RESEARCH ARTICLE

**Novel somatic single nucleotide variants within the RNA binding protein hnRNP A1 in multiple sclerosis patients**

[version 2; peer review: 3 approved]

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**Abstract**

Some somatic single nucleotide variants (SNVs) are thought to be pathogenic, leading to neurological disease. We hypothesized that heterogeneous nuclear ribonuclear protein A1 (hnRNP A1), an autoantigen associated with multiple sclerosis (MS) would contain SNVs. MS patients develop antibodies to hnRNP A1²⁹³-³⁰⁴, an epitope within the M9 domain (AA²⁶⁸-³⁰⁵) of hnRNP A1. M9 is hnRNP A1’s nucleocytoplasmic transport domain, which binds transportin-1 (TPNO-1) and allows for hnRNP A1’s transport into and out of the nucleus. Genomic DNA sequencing of M9 revealed nine novel SNVs that resulted in an amino acid substitution in MS patients that were not present in controls. SNVs occurred within the TPNO-1 binding domain (hnRNP A1²⁶⁸-²⁸⁹) and the MS IgG epitope (hnRNP A1²⁹³-³⁰⁴), within M9. In contrast to the nuclear localization of wild type (WT) hnRNP A1, mutant hnRNP A1 mis-localized to the cytoplasm, co-localized with stress granules and caused cellular apoptosis. Whilst WT hnRNP A1 bound TPNO-1, mutant hnRNP A1 showed reduced TPNO-1 binding. These data suggest SNVs in hnRNP A1 might contribute to pathogenesis of MS.

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Any reports and responses or comments on the article can be found at the end of the article.
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Competing interests: Drs. Michael Levin and Sangmin Lee have a United States patent pending titled "Biomarker for neurodegeneration in neurological disease".

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Amendments from Version 1

We appreciate the reviewer’s comments and have addressed them in the revised manuscript. None of the reviewers required further experiments. The reviewers requested none of the data indicating that SNVs may be the result of oxidative stress or clonal expansion of T-cells. They also suggested that SNVs in CNS targets other than neurons might be important in the pathogenesis of MS and that future studies should include more genes and control groups. All of these suggestions were specifically addressed in the discussion.

See referee reports.

Introduction

Multiple sclerosis (MS) is the most common autoimmune disease of the central nervous system (CNS) in humans, whose pathogenesis remains unknown. A number of genetic and immune studies indicate dysregulated immune responses as contributors to the pathogenesis of MS [27]. Genetic analyses show an association of major histocompatibility complex (MHC) Class II human leukocyte antigen (HLA)-DRB1 and single nucleotide polymorphisms (SNPs) related to immune function [12, 18]. Both Th1/Th17 CD4+ T-lymphocytes and immunoglobulins appear to have a causative role [2, 15]. Immunoglobulin G (IgG) responses to myelin and non-myelin targets have differentiated some MS patients from healthy controls [21, 22]. Non-myelin antigens that are targets for immunoglobulins isolated from MS patients include neurofilaments, axonal neurofascin and RNA binding proteins, including heterogeneous nuclear ribonucleoprotein A1 (hnRNP A1) [20, 21, 24].

Recently, mutations in RNA binding proteins have been shown to cause neurological disease [14]. For example, a mutation (p.D263V) in the prion-like domain (PrLD) of hnRNP A1 has been shown to cause familial amyotrophic lateral sclerosis (ALS) [12]. In addition to inherited mutations, somatic variants have also been shown to cause neurological disease [7]. hnRNP A1 performs a number of critical cellular functions related to transcription, nucleocytoplasmic transport of mRNA and translation [23, 24]. In addition to the PrLD, other important functional domains in hnRNP A1 include two RNA binding domains (RBDs) and M9, its nucleocytoplasmic shuttling domain [24]. M9 binds its nuclear receptor, transportin-1 (TPNO-1, also known as karyopherin β2) and the hnRNP A1:TPNO-1 complex is transported into and out of the nucleus [5, 9, 16, 26, 27].

Our lab has performed extensive studies on the role of autoimmunity to hnRNP A1 in MS and human T-lymphotropic virus type 1 (HTLV-1) associated myelopathy/tropical spastic paraparesis (HAM/TSP), a viral-induced model and clinical mimic of MS [28, 30]. Initially, we discovered that HAM/TSP patients develop antibodies to hnRNP A1 that cross-react with HTLV-1-tax, indicative of molecular mimicry [29, 31]. Next, the epitope of the HAM/TSP IgG response (AA293-304) was localized to M9 (AA298-304) [31]. M9 is a bipartite phenylalanine-tyrosine nuclear localization sequence (PY-NLS) that requires binding to TPNO-1 for hnRNP A1 to shuttle between the nucleus and cytoplasm [32, 33]. Because of the similarities between MS and HAM/TSP, we hypothesized that MS patients would also develop antibodies to hnRNP A1. In fact, antibodies isolated from MS patients, in contrast to healthy controls and Alzheimer’s patients, were also found to immunoreact with the identical hnRNP-A1-M9 epitope (AA293-304) [16]. Subsequent studies indicated that the IgG was biologically active and potentially pathogenic. For example, mono-specific antibodies to hnRNP A1 isolated from patients caused decreased neuronal firing using neuronal patch clamp in rat brain sections [11, 33]. Further, neurons exposed to anti-hnRNP A1-M9293-304 specific antibodies resulted in neurodegeneration and neuronal death [13, 14]. The anti-hnRNP A1-M9293-304 specific antibodies also caused changes in neuronal RNA expression that correlate with the clinical phenotype of MS and HAM/TSP patients (ie. spastic paraparesis), which was subsequently confirmed in neurons isolated from the brains of MS patients [16]. Additional studies showed that anti-hnRNP A1-M9293-304 specific antibodies entered neurons via clathrin-mediated endocytosis and caused apoptosis in a neuronal cell line [15]. Anti-hnRNP A1-M9293-304 specific antibodies also caused a redistribution of hnRNP A1 in neurons from nuclear to an equal distribution of nuclear and cytoplasmic localization, suggesting the antibodies interfered with M9, which is required for hnRNP A1 nuclear import [14]. Considering: (1) the role of hnRNP A1 in cellular function; (2) variant forms of hnRNP A1 cause neurodegenerative disease, and (3) hnRNP A1 is an autoimmune target in MS patients, we hypothesized that MS patients would contain novel genomic DNA single nucleotide variants (SNVs) in hnRNP A1-M9, which when expressed, would alter cellular function and contribute to cell death.

Methods

Patients

All blood samples were collected according to the approved Institutional Review Board protocols (Veterans Affairs Medical Center - Memphis, Study #317164, University of Tennessee Health Science Center, Study #98-06618-FB) with patient consent. The diagnosis of MS was made using published criteria (Supplement 1, see Results) [13].

Preparation of human peripheral blood monocytes (PBMCs) and isolation of genomic DNA

Human PBMCs were isolated from fresh blood by Ficoll-Paque gradient centrifugation and washed with PBS. Genomic DNA was isolated from PBMCs using the QIAmp blood kit (Qiagen Inc., Chatsworth, CA, U.S.A.) according to manufacturer’s protocol. All DNA samples were quantified using Nanodrop (Quawell) and restriction enzyme digestion methods.

