



# Flow CAST<sup>®</sup>

## Basophil Activation Test (BAT) Flow Cytometry

This product is for research use only  
It is not intended for use in diagnostic procedures

FK-CCR-U 100 tests

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## INTENDED USE

The BÜHLMANN Flow CAST® is a test for the assessment of basophil activation upon stimulation with specific allergens. The test employs flow cytometry to determine the basophil population expressing CD63 cell surface marker in K-EDTA whole blood samples.

For research use only. Not for use in diagnostic procedures.

## PRINCIPLE OF THE ASSAY

Flow CAST® is a flow cytometry-based basophil activation test. Whole blood from individuals is stimulated with specific allergens, as well as stimulation buffer and stimulation controls, to evaluate the sample's basophil degranulation *ex vivo*. The sample is stained using two fluorescently labeled monoclonal antibodies: one for basophil selection (anti-CCR3-PE) and one for basophil activation status determination (anti-CD63-FITC). CD63 is a transmembrane protein present on intracellular vesicles and only presented on the cell surface after basophil degranulation.

Erythrocytes from the sample are removed by a lysing reaction. Depending on the protocol, the cells are centrifuged, resuspended in wash buffer and fixed for later analysis by flow cytometry or analyzed directly after lysis. Basophils are gated from the leukocyte population as CCR3<sup>pos</sup>/SSC<sup>low</sup>. The activation status of the gated basophils is determined by their CD63 expression (activation marker). Samples who do not elicit IgE-mediated responses, so called non-responders, are identified based on the results of the positive controls. The readout of the assay is indicated as the ratio of CD63 positive basophils over all basophils (%CD63 activation).

## REAGENTS SUPPLIED AND PREPARATION

Reagents	Quantity	Code	Comments
<b>Stimulation Buffer</b> containing calcium, heparin and IL-3	1 vial lyophilized	B-CCR-STB	Reconstitute with 50 mL of water <sup>1)</sup>
<b>Stimulation Control</b> anti-FcεRI mAb	1 vial lyophilized	B-CCR-STCON	Reconstitute with 1.5 mL of B-CCR-STB
<b>Stimulation Control</b> fMLP <sup>2)</sup>	1 vial lyophilized	B-CCR-FMLP	Reconstitute with 1.5 mL of B-CCR-STB
<b>Staining Reagent</b> Mix of anti-CD63-FITC and anti-CCR3-PE mAb	1 vial 2.2 mL	B-CCR-SR	Ready to use
<b>Lysing Reagent</b> <sup>3)</sup> 10x concentrated	1 vial 25 mL	B-CCR-LYR	Dilute with 225 mL of deionized water
<b>Wash Buffer</b> with 0.1% formaldehyde	1 vial 100 mL	B-CCR-WB	Ready to use

Table 1

<sup>1)</sup> For required water quality, refer to chapter Technical Precautions

<sup>2)</sup> N-formyl-methionyl-leucyl-phenylalanine

<sup>3)</sup> Crystals may be formed during storage at 2-8°C and should be dissolved at 18-28°C prior to dilution

## STORAGE AND SHELF LIFE OF REAGENTS

Unopened reagents	
Store at 2-8 °C. Do not use the reagents beyond the expiration date printed on the labels.	
Opened reagents and reconstituted reagents	
Stimulation Buffer	Stable at -20°C for 6 months. Aliquot if repeated use is expected.
Stimulation Control	
Stimulation Control fMLP	
Lysing Reagent	Stable at 2-8°C for 6 months.
Staining Reagent	
Wash Buffer	

