# **w** abeomics

### 12-4304: Phospho-Aurora A (Thr288)/Aurora B (Thr232)/Aurora C (Thr198) (Clone: CC12) rabbit mAb PE Conjugate

Clonality :	Monoclonal
Clone Name :	AuroraABC-CC12
Application :	FACS
Reactivity :	Human
Conjugate :	PE
Format :	Conjugated
Alternative Name :	Aurora kinase A/B/C, AURKA, AURKB, AURKC
lsotype :	Rabbit IgG1k
Immunogen Information	A synthetic phospho-peptide corresponding to residues surrounding human Aurora A (Thr288)/Aurora B (Thr232)/Aurora C (Thr198)

#### Description

Aurora kinases (serine/threonine kinases) are essential requirement for the onset and progression of mitosis. These kinases share a similar protein structure as well as kinase activity, however each kinase display distinct cellular and subcellular localization. Each Aurora member is phosphorylated at specific residues upon co-factor binding during mitosis. Aurora kinases acquire active kinase conformations due to the activation loop. The active kinase conformation is acquired upon auto-phosphorylation through an intermolecular (trans)-reaction within Aurora kinase domain. Aurora Kinase A (Aurora A) is involved in G2/M transition. AuroraA promotes centrosome maturation and mitotic spindle assembly, whereas AuroraB and AuroraC act as chromosome-passenger complex proteins. They play a crucial role in chromosomal binding to kinetochores and segregation of chromosomes. Aurora B is widely distributed in the cell, while AuroraC is expressed mainly in the meiotically-active germ cells. Aurora kinases are auto-phosphorylated into active forms at conserved threonine residues (i.e. the Thr288 (AurA), Thr232 (AurB) and Thr195 (AurC) residues). AuroraA auto-phosphorylation is initiated by several co-factors acting at different steps of mitosis. AroraB and AruroaC auto-phosphorylation are mediated by survivin and Borealin proteins.

#### **Product Info**

Amount :	10 Tests / 100 Tests
Content :	1X PBS, 0.09% NaN3, 0.2% BSA
Storage condition :	Store at 2-8°C. Avoid repeated freeze and thaw cycles.

#### **Application Note**

For flow cytometric staining, the suggested use of this reagent is 5  $\tilde{A} \square \hat{A} \mu L$  per million cells or 5  $\tilde{A} \square \hat{A} \mu L$  per 100  $\tilde{A} \square \hat{A} \mu L$  of staining volume. It is recommended that the reagent be titrated for optimal performance for each application. See product image legends for additional information.

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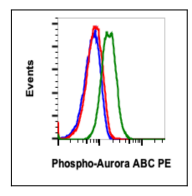


Fig-1: Flow cytometric analysis of HeLa cells untreated and unstained as negative control (blue) or untreated red) or treated with nocodazole (green) and stained using Phospho-Aurora A (Thr288)/Aurora B (Thr232)/Aurora C (Thr198) antibody AuroraABC-CC12 PE conjugate.