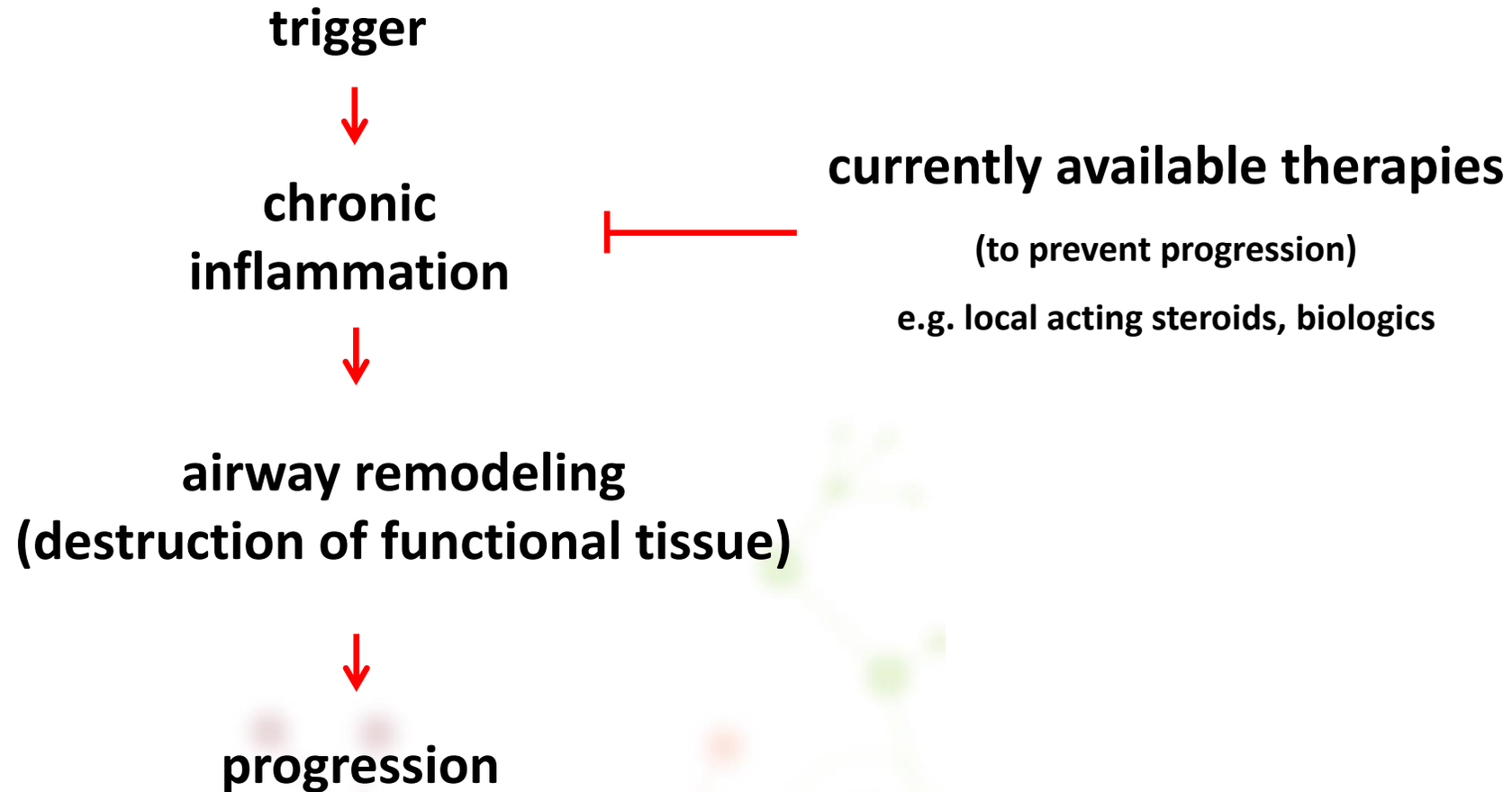


Manipulation of the immune response

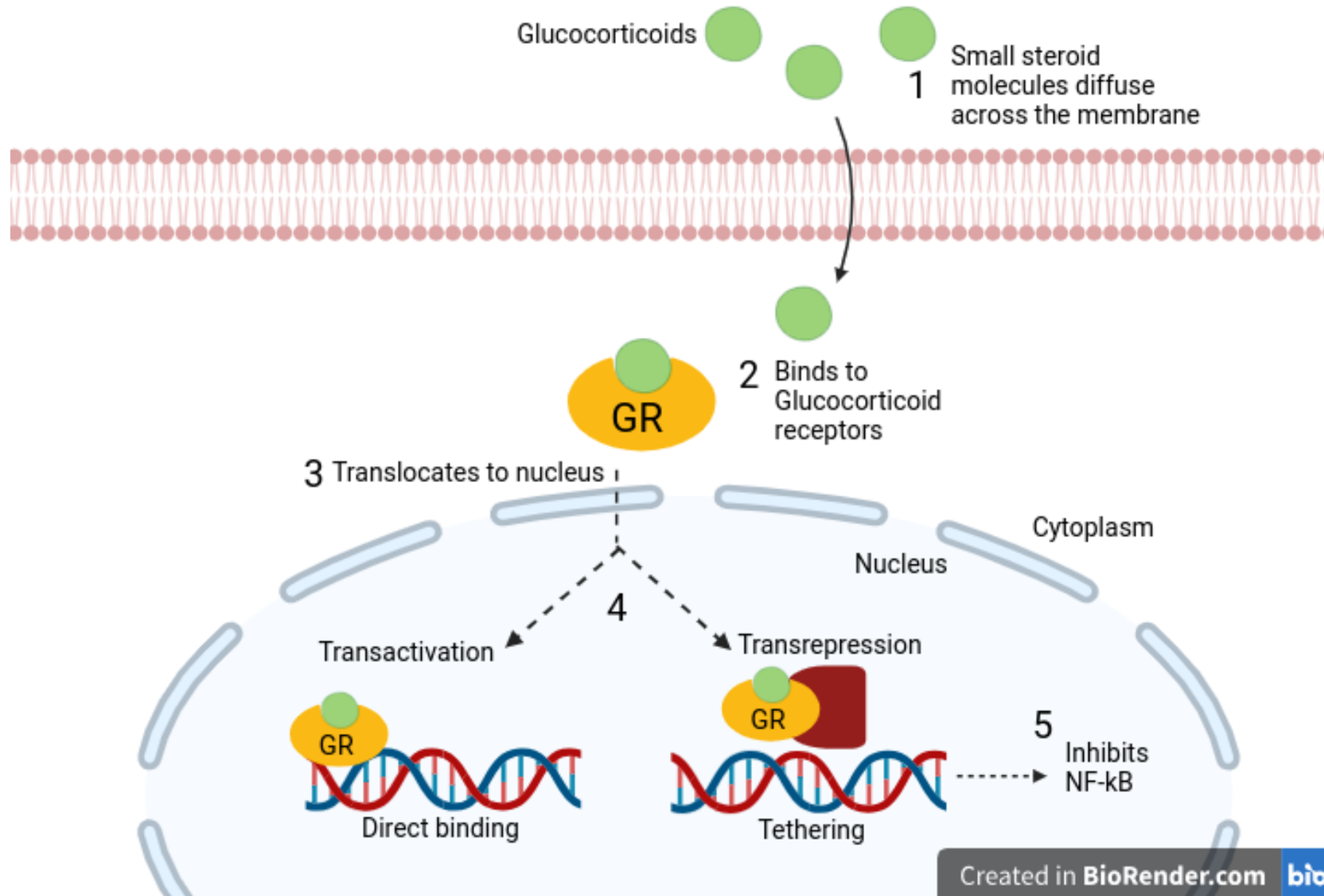
PD Dr. Jürgen Knobloch

Chronic inflammatory diseases of the upper and lower airways

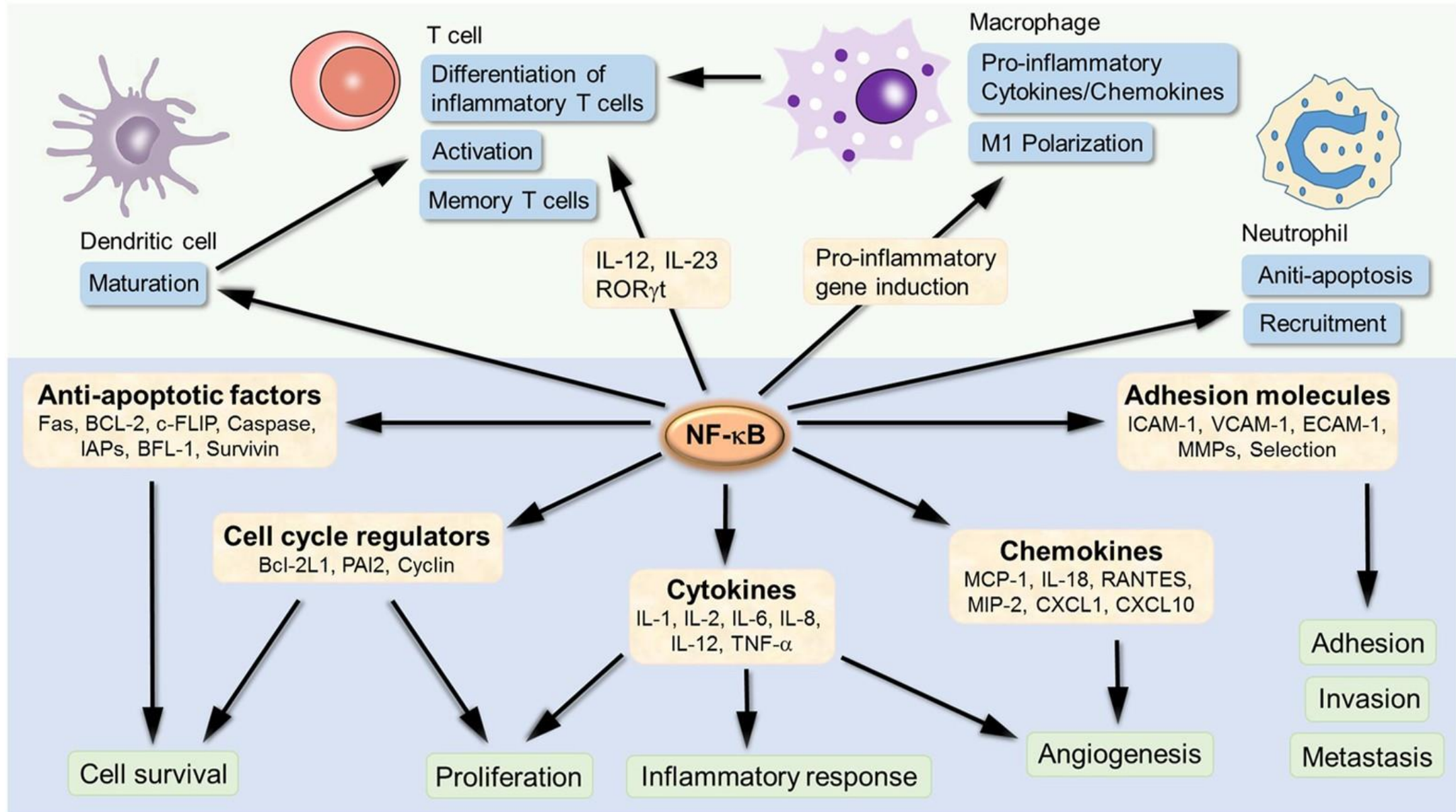
Asthma, COPD, CRS, and others



Steroids/NF- κ B blockade: mechanism of action

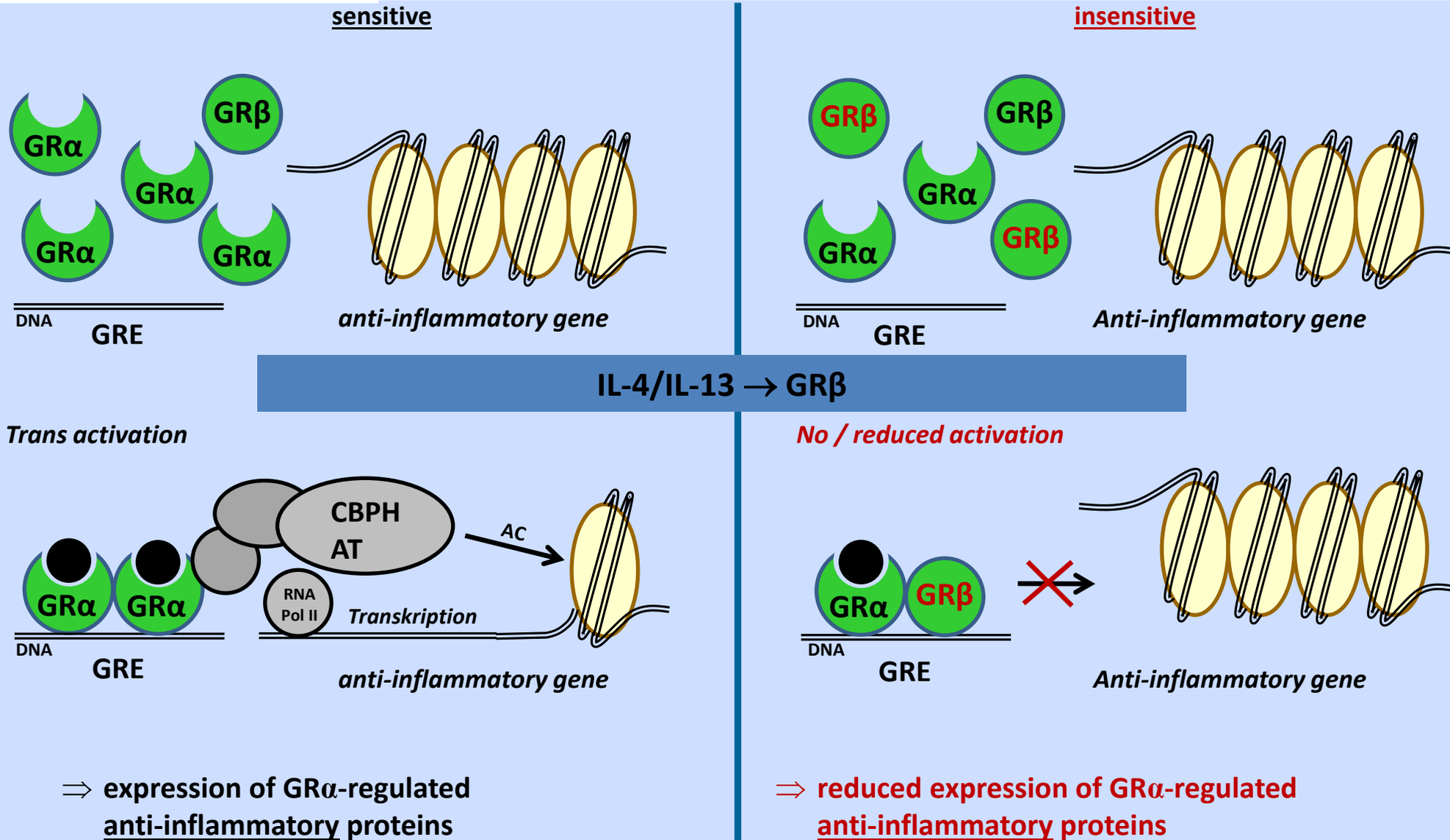


Steroids/NF-κB blockade: mechanism of action



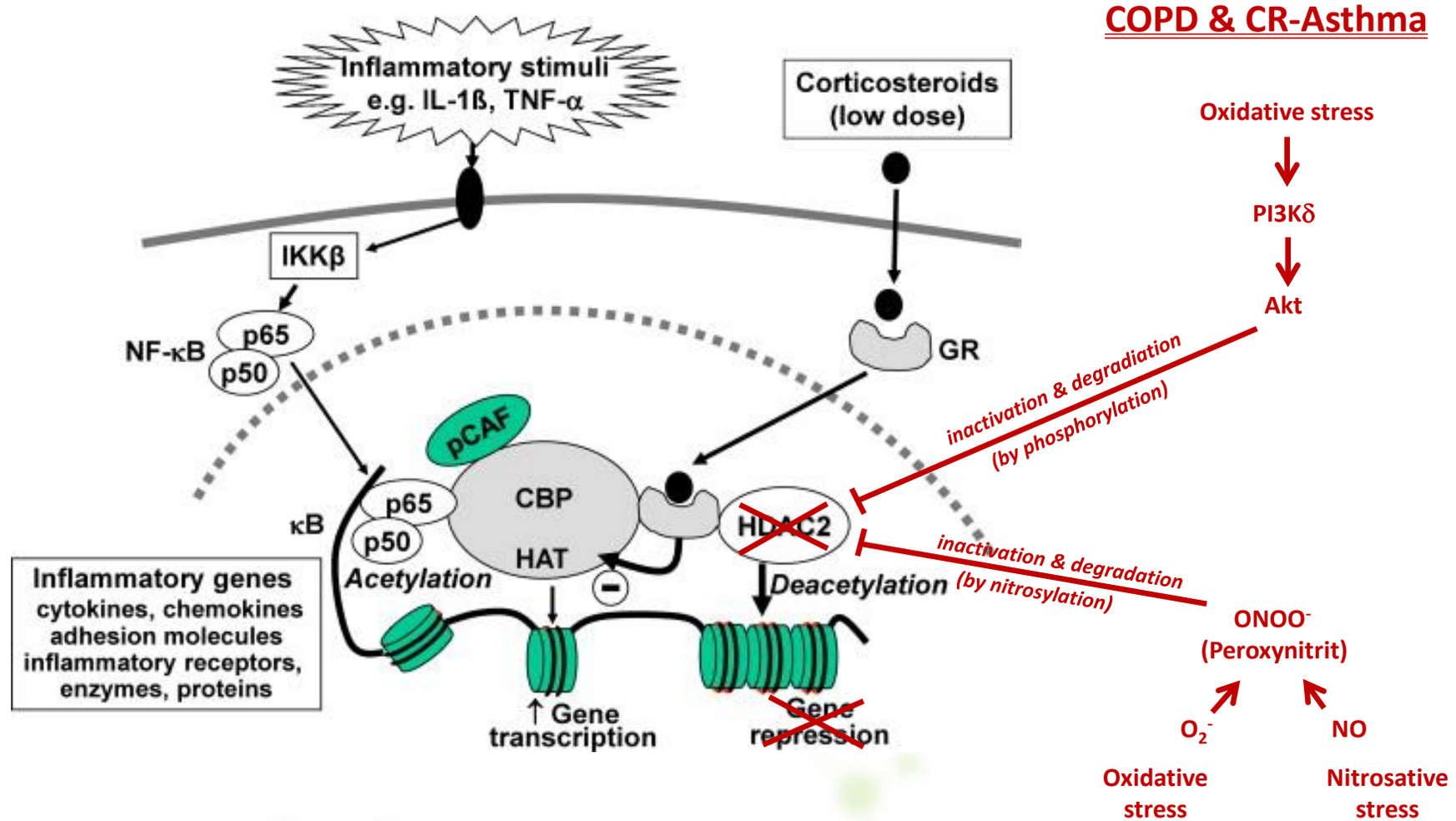
Mechanisms of steroid resistance

(in neutrophilic granulocytes)



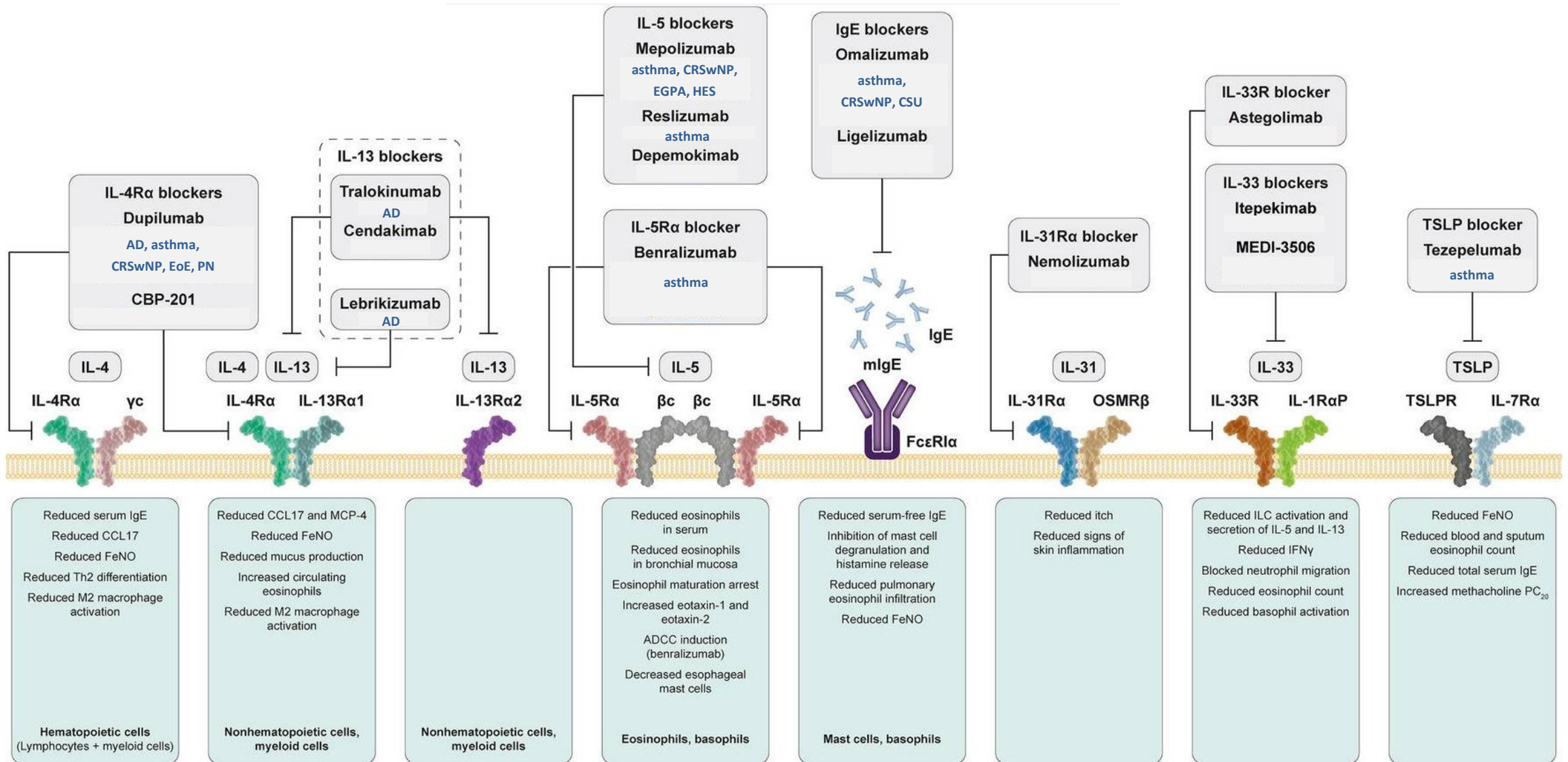
Mechanisms of steroid resistance: HDAC2 reduction

(in macrophages)