Table 2

## MATERIALS REQUIRED BUT NOT PROVIDED

- K-EDTA venipuncture tubes
- Centrifuge
- Disposable, pyrogen-free polypropylene or polystyrene flow cytometry test tubes
- Flow cytometry test tube racks for the stimulation
- Vortex mixer
- (Optional) tissue culture grade microtiter plates for cell stimulation and staining for the standard protocol
- (Optional) deep-well plates for cell stimulation, staining, lysis and flow-cytometry acquisition for the lyse-no-wash protocol
- Precision pipettes with disposable, pyrogen-free tips:
  - 10-100 µL, 100-1000 µL,
  - 1-5 mL adjustable pipette and
  - Optional: 10-50 µL adjustable dispenser
- 50 mL cylinder for stimulation buffer reconstitution
- Sterile, ultrapure and apyrogenic water for reconstitution of the stimulation buffer (refer to chapter Technical Precautions)
- Water bath (recommended) or incubator set at 37°C
- Distilled or deionized water as well as appropriate laboratory glassware for the dilution of lysing reagent
- Lids or parafilm to cover tubes during incubation steps
- Bottle-top dispensers for lysing reagent and wash buffer
- Flow cytometer equipped with a 488 nm (blue) laser source as well as emission filters for PE and FITC detection
- Flow cytometric analysis software (refer to chapter Flow Cytometric Data Acquisition)

## PRECAUTIONS

### Safety precautions

- The stimulation buffer (B-CCR-STB) of this test contains components of human origin. Although tested and found negative for HBV surface antigen, HCV and HIV1/2 antibodies, the reagents should be handled as if capable of transmitting infections and should be handled in accordance with Good Laboratory Practices (GLP) using appropriate precautions.
- Avoid contact of reagents with the skin, eyes, or mucous membranes. If contact does occur, immediately wash with generous amounts of water.

- Reagents and chemicals have to be treated as hazardous waste according to the national biohazard safety guideline or regulation.

### Technical precautions

- Recommended water quality for the Flow CAST®: The use of sterile, ultrapure and apyrogenic water to reconstitute stimulation buffer (B-CCR-STB) is essential for good and reproducible basophil stimulation. The following sources of water may be used: cell culture grade water, infusion grade water or deionized, double distilled water that is ultra-filtered in a periodically sanitized 10 kDa ultra filter.
- The lysing reagent (B-CCR-LYR) can be reconstituted with deionized, double distilled water or the same water quality that is used for the reconstitution of the stimulation buffer.
- Avoid allergen contamination during cell stimulation: Aeroallergens in the laboratory may contaminate open blood samples and cell suspensions, causing an elevated background. Blood samples and cell stimulation tubes must be carefully covered by lids or parafilm. Avoid house dust mites, pollinating plants, latex gloves or equipment potentially containing latex as well as open windows in the laboratory where the cell stimulation is performed. It is recommended to carry out the cell preparation and stimulation steps in a laminar flow hood.
- A water bath is recommended compared to an incubator, due to more efficient heat transfer. If using an incubator, verify that the temperature is 37°C. Lower or higher temperatures may affect results.
- Generally low level of basophil activation is expected for drug allergens. It is therefore crucial that optimal conditions during stimulation including temperature are achieved. The use of single tubes instead of deep-well plates is recommended for drug allergens.
- Components must not be used after the expiry date printed on the labels.
- Do not mix different lots of reagents.
- Avoid contamination of reagents.

### Test procedure

- Equilibrate the lysing reagent to room temperature (18-28 °C).
- Carefully read the instructions prior to carrying out the test. Test performance will be adversely affected, if reagents are incorrectly diluted, handled or stored under conditions other than those detailed in this instruction for use.
- Samples that are not properly handled may cause inaccurate results.
- Verify the preparations by eye to assess the efficacy of lysis. The erythrocytes may be incompletely lysed and appear on a light diffraction dot plot in the same location as the leucocytes.
- Prolonged lysis time can lead to cell loss. Ensure that you have at least 300 basophils for data acquisition. We recommend that acquisition of samples processed with the lyse-no-wash protocol is performed within an hour.
- Flow cytometry may produce false results, if: the cytometer is misaligned, the fluorescence emission has not been appropriately compensated, the gating regions have not been carefully positioned.

## SPECIMEN COLLECTION AND STORAGE

It is recommended that individuals avoid systemically administered antiallergenic drugs such as corticosteroids, chromoglycic acid (DSCG) for at least 24 hours prior to blood sampling.

Collect blood into **K-EDTA venipuncture tubes**, by filling the tubes up to the dedicated volume mark. Tubes must be filled at least half-way. One (1) mL of whole blood is sufficient for approximately 18 test tubes.

**Do not centrifuge or freeze blood samples.**

### Whole blood

Whole blood samples stored at 2-8 °C should be processed within 48 hours of collection.

