

# Anti-E Cadherin antibody [EP700Y] - BSA and Azide free ab256580

KO 評価済 リコンビナント RabMAb

2 References 画像数 21

### 製品の概要

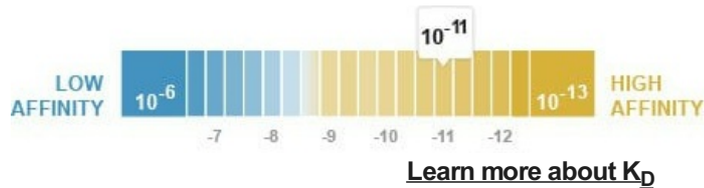
製品名	Anti-E Cadherin antibody [EP700Y] - BSA and Azide free
製品の詳細	Rabbit monoclonal [EP700Y] to E Cadherin - BSA and Azide free
由来種	Rabbit
アプリケーション	適用あり: Flow Cyt (Intra), WB, IHC-P, ICC/IF, mIHC
種交差性	交差種: Human
免疫原	Synthetic peptide. This information is proprietary to Abcam and/or its suppliers.
ポジティブ・コントロール	IHC-P: Human breast carcinoma, lung adenocarcinoma and colonic adenocarcinoma tissue. Human papillary carcinoma of thyroid gland and transitional cell carcinoma of kidney tissue. ICC/IF: MCF7, HT-29 and A431 cells. Flow Cyt (intra): A431 and MCF7 cells. WB: MCF-7, HT-29, HepG2 and PC-3 whole cell lysate. mIHC: Human endometrium tissue.
特記事項	<p>ab256580 is the carrier-free version of <a href="#">ab40772</a>.</p> <p>Our <b>carrier-free</b> antibodies are typically supplied in a PBS-only formulation, purified and free of BSA, sodium azide and glycerol. The carrier-free buffer and high concentration allow for increased conjugation efficiency.</p> <p>This conjugation-ready format is designed for use with fluorochromes, metal isotopes, oligonucleotides, and enzymes, which makes them ideal for antibody labelling, functional and cell-based assays, flow-based assays (e.g. mass cytometry) and Multiplex Imaging applications.</p> <p>Use our <b>conjugation kits</b> for antibody conjugates that are ready-to-use in as little as 20 minutes with &lt;1 minute hands-on-time and 100% antibody recovery: available for fluorescent dyes, HRP, biotin and gold.</p> <p>This product is compatible with the Maxpar<sup>®</sup> Antibody Labeling Kit from Fluidigm, without the need for antibody preparation. Maxpar<sup>®</sup> is a trademark of Fluidigm Canada Inc.</p> <p>This product is a recombinant monoclonal antibody, which offers several advantages including:</p> <ul style="list-style-type: none"><li>- High batch-to-batch consistency and reproducibility</li><li>- Improved sensitivity and specificity</li><li>- Long-term security of supply</li><li>- Animal-free production</li></ul> <p>For more information <a href="#">see here</a>.</p> <p>Our RabMAb<sup>®</sup> technology is a patented hybridoma-based technology for making rabbit</p>

monoclonal antibodies. For details on our patents, please refer to [RabMAb® patents](#).

Mouse, Rat: We have preliminary internal testing data to indicate this antibody may not react with these species. Please contact us for more information.

## 製品の特性

製品の状態	Liquid
保存方法	Shipped at 4°C. Store at +4°C. Do Not Freeze.
解離定数 (K <sub>D</sub> 値)	K <sub>D</sub> = 2.80 x 10 <sup>-11</sup> M



バッファー	pH: 7.20 Constituent: PBS
キャリア・フリー	はい
精製度	Protein A purified
ポリ/モノ	モノクローナル
クローン名	EP700Y
アイソタイプ	IgG

## アプリケーション

**The Abpromise guarantee**      **Abpromise保証は、次のテスト済みアプリケーションにおけるab256580の使用に適用されます**  
アプリケーションノートには、推奨の開始希釈率がありますが、適切な希釈率につきましてはご検討ください。

アプリケーション	Abreviews	特記事項
Flow Cyt (Intra)		Use at an assay dependent concentration.
WB		Use at an assay dependent concentration. Detects a band of approximately 80-120 kDa (predicted molecular weight: 97 kDa).
IHC-P		Use at an assay dependent concentration.
ICC/IF		Use at an assay dependent concentration.
mlHC		1/3000.

## ターゲット情報

**機能**      Cadherins are calcium-dependent cell adhesion proteins. They preferentially interact with themselves in a homophilic manner in connecting cells; cadherins may thus contribute to the

sorting of heterogeneous cell types. CDH1 is involved in mechanisms regulating cell-cell adhesions, mobility and proliferation of epithelial cells. Has a potent invasive suppressor role. It is a ligand for integrin alpha-E/beta-7.