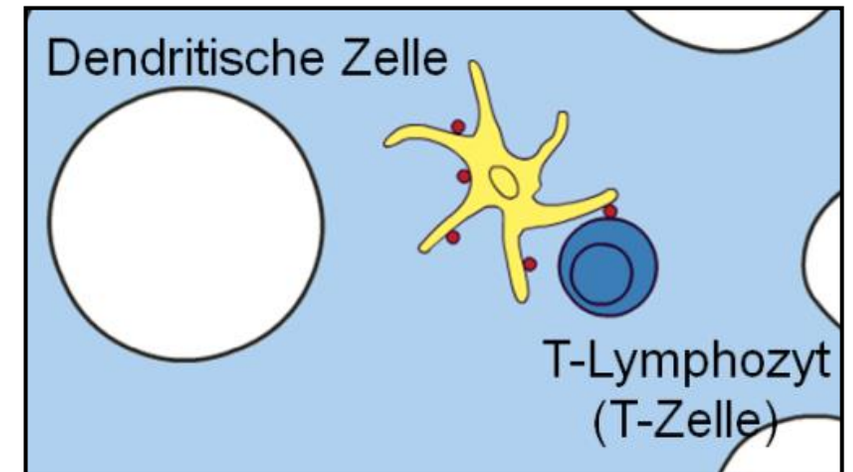
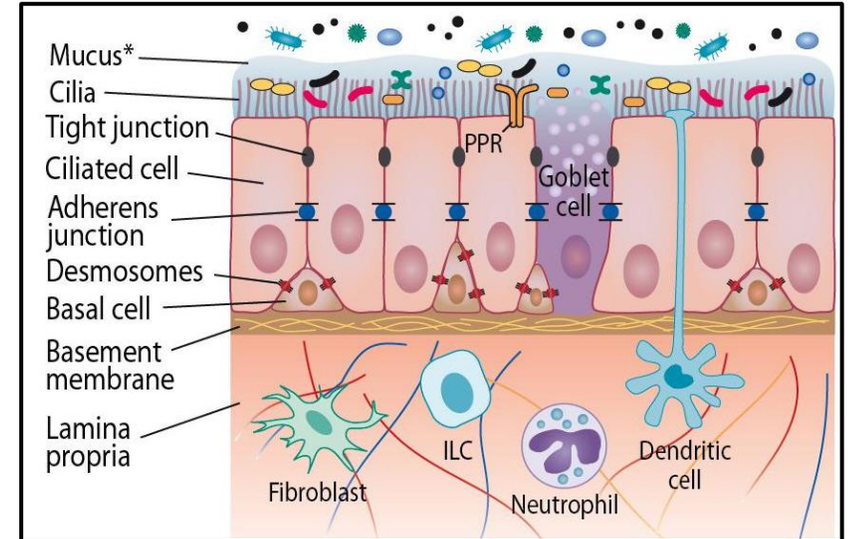
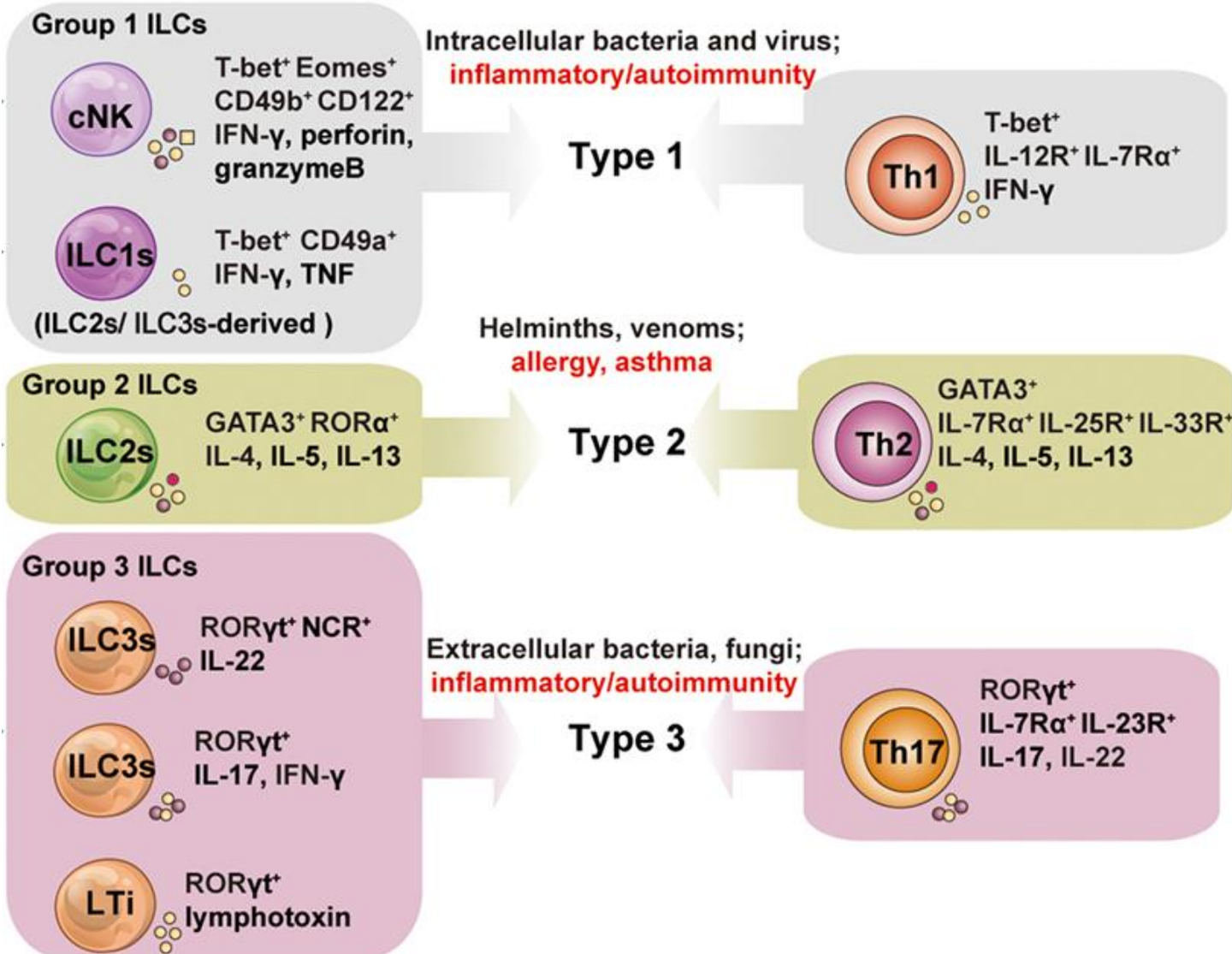


⇒ reduced inhibition of inflammatory gene expression by steroids

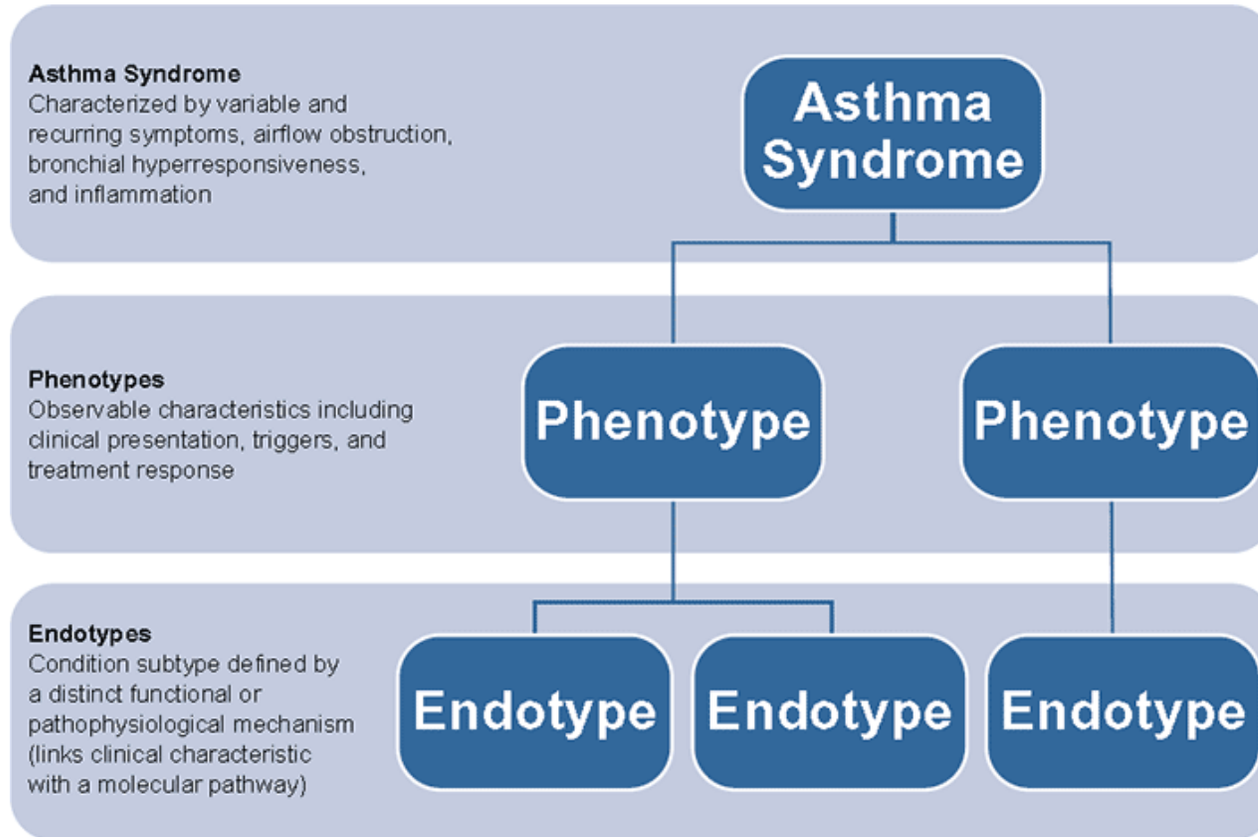
Biologics / monoclonal antibodies



Three types of inflammation



Concept of phenotyping and endotyping a disease



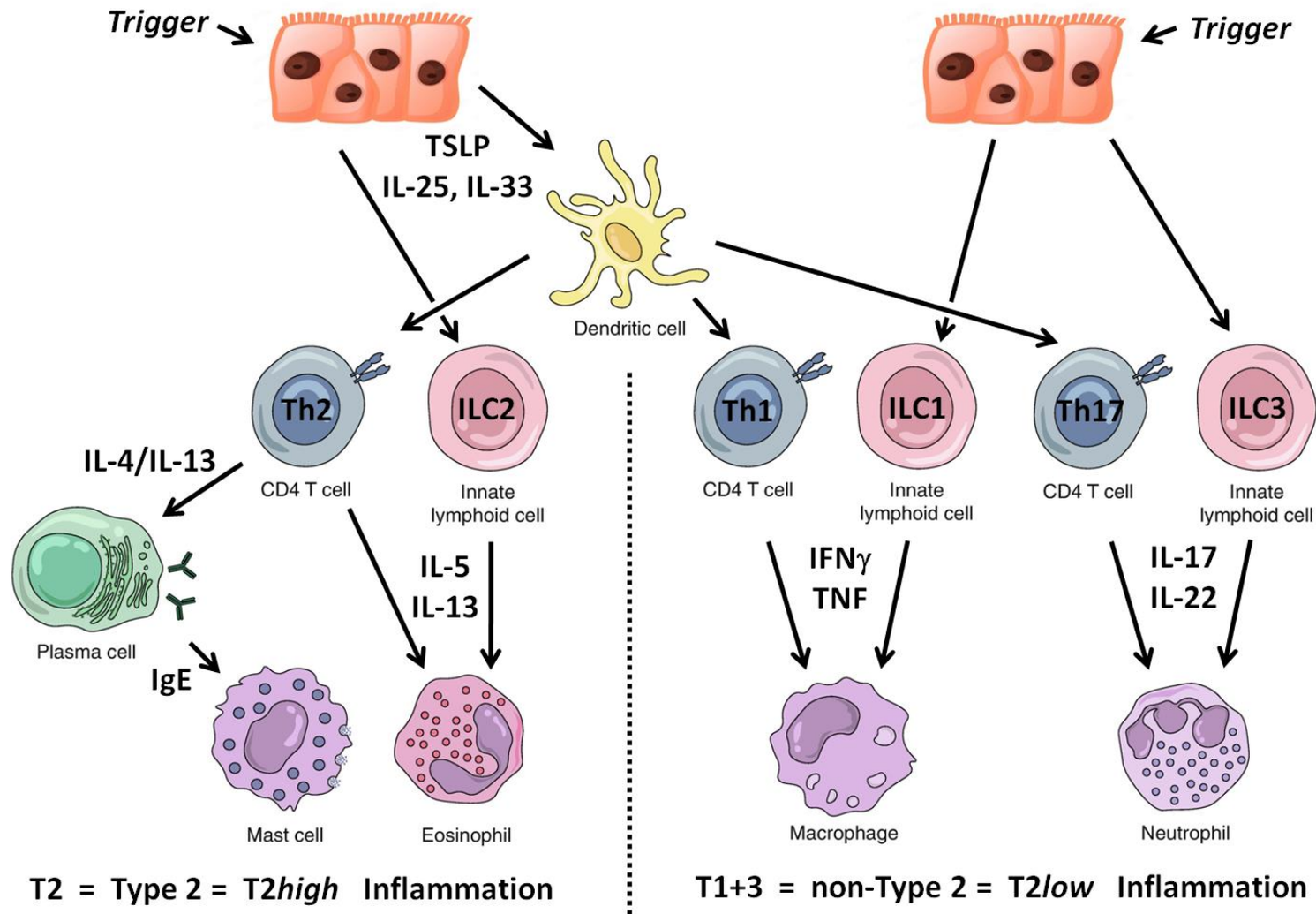
⇒ endotypes are key for the individual success of a therapy / efficiency of a drug

⇒ there are smooth transitions between the different endotypes

⇒ endotypes might change with age and disease progression

⇒ endotype research is the way to a personalized medicine – the ultimate goal to optimize individual therapy strategies

Inflammation endotypes T2 versus non-T2



Endotype:

T2 = Type 2 = T2^{high} Inflammation

T1+3 = non-Type 2 = T2^{low} Inflammation

created/mod. according to
 Brussels et al. Nature Medicine 2013
doi.org/10.3389/fimmu.2017.00695
[doi: 10.1007/s13555-022-00737-7](https://doi.org/10.1007/s13555-022-00737-7)
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[doi: 10.1007/s13555-022-00737-7](https://doi.org/10.1007/s13555-022-00737-7)
doi.org/10.3390/biomedicines10102486

Biomarker

Severe Asthma:

Biomarker for inflammatory patterns

Serum-IgE: ≥ 76 IU/mL

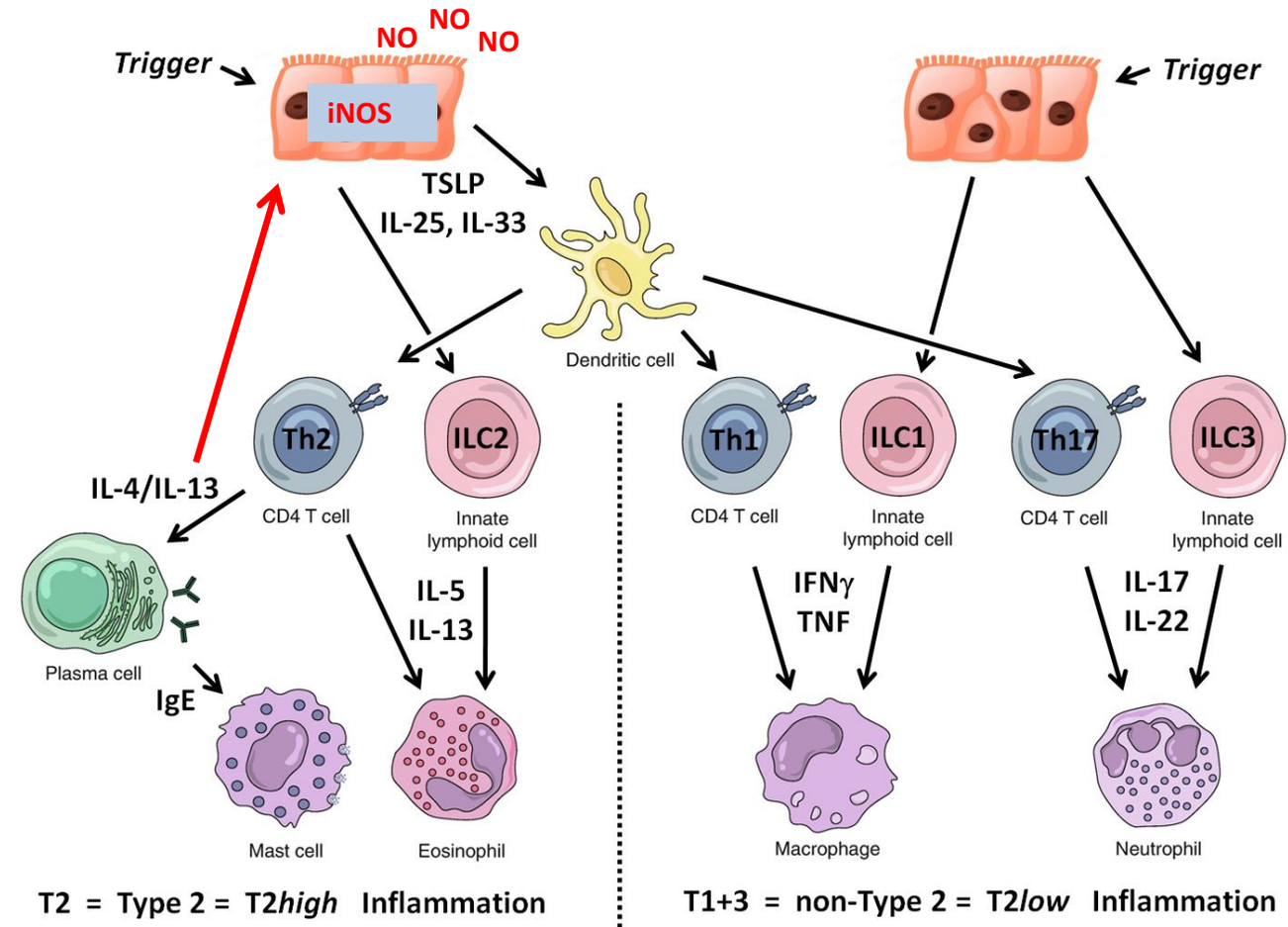
Bluteosinophile: $\geq 150/300$ Zellen/ μ L

FeNO: ≤ 25 ppb

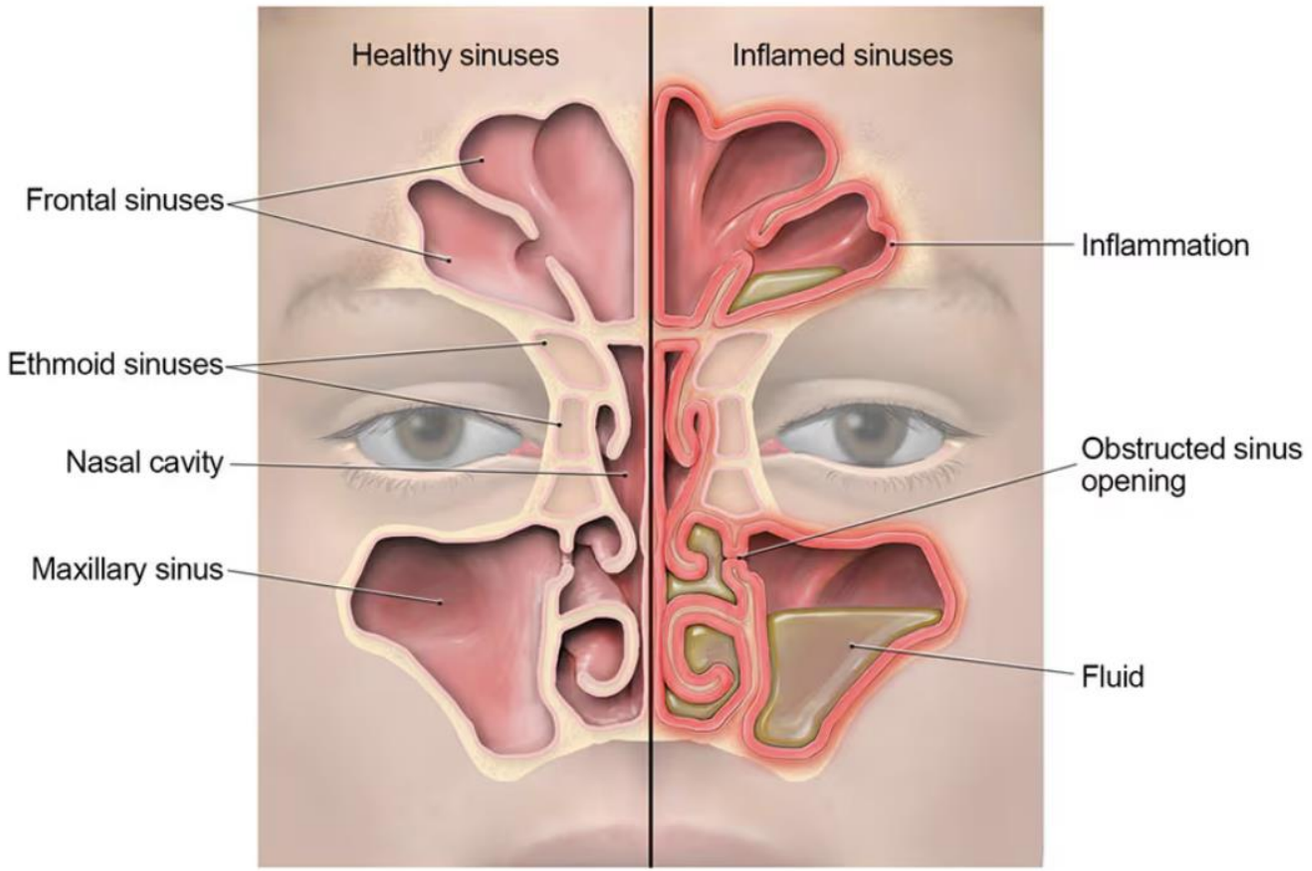
FeNO: ≥ 50 ppb

doi.org/10.3389/fmed.2022.921967

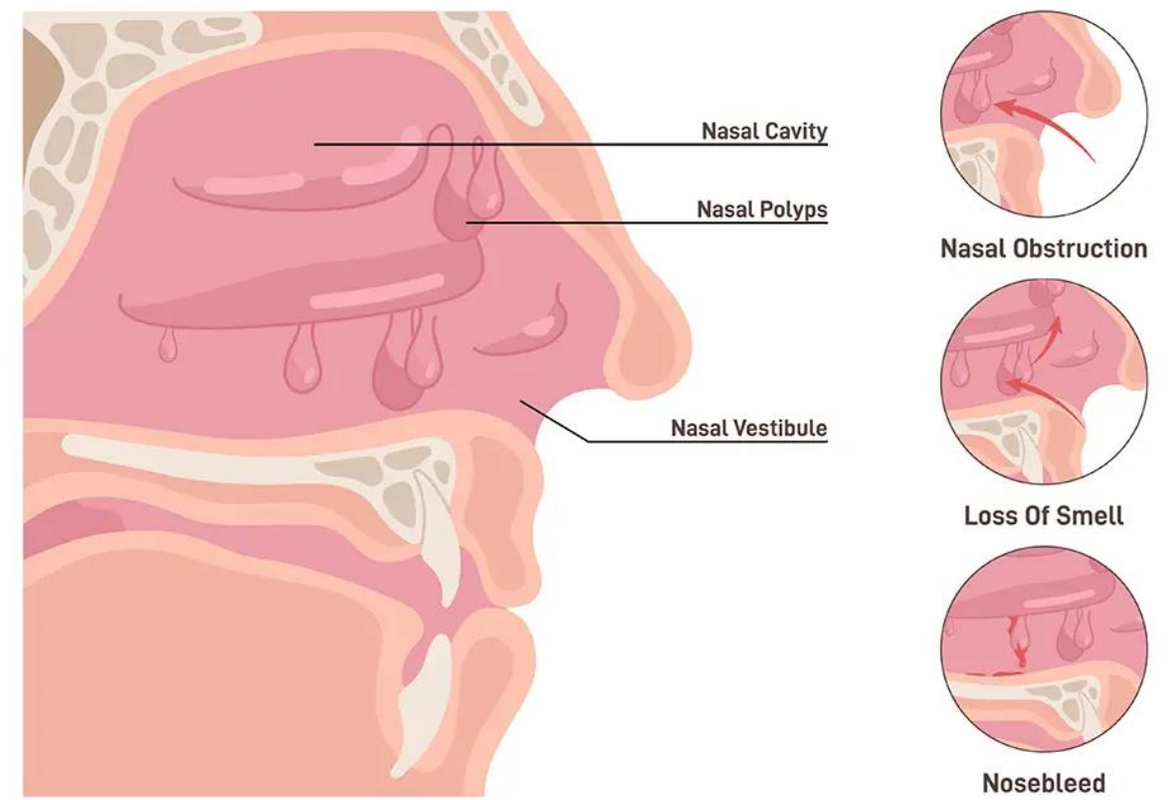
- T2 / allergic inflammation
- T2 / eosinophilic inflammation
- non-T2 / steroid-insensitive inflammation
- T2 / steroid-sensitive inflammation



