Overexpression of the ubiquitin-editing enzyme A20 in the brain lesions of Multiple Sclerosis patients: moving from systemic to central nervous system inflammation

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INTRODUCTION

Multiple Sclerosis (MS) is one of the most common chronic-inflammatory, immune-related disease of the central nervous system (CNS), characterized by demyelination, glia alterations, and neuro-axonal damage (16, 33). In this complex disorder, there is an interplay between inflammatory and neurodegenerative processes, leading to intermittent neurological disturbance followed by progressive accumulation of sensory-motor dysfunctions and cognitive disability.

Focal inflammatory demyelinating lesions in the white matter (WM), typically called “plaques,” have been historically considered the neuropathological hallmark of MS (33, 34). In the last years, pathological and MRI studies have revealed that lesions are also located and widely extended in the grey matter (GM), especially in the cerebral cortex (9, 11, 17).

Although the etiology of MS remains poorly understood, its complex multifactorial nature involving genetic susceptibility, epigenetic and environmental factors is widely known. Among the genomic regions associated to MS in genome-wide association studies (GWAS) (3) there is the locus containing the TNF associated protein 3 (TNFAIP3) gene, which encodes for the ubiquitin-editing enzyme A20.

Abstract

Multiple Sclerosis (MS) is a chronic demyelinating disease of the central nervous system (CNS) in which inflammation plays a key pathological role. Recent evidences showed that systemic inflammation induces increasing cell infiltration within meninges and perivascular spaces in the brain parenchyma, triggering resident microglial and astrocytic activation. The anti-inflammatory enzyme A20, also named TNF associated protein 3 (TNFAIP3), is considered a central gatekeeper in inflammation and peripheral immune system regulation through the inhibition of NF-κB. The TNFAIP3 locus is genetically associated to MS and its transcripts is downregulated in blood cells in treatment-naïve MS patients. Recently, several evidences in mouse models have led to hypothesize a function of A20 also in the CNS. Thus, here we aimed to unveil a possible contribution of A20 to the CNS human MS pathology. By immunohistochemistry/immunofluorescence and biomolecular techniques on post-mortem brain tissue blocks obtained from control cases (CC) and progressive MS cases, we demonstrated that A20 is present in CC brain tissues in both white matter (WM) regions, mainly in few parenchymal astrocytes, and in grey matter (GM) areas, in some neuronal populations. Conversely, in MS brain tissues, we observed increased expression of A20 by perivascular infiltrating macrophages, resident-activated astrocytes, and microglia in all the active and chronic active WM lesions. A20 was highly expressed also in the majority of active cortical lesions compared to the neighboring areas of normal-appearing grey matter (NAGM) and control GM, particularly by activated astrocytes. We demonstrated increased A20 expression in the active MS plaques, particularly in macrophages and resident astrocytes, suggesting a key role of this molecule in chronic inflammation.
A20 is an inducible cytoplasmic protein, whose functions make it a central gatekeeper in inflammation and immunity (14). Above all, A20 is considered the most potent negative regulator of the pro-inflammatory transcription factor NF-kB. A20 has proved to be necessary for the homeostasis of immune cells such as B cells, T cells, dendritic cells, and macrophages (64). Recently, new functions emerged, including the control of necroptosis and inflammasome activity in immune cells (15).

An A20 dysfunction has been involved in the pathogenesis of several autoimmune and inflammatory disorders, including MS. An expanding list of common coding and noncoding single-nucleotide polymorphism (SNPs) variants in the TNFAIP3 locus has been associated with an increased susceptibility to inflammatory disorders such as systemic lupus erythematosus (SLE), rheumatoid arthritis (RA), psoriasis, type 1 diabetes, coeliac disease, Crohn’s disease, and systemic sclerosis (6, 10, 32, 38, 48).

A decreased level of A20 transcript has been found in blood cells of patients with MS, RA, or SLE compared to healthy controls (19, 20, 35, 49, 52, 53), and recent evidences describe an altered A20 expression also in immune cells of patients with neurodegenerative disorders such as Parkinson’s disease (53). In mice, a specific A20 deletion in immune cells triggers inflammatory phenotypes mimicking those seen in the genetically associated human disorders (12, 24, 26, 31, 42, 60). Consistently, in humans the “Haploinsufficiency of A20” (HA20), a monogenic disease characterized by rare familial A20 mutations and a reduced A20 protein expression, leads to generalized inflammation in patients (59).

