CSF microRNAs discriminate MS activity and share similarity to other neuroinflammatory disorders

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Abstract

Objective
To perform a comprehensive multicompartment analysis of microRNA (miRNA) expression in multiple sclerosis (MS) linked to disease activity and compared with other neuroinflammatory diseases through a retrospective cross-sectional study.

Methods
One hundred twenty-seven miRNAs were measured by PCR arrays on pooled CSF, serum, and peripheral blood mononuclear cell (PBMC) samples of 10 patients with relapsing MS and 10 controls. Sixty-four miRNAs were then measured by quantitative PCR on individual CSF samples of patients with relapsing or remitting MS and controls (n = 68). Fifty-seven miRNAs were analyzed in the CSF from a second cohort (n = 75), including patients with MS, neuroinfectious, or neuroinflammatory diseases and controls. MiRNAs significantly dysregulated in the CSF were analyzed on individual serum/PBMC samples (n = 59/48) of patients with relapsing or remitting MS and controls. Post hoc analysis consisted of principal component analysis (PCA), gene set, and pathway enrichment analysis.

Results
Twenty-one miRNAs were differentially expressed, mainly upregulated in the CSF during MS relapses. Relapsing MS and neuroinfectious/inflammatory diseases exhibited a partially overlapping CSF miRNA expression profile. Besides confirming the association of miR-146a-5p/150-5p/155-5p with MS, 7 miRNAs uncharacterized for MS emerged (miR-15a-3p/124-5p/149-3p/29c-3p/33a-3p/34c-5p/297). PCA showed that distinct miRNA sets segregated MS from controls and relapse from remission. In silico analysis predicted the involvement of these miRNAs in cell cycle, immunoregulation, and neurogenesis, but also revealed that the signaling pathway pattern of remitting MS is more akin to controls rather than patients with relapsing MS.

Conclusions
This study highlights the CSF-predominant dysregulation of miRNAs in MS by identifying a signature of disease activity and intrathecal inflammation among neuroinflammatory disorders.
Glossary

**DMT** = disease-modifying treatment; **EAE** = experimental autoimmune encephalomyelitis; **EV** = extracellular vesicle; **FDR** = false discovery rate; **GEL** = gadolinium-enhancing lesion; **HC** = healthy control; **IgG** = immunoglobulin G; **Infect-ND** = infectious neurologic disorder; **Inflam-ND** = inflammatory neurologic disorder; **KEGG** = Kyoto Encyclopedia of Genes and Genomes; **miRNA** = microRNA; **mRNA** = messenger RNA; **MS** = multiple sclerosis; **OCB** = oligoclonal band; **PBMC** = peripheral blood mononuclear cell; **PCA** = principal component analysis; **qPCR** = quantitative PCR; **ROC** = receiver operating characteristic; **RRMS** = relapsing-remitting MS; **RT** = reverse transcription; **SC** = symptomatic control; **SiPa** = signaling pathway; **Treg** = regulatory T-cell.

Multiple sclerosis (MS) is characterized by multifocal inflammatory lesions inducing myelin sheath damage and axonal degeneration. MS results from a complex interplay between genetic susceptibility and environmental and epigenetic factors, but its molecular determinants remain elusive. A predominance of patients present initially with a relapsing-remitting disease course.1

MicroRNAs (miRNAs) are single-stranded noncoding RNAs regulating posttranscriptionally the expression of a large spectrum of genes during various biological processes.2 Several miRNAs have been associated with MS, but only few were linked to disease activity.3,4 Most studies were performed in small cohorts, in single biological compartments, rarely compared with other neurologic disorders or were not replicated, hence large heterogeneity between study populations and analysis methods. Few studies have focused on CSF miRNAs,5 although these might be more relevant to understanding disease regulation.

We sought to characterize miRNA expression in 3 biological compartments, i.e., CSF, serum, and peripheral blood mononuclear cells (PBMCs) of patients with MS according to disease activity compared with that of controls and other neurologic disorders in the CSF. MiRNAs known for their involvement in immunity, inflammation, neurodegeneration, and lipid metabolism were measured by quantitative PCR (qPCR). Finally, a principal component analysis (PCA) and a pathway enrichment analysis were performed, altogether leading to a comprehensive multicompartment characterization of the miRNA expression profile in MS.