For determination of IgE mediated responses to drugs it is advised to process samples immediately and no later than within 24 hours after sample collection.

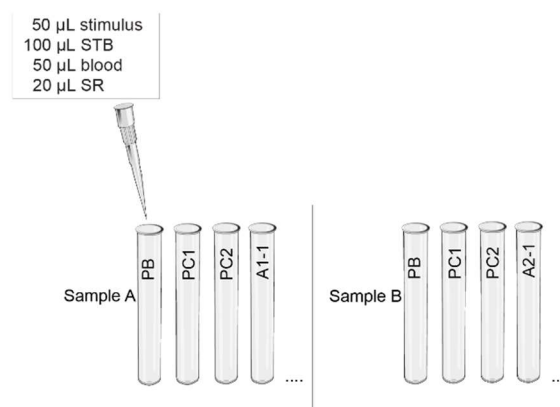
Whole blood samples may also be kept at room temperature (temperatures up to 28 °C). They must however be processed within 24 hours of collection using the standard protocol or on the day of collection using the lyse-no-wash protocol.

### Processed samples

Cells processed using the standard protocol are fixed. Fixed cells may be stored at 2-8 °C for 5 days protected from light for subsequent acquisition by flow cytometry.

## ASSAY PROCEDURE

1. Mix the anti-coagulated blood sample by inverting the venipuncture tube several times.
2. Prepare fresh and pyrogen-free standard polypropylene or polystyrene flow cytometry tubes.
3. For each sample, label the tubes e.g. the following:  
PB = sample background  
PC1 = stimulation control with anti-FcεRI Ab  
PC2 = stimulation control with fMLP  
A1-1 for allergen 1 with dilution 1  
A1-2 for allergen 1 with dilution 2 etc.



### Stimulation and staining

4. Add 50 µL of the corresponding stimulus to each tube:  
PB tube: 50 µL of **stimulation buffer** (sample background)  
PC1 tube: 50 µL of **stimulation control** anti-FcεRI mAb

PC2 tube: 50  $\mu$ L of **stimulation control** fMLP

Ax-y tube: 50  $\mu$ L of **allergen**

5. Add 100  $\mu$ L of stimulation buffer (STB) to each tube.
6. Add 50  $\mu$ L of sample's whole blood to each tube. Make sure that the side and top of the tube are free from blood.
7. Mix gently.
8. Add 20  $\mu$ L staining reagent (SR) to each tube.
9. Mix gently, cover the tubes and incubate for 15 minutes at 37°C in a **water bath**.

**Note:** If an incubator, instead of a water bath, is used the incubation time is prolonged to 25 minutes due to less efficient heat transfer.

## Lysing

**Note:** The lysing reagent must be equilibrated to room temperature (18-28 °C).

### Standard protocol: Lyse and wash

10. Add 2 mL equilibrated (18-28°C) lysing reagent to each tube, mix gently.
11. Incubate for 5-10 minutes at 18-28°C.
12. Centrifuge the tubes for 5 minutes at 500 x g.
13. Decant the supernatant by using blotting paper.
14. Resuspend the cell pellet with 300  $\mu$ L of wash buffer (a fixative is included in the wash buffer).

**Note:** The amount of wash buffer may be adapted to the specific flow cytometer instrumentation used, according to the dead volume and cell density compatible with the device.

15. Vortex gently.

**Either** 16a. acquire the samples on the flow cytometer.

**Or** 16b. if not acquired immediately, let the samples incubate for 30 min at RT and protected from light (fixation). Store the samples sealed and protected from light at 2-8°C until measurement. Fixed cells may be stored at 2-8°C for 5 days for subsequent acquisition by flow cytometry. Vortex gently the sample tubes prior to acquisition.

**Note:** Stored fixed samples can be acquired without any pre-treatment at any time. Please refer to section "Specimen Collection and Storage" for storage times. A slight decrease of fluorescence intensity and a lower basophil recovery  $\geq 80\%$  may be observed after longer storage.

### Alternative protocol: Lyse-no-wash protocol

New generation, high-performance flow cytometers can analyze lysed, unwashed samples. This procedure must be adapted to the flow cytometer instrumentation used and may require optimization. The protocol below is based on data acquired with an Attune NxT flow cytometer (Thermo Fisher).