Besides the role in immune cells, NF-kB pathway and its inhibitor A20 have a well-known role in non-hematogenous organ-resident cells, particularly in the CNS, in both physiological and pathological conditions. Neurons exhibit a high constitutive activity of NF-kB, which is involved in learning and memory and in the regulation of neurogenesis, neuritogenesis, and synaptogenesis (43, 44, 50). In contrast, in astrocytes and in microglia the NF-kB activity is promptly induced by pro-inflammatory molecules, accompanied by the production of a specific pattern of cytokines and chemokines and reactive nitrogen and oxygen species (58). In fact, it appears that the detrimental action of NF-kB in CNS disorders takes place mostly in glial cells and not in neurons (2, 27, 29, 37, 41, 57). A chronic activation of NF-kB pathway in infiltrating and resident glia-like cells in white matter (55). Furthermore, a recent molecular and neuropathological study of cortical lesions in post-mortem MS brain tissues has suggested that the overexpression of the A20 gene, together with HSP70, is related to the upregulation of the TNF/TNFR1-stimulated necrotic signaling in conditions of increased meningeal inflammation and severe and rapid disease progression (39), which could possibly suggest its role in the inhibition of survival/repair pathways. However, A20 protein expression has never been detected in brain tissues. On this basis, we aimed to perform a neuropathological characterization of A20 protein and gene expression in human healthy and pathological brain tissues, focusing on MS disease.

**MATERIALS AND METHODS**

**Human post-mortem snap-frozen brain tissues**

For this study, 38 brain tissue blocks obtained from five control cases (CC) with no neurological diseases and 11 secondary progressive (SP) and six primary progressive (PP) MS cases were used. Demographic and clinical features are reported in Table 1. Post-mortem brains were cut into blocks (2x2x1cm) and unfixed blocks were immediately frozen by immersing in isopentane, pre-cooled on a bed of dry ice and stored at −75°C. Human post-mortem brain tissues were obtained from the United Kingdom (UK) MS Tissue Bank at Imperial College via a UK prospective donor scheme with full ethical approval (08/MRE09/31). Neuropathological confirmation of the diagnosis of MS was carried out according to the International Classification of Diseases of the Nervous System criteria (www.ICDNS.org), and confounding pathologies were excluded, including Alzheimer-like changes (56).

The project has been approved by the University Bioethics Committee of the University of Turin on 28 February 2015.

**Neuropathological characterization of human post-mortem snap-frozen brain tissue by immunohistochemistry**

Air-dried 10-µm-thick cryosections cut from snap-frozen tissue blocks were characterized by immunohistochemistry (IHC) with specific antibodies. The complete list of used antibody, the working dilution and the postfixation method are reported in Table 2. Immunostaining for myelin-oligodendrocyte glycoprotein (MOG) was performed to discriminate presence of white matter (WM) and grey matter (GM) and for the occurrence and extension of demyelinated lesions in the spinal cord, due to an increased function of NF-kB (62). A20 deficiency in microglia determines an increase of the microglial cell number and exacerbates the EAE course, due to the uncontrolled activation of Nlrp3 inflammasome and the subsequent increased IL-1b secretion and CNS inflammation (45, 61).

Pranski et colleague observed the expression of A20 transcripts and proteins belonging to A20 ubiquitin-editing complex in several area and cell types of control cases post-mortem brain tissues, such as neurons in cerebral cortex and glia-like cells in white matter (55). Furthermore, a recent molecular and neuropathological study of cortical lesions in post-mortem MS brain tissues has suggested that the overexpression of the A20 gene, together with HSP70, is related to the upregulation of the TNF/TNFR1-stimulated necrotic signaling in conditions of increased meningeal inflammation and severe and rapid disease progression (39), which could possibly suggest its role in the inhibition of survival/repair pathways. However, A20 protein expression has never been detected in brain tissues. On this basis, we aimed to perform a neuropathological characterization of A20 protein and gene expression in human healthy and pathological brain tissues, focusing on MS disease.
in WM (WML) and GM (GML). Immunostaining for major histocompatibility complex class II (MHC II) molecules was performed to assess the stage of lesion activity. The polyclonal anti-A20 antibody was used to detect A20.

Primary antibodies indicated in Table 2 were diluted in PBS containing 0.1% of Triton X-100 and 10% of normal serum and incubated overnight at 4°C. Next, sections were incubated with the appropriate biotinylated secondary antibodies (Jackson ImmunoResearch Laboratories, West Grove, PA), followed by avidin-biotin peroxidase complex (ABC; Vector Laboratories Burlingame, CA). The immune reactions were visualized with 3′3-diaminobenzidine (DAB, Sigma, St. Louis, MO). Hematoxylin was used to counter-stain nuclei in the slices. All the sections were sealed with DPX (Sigma, St. Louis, MO).

Control blocks contained normal WM and GM areas, while MS blocks included normal-appearing WM (NA WM), normal-appearing GM (NAGM), and lesions. Pre-active (pre-AL), active (AL), and chronic active (CAL) WM lesions (WML) and active (aGML) or inactive (iGML) GM lesions (GML) were identified based on the anti-MHC II immunostaining according to Reynolds and colleagues (56).

