MAPT (Tau) expression is a biomarker for an increased rate of survival for low-grade glioma

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Abstract. The association of MAPT (Tau) with various tauopathies and other neurological disorders has long been established. However, the role of microtubule-associated protein Tau (MAPT) expression in brain cancer is largely unknown. To determine whether MAPT expression is related to low-grade glioma (LGG) survival rates, RNASeq data representing samples from the Cancer Genome Atlas (TCGA) were assessed. Results revealed that high expression of the MAPT gene is very strongly associated with increased overall and disease-free survival in LGG but not in breast cancer or melanoma. No such association was apparent for either amyloid precursor protein or α-synuclein gene expression. The expression levels of particular apoptosis- and pro-proliferative-effector genes were consistent with the Tau-associated increased survival rates. It has been well established that the Tau protein plays a neurodegenerative role, and in this study we identified, for the first time, a potential cell apoptosis function that Tau may play in cancers of the central nervous system.

Introduction

The MAPT gene (microtubule-associated protein Tau gene) encodes the Tau protein, which is involved in microtubule stabilization and tubulin polymerization into microtubules. It has long been established that aggregations of hyperphosphorylated neurofibrillary and gliofibrillary tangles of Tau protein are involved in several neurodegenerative disorders, known as tauopathies (1,2). However, the role of Tau in brain cancers, and specifically, low-grade glioma, has not been explored.

Gliomas are histologically graded from I to IV. Grade I gliomas are usually benign. Grade II [low-grade glioma (LGG)] has an average survival period of ~7 years. Grade II gliomas can progress into grade III (high-grade gliomas), and eventually grade IV (secondary glioblastoma). Recent research indicates that historically used clinical variables in LGG are inferior prognostic indicators relative to current genetic information, for example the IDH1 mutational status (3). As LGG is an early stage in the progression of gliomas from low grade to high grade, there is a need to understand the disease process from this early onset. Pinpointing key molecular features that are associated with improved outcomes will help elucidate the disease progression and possibly provide prognostic markers (4).

Given Tau's role in neurodegeneration in the brain and microtubule stabilization, it was of interest to explore Tau's connection to LGG. Colodner et al demonstrated that glial fibrillary tangle formation was accompanied by JAK/STAT-mediated apoptotic cell death of both glia and neurons in a Drosophila model of glial tauopathy (5). However, previous research on Tau in the cancer setting, for the most part, has focused on microtubule-inhibiting chemotherapy in breast cancer (6-8). For example, Zhou et al identified Tau expression as a factor to predict tumor sensitivity to microtubule-inhibiting compounds such as taxanes (9). In the present study, we present a novel look into the association of Tau with survival rates for LGG patients.

Materials and methods

MAPT, apoptosis-effector gene and pro-proliferative gene RNAseq analysis. RNASeq values for MAPT, for the panel of apoptosis-effector genes, and for the panel of pro-proliferative genes, were downloaded from cbioportal.org. Pearson correlation coefficients were obtained, using Microsoft Excel (10-12).

Kaplan-Meier analyses and RNASeq data. The barcodes (patient identifiers) representing 530 LGG samples,
1,105 BRCA (breast cancer) samples and 478 SKCM (skin cutaneous melanoma) samples, for which RNASeq data were available, were sorted from low to high based on the RNASeq values for each of the indicated genes in the Results section. Survival relationships were then obtained by matching barcode survival data, available from cbioportal.org, with the RNASeq data representing the top and bottom 20% of the RNASeq values (106 barcodes for LGG, 221 barcodes for BRCA and 94 barcodes for SKCM); and with the top and bottom 50, 40, 30 and 10% of the RNASeq values for MAPT (LGG only).

The clinical data (months of overall survival and disease-free survival) for the associated barcodes were then processed using IBM Statistics Package for the Social Sciences (SPSS), version 24, to provide the Kaplan-Meier (KM) curves and an average difference in survival rates between samples representing the top and bottom RNASeq value percentages (for the genes as indicated in Results).

Histological grade analysis. Histological grades for LGG barcodes were downloaded from cbioportal.org. Histological grade barcodes were then matched to the RNAseq data (106 barcodes for LGG). Comparison of proportions test for the data representing the top and bottom 20% of the MAPT RNASeq values was carried out using the MedCalc comparison of proportions calculator (https://www.medcalc.org/calc/comparison_of_proportions.php).

