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Milk Fat Globule Proteins Are Relevant Bovine Milk Allergens in Patients with alpha-Gal Syndrome

Original

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Milk fat globule proteins are relevant bovine milk allergens in patients with a-Gal syndrome

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13 14	4	Beatrice Aiuto ^{a,b} , Simona Cirrincione ^a *, Maria Gabriella Giuffrida ^a , Laura Cavallarin ^a , Chiara Portesi ^c ,
15 16	5	Andrea Mario Rossi ^c , Giorgio Borreani ^d , Giovanni Rolla ^e , Massimo Geuna ^e , Stefania Nicola ^e , Anna
17 18 19	6	Quinternetto ^e , Lucrezia Alessi ^e , Elena Saracco ^e , Luisa Brussino ^e , Cristina Lamberti ^a .
20 21	7	
22 23	,	
24 25	8	Institutional affiliation
26	0	
27 28 20	9	a. Institute of the Science of Food Production (ISPA) – National Research Council, Largo Braccini 2,
30 31	10	10095 Grugliasco, TO, Italy
32 33	11	b. Politecnico di Torino, Corso Castelfilardo 39, 10129 Torino, Italy
34 35 36	12	c. National Institute of Metrological Research (INRIM), Strada delle Cacce 91, 10135 Torino, Italy
37 38	13	d. University of Turin, Italy - Department of Agriculture, Forestry and Food Sciences (DISAFA)
39 40 41	14	e. Department of Medical Sciences, Allergy and Clinical Immunology Unit, University of Torino &
42 43	15	Mauriziano Hospital, Torino, Italy
44 45	16	
46 47	17	
48 49 50	18	*Corresponding author
51 52	19	Simona Cirrincione:
53 54 55	20	simona.cirrincione@ispa.cnr.it
56 57	21	Largo Braccini, 2
58 59 60	22	10095 Grugliasco (TO), Italy

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Scope. Alpha-gal syndrome (AGS) is a mammalian meat allergy associated with tick bites and specific IgE to the oligosaccharide galactose- α -1,3-galactose (α -gal). Recent studies have shown that 10– 20% of AGS patients also react to the whey milk proteins. Considering the already described role of the lipid fraction of meat in AGS clinical manifestations, the aim of this work has been to investigate whether the milk fat globule proteins (MFGP) could be involved in AGS.

Methods and results. The MFGP were extracted and their recognition by the IgE of AGS patients was proved through immunoblotting experiments with the sera of AGS patients. The identification of the immunoreactive proteins by LC-HRMS analysis allowed to demonstrate for the first time that butyrophillin, lactoadherin and xanthine oxidase are α -gal glycosylated. The role of xanthine oxidase seems to be prevalent since both the anti- α -gal antibody and AGS patient sera showed the highest immunoreactivity against it.

Conclusion. The results obtained in this study have confirmed the role of α -Gal carrying glycoproteins in AGS patients reacting to milk. Although additional factors are probably associated with the clinical manifestations, the consumption of milk and milk products should be limited or even avoided in individuals with AGS.

1. Introduction

Alpha-gal syndrome (AGS) is a mammalian meat allergy associated with tick bites and specific IgE antibodies to the oligosaccharide galactose- α -1,3-galactose (α -gal).^[1,2] Alpha-gal carbohydrate is missing in humans and some primates, since the α -1,3-galactosyltransferase is expressed in an enzymatically inactive form. It is instead present in most mammals, most pathogens such as bacteria and parasites and in the salivary glands of several tick species, including the most prevalent hard tick in Europe (*Ixodes ricinus*).^[3] Anti α-gal antibodies are the most abundant natural antibodies in humans and some primates constituting up to 1% of the circulating antibodies. These antibodies are mainly IgM and IgG class, but anti α-Gal IgEs are also produced and are responsible for the red meat allergy^[4] AGS symptoms vary from abdominal pain and diarrhea to urticaria and anaphylaxis, the latter being experienced by nearly 50% of patients.^[5,6] AGS shows several exclusive features that make it different from other food allergies: i) reactions are generally delayed, appearing 3 to 6 hours after meat consumption; ii) IgE antibodies react to a carbohydrate moiety rather than a protein epitope; iii) patients can develop AGS in late adulthood after a previous period of meat tolerance.^[7] This atypical food allergy was first described in the southeastern regions of the United States and in Australia, but it was also reported soon thereafter in Europe, Asia, Africa, and Central America.^[8] More than 5000 cases have been described to date in the United States.^[9] The frequency of positivity of specific IgE to α-Gal in Europe has been reported to be increasing in northern countries (Denmark, Sweden, etc.), where it was first investigated,^[10,11] but also in Spain [12] and in the rural areas of northeast Italy.^[13] AGS is characterized by reactions to mammalian meat and innards, including beef, pork, and lamb, as well as to food gelatins and some medications (cetuximab, antivenom, gelatin-containing vaccines).^[14] Unlike common food allergies, the allergic reactions may not occur at every exposure to the allergen. This variability depends on the amount of allergen ingested and on the nature of the biologic macromolecules within the α -gal-containing food. Lipid-rich mammalian

meats are associated with more consistent and severe reactions.^[15] Lipid-bound α -Gal appears to be able to cross the intestinal monolayer and to trigger an allergic reaction, thus suggesting that not only glycoproteins but also glycolipids should be investigated as potential allergenic molecules.^[16] Chakrapani et al.,^[17] have recently confirmed the involvement of glycolipids in the activation of AGS patient basophils, even if the major role played by glycoproteins, particularly those from pork kidneys and beef extracts, is already well established. Glycolipids extracted from these food matrices have shown a lower basophil activation capacity than their respective protein extracts.^[18] Not only red meat but also bovine milk might contain α -Gal-epitopes, although in smaller amounts.^[19] Some recent studies,^[7,20,21] including one involving the analysis of a large cohort of 2,500 AGS patients in the USA,^[22] have proved that 10–20% of AGS patients also react to milk. The most reported symptoms in AGS patients following bovine milk ingestion are abdominal pain and urticaria with a delayed onset of the symptoms.^[23] Unlike meat, where α -Gal-bearing proteins have long been extensively studied, sources containing α -Gal epitopes in dairy products have only recently been investigated. Perusko et al.^[24] demonstrated that bovine y-globulin (BGG), lactoferrin (LF), and lactoperoxidase (LPO) are α -Gal carrying proteins that have been recognized by the IgE of AGS patients and which are able to activate the basophils of patients. More recently, the same α -Gal glycosylated proteins were found in sheep milk by German-Sanchez et al.^[25]

Considering the involvement of milk proteins in AGS and the role of the lipid fraction in facilitating
clinical manifestations of AGS, the aim of this work has been to investigate whether the milk fat
globule protein (MFGP) fraction could also play a role in AGS.

2. Experimental Section

2.1 Characterization of the patients

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2 3 96 This observational study was carried out on 10 adults Italian AGS patients at the Allergy and Clinical 4 5 Immunology University Clinic in Turin (AO Ordine Mauriziano di Torino). All the patients were 97 6 7 8 diagnosed with AGS based on the history of at least one previous hypersensitivity reaction to 98 9 10 mammalian meat and/or its related products (food gelatine), and the presence of positive α -Gal 99 11 12 13 100 specific IgE antibodies. Only adult patients (≥18 years old) with an established diagnosis of AGS were 14 15 101 enrolled. The exclusion criteria were age<18 years old, ongoing anti-IgE biological therapy 16 17 18 102 (Omalizumab), or the lack of informed consent release. The study was approved by the local ethical 19 20 103 committee (Comitato Etico Interaziendale A.O.U. Città della Salute e della Scienza di Torino- A.O. 21 22 Ordine Mauriziano- A.S.L. Città di Torino, study number 0053278, date of approval: 24.05.2019) and 23 104 24 ²⁵ 105 conducted according to the Declaration of Helsinki. Data were collected between June 2019 and 26 27 ₂₈ 106 March 2023. 29 30 ₃₁ 107 The demographic data, the description of previous reactions (culprit food, clinical presentation, time 32 33 108 of symptoms onset, treatment, and the presence of co-factors) and their history of previous tick 34 35 36 109 bites are reported in Table 1. All the patients underwent blood tests to analyze the total serum IgE 37 and α -Gal specific IgE antibodies (Immunocap Fluorescence Enzyme Immunoassay Feia, by Thermo 38 110 39 40 111 Fisher). According to the manufacturer's recommendations, levels of total IgE below 205 KU/L and