PCR primers

Specific oligonucleotides were designed from the published genomic DNA sequence of the human hnRNP A1 gene. The upstream primer 5’-CAGATAAAGGC CCTCTTTCCC -3’ (3080–3100) and the downstream primer 5’-CTCAGCTACATTAGGGTTATTGGG -3’ (3667–3690) flank a 611 bp region of the human hnRNP A1 genomic DNA containing exon 8 and exon 9.

PCR amplification and subcloning

One microgram of genomic DNA was amplified in a reaction mixture containing the primers and KOD Hot Start DNA polymerase (Novagen). Use of this DNA polymerase has a mutation frequency of 0.10% . Before adding enzyme, the reaction mixture was heated at 95°C for 2 minutes. Amplification was carried out for 35 cycles of denaturation at 95°C for 20 s, annealing at 57°C for 10 s, and extension at 72°C for 15 s, followed by terminal elongation at 70°C for 20 s. The resulting PCR product was cloned into the pCR2.1-TOPO
blunt vector (Invitrogen), yielding pCR2.1-TOPO-Blunt-hnRNP A1-611 bp, and E. coli TOP10 was transformed with this plasmid. Purified plasmid was digested with EcoRI yielding either one band (no insert) or two bands, 3.9 kb (plasmid) and 611 bp (insert). Digests were subjected to electrophoresis on 1.5% agarose gel and visualized with the Gel Logic 200 Imaging System (Kodak). Clones that contained the 611 bp insert were sequenced (Supplement 3).

DNA sequencing and sequence analysis
Tue pCR2.1-TOPO-Blunt-hnRNP A1 611 bp clones were subjected directly to automated DNA sequencing (ABI 3130 X L) at the University of Tennessee Health Sciences Center Molecular Resource Center. Electropherograms were obtained and sequence quality was analyzed by Sequence Analysis Software (ABI). Sequence alignment was carried out by Nucleotide BLAST (National Center for Biotechnology Information). Called genomic DNA sequences were compared to mutations (SNVs, SNPs) listed in four different public databases: (1) Exome variant server (ESV): http://evs.gs.washington.edu/EVS/, (2) Catalogue of somatic mutations in cancer (COSMIC): http://cancer.sanger.ac.uk/cancergenome/projects/cosmic/, (3) 1000 genomes; a deep catalog of human genetic variation. http://www.1000genomes.org, (4) NCBI dbSNP: (http://www.ncbi.nlm.nih.gov/snp/).

Cloning and expression of hnRNP-A1
cDNA encoding the entire sequence of hnRNP A1 (WT) was cloned into the expression vector pTriEx™5 Ek/LIC vector (Novagen) and transfected into SK-N-SH cells, a neuroblastoma cell line (ATCC - American Type Culture Collection). The amplified open reading frame (ORF) of hnRNP A1 was subcloned into Bam HI and Hind III sites of modified pGEX-6p-1 vector to create recombinant E. coli expression vectors for glutathione S-transferase (GST) full down assay.

Primers and site-directed mutagenesis
The primers for mutagenesis by PCR were designed basically according to the manufacturer (QuikChange™ II XL Site-Directed Mutagenesis kit; Agilent Technologies, CA). Briefly, each pair of primers contained a primer-primer complementary (overlapping) sequence at the 3'- and 5'-terminus. The forward primers were used for mutagenesis of the target residues F273L, M276L and F281L in hnRNP A1. The primers for each of the variants were: (1) p.F273L - forward: CAG TCT TCA AAT CTT GGA CCC ATG AAG GGA GG, reverse: CCT CCC TTC AGG GGT GTT CCA AAA TTT GAA GAC TG; (2) p.M276L - forward: CAG TCT TCA AAT TTT GGA CCC CTG AAG GGA G, reverse: CCT CCC TTC ATG GGT CCA AGA TTT GAA GAC TG; (3) p.F281L - forward: C ATG AAG GGA GAA AAT CTT GGA GCC AGT GGC AGC TC, reverse: GA GCT TCT GCC TCC AAG ATT TCC TCC CTT CAT G. All variant sites were located in hnRNP A1-M9 and both forward and reverse primers shared the region in question. The melting temperature ($T_m$) was calculated using the formula provided by the manufacturer Agilent Technologies: $T_m = 81.5 + 0.41(\%GC) - 675/N - \%\text{mismatch}$. Here, $N$ is the primer length in bases. All the primers were synthesized by Genelink (Hawthorne, NY). Mutagenic reaction was performed in 50 µl of PCR mix containing 10 ng of pTriEx-5 Ek/LIC-hnRNP A1 (WT) or pGEX-6p-1-hnRNP A1 (WT) as template, 200 nM primer and 2.5 U Pfu DNA polymerase. The PCR temperature profile was: an initial denaturation at 95°C for 1 min, followed by 18 cycles with each at 95°C for 50 sec, 60°C for 50 sec and 68°C for 1 min, and a final extension at 68°C for 7 min. The PCR products of Site-Directed Mutagenesis were transformed into E. coli XL10-Gold competent cells and isolated using Qiagen miniprep kits (Qiagen, Germany).

Transfection
DNA complexes prepared using a DNA (µg) to Lipofectamine® 2000 (µl) ratio of 1:2.5 for SK-N-SH cell line. For hnRNP A1 relocalization experiments, the human hnRNP A1 (WT or variant) cDNA was transfected into SK-N-SH cells (70–80% confluence) using Lipofectamine 2000 (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. After 5 hours incubation, the transfection mixture was removed from each well and replaced with DMEM containing 10% FBS. Fresh medium was conditioned for 24 h before relocalization analysis of hnRNP A1 by immunocytochemistry.

Immunocytochemistry
SK-N-SH Cells (ATCC HTB-11) were grown on poly-L-lysine-coated cover slips and were transfected using Lipofectamine 2000. Cells were then rinsed with PBS, fixed with 4% paraformaldehyde, permeabilized with cold acetone, and blocked in PBS containing 5% BSA. Primary antibodies used were: rabbit anti-TDP-43 (1:1000, Millipore, catalog #ABN271), rabbit anti-active caspase-3 (1:50, Millipore, catalog #AB3623), rabbit anti-Neuron specific beta III tubulin (NTB3) (1:1000, Abcam, catalog #ab18207) and biotinylated mouse anti-strep-Tag II (1:1000, GenScript, catalog #A01737). Secondary antibodies were: Texas Red conjugated goat anti-rabbit IgG (1:300, Vector, catalog #TI-5000 and FITC conjugated streapavidin (1:300, Vector, catalog #SA-5001). Primary antibodies were diluted in blocking solution incubated with each coverslip for overnight at 4°C. Cells were then washed with PBS and incubated in secondary antibody for 1 hr. Cells were then washed with PBS and mounted in Prolong-Gold anti-fade reagent with DAPI (Invitrogen).

