Increased adenosine-to-inosine RNA editing in rheumatoid arthritis

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\textbf{A B S T R A C T}

\textbf{Objective:} Adenosine-to-ino sine (A-to-I) RNA editing of Alu retroelements is a primate-specific mechanism mediated by adenosine deaminases acting on RNA (ADARs) that diversifies transcriptome by changing selected nucleotides in RNA molecules. We tested the hypothesis that A-to-I RNA editing is altered in rheumatoid arthritis (RA).

\textbf{Methods:} Synovium expression analysis of ADAR1 was investigated in 152 RA patients and 50 controls. Peripheral blood mononuclear cells derived from 14 healthy subjects and 19 patients with active RA at baseline and after 12-week treatment were examined for ADAR1p150 and ADAR1p110 isoform expression by RT-qPCR. RNA editing activity was analysed by AluSx\textsuperscript{+} Sanger-sequencing of cathepsin S, an extracellular matrix degradation enzyme involved in antigen presentation.

\textbf{Results:} ADAR1 was significantly over-expressed in RA synovium regardless of disease duration. Similarly, ADAR1p150 isoform expression was significantly increased in the blood of active RA patients. Individual nucleotide analysis revealed that A-to-I RNA editing rate was also significantly increased in RA patients. Both baseline ADAR1p150 expression and individual adenosine RNA editing rate of cathepsin S AluSx\textsuperscript{+} decreased after treatment only in those patients with good clinical response. Upregulation of the expression and/or activity of the RNA editing machinery were associated with a higher expression of edited Alu-enriched genes including cathepsin S and TNF receptor-associated factors 1,2,3 and 5.

\textbf{Conclusion:} A previously unrecognized regulation and role of ADAR1p150-mediated A-to-I RNA editing in post-transcriptional control in RA underpins therapeutic response and fuels inflammatory gene expression, thus representing an interesting therapeutic target.

\section{1. Introduction}

Rheumatoid Arthritis (RA) is a chronic inflammatory rheumatic disease, which affects approximately 0.5–1% of the population worldwide [1]. To date RA remains an incurable disease, heavily affecting the quality of life of the patients, their ability to work, as well as their life expectancy overall [1]. Significant advances in the understanding of disease pathogenesis in the past decades, mainly the delineation of the pivotal role of tumor necrosis factor alpha (TNF\textalpha), have lead to the development of biological therapies, which have transformed the course of these patients [2]. However, in 2019 remission and drug-free remission rates among RA patients remain unsatisfactorily low [1,3], with more than half of the patients treated with biological therapies showing persisting clinically active disease [4]. Moreover, there are currently no established biomarkers that can predict response to therapy [5]. Therefore, there is a demanding need to explore novel aspects underlying the pathogenetic mechanisms involved in RA.

Adenosine-to-inosine (A-to-I) RNA editing is the most abundant substitutional RNA modification, catalysed by the enzymes adenosine deaminases acting on RNA-1 and -2 (ADAR1 and ADAR2) [6]. A-to-I
RNA editing takes place predominantly in repetitive non-coding regions of RNA, namely Alu elements, which have the ability to form double-stranded RNA regions, a pre-requisite for RNA editing [7–11]. Alu RNA editing is mainly, if not exclusively, catalysed by ADAR1 and is a pre-requisite for RNA editing [7–11]. Interestingly, presence of Alu elements in human gene sequences has recently emerged as a critical regulator of RNA metabolism dictating the fate of the RNA molecules including alternative splicing, mRNA stability and translation [10,12]. More importantly, Alu elements are conserved only in primates, therefore Alu RNA editing constitutes an additional level of gene regulation specifically for human disease [10,12,13]. Despite significant advances in the study of RNA editing with the introduction of second-generation RNA sequencing, the involvement of Alu RNA editing in human disease development and progression is currently poorly understood [12].

