

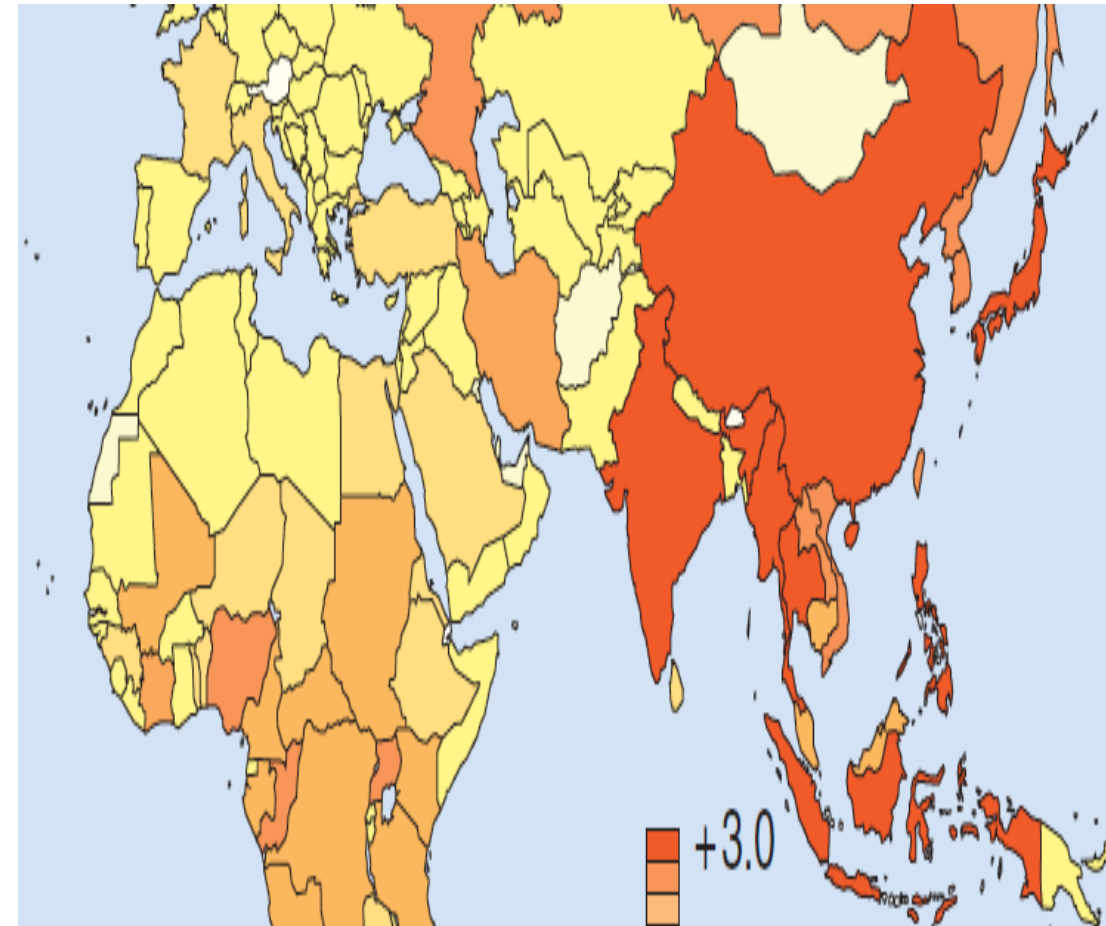
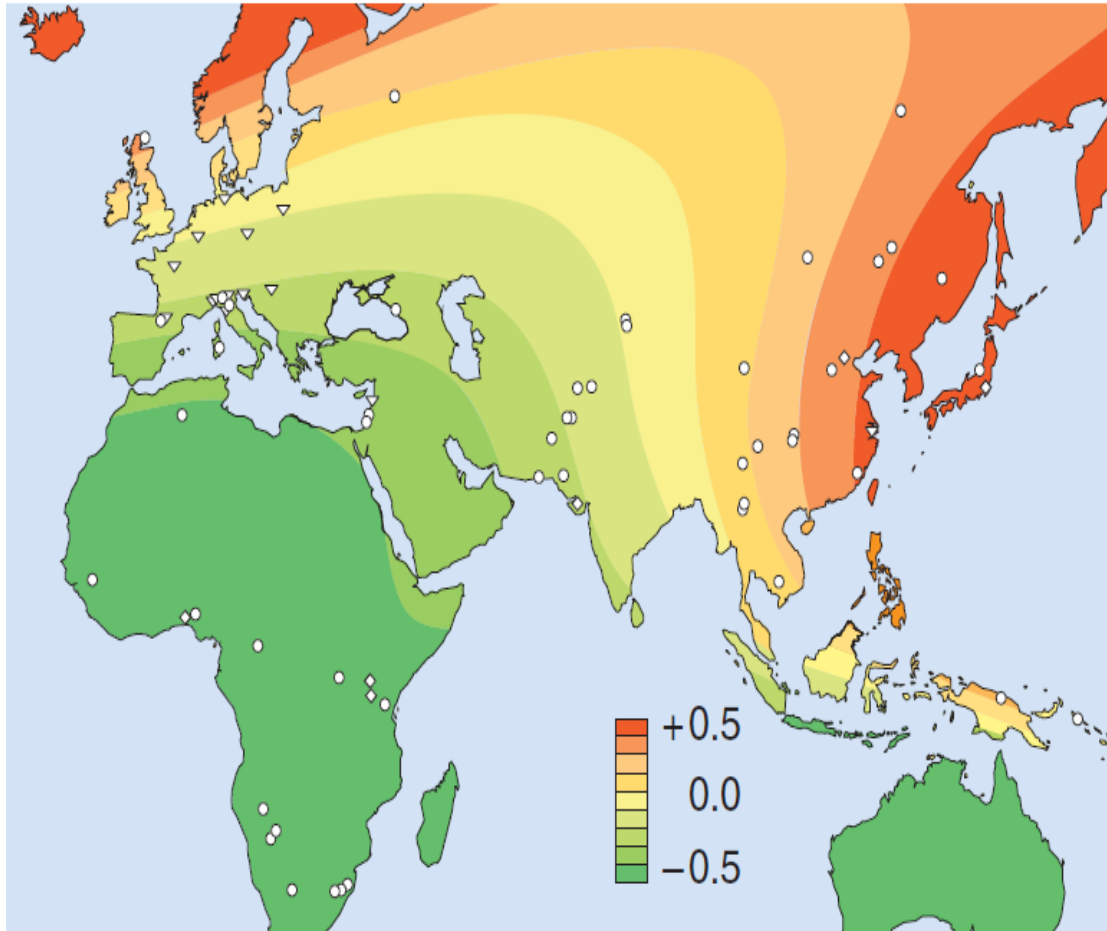
# New Research In IgA Nephropathy

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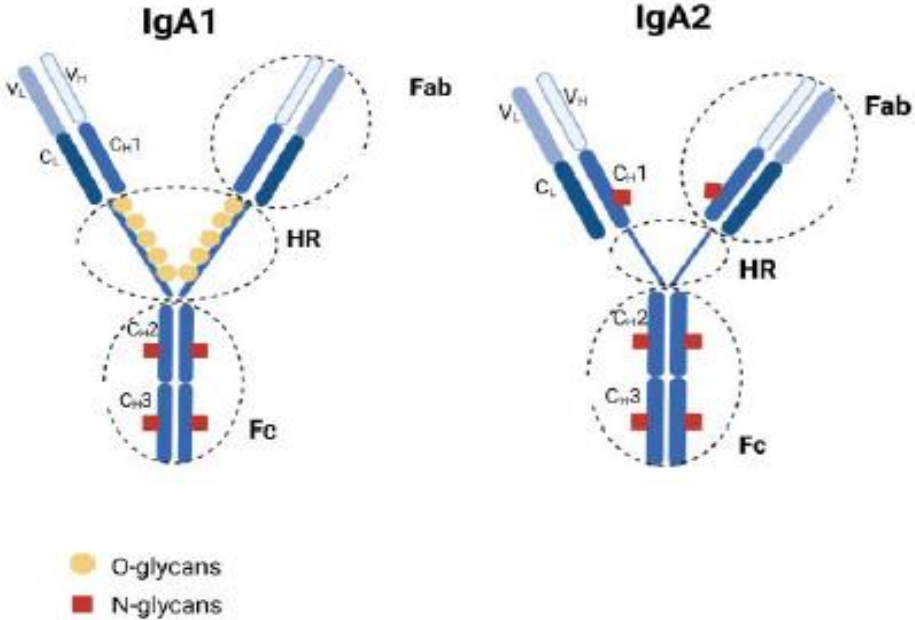
# Introduction

- IgA nephropathy (IgAN) is the most common primary glomerular disease with a marked heterogeneity in its clinical and pathological features.
- Most common cause of kidney failure in Asia, has lower prevalence in Europe, and is very infrequent among populations of African.
- Among patients with reduced renal function and proteinuria  $>1\text{g}/24\text{ h}$ , outcomes remain poor.
- Up to 50% of such patients will progress to ESRD over 10 years.

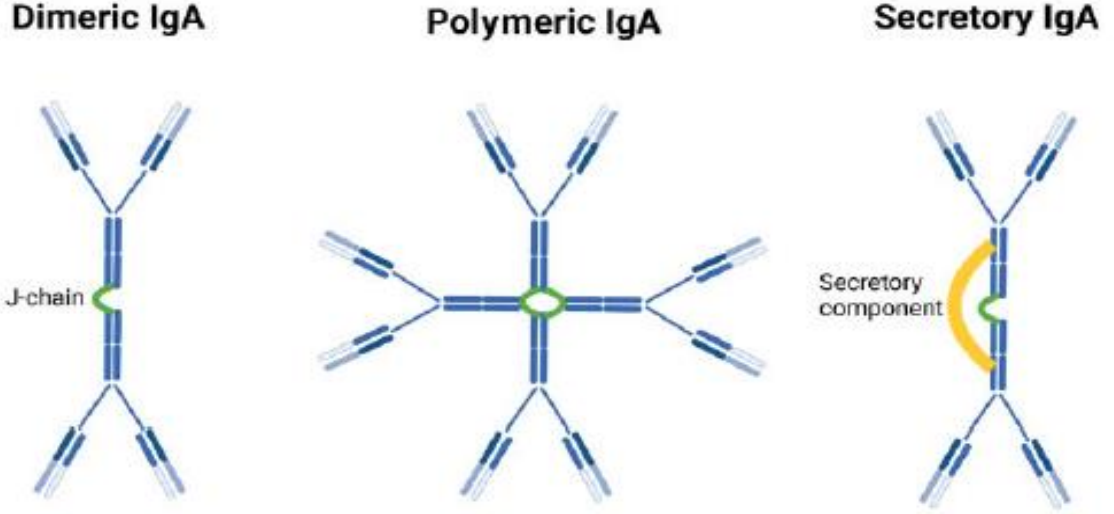
# Geospatial pattern of genetic risk for immunoglobulin A (IgA) nephropathy and worldwide map of helminth diversity.