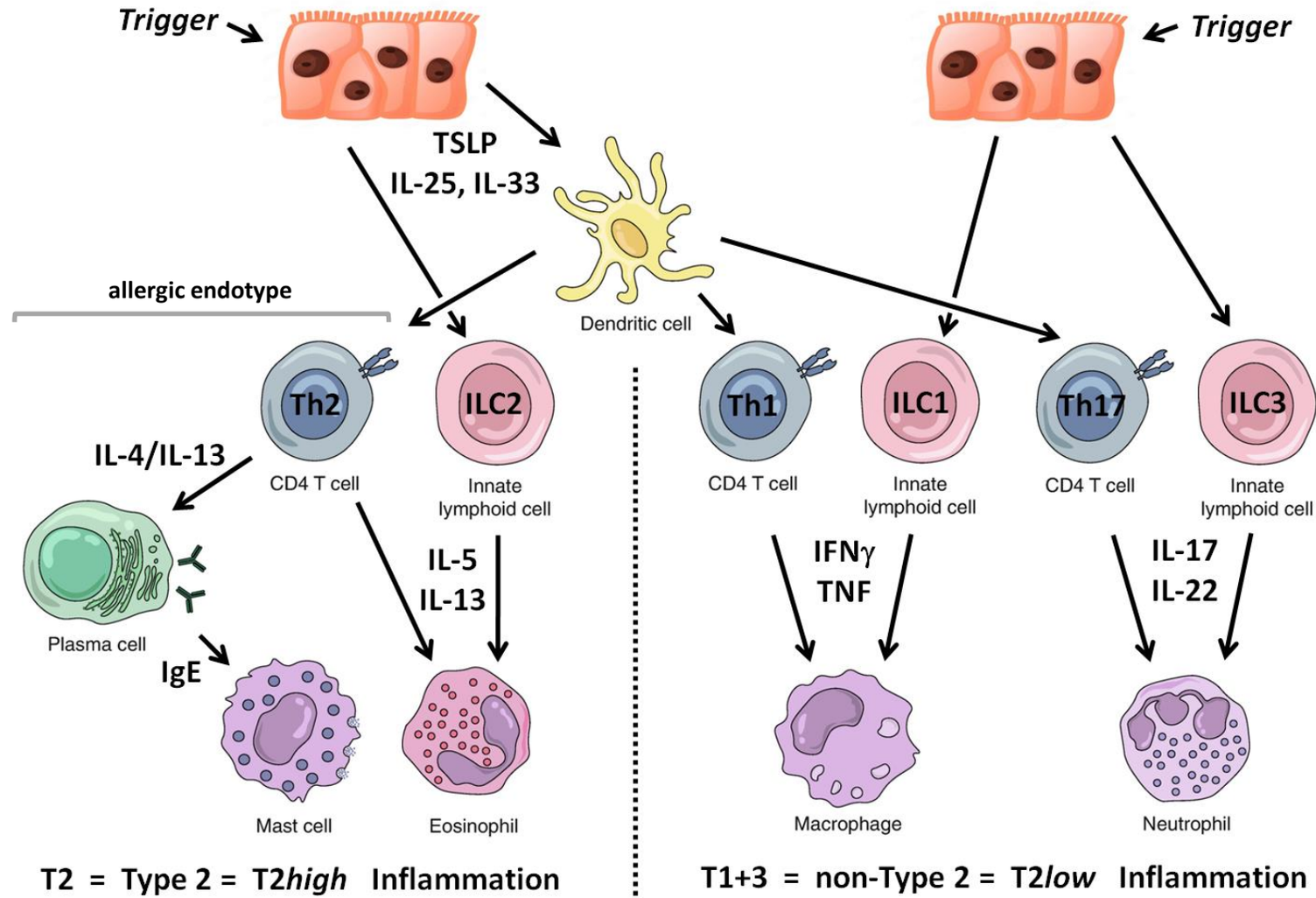
Chronic Rhinosinusitis



Nasal Polyps



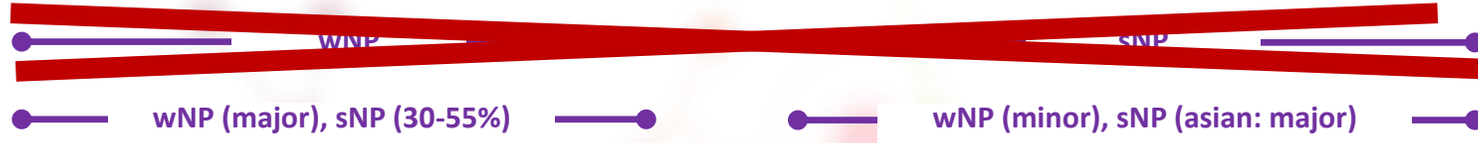
Chronische Rhinosinusitis : Endotypen



T2 = Type 2 = T2high Inflammation

T1+3 = non-Type 2 = T2low Inflammation

created/mod. according to
 Brussels et al. Nature Medicine 2013
doi.org/10.3389/fimmu.2017.00695
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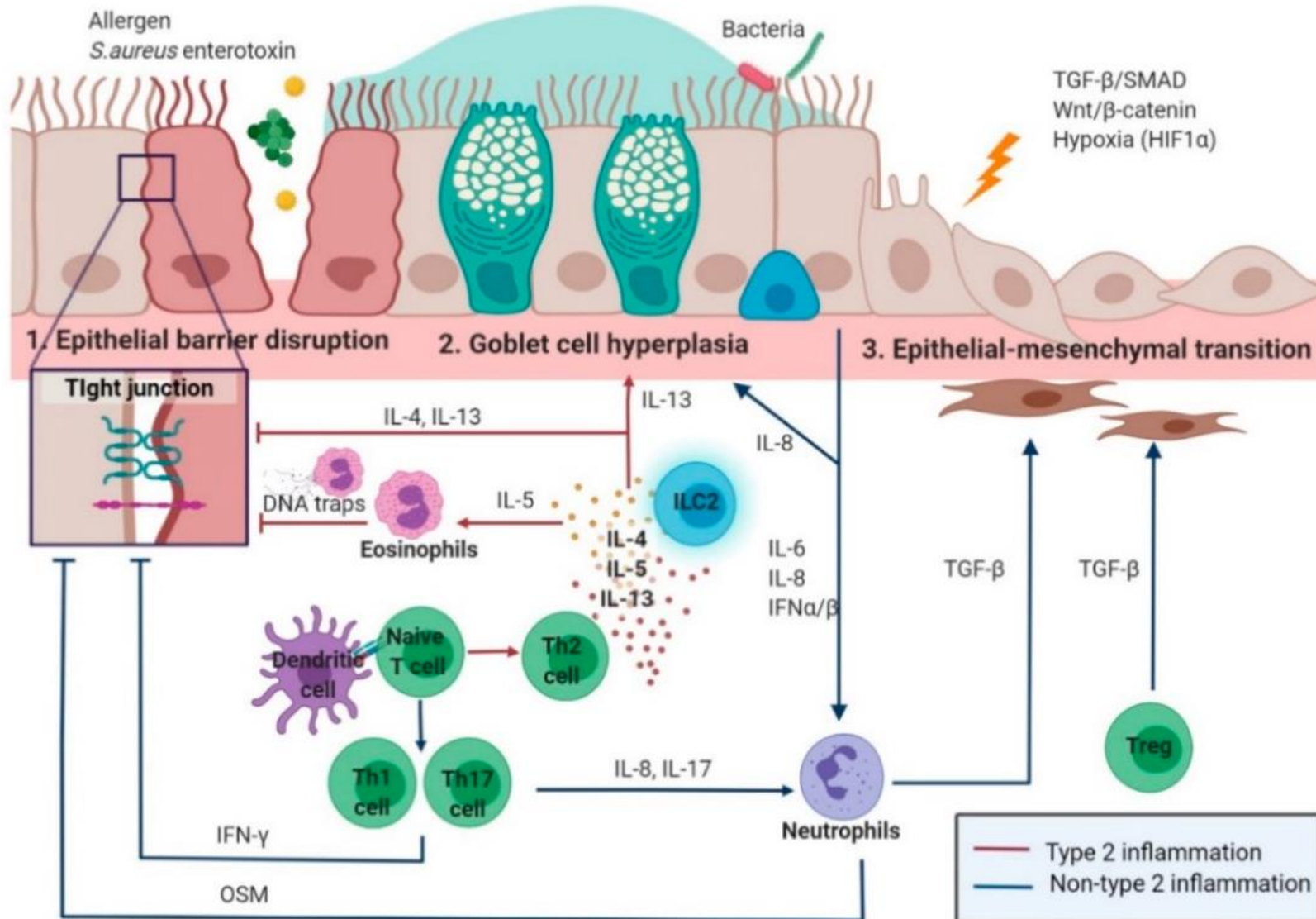
wNP (major), sNP (30-55%)

wNP (minor), sNP (asian: major)

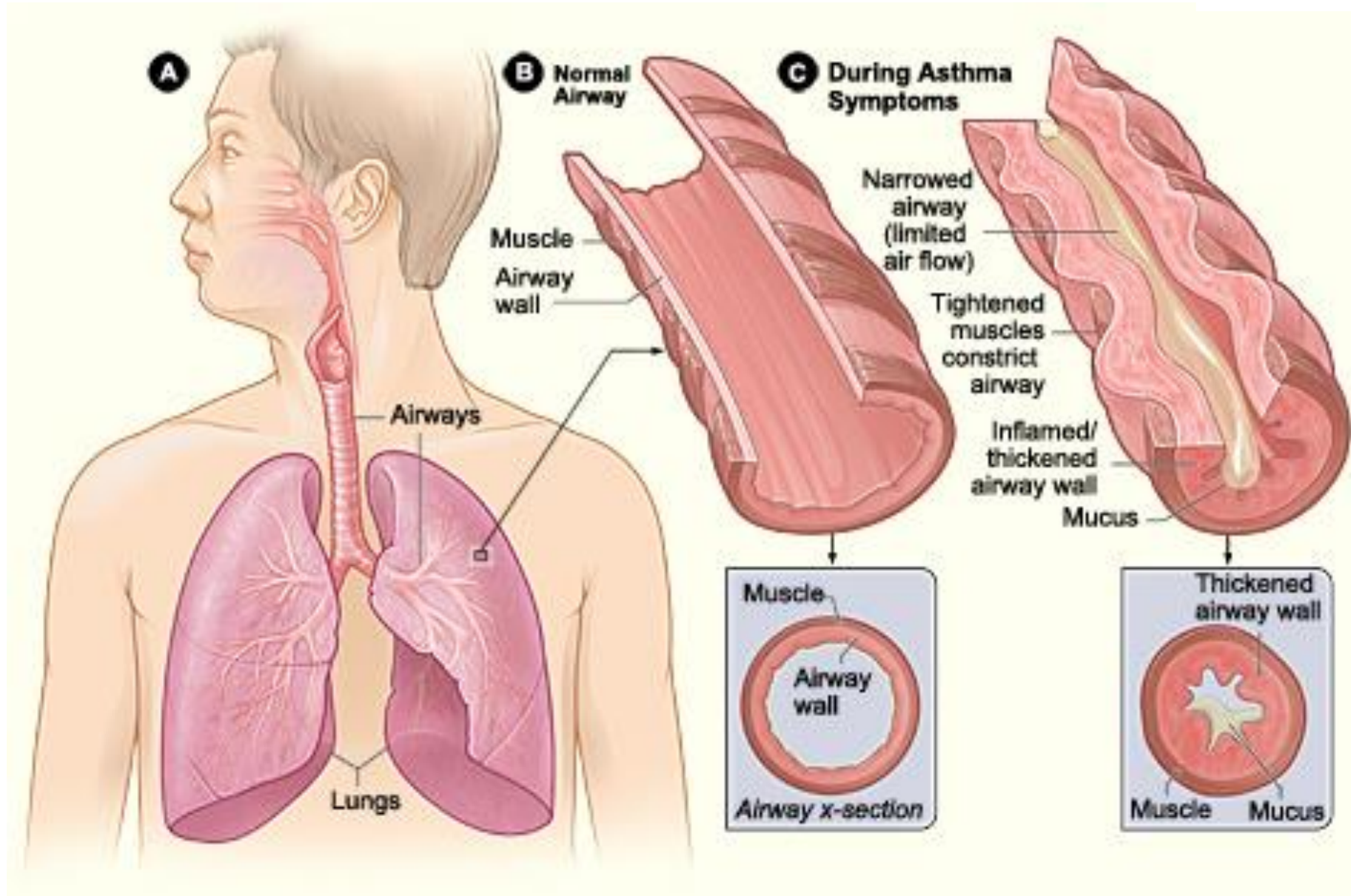
CRS

CRS

Chronic Rhinosinusitis: pathophysiology

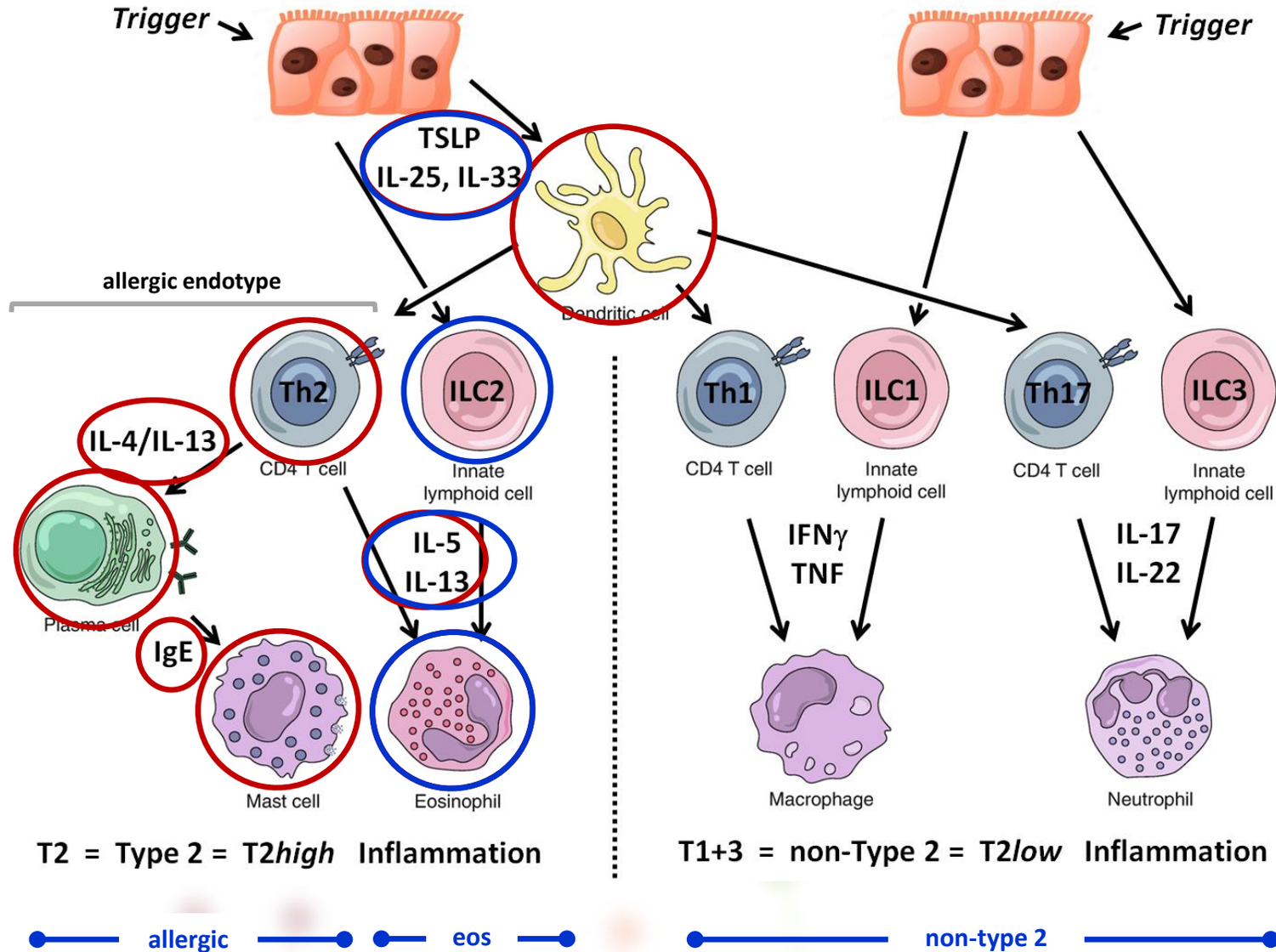


Asthma



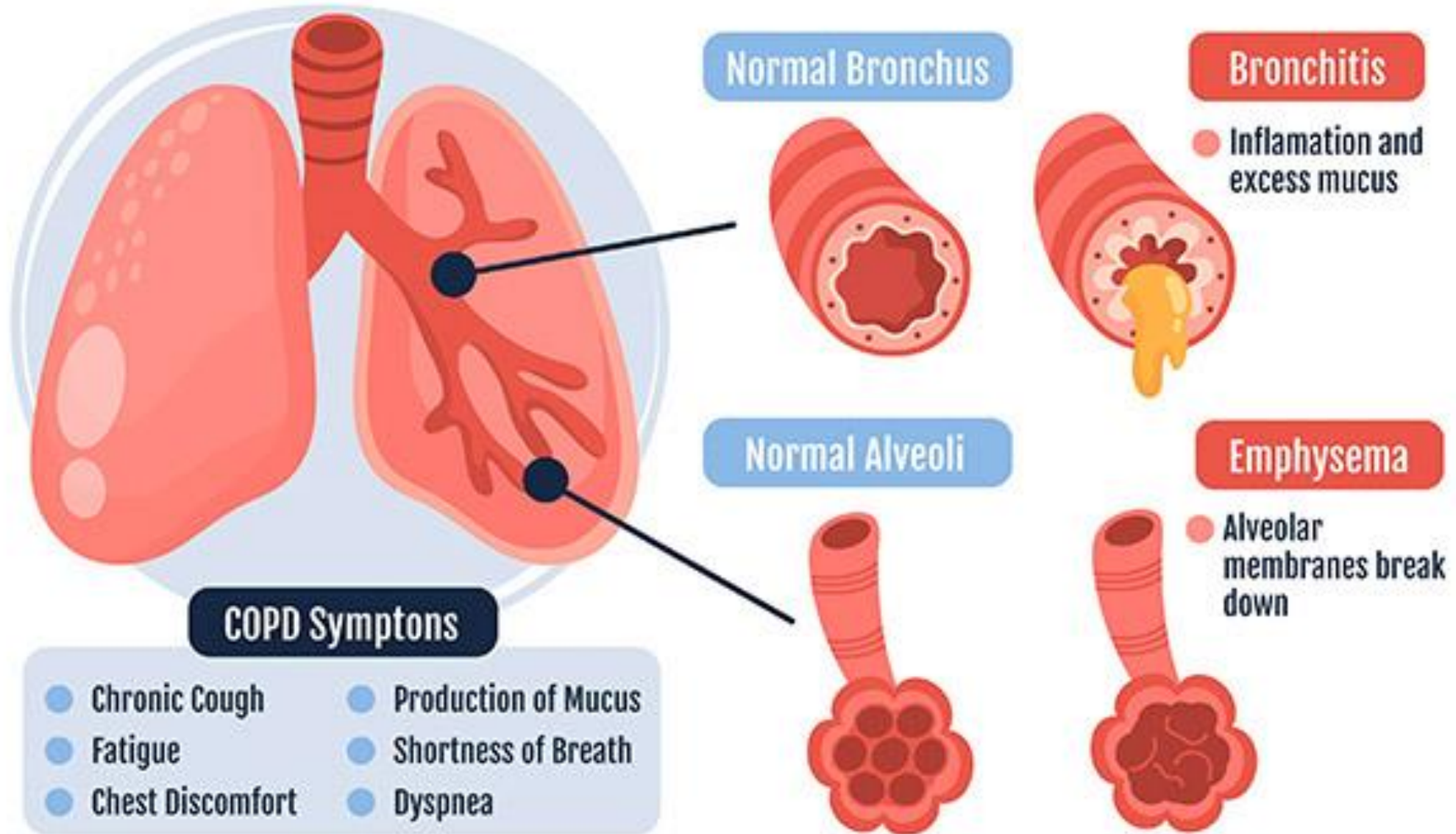
By United States-National Institute of Health: National Heart, Lung, Blood Institute - <http://www.nhlbi.nih.gov/health/health-topics/topics/asthma/>, Public Domain, <https://commons.wikimedia.org/w/index.php?curid=24760677>

Asthma: Endotypen

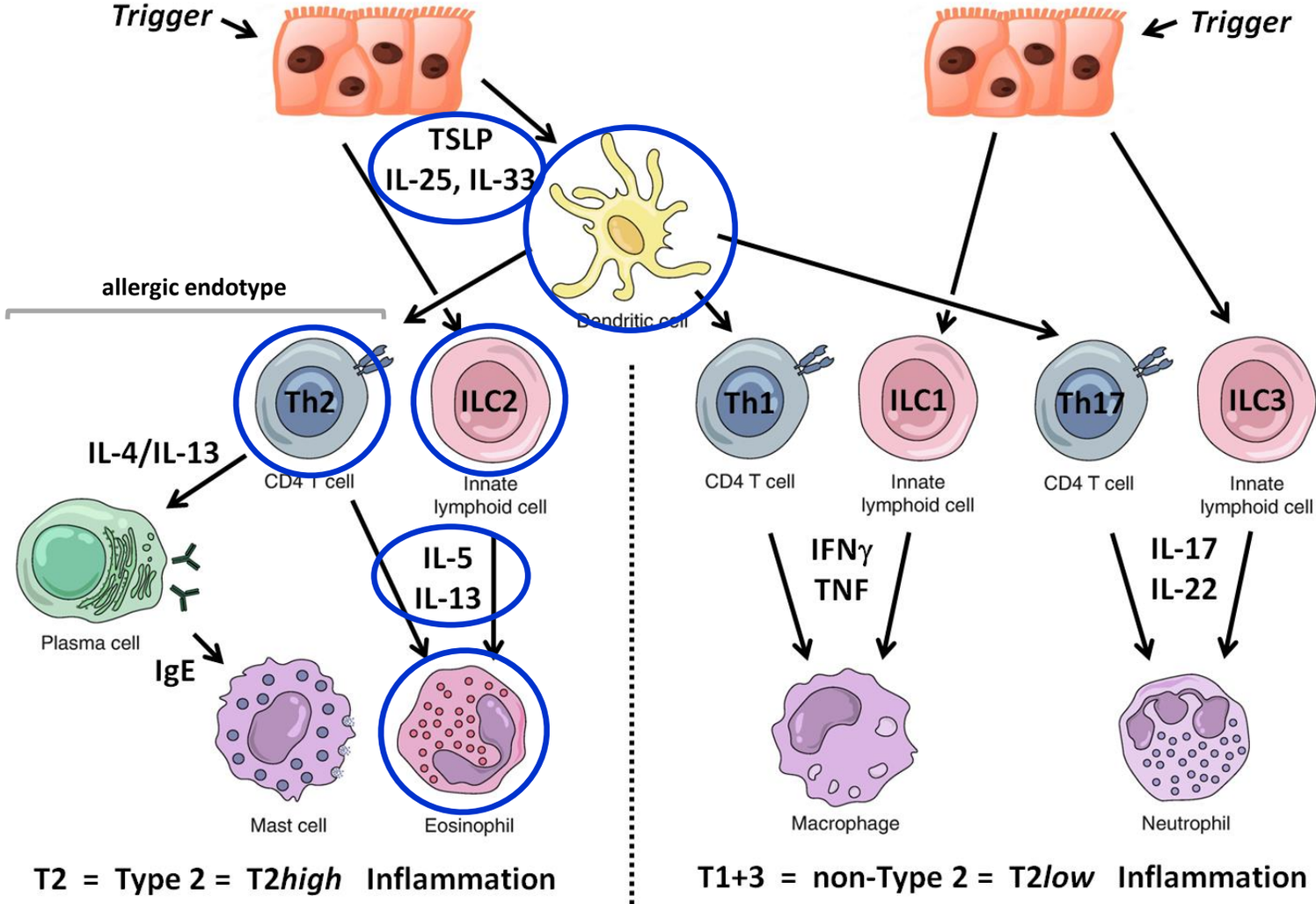


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doi.org/10.3390/biomedicines10102486

COPD (Chronic Obstructive Pulmonary Disease)