Herein, we tested the hypothesis that ADAR1 expression and A-to-I RNA editing are altered in the pro-inflammatory milieu of RA, probably influencing the expression of Alu-enriched pro-inflammatory genes. As an example, here, we focused on cathepsin S, a well-established target of ADAR1, whose mRNA stability and expression is controlled by Alu RNA editing [14]. Cathepsin S has an essential role in major histocompatibility complex (MHC)-II–mediated antigen presentation [15], autoantibody production [16] and extracellular matrix degradation [17], and has been previously reported to control experimental RA development in mouse [18], as well as to be increased in both the synovial fluid [19] and the systemic circulation of patients with RA [20].

2. Materials and methods

2.1. Analysis of RNA-sequencing dataset of synovial tissues

A large RNA-sequencing dataset (GSE89408) [21] including in total 202 synovial biopsy samples (normal joint = 28, osteoarthritis = 22, early RA-disease duration < 1 year = 57, established RA = 95) was accessed through the NCBI database. Aligned data on ADAR1 and cathepsin S expression were extracted through the Gene Expression Omnibus database.

2.2. Analysis of RNA-sequencing dataset for ADAR1p110 and ADAR1p150 isoform expression

RNA-sequencing data (read length 101 bp generated from total RNA) from synovial samples of normal (28), osteoarthritis (22), early rheumatoid arthritis (57) and established rheumatoid arthritis (95) patients, were downloaded from the NCBI Sequence Read Archive (SRP092408) in .sra format and converted in .fastq by means of fastq-dump program that is part of the SRA toolkit package. Quality-filtering with the NGS QC Toolkit [22] and default parameters (cutoff read length for HQ = 70%, cutoff quality score = 20). High quality cleaned reads were mapped against pre-indexed human genome GRCh37, transcriptome (pre-processed set of known splice junctions from Ensembl annotation), and dbSNP common release 144 using HISAT2 version 2.1.0 [23] and default parameters. Unique and concordant alignments in .sam format were converted in the binary .bam format, sorted by genomic coordinates, and indexed by SAMtools. Transcriptome quantification was performed for each sample with StringTie v1.3.6 release [24] and differential expression was tested with the DESeq2 1.2.0 [25] (FDR ≤ 0.05) R package. Reference human transcriptome was obtained from UCSC (http://hgdownload.cse.ucsc.edu/goldenpath/hg19/database/). Expression of ADAR1 isoforms, ADAR1p110 (NM_001111) and ADAR1p150 (NM_001025107), was determined for each sample.

2.3. Patient recruitment and follow-up

Peripheral blood was collected in EDTA tubes (BD Vacutainer) from 19 consecutive consenting patients with RA fulfilling the 2010 ACR/ EULAR criteria [26] and 14 apparently healthy controls (HC). All patients had active disease (DAS28-ESR: 3.4–7.1) at the time of sampling. Exclusion criteria included viral or bacterial infection during the past month and severe co-morbidities (cancer, heart or kidney failure). Clinical and laboratory RA features (28 tender/swollen joint count, hematocrit and white-blood cell count, erythrocyte sedimentation rate (ESR; mm/1st h.), C-reactive protein (CRP; mg/dl), visual analogue scale (VAS)-patient global) and rheumatoid factor (RF)/anti-cyclic citrullinated peptide (CCP) status were recorded at baseline (Table 1).

RA patients were re-examined 12 weeks after initiation of new treatment (scale-up of treatment, conventional synthetic disease-modifying antirheumatic drugs (csDMARDs) ± corticosteroids and/or
biological DMARDs) and were categorized in responders and moderate/non-responders according to EULAR’s response criteria [27] (treatment modalities and their effect on disease characteristics in EULAR responders vs moderate/non-responders are shown in Table 1). For schematic representation of study design and patient recruitment see Supplementary Fig. 1. All participants gave written informed consent in compliance with the Declaration of Helsinki, which had been previously approved by the Ethics Committee of Laiko Hospital, Athens, Greece (Protocol Nr.:1368/17-11-2016), as well as by the Hellenic Data Protection Authority (Protocol Nr.:Γ/Ε/7901-2/22-12-2016).

