Proliferation and aneusomy predict survival of young patients with astrocytoma grade II

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The clinical course of astrocytoma grade II (AII) is highly variable and not reflected by histological characteristics. As one of the best prognostic factors, higher age identifies rapid progressive A II. For patients over 35 years of age, an aggressive treatment is normally propagated. For patients under 35 years, there is no clear guidance for treatment choices, and therefore also the necessity of histopathological diagnosis is often questioned. We studied the additional prognostic value of the proliferation index and the detection of genetic aberrations for patients with A II. The tumour samples were obtained by stereotactic biopsy or tumour resection and divided into two age groups, that is 18–34 years (n = 19) and ≥35 years (n = 28). Factors tested included the proliferation (Ki-67) index, and numerical aberrations for chromosomes 1, 7, and 10, as detected by in situ hybridisation (ISH). The results show that age is a prognostic indicator when studied in the total patient group, with patients above 35 years showing a relatively poor prognosis. Increased proliferation index in the presence of aneusomy appears to identify a subgroup of patients with poor prognosis more accurately than predicted by proliferation index alone. We conclude that histologically classified cases of A II comprise a heterogeneous group of tumours with different biological and genetic constitution, which exhibit a highly variable clinical course. Immunostaining for Ki-67 in combination with the detection of aneusomy by ISH allows the identification of a subgroup of patients with rapidly progressive A II. This is an extra argument not to defer stereotactic biopsy in young patients with radiological suspicion of A II.

Keywords: astrocytoma grade II; prognosis; proliferation; aneusomy; Ki-67 index; in situ hybridisation

Controversy exists with respect to optimal treatment protocols for low-grade diffuse astrocytoma (astrocytoma WHO grade II; A II) because prospective studies comparing treatment strategies are rare (Karim et al., 1996; Shaw et al., 2002). Another reason for this controversy is that the interval after which malignant progression of A II occurs is difficult to predict. Clinical factors that may correlate with survival include those related to patient age, presenting symptoms, duration of symptoms, performance status, tumour volume, extent of resection, and timing and dose of radiotherapy (Berger et al., 1994; Lote et al., 1997). Patient age is the single consistent prognostic factor in these retrospective studies. Patient age under 35 years (Shaw et al., 1989; Arienti et al., 2001), or under 40 years in other series (Bauman et al., 1999) is often associated with a prolonged survival. Since the benefits of early treatment have to be balanced against the possibility of long-term side effects from radiation therapy, patient age is decisive in current treatment protocols for A II. For patients under 35 years of age, the benefit of early and aggressive treatment has never been proven sufficiently and therefore treatment is often deferred (Vecht, 1993). As a result, in young patients also a controversy exists concerning the usefulness of an immediate histological diagnosis, involving stereotactic biopsy (Recht et al., 1992).

Ultimately the majority of A II progresses to high-grade astrocytomas (astrocytoma WHO grade III and grade IV; A III/A IV), which is characterised by an increase in proliferation activity and an accumulation of genetic abnormalities. Proliferation and cytogenetic markers may therefore identify rapid progressive A II. Increased proliferation activity correlates with shorter survival in most series of astrocytomas, although the number of A II was often too small for separate analyses (Sallinen et al., 1994; Korkolopoulou et al., 1997). Numerical chromosomal aberrations have been reported in astrocytomas, such as aneusomy 1, trisomy 7 and monosomy 10. It has been shown that trisomy 7 correlates with shorter survival of A II patients (Wessels et al., 2002).

In the underlying study, the prognostic value of the proliferation (Ki-67) index and the detection of numerical aberrations for
chromosomes 1, 7, and 10 was evaluated. The correlation of these parameters with survival analysis was performed for patients aged 18–34 years and ≥35 years to assess whether these parameters allow the identification of rapid progressive A II in young patients.