COPD: Endotypen

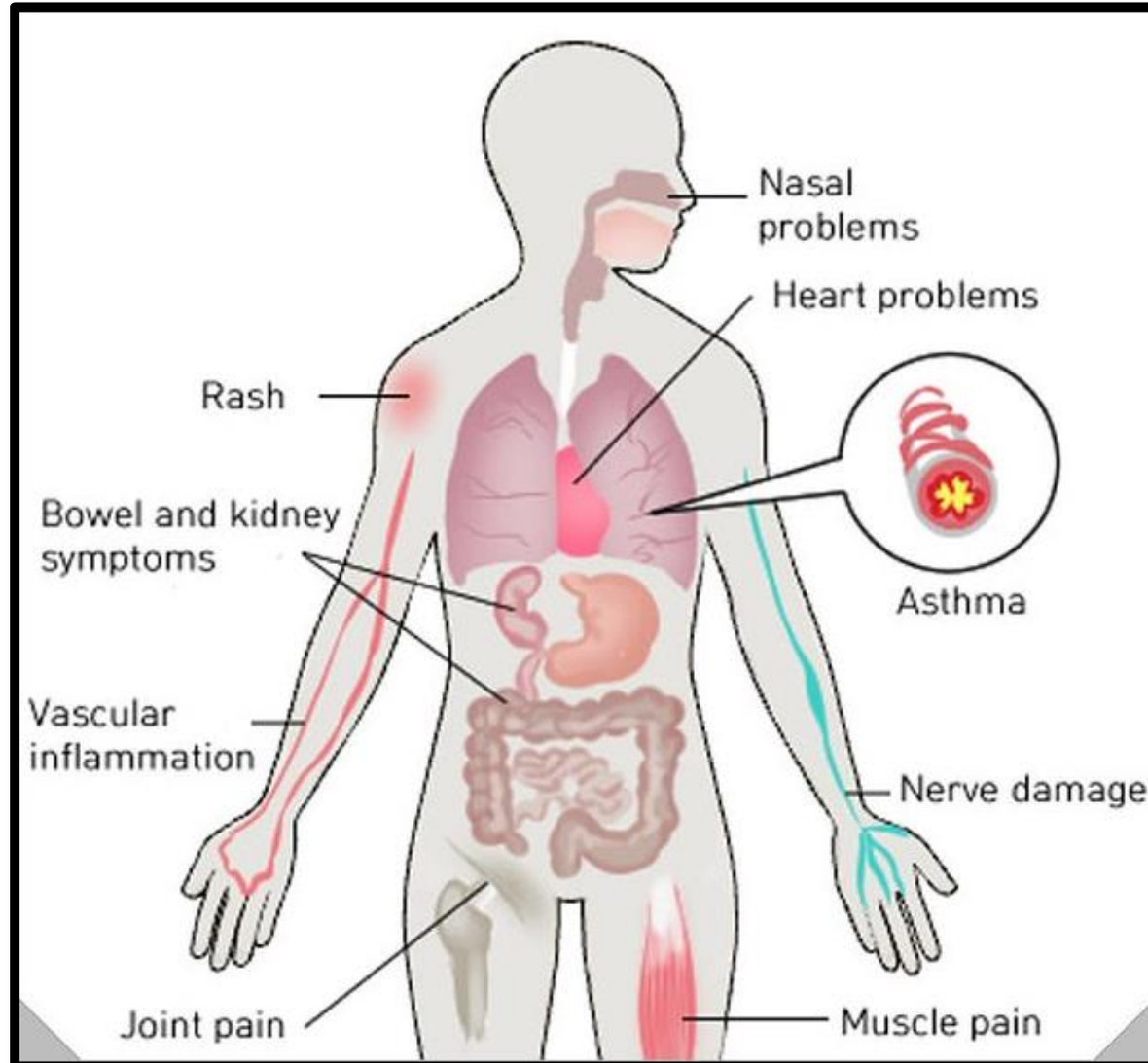


zusätzlich

— + eos —

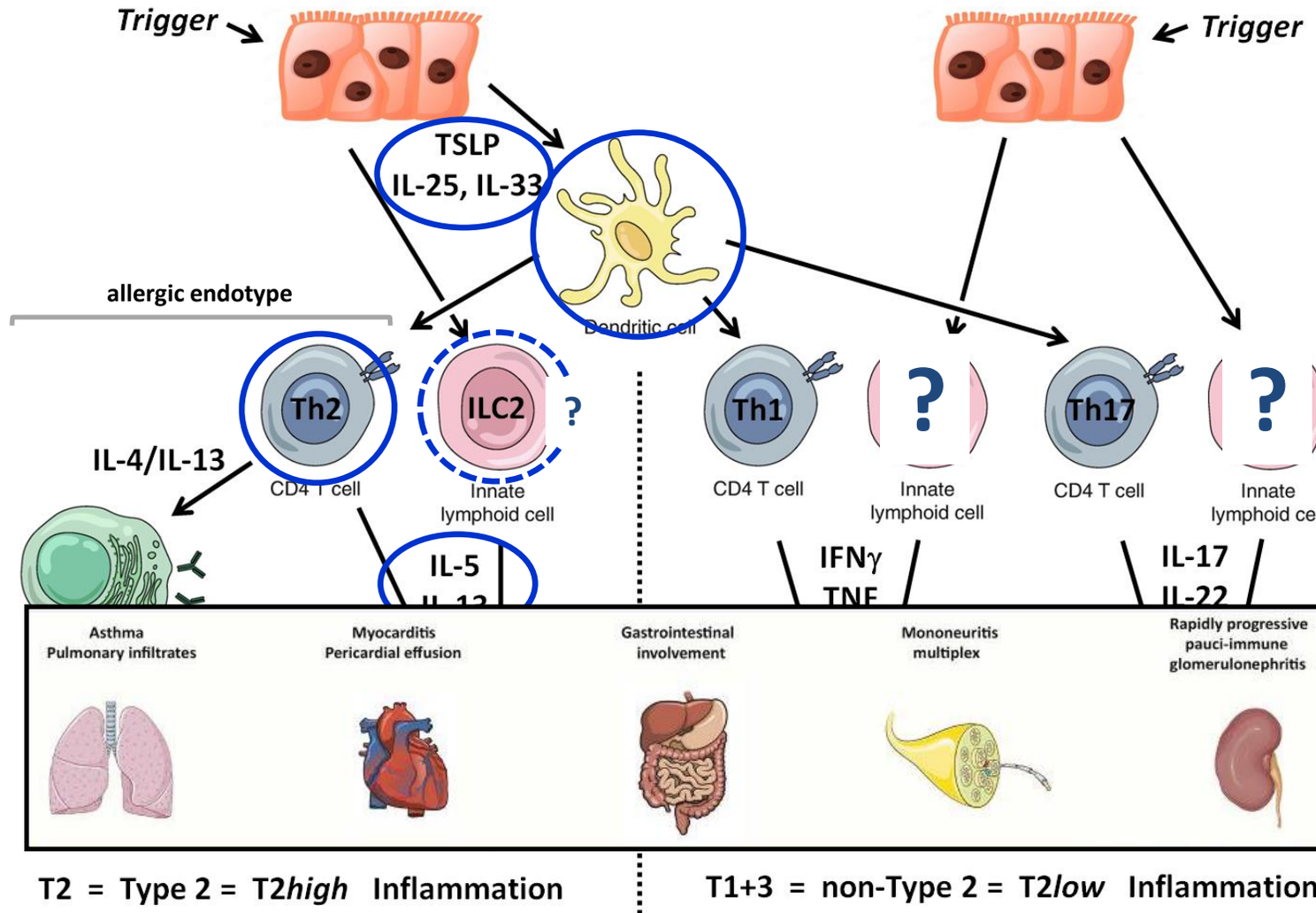
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[doi: 10.3389/fmed.2021.627776](https://doi.org/10.3389/fmed.2021.627776)
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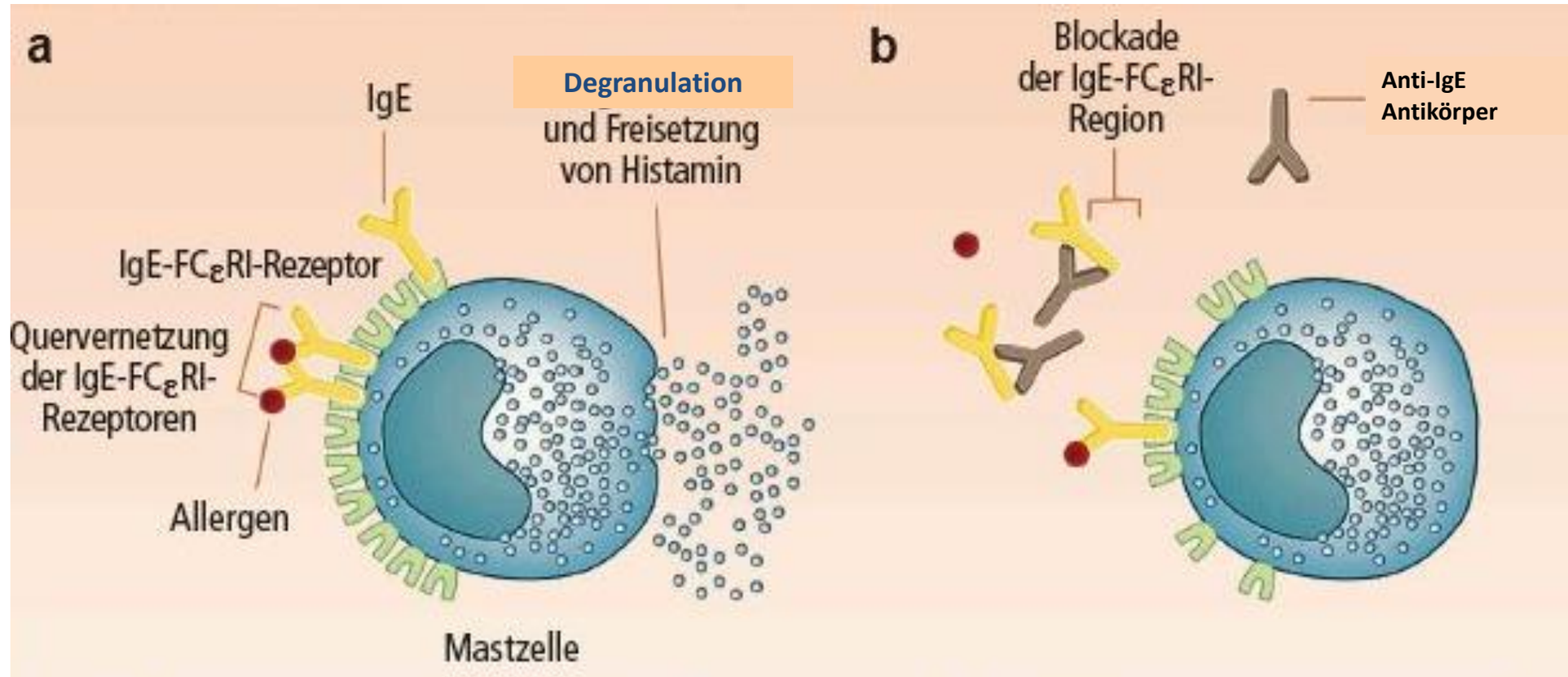
Eosinophile Granulomatose mit Polyangiitis (EGPA)



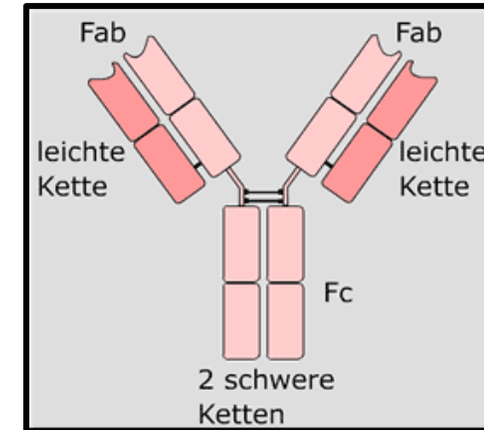
created/mod. according to
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doi.org/10.3389/fimmu.2017.00695
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doi.org/10.3390/biomedicines10102486
[doi: 10.3389/fmed.2021.627776](https://doi.org/10.3389/fmed.2021.627776)



Mechanisms of biologic drugs: anti-IgE antibody (Omalizumab)

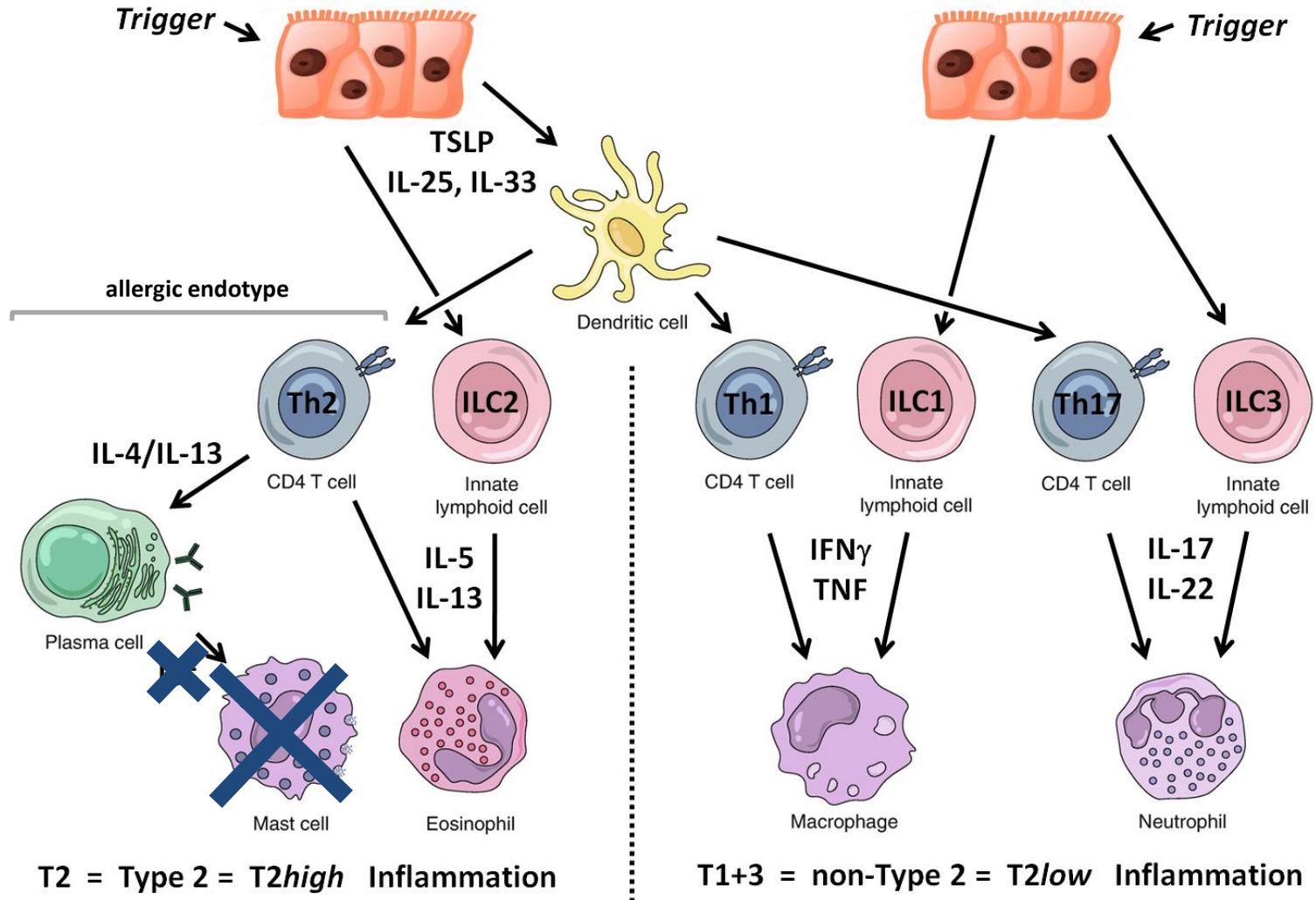


mod. nach Boushey, JACI, 2007



- der anti-IgE Antikörper bindet an die konstante Fc Region des IgE Antikörpers
- dadurch kann der IgE Antikörper nicht an die Mastzellen binden
- dadurch können die Mastzellen nicht durch das Allergen aktiviert werden

Mechanisms of biologic drugs: anti-IgE antibody (Omalizumab)



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doi.org/10.3389/fimmu.2017.00695
[doi: 10.1007/s13555-022-00737-7](https://doi.org/10.1007/s13555-022-00737-7)
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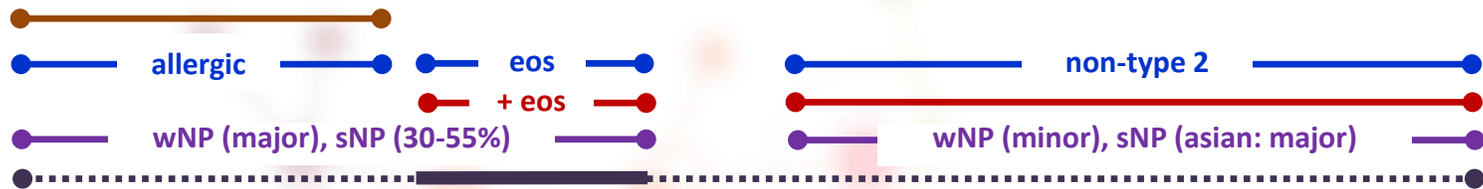
[Allergic rhinitis]

Asthma

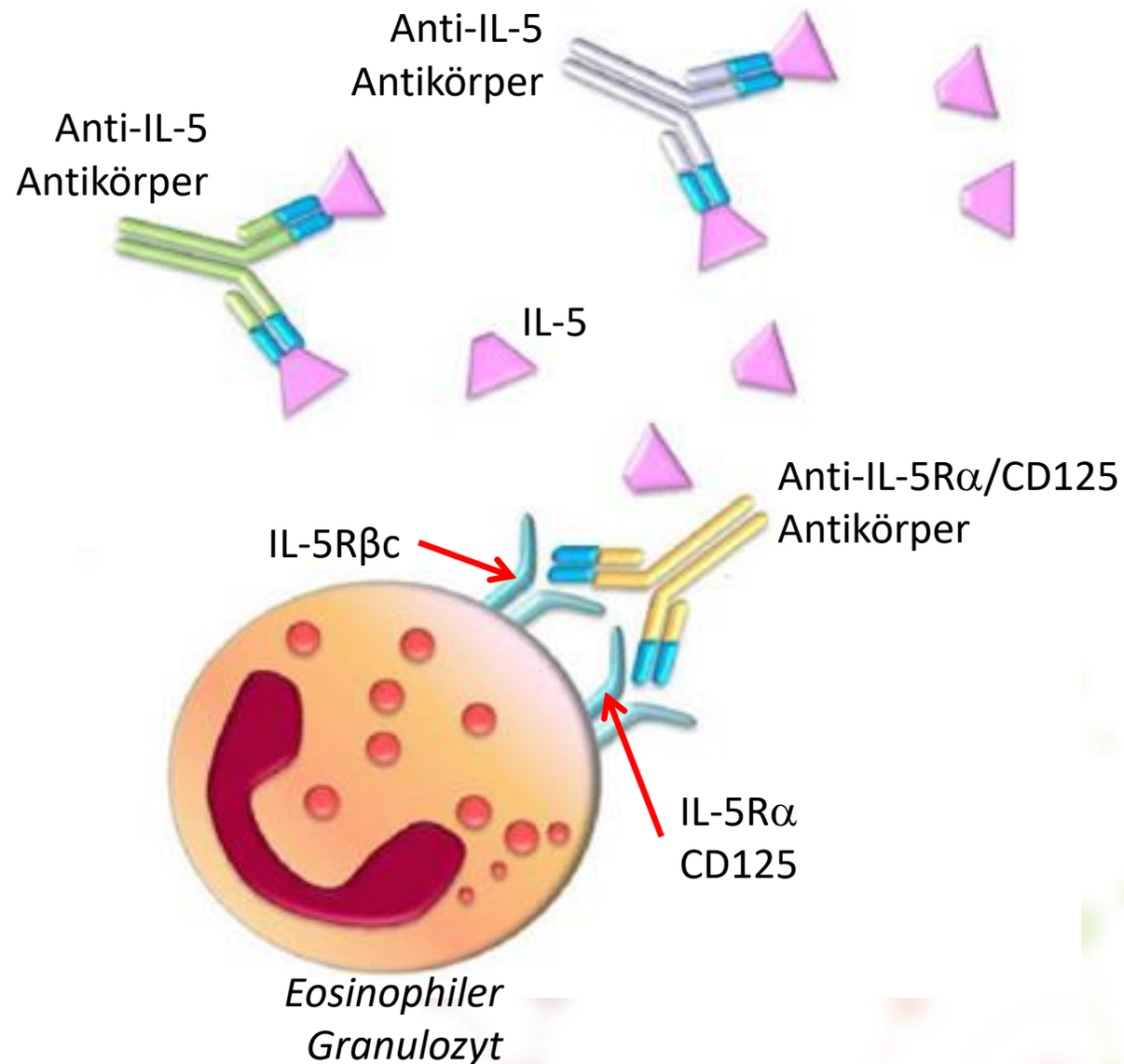
COPD

CRS

EGPA (role of ILCs is unclear)



Mechanisms of biologic drugs: anti-IL-5 und anti-IL-5R α /CD125 antibody (Mepolizumab, Reslizumab, Benralizumab)

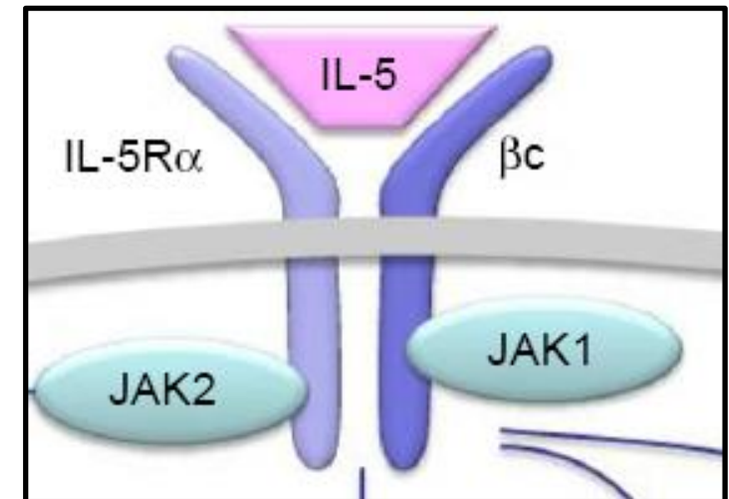


- anti-IL-5 Antikörper binden an IL-5
- anti-IL5R α Antikörper binden an und blockieren den IL-5 Rezeptor auf den Eosinophilen

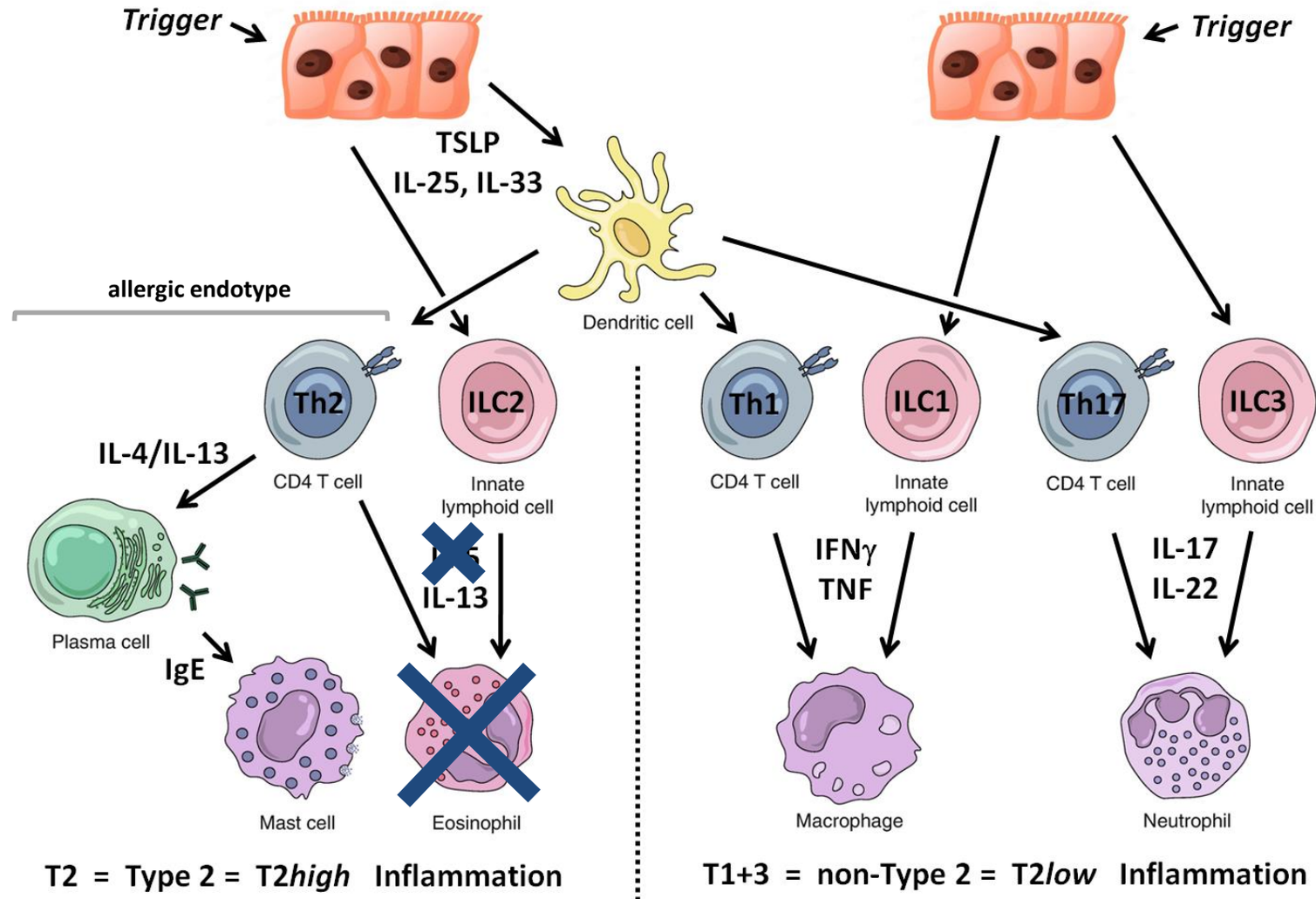
⇒ IL-5 kann nicht an den IL-5 Rezeptor auf den Eosinophilen binden