GST pull-down assay
SK-N-SH cells were cultured in Dulbecco’s Modified Eagle’s medium (BD Biosciences) supplemented with 10% fetal bovine serum, 100 U/mL penicillin G and 100 µg/mL streptomycin, at 37°C under 5% CO2. Cells were harvested and lysed with CytoBuster™ Protein Extraction Reagent (Millipore), containing inhibitor cocktail, homogenized for a few seconds with a handheld homogenizer and spun at 16,000 × g for 5 minutes. Supernatants were used for GST-pull down assays. Glutathione-Sepharose 4B beads coupled with GST-hnRNP A1 (WT or variant), which includes the Transportin 1-binding domain, were incubated for 1 h at 4°C with 600 µL of the cell lysates in CytoBuster™ Protein Extraction Reagent and protease inhibitors. After washing the beads three times with 600 µL of 10 mM PBS (10 mM Na2HPO4, 140 mM NaCl, 2.7 mM KCl, 1.8 mM KH2PO4, pH 7.4) and 4 protease inhibitors, proteins bound to the beads were analyzed by 8–16% SDS-PAGE followed by immunoblotting with rabbit polyclonal GST antibody (1:1000, Millipore, catalog #06-332), mouse monoclonal Transportin 1 antibody (1:1000, Millipore, catalog #05-1515) and mouse monoclonal TDP-43 antibody (1:1000, Millipore, catalog #MABN45). The immunoreactive bands were visualized using enhanced chemiluminescence.
Figure 1. Functional domains and sequence alignment of hnRNP A1. A: Domain schematic of hnRNP A1. The sequence shown is isoform A (NP_002127). Isoform B contains a 52 amino acid insert following AA 251, resulting in a 372 amino acid protein (not shown). The RNA binding domains (RBD 1 and 2) are contained within the N-terminal half of hnRNP A1. The prion-like domain (PrLD, AA 233-267), M9 (AA 268-305) and the C-terminus (AA 306-320) are shown. M9 (orange) contains the ‘core’ transportin-1 binding domain (TPNO-1, AA 268-289), red) and the MS IgG epitope (AA 293-304, yellow). Some literature indicates that the PrLD and TPNO-1 binding domain may overlap, such that the PrLD region includes AA 233-272 and the TPNO-1 binding domain includes AA 263-289, with the a resulting overlap of AA 263-272 (Supplement 2). The amplicon included DNA from exons 8 and 9 with the intervening intron. The expressed protein included the PrLD that contained ALS-associated mutations (p.D262V (familial), p.N267S (sporadic)), as well as M9 and the C-terminus of hnRNP A1. (bp base pair).

B: Sequence alignment of hnRNP A1-M9. Human sequences are 100% conserved in mammals, except for M. mulatta (rhesus). There is also high sequence conservation between orthologs (mammals, non-mammalian vertebrates (G. gallus - chicken, D. rerio - zebrafish, X. laevis - frog) and invertebrates (D. melanogaster - fruit fly). SNVs resulting in amino acid substitutions in the TPNO-1 core domain are highlighted in red boxes. SNVs resulting in amino acid substitutions in the MS IgG epitope are highlighted in yellow boxes. Colored amino acids are present for clarity identifying conserved sequences through all species. Black lines (gaps) are inserted between residues so that similar or identical amino acids are aligned in each column. (http://www.bioinformatics.org/strap/).

Results
Novel somatic DNA SNVs are contained within the TPNO-1 binding domain and MS IgG epitope of hnRNP A1-M9 in MS patients

We sequenced a 611 bp region of hnRNP A1 genomic DNA inclusive of exons 8 and 9 with intervening introns (NP_002127.1, Chromosome 12q13.1, DNA g.3080-3690, RNA (cDNA) c.752-963, protein AA 252-320) (Figure 1A) isolated from the PBMCs of patients with MS: relapsing remitting MS (RRMS, n=5), secondary progressive (SPMS, n=5) and primary progressive MS (PPMS, n=4) and healthy controls (HC, n=6) (Supplement 1). The expressed sequence included: the C-terminal of the PrLD (AA 252-267), M9 (AA 268-305) and the residual C-terminus of hnRNP A1 (AA 306-320) (Figure 1A). This region also includes the ‘core’ TPNO-1 binding domain (AA 286-299) and the MS IgG epitope (AA 293-304). Some literature indicates that the PrLD and TPNO-1 binding domain may overlap, such that the PrLD region includes AA 233-272 and the TPNO-1 binding domain includes AA 263-289, with the a resulting overlap of AA 263-272 (Supplement 2). A small percentage of clones from each individual contained genomic DNA SNVs, indicative of these being somatic SNVs derived from a small percentage or subset of PBMC. SNVs were compared to those found in four different databases (see Methods and Supplement 2).

Of the six HCs, zero SNVs resulted in an amino acid substitution within the TPNO-1 binding domain, MS IgG epitope or M9 from the 481 clones that were sequenced (Table 1, Supplement 2, Supplement 3). One individual had a likely benign variant which did not result in a change in the associated amino acid (c.900A>G, p.R300R) (Supplement 2), and three others had SNVs in the C-terminal region include DNA from exons 8 and 9 with the intervening intron. The expressed protein included the PrLD that contained ALS-associated mutations (p.D262V (familial), p.N267S (sporadic)), as well as M9 and the C-terminus of hnRNP A1. (bp base pair).
Table 1. hnRNP A1 single nucleotide variants that resulted in an amino acid substitution. HC: Healthy Control. RRMS: Relapsing Remitting Multiple Sclerosis. SPMS: Secondary Progressive Multiple Sclerosis. PPMS: Primary Progressive Multiple Sclerosis. TPNO-1: Transportin 1. PRLD: Prion-like Domain. AA: amino acid. SNV: single nucleotide variant.

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- **PPMS-1**: 793 A>G, 823 C>T, 950 G>A
- **PPMS-2**: 831 G>T
- **PPMS-3**: 835 A>G
- **PPMS-4**: 817 T>C, 850 A>G, 901 A>G, 922 T>G

- **SPMS-1**: 793 A>G, 850 A>G
- **SPMS-2**: 817 T>C, 841 T>C, 853 A>G
- **SPMS-3**: 817 T>C
- **SPMS-4**: 793 A>G, 823 C>T, 950 G>A, 958 T>C

- **HC-1**: 922 T>C
- **HC-2**: 922 T>C
- **HC-3**: 922 T>C
- **HC-4**: 922 T>C
- **HC-5**: 922 T>C
- **HC-6**: 922 T>C

- **DOMAIN**: 261-280
- **DOMAIN**: 261-280
- **DOMAIN**: 261-280
- **DOMAIN**: 261-280
Table 1

Table 1

Figure 2A

Drosophila melanogaster (http://www.ncbi.nlm.nih.gov/snp). These data, in which SNVs in hnRNP A1 are a rare event, are consistent with the finding in sporadic (1 of 305) and familial ALS (1 of 212) patients where most did not have mutations by whole exome sequencing.22 Of the five RRMS patients in which 358 clones were sequenced, one patient had two novel SNVs contained within the TPNO-1 binding domain that resulted in an amino acid substitution (c.826A>G, p.F273L). Although there was a somatic SNV contained within this codon in the COSMIC database (c.789T>G, p.V267D), which also aligned with a somatic SNV resulting in an amino acid substitution in SPMS predominated the five patients (c.884A>G, p.Y295C; c.855A>G, p.R285C) and one within the PrLD (c.775A>G, p.S259G). These SNVs are also novel (Table 1, Supplement 2, Data availability). There was a single SNV (c.963A>G) that did not alter the stop codon amino acid sequence (Supplement 2).