10. Perform assay procedure steps 1 to 9 (above), and then continue at step 10 here. Add 1.5  $\pm$  0.5 mL equilibrated (18-28°C) lysing reagent to each tube, mix gently (volume must be optimized depending on acquisition speed capabilities of the flow cytometer used).
11. Acquire the samples using a high throughput suitable flow cytometer device at high acquisition speed to keep analysis time minimal.

**Note:** Samples should be analyzed within 24 hours of receiving the sample. Please refer to section "Specimen Collection and Storage".

## FLOW CYTOMETRIC DATA ACQUISITION

Flow cytometric acquisition can be performed on any flow cytometer working with a 488 nm argon laser diode (blue-green excitation light).

The flow cytometer must be equipped to detect Forward Scatter (FSC), Side Scatter (SSC) and the two fluorochromes FITC and PE channels.

Ensure that the flow cytometer is properly aligned, and color compensation is set.

For the appropriate acquisition and characterization of resting and activated basophils create the following dot plot:

1. Create dot plot 1, as Forward Scatter vs Side Scatter to acquire the whole leukocyte population as shown in Figure 1. During acquisition of the samples, make sure that the leukocyte population is separated into three discrete populations (lymphocytes, monocytes and granulocytes) on the FSC/SSC dot plot. Adjust the amplification (gain) of FSC and SSC signals to obtain a distribution as shown in Figure 1. Refer to the flow cytometer product manuals for instructions.
2. Create dot plot 2, as CCR3-PE vs Side Scatter as shown in Figure 2. Set a gate (e.g., basophils) including the entire basophil population as CCR3<sup>pos</sup> and SSC<sup>low</sup> as shown with the rectangle gate in Figure 2. Eosinophils, that are also CCR3<sup>pos</sup>, must be excluded based on the high SSC.
3. Create dot plot 3, as CD63-FITC vs CCR3-PE, showing only the gated basophils, as shown in Figure 3. Use the non-stimulated, resting basophils of the sample background (PB) tube to set a quadrant gate including CD63 negative basophils cells in the lower right quadrant (CD63<sup>neg</sup> CCR3<sup>pos</sup>/SSC<sup>low</sup>) as shown in Figure 3. Basophils activated by the stimulation of positive controls and specific allergens will result in CD63 positive basophil population (CD63<sup>pos</sup>/CCR3<sup>pos</sup>/SSC<sup>low</sup>) identified in the upper right quadrant, as shown in Figure 4 with an example of positive control stimulation (STCON).

The readout of the assay is indicated as the ratio of CD63 positive basophils over all basophils (%CD63 activation) as identified in the quadrant gate of the histogram 3 for any of the stimulation tubes.

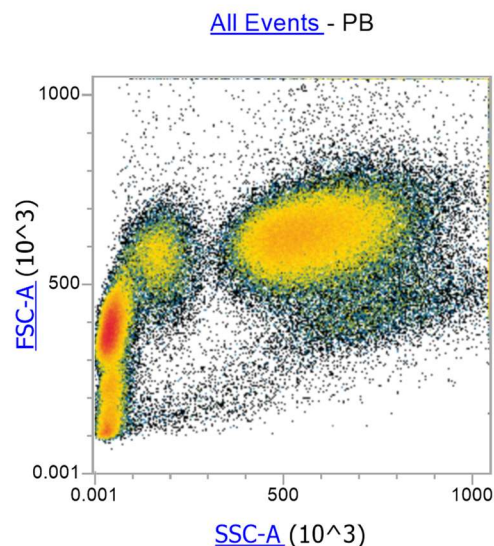


Figure 1: Three discrete populations (lymphocytes, monocytes and granulocytes) on an FSC/SSC dot plot

Acquire 500 or more basophilic cells for any stimulation tubes (gated as shown in dot plot 2, Figure 2 below). If less than 300 basophilic cells are acquired (e.g. in case of basopenia), test results cannot be evaluated.

## DATA ANALYSIS

Acquired data is analyzed with appropriate flow cytometry analysis software. Set similar dot plots and gates as done for the acquisition.

The gates that identify the basophils in dot plot 2 can be independently adapted in any of the different stimulations for the same sample.