⇒ die Eosinophilen werden nicht aktiviert

⇒ die Eosinophilen gehen in den Zelltod



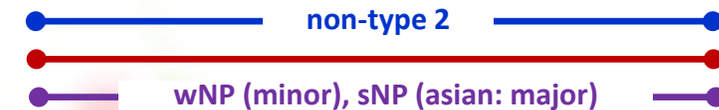
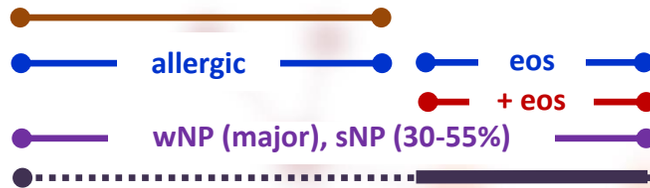
Mechanisms of biologic drugs: anti-IL-5 und anti-IL-5Rα/CD125 antibody (Mepolizumab, Reslizumab, Benralizumab)



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doi.org/10.3390/biomedicines10102486

T2 = Type 2 = T2^{high} Inflammation

T1+3 = non-Type 2 = T2^{low} Inflammation



[Allergic rhinitis]

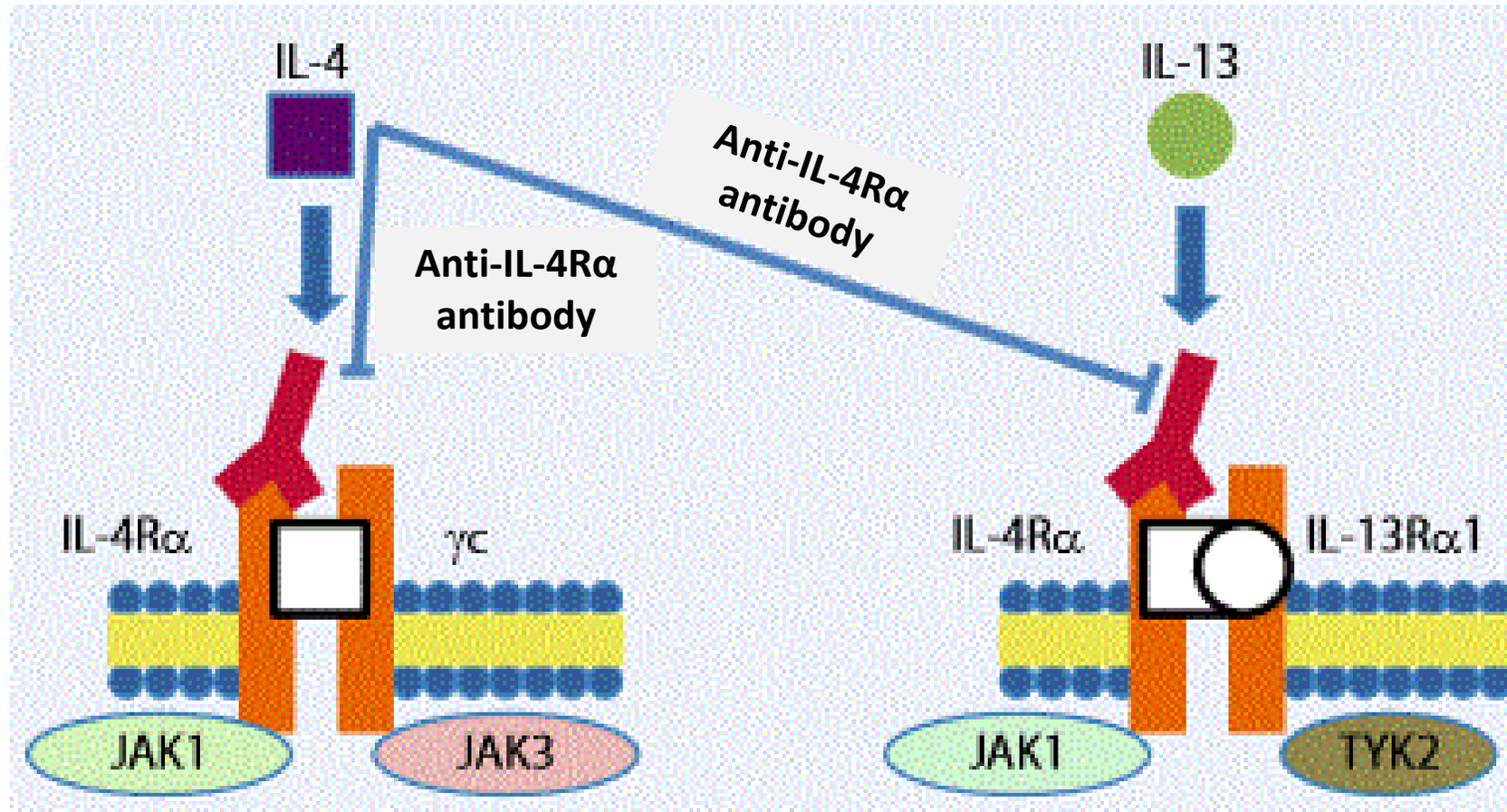
Asthma

COPD

CRS

EGPA (role of ILCs is unclear)

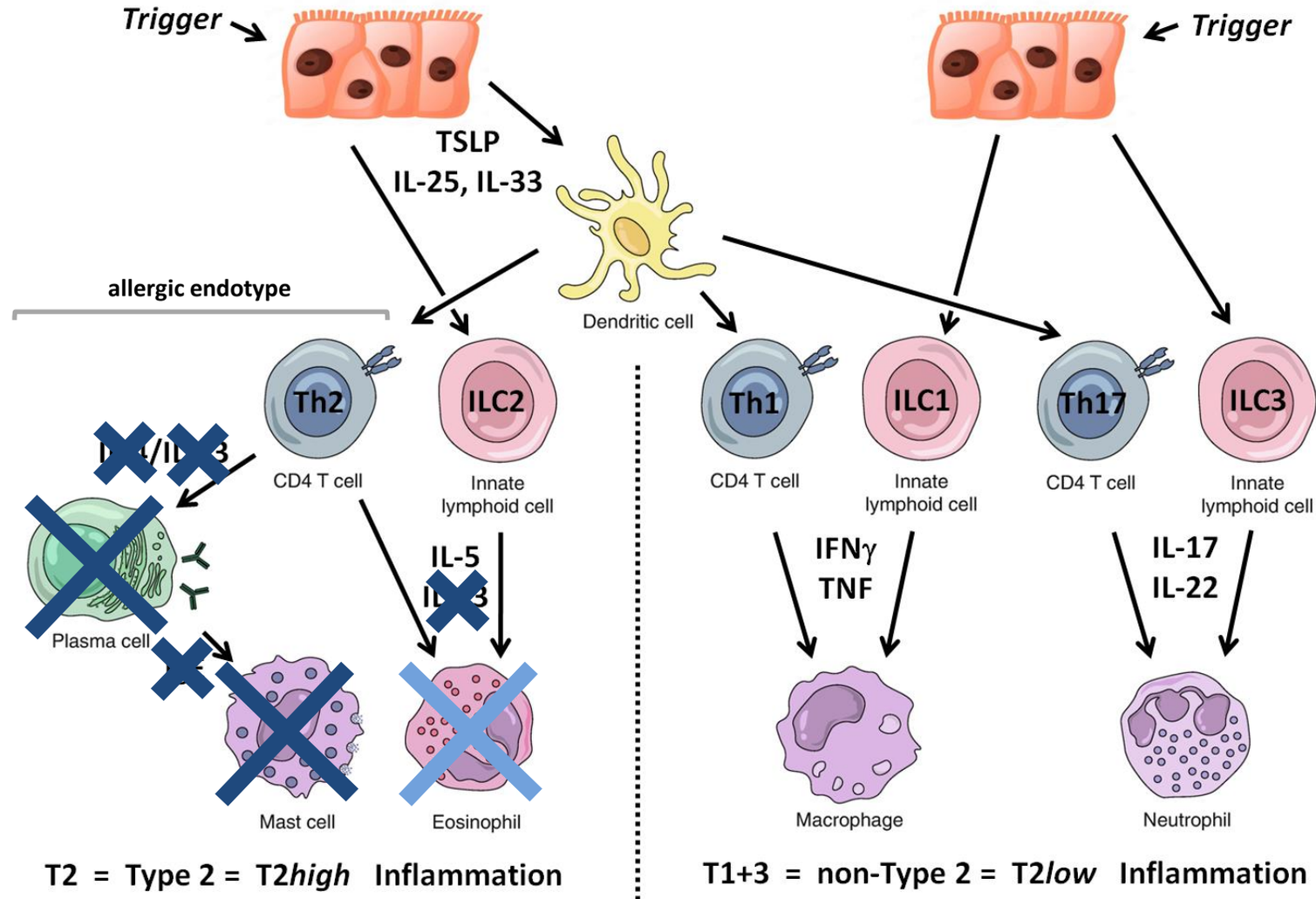
Mechanisms of biologic drugs: anti-IL-4R α antibody (Dupilumab)



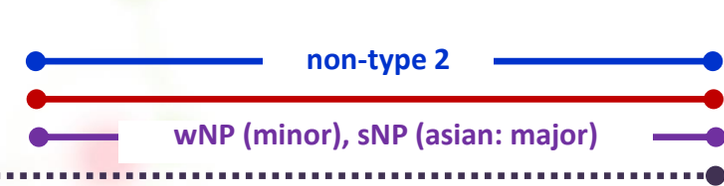
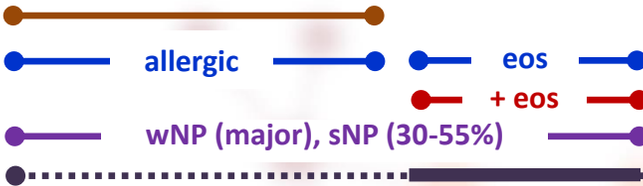
mod. according
to <http://www.springermedizin.at/artikel/50157>

- IL-4- and IL-13-receptors each are heterodimers
- both receptors contain IL-4R α
- ⇒ anti-IL4R α antibody binds and blocks both IL-4 and IL-13 receptors
- ⇒ B cells are not activated by IL-4 and IL-13 to produce IgE antibodies
- ⇒ this concept evades the redundant functions of IL-4 and IL-13

Mechanisms of biologic drugs: anti-IL-4R α antibody (Dupilumab)



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[Allergic rhinitis]

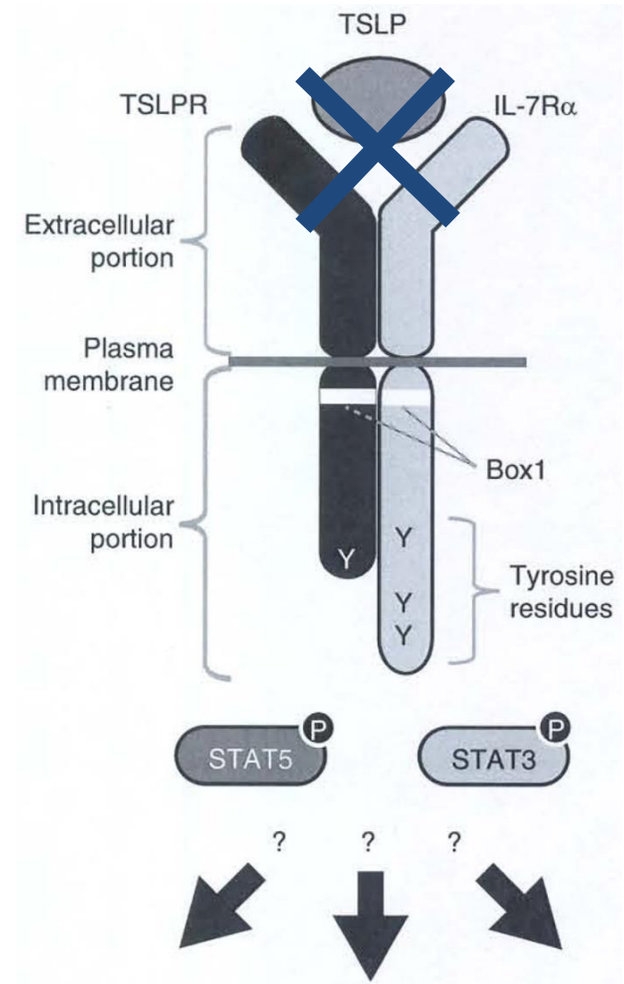
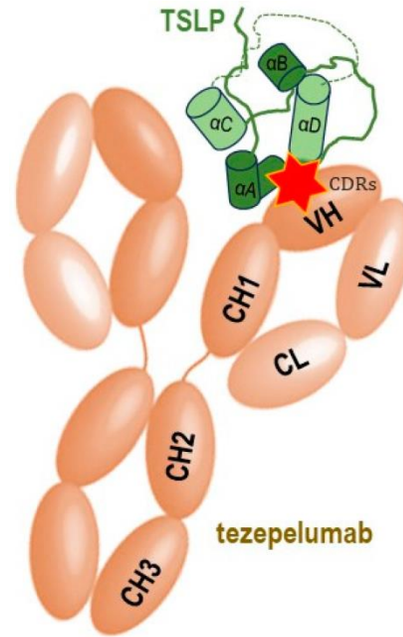
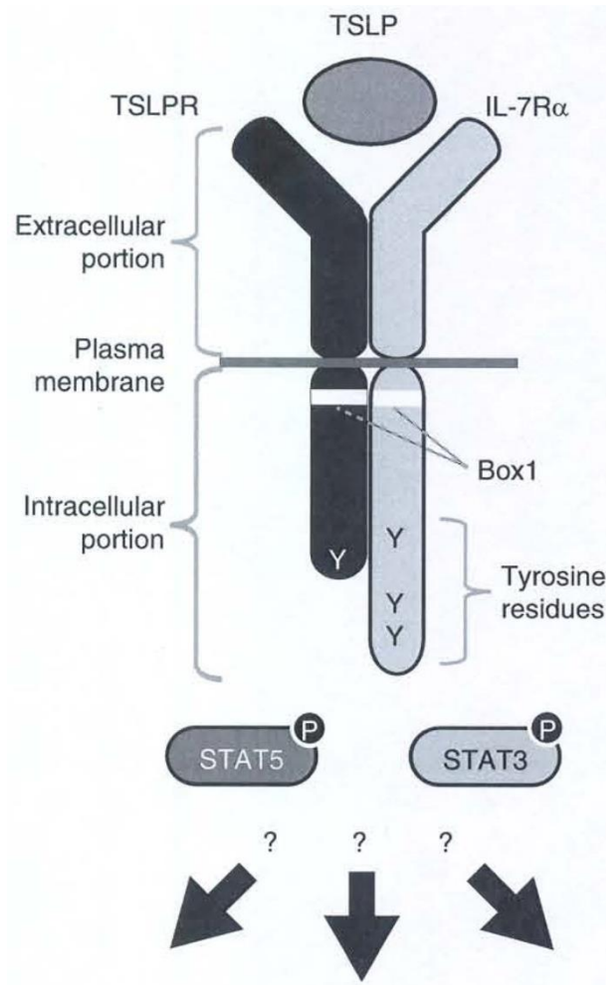
Asthma

COPD

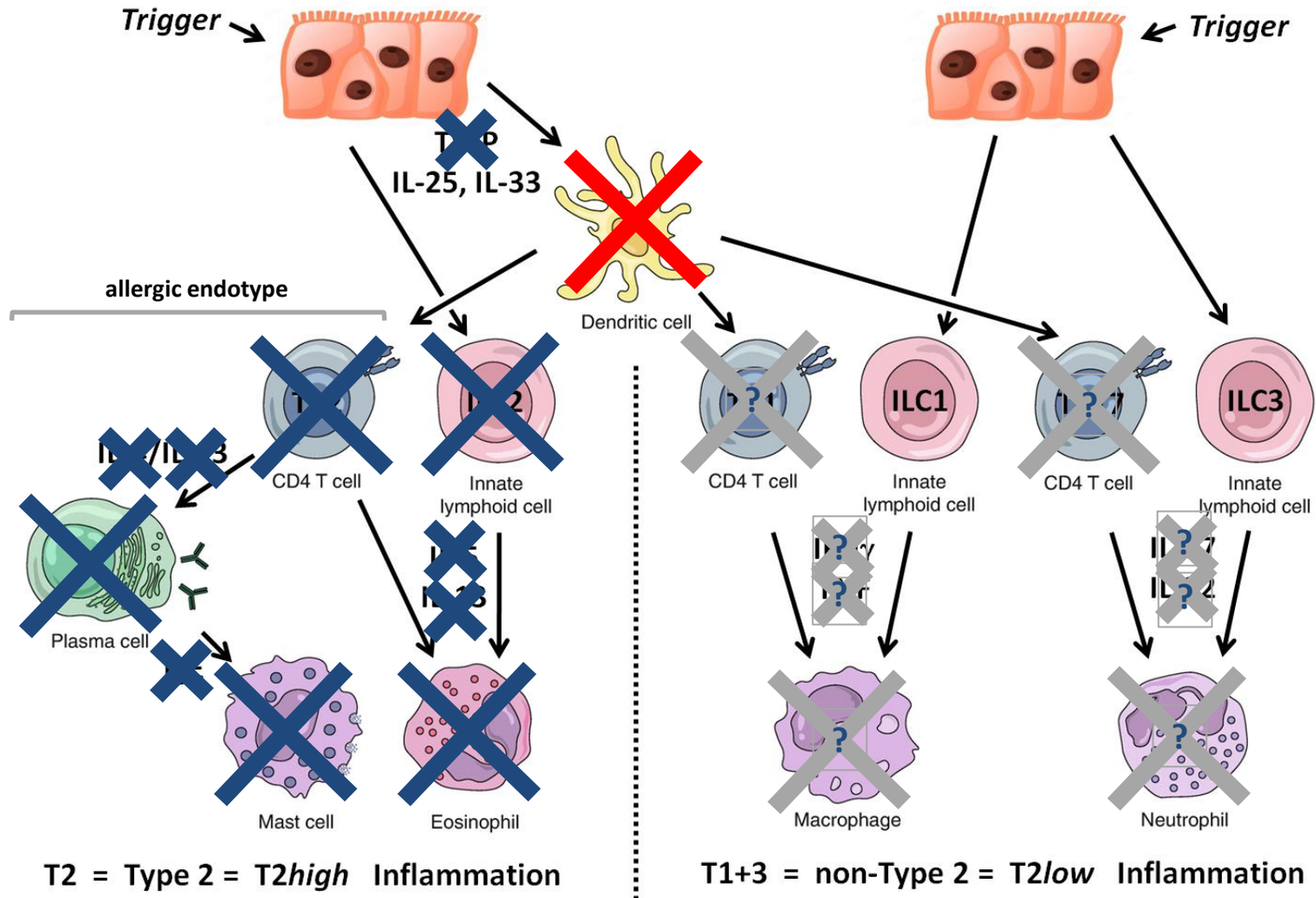
CRS

EGPA (role of ILCs is unclear)

Mechanisms of biologic drugs: anti-TSLP antibody (Tezepelumab)



Mechanisms of biologic drugs: anti-TSLP antibody (Tezepelumab)



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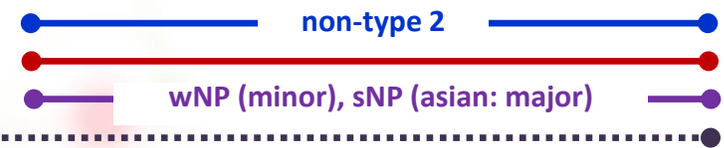
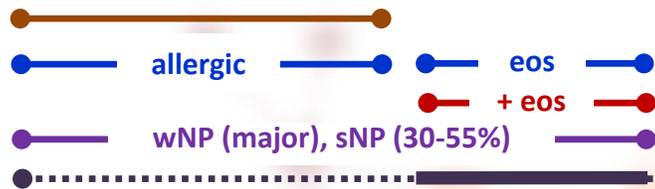
[Allergic rhinitis]

Asthma

COPD

CRS

EGPA (role of ILCs is unclear)



NUCALA 
mepolizumab

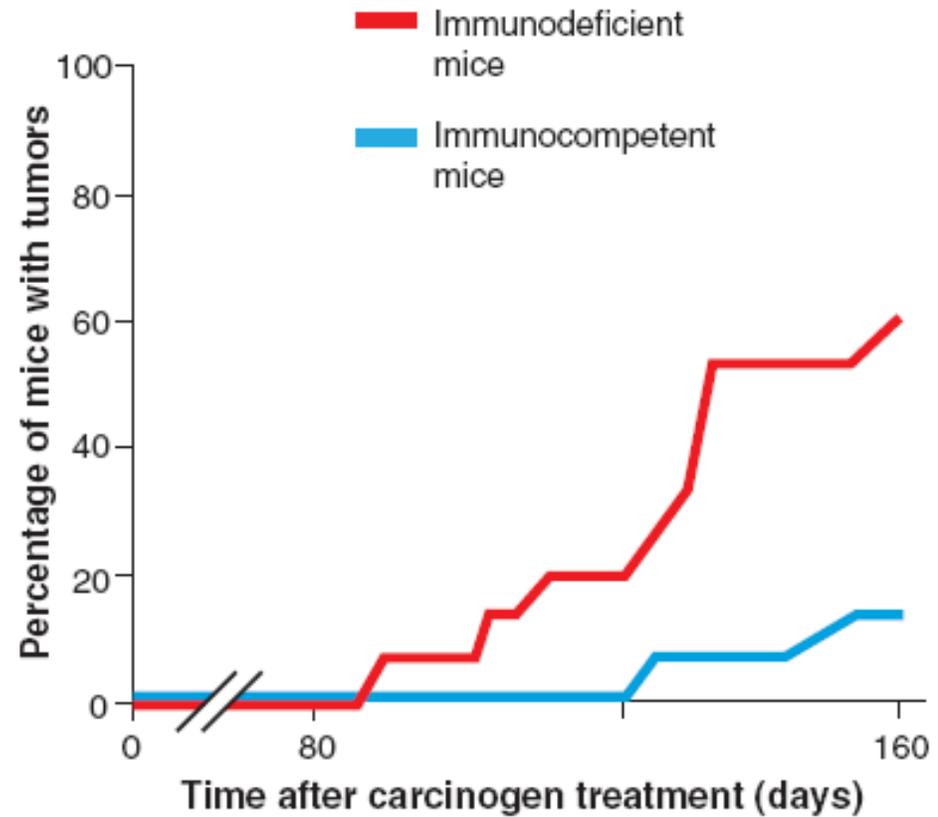
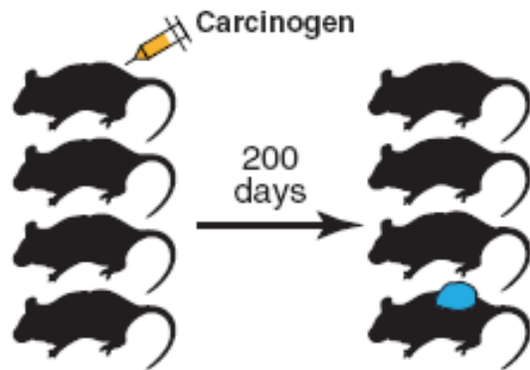
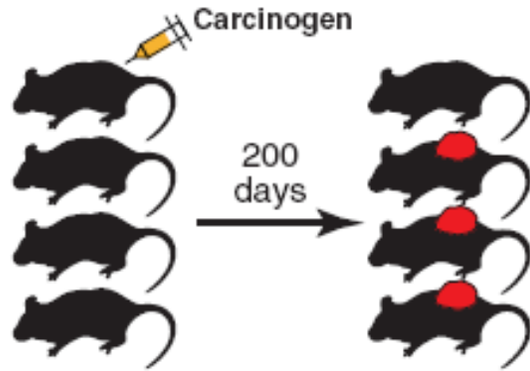
nexus 