Copy number variation (CNV) analysis. Copy numbers for the MAPT gene for LGG barcodes were downloaded from cbioportal.org. Copy numbers were then matched to barcode RNAseq data (530 barcodes for LGG). A Student's t-test for copy number data representing the top and bottom 20% of the MAPT RNASeq values was carried out using the Microsoft Excel function.

Methylation analysis. Methylation data for the MAPT gene for LGG barcodes were downloaded from cbioportal.org. Methylation data barcodes were then matched to RNASeq data (530 barcodes for LGG). A Student's t-test for the methylation data representing the top and bottom 20% of the MAPT RNASeq values was carried out using the Microsoft Excel function.

Diagnosis age analysis. Diagnosis age data for LGG barcodes were downloaded from cbioportal.org. Diagnosis age barcodes were then matched to RNASeq data (530 barcodes for LGG). A Student's t-test for diagnosis age representing the top and bottom 20% of the MAPT RNASeq values was carried out using the Microsoft Excel function.

R2: Genomics Analysis and Visualization Platform. MAPT gene expression and overall survival data for additional glioma datasets were assessed using the ‘R2: Genomics Analysis and Visualization Platform (http://r2.amc.nl)’. Single gene Kaplan scan analysis was conducted for MAPT expression (203929_s_at) for each dataset as indicated in Results. Median MAPT mRNA expression was used as a cut-off for overall survival comparisons. The R2 algorithm was used to generate Kaplan-Meier analyses to compare survival outcomes for the top half vs. bottom half of MAPT expressers in each indicated dataset.

Supplementary and supporting material. A large amount of supplementary and supporting documentation has been prepared for the generation of this report, including detailed data supporting for the Materials and methods section directly above. This information is available as a single PDF upon E-mail request to the corresponding author or by accessing: http://www.universityseminarassociates.com/Supporting_online_material_for_scholarly_pubs.php.

Results

MAPT (Tau) expression and survival rates. We first compared the rate of overall and disease-free survival for LGG patients representing the top and bottom 20% of MAPT expressers, based on the MAPT RNASeq values representing the TCGA-LGG data set (Fig. 1). This comparison indicated that high MAPT RNASeq values were associated with a significantly increased overall and disease-free survival rates for LGG. To substantiate this result, we compared the top 10, 30, 40 and 50% MAPT expressers with their corresponding lower level MAPT expressers for overall survival. In all cases, the initial result was confirmed.

To determine whether the above association of MAPT RNASeq levels with the markedly distinct survival rates was specific to LGG, data for overall survival and disease-free survival for BRCA and SKCM tumor barcodes were downloaded from cbioportal.org, and the RNAseq and KM analyses were conducted. Results indicated that MAPT expression levels did not correlate with either overall or disease-free survival rate differences in the cases of BRCA or SKCM (Fig. 1). The mean survival periods and P-values for the log-rank tests are given in the figure legend for Fig. 1. These data indicated that the higher overall and disease-free survival rates for high MAPT expressing patients was specific to LGG.

To determine whether the increased LGG survival rate associated with MAPT expression could be associated with other neurotoxic proteins, we assessed whether significantly increased survival rates could be associated with higher RNASeq values for APP or SNCA. Analyses indicated there were no differences in survival for the patients represented by the LGG barcodes representing the top and bottom quintiles, for RNASeq values for APP or SNCA, respectively (Fig. 2).

LGG apoptosis-effector gene RNASeq values are correlated with MAPT RNASeq values. As Tau is a neurotoxic protein associated with various tauopathies, and contributes to neuronal cell death, we sought to determine whether apoptosis-effector genes were expressed at higher levels in the LGG samples represented by the barcodes with higher levels of MAPT RNASeq values. We first evaluated a set of 28 apoptosis-effector genes (13) for RNASeq expression as represented by the LGG barcodes. Of those 28 genes, we identified a subset of five apoptosis-effector genes that had significantly increased RNASeq values associated with barcodes that represented the top 20% of MAPT expressers: CASP9, CAC1, CRADD, COX7A2L and UQCRCC2 (Table I).

