Reappraisal of the use of procarbazine in the treatment of lymphomas and brain tumors

Jean-Pierre Armand\textsuperscript{1}  
Vincent Ribrag\textsuperscript{1}  
Jean-Luc Harrousseau\textsuperscript{1}  
Lauren Abrey\textsuperscript{2}

\textsuperscript{1}Institut Gustave-Roussy, Desmolins, Villejuif Cedex, France; \textsuperscript{2}Memorial Sloan-Kettering Cancer Center, New York, New York, USA

\textbf{Abstract:} Procarbazine HCl is a ‘nonclassical’ oral alkylating anticancer agent that was first synthesized in the late 1950s. It has been used in the treatment of many cancers, but its main use is in the treatment of Hodgkin’s lymphoma and brain tumors and, to a lesser extent, Non-Hodgkin’s lymphoma and primary central nervous system lymphoma. Procarbazine is a prodrug that undergoes metabolic transformation into active intermediates that are thought to inhibit DNA, RNA, and protein synthesis. Early use of procarbazine in combination with mechlorethamine, vincristine, and prednisone (MOPP) was effective in the treatment of advanced Hodgkin’s lymphoma, but late toxic effects such as secondary cancer and infertility led to its replacement by other regimens. However, its recent reintroduction in the dose-intensified BEACOPP (bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristine, procarbazine, and prednisone) regimen has yielded very promising findings. Procarbazine alone, or more commonly combined in the PCV (procarbazine, lomustine [CCNU], and vincristine) regimen, is also effective in treating gliomas comprising astrocytomas, glioblastomas, and oligodendrogliomas. The most common side effects of procarbazine are gastrointestinal disturbances, myelosuppression, and central nervous system effects. In conclusion, the use of procarbazine in combination with other drugs means that it remains a major anticancer drug in the management of Hodgkin’s lymphoma and gliomas.

\textbf{Keywords:} procarbazine, Hodgkin’s lymphoma, non-Hodgkin’s lymphoma, lymphoma, brain tumor, glioma

\textbf{Introduction}

Procarbazine HCl is a ‘nonclassical’ oral alkylating agent that belongs to the same family as dacarbazine and hexamethylamine (Zeller et al 1963). It contains an N-methyl group that is essential for its activity, but not the chloroethyl group that is present in nitrogen mustard-type alkylating agents (Newell et al 1987). It was first synthesized in the late 1950s during a search for a new monoamine oxidase inhibitor, but was soon developed as an anticancer agent (Kenis et al 1966; Martin and Schubert 1966; Livingston and Carter 1970; Friedman 2001) and has since been used in the treatment of many cancers including Hodgkin’s lymphomas, non-Hodgkin’s lymphoma, brain tumors, multiple myeloma, primary central nervous system lymphoma, malignant melanoma, and lung cancer.

Procarbazine was first granted approval in 1965 and was subsequently marketed in around 84 countries. It has been widely deployed in the treatment of a number of cancers and a considerable body of clinical experience with this drug has accumulated. The introduction of safer combination therapies led to the gradual supplantation of procarbazine. During recent years, there has been a re-emergence of interest in procarbazine combinations with other chemotherapeutic agents, specifically for the treatment of Hodgkin’s lymphoma and gliomas and, to a lesser extent, non-Hodgkin’s lymphoma and primary central nervous system lymphoma. This is partly prompted by
the unique mechanism of action of procarbazine; this agent has multiple sites of action and is not cross-resistant with other alkylating agents, cytostatics or radiotherapy (Matche et al 1963; Brunner and Young 1965).

Reintroduction of procarbazine in the dose-intensified BEACOPP (bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristine, procarbazine, and prednisone) regimen has yielded impressive therapeutic results in the treatment of Hodgkin’s lymphoma and, in combination with lomustine (CCNU) and vincristine in the PCV (procarbazine, CCNU, and vincristine) regimen, it shows considerable promise in the treatment of gliomas.

**Preclinical studies**

**Pharmacology**

Procarbazine is a prodrug that undergoes a complex metabolic transformation into active intermediates. Although these intermediates have been characterized, their precise mechanism of action remains unclear. They are thought to inhibit DNA, RNA, and protein synthesis (Berneis et al 1963; Rutinshauser and Bollag 1963). Studies suggest that procarbazine may inhibit transmethylation of the methyl groups of methionine into t-RNA; the subsequent lack of functional t-RNA could cause cessation of protein synthesis, and consequently DNA and RNA synthesis. DNA may also be damaged directly by hydrogen peroxide, formed during auto-oxidation of procarbazine, which may attack the protein sulphydryl groups in residual protein that is tightly bound to DNA. These effects are specific for the S and G2 phases of the cell cycle. Drug resistance mechanisms to procarbazine also indicate that the methylating pathway is important in its mechanism of action and may help to develop new therapeutic strategies. Potential resistance mechanisms involve O6-alkylguanine DNA alkyltransferase (AGT)-mediated repair of O6-methylguanine activity and hMSH2 mismatch-repair deficiency (Pegg 1990; Friedman et al 1997).