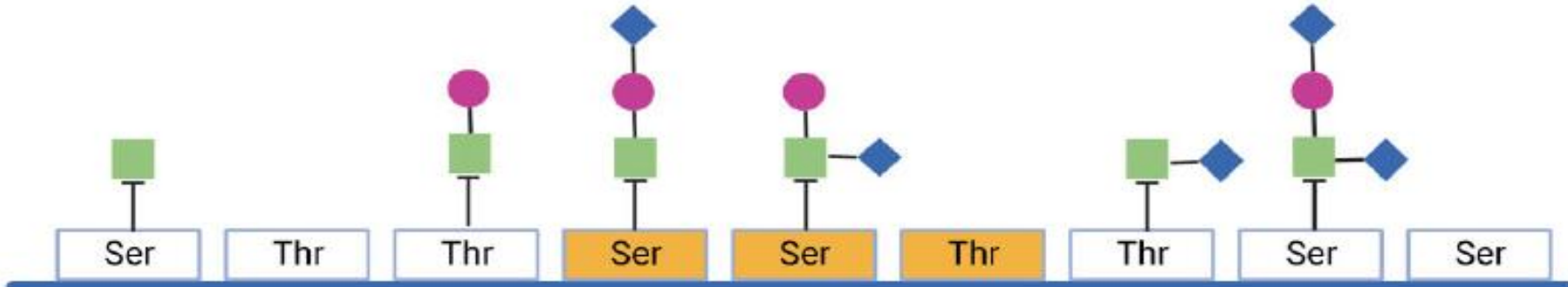
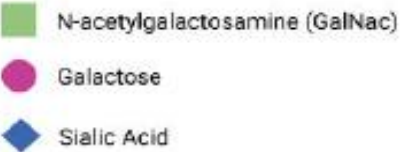
Hypothesis :higher genetic risk of IgAN in Asia represents an untoward consequence of protective adaptation to worm infections, the process that has likely occurred over thousands of years of human-parasite coevolution.



