Synovial Fluid Cell Proteomic Analysis Identifies Upregulation of Alpha-Taxilin Proteins in Rheumatoid Arthritis: A Potential Prognostic Marker

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Rheumatoid arthritis (RA) is a chronic autoimmune inflammatory disease affecting the joints and surrounding tissue. Identification of novel proteins associated with the progression of a disease is a prerequisite for understanding the pathogenesis of RA. The present study was undertaken to identify the potential biomarkers from a less explored biological sample such as synovial fluid (SF) cells which is specific for RA and to analyze their functional aspects using proteomic approach. Two-dimensional gel electrophoresis (2-DE) was performed using synovial fluid cells of RA and osteoarthritis (OA) patients, and 7 differentially expressed proteins were identified using matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS/MS). Alpha-Taxilin (α-Taxilin) has been found as one of the novel, significantly up regulated protein in RA. It has been validated in the synovium, synovial fluid (SF), SF cells, and plasma samples by Western blot, enzyme-linked immunosorbent assay (ELISA), fluorescence-activated cell sorting (FACS), immunohistochemistry (IHC), and real-time PCR. The identification of autoantibody against α-Taxilin and in silico studies has further helped us to understand its involvement in disease mechanism. The present study will therefore provide knowledge towards the etiology of RA that pave the way for suitable prognostic marker identification along with other clinical parameters.

1. Introduction

Rheumatoid arthritis (RA) is the most common chronic, systemic inflammatory autoimmune disease affecting around 1% of the population worldwide [1, 2]. It is often accompanied by systemic manifestation such as anemia, fatigue, and osteoporosis [3]. In spite of many efforts, the etiology of RA is still not clear [4, 5]. As the understanding about the disease is very limited, symptomatic treatment is mainly by the use of disease-modifying antirheumatic drugs (DMARDs), biologics, steroids, or combinations thereof [6, 7].

RA is believed to be mediated by an immune complex that does not clarify the acute inflammatory features of the disease [8]. Tumor Necrosis Factor (TNF), IL-6, anticyclic citrullinated peptide (anti-CCP), rheumatoid factor (RF), and antimannose binding lectin [9] (anti-MBL) are helpful in diagnosing the disease to a certain extent. However, these antigens are not specific whether they initiate autoimmune reactions, thus making the diagnosis critical [8, 10]. As the early stage of RA is characterized by nonspecific clinical symptoms, a delayed diagnosis leads to irreversible joint damage [11, 12]. Thus, there is an urgent and strong need for novel and more precise biomarkers with higher sensitivity and specificity for early diagnosis of disease. The study can therefore complement the conventional measures which may provide a more efficient marker to diagnose the disease for timely initiation of available therapies. Since the specific site of inflammation plays a vital role in the disease
progression, synovial fluid cell (SF cell) has been used as primary source to study the differentially expressed proteome profile. Additionally, synovium and plasma samples were also used to strengthen our findings.

We used two-dimensional gel electrophoresis (2-DE) followed by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS/MS) [13] to study the differential proteome profile of RASF compared to OA SF cells. OA SF cells were used as control SF cells since OA is an old age inflammatory joint disease but lacks autoimmune response [4]. OA is characterised by accumulation of synovial fluid and inflammation; thus, synovium and synovial fluid samples from OA patients have been widely used as control for RA study [14, 15]. Following the study, a total of 17 differentially expressed protein spots were observed. Among them, 7 spots have been identified successfully by mass spectrometry analysis. “TXLNA” or “Alpha-Taxilin/α-Taxilin” also known as interleukin-14 (IL-14) [16] or “High molecular growth factor” was found to be the most significantly upregulated protein in RA. Alpha-Taxilin belongs to the Taxilin family, which has three members including Alpha-(α)-, Beta-(β)-, and Gamma-(γ)- Taxilin. While α-Taxilin and γ-Taxilin are globally expressed, β-Taxilin (MDP77) is only expressed in the heart and skeletal muscles [16, 17]. α-Taxilin binds to proteins of the syntaxis family that are localized on the plasma membrane [18, 19]. The role of α-Taxilin has been reported earlier in exocytosis and cytokine activity which leads to inflammation via B-cell activation [20]; interestingly, literature support that an increased level of α-Taxilin has been found responsible for development of autoimmunity in transgenic mice [21], thus making our finding more relevant and important for understanding the etiology of RA.