The quantification of the A20 immunoreactivity in the lesions and in the normal-appearing areas was performed...

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**Table 1.** Demographic and clinical data of examined MS patients and CC, provided by the UK MS Tissue Bank. Abbreviations: F = female; M = male; CC = control case; MS = multiple sclerosis; SPMS = secondary progressive multiple sclerosis; PPMS = primary progressive multiple sclerosis.

| Case | Gender | Diagnosis | Disease length (years) | Age of death | Cause of death | Post-mortem delay (hours) |
|------|--------|-----------|------------------------|--------------|----------------|--------------------------|
| C08  | F      | CC        | -                      | 93           | Broncho-pneumonia, cerebrovascular accident | 9 |
| C14  | M      | CC        | -                      | 64           | Cardiac failure | 18 |
| C15  | M      | CC        | -                      | 82           | -              | 21 |
| C22  | F      | CC        | -                      | 69           | Lung cancer | 33 |
| C25  | M      | CC        | -                      | 35           | Carcinoma of tongue | 22 |
| MS104| M      | SPMS      | 11                     | 53           | Advanced MS, urinary tract infection | 12 |
| MS122| M      | SPMS      | 16                     | 44           | Broncho-pneumonia | 16 |
| MS154| F      | SPMS      | 10                     | 34           | Pneumonia | 12 |
| MS163| F      | SPMS      | 6                      | 45           | Multi-organ failure sepsis due to urinary tract infection, MS | 28 |
| MS166| F      | SPMS      | 36                     | 62           | Broncho-pneumonia, MS | 7 |
| MS187| F      | SPMS      | 30                     | 57           | Left ventricular failure, myocardial infarct, ischemic heart disease, sepsis, ms | 13 |
| MS197| F      | SPMS      | 27                     | 51           | Broncho-pneumonia, MS | 10 |
| MS200| F      | SPMS      | 19                     | 44           | Urinary tract infection, sepsis, MS | 20 |
| MS207| F      | SPMS      | 25                     | 46           | Pneumonia, MS | 10 |
| MS230| F      | SPMS      | 20                     | 42           | MS | 31 |
| MS245| M      | PPMS      | 26                     | 64           | Broncho-pneumonia, MS | 25 |
| MS248| F      | PPMS      | 16                     | 58           | C.O.P.D., multiple sclerosis | 17 |
| MS273| M      | PPMS      | 31                     | 61           | Urinary tract infection, septicemia, MS | 24 |
| MS325| M      | PPMS      | 2                      | 51           | Broncho-pneumonia | 13 |
| MS383| M      | PPMS      | 8                      | 42           | Aspiration pneumonia | 17 |
| MS398| F      | PPMS      | 28                     | 57           | Broncho-pneumonia, MS | 28 |
| MS473| F      | PPMS      | 13                     | 39           | Broncho-pneumonia, MS | 9 |

**Table 2.** List of antibodies used for neuropathological characterization of human post-mortem snap-frozen brain tissue by Immunohistochemistry (IHC) and Immunofluorescence (IF). Abbreviations: MOG = myelin-oligodendrocyte glycoprotein; MHC II = major histocompatibility complex class II molecules, GFAP = glial fibrillary acidic protein; NEFH = neurofilament heavy chain; PFA = paraformaldehyde.

| Antigen | Host | Clonality | Clone | Dilution for IHC/IF | Working post-fixation method | Code | Source | Marker |
|---------|------|-----------|-------|---------------------|-----------------------------|------|--------|--------|
| MOG     | Rabbit | Polyclonal | -     | IHC 1:200; IF 1:100 | Cold Methanol; IHC Ethanol; IF 4% PFA | 12690-1-AP | Proteintech | myelin antigen presenting cells |
| MHCII   | Mouse | Monoclonal | C3/43 | IHC 1:200; IF 1:100 | IHC 1:200; IF 1:100 | M0775 | Dako | antigen presenting cells |
| A20     | Rabbit | Polyclonal | -     | IHC 1:100; IF 1:50 | 4% PFA | ab47037 | Abcam | A20 |
| CD4     | Mouse | Monoclonal | MT310 | IF 1:50 | 4% PFA | sc-19641 | Santa Cruz Biotech. | T helper cells |
| CD8     | Mouse | Monoclonal | UCH-T4 | IF 1:50 | 4% PFA | sc-1181 | Santa Cruz Biotech. | cytotoxic T cells |
| CD68    | Mouse | Monoclonal | S14H12 | IF 1:50 | 4% PFA | MCA1851T | AbD Serotec | macrophages |
| CD20    | Mouse | Monoclonal | -     | IF 1:50 | 4% PFA | M0755 | Dako | B cells |
| GFAP    | Mouse | Monoclonal | S10C | IF 1:100 | 4% PFA | ab190288 | Abcam | astrocytes |
| NEFH    | Mouse | Monoclonal | N52.1.7 | IF 1:100 | 4% PFA | MAB9540 | Abnova | neurons |
Total RNA was extracted from the cryosections using the RNeasy Lipid Tissue Mini Kit (#74804, Qiagen, Hilden, Germany), according to the manufacturer’s instruction. Total RNA was reverse-transcribed at final concentration of 20 ng/μL using random hexamer primers (High-Capacity cDNA Reverse Transcription Kit, #4368814, Thermo Fisher Scientific). Gene expression analysis was performed by real-time PCR using Applied Biosystems’ TaqMan gene expression products (Thermo Fisher Scientific) for the following genes: A20, TNFR1, TNFR2, TNF, IL1B, CCL11, TGFβ, IL10, IL6, NF-κB, STAT3, TRAF6, CASP3, CASP8, BCL2, and XIAP. Transcriptional expression was normalized using GA3PDH as a housekeeping gene. Expression levels of the analyzed genes were calculated by the normalized comparative cycle threshold (Ct) method (2−ΔΔCt), using the Universal Human Reference RNA (Stratagene, Santa Clara, California) for calibration.