⁴¹ ¹¹¹ ¹¹¹

A statistical analysis was performed, using IBM SPSS Statistics for Windows, version 26 (IBM Corp., Armonk, NY, USA). A normality distribution of data was first tested using the Kolmogorov–Smirnov normality test, and a descriptive analysis of the variables was then performed. The baseline characteristics were evaluated over the whole cohort and expressed as the mean (standard characteristics were evaluated over the whole cohort and expressed as the mean (standard

1 2	
3 119 4	deviation, SD), unless otherwise specified, for the continuous variables, and as absolute and relative
6 120 7	frequencies for the categorical variables.
8 121 9	
10 11 122 12	2.2 Chemicals
13 14 123 15	Details pertaining to this topic are available in the Online Repository.
16 17 124 18	2.3 Milk fat globule membrane associated protein extraction
19 20 125 21	The MFGP was extracted according to Barello et al. ^[26] Details are available in the Online Repository.
23 126 24 25	2.4 Glycosylated milk fat globule membrane associated protein enrichment
26 127 27	Sixty μ l of BioMag Goat Anti-Human IgG beads (5,2mg/ml) (BioMag beads) were washed twice with
28 29 128 30	500 μl of PBS. The washed BioMag beads were blocked twice with TBS with 0,3% of Tween 20
31 129 32	(blocking solution) for 15 minutes under agitation at 4°C. After removing the BS, the BioMag beads
³³ 34 35	were incubated with 1:1 of α -gal-IgG Ab for 6 hours under rotation at 4°C. The α -gal-IgG Ab /BioMag
36 131 37	bead complexes were collected by means of a magnetic bar and were washed twice with 500 μl of
³⁸ 132 39	PBS. Sixty μg of MFGP were added to the $\alpha\mbox{-}gal\mbox{-}lgG$ Ab /BioMag bead complexe and incubated
40 41 133 42	overnight (O.N.) at 4°C. The α -gal-IgG antibody /BioMag bead / MFGP complexes were then
43 134 44	collected again and washed twice with 500 μl of PBS. The MFGP and $\alpha\text{-gal-IgG}$ antibodies were
45 46 135 47	released from the BioMag beads by incubating them with the elution buffer (1% (w/v) SDS, 100 mM $$
48 136 49	Tris HCl, pH 7.4, 10 mM DTT, 8M urea) for 10 min at 95°C. The proteins released from the beads
50 51 52	were then used in the subsequent experiments.
53 54 55	2.5 Milk fat globule membrane associated protein N-de-glycosylation
56 57 139 58	Enzymatic removal of the N-linked glycans was performed using PNGase F, a glycan-Asn-amidase

that specifically cleaves the innermost GlcNAc of all N-linked oligosaccharides, unless they carry $\alpha(1-60)$

3) core-bound fucose residues.^[27] The experiment was carried out under denaturing conditions: 40g
 of proteins were resuspended in a modified Laemmli buffer (60mM Tris-HCl pH 6.8, 0,25% SDS, 10%
 glycerol) and 1ul of 1M DTT was added. The sample was incubated at 95°C for 5 min. After cooling,
 2 μl of 10% NP-40 and a quantity of PNGase F (10U/μg) were added, in a 1:1 enzyme/substrate ratio,
 to the sample and incubated at 37 °C for 3 hours and overnight (ON) under slight shaking.

2.6 Protein separation and LC-HRMS analysis

The LDS-PAGE separation and LC-HRMS analysis were performed according to Cirrincione et al.^[28]
 Details are available in the Online Repository.

49 2.7 Protein identification strategy

All the Data Dependent Analysis (DDA) files were searched using MaxQuant (https://maxquant.org) v. 2.0.3.0 against the UniProt *Bos taurus* database (reviewed and unreviewed). The search was performed using a list of contaminants devoid of bovine proteins, because they were our target. The search parameters were set as follow: S-carbamidomethyl derivate on cysteine as a fixed modification, oxidation on methionine, Acetyl (N-term) as variable modifications and two missed cleavage sites for trypsin digestion. The possibility of Asn becoming Asp was added as a variable modification for bands derived from enzymatic de-glycosylation. The MS/MS fragment mass tolerance was set at 20 ppm. A minimum of 2 peptides, an FDR of 0.01% for both the protein and peptides, and a score of 20 for unmodified and modified peptides were set for the protein identification. Only proteins identified with a score >30 were listed in the tables, with the exception of the identification performed on unstained bands cut in the upper part of the gels where a score of >10 was allowed.

2.8 Whey and milk fat globule membrane associated protein immunoblotting

After LDS-PAGE, the protein bands were electro-transferred into Nitrocellulose Membranes (0.2 μ m) with an XCell II Blot Module, using a transfer buffer with 10% methanol (v/v). The membranes were blocked in TBS with 0.3% Tween 20 (blocking solution) for 30 min and incubated ON at 4 °C with 800 μ L of the HRP conjugated Human IgG1 anti α -Gal epitope antibody (Absolute Antibody) diluted 1:1000 in the incubation buffer (TBS, 0.05% Tween 20, 0.05% vegetal gelatin) or with the patients' sera diluted 1:10 in the incubation buffer. After incubation, the membranes were washed three times with TBS, 0.05% and Tween 20 (washing solution) for 10 min. The membranes incubated with the patient's sera were incubated again with the anti-Human IgE antibody (Sera Care Life Sciences Inc., Milford, Massachusetts) diluted 1:5000 in the incubation buffer. The membranes were washed three times and developed with different development kits according to the used primary antibody: an Alkaline Phosphatase Substrate Kit (Bio-Rad) for the patients' sera and an Opti 4 CN Kit (Bio-Rad) for the HRP conjugated Human IgG1 anti α -Gal epitope antibody.

2.9 Immunoprecipitation of the AGS patient sera

Immunoprecipitation experiments were performed with two glycosylated proteins: bovine thyroglobulin and the bovine xanthine oxidase from Sigma-Aldrich. The sera of three AGS patient ($\alpha 2$, $\alpha 3$, $\alpha 5$) were incubated for 1h at room temperature with four amounts of thyroglobulin (1, 3, 30, 60 µg) and other three patients ($\alpha 1$, $\alpha 2$, $\alpha 7$) were incubated at the same conditions with three amounts of xanthine oxidase (3, 30, 60 µg). Nitrocellulose membranes containing electrotransferred MFGP associated proteins were blocked with the blocking solution for 30 min and then incubated overnight at 4°C with the immunoprecipitated sera. The immunoblotting procedure was then performed as previously explained in Section 2.7.

60 185 **3. Results**

186 3.1 Study population

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Ten adult patients (4 females; 40.0%) with a mean age of 59.4 years (range 25-74 years) and a 187 diagnosis of α -gal syndrome (AGS) were enrolled in the experiment. One patient (M, 48 years) with 188 189 a history of non-IgE mediated milk hypersensitivity reaction was used as a healthy control.

₁₅ 190 3.1.1 Comorbidities

17 191 Three patients had arterial hypertension, one suffered from diabetes, one was HIV positive, and one suffered from atrial fibrillation. All the patients had a normal weight, and three of them were 192 22 193 smokers. As far as atopic diseases are concerned, 3 patients had allergic rhinitis, 2 patients showed 194 sensitization to lipid transfer protein (LTP) with mild food allergy manifestations, and one patient 27 195 reported nonsteroidal anti-inflammatory drug (NSAID) hypersensitivity (urticaria) as well ampicillin ²⁹ 196 hypersensitivity. One patient (male, 64 years old, α 6) was affected by systemic indolent 32 197 mastocytosis.