Of the five SPMS patients, one had a novel SNV that resulted in an amino acid substitution in the ‘core’ TPNO-1 binding domain (c.817T>C, p.F273L). Although there was a somatic SNV contained within this codon in the COSMIC database (in patients with cancer), its SNV and amino acid change were different (c.818T>G, p.F273C). A second patient had a novel SNV contained within the PrLD (c.755G>A, p.S252N), which also aligned with a somatic SNV in the COSMIC database (c.755G>T, p.S252I), but yielded different amino acids. A third patient had an SNV within the PrLD (c.787T>C, p.F263L), which also aligned with a somatic SNV contained within this codon in the COSMIC database (c.789T>G, p.F273L). In contrast to HC, RRMS or PPMS, novel SNVs that resulted in an amino acid substitution in SPMS predominated within the MS IgG binding epitope of M9 (c.884A>G, p.Y295C; c.886T>C, p.F296L; c.896C>T, p.P299L; c.898C>T, p.R300S; c.902A>G, p.N301S) (Table 1, Supplement 2, Supplement 3, Data availability). Overall, of the 355 clones that were sequenced from the five SPMS patients, 8 (2.25%) were contained within the PrLD and M9 domains of which 6 were within M9, and 5 were exclusive to the MS IgG epitope (Table 1, Supplement 2, Supplement 3). Like HC and RRMS, there were several SNVs contained in the C-terminal region of hnRNP A1 (c.941A>G, p.Y314C; c.952A>G, p.R318G) (Table 1, Supplement 2, Supplement 3).

In contrast to HC, all four PPMS patients, had novel somatic SNVs that resulted in an amino acid substitution and were contained within the ‘core’ TPNO-1 binding domain of hnRNP A1 as follows: patient 1 (c.823C>T, p.P275S), patient 2 (c.831G>T, p.K277N), patient 3 (c.850A>G, p.R284G), and patient 4 (c.817T>C, p.F273L; c.841T>C, p.F273L; c.853A>G, p.S285G) (Table 1, Supplement 2, Data availability). Patients 1 and 3 each had a novel SNV within the PrLD region (c.793A>G, p.N265D). Thus, of the 317 clones that were sequenced, 28.4% were contained within the M9 or PrLD domains, two-thirds of which were exclusive to the TPNO-1 binding domain (Supplement 3). Other SNVs were contained within the MS IgG binding epitope (c.901A>G, p.N301D) or the C-terminal region (c.922T>G, p.S308P; c.950G>A, p.G317D) (Table 1, Supplement 2, Supplement 3, Data availability). Only the c.817T>C, p.F273L SNV aligned with a somatic SNV within the same codon in the COSMIC database (c.818T>G, p.F273C), but again, both the SNV and amino acid substitution differed.

The TPNO-1 binding domain is highly conserved within mammals and evolutionarily conserved between species (Figure 1B). Specifically, the TPNO-1 binding domain is 100% conserved in mammals, except for five mutations contained only within the rhesus monkey (Macaca mulatta) (only one of which overlapped with the SNVs we discovered (AA100)). Further, amino acid sequences were highly conserved between species, as shown by the orthologs between mammals, bird (Gallus gallus), fish (Danio rerio) and frog (Xenopus laevis) as well as with the fruit fly (Drosophila melanogaster) (Figure 1B). Taken together, these data indicate that variants in this highly conserved domain may have pathological consequences, which might contribute to human disease.

Disease-associated SNVs of the TPNO-1 binding domain of hnRNP A1-M9 result in mis-localization of hnRNP A1 into cytoplasmic stress granules and cellular apoptosis A total of nine novel SNVs that resulted in an amino acid substitution were discovered in MS patients within the ‘core’ TPNO-1 binding domain of hnRNP A1-M9. hnRNP A1 has a number of functions, including the transport of nascent mRNA from the nucleus to the cytoplasm. hnRNP A1 shuttles between the nucleus and cytoplasm and binds TPNO-1, which is required for its nuclear import.6,20,22 At equilibrium, hnRNP A1 is predominantly found in the nucleus.17,24 Considering the disease-associated SNVs are contained within a highly conserved region of hnRNP A1 that plays a critical role in cellular function, we hypothesized that altered hnRNP A1 would change hnRNP A1 localization, TPNO-1 binding and induce cellular damage. To test this hypothesis, we performed a number of experiments. First, we manufactured three different hnRNP A1 mutations (by site-directed mutagenesis) contained within its TPNO-1 binding domain (F273L, M276L and F281L), transfected each mutant into SK-N-SH cells and examined the cells for hnRNP A1 localization relative to transfection of Wild Type (WT) hnRNP A1. As shown in Figure 2A (upper panel), WT hnRNP A1 almost completely localized to the nucleus of SK-N-SH cells. In contrast, mutant forms of hnRNP A1 localized to the cytoplasm of cells (Figure 2A, lower panel). Localization within the cytoplasm was not diffuse, but granular, suggestive of stress granule (SG) formation (Figure 2A, lower panel, arrows). There was also localization within cellular processes (Figure 2A, lower panel, arrowhead). To confirm that mutant hnRNP A1 was present in SGs, we double-labeled SK-N-SH cells that contained the transfected mutant hnRNP A1 with anti-TDP-43 antibodies. As shown in Figure 2B, like hnRNP A1, TDP-43 was localized to nuclei (without transfection). In addition, WT hnRNP A1 and TDP-43 co-localized within the nuclei of SK-N-SH cells. In contrast, mutant hnRNP A1 (F273L, M276L and F281L) co-localized with TDP-43 within the cytoplasm of cells (Figure 2B). Considering recent data indicating binding between hnRNP A1 and TDP-43; co-localization of mutant hnRNP A1 (p.D262V) to TDP-43 containing SGs; the role of TDP-43 in SG formation, and the localization of hnRNP A1 in SGs in stress activated cells, these experiments confirm that mutant hnRNP A1 is contained within TDP-43 positive SGs.19,20,22 Because TPNO-1 is required for hnRNP A1 nucleocytoplasmic transport, we hypothesized that mutant...
hnRNP A1 would alter its binding to TPNO-1. In these experiments, protein lysates purified from SK-N-SH cells were incubated with either WT or mutant GST-tagged-hnRNP A1 bound to Glutathione-Sepharose 4B beads. The resultant eluent was then probed for TPNO-1. As shown in Figure 3A, western blots showed immuno-reactivity for TPNO-1 protein with WT-hnRNP A1, indicative of TPNO-1's binding to hnRNP A1. In contrast, there was significantly reduced binding between mutant forms of hnRNP A1 and TPNO-1 (Figure 3A). These experiments show that mutations in the TPNO-1 binding domain of hnRNP A1-M9 alter TPNO-1 binding to hnRNP A1. To confirm protein binding between hnRNP A1 and TPNO-1, which were visualized by immunocytochemistry (Figure 2B), we probed the identical eluents with an anti-TPNO-1 antibody. Both WT and mutant hnRNP A1 bound TPNO-1 (Figure 3B), indicative of their interaction in both the nuclei and cytoplasmic SGs in the cell line. This is consistent with other reports indicative of an interaction between hnRNP A1 and TPNO-1.