2. Material and Methods

2.1. Sample Collection. Whole blood samples (RA = 100 (Age 50 y ± 5, male+female), Disease activity score (DAS) 28 = 6 ± 0.5), OA = 100 (Age 50 y ± 5, male+female) were collected hygienically in the EDTA-coated vacutainer (Ptech) from All India Institute of Medical Sciences (AIIMS, Ansari Nagar, New Delhi, India) and Army Hospital Research and Referral Hospital (Dhaulakuan, New Delhi, India). All enrolled participants had fulfilled the 2010 American College of Rheumatology criteria for RA and OA diagnosis, having radiological evidence and clinical history (detailed clinical parameter is given in “Supplementary Table 1”). Healthy volunteers (HC = 64) also required in this study having no radiographic evidence of joint degradation and other related clinical history. Synovial fluid (6 ml to 8 ml) were collected from RA (n = 16) and OA (n = 16) patients, and the synovium was collected after biopsy from RA (n = 6) and OA (n = 6), respectively. All the patients and healthy group provided signed written informed consent for these studies and were explained of all the associated risks before sample collection.

2.2. Isolation and Extraction of Proteins from Synovial Fluid Cells. Patient’s SF (=8 ml) was centrifuged at 4000 × g for 5 min in a swinging bucket rotor at 4°C. Cells were collected and washed 3 times with phosphate-buffered saline (PBS) at 300 × g for 5 min and incubated for 30 min at 4°C in SF cell lysis buffer (25 mM Tris, 1% Nonidet P-40, 150 mM Sodium Chloride (NaCl), 1.5 mM Ethylenediaminetetraacetic acid (EDTA), 0.5% Sodium dodecyl sulfate (SDS), 1 mM phenylmethyl sulfonyl fluoride (PMSF), and 1% v/v Protease Inhibitor cocktail (PI cocktail) followed by sonication at 20% amplitude for 5 min. The cell lysate was then centrifuged at 15000 × g for 30 min, and the supernatant was collected for further experiments.

2.3. Two-Dimensional Gel Electrophoresis (2-DE). Blood plasma samples (RA = 12, OA = 12, 50 y ± 5, male:female, 1:1) were pooled and quantified by the Bradford assay [22]. Three sets of 2-DE gels were run by using a pooled plasma sample from RA and OA after small modification [23]. Briefly, 150 μg protein was added to the rehydration buffer, loaded on immobilized pH gradient (IPG) strips (17 cm, pH 4–7 (Bio-Rad Laboratories), and was separated in the first dimension by isoelectric focusing (IEF) followed by the second dimension using sodium dodecyl sulfate-polyacrylamide gel electrophoresis [23] (SDS-PAGE).

2.4. Silver Staining. After 2-DE, gels were kept in fixative (50% methanol and 12% acetic acid) for an hour [24], followed by washing with 50% ethanol and 30% ethanol for 30 min each. Gels were then sensitized with 0.2% Na2S2O3 for 1 min, washed with Milli-Q thrice, and stained with 1 M AgNO3 and 700 μl/l formaldehyde (HCHO) solution for 20 min followed by the development of gels using a developer consisting Na2CO3 (60 gm/lit), HCHO (500 μl/lit), and Na2S2O3 (25 mm) [23]. The reactions were then stopped by adding 6% acetic acid.