³⁵ 198

3.1.2 Clinical presentation of AGS

37 ₃₈ 199 All the AGS patients reported at least one delayed reaction (average 3.40± 1.58 events/person) with 39 40 200 a mean onset time of 4.1 hours after eating red meat, innards, or meat-related food (Tab. 1). None 41 42 201 of the patients experienced reactions to cow's milk or dairy products. The most common culprit 43 44 food was pork meat. Urticaria was the most common clinical manifestation (100%), followed by 45 202 46 47 203 gastrointestinal symptoms (vomiting, diarrhea, and abdominal pain) (60%), hypotension (50%), 48 49 50 204 angioedema (50%) and dyspnea (30%). Nine patients (90%) had at least one episode of anaphylaxis, 51 ⁵² 205 diagnosed according to NIAID/FAAN criteria.^[29] No cofactor of anaphylaxis, including ethanol, or 53 54 ₅₅ 206 nonsteroidal anti-inflammatory drug consumption was identified, apart from one patient who 56 ⁵⁷ 207 reported anaphylaxis after red meat ingestion and physical exercise. None of our patients had 58

1 2		
2 3 4	208	previously been treated with cetuximab. Eight patients (80%) reported one or multiple tick bites
5 6 7	209	before AGS.
7 8 9	210	All the patients were positive to α -gal specific IgE (26,08 ± 35.87 KUA/I) with a mean serum total IgE
10 11	211	of 389.99±429.94 KU/I. Tryptase resulted normal in all the patients, with a mean value of 7,18±3.78
12 13 14	212	μg/I).
15 16	213	
17 18 19	214	3.1.3 AGS treatment
20 21	215	All the patients received corticosteroids and antihistamines for their hypersensitivity reactions.
22 23 24	216	Seven patients (70%) had been admitted to the intensive care unit for a total of 10 times. In five
25 26	217	cases, the reactions were treated with adrenaline.
27 28 29	218	
30 31 32	219	3.2 The Anti- α -gal antibody recognizes whey and milk fat globule proteins
33 34	220	The milk fat globule proteins (MFGP), whey proteins (WP), and caseins (CAS) were separated by
36 37	221	means of LDS page followed by immunoblotting analysis with anti- α -Gal IgG and a pool of sera from
38 39	222	10 AGS patients (Fig 1). Both the MFGP and WP extracts showed immunoreactive bands for anti- α -
40 41 42	223	Gal IgG: G1, G2, G3, G5, G6, G7 and W1, W2, W3, W5, W6, respectively (Fig 1, panel B). LC-HRMS
43 44	224	analysis (Tab. 2) allowed LF and LPO to be identified in band W3, and several Ig-like domain
45 46 47	225	containing proteins were identified in bands W2, W3, W5, W6. Xanthine Oxidase (XO) was identified
48 49	226	in W1 and in several reactive bands of MFGP (G1, G2 and G3), while the other reactive bands (G3,
50 51 52	227	G5, G6 and G7) mainly contained butyrophilin (BT) and lactadherin (LA).
53 54 55	228	The pool of AGS patient sera immunorecognized all the bands already recognized by anti- α -Gal IgG,
56 57	229	albeit with the addition of bands G4, G8, W4, W7, C1 and C2. Band G4 contained several proteins
58 59 60	230	including XO, BT and LA; G8, W4 and C1 contained already known α -Gal glycosylated proteins (BT,
	231	LPO and Ig-like domain-containing proteins); while bands W7 and C2 contained typical milk allergens

(β-lactoglobulin and caseins) and were probably recognized because the patients were sensitized to milk, although they tolerated it well, according to the study inclusion criteria (Fig. 1, panel C).

Band G1, which contained XO, was not visualized by colloidal Coomassie staining or even by silver staining (data not shown), but it was clearly recognized by anti- α -Gal IgG and by the AGS patient IgEs in the immunoblotting experiment.

3.3 Xanthine oxidase, butyrophilin and lactadherin are α -Gal-glycosylated proteins

In order to enrich the sample in α -Gal-glycosylated proteins, we isolated glycosylated MFGP using BioMag Goat Anti-Human IgG beads conjugated with the anti-α-gal IgG system. After the enrichment, the proteins were separated by means of LDS PAGE (Fig. 2, panel A, lane MFGPb). The thus isolated MFGP resulted to be high molecular weight proteins and as expected, they were recognized by the anti-α-gal IgG. However, the situation was different for bands G16, G17 and G25, as they contained heavy and light anti- α -gal IgG chains partially released from the beads during protein elution, and PNGase F, the enzyme used for de-glycosylation. In addition to the heavy anti-244 α -gal IgG chain, LA was identified in band G16, which is probably responsible for the corresponding 246 immunoreactivity, while the other two bands did not result to be immunoreactive. When the α -Galenriched protein sample was de-glycosylated with PNGase F, the anti- α -gal IgG did not recognize 248 any band, except for a slight recognition of G18 where XO was present (Fig. 2, Panel A, lane MFGPbDEG). This reactivity completely disappeared only after a more exhaustive overnight PNGase F de-glycosylation (Fig. 2, panel A, lane MFGPbDEGon). The analysis of the bands containing the Nde-glycosylated proteins that lost reactivity revealed which asparagine carried the α -gal moiety (Fig. 2 panel A and B, lane MFGPbDEG). The presence of new tryptic peptides with aspartic acid instead of the original asparagine was considered as proof of the presence of the α -gal sugar chain on the peptide before digestion. The LC-HRMS study of the G22 band showed a BT peptide with Asn₂₁₅ 59 254 60

The MFGP sample was also incubated with the serum of each single patient (Fig. 3). As in previous

experiments, the most recognized bands were G1 (recognized by 7/10 patients), G2 (8/10 patients)

and G4 (8/10 patients), which mainly contain XO and BP. Bands G5 and G8, which showed a reduced

recognition rate, were recognized by 2/10 patients, while G6 was recognized by 3/10 patients and

G9 by only 1 patient. Once again, these bands mainly contained XO, but also LA and β -lactoglobulin.

In order to verify that the patient immunorecognition was addressed to α -gal epitopes,

immunoprecipitation of three patients' sera ($\alpha 2$, $\alpha 3$, $\alpha 5$) was performed with four concentrations

of bovine thyroglobulin (1, 3, 30, 60 μ g) (Fig. 4, Panel A). Only patient α 3 needed 60 μ g of

thyroglobulin to completely inhibit the immunorecognition. Instead, for the other two patients, 3

or 30 μ g was sufficient. The same experiment was performed with bovine XO (patients α 1, α 2, α 7)

(Fig. 4, Panel B). In this case, 60 µg of XO was needed to immunoprecipitate the patients' sera.

Patient $\alpha 2$, who was tested in both inhibition experiments, needed 60 µg of XO and only 3 µg of

3 4	2	5	5
5 6	2	5	6
/ 8 9	2	5	7
10 11 12	2	5	8
13 14 15	2	5	9
16 17	2	6	0
18 19 20	2	6	1
21 22 23	2	6	2
23 24 25	2	6	3
26 27 28	2	6	4
29 30	2	6	5
31 32 33	2	6	6
34 35	2	6	7
36 37 38	2	6	8
39 40	2	6	9
41 42	2	7	0
45 44 45 46	2	7	1
47 48	2	7	2
49 50 51	2	7	3
52 53 54	2	7	4
55 56	2	7	5
57 58 59	2	7	6

thyroglobulin.

Discussion

4.

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255 modified to Asp₂₁₅ after the de-glycosylation protocol. The same was observed in band G24, where
256 LA showed an Asn₂₂₇ modified to Asp₂₂₇. All these results are summarized in Table 3.

3.4 The AGS patients' IgE antibodies recognize Xanthine Oxidase and Butirophilin

Patients with AGS have been known to report allergic manifestations associated with the ingestion

of dairy products, due to the presence of α -Gal carrying proteins, which have recently been

identified in bovine milk whey.^[7,19,22,24] In order to prove that these milk-induced allergic reactions

are due to IgE recognizing α -Gal, it is necessary to exclude other more common causes of reactions

to milk, including lactose intolerance and cow's milk allergy.^[30] In the present work, we have found that milk fat globule associated proteins contain α -Gal epitopes recognized by the specific IgE of patients with AGS. Specifically, we have demonstrated, for the first time, that BT, LA, and XO contained in milk fat globules are α -gal glycosylated. The pool of patients' sera also immune-13 282 recognized milk LF, LPO, and IgG-like proteins, as expected.

16 283 The α -gal-glycosylation of BT, LA, and XO was confirmed by means of an LC-HRMS approach, since new tryptic peptides containing Asp instead of Asn were generated after enzymatic de-21 285 glycosylation, and by means of immunoblotting experiment, since immunorecognition by the anti- α -gal IgG and by AGS patients' sera was lost after de-glycosylation. Although the glycosylation sites of BT and LA had previously been identified by Sato et al.^[31] and by Hvarregaard et al.,^[32] we have 26 287 identified, for the first time, the glycosylation site of XO (Asn₇₀₄ modified to Asp₇₀₄).

No correlations were found between the levels of α -gal sigE and the immuno reaction profile when 34 290 the serum of single patients was tested. This is not surprising, as the presence of elevated IgE levels ³⁶ 291 is indicative of sensitization to α -gal but is not necessarily predictive of a severe allergic reaction. In ₃₉ 292 fact, an allergic manifestation recognizes several triggers that can exacerbate or mask the reaction ⁴¹ 293 itself, thus giving rise to profoundly different clinical pictures.

