

CF150 Antibody (Center)
Affinity Purified Rabbit Polyclonal Antibody (Pab)
Catalog # AP10510c

Specification

CF150 Antibody (Center) - Product Information

Application	WB, FC,E
Primary Accession	O8N884
Other Accession	NP_612450.2
Reactivity	Human
Host	Rabbit
Clonality	Polyclonal
Isotype	Rabbit IgG
Antigen Region	266-295

CF150 Antibody (Center) - Additional Information

Gene ID 115004

Other Names

Cyclic GMP-AMP synthase, cGAMP synthase, cGAS, h-cGAS, Mab-21 domain-containing protein 1, MB21D1, C6orf150

Target/Specificity

This CF150 antibody is generated from rabbits immunized with a KLH conjugated synthetic peptide between 266-295 amino acids from the Central region of human CF150.

Dilution

WB~~1:1000

FC~~1:25

Format

Purified polyclonal antibody supplied in PBS with 0.09% (W/V) sodium azide. This antibody is purified through a protein A column, followed by peptide affinity purification.

Storage

Maintain refrigerated at 2-8°C for up to 2 weeks. For long term storage store at -20°C in small aliquots to prevent freeze-thaw cycles.

Precautions

CF150 Antibody (Center) is for research use only and not for use in diagnostic or therapeutic procedures.

CF150 Antibody (Center) - Protein Information

Name CGAS {ECO:0000303|PubMed:23258413, ECO:0000312|HGNC:HGNC:21367}

Function Nucleotidyltransferase that catalyzes the formation of cyclic GMP-AMP (2',3'-cGAMP)

from ATP and GTP and plays a key role in innate immunity (PubMed:[21478870](#), PubMed:[23258413](#), PubMed:[23707061](#), PubMed:[23707065](#), PubMed:[23722159](#), PubMed:[24077100](#), PubMed:[24116191](#), PubMed:[24462292](#), PubMed:[25131990](#), PubMed:[26300263](#), PubMed:[29976794](#), PubMed:[30799039](#), PubMed:[31142647](#), PubMed:[32814054](#), PubMed:[33273464](#), PubMed:[33542149](#), PubMed:[37217469](#), PubMed:[37802025](#)). Catalysis involves both the formation of a 2',5' phosphodiester linkage at the GpA step and the formation of a 3',5' phosphodiester linkage at the ApG step, producing c[G(2',5')pA(3',5')p] (PubMed:[28214358](#), PubMed:[28363908](#)). Acts as a key DNA sensor: directly binds double-stranded DNA (dsDNA), inducing the formation of liquid-like droplets in which CGAS is activated, leading to synthesis of 2',3'-cGAMP, a second messenger that binds to and activates STING1, thereby triggering type-I interferon production (PubMed:[28314590](#), PubMed:[28363908](#), PubMed:[29976794](#), PubMed:[32817552](#), PubMed:[33230297](#), PubMed:[33606975](#), PubMed:[35322803](#), PubMed:[35438208](#), PubMed:[35460603](#), PubMed:[35503863](#)). Preferentially recognizes and binds curved long dsDNAs of a minimal length of 40 bp (PubMed:[30007416](#)). Acts as a key foreign DNA sensor, the presence of double-stranded DNA (dsDNA) in the cytoplasm being a danger signal that triggers the immune responses (PubMed:[28363908](#)). Has antiviral activity by sensing the presence of dsDNA from DNA viruses in the cytoplasm (PubMed:[28363908](#), PubMed:[35613581](#)). Also acts as an innate immune sensor of infection by retroviruses, such as HIV-2, by detecting the presence of reverse-transcribed DNA in the cytosol (PubMed:[23929945](#), PubMed:[24269171](#), PubMed:[30270045](#), PubMed:[32852081](#)). In contrast, HIV-1 is poorly sensed by CGAS, due to its capsid that cloaks viral DNA from CGAS detection (PubMed:[24269171](#), PubMed:[30270045](#), PubMed:[32852081](#)). Detection of retroviral reverse-transcribed DNA in the cytosol may be indirect and be mediated via interaction with PQBP1, which directly binds reverse-transcribed retroviral DNA (PubMed:[26046437](#)). Also detects the presence of DNA from bacteria, such as M.tuberculosis (PubMed:[26048138](#)). 2',3'-cGAMP can be transferred from producing cells to neighboring cells through gap junctions, leading to promote STING1 activation and convey immune response to connecting cells (PubMed:[24077100](#)). 2',3'-cGAMP can also be transferred between cells by virtue of packaging within viral particles contributing to IFN- induction in newly infected cells in a cGAS-independent but STING1- dependent manner (PubMed:[26229115](#)). Also senses the presence of neutrophil extracellular traps (NETs) that are translocated to the cytosol following phagocytosis, leading to synthesis of 2',3'-cGAMP (PubMed:[33688080](#)). In addition to foreign DNA, can also be activated by endogenous nuclear or mitochondrial DNA (PubMed:[28738408](#), PubMed:[28759889](#), PubMed:[31299200](#), PubMed:[33031745](#), PubMed:[33230297](#)). When self-DNA leaks into the cytosol during cellular stress (such as mitochondrial stress, SARS-CoV-2 infection causing severe COVID-19 disease, DNA damage, mitotic arrest or senescence), or is present in form of cytosolic micronuclei, CGAS is activated leading to a state of sterile inflammation (PubMed:[28738408](#), PubMed:[28759889](#), PubMed:[31299200](#), PubMed:[33031745](#), PubMed:[33230297](#), PubMed:[35045565](#)). Acts as a regulator of cellular senescence by binding to cytosolic chromatin fragments that are present in senescent cells, leading to trigger type-I interferon production via STING1 and promote cellular senescence (By similarity). Also involved in the inflammatory response to genome instability and double-stranded DNA breaks: acts by localizing to micronuclei arising from genome instability (PubMed:[28738408](#), PubMed:[28759889](#)). Micronuclei, which are frequently found in cancer cells, consist of chromatin surrounded by their own nuclear membrane: following breakdown of the micronuclear envelope, a process associated with chromothripsis, CGAS binds self-DNA exposed to the cytosol, leading to 2',3'-cGAMP synthesis and subsequent activation of STING1 and type-I interferon production (PubMed:[28738408](#), PubMed:[28759889](#)). Activated in response to prolonged mitotic arrest, promoting mitotic cell death (PubMed:[31299200](#)). In a healthy cell, CGAS is however kept inactive even in cellular events that directly expose it to self-DNA, such as mitosis, when cGAS associates with chromatin directly after nuclear envelope breakdown or remains in the form of postmitotic persistent nuclear cGAS pools bound to chromatin (PubMed:[31299200](#), PubMed:[33542149](#)). Nuclear CGAS is inactivated by chromatin via direct interaction with nucleosomes, which block CGAS from DNA binding and thus prevent CGAS-induced autoimmunity (PubMed:[31299200](#), PubMed:[32911482](#), PubMed:[32912999](#), PubMed:[33051594](#), PubMed:[33542149](#)). Also acts as a suppressor of DNA repair in response to DNA damage: inhibits homologous recombination repair by interacting with PARP1, the CGAS-PARP1 interaction leading to impede the formation of the PARP1-TIMELESS complex (PubMed:[30356214](#), PubMed:[31544964](#)). In addition to DNA, also sense translation stress: in