2.5. In-Gel Digestion of Proteins and Preparation for MALDI-TOF MS/MS Analysis. For identification, differentially expressed protein spots were digested with trypsin gold (Promega, USA) as earlier [4]. Briefly, protein spots were excised into 1 mm cubes, washed with Milli-Q destained with 30 mM potassium ferricyanide and 100 mM sodium thiocyanate (1:1) at RT, and dehydrated by incubation with 1:1 acetonitrile (ACN)/water. Followed by washing with ACN, gel pieces were rehydrated (10 mM NH4HCO3 equal volume of ACN), vacuum dried, and digested at 37°C for 18 h with 20 μl of trypsin (20 ng/μl. Trypsin Promega). Peptides were extracted by varying concentrations of ACN and trifluoroacetic acid (TFA, Sigma-Aldrich, St. Louis, MO, USA). Peptide concentration (1 μl) and sinapinic acid (1 μl) were mixed and air dried and run using MALDI-TOF MS/MS (Applied Biosystems, Life Technologies, USA) as earlier [4]. MS/MS spectra were procured in reflector positive mode over the mass scope of 10,000-20,000 Da, and proteins having more than or equal to 2 unique peptides were considered.

2.6. Western Blot. Western blot was carried out as earlier [23] with little modification. Proteins (40 μg) were separated by running in 12% SDS gel and transferred to nitrocellulose (NC) membrane (Millipore, USA) at 20 V for 20 min using a semidry Western blot unit (Bio-Rad) followed by overnight
microtiter plates (Nunc, USA) were coated with 100 μl along with RASF (diluted sample (plasma/SF; dilution 1 : 200) and incubated with samples and healthy (1 : 500) individually as a source of primary antibodies followed by incubation with anti-human HRP-conjugated secondary antibody (dilution 1 : 5000) and developed by ECL substrate.

2.7. Autoantibody Detection. α-Taxilin protein was immuno precipitated using catch and release v2.0 (Millipore, Cat 17-500) and separated on 10% SDS-PAGE. The proteins were transferred on the NC membrane using the semidry western method (Bio-Rad). The presence of autoantibody against α-Taxilin in plasma was checked by incubating the blot with RA, OA, and healthy plasma (1 : 500) individually as a source of primary antibodies followed by incubation with anti-human HRP-conjugated secondary antibody (dilution 1 : 5000) and developed by ECL substrate.

2.8. Enzyme-Linked Immunosorbent Assay (ELISA). ELISA was performed using RA (n = 100) and OA (n = 100) plasma samples and healthy (n = 62) individual plasma samples along with RASF (n = 16) and OASF (n = 16) samples. ELISA microtiter plates (Nunc, USA) were coated with 100 μl diluted sample (plasma/SF; dilution 1 : 200) and incubated at 4°C overnight. After washing, wells were blocked with 1% BSA for 1 h at RT followed by 2 h incubation with 100 μl diluted (1 : 2000) anti-α-Taxilin (Santacruz, USA) at RT. Plates were then washed and incubated with 100 μl HRP-conjugated secondary anti-mouse antibody (dilution 1 : 1000) (Jackson, USA) for 1 h at RT. The reactions were then developed with orthophenylene dine (1 mg/ml) substrate for 15 min, terminated by adding 50 μl of 3 N H2SO4. The absorbance was measured at 492 nm (SpectraMax Plus 384) [25].

2.9. Fluorescence-Activated Cell Sorting (FACS). Fluorescence-activated cell sorting (FACS) was carried out using leukocytes and SF cells. Approximately 2 million cells were taken from each group and fixed with fixative (Cytoper BD) for 15 min and then washed and blocked with 1.5% BSA for 30 min in their respective vials. After washing, cells were incubated for 2 h with 200 μl diluted (1 : 2000) anti-α-Taxilin antibody followed by incubation with 200 μl anti mouse (Alexa Fluor 467) secondary antibody (1 : 1000) for 30 min. Cells were then washed again with PBS and resuspended in 500 μl of PBS, and 10000 cells were acquired in each run using FACS caliper (BD Biosciences) and analysed.