The role of XO seems to be prevalent, since it was identified in most of the immunoreactive bands, 47 295 especially those separated in the upper part of the gel where no Comassie Blue stained bands were detectable, but both anti- α -gal IgG antibody and AGS patient sera showed the highest ₅₂ 297 immunoreactivity. For this reason, bovine XO was used to perform immunoinhibition experiments 54 298 on three selected patients. XO was able to inhibit immunorecognition by the AGS patient sera as well as thyroglobulin, but a higher amount of protein was needed, and a smaller number of α -gal-59 300 glycosylated sites was indicated for XO than for thyroglobulin.

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5 6 7	302	reco	ognized by
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36 37	314		
38 39 40 41	315	5.	Reference
42 43	316	[1]	Commin
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49 50 51	319	[2]	Fischer J
52 53	320		delayed
54 55 56	321	[3]	Apostolo
50 57 58	322		Allergen
59	323		meat alle

we have found that milk fat globule associated proteins contain α -Gal epitopes the specific IgE of patients with AGS. Previously, Ròman-Carrasco et al.^[16] the presence of α -gal determinants in the lipidic fraction of milk and their ability to tinal monolayer, as well as the potential to trigger allergic reactions in patients with

the patients recruited in the present study, recognized several α -Gal carrying proteins hey and in milk fat globules, although those consuming milk and dairy products seem m. This is not surprising, as IgE reactivity to bovine milk has been reported in 70-90% ts [7,21,24], but the allergic manifestations triggered by dairy products only seem to ne third of patients [20]. Additional host factors are certainly associated with clinical s, and the role of α -Gal carrying glycolipids in reactions to milk and dairy products rther investigated.

erest statement: the authors have no conflicts of interest to disclose.

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s SP, Satinover SM, Hosen J, Mozena J, Borish L, Lewis BD, et al. Delayed anaphylaxis, ema, or urticaria after consumption of red meat in patients with IgE antibodies specific for $e-\alpha-1,3$ -galactose. Journal of Allergy and Clinical Immunology. 2009;123(2):426-433.e2.

J, Yazdi AS, Biedermann T. Clinical spectrum of α -Gal syndrome: From immediate-type to immediate-type reactions to mammalian innards and meat. Allergo J Int. 2016;25(2):55–62.

ovic D, Mihailovic J, Commins SP, Wijnveld M, Kazimirova M, Starkhammar M, et al. omics of the tick Ixodes ricinus reveals important α -Gal–carrying IgE-binding proteins in red ergy. Allergy: European Journal of Allergy and Clinical Immunology. 2020;75(1):217–20.

1		
2 3 324 4	[4]	Boussamet L, Montassier E, Soulillou JP, Berthelot L. Anti $lpha$ 1-3Gal antibodies and Gal content in gut
5 6 325		microbiota in immune disorders and multiple sclerosis. Vol. 235, Clinical Immunology. Academic Press
7 8 326		Inc.; 2022.
9 10 11 327	[5]	Kennedy JL, Stallings AP, Platts-Mills TAE, Oliveira WM, Workman L, James HR, et al. Galactose- α -1, 3-
12 13 328		galactose and delayed anaphylaxis, angioedema, and urticaria in children. Pediatrics. 2013;131(5).
14 15 16 329	[6]	Young I, Prematunge C, Pussegoda K, Corrin T, Waddell L. Tick exposures and alpha-gal syndrome: A
17 18 330		systematic review of the evidence. Ticks Tick Borne Dis. 2021;12(3):101674.
19 20 21 331	[7]	Kiewiet MBG, Apostolovic D, Starkhammar M, Grundström J, Hamsten C, van Hage M. Clinical and
22 23 332		Serological Characterization of the α -Gal Syndrome—Importance of Atopy for Symptom Severity in a
24 25 333 26		European Cohort. Journal of Allergy and Clinical Immunology: In Practice. 2020;8(6):2027-2034.e2.
27 28 334	[8]	van Nunen S. Galactose-Alpha-1,3-Galactose, Mammalian Meat and Anaphylaxis: A World-Wide
29 30 31 335		Phenomenon? Curr Treat Options Allergy. 2014;1(3):262–77.
32 ³³ 336	[9]	Cabezas-Cruz A, Hodžić A, Román-Carrasco P, Mateos-Hernández L, Duscher GG, Sinha DK, et al.
34 35 36 337		Environmental and molecular drivers of the α -Gal syndrome. Front Immunol. 2019;10(MAY):1–12.
37 38 338	[10]	Apostolovic D. Krstic M. Mihailovic J. Starkhammar M. Cirkovic Velickovic T. Hamsten C. et al.
39 40 41 339		Peptidomics of an in vitro digested α -Gal carrying protein revealed IgE-reactive peptides. Sci Rep.
42 43 340		2017;7(1):1–10.
44 45 46 341	[11]	Gonzalez-Quintela A. Dam Laursen AS. Vidal C. Skaaby T. Gude F. Linneberg A. IgE antibodies to alpha-
47 48 342		gal in the general adult population: Relationship with tick bites, atopy, and cat ownership. Clinical and
49 50 343 51		Experimental Allergy. 2014;44(8):1061–8.
52 53 344	[12]	Mateo-Borrega MB, Garcia B, Larramendi CH, Azofra J, González-Mancebo E, Alvarado MI, et al. Ige-
54 55 56 345		mediated sensitization to galactose- α -1,3-galactose (α -gal) in urticaria and anaphylaxis in spain:
57 58 346		Geographical variations and risk factors. J Investig Allergol Clin Immunol. 2019;29(6):436–43.
59 60		

1 2			
2 3 3 4	47	[13]	Villalta D, Cecchi L, Farsi A, Chiarini F, Minale P, Voltolini S, et al. Galactose- α -1,3-galactose syndrome:
53 67	48		An Italian survey. Eur Ann Allergy Clin Immunol. 2017;49(6):263–9.
8 9	49	[14]	Platts-Mills TAE, Commins SP, Biedermann T, van Hage M, Levin M, Beck LA, et al. On the cause and
10 11 12	50		consequences of IgE to galactose- α -1,3-galactose: A report from the National Institute of Allergy and
12 13 3 14	51		Infectious Diseases Workshop on Understanding IgE-Mediated Mammalian Meat Allergy. Journal of
15 3 16 17	52		Allergy and Clinical Immunology. 2020;145(4):1061–71.
18 3 19	53	[15]	Iweala OI, Choudhary SK, Addison CT, Batty CJ, Kapita CM, Amelio C, et al. Glycolipid-mediated
20 3 21	54		basophil activation in alpha-gal allergy. Journal of Allergy and Clinical Immunology. 2020;146(2):450-
²² 23 24	55		2.
²⁴ 25 26	56	[16]	Román-Carrasco P, Lieder B, Somoza V, Ponce M, Szépfalusi Z, Martin D, et al. Only α -Gal bound to
27 28 3	57		lipids, but not to proteins, is transported across enterocytes as an IgE-reactive molecule that can
29 30 3	58		induce effector cell activation. Allergy: European Journal of Allergy and Clinical Immunology.
31 32 3 33	59		2019;74(10):1956–68.
34 35 3 36	60	[17]	Chakrapani N, Fischer J, Swiontek K, Codreanu-Morel F, Hannachi F, Morisset M, et al. α -Gal present
37 3 38	61		on both glycolipids and glycoproteins contributes to immune response in meat-allergic patients.
³⁹ 3 40	62		Journal of Allergy and Clinical Immunology. 2022;150(2):396-405.e11.
41 42 43	63	[18]	Carson AS, Gardner A, Iweala OI. Where's the Beef? Understanding Allergic Responses to Red Meat in
44 45 3	64		Alpha-Gal Syndrome. The Journal of Immunology. 2022;208(2):267–77.
46 47 3	65	[19]	Commins SP Invited Commentary: Alpha-Gal Allergy: Tip of the Iceberg to a Pivotal Immune
48 ³ 49 50 ³	66	[13]	Response. Curr Allergy Asthma Rep. 2016:16(9):1–3.
50 ° 51 52			
53 53 54	67	[20]	Armstrong P, Binder A, Amelio C, Kersh G, Biggerstaff B, Beard C, et al. Descriptive Epidemiology of
55 3 56	68		Patients Diagnosed with Alpha-gal Allergy — 2010–2019. Journal of Allergy and Clinical Immunology.
573 58 59	69		2020;145(2):AB145.
60			