In the in-vitro experiments, SG formation in SK-N-SH cells formed within several hours of transfection. When we waited overnight (approximately 24 hours) the cells containing mutant hnRNP A1 developed apoptotic blebs, which contained hnRNP A1 (Figure 2C, arrows). Apoptosis was confirmed by active caspase-3 staining (Figure 2D). As shown in Figure 2D, SK-N-SH cells transfected with mutant hnRNP A1 showed a cytoplasmic hnRNP A1 distribution, stained positive for active caspase-3 and contained fragment nuclei, confirming apoptosis.

In summary, in contrast to WT hnRNP A1, mutant hnRNP A1 showed markedly reduced binding to its co-receptor TPNO-1, co-localized with TPNO-1 within cytoplasmic SGs of cells and caused apoptosis, indicative of the potential pathogenic nature of these disease-associated SNVs in MS patients.

Discussion
Recent studies indicate that in addition to cancer, somatic variants can cause neurological disease. In this study, we discovered novel somatic genomic DNA SNVs in MS patients. Nine were contained within the 'core' TPNO-1 binding domain of hnRNP A1-M9 (AA268-289). Three additional SNVs (c.793A>G, p.N265D (in two patients); c.787T>C, p.F263L) included amino acids within the PrLD - M9 overlap region (AA263-267), which also bind TPNO-1.
Figure 3. Western blots of hnRNP A1 binding with TPNO-1 and TDP-43. GST-tagged WT or mutant forms of hnRNP A1 were bound to Glutathione Sepharose 4B beads, protein lysates of SK-N-SH cells were applied to columns, and the resulting eluent was probed for either TPNO-1 (A) or TDP-43 (B). A. There was strong binding between WT hnRNP A1 and TPNO-1. In contrast, there was no binding between mutant hnRNP A1 (F273L) and TPNO-1, and markedly reduced binding between mutant hnRNP A1s (M276L, F281L) and TPNO-1. B. There was no change in binding between WT and mutant forms of hnRNP A1 and TDP-43.

These variants were in a region of hnRNP A1 that are adjacent to mutations shown to cause ALS (p.D262V, p.N267A). Interestingly, 8 of these 12 SNV’s that involved hnRNP A1-M9 binding to TPNO-1 occurred in PPMS patients. In addition, two hnRNP A1 SNVs were contained exclusively within the PrLD (c.755G>A, p.S252N; c.775A>G, p.S259R). There were also six novel SNVs that resulted in an amino acid substitution within the MS IgG epitope of M9 (AA198-208), five of which segregated to patients with SPMS. Finally, there were nine SNV’s in the C-terminal of hnRNP A1 (AA208-260), occurring with similar frequency in HCs and MS patients. The overall somatic SNV rate (based on the number of clones sequenced) for the M9 target sequence was: PPMS - 2.21%, SPMS - 1.69%, RRMS - 0.56%, HC - 0%. If one includes the PrLD (a domain shown to be critical to hnRNP A1 function), the rates increase in PPMS, SPMS and RRMS to 2.84%, 2.25% and 0.84% respectively. None were identical to somatic mutations in the COSMIC database (n = 981,720 samples, n = 1,292,597 unique variants). We utilized a PCR - cloning technique that has been fine-tuned for more than a decade and shows a mutation rate of approximately 0.1% in more than 46,000 clones that were examined. The rates in progressive MS patients exceed this error rate by more than a log. In addition, under identical conditions, there were no mutations in the M9 target sequence or the PrLD domain in the HCs we examined. Thus, these results are unlikely to be due to PCR errors. Importantly, there was little or no overlap with either SNVs or SNPs reported in four different databases.

hnRNP A1 was one of the first RNA binding proteins shown to shuttle into and out of the nucleus. Nucleocytoplasmic transport is dependent upon binding between the M9 domain (AA208-260) of hnRNP A1 to TPNO-1, in order for this complex to pass through the nuclear pore. M9 acts as both an NES and NLS. M9 is a bipartite domain, a basic residue (522R) and a C-terminal RX(3,5)PY motif, each connected by ‘linker’ residues6,27,43. Each epitope, as well as individual amino acids within an epitope, conveys varying degrees of binding activity to TPNO-126,14. For example, mutant P275A dramatically inhibits nucleocytoplasmic transport and substitution of 273-FGPM-276 with 273-AAAAA-276, completely abolishes TPNO-1 binding and nucleocytoplasmic transport12,43,44. Our data closely align with these findings. For instance, genomic SNVs in MS patients occurred at F273L, F275S, and M276L - all contained within epitope 1. Experimentally, we showed that transfection of F273L and M276L mutants caused mis-localization of hnRNP A1, S formation (co-localizing with TDP-43), cellular apoptosis and diminution of TPNO-1 binding. Mutant F281L caused similar results. Interestingly, F273, F281 and R284 all have two or more side chains that bind TPNO-1, thus are critical contact points between M9 and TPNO-1. MS patients also had an SNV at R284G. Although we did not test this variant, a parallel substitution in the RNA binding protein fused in sarcoma (FUS) (R522G) (Table 2) caused a five-fold decrease in TPNO-1 binding and cytoplasmic mis-localization of FUS45. Like hnRNP A1, FUS is an hnRNP, which at equilibrium localizes to the nucleus and contains a bipartite PY-NLS that binds TPNO-1. Interestingly, mutations in FUS have been shown to cause ALS and frontotemporal lobe dementia46.