2.10. Immunohistochemistry (IHC). Tissue samples RA (n = 6) and OA (n = 6) were fixed in 10% formalin for 1 h, paraffin-embedded, and cut into desired thickness (5 μm) using a microtome. Samples were then submerged in citrate buffer (pH 6.0), incubated at 40°C for 2 min periods repeatedly for three times, and then finally kept for 20 min in a microwave. The tissue sections were then equilibrated at RT in the buffer (3% formaldehyde) and rinsed with Milli-Q water. The anti-α-Taxilin antibody was then added to the sections at a 1 : 400 dilution and incubated overnight at 4°C. Anti-mouse HRP conjugated was used as a secondary antibody (1 : 200 dilutions) and incubated for 2 h. The sections were counterstained with hematoxylin for 20 s followed by incubation with freshly prepared 3, 3’-diaminobenzidine (DAB, Sigma-Aldrich, USA) till the section developed. The slides were then mounted with a cover slip and observed under a microscope [26].

2.11. RNA Extraction and Real-Time PCR. Total cellular RNA from SF cells were extracted using Trizol (Gene Mark), following the manufacturer’s instructions, and the cDNA was obtained by reverse transcriptase (Fermentas, source MULV). Quantitative real-time PCR was performed using ABI SYBR green PCR master mix and amplified by (Applied Biosystem StemOnePlus™) real-time PCR thermal cycling block. The primers for α-Taxilin were as follows: FP 5’-G GTTTTGGGAAAGGAGATCACG, RP 5’-GGAGCTTCATC TGCTTCGTGTG.

The fold change of mRNA was calculated using the comparative Delta threshold cycle (ΔCt) method with 18S as the loading control. All reactions were performed in triplicate for each sample.

2.12. String Pathway Analysis and Interaction Study. Online tool “STRING” pathway has been used to find out the interacting partner of α-Taxilin. Apart from collecting and reassessing available experimental data from database on protein-protein interactions, imported known biological pathways from curated database interaction predictions were made [27] by this tool.

2.13. Statistical Analysis and Software. The patient DAS-28 score was calculated using a freely available online DAS-28 calculator (Rheumakit) by providing the patient clinical parameter such as tender joint, swollen joints, ESR, and overall health status. Densitometric analysis was carried out using the “Image Lab v 3.00” (Bio-Rad Laboratories) analysis tool for all Western blots. A bar graph for all Western analysis was plotted using mean adjusted densitometric values obtained in Image Lab analysis with the help of Microsoft Excel (Microsoft Corporation). A dot plot for ELISA obtained by putting absolute absorbance for each group in “GraphPad Prism 7” and p values were obtained. The standard deviation was calculated for replicates and plotted on the bar graph. Data having a p value less than 0.05 was considered significant.

3. Results

3.1. Identification of Differentially Expressed Proteins by Two Dimensional Gel Electrophoresis (2-DE). 2-DE was carried out to compare the protein profile of RASF and OASF cells. MALDI-TOF MS/MS analysis successfully identified 7 proteins out of 17 marked differentially expressed protein spots from 2-DE (Figure 1). Among these identified spots, regulatory protein E2 (spot 1), shikimate kinase (spot 2), α-Taxilin (spot 3), protein kinase A type 1a regulatory subunit (spot 5), recombinant signal binding protein (spot 6), and putative enoyl (spot 15) were found to be upregulated in RA while
“chain A structure of Hop Tpr2a domain in complex” (spot 4), a stress-induced phosphoprotein, was observed to be downregulated in RASF cells (Table 1). After performing densitometric analysis of the 2-DE gel image, we found that α-Taxilin is one of the significantly upregulated (2.4) RASF cells compared to OASF cells.