1 2			
2 3 4	370	[21]	Wilson JM, Schuyler AJ, Workman L, Gupta M, James HR, Posthumus J, et al. Investigation into the α -
5 6	371		Gal Syndrome: Characteristics of 261 Children and Adults Reporting Red Meat Allergy. Journal of
7 8 9	372		Allergy and Clinical Immunology: In Practice. 2019;7(7):2348-2358.e4.
10 11 12	373	[22]	Commins SP. Diagnosis & management of alpha-gal syndrome:lessons from 2,500 patients. Expert Rev
13 14	374		Clin Immunol. 2020;16(7):667–77.
15 16 17	375	[23]	Choi JO, Lee MH, Park HY, Jung SC. Characterization of Fabry mice treated with recombinant adeno-
18 19	376		associated virus 2/8-mediated gene transfer. J Biomed Sci. 2010;17(1):1–10.
20 21 22	377	[24]	Perusko M, Apostolovic D, Kiewiet MBG, Grundström J, Hamsten C, Starkhammar M, et al. Bovine γ-
23 24	378		globulin, lactoferrin, and lactoperoxidase are relevant bovine milk allergens in patients with α -Gal
25 26 27	379		syndrome. Allergy: European Journal of Allergy and Clinical Immunology. 2021;76(12):3766–75.
28 29	380	[25]	German-Sanchez A, Alonso-Llamazares A, Latorre-Ibañez M, Bartolome-Zavala B, Antepara-Ercoreca
30 31 32	381		I. Sheep cheese allergy in Alpha-gal Syndrome. J Investig Allergol Clin Immunol. 2023;33(6):1–7.
33 34	382	[26]	Barello C, Garoffo LP, Montorfano G, Zava S, Berra B, Conti A, et al. Analysis of major proteins and fat
35 36 37	383		fractions associated with mare's milk fat globules. Mol Nutr Food Res. 2008;52(12):1448–56.
38 39	384	[27]	TRETTER V, ALTMANN F, MÄRZ L. Peptide-N4-(N-acetyl-β-glucosaminyl) asparagine amidase F cannot
40 41 42	385		release glycans with fucose attached $\alpha 1 \rightarrow 3$ to the asparagine-linked N-acetylglucosamine residue.
43 44	386		Eur J Biochem. 1991;199(3):647–52.
45 46 47	387	[28]	Cirrincione S, Aiuto B, Gosso E, Schiavone C, Portesi C, Rossi AM, et al. Proteomic study of walnut
48 49	388		oleosome and first evidence on oleosin sensitization in allergic patients. Journal of Food Composition
50 51 52	389		and Analysis. 2023;121(March):105386.
53 54	390	[29]	Loprinzi Brauer CE, Motosue MS, Li JT, Hagan JB, Bellolio MF, Lee S, et al. Prospective Validation of
55 56	391		the NIAID/FAAN Criteria for Emergency Department Diagnosis of Anaphylaxis. J Allergy Clin Immunol
57 58 59	392		Pract. 2016;4(6):1220–6.
60			

1 2			
3 4	393	[30]	Binder AM, Cherry-Brown D, Biggerstaff BJ, Jones ES, Amelio CL, Beard CB, et al. Clinical and laboratory
5 6	394		features of patients diagnosed with alpha-gal syndrome-2010-2019. Allergy: European Journal of
7 8 9	395		Allergy and Clinical Immunology. 2023;78(2):477–87.
10 11 12	396	[31]	Sato T, Takio K, Kobata A, Greenwalt DE, Furukawa K. Site-Specific Glycosylation of Bovine
12 13 14	397		Butyrophilin. The Journal of Biochemistry. 1995;117(1):147–57.
15 16 17	398	[32]	Hvarregaard J, Andersen MH, Berglund L, Rasmussen JT, Petersen TE. Characterization of glycoprotein
18 19	399		PAS-6/7 from membranes of bovine milk fat globules. Eur J Biochem. 1996;240(3):628–36.
20 21 22	400		
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402 **Figure legends**

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404 Figure 1. Investigation of the three bovine milk fractions: caseins (CAS), whey proteins (WP) and milk fat 11 405 globule associated proteins (MFGP). Panel A: LDS page of MGFP, WP and CAS. Panel B: immunoblotting of 406 MFGP, WP and CAS with the anti- α -Gal IgG antibody. Panel C: immunoblotting of MFGP, WP and CAS with 407 the sera of a pool of 10 AGS patients. M: molecular weight markers; C+: thyroglobulin; CII: secondary antibody 18 408 control.

20 21 409 **Figure 2.** Investigation of α -gal bovine milk proteins. Panel A: LDS page of milk fat globule associated proteins. 22 23 410 (MFGP). MFGP enriched by means of incubation with beads bound with anti- α -gal IgG (MFGPb) and MGFPb ²⁵ 411 de-glycosilated with PNGase for 3 hours (MFGPbDEG) and ON. (MFGPbDEGon). Panel B: immunoblotting of 28 412 MFGP. MFGPb. MFGPbDEG. and MFGPbDEGon with anti- α -gal IgG. M: molecular weight; C+: thyroglobulin; 30 413 CII: secondary antibody control.

32 ₃₃ 414 **Figure 3.** Recognition of milk fat globule associated proteins (MFGP) by α -gal syndrome (AGS) patients. 34 Immunoblotting of MFGP with the sera of 10 AGS patients (from $\alpha 1$ to $\alpha 10$). M: molecular weight marker; 35 415 36 37 416 C+: thyroglobulin. C-: patient not assuming meat. negative control CII: secondary antibody control. 38

40 417 Figure 4. Immunoprecipitation experiments of α -gal syndrome (AGS) patient's sera. Panel A: Immunoblotting 41 ⁴² 418 of milk fat globule associated proteins (MFGP) with the sera of three patients ($\alpha 2$. $\alpha 3$. $\alpha 5$) 43 44 419 immunoprecipitated with different concentrations of thyroglobulin (1. 3. 30. and 60 μ g). Panel B: 45 46 47 420 Immunoblotting of MFGP with the sera of three patients ($\alpha 1$. $\alpha 2$. $\alpha 7$) immunoprecipitated with different 48 49 421 concentrations of bovine xanthine oxidase (3. 30. and 60 μg). M: molecular weight marker; C-: patient not 50 ⁵¹ 422 assuming meat; CII: secondary antibody control.

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425 Table 1. Characterization of the patients.

5 7 3 9 10 11	ID PATIENT	SEX	AGE	CULPRIT FOOD	CLINICAL presentation	SYMPTOM S ONSET	TREATMENT	CO- FACTORS	TICK BITES	TOTAL SERUM IGE (KU/L)	ALPHA GAL IGEs (KUA/L)	TRYPTASE (ug/L)
13 14 15 16 17 18 19 20 21	Alpha1	M	37	Since 2020: always after eating red meat	Anaphylaxis: angioedema, urticaria, dyspnea	3-4 hours	Ebastine 10 mg at home Betamethas one 4mg at home	Physical exercise	Not known	117	3.08	5.50
22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43	Alpha2	F	74	9/2017: soup with egg pasta and veal kidney 3/2018: tripe	Anaphylaxis: diffuse erythema, hypotension Generalized urticaria	3 hours 3 hours	Chlorphena mine 10 mg (im) Betamethas one 4 mg (iv) Adrenaline 0.5 mg + 0.5 mg (im) Betamethas one 4mg at home Ebastine 10 mg at home	None	At least one tick bite in the past	117	0.37	6.19
14 15 16 17 18 19 50 51 52 53 54 55 56 57 58 59 50	Alpha3	F	69	9/2017: boiled meat, anchovies (parsley and garlic) and soup with beef broth 12/2018:	Anaphylaxis: generalized urticaria, palpebral oedema and vomiting Severe	5 hours 2 hours	Chlorphena mine 10 mg (im) Betamethas one 4 mg (iv) O2-support	None	Previous tick bites not known	702	86.50	6.30

