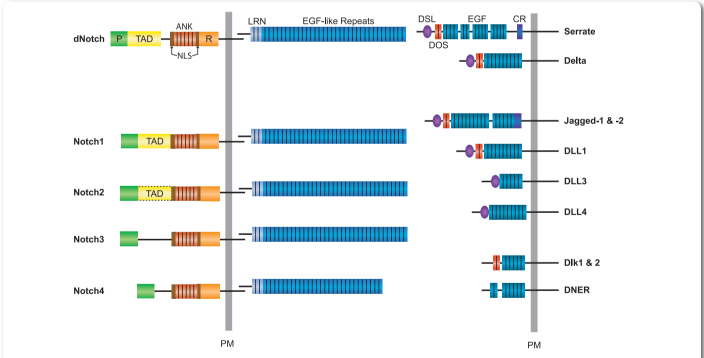


# **Notch Ligands**

The highly conserved Notch signaling pathway regulates many different cell fate decisions in both vertebrate and invertebrate species [1-4]. It is important for pattern formation during development such as neurogenesis, angiogenesis or myogenesis and regulates T cell development and stem cell maintenance [5-7]. But Notch signaling is also involved in cellular processes throughout adulthood [8]. Signaling via Notch occurs between neighbouring cells and both the receptor and its ligands are transmembrane proteins (see Figure 1).

Notch ligands are single-pass transmembrane proteins with a DSL (Delta, Serrate, LAG-2)-domain and varying numbers of EGF-like repeats. There are two classes of canonical Notch ligands, the Delta/Delta-like and the Serrate/Jagged class. The later has an additional domain of cysteine rich repeats close to the transmembrane domain. There are 5 canonical Notch ligands in mammals: Jagged-1, Jagged-2, DLL1, DLL3 and DLL4. These can bind to the four Notch receptors Notch 1-4. In contrast to mammals, *Drosophila* has only two Notch ligands - Delta and Serrate - that can activate the single Notch receptor [9-11].

#### **CONTINUED ON NEXT PAGE**



#### FIGURE 1: Notch Receptors and Their Ligands.

Drosophila contains one Notch receptor (dNotch) that is bound by two transmembrane DSL-ligands (Delta and Serrate). Mammals possess four Notch receptors (Notch1–4) and five ligands (Jagged-1 and -2, which are homologous to Serrate, and Delta-like (DLL) 1, 3 and 4, which are homologous to Delta). Additional noncanonical Notch ligands are Dlk1, Dlk2 and DNER.

ANK: ankyrin repeats, CR: cysteine-rich domain, DOS: Delta and OSM-11-like proteins domain, DSL: Delta, Serrate and LAG-2 domain, EGF: epidermal growth factor-like repeats, LNR: cysteine-rich Lin12-Notch repeats, NLS: nuclear localization signal, P: PEST domain, PM: plasma membrane, R: RAM domain, TAD: transactivation domain. Adapted from: *The intracellular region of Notch ligands: does the tail make the difference?* A. Pintar, et al.; Biol. Direct **2**, 19 (2007), *The canonical Notch signaling pathway: unfolding the activation mechanism:* R. Kopan & M.X. Ilagan; Cell **137**, 216 (2009)





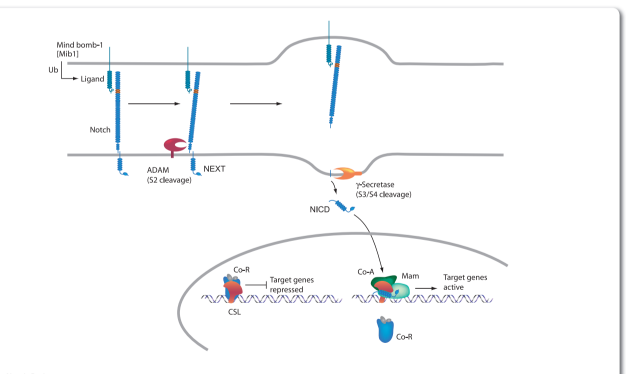
The noncanonical Notch ligands lack the DSL domain, among these are Dlk1, Dlk2 and DNER [12-15]. Other noncanonical ligands lack all typical Notch ligand domains and can have a completely different structure, some are not even membrane-tethered [16]. They are thought to act as co-ligands to enhance or inhibit Notch activation and might be important modulators of the Notch pathway [14, 16].