GSK



Animal experimental evidence of the "immunosurveillance Theory" by Burnet (1957)

RAG2-deficient mice

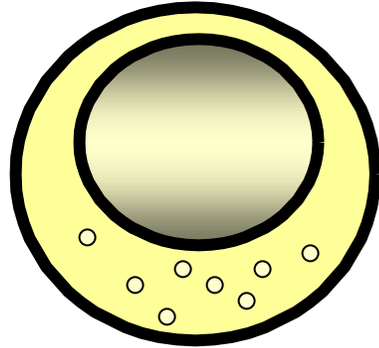


immunocompetent mice



*"Immunosurveillance" by natural killer cells
of the innate immune system*

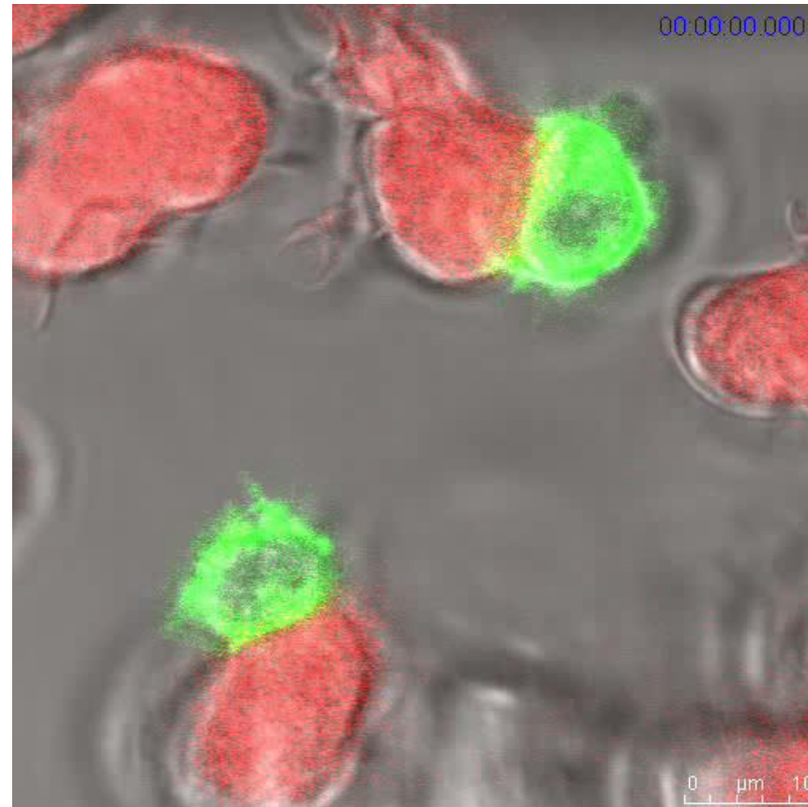
Natural Killer Cells (NK-cells)



function



**Release of Perforin and
Granzym which can kill
tumor cells**



NK cells recognise tumour cells: mechanism 1

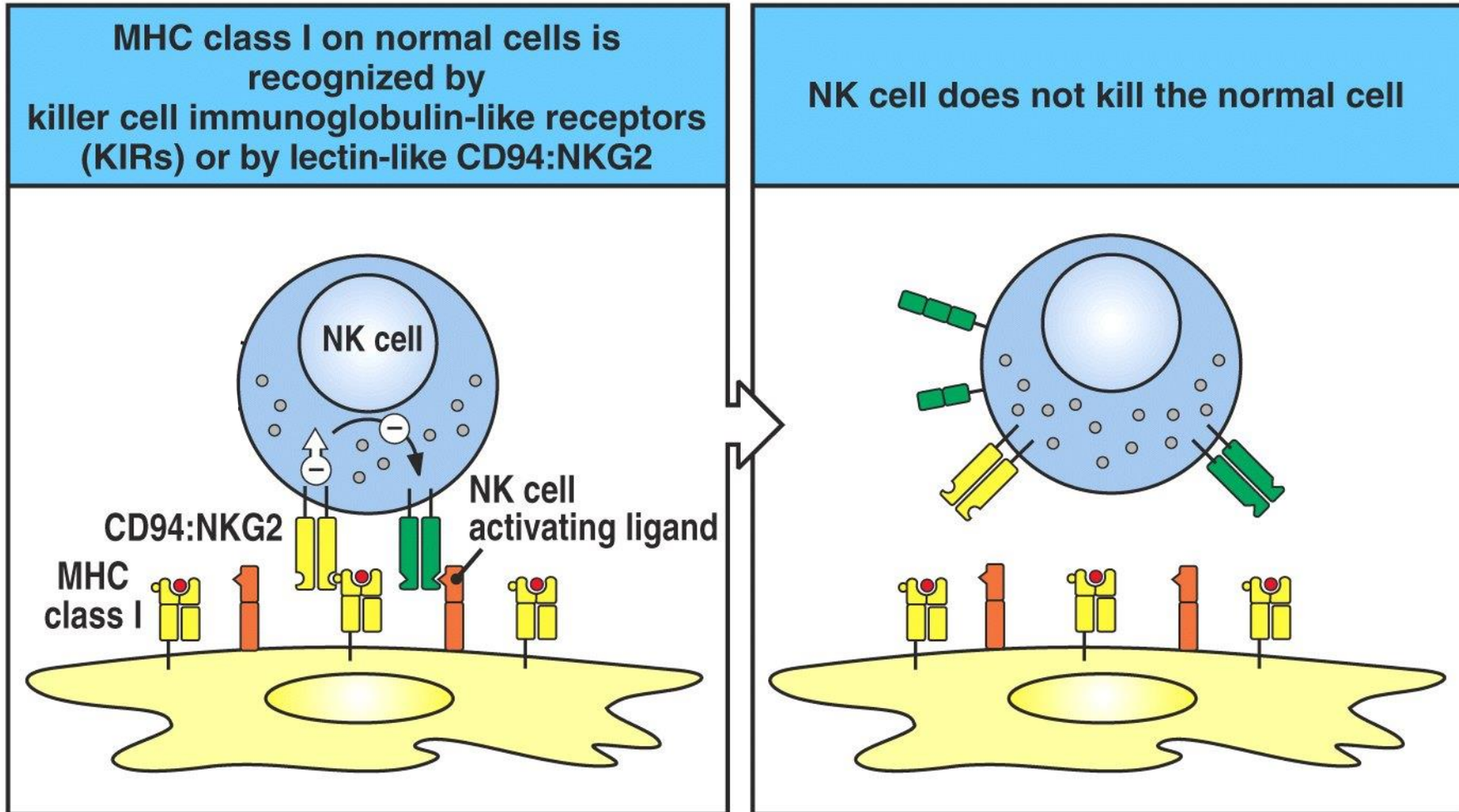
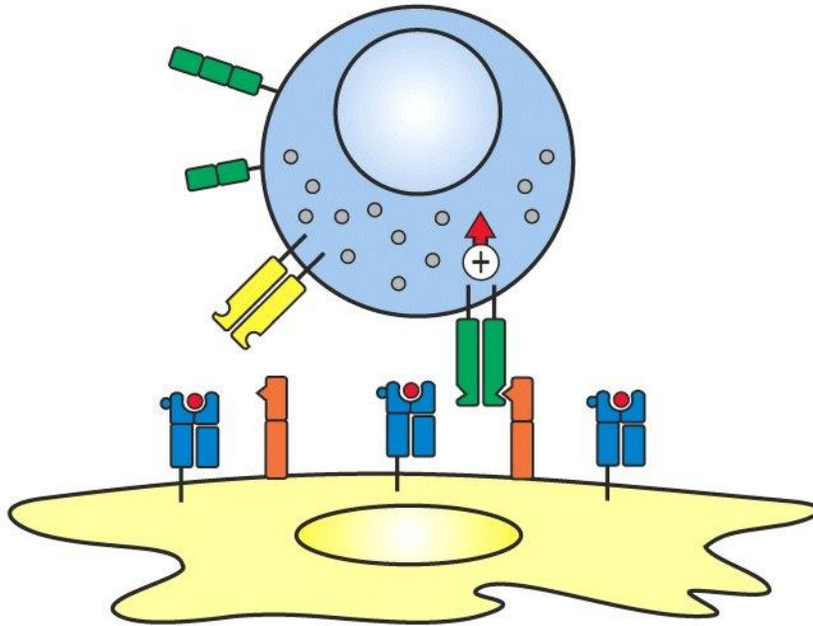


Figure 2-50 part 1 of 2 Immunobiology, 6/e. (© Garland Science 2005)

'Altered' or absent MHC class I cannot stimulate a negative signal. The NK cell is triggered by signals from activating



Activated NK cell releases granule contents, inducing apoptosis in target cell

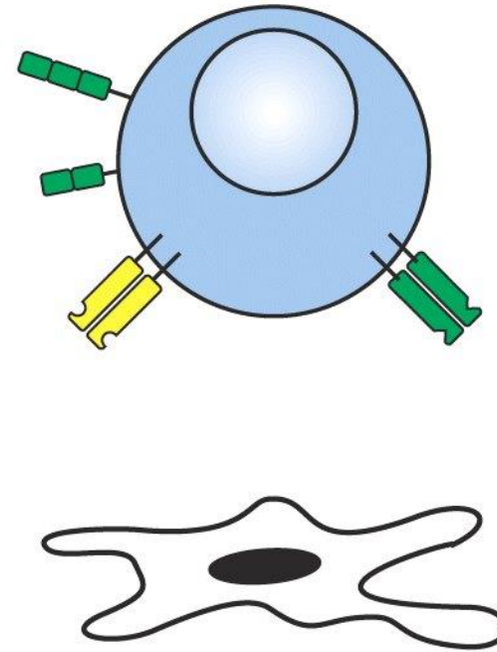


Figure 2-50 part 2 of 2 Immunobiology, 6/e. (© Garland Science 2005)



Tumour antigens against which an adaptive
immune response can evolve

Examples for the presentation of tumour antigens

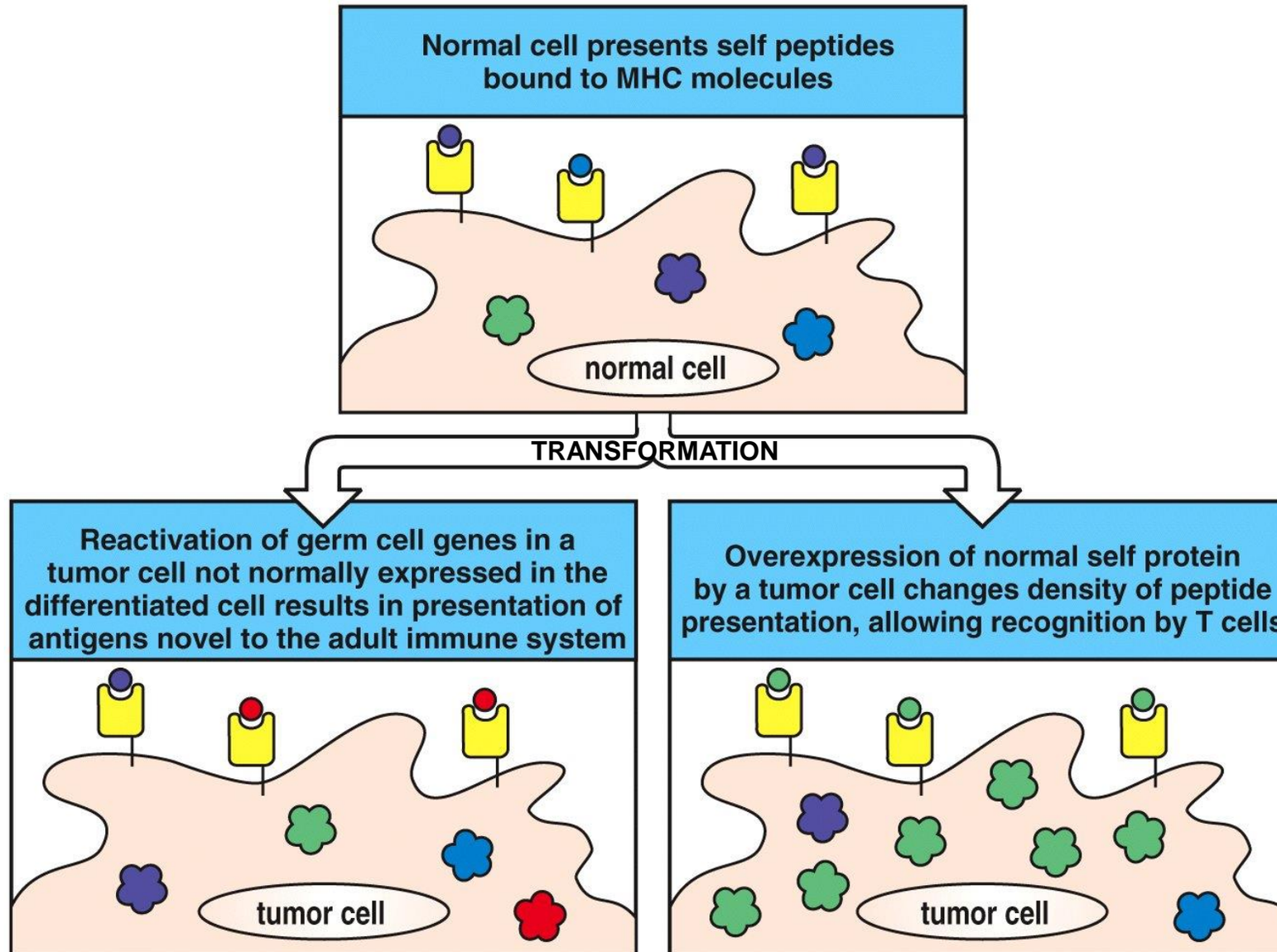


Figure 14-13 Immunobiology, 6/e. (© Garland Science 2005)

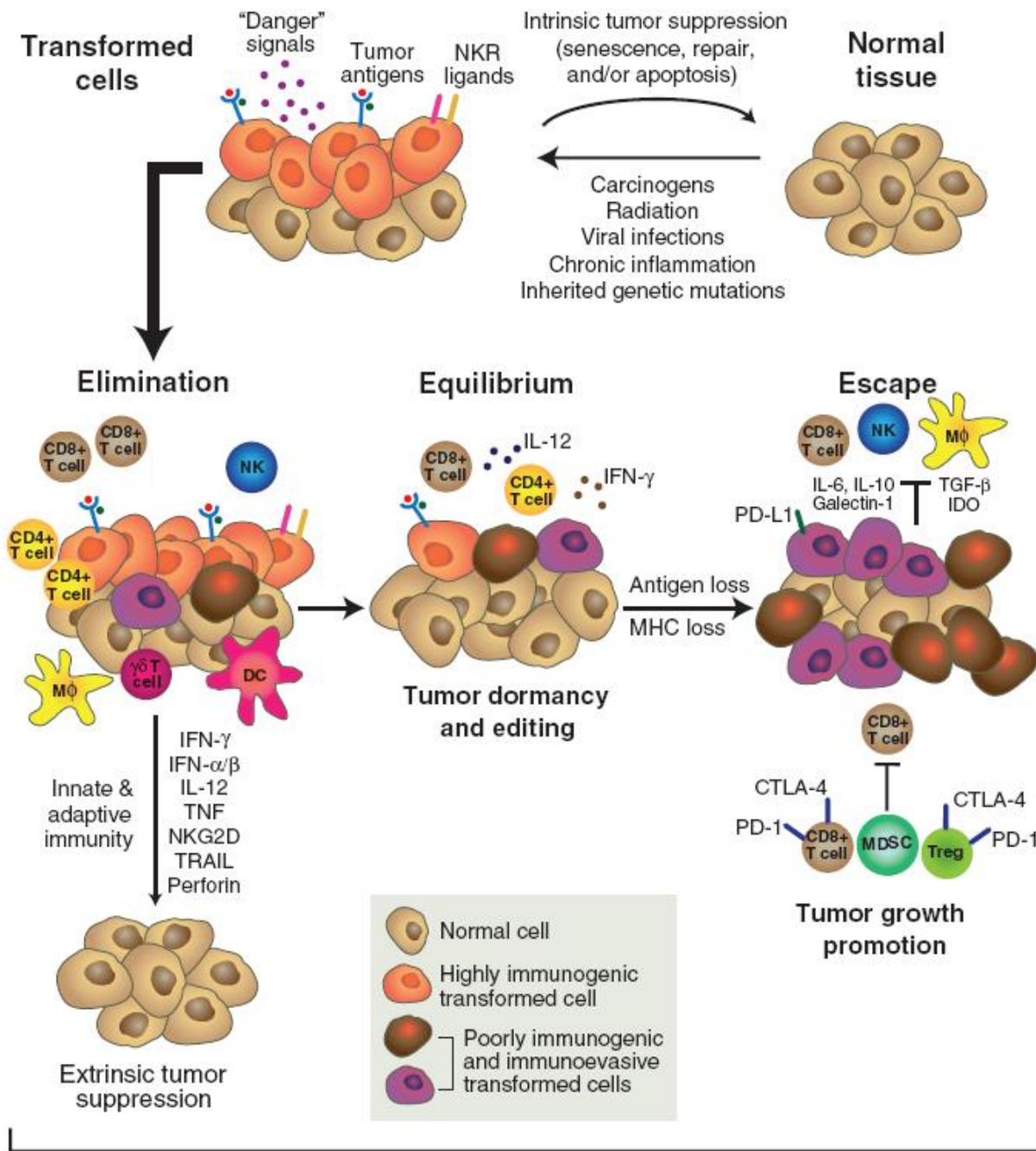
Potential tumor rejection antigens have a variety of origins

Class of antigen	Antigen	Nature of antigen	Tumor type
Differentiation	Tyrosinase	Enzyme in pathway of melanin synthesis	Melanoma
	Surface Ig	Specific antibody after gene rearrangements in B-cell clone	Lymphoma
Abnormal gene expression	HER-2/neu	Receptor tyrosine kinase	Breast Ovary
Abnormal post-translational modification	MUC-1	Underglycosylated mucin	Breast Pancreas
Oncoviral protein	HPV type 16, E6 and E7 proteins	Viral transforming gene products	Cervical carcinoma
Tumor-specific mutated oncogene or tumor suppressor	Cyclin-dependent kinase 4	Cell-cycle regulator	Melanoma
	β -Catenin	Relay in signal transduction pathway	Melanoma
	Caspase-8	Regulator of apoptosis	Squamous cell carcinoma
Germ cell	MAGE-1 MAGE-3	Normal testicular proteins	Melanoma Breast Glioma

MAGE=melanoma-associated antigen

HER-2= Receptor tyrosine-protein kinase erbB-2

MUC= Mucin



Cancer Immunoeediting

Mechanisms involved in the immune escape of tumor cells

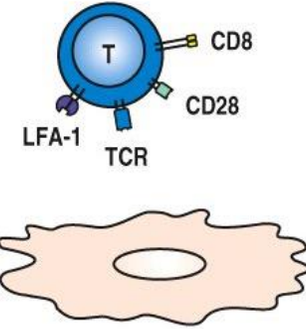
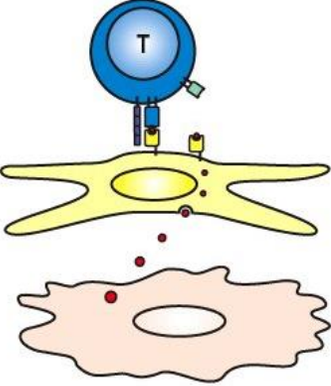
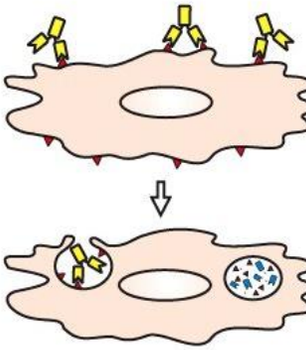
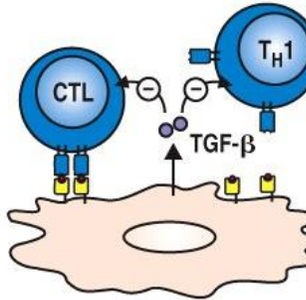
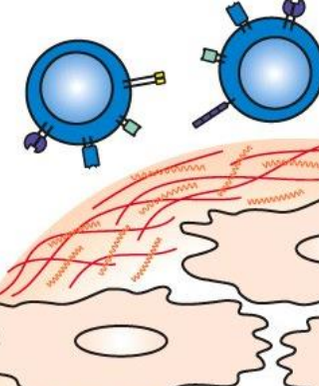
Low immunogenicity	Tumor treated as self antigen	Antigenic modulation	Tumor-induced immune suppression	Tumor-induced privileged site
<p>No peptide:MHC ligand No adhesion molecules No co-stimulatory molecules</p>	<p>Tumor antigens taken up and presented by APCs in absence of co-stimulation tolerize T cells</p>	<p>Antibody against tumor cell-surface antigens can induce endocytosis and degradation of the antigen. Immune selection of antigen-loss variants</p>	<p>Factors (eg, TGF-β) secreted by tumor cells inhibit T cells directly</p>	<p>Factors secreted by tumor cells create a physical barrier to the immune system</p>
				

Figure 14-14 Immunobiology, 6/e. (© Garland Science 2005)

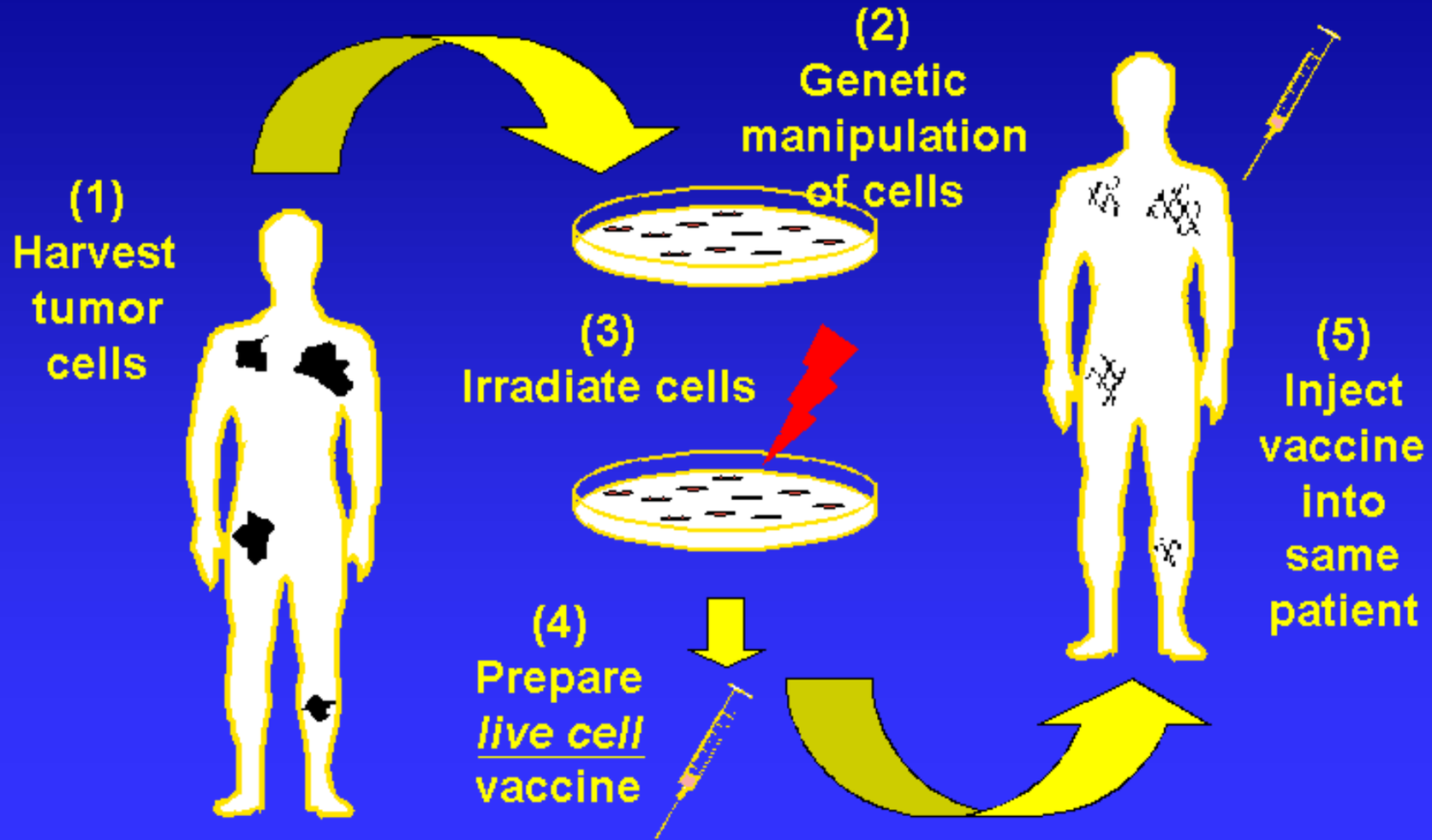


How can the immune system be activated so that
it actively fights tumor cells?