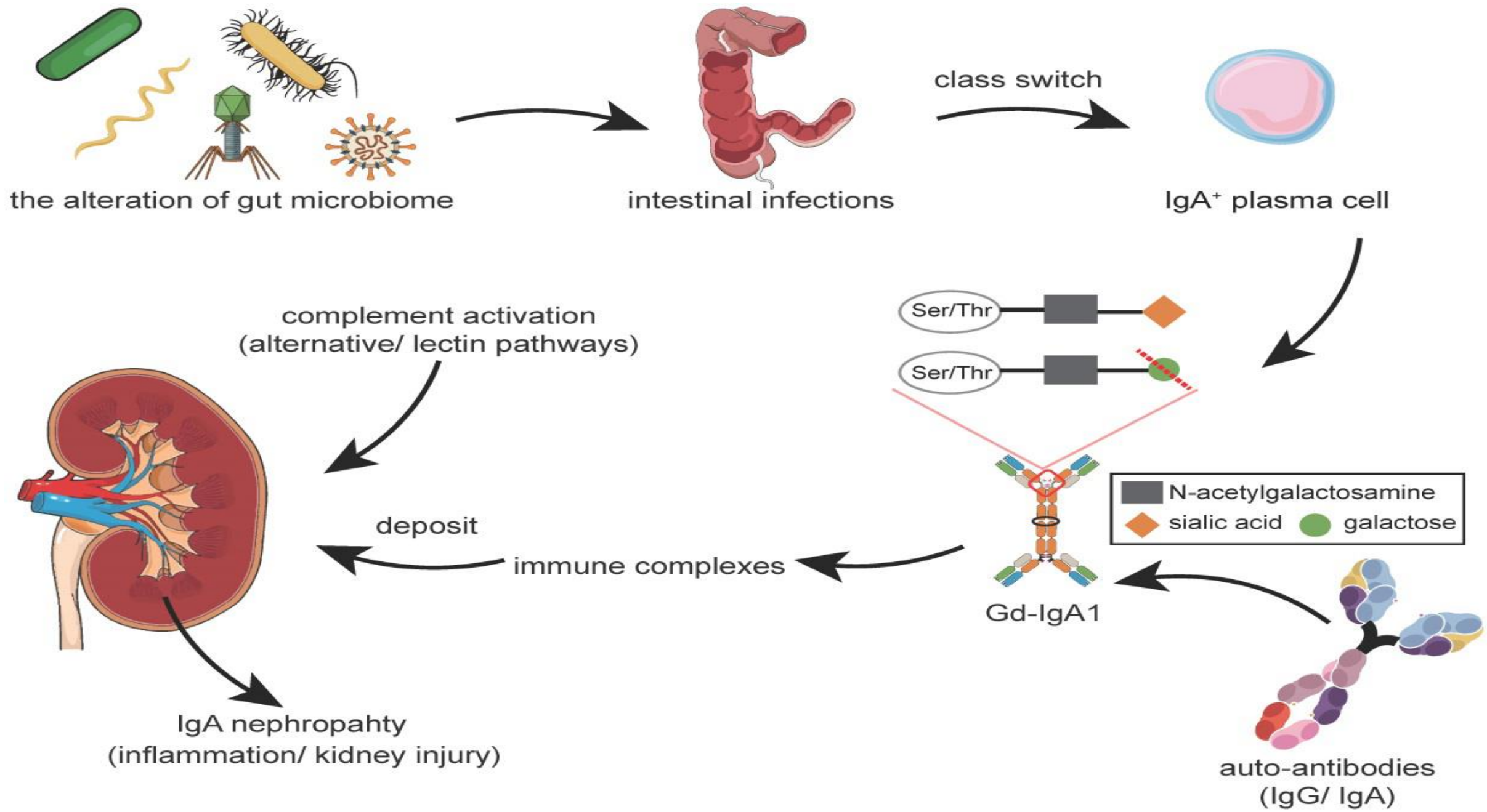
**A**



**B**

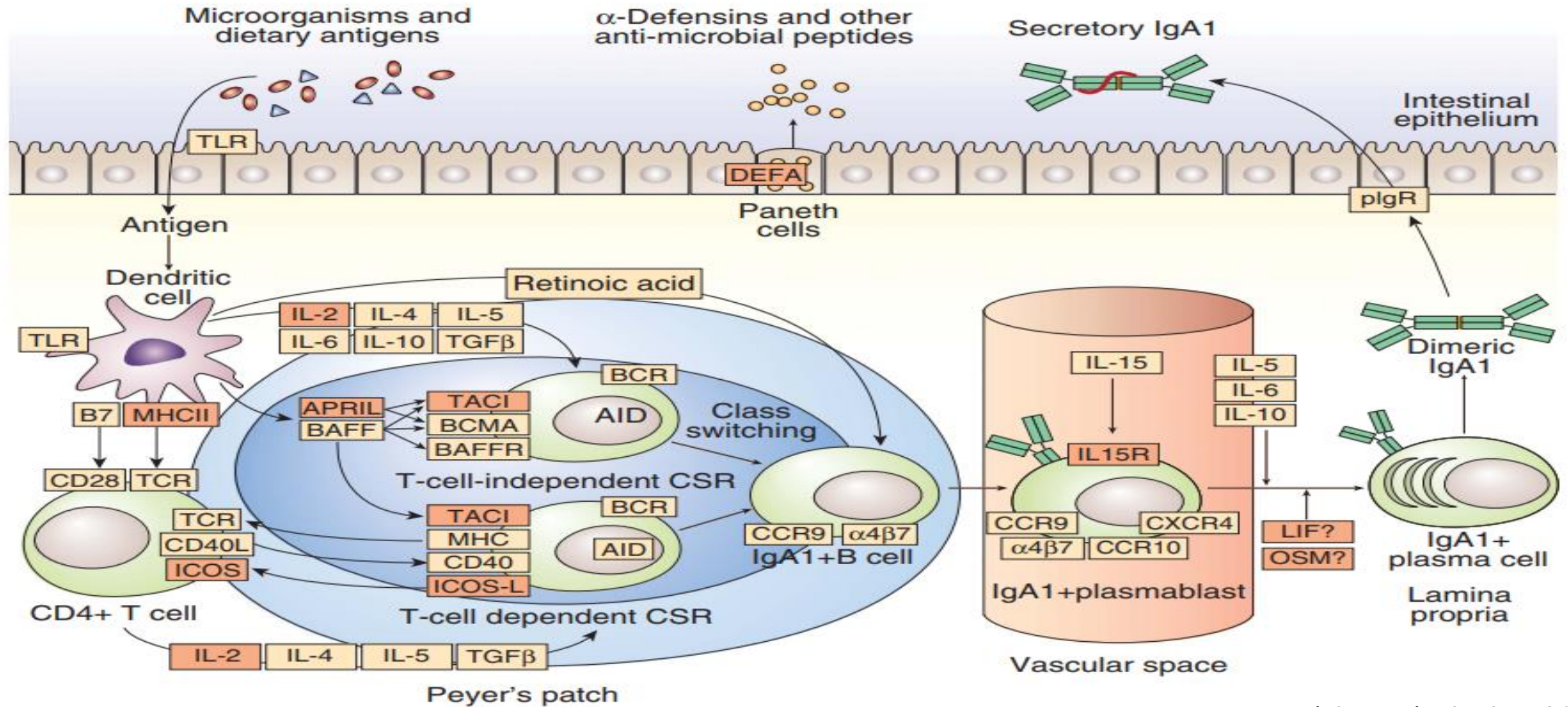


**C**

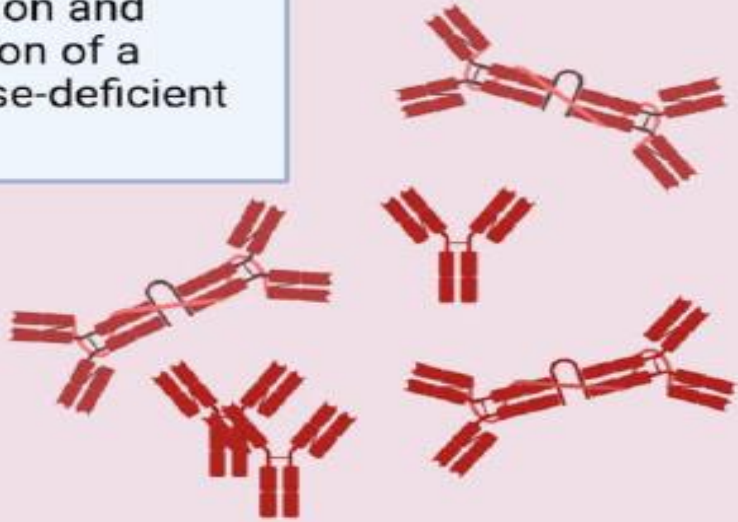




# Genetic hits to the Intestinal Immune Network for IgA Production



Hit 1: Increased production and circulation of a galactose-deficient IgA1



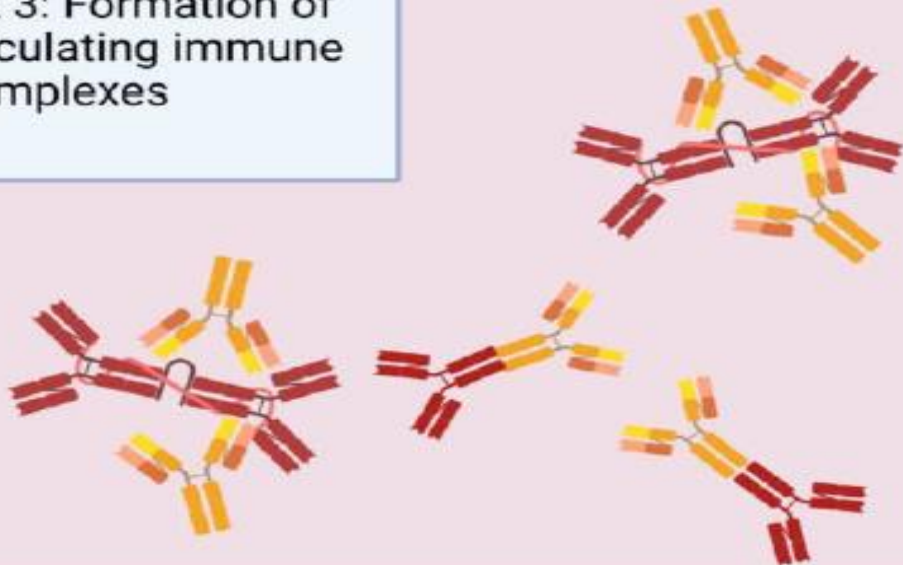
**A**

Hit 2: Production of auto-antibodies against galactose deficient IgA1



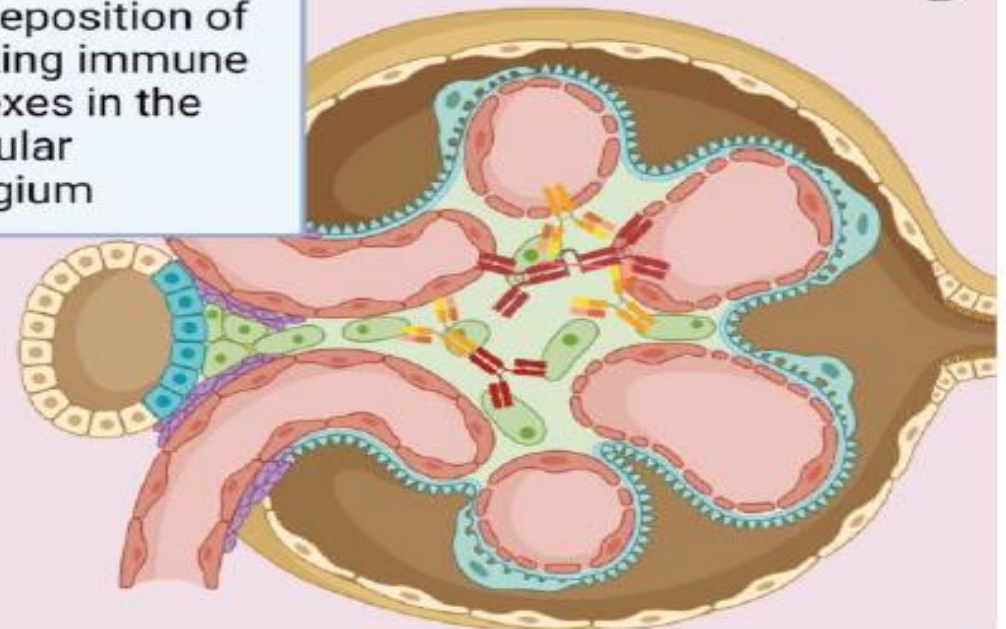
**B**

Hit 3: Formation of circulating immune complexes



**C**

Hit 4: Deposition of circulating immune complexes in the glomerular mesangium



**D**

# Risk Factors to disease progression

- Proteinuria > 1gram/ 24 hours
- Hypertension
- Reduced GFR
- Microscopic Hematuria
- Histologic Predictors ( MESTC Classification)



## Considerations for the prognostication of primary IgAN:

- Clinical and histologic data at the time of biopsy can be used to risk stratify patients.
- The International IgAN Prediction Tool is a valuable resource to quantify risk of progression and inform shared decision-making with patients.
- **Calculate by QxMD**
- **No validated prognostic serum or urine biomarkers** for IgAN other than eGFR and proteinuria.

  **International IgAN Prediction Tool  
at biopsy - Adults**

Determine prognosis in adults with IgA nephropathy

**Risk Assessment 1-2 year after biopsy**  
**Hematuria and crescent not included**  
**No recommendation for treatment**  
**based on calculator**  
**Needs more work on different ethnicity**  
**New adjustment for children**

## **Risk factors for progression**

- Estimated GFR
- Blood pressure
- Proteinuria
- Age
- Race/ethnicity (White, Japanese, Chinese, or other)
- Prior use of an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB)
- Oxford classification of IgAN MEST histology scores
- Immunosuppression use at or prior to biopsy

# Proteinuria Reduction as a Surrogate End Point in Trials of IgA Nephropathy

- A strong and consistent relationship between the level and duration of **proteinuria and loss of kidney function**.
- Trial-level analyses of data from 13 controlled trials (830 subjects) showed an association between treatment effects on percent reduction of **proteinuria** and treatment effects on a composite of time to **doubling of serum creatinine**, ESKD, or

Proteinuria reduction to under 1 g/d is a surrogate marker of improved kidney outcome in IgAN, and reduction to under 1 g/d is a reasonable treatment target. (KDIGO 2021)

## Supportive therapy in IgAN

### Blood pressure management

- target sitting systolic BP <120 mmHg
- preferred antihypertensives:
  - first choice: ACE inhibitors or ARBs (with dosage up-titration as tolerated) in all patients with proteinuria > 0.5 g/d; no combination therapy
  - non-dihydropyridine calcium channel blockers (e.g. verapamil, diltiazem)
  - aldosterone antagonists
  - beta blockers
- avoid dihydropyridine calcium-channel blockers (e.g. amlodipine, nifedipine)

### Dietary advices and fluid management

- restrict sodium intake to less than 2 g/d or 90 mmol/d and/or use diuretics
- control protein intake
- control fluid intake (less than 1.5 to 2 L/d)

### Lifestyle modifications

- quit smoking
- normalize body weight
- encourage regular endurance sports, avoid strenuous exercise

### Additional measures

- avoid NSAIDs
- avoid prolonged severe hyperkalemia
- consider hydroxychloroquine in proteinuric patients despite maximal dosage of RAS blocker
- SGLT-2 inhibitor (currently off-label; status 8/2021)



# High Risk of progression

- **Proteinuria >0.75–1 g/d** despite  $\geq$ 90 days of optimized supportive care.
- Insufficient evidence to support the use of the Oxford Classification MEST-C score in determining whether immunosuppression should be commenced in IgAN.
- Dynamic assessment of patient risk over time should be Performed
- Adverse treatment effects are more likely in patients with an eGFR <50 ml/min per 1.73 m<sup>2</sup>.

- Recommendation 2.3.1.1: We suggest that patients who remain at high risk of progressive CKD despite maximal supportive care be considered for a 6-month course of glucocorticoid therapy. The important risk of treatment-emergent toxicity must be discussed with patients, particularly those who have an eGFR <50 ml/min per 1.73 m<sup>2</sup> (2B).

Study	Medication	Start dose	Duration high dose	Taper	Total exposure
TESTING <sup>(1)</sup>	Methylprednisolone	0.6–0.8 mg/kg/d (per investigator), rounded to nearest 4 mg. Max 48 mg/d	2 months	8 mg/month	6–8 months
Manno <sup>(2)</sup>	Prednisone	1 mg/kg/d, max 75 mg/d	2 months	0.2 mg/kg/month	6 months
Lv <sup>(3)</sup>	Prednisone	0.8–1 mg/kg/d	8 weeks	5–10 mg/d every 2 weeks	8 months

Mycophenolate mofetil (MMF)	<p><b>Chinese patients</b> In those patients in whom glucocorticoids are being considered MMF may be used as a glucocorticoid-sparing agent</p>	In a single RCT conducted in China, MMF with low-dose glucocorticoids was noninferior to standard-dose glucocorticoids for the treatment of incident IgAN presenting with proliferative histologic lesions (E or C lesions with or without necrosis) on kidney biopsy and proteinuria >1.0 g/d. There were significantly fewer glucocorticoid-related side effects in the combination-therapy arm. <sup>(1, 5)</sup>
	<p><b>Non-Chinese patients</b> There is insufficient evidence to support the use of MMF</p>	In the RCTs of MMF in non-Chinese patients there was no evidence for efficacy of MMF monotherapy. <sup>(2-5)</sup>
Hydroxychloroquine	<p><b>Chinese patients</b> In those patients who remain at high risk of progression in spite of optimized supportive care</p>	In a small, short-term RCT conducted in China, hydroxychloroquine introduced to patients with proteinuria of 0.75–3.5 g/d despite optimized ACEi/ARB reduced proteinuria by 48% versus 10% in the placebo group at 6 months. <sup>(6)</sup>
	<p><b>Non-Chinese patients</b> There is insufficient evidence to support the use in those patients</p>	Hydroxychloroquine has not been evaluated in non-Chinese patients.