To verify the above positive correlations of apoptosis-effector gene RNASeq values with MAPT RNASeq values, we obtained the Pearson correlation coefficients for the
Figure 1. Kaplan-Meier curves representing distinct MAPT expression levels. Results here indicate that there were overall survival (OS) and disease-free survival (DFS) distinctions for low-grade glioma (LGG) based on the MAPT levels in the patient tumors. However, no such distinctions were detectable in BRCA (breast cancer) and SKCM (skin cutaneous melanoma) tumor samples. The following text provides mean survival times and P-values for the Kaplan-Meier (KM) analyses. (A) KM OS curve for LGG barcodes that are in the top quintile of MAPT expressers (n=96), compared to the OS for the bottom quintile of MAPT expressers (n=104). Mean OS for the top quintile of MAPT expressers, 109.04 months; mean OS for bottom quintile of MAPT expressers, 59.36 months. Log-rank comparison P-value, P=9.152E-10. (B) KM OS analysis for SKCM barcodes that are in the top quintile of MAPT expressers (n=90), compared to the OS for the bottom quintile of MAPT expressers (n=91). Mean OS for the top quintile of MAPT expressers, 131.09 months; mean OS for bottom quintile of MAPT expressers, 131.09 months. Log-rank comparison P-value, P=0.975. (C) KM OS analysis for BRCA barcodes that are in the top quintile of MAPT expressers (n=218), compared to the OS for the bottom quintile of MAPT expressers (n=218). Mean OS for the top quintile of MAPT expressers, 154.70 months; mean OS for bottom quintile of MAPT expressers, 164.95 months. Log-rank comparison P-value, P=0.302. Mean DFS for the top quintile of MAPT expressers, 74.72 months; mean DFS for bottom quintile of MAPT expressers, 34.80 months. Log-rank comparison P-value, P=2.215E-10. (D) KM DFS analysis for LGG barcodes that are in the top quintile of MAPT expressers (n=93), compared to the DFS for the bottom quintile of MAPT expressers (n=86). Mean DFS for the top quintile of MAPT expressers, 169.72 months; mean DFS for bottom quintile of MAPT expressers, 192.60 months. Log-rank comparison P-value, P=0.316. MAPT, microtubule-associated protein Tau gene.

Figure 2. Kaplan-Meier (KM) curves representing distinct APP and SNCA expression levels. Results here indicate that no survival distinctions are detectable for low-grade glioma (LGG) based on APP or SNCA expression levels. (A) KM overall survival (OS) analysis for LGG barcodes that are in the top quintile of SNCA expressers (n=105), compared to the OS for the bottom quintile of SNCA expressers (n=100). Mean OS for the top quintile of MAPT expressers, 89.44 months; mean OS for bottom quintile of MAPT expressers, 82.74 months. Log-rank comparison P-value, P=0.279. (B) KM OS analysis for LGG barcodes that are in the top quintile of APP expressers (n=102), compared to the OS for the bottom quintile of APP expressers (n=102). Mean OS for the top quintile of MAPT expressers, 134.17 months; mean OS for bottom quintile of MAPT expressers, 138.93 months. Log-rank comparison P-value, P=0.565. APP, amyloid precursor protein gene; MAPT, microtubule-associated protein Tau gene.

Table I. Average apoptosis-effector gene RNASeq values for LGG barcodes representing high and low expression of MAPT.

<table>
<thead>
<tr>
<th>Apoptosis-effector gene</th>
<th>Bottom 20% MAPT RNASeq values</th>
<th>Top 20% MAPT RNASeq values</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CASP9</td>
<td>511.0</td>
<td>901.2</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>COX7A2L</td>
<td>1,158.2</td>
<td>1,300.2</td>
<td>0.0011</td>
</tr>
<tr>
<td>CRADD</td>
<td>193.1</td>
<td>215.0</td>
<td>0.0054</td>
</tr>
<tr>
<td>CYC1</td>
<td>1,952.8</td>
<td>2,339.4</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>UQRC2</td>
<td>2,601.8</td>
<td>3,853.1</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

LGG, low-grade glioma; MAPT, microtubule-associated protein Tau gene; CASP9, caspase 9; COX7A2L, cytochrome c oxidase subunit 7A2 like; CRADD, CASP2 and RIPK1 domain containing adaptor with death domain; CYC1, cytochrome c1; UQRC2, ubiquinol-cytochrome c reductase core protein 2.
MAPT RNA expression levels and the RNA expression levels of CASP9, CYC1, CRADD, COX7A2L and UQCRC2 in LGG. In each case, there was a statistically significant correlation with MAPT expression (Fig. 3).