2												
3				offal and	anaphylaxis:		Chlorphena					
4 5				bread.	headache,		mine 10 mg					
6					loss of		(im)					
8					consciousness		Methylpredn					
9 10					, hives,		isolone 80					
11					vomiting,		mg (iv)					
12 13					respiratory		Adrenaline					
14					failure		0.5 mg (im)					
16												
17				5/2019: 1	Severe	2 hours	Chlorphena					
18				spoonful of	anaphylaxis:		mine 10 mg					
20				veal broth.	generalized		(im)					
22					urticaria, loss		Methylpredn					
23 24					of		isolone 80					
25					consciousness		mg (iv)					
26 27							Adrenaline					
28						\bigcirc	0.5 mg (im)					
29 30												
31		М	68	Since 2018:	Urticaria	5-6 hours	Betamethas	None	Many	130	15.01	<1
32 33				after eating			one 4mg at		tick bites			
34 35				offal			home		in the			
36	Alpha4			2022	Itching,	6 hours	Ebastine 10		past			
37 38				stuffed	urticaria		mg at home					
39				pasta				\sim				
40 41				(meat)				4				
42												
43 44		F	74	8/2017:	Anaphylaxis:	3-4 hours	Chlorphena	None	Previous	1383	2.54	8.13
45				lamb stew	generalized		mine 10 mg		tick bites			
40 47				and pasta	itching,		(im)		not			
48 49				with red	urticaria,		Methylpredn		known			
50				sauce	vomiting		isolone 100					
51 52	Alpha5						mg (iv)					
53							Adrenaline					
54 55							0.5 mg (im)					
56				7/2010	Canasali	4 h a	Theat's 10					
58				//2018:	Generalized	4 nours	Ebastine 10					
59 60				pasta with	urticaria and		mg at home					
00				tomato,								

2												
3				lamb liver	stomach-ache							
4 5				and lung								
6 7				(peppers								
8				and onions)								
9												
10 11		М	66	5/2018:	Severe	1 hour and	O2-support	Clonal	Many	309	11.60	10.20
12				carrots,	anaphylaxis:	half	Chlorphena	mast cell	tick bites			
13 14				"capocollo"	abdominal		mine 10 mg	disorder	in the			
15 16				and wine	pain, nausea,		(im)		past			
17					vomiting,		Methylpredn					
18 19					diarrhea,		isolone 250					
20					flushing, loss		mg (iv)					
21					of		Adrenaline					
23 24	Alpha6				consciousness		0.5 mg (im)					
25				3 more			Not available					
26 27				similar								
28 29				episodes								
30				after								
31 32				ingestion of								
33 34				pork or			2					
35				offal.								
36 37							Ľ					
38		М	57	2010-2014:	Recurrent	3-4 hours	Ebastine 10	None	Many	764	>100	14.60
39 40				gummy	urticaria		mg at home		tick bites			
41 42				bears				2	in the			
43				2016: red	Anaphylaxis:	4-5 hours	Chlorphena		past			
44 45	Alpha7			meat	urticaria,		mine 10 mg					
46 47					hypotension,		(im)					
47 48					diarrhea		Methylpredn					
49 50							isolone 250					
51				2017:	Urticaria,	6 hours	mg (iv)					
52 53				rabbit liver	angioedema							
54		M	58	In 2018:	Anaphylaxis:	Not known	Chlorphena	None	Many	162	31.50	9.12
56				always	angioedema,		mine 10 mg		tick bites			
57 58	Alpha8			after eating	urticaria,		(im)		in the			
59 60				meat	abdominal		Methylpredn		past			
00												

		_										
					pain,		isolone 250					
					hypotension		mg (iv)					
		F	26	Since 2015:	Recurrent	6-7 hours	Betamethas	None	At least	52.9	1.17	3.07
				after eating	urticaria and		one 4mg at		one tick			
	Alpha9			red meat	abdominal		home		bite in			
					pain		Ebastine 10		the past			
							mg at home					
		М	67	Since 2010:	Anaphylaxis:	5-6 hours	Chlorphena	None	Not	163	8.95	7.73
				always	urticaria,		mine 10 mg		known			
				after eating	ducanaa		(im)					
				arter eating	uyspnea,		(111)					
				meat	peripheral		Methylpredn					
	Alpha10				edema)		isolone 250					
							mg (iv)					
							Adrenaline					
							0.5 mg + 0.5					
							mg (im)					
		M	48	Since 2019:	Diffuse itching	2-12 hours	Cetirizine 10	None	Not	78	<0.10	3.80
				milk and	and small		mg at home		known			
	Healthy			mink and	dilu silidii		ing at nome		KHOWH			
	control			dairy	wheals							
	2011101			products,								
				vegetarian								
				-0				1				
26 ¹	F: female;	M: male	e; im: int	ramuscular; iv: i	ntravenous.	1			1	1	1	I

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Tab 2. Identification of the proteins immunorecognized by anti- α -Gal IgG and/or by the pool of α -

Gal syndrome patient's sera in the milk fat globule protein, whey protein and casein fractions.

5)		- .				N° of matching	Protein
0 1	N° Band	Entry	Protein name	[Da]	Protein Score	peptides	coverage [%]
12	G1	P80457	Xanthine dehydrogenase	300000/142330	130.480	16	16
3 4		Q8WNR8	Perilipin	300000/45251	52.718	8	27.9
15		Q27960	Sodium-dependent phosphate transport	300000/75825	52.365	5	6.6
17			protein 2B				
18 19		Q4GZT4	Broad substrate specificity ATP-binding	300000/72724	43.296	7	15
20			cassette transporter ABCG2				
22		P18892	Butyrophilin subfamily 1 member A1	300000/59231	25.794	4	11
23 24		A0A4W2HXW4	3-hydroxyacyl-[acyl-carrier-protein]	300000/268170	12.606	2	1.1
25			dehydratase				
20 27	G2	P80457	Xanthine dehydrogenase	130000/146790	317.140	30	30.1
28 29		P18892	Butyrophilin subfamily 1 member A1	130000/59231	59.231	10	24.7
30		A0A4W2I0L9	ATP-binding cassette sub-family G	130000/67774	34.445	4	6
31 32			member 2				
33	G3	G5E513	Ig-like domain-containing protein	80000/48107	31.553	9	23.2
35 35		G5E5T5	Ig-like domain-containing protein	80000/55968	129.030	10	22.4
36 37		A0A3Q1M193	Glycoprotein 2	80000/58465	260.530	8	17.3
38		P18892	Butyrophilin subfamily 1 member A1	80000/59276	145.460	10	24.1
39 10		C7FE01	Lactoferrin	80000/80278	55.906	8	12.8
41 12		P80457	Xanthine dehydrogenase	80000/142330	32.807	5	3.8
13	G4	P81265	Polymeric immunoglobulin receptor	68000/82434	211.620	18	35.1
14 15		A0A3Q1M193	Glycoprotein 2	68000/58465	92.215	10	23.8
16 17		P18892	Butyrophilin subfamily 1 member A1	68000/59276	106.110	15	41.3
+7 18		P26201	Glycoprotein IIIb	68000/46055	91.212	6	12.9
19 50		G5E513	Ig-like domain-containing protein	75000/48107	95.157	9	33.3
51		A0A3Q1LWT4	Acyl-CoA synthetase long chain family	68000/81442	79.564	10	18
52 53			member 1				
54		J7K1V4	Lactoferrin	68000/80278	75.774	12	18.6
55 56		F1MHI1	Perilipin	80000/45281	53.926	7	25.5
57 58		Q27960	Sodium-dependent phosphate transport	68000/75825	32.389	5	9.7
59			protein 2B				
50							