## A Randomized, Controlled Trial of Rituximab in IgA Nephropathy with Proteinuria and Renal Dysfunction

Richard A. Lafayette, Pietro A. Canetta, [...], and Fernando C. Fervenza

- Open label, multicenter study, 1-year follow-up, randomized
- **34 adult patients** proteinuria  $>1$  g/d, maintained on ACE or ARB, well controlled BP and  $eGFR < 90$  ml/min per  $1.73$  m<sup>2</sup>,
- Receive standard therapy or rituximab with standard therapy.
- **Rituximab did not alter the level of proteinuria** compared with that at baseline or in the control group.
- Serum levels of galactose-deficient IgA1 or antibodies against galactose-deficient IgA1 did not change however effectively depleted B cell.

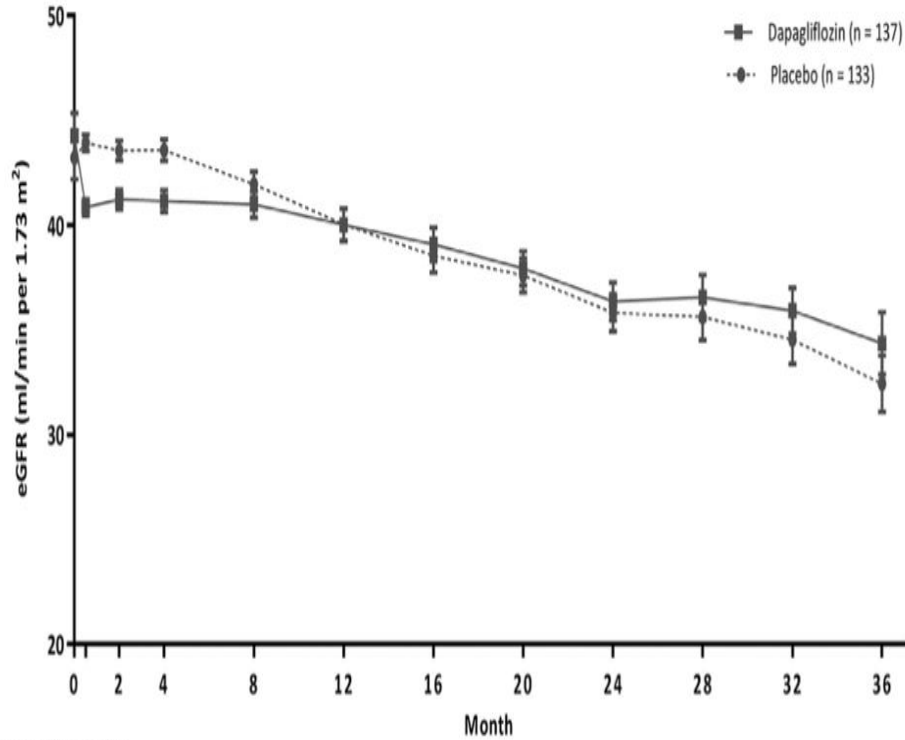


Agent	Suggested usage	Remarks
Antiplatelet agents	Not recommended	No documented evidence of efficacy
Anticoagulants	Not recommended	No documented evidence of efficacy
Azathioprine	Not recommended	No evidence for efficacy as monotherapy or when combined with glucocorticoids
Cyclophosphamide	Not recommended	Unless in the setting of rapidly progressive IgAN
Calcineurin inhibitors	Not recommended	No documented evidence of efficacy
Rituximab	Not recommended	No documented evidence of efficacy
Fish oil	Not recommended	Patients who wish to take fish oil should be advised of the dose and formulation used in the published clinical trials that reported efficacy.

- Practice Point 2.3.1.6: Tonsillectomy in IgAN: Tonsillectomy should not be performed as a treatment for IgAN in Caucasian patients.
- Tonsillectomy is suggested in some national guidelines for the treatment of recurrent tonsillitis in patients with IgAN.
- Multiple studies from Japan have reported improved kidney survival and partial or complete remission of hematuria and proteinuria following tonsillectomy alone or with pulsed glucocorticoids

**A pre-specified analysis of the DAPA-CKD trial demonstrates the effects of dapagliflozin on major adverse kidney events in patients with IgA nephropathy**

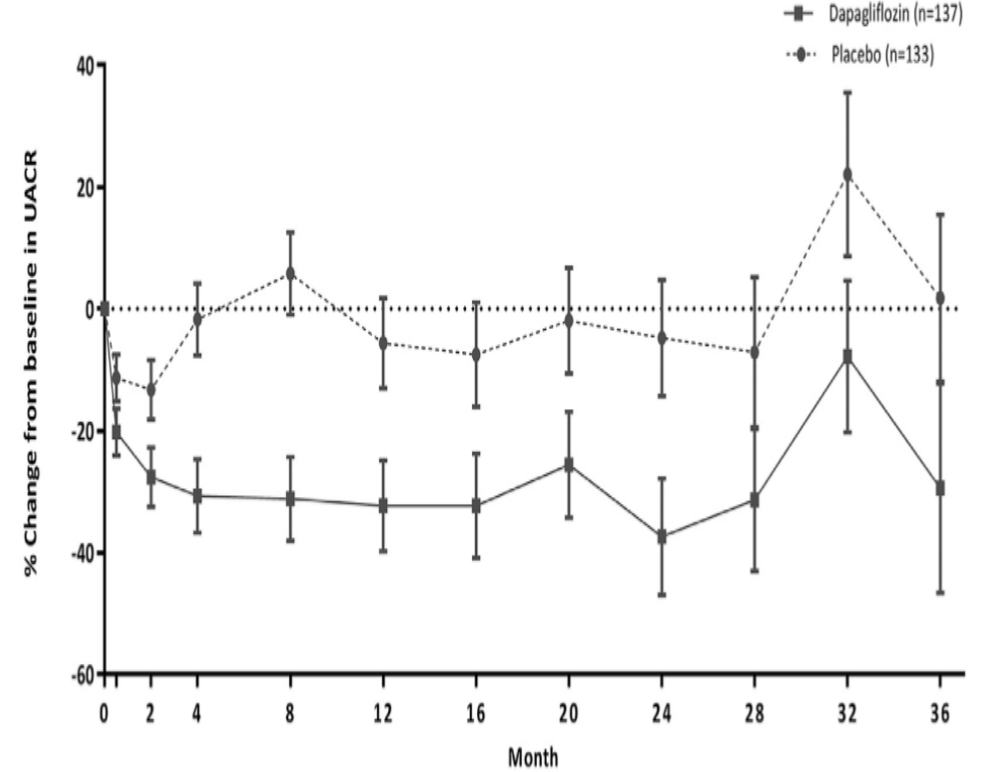
- **270 Participants** (386 study sites in 21 countries) with estimated (eGFR) 25-75 mL/min/1.73m<sup>2</sup> and urinary albumin-to-creatinine ratio 200-5000 mg/g were randomized to **Dapagliflozin 10mg or placebo**, as adjunct to standard care.
- The primary composite endpoint was a sustained decline in eGFR of 50% or more, end-stage kidney disease, or death from a kidney disease-related or cardiovascular cause.
- Mean rates of eGFR decline with **Dapagliflozin and placebo were 3.5 and 4.7 mL/min/1.73m<sup>2</sup> /year**, respectively. Dapagliflozin reduced the urinary **albumin-to-creatinine ratio by 26%** relative to placebo.
- Dapagliflozin significantly and substantially reduces the risk of CKD progression with a favorable safety profile.

**a**

Participants per time point

Dapagliflozin	137	130	121	111	95	99	98	95	77	53	34	10
Placebo	133	129	124	118	105	100	98	93	80	45	28	19

**Changes over time estimated  
glomerular filtration rate**

**b**

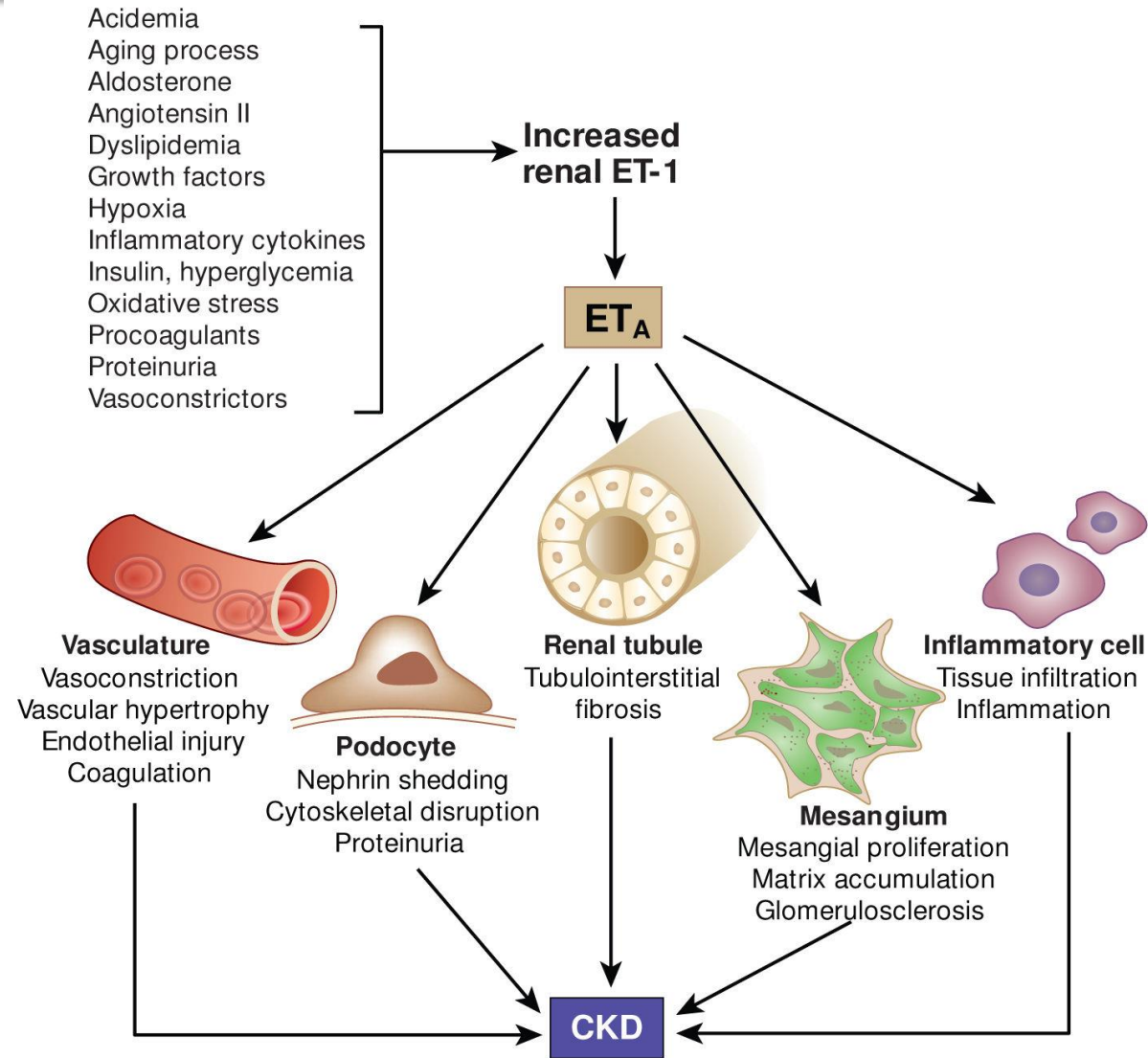
Participants per time point

Dapagliflozin	137	130	121	111	95	99	98	95	77	53	34	10
Placebo	133	129	124	118	105	100	98	93	80	45	28	19

**Changes over time Urine albumin to  
creatinine ratio**

# Pathophysiological role of Endothelin in CKD development

- ET-1 is the most biologically relevant to kidney function in health and disease.
- Endothelins bind to two receptor isoforms, ETA and ETB
- ETA promotes vasoconstriction, cell proliferation and matrix accumulation;
- ETB can promote tissue injury and scarring in pathological condition however in normal condition ,ETB activation is vasodilatory,antiproliferative and antifibrotic effect.