To determine whether any of the five above identified apoptosis-effector genes represented independent markers of survival rates, the overall survival data for LGG barcodes representing the highest 20% and lowest 20% of CASP9, CYC1, CRADD, COX7A2L and UQCRC2 expression levels were obtained for KM analysis. Results indicated that only CASP9 and UQCRC2 represented independent markers of survival, with barcodes representing the highest 20% of both CASP9 and UQCRC2 having significantly increased survival rates (Fig. 4).
expressed at higher levels in the samples with lower levels of the MAPT RNASeq values. We previously established a panel of 62 proliferation-effector genes (13). Of those 62 genes, we identified a subset of 28 proliferation-effector genes that have significantly increased RNASeq expression in barcodes that represent the bottom 20% of MAPT expressers.

To verify the above inverse correlations of proliferation-effector gene expression with MAPT expression, we obtained the Pearson correlation coefficients for MAPT RNASeq values and the RNASeq values for a histone subset of the above indicated, 28 proliferation-effector genes, for the LGG barcodes: H2AFX, HIST1H2BK, and HIST2H2BE. In each case, there was a statistically significant inverse correlation with MAPT expression (Fig. 5).

To determine whether any of the above indicated histone genes represented independent survival markers, we generated KM survival curves based on the top and bottom LGG expressers. From these analyses, we determined that the expression of all three of the histone genes independently represented survival distinctions (Table II).

**LGG histological grade is correlated with MAPT RNASeq values.** To determine whether MAPT expression correlates with histological features of LGG, we compared histological grades of the top quintile and bottom quintile of MAPT-expressing barcodes. All LGG barcodes in the dataset used were graded either G2 or G3. We determined that the low MAPT-expressing barcodes, that also represented decreased survival rates, also had a greater proportion of G3 histological grade than the high MAPT-expressing barcodes (Table III).

**CNV and methylation are correlated with MAPT RNASeq values in LGG.** To determine whether there may be any genetic or epigenetic associations with increased MAPT expression, we compared MAPT copy number variation (CNV) and methylation of the MAPT gene in the top quintile and bottom quintile of MAPT-expressing barcodes using data available on...
For CNV, a negative result indicates average copy number loss, while a positive result indicates average copy number gain among the barcodes analyzed. Our results indicated that there is an average increase in MAPT copy number among barcodes that are in the top quintile of MAPT expressers, while there is an average decrease in copy number among barcodes that are in the bottom quintile of MAPT expressers. Furthermore, our results indicated that there is an increase in MAPT gene methylation among barcodes that are in the bottom quintile of MAPT expressers (Table IV). Together, the CNV and methylation results indicated that there are likely straightforward genetic and epigenetic explanations for decreased MAPT expression.

Table III. Identification of histological grade associations with MAPT expression.

<table>
<thead>
<tr>
<th></th>
<th>Bottom quintile MAPT expressers</th>
<th>Top quintile MAPT expressers</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fraction G3 histological grade</td>
<td>0.78</td>
<td>0.22</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

MAPT, microtubule-associated protein Tau gene.

Table IV. Copy number variation and methylation associations with MAPT expression.

<table>
<thead>
<tr>
<th></th>
<th>Bottom quintile MAPT expressers</th>
<th>Top quintile MAPT expressers</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copy no. variation</td>
<td>-0.031</td>
<td>0.048</td>
<td>0.0002</td>
</tr>
<tr>
<td>Methylation</td>
<td>0.490</td>
<td>0.355</td>
<td>3.479E-13</td>
</tr>
</tbody>
</table>

MAPT, microtubule-associated protein Tau gene.

Table V. Diagnosis age associations with MAPT expression.

