RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2018, 8, 26837

The polysaccharide of Atractylodes macrocephala koidz (PAMK) alleviates cyclophosphamide-mediated immunosuppression in geese, possibly through novel_mir2 targeting of CTLA4 to upregulate the TCR-NFAT pathway

Wanyan Li, (10 † ab Danning Xu, † a Bingxin Li, a Nan Cao, a Sixuan Guo, a Qingyan Jiang*b and Yunbo Tian*a

The polysaccharide of Atractylodes macrocephala koidz (PAMK) has been proved to have antioxidant, antiinflammatory, antiviral, and immunity promoting effects. MicroRNAs (miRNAs) have also been shown to participate in the regulation of immune function by negatively regulating the expression of target genes. However, little is known about how PAMK alleviates the immunosuppression via the miRNA pathway in geese. The aim of this study is to evaluate the influence of PAMK on immunosuppression. Magang geese (1 day old, n = 200) were randomly divided into groups, namely, the control group (normal feeding), PAMK (fed 400 mg kg^{-1} PAMK), CTX (injected 40 mg kg^{-1} BW cyclophosphamide), and CTX + PAMK $(40 \text{ mg kg}^{-1} \text{ BW cyclophosphamide} + 400 \text{ mg kg}^{-1} \text{ PAMK})$ groups. Thymus development was examined by the thymus index, transmission electron microscopy and scanning electron microscopy. The T cell proliferation rate was stimulated by phytoagglutinin (PHA), and T cell activation related genes (CD28, CD96, MHC-II), and IL-2 levels in serum were detected. Differentially expressed miRNAs of geese to regulate T cell activation were found by miRNA sequencing technologies. The results showed that PAMK could alleviate thymus damage and the decrease in the T lymphocyte proliferation rate, T cell activation, and IL-2 levels that were induced by CTX. MiRNA sequencing found that the combination of PAMK and CTX significantly promoted T cell activation via upregulation of novel_mir2 (P < 0.05), which inhibited cytotoxic T lymphocyte antigen 4 (CTLA4) expressions, thereby promoting the TCR-NFAT signaling pathway. It can be concluded that PAMK, through novel_mir2 targeting of CTLA4 to upregulate TCR pathway, finally alleviated immunosuppression induced by CTX in geese.

Received 12th January 2018 Accepted 7th July 2018

DOI: 10.1039/c8ra00368h

rsc.li/rsc-advances

Introduction

Many polysaccharides from Chinese herbal medicine have been shown to promote immune function as well as alleviate cyclophosphamide-induced immunosuppression. The polysaccharide of *Atractylodes macrocephala koidz* (PAMK) is an immune enhancer that improves immune function and has anti-oxidant and anti-inflammatory effects. PAMK modulates immune function by promoting immune organ development and immune cell activation, as well as increasing lymphocyte proliferation and cytokine secretion. Pacent research also revealed that another important biological effect of PAMK is to resist heat stress and maintain the immune function at a stable

level.^{5,12,13} However, there has been no report about the effects of PAMK on alleviating immunosuppression.

MicroRNAs (miRNAs) are small non-coding RNAs that regulate gene expression after transcription by selectively degrading mRNA or inhibiting translation. Studies have shown that many miRNAs are involved in the regulation of immune functions, and some of them are cell-type specific. ¹⁴ MiR-15b, miR-181, miR-31, miR-126, miR-155, and miR146a are involved in the regulation of T cell activation and proliferation, secretion of cytokines, and other related functions. ¹⁵⁻¹⁸ MiRNAs regulate T cell activation mainly by targeting different genes in the T cell receptor (TCR) pathway. It was reported that miR-181 targeted multiple phosphatases, miR-21 targeted activator protein-1 (AP-1) and signal transducer and activator of transcription 3 (STAT3), miR-155 and miR-429 targeted cytotoxic T lymphocyte antigen 4 (CTLA4) in the TCR pathway and finally changed T cell activation. ^{17,19-22}

Although it has been confirmed that miRNAs are involved in regulating immune function, the role of miRNAs in the

^eGuangdong Province Key Laboratory of Waterfowl Healthy Breeding, Guangzhou 510225, P. R. China. E-mail: tyunbo@126.com

^bGuangdong Provincial Key Laboratory of Animal Nutrition Control, College of Animal Science, South China Agricultural University, Guangzhou 510642, P. R. China

[†] These authors contributed equally to this work.

mechanism of PAMK that promotes immune function, espe-

cially the alleviation of immunosuppression in geese, has not been clarified. In this study, an immunosuppressed model of geese was constructed by injecting cyclophosphamide (CTX), and the effects of PAMK on the alleviation of immunosuppression were studied in the combination of PAMK and CTX. We also focus on the difference in miRNAs between CTX and CTX + PAMK to find out the mechanisms of PAMK in the alleviation of immunosuppression.

Materials and methods

Animals

RSC Advances

One-day-old geese (Anser cygnoides) were purchased from Guangdong Qingyuan Jinyufeng Goose Co. Ltd., which is a professional goose hatching company, and were housed in a specific pathogen-free environment. The geese had free access to food (including the same amount of vegetables) and water, were treated humanely, and the experiments received prior ethical approval in accordance with Zhongkai University of Agriculture and Engineering, and under the approved protocol number SRM-11.

Reagents

PAMK (purity 70%) was purchased from Tianyuan (Xi'an, China), diluted with water and sprayed on the feed according to the concentration of 400 mg kg⁻¹. Cyclophosphamide (Jiangsu Hengrui Pharmaceutical, China), was dissolved in physiological saline and intraperitoneally injected in the geese at a concentration of 40 mg kg⁻¹ BW. Lymphocyte separation fluid was purchased from Tianjin Haoyang Biological Manufacturers. Phytoagglutinin (PHA; Sigma, USA), as the T cell mitogen, was dissolved in 15 μg mL⁻¹ with RPMI-1640. Calf serum (Corning, USA) was used to culture lymphocytes and an MTS Kit (Promega, USA) was used to detect the T cell proliferation rate. The enzyme-linked immunosorbent assay (ELISA) kit was purchased from Nanjing Jiancheng Bioengineering Institute. TRIzol reagent (Invitrogen, China) and a first-strand complementary (cDNA) kit (Invitrogen, China) were used for quantitative realtime PCR analysis.

Immunosuppression induction and treatment

Two hundred geese (1 day-old, with half of them being male and half female) were randomly divided into four groups (n = 50 per group). The control and CTX groups were fed with normal diets, whereas the PAMK and CTX + PAMK groups were fed with 400 mg kg⁻¹ PAMK. In addition, the control and PAMK groups were injected with 0.5 mL saline, while the CTX and CTX + PAMK groups were injected with 40 mg kg⁻¹ BW CTX per day at 12-14 days of age to construct immunosuppression models. Thymus and blood were collected at 28 days of age. All organs were immediately placed in liquid nitrogen and stored at -80 °C until used. IL-2 levels in serum were detected by ELISA.

Thymus index assay

The weights of the thymuses were measured immediately after the geese were slaughtered. The thymus index was calculated as the ratio of the thymus weight (g) to body weight (kg).

