) at outlet of CAD container (2) and allow platelets to flow into storage container (3).
- 4. Express air from storage container (3) into CAD container (2).
- 5. Ensure appropriate donor identification is attached to the filled platelet storage container (3).
- 6. Seal tubing to storage container (**3**).
- 7. Remove and discard empty CAD container (**2**) and tubing. Place platelets on platelet agitator.

The INTERCEPT treatment process is now complete.

Sampling Platelet Product (optional)

- 1. Mix INTERCEPT treated platelets by gently agitating platelet storage container.
- 2. Open clamp (**6**) to platelet sampling pouch and squeeze several times.
- 3. Allow sampling pouch to fill with platelets. Seal tubing.
- 4. Remove sample pouch.
- 5. Transfer sample to appropriate laboratory tube immediately.

STORAGE

Following processing, INTERCEPT Platelets should be stored in the storage container provided in the processing sets at 20-24°C with continuous agitation.

INTERCEPT Platelets suspended in PAS may be stored for up to 5 days from the day of collection: bacterial inactivation data, as provided in **Table 4**, measure inactivation immediately post illumination. *In vitro* studies, by Schmidt et al ¹⁴, Lin et al ¹⁵, and Nussbaumer et al ¹⁶, have demonstrated that representative bacteria are not detected in INTERCEPT Platelets following up to 5 days of storage.

INTERCEPT Platelets suspended in 100% plasma may be stored for up to 5 days from the day of collection.

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- 61. ">" refers to inactivation below the limit of detection of the assay.
- 62. "\geq" refers to inactivation equal to or below the limit of detection of the assay.

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