**Statistical analysis**

Normality of distributions was assessed by graphical inspection and by the Shapiro–Wilks test. A20 OD was compared between lesioned and neighboring normal-appearing areas of MS cases by means of the paired Mann–Whitney U test. Normalized OD signals of AL and CAL and of aGML and aGML were compared by the Mann–Whitney U test. The association between A20 levels and clinical and demographical variables was evaluated by regression models. Kruskal–Wallis with Dunn post hoc test was applied to compare gene expression data between CC samples and normal-appearing or lesioned areas of MS cases. Lesioned and neighboring normal-appearing areas of MS cases were compared by the paired Mann–Whitney U test. p-values were adjusted for multiple comparisons using the false discovery rate (fdr) correction. Spearman correlation was evaluated between gene expression data. Statistical analyses were performed using R software 3.5.3 version (www.r-project.org).

**RESULTS**

**Characterization of A20 protein expression in human post-mortem control brain tissues**

To investigate the A20 protein expression in brain, we performed IHC and double IF using A20-specific antibodies on brain slices from 5 CC. In both WM (Figure 1A-D) and GM (Figure 1E-J) of CC, the IHC revealed the A20 expression in cells morphologically resembling astrocytes (Figure 1C-C′ and I-I′) and the double confocal IF confirmed that A20 is basally expressed by scattered GFAP-positive (GFAP+) astrocytes (Figure 1D, J and Figure S1). In GM, the IHC also showed the A20 expression in the perinuclear area of cells morphologically resembling neurons (Figure 1G-G′), as confirmed by double confocal IF showing a population of NEFH+ neuronal cells expressing A20 (Figure 1H and Figure S1). We did not detect A20 in MHC II+ microglial cells in either WM or GM (data not shown).
We further evaluated A20 expression in human post-mortem brain tissues from progressive MS patients, both in normal-appearing and demyelinated areas of WM and GM. In NAWM (Figure 2A-E') of MS cases, we detected A20 expression on ramified cells resembling astrocytes (Figure 2C-C'). Double confocal microscopy analysis demonstrated that the A20+ cells were mostly GFAP+ astrocytes, although not all the astrocytes expressed A20 (Figure 2D-D' and Figure S2). However, differently from the CC WM, we observed A20 expressed also by few MHC II+ microglia cells (Figure 2E-E' and Figure S2). In the demyelinated area (Figure 2F-M'), including pre-AL, AL, and CAL, we detected A20 expressed by a huge number of highly ramified resident cells (Figure 2H-H' and L-L'). Particularly in CAL we observed a greater number of ramified cells expressing A20 in the active plaque rim compared to few A20+ cells in the lesion core (Figure 2L-L'). Double confocal microscopy clarified that in WML A20 is expressed by all the GFAP+ astrocytes (Figure 2I-I') and by rare cells with astrocytic morphology expressing GFAP astrocytic marker (J). The inserts in (C', G', and I') show A20-positive cells in WM and GM at high magnification. In D, H, and J, the arrows indicate the colocalization of the A20 and the specific cell marker signals, while the arrowheads show the lacking of colocalization. Original confocal microscopy magnification: 40x. [Colour figure can be viewed at wileyonlinelibrary.com]

Characterization of A20 protein expression in human post-mortem brain tissues

To quantify the A20 protein expression in both WML and GML, we performed a densitometric analysis on IHC images.
from 13 MS cases (10 SPMS and 3 PPMS). We excluded major differences in the A20 expression pattern between PPMS and SPMS by graphical inspection. When we examined all the MS cases together, for WML, we measured A20 optic density (OD) in 8 AL, 12 CAL, and 20 neighboring NA WM areas, classified based on MOG and MHC II immunostaining. For AL, we considered the entire area of the lesion, while for CAL only the active rim. A20 OD was significantly higher in both AL and CAL compared to their neighboring NA WM (paired Mann–Whitney U test, \( P = 0.0078 \) and \( P = 0.0005 \), respectively) (Figure 5A,B). Furthermore, we compared normalized OD signals of AL and CAL, obtained by subtracting the OD value of the neighboring NA WM area to that of the lesion. As a result, we did not observe significant differences (Mann–Whitney U test, \( P = 0.34 \)) (Figure 5C).