In vivo, significant antitumor activity has been demonstrated against Ehrlich carcinoma and Crocker sarcoma S180 in mice, and Walker carcinoma 256, Guerin uterine carcinoma T8, and intracerebral leukemia L1210 in rats (Bollag and Grunberg 1963).

**Toxicology**

Mutagenic effects of procarbazine have been demonstrated in a variety of bacterial and mammalian test systems, and it has been shown to inhibit fetal development in rats (Malek et al 2003). Unscheduled DNA synthesis in the testes of rabbits has also been reported (Zbinden 1980). As might be expected with an agent that inhibits DNA, RNA, and /or protein synthesis, reduced fertility has been seen in male mice given procarbazine (Chryssanthou et al 1983).

Carcinogenic effects of procarbazine have been reported in mice, rats, and monkeys.

**Pharmacokinetics and drug disposition**

Procarbazine is quickly and completely absorbed from the gastrointestinal tract following oral administration, with peak plasma levels achieved after 0.5–1 hour (Supko et al 2002; Preiss et al 2006). It readily crosses the blood-brain barrier and peak cerebrospinal fluid levels are reached after 0.5–1.5 hours.

The half-life of procarbazine is about 10 minutes and it is rapidly and extensively metabolized by the liver. Oxidation of procarbazine to azo-procarbazine occurs by microsomal cytochrome P450 oxidoreductase, as well as by mitochondrial monoamine oxidase enzymatic conversion. Plasma levels of azo-procarbazine are notably higher than those of procarbazine itself. Azo-procarbazine is microsomally oxidized to azoxyderivatives. Of these, methylazoxy-procarbazine appears to be the major cytotoxic compound, showing several-fold greater cytotoxicity than procarbazine, azo-procarbazine, and benzylazoxy-procarbazine (Eriksen et al 1989). Plasma levels of methylazoxy-procarbazine peak after about 1.5 hours and it has a half-life of around 1 hour.

About 70% of the dose is renally excreted in the first 24 hours; the major urinary metabolite is the biologically inactive N-isopropylterphthalamic acid. The majority of the remainder is excreted via the respiratory system, with only minimal fecal excretion. Impaired renal or hepatic function can delay elimination of procarbazine and its active metabolites.

**Clinical efficacy**

**Lymphomas**

**Hodgkin’s lymphoma**

Hodgkin’s lymphoma affects approximately 8000 new patients each year in the USA and around 1500 in the UK, occurring most commonly in young adults and in people who are more than 55 years old (Glaser and Jarrett 1996; ACS 2006). Although the management of the disease has improved considerably in recent years, around 20% to 30% of patients still experience disease progression or relapse.

Early studies reported overall response rates of 53% to 69% with single agent procarbazine in patients with advanced
Hodgkin’s lymphoma (Brunner and Young 1965; Livingston and Carter 1970). However, these responses were generally very short-lived and there were few complete responses. In the early 1970s, procarbazine was used successfully in combination with three other drugs in the regimen known as MOPP (mechlorethamine [nitrogen mustard], Oncovin® [vincristine], procarbazine, and prednisone) in the treatment of advanced Hodgkin’s lymphoma (DeVita et al 1970). A complete remission rate of 84% was attained, with relapse-free and overall survival rates of 66% and 48%, respectively, after more than 20 years. Unfortunately, although the acute toxic effects of MOPP are manageable, it is associated with significant long-term toxicity. For example, more than 90% of men, and 80%–100% of women over 25 years old, become irreversibly infertile (Pazdur et al 2003). There is also a risk of secondary acute leukemia, which is frequently resistant to treatment and has an extremely poor prognosis (Ng et al 2002).

A number of MOPP-like regimens were developed in an attempt to reduce this toxicity. It would appear that procarbazine plays an important role in MOPP, as lower complete remission rates of shorter duration were reported with MOPP-like regimens that did not contain procarbazine (Nicholson et al 1970; Luce et al 1971).

ABVD (doxorubicin, bleomycin, vinblastine, and dacarbazine) subsequently became the regimen of choice for advanced disease after showing superior efficacy and less long-term toxicity than MOPP and MOPP hybrid regimens (Canellos et al 1992; Duggan et al 2003). The combination of MOPP plus epirubicin, bleomycin, vinblastine, lomustine, doxorubicin, and vindesine (MOPPEBVCAD) has recently been suggested to be as effective as ABVD and more effective than the Stanford V schedule (doxorubicin, vinblastine, mechlorethamine, vincristine, bleomycin, etoposide, and prednisone) (Gobbi et al 2005, 2006). However, despite lower toxicity than normally expected with MOPP analogues, the incidence of second cancers remains somewhat higher than with ABVD. It is, nevertheless, a good option for patients who are not suitable for high-dose chemotherapy.