For the correct evaluation and standardization of the results a background setting for each single sample is defined using the sample background stimulation (PB). The quadrant gate set on the dot plot 3 must be defined set on the PB. To standardize the analysis, the gating is set to be between 2 and 2.5% activated basophils (see Figure 3).

This gate must be applied to all subsequent stimulations for the same samples (PC1, PC2, and all measured allergens) to calculate the percentage of CD63 positive cells in any stimulation (see Figure 4).

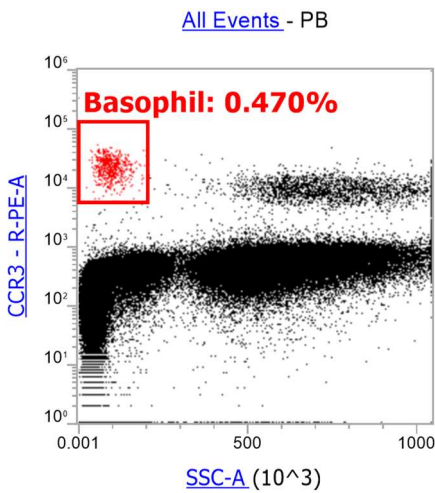
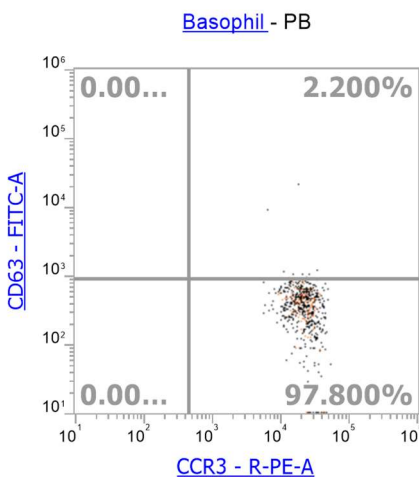
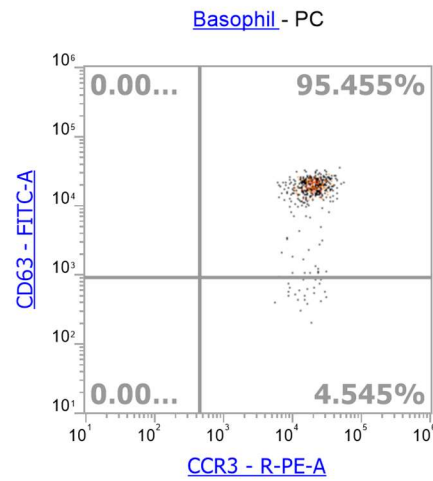


Figure 2: Selection of basophilic cells CCR3<sup>pos</sup> / SSC<sup>low</sup>



Gated Region	Count (n=)	%
Total	125'864	100.0
Basophil	591	0.47
Q2 (CD63 <sup>pos</sup> )	13	2.2
Q4 (CD63 <sup>neg</sup> )	578	97.8

Figure 3: Sample Background (PB) with STB only



Gated Region	Count (n=)	%
Total	130'926	100.0
Basophil	506	0.39
Q2 (CD63 <sup>pos</sup> )	483	95.5
Q4 (CD63 <sup>neg</sup> )	23	4.5

Figure 4: Stimulation Control (STCON)

## QUALITY CONTROL

The following criteria and quality control measures should be met for a valid result:

**Leukocyte populations:** Typically, three distinct leukocyte populations lymphocytes, monocytes and granulocytes should appear in the FSC/SSC plot (see Figure 1). Their occurrence can be regarded as a criterion for the quality of the blood sample (time frame between sample collection and assay execution, storage conditions). Test results cannot be evaluated if less than 300 basophils are acquired.

**Stimulation (positive) controls anti-FcεRI mAb and fMLP:** Anti-FcεRI mAb mimics the bridging of the receptor caused by the allergen *in vivo*. fMLP is a tripeptide causing basophil activation in a non-immunologic way.

- If the Anti-FcεRI mAb control exhibits a value of ≥ 10% activated basophils, the samples can be evaluated.
- If only the fMLP control shows a signal ≥ 10%, but the Anti-FcεRI mAb does not, the assay has been executed correctly, but the sample is considered an IgE non-responder.
- If both Anti-FcεRI mAb and fMLP exhibit values <10% activated basophils a technical error is likely. The test result should be considered invalid and the test should be repeated.