Methods

**Patients**

CSF and blood samples were collected at the Cliniques Universitaires Saint-Luc (Brussels) between March 2005 and March 2018, processed according to international guidelines within 1 hour before storage at −80°C until miRNA extraction.6,7 PBMCs paired with serum were processed as previously described.8 The study was conducted stepwise (figure 1). A group size of at least 9 patients was calculated using the CSF fold changes of relapsing MS vs controls in the 2 miRNA arrays, with alpha error at 0.05 and power of 0.80 (stat.ubc.ca/~rollin/stats/ssize/n2.html). Patient cohorts were classified according to the BioMS-eu consortium definitions (table e-1, links.lww.com/NXI/A188).9 The 2017 McDonald criteria10 were used for MS diagnosis. Relapse was defined clinically and/or by the presence of gadolinium-enhancing lesions (GELs) on MRI for 93%–100% of patients, depending on the cohort.11 Remitting patients were clinically stable for 1 month up to 21 years. To reduce bias, patients were age and sex matched to controls, and none of the patients with MS were under any disease-modifying treatment (DMT). None of the relapsing patients had received high-dose IV methylprednisolone before sample collection. Clinical and paraclinical features of the patients are summarized in tables 1 and e-1.12,13 Diagnoses of other disease categories are listed in table e-2. Data were collected between February 2015 and July 2018.

**MiRNA isolation**

MiRNAs were isolated from CSF (400 μL) and serum (200 μL) using the miRNeasy Serum/Plasma Kit, and from 11-13.10^6 PBMCs using the miRNeasy Mini Kit, following the manufacturer’s protocol (Qiagen, Hilden, Germany). Caenorhabditis elegans miR-39 mimic (Qiagen) was added to CSF and serum samples as spike-in control for relative quantification. MiRNA extraction, reverse transcription (RT), and PCR were performed sequentially.

**MiRNA PCR array**

MiRNA profiling was kindly performed by the manufacturer with “Inflammatory Response & Autoimmunity” and “T-cell & B-cell Activation” miScript miRNA PCR Arrays (Qiagen) on a pool of 10 patients with relapsing MS and 10 symptomatic controls (SCs, for CSF) or 10 healthy controls (HCs) (for serum and PBMCs).

**RT, preamplification, and qPCR**

RT was performed on 5 μL of miRNA using the miScript II RT Kit. One microliter of complementary DNA was preamplified using the miScript PreAmp PCR Kit following the manufacturer’s recommendations (Qiagen). The spike-in control C. elegans miR-39 mimic was not preamplified. qPCR reactions were performed in duplicate, except for miR-24-3p in CSF, a putative internal reference gene,14 which was assayed in triplicate. Patients’ subgroups and controls were equally distributed across each run to minimize interrun bias for a single miRNA target.

**qPCR analysis**

Sample selection was based on Ct variability within duplicates of a same sample and between samples, as well as melt curve
morphology. Ct was set at 40 in case of inefficient amplification. The miRNA transcripts were normalized using references (exogenous miR-39 for CSF/serum or intracellular RNU6B [small nuclear ribonucleoprotein U6B, later referred to as RNU] for PBMCs). Relative expression was determined by the comparative Ct method to the mean of the SCs or HCs (2−ΔΔCt with ΔΔCt = ΔCt [target-reference]sample − ΔCt [target-reference]mean of controls). Extraction efficiency was considered by comparing the mean Ct of miR-39 or RNU amplification between subgroups and miRNA extraction series. When extraction efficiency appeared variable, results were expressed relative to the mean of controls extracted using the same RT enzyme lot.

Gene target prediction and pathway enrichment analysis
Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways were investigated for miRNAs displaying a statistically significant relative expression difference above 1.5-fold between disease categories. MiRNAs were grouped in sets according to the disease subgroup and biological compartment. Two (table e-6, links.lww.com/NNX/A188) or three (figure S) databases were combined comprising predicted and validated miRNA targets, namely mirPath v.3 by DIANA TOOLS based on TarBase v7.0 (filtered with Pathway Union and false discovery rate [FDR] correction; snf-515788.vm.okeanos.grnet.gr/), miRNet (filtered with a 2.0 cutoff degree and using a hypergeometric Fisher exact test algorithm; mirnet.ca/), and enrichment using the Cytoscape stringApp (with data set from the filtered miRNet analysis and a confidence score cutoff set at 0.6). The miRNA-gene network was generated from the miRNet database, filtered (2.0 cutoff degree of analysis and 40.0 betweenness) and sketched in the Cytoscape stringApp to show the most representative genes.