	Q95114	Milk fat globule-EGF factor 8 protein	68000/37465	32.227	4	17
		(Lactadherin)				
	A0A3Q1MK38	Terpene cyclase/mutase family member	68000/74156	52.104	5	8.6
	P80457	Xanthine dehydrogenase	68000/142330	30.771	5	3.8
G5	P18892	Butyrophilin subfamily 1 member A1	60000/59276	252.710	19	31.4
	Q95114	Milk fat globule-EGF factor 8 protein	60000/43140	50.477	6	19
		(Lactadherin)				
G6	Q95114	Milk fat globule-EGF factor 8 protein	51000/43140	198.570	22	57.2
		(Lactadherin)				
	Q9TUM6	Perilipin-2	51000/49368	189.240	19	59.5
	P18892	Butyrophilin subfamily 1 member A1	51000/59276	31.353	5	13.3
G7	Q95114	Milk fat globule-EGF factor 8 protein	49000/43140	231.350	13	35.3
		(Lactadherin)				
	Q8HZM7	Perilipin	49000/45281	55.801	4	18
G8	P02663	Alpha-S2-casein	34000/26018	41.439	6	18.9
	P18892	Butyrophilin subfamily 1 member A1	34000/59231	47.768	5	12.5
G9	B5B0D4	Major allergen beta-lactoglobulin	19000/19969	116.590	11	65.2
	Q5E9I6	ADP-ribosylation factor 3	19000/20601	47.494	7	45.9
	Q8HZM7	Perilipin	19000/45281	35.690	5	18.9
G10	P80457	Xanthine dehydrogenase	300000/142330	37.778	5	6.4
	A0A3Q1MGL5	SRCR domain-containing protein	300000/35988	16.338	2	23.1
	A0A4W2HXW4	3-hydroxyacyl-[acyl-carrier-protein]	300000/268170	13.596	2	1.4
		dehydratase		\sim		
	Q27960	Sodium-dependent phosphate transport	300000/75825	14.233	2	2.7
		protein 2B				
G11	P80457	Xanthine dehydrogenase	170000/142330	97.052	11	13
G12	P80457	Xanthine dehydrogenase	130000/146790	167.660	20	21.4
G13	P80457	Xanthine dehydrogenase	116000/14233	103.510	11	9.4
	Q27960	Sodium-dependent phosphate transport	116000/75825	35.245	2	2.2
		protein 2B				
G14	G5E5T5	Immunoglobulin heavy constant mu	80000/56043	157.780	12	32.1
	F1MZQ4	Butyrophilin subfamily 1 member A1	80000/59231	65.440	7	15.8
G15	A0A4W2DWX4	Butyrophilin subfamily 1 member A1	60000/59245	94.962	13	25.9
G16	P0DOX5	Immunoglobulin gamma-1 heavy chain	53000/49328	97.277	10	31.2
	Q95114	Milk fat globule-EGF factor 8 protein	53000/37465	34.165	5	16.7
		(Lactadherin)				
L	1		1	1	1	1

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G17	P01834	Immunoglobulin kappa constant	28000/11765	59.743	5	67.3
	А5РК49	Ig-like domain-containing protein	28000/24592	30.416	4	12.8
G18	A0A4W2I0L9	ATP-binding cassette sub-family G	300000/67774	11.869	2	3.1
		member 2				
	P80457	Xanthine dehydrogenase	300000/146690	17.852	3	1.9
Q27960		Sodium-dependent phosphate transport	300000/75825	42.271	2	2.2
		protein 2B				
G19	P80457	Xanthine dehydrogenase	130000/146790	292.240	32	27
G20	G5E513	Immunoglobulin heavy constant mu	60000/56043	84.106	10	29.2
	P81265	Polymeric immunoglobulin receptor	60000/82434	65.441	9	17.6
	P18892	Butyrophilin subfamily 1 member A1	60000/59276	51.220	8	20
G21	F1MZQ4	Butyrophilin subfamily 1 member A1 57000/59231 63.366		7	17.5	
G22	P18892	Butyrophilin subfamily 1 member A1	55000/59231	143.590	16	35.6
G23	Q9TUM6	Perilipin-2	48000/49368	83.058	8	28.9
	P18892	Butyrophilin subfamily 1 member A1	48000/59276	45.368	6	12.7
G24	Q95114	Milk fat globule-EGF factor 8 protein	44000/43140	137.390	16	39.2
		(Lactadherin)	D.			
	P18892	Butyrophilin subfamily 1 member A1	44000/59231	31.275	5	9.9
G25	P21163.2	Peptide-N(4)-(N-acetyl-beta-D-	40000/39032	227.360	16	52.5
		glucosaminyl)asparagine amidase				
		PNGase F				
	P18892	Butyrophilin subfamily 1 member A1	40000/59276	49.619	5	11.2
W1	P80457	Xanthine dehydrogenase	130000/146790	323.310	20	18.2
W2	A0A4W2CZN6	C3 complement	110000/190950	308.810	32	20.3
	A0A3Q1M3L6	Ig-like domain-containing protein	110000/40475	106.250	7	29.7
W3	C7FE01	Lactoferrin	75000/76274	323.310	45	66.5
	G5E513	Ig-like domain-containing protein	75000/48107	307.500	16	54.2
	G3X6N3	Serotransferrin	75000/77738	117.080	22	39.9
	P80025	Lactoperoxidase	75000/71350	187.390	22	39.9
	A0A3Q1M3L6	Ig-like domain-containing protein	75000/40475	44.510	4	19.3
	B3VTM3	Lactotransferrin	75000/78056	45.075	7	13
	A0A3Q1LNN7	Albumin	75000/68198	32.956	5	9.2
W4	P81265	Polymeric immunoglobulin receptor	68000/82434	134.530	11	25
	A0A4W2DZ09	Serotransferrin	68000/77738	133.090	15	33.5
	E1BMJ0	Serpin family G member 1	68000/51772	95.139	5	17.9
	A0A4W2CZN6	C3-beta-c	68000/190950	79.946	10	9

A0A3Q1M032	Ig-like domain-containing protein	68000/40475	92.036	4	16.800
A0A4W2DDL5	Albumin	68000/68198	60.754	8	18.9
A0A4W2GX34	Lactoperoxidase	68000/71350	33.890	5	9.2
P02769	Albumin	60000/68198	323.310	41	64.7
A0A4W2CZN6	C3 complement	60000/190950	244.450	28	22.2
Q2KJF1	Alpha-1B-glycoprotein	60000/39566	75.560	9	36.2
A0A3Q1M3L6	Ig-like domain-containing protein	50000/40475	148.910	10	52.9
G3N0V0	Ig-like domain-containing protein	50000/35951	49.249	6	25.2
Q9TTE1	Serpin A3-1	50000/46236	75.075	7	25.3
A0A4W2HXY3	Serpin A3-1	50000/46815	33.062	5	17.9
A0A140T8A9	Kappa-casein	50000/21237	30.867	3	23.7
A0A3Q1NG86	Alpha-S1-casein	50000/23689	30.935	3	18
P02754	Beta-lactoglobulin	50000/19883	32.047	3	22.5
A0A4W2FAA0	Antithrombin-III	50000/52456	32.029	5	11.8
P08037-2	Isoform Short of Beta-1.4-	50000/43483	79.095	2	5.1
	galactosyltransferase 1				
P02754	Beta-lactoglobulin	15000/19883	31.959	4	32.1
P00711	Alpha-lactalbumin	15000/14156	144.240	3	24.4
P80195	Glycosylation-dependent cell adhesion	15000/17151	31.741	2	12.4
	molecule 1				
P24627	Lactotransferrin	75000/78056	323.310	47	61.90
P18892	Butyrophilin subfamily 1 member A1	75000/59276	41.743	3	12.20
G5E513	Ig-like domain-containing protein	75000/48106	32.696	3	9.30
P02662	Alpha-S1-casein	27000/23689	323.310	8	42.2
P80195	Glycosylation-dependent cell adhesion	27000/17151	93.318	2	12.4
	molecule 1				
A0A140T8A9	Kappa-casein	27000/21237	190.770	4	30.5
A0A452DHW7	Beta-casein	27000/29221	62.074	5	18.5
					I
P02754	Beta-lactoglobulin	27000/19883	61.784	5	37.1
	A0A3Q1M032 A0A4W2DDL5 A0A4W2CXN6 Q2KJF1 A0A4W2CZN6 Q2KJF1 A0A3Q1M3L6 G3N0V0 Q9TTE1 A0A4W2HXY3 A0A140T8A9 A0A3Q1NG86 P02754 A0A4W2FAA0 P08037-2 P08037-2 P02754 P00711 P80195 P24627 P18892 G5E513 P02662 P80195	A0A3Q1M032Ig-like domain-containing proteinA0A4W2DDL5AlbuminA0A4W2GX34LactoperoxidaseP02769AlbuminA0A4W2CZN6C3 complementQ2KJF1Alpha-1B-glycoproteinA0A3Q1M3L6Ig-like domain-containing proteinG3N0V0Ig-like domain-containing proteinQ9TTE1Serpin A3-1A0A4W2HXY3Serpin A3-1A0A4W2HXY3Serpin A3-1A0A4W2HXY3Serpin A3-1A0A4W2FAA0Altpha-S1-caseinP02754Beta-lactoglobulinA0A4W2FAA0Antithrombin-IIIP08037-2Isoform Short of Beta-1.4- galactosyltransferase 1P02754Beta-lactoglobulinP00711Alpha-lactalbuminP80195Glycosylation-dependent cell adhesion molecule 1P18892Butyrophilin subfamily 1 member A1G5E513Ig-like domain-containing proteinP02662Alpha-S1-caseinP80195Glycosylation-dependent cell adhesion molecule 1A0A140T8A9Kappa-casein	A0A3Q1M032Ig-like domain-containing protein68000/40475A0A4W2DL5Albumin68000/68198A0A4W2GX34Lactoperoxidase68000/71350P02769Albumin60000/68198A0A4W2CZN6C3 complement60000/190950Q2KJF1Alpha-1B-glycoprotein60000/39566A0A3Q1M3L6Ig-like domain-containing protein50000/40475G3N0V0Ig-like domain-containing protein50000/40475G3N0V0Ig-like domain-containing protein50000/46236A0A4W2HXY3Serpin A3-150000/46815A0A140T8A9Kappa-casein50000/21237A0A3Q1NG86Alpha-S1-casein50000/23689P02754Beta-lactoglobulin50000/52456P08037-2Isoform Short of Beta-14- galactosyltransferase 150000/19883P00711Alpha-lactalbumin15000/19883P00711Alpha-lactarbglobulin50000/17151molecule 175000/78056P18892Butyrophilin subfamily 1 member A175000/78056P18892Ig-like domain-containing protein75000/78056P02662Alpha-S1-casein27000/23689P80195Glycosylation-dependent cell adhesion molecule 127000/21237A0A140T8A9Kappa-casein27000/21237	A0A3Q1M032 Ig-like domain-containing protein 68000/04475 92.036 A0A4W2DDL5 Albumin 68000/68198 60.754 A0A4W2GX34 Lactoperoxidase 68000/71350 33.890 P02769 Albumin 60000/8198 323.310 A0A4W2CZN6 C3 complement 60000/190950 244.450 Q2KJF1 Alpha-1B-glycoprotein 60000/39566 75.560 A0A3Q1M3L6 Ig-like domain-containing protein 50000/40475 148.910 G3N0V0 Ig-like domain-containing protein 50000/40475 148.910 G3N0V0 Ig-like domain-containing protein 50000/40475 33.062 A0A4W2HXY3 Serpin A3-1 50000/46815 33.062 A0A4W2HXY3 Serpin A3-1 50000/21237 30.867 A0A3Q1NG86 Alpha-S1-casein 50000/21883 32.047 A0A4W2FAA0 Antithrombin-III 50000/23689 30.935 P02754 Beta-lactoglobulin 50000/19883 31.959 P02754 Beta-lactoglobulin 15000/19156 144.240	A0A3Q1M032 Ig-like domain-containing protein 68000/40475 92.036 4 A0A4W2DDL5 Albumin 68000/68198 60.754 8 A0A4W2GX34 Lactoperoxidase 68000/71350 33.890 5 P02769 Albumin 60000/68198 323.310 41 A0A4W2CZN6 C3 complement 60000/190950 244.450 28 Q2XIF1 Alpha-1B-glycoprotein 50000/40475 148.910 10 G3N0V0 Ig-like domain-containing protein 50000/46236 75.075 7 A0A4W2HXY3 Serpin A3-1 50000/46236 75.075 7 A0A40W2HXY3 Serpin A3-1 50000/46236 33.062 5 A0A140T8A9 Kappa-casein 50000/21237 30.867 3 A0A3Q1M3E6 Alpha-51-casein 50000/21237 30.867 3 A0A4W2FAA0 Antithrombin-III 50000/24883 32.047 3 A0A4W2FAA0 Antithrombin-III 50000/24883 31.959 4 P00711 Alpha-lactal