E-Cad/CTF2 promotes non-amyloidogenic degradation of Abeta precursors. Has a strong inhibitory effect on APP C99 and C83 production.

#### 組織特異性

Non-neural epithelial tissues.

#### 関連疾患

Defects in CDH1 are the cause of hereditary diffuse gastric cancer (HDGC) [MIM:137215]. An autosomal dominant cancer predisposition syndrome with increased susceptibility to diffuse gastric cancer. Diffuse gastric cancer is a malignant disease characterized by poorly differentiated infiltrating lesions resulting in thickening of the stomach. Malignant tumors start in the stomach, can spread to the esophagus or the small intestine, and can extend through the stomach wall to nearby lymph nodes and organs. It also can metastasize to other parts of the body. Note=Heterozygous germline mutations CDH1 are responsible for familial cases of diffuse gastric cancer. Somatic mutations in the has also been found in patients with sporadic diffuse gastric cancer and lobular breast cancer.

Defects in CDH1 are a cause of susceptibility to endometrial cancer (ENDMC) [MIM:608089].

Defects in CDH1 are a cause of susceptibility to ovarian cancer (OC) [MIM:167000]. Ovarian cancer common malignancy originating from ovarian tissue. Although many histologic types of ovarian neoplasms have been described, epithelial ovarian carcinoma is the most common form. Ovarian cancers are often asymptomatic and the recognized signs and symptoms, even of late-stage disease, are vague. Consequently, most patients are diagnosed with advanced disease.

#### 配列類似性

Contains 5 cadherin domains.

#### 翻訳後修飾

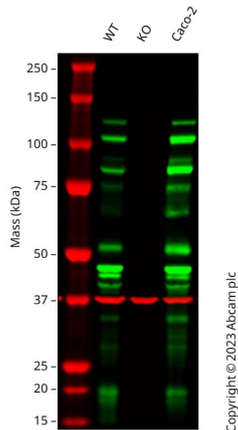
During apoptosis or with calcium influx, cleaved by a membrane-bound metalloproteinase (ADAM10), PS1/gamma-secretase and caspase-3 to produce fragments of about 38 kDa (E-CAD/CTF1), 33 kDa (E-CAD/CTF2) and 29 kDa (E-CAD/CTF3), respectively. Processing by the metalloproteinase, induced by calcium influx, causes disruption of cell-cell adhesion and the subsequent release of beta-catenin into the cytoplasm. The residual membrane-tethered cleavage product is rapidly degraded via an intracellular proteolytic pathway. Cleavage by caspase-3 releases the cytoplasmic tail resulting in disintegration of the actin microfilament system. The gamma-secretase-mediated cleavage promotes disassembly of adherens junctions.

#### 細胞内局在

Cell junction. Cell membrane. Endosome. Golgi apparatus > trans-Golgi network. Colocalizes with DLGAP5 at sites of cell-cell contact in intestinal epithelial cells. Anchored to actin microfilaments through association with alpha-, beta- and gamma-catenin. Sequential proteolysis induced by apoptosis or calcium influx, results in translocation from sites of cell-cell contact to the cytoplasm. Colocalizes with RAB11A endosomes during its transport from the Golgi apparatus to the plasma membrane.

#### 画像

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Western blot - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

**All lanes :** Anti-E Cadherin antibody [EP700Y] - Intercellular Junction Marker ([ab40772](#)) at 1/1000 dilution

**Lane 1 :** Wild-type A431 cell lysate

**Lane 2 :** CDH1 knockout A431 cell lysate

**Lane 3 :** Caco-2 cell lysate

Lysates/proteins at 20 µg per lane.

Performed under reducing conditions.

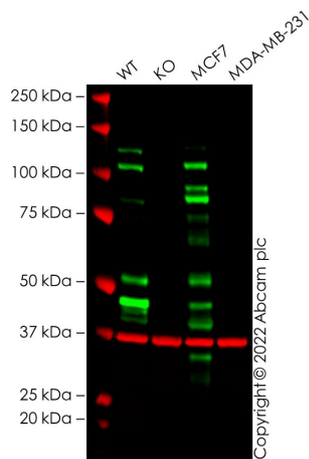
**Predicted band size:** 97 kDa

**Observed band size:** 110,130,40,55,80 kDa

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).

Western blot: Anti-CDH1 antibody [EP700Y] ([ab40772](#)) staining at 1/1000 dilution, shown in green; Mouse anti-GAPDH antibody [6C5] ([ab8245](#)) loading control staining at 1/20000 dilution, shown in red.