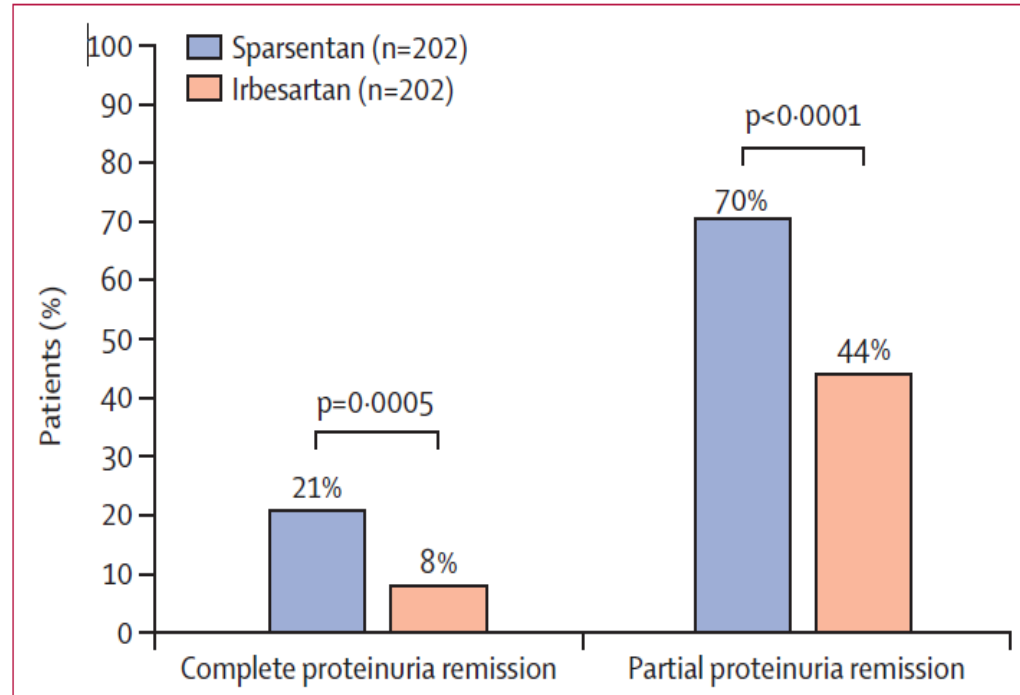
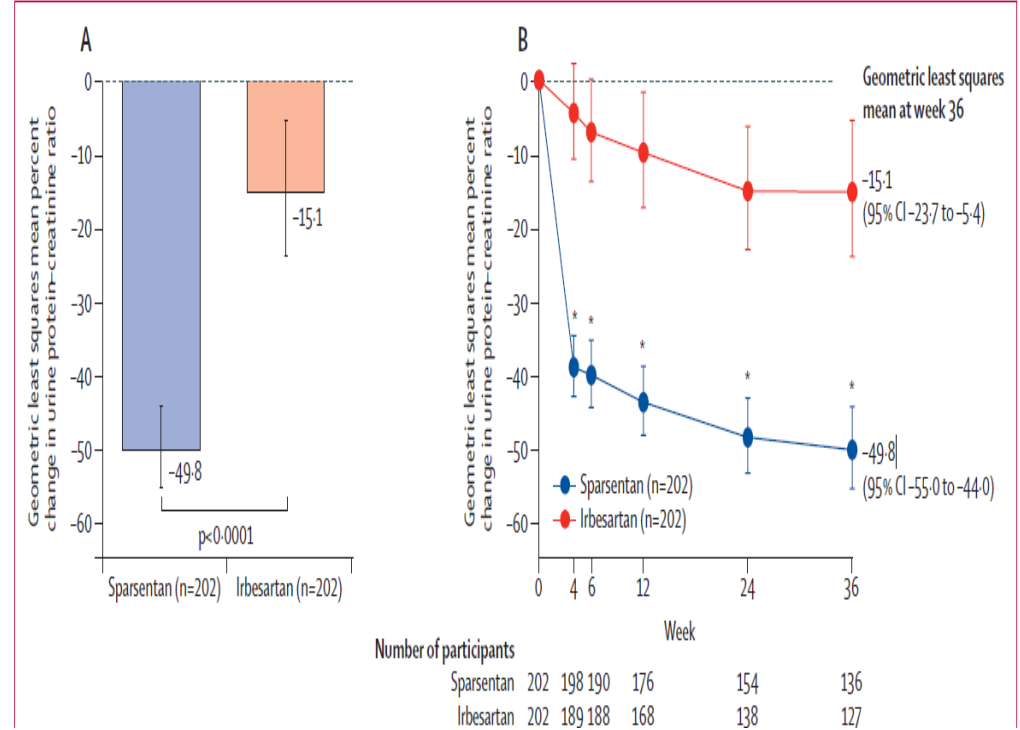




# Sparsentan in patients with IgA nephropathy: a prespecified interim analysis from a randomised, double-blind, active-controlled clinical trial

- **PROTECT** is an international, randomised, double-blind, active-controlled study, being conducted in **134 clinical practice sites in 18 countries**
- Between **Dec 20, 2018, and May 26, 2021, 404 participants** were randomly assigned to sparsentan 400 mg (n=202) or irbesartan 300 mg (n=202) and received treatment.
- **At week 36**, the geometric least squares mean percent change from baseline in urine protein–creatinine ratio was statistically significantly greater in the sparsentan group (–49·8%) than the irbesartan group (–15·1%), resulting in a **between-group relative reduction of 41% (least squares mean ratio=0·59; 95% CI 0·51–0·69; p<0·0001)**.

- Sparsentan, a single molecule with dual antagonism of ETA receptors and AT1 receptors.
- Once-daily treatment with sparsentan produced meaningful reduction in proteinuria compared with irbesartan in adults with IgA nephropathy.
- Safety of sparsentan was similar to irbesartan.
- Completion of the will show whether these beneficial effects translate into a long-term nephroprotective potential of Sparsentan.



# Anti Endothelin A receptors side effects

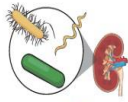
- Edema : 14% vs 9% with *irbesartan*
- Hyperkalemia : 13% vs 10% with *irbesartan*
- Dizziness (13% vs 5% with *irbesartan*)
- Hypotension: (including orthostatic hypotension; 14% vs 6% with *irbesartan*)
- Sulfonamide-based ERAs can cause hepatotoxicity
- Absolutely contraindicated during pregnancy
- Potentially testicular toxicity
- In CKD and CHF needs careful prescription due to edema

# The Intestine – Renal Connection In Iga Nephropathy

- A gross hematuria follows mucosal infection
- Association of celiac disease , dermatitis herpetiformis, IBD and with IgA nephropathy.
- High association of IgA against gliadin ,bovine serum albumin and lactoglobulin in 20-30 % of cases

# Mucosal Infections and Immune Responses

## A specific pathogens



specific pathogens deposit in kidney — direct contact & kidney injury

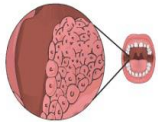


*Helicobacter pylori* — cytotoxin associated gene A protein — Gd-IgA1 ↑



poliovirus — serum levels of IgA ↑ / IgA deposits / mesangial proliferation

## B chronic and persistent infections



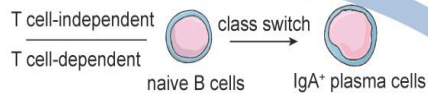
tonsils — close relationship between gross hematuria and tonsillitis / producing sites of Gd-IgA1 ↑ / microbiome analysis / *HORMAD2* / population-based tonsillectomy

## C alterations of gut microbiome

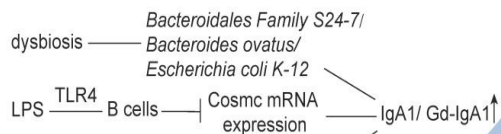


persistent antigenic stimulation — aberrant mucosal immune responses — IgA1 ↑

· B cells class switch



· microbes & IgA production



· metabolites & IgA production uremic toxins — pro-inflammatory cytokines

## A- Specific pathogens :

believed to be involved in the initiation and progression of IgAN.