Importantly, none of the SNVs contained within hnRNP A1 - M9 or the PrLD has been reported previously. For decades, the only certain genetic risk factor for MS was with MHC Class II HLA-DRB1*. Genome Wide Association Studies (GWAS) have uncovered novel genetic associations with MS12,45 including with the interleukin-2 receptor-α and interleukin-7 receptor genes45. Subsequent studies using several thousand MS cases and controls, which analyzed hundreds of thousands of autosomal SNPs, confirmed the association of MS with major MHC Class II HLA-DRB1 (DRB1 *15:01, *15:03, *13:03) and the protective effect MHC Class I HLA-A*02,6,13. Additional studies showed a total of 48 new and 49 known non-MHC SNPs associated with MS4. Interestingly, the functions of the vast majority of the SNP’s were related to CD4+ T-lymphocyte and immune regulation4. This is important, considering the role that T-lymphocytes and the immune response play in the pathogenesis of MS. A few were potentially associated with neurodegeneration46. Further, >95% of the SNPs were intronic or intergenic, with only a few SNPs involving exons, in contrast to the somatic SNVs discovered here12. In addition to GWAS, whole exome sequencing (WES) is being used to examine differential gene expression in MS patients. In contrast to GWAS, which detects known SNPs and utilizes statistical analyses designed to reveal common variants, WES is designed to discover novel, rare pathologic variants1, One
Table 2. Binding epitopes of the PY-NLS's of hnRNP A1 and fused in sarcoma (FUS). Aligned PY-NLS sequences of hnRNP A1-M9 and FUS-NLS are shown. The three binding epitopes of the PY-NLS are shaded yellow (epitope 1 - hydrophobic), blue (epitope 2 - basic) and red (epitope 3 - C-terminal)

| hnRNP A1 AA sequence # | 262 | 263 | 264 | 265 | 266 | 267 | 268 | 269 | 270 | 271 | 272 | 273 | 274 | 275 | 276 | 277 | 278 | 279 | 280 | 281 | 282 | 283 | 284 | 285 | 286 | 287 | 288 | 289 |
|-------------------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| hnRNP A1 AA SNV (MS)    | L   | D   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| FUS AA sequence         | G   | P   | G   | K   | M   | D   | S   | R   | G   | E   | H   | R   | Q   | D   | R   | R   | E   | R   | P   | Y   |     |     |     |     |     |     |     |
| FUS AA mutation (ALS)   | D   | E/R | P   | G/S^{17,41} | C^* | V^* | P/Q | G/K | G/H/L | G | S/T/W | L |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| FUS AA sequence #       | 507 | 508 | 509 | 510 | 511 | 512 | 513 | 514 | 515 | 516 | 517 | 518 | 519 | 520 | 521 | 522 | (-) | 523 | 524 | 525 | 526 |     |     |     |     |     |     |     |

SNVs resulting in an amino acid substitution (hnRNP A1) and missense mutations (FUS) are shown by gray shading of the amino acid sequence number. Amino acid SNVs associated with multiple sclerosis (MS) in hnRNP A1 are shown above the primary sequence and mutations associated with amyotrophic lateral sclerosis (ALS) in FUS are shown below the primary sequence. Two SNVs (*) in hnRNP A1-M9 align with mutations in the FUS NLS: hnRNP A1-M9 AA^{275} with FUS NLS AA^{510} and hnRNP A1-M9 AA^{284} with FUS NLS AA^{522}. Stop codons, frame shift mutations, insertions and deletions not included. Notes: R514G/S, R514S, G515C, R514S, E516V.

Notes:
1. R514G/S
2. R514S, G515C
3. R514S, E516V

of the genes identified by GWAS was CYP27B1, which encodes an enzyme of the same name that converts 25-hydroxyvitamin D to 1,25 dihydroxyvitamin D, the biologically active form of vitamin D. A single individual from 43 MS families was found to have a rare p.R389H genetic variant in CYP27B1, which resulted in complete loss of enzyme activity. However, unaffected relatives of the individual also carried the identical variant, which has a high frequency in the general population. Thus, it is clear that an individual’s genetic background makes an important contribution to the pathogenesis of MS. This supports the tripartite hypothesis that an environmental trigger in a genetically susceptible individual causes an autoimmune response to CNS antigens that result in the pathology observed in the brain and spinal cord of MS patients. Potential environmental triggers include viral infection, low vitamin D levels and sun exposure. During the relapsing phase of MS, Th1 and Th17 CD4+ T-lymphocyte responses appear to predominate and correlate with focal MS plaque formation in the CNS. As MS evolves into a secondary progressive phase, CNS damage becomes more diffuse. Immune cells also become more diffuse and IG containing plasma cells, B-lymphocytes and macrophage/microglia response predominates. Many of these latter features are also characteristic of primary progressive MS. MS is also characterized by increased oxidative stress (in PBMC and brain), which can cause DNA damage and somatic mutations. Further, T-lymphocytes isolated from MS patients that contain somatic mutations have been shown to react to myelin peptides. Specifically, T-cell clones mutant for the hypoxanthine guanine phosphoribosyltransferase (HPRT) gene were found only in MS patients and not in HCs. The MS patients studied had ‘chronic progressive MS’, clinically similar to the PPMS patients in this study. The T-cell clones recognized and proliferated to myelin basic protein peptides. These data suggest that T-cells that undergo clonal expansion are more prone to somatic mutations and that clones containing somatic mutations contribute to the pathogenesis of MS. How might variants in hnRNP A1 contribute to the pathogenesis of MS? Although direct evidence is not yet available and requires further study, the molecular consequences of abnormal forms of hnRNP A1 on cellular function may have profound effects on the immune system. For example, as it relates to the environment, hnRNP A1 regulates the synthesis of several viruses including human immunodeficiency virus, HTLV-1, and human rhinovirus. Immunologically, hnRNP A1’s nucleocytoplasmic shuttling and RNA binding specificity is required for myelopoiesis and modulation of immune-mediated programmed cell death. Further, apoptotic blebs (which we showed in this study to contain hnRNP A1) have profound effects on the immune response, as they are believed to initiate and perpetuate autoimmune diseases such as systemic lupus erythematosus.

In summary, we discovered novel SNVs in MS patients. The SNVs involve the M9 nucleocytoplasmic binding domain of hnRNP A1, which when transfected into a cell line, resulted mis-localization of hnRNP A1 to cytoplasmic stress granules and cellular apoptosis. Future studies are required to replicate this data, expand it to include a broader spectrum of genes, a greater number of MS patients and patients with other chronic inflammatory diseases such as rheumatoid arthritis. In addition, we also plan to perform functional studies of the SNVs in immune and nervous system cells of MS patients.

### Data availability

Single nucleotide variants (SNVs) (somatic) in MS patients submitted to dbSNP.

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Supplementary materials

Supplement 1: Patient demographics (xlsx)
Supplement 2: Single nucleotide variants in the target sequence of hnRNP A1 (xlsx)
Supplement 3: Number of clones containing single nucleotide variants (xlsx)

References


Grant information

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Version 2

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Hans Lassmann
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I have no further comments. The authors have addressed my points and have pointed out their view on this issue and that some of the points have to be addressed in future studies.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 10 September 2014

https://doi.org/10.5256/f1000research.4747.r5807

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Elliot M. Frohman
Department of Neurology, University of Texas Southwestern Medical Center at Dallas, Dallas, TX, USA

My distinguished colleagues from the University of Tennessee in Memphis have performed an innovative set of complex studies seeking to elucidate somatic mutation variants in patients with MS; across the various clinical subtypes, and compared to normal subjects. Quite surprising to this reviewer, the authors identified a TPNO-1 binding domain change of hnRNPA1-M9; an RNA
binding protein that is among the first to be shuttled into and out of the nucleus. Perhaps even more conspicuous is that these somatic modifications are directly adjacent to well characterized mutations that are associated with the development of ALS (i.e. accelerated motor neuron disease of the upper and lower motor neuron compartments).