In Western blot, [ab40772](#) was shown to bind specifically to CDH1. A band was observed at 130, 110, 80, 55, 40 kDa in wild-type A431 cell lysates with no signal observed at this size in CDH1 knockout cell line [ab273747](#) (knockout cell lysate [ab273781](#)). To generate this image, wild-type and CDH1 knockout A431 cell lysates were analysed. First, samples were run on an SDS-PAGE gel then transferred onto a nitrocellulose membrane. Membranes were blocked in 3 % milk in TBS-0.1 % Tween<sup>®</sup> 20 (TBS-T) before incubation with primary antibodies overnight at 4 °C. Blots were washed four times in TBS-T, incubated with secondary antibodies for 1 h at room temperature, washed again four times then imaged. Secondary antibodies used were Goat anti-Rabbit IgG H&L 800CW and Goat anti-Mouse IgG H&L 680RD at 1/20000 dilution.



Western blot - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

**All lanes** : Anti-E Cadherin antibody [EP700Y] - Intercellular Junction Marker (**ab40772**) at 1/10000 dilution

**Lane 1** : Wild-type Raji cell lysate

**Lane 2** : CDH1 knockout Raji cell lysate

**Lane 3** : MCF7 cell lysate

**Lane 4** : MDA-MB-231 cell lysate

Lysates/proteins at 20 µg per lane.

### Secondary

**All lanes** : Goat anti-Rabbit IgG H&L 800CW and Goat anti-Mouse IgG H&L 680RD at 20000 µg

Performed under reducing conditions.

**Predicted band size:** 97 kDa

**Observed band size:** 105,130 kDa

This data was developed using the same antibody clone in a different buffer formulation (**ab40772**).

False colour image of Western blot: Anti-E Cadherin antibody [EP700Y] - Intercellular Junction Marker staining at 1/10000 dilution, shown in green; Mouse anti-GAPDH antibody [6C5] (**ab8245**) loading control staining at 1/20000 dilution, shown in red. In Western blot, **ab40772** was shown to bind specifically to E Cadherin. A band was observed at 105/130 kDa in wild-type Raji cell lysates with no signal observed at this size in CDH1 knockout cell line **ab273747** (knockout cell lysate **ab273781**). To generate this image, wild-type and CDH1 knockout Raji cell lysates were analysed. First, samples were run on an SDS-PAGE gel then transferred onto a nitrocellulose membrane. Membranes were blocked in 3 % milk in TBS-0.1 % Tween® 20 (TBS-T) before incubation with primary antibodies overnight at 4 °C. Blots were washed four times in TBS-T, incubated with secondary antibodies for 1 h at room temperature, washed again four times then imaged. Secondary antibodies used were Goat anti-Rabbit IgG H&L 800CW and Goat anti-Mouse IgG H&L 680RD at 1/20000 dilution.

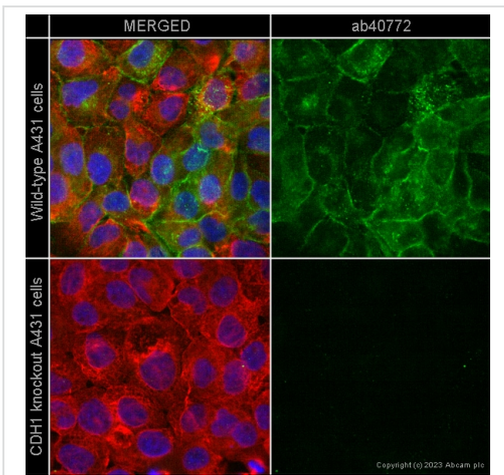
Tissue Microarray (TMA) data for ab40772			
Normal tissue samples		Malignant tissue samples	
Human cardiac muscle	✗	Human placenta	✓
Human cerebrum	✗	Human skeletal muscle	✗
Human colon	✓	Human skin	✓
Human endometrium	✓	Human spleen	✗
Human kidney	✓	Human stomach	✓
Human liver	✓	Human testis	✗
Human lung	✓	Human thyroid	✓
Human mammary gland	✓	Human tonsil * (epithelial cells*)	✗
Human pancreas	✓		
		Clear cell carcinoma of human kidney	✓
		Human bladder cancer	✓
		Human breast carcinoma	✓
		Human cervical carcinoma	✓
		Human colon carcinoma	✓
		Human endometrial carcinoma	✓
		Human gastric adenocarcinoma	✓
		Human glioma	✗
		Human hepatocellular carcinoma	✓
		Human lung carcinoma	✓
		Human ovarian carcinoma	✓
		Human pancreatic carcinoma	✓
		Human prostatic hyperplasia	✓
		Human thyroid carcinoma	✓

Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).