Ultrastructural observation

Transmission electron microscopy (TEM). The thymus was divided into small blocks of about 1 mm3 and then fixed with 2.5% glutaraldehyde at 4 °C. Ultra-thin sections of 50-70 nm thickness were prepared and stained with uranyl acetate and lead citrate following the conventional protocol. The material was observed using TEM (JEM-1400, Japan).

Scanning electron microscopy (SEM). The thymus was fixed with 2.5% glutaraldehyde, as above, for 2 hours, and observed using SEM (Hitachi S3000N, Japan).

T cell proliferation assay

Thymuses of the four groups were collected and ground separately into cell suspensions, which were then filtered using a 40 µm cell sieve. Lymphocytes were isolated using lymphocyte separation fluid and collected. The lymphocytes were washed twice with phosphate-buffered saline (PBS) and re-suspended in RPMI-1640 complete medium containing 10% calf serum. Trypan blue staining requires more than 98% cell viability for each group. The cell concentration was adjusted to 1×10^6 cells per mL. PHA was added to the culture medium to activate T cell proliferation. After culturing for 44 hours, an MTS Kit was used to detect the T cell proliferation rate. The absorbance was measured at 490 nm by an enzymelinked immunosorbent assay reader (Biotek Epoch, USA). The ratio of the absorbance of cells treated with PHA to the absorbance without PHA was used as an indicator of the T cell proliferation rate.

Serum IL-2 assay

Venous blood was collected to separate the serum. The levels of IL-2 secreted by activated T cells in the serum were measured using an ELISA kit according to the manufacturer's instruction.

Quantitative real-time PCR assay

Specific primers for miRNAs and target genes were designed (Table 1). U6 was used to normalize the miRNAs and β-actin was used to normalize the genes. The procedure used for qPCR was similar to that used by Yao.23 Total RNA was isolated from individual thymuses using the TRIzol reagent according to the manufacturer's instructions. First-strand complementary (cDNA) was synthesized using oligo dT primers and SuperScript II reverse transcriptase, according to the manufacturer's instructions. QPCR was performed on an ABI PRISM 7500 detection system (Applied Biosystems, USA). The PCR procedure steps included 95 °C for 30 s, followed by 40 cycles of 95 °C for 15 s, 60 °C for 30 s, and 60 °C for 30 s.

Table 1 Primer sequences of mRNA for qPCR

Gene	Primer sequence (5'-3')
MHC-II	F: CCGAGATCGAGGTGAAGTG
	R: GGGTGCTTTCCAGCATCA
CD28	F: ATCTGGACACCCCTCAACAT
	R: TGAACTGGATGCTGTAGGAA
CD96	F: CCAGAGAGGATCCGGAAGAG
	R: ATGCGTTCCTCACAATGCA
CTLA4	F: CCCTAGCCGAAACAATGTG
	R: GCTCCATCTTGCAGACATAA
CD247	F: CGTGGGCACAGAGATGAATA
	R: CCGTCTCTGATGGTCTCCTT
NFAT	F: GCCAAGCTCTACCACCAATG
	R: CAGCACTTGGGCAGTGGTAT
β-Actin	F: GCACCCAGCACGATGAAAAT
	R: GACAATGGAGGGTCCGGATT

Sequencing and differential expression analysis of miRNA

Thymuses were collected for miRNA sequencing. Each group possessed three biological replicates. All samples from the geese were sent to the Beijing Genomics Institute (BGI) for miRNA sequencing and were sequenced using Hiseq technology, referring to Anser cygnoides. The results were filtered to obtain clean tags, which were mapped to sRNA databases such as miRBase, Rfam, siRNA, piRNA, small nucleolar RNAs (snoRNAs) to annotate miRNAs, piRNAs, ribosomal RNAs (rRNAs) and other non-coding RNA (ncRNA) sequences, and then aligned to exons and introns of mRNAs to screen and remove degraded fragments. To obtain unique small RNAs that mapped to only one annotation, the following priority rules were used: MiRNA > piRNA > snoRNA > Rfam > other sRNA. MiRDeep2 was used to identify the known and predict novel miRNAs. Small RNAs with reads less than 100 were discarded, and the miRNA level was calculated using transcripts per kilobase million (TPM) values (TPM = (miRNA total reads/total clean reads) \times 10⁶), which eliminates the influence of sequencing discrepancy on the calculation of small RNA expression.

To validate the effects of PAMK under the treatment of CTX, the differentially expressed miRNAs (|log2(fold change)| > 1 and P-value \leq 0.05) between the control group and PAMK group (control vs. PAMK), and between the CTX and CTX + PAMK groups (CTX vs. CTX + PAMK) were identified. To identify the possible targets of the miRNAs, RNAhybrid, miRanda, and TargetScan software were used. To understand the functions of the predicted target genes, gene ontology (GO) and KEGG enrichment analyses on differentially expressed miRNAs were performed using a hypergeometric test with a false discovery rate \leq 0.05. The differentially expressed miRNAs of control vs. PAMK and CTX vs. CTX + PAMK were mapped to KEGG and the most significant changed pathway related to T cell activation was identified. In addition, detection and interaction analyses of differentially expressed miRNAs and target mRNAs in this pathway were performed based on the target prediction, functional annotation, and the negative regulatory mechanism of mRNAs and miRNAs.

Table 2 Primer sequences of miRNA for qPCR

miRNA	Primer sequence (5′–3′)
novel_mir457	ACACTCCAGCTGGGAGGATGCAGGCAGGAGG
novel_mir19	ACACTCCAGCTGGGCATGTCGCACACCTCTCCTG
novel_mir46	ACACTCCAGCTGGGAGGCAGTGTATTGTTAGTTAG
miR-1805-3p_1	ACACTCCAGCTGGGTGTATTGGAACACTACAG
novel_mir66	ACACTCCAGCTGGGTGTATTGGAACACTACAG
miR_218-3p_2	ACACTCCAGCTGGGATGGTTCTGTCAAGCAC
miR-183	ACACTCCAGCTGGGTATGGCACTGGTAGAATTC
novel_mir443	ACACTCCAGCTGGGTTGGGAACTTGCTGTC
novel_mir2	ACACTCCAGCTGGGTTAGTGCGCAGTAAGCTAG
miR-122	ACACTCCAGCTGGGTGGAGTGTGACAATGGTGT
^a miRNA-R	CTCAACTGGTGTCGTGGA
U6-F	CTCGCTTCGGCAGCACA
U6-R	AACGCTTCACGAATTTGCGT

miRNA-R is the universal reverse primer.

Verification of miRNAs and target genes

Specific primers for the miRNAs and target genes were designed (Table 2). The qPCR process was the same as that detailed above.

Statistics

The results were expressed as means \pm SD. All the qPCR reactions were performed in triplicate and the relative levels were measured in terms of the threshold cycle value (C_t) and were normalized using the equation $2^{-\Delta\Delta C_t}$. Statistical analysis of all data was performed using SPSS for Windows (version 16, SPSS Inc.). The statistical significance (P < 0.05) was evaluated by oneway ANOVA with simultaneous multiple comparisons among different groups.