## STANDARDIZATION

Flow CAST® detects the population of basophils expressing the CD63 cells surface marker as % of total basophils. There are no internationally or nationally recognized reference materials or reference measurement procedures for this analyte.

Batch-to-batch reproducibility is guaranteed by titration of anti-CD63-FITC and anti-CCR3-PE monoclonal antibody conjugates against calibration beads. For an estimate of batch-to-batch variation please refer to reproducibility results in section "performance characteristics".

## TECHNICAL CUT-OFF AND REFERENCE INTERVAL

A technical cut-off of 5% activated basophils has been established, where results  $\geq 5\%$  CD63<sup>pos</sup> indicate basophil activation.

Reference intervals were established according to CLSI C28-A3. One hundred and twenty (120) blood samples from a blood donation center were stimulated with stimulation buffer or Anti-Fc $\epsilon$ RI mAb and tested according to standard and lyse-no-wash protocols. Testing was performed over the course of 26 days by three operators with two Flow CAST<sup>®</sup> reagent lots.

Control	Assay Protocol	Reference Interval (90% CI) [% CD63 <sup>pos</sup> ]	
		2.5th percentile	97.5th percentile
Stimulation Buffer	standard	0.8 (0.5 - 1.2)	4.6 (4.1 - 6.4)
	lyse-no-wash	0.9 (0.6 - 1.0)	4.2 (3.9 - 5.3)
Anti-Fc $\epsilon$ RI mAb	standard	18.0 (11.5 - 26.0)	97.7 (96.0 - 98.5)
	lyse-no-wash	13.2 (11.4 - 21.2)	96.4 (94.3 - 97.3)

Table 3

## PERFORMANCE CHARACTERISTICS

### Within-laboratory precision: $\leq 25\%$ CV for stimulus

Repeatability (within-run) and within-laboratory precision were established based on the CLSI guideline EP05-A3 and ISO standard 15197:2013. Four donor blood samples were stimulated with stimulation buffer or stimulation control anti-Fc $\epsilon$ RI mAb. For the standard procedure, a 2 operators x 4 days x 1 run x 4 replicates study design was used. For the lyse-no-wash protocol, a 2 operators x 1 day x 4 runs x 4 replicates study design was applied. A replicate corresponds to an independent stimulation reaction and a full assay procedure. The results for stimulation control anti-Fc $\epsilon$ RI mAb are summarized in Table 4.

Assay Protocol	Donor	Mean [%CD63]	n	Within-run [%CV]	Between-day (A)	
					Between-run (B)	Total [%CV]
Standard (A)	A	34.7	32	8.8%	0.0%	15.9%
	B	90.3	32	1.3%	2.0%	3.6%
	C	82.4	32	1.8%	0.0%	5.1%
	D	91.4	32	1.1%	4.5%	5.0%
lyse-no-wash (B)	E	89.5	32	1.5%	1.1%	1.9%
	F	74.0	32	2.7%	3.5%	6.0%
	G	68.2	32	4.2%	12.5%	15.5%
	H	73.9	32	3.3%	2.9%	5.2%

Table 4

### Reproducibility: $\leq 25\%$ CV for stimulus

Reproducibility was established based on the CLSI guideline EP05-A3 and ISO standard 15197:2013. Four donor blood samples were stimulated with stimulation buffer or stimulation control anti-Fc $\epsilon$ RI mAb. Samples were assayed at two laboratory sites according to the standard protocol. A 3 instruments/lots x 2 operators x 1 day x 5 replicates study design was applied. A replicate corresponds to an independent stimulation reaction and a full assay procedure. The results for stimulation control anti-Fc $\epsilon$ RI mAb are summarized in Table 5.