For GML, we measured A20 OD in 27 aGML, 9 iGML, and 36 neighboring NAGM areas, classified through MOG and MHC II immunostaining. We found higher A20 immunoreactivity in the aGML as compared to the closest NAGM areas (paired Mann–Whitney U test, \( P < 0.0001 \)) (Figure 5D). In contrast, we did not observe significant differences between A20 immunoreactivity in iGML and neighboring NAGM (paired Mann–Whitney U test, \( P = 0.10 \)) (Figure 5E).

As expected, A20 normalized expression was higher in the aGML compared to iGML (Mann–Whitney U test, \( P < 0.0001 \)) (Figure 5F).

Also, we did not highlight relevant correlations between A20 immunoreactivity in both WML and aGML and plaque dimension, demographical and clinical features of subjects, such as gender, MS type, disease length, age and cause of death, and post-mortem delay (data not shown).

**Gene expression analysis of A20 and related molecules in human post-mortem CC and MS brain tissues**

We further investigated the snap-frozen brain tissues of CC and MS cases by quantifying the gene expression level of A20 (Figure 6) and other 13 genes belonging to the A20 pathways and involved in the regulation of inflammation and apoptosis (Figure S4). In particular, we analyzed 4 WM and 4 GM samples from 4 CC and 12 NAWM, 9 WML, 12 NAGM, and 12 GML samples from 10 progressive MS patients. Given the high variability between-subjects in the gene expression data, we first compared CC samples (WM and GM) with normal-appearing and lesioned areas of MS cases (Figure 6A,C), then, we restricted the
comparison of normal-appearing and lesioned areas of MS cases to matched samples from the same slice of brain tissue blocks (Figure 6B,D).

The A20 expression level was significantly higher in WML compared to both WM of CC and matched NAWM (WM vs. WML: Kruskal–Wallis with Dunn post hoc test, \( P = 0.048 \); NAWM vs. WML: paired Mann–Whitney \( U \) test, \( P = 0.008 \)) (Figure 6A,B). A20 was more expressed also by GML compared to matched NAGM, although the difference was less pronounced than in the WM (paired Mann–Whitney \( U \) test, \( P = 0.02 \)) (Figure 6D). On the contrary, we did not observe significant differences between NAGM or aGML and GM of CC (Kruskal–Wallis test, \( P = 0.27 \)) (Figure 6C).

Figure 3. Characterization of A20 expression in perivascular infiltrates in human post-mortem MS brain tissue by IHC and IF. In the area of demyelination, A20 was found in perivascular infiltrates (C, C’), not expressing CD4 T helper (D-G’), CD8+ T cytotoxic (H-K’), or CD20 B lymphocytes markers (L-O’). Conversely, A20 was expressed by a high number of CD68+ macrophages infiltrating the active demyelinated lesions in WM (P-S’). The arrows indicate the colocalization between A20 and CD68 macrophage marker, while the arrowheads indicate the lack of A20 colocalization with cell-specific markers (CD4, CD8 or CD20). Original confocal microscopy magnification: 40x. [Colour figure can be viewed at wileyonlinelibrary.com]
None of the genes belonging to the A20 pathways and involved in the regulation of inflammation and apoptosis showed significant differences among GM tissues (Figure S4). However, most of them showed differences between WM tissues. TNF receptors 1 (TNFR1) and 2 (TNFR2), whose stimulation by TNF-\(\alpha\) induces an A20 upregulation, were higher expressed in the WML compared to the matched NA WM (paired Mann–Whitney \(U\) test, \(P = 0.02\) and \(P = 0.02\), respectively) (Figure S4A-B). The pro-inflammatory cytokines TNF\(\alpha\) and IL-1\(\beta\), both able to upregulate A20 expression through the binding to their receptors, showed higher levels in the WML compared to the neighboring matched NAWM (paired Mann–Whitney \(U\) test, \(P = 0.02\) and \(P = 0.02\), respectively) (Figure S4C-D). However, we did not observe any difference in the expression of the pro-inflammatory cytokine IL6 (Figure S4E). The anti-inflammatory/regulatory cytokines IL10 and TGF\(\beta\) were highly expressed in the WML compared to the matched NAWM (paired Mann–Whitney \(U\) test, \(P = 0.04\) and \(P = 0.008\), respectively) and to WM of CC, although for TGF\(\beta\) this difference was not significant after adjusting for multiple comparisons (Kruskal–Wallis with Dunn post hoc test, \(P = 0.03\) and \(P = 0.11\), respectively) (Figure S4F-G).