Endocytosis of Notch ligands in the signal-producing cells is absolutely required for the initiation of Notch signaling. Two structurally distinct E3 ubiquitin ligases, Neuralized (Neur) and Mind bomb (Mib), are known to regulate the endocytosis of Notch ligands in Drosophila and zebrafish, respectively [42, 43]. In mammals, 2 Neur homologs, Neur1 [44, 45] and Neur2 [46], and 2 Mib homologs, Mind bomb-1 (Mib1) [47] and Mib2 [48], have been identified. Although all four E3 ubiquitin ligases are known to induce the endocytosis of Notch ligands in vitro, only Mib1 has an obligatory role in the activation of Jag- as well as DLL-mediated Notch signaling in mammalian development, while Neur1, Neur2, and Mib2 are dispensable [49].

Upon ligand binding, the Notch receptor is cleaved proteolytically (see Figure 2). This cleavage occurs in a two stepprocess releasing the extracellular and the intracellular part of Notch from the membrane. First the extracellular domain with the bound ligand attached to it is released by the action of a ADAM family metalloprotease creating a membranebound intermediate called Notch extracellular truncation (NEXT) (S2 cleavage). In a second step, NEXT is cleaved within the transmembrane domain by y-secretase (S3/S4 cleavage), releasing the Notch intracellular domain (NICD). NICD translocates to the nucleus and with the help of additional enhancers and co-activators activates target genes [10, 17, 18].

Each Notch molecule signals only once without signal amplification by second messengers. This allows a rapid and highly responsive downregulation and reactivation of the signaling pathway. The activity of Notch and its ligands requires endocytosis and is regulated through glycosylation, ubiquitinylation and microRNAs by mechanisms not yet fully understood [19-27].

The Notch pathway plays an important role in many different processes in a wide range of tissues, this is why aberrations in Notch signaling components have been associated with various human disorders such as cancer, immune disorders, developmental syndromes, stroke and cognitive symptoms [10, 28-36]. A mutation in JAG1 can cause the Alagille syndrome [37-39] and other cognitive dysfunctions such as CADASIL or schizophrenia have been associated with mutations of the Notch receptors [40, 41].



#### FIGURE 2: The Notch Pathway.

Binding of the Notch ligand (green) on one cell to the Notch receptor (blue) on another cell results in two proteolytic cleavages of the receptor. First a protease of the ADAM family cleaves off most of the extracellular part of Notch leaving a Notch extracellular truncation (NEXT) (S2 cleavage). Second  $\gamma$ -secretase cleaves NEXT at positions S3 and S4 within the transmembrane domain releasing the Notch intracellular domain (NICD). NICD enters the nucleus and interacts with the DNA-binding CSL protein (red), Mastermind (Mam; green) and additional co-activators (Co-A; green). Co-repressors (Co-R; blue and grey) are released and target genes become active.

Adapted from: Notch signalling: a simple pathway becomes complex: S.J. Bray; Nat. Rev. Mol. Cell. Biol. 7, 678 (2006). The canonical Notch signaling pathway: unfolding the activation mechanism: R. Kopan & M.X. Ilagan; Cell 137, 216 (2009)

## DLL1 [Delta-like Protein 1; Delta1]

### DLL1 (human), (rec.)

ALX-201-382-C010	10 µg
ALX-201-382-C050	50 µg
Produced in HEK 293 cells. Signal peptide and extracellular	domain of recom-

binant human DLL1 (aa 1-545) is fused at the C-terminus to a FLAG®-tag.