Donor	Mean [%CD63]	n	Within-run [%CV]	Between-operator [%CV]	Between-lot/instrument [%CV]	Total [%CV]
A	91.6	30	1.4%	2.1%	1.9%	3.2%
B	87.6	30	1.7%	1.2%	3.3%	3.9%
C	91.9	30	0.8%	0.9%	2.1%	2.5%
D	96.5	30	0.5%	0.0%	0.8%	0.9%

Table 5

## INTERFERING SUBSTANCES

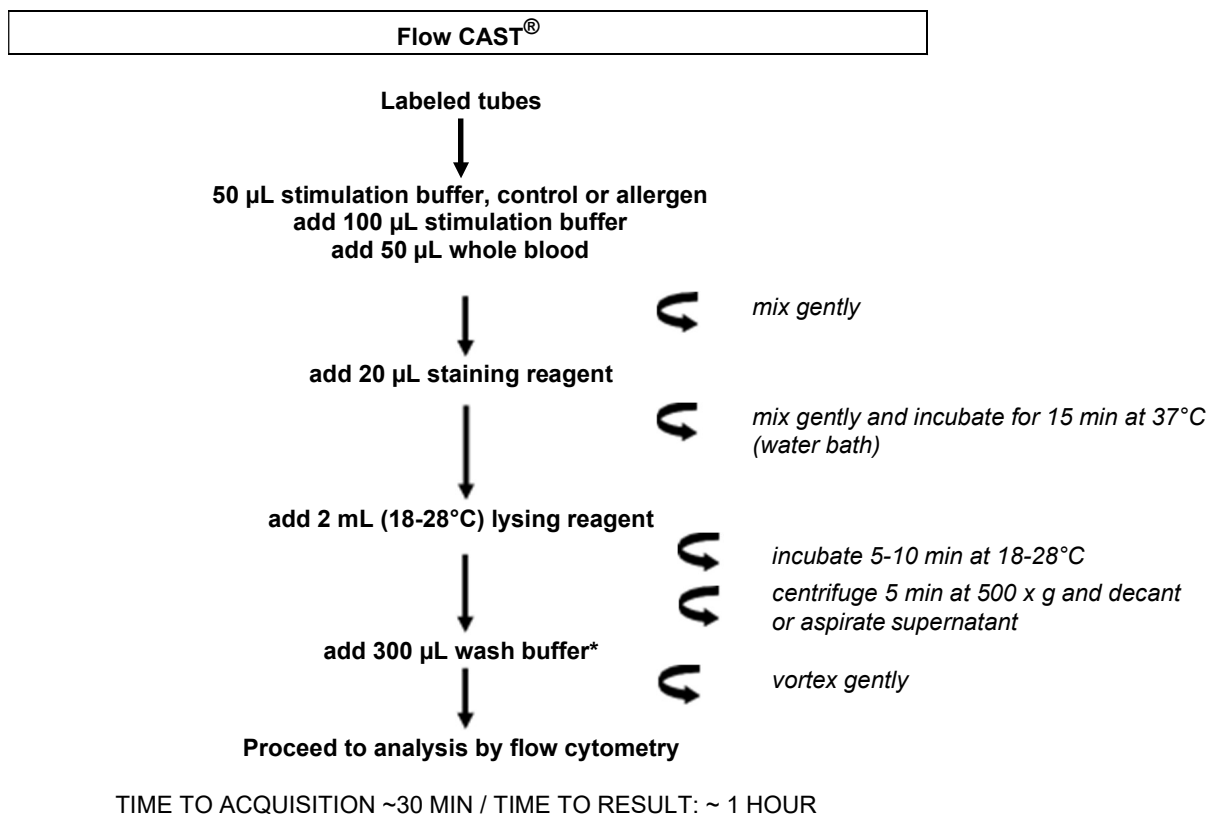
The susceptibility of the Flow CAST<sup>®</sup> assay to pharmaceuticals, abnormal blood conditions and the K-EDTA sample additive was assessed according to the CLSI guideline EP07-A2. Bias in results exceeding 20% for stimulation Control anti-Fc $\epsilon$ RI mAb and 20% CD63<sup>pos</sup> (absolute) for stimulation control fMLP was considered interference. No interferences were detected at the stated concentrations with the substances listed in Table 6 at the listed concentrations. Interference was detected with K-EDTA at double K-EDTA venipuncture tube concentration for one donor.