Statistical analysis
For statistical analysis, a nonparametric 1-way analysis of variance Kruskal-Wallis test of all relative miRNA expression levels across subgroups was performed using GraphPad Prism 5.0, followed by a Dunn post-test for a 2-by-2 comparison of the subgroups. Dunn p values, with a significance threshold set at ≤0.05, were calculated and corrected by the Benjamini-Hochberg FDR method (astatsa.com/KruskalWallisTest/). Differential expression of miRNAs was considered only if statistically significant for both post-tests. PCA of continuous variables obtained from the CSF miRNAs screening was performed using the R FactoMineR program. The presence/absence of oligoclonal bands (OCBs), MRI lesion load, presence/absence of GELs, and diagnostic category were analyzed as illustrative qualitative variables. The

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AUC = area under the curve; HC = healthy control; Infect/Inflam-ND = patients with infectious or inflammatory neurologic disorders; KEGG = Kyoto Encyclopedia of Genes and Genomes; mRNA = messenger RNA; miRNA = microRNA; PBMC = peripheral blood mononuclear cell; PCA = principal component analysis; qPCR = quantitative PCR; Rel MS = relapsing MS; Rem MS = remitting MS; ROC = receiver operating characteristic; RRMS = relapsing-remitting MS; SC = symptomatic control.
missing values were replaced by the mean value of the subgroup to which the sample belonged to. Results were plotted using Rcmdr (cran.r-project.org/web/packages/Rcmdr/index.html). For miR-146a-5p, -150-5p, and -155-5p in CSF, the receiver operating characteristic (ROC) curves were calculated providing a cutoff with an optimal compromise between sensitivity and specificity to discriminate MS from SC.

Standard protocol approvals, registrations, and patient consents
Patients sign a charter upon hospital admission stating that the residual CSF and serum samples, collected for routine diagnostic procedures or patient care, can be used for retrospective academic research, without an additional informed consent (ethical approval 2007/10SEP/233). PBMCs and paired serum were collected separately under a distinct protocol.

Table 1 Baseline characteristics of patients and controls included in the CSF microRNA study

<table>
<thead>
<tr>
<th>CSF</th>
<th>MS vs SC</th>
</tr>
</thead>
<tbody>
<tr>
<td>n (% ♀)</td>
<td>68 (64.7)</td>
</tr>
<tr>
<td>Mean age (±SD)</td>
<td>35.1 (±10.8)</td>
</tr>
<tr>
<td>n (% ♀)</td>
<td>40 (72.5)</td>
</tr>
<tr>
<td>Mean age (±SD)</td>
<td>34.5 (±11.7)</td>
</tr>
<tr>
<td>Disease duration in months: mean (±SD)</td>
<td>5 (±16)</td>
</tr>
</tbody>
</table>

Relapse phenotype (%)

1: Spinal
2: Brainstem/cerebellum
3: Optic neuritis
4: Supratentorial
5: Multifocal

EDSS score: median [range]

No. of relapses since disease onset: median [range]

Proportion of patients with a 1st clinical event (%)

Barkhof criteria* scores 3 and 4 (%)

GELs on brain MRI: mean (±SD)

Proportion of patients with spinal cord lesions on MRI (%)

GELs on spinal cord MRI: mean (±SD)

Proportion of patients with at least 1 GEL on brain or spinal cord MRI (%)

No. of CSF cells/mm³: mean (±SD) [range]

Proportion of patients with CSF-specific IgG OCBs or CSF FKLCs (%)

Proportion of patients with IgG intrathecal synthesis (%)

% IgG intrathecal synthesisb: mean (±SD)

Proportion of patients with IgM intrathecal synthesis (%)

% IgM intrathecal synthesisb: mean (±SD)

Abbreviations: % ♀ = percentage of female; EDSS = Expanded Disability Status Scale; FKLC = free kappa light chain; GEL = gadolinium-enhancing lesion; IgG = immunoglobulin G; IgM = immunoglobulin M; NA = not applicable; OCB = oligoclonal band; SC = symptomatic control.

Number of patients for which data are available is indicated by “n.”

* Barkhof criteria was assessed to reflect the lesion load (for definition, see reference 13).

b IgG and IgM intrathecal fractions were calculated using Reiber formulas (see reference 12, normal value is 0%). Patients with relapsing-remitting MS were disease-modifying treatment naive. Furthermore, none of the patients had received high-dose IV methylprednisolone. According to the consensus definition (see reference 9), SCs are patients with neurologic symptoms, but without any objective clinical or paraclinical findings to support the diagnosis of a specific neurologic disease.
research protocol upon signature of an informed consent (ethical approval 2007/31AOUT/222).

Data availability
Raw PCR files and detailed in silico and statistical outputs are not included in this article. Any unpublished and anonymized data will be shared upon request from a qualified investigator. Strengthening the Reporting of Observational Studies in Epidemiology guidelines have been respected.