My colleagues further show that 8/12 SNVs are shown to occur in primary progressive MS (a slowly and insidiously progressive loss of the upper motor neuron compartment of the CNS. They further elucidated six novel SNVs; with substitutions within the MS IgG epitope of M9 (AA 293-304).

The authors proceed to provide a SNV rate stratification across normal subjects vs clinical subtypes of MS. Here they found the following corresponding rates:

PPMS- 2.21%
SPMS- 1.69%
RRMS- 0.56%
Normal human subjects = 0%

Much has been achieved with the application of genome wide association studies (GWAS), such as the identification of a linkage between the IL-2 and IL-7 receptor genes; immune regulatory elements that dovetail nicely with the higher risk of MS developing in subjects who are also positive for the MHC Class II HLADR21501B1 (2 copies > 1 copy with respect to relative risk). Alternately, MHC Class I HLA-A*02 confers a protective influence. Indeed, it is intriguing that most of the SNPs are associated with CD4+T cell regulation.

Contemporary screening techniques have further evolved with even greater refinement in genetic element identification conferring risk of MS by virtue of the utilization of the powerful technique of whole exome sequencing (WES), that provides important perspective on differential gene expression mechanisms.

Finally, the GWAS discovery associated with MS of the CYP27B1 gene, has particular relevance to an epigenetic disorder such as MS, given that this gene encodes for the critical enzyme that serves to transform the 25-OH-D form to the 1,25-OH-D active form of vitamin D.

With respect to the pathophysiologic relevance of the hnRNPA1 variants, these appear to alter neuronal function, and ultimately culminate in neurodegeneration, in conjunction with serving a role in the important regulation of viral synthesis (another highly conspicuous epigenetic factor with special relevance to MS and related disorders). As the authors rightly point out, to date the somatic mutations in question and under investigation have only been elucidated from PBMCs of MS patients.

This work, if confirmed, may have special and critical relevance to the highly enigmatic phenomenology of what determine the transition from a relapsing to a progressive form of MS (or a progressive form from the inception without an antecedent relapsing and adaptive immune phase of the disorder), and such studies might shed light on the pathobiological underpinnings of axonal and neuronal degeneration; the histopathological signatures of what we currently believe to constitute the signature of irreversible (at least so far) neurologic disability with corresponding loss of functional capabilities.
This work, contributes to our deeper understanding of the potential mechanisms of disease that figure prominently in influencing progression and neurodegeneration for MS in particular, but may also shed light on those mechanisms that underly neurodegenerative disorders in general.

The authors are to be congratulated for this fine contribution to the literature.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

**Author Response 11 Sep 2014**

**Michael Levin**, Veterans Affairs Medical Center, Memphis, TN, USA

We appreciate Professor Frohman's insightful commentary about our work. We concur that to date, we have only found somatic nucleotide variants (SNVs) in PBMCs of MS patients. Further studies are warranted about the contribution of SNVs to axonal and neuronal degeneration in the CNS of MS patients. This is addressed in the last two paragraphs of the discussion.

**Competing Interests:** No competing interests.

**Reviewer Report 23 July 2014**

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Michael K. Racke

Department of Neurology and Multiple Sclerosis Center, The Ohio State University, Columbus, OH, USA

This manuscript by Lee and Levin continues work by Levin on the role of an immune response to hnRNP-A1 in human demyelinating disorders, including multiple sclerosis. Previously, Levin had shown that an antibody response to HTLV-I Tax cross-reacted with hnRNP-A1. Now, they go on to show that there are mutations in hnRNP-A1 that affect the localization of this protein and how it could affect neuronal survival in multiple sclerosis. In general, the experiments appear to be well performed. One might be interested to know whether testing in another cell line such as a oligodendroglial cell line would have similar effects on cell survival as in the neuroblastoma cell line. This could also be highly relevant in MS, as in addition to neuronal loss, there is demyelination and loss of oligodendrocytes. This work attempts to address whether mutations in
hnRNP-A1 could contribute to MS pathogenesis. It would be interesting to note whether antibody responses to the protein correlated with mutations (i.e. does the mutation in the protein affect tolerance). Another issue that needs to be addressed is whether the mutations in the hnRNP-A1 gene are due to increased mutational frequency that can be observed in replicating cells, similar to the observation of increased hprt mutations made by Allegretta many years ago in MBP-specific T cells. Overall, this is an interesting study which certainly increases the interest in non-myelin targets in diseases such as MS.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 11 Sep 2014

Michael Levin, Veterans Affairs Medical Center, Memphis, TN, USA

We appreciate Professor Racke's critique, including that the studies were “well performed.” The following concerns expressed by Professor Racke were addressed:

1. **One might be interested to know whether testing in another cell line such as a oligodendroglial cell line would have similar effects on cell survival as in the neuroblastoma cell line. This could also be highly relevant in MS, as in addition to neuronal loss, there is demyelination and loss of oligodendrocytes.**

   We used SK-N-SH neurons as a model system to examine the effect that mutant forms of hnRNP A1 might have on target cell function. Considering that cells other than neurons are clearly involved in the pathogenesis of MS, we plan to add oligodendrocyte cell lines to future studies of mutant hnRNP A1. We addressed this in the sixth paragraph of the discussion.

2. **This work attempts to address whether mutations in hnRNP-A1 could contribute to MS pathogenesis. It would be interesting to note whether antibody responses to the protein correlated with mutations (i.e. does the mutation in the protein affect tolerance).**

   We agree that this would be in important and insightful experiment, one that we think is beyond the scope of the current study. Our previous work showed that MS patients develop antibodies to hnRNP A1 using Western blots. We have not yet developed the technology that would allow us to quantitate the amount of antibodies to hnRNP A1 in individual patients. We will consider this in the future as we examine the potential connection between antibodies to and somatic mutations within hnRNP A1.

3. **Another issue that needs to be addressed is whether the mutations in the hnRNP-A1 gene are due to increased mutational frequency that can be observed in replicating cells, similar to the observation of increased hprt mutations made**
by Allegretta many years ago in MBP-specific T cells.

We appreciate this insight offered by Professor Racke, which is very relevant to our work and have included it in paragraph 4 of the discussion.

**Competing Interests:** No competing interests.
disease. The key message from the authors that this is an important finding for MS pathogenesis is not substantiated by the data. I agree with the authors that further experiments (possibly beyond the scope of a new version of this article) are needed. For example:

1. To look for somatic mutations for other genes, to see whether the paper just shows that in PBMCs of MS patients there is an increased rate of somatic mutations in many different genes or whether the mutations in the specific gene are MS specific.

2. To expand the sample of control patients, in particular by including patients with other chronic inflammatory diseases (e.g. patients with chronic rheumatoid arthritis).

3. To search for similar mutations in neurons and glia of the MS brain to establish a link to neurodegeneration in this disease.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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Author Response 11 Sep 2014

**Michael Levin**, Veterans Affairs Medical Center, Memphis, TN, USA

We appreciate Professor Lassmann's comments including that the study was “well performed” and the acknowledgement that “increased incidence of somatic mutations in PBMCs in MS presented in this article is a first...”. We agree with his constructive critiques, which we have addressed below.