MATERIALS AND METHODS

Patient material

Tissue specimens from 47 adult patients diagnosed with supratentorial astrocytoma grade II were collected from the data files of the Departments of Pathology of the University Hospitals of Maastricht and Groningen, and the Atrium Hospital in Heerlen, The Netherlands. Histopathological revision according to the WHO classification (Kleihues and Cavenee, 2000) revealed 44 fibrillary astrocytomas and three gemistocytic astrocytomas (all WHO grade II). Astrocytomas with mitotic figures were not included in this series. In order to diminish the probability of sampling error A II in which the neuroradiologists suspected high-grade astrocytoma and which showed radiological characteristics for tumour bleeding, as well as extensive contrast enhancement, were not included in this series. Patient records were examined with regard to the first neurological symptoms, radiological findings, neurological procedure, dose and timing of radiotherapy, and survival interval.

Mean and median ages were respectively, 38 and 41 years (range, 18–69 years). The study included 22 women (47%) and 25 men (53%). Seizures were the most frequent presenting symptom (77%), followed by focal neurological deficit (26%), mental changes (13%) and signs of raised intracranial pressure (17%). The median duration of preoperative symptoms was 3 months (range: 1 week–157 months). Most tumours were located in the frontal (51%) and temporal/parietal (45%) lobes and less frequently in the occipital lobe (4%). The majority of patients (n = 41) underwent neurosurgery immediately after coming for medical attention and had their first neuroimaging test. In six patients, neurosurgical intervention was extended for 36–150 months. No association was found, using Pearson’s correlation coefficient, between preoperative interval and proliferation index or chromosomal status. For this reason all cases were included in this study.

Neurosurgical procedures consisted of a biopsy in 29 (62%) and resection in 18 (38%) patients. Radiotherapy was given immediately postoperative in 28 (60%) patients, delayed in 10 (21%) patients, and nine (19%) patients were still not irradiated at the last follow-up.

Proliferation index: Ki-67 immunohistochemistry

Paraffin sections (5-μm thick) were preincubated in methanol with 0.3% H2O2. Tissues known to be negative and positive for Ki-67 were used as controls. Antigen retrieval was achieved by incubation with 10 mM citrate buffer (pH 6.0) in a domestic microwave oven at 700 W for 10 min. The sections were incubated with the mouse monoclonal antibody MIB-1 directed against Ki-67 (Immunotech S.A., Marseille, France) at 1:12 solution in PBS containing 4% normal goat serum for 60 and 45 min, respectively. Peroxidase activity was detected using diaminobenzidine in PBS/ imidazol buffer with 0.02% H2O2. Nuclei were scored for positivity in 500 cells in each sample in the areas with highest immunopositivity. The mean proliferation (Ki-67) index was 2.7% (range 0.0–9.2%).

Statistical analyses

The following factors were considered as possible prognostic parameters for survival: patient age, sex, presenting symptoms, preoperative duration of symptoms, tumour location, neurosurgical procedure, timing of radiotherapy, proliferation index, and chromosomal status.

The influence of these factors on survival was tested by univariate analysis using log-rank tests. Possible prognostic factors from the univariate analyses with P-values less than 0.10 (according to recommendation in the literature (Hosmer and Lemeshow, 2000)) were entered into the multivariate analyses using a forward stepwise method in order to identify independent prognostic parameters (Cox, 1972). Univariate and multivariate associations between factors and survival were assessed using a Cox regression model. Subsequently, tumour-related factors were used for analysis in age-stratified groups.
RESULTS

The median survival interval for patients with astrocytoma grade II as estimated by the Kaplan–Meier method was 90 months (95% confidence interval 72–108 months). When determining the influence of age on the period of survival after the first diagnosis of A II, it becomes obvious that the group aged ≥35 years at the time of diagnosis exhibits a significantly shorter survival period as compared to the 18–34 year-old patient group (Figure 1; log-rank; *P*-value = 0.05). Other factors significantly associated with shorter survival were focal neurological deficit at presentation, proliferation (Ki-67) index >1%, and aneusomy. Upon multivariate analysis proliferation index >1%, patient age ≥35 years, and aneusomy were independently correlated with shorter survival (Table 1).