C-MOPP/ABV, a hybrid chemotherapy in which cyclophosphamide is substituted for mechlorethamine (which has been implicated as the cause of secondary malignancies) has shown beneficial effects (Monoto et al 2000). In 73 patients with advanced disease, 57 (78%) achieved a complete response and 7 further patients who achieved partial response reached complete response after radiotherapy (Monoto et al 2000). The overall survival rate after 4 years was 92%. In another study in which dacarbazine was added to the regimen (C-MOPP-ABV), a response rate of 93% and a complete response rate of 84% was achieved amongst the 67 patients (Takenaka et al 2000).

A recent report of the long-term follow-up of 533 patients with advanced Hodgkin’s lymphoma indicated that ABVPP (doxorubicin, bleomycin, vinblastine, procarbazine, and prednisone) was associated with a better survival probability than MOPP-ABV (Ferme et al 2006). The 10-year overall survival estimates were 90% for ABVPP, 78% for MOPP-ABV, 82% for MOPP-ABV and radiotherapy, and 77% for ABVPP and radiotherapy (p = 0.03).

The German Hodgkin’s Study Group developed a new combination regimen known as BEACOPP, based on a statistical model that considers tumor growth and chemotherapeutic effects (Loeffler et al 1998). The dose of procarbazine is 100 mg/m² over 7 days, which is half that delivered over 14 days in the MOPP regimen. Both regimens are given on a 21-day cycle. In a randomized study, 1195 patients with newly diagnosed Hodgkin’s lymphoma in an unfavorable stage (IIIB, IIIA, IIIB, or IV) were randomized to receive eight cycles of COPP-ABVD (alternating cyclophosphamide, vincristine, procarbazine, and prednisone with doxorubicin, bleomycin, vinblastine, and dacarbazine), standard-dose BEACOPP, or increased-dose BEACOPP (Table 1), each followed by local radiotherapy when indicated (Diehl et al 2003). The outcome of treatment and five-year survival rates are shown in Table 2. The increased-dose BEACOPP regimen was significantly (pd ≤ 0.002) more effective than COPP-ABVD in terms of freedom from treatment failure (87% vs 69%) and overall survival (91% vs 83%) at 5 years, as well as early progression (2% vs 10%). Standard-dose BEACOPP was also significantly (p = 0.04) more effective than COPP-ABVD for freedom from treatment failure at 5 years (76% vs 69%). When the two BEACOPP regimens were compared with each other, the increased-dose regimen resulted in a significantly lower rate of early progression (2% vs 8%) and a significantly (p < 0.001) greater freedom from treatment failure at 5 years (87% vs 76%). Acute hematological effects were more common with the increased-dose BEACOPP regimen. The actuarial rate of secondary acute leukemia five years after diagnosis of Hodgkin’s disease was also significantly (p = 0.03) higher with increased-dose BEACOPP (2.5%) compared with standard-dose (0.6%), and COPP-ABVD (0.4%).

The acute hematological toxicity of standard-dose and dose-intensified BEACOPP was investigated in 858 patients
Armand et al

with advanced Hodgkin’s lymphoma (Engel et al 2000). Dose-limiting toxicity occurred in 25% of cycles of dose-intensified treatment, but severe thrombocytopenia and leukocytopenia were rare. Myelosuppression was the most important dose-limiting factor. The findings indicated that, despite some increase in hemotoxicity, moderate dose escalation was safe for most patients receiving granulocyte colony-stimulating factor (G-CSF) and standard supportive treatment.

BEACOPP has also been used successfully as salvage treatment in patients with early stage Hodgkin’s lymphoma who relapsed after extended field radiotherapy. A retrospective analysis of 107 patients treated mainly with COPP-ABVD-like regimens (n = 74) or BEACOPP (n = 24) showed that BEACOPP was significantly (p = 0.02) superior to the other regimens, with 100% of patients achieving complete remission compared with 85% of those given COPP-ABVD-like regimens, and 67% given other regimens including radiotherapy (Rüffer et al 2005). BEACOPP was also a significant (p = 0.03) prognostic factor for freedom from second failure.

In a study in children and adolescents with advanced disease, induction treatment with dose-intensified BEACOPP improved early tumor control, with a 45% response rate (>70% reduction in disease burden) after 2 cycles and a 72% response rate after 4 cycles (Kelly et al 2002).