Results

Multicompartment screening of immune-related miRNAs from a pool of patients with relapsing MS vs SCs/HCs

To investigate whether miRNA expression was modified in MS, particularly during relapses, we first performed a screening on a pool of CSF, serum, or PBMCs obtained from 10 patients with MS in relapse and 10 age- and sex-matched SCs or HCs (table e-1, links.lww.com/NXI/A188). The expression of 164 miRNAs was assessed using the “Human Inflammatory Response & Autoimmunity” and the “T-Cell & B-Cell Activation” miScript miRNA PCR Arrays. Thirty-seven miRNAs were assayed in both arrays. Absolute expression differences ranging between 3- (ΔCt = 1.59) and 148-fold (ΔCt = 7.21) were observed for 49 miRNAs, of which 35 originated from CSF, 9 from serum, and 8 from PBMCs. Of note, some miRNAs were found dysregulated in 2 compartments (miR-181c-5p and miR-210-3p in CSF and PBMCs; miR-34c-5p and miR-184 in PBMCs and serum). Most CSF miRNAs were overexpressed in the relapsing MS pool in comparison to the SC pool (ΔCt < 0), except miR-101-3p and miR-17-5p. Twenty-three miRNAs were discarded from subsequent analysis because of low expression levels (Ct ≥ 30) (figure e-1).

Quantification of miRNAs in the CSF of patients with relapsing and remitting MS vs SCs

As most miRNAs were found dysregulated in the CSF, we aimed to confirm these trends in individual CSF samples of patients with MS, according to disease activity (n = 40 in relapse and n = 13 in remission) and compared with age- and sex-matched SCs (n = 15) (table 1). In total, 64 miRNAs were amplified from individual CSF samples (table e-3, links.lww.com/NXI/A188). Those were either selected from the differentially expressed targets identified in the screening arrays, from previously published studies in MS or for their involvement in immunity, inflammation, lipid metabolism, or neurodegeneration.

Eighteen miRNAs were differentially expressed in at least 1 subgroup, with statistically significant (p ≤ 0.05) median fold changes ranging from −4.17 to 5.41 (figure 2A, figure e-2 and table e-4A, links.lww.com/NXI/A188). MiR-150-5p and miR-155-5p were upregulated, whereas miR-15a-3p, -34c-5p, and -297 were downregulated in patients with MS, irrespective of disease activity in comparison to SCs. Most miRNAs (n = 8) were upregulated in patients with relapsing MS compared with patients with remitting MS or SCs, except miR-124-5p, which was downregulated compared with SCs. Three miRNAs (miR-20a-5p, -33a-3p, and -214-3p) were downregulated in patients with relapsing MS compared with patients with relapsing MS and/or SCs, whereas miR-149-3p was upregulated. In addition, miR-24-3p was differentially regulated in patients with MS and could therefore not be used as an internal reference gene in this setting.

Specificity of the differentially expressed CSF miRNAs for MS

qPCR was performed for 57 miRNAs (table e-3, links.lww.com/NXI/A188) on individual CSF samples of 14 patients having infectious neurologic disorders (Infect-NDs), 12 patients having inflammatory neurologic disorders (Inflam-NDs), 15 patients with relapsing MS, 9 patients with remitting MS, and 25 SCs.

Twenty-four miRNAs were differentially expressed in at least 1 subgroup reaching a statistically significant threshold of 0.05 following Benjamini-Hochberg correction (table e-4B, links.lww.com/NXI/A188). These were all upregulated in Infect-ND compared with SC except miR-22-3p.

The levels of miR-21-5p, -142-3p, -223-3p, -342-3p, -423-5p, and let-7f-5p were increased in the CSF of both Infect- and Inflamm-ND vs SC, whereas miR-15b-5p, -29b-3p, and -29c-3p were upregulated in Infect-ND, Inflam-ND, and relapsing MS compared with SC. miR-150-5p was the only miRNA upregulated in all disease subgroups in comparison to SC. The profile of overexpressed miRNAs in these disease categories was strikingly similar except for the expression levels that were 2- to 6-fold higher in the Infect-ND subgroup and even 14-fold higher for miR-150-5p compared with relapsing MS (figure 3).

In addition, 46 of these miRNAs were investigated on CSF samples of 16 patients having neurodegenerative disorders and 17 patients with peripheral nervous system disorders, but no differences in expression levels were found, except compared with Infect-ND (data not shown).

Quantification of miRNAs in the serum of relapsing and remitting MS vs SCs

To determine whether the dysregulated miRNAs identified in the CSF were also differentially expressed in peripheral compartments, qPCR was performed for 24 miRNAs (table e-3, links.lww.com/NXI/A188) on individual serum samples of 20 patients with relapsing MS, 19 patients with remitting MS, and 20 HCs.