Regarding the apoptotic pathways, we observed higher expression levels of the caspases CASP3 and CASP8 in the WML compared to the neighboring NAWM (paired Mann–Whitney \(U\) test, \(P = 0.02\) and \(P = 0.008\), respectively) and, for CASP8 only, also compared to WM of CC (Kruskal–Wallis with Dunn post hoc test, \(P = 0.04\)) (Figure S4H-I). The anti-apoptotic molecule XIAP, but not BCL2, showed an increased expression in WML compared to matched NAWM (paired Mann–Whitney \(U\) test, \(P = 0.02\)) (Figure S4J-K).

In the correlation plots, A20 expression was significantly correlated with the expression level of most of the genes in its pathway, in both WML and GML (Spearman correlation coefficient, \(P < 0.05\)) (Figure S4N-O). Notably, in WML the strongest correlation was observed between A20 and the neuronal nuclei in the NAGM (Figure S4L-M). As shown in the correlation plots, A20 expression was significantly correlated with the expression level of most of the genes in its pathway, in both WML and GML (Spearman correlation coefficient, \(P < 0.05\)) (Figure S4N-O). Notably, in WML the strongest correlation was observed between A20 and the neuronal nuclei in the NAGM (Figure S4L-M). As shown in the correlation plots, A20 expression was significantly correlated with the expression level of most of the genes in its pathway, in both WML and GML (Spearman correlation coefficient, \(P < 0.05\)) (Figure S4N-O).

DISCUSSION

The key role of A20 in immune system regulation and in restricting the inflammatory processes in periphery (38, 40), as well as its involvement in inflammatory and autoimmune diseases such as MS, is widely known (10, 14, 38, 39).
However, recent data suggested an involvement of A20 also in the pathophysiology of CNS, especially in mouse model of MS (1, 22, 36, 45, 51, 61, 62). In humans, the distribution of some components of the A20 ubiquitin-editing complex has been described in post-mortem brain tissues by Pranski and coauthors, but the study of A20 has been limited by the difficulty to find an anti-A20 primary antibody working on this type of tissues (55). Undoubtedly, close to the identification a suitable antibody working on snap-frozen human brain samples, the high quality of the tissues obtained by the UK tissue bank, probably due to the standardized procedure of preservation and freezing adopted, eased to detect the protein.

Here, we aimed to perform, for the first time to our knowledge, a neuropathological characterization of the A20 expression in post-mortem brain tissues obtained from CC and MS patients, with both PP and SP form of the disease. We observed that A20 protein and transcript are expressed in brain tissues of healthy donors, both in the GM, by several neurons, and in the WM, by few parenchymal astrocytes. These data are in accordance with those of Pranski and colleagues (55), which described the presence of components of the A20 ubiquitin-editing complex prevalently in neurons, in which NF-kB pathway is constitutively activated and involved in cell homeostasis and survival, and more rarely in glia cells. Unlike neurons, in glia cells as well as in the immune cells, A20 expression is minimal at the steady state. Conversely, A20 rapidly increases its expression upon NF-kB activation following stimulation with different factors including inflammatory cytokines and microbial derived products. In this way, A20 controls and restricts the NF-kB activity by a negative feedback loop mechanism (28).

In MS brain tissues, we observed a similar pattern of A20 expression in PP and SPMS, suggesting that they share, at least partially, some common inflammatory and/or regulatory mechanisms of responses to pathological conditions.

**Figure 5.** Densitometric analysis of A20 immunoreactivity measured on IHC images of human post-mortem brain tissues of MS cases. The A20 OD was compared between NAWM and AL (A) or CAL (B) and between NAGM and aGML (D) or iGML (E) (paired Mann–Whitney U-test, *P < 0.05, **P < 0.001, ***P < 0.0001). Normalized A20 OD signals, calculated by subtracting the OD value of the normal-appearing area to that of the matched lesion area, were compared between AL and CAL (C) and between aGML and iGML (F) (Mann–Whitney U test, *P < 0.05, **P < 0.001, ***P < 0.0001). [Colour figure can be viewed at wileyonlinelibrary.com]
In addition, the distribution of the A20 protein in CNS resident cells of no-lesioned area was similar to that identified in the CC. Particularly, in the NAGM A20 was expressed by several neurons in the soma, while in the NAWM it was expressed by some highly ramified astrocytes and by few microglia cells.