Tissue Microarrays stained for Anti-E Cadherin antibody [EP700Y] - Intercellular Junction Marker using **ab40772** in immunohistochemical analysis. This table provides a detailed overview of positive (tick mark) and negative (cross mark) staining per sample type tested. The section was incubated with **ab40772** for 30 mins at room temperature followed by a ready to use Rabbit specific IHC polymer detection kit HRP/DAB (**ab209101**). The immunostaining was performed on a Leica Biosystems BOND® RX instrument.

Heat mediated antigen retrieval was performed with Tris-EDTA buffer (pH 9.0, Epitope Retrieval Solution2) for 20 mins.

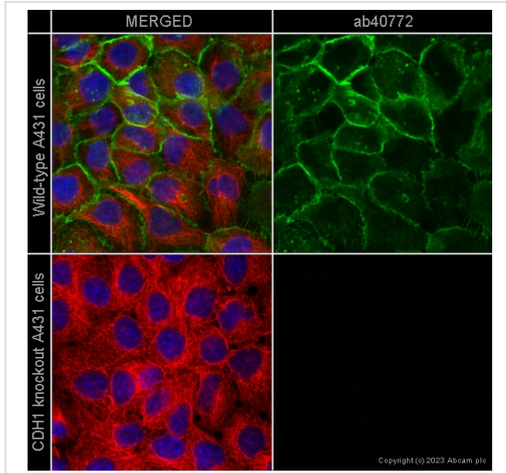


Immunocytochemistry/ Immunofluorescence - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Immunofluorescence staining of E-Cadherin using **ab40772** in wild-type A431 cells (top panel) and CDH1 knockout A431 cells (bottom panel). The cells were fixed with 4% formaldehyde (10 min), permeabilized with 0.1% Triton-X-100 for 5 minutes and then blocked with 1% BSA/10% normal goat serum/0.3M glycine in 0.1% PBS-Tween for 1h. The cells were then incubated with **ab40772** at 1 µg/mL and **ab7291** at 1 µg/mL overnight at +4°C, followed by a further incubation at room temperature for 1h with Goat Anti-Rabbit IgG H&L (Alexa Fluor® 488) preadsorbed (**ab150081**) (shown in green) and Goat Anti-Mouse IgG H&L (Alexa Fluor® 594) preadsorbed (**ab150120**) (shown in red), both at 1/1000. Nuclear DNA was labelled with DAPI (shown in blue).

Image was acquired with a high-content analyser (Operetta CLS, Perkin Elmer) and a confocal section is shown.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).

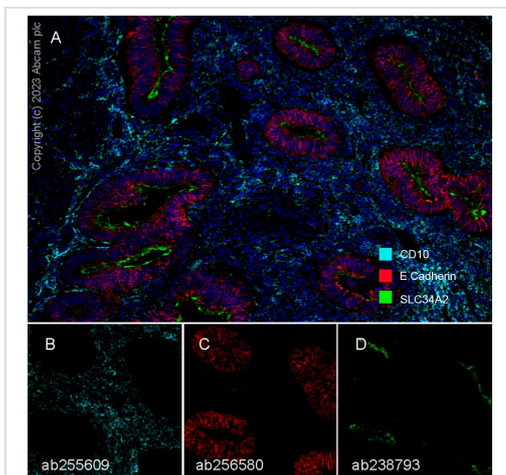


Immunocytochemistry/ Immunofluorescence - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Immunofluorescence staining of E-Cadherin using **ab40772** in wild-type A431 cells (top panel) and CDH1 knockout A431 cells (bottom panel). The cells were fixed with 100% methanol (5 min), permeabilized with 0.1% Triton-X-100 for 5 minutes and then blocked with 1% BSA/10% normal goat serum/0.3M glycine in 0.1% PBS-Tween for 1h. The cells were then incubated with **ab40772** at 0.2 µg/mL and **ab7291** at 1 µg/mL overnight at +4°C, followed by a further incubation at room temperature for 1h with Goat Anti-Rabbit IgG H&L (Alexa Fluor® 488) preadsorbed (**ab150081**) (shown in green) and Goat Anti-Mouse IgG H&L (Alexa Fluor® 594) preadsorbed (**ab150120**) (shown in red), both at 1/1000. Nuclear DNA was labelled with DAPI (shown in blue).

Image was acquired with a high-content analyser (Operetta CLS, Perkin Elmer) and a confocal section is shown.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).



Multiplex immunohistochemistry - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Fluorescence multiplex immunohistochemical analysis of the human endometrium (Formalin/PFA-fixed paraffin-embedded sections).