Relative quantification of 6 miRNAs reached statistical significance, with fold change in the median miRNA expression level ranging between −3.23 and −1.47 (figure 2B and table e-4A, links.lww.com/NXI/A188). Five miRNAs (miR-15a-3p, -24-3p, -126-3p, -146a-5p, and -181c-5p) were downregulated in both relapsing and remitting MS compared with HC, whereas miR-214-3p was downregulated in relapsing MS compared with HC only.
Quantification of miRNAs in PBMCs of patients with relapsing and remitting MS vs HCs

qPCR was performed for 24 miRNAs (table e-3, links.lww.com/NXI/A188) on individual PBMC samples of 18 patients with relapsing MS, 16 patients with remitting MS, and 14 HCs. Only 1 miRNA, miR-34a-5p, was significantly downregulated in relapsing MS compared with remitting MS with a fold change of $-1.89$ (table e-4A).

PCA and correlation with CSF parameters

To identify the combination of variables that best explain the differences between patients with MS and SCs, we performed a PCA on upregulated CSF miRNAs that showed more than 1.5-fold variation between these groups. Significantly upregulated miRNAs in patients with MS (miR-150-5p, -155-5p, -146a-5p, -21-5p, and -149-3p) accounted for 40.32% of the sample population variance, whereas other numerical variables such as the immunoglobulin G (IgG) or immunoglobulin M intrathecal fractions and CSF pleocytosis accounted for 18.43% of the variance, allowing an excellent discrimination between patients with MS and SCs (figure 4, A and B). The relative contribution of selected miRNAs to the variability of the population was significant as indicated by their $\cos^2$ value, with the most significant miRNAs being miR-146a-5p and miR-150-5p (table e-5A, links.lww.com/NXI/A188). Conversely, the analysis performed with the miRNAs that were significantly downregulated in MS vs SC (miR-15a-3p, -20a-5p, -33a-3p, -34c-5p, -124-5p, -214-3p, and -297) accounted only for 31.95% of the sample population variance (data not shown). ROC curves were plotted for miR-150-5p, -146a-5p, and -155-5p, with an area under the curve of 0.88, 0.82, and 0.8, respectively ($p < 0.0001$). Relative cutoff value for miR-150-5p was 1.9, with a sensitivity of 80% and a specificity of 81% for MS diagnosis (figure 4C). In addition, miR-150-5p expression levels were moderately correlated with CSF pleocytosis, but not with other CSF parameters, both in the MS vs control cohort and in the cohort of different Infect-NDs/Inflam-NDs (Spearman $r = 0.47$ and $r = 0.58$, respectively, $p < 0.0001$). miR-150-5p levels were also significantly upregulated in patients with CSF-specific IgG OCBs (data not shown).

PCA on miRNAs whose expression was modified in patients with relapsing vs remitting MS (miR-20a-5p, -145-5p, -214-3p, -27a-3p, -27b-3p, -24-3p, and -29c-3p), using the presence/absence of GELs as illustrative variable for disease activity, showed a good separation between patients with relapsing and remitting MS. This miRNA combination accounted for 77.52% of the population variance (figure 4, D and E), with the most representative miRNAs being miR-24-3p, -27a-3p, and -145-5p (table e-5B, links.lww.com/NXI/A188). Adding miR-149-3p, which is decreased 2-fold in relapsing vs remitting MS, did not improve disease activity discrimination (data not shown).

Figure 2 miRNA expression fold change in MS according to disease activity

Differences in CSF (A) and serum (B) miRNA expression levels. The fold change is calculated as the ratio between the median of the relative miRNA expression level of the respective subgroups. HC = healthy control; miRNA = microRNA; Rel = relapsing MS; Rem = remitting MS; SC = symptomatic control.
Biological significance and targeted genes of the differentially expressed miRNAs in MS

To obtain insight regarding the functions of the dysregulated miRNAs, we performed pathway enrichment analysis for the differentially expressed miRNA sets (more than 1.5-fold difference with corrected p value ≤ 0.05) between subgroups, as defined below and in figure 5. Some pathways were affected, independently from disease activity, such as cancer (p53 signaling pathway [SiPa], glioma, or chronic myeloid leukemia), cell cycle and proliferation (Wnt), immune (transforming growth factor beta and FoxO), cellular adhesion (focal adhesion and adherens junction) SiPas. We nevertheless observed that the enrichment score (defined as the -log₁₀[p value]) was always stronger for the relapsing vs remitting (Rel/Rem) subgroup compared with the 2 other subgroups. More strikingly, some pathways were found specifically in relapsing patients (neurotrophin, mTOR, and ErbB) compared with the 2 other groups, suggesting some relevance of these pathways in modulating disease activity. On the other hand, nuclear factor kappa-light-chain-enhancer of activated B cells was shown as general signaling for relapsing-remitting MS (RRMS). Some pathways were only affected in the Rel/Rem subgroup (axon guidance and insulin) or specifically in the Rel/SC subgroup (tumor necrosis factor alpha) (figure 5A). The comparison between up- and downregulated miRNAs in the different compartments showed a parallelism between pathways targeted by miRNAs upregulated in the CSF of relapsing MS and downregulated in the CSF of remitting MS (table e-6, links.lww.com/NXI/A188). In addition, the most representative genes retrieved from the analysis of all differentially expressed miRNAs in the CSF were introduced in a miRNA-gene interaction network. Several genes (MYC, VEGFA, MKN2, BCL2, PMAIP1, KIAA1551, XIAP, NFAT5, E2F3, WEE1, STAT3, CCND1, SMAD4, TMEM167A, and LRRC58) were targeted by at least 4 miRNA species from the 3 different sets, revealing possibly important miRNA-related genes affected in MS (figure 5B).