Non-Hodgkin’s lymphoma

Single-agent procarbazine is less active in non-Hodgkin’s lymphoma than in Hodgkin’s lymphoma, resulting in overall response rates of only around 36%–40% (Livingston and Carter 1970), although it has a role to play for patients who fail standard regimens or who cannot tolerate or refuse such treatment (Chaar et al 2006). Combination regimens containing procarbazine, such as MOPP and COPP (cyclophosphamide, Oncovin®[vincristine], procarbazine, and prednisone), have resulted in more beneficial effects, particularly in patients with advanced stage nodular mixed or lymphocytic poorly differentiated disease (Lowenbraun et al 1970; DeVita et al 1975; Longo et al 1984; Ezdinli et al 1985). The best response rates to regimens containing procarbazine are seen with intensive cyclic multi-drug combinations such as ProMACE-MOPP (procarbazine, methotrexate with leucovorin, doxorubicin, cyclophosphamide, and etoposide).

In a randomized, controlled trial, 221 patients with diffuse intermediate- to high-grade non-Hodgkin’s lymphoma received ProMACE-MOPP for 6 cycles or the newer generation more aggressive MACOP-B (methotrexate with leucovorin, doxorubicin, cyclophosphamide, and etoposide). In 42 patients with intermediate- or high-grade non-Hodgkin’s lymphoma, THPCOPBLM (pirarubicin, cyclophosphamide, vincristine, prednisone, bleomycin, and procarbazine) combined with G-CSF resulted in a complete remission rate of 88% and a partial remission rate of 12% (Niitsu and Umeda 1998). In 51 older patients (≥55 years old) with intermediate-grade non-Hodgkin’s lymphoma, COPBLAM-I (cyclophosphamide, vincristine, prednisone,

| Table 1 COPP-ABVD, standard-dose BEACOPP and increased-dose BEACOPP dosage regimens (Diehl et al 2003) |
| BEACOPPb | COPP-ABVDa | Standard-dose BEACOPPb | Increased-dose |
|---|---|---|---|
| | Single dose (mg/m²) | Days given | Single dose (mg/m²) | Days given | Single dose (mg/m²) | Days given |
| Bleomycin | 10 iv | 29 and 43 | 10 iv | 8 | 10 iv | 8 |
| Etoposide | - | - | - | - | - | - |
| Doxorubicin | 25 iv | 29 and 43 | 25 iv | 1 | 35 iv | 1 |
| Cyclophosphamide | 650 iv | 1 and 8 | 650 iv | 1 | 1200 iv | 1 |
| Vincristine | 1.4 iv | 1 and 8 | 1.4 iv | 8 | 1.4 iv | 8 |
| Procarbazine | 100 po | 1–14 | 100 po | 1–7 | 100 po | 1–7 |
| Prednisone | 40 po | 1–14 | 40 po | 1–14 | 40 po | 1–14 |
| Vinblastine | 6 iv | 29 and 43 | - | - | - | - |
| Dacarbazine | 375 iv | 29 and 43 | - | - | - | - |

Note: a regimen repeated on day 57; b regimen repeated on day 22.
Abbreviations: BEACOPP, bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristine, procarbazine, and prednisone; COPP-ABVD, alternating cyclophosphamide, vincristine, procarbazine, and prednisone with doxorubicin, bleomycin, vinblastine, and dacarbazine; iv, intravenous; po, oral.
bleomycin, doxorubicin, and procarbazine) resulted in a complete remission rate of 65% after a median follow-up of 2.4 years (Shpilberg et al 1994). The projected 3-year overall survival and progression-free survival rates were 58% and 77%, respectively.

Primary central nervous system lymphoma

Primary central nervous system lymphoma is an aggressive non-Hodgkin’s lymphoma arising in the brain, eyes, spinal cord, or leptomeninges. The incidence has increased markedly over the last two decades, both in patients with and without immunodeficiency (Corn et al 1997).

Although the use of radiotherapy prior to methotrexate improves survival in these patients, it is also associated with neurotoxic effects, such as severe cognitive dysfunction and ataxia (DeAngelis et al 1989). This is a particular problem because at least half of all patients are at least 60 years old at the time of diagnosis. To avoid this problem, a number of chemotherapeutic regimens have been investigated as sole therapy. Procarbazine crosses the blood-brain barrier and has therefore been included in some of these regimens.

In a study of 52 patients treated with 5 cycles of high-dose methotrexate, procarbazine, and vincristine, with or without whole-brain radiotherapy, the objective response rate was 90% and the median survival 60 months (Abrey et al 2000). The median survival was similar amongst the older patients (>60 years old), irrespective of whether or not they received radiotherapy, but late neurotoxicity was significantly more common in those who received radiotherapy. Another study, conducted by the European Organisation for Research and Treatment of Cancer (EORTC) Brain Tumour Group in 50 elderly patients (median age 72 years), investigated the use of high-dose methotrexate, lomustine, procarbazine, methylprednisolone, and intrathecal chemotherapy with methotrexate and cytarabine (Hoang-Xuan et al 2003). The objective response rate was 48% (42% complete and 6% partial responses); the median duration of the response in the patients with complete response was 27 months; the median survival was 14.3 months; and the one-year progression-free survival was 40%. The majority of patients had improved or preserved cognitive function and Karnofsky performance scores (KPS) until relapse. A decline in cognitive function and KPS attributed to late neurotoxicity occurred in only 8% and 12% of patients, respectively. In the long-term responders who were in remission 2 years after the start of therapy, only one patient showed a decline in cognitive function or performance status due to delayed neurotoxicity. This outcome was superior to that expected with radiotherapy alone in elderly patients and considerably reduced the risk of delayed neurotoxicity.