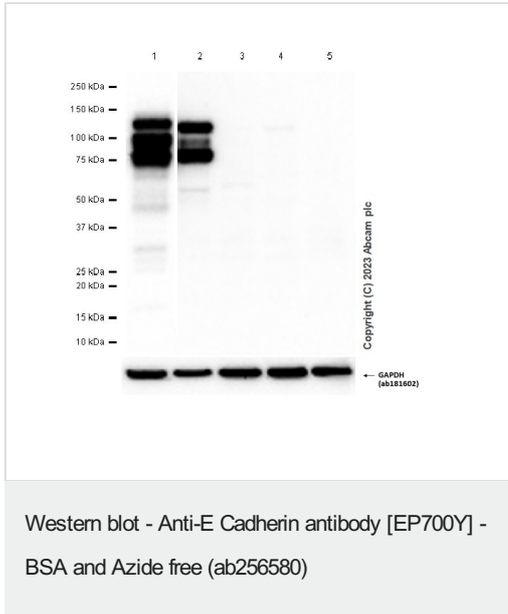
Panel A: merged staining of anti-E Cadherin (**ab256580**, red; Opal™690), anti-SLC34A2 (**ab238793**, green; Opal™520) and anti-CD10 (**ab255609**, cyan; Opal™570) on human endometrium.

Panel B: anti-CD10 stained on stromal cells. Panel C: anti-E Cadherin stained on glandular cells. Panel D: anti-SLC34A2 stained on apical membrane of glandular cells. Opal Polymer HRP Ms + Rb was used as a secondary antibody.

The section was incubated in three rounds of staining: in the order of **ab256580** at 1/3000 dilution (0.324 µg/ml) for 30mins, **ab238793** at 1/1000 dilution (2.26 µg/ml) for 10mins and **ab255609** at 1/1000 dilution (0.615 µg/ml) for 30 mins at room temperature. Each round was followed by a separate fluorescent tyramide signal amplification system.

The immunostaining was performed on a Leica Biosystems BOND® RX instrument with an Opal™ 4-color kit. Image acquisition was performed with Leica SP8 confocal microscope.

Heat mediated antigen retrieval with Tris-EDTA buffer (pH 9.0, epitope retrieval solution2) for 20 mins. DAPI (blue) was used as a nuclear counter stain.



**All lanes** : Anti-E Cadherin antibody [EP700Y] - Intercellular Junction Marker ([ab40772](#)) at 1/1000 dilution

**Lane 1** : MCF7 (Human breast adenocarcinoma epithelial cell) whole cell lysate

**Lane 2** : HepG2 (Human hepatocellular carcinoma epithelial cell) whole cell lysate

**Lane 3** : A375 (Human malignant melanoma epithelial cell) whole cell lysate

**Lane 4** : HeLa (Human cervix adenocarcinoma epithelial cell) whole cell lysate

**Lane 5** : HT-1080 (Human fibrosarcoma epithelial cell) whole cell lysate

Lysates/proteins at 20 µg per lane.

### Secondary

**All lanes** : Goat Anti-Rabbit IgG H&L (HRP) ([ab97051](#)) at 1/20000 dilution

**Predicted band size:** 97 kDa

**Observed band size:** 80-125 kDa

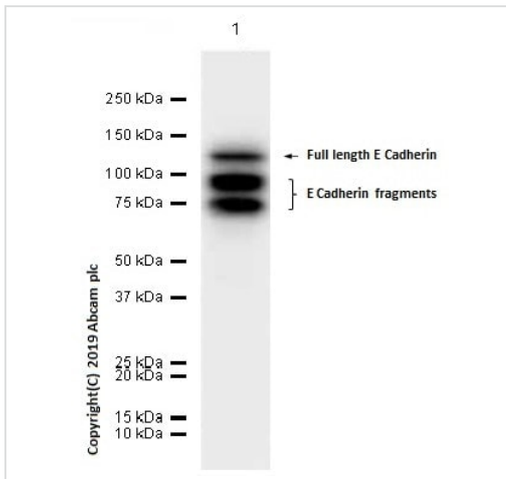
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).

Blocking and diluting buffer and concentration: 5% NFDm/TBST.

[ab181602](#) was as GAPDH loading control.

Exposure time: Lane1: 3 seconds; Lane 2-5: 40 seconds.

A375, HeLa and HT-1080 were reported as negative or express low level of E cadherin (PMID: 30393081, PMID: 16980628, PMID: 34715746), PMID: 25411788).