2.4. Alu Sanger sequencing and RNA editing analysis
cDNA from peripheral blood mononuclear cells (PBMCs) was subjected to PCR with KOD Hot-start DNA polymerase (Millipore) for the amplification of the most edited Alu region in cathepsin S 3’ untranslated region (UTR), namely AluSx⁺ (PCR product size: 442bp) [14]. Gel-extracted PCR product was subjected to Sanger sequencing and editing rates of individual adenosines were determined following the analysis of the chromatopherograms as we have previously described [14].

2.5. Statistical analysis
Statistical analysis was conducted with SPSS 24.0. Normality of continuous variables was graphically assessed by histograms and P-P plots, as well as by Kolmogorov-Smirnov and Shapiro-Wilk tests. Pairwise differences were evaluated with two independent samples Student’s t-test or the non-parametric Mann-Whitney U test for continuous variables between groups of RA and controls and chi-squared test for nominal variables. The Kruskal-Wallis test was used when comparing more than 2 groups. In order to account for multiple comparisons, Bonferroni correction was performed where applicable. Linear regression analysis was used to control for the effect of confounding factors. Correlations between continuous variables were explored with Pearson’s test when normal distribution applied and Spearman’s rank correlation analysis was used to control for the effect of confounding factors. Correlations between continuous variables following treatment (same patients) were examined with the use of paired t-test and Wilcoxon signed-rank test for non-normal variables. Results were considered statistically significant when P < 0.05.

More details on PBMC isolation, RNA extraction, gene expression analysis, sample size calculation and patient and public involvement are given in Supplementary methods.

3. Results
3.1. The RNA editor ADAR1 is increased in RA synovium and peripheral blood
Expression analysis of the RNA-sequencing dataset GSE89408 revealed significant upregulation of total ADAR1 in synovium of patients with either early RA or established RA (P < 0.001 for each comparison: early or established RA vs. normal synovium or osteoarthritis, Fig. 1A). Interestingly there was a trend upregulation of ADAR1 in early RA when compared to established RA (P = 0.06 early vs. established RA). ADAR1 has two isoforms, namely ADAR1p110 and ADAR1p150 [6,28]. ADAR1p110 is constitutively expressed [6,28], while the interferon-inducible ADAR1p150 was reported to be also induced by the pro-inflammatory cytokines TNFα and interferon-γ [14]. Expression of ADAR1p150 isoform was significantly increased in rheumatoid synovium as well as in PBMCs of active RA patients (Fig. 1B,D), while mRNA levels of the constitutively expressed ADAR1p110 isoform did not differ between patients and controls (Fig. 1C,E). The expression levels of the second RNA editor ADAR2 (ADARB1) were similar in RA patients and healthy controls (Supplementary Fig. 2) indicating that ADAR1p150 is the only RNA editing enzyme induced under pro-inflammatory conditions in RA patients. Taken together these results suggest that total ADAR1 and particularly its pro-inflammatory isoform ADAR1p150 is increased in RA in both PBMCs and synovial tissue.