## B- Chronic and persistent infections:

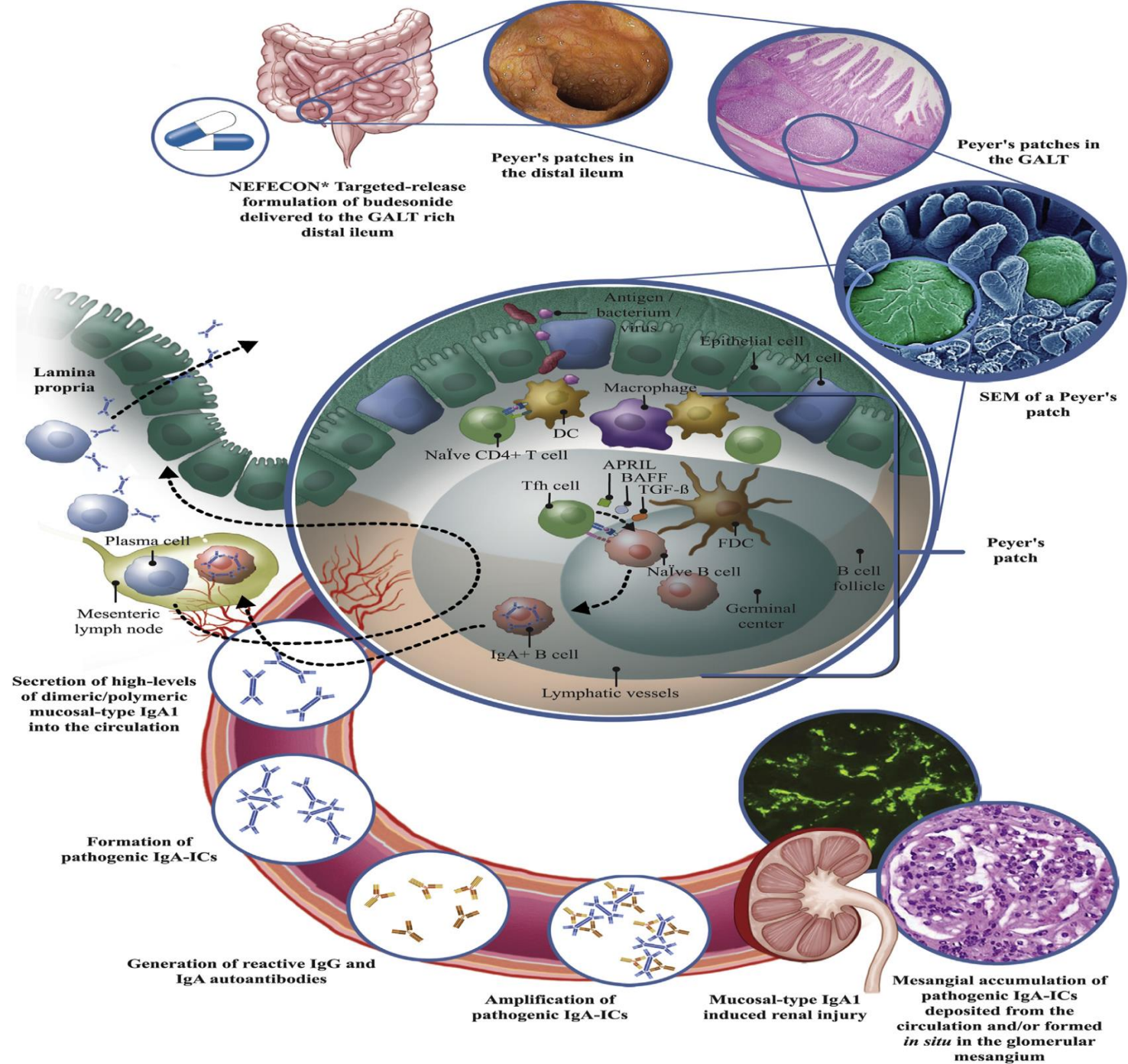
The occurrence of tonsillitis is believed to be related to IgAN. Clinically, there is a close relationship between upper respiratory infections and hematuria

## C- Intestinal infections :

caused by the alterations of the gut microbiome and Persistent antigenic stimulation causes aberrant mucosal immune responses.



# The Peyer's patch, mucosal IgA synthesis, IgA nephropathy, and a role for NEFECON in the treatment of IgAN

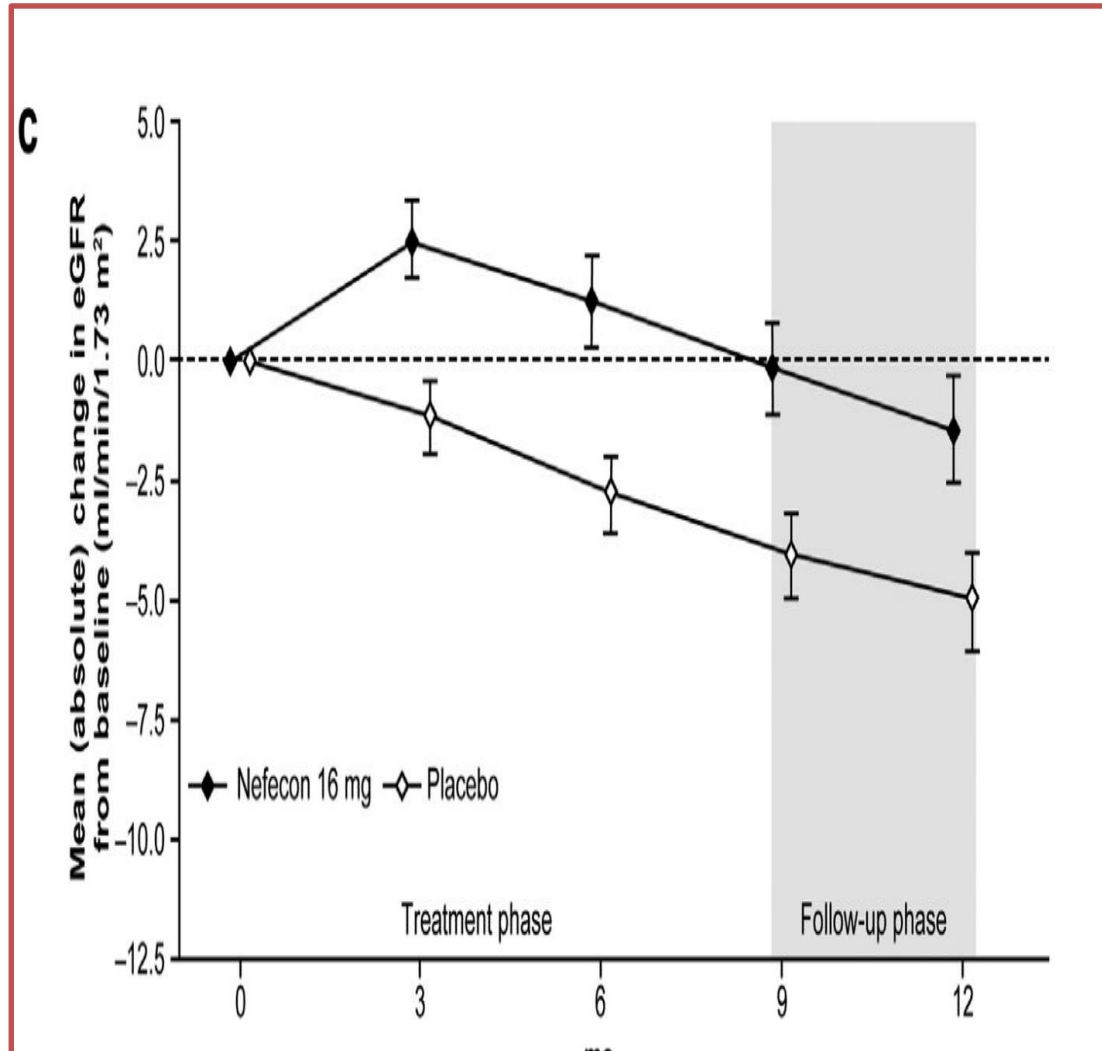


\*NEFECON is an investigational treatment for IgAN and is not FDA approved

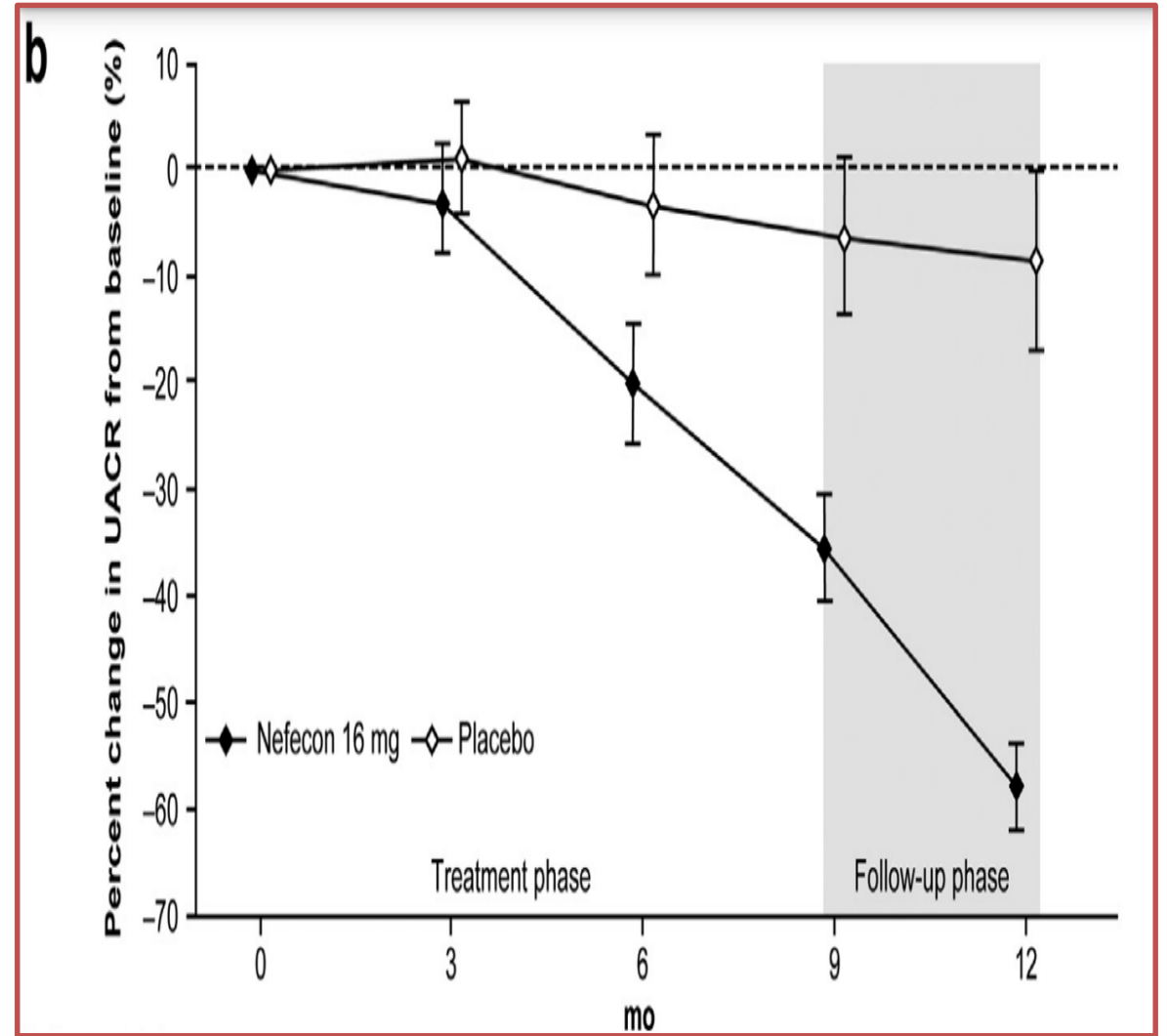
**Results from part A of the multi-center, double-blind, randomized, placebo-controlled NeflgArd trial, which evaluated targeted-release formulation of budesonide for the treatment of primary immunoglobulin A nephropathy**

- NeflgArd was a multicenter, randomized, double-blind, placebo-controlled two-part trial
- In Part A, **199 patients** with IgAN were treated with **Nefecon 16 mg or placebo** for **nine months** and observed for an additional three months.
- At nine months, **UPCR was 27% lower in the Nefecon group** compared with placebo, along with a benefit in eGFR preservation corresponding to a 3.87 ml/min/1.73 m<sup>2</sup> difference versus placebo (both significant).