Results

PAMK improves the thymus index in geese

To confirm the effects of PAMK on the thymus of geese, the thymus indexes of four groups were analyzed (Fig. 1). Compared with the control group, the thymus index of the CTX group was significantly decreased (P < 0.05), but the PAMK group was significantly increased (P < 0.05), which indicated that

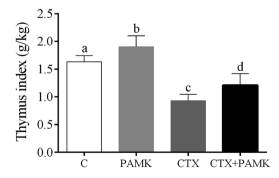


Fig. 1 Effects of PAMK on the thymus index of CTX-induced geese. Means with the different letters represent statistically significant values (P < 0.05); the bars with a common letter are not significantly different (P > 0.05)

RSC Advances

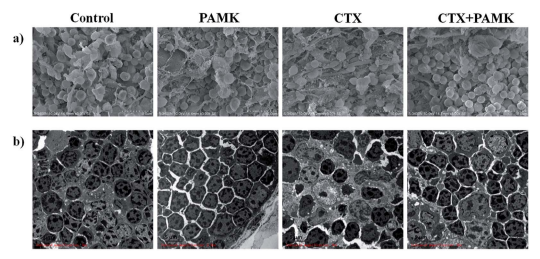


Fig. 2 (a) Scanning electron microscopy (SEM \times 3000) of goose thymus at 28 days of age. (b) Transmission electron microscopy (TEM \times 3000) of goose thymus at 28 days of age.

immunosuppression induced by CTX inhibited thymus development but PAMK promoted it. The thymus index of the group treated with PAMK and CTX (CTX + PAMK group) was significantly lower than that of the control group (P < 0.05), but significantly higher than the CTX group (P < 0.05). Thus, PAMK could not only improve the thymus index of the control group, but also improve the decreased thymus index due to CTX treatment; however, PAMK could not restore the index to normal levels.

PAMK protects the development and cell morphology of the thymus

In the SEM of the thymus, the densities of thymocytes in the control and PAMK groups were normal, the morphology of cells was regular, and no pathological changes could be observed. In contrast, the CTX group showed low cell density, increased connective tissue, irregular cell morphology, and shrinkage of thymocytes. Compared with the CTX group, the CTX + PAMK group exhibited increased cell density and regular thymocyte morphology (Fig. 2a).

TEM of the thymocytes showed that the thymocytes of the CTX group had undergone vacuolization and chromatin margination was observed in the nucleus. In addition, apoptotic bodies were observed. The morphology of the cells of the CTX + PAMK group was improved compared with that of the CTX group. The chromatin was still loose in the nucleus, but chromatin margination was reduced, indicating that thymocytes of geese could be protected from the damaging effects of CTX by feeding with PAMK (Fig. 2b).

The results showed that PAMK could not only increase the thymus index, but could also restore the structure of the thymus to a more normal phenotype, indicating that PAMK could promote the development of the thymus but had no obvious effects on the control group. After treatment with CTX, the immune organ index decreased significantly, the thymus tissue was loose, and cell shrinkage and apoptosis were observed. The results of the CTX + PAMK group showed that PAMK could

alleviate the CTX-induced damage to thymus development and cell morphology.

PAMK alleviated CTX-mediated inhibition of T cell proliferation

Compared with the control group, the T cell proliferation rate of the CTX group was significantly decreased (P < 0.05) and the PAMK group showed no significant difference (P > 0.05), indicating that CTX had immunosuppressive effects on T cell proliferation. However, the T cell proliferation rate of the CTX + PAMK group was significantly higher than that of the CTX group (P < 0.05), but remained significantly lower than that of the control and PAMK groups (P < 0.05). The results indicated PAMK had no significant effects on T cell proliferation in normal geese, but increased the CTX-inhibited T cell proliferation, albeit not up to the normal level (Fig. 3).

PAMK increased T cell activation-related gene expression

The mRNA expression of CD28, CD96, and MHC-II, which are associated with the activation and maturation of T cells, were

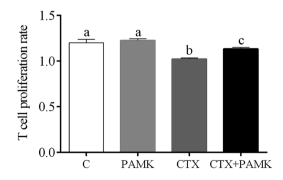
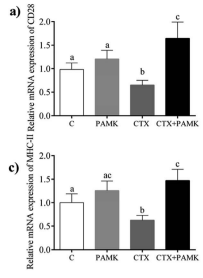
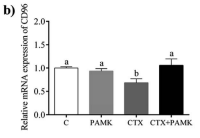


Fig. 3 Effects of PAMK on the T cell proliferation rate in CTX -treated geese. Means with the different letters represent statistically significant values (P < 0.05); the bars with a common letter are not significantly different (P > 0.05).

Paper RSC Advances





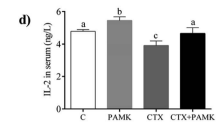


Fig. 4 Effects of PAMK on T cell activation-related genes in CTX-treated geese. (a) Relative mRNA expression of CD28. (b) Relative mRNA expression of CD96. (c) Relative mRNA expression of MHC-II. (d) Effects of PAMK on the IL-2 content of CTX-treated geese. Means with the different letters represent statistically significant values (P < 0.05); the bars with a common letter are not significantly different (P > 0.05).

tested in the homogenized entire thymus. As shown in Fig. 4, the relative mRNA expressions of CD28, CD96, and MHC-II in the CTX group decreased significantly (P < 0.05) compared with those in the control group, indicating that CTX significantly inhibited T cell activation. The relative mRNA expressions of CD28, CD96, and MHC-II in the PAMK group showed no significant differences compared with those of the control group (P > 0.05), while in the CTX + PAMK group, their expressions were significantly higher than those of the CTX group (P < 0.05). It should be noted that the relative mRNA expression of CD96 in the CTX + PAMK group showed no significant difference compared with that of the control group (P > 0.05), whereas the relative mRNA levels of CD28 and MHC-II in the CTX + PAMK group were significantly higher than those in the control group (P < 0.05). Thus, PAMK could not significantly affect the expressions of T cell activation related genes in normal geese but increased the expressions of mRNA downregulated by CTX to normal or higher levels.

PAMK increased the IL-2 content in serum

Compared with the control group, the IL-2 level in the serum of the CTX group was significantly decreased (P < 0.05), but the PAMK group increased significantly (P < 0.05), indicating that CTX inhibited but PAMK promoted IL-2 secretion in geese (Fig. 4d). However, the IL-2 level in the serum of the CTX + PAMK group was significantly higher than that in the CTX group (P < 0.05), and had no significant difference compared with that of the control group (P > 0.05). The results indicated that PAMK not only increased the IL-2 levels in normal geese but also increased the IL-2 level in geese treated with CTX.

Bioinformatics analyses of miRNA sequencing

After removing low-quality reads and adaptor sequences, clean reads of each sample were obtained, respectively (Table 3). The percentage of clean reads among the raw tags was greater than

96% for each sample. Clean reads were used to summarize the length distribution. The results showed that most clean reads were in 21–24 nt length. Generally, miRNAs are 21 nt or 22 nt, short interfering RNAs (siRNAs) are 24 nt, and piwi-interacting RNAs (piRNAs) are between 28 to 30 nt.