3.2. Increased Alu A-to-I RNA editing rate in active RA
In order to evaluate the RNA editing activity of ADAR1, we sequenced the AluSx⁺ that is located in cathepsin S mRNA 3’ UTR (Fig. 2A), which is edited only by ADAR1 and is responsible for the post-transcriptional regulation of cathepsin S mRNA stability and expression [14]. Since inosines are recognized by the reverse transcriptase as guanosines, we could detect A-to-I RNA editing levels of modified adenosines as A-to-G nucleotide mismatches when we compare the RNA sequence with the genome. We were able to detect more than 20 edited adenosines (Supplementary Table 1). Both the average of all edited adenosines (P = 0.02) in AluSx⁺ as well as the individual RNA editing rate of 8 specific adenosines (Fig. 2B) were significantly higher in RA compared to controls (6–47% increase per adenosine, P < 0.05 for all, Fig. 2B). The RNA editing rates of 16 individual adenosines within cathepsin S 3’ UTR AluSx⁺ strongly correlated with ADAR1p150 mRNA levels (r = range 0.454–0.754, P ≤ 0.05 for all-Supplementary Table 1) The average RNA editing rate of the 8 significantly higher edited nucleotides (depicted in Fig. 2B) correlated with the expression of the pro-inflammatory inducible ADAR1p150 isoform (n = 19, r = 0.623, P = 0.004, Fig. 2C), but not with the constitutively expressed ADAR1p110 isoform (r = 0.265, P = 0.27).

3.3. Treatment of RA decreases A-to-I RNA editing levels only in responding patients
Next, we investigated whether the observed upregulation of ADAR1p150 and AluSx⁺ RNA editing rate in active RA was reversible after 12 weeks of antirheumatic treatment with csDMARDS, corticosteroids and/or bDMARDs. Of note, both ADAR1p150 expression levels and average AluSx⁺ RNA editing rate decreased post-treatment only in EULAR responders (P = 0.008 and P = 0.02, respectively) (Fig. 3A and B), while they remained unaffected in those patients with moderate to no response (Fig. 3C and D). These results indicate that the effect of the antirheumatic treatment on RNA editing machinery depends on the clinical response to the given therapy, thus suggesting that the dynamic regulation of Alu A-to-I RNA editing reflects the course of RA.

3.4. Increased A-to-I RNA editing is associated with pro-inflammatory gene expression
A-to-I RNA editing of Alu elements disrupts the double-stranded RNA structure through the production of weak I–U bonds, unwinding double-stranded RNA secondary structure [10]. In this way, double-stranded RNA is regionally converted into a more single-stranded structure enabling the binding of single-strand RNA-binding proteins, such as HuR (ELAVL1), which can stabilize mRNAs and, thus, increase their expression, as we have previously described for cathepsin S mRNA [14]. Herein, we used cathepsin S, not only as a well-established target of ADAR1 [14] but also as an important molecule in RA development [18] to test whether the observed upregulation of ADAR1-mediated Alu RNA editing rate was associated with increased cathepsin S mRNA expression in RA. First, we confirm previous reports showcasing that cathepsin S mRNA expression was significantly increased in PBMCs of active RA patients (1.34-fold increase compared to healthy controls, P < 0.05, Fig. 4A), as well as at the synovium of RA patients (6-fold increase compared to normal synovium, P < 0.001, Fig. 4B). Most importantly, cathepsin S expression significantly correlated with the pro-inflammatory inducible ADAR1p150 isoform (r = 0.623, P = 0.004), as well as with the RNA editing rate of 12 individual adenosines within cathepsin S AluSx⁺ (r = range 0.461–0.716, P ≤ 0.05 for all; Supplementary Table 1) and mean RNA editing rate
(r = 0.589, P = 0.008, Fig. 4C) in active RA. In contrast, no association was observed between the constitutively expressed ADAR1p110 isoform and cathepsin S mRNA levels (r = 0.268, p = 0.27). Next, we studied the association between the expression of ADAR1 and cathepsin S in a large cohort of RA synovial tissue. Interestingly, a similar correlation between ADAR1 and cathepsin S mRNA was also observed in synovial tissue derived from RA patients (n = 152, r = 0.516, P < 0.001, Fig. 4D) indicating that inflammation-induced Alu A-to-I RNA editing may augment cathepsin S mRNA.