## DLL1 (human):Fc (human), (rec.)

ALX-201-425-C010	
ALX-201-425-C050	

Produced in HEK 293 cells. Signal peptide and extracellular domain of recombinant human DLL1 (aa 1-545) is fused at the C-terminus to the Fc portion of human IgG. Interacts with human Notch1 (as confirmed by flow cytometry).

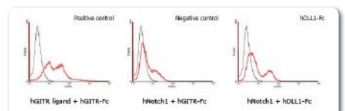


FIGURE: Interaction of human Notch1 with DLL1 (human):Fc (human), (rec.) (Prod. No. ALX-201-425).

**METHOD:** HEK 293 cells transfected with a human Notch 1 or a control vector were incubated with 25µg/ml of GITR (human):Fc (human), (rec.) or DLL1 (human):Fc (human), (rec.) (Prod. No. ALX-201-425). Cells were stained with human IgG (Fc specific) FITC conjugate for DLL1-Fc binding.

## DLL1 (mouse):Fc (human), (rec.)

ALX-201-455-C010	10 µg
ALX-201-455-C050	50 µg

Produced in HEK 293 cells. Signal peptide and extracellular domain of recombinant mouse DLL1 (aa 1-545) is fused at the C-terminus to the Fc portion of human IgG. Interacts with Notch receptors.

## DLL1 (human), mAb (D1L165-6)

ALX-804-758-C050	50 µg
ALX-804-758-C100	100 µg
CLONE: D1I 165-6, ISOTYPE: MOUSE IgG1, IMMUNOGEN:	Recombinant human

CLONE: D1L165-6. ISOTYPE: Mouse IgG1. IMMUNOGEN: Recombinant human DLL1. SPECIFICITY: Recognizes human DLL1. APPLICATION: WB.

## DLL1 (human), pAb

#### ALX-210-466-C100

From rabbit. IMMUNOGEN: Recombinant human DLL1. SPECIFICITY: Recognizes human DLL1. APPLICATION: IHC (PS), WB.

# DLL1 (human), pAb

ALX-210-483-C100

100 µg

100 ua

From rat. IMMUNOGEN: Recombinant human DLL1. SPECIFICITY: Recognizes human DLL1. APPLICATION: WB.

# DLL3 [Delta-like Protein 3; Delta3]

## DLL3 (human):Fc (human), (rec.)

ALX-201-422-C010	10 µg
ALX-201-422-C050	50 µg
Produced in HEK 293 cells. Signal peptide and extracellular domain	n of re-

combinant human DLL3 (aa 1-466) is fused at the C-terminus to the Fc portion of human IgG1.

# DLL4 [Delta-like Protein 4; Delta4]

## DLL4 (human):Fc (human), (rec.)

ALX-201-386-C010	10 µg
ALX-201-386-C050	50 µg

Produced in HEK 293 cells. Signal peptide and extracellular domain of recombinant human DLL4 (aa 1-529) is fused at the C-terminus to the Fc portion of human IgG. Interacts with human Notch1 (as confirmed by flow cytometry).

DLL4 (mouse):Fc (human), (rec.) NEW

ALX-201-453-C010

10 µg

50 µg

NEW

Produced in HEK 293 cells. Signal peptide and extracellular domain of recombinant mouse DLL4 (aa 1-532) is fused at the C-terminus to the Fc portion of human IgG.

DLL4 (human), mAb (DL86-3AG)	NEW
ALX-804-763-C050	50 µg

ALX-804-763-C1	00		100 µg
		Maura InOd	 Decembined by

CLONE: DL86-3AG. ISOTYPE: Mouse IgG1. IMMUNOGEN: Recombinant human DLL4 (ectodomain). SPECIFICITY: Recognizes human DLL4. APPLICA-TION: WB.

# DLL4 (human), pAb

ALX-210-484-C100

100 µg

10 µg

From rat. IMMUNOGEN: Recombinant human DLL4. SPECIFICITY: Recognizes human DLL4. APPLICATION: WB.