Western blot - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Anti-E Cadherin antibody [EP700Y] - Intercellular Junction Marker ([ab40772](#)) at 1/1000 dilution + MCF7 (Human breast adenocarcinoma epithelial cell) whole cell lysate at 20 µg

### Secondary

Goat Anti-Rabbit IgG H&L (HRP) ([ab97051](#)) at 1/20000 dilution

**Predicted band size:** 97 kDa

**Observed band size:** 80-125 kDa

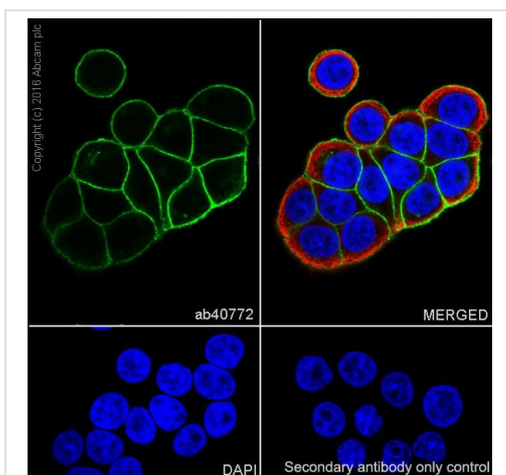
**Exposure time:** 3.25 seconds.

**Blocking and diluting buffer:** 5% NFDM/TBST.

Full-length E Cadherin has a molecular weight of approximately 125 kDa. Other molecular weights between 80-100 kDa could also be observed depending on cell types or cell conditions.

PMID: 27274359, PMID: 26983597, PMID: 18478055, PMID: 22375065.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).

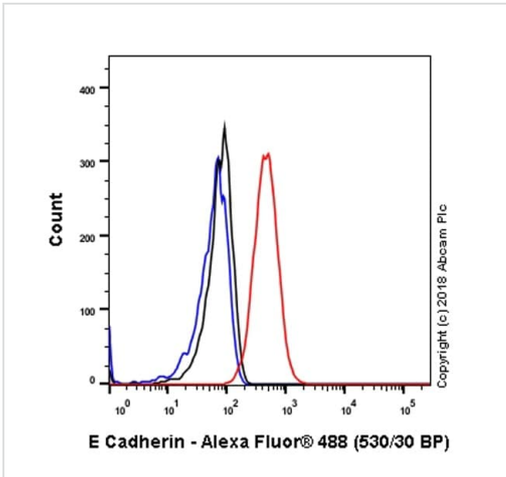


Immunocytochemistry/ Immunofluorescence - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

[ab40772](#) staining E Cadherin in HT-29 (Human colorectal adenocarcinoma) cells by ICC/IF

(Immunocytochemistry/Immunofluorescence). Cells were fixed with 4% paraformaldehyde and permeabilized with 0.1% tritonX-100. Samples were incubated with primary antibody at 1/500 dilution. An Alexa Fluor® 488 Goat anti-Rabbit ([ab150077](#)) was used as the secondary antibody at 1/1000 dilution. Anti-alpha Tubulin antibody [DM1A] - Microtubule Marker (Alexa Fluor® 594) ([ab195889](#)) at 1/200 dilution was used as a counterstain. DAPI was used as a nuclear counterstain. This is a confocal image showing membranous staining on HT-29 cell line.

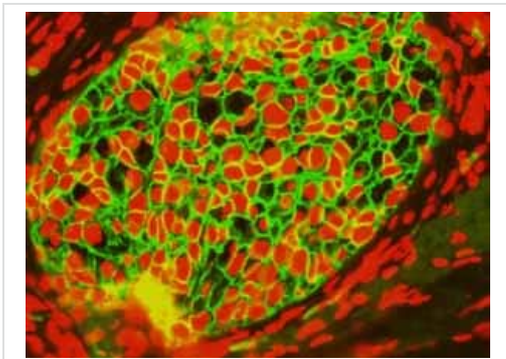
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).



Flow Cytometry (Intracellular) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Intracellular Flow Cytometry analysis of MCF7 (Human breast adenocarcinoma epithelial cell) cells labeling E Cadherin with purified **ab40772** at 1/30 dilution (10µg/ml) (red). Cells were fixed with 4% Paraformaldehyde. A Goat anti rabbit IgG (Alexa Fluor® 488) secondary antibody was used at 1/2000 dilution. Isotype control - Rabbit monoclonal IgG (Black). Unlabeled control - Cell without incubation with primary antibody and secondary antibody (Blue).

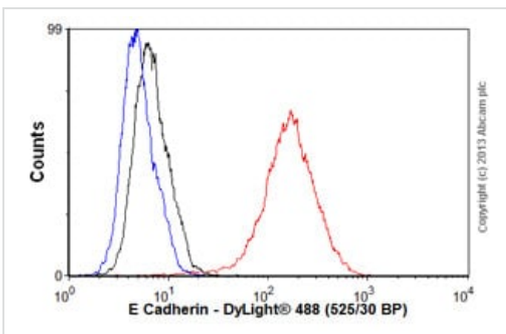
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

This IHC data was generated using the same anti-E Cadherin antibody clone, EP700Y, in a different buffer formulation (**ab40772**).