On the contrary, A20 showed a very different pattern of expression in the demyelinated active lesions in both WM and GM. A20 protein was highly expressed in all the WML examined. In particular, in the CAL A20 was mostly expressed by a very dense network of reactive astrocytes in the active rim of the lesions, while in the AL it was mainly expressed by infiltrating macrophages along the whole area of injury, but not by B and T lymphocytes. In AL, as in the CAL, A20 was also expressed by astrocytes characterized by massive enlargement of the cell soma and reduced prove density, a hypertrophic morphology typical of astrogliosis reaction. We also observed A20 expression in some population of

Figure 6. A20 gene expression analysis in human post-mortem brain tissues of MS and CC. The A20 expression was compared between WM of CC and NAWM and WML of MS cases (A), and between GM of CC and NAGM and aGML of MS cases (C) (Kruskal–Wallis with Dunn post hoc test, *P < 0.05, **P < 0.001, ***P < 0.0001). In addition, the A20 expression was compared between matched samples of normal-appearing and lesioned areas of MS cases (B, D) (paired Mann–Whitney U test, *P < 0.05, **P < 0.001, ***P < 0.0001). Relative gene expression was calculated by the normalized comparative cycle threshold (Ct) method 2−ΔΔC. [Colour figure can be viewed at wileyonlinelibrary.com]
microglia, but the number of A20-positive microglia cells could be underestimated due to the massive presence of A20-positive reactive astrocytes in the same areas. Corroborating these data, Voet and colleagues described an overexpression of the A20 transcript in WML compared to NAWM control tissue of human post-mortem MS brain tissues (61).

Similarly to WML, in the majority of aGML a tick mantle of reactive astrocytes expressed A20 at high levels. We did not observed the A20 expression in other cell types, but, as for the WML, we cannot exclude it could have been hidden by the strong and compact A20 immunofluorescence signal in astrocytes.

To this regard, neurons present in cortical lesions were previously demonstrated to overexpress A20 in combination with TNFR1. This profile of expression was associated to a gradient of neuronal loss from the CSF/pial surface toward the inner cortical layers (54).

In addition, we were not able to detect A20 in oligodendrocytes, due to the lack of validated protocols for oligodendrocytes markers working after PFA fixation in frozen brain tissues (39).

Finally, we described the upregulation of the gene expression of several A20-related molecules in the WML compared to NAWM, including upstream receptors inducing A20 expression such as TNFR1 and TNFR2, pro-inflammatory cytokines such as IL-1b and TNF-a, the transcription factor NF-kB and STAT3, and the pro-apoptotic-molecules CASP3 and CASP8. We also observed the upregulation of regulatory cytokines such as TGFβ and IL10 and anti-apoptotic enzymes such as XIAP, suggesting that negative feedback processes might possibly occur in some active lesioned areas. However, in the GML, we did not observe significant modulation of these molecules.

The high levels of A20 expression that we observed in the macrophages infiltrating the WML is line with the well-known regulatory role of the enzyme in the immune system and in particular in the circulating myeloid cells, where it controls cell homeostasis and activation to prevent an abnormal pro-inflammatory response (15, 40). Macrophages are the most predominant immune cell type in inflammatory demyelinating MS lesions. It is well established that foamy macrophages primarily promote MS disease progression by internalizing myelin debris, presenting brain-derived autoantigens, and secreting pro-inflammatory mediators (13, 25). However, early studies show that they also promote axonal sprouting and remyelination (21, 30). This dual role can be explained by the fact that macrophages comprise several different phenotypical and functional subpopulations (46, 47), thus, further experiments are needed to clarify the role of the enzyme in these cells.

Glia cells as well display different phenotypes and functions. On the one hand, the microglia promotes neuroinflammatory and neurodegenerative events in MS by releasing inflammatory mediators. On the other hand, it supports the CNS repair through the production of neurotrophic factors and the clearance of myelin debris (4). Also astrocytes are early and active players in the lesion pathology (7, 54) and could exert both detrimental and beneficial effects (54). In active lesions, astrocytes assume a hypertrophic morphology and may sustain damage through the production of pro-inflammatory cytokines and reactive oxygen species (7). In a later stage of lesion development, astrocytes may dampen inflammation through the secretion of BDNF and the expression of TLRs promoting neuroprotection and lesion repair (8, 18).

Contrary to the vast literature on A20 in blood peripheral cells, less is known on its role in glia cells. The completely germinal deficiency of A20 in mice causes spontaneous neuroinflammation in puppies, with remarkable reactive microgliosis and astrogliosis (22). Consistently, the selective ablation of A20 in astrocytes resulted in a more severe clinical course of the EAE, as a result of increased chemokines production and enhanced recruitment of pro-inflammatory T lymphocytes in the spinal cord (62). The specific A20 deletion in microglia increases microglial cell number, affects microglial regulation of neuronal synaptic functions, and also exacerbates MS-like disease (45, 61).