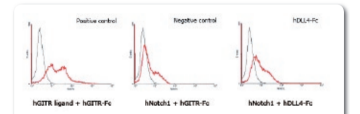


FIGURE: Interaction of human Notch1 with human DLL4 (Prod. No. ALX-204-386). **METHOD:** HEK 293 cells transfected with a human Notch1 or a control vector were incubated with 25µg/ml of GITR (human):Fc (human), (rec.) or DLL4 (human):Fc (human), (rec.) (Prod. No. ALX-204-386). Cells were stained with human IgG (Fc specific) FITC conjugate for DLL4-Fc binding.

# DLL1, Soluble (human) ELISA Kit

AG-45A-0027EK-KI01 AG-45A-0027TP-KI01 AG-45A-0027PP-KI01

Twin Plex Penta Plex 96 wells 2 x 96 wells 5 x 96 wells



For the quantitative determination of soluble DLL1 in human serum, plasma or cell culture supernatant. **SENSITIVITY**: 120pg/ml (range 0 to 16ng/ml).



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# Selected Latest Review Articles

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# Dlk1 [Delta-like Protein; Pref-1]

### Dlk1 (human), (rec.)

ALX-201-442-C010	10 µg
ALX-201-442-C050	50 µg
Produced in HEK 293 cells. The signal peptide and the extracellula	ır domain

of human Dlk1 (aa 1-303) is fused at the C-terminus to a FLAG®-tag.

## Dlk1 (human):Fc (human), (rec.)

ALX-201-427-C010	10 µg
ALX-201-427-C050	50 µg
Produced in HEK 293 cells The	signal pentide and the extracellular do-

main of human Dlk1 (aa 1-303) is fused at the C-terminus to the Fc portion of human lqG.

## Dlk1 (mouse):Fc (human), (rec.)

ALX-201-416-C010			10 µg
ALX-201-416-C050			50 µg

Produced in HEK 293 cells. The signal peptide and the extracellular domain of mouse Dlk1 (aa 1-305) is fused at the C-terminus to the Fc portion of human IgG.

## Dlk1 (human), mAb (PF13-3)

ALX-804-754-C050	50 µg
ALX-804-754-C100	100 µg

CLONE: PF13-3. ISOTYPE: Mouse IgG1. IMMUNOGEN: Recombinant human Dlk1 (extracellular domain). SPECIFICITY: Recognizes human Dlk1. AP-PLICATION: ELISA, FC, IHC, WB.

# Dlk1 (human), mAb (PF299-1

	 	_	 
ALX-804-755-C100			100 µg
ALX-804-755-C050			50 µg

CLONE: PF299-1. ISOTYPE: Mouse IgG1. IMMUNOGEN: Recombinant human Dlk1 (extracellular domain). SPECIFICITY: Recognizes human Dlk1. APPLI-CATION: ELISA, FC, IHC, WB.

# Dlk1 (mouse), mAb (PF105B)

	 -		
ALX-804-742-C100			100 µg
ALX-804-742-C050			50 µg

CLONE: PF105B. ISOTYPE: Rat IgG2. IMMUNOGEN: Recombinant mouse DIk1 (extracellular domain). SPECIFICITY: Recognizes mouse Dlk1. APPLICATION: ELISA (1:2'000-1:10'000), WB.

## Dlk1 (mouse), mAb (PF183E)

ALX-804-743-C050	50 µg
ALX-804-743-C100	100 µg

CLONE: PF183E. ISOTYPE: Rat IgG2. IMMUNOGEN: Recombinant mouse DIk1 (extracellular domain). SPECIFICITY: Recognizes mouse Dlk1. APPLICATION: WB.

#### Dlk1 (human), pAb

NEW

ALX-210-495-C100

100 µg

From rabbit. IMMUNOGEN: Recombinant human Dlk1 (extracellular domain). SPECIFICITY: Recognizes human Dlk1. Weakly cross-reacts with mouse DIk1. APPLICATION: WB.