## Estimated glomerular filtration rate (CKD-EPI)



## Urine albumin-to-creatinine ratio (UACR) (g/g)



# Role of Complement in IgA nephropathy

**Hit 1** :Increased circulating galactose-deficient IgA1

**Hit 2** :Production of unique antiglycan antibodies

Activation in formation of C3 participates in the formation of pathogenic immune complexes

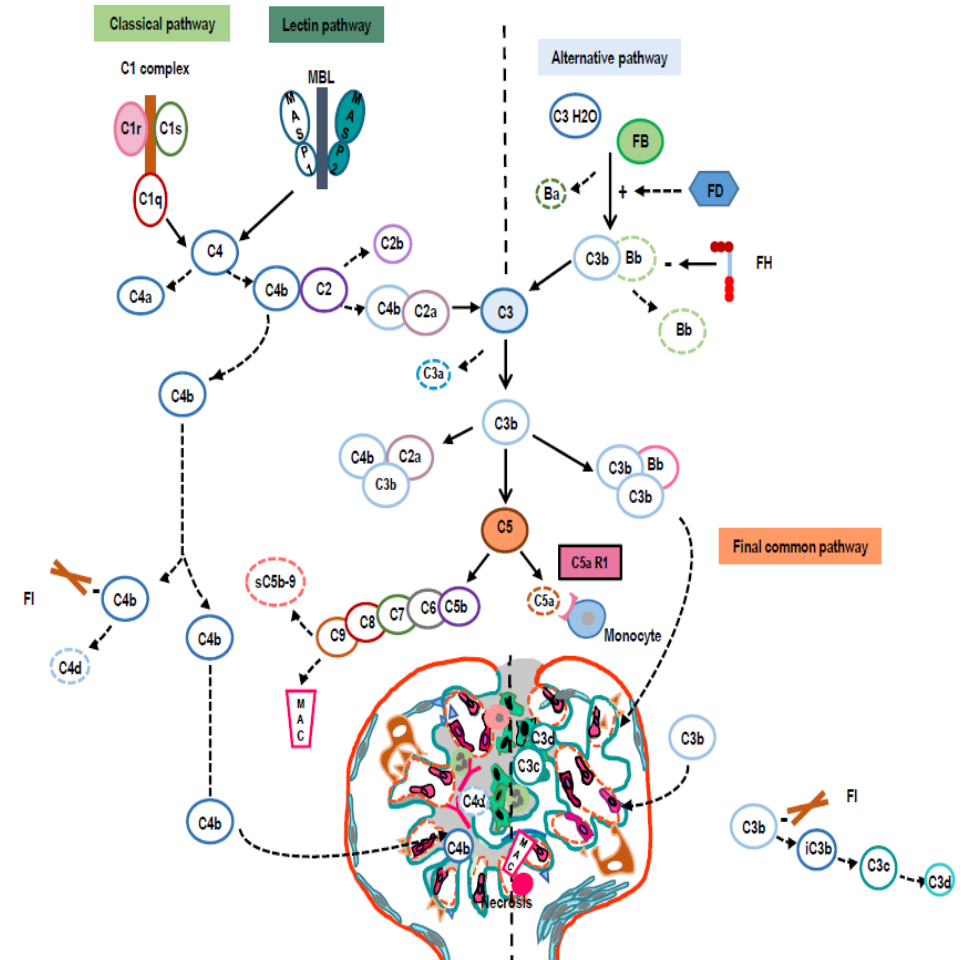
**Hit 3** :Formation of pathogenic IgA1-containing circulating immune complexes

**Hit 4**:Mesangial deposition and mesangial cells activation leading to glomerular injury

Mesangial cells have an active role in complement activation  
Lectin and alternative pathways are activated and contribute to tissue injury

# Complement Activation In IgA Nephropathy

- C4d, MBL, and C5b-9 deposits are characteristic pathologic features of IgA nephropathy.
- Markers of glomerular activation of the lectin pathway (MBL, L-ficolin, MASP2, MASP1/3, and C4d) have been associated with a worse outcome of IgA Nephropathy.
- Variations in complement genes have been associated with better (CFHR3,1deletion) or worse outcome (CFH, CFHR5) of IgA nephropathy.
- Plasma levels of FHR-1 and FHR-1/FH ratio are associated with a progressive course of IgA nephropathy.
- C3a receptor/C5a receptor deficiency in mice alleviates IgA nephropathy in mice.





# C5a receptor inhibitor avacopan in IgA nephropathy – an open-label pilot study

This study evaluates the safety and efficacy of avacopan in patients with IgAN with persistent proteinuria despite RASi blockage

## Methods

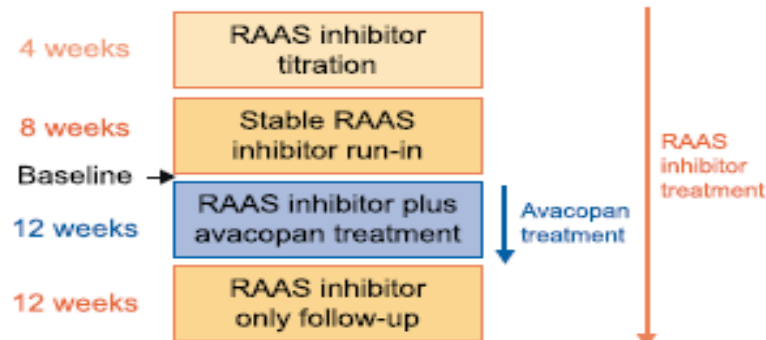
### Open-label pilot trial

- ✓ UPCR > 1g/g
- ✓ eGFR > 60 mL/min/1.73 m<sup>2</sup>



OR

- ✓ eGFR > 45 mL/min/1.73 m<sup>2</sup> (if eGFR has not declined > 10 mL/min/1.73 m<sup>2</sup> in 24w)



## Results

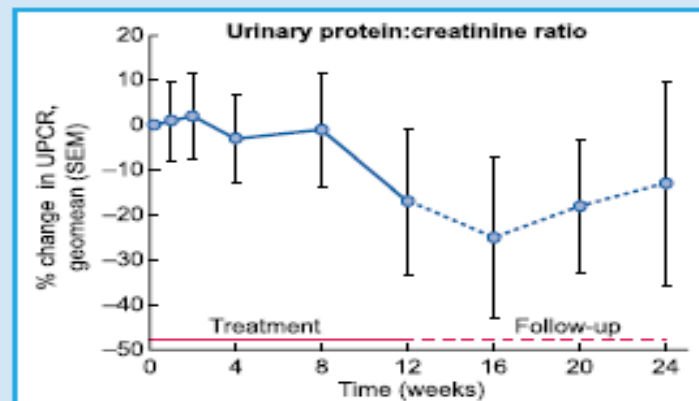


Run-in period of 8 w

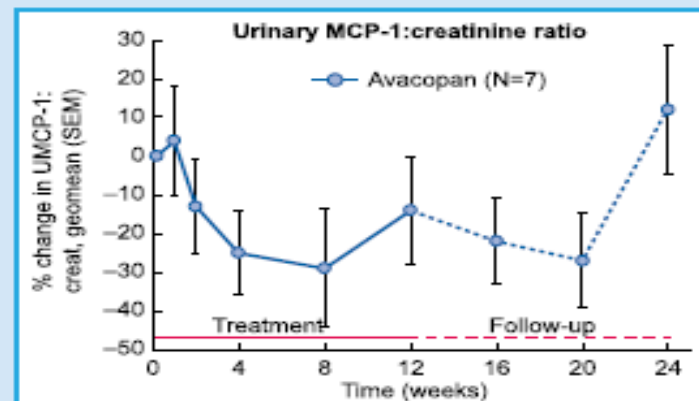
UPCR > 1g/g



Avacopan  
30 mg × 2 day



Improvement in UPCR during treatment



Urinary MCP-1: creatinine decreased 30%



1 event: unstable angina – unrelated to avacopan

**Conclusion:** This short-term trial showed an improvement in the slope of UPCR in 6 out of 7 patients, with ~ 50% improvement in 3 out of 7 patients with IgAN. Longer avacopan treatment duration may be indicated for maximal benefit.