Since RNA editing controls cathepsin S expression through increased HuR binding to AluSx+ [14], we statistically controlled for the potential involvement of HuR in regulation of cathepsin S expression by ADAR1p150-mediated RNA editing in RA. Towards this goal, we used a linear regression analysis for cathepsin S mRNA expression levels (dependent variable) and ADAR1p150 or average cathepsin S AluSx+ RNA editing rate controlling for the effect of HuR expression in these samples. Indeed, controlling for HuR abolished the observed relationship between cathepsin S mRNA expression and ADAR1p150 (P = 0.58) or average AluSx+ RNA editing (P = 0.29) suggesting that regulation of cathepsin S expression by RNA editing in RA is mainly mediated by HuR.

Of note, cathepsin S is only an example among several Alu-enriched
Fig. 2. Increased Alu A-to-I RNA editing rate in active RA A.
Schematic representation of cathepsin S 3' UTR and cDNA sequence of cathepsin S AluSx+. All detected edited adenosines are depicted in green, while individual adenosines with significantly increased RNA editing rates in RA PBMCs are depicted in red. HuR binding sites are underlined. B. Chromatopherograms depict the genomic sequence (upper part) and the cDNA sequence (lower part) showing the RNA editing events (A-to-G mismatches). Scatter-plots depict RNA editing rate of 8 individual adenosines in PBMCs of active RA patients (n = 19) vs. healthy controls (n = 14). RNA editing rates of individual adenosines were determined following analysis of chromatopherograms. C. Scatter-plot (left) of mean cathepsin S AluSx+ RNA editing rate of the 8 edited adenosines in PBMCs from patients with active RA (n = 19) vs. controls (n = 14) and scatter-plot (right) showing correlation of individual RNA editing rates with ADAR1p150 mRNA levels in RA patients. Lines in the scatter-plots represent mean ± SEM. P-values were produced by Mann-Whitney U test/independent-samples t-test or Spearman's rank test, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
pro-inflammatory genes, which are predicted to be highly edited according to the RNA editing database RADAR [29] and whose regulation by RNA editing in inflammatory disease warrants further investigation. For example, TNF receptor-associated factors TRAF1, TRAF2, TRAF3, TRAF5 have 10–117 Alu elements and 18-1135 predicted RNA editing sites, while when we analysed their expression in rheumatoid synovium, we found a significant correlation with ADAR1 expression (n = 152, r = range 0.356–0.743, Supplementary Table 2).

Taken together, our findings imply that Alu RNA editing may be a global primate-specific mechanism controlling Alu-enriched inflammatory mediators at post-transcriptional level through HuR-mediated RNA processing/stability in patients with RA. The proposed mechanism is summarized in Fig. 5.

**4. Discussion**

The present study provides first evidence that: 1) the RNA editor ADAR1, and specifically ADAR1p150 isoform, is increased in diseased synovium, the target tissue in RA, 2) the pro-inflammatory ADAR1p150 isoform as well as the A-to-I RNA editing rates of repetitive Alu elements are increased in circulating mononuclear cells from patients with active RA, 3) after a 12-week antirheumatic treatment reductions of ADAR1p150 expression and Alu RNA editing rates are prominent only in patients with good clinical response and 4) increased ADAR1 RNA editing rate is associated with increased pro-inflammatory activity of ADAR1 isoform ADAR1p150 is a type I interferon-inducible gene [28]. Accordingly, ADAR1 expression was found to be increased in type I interferon-associated autoimmune diseases [34–39], but also in other inflammatory diseases including acute myocardial infarction, atherosclerosis, cancer and viral infections [14,40,41]. More importantly, ADAR1 expression and activity are increased after stimulation with TNFα [14], the major cytokine that regulates the dynamics of transcriptome in RA [42], due to a significant increase in levels of ADAR1p150 isoform [14]. Our results may suggest that ADAR1 and specifically the ADAR1p150 isoform is increased in RA through a synergistic effect driven by TNF and type I interferon signaling. The up-regulation of ADAR1 was found at the inflammatory tissues as well as in the circulation, and therefore, ADAR1 may have an impact on other autoimmune diseases besides RA. Further studies are warranted to elucidate the systematic and tissue-specific regulation and role of ADAR1, and more specifically of the pro-inflammatory ADAR1p150 isoform, in systemic autoimmunity.