SPECIFICITY: Recognizes human Dlk1. APPLICATION: FC, WB.

## Dlk1 (human), pAb

ALX-210-496-C100

100 µg From rat. IMMUNOGEN: Recombinant human Dlk1 (extracellular domain).

# Latest Insight

# Notch Ligands Inhibit Adipocyte Differentiation

Not only Dlk1/Pref-1 [Y. Wang & H.S. Sul; Cell Metab. 9, 287 (2009)] but also the human Notch ligands DLL1, DLL4, DNER have been shown to inhibit adipocytes differentiation (adipogenesis) [unpublished data], therefore being interesting tools for stem cell research.

### **Experimental: Adipogenesis Inhibition**

3T3L1 cells were maintained in DMEM supplemented with 10% FBS and penicillin-streptomycin. When the cells were confluently grown, adipogenesis was initiated by adding IBMX, dexamethasone, and insulin to 0.5mM, 1µM, and 10µg/ml, respectively and continued for 2 days (day 0). The medium was replaced every 2 days with new medium containing insulin in the presence or absence of 5µg/ml of each human recombinant Notch ligand-Fc fusion protein (human DLL1-Fc, human DNER-Fc, or human Dlk1/Pref-1) and human CD137-Fc as a control-Fc. Staining with Oil Red O was typically performed on day 7.











IBMX+Desam cone+Inculin +TNF-alpha

FIGURE 1: Adipogenesis inhibition of 3T3L1 cells by DLL1 (human):Fc (human), (rec.) (Prod. No. ALX-201-425).

+hCD137-Fc ( Fc control)

shasone+Insulin



DBMX+Dexamethasone+Insulin +TNF-alpha

FIGURE 2: Adipogenesis inhibition of 3T3L1 cells by DNER (extracellular domain) (human):Fc (human), (rec.) (Prod. No. ALX-201-428). IBMX+Dexamethasone+Insulin +hCD137-Fc (Fc control)

Purified (PF) = Purified (Preservative free); FC = Flow Cytometry; ICC = Immunocytochemistry; IP = Immunoprecipitation; IHC = Immunohistochemistry (FS = Frozen Sections, PS = Paraffin Sections); WB = Western blot; BP = Blocking Peptide

## Jagged-1 (human):Fc (human), (rec.)

ALX-201-390-C010	10 µg
ALX-201-390-C050	50 µg

Produced in HEK 293 cells. Signal peptide and extracellular domain of human Jagged-1 (HJ1: CD339) (aa 1-1067) are fused at the C-terminus to the Fc portion of human lgG.

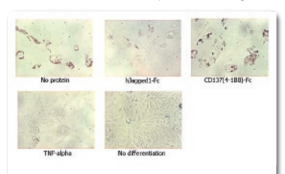


FIGURE: Differentiation of human mesenchymal stem cells (MSCs) into adipocytes in the presence or absence of Notch ligands that inhibit adipogenic differentiation of MSCs.

# Jagged-1 (human), mAb (J1G74-7)

ALX-804-735-C100

100 ua

100 µg

10 µg

50 µg

CLONE: J1G74-7. ISOTYPE: Mouse IgG1. IMMUNOGEN: Recombinant human Jagged-1. SPECIFICITY: Recognizes human Jagged-1. APPLICATION: ELISA, FC, WB.

### Jagged-1 (human), pAb

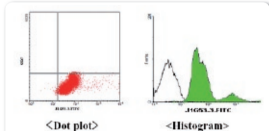
ALX-210-485-C100

From rat. IMMUNOGEN: Recombinant human Jagged-1 (ECD). SPECIFICITY: Recognizes human Jagged-1. APPLICATION: ELI-SA, WB.