1. **To look for somatic mutations for other genes, to see whether the paper just shows that in PBMCs of MS patients there is an increased rate of somatic mutations in many different genes or whether the mutations in the specific gene are MS specific.**

Our ongoing studies are designed to evaluate more genes. Using next-generation sequencing, which is beyond the scope of this paper, we may be able to assess a broader number of genes, potentially including coding exomes of the human genome. In this study, we evaluated three different regions of hnRNP A1: the C-terminal of PrLD, M9 and hnRNP A1’s C-terminus. Within this subset of genes, we saw a segregation of SNVs. For example, there were similar numbers of SNVs in hnRNP A1’s C-terminus in all groups examined (Table 1, Supplement 2). In contrast, in PPMS SNVs segregated to the TPNO-1 binding domain of M9 and SPMS to the MS IgG epitope of M9. Although more genes are required to assess their role in MS compared to other diseases, in this study, we have already observed a differential expression of SNVs between types of MS and HCs. This is addressed in the last paragraph of the discussion.

2. **To expand the sample of control patients, in particular by including patients**
with other chronic inflammatory diseases (e.g. patients with chronic rheumatoid arthritis).

We will add patients with other chronic inflammatory diseases such as rheumatoid arthritis to our future studies. Fortunately, there is a large rheumatoid arthritis research group at our institution, which allow access to these samples. We addressed this in the last paragraph of the discussion.

3. **To search for similar mutations in neurons and glia of the MS brain to establish a link to neurodegeneration in this disease.**

Our ongoing studies will apply the techniques used in this manuscript in central nervous system tissues isolated from MS patients at autopsy. This is addressed in paragraph 6 of the discussion. As we complete these studies, as well as studies examining the role of SNVs on the immune response in MS patients, this will address Professor Lassmann's concern about these findings being relevant to the pathogenesis of MS.

4. **Professor Lassmann also suggested that oxidative stress might be a cause of SNVs in the PBMCs of MS patients.**

We added this insight with references to the fourth paragraph of the discussion: “MS is also characterized by increased oxidative stress (in PBMC and brain), which can cause DNA damage and somatic mutations.”

**Competing Interests:** No competing interests.

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**Comments on this article**

**Version 2**

**Author Response 10 Jan 2017**

**Michael Levin**, Veterans Affairs Medical Center, Memphis, TN, USA

We already acknowledged the possibility of cloned PCR products causing artifactual results in the manuscript (page 9). There are several reasons why we do not think this had significant effects on the results and conclusions of these experiments. The technical nature of PCR errors makes them random. If the data were the result of PCR errors, the DNA mutations would be equally distributed (both in location and frequency) throughout the four groups of patients we examined (PPMS, SPMS, RRMS, HCs).

In terms of location, we found a similar mutation rate between the four groups in the C-terminal
region of hnRNP A1 (Table 1, Supplements 2 and 3). These could be true mutations or PCR errors. In contrast, there were distinct differences between the four groups in the M9 region of hnRNP A1. In M9, the frequencies of mutations (from over 1500 individual clones that were sequenced) were PPMS 2.21%, SPMS 1.69%, RRMS 0.56%, HC 0.00%. The differences in these values were highly statistically significant when comparing either PPMS or SPMS (individually or in combination) compared to HCs (p-value 0.0013 – 0.0078, see tables). This was true when using the percentage of mutations relative to the number of clones or base pairs within M9 in each group. This further supports our hypothesis that somatic mutations are important in progressive forms of MS.

Thus, the somatic mutations we discovered were different in both location and frequency within hnRNP A1. Taken together, these data are inconsistent with PCR artifacts. Consistent with our observation, other groups later confirmed the presence of somatic mutations in the lymphocytes of MS patients (Clinical Immunology, doi:10.1016/j.clim.2016.11.018), indicative of their potential importance in the pathogenesis of MS.

I. MUTATION RATE COMPARISONS IN MS PATIENTS BASED ON CLONES

<table>
<thead>
<tr>
<th>Group</th>
<th>Mutation Rate</th>
<th>Lower Bound*</th>
<th>Upper Bound*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPMS</td>
<td>7/317=2.21%</td>
<td>0.89%</td>
<td>4.5%</td>
</tr>
<tr>
<td>SPMS</td>
<td>6/355=1.69%</td>
<td>0.62%</td>
<td>3.64%</td>
</tr>
<tr>
<td>RRMS</td>
<td>2/358=0.56%</td>
<td>0.07%</td>
<td>2.00%</td>
</tr>
<tr>
<td>HC</td>
<td>0/481=0.00%</td>
<td>0.00%</td>
<td>0.076%</td>
</tr>
</tbody>
</table>

*Clopper-Pearson (Exact) 95% Confidence Intervals

Statistical Comparisons: Fisher's Exact Test P-value
PPMS vs. HC          0.0015
SPMS vs. HC          0.0057
RRMS vs. HC          0.18
PPMS + SPMS vs. RRMS 0.17
PPMS + SPMS vs. HC   0.0013

II. MUTATION RATE COMPARISONS IN MS PATIENTS BASED ON BASE PAIRS

<table>
<thead>
<tr>
<th>Group</th>
<th>Mutation Rate</th>
<th>Lower Bound*</th>
<th>Upper Bound*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPMS</td>
<td>7/36138=0.019%</td>
<td>0.008%</td>
<td>0.041%</td>
</tr>
<tr>
<td>SPMS</td>
<td>6/40470=0.015%</td>
<td>0.006%</td>
<td>0.032%</td>
</tr>
<tr>
<td>RRMS</td>
<td>2/40812=0.005%</td>
<td>0.000%</td>
<td>0.018%</td>
</tr>
<tr>
<td>HC</td>
<td>0/54834=0.000%</td>
<td>0.000%</td>
<td>0.0007%</td>
</tr>
</tbody>
</table>

*Clopper-Pearson (Exact) 95% Confidence Intervals

Statistical Comparisons: Fisher's Exact Test P-value
PPMS vs. HC          0.0016
SPMS vs. HC          0.0059
RRMS vs. HC          0.18
PPMS + SPMS vs. RRMS 0.19
PPMS + SPMS vs. HC   0.0078

Sangmin Lee and Michael C. Levin
Competing Interests: No conflicts

Reader Comment 02 Aug 2016
Pentti Tienari, University of Helsinki and Helsinki University Hospital, Finland

This study reports the discovery of somatic (acquired) mutations in the hnRNPA1 gene in multiple sclerosis patients. Unfortunately, the method for detecting somatic mutations is not valid. The authors discovered the mutations using a method that is artefact prone. They first created PCR-product of the target region, then cloned these PCR products into a plasmid vector and then sequenced bacterial colonies carrying these vectors. It is a well-known problem to all, who have done PCR-cloning that artefact mutations, created by PCR amplification, are commonly encountered. There was no validation of the mutation by another method. Therefore, there is no convincing evidence for the existence of somatic mutations in the multiple sclerosis patients in the hnRNPA1 gene. These mutations are most likely PCR-artefacts.

Competing Interests: No competing interests.