The statistics for the alignment of the tags to the reference genome for the control group, PAMK group, CTX group, and CTX + PAMK group were 93.05%, 93.50%, 83.35%, and 80.43%, respectively. A higher ratio of alignment indicates a closer genetic relationship between the sample and reference species. A lower rate may be caused by low similarity with the reference species or by contamination.

Small RNAs (sRNAs) prediction

After sRNA annotation, unknown tags were used to predict novel sRNAs and novel miRNAs, based on their architectural features. Ultimately, 490 novel miRNAs and 662 novel siRNAs were predicted from among 1048576 unknown sRNAs, accounting for 0.048% and 0.063%, respectively.

Table 3 Summary of sequencing data for each sample

Sample name	Raw tag count	Clean reads count	Percentage (%)
Control-1	12 268 083	11 987 864	97.72
Control-2	11 576 318	11 351 789	98.06
Control-3	11 849 430	11 607 797	97.96
PAMK-1	12 359 670	12 069 858	97.66
PAMK-2	11 550 925	11 190 488	96.88
PAMK-3	12 388 333	12 103 559	97.70
CTX-1	11 545 051	11 268 052	97.60
CTX-2	11 832 741	11 487 523	97.08
CTX-3	12 509 679	12 080 906	96.57
CTX + PAMK-1	11 931 196	11 585 937	97.11
CTX + PAMK-2	12 116 956	11 749 233	96.97
CTX + PAMK-3	12 029 029	11 656 445	96.90

RSC Advances Paper

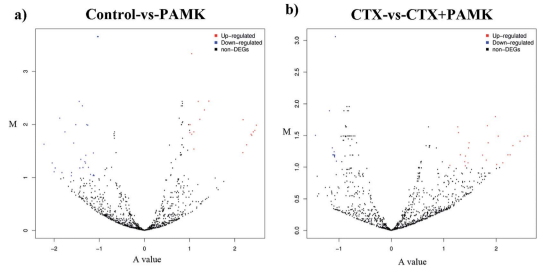


Fig. 5 (a) Volcano plot of the differentially expressed miRNAs of the control vs. PAMK. (b) Volcano plot of the differentially expressed miRNAs of CTX vs. CTX + PAMK. The A value on the x-axis represents the average level after log2 conversion; the M value on the y-axis represents the difference multiple after log2 conversion. Red means upregulated; blue means downregulated; and black means no significant difference.

Differentially expressed miRNAs

After identifying the differentially expressed miRNAs, volcano plots were used to show the distribution of the differentially expressed miRNAs in the control vs. PAMK and CTX vs. CTX + PAMK comparisons (Fig. 5a and b). After normalization and Ttesting of the sequencing data, 37 differentially expressed miRNAs were found between the control group and the PAMK group (20 upregulated and 17 downregulated); of these, 13 were known miRNAs and 24 were novel miRNAs (Table 4). In addition, 42 differentially expressed miRNAs were found between the CTX group and the CTX + PAMK group (32 upregulated and 10 downregulated); of these, 14 were known miRNAs and 28 were novel miRNAs (Table 5). MiR-218-3p_2, novel_mir46, and novel_mir19 were differentially expressed in both comparisons. Among them, novel_mir46 was upregulated in both results, while miR-218-3p_2 and novel_mir19 had opposite effects in the two results.

Validation of the expression of partial difference miRNAs

To validate the sequencing result, 9 miRNAs showing significant differences in the comparisons were selected for quantitative real-time PCR (qPCR) detection. The levels of novel_mir443, novel_mir19, miR-218-3p_2 were upregulated, while novel_mir46, miR-122, and novel_mir66 were downregulated in the control vs. PAMK. The levels of miR-1805-3p_1 and miR-183 were upregulated, while miR-218-3p_2, novel_mir46, novel_mir19, and novel_mir457 were downregulated in CTX vs. CTX + PAMK. All the qPCR results were in agreement with the miRNA sequencing results (Fig. 6a and b).

Novel mir2 is involved in the immunomodulation of PAMK

To obtain more accurate data, the RNAhybrid and miRanda software were used to predict the target genes of the differentially expressed miRNAs. Combined with the filter conditions,

such as free energy and score values for filtering, the filtered results showed that RNAhybrid predicted 41 169 and miRanda predicted 16 726 target genes, among which 16 334 target genes were predicted by both algorithms. Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis of these target genes revealed that they were related to signal transduction, cancers (overview), immune system, and global and overview maps signal pathways (Fig. 7a and b). The fold change in miRNA expression was used to represent the difference in the expression of the target genes, and then we summed the total miRNA fold change related to the KEGG pathways. The major enriched KEGG pathways in the immune system in the CTX vs. CTX + PAMK comparison were the chemokine signaling pathway, complement and coagulation cascades, platelet activation, tolllike receptor signaling pathway, NOD-like receptor signaling pathway, hematopoietic cell lineage, T cell receptor (TCR) signaling pathway, and natural killer cell mediated cytotoxicity (Table 6). T cell activation can be regulated by the TCR signaling pathway; therefore, the TCR signaling pathway was selected for further analysis of the differentially expressed miRNAs and their target mRNAs.

We chose mRNAs that were most highly regulated by miR-NAs in the CTX ν s. CTX + PAMK and that encoded members of the TCR signaling pathway and then identified the corresponding miRNA. Target genes with the highest fold change in the TCR signaling pathway are shown in Table 7. The results for novel_mir2 in the control vs. PAMK and CTX vs. CTX + PAMK comparisons showed that the differential expression of novel_mir2 was not significant in control vs. PAMK, but its expression increased significantly in CTX vs. CTX + PAMK (Table 8). The expression of novel_mir2 has the same tendency as the results of thymus development and T cell activation above. Thus, the PAMK-mediated alleviation of immune damage in geese is closely related to the expression of novel mir2.

Table 4 Differential expression of miRNAs in the thymus of geese between the control and PAMK groups^a