<table>
<thead>
<tr>
<th></th>
<th>Bottom quintile MAPT expressers</th>
<th>Top quintile MAPT expressers</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average age of diagnosis</td>
<td>48.56</td>
<td>39.38</td>
<td>4.248E-07</td>
</tr>
</tbody>
</table>

MAPT, microtubule-associated protein Tau gene.

Table VI. Identifying MAPT expression associations with survival in additional glioma datasets, using the R2: Genomics Analysis and Visualization Platform.

<table>
<thead>
<tr>
<th>Survival distinctions</th>
<th>Glioma (French) n=284</th>
<th>Glioma (Kawaguchi) n=50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increased/decreased survival for the top half of MAPT expressers in the indicated dataset</td>
<td>Increased</td>
<td>Increased</td>
</tr>
<tr>
<td>Increased/decreased survival for the bottom half of MAPT expressers in the indicated dataset</td>
<td>Decreased</td>
<td>Decreased</td>
</tr>
<tr>
<td>KM, log-rank P-value</td>
<td>&lt;0.0001</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

MAPT, microtubule-associated protein Tau gene; KM, Kaplan-Meier.

Diagnosis age is correlated with MAPT RNASeq values in LGG. To determine whether MAPT expression may have an effect on diagnosis age, we compared the average age of diagnosis in patient barcodes representing the top and bottom quintile of MAPT RNASeq expression using data available on cbioportal.org. Results indicated that increased MAPT expression was associated with an earlier diagnosis age. The top quintile of MAPT expressers had an average diagnosis age of 39, while the lowest quintile of MAPT expressers had an average diagnosis age of 48 (Table V).

Analysis of MAPT expression and survival in additional glioma datasets from the R2: Genomics Analysis and
visually correlates with Tau expression. In addition, the histological grade of the tumor was inversely correlated with Tau expression. Several studies have noted the link between DNA damage and Tau aggregation, ultimately leading to cell cycle activation and apoptosis in mature neurons (14). Various studies indicate that Tau forms hyperphosphorylated aggregates which causes the loss of heterochromatin, allowing genes that are normally silenced by heterochromatin to be transcribed, ultimately leading to cell cycle activation and apoptosis in mature neurons (14). Several studies have noted the link between DNA damage and neurodegeneration in Alzheimer's disease (15-19).

While much has been explored regarding the role of Tau pathology in neurons, there has been less focus on investigating the role of Tau aggregation in glial cells. Several studies have demonstrated the functional consequences of Tau aggregation in glial cells (5,20). Various studies indicate that Tau pathology in glial cells is similar to that in neurons, to an extent, but is not identical (21,22). If we apply the model of Tau oligomerization impact on neural cells (in tauopathies), to glial cells, Tau accumulation in glial cells (in LGG) could trigger a DNA damage response and apoptosis. Our results would be consistent with this hypothesis through increased expression of several apoptosis-effector genes in the population of patients that have high MAPT expression.

Tau is crucial in stabilizing and organizing axonal microtubules in neural cells (23). However, in various tauopathies, Tau forms hyperphosphorylated aggregates which causes the disruption of the microtubule network (24). While the role of Tau in glial cells remains less-investigated, if we apply the previously established model of Tau aggregation and phosphorylation contributing to microtubule destabilization in neural cells to glial cells, it becomes evident that decreased Tau aggregation may allow increased microtubule stabilization, and thus increased cell proliferation. The above data are consistent with this idea as LGG samples with decreased MAPT expression expressed higher proliferation-effector genes, and specifically expressed more histone genes.

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Availability of data and materials
The supporting online material files indicated in the article are available upon E-mail request to the corresponding author.

Authors' contributions
SZ made the initial discovery connecting Tau expression to LGG survival outcomes and conducted the majority of the subsequent analyses. BIC conducted the analyses regarding the proliferation-effector genes. DS contributed to the design of particular aspects of RNAseq interpretations. GB supervised the project, provided guidance for the design of the project, conducted some of the analyses directly, spot-checked much of the analyses, wrote most of the drafts of the manuscript and takes full responsibility for the content. All authors read and approved the manuscript and agree to be accountable for all aspects of the research in ensuring that the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Ethics approval and consent to participate
Not applicable

Patient consent for publication
Not applicable.

Competing interests
The authors state that they have no competing interests.

References