response to translation stress, translocates to the cytosol and associates with collided ribosomes, promoting its activation and triggering type-I interferon production (PubMed:[34111399](#)). In contrast to other mammals, human CGAS displays species-specific mechanisms of DNA recognition and produces less 2',3'-cGAMP, allowing a more fine-tuned response to pathogens (PubMed:[30007416](#)).

Cellular Location

Nucleus. Chromosome. Cell membrane; Peripheral membrane protein. Cytoplasm, cytosol. Note=Mainly localizes in the nucleus, and at low level in the cytosol (PubMed:31544964, PubMed:31808743). On chromosomes, enriched on centromeric satellite and LINE DNA repeat elements (PubMed:30811988). Exported from the nucleus to the cytosol in a XPO1/CRM1 via the nuclear export signal in response to DNA stimulation (PubMed:33406424). Outside the nucleus, localizes at the cell membrane as a peripheral membrane protein in resting conditions: association to the cell membrane is mediated via binding to phosphatidylinositol 4,5-bisphosphate (PtdIns(4,5)P2) (PubMed:30827685). Localization at the cell membrane is required to limit the recognition of self-DNA (PubMed:30827685). Following detection of double-stranded DNA (dsDNA), released from the cell membrane into the cytosol in order to signal (PubMed:30827685). Upon transfection with dsDNA forms punctate structures that co-localize with DNA and Beclin-1 (BECN1) (PubMed:26048138). Phosphorylation at Tyr-215 promotes cytosolic retention (PubMed:30356214). In response to translation stress, translocates to the cytosol and associates with collided ribosomes (PubMed:34111399).

Tissue Location

Expressed in the monocytic cell line THP1.

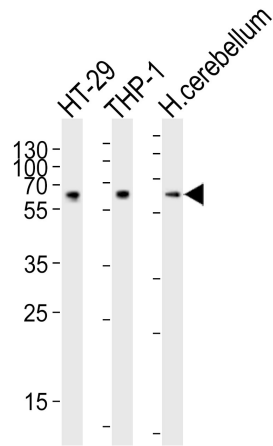
CF150 Antibody (Center) - Protocols

Provided below are standard protocols that you may find useful for product applications.