miR-name	Control read	PAMK read	log2 ratio	Up/down-regulation	<i>P</i> -value	Significanc level
novel_mir66	0	15	5.003986249	Up	4.91×10^{-5}	**
novel_mir209	0	14	4.904450575	Up	9.19×10^{-5}	**
novel_mir358	0	12	4.682058154	Up	0.000327404	**
novel_mir327	0	12	4.682058154	Up	0.000327404	**
novel_mir467	0	11	4.556527272	Up	0.000623823	**
novel_mir456	0	11	4.556527272	Up	0.000623823	**
miR-122_1	80	1255	4.068639207	Up	3.78×10^{-276}	**
miR-122-3p_2	85	1302	4.03421845	Up	$5.65 imes 10^{-285}$	**
miR-122-3p_4	90	1278	3.924914678	Up	4.57×10^{-275}	**
miR-122	87	1198	3.88056435	Up	4.16×10^{-256}	**
miR-122-5p	96	1252	3.802152	Up	5.89×10^{-264}	**
miR-122_2	95	1236	3.798703073	Up	1.89×10^{-260}	**
miR-122-3p_3	100	1263	3.755878387	Up	4.93×10^{-264}	**
novel mir404	78	303	2.054867418	Up	8.34×10^{-36}	**
novel mir28	12	46	2.035695109	Up	1.31×10^{-6}	**
novel_mir46	48	128	1.512133153	Up	5.26×10^{-11}	**
niR-218a	1038	2530	1.382426595	Up	6.47×10^{-164}	**
niR-206-3p	2266	4800	1.179982198	Up	4.76×10^{-242}	**
miR-206	2303	4826	1.164409158	Up	2.70×10^{-238}	**
novel mir16	41	77	1.006330189	Up	0.000217115	**
niR-218-3p_2	1094	505	-1.018161792	Down	3.53×10^{-42}	**
novel_mir8	440	202	-1.026052577	Down	3.76×10^{-18}	**
novel_mir12	160	72	-1.05490744	Down	8.41×10^{-8}	**
novel mir460	84	35	-1.165938752	Down	2.56×10^{-5}	**
niR-19b-2-5p	8028	3275	-1.196450057	Down	0	**
niR-16b-3p_2	47	18	-1.287568197	Down	0.000662984	**
novel mir393	69	22	-1.551997185	Down	2.04×10^{-6}	**
novel mir20	110	27	-1.929376558	Down	2.89×10^{-12}	**
novel mir399	29	7	-1.95353042	Down	0.000300161	**
novel mir87	25	6	-1.961798036	Down	0.000762743	**
novel mir177	131	28	-2.128972426	Down	4.55×10^{-16}	**
novel_mir262	29	6	-2.175922841	Down	0.000106849	**
novel_mir448	39	8	-2.188306566	Down	6.51×10^{-6}	**
novel_mir366	19	2	-3.15083186	Down	0.000125388	**
novel_mir426	21	1	-4.295221769	Down	6.55×10^{-6}	**
novel_mir19	66	3	-4.362335965	Down	1.03×10^{-15}	**
novel_mir443	13	0	-4.603344065	Down	0.000313854	**

Novel_mir2 targets CTLA4 to regulate the TCR pathway

Novel_mir2 is a newly annotated miRNA that is located on chromosome NW_013185761. The mature and precursor sequences of novel_mir2 are shown in Table 7. The stem loop structure of novel_mir2 was predicted and is shown in Fig. 8. It showed that novel_mir2 is a miRNA, not a fragment of an mRNA (Table 9).

To confirm the differences in novel_mir2 expression between the groups, qPCR was used to detect novel_mir2 expression. The results showed that the relative mRNA expression of novel_mir2 between the control and PAMK groups was not significantly different (P > 0.05). However, its relative expression in the CTX + PAMK group was significantly higher than that in the CTX group (P < 0.05) (Fig. 9a), which is consistent with the sequencing results.

Based on the target gene prediction results of miRNA sequencing, novel_mir2 could target multiple mRNAs in the TCR signaling pathway, such as CD28, CTLA4, and PI3K. CTLA4

can only be targeted by novel_mir2, while CD28 and PI3K can also be targeted by other miRNAs. In addition, CD28 and CTLA4 are very important genes in the TCR pathway. Their encoded proteins competitively bind to the same ligands, CD86 and CD80, during T cell activation. When CD28 binds to the ligands, CD28 is activated and T cell activation is promoted. When CTLA4 binds to the ligands, the TCR-CD3 pathway is inhibited so that T cell activation is inhibited. The qPCR results for CTLA4 showed that its expression had the opposite tendency to that of novel_mir2 (Fig. 9b), while the expression of CD28 showed the same expression pattern as novel_mir2 (Fig. 4a). Thus, novel_mir2 targets the expression of CTLA4. To further confirm whether novel_mir2 regulates T cell activation by targeting CTLA4, the mRNA levels of CD247 (CD3 zeta) and the nuclear factor of activated T-cells (NFAT) in the TCR signaling pathway were detected. The results showed that the mRNA expression patterns of CD247 and NFAT were the opposite to that of CTLA4 (Fig. 9c and d). CTLA4 expression showed no significant

Table 5 Differential expression of miRNAs in the thymus of geese between the CTX and CTX + PAMK groups^a

miR-name	CTX read num	CTX + PAMK read num	log2 ratio	Up–down regulation	<i>P</i> -value	Significance level
novel mir457	0	19	5.295854409	Up	5.18×10^{-6}	**
novel_mir254	1	16	4.047926896	Ор Up	7.24×10^{-5}	**
novel_mir331	1	16	4.047926896	Ор Up	7.24×10 7.24×10^{-5}	**
novel_mir168	4	37	3.257380262	Ор Up	2.56×10^{-8}	**
miR-103a-5p	3	20	2.78489249	Up	0.000145192	**
novel mir490	3	18	2.632889397	Ор Up	0.000143192	**
novel_mir224	3 31	181	2.593576473	•	3.71×10^{-28}	**
novel_mir13	18	96	2.462964395	Up	7.04×10^{-15}	**
novel_mir19	5	26	2.426438519	Up Up	5.92×10^{-5}	**
novel_mir33	4	20	2.369854991	•	0.000512685	**
novel_mir64	4 14	58	2.098552969	Up	4.00×10^{-8}	**
_	6			Up		**
miR-2188-5p_1		24	2.047926896	Up	0.000506439	**
novel_mir80	7	27	1.995459476	Up	0.000287185 1.68×10^{-7}	**
novel_mir361	15	57	1.973926314	Up		**
novel_mir46	37	125	1.804257815	Up	3.03×10^{-13}	**
novel_mir392	48	162	1.802814398	Up	1.06×10^{-16}	**
novel_mir113	9	29	1.73598289	Up	0.000625848	**
novel_mir263	38	122	1.73073672	Up	2.56×10^{-12}	**
novel_mir451	12	37	1.672417761	Up	0.000167905	**
novel_mir133	20	58	1.583979796	Up	5.87×10^{-6}	
miR-218b_3	251	650	1.42067925	Up	1.66×10^{-44}	**
novel_mir71	18	44	1.337433513	Up	0.000505705	**
miR-218-5p	260	628	1.320179832	Up	1.10×10^{-38}	**
novel_mir278	53	126	1.297286365	Up	9.02×10^{-9}	**
miR-218-3p_3	284	646	1.233570131	Up	6.26×10^{-36}	**
miR-218-2-3p	293	658	1.215113815	Up	9.53×10^{-36}	**
miR-218-3p_2	294	647	1.185876453	Up	6.53×10^{-34}	**
novel_mir4	177	383	1.161521928	Up	4.05×10^{-20}	**
miR-218-3p_1	305	638	1.112674077	Up	2.22×10^{-30}	**
novel_mir273	90	185	1.08745526	Up	1.41×10^{-9}	**
novel_mir2	1643	3238	1.026697401	Up	4.57×10^{-129}	**
novel_mir9	102	200	1.019357744	Up	2.35×10^{-9}	**
novel_mir363	65	30	-1.067550322	Down	0.000506556	**
miR-183_1	8203	3532	-1.16773939	Down	0	**
miR-183	8066	3472	-1.168159562	Down	0	**
miR-183_2	8184	3423	-1.209617957	Down	0	**
miR-183-5p	8391	3507	-1.210678428	Down	0	**
miR-183-3p_1	8220	3379	-1.234615148	Down	0	**
miR-183_3	8364	3436	-1.23553613	Down	0	**
novel_mir126	209	79	-1.355651488	Down	4.98×10^{-14}	**
novel_mir37	47	16	-1.506661956	Down	0.000110035	**
miR-1805-3p_1	23	5	-2.153706965	Down	0.000546189	**

^a *p < 0.05, **p < 0.001.