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Tab 3. Analysis of the xanthine oxidase. butyrophilin and lactadherin glycosylation sites by means

433 of LC-HRMS.

		THEORET	ICAL DATA	LC-HRMS EXPERIMENTAL DATA		
α	t-gal MFGP	N-glycosylated triplets	triplets already known	Peptide-containing triplet before	Peptide-containing modified	
			from literature	enzymatic de-glycosylation	triplet (N>D)	
		N ₆₄₄ ET	not	not found	not found	
		N ₇₀₄ NS	not	not found	704-713 (D ₇₀₄ NS)	
x	O (P80457)	N ₉₀₄ LS	yes (in goat)	903-912 (N ₉₀₄ LS)	not found	
		N ₁₀₇₃ SS	yes (in human)	not found	not found	
		N ₁₂₈₈ NT	not	1283-1290 (N ₁₂₈₈ NT)	not found	
		N55VS	yes (in cow)	not found	not found	
В	ST (P18892)	N215VS	yes (in cow)	not found	215- 221 (D215VS)	
		N337MT	not	not found	not found	
		N ₅₉ ET	yes (in cow)	not found	not found	
	A (OOE114)	N ₁₄₄ NS	not	138-149 (N ₁₄₄ NS)	not found	
	A (U95114)	N ₂₂₇ NS	yes (in cow)	not found	221-232 (D ₂₂₇ NS)	
		N ₃₉₀ NS	not	382-395 (N ₃₉₀ NS)	not found	

36 435 N: asparagine; D: aspartic acid; MFGP: milk fat globule protein; XO: xanthine oxidase; BT: butyrophilin; LA: lactadherin.

³⁸ 39 436



200 116 97 W4 66 W5 G5 55 G7 36 31 21 14 М MFGP WP CAS T (+) MFGP WP CAS C+ CII MGFP WP CAS CII

Figure 1. Investigation of the three bovine milk fractions: caseins (CAS), whey proteins (WP) and milk fat globule associated proteins (MFGP). Panel A: LDS page of MGFP, WP and CAS. Panel B: immunoblotting of MFGP, WP and CAS with the anti-α-Gal IgG antibody. Panel C: immunoblotting of MFGP, WP and CAS with the sera of a pool of 10 AGS patients. M: molecular weight markers; C+: thyroglobulin; CII: secondary antibody control.

157x222mm (300 x 300 DPI)

В

MFGPbDEGon

₽

t

MFGPbDEG

MFGPb

MFGP

А

Σ

G16

G25

MFGPbDEGon MFGPbdeg

ţ

Figure 2. Investigation of a-gal bovine milk proteins. Panel A: LDS page of milk fat globule associated

proteins (MFGP). MFGP enriched by means of incubation with beads bound with anti-a-gal IgG (MFGPb) and

MGFPb de-glycosilated with PNGase for 3 hours (MFGPbDEG) and ON. (MFGPbDEGon). Panel B:

immunoblotting of MFGP. MFGPb. MFGPbDEG. and MFGPbDEGon with anti-a-gal IgG. M: molecular weight;

C+: thyroglobulin; CII: secondary antibody control.

157x222mm (300 x 300 DPI)

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MFGPb

MFGP







Figure 3. Recognition of milk fat globule associated proteins (MFGP) by a-gal syndrome (AGS) patients. Immunoblotting of MFGP with the sera of 10 AGS patients (from a1 to a10). M: molecular weight marker; C+: thyroglobulin. C-: patient not assuming meat. negative control CII: secondary antibody control.

175x222mm (300 x 300 DPI)

1 2 3 4 5 6 7 8 9	200 A Pattern #2
11 12 13 14 15 16 17 18 19 20	116
21 22 23 24 25 26 27 28 29 30	Figure 4. Immunop Immunoblotting of milk fa immunoprecipitated v Immunoblotting of MFGI concentrations of bovine x
30 31 32 33 34 35 36 37 38 20	
 39 40 41 42 43 44 45 46 47 	
48 49 50 51 52 53 54 55 56	
57 58 59	

Patient as Patient al Patient a7 TQ 30 60 TQ 1 30 TQ 3 30 60 τQ 3 30 60 то 3 30 60 C-CI 1 3 3

Figure 4. Immunoprecipitation experiments of α-gal syndrome (AGS) patient's sera. Panel A: Immunoblotting of milk fat globule associated proteins (MFGP) with the sera of three patients (α2. α3. α5) immunoprecipitated with different concentrations of thyroglobulin (1. 3. 30. and 60 μg). Panel B: Immunoblotting of MFGP with the sera of three patients (α1. α2. α7) immunoprecipitated with different concentrations of bovine xanthine oxidase (3. 30. and 60 μg). M: molecular weight marker; C-: patient not assuming meat; CII: secondary antibody control.

157x88mm (300 x 300 DPI)

Graphical Abstract Text

- 10-20% of patients affected by α -Gal syndrome (AGS) also react towards bovine milk.
- Alpha-gal glycosylated milk proteins were recognized by the IgE of AGS patients.
- Xanthine oxidase, butyrophillin and lactoadherin were found to be α -gal glycosylated.
- Xanthine oxidase is the milk protein most immunorecognized by the IgE of AGS patients.

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769x333mm (130 x 130 DPI)