ADAR1-induced A-to-I RNA editing takes place primarily in repetitive Alu elements, therefore comprising an upstream regulatory mechanism of RNA metabolism specific for primates [8–13]. The widespread Alu elements have recently emerged as critical regulators of inflammation [43] and an enrichment of Alu elements has been recently reported in autoimmune diseases, potentially contributing to type I interferon pathway activation [43,44]. Data in systemic autoimmune disease are yet controversial, as studies have shown up-regulation of global Alu RNA editing index in peripheral blood of patients with systemic lupus erythematosus [38], but downregulation in keratinocytes of patients with psoriasis [39]. Clearly, comparative studies at transcriptome-wide level and especially at a single nucleotide or Alu level of individual transcripts in inflammatory and autoimmune diseases are lacking ADAR1 or having an editing-deficient knock-in mutation die in utero [30–32]. In humans, mutations in ADAR1 cause Aicardi-Goutières Syndrome associated with a type I interferon signature [33]. Further, ADAR1 and especially its isoform ADAR1p150 is a type I interferon-inducible gene [28]. The effect of antirheumatic treatment on ADAR1p150 and RNA editing is dependent on clinical response.
warranted to enlighten the tissue-specific and disease-specific effects of Alu A-to-I RNA editing. The difference between global and transcript-specific RNA editing levels among the inflammatory and especially the autoimmune diseases may reflect not only the tissue-specificity of RNA editing [45], but also the inflammatory microenvironment and milieu in each disease [46]. We found that effective anti-inflammatory

Fig. 4. Cathepsin S mRNA expression in RA A. Cathepsin S mRNA expression in PBMCs of patients with active RA (n = 19) and controls (n = 14) quantified by RT-qPCR. B. Expression analysis of cathepsin S mRNA levels at synovial biopsies derived from the RNA sequencing dataset GSE89408. Aligned RNA-seq data were downloaded from Gene Expression Omnibus. C. Scatterplot showing correlation of individual cathepsin S mRNA levels with RNA editing rate in active RA PBMCs. D. Scatterplot showing correlation of ADAR1 and cathepsin S mRNA levels at synovial tissue of RA patients from RNA-sequencing dataset GSE89408; values are shown after logarithmic transformation. Lines in the scatter-plots represent mean ± SEM. P-values were produced by Mann-Whitney U test or Spearman’s rank test/Pearson’s test, respectively. Est. RA: established rheumatoid arthritis (> 1 year disease duration).

Fig. 5. Proposed mechanism: Increased Alu A-to-I RNA editing fuels inflammatory gene expression in RA through post-transcriptional regulation of RNA metabolism. Synovial or systemic inflammation induces the expression of the pro-inflammatory inducible ADAR1p150 isoform and consequently A-to-I RNA editing of Alu elements. Increased Alu RNA editing may fuel inflammation by controlling the expression of inflammatory mediators through post-transcriptional HuR-dependent mRNA stability/processing. Certain items on this figure have been adapted from Servier Medical Art by Servier (https://smart.servier.com) – licensed under Creative Commons Attribution 3.0 Unported License). p110: ADAR1p110, p150: ADAR1p150, HuR: human antigen R, TRAFs: TNF receptor-associated factors.
treatment was able to decrease RNA editing levels in our patients. To the best of our knowledge this is the first report showing the effect of anti-inflammatory treatment on RNA editing levels in association with clinical response. Whether RNA editing is a predictive biomarker of clinical response and/or involved in the response to therapy remains to be investigated in future studies.