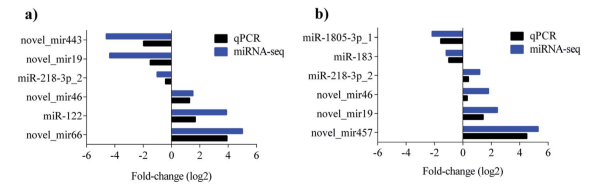
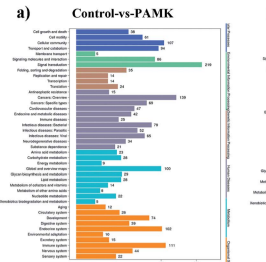


Fig. 6 (a) Comparison results of miRNA sequencing and qPCR of control vs. PAMK. (b) Comparison results of miRNA sequencing and qPCR of CTX vs. CTX + PAMK. Blue represents the results of miRNA sequencing; black represents the results of miRNA qPCR.

Paper



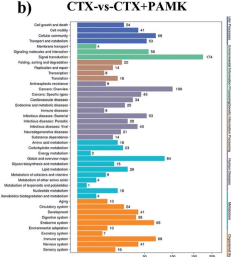


Fig. 7 (a) Enrichment of KEGG pathways based on target genes of control vs. PAMK. (b) Enrichment of KEGG pathways based on target genes of CTX vs. CTX + PAMK. The x-axis is the target gene number of the miRNA; the right y-axis shows the pathway terms, divided into six categories with different colors.

Table 6 The majority KEGG enriched in the immune system in CTX vs. CTX + PAMK

Pathway	DEGs genes with pathway annotation (23 476)	All genes with pathway annotation (28 984)	P value	Q value	Pathway ID
Platelet activation	750 (3.19%)	882 (3.04%)	0.000846	2.94×10^{-2}	ko04611
Fc gamma R-mediated phagocytosis	287 (1.22%)	337 (1.16%)	0.026657	2.78×10^{-1}	ko04666
Hematopoietic cell lineage	358 (1.52%)	424 (1.46%)	0.037209	3.10×10^{-1}	ko04640
B cell receptor signaling pathway	212 (0.9%)	252 (0.87%)	0.115164	5.63×10^{-1}	ko04662
Fc epsilon RI signaling pathway	200 (0.85%)	238 (0.82%)	0.131098	5.78×10^{-1}	ko04664
NOD-like receptor signaling pathway	179 (0.76%)	213 (0.73%)	0.146814	6.13×10^{-1}	ko04621
Antigen processing and presentation	81 (0.35%)	95 (0.33%)	0.176778	6.83×10^{-1}	ko04612
Toll-like receptor signaling pathway	234 (1%)	282 (0.97%)	0.220589	7.94×10^{-1}	ko04620
Leukocyte transendothelial migration	744 (3.17%)	919 (3.17%)	0.532457	1.00×10^{00}	ko04670
Natural killer cell mediated cytotoxicity	270 (1.15%)	336 (1.16%)	0.648932	1.00×10^{00}	ko04650
Intestinal immune network for IgA production	80 (0.34%)	103 (0.36%)	0.838767	1.00×10^{00}	ko04672
T cell receptor signaling pathway	330 (1.41%)	418 (1.44%)	0.871833	1.00×10^{00}	ko04660
RIG-I-like receptor signaling pathway	120 (0.51%)	156 (0.54%)	0.917	1.00×10^{00}	ko04622
Chemokine signaling pathway	408 (1.74%)	523 (1.8%)	0.963163	1.00×10^{00}	ko04062
Cytosolic DNA-sensing pathway	66 (0.28%)	95 (0.33%)	0.997703	1.00×10^{00}	ko04623
Complement and coagulation cascades	168 (0.72%)	247 (0.85%)	1	1.00×10^{00}	ko04610

difference between the control group and the PAMK group (P > 0.05); however, in the CTX + PAMK group, its expression was significantly higher than that in the CTX group (P < 0.05). The trend of CTLA4 expression was the opposite to that of novel_mir2, which suggested that novel_mir2 regulates CTLA4. In addition, the expression patterns of CTLA4, CD247, and NFAT were in line with the mechanism of the TCR signaling pathway, indicating that CTLA4 is involved in the regulation of the TCR signaling pathway.

Discussion

Cyclophosphamide (CTX) is often used to construct immunosuppressed models to study immune function in mammals and poultry because of the inhibition of humoral and cellular immunity.^{2,24} Therefore, in this research an immunosuppressed model of geese was prepared with the injection of CTX. The results showed that CTX reduced the thymus index, inhibited thymus development, led to thymocyte shrinkage and

Table 7 Partial miRNAs and target genes in the TCR pathway

Target gene	miRNA
CD 28 CTLA4 PI3K	novel_mir2; novel_mir4; miR-122 novel_mir2 novel_mir224; novel_mir2; novel_mir3; novel_mir4

RSC Advances

Table 8 Comparison of novel mir2 in the control vs. PAMK and CTX vs. CTX + PAMK

Comparison group (A vs. B)	Read num A	Read num <i>B</i>	log2 ratio (A/B)	Up–down regulation	<i>P</i> -value	Significance level
Control vs. PAMK CTX vs. CTX + PAMK	1562 1643	1427 3238	-0.382 1.027	Down Up	$0.485 \\ 4.57 \times 10^{-129}$	Not significance

Provisional ID	: 1	1609T-1197681810_AA		
Score total	0	16657.7		
Score for star read(s)	1	3.9		5' C
Score for read counts	0	16652	31	U U A G U G C G A G U AAG C U A G G U C H
Score for mfe	:	2.3	NT.	AAUCACCU UCCUCCAUC LIII U
Score for randfold	:		0	e e
Score for cons. seed	:	-0.6		C-A
Total read count	0	32674		
Mature read count	:	31718		
Loop read count	0	0		
Star road count		056		

Fig. 8 The stem loop structure of novel_mir2

Characteristics of novel_mir2

miRNA ID	Novel_mir2
Chromosome Sequence(mature) Sequence(star) Sequence(precursor)	NW_013185761 uuagugcgcaguaagcuagggugu cgcuagcugcucugcacuaacu uuagugcgcaguaagcuagggugugaauugacagcacgcuagcugcucugcacuaacu

apoptosis, inhibited T cell proliferation and T cell activation, revealing that immunosuppression was successfully constructed in geese.