Fluorescent immunohistochemical analysis of paraffin-embedded human breast carcinoma tissue using **ab40772**. Green-E-Cadherin red-PI.

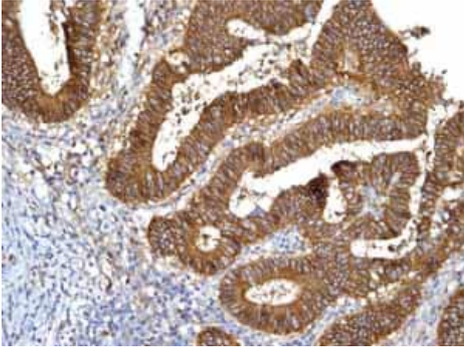


Flow Cytometry (Intracellular) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Overlay histogram showing A431 (Human epidermoid carcinoma cell line) cells stained with unpurified **ab40772** (red line). The cells were fixed with 80% methanol (5 minutes) and incubated in 1x PBS / 10% normal goat serum / 0.3M glycine to block non-specific protein-protein interactions. The cells were then incubated with the antibody (**ab40772**, 1/1000 dilution) for 30 minute at 22°C. The secondary antibody used was DyLight® 488 goat anti-rabbit IgG (H+L) (**ab96899**) at 1/500 dilution for 30 minutes at 22°C. Isotype control antibody (black line) was rabbit IgG (monoclonal) (0.1 µg/1x10<sup>6</sup> cells) used under the same conditions. Unlabeled sample (blue line) was also used as a control. Acquisition of >5,000 events were collected using a 20mW Argon ion laser (488nm) and 525/30 bandpass filter.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and

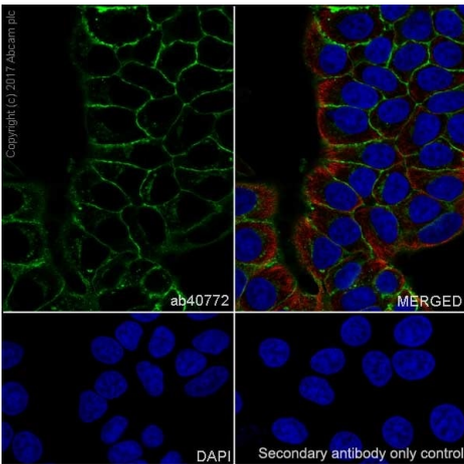
sodium azide ([ab40772](#)).



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free ([ab256580](#))

Formalin/PFA-fixed paraffin-embedded human colonic adenocarcinoma tissue stained for E Cadherin with unpurified [ab40772](#) at a 1/500 dilution in immunohistochemical analysis.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).

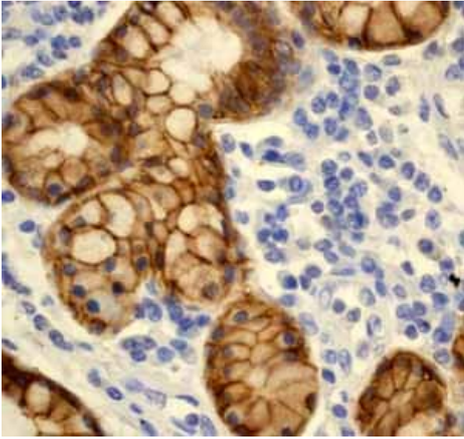


Immunocytochemistry/ Immunofluorescence - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free ([ab256580](#))

Immunocytochemistry/Immunofluorescence analysis of MCF7 (human breast adenocarcinoma epithelial) cells labeling E Cadherin with [ab40772](#). Cells were fixed with 4% paraformaldehyde and permeabilized with 0.1% tritonX-100. Samples were then incubated with the primary antibody at a 1/500 dilution, followed by Goat anti-rabbit IgG (Alexa Fluor<sup>®</sup> 488) ([ab150077](#)) secondary antibody at a 1/1000 dilution (green). The nuclear counter stain is DAPI (blue). Counterstained with [ab195889](#) anti-alpha Tubulin antibody [DM1A] - Microtubule Marker (Alexa Fluor<sup>®</sup> 594) at a 1/200 dilution (red).

Confocal image shows membranous staining on MCF7 cell line.