Active component	Test Concentration [ $\mu$ g/mL]
Fexofenadine hydrochloride	1.6
Cetirizine dihydrochloride	4.35
Hydroxyzine dihydrochloride	0.27
Ketotifen	0.6
Montelukast	3.84
Prednisone	1.2
N-Acetyl-L-tryptophan	30
Triglyceride (Intralipid)	20'000
Bilirubin conjugated	400
Bilirubin unconjugated	400
Hemolysis	56'100

Table 6

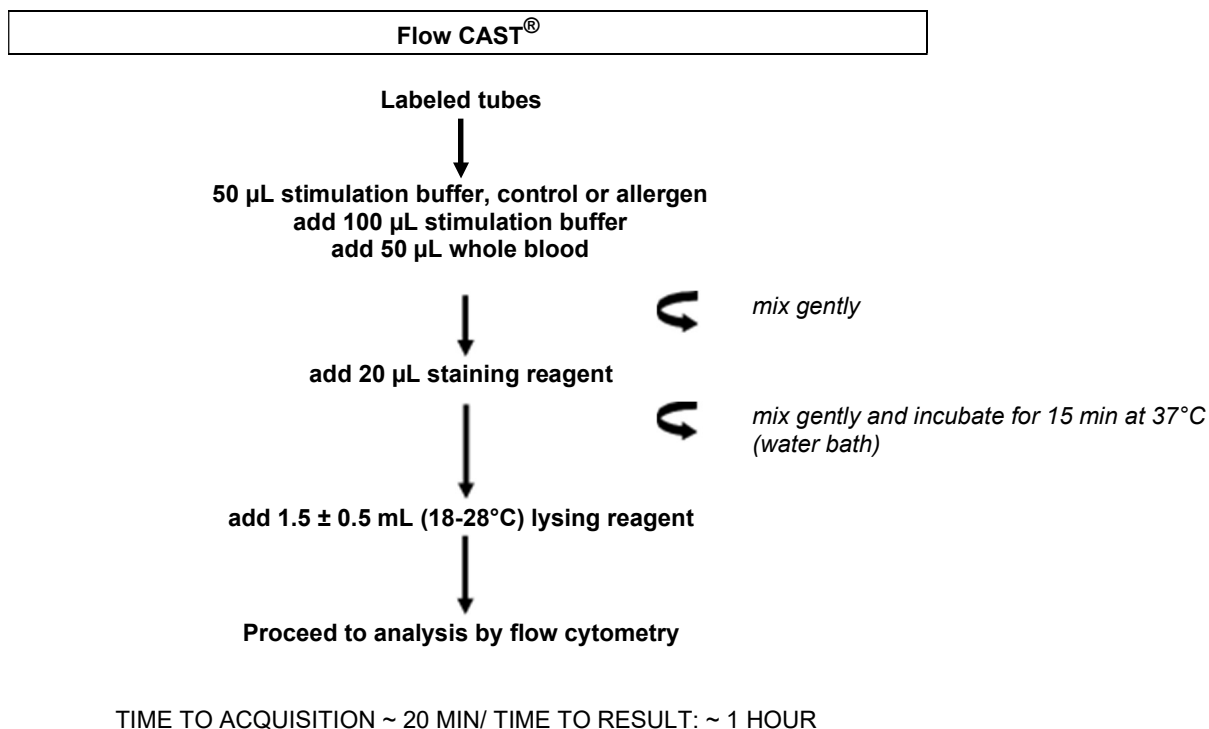
## SHORT PROTOCOLS

### STANDARD PROTOCOL: LYSE AND WASH



\* Note: Depending on the flow cytometer instrumentation used, the amount of wash buffer should be adapted in regard on dead volume and cell density compatible with the instrument.

### ALTERNATIVE PROTOCOL: LYSE-NO-WASH PROCEDURE



## CHANGELOG

Date	Version	Change
2024-04-04	A3	Correction of intended use Removal of reference to CAST® Allergens

## SHIPPING DAMAGE

Please notify your distributor, if this product was received damaged.

## SYMBOLS

BÜHLMANN uses symbols and signs listed and described in ISO 15223-1. In addition the following symbols and signs are used:

Symbol	Explanation
BUF   STIM	Stimulation Buffer
CONTROL   STIM	Stimulation Control
CONTROL   FMLP	Stimulation Control fMLP
REAG   STAIN	Staining Reagent
REAG   LYS	Lysing Reagent
BUF   WASH	Wash Buffer