As reported, the immune enhancing effects of PAMK were mainly in promoting immune organ development, lymphocyte proliferation, antioxidant ability and secretion of cytokines in

mice and chickens.5,10,25 Here, we found that PAMK also promoted thymus development, T lymphocyte proliferation in geese, suggesting that there was no species difference in waterfowl, poultry and mammals. However, PAMK had no significant pathological effects on the thymus structure of normal geese, as well as the spleen structure of chickens.5

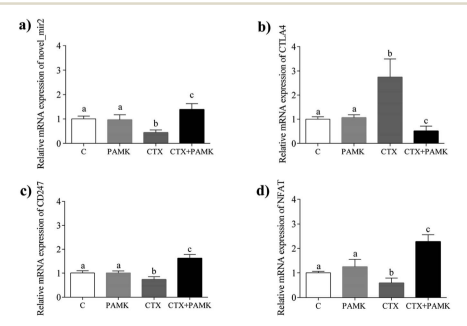


Fig. 9 Relative mRNA expression of novel_mir2 and partial TCR pathway genes. (a) Relative mRNA expression of novel_mir2. (b) Relative mRNA expression of CTLA4. (c) Relative mRNA expression of CD247. (d) Relative mRNA expression of NFAT. Means with the different letters represent statistically significant values (P < 0.05); the bars with a common letter are not significantly different (P > 0.05).

Paper **RSC Advances**

Besides, PAMK was first claimed to alleviate the immunosuppressive effects induced by CTX in this study. Lots of polysaccharides of traditional Chinese herbs have been reported as having the effects of alleviating immune suppression, similar to PAMK. Chuanminshen violaceum polysaccharide (CVP) was proven to overcome the CTX-induced immunosuppression of mice by raising IL-2 levels in serum, while Cordyceps militaris polysaccharides (CMP) were mainly via the function of macrophages and anti-oxidative activity in immunosuppressed mice.2,3 However, we found that PAMK alleviated CTX-induced immunosuppression closely related to T cell activation according to the changes in CD28, CD96, MHC II, and IL-2. T cell activation requires a dual-signal in the TCR pathway: the TCR-CD3 complex binds to the antigen peptide-MHC molecule, and the CD molecule on the T cell surface binds to the corresponding ligand on the surface of the antigen presenting cell (APC), which is a CD28 activation signal and a CD152 inhibitory signal.26 Besides, CD96 expression was positively associated with CD4⁺ T cell counts and was upregulated during T cell activation (23).27 In addition, Curtsinger28 suggested a third signal for T cell activation, which stimulated CD8+ T cell proliferation and cytokine secretion (IL-2, IL-12). IL-2 is also an important cytokine secreted by activated T cells.29 For these reasons, CD28, CD152, CD96, MHC II, and IL-2 are the typical molecular signs of T cell activation, and PAMK alleviated CTXinduced immunosuppression might through the TCR pathway.

It has been reported that miRNAs could regulate the target genes in the TCR pathway, thereby regulating T cell activation.17,19 Although each miRNA targets different genes and regulates different pathways, most of them eventually alter the expression of NF-κB, thereby regulating the activation of T cells. The present study has shown that PAMK alleviated the inhibition of T cell activation induced by CTX mainly through the upregulation of novel_mir2 to degrade or inhibit the translation of CTLA4. Coincidentally, it was reported that miR-155 promotes T cell activation by targeting CTLA4, which is the negative regulator of T cell activation in humans.20 CTLA4 is also called CD152, which inhibits T cell activation by inhibiting the expression of the TCR-CD3 complex or ZAP70 and finally inhibits the TCR pathway. Thus, the inhibition of T cell activation induced by CTX was due to the high expression of CTLA4 and inhibited the downstream activation of the TCR pathway, and the alleviation of PAMK on immunosuppression was due to the low expression of CTLA4 and the activation of the TCR pathway. According to the results, the changes in CD28 and CTLA4 in the CTX and CTX + PAMK groups are all opposite. The novel_mir2 did not directly target CD28 to regulate T cell activation but targeted CTLA4 so that the ability of CTLA4 to compete with CD28 to bind CD80 and CD86 was inhibited. Thus, the increased expression of CD28 might further promote the TCR pathway because it binds more ligands. However, some reports found that the changes in CD28 and CTLA4 are the same in the regulation of T cell activation in vitro. 30 The simultaneous increase in CD28 and CTLA4 is inconsistent with the regulatory mechanism of T cell activation. The reason may be that there are no APCs with ligands cultured in vitro for CD28 to compete with CTLA4. In addition, NFAT is an important nuclear

transcription factor of T cell activation that is also regulated by miRNAs.31,32 In the present study, we found that PAMK treatment significantly increased the expression of NFAT in the presence of CTX, but not NF-kB, indicating that PAMK might upregulate the expression of novel mir2 to inhibit CTLA4, leading to increased levels of nuclear transcription factor NFAT, which ultimately changes the expression of cytokines and promotes T cell activation. This means that the mechanism by which PAMK promotes T cell activation, inhibited by CTX in geese, is not the same as the mechanism by which CMP promotes macrophage activation.2

Conclusions

In conclusion, PAMK upregulates novel_mir2 expression, which targets and inhibits CTLA4 expression, leading to the promotion of T cell activation through the TCR-NFAT pathway, ultimately alleviating the immune suppression induced by CTX in geese. This study is the first to systematically analyze the effects of PAMK on immunosuppressed geese.

Conflicts of interest

The authors declare that there are no conflicts of interest.

Abbreviations

AP-1 Activator protein-1 APC Antigen presenting cell

CMP Cordyceps militaris polysaccharides Cytotoxic T lymphocyte antigen 4 CTLA4

CTX Cyclophosphamide

CVP Chuanminshen violaceum polysaccharide

GO Gene ontology

Kyoto Encyclopedia of Genes and Genomes **KEGG**

miRNAs MicroRNAs ncRNA Non-coding RNA

NFAT Nuclear factor of activated T-cells

PAMK Polysaccharide of atractylodes macrocephala koidz

PBS Phosphate-buffered saline

PHA Phytoagglutinin rRNAs Ribosomal RNAs

SEM Scanning electron microscopy

Small nucleolar RNAs snoRNAs

STAT3 Signal transducer and activator of transcription 3

TCR T cell receptor

TEM Transmission electron microscopy Transcripts per kilobase million **TPM**

Acknowledgements

The authors extend their sincere thanks to Inter-governmental S & T Exchange Project between China and Belarus (No. CB01-13); Science & Technology Planning Project of Guangzhou (No. 201604020061); Modern Agriculture Industry Technology System of Guangdong (No. 2016LM1113).