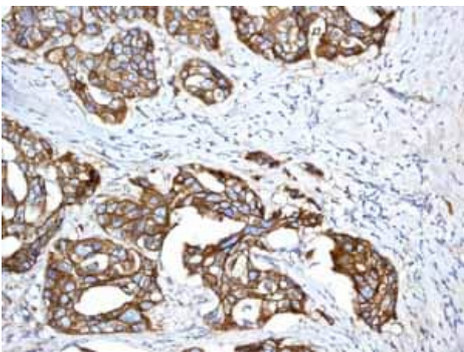
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Formalin-fixed, paraffin-embedded human lung adenocarcinoma tissue stained for E Cadherin with unpurified [ab40772](#) at a 1/500 dilution in immunohistochemical analysis.

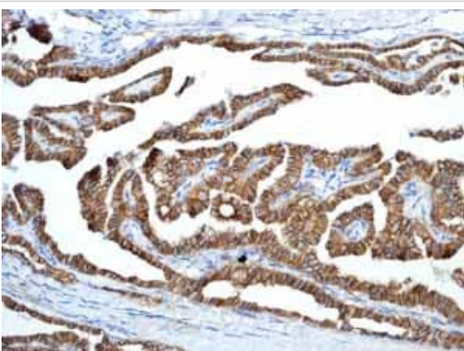
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Formalin-fixed, paraffin-embedded human breast carcinoma tissue stained for E Cadherin with unpurified [ab40772](#) at a 1/500 dilution in immunohistochemical analysis.

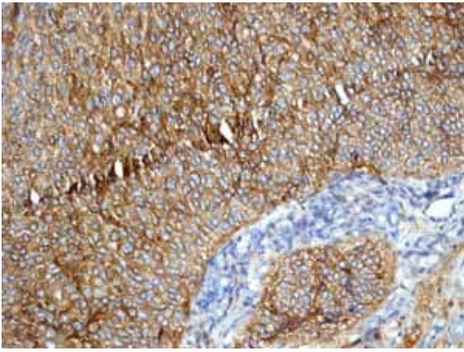
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Formalin-fixed, paraffin-embedded human papillary carcinoma of thyroid gland tissue stained for E Cadherin with unpurified [ab40772](#) at a 1/500 dilution in immunohistochemical analysis.

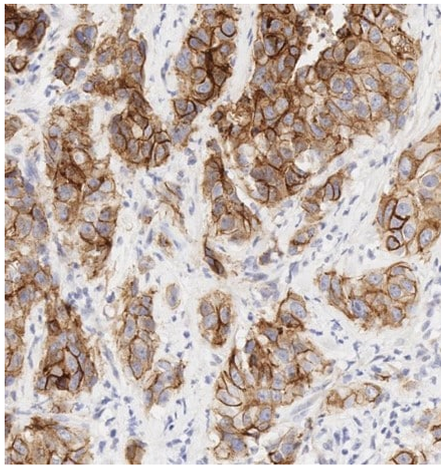
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide ([ab40772](#)).



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Formalin-fixed, paraffin-embedded human transitional cell carcinoma of kidney tissue stained for E Cadherin with unpurified **ab40772** at a 1/500 dilution in immunohistochemical analysis.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).

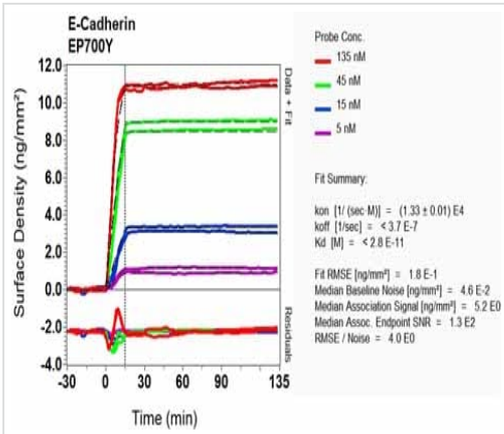


Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections) - Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

Immunohistochemistry of breast carcinoma staining E Cadherin with **ab40772** at 1 µg/ml.

Perform heat mediated antigen retrieval with citrate buffer pH 6 before commencing with IHC staining protocol.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).



OI-RD Scanning - Anti-E Cadherin antibody  
 [EP700Y] - BSA and Azide free (ab256580)

Produced using unpurified **ab40772**

Equilibrium disassociation constant (KD)

Learn more about KD

**[Click here to learn more about KD](#)**

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab40772**).

### Why choose a recombinant antibody?

 <b>Research with confidence</b> Consistent and reproducible results	 <b>Long-term and scalable supply</b> Recombinant technology
 <b>Success from the first experiment</b> Confirmed specificity	 <b>Ethical standards compliant</b> Animal-free production

Anti-E Cadherin antibody [EP700Y] - BSA and Azide free (ab256580)

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