According to the literature, here, we demonstrated the overexpression of A20 protein and transcript in glia cells, predominantly astrocytes, in both WML and GML of MS patients. In particular, we found reactive astrocytes expressing A20 in the active margin of demyelinating lesion extending into neighboring NAWM, suggesting possible contribution to lesion development. It would be of interest to investigate which type of astrocytes expresses the enzyme through double staining for A20 and astrocytic markers as well as in vitro or in vivo experiments on mouse models of demyelination.

CONCLUSION

Overall, our results support the hypothesis of a specific role of A20 in activated astrocytes and microglia, close to the well know role in the myeloid cells. However, they do not allow to determine whether it is beneficial or a detrimental, since glia cells display spatial- and temporal-dependent identities in inflamed CNS. A20 could be involved in an attempt to regulate inflammation, or could actively induce demyelination and cell death sustaining the chronic-inflammatory state, according to the inflammatory stimuli. Functional studies are necessary to shed light on this debate.

The knowledge of the role of the A20 pathway in these cells and in the ongoing inflammation during the progressive stage of MS could be crucial to design and address new therapeutic approaches.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS
Study concept and design: Simona Perga, Francesca Montarolo, Antonio Bertolotto, and Roberta Magliozzi. Analysis and interpretation of data: Simona Perga, Serena Martire, and Francesca Montarolo. Drafting of the manuscript: Simona Perga. Critical revision of the manuscript: Simona Perga, Serena Martire, Francesca Montarolo, and Roberta Magliozzi. Statistical analysis: Serena Martire. Obtained funding: Serena Martire. Study supervision: Simona Perga and Antonio Bertolotto.

DATA AVAILABILITY STATEMENT
The authors confirm that the data supporting the findings of this study are available within the article and its supplementary materials.

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of this article at the publisher’s web site:

Figure S1. Identification of cell types expressing A20 in human post-mortem CC brain tissues by IHC and IF. A20 was expressed by rare astrocytes in WM and GM, as validated by double confocal immunofluorescence and confocal microscopy with GFAP astrocytic marker (A-D′ and I-L′). Additionally, in GM, A20 was also expressed by some population of NEFH+ neurons, with a perinuclear distribution, as shown by the colocalization of A20 with the DAPI nuclear staining (E-H′). The inserts in (D′, H′, L′, and P′) show A20-positive cells in WM and GM at high magnification. Original magnification: 40X. The arrows indicate the colocalization between A20 and the cell-specific markers; the arrowheads indicate cells that do not express A20.

Figure S2. Identification of cell type expressing A20 in human post-mortem MS brain tissues by IHC and IF. In NAGM, A20 was expressed in the perinuclear area of some population of neurons, identified with anti-NEFH neuronal marker (A-D′), and by some GFAP+ astrocytes (I-L′). In the area with aGML, A20 is expressed by neurons (E-H′) and by a fit network of astrocytes, identified with GFAP-specific marker (M-P′) similarly to NAGM). The inserts in (D′, H′, L′, and P′) show A20-positive cells in NAGM and aGML at high magnification. Original magnification: 40X. The arrows indicate the colocalization between A20 and the cell-specific markers; the arrowheads indicate cells that do not express A20.

Figure S3. Identification of cell type expressing A20 in human post-mortem MS brain tissues by IHC and IF. In NAGM, A20 was expressed in the perinuclear area of some population of neurons, identified with anti-NEFH neuronal marker (A-D′), and by some GFAP+ astrocytes (I-L′). In the area with aGML, A20 is expressed by neurons (E-H′) and by a fit network of astrocytes, identified with GFAP-specific marker (M-P′) similarly to NAGM). The inserts in (D′, H′, L′, and P′) show A20-positive cells in NAGM and aGML at high magnification. Original magnification: 40X. The arrows indicate the colocalization between A20 and the cell-specific markers; the arrowheads indicate cells that do not express A20.

Figure S4. Gene expression analysis of molecules in the A20 pathway: TNF receptors (A-B), pro- and anti-inflammatory cytokines (C-G), molecules involved in the apoptosis regulation (H-K) and pro-inflammatory transcription factors (L-M). The expression levels were compared between WM of CC and NAWM and WML of MS cases, and between GM of CC and NAGM and aGML of MS cases (Kruskal–Wallis with Dunn post hoc test, *P* < 0.05, **P* < 0.001, ***P* < 0.0001). In addition, the A20 expression was compared between matched samples of normal-appearing and lesioned areas of MS cases (paired Mann–Whitney *U* test, *P* < 0.05, **P* < 0.001, ***P* < 0.0001). Relative gene expression was calculated by the normalized comparative cycle threshold (Ct) method 2−ΔΔCt. Correlation plot of genes in WM (N) and GM (O) (Spearman correlation coefficient, *P* < 0.05).