RSC Advances

References

- 1 C. Ma, Q. X. Yue, S. H. Guan, W. Y. Wu, M. Yang, B. H. Jiang, X. Liu and D. A. Guo, *Food Chem. Toxicol.*, 2009, 47, 1841–1847.
- 2 M. Wang, X. Y. Meng, R. L. Yang, T. Qin, X. Y. Wang, K. Y. Zhang, C. Z. Fei, Y. Li, Y. Hu and F. Q. Xue, *Carbohydr. Polym.*, 2012, 89, 461–466.
- 3 X. Zhao, Y. Zhang, X. Song, Z. Yin, R. Jia, X. Zhao, X. Lai, G. Wang, X. Liang, C. He, L. Yin, C. Lv, L. Zhao, G. Shu, G. Ye and F. Shi, *Int. J. Clin. Exp. Med.*, 2015, 8, 558–568.
- 4 T. Wang, F. Long, X. Zhang, Y. Yang, X. Jiang and L. Wang, *Oncotarget*, 2017, 8, 77500–77514.
- 5 D. Xu, B. Li, N. Cao, W. Li, Y. Tian and Y. Huang, *Oncotarget*, 2017, **8**, 70394–70405.
- 6 L. Guo, Y. L. Sun, A. H. Wang, C. E. Xu and M. Y. Zhang, *Mol. Biol. Rep.*, 2012, 39, 9285–9290.
- 7 W. Fan, S. Zhang, P. Hao, P. Zheng, J. Liu and X. Zhao, Carbohydr. Polym., 2016, 153, 631–640.
- 8 G. Q. Ji, R. Q. Chen and J. X. Zheng, *Pharm. Biol.*, 2015, 53, 512–517.
- 9 W. Sun, K. Meng, C. Qi, X. Yang, Y. Wang, W. Fan, Z. Yan, X. Zhao and J. Liu, *Carbohydr. Polym.*, 2015, **126**, 91–96.
- 10 J. Liu, X. Chen, C. Yue, R. Hou, J. Chen, Y. Lu, X. Li, R. Li, C. Liu, Z. Gao, E. Li, Y. Li, H. Wang, Y. Yan, H. Li and Y. Hu, *Int. J. Biol. Macromol.*, 2015, 72, 1435–1440.
- 11 G.-Q. Ji, R.-Q. Chen and J.-X. Zheng, *Pharm. Biol.*, 2014, 53, 512–517.
- 12 D. Xu, W. Li, Y. Huang, J. He and Y. Tian, *Biol. Trace Elem. Res.*, 2014, **160**, 232–237.
- 13 D. Xu and Y. Tian, Biol. Trace Elem. Res., 2015, 168, 235-241.
- 14 F. Allantaz, D. T. Cheng, T. Bergauer, P. Ravindran, M. F. Rossier, M. Ebeling, L. Badi, B. Reis, H. Bitter, M. D'Asaro, A. Chiappe, S. Sridhar, G. D. Pacheco, M. E. Burczynski, D. Hochstrasser, J. Vonderscher and T. Matthes, *PLoS One*, 2012, 7, e29979.
- B. J. Kroesen, N. Teteloshvili, K. Smigielska-Czepiel,
 E. Brouwer, A. M. Boots, A. van den Berg and J. Kluiver,
 Immunology, 2015, 144, 1–10.
- 16 I. S. Khan, C. Y. Park, A. Mavropoulos, N. Shariat, J. L. Pollack, A. J. Barczak, D. J. Erle, M. T. McManus, M. S. Anderson and L. T. Jeker, *PLoS One*, 2015, 10, e0135440.
- 17 Q.-J. Li, J. Chau, P. J. R. Ebert, G. Sylvester, H. Min, G. Liu, R. Braich, M. Manoharan, J. Soutschek, P. Skare, L. O. Klein, M. M. Davis and C.-Z. Chen, *Cell*, 2007, 129, 147–161.

- 18 J. Henao-Mejia, A. Williams, L. A. Goff, M. Staron, P. Licona-Limon, S. M. Kaech, M. Nakayama, J. L. Rinn and R. A. Flavell, *Immunity*, 2013, 38, 984–997.
- 19 D. Loffler, K. Brocke-Heidrich, G. Pfeifer, C. Stocsits, J. Hackermuller, A. K. Kretzschmar, R. Burger, M. Gramatzki, C. Blumert, K. Bauer, H. Cvijic, A. K. Ullmann, P. F. Stadler and F. Horn, *Blood*, 2007, 110, 1330–1333.
- E. Sonkoly, P. Janson, M. L. Majuri, T. Savinko, N. Fyhrquist,
 L. Eidsmo, N. Xu, F. Meisgen, T. Wei, M. Bradley, J. Stenvang,
 S. Kauppinen, H. Alenius, A. Lauerma, B. Homey,
 O. Winqvist, M. Stahle and A. Pivarcsi, J. Allergy Clin. Immunol., 2010, 126(581–589), e581–520.
- 21 K. Yao, Q. Wang, J. Jia and H. Zhao, *Tumour Biol.*, 2017, 39, 1–13
- 22 T. B. Huffaker, S.-H. Lee, W. W. Tang, J. A. Wallace, M. Alexander, M. C. Runtsch, D. K. Larsen, J. Thompson, A. G. Ramstead, W. P. Voth, R. Hu, J. L. Round, M. A. Williams and R. M. O'Connell, *J. Biol. Chem.*, 2017, 292, 18530–18541.
- 23 H. D. Yao, Q. Wu, Z. W. Zhang, S. Li, X. L. Wang, X. G. Lei and S. W. Xu, *Biochim. Biophys. Acta*, 2013, **1830**, 3112–3120.
- 24 L. Guo, J. Liu, Y. Hu, D. Wang, Z. Li, J. Zhang, T. Qin, X. Liu, C. Liu, X. Zhao, Y. P. Fan, G. Han and T. L. Nguyen, *Carbohydr. Polym.*, 2012, 90, 1055–1060.
- 25 W. Sun, K. Meng, C. Qi, X. Yang, Y. Wang, W. Fan, Z. Yan, X. Zhao and J. Liu, *Carbohydr. Polym.*, 2015, **126**, 91–96.
- 26 P. Bretscher and M. Cohn, Science, 1970, 169, 1042-1049.
- 27 E. M. Eriksson, C. E. Keh, S. G. Deeks, J. N. Martin, F. M. Hecht and D. F. Nixon, *PLoS One*, 2012, 7, e51696.
- 28 J. M. Curtsinger, C. S. Schmidt, A. Mondino, D. C. Lins, R. M. Kedl, M. K. Jenkins and M. F. Mescher, *J. Immunol.*, 1999, 162, 3256–3262.
- 29 I. Panagoulias, T. Georgakopoulos, I. Aggeletopoulou, M. Agelopoulos, D. Thanos and A. Mouzaki, *J. Biol. Chem.*, 2016, 291, 26707–26721.
- 30 Z. Y. Zhuge, Y. H. Zhu, P. Q. Liu, X. D. Yan, Y. Yue, X. G. Weng, R. Zhang and J. F. Wang, *PLoS One*, 2012, 7, e29320.
- 31 K. Kang, X. Peng, X. Zhang, Y. Wang, L. Zhang, L. Gao, T. Weng, H. Zhang, R. Ramchandran, J. U. Raj, D. Gou and L. Liu, *J. Biol. Chem.*, 2013, 288, 25414–25427.
- 32 C. H. Gabriel, F. Gross, M. Karl, H. Stephanowitz, A. F. Hennig, M. Weber, S. Gryzik, I. Bachmann, K. Hecklau, J. Wienands, J. Schuchhardt, H. Herzel, A. Radbruch, E. Krause and R. Baumgrass, J. Biol. Chem., 2016, 291, 24172–24187.