To investigate the interactions between the proliferation index and aneusomy with age, we analysed the influence of these two factors on survival stratified for age. When correlating the proliferation index to survival in the two age groups separately, it appears that in the group of 18–34-year-old patients, a clearcut proliferation index to survival in the two age groups separately, factors on survival stratified for age. When correlating the proliferation index and aneusomy with age, we analysed the influence of these two factors on survival stratified for age. When correlating the proliferation index to survival in the two age groups separately, it appears that in the group of 18–34-year-old patients, a clearcut distinction can be made between those with a proliferation index ≤1%, showing long-term survival, and those with a proliferation index >1%, showing more progressive AII (Figure 2A; log-rank; *P*-value = 0.02). Also in the ≥35-year-old patients, the proliferation index proved to have additional prognostic value, although less apparent as compared to the younger group (Figure 2B; *P*-value = 0.03).

Using the ISH protocol with probes for chromosomes 1,7, and 10, cases of A II with an apparently normal (disomic) chromosomal content can be separated from aneusomic cases. In the whole group of patients with A II the detection of aneusomy has additional value in distinguishing between rapid and slow progressive A II. In the stratified age groups an aberrant chromosomal constitution is not associated with shorter survival in both the 18–34-year-old (Figure 3A; *P*-value = 0.36) and the ≥35-year-old (Figure 3B; *P*-value = 0.09) groups.

However, the combination of a high proliferation index and aneusomy very accurately identified patients with an unfavourable outcome in both the 18–34-year-old (Figure 4A; *P*-value = 0.01) and the ≥35-year-old patient groups (Figure 4B; *P*-value = 0.001). The combination of both factors reclassified three (27%) of 11 patients of 18–34 years into the group with relatively good prognosis (compare Figures 4A to 2A). In five of 18 patients in the
35 years group, the high proliferation index with disomy now correlated with a more favourable course (compare Figures 4B to 2B).

Figure 5 illustrates a case of A II, localised in the frontal lobe of a 25-year-old patient with epileptic seizures as the sole manifestation. The proliferation (Ki-67) index was 4.5% and a high percentage (35%) of nuclei with trisomy for chromosome 7 was detected by ISH. Despite the favourable clinical features (young age, no focal deficit), the patient rapidly progressed to astrocytoma grade IV, and the survival interval after histological diagnosis was only 36 months.

**DISCUSSION**

A 'wait and see' policy has been propagated for patients under 35 years of age, suffering from epileptic seizures and radiological suspicion of astrocytoma grade II (A II) (Vecht, 1993). An important argument for not performing a stereotactic biopsy in these patients is that the histological diagnosis of A II does not alter treatment strategy. In the current study, we demonstrate that the detection of a high proliferative activity in combination with chromosomal aberrations identifies a subgroup of young A II patients with a rapid malignant course.

Our study confirms previous studies of A II showing the prognostic value of patient age and symptoms at presentation (Lote et al, 1997; Bauman et al, 1999). In multivariate analyses, only the age of patients at diagnosis remained significantly associated with survival as an independent clinical factor, when comparing patient group under and over 35 years of age. However, when including biological factors describing the proliferative

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**Figure 3** Relation of aneusomy with shorter survival in (A) A II patients aged 18–34 (P-value = 0.36) and (B) patients ≥ 35 years (P-value = 0.09). Kaplan–Meier pooled log-rank test, P = 0.05. —— Aneusomy (n = 15). — Disomy (n = 32).

**Figure 4** The combination of both high proliferation (Ki-67) index and aneusomy is strongly associated with a shorter survival in A II patients (A) aged 18–34 (P = 0.01) and (B) aged ≥ 35 years (P = 0.001). Kaplan–Meier pooled log-rank test, P < 0.001. —— Proliferation index ≤ 1% or Disomy (n = 26). —— Proliferation index > 1% and Aneusomy (n = 21).
et al, 2002). In our series, the discriminative power of the proliferation (Ki-67) index diminished when using a higher cutoff level than 1% (data not shown). These differences in cutoff levels, among different studies, may be explained by differences in the immunohistochemical procedures applied, such as the method of antigen retrieval, the immunolabelling protocol, and by differences in the scoring criteria of (clustered) Ki-67-positive cells. Therefore, an additional, but independent prognosticator is needed.