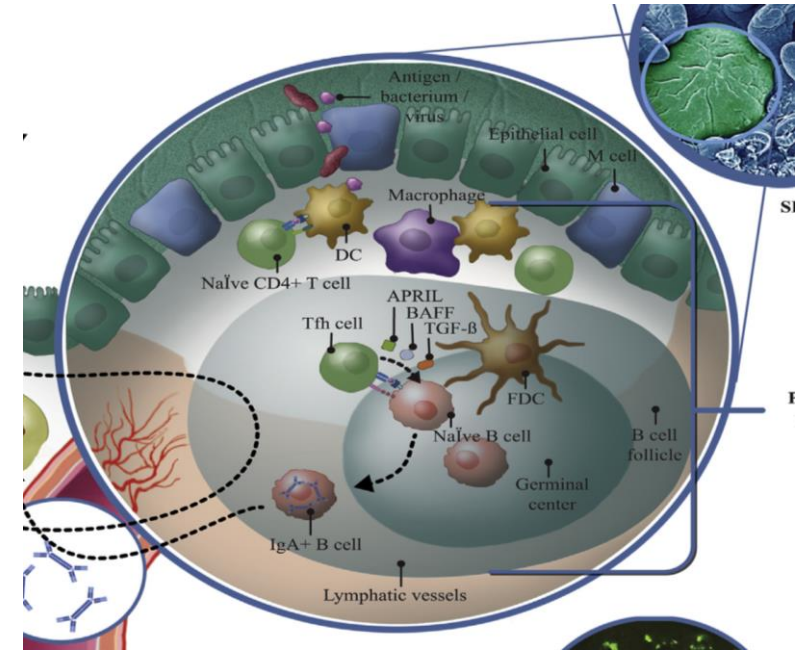
Bruchfeld, A. et al.  
Clinical Kidney Journal (2021)  
annette.bruchfeld@liu.se  
@CKJsocial

# Main complement inhibitors undergoing development in kidney diseases

Target in the complement cascade	Mechanism of action	Drug	Pharmaceutical company	Type of inhibitor	Mode of administration	Phases of drug development	Potential indications in kidney diseases
C5	Inhibition of the release of C5a and C5b, and ultimately of the formation of C5b9	Eculizumab	Alexion Pharma/ AstraZeneca	mAb	i.v.	Commercialized	aHUS
		Ravulizumab	Alexion Pharma/ AstraZeneca	mAb	i.v.	Commercialized, phase III	aHUS
		Crovalimab	Roche	mAb	s.c.	Phases II–III	aHUS
C3	Inhibition of the binding of C3 to the C3bBb and thus of the cleavage of C3	Pegcetacoplan	Apellis Pharma/ SOBI	Pegylated peptide	s.c.	Phase III	C3G, IgAN, MN
Factor B	Inhibition of the serine protease FB and thus of the cleavage of C3 and C5	Iptacopan	Novartis	Small molecule	Oral	Phases II–III	aHUS, C3G, MN, IgAN
Factor D	Inhibition of the cleavage of FB	Danicopan	Alexion Pharma/ AstraZeneca	Small molecule	Oral	Phases II–III	C3G
MASP2	Inhibition of the serine protease MASP2	Narsoplimab	Omeros	mAb	i.v.	Phase II	IgAN
C5a receptor	Inhibition of the binding of C5a to its receptor	Avacopan	Chemocentrix	Small molecule	Oral	Phase III	ANCA-associated vasculitis aHUS

# BAFF AND APRIL in IgA nephropathy

- B-cell-activating factor (BAFF) and A proliferation-inducing ligand (APRIL), produced by antigen-exposed dendritic cells and intestinal epithelial cells.
- Increased serum levels of BAFF and APRIL in patients with IgAN that correlate with gd-IgA1 levels and disease severity.
- Elevated BAFF levels have been associated with specific fecal metabolites, especially phenols, in individuals consuming high beef diet who had increased populations of phenol-producing anaerobic *Bacteroides*.



2018

## Bortezomib for Reduction of Proteinuria in IgA Nephropathy



Choli Hartono<sup>1,2</sup>, Miriam Chung<sup>3</sup>, Alan S. Perlman<sup>1,2</sup>, James M. Chevalier<sup>1,2</sup>, David Serur<sup>1,2</sup>, Surya V. Seshan<sup>4</sup> and Thangamani Muthukumar<sup>1</sup>

- Bortezomib is a proteasome inhibitor that targets plasma cells.
- **Eight** consecutive subjects from **July 2011 until March 2016** with **4** doses of **bortezomib every two weeks**. All subjects had biopsy-proven IgA nephropathy and proteinuria of greater than 1 g per day.
- The **3 subjects who had complete remission** had Oxford classification T scores of 0 before enrollment. Of the remaining 5 subjects, 1 was lost to follow-up within 1 month of enrollment and **4 (50%) did not have any response** or had progression of disease.

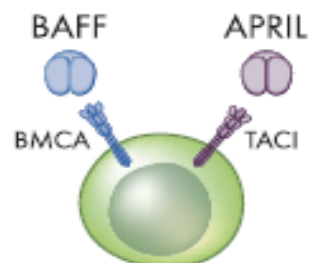
# B-cell directed therapies currently being evaluated in IgAN

Drug	Mechanism	Phase	Design	Primary Outcome	Identifier
Mezagitamab	mAb against CD38	I	Open label single arm trial	Incidence of adverse events	NCT05174221
BION-1301	mAb against APRIL	I/II	Double blinded, placebo controlled single ascending dose study	Incidence and severity of adverse events	NCT03945318
Ataccept	TACI fusion protein, acts as a BAFF and APRIL inhibitor	II	Open label multiple dose study Double blinded placebo-controlled trial	Proteinuria Reduction	NCT04716231
Belimumab	mAb against BAFF	II	Double blinded placebo-controlled trial	Proteinuria reduction, Change in eGFR, Adverse events	EudraCT: 2017-004366-10
Felzartamab	mAb against CD38	II	Double blinded, placebo-controlled trial	Proteinuria reduction	NCT05065970
Telitaccept	TACI fusion protein, acts as a BAFF and APRIL inhibitor	II	Double blinded placebo-controlled trial	Proteinuria Reduction	NCT04905212
Blisibimod	Peptibody inhibitor of BAFF	II/III	Double blinded, placebo-controlled trial	Proteinuria Reduction	NCT02062684
Sibeprenlimab	mAb against APRIL	II	Double-blinded, placebo-controlled, multiple dose trial	Proteinuria Reduction	NCT04287985
		III	Double-blinded, placebo-controlled trial	Proteinuria Reduction	NCT05248646
Rituximab	mAb against CD20	IV	Single blinded randomized trial against supportive care	Proteinuria reduction	NCT04525729



# Immune abnormalities in IgA nephropathy

## B cells



Increased BAFF and APRIL serum levels

Higher expression of BMCA and TACI

Higher levels of gut-homing (CCR9<sup>+</sup> β7integrin<sup>+</sup>) B cells

## T cells



Th1/Th2 imbalance

Decreased Treg

Increased Th22 and Th17

Positive correlation between CXCR5<sup>+</sup> PD-1<sup>+</sup>Tfh and serum gd-IgA1

## Toll-like receptors



Higher expression of TLRs in the kidney

Increased expression of TLR mRNA in PBMCs with positive correlation with proteinuria

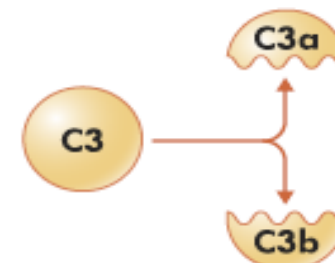
## Monocytes/macrophages



Increase in non-classical monocytes

Increased expression of TIM-3<sup>+</sup> with positive correlation with proteinuria

## Complement



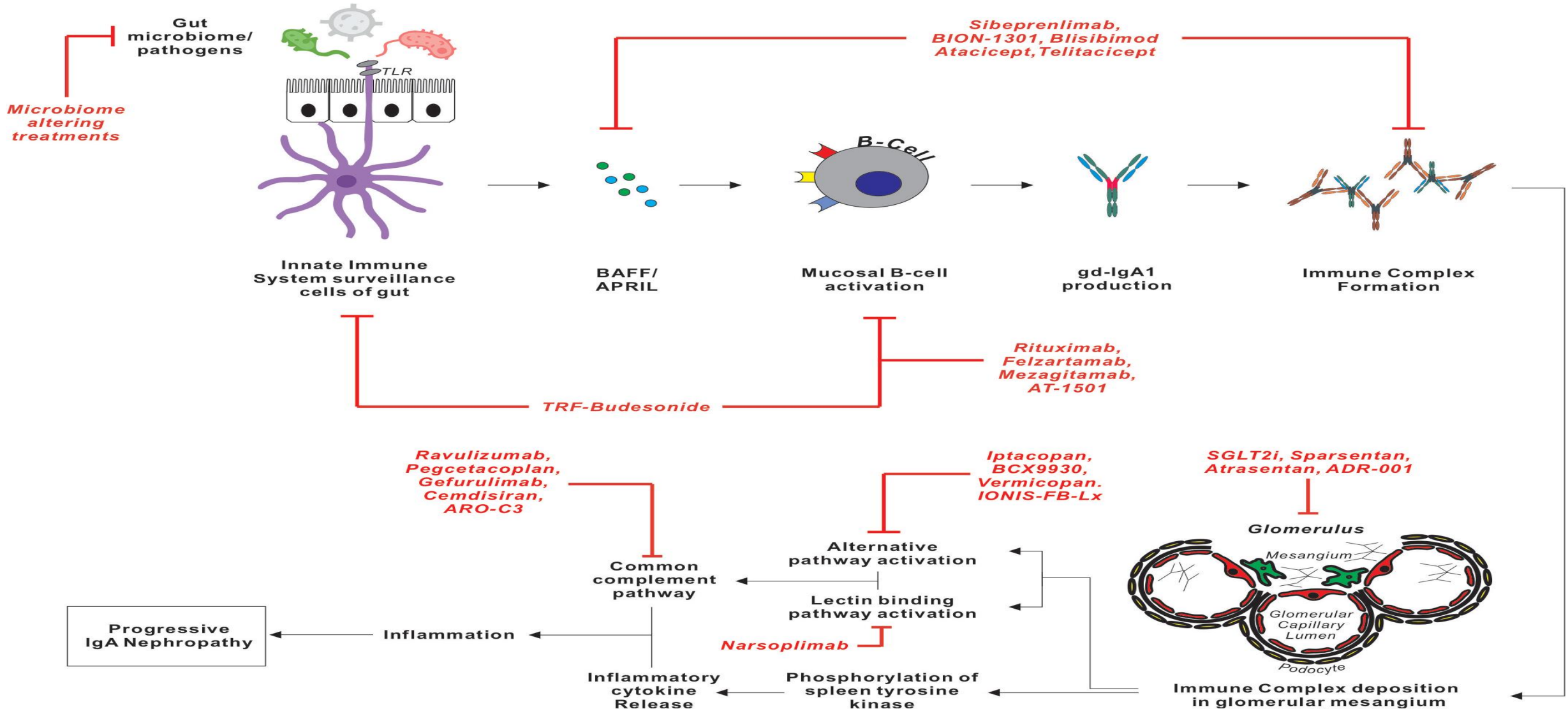
Activation of alternative and lectin pathways

Glomerular C3b deposition correlates with the progression of IgAN

**Conclusion:** Although several key questions about the production of gd-IgA1 and the formation of anti-gd-IgA1 antibodies remain unanswered, a growing body of evidence is shedding light on the innate and adaptive immune mechanisms involved in this complex pathogenic process and how they could be therapeutically targeted.

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 Clinical Kidney Journal (2023)  
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# Pathophysiology Of IgA Nephropathy And Directed Treatment Strategies



# Current Clinical Trials In Iga Nephropathy

Selvaskandan et al  
 EXPERT OPINION ON  
 INVESTIGATIONAL DRUGS  
 2022, VOL. 31, NO. 12,  
 1321–1338