Discussion

CSF-predominant miRNA dysregulation and miRNA species newly associated with MS

This study sought to perform a comprehensive analysis of the differential expression of carefully preselected miRNAs in
patients with RRMS, according to disease activity, vs controls in 3 different biological compartments. Patients were free of DMT and devoid of high-dose steroids. A second CSF cohort was analyzed to compare the MS miRNA profile with that of other neurologic diseases. Strikingly, we observed 18 significantly dysregulated miRNAs in the CSF vs 6 in the serum and only 1 in PBMCs, pointing toward specific intrathecal processes ongoing in MS and confirming the results of the preliminary miRNA array analysis performed on pooled samples. We have identified 7 miRNAs, namely miR-15a-3p, -124-5p, -149-3p, -29c-3p, -33a-3p, -34c-5p, and -297 that have not yet been described in MS to the best of our knowledge, but rather found in cancer either as tumor suppressors or as oncogenes. Other miRNA species (miR-21-5p, -27b-3p, -145-5p, -146a-5p, -150-5p, -155-5p, -20a-5p, -214-3p, -24-3p, -27a-3p, and -125b-5p) were previously described as differentially expressed in RRMS in at least 1 of the following sample types: PBMCs or immune cell subpopulations, serum, whole blood, or active brain lesions. Moreover, this

**Figure 4** Patient stratification according to diagnosis or disease activity by PCA

PCA was applied using the significantly upregulated CSF miRNAs in patients with MS and other CSF parameters (IgG or IgM IF, CSF pleocytosis). PCA can distinctly separate MS from SC (A) in a 2-dimensional representation comprising in total 58.75% of sample population variability. The upregulated miRNA panel (i.e., miR-21-5p, -146a-5p, -149-3p, -150-5p, and -155-5p) segregates from other diagnostic criteria (Barkhof imaging criteria, IgM and IgG IF). Cos² values for each variable are plotted in abscissa and ordinate (B). They represent the strength of the individual variables in indicating population variability according to each dimension (significant above 0.5). ROC curves are plotted (C) for miR-150-5p, -146a-5p, and -155-5p as diagnostic biomarkers for MS with respective areas under the curve of 0.87, 0.82, and 0.8, respectively (p < 0.0001). PCA can also partially separate patients with Rel MS from patients with Rem MS (D) in a 2-dimensional representation comprising in total 77.52% of sample population variability. MiR-20a-5p, -24-3p, -27a-3p, -27b-3p, and -145-5p segregate from miR-29c-3p and miR-214-3p (E). AUC = area under curve; GEL = gadolinium-enhancing lesion; IF = intrathecal fraction; IgG = immunoglobulin G; IgM = immunoglobulin M; miRNA = microRNA; PCA = principal component analysis; Rel = relapsing MS; Rem = remitting MS; ROC = receiver operating characteristic; SC = symptomatic control.
study suggests that a specific set of miRNAs could serve as potential biomarkers for MS disease activity.

The CSF miRNA profile parallels the extent of intrathecal inflammation, but is not specific to MS

Although studies on CSF miRNAs in MS are sparse, several miRNAs identified in this study were previously described by other groups either in the CSF or other sample types. Our work emphasizes, however, the predominance of CSF miRNA dysregulation in MS, in concordance with its pathogenesis. Furthermore, our results highlight the partially overlapping profile of CSF-upregulated miRNAs in infectious and inflammatory CNS disorders compared with relapsing MS. The CSF miRNA expression level was related to pleocytosis, as it was higher in Infect-NDs than in relapsing MS, especially regarding miR-150-5p expression, which was also associated with the presence of OCBs.