Blood-brain barrier disruption (BBBD) results in enhanced chemotherapy delivery. In 74 patients who underwent BBBD preceded by systemic cyclophosphamide and followed by intra-arterial methotrexate in combination with procarbazine and dexamethasone, a complete response

| Table 2 Outcome of treatment and five-year survival rates with COPP-ABVD, standard-dose BEACOPP, and increased-dose BEACOPP (Diehl et al 2003) |
|---------------------------------------------------------------|
| **COPP-ABVD** *(n = 260)*                                      | **Standard-dose BEACOPP** *(n = 469)*                          | **Increased-dose BEACOPP** *(n = 466)*                          |
| Complete remission                                           | 85 (80–89)                                                  | 88 (85–91)                                                  | 96 (93–97)                                                  |
| Early progression                                            | 10 (7–15)                                                   | 8 (5–10)                                                    | 2 (1–4)**†                                                  |
| Freedom from treatment failure at 5 years                   | 69 (63–75)                                                  | 76 (72–80)**                                                | 87 (83–91)**                                                |
| Overall survival at 5 years                                  | 83 (78–87)                                                  | 88 (85–91)                                                  | 91 (88–94)**                                                |

**Note:** Values given as percentage (95% confidence interval); Early progression defined as progression during treatment or within 3 months after the end of treatment; **p < 0.05 vs COPP-ABVD; †p < 0.001 vs standard-dose BEACOPP.

**Abbreviations:** BEACOPP, bleomycin, etoposide, doxorubicin, cyclophosphamide, vincristine, procarbazine, and prednisone; COPP-ABVD, alternating cyclophosphamide, vincristine, procarbazine, and prednisone with doxorubicin, bleomycin, vinblastine, and dacarbazine.

Table 3 PCV and intensified PCV regimens

| PCV      | Intensified PCV | Days of administration | Frequency       |
|----------|-----------------|------------------------|-----------------|
| Procarbazine 60 mg/m² po | 75 mg/m² po | 8–21                  | Every 6–8 weeks |
| Lomustine 110 mg/m² po | 130 mg/m² po | 8 and 29              |                 |
| Vincristine 1.4 mg/m² iv | 1.4 mg/m² iv | 1                     |                 |

**Abbreviations:** PCV, procarbazine, lomustine (CCNU), vincristine.
was achieved in 65% (n = 48) and a partial response in 19% (n = 14) (McAllister et al 2000). Amongst the 36 patients who continued to show a complete response after one year, none had any evidence of cognitive loss.

Procarbazine has also been used as part of the PCV (procarbazine, CCNU, and vincristine) regimen. Although the prognosis at relapse is often poor for patients suffering from primary central nervous system lymphoma, treatment of 7 relapsed patients with PCV resulted in a complete response in 4 and a partial response in 2 (Herrlinger et al 2000).

**Brain tumors—glioma/astrocytoma**

Gliomas comprise about half of all primary brain tumors and include astrocytomas, glioblastomas, and oligodendrogliomas. The dosage details for the PCV and intensified PCV regimens are shown in Table 3. A summary of the main clinical trials into the use of single-agent procarbazine and procarbazine-containing regimens in patients with glioma is provided in Table 4.

Interest in the use of procarbazine emerged around 1990 when data from the Northern California Oncology Group was reanalyzed to compare the effects of carmustine (BCNU) with that of PCV following radiation/oral hydroxyurea in patients with glioblastoma multiforme or other anaplastic gliomas (Levin et al 1990). PCV treatment resulted in longer survival and time to tumor progression than BCNU, with the difference achieving statistical significance in the group of patients with anaplastic gliomas.

In a randomized study of 171 patients with grade III/IV astrocytoma given PCV, with or without radiotherapy, the median time to progression was 21 weeks and the median survival time was 53 weeks (Sandberg-Wollheim et al 1991). Overall, 18% of patients survived for at least two years. Survival analysis showed that the best responses were achieved in patients less than 50 years old who received both PCV and radiotherapy (median time to progression and survival time of 81 and 124 weeks, respectively).