Alu RNA editing can control various aspects of RNA metabolism including splicing and mRNA stability, which can ultimately affect expression levels of edited genes [10,12,47,48]. One of the major mechanisms leading to increased mRNA stability is the binding of the stabilizing RNA-binding protein HuR to its target motifs TTTTG, TTTTT and ATTATA located within the Alu elements [47,49]. Therefore, we also checked whether increased RNA editing observed in active RA patients led to aberrant gene expression, serving probably as an additional, primate-specific regulatory mechanism at the post-transcriptional level. Towards this goal, we used as exemplar the well-established ADAR1-target cathepsin S [14]. Cathepsin S belongs to the family of lysosomal cysteine proteases, which are critically involved in antigen presentation and immune response [50]. In mice, knockout of cathepsin S prevents collagen-induced arthritis [18], whereas increased circulating cathepsin S protein levels have also been detected in RA patients [20]. Moreover, cathepsin S may contribute to autoantibody production since it specifically mediates degradation of the invariant chain β2, thereby promoting MHC-II antigen binding [15,16]. Previous studies have shown a significant upregulation of cathepsin S in RA synovial fluid [19], suggesting a role in synovial inflammation by its elastolytic properties and/or by enhancing antigen presentation and autoantibody production.

Our findings show a significant increase of cathepsin S both at the synovium and PBMCs of patients with RA which is significantly associated with the expression of ADAR1 and especially of ADAR1p150, as well as with the individual RNA editing rate of the adenosines located within the AluSx” of cathepsin S 3’ UTR. These associations imply that the disruption of AluSx” double-stranded RNA structure to a more single-stranded one by A-to-I RNA editing reveals the HuR target-sequences enabling the single-strand RNA-binding protein HuR to bind to its target cathepsin S, thus increasing its mRNA stability and expression, as previously evidenced in a series of mechanistic studies [14]. Of interest, cathepsin S is only one of the many edited Alu-enriched genes, e.g. TRAFs, whose expression was found to be associated with ADAR1 in a large cohort of RA patients, suggesting that our proposed mechanism (Fig. 5) may be applicable to a large number of pro-inflammatory genes in RA or other inflammatory diseases. Further studies are warranted to evaluate the role of A-to-I RNA editing in transcriptome metabolism and cellular function in RA.

5. Conclusions

To conclude, our data reveal a previously unrecognized dynamic regulation of Alu A-to-I RNA editing in RA that underpins therapeutic response and fuels inflammatory gene expression. The pro-inflammatory inducible ADAR1p150 isoform acts as a transcriptome-wide regulator of human-specific Alu-enriched inflammatory gene expression and, thus, may comprise an interesting predictive biomarker and a potential therapeutic target in patients with chronic inflammatory (auto)immune-mediated diseases, as supported by recent promising results in preclinical cancer models [51,52].

Declarations of interest

None.

Conflicts of interest

There are no competing interests for any author.

Contributorship

K.Stellos and P.P.S. designed and guided research; N.I.V., M.G.T. and P.P.S. recruited the PBMC cohort; N.I.V. and A.Gatsiou performed research; D.A.S. and A.Gallo performed the analysis of RNA-seq data for ADAR1 isoform expression; N.I.V., K.Stamatelopoulos, P.P.S. and K.Stellos analyzed data; A.Gatsiou, K.Stamatelopoulos, M.G.T., P.P.S and K.Stellos gave conceptual advice; and N.I.V. drafted the paper with input from all authors.

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Ethical approval information

All participants gave written informed consent, which had been previously approved by the Ethics Committee of Laiko Hospital, Athens, Greece (Protocol Nr.:1368/17-11-2016), as well as by the Hellenic Data Protection Authority (Protocol Nr.:1/EN/ES/7901-2/22-12-2016).

Data sharing statement

RNA-sequencing dataset GSE89408 is publicly available at GEO database (NCBI). Raw data of the qPCR experiments and RNA editing analysis are available by the corresponding author upon reasonable request.

Patient and public involvement

We did not involve patients or the public in our work.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jaut.2019.102329.

References