The upregulation of CSF miR-150-5p in MS is in agreement with the previously published studies by Quintana et al. and Bergman et al. The expression of this miRNA is also increased in serum and exosomes from activated lymphocytes and is therefore considered as a sensor for activation of adaptive immunity. In this study, however, miR-150-5p was not upregulated in peripheral compartments, again suggesting the predominance of intrathecally mediated immune processes. We illustrated by PCA that miR-150-5p, -146a-5p, and -155-5p could significantly segregate patients with RRMS from controls. ROC curve analysis was the strongest for miR-150-5p, with a sensitivity and specificity of 80% and 81%, respectively, an even stronger result than previously shown and comparable to ROC curves of kappa free light chains for MS diagnosis. Further studies, in larger independent cohorts, should investigate whether these miRNAs could serve as biomarkers of disease activity.
MiRNA profile in the serum and PBMCs of patients with MS

Five miRNAs were downregulated in the serum of both relapsing and remitting MS (miR-15a-3p, -24-3p, -126-3p, -146a-3p, and -181c-5p), whereas miR-214-3p was downregulated in the serum of relapsing MS only. Remarkably, only downregulation of miR-15a-3p was consistent in both serum and CSF, whereas miR-24-3p, -146a-5p, and -214-3p were found upregulated in the CSF. Comparably, miR-126-3p was downregulated in whole blood MS samples.23 Other miRNAs (miR-122-5p, -196b-5p, -301a-3p, and -532-5p) found in extracellular vesicles (EVs) were also decreased in the serum of patients with relapsing MS.23 This discrepancy between the CSF and peripheral compartments was also shown in Alzheimer disease and intracerebral hemorrhage, but is yet not understood.24,25 The expression patterns of miR-181c-5p remain controversial, as some have described its upregulation in the CSF and in active MS lesions.5,26 In our hands, the levels of miR-181c-5p were however too low to allow quantification in the CSF, but it was downregulated in the serum of patients with MS compared with controls. Finally, in agreement with the decrease of miR-34a-5p levels in CD4+ T cells of patients with MS, we found its expression significantly downregulated in the PBMCs of patients with relapsing MS.27

Target analysis of miRNAs dysregulated in MS

To generate hypothesis regarding the functional role of the identified miRNAs in CSF, we performed a pathway enrichment analysis on several KEGG pathway databases to reveal the most affected SiPas according to disease activity, but also in serum and PBMCs. As a single miRNA species is not sufficient to account solely for the disease, as discussed above, we grouped the different miRNAs in sets. For CSF miRNAs, we first observed the highest enrichment score for several cancer SiPas, but we only presented the results for glioma and chronic myeloid leukemia because (1) these pathways are relevant for the (neuro)immunopathologic aspect of MS disease and (2) most of the miRNAs were extensively investigated in cancer research, which naturally biases most analysis toward cancer-related pathways. Besides cancer-related SiPas, the score comparison of the different subgroups of patients (relapsing, remitting MS and SC vs each other) and the Cytoscape miRNA-messenger RNA (mRNA) network representation obtained from CSF showed pathways consistent with MS pathogenesis such as cell cycle or apoptosis, immunoregulation, neuroprotective, and neurodegenerative processes.

MiRNA molecular function prediction by gene ontology provided the highest score to several negative regulators of transcriptional activity through targeting of CCND1, STAT3, the E2F family, MYC, SPI, ZBTB18, and PLAGL2 among others (findings obtained by miRNet; data not shown). This result confirms previous studies on the impact of miRNAs on the transcriptional network in MS pathogenesis.23,28–31