A durable response to PCV was reported by Kim and colleagues (1996) in 32 patients with grade III/IV oligoastrocytoma or anaplastic oligodendroglioma. The PCV therapy was administered every 6 weeks for a total of at least 124 cycles. The median duration of follow-up review from the start of chemotherapy was 19.3 months. Grade 3 or 4 hematological toxicity was experienced by nine (31%) of 29 patients. Ten patients had delayed treatment due to treatment-related toxicities (34.5%). Ninety-one percent of the 32 patients responded to the therapy. These included 10 patients with a complete response and 19 with a partial response. The median time to progression was 15.4 months for all patients and 23.2 months for those with Grade III tumors. The median time to progression for patients with Grade III oligoastrocytomas was 13.8 months; for those with Grade IV oligoastrocytoma it was 12.4 months, and for those with anaplastic oligodendrogliomas it was 63.4 months (p = 0.0348). These patients survived a median of 49.8 months, 16 months, and 76 or more months, respectively, from the start of chemotherapy (p = 0.0154), thus showing that PCV therapy provides durable responses in patients with Grade III or IV oligoastrocytomas.

A number of other studies found no marked beneficial effects of PCV therapy on survival (Prados et al 1999; Kappelle et al 2001; MRC 2001) and so there has been an increasing tendency to use procarbazine as a salvage option. However, little data are available in the literature regarding this strategy (Rodriguez et al 1989; Newton et al 1990; Brandes et al 1999; Newlands et al 2003). Response rates of 15% to 25% were reported with single-agent procarbazine in patients with recurrent glioblastoma or anaplastic astrocytoma (Rodriguez et al 1989; Newton et al 1990). An overall response rate of 30% was achieved with combined procarbazine and tamoxifen in patients with high-grade gliomas that recurred after surgery and radiotherapy (Brandes et al 1999). Compared with patients suffering from glioblastoma, those with anaplastic astrocytoma had a significantly longer median time to progression (33 vs 13 weeks) and median survival time (57 vs 27 weeks).

Recent results of adjuvant PCV in newly diagnosed patients with anaplastic oligodendroglioma are extremely encouraging (Buckner et al 2003; Stege et al 2005; van den Bent et al 2005, 2006; Yamamoto et al 2005). PCV resulted in tumor regression in a meaningful proportion of newly diagnosed low-grade oligodendroglioma, with or without loss of 1p/19q. In a phase III EORTC study (van den Bent et al 2005, 2006), 6 cycles of adjuvant PCV following radiotherapy in 368 patients with highly anaplastic oligodendroglioma prolonged the progression-free survival from 13 to 23 months compared with radiotherapy alone (p = 0.0018) (Figure 1). Overall survival was also longer with adjuvant PCV, but the difference did not reach statistical significance (Figure 1). The percentage of patients with progression-free survival after 2 years and overall survival after 5 years also showed advantages for adjuvant PCV treatment (Figure 2). Recent evidence suggests that PCV may also be beneficial in patients with symptomatic low-grade oligodendroglioma (Catenois et al 2006).

Beneficial effects were reported with 6-thioguanine, procarbazine, dibromodulcitol, lomustine, and vincristine.
Figure 1 Median progression-free survival and overall survival in patients with newly diagnosed anaplastic oligodendroglioma treated with radiotherapy alone or radiotherapy plus PCV (procarbazine, CCNU, and vincristine) (van den Bent et al 2006).