3.2. Validation of α-Taxilin by Western Blot. Western blot analysis of RASF cells showed 2.22 fold and 3.6 fold up regulation of α-Taxilin as compared to OASF cells and OASF, respectively (Figures 2(a) and 2(d)). Similarly, the expression of α-Taxilin was found to be 1.68-fold higher in RA synovium as compared to OA synovium (Figure 2(b)). Furthermore, the expression was also compared in plasma samples of RA, OA, and HC (Figure 2(c)). Expression of α-Taxilin has been found to increase by 1.52-fold and 1.35-fold in RA plasma compared to HC plasma and OA plasma ($p \leq 0.048$), respectively.

3.3. Validation by Enzyme-Linked Immunosorbent Assay (ELISA). The ELISA results revealed 1.31-fold and 1.21-fold upregulation of α-Taxilin expression in RA plasma compared to HC and OA plasma, respectively ($p \leq 0.026$). Further, the ELISA results of RASF revealed 1.50-fold higher expression of α-Taxilin in RA compared to OA with significant $p$ value ($p \leq 0.0021$) (Figure 3(b)). However, we do not found a much significant difference of α-Taxilin expression between OA plasma and HC plasma (Figure 3(a)) which supports specificity of α-Taxilin for RA.

3.4. Validation by Fluorescence-Activated Cell Sorting (FACS). There were 88.13% positively stained leukocytes with α-Taxilin in RA as compared to 28.57% in OA and 19.7% in HC (Figures 4(a) and 4(b)). The expression of α-Taxilin in RA blood cells was 4.4 and 3.0 times higher as compared to HC and OA, respectively, whereas the expression of α-Taxilin was found to be 1.4 times higher in OA as compared to HC.

FACS analysis on synovial fluid cells was found to follow similar trends as observed in FACS performed on leukocytes. The data revealed that the total percentage of positively stained SF cells with α-Taxilin in RA was 62.3% and in OA was 15.4% (Figure 4(c)), suggesting 4 times higher expression of α-Taxilin in RASF cells compared to OASF (Figure 4(d)).

3.5. Validation by Immunohistochemistry (IHC) and Real-Time PCR. The expression of α-Taxilin was further checked in the synovium by the IHC method. Higher expression of α-Taxilin was observed in RA synovium at pannus formation as compared to a section of OA synovium where no pannus formation was observed (Figure 5(a)). The results were validated at the mRNA level by real-time PCR (data not shown).

3.6. Autoantibody Detection. The level of circulating autoantibody against α-Taxilin in the RA, OA, and HC plasma was
analysed. The autoantibody level present in RA against α-Taxilin is much lower than that in HC and OA; however, there is no significant increase in OA compared to HC. The expression level of autoantibody in RA patients has been found to be significantly downregulated (0.396-fold downregulated) compared to OA and HC (Figure 5(b)).

3.7. String Pathway Analysis and Interaction Study. The in silico pathway analysis was carried out by protein-protein analysis online tool “STRING” and revealed that α-Taxilin is interacting with ten proteins. The results from the association score of proteins showed that α-Taxilin is strongly associated with five proteins (Figure 6), namely, “Nascent polypeptide associated complex subunit alpha” (NACA), syntaxin-4 (STX4), syntaxin-3 (STX3), syntaxin 1A (STX1A), and Gamma Taxilin (TXLNG).