Immunization with tumor antigens

Immunization with gene-transfected tumor cells

Recombinant (DNA) Cell Vaccines



Immunization with gene-transfected tumor cells

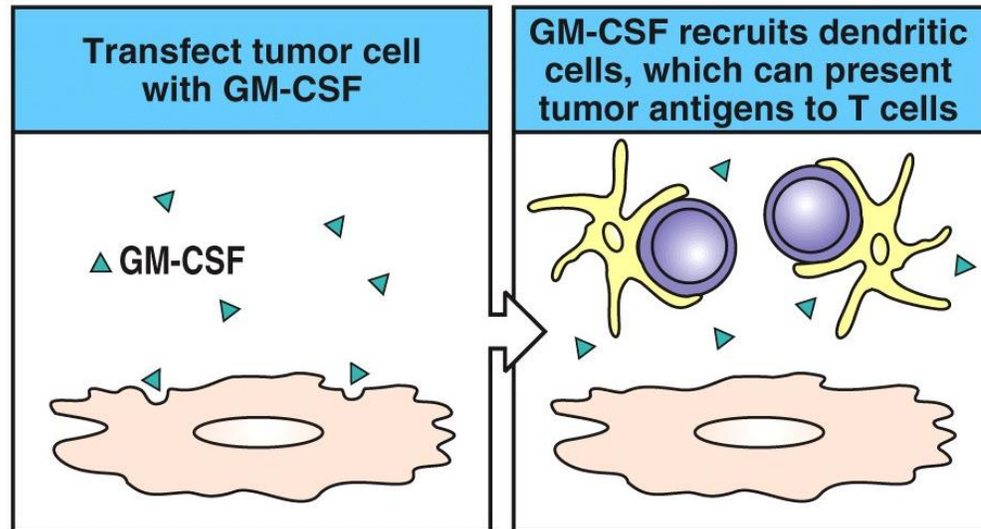


Figure 14-20 part 2 of 3 Immunobiology, 6/e. (© Garland Science 2005)

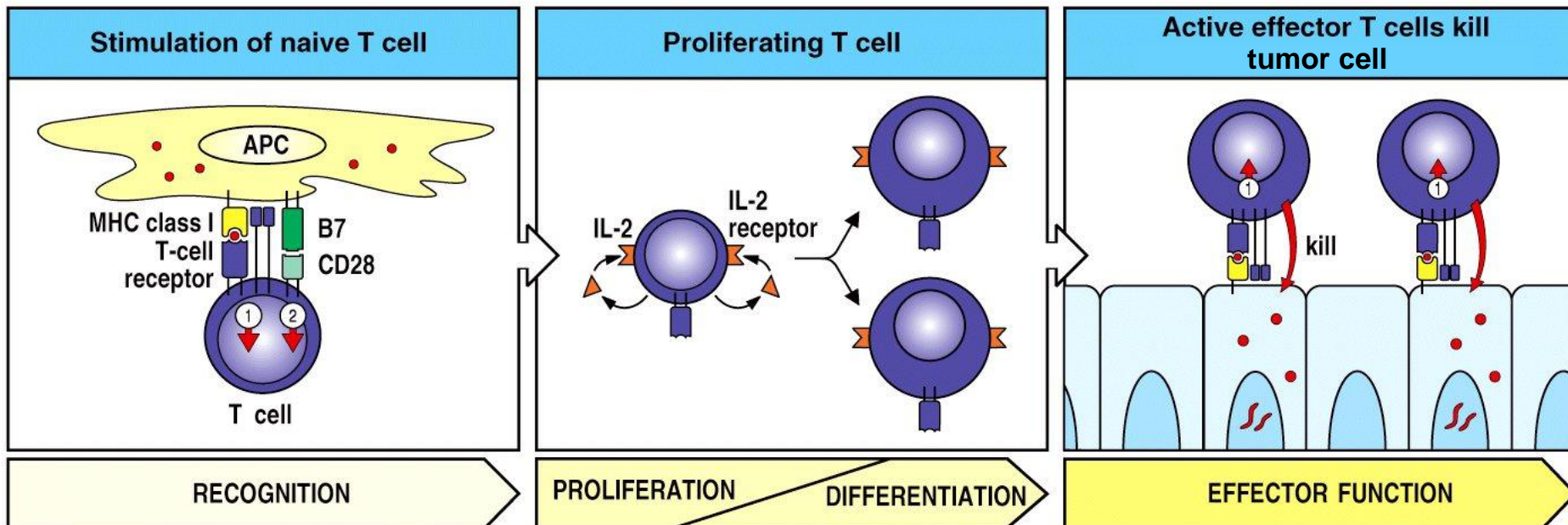
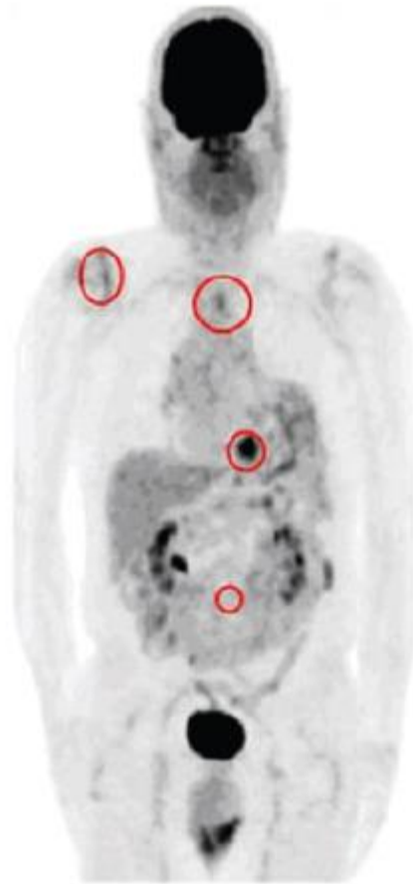
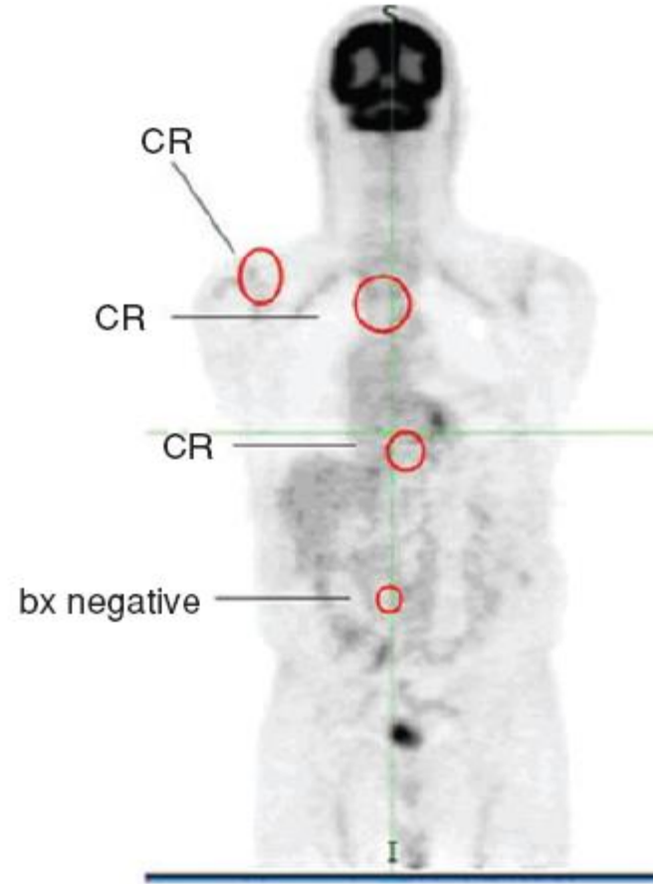


Figure 8-22 Immunobiology, 6/e. (© Garland Science 2005)

"Complete response" through vaccination in a melanoma patient



07/07/08



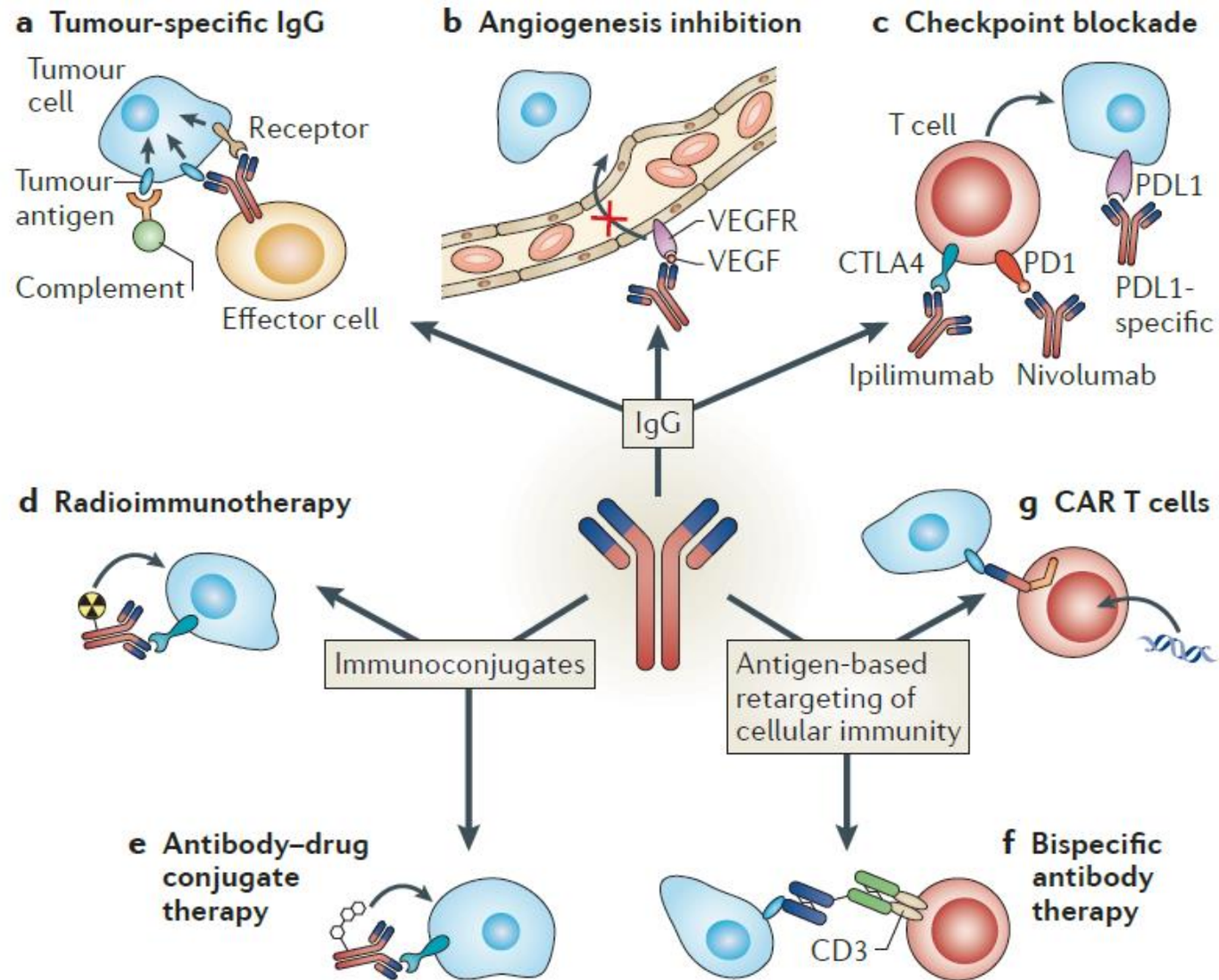
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How can the immune system be activated so that it actively fights tumor cells?

Use of antibodies and tumor antigen specific immune cells

Use of antibodies (conjugates) for the treatment of tumours



NK cells recognise tumour cells: mechanism 2

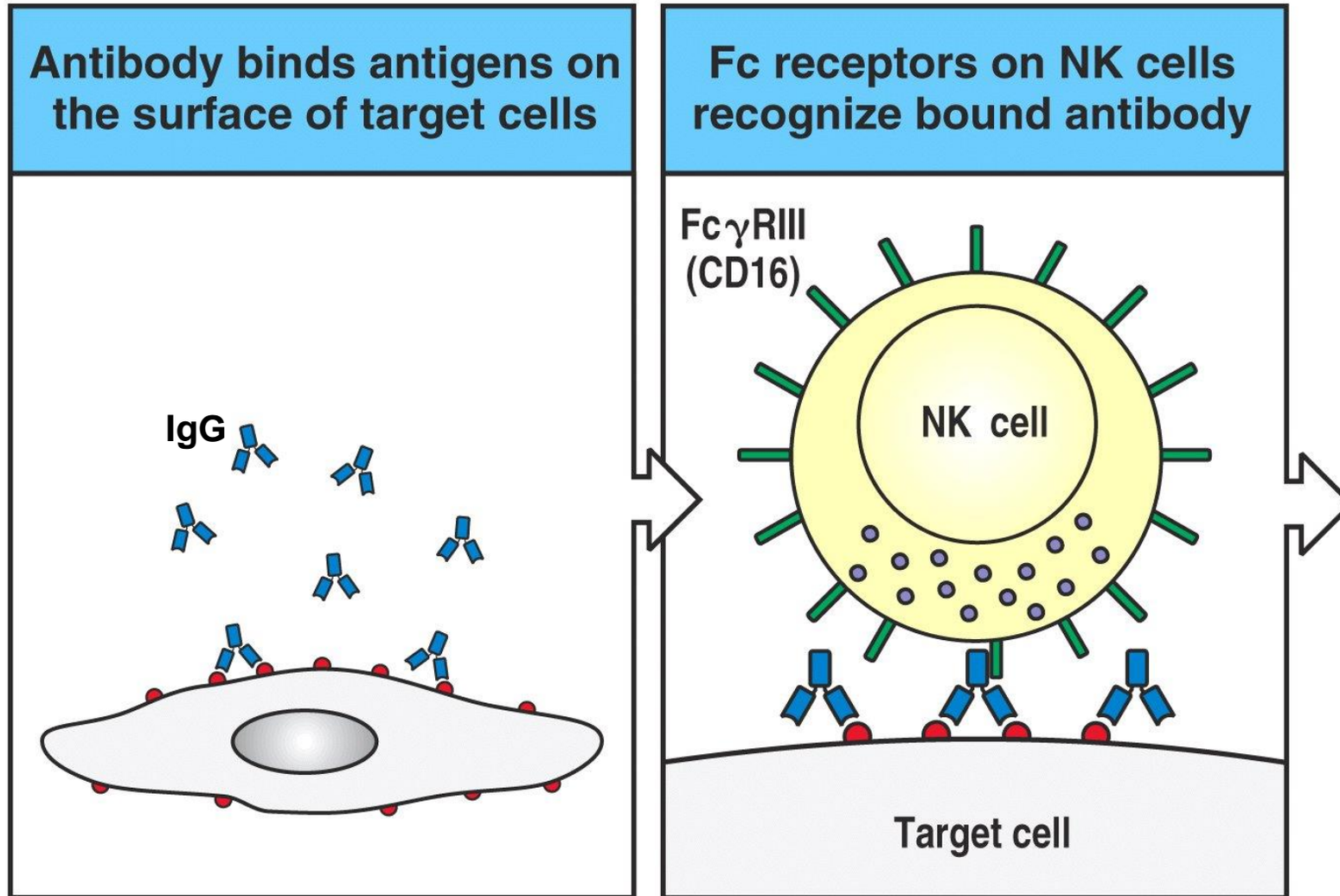
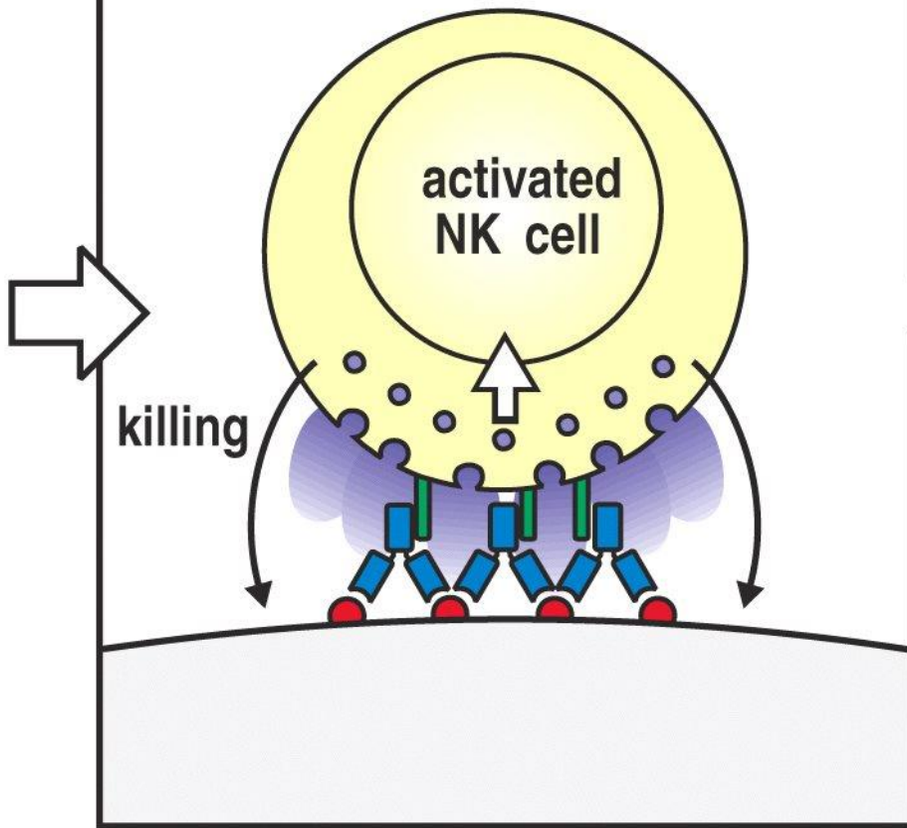


Figure 9-34 part 1 of 2 Immunobiology, 6/e. (© Garland Science 2005)

Cross-linking of Fc receptors signals the NK cell to kill the target cell



Target cell dies by apoptosis



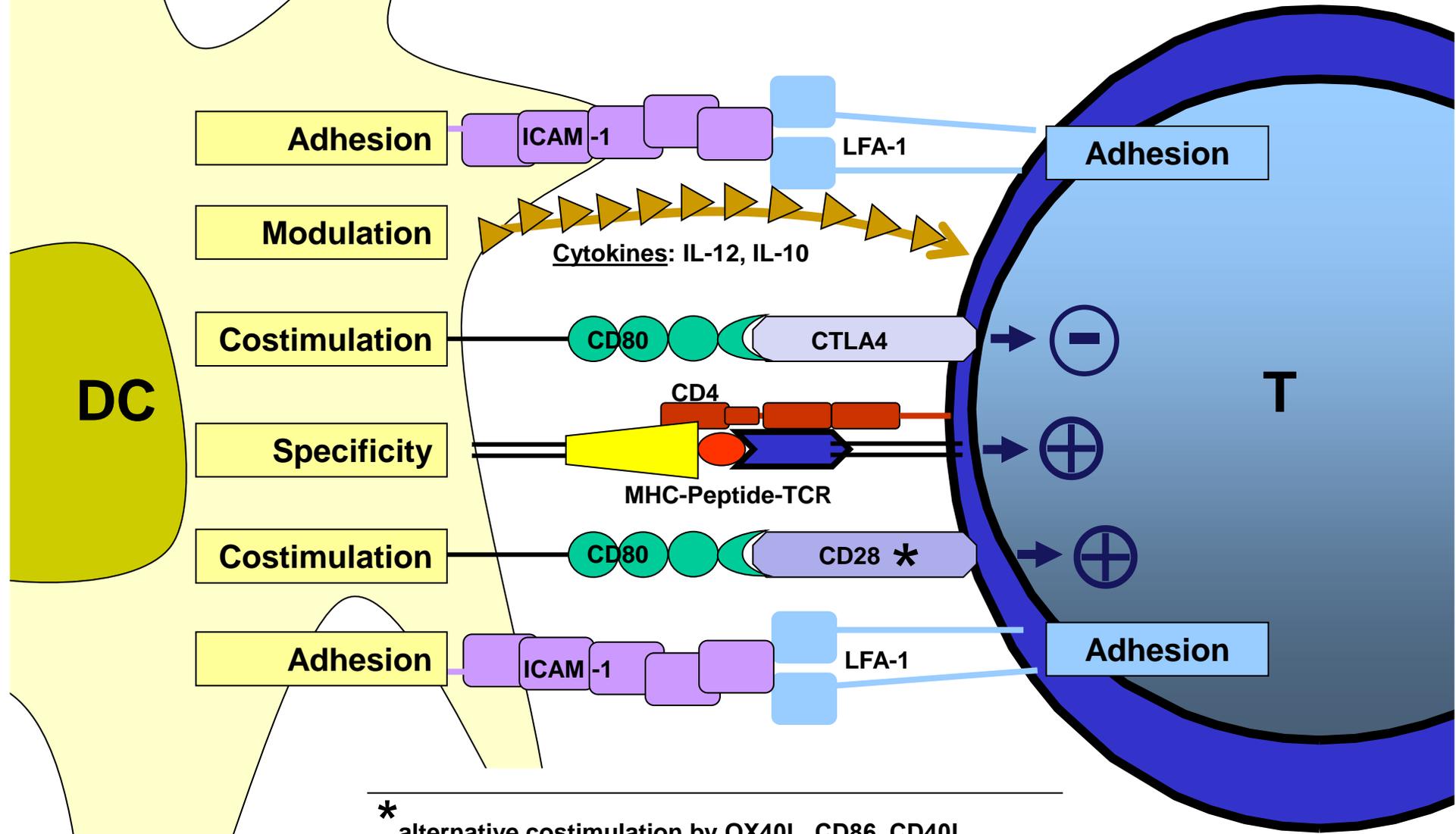
Figure 9-34 part 2 of 2 Immunobiology, 6/e. (© Garland Science 2005)