## Jagged-1 (human), mAb (J1G53-3)

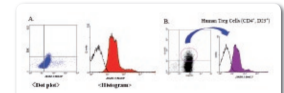
ALX-804-734-C100		100 µg
ALX-804-734PC-C050	PerCP	50 µg
ALX-804-734F-C050	FITC	50 µg
CLONE: .11G53-3 ISOTYPE:	Mouse IaG1	IMMUNOGEN: Be-

combinant human Jagged-1. SPECIFICITY: Recognizes human Jagged-1. APPLICATION: ELISA, FC, WB.



<Dot plot>

FIGURE: Flow Cytometry. Surface staining method was used to stain normal human peripheral blood cells (CD4+ selections) with Jagged-1 (human), mAb (J1G53-3). An appropriate isotype control was used for FITC mouse IgG1.



#### FIGURE: Flow Cytometry.

A: Surface staining method was used to stain normal human peripheral blood cells (CD4+ selections) with Jagged-1 (human), mAb (J1G53-3) (PerCP). An appropriate isotype control was used for PerCP mouse laG1.

B: Human CD4+ (selected via DYNAL113.03) cells were surfacestained with cocktail of CD4-FITC and CD25-PE subsequently with 20µl/million cells Jagged-1 (human), mAb (J1G53-3) (PerCP) or PerCP mouse IgG1 isotype control using the human regulatory T cell staining set. The dot blot on the left demonstrates co-staining of CD4 and CD25, while the histogram on the right demonstrates J1G35-3 staining of gated CD4+ CD25+ lymphocytes (Regulatory T cells / Treg) versus isotype control. Cells in the lymphocyte gate were used for flow cytometric analysis.

## DNER

### DNER (extracellular domain) (human), (rec.)

Produced in HEK 293 cells. The signal peptide and the extracellular domain of human DNER (aa 1-640) are fused at the C-terminus with a FLAG®-tag.

## DNER (extracellular domain) (human):Fc (human), (rec.)

ALX-201-428-C010	10 µg
ALX-201-428-C050	50 µg

Produced in HEK 293 cells. The signal peptide and the extracellular domain of human DNER (aa 1-637) is fused at the C-terminus to the Fc portion of human IgG.

## DNER (human), mAb (DR324-4)

ALX-804-761-C050

ALX-804-761-C100

CLONE: DR324-4. ISOTYPE: Mouse IgG2. IMMUNOGEN: Recombinant human DNER (ectodomain). SPECIFICITY: Recognizes human DNER. APPLICATION: WB.

### DNER (human), pAb

ALX-213-004-C100

From rabbit. IMMUNOGEN: Recombinant human DNER (ectodomain). SPECIFICITY: Recognizes human DNER. APPLICA-TION: WB.



#### Switzerland & Rest of Europe ENZO LIFE SCIENCES AG

Industriestrasse 17. Postfach CH-4415 Lausen / Switzerland Tel. + 41/0 61 926 89 89 Fax + 41/0 61 926 89 79 info-ch@enzolifesciences.com

#### North/South America

ENZO LIFE SCIENCES INTERNATIONAL, INC. 5120 Butler Pike

Plymouth Meeting, PA 19462-1202 / USA Tel. 1-800-942-0430/(610) 941-0430 Fax (610) 941-9252 info-usa@enzolifesciences.com

#### Benelux

ENZO LIFE SCIENCES BVBA Melkerijweg 3 BE-2240 Zandhoven / Belgium Tel. +32/0 3 466 04 20

Fax +32/0 3 466 04 29 info-be@enzolifesciences.com

#### France

ENZO LIFE SCIENCES FRANCE c/o Covalabis a s

13, avenue Albert Einstein, 69100 Villeurbanne / France Tel. +33/0 472 440 655 Fax +33/0 437 484 239 info-fr@ enzolifesciences.com

#### Germany

ENZO LIFE SCIENCES GmbH Marie-Curie-Strasse 8 DE-79539 Lörrach / Germany Tel. +49/0 7621 5500 526 Toll Free: 0800 6649518 Fax +49/0 7621 5500 527 info-de@enzolifesciences.com

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