4. Discussion

Proteomics is a promising approach in identifying the disease-associated proteins. We have focused our study to find out the differentially expressed proteins in RA using proteomic approach and identified 7 differentially expressed proteins from RASF cells (Table 1). Among these identified
proteins, 6 proteins: regulatory protein E2, shikimate kinase, α-Taxilin, protein kinase A type 1α regulatory subunit, recombinant signal binding protein, and putative enoyl were found upregulated, while chain A structure of Hop Tpr2α domain protein was observed to be downregulated in RASF cells compared to OASF cells. Regulatory protein E2 plays a role in the initiation of viral DNA replication and is essential for the survival of the tubercle bacillus [29]. Downregulation of protein kinase A regulatory subunit was found to be responsible for endocrine and other tumors [30]. Putative enoyl reductase has a role in very long-chain fatty acid metabolic process, and chain A structure of Hop Tpr2α domain has role in protein-protein interactions [31]. Since RA is an autoimmune disease, we found that two proteins, namely, "α-Taxilin" and "recombinant signal binding protein" directly linked with autoimmune response. Both of these proteins have a role in B-cell differentiation, immune responses, and interleukin secretion [21]. Upregulated levels of α-Taxilin have been observed in renal cell carcinoma [19] and in development of autoimmunity in IL-14α-transgenic mice [21]. It also participates in the transferrin receptor recycling pathway by interacting with sorting nexin 4, involved in cytokine activity [20], B-cell activation, and exocytosis [21]. We also have found that α-Taxilin is one of the significant upregulated proteins in RA in our preliminary screening and moreover, its role in RA is yet not explored.

The level of α-Taxilin was also found upregulated in other diseases also such as in hepatocellular carcinomas, renal cell carcinoma, and astrocytic tumor and in proliferating neural stem cells during embryonic development in rat [32, 33]. Taxilin interact with the syntaxin family which is well known for their role in intracellular membrane fusion and regulation of exocytosis. It has been seen that the inflammatory marker has correlation with cancer [34]. Moreover, remodeling of tissues takes place in both RA and cancer [35, 36]. However, it is not clear to date whether Alpha-Taxilin takes part in tissue remodeling or not. It may be possible that Taxilin may act as a common pathway regulator for inflammation and cancer response.

In order to strengthen our finding and to find out the role of α-Taxilin in the pathogenesis of RA, validation assays were extensively carried out by Western blot and ELISA. SF cells, synovium, SF, and plasma sample were checked for the presence of α-Taxilin and has been found upregulated in RA compared to their respective controls. This was the first report where upregulation of α-Taxilin has been seen in RA plasma, RASF cell, and RASF. After analyzing the Western blot, we found a higher (3.6-fold and 1.52-fold) expression of α-Taxilin in RASF and RAPL as compared to OASF and HC plasma, respectively, while the results from ELISA revealed that there is 1.5-fold and 1.31-fold increase in the α-Taxilin level in RASF and RAPL, respectively. This was further strengthened by FACS analysis where we found a nearly 4-fold increased level in RAPL compared to OA and HC (Figure 4(b)). The significant upregulated expression of α-Taxilin in FACS along with ELISA from SF (Figure 3(b)) revealed that 89% of RA patients gave positive expression of α-Taxilin compared to OA and HC. The results were therefore further strengthened by IHC results using the synovium. The higher expression of α-Taxilin at pannus formation indicated its involvement in the pathogenesis of RA compared to OA. As per the literature, specificity of anti-CCP and RF factor was found to be 50% and 40%, respectively; however, the specificity was found 96% and 55%, respectively [37]. The cytokines have many limitations in RA diagnosis due to the effect of age, gender, and diet, and variability [38] thus has not been evaluated. We have compared anti-CCP and RF factor with the current marker in our biological sample. The sensitivity and specificity for Alpha-Taxilin has been found to be 79% and 50%, respectively, compared to RF and Anti-CCP. Though the sensitivity of Taxilin is higher than both the anti-CCP and RF, the specificity remains below that of the anti-CCP. Thus, addition of the Alpha-Taxilin level can be helpful for the evaluation of RA diagnosis and pathogenicity. It is reported that
overexpression of α-Taxilin induces autoimmunity in transgenic mice [21] where IFN-γ was observed to show initiation of autoimmune response. Similarly, T-cell cytokines such as IL-2 and IFN-γ were also found to be involved in RA pathogenesis [3, 39]. The activation of B cell by Taxilin leads to secretion of inflammatory cytokines. The pathway followed