Figure 2 Percentage of patients with progression-free survival after 2 years and overall survival after 5 years in patients with newly diagnosed anaplastic oligodendroglioma treated with radiotherapy alone or radiotherapy plus PCV (procarbazine, CCNU, and vincristine) (van den Bent et al 2006).
| Study design | N  | Diagnosis                                                                 | Treatment                                                                 | Results                                                                                           |
|-------------|----|----------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------|
| Rodriguez et al 1989 | Open | 99 | Recurrent malignant central nervous system tumors | Procarbazine | Glioblastoma multiforme: response + stabilization rate 27% and MTP 30 weeks. Other anaplastic gliomas: response + stabilization rate 28% and MTP 49 weeks |
| Newton et al 1990 | Open | 35 | Recurrent glioblastoma or anaplastic astrocytoma | Procarbazine | After 2 courses, 2 patients had CR, 7 had PR, 11 had stable disease and 15 had progression. |
| Yung et al 2000 | RCT | 225 | Glioblastoma multiforme at first relapse | Procarbazine vs temozolamidine | 6-month progression-free survival 8% with procarbazine vs with temozolamide, median progression-free survival 8.32 vs. 12.4 weeks, and 6-month overall survival 44% vs. 60% |
| Brandes et al 1999 | Open | 51 | Recurrent high-grade glioblastoma or anaplastic astrocytoma | Procarbazine + high-dose tamoxifen | 4% CR, 26% PR, 32% stable disease. Overall response rate 30%. MTP 13 weeks for glioblastoma and 33 weeks for anaplastic astrocytoma. |
| Newlands et al 2003 | - | 28 | Anaplastic astrocytoma, glioblastoma multiforme or low-grade glioma | Temozolamidine alone for course 1 followed by procarbazine + temozolamide for subsequent courses | Overall, there were 10 (36%) responses lasting from 2 to 17+ months. Clinical impression of the investigators that the combination was somewhat more effective. |
| Levin et al 1985 | Open | 148 | Glioblastoma multiforme, other anaplastic gliomas | BCNU vs PCV | Glioblastoma multiforme: MTP 31 vs 32 weeks for PCV vs BCNU. Other gliomas: MTP 123 and 77 weeks for PCV and BCNU. |
| Levin et al 1990 | Reanalysis of RCT | | Glioblastoma or other anaplastic gliomas | RT/hydroxyurea + PCV or BCNU | Longer survival and time to tumor progression with PCV vs BCNU. |
| Sandberg-Wollheim et al 1991 | RCT | 171 | Grade III/IV astrocytoma | PCV with or without RT | MTP 21 weeks, median survival 53 weeks, 18% survived ≥ 2 years |
| Jeremic et al 1992 | Open | 133 | Anaplastic astrocytoma, glioblastoma multiforme | modified PCV vs VM-26 +CCNU | MTP and survival 2-fold higher with PCV at 50th and 25th percentiles for anaplastic astrocytoma |
| Kim et al 1996 | Open | 32 | Grade III/IV oligoastrocytoma or anaplastic oligodendroglioma | PCV | 31% CR and 59% PR. MTP 15.4 months overall and 23.2 months for those with grade III tumors |
| Prados et al 1999 | Retrospective review | 432 | Anaplastic astrocytoma | PCV vs BCNU | No difference in survival between groups |

*Continued...*
| Study design | N | Diagnosis | Treatment | Results |
|--------------|---|-----------|-----------|---------|
| Levin, Uhm, et al 2000 | Randomized | 2.72 | Glioblastoma | PCV vs. PCV + alpha-difluoromethylornithine | Overall survival 14.2 vs 3.3 months (median) and 8.7% vs 6.2% at 5 years |
| Kappelle et al 2001 | Open | 63 | Recurrent glioblastoma multiforme | PCV | 3%CR, 8% PR and 25% stable disease. MTP 13 weeks. Median survival 33 weeks. |
| MRC 2001 | RCT | 674 | Grade III/IV astrocytoma | RT vs RT + PCV | After median follow-up of 3 years, 310 patients given RT and 307 given RT + PCV died. Median survival 9.5 months with RT and 10 months for with RT + PCV |
| Buckner et al 2003 | Open | 28 | Newly diagnosed low-grade oligodendroglia or oligoastrocytoma | PCV | 8/28 (29%) and 13/35 (52%) eligible patients had tumor regression as assessed by the physician and a blinded neuron-radiology reviewer, respectively. |
| Brandes et al 2004 | Open | 37 | Recurrent anaplastic oligodendroglia or oligoastrocytoma | PCV | 30% CR, 30% PR, 22% stable disease. MTP 12.3 months. |
| Stege et al 2005 | Open | 21 | Newly diagnosed and recurrent oligodendroglia and mixed oligoastrocytomas | PCV | 3/5 recurrent and 13/16 newly diagnosed responded. MTP in the newly diagnosed >24 months. |
| van den Bent et al 2006 | RCT | 368 | Newly diagnosed anaplastic oligodendroglia or anaplastic oligoastrocytoma | RT vs RT + PCV | Progression free survival significantly increased with RT + PCV vs. RT (hazard ratio 0.68; 95% CI 0.53; 0.87, p = 0.0018). No difference in overall survival (hazard ratio 0.85; 95% CI 0.65; 1.11). |
| Coyle et al 1990 | Open | 31 | Recurrent glioma | mechloretamine, vincristine, procarbazine | Response rate 52%. Median duration of response 42 weeks. |
| Kyritsis et al 1996 | Open | 77 | Recurrent malignant glioma | 6-thioguanine, procarbazine, lomustine, hydroxyurea | MTP 9 weeks glioblastoma, 24 weeks anaplastic glioma. PR 2/30 and 11/47. |
| Hildebrand et al 1998 | Open | 37 | Glioblastoma, anaplastic astrocytoma | Dibromodulcitol, BNCU, procarbazine | Response rate 55% with anaplastic astrocytoma, 12% with glioblastoma |
| Levin, Lamborn et al 2000 | Open | 59 | Astrocytoma, ependymoma | 6-thioguanine, procarbazine, dibromodulcitol, lomustine, vincristine | Survival: 4/17 glioblastoma multiforme, 11/25 nonglioblastoma astrocytoma, 6/17 ependymoma |