Our research demonstrates for the first time that the CSF profile of remitting MS (Rem) is more closely related to the profile of controls than relapsing MS (Rel) because (1) the SiPas target analysis scores were the lowest (if not absent) for the miRNAs differentially expressed between Rem vs SC (Rem/SC), whereas the scores were the highest for the ones between Rel/Ren and (2) that no common SiPas specifically stood out for the dysregulated miRNAs in Rel and Rem compared with controls. At the miRNA level, it is worth noting that, unlike most of the genes regulated by miRNAs from all groups, PLAGL2 and LMNB2 were specifically clustered to the Rel/Rem and Rem/SC subgroups. Of interest, LMNB2 was related to 2 miRNAs not previously associated with MS, namely miR-149-3p and miR-33a-3p. PLAGL2 is a transcription factor regulating Wnt signaling, and thus leading to the inhibition of neural stem cell differentiation.32 It could therefore be relevant for reparative processes during remission. LMNB2 codes for a member of the lamin family and was found 5-fold downregulated in CD4+ T cells of patients with remitting MS compared with HCs.33 Altogether, the pathway enrichment analysis with the miRNA sets identified in this study remarkably pinpoints to important pathways and miRNA-gene interactions pertinent to MS pathogenesis, as evidenced in earlier studies. The differentially expressed miRNAs have relevant mRNA interactions, especially in regard to the modulation of the immune response. For instance, miR-155-5p promotes Th17 and regulatory T-cell (Treg) differentiation, Th17 function, and microglia-mediated immune responses, through expression of interleukin (IL-) 6, tumor necrosis factor alpha, and iNOS, by targeting SOCS-1.34,35 MiR-155-5p also alters blood-brain barrier permeability and enhances leukocyte adhesion to endothelial cells, whereas miR-126-3p and miR-146a-5p counteract this by modulating the expression of brain endothelial ICAM1, VCAM1.36–38 Moreover, miR-146a-5p exerts its immunoregulatory effect by blocking Th17 differentiation through repression of TRAF6 and IRAK1, which are transducers of nuclear factor kappalight-chain-enhancer of activated B cells and thus reduces IL-6 and IL-21 expression.39 MiR-27b-3p inhibits Treg differentiation but not their function, by downregulating SMAD4 and TGFBR1.40 MiR-21-5p represses lymphocyte migration through STAT3 regulation.41 The deletion of miR-150-5p in experimental autoimmune encephalomyelitis (EAE) mice affects CD3+, CD4+, and CD8+ cells and miRNA expression of proinflammatory cytokines compared with wild-type EAE, likely by regulating CYMB.42

Cellular source and carriers of CSF miRNAs

Although our study did not investigate the cellular source of the CSF-associated miRNAs, according to the cell-specific miRNA catalogue, nearly all immune cell subpopulations express the miRNAs found in our study.43 Yet, some miRNA species might be more specific of certain cell types, as EVs of Treg cells were specifically enriched with miR-21-5p, -146-5p, and -150-5p, whereas those of Th1 and Th17 cells preferentially expressed miR-155-5p.44
Whether miRNAs are carried exclusively by EVs such as exosomes in the CSF is still an open question. However, miRNAs were enriched in exosomal fractions of CSF.45 The number of CSF EVs increases during MS relapses and GELs on brain or spinal cord MRIs were associated with an increase in CSF EVs expressing markers of CD8+ memory T cells, Th2 and Th1 cells.46 The cellular source of miRNAs from the CSF and their carriers should be then further investigated in the setting of MS.

**Study limitations**
The strength of this study resides in the comprehensive stepwise strategy used, the large CSF sample size, the use of a confirmatory cohort for the CSF, the comparison between compartments, and finally the comparison with miRNA expression in other disease categories. Moreover, patients were systematically sex- and age-matched to controls. However, there is inevitable heterogeneity between reports on miRNA expression analysis in MS due to discrepancies in sample numbers, collection and storage procedures, patient characteristics, study design, molecular biology techniques (microarray or qPCR, with or without preamplification, TaqMan vs SYBR Green), data analysis (quantification method, choice of reference gene), and finally the type of statistical tests. These factors, altogether or separately, possibly affect results. In this study, qPCRs from CSF and serum were normalized with an external reference, as an internal reference could not be identified. Future studies could possibly combine a panel of nonsignificantly altered miRNAs as normalizers. Some selected miRNAs were below the PCR detection level, preventing further analysis. The targeted selection of miRNAs based on the literature and immune-related miRNA PCR arrays does not exclude that other miRNA species are also dysregulated in MS. Finally, the patient cohort from a single academic center may not be representative of a more global MS cohort. An international and multicentric study that includes miRNA sequencing could provide novel miRNA targets that would refine our current results.

In conclusion, we have demonstrated that 21 miRNAs were differentially expressed in RRMS, mostly in the CSF, which is at the core of the inflammation, 7 of which were never associated with MS so far. The miRNA expression profile varied according to disease activity. It was associated with the extent of intrathecal inflammation, notably CSF pleocytosis, not only in MS but also in neuroinfectious and other neuro-inflammatory diseases. Considering this, a single miRNA is unlikely to be considered as a MS biomarker, but a set of miRNAs could rather be investigated for that purpose. The strength of this work resides in (1) the study of different disease categories, (2) the comprehensive miRNA analysis performed on 3 biological compartments in patients with MS, and (3) the validation of CSF results by our second cohort and some miRNAs already described in other studies. Our work emphasizes the potential of a miRNA set to reflect intrathecal inflammation and MS disease activity. The miRNA expression profile was in concordance with their known effects on immune and neuroinflammatory processes at play in MS. In silico analysis provided a view on affected SiPas, warranting further studies into their function, cellular source, and carriers, especially for the newly found miRNAs associated with MS. Ultimately, the present report will help fundamental research on MS by paving the way for the functional characterization of miRNAs and unraveling novel pathways for possible therapeutic intervention.

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References


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