Use of antibodies as Checkpoint inhibitors

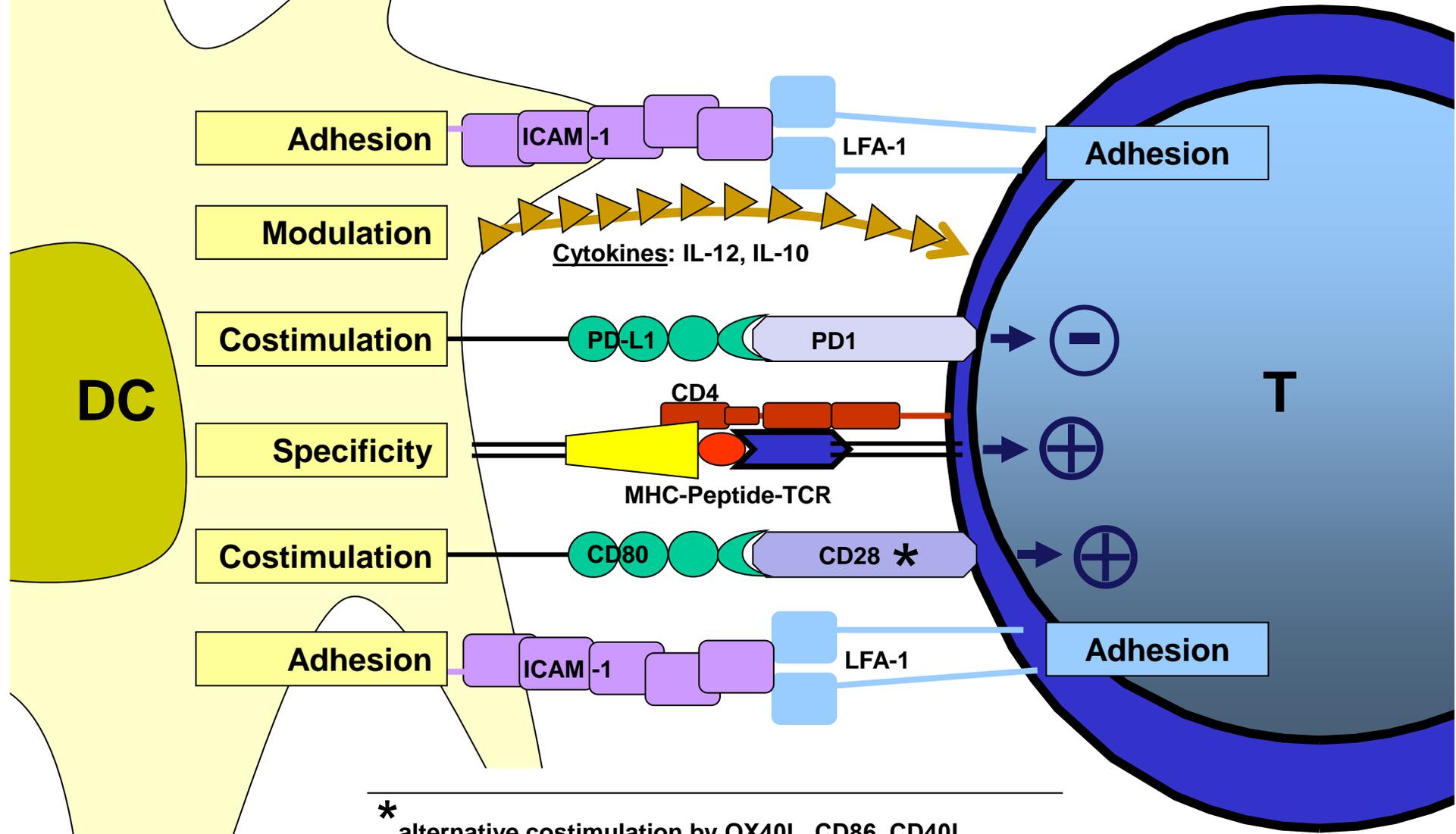
Blocking the interaction of:
CD80 --- CTLA4
PDL1 --- PD1

The Immunological Synapse

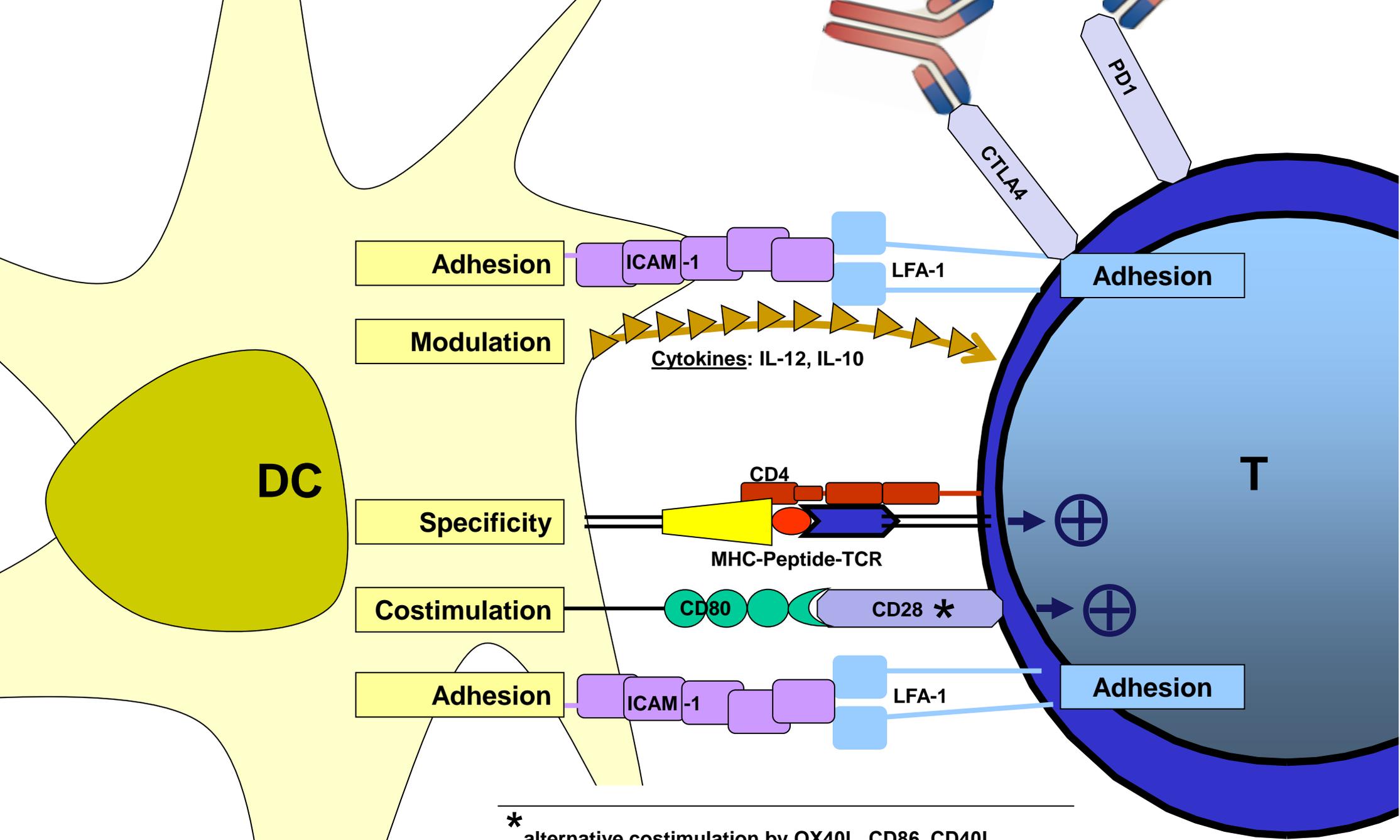


* alternative costimulation by OX40L, CD86, CD40L....

The Immunological Synapse



* alternative costimulation by OX40L, CD86, CD40L....



* alternative costimulation by OX40L, CD86, CD40L....

Treatment with anti-PD1 without anti-CTLA-4

Drug	Condition	Treatment regimen	Treatment in control group	Objective response rate	Complete response rates	Overall survival (months)	Progression-free survival (months)	Grade 3-5 adverse events	Participants treated (and controls)
Nivolumab (IgG4a)	Melanoma (stage III/IV)	3 mg/kg/2 weeks	(vs combination therapy)	43.7%	8.9%	n/a	6.9	16.3%	316
	Renal cell carcinoma (metastatic)	3 mg/kg/2 weeks	10 mg/day Everolimus	25% (4% control)	1% (<1% control)	25.0 (19.6 control)	4.6 (4.4 control)	19% (27% control)	406 (397 control)
	Hodgkin's lymphoma (relapsed/refractory)	3 mg/kg/2 weeks	n/a	87%	17%	n/a	86% at 24 weeks	22%	23
	Squamous-cell carcinoma of the head and neck (recurrent)	3 mg/kg/2 weeks	Single-agent systemic therapy (methotrexate, docetaxel, or cetuximab)	13.3% (5.8% control)	2.5% (0.8% control)	36.0%/1 year (16.6% control)	19.7% at 6 months (9.9% control)	13.1% (35.1%)	240 (121 control)
	Non-small cell lung cancer	3 mg/kg/2 weeks	Docetaxel	19% (12% control)	1% (<1% control)	12.2 (9.4 control)	2.3 (4.2 control)	10% (54% control)	292 (290 control)
			Docetaxel	20% (9% control)	1% (0% control)	9.2 (6 control)	3.5 (2.8 control)	7% (55% control)	135 (137 control)
	Ovarian cancer (platinum-resistant)	1 or 3 mg/kg/2 weeks	n/a	15%	10%	20	3.5	40%	20
Pembrolizumab (IgG4a)	Melanoma (stage III/IV)	10 mg/2 weeks or 3 weeks	(vs ipilimumab)	33.7-32.9%	5.0-6.1%	n/a	5.5-4.1	13.3-10.1%	279-277
	Merkel cell carcinoma	2 mg/kg/3 weeks	n/a	56%	16%	n/a	65% at 6 months	15%	26
	Non-small cell lung cancer	2 mg/kg/3 weeks 10 mg/kg/3 weeks 10 mg/kg/2 weeks	n/a	19.4%	n/a	12	3.7	9.5%	495
		200 mg/2 weeks (PD-L1 + patients only)	Platinum-based chemotherapy	44.8 (27.8% control)	n/a	80.2% at 6 months (72.4% control)	10.3 (6 control)	26.6% (53.3% control)	154 (154 control)
		2 or 10 mg/kg/3 weeks (PD-L1 + patients only)	Docetaxel	18/18% (9% control)	0/0% (0% control)	10.4/12.7 (8.5 control)	3.9/4.0 (4.0 control)	13/16% (35% control)	345/346 (343 control)

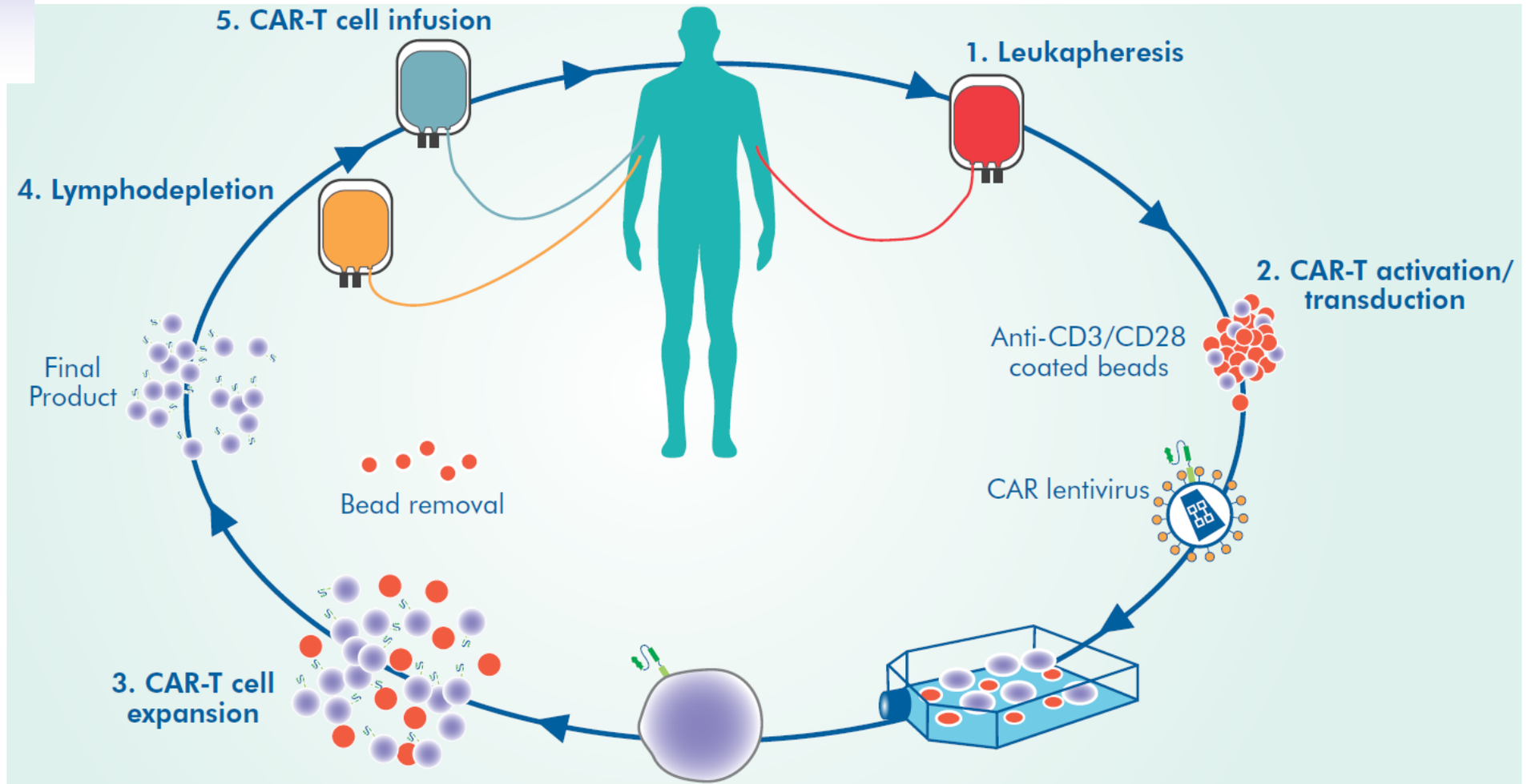
Treatment with anti-CTLA4 +/- Anti PD1

	Drug	Condition	Treatment regimen	Treatment in control group	Objective response rate	Complete response rates	Overall survival (months)	Progression-free survival (months)	Grade 3-5 adverse events	Participants treated (and controls)
CTLA-4	Ipilimumab (IgG1)	Melanoma (stage III/IV)	10 mg/kg plus decarbazine	Decarbazine alone	15.2% (10.3% control)	1.6% (0.8% control)	11.2 (9.1 control)	n/a	56.3% (27.5%)	250 (252 control)
			3 mg/kg/3 weeks	(vs Pembrolizumab)	11.9%	1.4%	n/a	2.8	19.9%	278 (315 control)
			3 mg/kg/3 weeks	(vs combination with nivolumab)	19%	2.2%	n/a	2.9	27.3%	311
	Tremelimumab (IgG2)	Melanoma (stage III/IV)	15 mg/kg/90 days	chemotherapy (temozolomide or dacarbazine)	10.7% (9.8% control)	3% (2% control)	12.6% (10.7 control)	20.3% at 6 months (18.1% control)	52% (37% control)	328 (327 control)
CTLA4 +PD1	Nivolumab + Ipilimumab	Melanoma (stage III/IV)	3 mg/kg/2 weeks Nivolumab 3 mg/kg/3 weeks Ipilimumab	(vs single)	57.6%	11.5%	n/a	11.5	55%	314
		Non-small cell lung cancer	Nivo + Ipi: 1 + 3 or 3 + 1 mg/ml	Nivolumab alone	23/19% (10% control)	2/0% (0%)	7.7/6 (4.4)	2.6/1.4 (1.4 control)	30/19% (13% control)	61/54 (98 control)

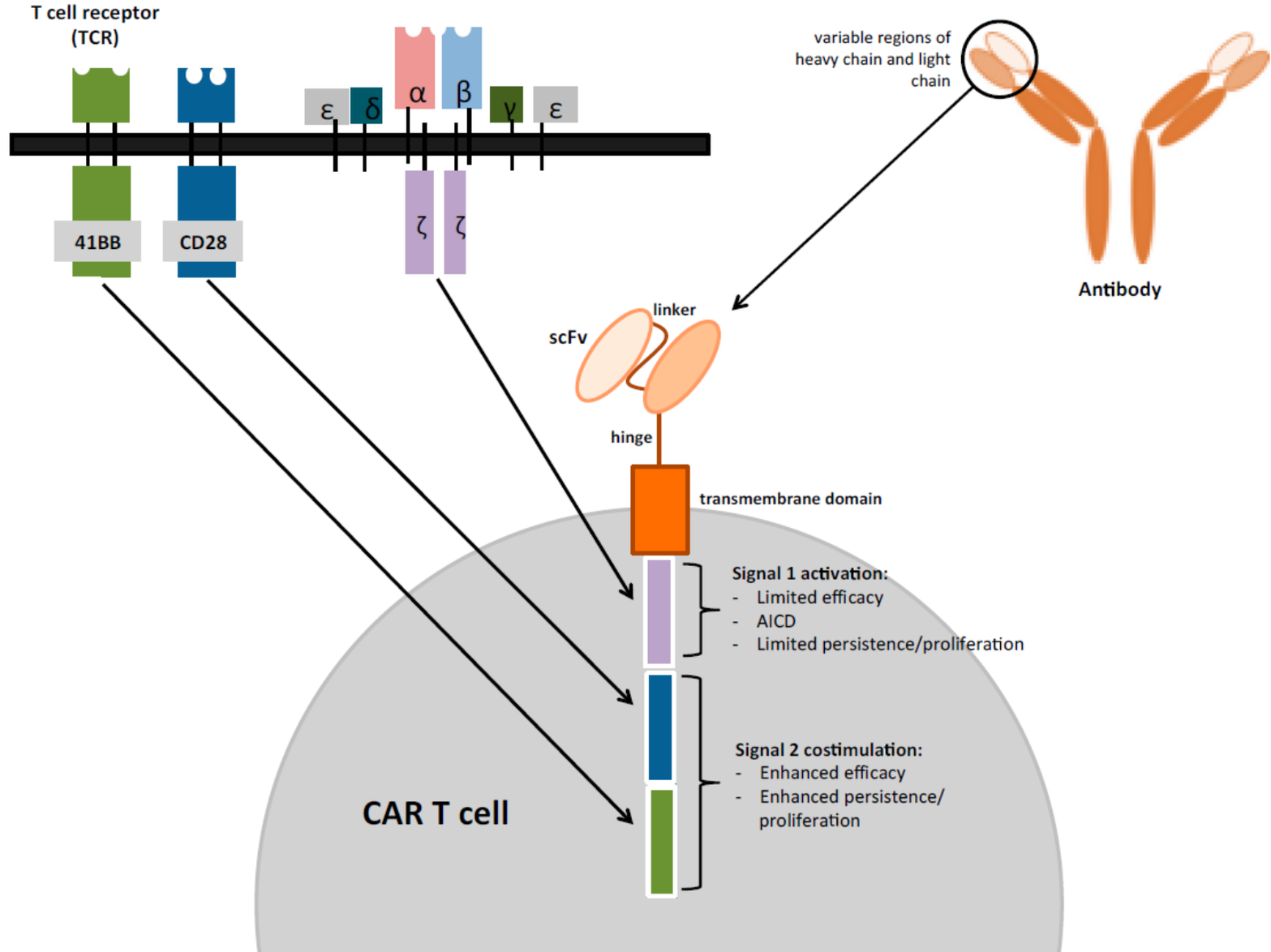


Chimeric Antigen Receptor (CAR)-T Cell Immunotherapy

Chimeric Antigen Receptor (CAR) T-cells

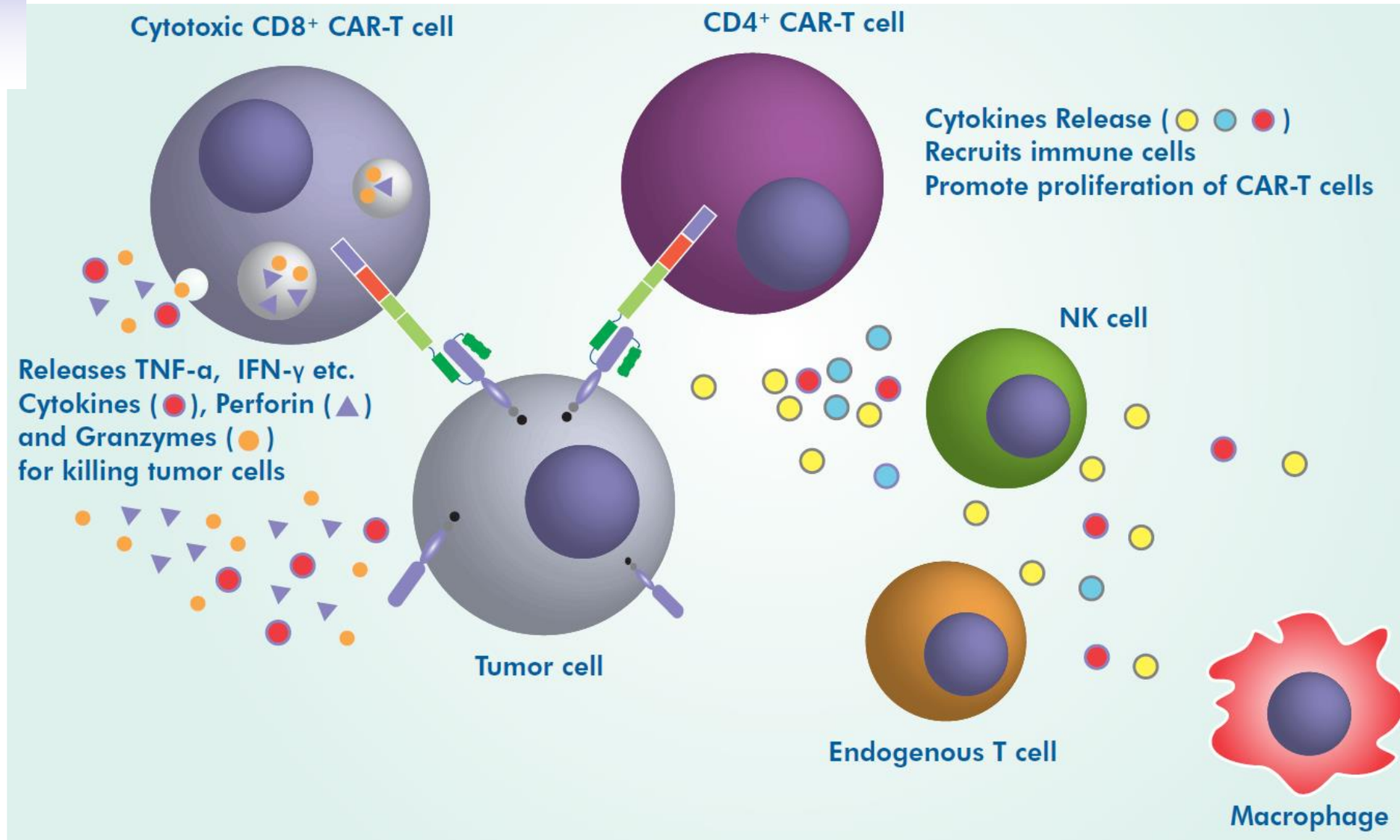


Chimeric Antigen Receptor (CAR) specific T-cells



scFV: single chain variable fragment

Effector mechanism of CAR T-cells



Key targets for CAR-T cell therapy and established clinical results

Disease	Target	Effector	Efficacy	Toxicity	Citation
B-cell Acute Leukemia (pediatric-young adult)	CD19	CTL019 (4-1BB)	CR 81% [61/75] RFS (12 months) 59%	CRS 77% Gr. 4 CRS 25% Neurotoxicity 40%	Maude <i>et al.</i> , 2018 [NEJM; PMID: 29385370]
B-cell Acute Leukemia (adult)	CD19	JCAR017 (CD4/CD8; 4-1BB)	CR 93% [27/29] RFS: n/a	CRS 83% Severe CRS (Doesn't Specify) 23% Neurotoxicity 50%	Turtle <i>et al.</i> , 2017 [JCI; PMID: 27111235]
B-cell Lymphoma (DLBCL, FL)	CD19	CTL019 (4-1BB)	CR 57% [16/28] PFS (28.6 months) 57%	CRS 57% Severe CRS: (Grade 3 or higher): 18% Neurotoxicity 39%	Schuster <i>et al.</i> , 2017 [NEJM; PMID: 9226764]
B-cell Chronic Leukemia	CD19	CTL019 (4-1BB)	CR 29% [4/14] PFS (18 months) 29%	CRS 64% Severe CRS 29% (43%) Neurotoxicity 37% [6/16]	Porter <i>et al.</i> , 2015 [STM; PMID: 26333935]
B-cell Acute Leukemia (pediatric-young adult)	CD22	CAR-T-22 (4-1BB)	CR 57% [12/21] PFS (6 months) 50%	CRS 76% Severe CRS 0% Neurotoxicity 37% [6/16]	Fry <i>et al.</i> 2017, [Nat Med; PMID: 29155426]
Multiple Myeloma	BCMA	BCMA-CAR-T (4-1BB)	ORR 100% [15/15] sCR+VGPR 73% [11/15]	CRS 78% [14/18] Severe CRS 11% [2/18]	Bergdeja <i>et al.</i> , 2017 [ASH; Paper 107984, Session: 653]

CR = complete remission; PFS = progression-free survival; DFS = disease-free survival; CRS = cytokine release syndrome; n/a = not applicable sCR = stringent CR; VGPR = very good partial remission; ORR = overall response rate