**Abbreviations:** BNCU, carmustine; CR, complete response; MTP, median time to progression; PCV, procarbazine, lomustine (CCNU), and vincristine; PR, partial response; RCT, randomized controlled trial; RT, radiotherapy; VM-26, teniposide.
(TPDCV) in 42 children and young people with malignant astrocytomas (Levin et al 2000). Of the 17 patients with glioblastoma multiforme, 13 died, the median time to progression was 49 weeks and median survival was 85 weeks. Amongst the remaining 25 patients with nonglioblastoma malignant astrocytoma, 14 died and the median time to progression was 224 weeks. The median survival was not reached in this group after a median follow-up of 494 weeks. The BBSFOP protocol, which comprises seven cycles of three drug pairs (carboplatin-procarbazine, cisplatin-etoposide and vincristine-cyclophosphamide), was used as first line treatment to avoid radiotherapy in 21 children (<5 years old) with high-grade glioma (Dufour et al 2006). The median time to progression was 10.7 months and the median survival time of all patients was 15.0 months. The treatment was safe and well tolerated.

Procarbazine, nimustine, and vincristine given in combination with hyperfractionated radiotherapy resulted in an objective response in 3 out of 8 patients with high-grade glioma (Ogawa et al 2006). The median time to progression was 10.7 months and the median survival time of all patients was 15.0 months. The treatment was safe and well tolerated.

Recent evidence indicates that enzyme-inducing anti-seizure medications have no effect on the pharmacokinetics of procarbazine in patients with glioma, but that there may be an increased risk of hepatic dysfunction with this drug combination (Grossman et al 2006).

Safety and tolerability

Gastrointestinal disturbances (eg, nausea and vomiting), myelosuppression (eg, leukopenia, anemia, and thrombocytopenia) and central nervous system effects (eg, headache, depression, nervousness, and insomnia) are the most common side-effects seen with procarbazine, followed by neuromuscular problems such as tremor, weakness, and paraesthesia. Procarbazine has also been associated with secondary nonlymphoid malignancies, including acute leukemia and solid tumors such as lung cancer, when used in combination therapy. Azoospermia and adverse effects on female gonadal function may also be a problem. Rare side-effects include hemolysis in patients with glucose 6-phosphate dehydrogenase deficiency, hypersensitivity reactions, eosinophilia, and pulmonary infiltrates.

Since procarbazine is a weak monoamine oxidase inhibitor, there is a possibility of interactions with food containing high levels of tyramine (eg, beer, cheese, wine, bananas) and with certain drugs such as sympathomimetics and decongestants. Possible effects include hypertensive crisis, intracranial bleeding, and headache. Procarbazine can also interact with other medications. For example, central nervous system depressants (eg, anesthetics, barbiturates, narcotic analgesics), drugs with anticholinergic effects (eg, tricyclic antidepressants), and antihypertensive agents should all be used with caution and in reduced dosage in patients taking procarbazine due to possible potentiation effects. Alcohol should also be avoided as disulfiram-like effects such as sweating and facial flushing may occur.

Discussion and place in therapy

Procarbazine is a long established anticancer medicine. Although there is renewed interest in this agent, it is worthwhile to remember the many patient-years of experience in many different oncological indications which exist; few ‘new’ cancer therapies have the benefit of such experience.

Although there was a long period in which interest in procarbazine waned due to safety concerns and the development of other cancer treatments, interest is currently re-emerging, mainly in combination with other agents for the treatment of Hodgkin’s lymphoma and gliomas, for which procarbazine activity is high. Some of the early combination therapies which used procarbazine (such as MOPP) were effective, but their use was limited by late toxicity. The new generation of procarbazine-containing combination therapies have been optimized to maximize efficacy and minimize the late-toxic effects seen with the first combinations.

Procarbazine looks likely to continue in the anticancer armamentarium for some time to come and, as a component of the BEACOPP combination, become part of the proposed ‘gold standard’ treatment for Hodgkin’s lymphoma (Massoud et al 2004). Recent results of adjuvant PCV in newly diagnosed patients with anaplastic oligodendroglioma are also extremely encouraging. Despite its long history, procarbazine is still an important drug in the treatment of Hodgkin’s lymphoma and gliomas.

References

Abrey LE, Yahalom J, DeAngelis LM. 2000. Treatment for primary CNS lymphoma: the next step. J Clin Oncol, 18:3144–50.

[ACS] American Cancer Society. 2006. Cancer facts and figures 2006 [online]. Accessed on 28 December 2006. URL: http://www.cancer.org/downloads/STT/CAFF2006PWSecured.pdf.

Berneis K, Kofer M, Bollag W, et al. 1963. The degradation of DNA by a new tumor inhibiting compound. Experientia, 19:132–3.

Bollag W, Grunberg E. 1963. The degradation of deoxyribonucleic acid by new tumour inhibiting compounds: the intermediate formation of hydrogen peroxide. Experientia, 19:130–1.