Figure 4: Validation by FACS. Blood cells or SF cells from RA, OA, and healthy blood cells were isolated and proceeded for FACS analysis with α-Taxilin primary antibody followed by Alexa Fluor 467-tagged secondary antibody. (a) Data captured using FACS caliber showing percentage of cells positively stained with Alexa Fluor 467 in H, OA, and RA blood cells. (b) Graphical representation of FACS data in terms of cells stained with Alexa Fluor for H, OA, and RA samples. (c) Data captured using FACS caliber showing the percentage of cells positively stained with Alexa Fluor 467 in SF cells. (d) Graphical representation of FACS data in terms of cells stained with Alexa Fluor for OA and RA SF cells.
by Taxilin to induce inflammation is not clearly understood, but IFN-γ may be one of the key factors involved for initiation of autoimmune response. Report shows that the IFN-γ level has a correlation with Alpha-Taxilin upregulation in a mouse model [21]. As RA is an autoimmune disease, findings of similar biological disease-associated activity in the progression of autoimmunity in the development of RA, indicating that α-Taxilin may be one of the key role players responsible for the onset of RA.

To get further insight on the interacting partner involved in signaling pathways with α-Taxilin, in silico interaction study was carried out. The study revealed five interacting partners such as NACA, STX4, STX3, STX1A, TXLNG. Binding of α-Taxilin with the syntaxin family (STX4, STX3, and STX1A) is reported to play a primary role in the regulation of vesicle exocytosis and cytokine-mediated signaling pathway and regulation of immunoglobulin. An IgE secretion implicates its possible role in pathogenesis of RA. The
“NACA” also interact with α-Taxilin that is known to have a preventive role in appropriate targeting of nonsecretary polypeptide and regulation of cell proliferation and is responsible for muscle fiber development [40]. Furthermore, interestingly, we found a downregulation (0.396-fold, Figure 5(b)) of autoantibody levels in RA plasma compared to HC. The decreased levels of autoantibody thus drew our attention towards the Paul Eherlich’s statement about the existence of anti-autotoxin antibodies. It is reported that the decreased level of anti-autotoxin antibodies may lead to the disturbance of homeostasis [41]. Studies have also shown that the central nervous system (CNS) of trauma spontaneously evolves a neuronal loss [41] indicating that the presence of autoantibody towards the Paul Eherlich autoantigen in the synovial fluid of rheumatoid arthritis patients using an immunoproteomics approach, “PloS One, vol. 8, no. 2, article e56246, 2013.

5. Conclusion

Our study therefore concludes that α-Taxilin might be one of the prerequisite factors responsible for the onset of immune response in RA while considering our finding of significantly upregulated expression of α-Taxilin in RA against HC and OA. The study will be helpful towards the development of a novel prognostic marker to understand further understanding for the development of autoimmune response in RA.

Data Availability

Data supporting the findings of this study are available from the corresponding author [Dr. Sagarika Biswas] on request.

Ethical Approval

Patients (RA/OA) with detail history were recruited at the hospital. The study protocol was approved by medical Ethics Committee of All India Institute of Medical Sciences, Department of Orthopedics, New Delhi, India, (Ref no. IEC/NP-7412010) and CSIR-Institute of Genomics and Integrative Biology, Delhi, India. Written consent was taken from all the patients and healthy volunteers. All methods were performed in accordance with the principles of the Helsinki declaration and were approved by the Ethics Committee of All India Institute of Medical Sciences (AIIMS) and CSIR-Institute of Genomics and Integrative Biology (CSIR-IGIB).

Conflicts of Interest

The authors confirm that this article content has no conflicts of interest.

Authors’ Contributions

The study was initiated and designed by A.S and S.B. The experiments were performed by A.S., S.S., T.S, P.A., and P.K. The statistical analysis and the figures were performed by A.S., S.S., and P.A. The manuscript was written by A.S and S.B. and critically reviewed and edited by S.B. All authors reviewed and approved the final version of the article submitted for publication.

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Supplementary Materials

Supplementary table 1: summary of the clinical measures of 200 patients (RA = 100, OA = 100) and 64 healthy control people. (Supplementary Materials)

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