The detection of numerical genomic aberrations has such additional prognostic value to the proliferation (Ki-67) index. For astrocytomas of all grades, it appears that certain genetic changes are associated with an unfavourable clinical course. Astrocytomas with chromosomal abnormalities as detected by karyotyping lead to a shorter survival as compared to astrocytomas without these abnormalities (Kimmel et al, 1992). An ISH study showed that monosomy for chromosome 10, harbouring the tumour suppressor gene PTEN/MMAC1, results in shorter survival (Cianciulli et al, 2000). However, in these series, the majority of tumours represented high-grade astrocytoma (grades III and IV).

In A II, the most frequently reported genetic aberration is loss or mutation of the p53 gene, which, however, shows no association with clinical course (Kraus et al, 1994; al-Sarraj and Bridges, 1995).

One of the very few additional studies correlating genetic aberrations and clinical course includes a comparative genomic hybridisation (CGH) analysis, which showed that A II with rapid malignant progression exhibit a significantly higher number of chromosomal aberrations as compared to A II with an indolent behaviour (Sallinen et al, 1997). However, the sensitivity of this CGH technique is relatively low due to contamination with normal/reactive cells, which are often present in A II (Kallioniemi et al, 1994).

In contrast, the ISH procedure is routinely applicable to paraffin-embedded samples of central nervous system tumours (Arnoldus et al, 1992). Our study demonstrates that the detection of chromosomal aberrations by ISH, using a panel of probes for chromosomes 1, 7, and 10, offers an extra independent predictor for survival of patients with A II. After stratification for age, the presence of chromosomal aberrations alone is not significantly associated with shorter survival, which may be caused by the small size of both groups. However, in both age groups, the detection of aneusomy by ISH adds value to the proliferation index, in that increased proliferation in the presence of chromosomal aberrations is associated with a poor prognosis.

A correlation between aneusomy and proliferative activity was demonstrated in astrocytomas of all grades, containing aneuploides for chromosomes 7 and 10, particularly in Ki-67-positive cells (Steilen-Gimbel et al, 1996). This suggests that an accumulation of chromosomal aberrations in proliferating cells plays an important role in the early stages of astrocytoma progression. This also suggests that the combination of both these parameters is therefore very useful in identifying A II with a rapidly malignant clinical course.

Although imaging studies were used as adjunct to histology, A III/IV may also present as nonenhancing lesions (Chamberlain et al, 1988). Therefore, one could argue that in the stereotactic specimen, sampling error from high-grade astrocytomas may have biased the results. This is contradicted, however, by the relatively long median survival of 90 months, also compared to other A II studies (Leighton et al, 1997; McCormack et al, 1992). Another argument against underscoring of the tumours is given by the fact that only one of the biopsy specimens exhibited monosomy 10, which is characteristic at low-grade areas in high-grade astrocytomas (Cheng et al, 1999).

The additional value of genetic and biological parameters to the current histological WHO classification was also seen in the three gemistocytic variants of A II. Previous studies suggest that A II
with high percentages (>60%) of gemistocytes, in fact, behave in a more similar manner to A III (Krouwer et al., 1991). Although two of our gemistocytic samples contained lower percentages of gemistocytes, the increased proliferation activity and the presence of aneusomy in all three samples was associated with a relatively short survival (range 41 – 43 months).

We postulate that identification of subtypes of astrocytomas on the basis of genetic and biological factors has additional prognostic value to the current histological classification. Tissue should be obtained in all patients with A II in order to assess the proliferation activity and, if possible, the presence of trisomy for chromosome 7. In the future, these markers may help in optimising treatment strategy, in particular in young patients with astrocytoma grade II, for whom optimal treatment is now controversial.

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