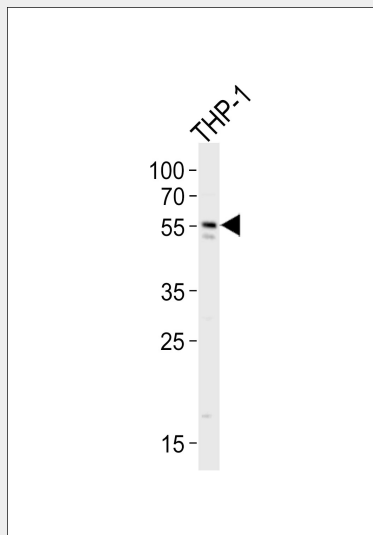
- [Western Blot](#)
- [Blocking Peptides](#)
- [Dot Blot](#)
- [Immunohistochemistry](#)
- [Immunofluorescence](#)
- [Immunoprecipitation](#)
- [Flow Cytometry](#)
- [Cell Culture](#)

CF150 Antibody (Center) - Images

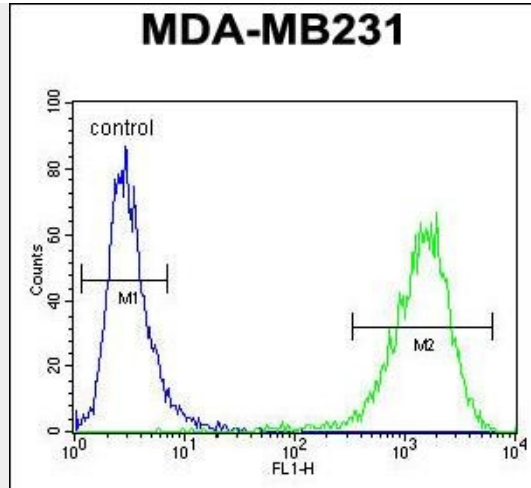




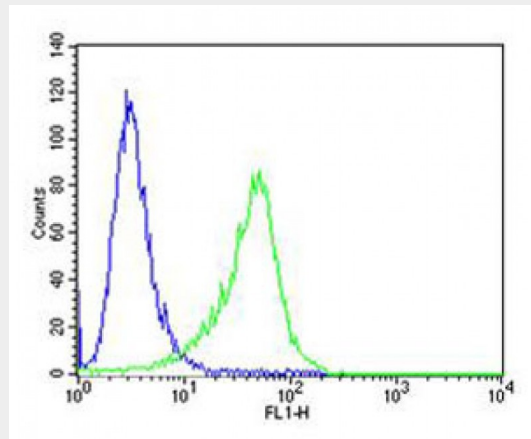
Western blot analysis of lysates from HT-29, THP-1 cell line, human cerebellum tissue lysate (from left to right), using CF150 Antibody (Center)(Cat. #AP10510c). AP10510c was diluted at 1:1000 at each lane. A goat anti-rabbit IgG H&L(HRP) at 1:10000 dilution was used as the secondary antibody. Lysates at 20ug per lane.



Western blot analysis of lysate from THP-1 cell line, using CF150 Antibody (Center)(Cat. #AP10510c). AP10510c was diluted at 1:1000. A goat anti-rabbit IgG H&L(HRP) at 1:10000 dilution was used as the secondary antibody. Lysate at 20ug.



CF150 Antibody (Center) (Cat. #AP10510c) flow cytometric analysis of MDA-MB231 cells (right histogram) compared to a negative control cell (left histogram). FITC-conjugated goat-anti-rabbit secondary antibodies were used for the analysis.



Flow cytometric analysis of A549 cells using CF150 Antibody (Center) (green, Cat#AP10510c) compared to an isotype control of rabbit IgG (blue). AP10510c was diluted at 1:25 dilution. An Alexa Fluor® 488 goat anti-rabbit IgG at 1:400 dilution was used as the secondary antibody.

CF150 Antibody (Center) - Background

The exact function of C6orf150 remains unknown.

CF150 Antibody (Center) - References

Rush, J., et al. Nat. Biotechnol. 23(1):94-101(2005)
 Rush, J., et al. Nat. Biotechnol. 23(1):94-101(2005)
 Mungall, A.J., et al. Nature 425(6960):805-811(2003)

CF150 Antibody (Center) - Citations

- [Cellular sensing of extracellular purine nucleosides triggers an innate IFN-β response](#)
- [cGAS-STING Signaling Regulates Initial Innate Control of Cytomegalovirus Infection.](#)
- [The DNA Sensor, Cyclic GMP-AMP Synthase, Is Essential for Induction of IFN-β during Chlamydia trachomatis Infection.](#)